

ShrinkBayes: Bayesian analysis of high-dimensional omics data

Mark A. van de Wiel

October 31, 2016

Department of Epidemiology & Biostatistics
VU University Medical Center
Amsterdam, The Netherlands

`mark.vdwiel@vumc.nl`

Contents

1	Overview	2
2	Pre-amble: Accounting for different library sizes	4
3	Example 0, Easy Start: ShrinkBayes in one line	5
4	Example 1: Gaussian simulation setting, 2-group setting	7
4.1	Workflow	7
4.2	Results when using the correct prior	12
5	Example 2: Gaussian setting, HT RNAi data	14
6	Example 3: CAGE data	24
7	Practical considerations to guide choices in the analysis workflow	32
7.1	Trying ShrinkBayes on your data	32
7.2	To shrink or not to shrink?	33
7.3	Paired data	34
7.4	Choice of the prior	34
7.5	When should I use <code>mixtdisp=TRUE</code> ?	35

7.6	Shape of the nonparametric prior: logconcave, unimodal, symmetric	35
7.7	What threshold to use for testing (lfdr and BFDR computation)?	36
7.8	How to deal with extremely large data sets?	36
7.9	Tips for speeding up computations	36
8	Future topics	37
9	Appendix	38
9.1	Code used for generating the simulated data set ‘datSIM’ . . .	38
9.2	Code for simulated example	38
9.3	Code for HTRNAi example	38
9.4	Code for CAGE example	39
9.5	Code for running ShrinkBayes on very large data sets: simulated example	40
9.6	Code used for True FDR in simulation setting	42

1 Overview

ShrinkBayes is a package for Bayesian (differential) expression analysis of high-dimensional -omics data. It applies Empirical Bayes-type *multi*-parameter shrinkage to improve parameter estimation and inference. You should use it because it

- Is very flexible in terms of study designs
- Allows for random effects in a GLM setting
- Applies to many different data types, including Gaussian (e.g. miRNA, mRNA arrays, HT RNAi), counts (e.g. label-free proteomics) and zero-inflated counts (e.g. AGE, RNAseq)
- Was demonstrated to be more reproducible and powerful than other methods in small-sample settings (??)
- Is much more computationally efficient than MCMC, because it makes use of INLA (www.r-inla.org; (??))
- Provides (Bayesian) False Discovery Rates.

This document provides an overview on the usage of the ShrinkBayes package. For more detailed information on the methodology, performance and assumptions we refer to the articles (???). As example data we attached four data sets: 1) Simulated Gaussian data with 1500 rows and 284 columns; 2) HTRNAi, a data set with 960 rows and 6 columns containing Gaussian normalized HT RNAi data; 3); and mirseqnorm, a data set containing miRNA sequencing-based counts with 2,060 rows (features) and 55 columns (samples); and 4) CAGEdata1000, with 10,000 rows (features) and 25 columns (samples) containing normalized sequencing-based counts.

The simulated data set is used to illustrate the following aspects of ShrinkBayes:

- Ease of use in plain 2-group setting
- Proof of concept: ShrinkBayes correctly identifies the prior effect size distribution
- How to use a parametric mixture prior with point mass at zero?

The HT RNAi data set is used to illustrate the following aspects of ShrinkBayes:

- Analysis of Gaussian data (hence this example also covers most microarray data)
- Use of an offset in your model
- How to shrink multiple fixed effects and error variance?
- Use of a non-symmetric prior

The CAGE data set is used to illustrate the following aspects of ShrinkBayes:

- Analysis of (zero-inflated) count data (example also covers other sequencing data like RNAseq)
- Dealing with a complex design, including nuisance factors and blocked individuals
- Modeling and analysing random effects
- Estimation and inference for multiple pair-wise comparisons
- Joint shrinkage of fixed effect, random effect and overdispersion

- Using a mixture distribution on overdispersion

The miRNA sequencing data set was added to illustrate simple, one-line use of ShrinkBayes. In addition, it illustrates K -sample testing for a factor with more than 2 levels.

All examples use a fairly similar same work flow:

1. Load the normalized data
2. Set up the model and study design, possibly including multiple comparisons
3. Apply joint iterative procedure to shrink multiple parameters
4. Fit models for all data rows using the shrunken priors
5. Combine posteriors when multiple models have been applied to the same data (e.g. Poisson and Negative Binomial)
6. Update priors for crucial parameters to non-parametric or mixture forms
7. Update posteriors accordingly using the fitted non-parametric or mixture priors
8. Compute summaries like posterior means (parameter estimates) or posterior tail probabilities (local false discovery rates)
9. Compute Bayesian False Discovery Rates

For the analysis of the HTRNAi and CAGE data sets we focus on illustrating the settings as used in (?, HT RNAi data) and (?, CAGE data), but will also discuss alternatives when appropriate.

2 Pre-amble: Accounting for different library sizes

ShrinkBayes does NOT automatically account for different library sizes. There are two solutions.

1. Create normalized (pseudo-)counts from the $p \times n$ counts matrix and apply ShrinkBayes to those:

```
> libsize <- colSums(counts)
```

Calculate library size relative to geometric mean

```
> rellibsize <- libsize/exp(mean(log(libsize)))
```

```
> pseudocounts <- round(sweep(counts, 2, rellibsize, "/"))
```

2. Use the original counts, but include sample specific offsets that relate to the library sizes in the model (e.g. a simple model with an intercept and a group variable, see further on for other examples):

```
> libsize <- colSums(counts)
```

```
> rellibsize <- libsize/exp(mean(log(libsize)))
```

```
> myoffsets <- log(rellibsize)
```

```
> form <- ~ 1 + group + offset(myoffsets)
```

We prefer the second solution, although the first may sometimes be convenient when the data is also used for other purposes than testing (e.g. classification or prediction), which would also require normalized data.

3 Example 0, Easy Start: ShrinkBayes in one line

Please try the following example.

```
> library(ShrinkBayes)
```

```
> data(mirseqnorm)
```

```
> head(mirseqnorm[,1:10])
```

	M1	M10	M11	M12_1	M12_2	M13	M14_1	M14_2	M15	M16
1	0	0	0	0	0	3	0	0	2	0
2	0	0	0	0	0	0	0	0	0	0
3	0	0	0	0	0	0	0	0	0	0
4	0	0	0	0	0	0	0	0	0	0
5	1	0	0	0	0	0	0	3	1	0
6	0	0	0	0	0	0	0	0	0	0

Loads the package and the data.

```
> data(designmirseq)
```

```
> head(designmirseq)
```

	PM	indiv	timepos	chemo	organ1	organ2	organ3	organ
1	M	1	1	1	0	1	0	organ2
2	M	10	0	0	0	0	1	organ3
3	M	11	0	0	0	0	0	organ4
4	M	12	1	0	0	0	0	organ4
5	M	12	1	0	0	1	0	organ2
6	M	13	1	0	0	1	0	organ2

Show the design of the study.

```
> PM <- designmirseq$PM
> indiv <- designmirseq$indiv
> timepos <- designmirseq$timepos
> chemo <- designmirseq$chemo
> organ <- designmirseq$organ
```

Retrieve relevant covariates

```
> form = ~ 1 + PM + timepos + chemo + organ + f(indiv)
```

Specifies the regression formula. Note that `f()` is used to define a random effect. Then, a one-line application of `ShrinkBayes` is (but try the one below for a quick test):

```
> SBmir <- ShrinkBayesWrap(mirseqnorm, form)
```

This performs shrinkage and testing for the FIRST variable in `form`, PM. If testing is desired for a different variable this may be specified in the `paramtotest` argument (see below). The result is a list object with for slots. The FDRs slot contains the FDRs for the PM factor, the local fdrs and the shrunk effect size estimate (P-M). The `nsigsFDR01` slot contains the number of significant results at (B)FDR threshold 0.1. The other two slots contain information on the (shrunk) priors.

For a relatively quick try, try the following:

```
> SBmirsmall <- ShrinkBayesWrap(mirseqnorm[1:100,], form,
+ ntag=c(25,50),maxiter=2,priorsimple=TRUE, approx0=TRUE)
```

Here, using a smaller data set, less iterations, a simpler prior (pointmass-Gauss) and an approximation of the null-model speeds up the computations.

If you wish to shrink additional parameters as well (can render more powerful testing, in particular when sample size is small (?)), you may do so by specifying these:

```
> SBmirshrink <- ShrinkBayesWrap(mirseqnorm,form,  
+ shrinkadddfixed=c("organ","chemo","timepos"))
```

A (marginal)-likelihood-ratio-based test for `organ` (which has four levels) is the default for factor variables with more than 2 levels:

```
> SBmirorgan <- ShrinkBayesWrap(mirseqnorm,form,paramtotest="organ")
```

What does `ShrinkBayesWrap()` do? It is just a wrapper for functions that are explained in more detail in the remainder of this document. These functions are `ShrinkSeq()` and `ShrinkGauss()` for simultaneous shrinkage (Count and Gaussian data, respectively); `FitShrinkAll()` for applying INLA to all data rows; `MixtureUpdatePrior()`, `MixtureUpdatePosterior()` and `BFUpdatePosterior()` for computation of a prior with point mass and posteriors; `SummaryTable()` for computing summaries (including FDRs) from the posteriors.

For a factor variable with more than 2 levels, it performs (marginal)-likelihood-ratio-based K -sample testing by default, but multiple comparisons can be performed by use of either of the arguments `allcontrasts` or `multvscontrol`. See also `?ShrinkBayesWrap` for another example.

4 Example 1: Gaussian simulation setting, 2-group setting

4.1 Workflow

The example data set contains 1500 rows (features) and 8 samples (divided in two groups of 4). The first 1000 represent non-differential features (so proportion non-differential equals $2/3$), the last 500 represent differential features with mean group difference simulated from a $N(0,1)$ distribution. The noise is simulated from a $N(0,0.5^2)$ distribution. See Section 9.1 for the code used to simulate the data.

```
> library(ShrinkBayes)
> data(datsim)
> head(datsim[,1:5])
```

```
      [,1]      [,2]      [,3]      [,4]      [,5]
[1,] -0.04191917  0.070037754  0.8198383 -0.35443640  0.1128436
[2,] -1.01134453 -0.235494343 -0.3670104 -0.53177746  0.3673819
[3,]  0.19333209 -0.008211341 -0.1544538  0.29010757 -0.1937141
[4,] -1.03722998 -0.391160705 -0.2452836 -0.98763216 -0.8069727
[5,] -0.40153756  0.040684804  0.5219447 -0.09762872  0.1809304
[6,] -0.82760690  0.706451170  1.0892873  0.55709375  0.3335727
```

Loads the package and the data.

```
> ncpus2use <- 10
```

Sets the number of cpus to use for parallel computations. Combined with the `ncpus=ncpus2use` argument in the functions this fixes the number of cpus used. Note that for all functions the default is 2. If `ncpus2use` is larger than the actual number of available cpus, computations will still run.

```
> group <- factor(c(rep("group1",4),c(rep("group2",4))))
```

Defines covariate “group”, corresponding to the columns of the `datsim` data.

```
> form = y ~ 1 + group
```

Define the model formula. Specification should be according to the `inla` `formula` argument.

```
> shrinksimul <- ShrinkGauss(form=form, dat=datsim, shrinkfixed="group",
+ ncpus=ncpus2use)
```

This function simultaneously shrinks the fixed effect parameter ‘group’ and the Gaussian error standard deviation (default), using standard parametric priors. The function may take considerable computing time. See Section 5 for discussion on a) some arguments of the `ShrinkGauss` function b) retrieving various types of information from `shrinksimul`.


```
> fitg <- FitAllShrink(form,dat=datsim,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
```

Applies `inla` to compute posteriors for all data rows using the priors stored in `shrinksimul`. IMPORTANT: for better performance in the next function (`MixtureUpdatePrior`), the prior for the `shrinkfixed` parameter as used in `ShrinkGauss` (here: “treatment”) is by default dispersed by a factor 10 (hence a vaguer prior is used), see Section 5.

```
> form0 = y ~ 1
```

Defines the null-model formula. Specification should be according to the `inla formula` argument.

```
> fitg0 <- FitAllShrink(form0,dat=datsim,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
```

Fits the null-model. Only needed when inference for a point null hypothesis is desired. Please note that, in principle, the several `UpdatePrior` functions also work without a null-fit; Then, the Savage-Dickey approximation is used to approximate the marginal likelihood under the null- model. This saves computing time, but can be less accurate.

Below we estimate a mixture prior, namely a mixture of a zero point mass and two non-central Normals: $\pi(\beta) = (1 - p_0) * p_{\min} * N(\beta; -\mu, \tau^2) + p_0\delta_0 + (1 - p_0) * (1 - p_{\min}) * N(\beta; \mu, \tau^2)$. Note that this prior is misspecified w.r.t. to true effect size distribution (pointmass + central Normal).

```
> mixtprior2gauss <-
+ MixtureUpdatePrior(fitall=fitg,fitall0=fitg0, modus="mixt",
+ shrinkpara="group",ncpus=ncpus2use)
```

```
> bestfinal <- mixtprior2gauss$allpara[1,]
> bestfinal
```

pmin	p0	mu	stdev	sumloglik
0.3666667	0.6333333	0.1197843	0.9280651	1011.0476812

```
> mixtpostshr <- MixtureUpdatePosterior(fitg,mixtprior2gauss,fitg0,
+ ncpus=ncpus2use)
```

Update the posteriors of the “group” parameter for all data rows using the mixture prior.

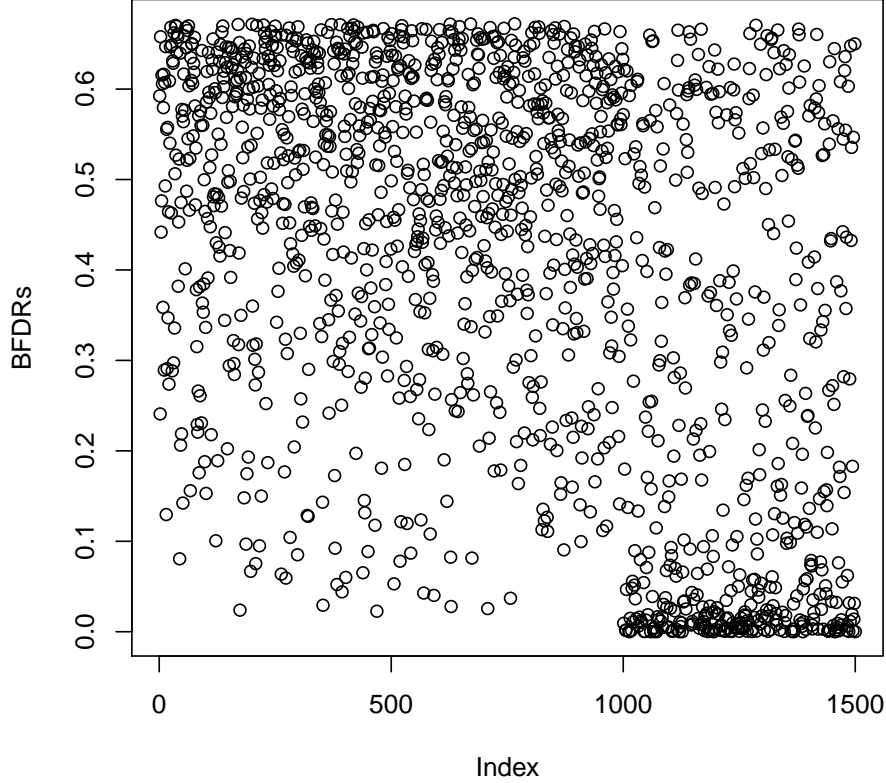
```
> lfdrless <- SummaryWrap(mixtpostshr, thr = 0, direction="lesser")
> lfdrgreat <- SummaryWrap(mixtpostshr, thr = 0, direction="greater")
```

Compute posterior (tail-)probabilities under shrinkage prior. Here `lfdrless` = $P(\beta_{\text{group}} \leq 0|Y)$ and `lfdrgreat` = $P(\beta_{\text{group}} \geq 0|Y)$ are computed. These can be interpreted as local false discovery rates corresponding to a one-sided interval null-hypothesis $H_0 : \beta_{\text{group}} \leq 0$ (for `lfdrless`) or $H_0 : \beta_{\text{group}} \geq 0$ (for `lfdrgreat`). See (?). Practically: if for a given feature `min(lfdrless, lfdrgreat)` is small, it means that the posterior is largely concentrated on either side of zero, which indicates that the group effect is different from 0. This fact is used when computing our *two-sided* version of Bayesian False Discovery Rate (??, BFDR):

```
> BFDRs <- BFDR(lfdrless, lfdrgreat)
```

Computes two-sided BFDRs. These can be interpreted as false discovery rates (in a cumulative sense) when one would select all features with smaller or equal null-probabilities (lfdrs) than the given feature.

```
> plot(BFDRs)
```



Plots the BFDRs against the feature index. Here we know that the first 1000 are generated from the null-hypotheses, the latter 500 from the alternative (see Section 9.1).

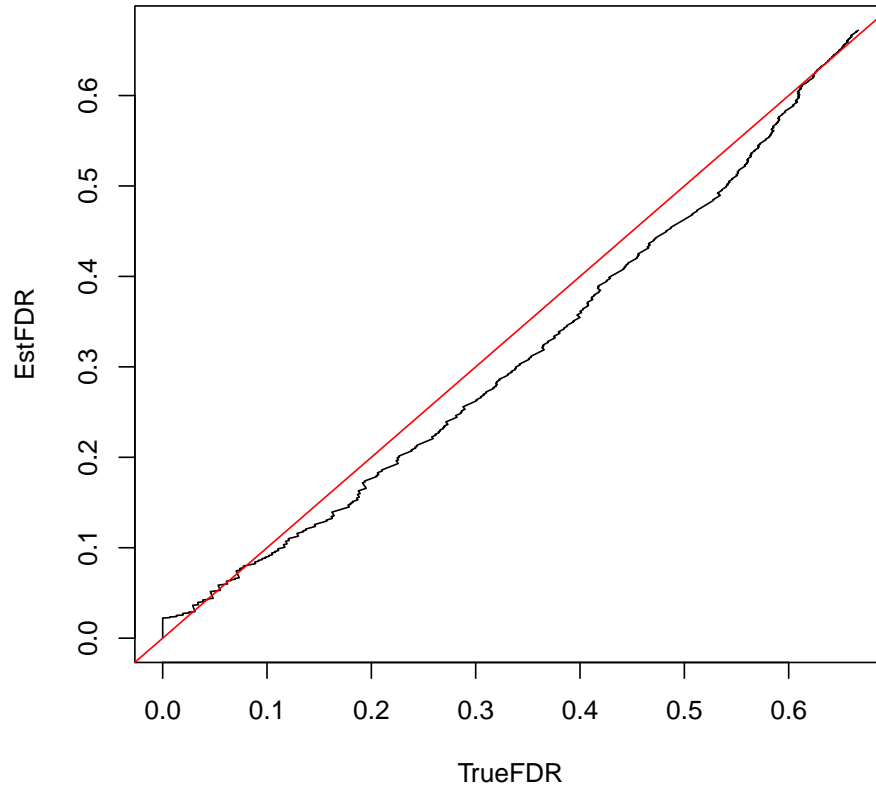
The computations above use a misspecified prior for β_{group} (because it is actually simulated from (1), the assumed model). Although it is fairly common to assume a fixed specific model in statistical computations, it is worthwhile to explore alternatives. **ShrinkBayes** offers two: 1) use a variety of parametric mixture models and study the robustness of the results against the parametric shape of the prior; 2) use a nonparametric prior. 1) is explored in the next Section, 2) in Sections 5 and 6.

In a simulation setting we can check the accuracy of the BFDR estimation to estimate the true FDR with the function `fdrcomp` (see Section 9.6):

```

> res <- fdrcomp(1:1000,BFDRs)
> plot(res,type="l")
> abline(a=0,b=1,col="red")

```



4.2 Results when using the correct prior

```

> prior1Normalp0 <- MixtureUpdatePrior(fitall=fitg,fitall=fitg0,
+ modus="gauss", shrinkpara="group",ncpus=ncpus2use)

```

Finds the (approximately) best mixture prior of the form

$$\pi(\beta) = p_0\delta_0 + (1 - p_0)N(\beta; 0, \tau^2), \quad (1)$$

so a mixture of zero-point mass (group effect equals zero) and a central Normal distribution. Grid search is used to maximize total log marginal

likelihood. Inclusion of a point mass may be attractive for the purpose of statistical inference (?).

```
> bestfinal <- prior1Normalp0$allpara[1,]
> bestfinal
```

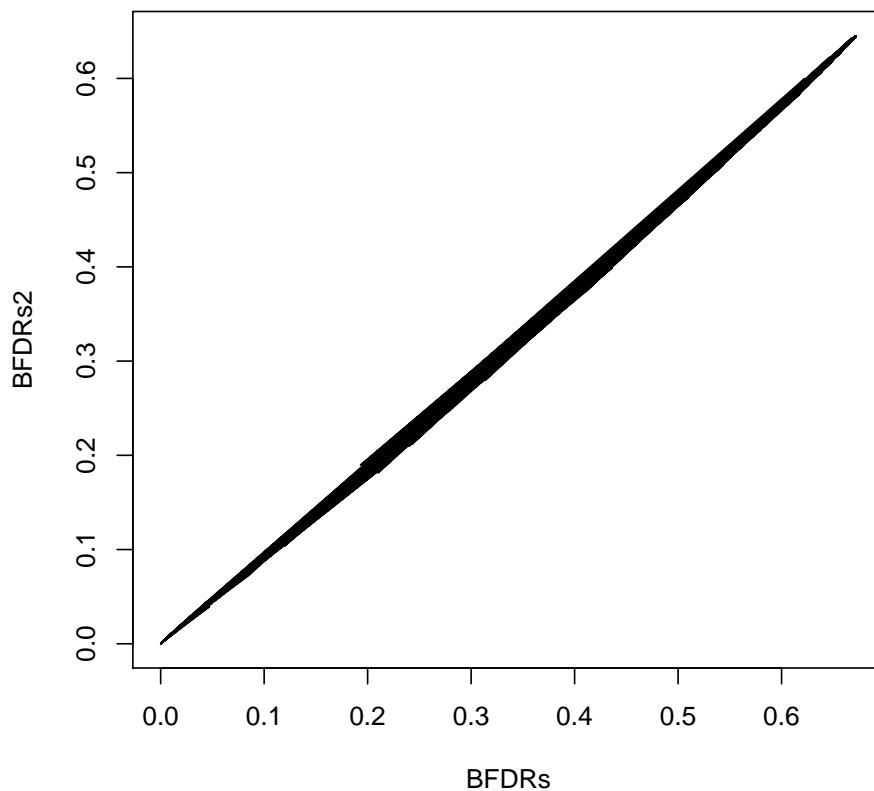
p0	stdev	sumloglik
0.6000000	0.8460445	1010.0916223

The best values of the parameters. Note the good performance of the algorithm: the true values are $p0 = 0.67$ and $stdev = 1$. Let us now recompute the posteriors, lfdRs and BFDRs of β_{group} under this prior:

```
> post1Normalp0shr <- MixtureUpdatePosterior(fitg,prior1Normalp0,fitg0,
+ ncpus=ncpus2use)
> lfdRless2 <- SummaryWrap(post1Normalp0shr, thr = 0, direction="lesser")
> lfdRgreat2 <- SummaryWrap(post1Normalp0shr, thr = 0, direction="greater")
> BFDRs2 <- BFDR(lfdRless2,lfdRgreat2)
```

We plot the BFDRs as computed under the misspecified prior (see previous Section; BFDRs) and under the correct prior (BFDRs2).

```
> plot(BFDRs,BFDRs2,type="l")
```



We observe they are very similar in this setting, hence the inference results are robust against misspecification of the prior in this setting.

5 Example 2: Gaussian setting, HT RNAi data

```
> library(ShrinkBayes)
> data(HTRNAi)
> head(HTRNAi)
```

	s1	s2	s3	s4	s5	s6
siRNA1	0.54317601	0.1953989	0.3883244	-0.03428689	0.6049207	-0.1790861
siRNA2	0.63533948	0.5057837	0.3928841	0.25338641	0.6935488	0.4185978
siRNA3	-0.91691878	-1.0838714	-0.6184123	-1.31090945	-0.2335765	-1.0377135

```

siRNA4 -0.74854963 -0.7619164 -0.2532474 -1.06233410 -0.5529374 -0.8189226
siRNA5  0.01110011 -0.6777245  0.3444957 -0.78924657  0.3554455 -0.8163929
siRNA6  0.59106644  0.5445666  0.4172433  0.31420140  0.6532168  0.5535300

```

Loads the package and the data

```
> ncpus2use <- 10
```

Sets the number of cpus to use for parallel computations.

```

> treatment <- factor(rep(c("untreated", "treated"), 3))
> assay <- factor(rep(1:3, each=2))

```

Defines covariates “treatment” and “assay”, corresponding to the columns of the HTRNAi data.

```

> offsetvalue <- c(0.1703984, -0.6958495,  0.3079694, -0.5582785,
+ 0.2251210, -0.6411269)

```

Defines an offset (often not needed). In this particular example, the offsets are computed from HT RNAi data with a positive control which is run for the same 6 screens. The offsets are posterior mean estimates from the same model as below (from many technical repeats per screen), but without shrinkage (because only one feature, the positive control, is involved). Use of an offset guarantees that the parameter estimates can be interpreted as deviation from the positive control.

```
> form = y ~ offset(offsetvalue) + 1 + treatment + assay
```

Define the model formula. Specification should be according to the `inla` formula argument.

```

> shrinksimul <- ShrinkGauss(form=form, dat=HTRNAi, shrinkfixed="treatment",
+ shrinkaddfixed="assay", fixedmeanzero = FALSE, ncpus=ncpus2use)

```

This function simultaneously shrinks the fixed effect parameters ‘treatment’ and ‘assay’. Note that in this particular setting it is likely that many treatment parameters are negative (since this concerns an effect with respect to a positive control; see (?)). Therefore, we set `fixedmeanzero = FALSE`,

so that we do not fix the mean of the prior to zero (which is the logical default in many microarray settings). In addition, the Gaussian error standard deviation is shrunken by default (`shrinksigma = TRUE`). This function may take considerable computing time.

The argument `shrinkfixed` contains the parameter of primary interest (“treatment”), while `shrinkaddfixed` contains an additional (nuisance) parameter (“assay”). Currently, maximally two fixed effect parameters can be shrunken simultaneously using Gaussian priors. `ShrinkGauss` has many additional arguments. We discuss a few crucial ones.

- `ntag`: Consecutive number of features used for shrinking. Important for computing time. Default is `ntag = c(100, 200, 500, 1000)`. You may consider using smaller values (e.g. `ntag = c(50, 100)`) for trying out, but we recommend to use at least 500 for final computations.
- `maxiter`: Maximum number of iteration per value of `ntag`. Important for computing time. Default is 10. Consider using a smaller value for trying out, e.g. `maxiter = 3`, but use at least `maxiter=10` for final computations.
- `fixedmeanzero` (`addfixedmeanzero`): Should the Gaussian mean of the `shrinkfixed` (`shrinkaddfixed`) prior be fixed to 0? Set to `FALSE` when this is undesirable/unrealistic.

```
> shrinksimul$pmlist
```

```
$mufixed
```

```
[1] -0.4220818
```

```
$precfixed
```

```
[1] 15.22325
```

```
$muaddfixed
```

```
[1] 0
```

```
$precaddfixed
```

```
[1] 88.85478
```

```
$shaperand
```



```
[1] 0.001
```

```
$raterand
```

```
[1] 0.001
```

```
$mixp
```

```
[1] 0.2 0.8
```

```
$shapeerr
```

```
[1] 14.80563
```

```
$rateerr
```

```
[1] 0.9963235
```

Shows the final parameter values. Note that the second parameter of Gaussian priors is a precision (1/variance) and the second parameter of Gamma priors (here, used for the precision of the error variance) is a rate.

```
> round(shrinksimul$paraall[,1:6],3)
```

	mufixed	precfixed	muaddfixed	precaddfixed	shaperand	raterand
paraall	0.000	0.100	0	0.100	0.001	0.001
paranew	-0.399	5.064	0	6.797	0.001	0.001
paranew	-0.405	7.761	0	13.148	0.001	0.001
paranew	-0.408	9.568	0	19.455	0.001	0.001
paranew	-0.407	10.626	0	25.331	0.001	0.001
paranew	-0.406	11.233	0	31.306	0.001	0.001
paranew	-0.406	11.657	0	36.715	0.001	0.001
paranew	-0.404	11.705	0	42.435	0.001	0.001
paranew	-0.406	11.478	0	47.533	0.001	0.001
paranew	-0.402	11.127	0	52.698	0.001	0.001
paranew	-0.406	10.904	0	57.092	0.001	0.001
paranew	-0.431	12.320	0	62.505	0.001	0.001
paranew	-0.444	13.260	0	66.303	0.001	0.001
paranew	-0.447	14.138	0	70.433	0.001	0.001
paranew	-0.447	14.877	0	74.258	0.001	0.001
paranew	-0.423	14.761	0	78.835	0.001	0.001
paranew	-0.420	15.117	0	82.779	0.001	0.001
paranew	-0.422	15.223	0	88.855	0.001	0.001

```
> round(shrinksimul$paraall[,-(1:6)],3)
```

	mixp1	mixp2	shapeerr	rateerr	
paraall	0.2	0.8	0.001	0.001	NA
paranew	0.2	0.8	3.258	0.312	100
paranew	0.2	0.8	7.238	0.787	100
paranew	0.2	0.8	9.331	0.960	100
paranew	0.2	0.8	10.760	1.006	100
paranew	0.2	0.8	11.647	1.007	100
paranew	0.2	0.8	12.578	1.011	100
paranew	0.2	0.8	13.350	1.012	100
paranew	0.2	0.8	13.944	1.005	100
paranew	0.2	0.8	14.487	1.004	100
paranew	0.2	0.8	15.039	1.005	100
paranew	0.2	0.8	14.440	0.982	200
paranew	0.2	0.8	13.957	0.974	200
paranew	0.2	0.8	13.713	0.975	200
paranew	0.2	0.8	13.309	0.970	200
paranew	0.2	0.8	13.961	0.992	500
paranew	0.2	0.8	14.388	0.994	500
paranew	0.2	0.8	14.806	0.996	960

Shows the consecutive estimates of the parameters. Note that parameters not involved in the shrinkage are not updated. The last columns shows the number of features used to estimate the parameters of the priors.

```
> fitg <- FitAllShrink(form,dat=HTRNAi,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
```

Applies `inla` to compute posteriors for all data rows using the priors stored in `shrinksimul`. IMPORTANT: the prior for the `shrinkfixed` parameter as used in `ShrinkGauss` (here: “treatment”) is by default dispersed by a factor 10 (hence a vaguer prior is used). This is done, because we experienced that it leads to better results when updating this prior to a nonparametric one (using `NonParaUpdatePrior`) or a mixture one (using `MixtureUpdatePrior`). If you do not want to disperse the prior of the `shrinkfixed` parameter, use the argument `dispersefixed=1`. Similar function arguments are available to disperse the prior of the `shrinkaddfixed` parameter or (when available) the random effects parameter. Additional arguments you may want to consider are `showupdate` (default: `FALSE`): when set to `TRUE` progression updates are shown for each `updateby` finished features. However, this may slow down the computations somewhat.

NOTE: you may see a lot of warnings from the call to `inla`. These usually occur for difficult data rows; this may either lead to missing results (which the subsequent function can cope with) or very wide posteriors. In both cases, it is unlikely that this leads to false positives or false negatives.

The result of `FitAllShrink` is a 2-component list. The first component `$res` is a list of `inla`-output objects of length `nr = nrow(dat)`, the number of data rows. When `nr` is large, the result may be a large object. Hence, for memory-efficiency not all `inla`-output is included. See the function arguments `effoutput`, `keepmargrand` and `keepmarghyper` for other options. The second component `$priors` contains the input prior parameters.

```
> fit1 <- fitg$res[[1]]
```

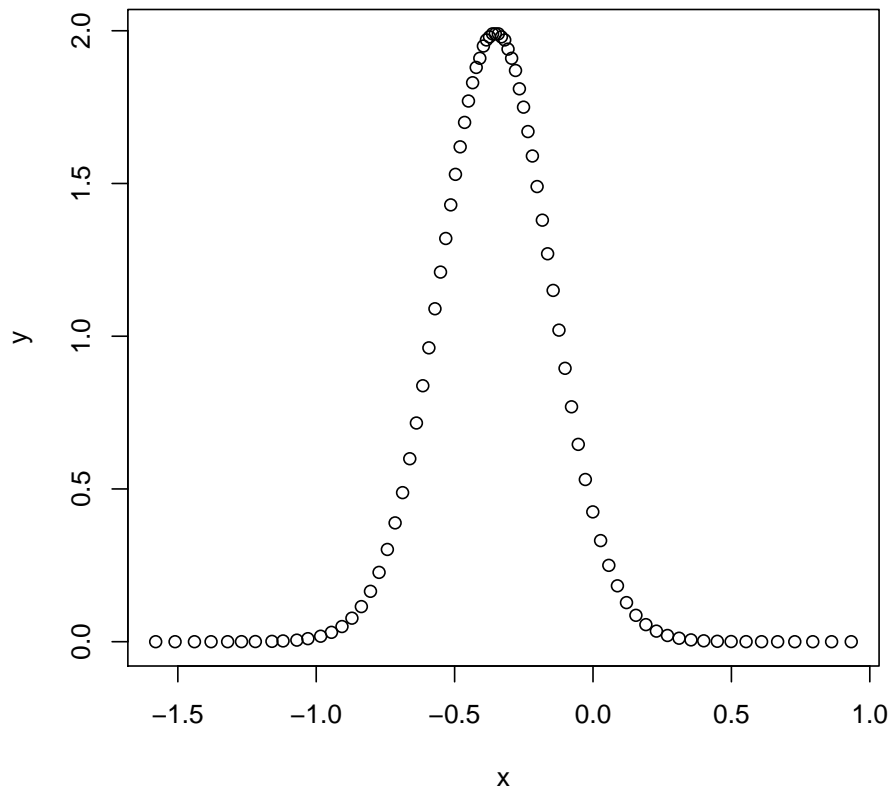
Retrieves the `inla`-output for the first feature. This contains a lot of information, including marginal posteriors of all relevant parameters in numeric format; summaries of the posteriors; model fit summaries; input arguments used by `inla`. Below we illustrate a few useful outputs.

```
> fit1$summary.fixed
```

	mean	sd	0.025quant	0.5quant	0.975quant	kld
(Intercept)	0.758	0.192	0.378	0.758	1.1300	7.40e-32
treatmentuntreated	-0.353	0.203	-0.752	-0.353	0.0463	0.00e+00
assay2	-0.247	0.223	-0.683	-0.248	0.1950	1.23e-32
assay3	-0.142	0.223	-0.578	-0.143	0.2990	0.00e+00

Shows the summaries of the fixed effect parameters.

```
> marginal <- fit1$marginals.fixed$treatmentuntreated
> plot(marginal)
```



Plots the marginal posterior (Use `type="l"` to obtain a curve instead of points). Note that this is obtained under a prior the variance of which is dispersed with a factor 10.

```
> fit1$summary.hyper
```

	mean	sd	0.025quant	0.5quant	0.975quant
Precision for the Gaussian observations	15.6	3.88	9.12	15.2	24.3

Shows the summaries of the hyper-parameters (parameters not involved in the regression formula). In this case only the error variance is a hyper-parameter.

```
> fit1$mlik
```

```

                                [,1]
log marginal-likelihood (integration) -6.7
log marginal-likelihood (Gaussian)    -6.7

```

Estimation of the marginal likelihood. Can be used to compare models.

```

> npprior <- NonParaUpdatePrior(fitall=fitg,modus="fixed",
+ shrinkpara="treatment",ncpus=ncpus2use, includeP0=FALSE,
+ allow2modes=FALSE, symmetric=FALSE, logconcave=TRUE)

```

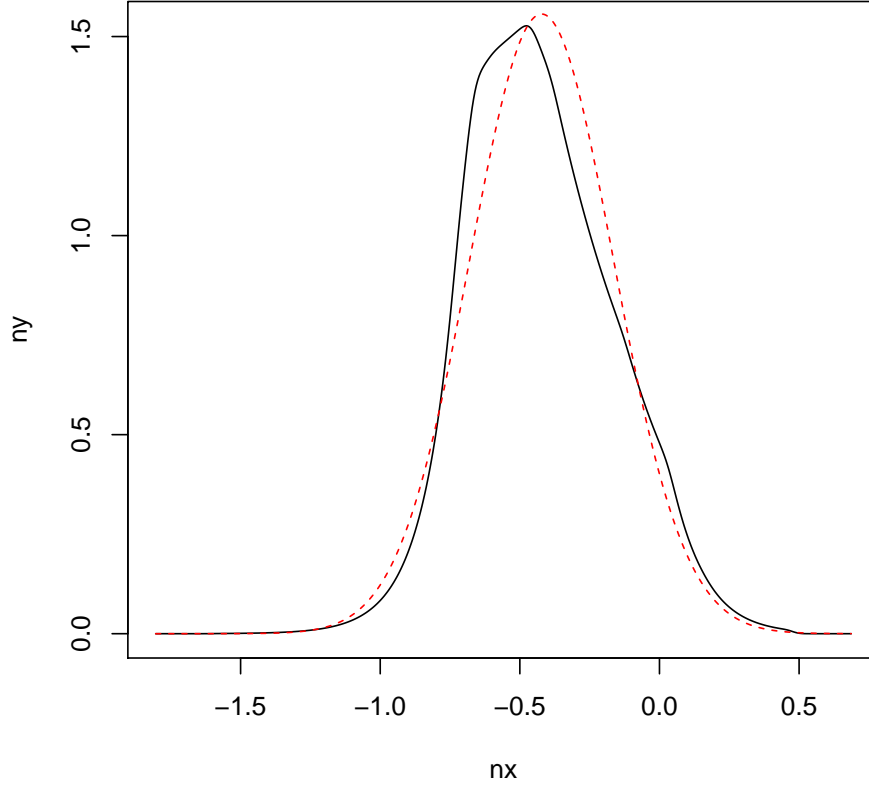
Updates the (dispersed) Gaussian prior for "treatment" to a smooth, log-concave non-parametric one. Note here we specifically allow an asymmetric prior (default is `symmetric=TRUE`). Also, in this example, we do not include a point mass on zero, because the main mass of effect sizes may well be concentrated elsewhere. This is context specific. See ? for explanation. Dropping the symmetry requirement may cause the non-parametric prior to become more unstable (its tails may become too dependent on specific data). Adding `logconcave=TRUE`, which forces a log-concave prior (?), aids in increasing the stability.

Let us now compare the resulting non-parametric prior with the original Gaussian one.

```

> plot(npprior$priornew,type="l")
> supp <- npprior$priornew[,1]
> points(supp,dnorm(supp,mean=shrinksimul$pmlist$mufixed,
+ sd = 1/sqrt(shrinksimul$pmlist$precfixed)),type="l",col="red",lty=2)

```



We

observe a clear asymmetry and a difference in the tails.

```
> nppostshr <- NonParaUpdatePosterior(fitg, npprior, ncpus=ncpus2use)
```

Updates the posteriors of the "treatment" parameter using the new, non-parametric prior.

```
> lfdr <- SummaryWrap(nppostshr, thr = 0, direction="lesser")
```

Here, we compute one-sided posterior null-probabilities of the kind $\text{lfdr} = P(\beta_{\text{treat}} \leq 0 | Y)$, which can be interpreted as a local false discovery rate. One-sided, because, when comparing with a positive control, we are mainly interested in siRNAs with larger (hence positive) effects w.r.t. the positive control, so lfdr should be small.

```
> BFDRs <- BFDR(lfdr)
```

Computes Bayesian False Discovery Rates (?) from the lfdrs.

```
> whsig <- which(BFDRs <= 0.1)
```

```
> whsig
```

```
[1] 176 608 749
```

```
> BFDRs[whsig]
```

```
[1] 0.07658634 0.01426447 0.03969088
```

```
> layout(matrix(1:3,nrow=1))
```

```
> plot(nppostshr[[whsig[1]]][[1]][[1]],xlim=c(-0.5,0.7),type="l")
```

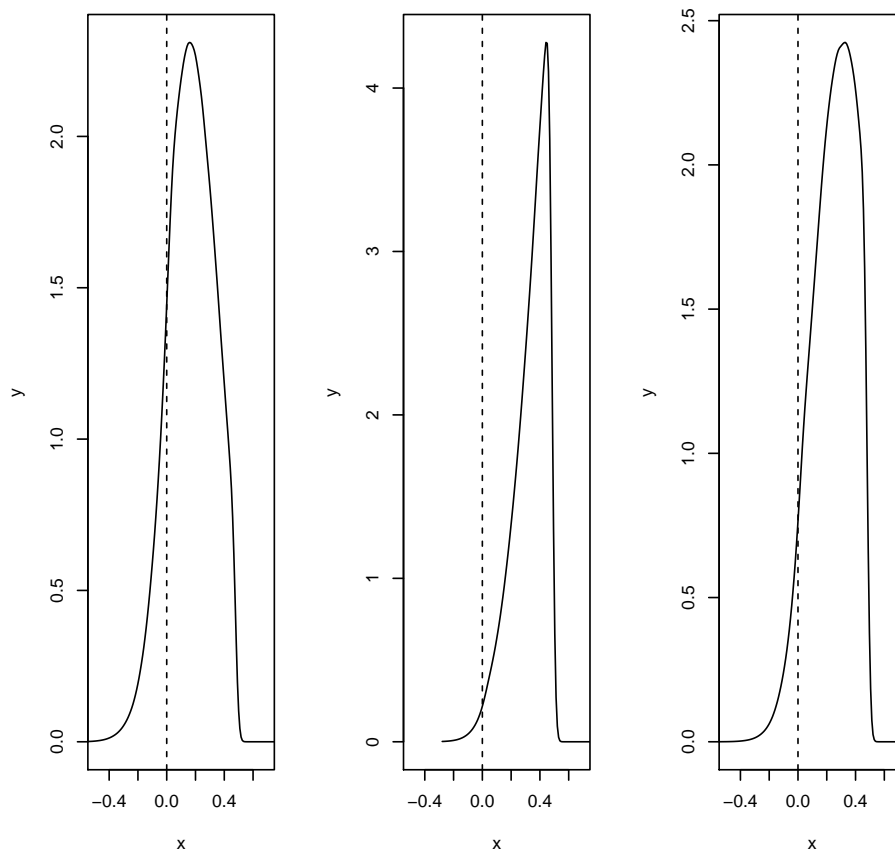
```
> abline(v=0,lty=2)
```

```
> plot(nppostshr[[whsig[2]]][[1]][[1]],xlim=c(-0.5,0.7),type="l")
```

```
> abline(v=0,lty=2)
```

```
> plot(nppostshr[[whsig[3]]][[1]][[1]],xlim=c(-0.5,0.7),type="l")
```

```
> abline(v=0,lty=2)
```



Displays the posteriors of the significant siRNAs. Note that use of a Gaussian prior rather than a non-parametric one would have rendered only the second siRNA (id: 608) to be significant (see (?)).

6 Example 3: CAGE data

The CAGE data set below consists of normalized sequencing (count) data for 10,000 tag clusters and 25 brain samples. For illustration purposes we limit ourselves to the analysis of the first 1,000 tag clusters. Details on the analysis are available in ? which present the results on 10,000 tag clusters.

```
> library(ShrinkBayes)
> data(CAGEdata10000)
```



```

> CAGEData <- CAGEData10000
> CAGEData <- CAGEData[1:1000,]
> CAGEData[1:2,]

```

	raw.0325_Frontal	raw.0325_Hippocampus	raw.0325_Putamen	raw.0325_Temporal	raw.0325_Temporal ra
Tag.1	0	0	0	0	
Tag.2	0	0	0	0	
	raw.05217_Frontal	raw.05217_Hippocampus	raw.05217_Putamen	raw.05217_Temporal	raw.05217_Temporal ra
Tag.1	68	73	49	57	
Tag.2	7	4	8	6	
	raw.05269_Temporal	raw.07319_Frontal	raw.07319_hippocampus	raw.07319_Temporal	raw.07319_Temporal r
Tag.1	0	0	0	0	
Tag.2	0	0	0	0	

Load ShrinkBayes, the data, select the first 1,000 rows and display the first 2.

```

> data(design_brain)
> design_brain

```

	pers	batch	groupfac
1	1	0	2
2	1	0	3
3	1	0	4
4	1	0	5
5	2	1	1
6	2	0	2
7	2	0	3
8	2	0	5
9	3	1	1
10	3	1	2
11	3	1	3
12	3	1	4
13	3	1	5
14	4	0	1
15	4	0	2
16	4	1	3
17	4	0	4
18	4	0	5
19	5	0	2
20	5	0	3

21	5	0	5
22	6	1	1
23	6	1	4
24	7	1	1
25	7	1	4

Loads the design of the brain study.

```
> pers <- design_brain$pers #persons
> batch <- design_brain$batch #batch
> groupfac <- design_brain$groupfac #group (= brain region)
```

Retrieves covariates from the design matrix.

```
> ncpus2use <- 10
```

Number of cpus to use in (parallel) computations. Note that this should be specified separately for functions that allow parallel computations by the `ncpus =` argument.

```
> groupfac <- BaselineDef("groupfac",baselinegroup="1")
```

The brain regions, coded by variable `groupfac`, are our main parameters of interest. As the design displays, we are in a multiple (>2) group setting. The function `BaselineDef` allows the user to set the baseline group, rather than let `INLA` decide. Here, the second argument is the character equivalent of the current level of the desired baseline group.

By default only comparisons with the baseline are included in the computations of posteriors. Use the following convenience function to create the other pair-wise comparisons for a given factor when this contains 3 or more levels (groups):

```
> lincombvec <- AllComp("groupfac")

> form = y ~ 1 + groupfac + batch + f(pers,model="iid")
```

Define the model formula. Specification should be according to the `inla` formula argument. Here `batch` and `group` are fixed effects, `pers` is a random effect.

```
> shrinksimul <- ShrinkSeq(form=form, dat=CAGEdata,shrinkfixed="groupfac",
+ shrinkrandom="pers",mixtdisp=TRUE,ncpus=ncpus2use)
```

Simultaneous shrinkage for `groupfac` and `pers`. In addition, the negative binomial overdispersion is shrunk by default (see `shrinkdisp` argument). Here, `batch` is not shrunk, because the effect of `batch` may not be uniform across the range of counts. In the example we allow a mixture prior for overdispersion (`mixtdisp=TRUE`). This increases the computing time by a factor 2, because `inla` has to fit the model under zero overdispersion (leading to a (zero-inflated) Poisson instead of a zero-inflated negative binomial.) `ShrinkSeq` default uses the zero-inflated negative binomial to fit sequencing data (see (?) for argumentation) by setting `fams="zinb"`. However, other options are `fams="nb"`, `fams="poisson"`, `fams="zip"` for Negative Binomial, Poisson and zero-inflated Poisson, respectively. In case one opts for `fams="nb"` it may be wise to explicitly account for the relationship between the mean and the overdispersion when shrinking (see e.g. (?)). In `ShrinkSeq` this is effectuated by setting `curvedisp=TRUE`. See the discussion on the `ShrinkGauss` function in Section 5 for other important arguments of the function.

```
> shrinksimul$pmlist$mixp
```

```
[1] 0.08376754 0.91623246
```

This returns the prior mass on the point mass of the mixture prior for overdispersion. If this would be close to zero (which we have observed in some other RNAseq applications), then it is sufficient to fit the model on all data for the (zero-inflated) Negative Binomial only. Otherwise, fits under the (zero-inflated) Poisson are also needed, like for this data set.

```
> fitzip <- FitAllShrink(form,dat=CAGEdata,fams="zip",shrinksimul,
+ ncpus=ncpus2use,lincomb=lincombvec)
```

Fits the model on all data using the priors resulting from `ShrinkSeq` and Zero-Inflated-Poisson likelihood. Note that, as discussed in Section 5, by default the variance of the prior for the main parameter of interest (here "`groupfac`") is increased by a factor 10. The `lincomb=lincombvec` argument specifies that the function should also generate posteriors for the linear combinations defined above.

```
> fitzinb <- FitAllShrink(form,dat=CAGEdata,fams="zinb",shrinksimul,
+ ncpus=ncpus2use,lincomb=lincombvec)
```

As above, but now under the Zero-Inflated-Negative Binomial likelihood (using the shrunk Gaussian prior for log-overdispersion).

```
> cp <- CombinePosteriors(fitzip,fitzinb,shrinksimul,para="groupfac",
+ ncpus=ncpus2use)
```

Combines the posteriors of the two fits (as outlined in the Suppl. Mat. of (?) for the "groupfac" parameters and the linear combinations (included by default). Hence, `cp` contains the posteriors of all pairwise comparisons under 1) a shrunk mixture prior for overdispersion; 2) a shrunk Gamma prior for the precision of `pers`; and 3) a (deliberately too wide) Gaussian prior for "groupfac". The latter will now be updated to a non-parametric prior.

```
> npprior <- NonParaUpdatePrior(fitall=cp,modus="fixed", shrinkpara="groupfac",
+ shrinklc=names(lincombvec),lincombs=lincombvec,ncpus=ncpus2use, maxiter=3,
+ includeP0 = FALSE, symmetric = TRUE, allow2modes=FALSE)
```

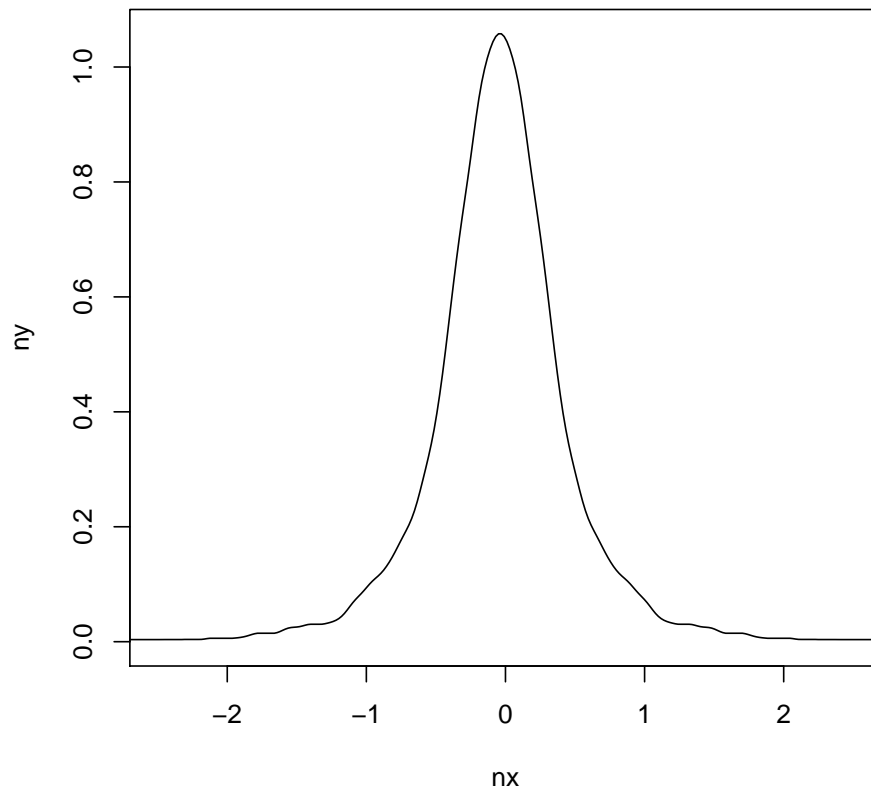
Find one common nonparametric prior for all group-wise differences, including the contrasts defined in `lincombvec`. Note that if `shrinklc` contains multiple contrasts, it is assumed they have the same mean and variance. If you do not want to include those contrasts (e.g. because you are focusing on comparison with the baseline group), set `shrinklc=NULL` (default). Note that we advise to increase `maxiter` for obtaining final results (default = 15). Important other arguments and defaults of the `NonParaUpdatePrior` function:

- `includeP0 = TRUE`: Include a point mass?
- `symmetric = TRUE`: Force the prior to be symmetric?
- `unimodal = TRUE`: Force the prior to be unimodal?
- `allow2modes = FALSE`: Can the prior have one mode on both the positive and negative halfplane (so unimodal on both half planes)? Only relevant when `unimodal = TRUE`.
- `logconcave = FALSE`: Force the prior to be logconcave?

Note that here we use the default setting for the shape of prior (symmetric, unimodal, but not necessarily logconcave). See also Section 7 for guidelines on how to use these settings.

Let's have a look at the prior.

```
> theprior <- npprior$priornew
> plot(theprior,type="l",xlim=c(-2.5,2.5))
```



```
> quantiles <- inla.qmarginal(p=c(0.05,0.25,0.5,0.75,0.95), theprior)
> quantiles

[1] -0.92262861 -0.30907530 -0.04458935  0.21989661  0.83344991

> expect <- inla.emarginal(function(x) x, theprior)
> expect

[1] -0.03955675

> sd <- sqrt(inla.emarginal(function(x) x^2, theprior) - expect^2)
> sd

[1] 0.7232569
```

Computes quantiles, the mean and standard deviation of the prior. These functions can also be applied to posteriors. See `?inla.qmarginal` and `?inla.emarginal` in your R-console for other options.

```
> nppostshr <- NonParaUpdatePosterior(cp, npprior, ncpus=ncpus2use)
```

Updates the posteriors of the "groupfac" parameters and the contrasts involved in `lincombvec` using the new, non-parametric prior.

```
> lfdr <- SummaryWrap(nppostshr, thr = log(1.5))
```

Compute *two-sided* local fdrs which equals $\text{lfdr} = \min(P(\beta_{\text{contrast}} \leq \text{thr}|Y), P(\beta_{\text{contrast}} \geq -\text{thr}|Y))$. The above `lfdr` introduces a desirable conservativeness with respect to an alternative posterior null-probability: $\text{lfdr}' = P(-\text{thr} \leq \beta_{\text{contrast}} \leq \text{thr}|Y)$. The latter, `lfdr'`, can not properly deal with wide posteriors that have a lot of probability mass outside $(-\text{thr}, \text{thr})$. See `?` for further discussion. The threshold `thr = log(1.5)` implies that we only wish to detect effects larger than 1.5 fold (see Section 7).

```
> BFDRs <- BFDR(lfdr)
```

```
> head(BFDRs)
```

	groupfac2	groupfac3	groupfac4	groupfac5	groupfac3mingroupfac2	groupfac4mingroupfac3
[1,]	0.7594013	0.7955304	0.8160210	0.8068468	0.8221437	0.8219
[2,]	0.8278324	0.8494276	0.8345920	0.8368664	0.8288337	0.7963
[3,]	0.8425993	0.1897167	0.7670844	0.8221101	0.1704118	0.5758
[4,]	0.4884483	0.7963692	0.6165695	0.7589679	0.8493518	0.8409
[5,]	0.8526093	0.8588785	0.8460374	0.8623392	0.8268282	0.7577
[6,]	0.8008360	0.7607720	0.6496556	0.7627002	0.7307867	0.6133

	groupfac5mingroupfac4
[1,]	0.8296039
[2,]	0.8355238
[3,]	0.8292926
[4,]	0.8396536
[5,]	0.8504250
[6,]	0.7984073

Compute (two-sided) Bayesian FDRs for all comparisons. Hence, a matrix is returned which features as rows and comparisons as columns.

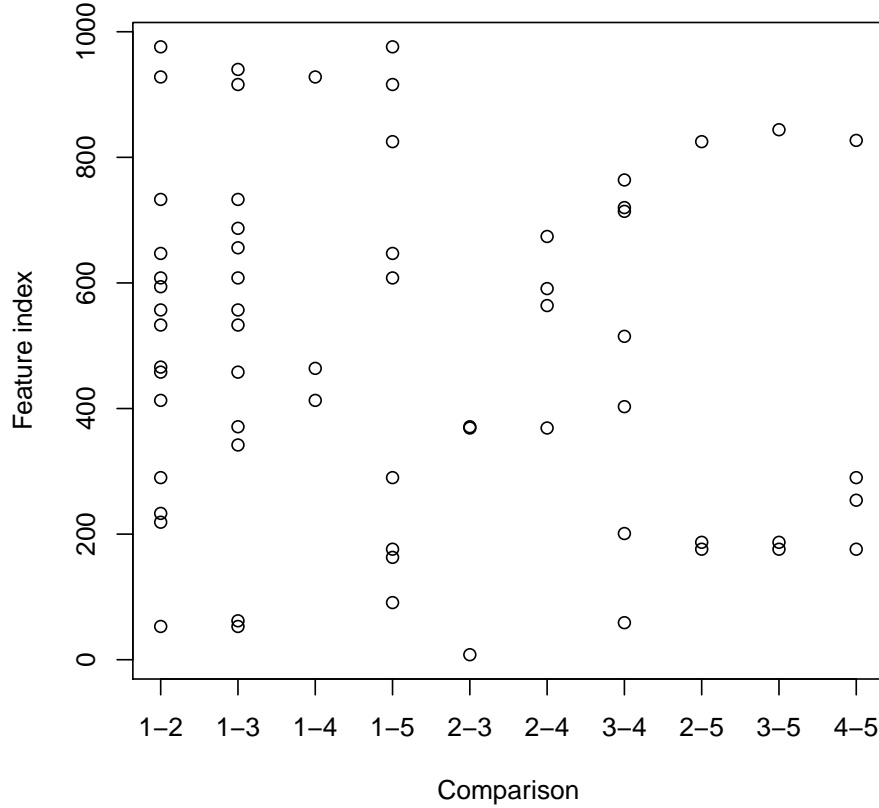
```
> BFDRmult <- BFDR(lfdr, multcomp=TRUE)
```

Compute (two-sided) Bayesian FDRs for testing the hypothesis: all comparisons parameters belong to the null domain. This allows one to perform K -group inference (like in ANOVA or an F-test) using a multiple comparison set-up. Hence, `BFDRmult` may be used to discover features showing at least one difference between groups (but not where: for that `BFDRs` is needed.)

```
> wh <- which(BFDRmult <= 0.1)
> length(wh)

[1] 43

> whcomp <- which(BFDRs[wh,]<= 0.1,arr.ind=TRUE)
> plot(cbind(whcomp[,2],wh[whcomp[,1]]),type="p",xlab="Comparison",
+ ylab="Feature index",xaxt="n")
> axis(1,at=1:10,labels=c("1-2","1-3","1-4","1-5","2-3","2-4","3-4",
+ "2-5","3-5","4-5"))
```



A plot that visualizes which features are significant for at least one comparison (hence according to `BFDReult`) and which comparison is significant for those features (according to `BFDRe`).

7 Practical considerations to guide choices in the analysis workflow

7.1 Trying `ShrinkBayes` on your data

Analysing data should not contain too many ‘trial-and-error’ steps, so ideally data analysis settings should be fixed a priori. Nevertheless, in particular in complex data analysis problems, it is often unavoidable that the researcher needs to try a few settings to understand what are sensible choices for the

data at hand. We advise to use a limited number of *randomly* selected features to try the various functions and setting in **ShrinkBayes**. The random selection is important, because many data sets contain structures on the features (e.g. genomic ordering). In our experience, 1,000 features is a safe choice for the algorithms to work well, for the results to be representative of the entire data set and computing times to be reasonable (in particular when processing on a multi-core computer).

We realize that the flexibility created by the many options in some of the functions in **ShrinkBayes** comes at a price: the (unexperienced) user may find it hard to make informed choices. Naturally, we assist the user on this issue by setting sensible defaults. Here, we share practical tips that should further aid in practical use of **ShrinkBayes**.

7.2 To shrink or not to shrink?

Shrinkage of a parameter can be beneficial in three ways. In small sample settings dispersion parameters are generally difficult to estimate. Hence, there is reasonable consensus on the benefit of shrinking dispersion-related parameters. This is effectuated in the functions **ShrinkSeq** and **ShrinkGauss** by the defaults `shrinkdisp = TRUE` and `shrinksigma = TRUE`, respectively. For the main parameter of interest we argued that shrinkage is important to come to proper inference in a Bayesian multiple testing setting (?). Hence, we strongly advise to shrink this parameter or these parameters as well.

For additional nuisance parameters in the regression the potential beneficial effect is more subtle. In ? we showed that shrinkage of the nuisance parameter "assay" was very beneficial for more powerful inference for the parameter of interest "treatment". The rationale for this was that in this experiment "assay", which had three levels, consumed 2 'degrees-of-freedom' (dof)¹, which is a lot in a 2*3 experiment. We found that the fitted prior for "assay" was very concentrated around zero, which basically renders more dof for the "treatment" parameter. Hence, in very small experiments with a nuisance parameter that has relatively many levels, it may certainly pay off to shrink the nuisance parameter. However, sometimes the effect of the nuisance parameter is not uniform across the range of data, even after normalization. This may be assessed by plotting a (naive) estimate of the nuisance parameter against the mean of the data ($\text{mean}(\log(x+1))$ for when x is a count). In the HTRNAi example, Section 5, the batch effect was

¹between quotes, because, strictly, dof is not a Bayesian concept

not uniform and so we decided not to shrink this. Since the zero-inflation parameter is also not uniform over the log-count range, we recommend to not shrink it either (hence `shrinkp0 = FALSE` is the default in `ShrinkSeq`; see also ?).

7.3 Paired data

Paired data may be modeled in various ways. Currently, we recommend to model directly the data (so not the pairwise differences) and account for the pairing by use of a random effect, similar to the use of `pers` in the CAGE example of Section 6. So for 5 individuals with a paired measurement this would be coded as: `indiv = c(1,1,2,2,3,3,4,4,5,5)` and `f(indiv,model="iid")`, assuming the columns of the data are ordered with respect to individuals.

When the pairs represent individuals, we suggest replacing the (zero-inflated) Negative Binomial by a (zero-inflated) Poisson for count data, by setting `fams="zip"` (or `fams="poisson"`) in the functions `ShrinkSeq` and `FitAll-Shrink`. The reason for this is that the between individual variation (which is the motivation for overdispersing the Poisson) is now accounted for by the Gaussian random effect in the regression.

7.4 Choice of the prior

`ShrinkBayes` has many options for the prior, in particular when it concerns the main parameter of interest. In general, we recommend to use

- `MixtureUpdatePrior` with `modus = "mixt"` for testing the point null-hypothesis $H_0 : \beta = 0$. Extensive simulations have shown that this leads to rather accurate (B)FDR estimation for a wide variety of true effect size distributions, while rendering good power properties. Alternative: `NonParaUpdatePrior` with `includeP0 = TRUE`, a non-parametric prior with point mass. Slightly worse (B)FDR estimation. Possibly somewhat more power than `MixtureUpdatePrior` for very small sample size (e.g. $N = 2 * 3$).
- `NonParaUpdatePrior` with `includeP0 = FALSE` when testing an interval null-hypothesis, e.g. using thresholds like `thr = log(1.25)`, `thr = log(1.5)`, `thr = log(2)` (or the negatives hereof). The non-parametric prior provides the best adaptivity to the data. For example, in Section 5 we illustrate the benefit of using a nonparametric prior

over a Gaussian one in a small-sample setting. Also we have shown in simulation settings that a nonparametric prior can accurately recover smooth parametric shapes (?).

- **ShrinkSeq** or **ShrinkGauss** using the option `excludeformull` in combination with **BFUpdatePosterior** to test two nested models that differ by more than one variable.

We, and others (?), noted that for some data sets, the point mass on zero in mixture priors is estimated to be very low, irrespective of the shape of continuous components. In such cases we advise to use the default lower bound 0.5 on the point mass component `p0lower = 0.5` in **NonParaUpdatePrior** to force some shrinkage towards 0, in combination with using a threshold `thr` different from zero in **SummaryWrap**.

7.5 When should I use `mixtdisp=TRUE`?

The **ShrinkSeq** function allows one to use a mixture of Poisson and Negative Binomial likelihood (or zero-inflated versions thereof) by setting `mixtdisp=TRUE`. When the experiment involves sequencing data on independent individuals our experience (and the consensus in the community) is that overdispersion (to allow more variance between samples than the Poisson does) is really needed, so for such studies we expect that the majority of features prefers the Negative Binomial, so in such a case we recommend to simply use the default `mixtdisp=FALSE`. However, if the experiment involves cell lines (hence less variability is likely) or when the between-individual variation is modeled by a random effect at the regression level (see Section 6) we advise to use `mixtdisp=TRUE`. If the estimate of the mixture proportion (`shrinksimul$pmlist$mixp`, where `shrinksimul` is the object containing the output of **ShrinkSeq**) is close to 0, proceed with (zero-inflated) Negative Binomial. If it is close to 1, proceed with (zero-inflated) Poisson. Otherwise, fit both and use the **CombinePosteriors** function (see Section 6).

7.6 Shape of the nonparametric prior: `logconcave`, `unimodal`, `symmetric`

In the function **NonParaPriorUpdate** the options `logconcave`, `unimodal` and `symmetric` allow one to increase stability of the prior. Setting all these options to `FALSE` renders the completely non-restrictive setting, where all `TRUE` gives the most restrictive setting. We recommend to use either of the

following two settings: a) The default: `logconcave = FALSE`, `unimodal = TRUE` and `symmetric = TRUE`, where enforcing symmetry and unimodality creates stability for the tails of the prior. b) `logconcave = TRUE`, `unimodal = TRUE` and `symmetric = FALSE`, if asymmetry is considered realistic; the log-concavity restriction may then help to stabilize both tails of the prior.

7.7 What threshold to use for testing (lfdR and BFDR computation)?

In the function `SummaryWrap` the threshold argument `thr` is important for determining the significant features by lfdR or BFDR. To a large extent the discussion on minimal relevant effect size (which is what `thr` represents, on log level) is a biological one. The more `thr` deviates from 0 the less features are detected, but those that are detected are likely to be (relatively) relevant. We advise to be careful with using `thr = 0` in the two-sided, non-parametric setting, because if the truth is sparse (or near-sparse: many very small effects) it may render too many false positives. In general, we believe it is wise to report results for at least two values of `thr`, e.g. `thr=log(1.25)` and `thr=log(2)`, representing 1.25 and 2-fold changes.

7.8 How to deal with extremely large data sets?

For very large data sets, the internal memory of the computer may not suffice to perform the fits for all features in one call. Hence, we advise to split the three major tasks of `ShrinkBayes`: a) Finding the correct priors; b) Computing the posteriors under those priors; and c) Inference (lfdR and BFDR computations). a) is usually accurate enough when using a random subset of features of size, say 10,000. For those computations, internal memory is usually sufficient. Then, for the computation of posteriors we advise to do this in batches and save intermediate results on the hard-disk. Then, perform inference on summaries of those posteriors using `SummaryWrap` and BFDR.

An example is included in the Appendix, Section 9.5.

7.9 Tips for speeding up computations

Several functions in `ShrinkBayes` may take a while to run. Therefore, we parallelized some of the most time-consuming computations, so your computations will obviously be faster when increasing the number of available. Note that computing time increases linearly with the number of features.

Some more tips for speeding up computations:

- When starting to use **ShrinkBayes** clean up the R-memory: either by re-starting R or by using `rm(list=ls());gc();`.
- If you use Windows OS: consider switching to Linux. Due to the intensive interaction of **inla** with the hard-disk, we experienced that use of Linux instead of Windows (on the same computer) may speed up computations by a factor of 3-4.

8 Future topics

We plan to extent **ShrinkBayes** in many directions.

- Other data types, in particular fractional data (on 0-1 scale), e.g. methylation
- Longitudinal settings (multiple measurements over time)
- Modelling and reporting joint posteriors rather than only marginal ones
- K -sample inference (other than through multiple comparisons, which was illustrated in the Section 6)
- “Small” multivariate GLM settings. Where “small” means: non-high-dimensional (at the level of the model)
- “Large” multivariate penalized regression settings
- Inference with point mass on zero and *nonparametric* continuous component
- Using Bayes’ Factors for inference in a multiplicity setting
- ...

If you are interested in any of these topics, or when you feel these are essential for your application, please let us know. We might have progressed and be able to share (preliminary) code with you.

9 Appendix

9.1 Code used for generating the simulated data set ‘datSIM’

```
> #1500 rows (siRNAs), 8 samples, pi0 = 2/3
> datSIM1 <- matrix(rnorm(8000,mean=0,sd=0.5),nrow=1000)
> meanvec <- matrix(rep(rnorm(500,0,1),4),nrow=500)
> datSIM2 <- cbind(matrix(rnorm(2000,mean=0,sd=0.5),nrow=500),
+ matrix(rnorm(2000,mean=0,sd=0.5),nrow=500) + meanvec)
> datSIM <- rbind(datSIM1,datSIM2)
```

9.2 Code for simulated example

```
> library(ShrinkBayes)
> data(datSIM)
> ncpus2use <- 10
> group <- factor(c(rep("group1",4),c(rep("group2",4))))
> form = y ~ 1 + group
> form0 = y ~ 1
> shrinksimul <- ShrinkGauss(form=form, dat=datSIM,shrinkfixed="group",
+ ncpus=ncpus2use)
> fitg <- FitAllShrink(form,dat=datSIM,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
> fitg0 <- FitAllShrink(form0,dat=datSIM,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
> mixtprior2gauss <- MixtureUpdatePrior(fitall=fitg,fitall0=fitg0,
+ modus="mixt", shrinkpara="group",ncpus=ncpus2use)
> mixtpostshr <- MixtureUpdatePosterior(fitg,mixtprior2gauss,fitg0,
+ ncpus=ncpus2use)
> lfdrless <- SummaryWrap(mixtpostshr, thr = 0, direction="lesser")
> lfdrgreat <- SummaryWrap(mixtpostshr, thr = 0, direction="greater")
> BFDRs <- BFDR(lfdrless,lfdrgreat)
```

9.3 Code for HTRNAi example

```
> library(ShrinkBayes)
> data(HTRNAi)
> ncpus2use <- 10
> treatment <- factor(rep(c("untreated","treated"),3))
> assay <- factor(rep(1:3,each=2))
> offsetvalue <- c(0.1703984, -0.6958495, 0.3079694, -0.5582785,
```

```

+ 0.2251210, -0.6411269)
> form = y ~ offset(offsetvalue) + 1 + treatment + assay
> shrinksimul <- ShrinkGauss(form=form, dat=HTRNAi, shrinkfixed="treatment",
+ shrinkaddfixed="assay", fixedmeanzero = FALSE, ncpus=ncpus2use)
> fitg <- FitAllShrink(form, dat=HTRNAi, fams="gaussian", shrinksimul,
+ ncpus=ncpus2use)
> npprior <- NonParaUpdatePrior(fitall=fitg, modus="fixed",
+ shrinkpara="treatment", ncpus=ncpus2use, includePO = FALSE,
+ logconcave=TRUE, allow2modes=FALSE)
> nppostshr <- NonParaUpdatePosterior(fitg, npprior, ncpus=ncpus2use)
> lfdr <- SummaryWrap(nppostshr, thr = 0, direction="lesser")
> BFDRs <- BFDR(lfdr)

```

9.4 Code for CAGE example

```

> data(CAGEdata10000)
> CAGEdata <- CAGEdata10000[1:1000,]
> data(design_brain)
> pers <- design_brain$pers ; batch <- design_brain$batch;
> groupfac <- design_brain$groupfac
> ncpus2use <- 10
> groupfac <- BaselineDef("groupfac", baselinegroup="1")
> lincombvec <- AllComp("groupfac")
> form = y ~ 1 + groupfac + batch + f(pers, model="iid")
> shrinksimul <- ShrinkSeq(form=form, dat=CAGEdata, shrinkfixed="groupfac",
+ shrinkrandom="pers", mixtdisp=TRUE, ncpus=ncpus2use)
> fitzip <- FitAllShrink(form, dat=CAGEdata, fams="zip", shrinksimul,
+ ncpus=ncpus2use, lincomb=lincombvec)
> fitzinb <- FitAllShrink(form, dat=CAGEdata, fams="zinb", shrinksimul,
+ ncpus=ncpus2use, lincomb=lincombvec)
> cp <- CombinePosteriors(fitzip, fitzinb, shrinksimul, para="groupfac",
+ ncpus=ncpus2use)
> npprior <- NonParaUpdatePrior(fitall=cp, modus="fixed",
+ shrinkpara="groupfac", shrinklc=TRUE, ncpus=ncpus2use, maxiter=3,
+ includePO = FALSE, symmetric=TRUE, logconcave=FALSE, allow2modes=FALSE)
> nppostshr <- NonParaUpdatePosterior(cp, npprior, ncpus=ncpus2use)
> lfdr <- SummaryWrap(nppostshr, thr = log(1.5))
> BFDRs <- BFDR(lfdr)
> BFDRmult <- BFDR(lfdr, multcomp=TRUE)

```

9.5 Code for running ShrinkBayes on very large data sets: simulated example

```
> datsim1 <- matrix(rnorm(400000*8,mean=0,sd=0.5),nrow=400000)
> meanvec <- matrix(rep(rnorm(200000,0,1),4),nrow=200000)
> datsim2 <- cbind(matrix(rnorm(200000*4,mean=0,sd=0.5),nrow=200000),
+ matrix(rnorm(200000*4,mean=0,sd=0.5),nrow=200000) + meanvec)
> datsimlarge <- rbind(datsim1,datsim2)
> save(datsimlarge,file="datsimlarge.Rdata")
```

Simulates a data set with 600,000 rows, 4*2 samples, $\pi_0 = 2/3$. See Section 4.

```
> tuningsize <- 10000
> batchsize <- 50000
```

`tuningsize` determines the maximum number of features used in the tuning phase of ShrinkBayes. The tuning phase determines the priors for those parameters for which shrinkage is desired. Then, the actual fitting and computation of posteriors for all features is performed in batches of size 'batchsize'. Results are written to the hard-disk in batches. All this to avoid having to store many posterior distributions in the internal memory.

```
> whrows <- sample(1:nrow(datsimlarge),tuningsize)
> datsimtune <- datsimlarge[whrows,]
> save(datsimtune,file="datsimtune.Rdata")
> rm(datsimlarge);gc()
```

Creates the tuning data set on a random set of features. `datsimtune` is used to fit the priors

```
> ncpus2use <- 6
> group <- factor(c(rep("group1",4),c(rep("group2",4))))
> form = y ~ 1 + group
> form0 = y ~ 1
> library(ShrinkBayes)
```

Number of cpus to use, covariate `group`, formula and library loading

```
> shrinksimul <- ShrinkGauss(form=form, dat=datsimtune,shrinkfixed="group",
+ ncpus=ncpus2use)
```



```

> fitg <- FitAllShrink(form,dat=datsimtune,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
> fitg0 <- FitAllShrink(form0,dat=datsimtune,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
> mixtprior<- MixtureUpdatePrior(fitall=fitg, fitall0=fitg0,modus="gauss",
+ shrinkpara="group",ncpus=ncpus2use)
> save(shrinksimul,mixtprior,file="priorstuned.Rdata")

```

Determining the priors using the exact same code as in Section 4, but now on the tuning data set `datsimtune`. Now, we start with fitting on the large data set.

```

> saveFits <- FALSE

```

Do you want to save the `inla`-output of the function `FitShrinkAll`? Note that these are large objects that contain a lot of information, including marginal posteriors of all parameters.

```

> savePosteriors <- TRUE

```

Do you want to save output of the update function (either `MixtureUpdatePosterior` or `NonParaUpdatePosterior`)? This output contains the marginal posteriors of the parameter of interest only. May still be a fairly large object, but generally much smaller than the fit object. Usually wise to store these in case you decide to change inference in the `SummaryWrap` and `BFDR` functions (e.g. a different threshold).

```

> load("priorstuned.Rdata")
> load("datsimlarge.Rdata")

```

Load the R-objects that contain the information on the priors and the large data set.

```

> nr <- nrow(datsimlarge)
> nloop <- ceiling(nr/batchsize)
> lfdrless <- c()
> lfdrgreat <- c()

```

Determines the number of features, the number of loops and initializes the `lfdrs`

```

> for(k in 1:nloop){
+ if(k>1) load("datSIMlarge.Rdata")
+ rangek <- ((k-1)*batchsize+1):min(nr,k*batchsize)
+ datk <- datSIMlarge[rangek,]
+ print(paste("Computing posteriors for features",rangek[1],"to",
+ rangek[length(rangek)]))
+ rm(datSIMlarge);gc()
+ fitgk <- FitAllShrink(form,dat=datk,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
+ fitg0k <- FitAllShrink(form0,dat=datk,fams="gaussian",shrinksimul,
+ ncpus=ncpus2use)
+ if(saveFits) {
+   save(fitgk,file=paste("fitg_batch_",k,".Rdata",sep=""))
+   save(fitg0k,file=paste("fitg0_batch_",k,".Rdata",sep=""))
+ }
+ mixtpostshrk <- MixtureUpdatePosterior(fitgk,mixtprior,fitg0k,
+ ncpus=ncpus2use)
+ if(savePosteriors) save(mixtpostshrk,
+ file=paste("mixtpostshr_batch_",k,".Rdata",sep=""))
+ lfdrllessk <- SummaryWrap(mixtpostshrk, thr = 0, direction="lesser")
+ lfdrgreatk <- SummaryWrap(mixtpostshrk, thr = 0, direction="greater")
+ lfdrlless <- rbind(lfdrlless,lfdrllessk)
+ lfdrgreat <- rbind(lfdrgreat,lfdrgreatk)
+ save(lfdrlless,lfdrgreat,file="lfdrs.Rdata")
+ rm(fitgk,mixtpostshr);gc()
+ }

```

The loop that determines posteriors for all features per batch of size `batchsize`. `FitAllShrink` determines the posteriors under the initial priors stored in `simulshrink`, `MixtureUpdatePosterior` updates these posteriors for the `mixtprior` on "group". Then `lfdrs` are computed for batch `k` and all `lfdrs` are stored in `lfdrlless` and `lfdrgreat`.

```

> BFDRs <- BFDR(lfdrlless,lfdrgreat)

```

Finally, BFDRs are computed. Note that these need the input of *all* features, so BFDR cannot be computed inside the loop above.

9.6 Code used for True FDR in simulation setting

negatives: Indices of true negatives , **sig:** BFDR measure of significance.

```

fdrcomp <- function(negatives,sig){
#True FDR = FP/N_P. Est FDR = \sum (p0s*I) / N_P
sortsig <- sort(sig,index.return=T)
sortind <- sortsig$ix
n <- length(sig)
arr <- rep(0,n)
wh <- which(sapply(sortind,is.element,set=negatives))
arr[wh] <- 1
FP <- cumsum(arr)
TrueFDR <- FP/1:n
EstFDR <- sortsig$x
return(cbind(TrueFDR,EstFDR))
}

```