# seqMeta: an R Package for meta-analyzing region-based tests of rare DNA variants

December 2, 2013

#### Abstract

Region-based tests are becoming a popular tool for analyzing rare genetic variants. In order for these tests to have adequate power, it is often necessary to meta-analyze information from multiple contributing studies, where consent restrictions make it difficult or impossible to share individual level data. We present the R package **seqMeta** for meta-analyzing region based tests, such as SKAT, SKAT-O, and burden tests, as well as single variant tests. The package can accommodate continuous, binary, and survival outcomes for unrelated individuals, and continuous outcomes for related individuals. We also provide convenient functions for conditional analyses and parallelization.

## 1 Introduction

In association studies of rare genetic variation it is common to pool variants across a gene, region, or pathway to improve statistical power and performance of tests. In order for these test to have adequate power to detect rare variants with modest effects, it is often necessary to use data from multiple contributing studies using a meta-analysis. While meta-analyzing single-variant effect estimates and their standard errors is relatively straightforward, implemented in e.g. METAL, meta-analyzing region-based tests is somewhat more complicated. One popular region based test is SKAT, which is based on a weighted sum of single-variant score tests, using the covariance matrix of the scores to calculate p-values (Wu et al., 2011). Meta-analyzing SKAT requires meta-analysis of score vectors and their covariance matrices. However, meta-analyzing covariance matrices is cumbersome, as they are 2-dimensional structures of various sizes and cannot be easily stored for a whole study in an array. The purpose of the **seqMeta** package is to facilitate meta-analysis of region-based score tests, such as SKAT, through convenient data structures for handling the components of the tests. There is some software available for similar analyses, for instance the MASS program (http://dlin.web.unc.edu/software/mass/). Our implementation is written in R (R Development Core Team, 2009) and supports continuous, binary and survival outcomes for unrelated individuals, continuous outcomes for related individuals, and performs individual variant tests, as well as burden tests, SKAT and SKAT-O.

Use of the package proceeds in two steps. The first is running the study level analyses with the function prepScores. This function need only be called once in each study for all genes under consideration. The output is a an R 'list' object where each element corresponds to a gene, and contains the scores and MAFs for that gene, as well as a matrix of the covariance between the scores at all pairs of SNPs within the gene. This list of output is labeled as a prepScores object, is saved as an .Rdata file, and passed to a central location. The second step combines the output from the study-level analyses, i.e. doing meta-analysis. The meta-analysis functions can create several

different tests including single variant, T1 count (Li and Leal, 2008), Madsen-Browning (Madsen and Browning, 2009), and of course SKAT (Wu et al., 2011).

Meta-analysis functions can be run on a single study alone, and this will produce study level test results. These can be useful for quality control. But the meta-analysis across studies does require the full output from prepScores; it cannot be done with just study level test results.

The rest of this document is organized as follows. Section 2 details the functions used at the study level for continuous outcomes in unrelated individuals (Section 2.1), and related individuals (Section 2.2). Section 3 describes the functions used at the meta-analysis level for SKAT (Section 3.1), burden tests (Section 3.2) and single SNP tests (Section 3.4). Section 4 describes modifications for binary and survival outcomes. Section 6 describes how to perform conditional analyses. Section 7 gives some guidance for parallelization on large datasets, and finally Section 8 describes the mathematical details of the implementation. All sections contain example code which should be directly executable in R using the package seqMeta available at http://cran.r-project.org/web/packages/seqMeta/

## 2 Study level analysis

We are interested in testing the association of groups of SNPs with a particular outcome. A fundamental ingredient for region-based tests is a common definition across studies of which variants are in which regions. For convenience, we will refer to these regions of SNPs as genes, though in principal they can be any unit of aggregation. All functions in the package require a SNP Information file (argument SNPInfo), which links SNPs to genes. By default, the package looks within SNP Information file (a data frame) for the fields Name and gene for SNP and gene identification respectively. If other fields are desired to indicate SNP names or genes, these must be specified explicitly via the arguments snpNames and aggregateBy.

Using prepScores, users are strongly recommended to include all SNPs in the analysis, even if they are monomorphic in one study. This is for two reasons; firstly, monomorphic SNPs provide information about MAF across all studies; without providing the information we are unable to tell if a missing SNP data was monomorphic in a study, or simply failed to genotype adequately in that study. Second, even if some SNPs will be filtered out of a particular analysis (e.g., because they are intronic or common) constructing prepScores objects using all available SNPs allows greater flexibility in the meta-analysis.

## 2.1 Unrelated subjects

To create a prepScores object containing one study's contributions, use e.g.

where;

• Z is a NSNP rows × NSAMPLE columns matrix containing the additively coded (typically 0-1-2 for number of minor alleles) genotypes, with column names which match the SNP names used in the SNPInfo file. Rows do not have to be named, but must be in the same order as in the phenotype file (the data argument). The Z matrix can contain missing genotypes (coded as NA); missing genotypes are imputed to the mean value of the genotype in that study. Because there is no allele-checking functionality, all genotypes should be coded the same way for all studies before running.

- formula is a formula object for the null model of the form outcome ~ covariates. In the example below, we use the formula y~sex+bmi in the example below to analyze an outcome y, adjusted for covariates sex and bmi.
- SNPInfo is a matrix or data-frame with columns titled Name and gene. These columns specify the SNPs, and their associated genes, to be used in the analysis. One can use different columns to designate SNP names and genes with the arguments Name and aggregateBy respectively.
- data is a data frame containing the phenotype information. Phenotypes and genotypes must have the same number of rows and be ordered in the same way, because prepScores matches them simply on row order and not by any form of ID variable. The phenos data frame must contain no missing data in the columns used in the analysis.

For example, analysis at a single study, using the example data in the package, might proceed as below.

```
> rm(list=ls())
> library(seqMeta)
> ######## load example data:
> # contains SNPInfo, phenotyes (pheno1, pheno2)
> # genotypes (Z1, Z2), and pedigree information (kins) for pheno2
> data(seqMetaExample)
> 1s()
[1] "SNPInfo" "Z1"
                        "Z2"
                                   "kins"
                                             "pheno1"
                                                       "pheno2"
> #Perform study-level analysis:
> c1 <- prepScores(Z1, y~bmi+sex, SNPInfo = SNPInfo, data = pheno1)
> ###save the output, which can be passed to a central location.
> study1.out.file <- tempfile()
> save(c1, file = study1.out.file)
```

In this section we described analysis for continuous outcomes. Binary and survival outcomes for unrelated individuals require only slight modification, and are described in Section 4.

#### 2.2 Related subjects

This analysis proceeds in essentially the same was as for related subjects, but requires pair-wise kinship information in stored in a matrix. This is supplied to prepScores via the kins argument

Family structure information is usually stored on disk in a 'makeped' linkage format. To convert this format into to the required kinship matrix, the makekinship function in the kinship2 R! package is recommended; see these packages documentation for details (Therneau et al., 2012). Note that family data is currently only supported for continuous outcomes.

# 3 Meta-analysis

The results of the one or more individual studies can be meta-analyzed using the functions skatMeta, skatOMeta, burdenMeta, or singlesnpMeta for SKAT, SKAT-O, burden tests and individual SNPs respectively. Regardless of the individual analysis (related or unrelated individuals with continuous, binary or survival outcomes) the meta-analysis procedure is the same. Each of these meta-analysis functions has a similar syntax, and contains the arguments

- ... One or more prepScores objects to be meta-analyzed.
- SNPInfo The SNP Info file, listing genes and SNPs to be included in the meta-analysis. This is in the same format as the SNPInfo file in prepScores, and should use the same naming convention for SNPs and genes.
- snpNames The field of SNPInfo where the SNP identifiers are found. Default is Name
- aggregateBy The field of SNPInfo on which the skat results were aggregated. Default is gene. For single snps which are intended only for single variant analyses, it is recommended that they have a unique identifier in this field.
- mafRange A range of minor allele frequencies to be included in the analysis. These are based on the pooled MAFs over all studies. By default this is c(0,0.5), which includes all SNPs.
- verbose logical, whether or not to print progress bars. Defaults to FALSE.

Each of the functions permits additional arguments, which we detail in their corresponding sections.

## 3.1 Meta-analyzing SKAT

The SKAT test is a weighted sum of individual score statistics

$$Q = \sum_{j} w_j^2 U_j^2$$

where  $w_j$  is a weight and  $U_j$  is the score statistic for for the association between phenotype and variant j. Details are given in Section 8. In the function skatMeta! these weights are specified via the wts argument, which gives either a function to be applied to the minor allele frequencies or a column of the SNP Information file which can be coerced to a numeric vector.

The additional argument method allows the user to specify the method of p-values calculation. The default is method'saddlepoint'=, which appears to work well in practice. See the documentation of pchisqsum and Section 8 for more details.

For example, below is a meta-analysis of the SKAT test. Here we have use the default testing weights wts dbeta(maf,c(1,25))=, often referred to as the 'Wu' weights, which are also the default in the **SKAT** package (Lee *et al.*, 2013).

- > load(study1.out.file)
- > load(study2.out.file)
- > meta.results <- skatMeta(c1, c2, SNPInfo = SNPInfo)
- > head(meta.results)

	gene	p	Qmeta	cmaf	nmiss	nsnps
1	gene1	0.858651041	97982.36	0.2050000	600	15
2	gene10	0.385906585	236430.85	0.3112500	1200	21
3	gene100	0.310106989	201665.04	0.2537500	600	16
4	gene11	0.348914794	298153.58	0.3766667	1800	26
5	gene12	0.005833266	364830.51	0.2175000	0	15
6	gene13	0.716506668	193644.56	0.3095833	600	22

Here, skatMeta returns the following information

- gene The gene name, or other unit of aggregation.
- p The p-value from the SKAT test
- Qmeta The SKAT Q statistic, defined as  $\sum_{j} w_{j}U_{j}^{2}$ , where  $w_{j}$  is the weight given to SNP j, and  $U_{j}^{2}$  is associated score statistic.
- cmaf The cumulative minor allele frequency. That is  $\sum_{j} MAF_{j}$ , where  $MAF_{j}$  is the minor allele frequency of variant j, and the sum is over all variants in the gene.
- nmiss The number of missing SNPs. For a gene with a single SNP this is the number of individuals which do not contribute to the analysis, due to studies that did not report results for that SNP. For a gene with multiple SNPs, nmiss is summed over the gene.
- nsnps The number of SNPs in the gene.

It will often be useful to examine results within a study before meta analysis. This can be done by 'meta-analyzing' a single study's results:

```
> study1.results <- skatMeta(c1, SNPInfo = SNPInfo)
```

#### 3.2 Meta-analyzing burden tests

Another commonly used family of tests are the 'burden' tests which regress the phenotype on a weighted sum of genotypes, within each gene. The score test for a weighted sum of genotypes has the form

$$T = \sum_{j} w_{j} U_{j},$$

where  $w_j$  is a weight for SNP j and  $U_j$  is the score for SNP j. For instance if  $MAF_j$  is the minor allele frequency for SNP j, the Madsen-Browning (Madsen and Browning, 2009) test uses  $w_j = (MAF_j(1 - MAF_j))^{-1}$ , while the and the T1 count test (Li and Leal, 2008) uses  $w_j = \mathbf{1}(MAF_j < 0.01)$ .

These types of tests can be computed with the burdenMeta function. This has the argument wts, which like skatMeta, gives either a function to be applied to the minor allele frequencies or a column of the SNP Information file which can be coerced to a numeric vector.

To perform the T1 test one could use the command

```
> meta.t1.results <- burdenMeta(c1, c2, wts = function(maf){maf < 0.01},
+ SNPInfo = SNPInfo)</pre>
```

Equivalently, we could use constant weights and limit the analysis to those SNPs with MAF < 0.01:

Likewise, the Madsen-Browning test could be performed with the command

Regardless of the weights used, these tests give output of the form

> format(head(meta.t1.results),digits=2)

	gene	p	beta	se	${\tt cmafTotal}$	${\tt cmafUsed}$	nsnpsTotal	${\tt nsnpsUsed}$	nmiss
1	gene1	0.888	0.029	0.20	0.20	0.0096	15	1	0
2	gene10	0.064	-0.376	0.20	0.31	0.0096	21	1	0
3	gene100	NA	NA	Inf	0.25	0.0000	16	0	0
4	gene11	NA	NA	Inf	0.38	0.0000	26	0	0
5	gene12	NA	NA	Inf	0.22	0.0000	15	0	0
6	gene13	0.214	0.211	0.17	0.31	0.0163	22	2	600

This output is similar to the gene-level summaries reported by skatMeta, but also includes additional information. Though we employ a score test, which does not explicitly estimate genetic effects, we do report estimated effects beta and their standard error se. These can be thought of as one-step approximations to standard maximum likelihood estimates, which may differ slightly when effect sizes are very large (Voorman et al., 2012). The additional columns cmafTotal and cmafUsed, nsnpsTotal and nsnpsUsed distinguishing the total cumulative minor allele frequency and number of SNPs, from those that are used in the test. Also, note the genes for which no p-value is returned. In these genes, no SNPs met the inclusion criteria, and thus burden is zero for all individuals, and the test is undefined.

#### 3.3 Meta-analyzing SKAT-O

Recently, the 'optimal' SKAT (SKAT-O, Lee et al., 2012) was proposed to test weighted averages of SKAT and burden tests of the form

$$Q_o(\rho) = (1 - \rho) \left( \sum_{j=1}^p w_j^{skat} U_j^2 \right) + \rho \left( \sum_{j=1}^p w_j^{burden} U_j \right)^2$$

When  $\rho=0$  this gives the SKAT test  $Q_o=\sum_{j=1}^p w_j U_j^2$ , and when  $\rho=1$  it gives the burden test  $Q_o=\left(\sum_{j=1}^p w_j U_j\right)^2$ . Note that unlike the version of SKAT-O implemented in the **SKAT** package, we do not assume that the weights used in the burden test are the same as those used in SKAT.

When  $\rho$  is fixed, computation of a p-value can be obtained as with SKAT, using that the test statistic  $Q_o(\rho)$  is a quadratic form of normally distributed score vector U. Lee et al. (2012) propose choosing the value of  $\rho$  which minimizes the p-value of the test over a set values of  $\rho$  in the interval [0, 1]. That is, they find the 'optimal' linear combination of SKAT and burden tests. However, when  $\rho$  is chosen to minimize the p-value, the actual p-value reported must reflect the flexibility afforded by this choice. For example, in the simple case where we allow  $\rho$  to be either 0 or 1, we would

perform both a burden test and a SKAT test and record the minimum of the p-values. We might correct for multiple testing using a Šidák or Bonferroni correction. However, this is not optimal, since burden tests and SKAT are correlated with each other, especially when the number of variants in a region is small. As shown by Lee  $et\ al.\ (2012)$  it is possible to calculate the distribution of the minimum p-value over any sequence of  $\rho$ 's, accounting for this correlation between tests. The mathematics are given in the Appendix of that paper, which require individual level information. In Section 8 we give an alternate derivation of the calculations which can be performed at the meta-analysis level.

The syntax of skat0Meta is similar to that of skatMeta and burdenMeta, with the only difference being that weights for SKAT and the burden test must be distinguished, and values of  $\rho$  must be specified. These are given in the arguments

- skat.wts and burden.wts Either a function, or a character string specifying a column in the SNPInfo file, which gives the weights to be used in SKAT and the burden test, respectively. The default is to use the 'beta' weights in SKAT and T1 weights for the burden test.
- rho The values of  $\rho$  to be used in SKAT-O. The default is c(0,1), which computes SKAT and the burden test, and reports the minimum p-value adjusted for multiple testing.

Here, we illustrate using SKAT-O using Wu' weights in both the SKAT and the burden test, and choose a sequence of 11  $\rho$ 's in [0, 1]

```
> meta.skato.results <- skatOMeta(c1, c2, rho=seq(0,1,length=11),
+ burden.wts = function(maf){dbeta(maf,1,25)}, SNPInfo = SNPInfo, method = "int")
> format(head(meta.skato.results),digits=2)
```

```
pmin rho cmaf nmiss nsnps errflag
     gene
1
    gene1 0.7878 0.5805 1.0 0.20
                                     600
                                            15
  gene10 0.5893 0.3859 0.0 0.31
                                    1200
                                            21
                                                      0
3 gene100 0.4045 0.2530 0.5 0.25
                                                      0
                                     600
                                            16
   gene11 0.5416 0.3490 0.0 0.38
                                    1800
                                            26
                                                      0
  gene12 0.0098 0.0058 0.0 0.22
                                            15
                                                      0
                                       0
   gene13 0.9043 0.7165 0.0 0.31
                                     600
                                            22
```

The format of the results is similar to the other region-based tests, with the addition of the fields pmin, which specifies the minimum p-value among the tests considered, and rho, which gives the value of  $\rho$  which resulted in pmin. Note that p should always be at least as large as pmin, since SKAT-O corrects for the multiple tests considered. If there is a single SNP in a gene, or either SKAT or the burden test is undefined, pmin will be identical to p.

The additional column errflag indicates a possibly inaccurate p-values. A value of 0 indicates no error, values larger than 0 indicate potentially inaccurate p-values. Genes where errflag is not zero can be re-run with more accurate calculation method, such as saddlepoint, which is slower, but more accurate, than integration.

```
> table(meta.skato.results$rho)
```

```
0 0.1 0.2 0.4 0.5 0.6 1
56 2 2 1 2 1 36
```

The table above displays the value of  $\rho$  which gave the smallest p-value among the 11 combinations of SKAT and the burden test. Note that the smallest p-value is typically given by either SKAT or the burden test ( $\rho = 0$  or  $\rho = 1$ ), rather than a proper combination of them. For this reason, the default is simply  $\mathbf{rhoc}(0,1)$ = which performs only SKAT and the burden test, and is typically much faster.

Though skat0Meta in some sense supersedes skatMeta and burdenMeta, we recommended running SKAT and burden tests on their own for quality control. For instance, here we calculate SKAT and the burden test that are used SKAT-O, and plot the SKAT-O p-value, to the minimum p-value from SKAT and the burden test.

We see that the SKAT-O *p*-values are always larger than the minimum of the SKAT and burden test *p*-values (the black line), but smaller than the Šidák correction for two independent tests (the red line). In addition to the errflag column, plots like these provide a rough check of the SKAT-O *p*-value accuracy, which can occasionally be hard to achieve for small *p*-values.

We also note here that the *p*-values for SKAT-O are close to Sidák correction for the minimum of SKAT and burden tests. As the number of variants increases, the SKAT and burden tests become independent. Thus, in the relatively large genes in this example, little efficiency is gained using the SKAT-O correction relative to the Šidák correction.

## 3.4 Meta-analyzing single SNPs

While skatMeta and burdenMeta perform region-based tests, the function singlesnpMeta can be used to perform score tests for single SNP associations. The only additional option available is whether or not to report study-specific effects and standard errors (argument studyBetas)

```
> meta.ss.results <- singlesnpMeta(c1, c2, SNPInfo = SNPInfo,
                                    studyBetas = TRUE)
> format(head(meta.ss.results),digits=2)
  gene
                        maf nmiss ntotal
                                            beta
                                                   se beta.c1 se.c1 beta.c2
           Name
1 gene1 1000001 0.83 0.0129
                                 0
                                     1200 -0.038 0.18
                                                       -0.066
                                                                0.25
                                                                      -0.010
2 gene1 1000002 0.41 0.0146
                                 0
                                     1200 -0.137 0.17
                                                       -0.268
                                                                0.23
                                                                       0.013
                                                                       0.354
3 gene1 1000003 0.89 0.0096
                                 0
                                     1200 0.029 0.20
                                                       -0.284
                                                                0.28
4 gene1 1000004 0.84 0.0158
                                 0
                                     1200 -0.032 0.16
                                                       -0.130
                                                                0.24
                                                                       0.043
5 gene1 1000005 0.50 0.0121
                                 0
                                     1200 0.124 0.18
                                                       -0.112
                                                               0.28
                                                                       0.292
6 gene1 1000006 0.47 0.0142
                                     1200 0.122 0.17
                                                        0.111
                                                               0.23
                                                                       0.134
  se.c2
  0.25
2
  0.24
3 0.29
```

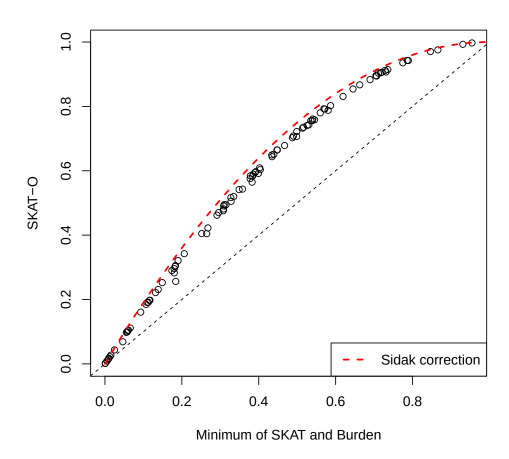


Figure 1: Comparison between SKAT-O p-values, and the minimum of SKAT and burden test p-values. The Šidák correction for two independent tests is  $1 - (1 - p)^2$ .

```
4 0.21
5 0.24
6 0.25
```

The output is similar to that of burdenMeta and skatMeta, but now includes effect estimates for single studies, suffixed by the name given to the corresponding prepScores object.

As with the burden test, the coefficients and standard errors can be thought of as one-step approximations to maximum likelihood estimates.

## 4 Binary and survival outcomes

The **seqMeta** package can also handle binary data and survival outcomes for unrelated populations.

For binary outcomes, this is specified with the family argument in the function prepScores, which accepts a family object as in the function glm.

```
> c1.bin <- prepScores(Z2, ybin~bmi+sex, family = binomial(),
+ SNPInfo = SNPInfo, data = pheno1)</pre>
```

For survival outcomes this fitting process is somewhat more involved, and handled with a separate function prepCox. The formula syntax is the same as that of coxph:

```
> c1.cox <- prepCox(Z=Z1, Surv(time, status) ~ bmi + strata(sex),
+ SNPInfo = SNPInfo, data =pheno1)</pre>
```

Regardless of model choice, the scores are meta-analyzed in exactly the same way; no modification is necessary of the skatMeta function:

# 5 Filtering

Sometimes it will be desirable to remove SNPs from the analysis, based on minor allele cutoffs or other criteria. Some 'static' criteria, such as SNP annotation, can be handled by subsetting the SNP Info file given to skatMeta, burdenMeta or singlesnpMeta.

A common filtering criteria is minor allele frequency, which in general depend on the individuals contributing to an analysis. For this reason, skatMeta, burdenMeta or singlesnpMeta have an option mafRange which includes SNPs within a specified range of minor allele frequencies. For instance, if one only wishes to use only variants with MAF between 0 and 0.05, one would perform the previous analysis by

```
> meta.results <- skatMeta(c1, c2, SNPInfo = SNPInfo,
+ aggregateBy = "gene", mafRange = c(0,0.05))
> head(meta.results)
```

```
Qmeta
                                       cmaf nmiss nsnps
     gene
1
    gene1 0.858651041
                       97982.36 0.2050000
                                              600
                                                     15
  gene10 0.385906585 236430.85 0.3112500
                                             1200
                                                     21
3 gene100 0.310106989 201665.04 0.2537500
                                              600
                                                     16
   gene11 0.348914794 298153.58 0.3766667
                                             1800
                                                     26
   gene12 0.005833266 364830.51 0.2175000
                                                0
                                                     15
   gene13 0.716506668 193644.56 0.3095833
                                              600
                                                     22
```

Note that the number of SNPs now included is lower than the previous analysis.

## 6 Conditional analyses

In many cases, it will be desirable to adjust for SNPs in a conditional analysis. This can be done in the package with the function prepCondScores. Note that these adjustments are required by the individual studies, and are not performed at the meta-analysis. The syntax and output are identical to prepScores, but requires the additional argument adjustments. The adjustments should be in the same format as the SNPInfo file (i.e. containing a 'gene' and 'Name' column specifying SNP names, or designated alternative columns), and specifies which genes to adjust for which SNPs. For instance, in the below example, we condition 'gene1' analyses on the SNPs labeled 1000001, 1000002, and 1000003, 'gene2' on SNP 1000020, and 'gene13' on SNP 1000100.

```
> adjustments \leftarrow SNPInfo[c(1:3, 20,100), ]
> adjustments
       Name
              gene
    1000001
1
             gene1
2
    1000002
             gene1
3
    1000003
             gene1
    1000020 gene10
100 1000100 gene13
> ####run on each study:
> c1.adj <- prepCondScores(Z=Z1, y~sex+bmi, SNPInfo = SNPInfo,
          adjustments=adjustments, data =pheno1)
 c2.adj <- prepCondScores(Z=Z2, y~sex+bmi, SNPInfo = SNPInfo,
          adjustments=adjustments, kins=kins, data=pheno2)
> SNPInfo.sub <- subset(SNPInfo, (SNPInfo$gene %in% adjustments$gene) &
+
          !(SNPInfo$Name %in% adjustments$Name) )
> #skat
> out.skat <- skatMeta(c1.adj,c2.adj, SNPInfo = SNPInfo.sub)</pre>
> head(out.skat)
                        Qmeta
                                   cmaf nmiss nsnps
    gene
                 p
1 gene1 0.7638026 86158.87 0.1679167
                                           600
                                                  12
                                                  20
2 gene10 0.3235175 235903.90 0.2933333
                                          1200
                                                  21
3 gene13 0.7966422 167332.89 0.2920833
                                           600
```

The output of prepCondScores have class prepScores, and contain *only* the genes listed in the adjustments argument. Any of the meta-analysis functions can be applied to these objects as before. In the above example, we performed SKAT, using the subset of the full SNPInfo file relevant to the conditional analyses.

# 7 Parallelization

In most cases, results for individual studies can be computed for whole-exome and exome-chip on several thousand subjects in a few minutes, without use of specialized data structures or parallel processing. However, if either memory limitations or processing speed make this cumbersome, we provide a generic concatenation function c(...) to combine multiple prepScores objects. This can be employed to break up computation over, e.g. chromosomes. For instance, we can separate computation for the first study over the first and second set of 50 genes. We perform this example in sequence, but it would be a simple matter to run each subset of genes in parallel.

```
> ##subset SNPInfo file to first 50 genes, and second 50 genes:
> SNPInfo1 <- subset(SNPInfo, SNPInfo$gene %in% unique(SNPInfo$gene)[1:50])
> SNPInfo2 <- subset(SNPInfo, !(SNPInfo$gene %in% unique(SNPInfo$gene)[1:50]))
> ##subset corresponding genotype files:
> Z1.1 <- subset(Z1, select = colnames(Z1) %in% SNPInfo1$Name)
> Z1.2 <- subset(Z1, select = colnames(Z1) %in% SNPInfo2$Name)
> ##run prepScores separately on each chunk:
> c1.1 <- prepScores(Z1.1, y~bmi+sex, SNPInfo = SNPInfo1, data = pheno1)
> c1.2 <- prepScores(Z1.2, y~bmi+sex, SNPInfo = SNPInfo2, data = pheno1)
> ##combine results:
> c1 <- c(c1.1, c1.2)
> class(c1)
[1] "seqMeta"
```

We do not provide a method for computing the meta-analysis in parallel. Individual prepScores objects are usually under 20 megabytes, and can be processed sufficiently quickly that parallelization is typically not warranted.

## 8 Method details

In this section we describe the method used in the meta-analysis. The theory for meta-analyzing score tests can be found in e.g. Lin and Zeng (2010).

We first focus on continuous and binary outcomes; modifications for family data and survival outcomes are described subsequently. For study k, denote  $y^{(k)}$  as the n-vector of outcomes,  $\hat{y}^{(k)}$  as the corresponding vector of fitted values under the null-model (i.e. without genotype), and  $G^{(k)}$  as the  $n \times p$  matrix of genotypes in one gene, and  $X^{(k)}$  as the  $n \times q$  matrix of adjustment variables.

#### 8.1 Study level functions

For each gene under consideration, prepScores computes the vector of scores  $U^{(k)}$  and their corresponding variances  $V^{(k)}$ . For unrelated individuals these are given by

$$U^{(k)} = G^{(k)T}(y^{(k)} - \hat{y}^{(k)}) / \sigma_k^2 \in \mathbb{R}^p$$

$$V^{(k)} = (W^{(k)}G^{(k)})^T (I - H^{(k)}) W^{(k)}G^{(k)} / \sigma_k^2 \in \mathbb{R}^{p \times p},$$

where  $\sigma_k^2$  is the the residual variance for continuous data and is 1 for binary data,

$$W^{(k)} = \begin{cases} \operatorname{diag}(\sqrt{\hat{y}^{(k)}(1-\hat{y}^{(k)})}) & \text{for binary data} \\ I_n & \text{for continuous data} \end{cases},$$

and

$$H^{(k)} = X^{(k)} W^{(k)} ((W^{(k)} X^{(k)})^T W^{(k)} X^{(k)})^{-1} (W^{(k)} X^{(k)})^T,$$

is a projection matrix. These can easily be obtained from the output of the glm function.

The prepScores function computes these for each gene under consideration, and concatenates them in a list. If genotype is missing, it is imputed to the study-specific minor allele frequency. This is done one gene at a time as scores and information are computed, rather than modifying the genotype argument Z, which prevents R! from copying potentially large genotype matrices. In addition, the sample size and minor allele count are also stored.

## 8.2 Meta-analysis

Each function skatMeta, burdenMeta, and singlesnpMeta first begins by meta-analyzing the scores and their variances across all studies, for each unique gene name in the SNPInfo file. This follows the simple formulas

$$U = \sum_{k} U^{(k)}, \quad V = \sum_{k} V^{(k)}.$$

Under the null hypothesis of no genetic effect, we have that

$$V^{-1/2}U \rightarrow_d N_p(0, I_p)$$

or that U is approximately  $N_p(0, V)$ 

Each of the tests statistics computed computed (SKAT, burden, and single snp) are functions of the vector U.

**skatMeta** For a vector of weights  $w = (w_1, \dots, w_p)^T$  denote  $R = \text{diag}(\sqrt{w_1}, \dots, \sqrt{w_p})$  The SKAT statistic Q is given by

$$Q = \sum_{j=1}^{p} w_j U_j^2 = ||RU||_2^2,$$

where  $S_j = \sum_k g_j^{(k)T}(y^{(k)} - \hat{y}^{(k)})/\sigma_k^2$  and  $g_j^{(k)}$ , is the genotype for SNP j in study k. Asymptotically

$$Q \sim \sum \lambda_i \chi_1^2$$
,

where the  $\lambda_i$  are the eigenvalues of RVR. The distribution function of a sum of eigenvalues can be approximated using the function pchisqsum in the survey package (Lumley, 2004). By default, we use method= 'saddlepoint' for the saddle point approximation, which is implemented in pure R. A slightly faster option is the Davies method, which inverts the characteristic function, and requires the CompQuadForm package (Duchesne and de Micheaux, 2010).

**burdenMeta and singlesnpMeta** The computation for burden tests is much more straightforward. For a weight vector  $w = (w_1, \dots, w_p)^T$ , the vector of burdens is given by  $G^{(k)}w$ , and the score for its association by  $w^T G^{(k)T}(y^{(k)} - \hat{y}^{(k)})/\sigma_k^2$ . Summing the scores across studies, we get the test statistic

$$T = \sum_{k} w^{T} G^{(k)T} (y^{(k)} - \hat{y}^{(k)}) / \sigma_{k}^{2} = w^{T} U$$

which is approximately  $N_1(0, w^T V w)$ . The function singlesnpMeta is a special case of this, where  $w_j = 1$  for the variant of interest, and is otherwise zero.

In order to approximate the coefficients and standard errors, we note that the first step of the Fisher scoring algorithm estimates the coefficients and their standard errors from the first and second derivatives of the likelihood, which are given by the score and its variance. We plug in the meta-analyzed score and variance to obtain the one-step approximations

$$\hat{\beta} = \frac{w^T U}{w^T V w}$$
, and  $\hat{se}(\hat{\beta}) = \frac{1}{\sqrt{w^T V w}}$ .

If we apply the Wald test to these coefficients, the resulting  $\chi^2$  test statistic is exactly the  $\chi^2$  test statistic for the score test.

#### **SKAT-O** Denote the SKAT-O statistics

$$Q_o(\rho) = (1 - \rho) ||UR||_2^2 + \rho(w^T U).$$

Our goal is to find the value of  $\rho$  which gives the smallest p-value among a sequence of  $\rho$ 's (given in **rho**), and correct for the flexibility afforded by this choice.

Denote the sequence of  $\rho$ 's  $0 \le \rho_1 < \cdots < \rho_q \le 1$  at which we calculate p-values for  $Q_o(\rho)$ . In the simplest case, we can choose  $\rho_1 = 0$  and  $\rho_2 = 1$  to perform the SKAT and burden tests. We then calculate the p-values for each of these tests. For fixed  $\rho$ ,  $Q_o(\rho_i)$  follows a mixture of  $\chi_1^2$  variables, where the mixing parameters are given by the eigenvalues of  $W^{1/2}VW^{1/2}$ , where  $W = ((1-\rho) \cdot RR + \rho \cdot ww^T)$ . To calculate  $W^{1/2}$  we use the eigenvalue decomposition of the matrix  $((1-\rho) \cdot RR + \rho \cdot ww^T)$ .

Now, let  $p_i$  be the p-value for  $Q_o(\rho_i)$ . We then select the minimum p-value  $p_{min} = \min\{p_1, \dots, p_q\}$ . To find the actual p-value we wish to report, we must compute the probability that of observing a  $p_{min}$  smaller than what was observed under the null hypothesis. Unfortunately, it is difficult to directly write down a distribution of  $p_{min}$  under the null hypothesis using well-known distributions. However, the distribution of  $p_{min}$  is numerically tractable if we break the distribution of  $Q_o(\cdot)$  into the conditional distribution of the SKAT statistic given the value of the burden test, and then average over values of the burden test. In order to do this, first note that if  $T_i$  is the  $1 - p_{min}$  quantifier of the distribution of  $Q_o(\rho_i)$ , we know that the test using  $\rho_i$  will give a p-value less than  $p_{min}$  when  $(1 - \rho_i) ||UR||_2^2 + \rho_i(w^T U)^2 > T_i$ , i.e. when

$$||UR||_2^2 > \frac{T_i - \rho_i(w^T U)^2}{1 - \rho_i}.$$

Thus, given the value of the burden test, we observe a smaller p-value than  $p_{min}$  when

$$||UR||_2^2 > \min_{i=1...q} \left\{ \frac{T_i - \rho_i(w^T U)^2}{1 - \rho_i} \right\}.$$

So, it suffices to find the conditional distribution of UR given  $w^TU$ , calculate  $Pr(\|UR\|_2^2 > \min_{i=1...q} \left\{ \frac{T_i - \rho_i(w^TU)^2}{1 - \rho_i} \right\}$ ), and integrate it over the distribution of  $w^TU$ . Since UR and  $w^TU$  are linear combinations of a Normal vector, we can write down their joint distribution. Standard algebra gives that

$$UR \mid \{w^T U = a\} \sim N_p \left(\frac{RVw}{w^T Vw}a, \quad R^T VR - \frac{RVww^T VR}{w^T Vw}\right).$$

We can again determine the distribution of  $||UR||_2^2$  using normal quadratic forms. These differ slightly from the calculations used in SKAT, due to the presence of the mean vector  $\frac{RVw}{w^TVw}a$ .

Instead, the distribution of  $||UR||_2^2 | \{w^TU = a\}$  is the sum of non-central  $\chi^2$  distributions, whose distribution functions are available in any of the functions in the **CompQuadForm** package (Duchesne and de Micheaux, 2010). For speed, we recommend method=`integration', which first attempts the Davies method, and in the case where this gives an error, it uses the farebrother method. Using method=`saddlepoint' is slower, but has higher relative accuracy.

Thus, if we denote  $\phi(\cdot)$  as the marginal distribution of  $w^T U$ , which is  $N(0, w^T V w)$ , we can find the true significance of  $p_{min}$  using the following algorithm

- 1. For each  $\rho_i$ , calculate  $p_i$  based on the quadratic forms  $Q_o(\rho_i)$ .
- 2. Calculate  $p_{min} = \min\{p_1, \dots, p_q\}$ .
- 3. Calculate  $T_i$ 's, the  $(1-p_{min})$  quantiles of the  $Q_o(\rho_i)$ . Denoting  $F_{Q_i}$  as the distribution function of  $Q_o(\rho_i)$  found in Step 1, we do this by solving  $F_{Q_i}(x) = p_{min}$  for x using uniroot. Potential inaccuracies here result in errflag=2.
- 4. Form the conditional distribution  $UR \mid \{w^T U = a\}$ , and the conditional probability

$$f_S(a) = Pr\left(\|UR\|_2^2 > \min_{i=1...q} \left\{ \frac{T_i - \rho_i(w^T U)^2}{1 - \rho_i} \right\} \mid w^T U = a \right).$$

5. Numerically integrate

$$p_{actual} = \int_{-\infty}^{\infty} f_S(a)\phi(a)da$$

and report  $p_{actual}$ .

## 8.3 Family data

This framework extends easily to the case of family data. A more detailed description of this method is given in Chen *et al.* (2013). In the pooled-data analysis, the related individuals would have a different form of the likelihood, corresponding to a mixed-model. But, since likelihoods are on the same scale, the scores from related and un-related data can be combined as usual. We can think of all studies using a mixed-model, but where un-related individuals have no random effects.

Here, instead of outcomes y having variance  $\sigma_y^2 I_n$ , they have variance  $\sigma_y^2 \Omega = \sigma_y^2 (\theta_1 \Theta + \theta_2 I_n)$ , where  $\Theta$  is twice the kinship matrix, and  $\theta_1 + \theta_2 = 1$ . In unrelated individuals  $\Theta = I_n$ . The scores for this model are

$$U^{(k)} = G^{(k)^T} \Omega^{-1} (y^{(k)} - \hat{y}^{(k)}).$$

If the matrix of covariates is X, the variance of the score is

$$V^{(k)} = G^{(k)T} \left( \Omega^{-1} - \Omega^{-1} X (X^T \Omega^{-1} X)^{-1} X^T \Omega^{-1} \right) G^{(k)}.$$

We calculate these using the lmekin function in the **coxme** package (Therneau, 2012). These can be combined as usual with scores and variances from other studies.

Unfortunately, the score test for generalized linear mixed models does not exist in closed form, and no available implementation of generalized linear mixed models allows large used-defined variance components, such as a kinship matrix. We therefore leave the extension of binary traits for family data for future work.

#### 8.4 Survival outcomes

We apply a similar procedure to obtain test statistics for Cox proportional hazards regression. In this case, the score test is anti-conservative, so we instead use a signed likelihood ratio statistic Lumley and Scott (2013). Specifically, let  $S_j^2$  is the likelihood ratio test statistic for the including of the single variant j in the null model,  $\beta_j$  be the corresponding maximum likelihood estimate, and V be the variance of the usual score test. we use

$$U_j = \operatorname{sign}(\beta_j) S_j / \sqrt{V_{jj}},$$

which is asymptotically equivalent to the score test, but enjoys superior finite sample performance. In order to implement this procedure efficiently, we use a modified version of the coxph.fit function in the survival package (Therneau, 1999).

## 9 Summary

In this paper, we have described the basic usage and syntax of the **seqMeta** R package. This includes functions for computing study level results with the functions **prepScores** for continuous and binary traits and **prepCox** for survival outcomes. We also describe the functions that pool study level results for meta-analysis, including **skatMeta**, **burdenMeta**, **skatOMeta** and **singlesnpMeta**.

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