

# Package ‘mashr’

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**Type** Package

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**Title** Multivariate Adaptive Shrinkage

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**Description** Implements the multivariate adaptive shrinkage (mash) method of Uribarri et al (2019) <[DOI:10.1038/s41588-018-0268-8](https://doi.org/10.1038/s41588-018-0268-8)> for estimating and testing large numbers of effects in many conditions (or many outcomes). Mash takes an empirical Bayes approach to testing and effect estimation; it estimates patterns of similarity among conditions, then exploits these patterns to improve accuracy of the effect estimates. The core linear algebra is implemented in C++ for fast model fitting and posterior computation.

**URL** <https://github.com/stephenslab/mashr>

**BugReports** <https://github.com/stephenslab/mashr/issues>

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**SystemRequirements** C++11

**Depends** R (>= 3.3.0), ashR (>= 2.2-22)

**Imports** assertthat, utils, stats, plyr, rmeta, Rcpp (>= 0.12.11), mvtnorm, abind, softImpute

**LinkingTo** Rcpp, RcppArmadillo, RcppGSL (>= 0.3.8)

**Suggests** MASS, REBayes, corrplot (>= 0.90), testthat, kableExtra, knitr, rmarkdown, profmem, flashier, ebnm

**Additional\_repositories** <https://stephenslab.github.io/flashier-drat>

**VignetteBuilder** knitr

**NeedsCompilation** yes

**Biarch** true

**RoxygenNote** 7.1.2

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---

contrast\_matrix      *Create contrast matrix*

---

**Description**

Create contrast matrix

**Usage**

```
contrast_matrix(R, ref, name = 1:R)
```

**Arguments**

R	the number of column for the contrast matrix
ref	the reference group. It could be a number between 1,..., R, R is number of conditions, or the name of reference group. If there is no reference group, it can be the string 'mean'.
name	a length R vector contains the name for conditions

**Examples**

```
contrast_matrix(5, 'mean')
```

---

cov\_canonical      *Compute a list of canonical covariance matrices*

---

**Description**

Compute a list of canonical covariance matrices

**Usage**

```
cov_canonical(
  data,
  cov_methods = c("identity", "singletons", "equal_effects", "simple_het")
)
```

**Arguments**

data	a mash data object, eg as created by mash_set_data
cov_methods	a vector of strings indicating the matrices to be used: "identity" for the identity (effects are independent among conditions); "singletons" for the set of matrices with just one non-zero entry $x_{jj} = 1$ ( $j=1,\dots,R$ ); (effect specific to condition $j$ ); "equal_effects" for the matrix of all 1s (effects are equal among conditions); "simple_het" for a set of matrices with 1s on the diagonal and all off-diagonal elements equal to 0.25, 0.5 or 0.75; see cov_simple_het for details; (effects are correlated among conditions).

**Details**

The default is that this function computes covariance matrices corresponding to the "bmalite" models.

**Value**

a list of covariance matrices

**Examples**

```
data = mash_set_data(Bhat = cbind(c(1,2),c(3,4)), Shat = cbind(c(1,1),c(1,1)))
cov_canonical(data)
cov_canonical(data,"singletons")
cov_canonical(data,c("id","sing")) # can use partial matching of names
```

---

cov_ed	<i>Perform "extreme deconvolution" (Bovy et al) on a subset of the data</i>
--------	---

---

**Description**

Perform "extreme deconvolution" (Bovy et al) on a subset of the data

**Usage**

```
cov_ed(data, Ulist_init, subset = NULL, algorithm = c("bovy", "teem"), ...)
```

**Arguments**

data	a mash data object
Ulist_init	a named list of covariance matrices to use to initialize ED; default is to use matrices from PCs
subset	a subset of data to be used when ED is run (set to NULL for all the data)
algorithm	algorithm to run ED
...	other arguments to be passed to ED algorithm, see <a href="#">extreme_deconvolution</a> for algorithm 'bovy', or <a href="#">teem_wrapper</a> for algorithm 'teem'

**Details**

Runs the extreme deconvolution algorithm from Bovy et al (Annals of Applied Statistics) to estimate data-driven covariance matrices. It can be initialized with, for example running `cov_pca` with, say, 5 PCs.

**Examples**

```

data = mash_set_data(Bhat = cbind(c(1,2),c(3,4)), Shat = cbind(c(1,1),c(1,1)))
U_pca = cov_pca(data,2)
U_x = apply(data$Bhat, 2, function(x) x - mean(x))
U_xx = t(U_x) %*% U_x / nrow(U_x)
cov_ed(data,c(U_pca, list(xx = U_xx)))

```

---

cov_flash	<i>Perform Empirical Bayes Matrix Factorization using flashr, and return a list of candidate covariance matrices</i>
-----------	--

---

**Description**

Perform Empirical Bayes Matrix Factorization using flashr, and return a list of candidate covariance matrices

**Usage**

```

cov_flash(
  data,
  factors = c("default", "nonneg"),
  subset = NULL,
  remove_singleton = FALSE,
  tag = NULL,
  output_model = NULL,
  greedy_args = list(),
  backfit_args = list()
)

```

**Arguments**

data	A “mash” data object.
factors	If factors = "default", the factors and loadings are both unconstrained. If factors = "nonneg", the factors are constrained to be non-negative, and the loadings are unconstrained.
subset	Data samples (rows) used to estimate the covariances. Sset to NULL to use all the data.
remove_singleton	If remove_singleton = TRUE, factors corresponding to singleton matrices will be removed from the output.
tag	How to name the covariance matrices.
output_model	The fitted flash model will be saved to this file (using <a href="#">saveRDS</a> ).
greedy_args	List containing additional parameters passed to flashier::flash.add.greedy.
backfit_args	List containing additional parameters passed to flashier::flash.backfit.

**Value**

A list of covariance matrices.

**Examples**

```
# See https://stephenslab.github.io/mashr/articles/flash\_mash.html  
# for an example
```

---

cov_pca	<i>Perform PCA on data and return list of candidate covariance matrices</i>
---------	---

---

**Description**

Perform PCA on data and return list of candidate covariance matrices

**Usage**

```
cov_pca(data, npc, subset = NULL)
```

**Arguments**

data	a mash data object
npc	the number of PCs to use
subset	indices of the subset of data to use (set to NULL for all data)

**Value**

Returns a list of covariance matrices: the npc rank-one covariance matrices based on the first npc PCs, and the rank npc covariance matrix

**Examples**

```
data = mash_set_data(Bhat = cbind(c(1,2),c(3,4)), Shat = cbind(c(1,1),c(1,1)))  
cov_pca(data,2)
```

---

cov_udi	<i>Compute a list of covariance matrices corresponding to the "Unassociated", "Directly associated" and "Indirectly associated" models</i>
---------	--

---

## Description

Compute a list of covariance matrices corresponding to the "Unassociated", "Directly associated" and "Indirectly associated" models

## Usage

```
cov_udi(data, model = udi_model_matrix(n_conditions(data)))
```

## Arguments

data	a mash data object, eg as created by mash_set_data
model	a model matrix with R columns, where R is the number of conditions in the data; each row should be a vector of length R with elements "U", "D" and "I" indicating whether each effect is Unassociated, Directly associated or Indirectly associated

## Details

If model is specified then this returns the covariance matrices for those models. The default creates all possible models. For a description of the "Unassociated", "Directly associated" and "Indirectly associated" models see Stephens M (2013), A unified framework for Association Analysis with Multiple Related Phenotypes, PLoS ONE.

## Value

a named list of covariance matrices

## Examples

```
data = mash_set_data(Bhat = cbind(c(1,2),c(3,4)), Shat = cbind(c(1,1),c(1,1)))
cov_udi(data)
cov_udi(data,c('I','D'))
```

---

```
estimate_null_correlation_simple
```

*Estimate null correlations (simple)*

---

### Description

Estimates a null correlation matrix from data using simple z score threshold

### Usage

```
estimate_null_correlation_simple(data, z_thresh = 2, est_cor = TRUE)
```

### Arguments

data	a mash data object, eg as created by <code>mash_set_data</code>
z_thresh	the z score threshold below which to call an effect null
est_cor	whether to estimate correlation matrix (TRUE) or the covariance matrix (FALSE).

### Details

Returns a simple estimate of the correlation matrix (or covariance matrix) among conditions under the null. Specifically, the simple estimate is the empirical correlation (or covariance) matrix of the z scores for those effects that have (absolute) z score  $< z\_thresh$  in all conditions.

### Examples

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
estimate_null_correlation_simple(data)
```

---

extreme\_deconvolution *Density estimation using Gaussian mixtures in the presence of noisy, heterogeneous and incomplete data*

---

### Description

We present a general algorithm to infer a d-dimensional distribution function given a set of heterogeneous, noisy observations or samples. This algorithm reconstructs the error-deconvolved or 'underlying' distribution function common to all samples, even when the individual samples have unique error and missing-data properties. The underlying distribution is modeled as a mixture of Gaussians, which is completely general. Model parameters are chosen to optimize a justified, scalar objective function: the logarithm of the probability of the data under the error-convolved model, where the error convolution is different for each data point. Optimization is performed by an Expectation Maximization (EM) algorithm, extended by a regularization technique and 'split-and-merge' procedure. These extensions mitigate problems with singularities and local maxima, which are often encountered when using the EM algorithm to estimate Gaussian density mixtures.



**Usage**

```

extreme_deconvolution(
  ydata,
  ycovar,
  xamp,
  xmean,
  xcovar,
  projection = NULL,
  weight = NULL,
  fixamp = NULL,
  fixmean = NULL,
  fixcovar = NULL,
  tol = 1e-06,
  maxiter = 1e+09,
  w = 0,
  logfile = NULL,
  splitnmerge = 0,
  maxsnm = FALSE,
  likeonly = FALSE,
  logweight = FALSE
)

```

**Arguments**

ydata	[ndata,dy] matrix of observed quantities
ycovar	[ndata,dy] / [ndata,dy,dy] / [dy,dy,ndata] matrix, list or 3D array of observational error covariances (if [ndata,dy] then the error correlations are assumed to vanish)
xamp	[ngauss] array of initial amplitudes (*not* [1,ngauss])
xmean	[ngauss,dx] matrix of initial means
xcovar	[ngauss,dx,dx] list of matrices of initial covariances
projection	[ndata,dy,dx] list of projection matrices
weight	[ndata] array of weights to be applied to the data points
fixamp	(default=None) None, True/False, or list of bools
fixmean	(default=None) None, True/False, or list of bools
fixcovar	(default=None) None, True/False, or list of bools
tol	(double, default=1.e-6) tolerance for convergence
maxiter	(long, default= 10**9) maximum number of iterations to perform
w	(double, default=0.) covariance regularization parameter (of the conjugate prior)
logfile	basename for several logfiles (_c.log has output from the c-routine; _loglike.log has the log likelihood path of all the accepted routes, i.e. only parts which increase the likelihood are included, during splitnmerge)
splitnmerge	(int, default=0) depth to go down the splitnmerge path
maxsnm	(Bool, default=False) use the maximum number of split 'n' merge steps, $K*(K-1)*(K-2)/2$

likeonly (Bool, default=False) only compute the total log likelihood of the data  
 logweight (bool, default=False) if True, weight is actually log(weight)

**Value**

avgloglikedata avgloglikedata after convergence  
 xamp updated xamp  
 xmean updated xmean  
 xcovar updated xcovar

**Author(s)**

Jo Bovy, David W. Hogg, & Sam T. Roweis

**References**

Inferring complete distribution functions from noisy, heterogeneous and incomplete observations  
 Jo Bovy, David W. Hogg, & Sam T. Roweis, Submitted to AOAS (2009) [arXiv/0905.2979]

**Examples**

```
ydata <-
c(2.62434536, 0.38824359, 0.47182825, -0.07296862, 1.86540763,
-1.30153870, 2.74481176, 0.23879310, 1.31903910, 0.75062962,
2.46210794, -1.06014071, 0.67758280, 0.61594565, 2.13376944,
-0.09989127, 0.82757179, 0.12214158, 1.04221375, 1.58281521,
-0.10061918, 2.14472371, 1.90159072, 1.50249434, 1.90085595,
0.31627214, 0.87710977, 0.06423057, 0.73211192, 1.53035547,
0.30833925, 0.60324647, 0.31282730, 0.15479436, 0.32875387,
0.98733540, -0.11731035, 1.23441570, 2.65980218, 1.74204416,
0.80816445, 0.11237104, 0.25284171, 2.69245460, 1.05080775,
0.36300435, 1.19091548, 3.10025514, 1.12015895, 1.61720311,
1.30017032, 0.64775015, -0.14251820, 0.65065728, 0.79110577,
1.58662319, 1.83898341, 1.93110208, 1.28558733, 1.88514116,
0.24560206, 2.25286816, 1.51292982, 0.70190717, 1.48851815,
0.92442829, 2.13162939, 2.51981682, 3.18557541, -0.39649633,
-0.44411380, 0.49553414, 1.16003707, 1.87616892, 1.31563495,
-1.02220122, 0.69379599, 1.82797464, 1.23009474, 1.76201118,
0.77767186, 0.79924193, 1.18656139, 1.41005165, 1.19829972,
1.11900865, 0.32933771, 1.37756379, 1.12182127, 2.12948391,
2.19891788, 1.18515642, 0.62471505, 0.36126959, 1.42349435,
1.07734007, 0.65614632, 1.04359686, 0.37999916, 1.69803203,
0.55287144, 2.22450770, 1.40349164, 1.59357852, -0.09491185,
1.16938243, 1.74055645, 0.04629940, 0.73378149, 1.03261455,
-0.37311732, 1.31515939, 1.84616065, 0.14048406, 1.35054598,
-0.31228341, 0.96130449, -0.61577236, 2.12141771, 1.40890054,
0.97538304, 0.22483838, 2.27375593, 2.96710175, -0.85798186,
2.23616403, 2.62765075, 1.33801170, -0.19926803, 1.86334532,
0.81907970, 0.39607937, -0.23005814, 1.55053750, 1.79280687,
0.37646927, 1.52057634, -0.14434139, 1.80186103, 1.04656730,
0.81343023, 0.89825413, 1.86888616, 1.75041164, 1.52946532,
```

```

1.13770121, 1.07782113, 1.61838026, 1.23249456, 1.68255141,
0.68988323, -1.43483776, 2.03882460, 3.18697965, 1.44136444,
0.89984477, 0.86355526, 0.88094581, 1.01740941, -0.12201873,
0.48290554, 0.00297317, 1.24879916, 0.70335885, 1.49521132,
0.82529684, 1.98633519, 1.21353390, 3.19069973, -0.89636092,
0.35308331, 1.90148689, 3.52832571, 0.75136522, 1.04366899,
0.77368576, 2.33145711, 0.71269214, 1.68006984, 0.68019840,
-0.27255875, 1.31354772, 1.50318481, 2.29322588, 0.88955297,
0.38263794, 1.56276110, 1.24073709, 1.28066508, 0.92688730,
2.16033857, 1.36949272, 2.90465871, 2.11105670, 1.65904980,
-0.62743834, 1.60231928, 1.42028220, 1.81095167, 2.04444209)

ydata <- matrix(ydata,length(ydata),1)
N      <- dim(ydata)[1]
ycovar <- ydata*0 + 0.01
xamp   <- c(0.5,0.5)
xmean  <- matrix(c(0.86447943, 0.67078879, 0.322681, 0.45087394),2,2)
xcovar <-
  list(matrix(c(0.03821028, 0.04014796, 0.04108113, 0.03173839),2,2),
        matrix(c(0.06219194, 0.09738021, 0.04302473, 0.06778009),2,2))
projection <- list()
for (i in 1:N)
  projection[[i]] = matrix(c(i%2,(i+1)%2),1,2)
res <- extreme_deconvolution(ydata, ycovar, xamp, xmean, xcovar,
  projection=projection, logfile="ExDeconDemo")

```

---

get\_estimated\_pi      *Return the estimated mixture proportions*

---

## Description

Return the estimated mixture proportions

## Usage

```
get_estimated_pi(m, dimension = c("cov", "grid", "all"))
```

## Arguments

m	the mash result
dimension	indicates whether you want the mixture proportions for the covariances, grid, or all

## Details

If the fit was done with ‘usepointmass=TRUE’ then the first element of the returned vector will correspond to the null, and the remaining elements to the non-null covariance matrices. Suppose the fit was done with  $K \times K$  covariances and a grid of length  $L$ . If ‘dimension=cov’ then the

returned vector will be of length  $K$  (or  $K+1$  if 'usepointmass=TRUE'). If 'dimension=grid' then the returned vector will be of length  $L$  (or  $L+1$ ). If 'dimension=all' then the returned vector will be of length  $LK$  (or  $LK+1$ ). The names of the vector will be informative for which combination each element corresponds to.

### Value

a named vector containing the estimated mixture proportions.

### Examples

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
get_estimated_pi(m)
```

---

get\_log10bf

*Return the Bayes Factor for each effect*

---

### Description

Return the Bayes Factor for each effect

### Usage

```
get_log10bf(m)
```

### Arguments

**m** the mash result (from joint or 1by1 analysis); must have been computed using usepointmass=TRUE

### Value

if **m** was fitted using usepointmass=TRUE then returns a vector of the  $\log_{10}(\text{bf})$  values for each effect. That is, the  $j$ th element  $\text{lbf}[j]$  is  $\log_{10}(\text{Pr}(B_j | g=\text{ghat-nonnull})/\text{Pr}(B_j | g = 0))$  where  $\text{ghat-nonnull}$  is the non-null part of  $\text{ghat}$ . Otherwise returns NULL.

### Examples

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
get_log10bf(m)
```

---

get\_n\_significant\_conditions  
*Count number of conditions each effect is significant in*

---

**Description**

Count number of conditions each effect is significant in

**Usage**

```
get_n_significant_conditions(  
  m,  
  thresh = 0.05,  
  conditions = NULL,  
  sig_fn = get_lfsr  
)
```

**Arguments**

m	the mash result (from joint or lbyl analysis)
thresh	indicates the threshold below which to call signals significant
conditions	which conditions to include in check (default to all)
sig_fn	the significance function used to extract significance from mash object; eg could be ashrr::get_lfsr or ashrr::get_lfdr

**Value**

a vector containing the number of significant conditions

**Examples**

```
simdata = simple_sims(50,5,1)  
data = mash_set_data(simdata$Bhat, simdata$Shat)  
m = mash(data, cov_canonical(data))  
get_n_significant_conditions(m)
```

---

get\_pairwise\_sharing *Compute the proportion of (significant) signals shared by magnitude in each pair of conditions, based on the posterior mean*

---

**Description**

Compute the proportion of (significant) signals shared by magnitude in each pair of conditions, based on the posterior mean

**Usage**

```
get_pairwise_sharing(m, factor = 0.5, lfsr_thresh = 0.05, FUN = identity)
```

**Arguments**

m	the mash fit
factor	a number in [0,1] the factor within which effects are considered to be shared
lfsr_thresh	the lfsr threshold for including an effect in the assessment
FUN	a function to be applied to the estimated effect sizes before assessing sharing. The most obvious choice beside the default 'FUN=identity' would be 'FUN=abs' if you want to ignore the sign of the effects when assessing sharing.

**Details**

For each pair of tissues, first identify the effects that are significant (by  $lfsr < lfsr\_thresh$ ) in at least one of the two tissues. Then compute what fraction of these have an estimated (posterior mean) effect size within a factor 'factor' of one another. The results are returned as an R by R matrix.

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
get_pairwise_sharing(m) # sharing by magnitude (same sign)
get_pairwise_sharing(m, factor=0) # sharing by sign
get_pairwise_sharing(m, FUN=abs) # sharing by magnitude when sign is ignored
```

---

```
get_pairwise_sharing_from_samples
```

*Compute the proportion of (significant) signals shared by magnitude in each pair of conditions*

---

**Description**

Compute the proportion of (significant) signals shared by magnitude in each pair of conditions

**Usage**

```
get_pairwise_sharing_from_samples(
  m,
  factor = 0.5,
  lfsr_thresh = 0.05,
  FUN = identity
)
```

**Arguments**

m	the mash fit with samples from posteriors
factor	a number in [0,1] the factor within which effects are considered to be shared
lfsr_thresh	the lfsr threshold for including an effect in the assessment
FUN	a function to be applied to the estimated effect sizes before assessing sharing. The most obvious choice beside the default 'FUN=identity' would be 'FUN=abs' if you want to ignore the sign of the effects when assessing sharing.

**Details**

For each pair of conditions, compute the fraction of effects that are within a factor 'factor' of one another. The results are returned as an R by R matrix.

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data), posterior_samples=5, algorithm='R')
get_pairwise_sharing_from_samples(m) # sharing by magnitude (same sign)
get_pairwise_sharing_from_samples(m, factor=0) # sharing by sign
get_pairwise_sharing_from_samples(m, FUN=abs) # sharing by magnitude when sign is ignored
```

---

get_samples	<i>Return samples from a mash object</i>
-------------	--

---

**Description**

Return samples from a mash object

**Usage**

```
get_samples(m)
```

**Arguments**

m	The mash fit.
---	---------------

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data), posterior_samples=5, algorithm='R')
get_samples(m)
```

---

`get_significant_results`*Find effects that are significant in at least one condition*

---

**Description**

Find effects that are significant in at least one condition

**Usage**

```
get_significant_results(m, thresh = 0.05, conditions = NULL, sig_fn = get_lfsr)
```

**Arguments**

<code>m</code>	the mash result (from joint or 1by1 analysis)
<code>thresh</code>	indicates the threshold below which to call signals significant
<code>conditions</code>	which conditions to include in check (default to all)
<code>sig_fn</code>	the significance function used to extract significance from mash object; eg could be <code>ashr::get_lfsr</code> or <code>ashr::get_lfdr</code> . (Small values must indicate significant.)

**Value**

a vector containing the indices of the significant effects, by order of most significant to least

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
get_significant_results(m)
```

---

`mash`*Apply mash method to data*

---

**Description**

Apply mash method to data



**Usage**

```

mash(
  data,
  Ulist = NULL,
  gridmult = sqrt(2),
  grid = NULL,
  normalizeU = TRUE,
  usepointmass = TRUE,
  g = NULL,
  fixg = FALSE,
  prior = c("nullbiased", "uniform"),
  nullweight = 10,
  optmethod = c("mixSQP", "mixIP", "mixEM", "cxxMixSquarem"),
  control = list(),
  verbose = TRUE,
  add.mem.profile = FALSE,
  algorithm.version = c("Rcpp", "R"),
  pi_thresh = 1e-10,
  A = NULL,
  posterior_samples = 0,
  seed = 123,
  outputlevel = 2,
  output_lfdr = FALSE
)

```

**Arguments**

<code>data</code>	a mash data object containing the Bhat matrix, standard errors, alpha value; created using <code>mash_set_data</code> or <code>mash_set_data_contrast</code>
<code>Ulist</code>	a list of covariance matrices to use (see <code>normalizeU</code> for rescaling these matrices)
<code>gridmult</code>	scalar indicating factor by which adjacent grid values should differ; close to 1 for fine grid
<code>grid</code>	vector of grid values to use (scaling factors $\omega$ in paper)
<code>normalizeU</code>	whether or not to normalize the U covariances to have maximum of 1 on diagonal
<code>usepointmass</code>	whether to include a point mass at 0, corresponding to null in every condition
<code>g</code>	the value of $g$ obtained from a previous mash fit - an alternative to supplying <code>Ulist</code> , <code>grid</code> and <code>usepointmass</code>
<code>fixg</code>	if $g$ is supplied, allows the mixture proportions to be fixed rather than estimated; e.g., useful for fitting mash to test data after fitting it to training data
<code>prior</code>	indicates what penalty to use on the likelihood, if any
<code>nullweight</code>	scalar, the weight put on the prior under "nullbiased" specification, see "prior".
<code>optmethod</code>	name of optimization method to use
<code>control</code>	A list of control parameters passed to <code>optmethod</code> .
<code>verbose</code>	If TRUE, print progress to R console.

<code>add.mem.profile</code>	If TRUE, print memory usage to R console (requires R library ‘ <code>profmem</code> ’).
<code>algorithm.version</code>	Indicates whether to use R or Rcpp version
<code>pi_thresh</code>	threshold below which mixture components are ignored in computing posterior summaries (to speed calculations by ignoring negligible components)
<code>A</code>	the linear transformation matrix, $Q \times R$ matrix. This is used to compute the posterior for $Ab$ .
<code>posterior_samples</code>	the number of samples to be drawn from the posterior distribution of each effect.
<code>seed</code>	A random number seed to use when sampling from the posteriors. It is used when <code>posterior_samples &gt; 0</code> .
<code>outputlevel</code>	controls amount of computation / output; 1: output only estimated mixture component proportions, 2: and posterior estimates, 3: and posterior covariance matrices, 4: and likelihood matrices
<code>output_lfdr</code>	If <code>output_lfdr = TRUE</code> , output local false discovery rate estimates. The lfdr tends to be sensitive to mis-estimated covariance matrices, and generally we do not recommend using them; we recommend using the local false sign rate (lfsr) instead, which is always returned, even when <code>output_lfdr = TRUE</code> .

## Value

a list with elements `result`, `loglik` and `fitted_g`

## Examples

```
Bhat = matrix(rnorm(100),ncol=5) # create some simulated data
Shat = matrix(rep(1,100),ncol=5)
data = mash_set_data(Bhat,Shat, alpha=1)
U.c = cov_canonical(data)
res.mash = mash(data,U.c)

# Run mash with penalty exponent on null term equal to 100.
# See "False discovery rates: a new deal" (M. Stephens 2017),
# supplementary material S.2.5 for more details.
set.seed(1)
simdata = simple_sims(500,5,1)
data = mash_set_data(simdata$Bhat,simdata$Shat)
U.c = cov_canonical(data)
res0 = mash(data,U.c)
res1 = mash(data,U.c,prior = "nullbiased",nullweight = 101)
plot(res0$fitted_g$pi,res1$fitted_g$pi,pch = 20)
abline(a = 0,b = 1,col = "skyblue",lty = "dashed")
```

---

mash_1by1	<i>Perform condition-by-condition analyses</i>
-----------	--

---

**Description**

Performs simple "condition-by-condition" analysis by running ash from package ash on data from each condition, one at a time. May be a useful first step to identify top hits in each condition before a mash analysis.

**Usage**

```
mash_1by1(data, alpha = 0, ...)
```

**Arguments**

data	A list with the following two elements: Bhat an n by R matrix of observations (n units in R conditions); and Shat, an n by R matrix of standard errors (n units in R conditions),
alpha	Numeric value of alpha parameter in the model. alpha = 0 for Exchangeable Effects (EE), alpha = 1 for Exchangeable Z-scores (EZ).
...	optionally, other parameters to be passed to ash

**Value**

A list similar to the output of mash, particularly including posterior matrices.

**Examples**

```
simdata = simple_sims(50,5,1)
mash_1by1(simdata)
```

---

mash_compute_loglik	<i>Compute loglikelihood for fitted mash object on new data.</i>
---------------------	--

---

**Description**

Compute loglikelihood for fitted mash object on new data.

**Usage**

```
mash_compute_loglik(g, data, algorithm.version = c("Rcpp", "R"))
```

**Arguments**

`g` A mash object or the fitted\_g from a mash object.  
`data` A set of data on which to compute the loglikelihood.  
`algorithm.version` Indicate R or Rcpp version

**Details**

The log-likelihood for each element is  $p(Bhat_j|Shat_j, g, \alpha)$  where  $Bhat_j|B_j, Shat_j \sim N(B_j, Shat_j)$  and  $B_j/Shat_j^\alpha|Shat_j \sim g$ .

**Value**

The log-likelihood for data computed using `g`.

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
mash_compute_loglik(m, data)
```

---

mash\_compute\_posterior\_matrices

*Compute posterior matrices for fitted mash object on new data*

---

**Description**

Compute posterior matrices for fitted mash object on new data

**Usage**

```
mash_compute_posterior_matrices(
  g,
  data,
  pi_thresh = 1e-10,
  algorithm.version = c("Rcpp", "R"),
  A = NULL,
  output_posterior_cov = FALSE,
  posterior_samples = 0,
  seed = 123
)
```

**Arguments**

<code>g</code>	a mash object or the fitted_g from a mash object.
<code>data</code>	a set of data on which to compute the posterior matrices
<code>pi_thresh</code>	threshold below which mixture components are ignored in computing posterior summaries (to speed calculations by ignoring negligible components)
<code>algorithm.version</code>	Indicates whether to use R or Rcpp version
<code>A</code>	the linear transformation matrix, Q x R matrix. This is used to compute the posterior for Ab.
<code>output_posterior_cov</code>	whether or not to output posterior covariance matrices for all effects
<code>posterior_samples</code>	the number of samples to be drawn from the posterior distribution of each effect.
<code>seed</code>	a random number seed to use when sampling from the posteriors. It is used when <code>posterior_samples &gt; 0</code> .

**Value**

A list of posterior matrices

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
mash_compute_posterior_matrices(m,data)
```

---

`mash_compute_vloglik` *Compute vector of loglikelihood for fitted mash object on new data*

---

**Description**

Compute vector of loglikelihood for fitted mash object on new data

**Usage**

```
mash_compute_vloglik(g, data, algorithm.version = c("Rcpp", "R"))
```

**Arguments**

<code>g</code>	A mash object.
<code>data</code>	A set of data on which to compute the loglikelihood.
<code>algorithm.version</code>	Indicate R or Rcpp version

**Details**

The log-likelihood for each element is  $p(Bhat_j|Shat_j, g, \alpha)$  where  $Bhat_j|B_j, Shat_j \sim N(B_j, Shat_j)$  and  $B_j/Shat_j^\alpha|Shat_j \sim g$ . Here the value of  $\alpha$  is set when setting up the data object in 'mash\_set\_data'. If  $g$  is a mash object (safest!) then the function will check that this value matches the  $\alpha$  used when fitting 'mash'. Note: as a convenience, this function can also be called with  $g$  a mixture distribution with same structure as the fitted\_ $g$  from a mash object. This is mostly useful when doing simulations, where you might want to compute the likelihood under the "true"  $g$ . When used in this way the user is responsible for making sure that the  $g$  makes sense with the alpha set in data.

**Value**

The vector of log-likelihoods for each data point computed using  $g$ .

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
mash_compute_vloglik(m,data)
```

---

mash\_estimate\_corr\_em *Fit mash model and estimate residual correlations using EM algorithm*

---

**Description**

Estimates a residual correlation matrix from data using an ad hoc EM algorithm.

**Usage**

```
mash_estimate_corr_em(
  data,
  Ulist,
  init,
  max_iter = 30,
  tol = 1,
  est_cor = TRUE,
  track_fit = FALSE,
  prior = c("nullbiased", "uniform"),
  details = TRUE,
  ...
)
```

**Arguments**

data	a mash data object, eg as created by <code>mash_set_data</code>
Ulist	a list of covariance matrices to use
init	the initial value for the residual correlation. If it is not given, we use result from <code>estimate_null_correlation_simple</code>
max_iter	maximum number of iterations to perform
tol	convergence tolerance
est_cor	whether to estimate correlation matrix (TRUE) or the covariance matrix (FALSE)
track_fit	add an attribute trace to output that saves current values of all iterations
prior	indicates what penalty to use on the likelihood, if any
details	whether to return details of the model, if it is TRUE, the mash model, the number of iterations and the value of objective functions will be returned
...	other parameters pass to mash

**Details**

Returns the estimated residual correlation matrix among conditions. We estimate the residual correlation matrix using an ad hoc em algorithm. The update in the ad hoc M step is not guaranteed to increase the likelihood, therefore, the EM algorithm is stopped before the likelihood drops. The residual correlation matrix  $V$  is estimated using the posterior second moment of the noise.

Warning: This method could take some time. The `estimate_null_correlation_simple` gives a quick approximation for the null correlation matrix.

**Value**

the estimated correlation matrix and the fitted mash model

V	estimated residual correlation matrix
mash.model	fitted mash model

**Examples**

```
simdata = simple_sims(100,5,1)
m.1by1 = mash_1by1(mash_set_data(simdata$Bhat,simdata$Shat))
strong.subset = get_significant_results(m.1by1,0.05)
random.subset = sample(1:nrow(simdata$Bhat),20)
data.strong = mash_set_data(simdata$Bhat[strong.subset,], simdata$Shat[strong.subset,])
data.tmp = mash_set_data(simdata$Bhat[random.subset,], simdata$Shat[random.subset,])
U_pca = cov_pca(data.strong, 3)
U_ed = cov_ed(data.strong, U_pca)
Vhat = mash_estimate_corr_em(data.tmp, U_ed)
```

---

mash_plot_meta	<i>Plot metaplot for an effect based on posterior from mash</i>
----------------	---

---

**Description**

Plot metaplot for an effect based on posterior from mash

**Usage**

```
mash_plot_meta(m, i, xlab = "Effect size", ylab = "Condition", ...)
```

**Arguments**

m	the result of a mash fit
i	index of the effect to plot
xlab	Character string specifying x-axis label.
ylab	Character string specifying y-axis label.
...	Additional arguments passed to <a href="#">metaplot</a> .

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
m = mash(data, cov_canonical(data))
mash_plot_meta(m,1)
```

---

mash_set_data	<i>Create a data object for mash analysis.</i>
---------------	--

---

**Description**

Create a data object for mash analysis.

**Usage**

```
mash_set_data(
  Bhat,
  Shat = NULL,
  alpha = 0,
  df = Inf,
  pval = NULL,
  V = diag(ncol(Bhat)),
  zero_check_tol = .Machine$double.eps,
  zero_Bhat_Shat_reset = 0,
  zero_Shat_reset = 0
)
```



**Arguments**

Bhat	An N by R matrix of observed estimates.
Shat	An N by R matrix of corresponding standard errors. Shat can be a scalar if all standard errors are equal. This is most useful if Bhat is a matrix of Z scores, so elements of Shat are all 1. Default is 1.
alpha	Numeric value of alpha parameter in the model. alpha = 0 for Exchangeable Effects (EE), alpha = 1 for Exchangeable Z-scores (EZ). Default is 0. Please refer to equation (3.2) of M. Stephens 2016, Biostatistics for a discussion on alpha.
df	An N by R matrix of corresponding degrees of freedom of the t-statistic Bhat/Shat. Can be a scalar if all degrees of freedom are equal. Default is inf (for large samples).
pval	An N by R matrix of p-values of t-statistic Bhat/Shat. Shat and df should not be specified when pval is provided.
V	an R by R matrix / [R x R x N] array of effect specific correlation matrix of error correlations; must be positive definite. [So Bhat_j distributed as $N(B\_j, \text{diag}(Shat\_j) V[,j] \text{diag}(Shat\_j))$ where $\_j$ denotes the jth row of a matrix]. Defaults to identity.
zero_check_tol	a small positive number as threshold for Shat to be considered zero if any Shat is smaller or equal to this number.
zero_Bhat_Shat_reset	Replace zeros in Shat matrix to given value if the corresponding Bhat are also zeros.
zero_Shat_reset	Replace zeros in Shat matrix to given value.

**Value**

A data object for passing into mash functions.

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
```

---

mash_update_data	<i>Update the data object for mash analysis.</i>
------------------	--

---

**Description**

This function can update two parts of the mash data. The first one is setting the reference group, so the mash data can be used for commonbaseline analysis. The other one is updating the null correlation matrix.

**Usage**

```
mash_update_data(mashdata, ref = NULL, V = NULL)
```

**Arguments**

mashdata	mash data object containing the Bhat matrix, standard errors, V; created using mash_set_data
ref	the reference group. It could be a number between 1,..., R, R is number of conditions, or the name of reference group. If there is no reference group, it can be the string 'mean'.
V	an R by R matrix / [R x R x N] array of correlation matrix of error correlations

**Value**

a updated mash data object

**Examples**

```
simdata = simple_sims(50,5,1)
data = mash_set_data(simdata$Bhat, simdata$Shat)
mash_update_data(data, 'mean')
```

---

simple\_sims

*Create some simple simulated data for testing purposes*

---

**Description**

Create some simple simulated data for testing purposes

**Usage**

```
simple_sims(nsamp = 100, ncond = 5, err_sd = 0.01)
```

**Arguments**

nsamp	number of samples of each type
ncond	number of conditions
err_sd	the standard deviation of the errors

**Details**

The simulation consists of equal numbers of four different types of effects: null, equal among conditions, present only in first condition, independent across conditions

**Examples**

```
simple_sims(100, 5)
```

---

simple_sims2	<i>Create some more simple simulated data for testing purposes</i>
--------------	--

---

**Description**

Create some more simple simulated data for testing purposes

**Usage**

```
simple_sims2(nsamp = 100, err_sd = 0.01)
```

**Arguments**

nsamp	number of samples of each type
err_sd	the standard deviation of the errors

**Details**

The simulation consists of five conditions with two types of effects those present (and identical) in first two conditions and those present (and identical) in last three conditions

**Examples**

```
simple_sims2(100, 5)
```

---

sim_contrast1	<i>Create simplest simulation, <math>c_j = \mu_1</math> data used for contrast analysis</i>
---------------	---

---

**Description**

Create simplest simulation,  $c_j = \mu_1$  data used for contrast analysis

**Usage**

```
sim_contrast1(nsamp = 100, ncond = 5, err_sd = sqrt(0.5))
```

**Arguments**

nsamp	number of samples of each type
ncond	number of conditions
err_sd	the standard deviation of the errors

**Details**

There is no true deviation exists in this case

**Examples**

```
sim_contrast1(100,5)
```

---

sim_contrast2	<i>Create simulation with signal data used for contrast analysis.</i>
---------------	---

---

**Description**

Create simulation with signal data used for contrast analysis.

**Usage**

```
sim_contrast2(nsamp = 1000, ncond = 5, err_sd = sqrt(0.5))
```

**Arguments**

nsamp	Number of samples of each type.
ncond	Number of conditions.
err_sd	The standard deviation of the errors.

**Details**

The first condition is the reference group. The deviations are the difference between the subsequent conditions with the reference group. The simulation consists of 90 10 different types of deviations: equal among conditions, present only in the first subsequent condition, independent across conditions.

**Examples**

```
sim_contrast2(100,5)
```

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