Computing q-values conditioned on covariates using swfdr

1 May 2024

Contents

1	Introduction	2
2	Installation	2
3	q-values without covariates	3
	3.1 Conceptual datasets	3
	3.2 pi0	5
	3.3 q-value	6
4	Conditioning on covariates	8
	4.1 Conditioned pi0.	В
	4.2 Conditioned q-values	9
5	Conditioning on multiple covariates	0
	5.1 Conditioned pi0s with multiple covariates	1
	5.2 Conditioned q-values with multiple covariates	2
	5.3 Comparison with qvalue	2
6	Discussion	4
Α	Technical notes	4
	A.1 Modeling resolution (argument Lambda)	5
	A.2 Covariate matrix (argument X)	5
	A.3 Model type (argument type)	5
	A.4 Thresholding (argument threshold)	5
	A.5 Smoothing (argument smoothing)	6
	A.6 Smoothing degrees of freedom (argument smooth.df) 10	6
	References	6

1 Introduction

Multiple hypothesis testing is a common concern in modern data science workflows: after performing several - in some cases, thousands - statistical tests on a dataset, many of the resultant p-values are expected to be small purely by chance. So it is imperative to interpret the results in a manner that limits spurious associations. At the same time, a parallel requirement is to identify significant hits despite having performed many tests.

In addition to the classic multiple hypothesis testing problem described above, analyses of complex datasets are also complicated by covariates. Given p-values calculated using a particular statistical test, background knowledge may suggest that the test may be more appropriate for some data instances than for others, or that some results may be more or less expected a-priori. A tempting analysis strategy in such situations might be to partition the dataset and to study each part separately. However, direct stratification is not trouble-free. It relies on splitting data into discrete groups, which can introduce challenges of its own, including in the choice of appropriate thresholds. It also leads to further statistical testing, which is redundant and magnifies the burden of multiple hypothesis testing. Thus, an alternative, more systematic, approach is needed to utilize background knowledge in statistical analyses.

There are, in principle, many possible ways to use covariate variables when tackling the multiple hypothesis testing problem. This vignette does not attempt an unbiased exposition of the available methods; some of them have been summarized and benchmarked by Korthauer et al. (2018). Instead, this vignette provides a practical guide to using the swfdr package, which implements algorithms for modeling the effect of covariates on a quantity called π_0 , which can be interpreted as the probability that a null hypothesis is true. The theoretical aspects and modeling details are provided by Jager and Leek (2013) and Boca and Leek (2018). The approach extends methods by Storey (2002) and Storey and Tibshirani (2003). In practice, these methods take in a set of p-values and output another set of quantities, q-values, that can be interpreted in the context of a multiple hypothesis analysis.

The vignette begins with notes on how to install the swfdr software. Next, it provides a review of q-values using the classic qvalue package. This section also introduces some conceptual datasets and motivates the use of covariates to augment standard q-value analysis. Subsequent sections explain how to condition q-values on covariates using swfdr. The main points are summarized in the discussion and some technical details are discussed in an appendix.

2 Installation

The swfdr package can be installed via the Bioconductor package manager, Morgan (2018). If you don't already have the package manager, you'll first have to install it.

```
install.packages("BiocManager")
```

This might show a prompt to select a package repository, and you can choose any one that seems convenient for you.

To perform the package installation, load the Bioconductor manager and request to install swfdr.

```
library("BiocManager")
BiocManager::install("swfdr")
```

After the above command completes, we can load the package.

library(swfdr)

For this vignette, we will also use Bioconductor package qvalue, Storey et al. (2018). This can be installed also via Bioconductor

BiocManager::install("qvalue")
library(qvalue)

3 q-values without covariates

Suppose we have measurements from two groups. To perform a statistical analysis, we would formulate a null hypothesis, compute a p-value against that hypothesis, and then use some criteria to call the result as significant or not. The practical details may vary from application to application. For example, a common use case is to assume that measuremets come from distributions with equal variances, formulate the hypothesis that the distributions have equal means, and compute p-values using a t-test. An interpretation strategy might consist of calling a result a 'hit' or 'significant' if the p-value is below a threshold, often 0.05.

In this section, we will look at how to interpret results from a large set of such statistical tests and control the rate of false discoveries using q-values. While the technical definitions for p-values and q-values are given in textbooks and original references, it is useful to highlight some of their properties here. Consider a statistical test and an associated numerical result v. The interpretation with respect to p-values and q-values would be as follows (Storey and Tibshirani (2003)).

- *p-value:* Assuming that the original data is consistent with the null hypothesis, the probability that the single result with *p* = *v* is a false positive is *v*.
- *q-value:* Among the tests assigned q-values below v during an analysis, the proportion
 of false positives is v.

There are some important conceptual and practical differences between these definitions.

- The interpretation of the p-value is focused on a single test instance, so any adjustments motivated by multiple hypothesis testing must be performed *post-hoc*. In contrast, the q-value is defined relative to an ensemble of tests and thus intrinsically refers to a multiple-hypothesis context.
- The p-value is defined from the perspective of a hypothetical ensemble of data that is
 consistent with the null hypothesis. The q-value is instead formulated with respect to
 the ensemble of performed measurements. (Technically, the properties of the q-value are
 valid in the limit of a large ensemble of measurements, but it is possible to replace that
 hypothetical ensemble by an actual set of measurements and thus work with q-value
 estimates.)

3.1 Conceptual datasets

To demonstrate calculations of q-values, let's work with some practical datasets. To cover various scenarios, let's define three synthetic datasets and some helper functions. Each dataset will consist of a moderate number of examples, i.e. instances requiring a statistical test.

N < -1000

In the first dataset, dataset0, let's consider two groups with three measurements each and formulate a null hypothesis that there is no difference in means between the two groups. Moreover, let's construct the data so that all measurements come from the same generative process. In other words, despite fluctuations in individual measurements, all instances are concordant with the null hypothesis.

```
set.seed(12345)
dataset0 <- matrix(rnorm(6*N), ncol=6)
head(dataset0, 3)
## [,1] [,2] [,3] [,4] [,5] [,6]
## [1,] 0.5855288 1.67751179 -0.6078411 0.5098921 -1.0426910 -0.6924722
## [2,] 0.7094660 0.07947405 1.0762231 -0.7153643 -0.1839484 1.1246956
## [3,] -0.1093033 -0.85642750 -0.5764258 -0.4055215 0.1792667 0.1745351</pre>
```

From this data, we can compute p-values for each set of measurements (rows) using a t-test. Because a similar calculation will be required again later, it is best to define a helper function.

```
# compute t-tests on each row of a matrix
dataset_p <- function(D, groupl=c(1,2,3), group2=c(4,5,6)) {
   t.test.groups <- function(x) {
     t.test(x[group1], x[group2])$p.value
   }
   round(apply(D, 1, t.test.groups), 5) # rounding for legibility
}
D0 <- data.frame(p=dataset_p(dataset0), truth=0)
summary(D0$p)
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.00527 0.26459 0.48864 0.50197 0.75112 0.99992</pre>
```

After the helper function, the above block defines a data frame with p-values and an integer code truth; we will use this code later. The p-values range from 0.00527 to 0.99992. Thus, although by construction all instances in dataset0 are consistent with the null hypothesis, some p-values are nominally quite small.

For the second dataset, dataset1, let's again consider two groups of three measurements each, but give one the groups a shifted mean.

```
dataset1 <- matrix(c(rnorm(3*N, 0, 1), rnorm(3*N, 2, 1)), ncol=6)</pre>
head(dataset1, 3)
##
              [,1]
                         [,2]
                                    [,3]
                                             [,4]
                                                       [,5]
                                                                   [,6]
## [1,] -0.3487804 -0.6790374 -1.486991 1.180370 2.1612775 2.7892407
## [2,] -1.0257007 -0.5723894 -2.205152 3.375843 2.5319214 -0.4168186
## [3,] -0.9074263 0.8232486 1.405955 1.701138 0.7020966 1.5649241
D1 <- data.frame(p=dataset_p(dataset1), truth=1)</pre>
summary(D1$p)
##
     Min. 1st Qu. Median
                              Mean 3rd Qu.
                                               Max.
## 0.00009 0.03287 0.07704 0.13351 0.17774 0.97222
```

The resulting p-values span a similar range as before, but are skewed toward lower values. It is important to note that although all instances are actually inconsistent with the null hypothesis, some p-values are nominally large.

To conclude this section, let's create a final dataset that is a concatenation of the previous two. We will only need the p-values, so we can omit creating the raw data and only define the summary data frame.

```
D2 <- rbind(D0, D1)
dim(D2)
## [1] 2000 2
table(D2$truth)
##
## 0 1
## 1000 1000</pre>
```

This combined dataset has 2000 instances. The first half is concordant with the null hypothesis (truth equal to zero) and the second half is not (truth equal to 1).

3.2 pi0

Given p-values from individual statistical tests, we can begin looking at their properties as a set of multiple hypotheses. Let's visualize the distributions of p-values.

```
par(mfrow=c(1,3), las=1, mar=c(3,4,2,1))
p_hist <- function(p, breaks=20, freq=F, ylim=c(0, 7.2), ...) {
    hist(p, breaks=breaks, freq=freq, ylim=ylim, xlab="", ...)
}
p_hist(D0$p, col="#6666666", main="p-values in D0")
p_hist(D1$p, col="#cccccc", main="p-values in D1")
p_hist(D2$p, col="#999999", main="p-values in D2")</pre>
```



In the first histogram based on D0, the distribution of p-values is flat. This is the expected pattern for data consistent with the null hypothesis. In the second panel based on dataset D1, the distribution is skewed and indicates deviation from the null hypothesis. The last histogram, which is based on the composition of D0 and D1, shows an intermediate skew.

Thus, the shape of the histogram is an indicator for the extent to which a dataset is consistent with the null hypothesis. This intuition can be codified into a probability that the null hypothesis is true, denoted as π_0 or pi0. It is computed from the histogram by considering the number of test instances that give p-values below various thresholds, $p < \lambda$, where λ might be 0.05, 0.1, etc. See e.g. Storey (2002) and Storey and Tibshirani (2003) for details. We can compute π_0 for each dataset using the qvalue package.

```
library(qvalue)
D0q <- qvalue(D0$p)
D1q <- qvalue(D1$p)
D2q <- qvalue(D2$p)
c(D0=D0q$pi0, D1=D1q$pi0, D2=D2q$pi0)</pre>
```

D0 D1 D2 ## 1.00000000 0.01388784 0.51395921

The π_0 estimate for D0 is unity, which is consistent with how we generated the dataset. For D1, π_0 is much lower. At 0.014, it is greater than the true value (zero; none of the instances in D1 were generated in a way consistent with the null hypothesis). However, this estimate is nonetheless closer to the truth. Unsurprisingly, the last π_0 estimate - for the combined dataset D2 - lies between the previous values.

3.3 q-value

While π_0 is an aggregate summary of an entire dataset, we are often interested in assigning significance calls to individual test instances. The q-value is a measure that controls for false positive calls and internally uses the π_0 to allow for true positive calls.

We already computed q-values when we ran the function qvalue in the previous section. We can display a summary for the D0 dataset as follows.

```
summary(D0q)
##
## Call:
## qvalue(p = D0$p)
##
## pi0: 1
##
## Cumulative number of significant calls:
##
##
             <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
                                                      <1
                  0
                         0
                            3 15
                                           36
## p-value
                                                90 1000
## q-value
                  0
                         0
                               0
                                      0
                                            0
                                                 0 1000
## local FDR
                  0
                         0
                               0
                                      0
                                            0
                                                 0
                                                      0
```

The table displays hit counts at various thresholds. To simplify the discussion, let's just look at the rows labeled 'p-value' and 'q-value' and the threshold of 0.05. For the following, a helper function will be useful to extract a subset of the relevant results.

```
# extract number of hits with values below a threshold
hit_counts <- function(x, threshold=0.05) {
   counts <- c(sum(x$pvalues<threshold), sum(x$qvalues<threshold))
   names(counts) <- paste0(c("p<", "q<"), threshold)
   counts
}
hit_counts(D0q)
## p<0.05 q<0.05
## 36 0
```

If we were to call significance using unadjusted p-values at a threshold of 0.05, we would have 36 hits. Because we explicitly constructed the dataset to be consistent with the null hypothesis, all of these hits would be false positives. In contrast, the number of hits using the q-value would drop to 0, which in this case is the appropriate answer.

As an aside, note that if we were to adjust p-values for multiple-hypothesis testing say with a Bonferroni correction, the number of hits would also drop to 0. To check that, evaluate sum(D0\$p < 0.05/N).

Let's move on to the second dataset, in which all instances are inconsistent with the null hypothesis.

```
hit_counts(D1q)
## p<0.05 q<0.05
## 365 1000</pre>
```

Here, the number of hits obtained by thresholding p-values is higher. However, at 365, it leaves a large number of instances of the dataset as false negatives. In contrast, the procedure based on q-values captures the entire dataset. Again, this is a satisfactory result for this conceptual dataset.

Continuing the previous side note, how many hits would we have using a Bonferroni correction? The answer is 0 hits, thus producing an even larger number of false negatives. This shows that procedures that use adjustments of p-values are quite distinct from q-values.

Finally, let's consider the combined dataset.

```
hit_counts(D2q)
## p<0.05 q<0.05
## 401 0</pre>
```

The number of hits based on p-values is simply the sum of the previous two results, 401 = 36 + 365. To look at this result from a different perspective and in preparation for subsequent calculations, we can display this information as a confusion matrix comparing expected versus computed hit counts.

}

We can now get a summary of performance based on thresholding on p-values.

The 2000 instances are split into true positives (bottom-right), false-positives (top-right), true negatives (top-left), and false negatives (bottom-left). Overall, there are many false negatives and quite a few false positives.

The confusion matrix based on q-values is as follows.

```
confusion_matrix(D2$truth, D2q, "q")
## q<0.05
## truth FALSE
## 0 1000
## 1 1000</pre>
```

There are no positive hits - we have lost all signal. However, we know that calling hits based on q-values produced correct result based on data from D0 and D1 separately. Thus, it might be possible to improve the calculation for the joint dataset. This leads to using background knowledge to condition calculations of π_0 and q-values in the next section.

4 Conditioning on covariates

In addition to a core set of data, many datasets come with additional descriptive variables - metadata. While these descriptive variables might not enter into a primary statistical analysis, it may nonetheless be fruitful to incorporate them in a larger analysis pipeline. Although many numerical recipes are possible in this direction. Package swfdr provides two functions, lm_pi0 and lm_qvalue, that use linear models to condition estimates for π_0 and q-values.

4.1 Conditioned pi0

Following from the datasets and examples of the previous section, let's compute π_0 estimates for D2 conditioned on the dataset of origin.

D2_lm_pi0 <- lm_pi0(D2\$p, X=D2\$truth)</pre>

The output is a composite object (list) with several components.

names(D2_lm_pi0)
[1] "call" "lambda" "X.names" "pi0.lambda" "pi0"

For some follow-up calculations, we will retrieve data from these components. However, it is instructive to first examine a summary of the object, via summary(D2_lm_pi0) or print(D2_lm_pi0) or by simply executing it in the console.

D2_lm_pi0

This should produce output with multiple sections. The first few summarize technical details.

```
##
## Call:
## lm_pi0(p = D2$p, X = D2$truth)
##
## lambda:
## (Length)
                                               Max
                  Min
                           Mean
                                  Median
##
         19
                 0.05
                            0.5
                                      0.5
                                              0.95
```

The top part reiterates the function call and the section on Lambda provides details on how the results were estimated, see Storey (2002) and Storey and Tibshirani (2003) for details. The next section summarizes the covariates.

Computing q-values conditioned on covariates using swfdr

##
covariates:
(Length)
1

In this case, there is only one covariate. The last section summarizes π_0 estimates.

pi0: ## (Length) Min Mean Median Max ## 2000 0.0177 0.508 0.508 0.9983

In contrast to the π_0 estimate from function qvalue, we now have one estimate per instance in the dataset, ranging from 0.018 to 0.998. Internally, the function used logistic regression to estimate how the distribution of input p-values depends on the covariate, see Boca and Leek (2018) for details.

4.2 Conditioned q-values

Let's now compute q-values conditioned on the same covariate.

```
D2_lm_q <- lm_qvalue(D2$p, X=D2$truth)</pre>
```

Again, the output is a composite object. A summary is available by evaluating it in the console.

D2_lm_q

The majority of the output is analogous to what was displayed for D2_lm_pi0. The new section summarizes hit counts at various thresholds.

##							
##	Cumulative	number	of signi	ficant o	calls:		
##		<1e-4	<1e-3	<0.01	<0.05	<0.1	<1
##	p-value	1	7	96	401	671	2000
##	q-value	Θ	0	913	1000	1000	2000

In this case, we have 1000 hits based on q-values at a threshold of 0.05. We can further compute a confusion matrix.

This reveals a strong presence of true positives with 0 false positives and 0 false negatives. The analysis of the combined dataset with covariate thus reproduces the performance we obtained when we analyzed D0 and D1 separately.

Of course, the above calculation is idealized in that the covariate perfectly stratifies the instances. To finish this section, we can repeat the calculation in a more difficult setting where the covariate carries imperfect signal. To this end, let's define an object similar to D2.

```
D2_noisy <- data.frame(p=D2[, "p"],
truth.noisy=D2[, "truth"]+rnorm(2*N, 0, 0.4))
head(D2_noisy, 3)
## p truth.noisy
## 1 0.30806 0.3076642
## 2 0.44155 -0.4250800
## 3 0.16487 -0.4386329
```

The new variable truth.noisy is now a real number based on the previous truth, but carries considerable noise. We can repeat the lm_qvalue calculations using this as a covariate.

D2_noisy_lm_q <- lm_qvalue(D2_noisy\$p, X=D2_noisy\$truth.noisy)</pre>

The hit summary for this object is as follows.

##							
##	Cumulative	number	of signit	ficant ca	alls:		
##		<1e-4	<1e-3	<0.01	<0.05	<0.1	<1
##	p-value	1	7	96	401	671	2000
##	q-value	Θ	Θ	5	442	837	2000

Let's inspect the confusion matrix at various thresholds.

```
confusion_matrix(D2$truth, D2_noisy_lm_q)
##
       q<0.05
## truth FALSE TRUE
##
   0 999 1
        559 441
##
      1
confusion_matrix(D2$truth, D2_noisy_lm_q, threshold=0.1)
##
       q<0.1
## truth FALSE TRUE
      0 976
              24
##
##
      1 187 813
```

In each case, the confusion matrix reveals a moderate or large number of true positives with a limited number of false positives. Thus, even in this noisy setting, we have a high hit rate while restraining false discovery.

Although the last example introduced some noise, all calculations in this section are still based on synthetic data and are thus quite artificial. We have also considered analyses based on just one covariate. Nonetheless, the examples are instructive because the analytical workflow and syntax also apply in more complex scenarios, as demonstrated in the next section.

5 Conditioning on multiple covariates

The package comes with a prepared dataset, BMI_GIANT_GWAS_sample, that summarizes a portion of a genome-wide association study by Locke et al. (2015). The dataset holds associations between simple-nucleotide polymorphisms (SNPs) and body-mass-index (BMI).

```
GWAS <- BMI_GIANT_GWAS_sample
dim(GWAS)
## [1] 50000 9</pre>
```

```
head(GWAS)
## # A tibble: 6 x 9
##
    SNP
             A1
                    A2
                         Freq_MAF_Hapmap
                                              b
                                                    se
                                                          p
                                                                 Ν
##
    <chr>
              <chr> <chr> <dbl>
                                          <dbl> <dbl> <dbl> <dbl>
## 1 rs10510371 T C
                                0.025
                                          0.0147 0.0152 0.334 212965
## 2 rs918232 A
                    G
                                0.342 -0.0034 0.0037 0.358 236084
## 3 rs4816764 A
                   С
                                 0.00830 0.0163 0.0131 0.213 221771
## 4 rs17630047 A
                 G
                                 0.167
                                         0.0004 0.0048 0.934 236177
## 5 rs4609437 C
                  G
                                 0.25
                                          0.0011 0.0042 0.793 236028
## 6 rs11130746 G
                                 0.233 -0.0006 0.0042 0.886 235634
                   Α
## # i 1 more variable: Freq_MAF_Int_Hapmap <fct>
```

This is a table with 50,000 rows. Column SNP contains an identifier for a genetic polymorphism. Column p contains the p-value from a statistical test linking that polymorphism and BMI. The other columns contain metadata about the polymorphism - see the original article Locke et al. (2015) or the data set annotation (help(BMI_GIANT_GWAS_sample)).

In this vignette, we will use two of the metadata columns, N and Freq_MAF_Hapmap. The first is an integer field conveying the number of patients in the study for which the SNP was measured. This impacts on the power of the individual statistical tests. The second feature is the prevalence of the polymorphism in a background population. This impacts the likelihood that the SNP is associated with an unhealthy phenotype.

5.1 Conditioned pi0s with multiple covariates

The calculation for π_0 proceeds just like in the previous section. In this case, however, argument X should be a matrix or data frame containing both covariates of interest.

GWAS_lm_pi0 <- lm_pi0(GWAS\$p, X=GWAS[, c("N", "Freq_MAF_Hapmap")])</pre>

We can view a summary of the output object by executing it in the console.

```
GWAS_lm_pi0
##
## Call:
## lm_pi0(p = GWAS$p, X = GWAS[, c("N", "Freq_MAF_Hapmap")])
##
## lambda:
## (Length)
                 Min
                         Mean
                              Median
                                             Max
                0.05
                         0.5
                                 0.5
                                            0.95
##
       19
##
## covariates:
## (Length)
          2
##
##
## pi0:
## (Length)
                 Min
                                Median
                                             Мах
                         Mean
##
      50000
              0.7439
                       0.9424
                                 0.9524
                                               1
```

The output structure is by now familiar. One interesting section here summarizes the covariates and confirms that the function used two variables in the calculation.

The final table summarizes the distribution of π_0 values. Recall that π_0 estimate the probability that a null hypothesis is true. Here, the estimates range from 0.7439 to 1. Because the values are all fairly close to unity, they suggest that the test instances are largely concordant with no association between SNPs and BMI. This is an expected property for a GWAS study.

5.2 Conditioned q-values with multiple covariates

Next, we can evaluate whether or not individual instances in the dataset are significant.

```
GWAS_lm_qvalue <- lm_qvalue(GWAS$p, X=GWAS[, c("N", "Freq_MAF_Hapmap")])</pre>
```

We can preview the contents of the output object by executing it on the console.

GWAS_lm_qvalue

Most of the output should be the analogous to that of GWAS_lm_pi0. The new section is a summary of hits at various thresholds.

```
##
## Cumulative number of significant calls:
##
                <1e-4
                        <1e-3
                                 <0.01
                                          <0.05
                                                    <0.1
                                                               <1
                  186
                           405
                                  1388
                                           3771
                                                    6468
                                                           49619
##
     p-value
##
     q-value
                   49
                            70
                                   126
                                            254
                                                     374
                                                           49912
```

We can use our helper function to focus just on the 0.05 threshold.

```
hit_counts(GWAS_lm_qvalue)
## p<0.05 q<0.05
## 3771 254</pre>
```

There are 3771 SNPs that have a nominal p-value lower than 0.05. (That hit count is unadjusted for multiple-hypothesis testing and is likely to include many false positives. If we were to correct for multiple testing via a Bonferroni correction, we would instead impose a threshold of 0.05/50000. The result would be 64 hits.) In contrast, using a q-value threshold, we have 254 hits.

5.3 Comparison with qvalue

In the above calculations, we computed q-values conditioned on two covariates. Let's now see how those results compare with q-values without conditioning. We can compute those by repeating the same steps as before, but without specifying argument X.

```
GWAS_lm_qvalue_0 <- lm_qvalue(GWAS$p)
## Warning: X is missing or NULL - modeling will have no effect</pre>
```

In this calculation, <code>lm_qvalue</code> displays a warning that raises attention to the fact that a function meant to model the effect of covariates is actually used without any covariates at all. This message is useful to detect coding errors, e.g. unintentionally providing an empty object for modeling. Here, omitting the covariates was the intended action, so we can look past the warning and inspect the output.

pi0:

##	(Length)	Min	Mean	Median	Max			
##	50000	0.9564	0.9564	0.9564	0.9564			
##								
##	Cumulative	number	of signif	icant ca	lls:			
##		<1e-4	<1e-3	<0.01	<0.05	<0.1	<1	
##	p-value	186	405	1388	3771	6468	49619	
##		40	60	124	247	254	E0000	

This reveals that although the function computes 50,000 estimates for pi0, all the estimates are equal. The hit table is slightly different than before.

When there are no covariates, the calculations performed by lm_qvalue are conceptually equivalent to those in function qvalue from the qvalue package. We can check this explicitly.

```
GWAS_qvalue = qvalue(GWAS$p)
summary(GWAS_qvalue)
##
## Call:
## qvalue(p = GWAS$p)
##
## pi0: 0.9559508
##
## Cumulative number of significant calls:
##
            <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
##
                                                      <1
                     405 1388
## p-value
               186
                                  2404 3771 6468 49619
                49
## g-value
                       69
                           124
                                   185
                                        247 354 50000
## local FDR
                42
                       50
                            75
                                   107
                                         148 203 27294
```

Apart from the stylistic discrepancies, the final q-value results in the hits table are the same as those output by lm_qvalue. There is, however, a small discrepancy in the estimate for π_0 . This is due to a slight difference in an internal algorithm, which is elaborated upon in the appendix. Here, we note that the discrepancy is <1% so it is likely to be inconsequential in most workflows. However, it is possible to obtain exact parity by calling lm_qvalue with an additional argument smoothing set to smooth.spline. We can verify this by repeating the calculation with this modified setting.

GWAS_lm_qvalue_smooth <- lm_qvalue(GWAS\$p, smoothing="smooth.spline")
Warning: X is missing or NULL - modeling will have no effect</pre>

The same warning appears, but we can again skip it and inspect the output.

```
table(GWAS_lm_qvalue_smooth$pi0)
##
## 0.955950835227913
## 50000
```

This reveals all the π_0 estimates are equal to the output from qvalue - we have exact parity with the value package. This is a useful technical control and it builds confidence in the correctness of the results. However, running this code in the console, there will be a noticeable difference in execution time. Thus, for large datasets, it is more convenient to use the default settings.

6 Discussion

Interpreting statistical results can be challenging, especially when an analysis involves a large number of tests and p-values. When the aim is to call significant hits while controlling the rate of false discoveries, it can be useful to re-frame the interpretation in terms of q-values. The influencial package qvalue provides functions for this purpose, but it is only applicable in contexts when all the statistical tests in an analysis are *a-priori* similar and can be weighted equally. These constraints are at odds with the analysis of complex datasets for which metadata may hold signals about the tested relationships. The swfdr package provides functions for using such metadata as covariates while controlling false discovery rates.

In this vignette, we saw how to install the software and walked through the analysis of some small datasets. These datasets, being synthetically generated, do not capture the nuances and complexities of a real-word analysis. Instead, the hope is that their they capture the value of using q-values for controlling false discovery and the potential benefits of using informative covariates. In particular, they demonstrate the utility of q-values for interpret datasets with a very high proportion of instances that are significant, as well as for modeling with covariates to identify hits when they are surrounded by data that is concordant with the null hypothesis. These are illustrative examples; further evaluations are described in the original article, Boca and Leek (2018), as well as in an independent review, Korthauer et al. (2018), which concluded that swfdr consistently performs well in a range of scenarios.

All the analyses included a discussion of π_0 , the probabilities that a null hypothesis is true, before going on to compute q-values. Given the outputs of functions lm_pi0 and lm_qvalue , it should be apparent that the second calculation is an extension of the first. This duplication of effort is included in the vignette to emphasize the central role π_0 plays in computing q-values and in conditioning on covariates. Practical pipelines, however, may rely on the single function lm_qvalue .

If you use package swfdr, please cite:

- the article describing the method for conditioning false-discovery rates on covariates via π_0 , Boca and Leek (2018)
- the R swfdr software, Leek, Jager, and Boca (2018)

To view the package source, report a bug, or contribute bug fixes, please see the github development page https://github.com/leekgroup/swfdr.

A Technical notes

The main vignette described a complete, coherent workflow for controlling false discovery rates using functions lm_pi0 and lm_qvalue from the swfdr package. The focus was on demonstrating the core features of those functions and on interpreting their outputs. But those same functions also support other arguments which can fine-tune aspects of the calculations.

These additional arguments have default values that should be suitable for common use cases. However, (rare) situations may occur when it may be desirable to adjust the default settings. This section quickly summarizes those options and their implications for the interpretation of the results and computational performance. The arguments are applicable to both lm_pi0 and lm_qvalue, unless specified otherwise.

A.1 Modeling resolution (argument lambda)

As hinted in one of the early sections, estimates for π_0 are computed from the distribution of the input p-values. The details for the algorithm are described e.g. in Storey (2002) and Storey and Tibshirani (2003). Briefly, the calculations take a threshold, say $\lambda = 0.05$, and evaluate the proportion of p-values below that threshold, and then repeat for several thresholds, say $\lambda = 0.1, 0.15, \ldots$. By using such estimates together with a spline fit, the calculations arrive at a single estimate for π_0 . Argument Lambda provides the series of thresholds used in this process.

A large number of thresholds, i.e. a long vector lambda, is needed to allow the spline to fit the trend in proportions in a reliable manner. However, excessive length can be counter-productive if the input set of p-values is short. Long lambda vectors can also increase running time.

In many common applications, the default lambda vector will be appropriate and it is recommended to leave it unchanged. For small datasets, say below 100 items, it may be advisable to provide a shorter vector, e.g. seq(0.05, 0.95, 0.1).

A.2 Covariate matrix (argument x)

One of the key inputs to lm_pi0 and lm_qvalue is an object, X, capturing covariates. When there is only one covariate, X can be specified as a vector. Otherwise, it should be either a matrix or a data frame.

Covariate data is used internally in relatively simple linear models and thus object X cannot contain missing (NA) or non-finite values (NULL, Inf, NaN). Datasets with such components should either omit incompatible parts before processing with swfdr, or impute values.

Covariate data should contain only numeric data. Factors and other categorical data are not supported.

A.3 Model type (argument type)

Modeling the effect of covariates on π_0 is controlled via argument type. The default value for this argument is logistic. In this configuration, functions lm_pi0 and lm_qvalue use logistic regression with one term for each given covariate. The other supported value for the type argument is linear, which uses a standard linear regression model instead.

The run time of linear regression is typically faster than logistic regression. However, linear regression is uninformed about the fact that the output variable is meant to be a probability in a finite range. Thus, logistic is usually the more appropriate approach.

It is recommended to leave type at its default value. The argument is provided primarily for backward compatibility and as a inerface for exploring other modeling approaches.

A.4 Thresholding (argument threshold)

Internally, lm_pi0 and lm_qvalue train models on the input data and then predict individual pi0 estimates. Because these predictions represent probabilities, they are expected to be in the range [0, 1]. However, some predictions may fall outside this range. By default, the functions threshold spurious values and impose the expected lower and upper bounds. Setting argument threshold to FALSE allows the raw prediction values to propagate to the output.

The value of the threshold argument has almost no impact on performance. Changing it from its default value can make interpretation of the output less intuitive, although unusual values can in principle provide insight on the quality of the covariate modeling.

It is recommended to leave threshold at its default value of TRUE. The argument is provided primarily for testing and research purposes, as well as for backward compatibility.

A.5 Smoothing (argument smoothing)

As described above, the calculations internally use splines to model the effect of λ on π_0 . There exist several algorithms for fitting splines, and argument smoothing toggles between two approaches. Both procedures are motivated by the same conceptual reasoning, but differ in the implementation, computational performance, and provided output.

Setting smoothing to smooth.spline instructs the algorithm to use base R's smooth.spline to fit the spline. This is the spline-fitting approach that is used within the qvalue package and in early versions of the swfdr package. In the context of covariate modeling, this approach is, however, computationally inefficient. Setting smoothing to unit.spline shifts the spline calculations to a bespoke implementation that exploits the fact that multiple splines are fitted for the same series of λ and the fact that the objective of the calculation is to obtain a single value π_0 and not the actual smoothing spline. The bespoke approach can reduce the running time many-fold with only small numerical discrepancies with the original algorithm.

It is important to stress that these implementations are provided for performance reasons only; you are welcome to pick the smoothing method of your choice.

A.6 Smoothing degrees of freedom (argument smooth.df)

Spline-fitting can be further tuned by setting the spline degrees of freedom via argument smooth.df. The value of smooth.df must be ≥ 3 to fit p-value distributions such as in the examples in the vignette. Higher values would allow the internal model to accommodate more complex distributions. However, if the p-value distribution deviates substantially from the examples, the premise of the π_0 calculation comes into question. Thus, there is rarely a good reason to increase the value of smooth.df.

It is recommended to leave this argument at its default value. The argument is provided primarily for testing, diagnostics, and backward compatibility.

References

Boca, Simina M, and Jeffrey T Leek. 2018. "A Direct Approach to Estimating False Discovery Rates Conditional on Covariates." *PeerJ*, 6:e6035.

Jager, Leah R, and Jeffrey T Leek. 2013. "Empirical Estimates Suggest Most Published Medical Research Is True." *Biostatistics*.

Korthauer, Keegan, Patrick K Kimes, Claire Duvallet, Alejandro Reyes, Ayshwarya Subramanian, Mingxiang Teng, Chinmay Shukla, Eric J Alm, and Stephanie C Hicks. 2018. "A Practical Guide to Methods Controlling False Discoveries in Computational Biology." *bioRxiv*, 458786.

Leek, Jeffrey T., Leah Jager, and Simina M. Boca. 2018. *Swfdr: Science-Wise False Discovery Rate and Proportion of True Null Hypotheses Estimation.*

Locke, Adam E, Bratati Kahali, Sonja I Berndt, Anne E Justice, Tune H Pers, Felix R Day, Corey Powell, et al. 2015. "Genetic Studies of Body Mass Index Yield New Insights for Obesity Biology." *Nature* 518 (7538): 197–206.

Morgan, Martin. 2018. *BiocManager: Access the Bioconductor Project Package Repository*. https://CRAN.R-project.org/package=BiocManager.

Storey, John D. 2002. "A Direct Approach to False Discovery Rates." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)* 64 (3): 479–98.

Storey, John D, Andrew J Bass, Alan Dabney, and David Robinson. 2018. *Qvalue: Q-Value Estimation for False Discovery Rate Control.* http://github.com/jdstorey/qvalue.

Storey, John D, and Robert Tibshirani. 2003. "Statistical Significance for Genomewide Studies." *Proceedings of the National Academy of Sciences* 100 (16): 9440–5.