

# plw

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AffySpikeU95Subset *Spike-in data*

---

### Description

This `AffyBatch-class` represents part of the Affymetrix spike-in data set.

### Usage

```
data(AffySpikeU95Subset)
```

### Format

An `AffyBatch-class` containing 6 arrays.

### Source

Array 1, 2, 21, 22, 41, and 42 of the Affymetrix U95 Latin square data set, thus the 6 arrays of group A and B, and a random subset of 1000 probe-sets together with the 16 spike-in probe-sets.

---

estimateMVbeta      *Zero mean multivariate t-dist. with covariate dependent scale.*

---

### Description

Estimate the parameters  $m$  and  $\nu$  of the multivariate t-distribution with zero expectation, where  $\nu$  is modeled as smooth function of a covariate. The covariance matrix  $\Sigma$  is assumed to be known.

### Usage

```
estimateMVbeta(y, x, Sigma, maxIter = 200, epsilon = 1e-06,
               verbose = FALSE, nknots = 10, nOut = 2000, nIn = 4000,
               iterInit = 3, br = NULL)
```

### Arguments

y	Data matrix
x	Covariate vector
Sigma	Covariance matrix
maxIter	Maximum number of iterations
epsilon	Convergence criterion
verbose	Print computation info or not
nknots	Number of knots of spline for $\nu$
nOut	Parameter for calculating knots, see getKnots
nIn	Parameter for calculating knots, see getKnots
iterInit	Number of iteration in when initiating $\Sigma$
br	Knots, overrides nknots, n.out and n.in

### Details

The multivariate t-distribution is parametrized as:

$$y|c \sim N(\mu, c\Sigma)$$

$$c \sim \text{InvGamma}(m/2, m\nu/2)$$

where  $\nu$  is function of the covariate  $x$ :  $\nu(x)$  and  $N$  denotes a multivariate normal distribution,  $\Sigma$  is a covariance matrix and  $\text{InvGamma}(\alpha, \beta)$  is the inverse-gamma distribution with density function

$$f(x) = (\beta)^\alpha \exp\{-\beta/x\} x^{-\alpha-1} / \Gamma(\alpha)$$

A cubic spline is used to parameterize the smooth function  $\nu(x)$

$$\nu(x) = \exp\{H(x)^T \beta\}$$

where  $H : R \rightarrow R^{2p-1}$  is a set B-spline basis functions for a given set of  $p$  interior spline-knots, see chapter 5 of Hastie (2001). In this application  $\mu$  equals zero, and  $m$  is the degrees of freedom.

**Value**

sigma	The input covariance matrix for y
m	Estimated shape parameter for inverse-gamma prior for gene variances
v	Estimated scale parameter curve for inverse-gamma prior for gene variances
converged	TRUE if the EM algorithms converged
iter	Number of iterations
modS2	Moderated estimator of gene-specific variances
histLogS2	Histogram of log(s2) where s2 is the ordinary variance estimator
fittedDensityLogS2	The fitted density for log(s2)
logs2	Variance estimators, logged with base 2.
beta	Estimated parameter vector $\beta$ of spline for $\nu(x)$
knots	The knots used in spline for $\nu(x)$
x	The input vector covariate vector x

**Author(s)**

Magnus Åstrand

**References**

- Hastie, T., Tibshirani, R., and Friedman, J. (2001). The Elements of Statistical Learning, volume 1. Springer, first edition.
- Kristiansson, E., Sjögren, A., Rudemo, M., Nerman, O. (2005). Weighted Analysis of Paired Microarray Experiments. Statistical Applications in Genetics and Molecular Biology 4(1)
- Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. Journal of Computational Biology, Accepted.
- Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. Bioinformatics, Submitted 1 October 2007.

**See Also**

plw, lmw, estimateSigmaMVbeta

---

estimateSigmaMVbeta

*Zero mean multivariate t-dist. with covariate dependent scale.*

---

**Description**

Estimate the parameters  $\Sigma$ ,  $m$  and  $\nu$  of the multivariate t-distribution with zero expectation, where  $\nu$  is modeled as smooth function of a covariate.

**Usage**

```
estimateSigmaMVbeta(y, x, maxIter = 200, epsilon = 1e-06,
  verbose = FALSE, nknots = 10, nOut = 2000, nIn = 4000,
  iterInit = 3, br = NULL)
```

**Arguments**

y	Data matrix
x	Covariate vector
maxIter	Maximum number of iterations
epsilon	Convergence criteria
verbose	Print computation info or not
nknots	Number of knots of spline for $\nu$
nOut	Parameter for calculating knots, see getKnots
nIn	Parameter for calculating knots, see getKnots
iterInit	Number of iteration in when initiating $\Sigma$
br	Knots, overrides nknots, n.out and n.in

**Details**

The multivariate t-distribution is parametrized as:

$$y|c \sim N(\mu, c\Sigma)$$

$$c \sim \text{InvGamma}(m/2, m\nu/2)$$

where  $\nu$  is function of the covariate  $x$ :  $\nu(x)$  and  $N$  denotes a multivariate normal distribution,  $\Sigma$  is a covariance matrix and  $\text{InvGamma}(a, b)$  is the inverse-gamma distribution with density function

$$f(x) = (b)^a \exp\{-b/x\} x^{-a-1} / \Gamma(a)$$

A cubic spline is used to parameterize the smooth function  $\nu(x)$

$$\nu(x) = \exp\{H(x)^T \beta\}$$

where  $H : R \rightarrow R^{2p-1}$  is a set B-spline basis functions for a given set of  $p$  interior spline-knots, see chapter 5 of Hastie et al. (2001). In this application  $\mu$  equals zero, and  $m$  is the degrees of freedom.

For details about the model see Kristiansson et al. (2005), Åstrand et al. (2007a,2007b).

**Value**

Sigma	Estimated covariance matrix for y
m	Estimated shape parameter for inverse-gamma prior for gene variances
v	Estimated scale parameter curve for inverse-gamma prior for gene variances
converged	T if the EM algorithms converged
iter	Number of iterations
modS2	Moderated estimator of gene-specific variances
histLogS2	Histogram of log(s2) where s2 is the ordinary variance estimator
fittedDensityLogS2	The fitted density for log(s2)
logs2	Variance estimators, logged with base 2.
beta	Estimated parameter vector $\beta$ of spline for $\nu(x)$
knots	The knots used in spline for $\nu(x)$
x	The input vector covariate vector x

**Author(s)**

Magnus Åstrand

**References**

Hastie, T., Tibshirani, R., and Friedman, J. (2001). The Elements of Statistical Learning, volume 1. Springer, first edition.

Kristiansson, E., Sjögren, A., Rudemo, M., Nerman, O. (2005). Weighted Analysis of Paired Microarray Experiments. Statistical Applications in Genetics and Molecular Biology 4(1)

Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. Journal of Computational Biology, Accepted.

Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. Bioinformatics, Submitted 1 October 2007.

**See Also**

plw, lmw

---

 estimateSigmaMV     *Fit zero mean multivariate t-distribution*


---

**Description**

estimate the parameters  $\Sigma$ ,  $m$  and  $\nu$  of the multivariate t-distribution with zero expectation.

**Usage**

```
estimateSigmaMV(y, maxIter=100, epsilon=0.000001, verbose=FALSE)
```

**Arguments**

<code>y</code>	data matrix
<code>maxIter</code>	maximum number of iterations
<code>epsilon</code>	convergence criteria
<code>verbose</code>	print computation info or not

**Details**

The multivariate t-distribution is parametrized as:

$$y|c \sim N(\mu, c\Sigma)$$

$$c \sim \text{InvGamma}(m/2, m\nu/2)$$

Here  $N$  denotes a multivariate normal distribution,  $\Sigma$  is a covariance matrix and  $\text{InvGamma}(\alpha, \beta)$  is the inverse-gamma distribution with density function

$$f(x) = (\beta)^\alpha \exp\{-\beta/x\}x^{-\alpha-1}/\Gamma(\alpha)$$

In this application  $\mu$  equals zero, and  $m$  is the degrees of freedom.

**Value**

<code>Sigma</code>	Estimated covariance matrix for $y$
<code>m</code>	Estimated shape parameter for inverse-gamma prior for gene variances
<code>v</code>	Estimated scale parameter for inverse-gamma prior for gene variances
<code>converged</code>	T if the EM algorithms converged
<code>iter</code>	Number of iterations
<code>modS2</code>	Moderated estimator of gene-specific variances
<code>histLogS2</code>	Histogram of $\log(s^2)$ where $s^2$ is the ordinary variance estimator
<code>fittedDensityLogS2</code>	The fitted density for $\log(s^2)$

**Author(s)**

Magnus Åstrand

**References**

Hastie, T., Tibshirani, R., and Friedman, J. (2001). The Elements of Statistical Learning, volume 1. Springer, first edition.

Kristiansson, E., Sjögren, A., Rudemo, M., Nerman, O. (2005). Weighted Analysis of Paired Microarray Experiments. Statistical Applications in Genetics and Molecular Biology 4(1)

Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. Journal of Computational Biology, Accepted.

Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. Bioinformatics, Submitted 1 October 2007.

**See Also**

estimateSigma

---

estimateSigma      *Fit zero mean multivariate t-distribution, known df*

---

**Description**

Estimate the covariance matrix  $\Sigma$  of the multivariate t-distribution with zero expectation assuming the degrees of freedom is known.

**Usage**

```
estimateSigma(y, m, v, maxIter = 100, epsilon = 1e-06, verbose = FALSE)
```

**Arguments**

<code>y</code>	data matrix
<code>m</code>	degrees of freedom
<code>v</code>	scale parameter
<code>maxIter</code>	maximum number of iterations
<code>epsilon</code>	convergence criteria
<code>verbose</code>	print computation info or not

**Details**

The multivariate t-distribution is parametrized as:

$$y|c \sim N(\mu, c\Sigma)$$

$$c \sim \text{InvGamma}(m/2, m\nu/2)$$

Here  $N$  denotes a multivariate normal distribution,  $\Sigma$  is a covariance matrix and  $\text{InvGamma}(\alpha, \beta)$  is the inverse-gamma distribution with density function

$$f(x) = (\beta)^\alpha \exp\{-\beta/x\}x^{-\alpha-1}/\Gamma(\alpha)$$

In this application  $\mu$  equals zero, and  $m$  is the degrees of freedom.

**Value**

<code>Sigma</code>	Estimated covariance matrix for y
<code>iter</code>	Number of iterations

**Author(s)**

Magnus Åstrand

**References**

- Hastie, T., Tibshirani, R., and Friedman, J. (2001). The Elements of Statistical Learning, volume 1. Springer, first edition.
- Kristiansson, E., Sjögren, A., Rudemo, M., Nerman, O. (2005). Weighted Analysis of Paired Microarray Experiments. *Statistical Applications in Genetics and Molecular Biology* 4(1)
- Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. *Journal of Computational Biology*, Accepted.
- Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. *Bioinformatics*, Submitted 1 October 2007.

**See Also**

`estimateSigmaMV`

---

`getKnots`

*Spline-knots for plw and lmw*

---

**Description**

Computes a set of `nKnots` interior knots( if possible) plus 2 boundary knots so that:

- 1) the `nOut` smallest and highest data points (in `x`) lies below and above the lower and upper boundary knots respectively.
- 2) there is at least `nIn` data points between all knots.

**Usage**

```
getKnots(x, nKnots=10, nOut=2000, nIn=4000)
```

**Arguments**

<code>x</code>	Data vector
<code>nKnots</code>	Number of interior knots
<code>nOut</code>	Number of data points below and above the lower and upper boundary knots respectively.
<code>nIn</code>	Number of data points between knots.

**Details**

See the definition (R-code) for details.

**Value**

A vector of knots.

**Author(s)**

Magnus Åstrand

**See Also**

`plw`, `lmw`, `estimateSigmaMVbeta`

---

HowToPLW

*View HowToPLW*

---

**Description**

Finds the location of the vignette HowToPLW and optionally opens it.

**Usage**

```
HowToPLW(view=TRUE)
```

**Arguments**

<code>view</code>	logical, should the document be opened using the default PDF document reader?
-------------------	---

**Details**

If the operating system is other than Windows, then the PDF viewer used is that given by `Sys.getenv("R_PDFVIEWER")`. The PDF viewer can be changed using `Sys.putenv(R_PDFVIEWER=)`.

This function is used by drop-down Vignettes menu when the Rgui interface for Windows is used.

**Value**

Character string giving the file location.

**Author(s)**

Magnus Astrand



**See Also**

[vignette](#), [openPDF](#), [openVignette](#), `Sys.getenv`, `Sys.putenv`

**Examples**

```
HowToPLW(view=FALSE)
```

---

lmw

*Locally Moderated Weighted-t.*


---

**Description**

Computes Locally Moderated Weighted t-test for microarray data.

**Usage**

```
lmw(x, design=rep(1, ncol(x)), contrast=matrix(1), meanX=NULL,
     maxIter = 200, epsilon = 1e-06, verbose = TRUE,
     nknots = 10, nOut = 2000, nIn = 4000, knots = NULL,
     checkRegulation = TRUE)
```

**Arguments**

<code>x</code>	Data, log2 expression indexes.
<code>design</code>	design matrix
<code>contrast</code>	contrast matrix
<code>meanX</code>	Covariate used to model scale parameter, default=NULL (see details)
<code>maxIter</code>	maximum number of iterations
<code>epsilon</code>	convergence criteria
<code>verbose</code>	print computation info or not
<code>nknots</code>	Number of knots of spline for $\nu$
<code>nOut</code>	Parameter for calculating knots, see <code>getKnots</code>
<code>nIn</code>	Parameter for calculating knots, see <code>getKnots</code>
<code>knots</code>	Knots, if not NULL it overrides <code>nknots</code> , <code>nOut</code> and <code>nIn</code>
<code>checkRegulation</code>	If TRUE, data is checked for a correct specified contrast (see details)

**Details**

This function computes the Locally Moderated Weighted-t statistic (LMW) described in Åstrand (2007b), thus calculating locally moderated weighted t-statistic, p-value and log2(FC) for each row of the data matrix `x`.

Each gene  $g$  (row of `x`) is modeled as:

$$y_g | c_g \sim N(\mu_g, c_g \Sigma)$$

$$c_g \sim \text{InvGamma}(m/2, m\nu/2)$$

where  $\nu$  is function of the mean intensity:  $\nu(\bar{\mu}_g)$ ,  $N$  denotes a multivariate normal distribution,  $\Sigma$  is a covariance matrix and  $\text{InvGamma}(a, b)$  is the inverse-gamma distribution with density function

$$f(x) = (b)^a \exp\{-b/x\} x^{-a-1} / \Gamma(a)$$

Given the design matrix  $D$ ,  $\mu_g$  equals  $D\gamma_g$ , and given the contrast matrix  $C$  the hypothesis  $C\gamma_g = 0$  is tested.  $C$  should be a one row matrix of same length as the column vector  $\gamma_g$ .

See examples on how to specify the design and contrast matrices.

A cubic spline is used to parameterize the smooth function  $\nu(x)$

$$\nu(x) = \exp\{H(x)^T \beta\}$$

where  $H : R \rightarrow R^{2p-1}$  is a set B-spline basis functions for a given set of  $p$  interior spline-knots, see chapter 5 of Hastie et al. (2001).

For details about the model see Kristiansson et al. (2005), Åstrand et al. (2007a,2007b).

As specified above,  $\nu$  is modeled as a function of mean intensity:  $\nu(\bar{\mu}_g)$ . If the parameter meanX is not NULL, meanX is used instead of the mean intensity when modeling  $\nu$ . Thus, if meanX is not NULL, meanX must be a vector of length equal to the number of rows of the data matrix  $x$ .

The parameter estimation procedure is based on the assumption that the specified contrast is close to zero for most genes, or at least that the median contrast over all genes is close to zero. A check is run on data to validate this assumptions. If the checking fails, with the error message "warning: most genes appears to be regulated..." and if YOU ARE SURE that the design and contrast is correct, use `checkRegulation=FALSE`.

## Value

<code>Sigma</code>	Estimated covariance matrix for $y = P^T x$
<code>m</code>	Estimated shape parameter for inverse-gamma prior for gene variances
<code>v</code>	Estimated scale parameter curve for inverse-gamma prior for gene variances
<code>converged</code>	T if the EM algorithms converged
<code>iter</code>	Number of iterations
<code>modS2</code>	Moderated estimator of gene-specific variances
<code>histLogS2</code>	Histogram of $\log(s2)$ where $s2$ is the ordinary variance estimator
<code>fittedDensityLogS2</code>	The fitted density for $\log(s2)$
<code>logs2</code>	Variance estimators, logged with base 2.
<code>t</code>	Moderated t-statistic
<code>coefficients</code>	Estimated contrast
<code>p.value</code>	P-value from the moderated t-statistic
<code>dfT</code>	Degrees of freedom of the moderated t-statistic
<code>weights</code>	Weights for estimating the contrast
<code>P</code>	Transformation matrix
<code>beta</code>	Estimated parameter vector $\beta$ of spline for $\nu(x)$
<code>knots</code>	The knots used in spline for $\nu(x)$

## Author(s)

Magnus Åstrand

## References

Hastie, T., Tibshirani, R., and Friedman, J. (2001). The Elements of Statistical Learning, volume 1. Springer, first edition.

Kristiansson, E., Sjögren, A., Rudemo, M., Nerman, O. (2005). Weighted Analysis of Paired Microarray Experiments. Statistical Applications in Genetics and Molecular Biology 4(1)

Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. Journal of Computational Biology, Accepted.strand

Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. Bioinformatics, Submitted 1 October 2007.

## See Also

estimateSigmaMVbeta, plw

## Examples

```
# -----
# Example analyzing the 6 arrays in the
# AffySpikeU95Subset data set

# Loading the data
data(AffySpikeU95Subset)

# Defining design and contrast matrix
group<-factor(rep(1:2,each=3))
design<-model.matrix(~group-1)
contrast<-matrix(c(1,-1),1,2)

# Computing RMA expression index
data.rma<-exprs(rma(AffySpikeU95Subset))

# Analyzing
modell<-lmw(data.rma,design=design,contrast=contrast,epsilon=0.01)

## Look at fitted vs observed density for log(s2)
varHistPlot(modell)

## Look at fitted curve for scale parameter
scaleParameterPlot(modell)
```

---

logitT

*logit-t and t-test by row*

---

## Description

Functions for the logit-t test (Lemon et al. 2003) and the ordinary t-test computed for each row of an matrix.

**Usage**

```
logitTTransform(pm)
logitTStat(affy.batch, group)
studenttTTest(x, group)
```

**Arguments**

pm	A matrix of Pm intensities
affy.batch	An AffyBatch object
group	A group indicator vector, should have values 1 and 2 only.
x	A matrix

**Details**

See the definition (R-code) of each function for details.

**Value**

logitTTransform returns a matrix  
 logitTStat returns a vector with the logit-t statistic for each probe set.  
 studenttTTest returns a vector with t-statistic for each row of x.

**Author(s)**

Magnus Åstrand

**References**

Lemon et al. (2003). A high performance test of differential gene expression for oligonucleotide arrays. *Genome Biol.* 2003; 4(10):R67

**Examples**

```
# -----
# Example analyzing the 6 arrays in the
# AffySpikeU95Subset data set

# Loading the data
data(AffySpikeU95Subset)

# Vector with groups assignment
group<-factor(rep(1:2, each=3))

# logit-T statistic
logitT<-logitTStat(AffySpikeU95Subset,
                  as.numeric(group))

# Computing RMA expression index
data.rma<-exprs(rma(AffySpikeU95Subset))

# Ordinary t-test by row/gene
```

```

studentT<-studenttTTest(data.rma, as.numeric(group))

# Comparing genes ranked top-20
logitTTop20 <- rank(-abs(logitT)) < 21
studentTTop20<- rank(-abs(studentT)) < 21
table(logitTTop20,studentTTop20)

```

---

plw-package	<i>plw</i>
-------------	------------

---

### Description

Locally moderated weighted analysis of microarray data, at the probe level (PLW) or at the level of expression indexes (LMW).

### Author(s)

Magnus Åstrand

### References

Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. *Journal of Computational Biology*, Accepted.

Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. *Bioinformatics*, Submitted 1 October 2007.

### See Also

plw, lwm

---

plw	<i>Probe level Locally moderated Weighted median-t.</i>
-----	---

---

### Description

Computes locally moderated weighted median t-test for microarray data.

### Usage

```

plw(x, design=rep(1, ncol(x)), contrast=matrix(1),
    probenames = unlist(ifelse(class(x) == "AffyBatch",
                               list(p = probeNames(x)),
                               list(p = NULL))),
    maxIter = 200, epsilon = 1e-06, verbose = TRUE,
    nknots = 10, nOut = 2000, nIn = 4000, knots = NULL,
    checkRegulation = TRUE)

```

## Arguments

<code>x</code>	Data, log2(PM) intensities or an AffyBatch object, see details
<code>design</code>	design matrix
<code>contrast</code>	contrast matrix
<code>probenames</code>	If not null, it is used to group PM probes into probe sets, see details.
<code>maxIter</code>	maximum number of iterations
<code>epsilon</code>	convergence criteria
<code>verbose</code>	print computation info or not
<code>nknots</code>	Number of knots of spline for $\nu$
<code>nOut</code>	Parameter for calculating knots, see <code>getKnots</code>
<code>nIn</code>	Parameter for calculating knots, see <code>getKnots</code>
<code>knots</code>	Knots, if not NULL it overrides <code>nknots</code> , <code>nOut</code> and <code>nIn</code>
<code>checkRegulation</code>	If TRUE, data is checked for a correct specified contrast (see details)

## Details

This function computes the Probe level Locally moderated Weighted median-t statistic (PLW) described in Åstrand (2007b), specially design for Affymetrix type data, or other microarray data with multiple probes.

The data object `x` should be either a matrix of perfect match (PM) intensities, or an object of class `AffyBatch`. When `x` is a matrix of PM intensities, the intensities should be background corrected, normalized, and logged (with base 2). If `x` is an `AffyBatch` object, the default background correction and normalization of RMA is applied to `x`.

When `probenames` is not null, it should be a vector of length equal to the number rows in the matrix `x`, giving the probe-set identity for each PM probe. Use the function `probeNames` in the `affy` package to get `probenames` when `x` is a matrix of log2(PM) intensities.

Inference is done for each PM probe, thus moderated t-statistic, p-value and log2(FC) is calculated for each probe. The median t-statistics for each probe-set is also computed.

Each PM probe  $g$  (row of `x`) is modeled as:

$$y_g | c_g \sim N(\mu_g, c_g \Sigma)$$

$$c_g \sim \text{InvGamma}(m/2, m\nu/2)$$

where  $\nu$  is function of the mean intensity:  $\nu(\bar{\mu}_g)$ ,  $N$  denotes a multivariate normal distribution,  $\Sigma$  is a covariance matrix and  $\text{InvGamma}(a, b)$  is the inverse-gamma distribution with density function

$$f(x) = (b)^a \exp\{-b/x\} x^{-a-1} / \Gamma(a)$$

Given the design matrix  $D$ ,  $\mu_g$  equals  $D\gamma_g$ , and given the contrast matrix  $C$  the hypothesis  $C\gamma_g = 0$  is tested.  $C$  should be a one row matrix of same length as the column vector  $\gamma_g$ .

See examples on how to specify the design and contrast matrices.

A cubic spline is used to parameterize the smooth function  $\nu(x)$

$$\nu(x) = \exp\{H(x)^T \beta\}$$

where  $H : R \rightarrow R^{2p-1}$  is a set B-spline basis functions for a given set of  $p$  interior spline-knots, see chapter 5 of Hastie et al. (2001).

The parameter estimation procedure is based on the assumption that the specified contrast is close to zero for most genes, or at least that the median contrast over all genes is close to zero. A check is run on data to validate this assumptions. If the checking fails, with the error message "warning: most genes appears to be regulated..." and if YOU ARE SURE that the design and contrast is correct, use `checkRegulation=FALSE`.

### Value

<code>Sigma</code>	Estimated covariance matrix for $y = P^T x$
<code>m</code>	Estimated shape parameter for inverse-gamma prior for probe variances
<code>v</code>	Estimated scale parameter curve for inverse-gamma prior for probe variances
<code>converged</code>	T if the EM algorithms converged
<code>iter</code>	Number of iterations
<code>modS2</code>	Moderated estimator of probe-specific variances
<code>histLogS2</code>	Histogram of $\log(s2)$ where $s2$ is the ordinary variance estimator
<code>fittedDensityLogS2</code>	The fitted density for $\log(s2)$
<code>logs2</code>	Variance estimators, logged with base 2.
<code>medianT</code>	Median moderated t-statistic for each probe-set
<code>t</code>	Moderated t-statistic for each PM probe
<code>coefficients</code>	Estimated contrast for each PM probe
<code>p.value</code>	P-value from the moderated t-statistic for each PM probe
<code>dfT</code>	Degrees of freedom of the moderated t-statistic
<code>weights</code>	Weights for estimating the contrast
<code>P</code>	Transformation matrix
<code>beta</code>	Estimated parameter vector $\beta$ of spline for $\nu(x)$
<code>knots</code>	The knots used in spline for $\nu(x)$
<code>x</code>	The input vector covariate vector $x$

### Author(s)

Magnus Åstrand

### References

- Hastie, T., Tibshirani, R., and Friedman, J. (2001). The Elements of Statistical Learning, volume 1. Springer, first edition.
- Kristiansson, E., Sjögren, A., Rudemo, M., Nerman, O. (2005). Weighted Analysis of Paired Microarray Experiments. Statistical Applications in Genetics and Molecular Biology 4(1)
- Åstrand, M. et al. (2007a). Improved covariance matrix estimators for weighted analysis of microarray data. Journal of Computational Biology, Accepted.
- Åstrand, M. et al. (2007b). Empirical Bayes models for multiple-probe type arrays at the probe level. Bioinformatics, Submitted 1 October 2007.

### See Also

`estimateSigmaMVbeta`, `lmw`

**Examples**

```

# -----
# Example analyzing the 6 arrays in the
# AffySpikeU95Subset data set

# Loading the data
data(AffySpikeU95Subset)

# Defining design and contrast matrix
group<-factor(rep(1:2,each=3))
design<-model.matrix(~group-1)
contrast<-matrix(c(1,-1),1,2)

# Analyzing with an AffyBatch object as input
modell<-plw(AffySpikeU95Subset,design=design,contrast=contrast,
           epsilon=0.01)

## Look at fitted vs observed density for log(s2)
varHistPlot(modell)

## Look at fitted curve for scale parameter
scaleParameterPlot(modell)

## Selecting top genes
topRankSummary(modell,nGenes=10)

## Plotting t-statistics and log2FC for top genes
par(mfrow=c(1,2))
plotSummaryT(modell,nGenes=20)
plotSummaryLog2FC(modell,nGenes=20)

###-----
# Analyzing with BG-adjusted and normalized PM data
pm1<-pm(bg.correct.rma(AffySpikeU95Subset, bgtype = 2))
pm2<-matrix(.C("qnorm_c", as.double(as.vector(pm1)),
              as.integer(nrow(pm1)),
              as.integer(ncol(pm1)))[[1]],
            nrow(pm1),ncol(pm1))
data<-log2(pm2)

probenames<-probeNames(AffySpikeU95Subset)
model2<-plw(data,design=design,contrast=contrast,
            probenames=probenames,epsilon=0.01)

###-----
# Modell and model2 should give identical result
# For example identical top ranking:
range(topRankSummary(modell)$t-
      topRankSummary(model2)$t,na.rm=TRUE)

```

---

scaleParameterPlot *Scale parameter plotted against mean intensity*

---



**Description**

Will produce a scatter plot of variance estimators (logged) for each probe (probe set) against the corresponding mean intensity together with the fitted scale-parameter curve and points showing the knots of the used spline.

**Usage**

```
scaleParameterPlot(model, main="Scale parameter curve",
                   col=1, pch='.', lty=1, curveCol=2, knotsPch=19, knotsCol=3)
```

**Arguments**

model	On object obtained from the function plw or lmw.
main	Main title of plot.
col	Color for individual points (mean,logs2).
pch	Plot symbol for individual points (mean,logs2).
lty	Line type for fitted scale parameter curve.
curveCol	Line color for fitted scale parameter curve.
knotsPch	Plot symbol for spline knots.
knotsCol	Plot color for spline knots.

**Author(s)**

Magnus Åstrand

**See Also**

plw, lmw

**Examples**

```
# -----
# Example using the result of the analysis of
# the 6 arrays in the AffySpikeU95Subset data set

# Loading the data
data(AffySpikeU95Subset)

# Defining design and contrast matrix
group<-factor(rep(1:2,each=3))
design<-model.matrix(~group-1)
contrast<-matrix(c(1,-1),1,2)

# Analyzing using plw
modell<-plw(AffySpikeU95Subset,design=design,contrast=contrast,
           epsilon=0.01)

## Look at fitted curve for scale parameter
scaleParameterPlot(modell)
```

---

statByIndex                      *Computes statistics by index or by row*

---

### Description

These function give the same result as `by(x,index, mad)` `by(x,index, mean)` `by(x,index, median)` but are much faster. NOTE: The index vector is assumed to be SORTED and should contain INTEGER values only.

The function `meanSdByRow` computes mean and standard deviation for each row of the matrix `mat`. A list with mean and sd is returned and gives the the same result as:

```
list(mean=apply(mat,1,mean),sd=apply(mat,1,sd))
```

### Usage

```
madByIndex(x, index)
meanByIndex(x, index)
medianByIndex(x, index)
orderStatByIndex(x, index, orderStat)
sdByIndex(x, index)
meanSdByRow(mat)
```

### Arguments

<code>x</code>	Data vector
<code>index</code>	Index vector
<code>orderStat</code>	Which order statistic to compute
<code>mat</code>	Matrix

### Details

See the definition (R-code) of each function for details.

### Value

All but the last function: A vector with the statistic for each level if `index`. `meanSdByRow`: A list with items mean and sd.

### Author(s)

Magnus Åstrand

### See Also

`by`, `apply`

**Examples**

```

## Example 1
## Computing, mad, mean and median by index.
## Compares with the result obtained using by(...)

n<-10000
x<-rnorm(n)
index<-sort(round(runif(n,0.5,10.5)))

mad1<-madByIndex(x,index)
mad2<-by(x,index,mad)

mean1<-meanByIndex(x,index)
mean2<-by(x,index,mean)

median1<-medianByIndex(x,index)
median2<-by(x,index,median)

par(mfrow=c(2,2),mar=c(4,4,1.5,.5),mgp=c(1.5,.25,0))
plot(mad1,mad2,main="Comparing mad",pch=19)
abline(a=0,b=1,col=2)
plot(mean1,mean2,main="Comparing mean",pch=19)
abline(a=0,b=1,col=2)
plot(median1,median2,main="Comparing median",pch=19)
abline(a=0,b=1,col=2)

## Example 2
## Computing, median by index
## Compares with the running time when using by(...)
n<-200000
x<-rnorm(n)
index<-sort(round(runif(n,0.5,10.5)))

system.time(median1<-medianByIndex(x,index))

system.time(median2<-by(x,index,median))

## Example 3
## Computing, mean and sd by row
## Compares with using apply
nrow<-5000
ncol<-20
mat<-matrix(rnorm(ncol*nrow),nrow,ncol)

system.time(res1<-meanSdByRow(mat))
system.time(res2<-list(mean=apply(mat,1,mean),sd=apply(mat,1,sd)))

par(mfrow=c(1,2),mar=c(4,4,1.5,.5),mgp=c(1.5,.25,0))
plot(res1$mean,res2$mean,pch='.')
plot(res1$sd,res2$sd,pch='.')

```

**Description**

Returns (or plots) t-statistic and/or log2FC for each probe and median for each probe set. P

**Usage**

```
plotSummaryLog2FC(model, nGenes=50, genesOfRank=1:nGenes, genes=NULL)
plotSummaryT(model, nGenes=50, genesOfRank=1:nGenes, genes=NULL)
topRankSummary(model, nGenes=50, genesOfRank=1:nGenes, genes=NULL)
```

**Arguments**

model	On object obtained from the function plw.
nGenes	Gives summary for the nGenes top ranking genes
genesOfRank	Gives summary for genes ranked genesOfRank
genes	Gives summary for specific genes.

**Author(s)**

Magnus Åstrand

**See Also**

plw

**Examples**

```
# -----
# Example using the result of the analysis of
# the 6 arrays in the AffySpikeU95Subset data set

# Loading the data
data(AffySpikeU95Subset)

# Defining design and contrast matrix
group<-factor(rep(1:2, each=3))
design<-model.matrix(~group-1)
contrast<-matrix(c(1, -1), 1, 2)

# Analyzing using plw
modell<-plw(AffySpikeU95Subset, design=design, contrast=contrast,
           epsilon=0.01)

## Selecting top genes
topRankSummary(modell, nGenes=10)

## Plotting t-statistics and log2FC for top genes
par(mfrow=c(1, 2))
plotSummaryT(modell, nGenes=20)
plotSummaryLog2FC(modell, nGenes=20)
```

---

varHistPlot	<i>Variance histogram and density</i>
-------------	---------------------------------------

---

**Description**

Will produce a histogram of observed variance estimators (logged) together with the fitted density.

**Usage**

```
varHistPlot(model, main="Histogram variance estimators",
             histCol=8, densityCol=2, drawLegend=TRUE)
```

**Arguments**

model	On object obtained from the function plw or lmw.
main	Main title of plot.
histCol	Color for histogram bars.
densityCol	Color for density function.
drawLegend	To draw a legend or not.

**Author(s)**

Magnus Åstrand

**See Also**

plw, lmw

**Examples**

```
# -----
# Example using the result of the analysis of
# the 6 arrays in the AffySpikeU95Subset data set

# Loading the data
data(AffySpikeU95Subset)

# Defining design and contrast matrix
group<-factor(rep(1:2, each=3))
design<-model.matrix(~group-1)
contrast<-matrix(c(1, -1), 1, 2)

# Analyzing using plw
modell<-plw(AffySpikeU95Subset, design=design, contrast=contrast,
           epsilon=0.01)

## Look at fitted vs observed density for log(s2)
varHistPlot(modell)
```

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