

# Package ‘stepNorm’

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**Author** Yuanyuan Xiao <yxiao@itsa.ucsf.edu>, Yee Hwa (Jean) Yang <jean@biostat.ucsf.edu>

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**Maintainer** Yuanyuan Xiao <yxiao@itsa.ucsf.edu>

**Description** Stepwise normalization functions for cDNA microarray data.

**License** LGPL

**URL** <http://www.biostat.ucsf.edu/jean/>

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 calcAIC
 

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*Extract AIC from a Fitted Model*


---

### Description

Computes the Akaike Information Criterion for a fitted parametric model.

### Usage

```
calcAIC(fit, subset=TRUE, scale = 0, enp, loss.fun = square)
```

### Arguments

fit	fitted model; see details below
scale	optional numeric specifying the scale parameter of the model; see <a href="#">scale</a> in <a href="#">step</a> .
subset	A "logical" or "numeric" vector indicating the subset of points used to compute the fitted model.
enp	equivalent number of parameters in the fitted model. If missing, the enp component from fit will be used.
loss.fun	the loss function used to calculate deviance; default uses the squared deviations from the fitted values; one could also use, for example, absolute deviations ( <a href="#">abs</a> ).

### Details

The argument `fit` can be an object of class `marrayFit`, in which case the residuals component from the `marrayFit` object will be extracted to calculate the deviance; the user can also pass in a numeric vector, in which case it will be interpreted as the residuals and the user needs to specify the argument `enp`.

The criterion used is

$$AIC = -2 * \log L + k * enp,$$

where  $L$  is the likelihood and `enp` the equivalent number of parameters of `fit`. For linear models (as in `marrayFit`),  $-2\log L$  is computed from the deviance.

$k = 2$  corresponds to the traditional AIC and is the penalty for the number of parameters.

### Value

A numeric vector of length 4, giving

Dev	the deviance of the <code>fit</code> .
enp	the equivalent number of parameters of the <code>fit</code> .
penalty	the penalty for number of parameters.
Criterion	the Akaike Information Criterion for <code>fit</code> .

**Author(s)**

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

**See Also**

[AIC](#), [deviance](#), [calcBIC](#).

**Examples**

```
## load in swirl data
data(swirl)

## fit a model
fit <- fitWithin(fun="medfit")
## res is an object of class marrayFit
res <- fit(swirl[,1])

## calculate AIC
calcAIC(res)
## or could pass in the residual vector, but then argument "enp" needs to be specified
calcAIC(res$residual, enp=1)
```

---

calcBIC

*Extract BIC from a Fitted Model*

---

**Description**

Computes the Bayesian Information Criterion for a fitted parametric model.

**Usage**

```
calcBIC(fit, subset=TRUE, scale = 0, enp, loss.fun = square)
```

**Arguments**

fit	fitted model; see details below
subset	A "logical" or "numeric" vector indicating the subset of points used to compute the fitted model.
scale	optional numeric specifying the scale parameter of the model; see <a href="#">scale</a> in <a href="#">step</a> .
enp	equivalent number of parameters in the fitted model. If missing, the enp component from fit will be used.
loss.fun	the loss function used to calculate deviance; the default uses the squared deviation from the fitted values; one could also use absolute deviations ( <a href="#">abs</a> ).

**Details**

The argument `fit` can be an object of class `marrayFit`, in which case the residuals component from the `marrayFit` object will be extracted to calculate the deviance; the user can also pass in a numeric vector, in which case it will be interpreted as the residuals and the user needs to specify the argument `enp`.

The criterion used is

$$BIC = -2 * \log L + k * enp,$$

where  $L$  is the likelihood and `enp` the equivalent number of parameters of `fit`. For linear models (as in `marrayFit`),  $-2\log L$  is computed from the deviance.

$k = \log(n)$  corresponds to the BIC and is the penalty for the number of parameters.

**Value**

A numeric vector of length 4, giving

Dev	the deviance of the fit.
enp	the equivalent number of parameters of the fit.
penalty	the penalty for number of parameters.
Criterion	the Akaike Information Criterion for fit.

**Author(s)**

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

**See Also**

[AIC](#), [deviance](#), [calcAIC](#).

**Examples**

```
## load in swirl data
data(swirl)

## fit a model
fit <- fitWithin(fun="medfit")
## res is an object of class marrayFit
res <- fit(swirl[,1])

## calculate BIC
calcBIC(res)
## or could pass in the residual vector, but then argument "enp" needs to be specified
calcBIC(res$residual, enp=1)
```

---

fit2DWithin

*Bivariate location normalization function for cDNA microarray data*


---

### Description

This function performs 2D location normalization on cDNA microarray. It operates on class `marrayRaw` or class `marrayNorm`. It allows the user to choose from a set of four basic normalization procedures.

### Usage

```
fit2DWithin(x1.fun = "maSpotRow", x2.fun = "maSpotCol", y.fun = "maM",
            subset=TRUE, fun = aov2Dfit, ...)
```

### Arguments

<code>x1.fun</code>	Name of accessor method for spot row coordinates, usually <code>maSpotRow</code> .
<code>x2.fun</code>	Name of accessor method for spot column coordinates, usually <code>maSpotCol</code> .
<code>y.fun</code>	Name of accessor method for spot statistics, usually the log-ratio <code>maM</code> .
<code>subset</code>	A "logical" or "numeric" vector indicating the subset of points used to compute the normalization values.
<code>fun</code>	Character string specifying the normalization procedures: <b>rlm2Dfit</b> for robust linear regression using the <code>rlm</code> function <b>loess2Dfit</b> for robust local regression using the <code>loess</code> function <b>aov2Dfit</b> for linear regression using the <code>lm</code> function <b>spatialMedfit</b> for spatial median normalization
<code>...</code>	Misc arguments for <code>fun</code>

### Details

The spot statistic named in `y` is regressed on spot row and column coordinates, using the function specified by the argument `fun`. Typically, `rlm2Dfit` and `loess2Dfit`, which treat row and column coordinates as numeric vectors, require a lot fewer parameters than `aov2Dfit` which specifies these two variables as categorical. `spatialMedfit` could yet fit the most complicated model, depending on size of the smoothing window specified; details see [Wilson et al \(2003\)](#).

### Value

The function `fit2DWithin` returns a function ( $F$ ) with bindings for `x1.fun`, `x2.fun`, `y.fun`, `subset` and `fun`. When the function  $F$  is evaluated with an object of class `marrayNorm` or `marrayRaw`, it carries out normalization and returns an object of class `marrayFit` that contains the normalization information as a list with the following components:

<code>varfun</code>	: A character vector of names of predictor variables.
<code>x</code>	: A numeric matrix of predictor variables.

y : A numeric matrix of responses.  
 residuals : A numeric matrix of normalized values (typically log ratios ( $M$ )).  
 fitted : A numeric matrix of the fitted values.  
 enp : The equivalent number of parameters; see [loess](#).  
 df.residual : The residual degrees of freedom.  
 fun : A character string indicating the name of the function used for normalization.

Note that the residuals component stores the normalized ratios.

### Author(s)

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>,  
 Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

### References

Y. H. Yang, S. Dudoit, P. Luu, and T. P. Speed (2001). Normalization for cDNA microarray data. In M. L. Bittner, Y. Chen, A. N. Dorsel, and E. R. Dougherty (eds), *Microarrays: Optical Technologies and Informatics*, Vol. 4266 of *Proceedings of SPIE*.

D. L. Wilson, M. J. Buckley, C. A. Helliwell and I. W. Wilson (2003). New normalization methods for cDNA microarray data. *Bioinformatics*, Vol. 19, pp. 1325-1332.

### See Also

[fitWithin](#)

### Examples

```

## use the swirl data as example
data(swirl)

## 2D rlm normalization
rlm2D <- fit2DWithin(fun="rlm2Dfit")
swirl1.rlm <- rlm2D(swirl[,1])
norm.M <- swirl1.rlm$residuals ## matrix of normalized ratios

## 2D loess normalization, default span=0.2
loess2D <- fit2DWithin(fun="loess2Dfit")
swirl1.loess <- loess2D(swirl[,1])
## 2D loess normalization, span=0.4
## Not run:
loess2D.1 <- fit2DWithin(fun="loess2Dfit", span=0.4)
swirl1.loess.1 <- loess2D.1(swirl[,1])
## End(Not run)

## 2D aov normalization
aov2D <- fit2DWithin(fun="aov2Dfit")
swirl1.aov <- aov2D(swirl[,1])

## 2D spatial median normalization, default window width=3

```

```

spatialMed2D <- fit2DWithin(fun="spatialMedfit")
swirl1.spatialMed <- spatialMed2D(swirl[,1])
## 2D loess normalization, window width=9
## Not run:
spatialMed2D.1 <- fit2DWithin(fun="spatialMedfit", width=9)
swirl1.spatialMed.1 <- spatialMed2D.1(swirl[,1])
## End(Not run)

```

---

fitWithin

*Simple location normalization function for cDNA microarray data*


---

### Description

This function performs location normalization on cDNA microarray. It operates on class `marrayRaw` or class `marrayNorm`. It allows the user to choose from a set of three basic normalization procedures.

### Usage

```
fitWithin(x.fun = "maA", y.fun = "maM", z.fun = TRUE, subset=TRUE, fun = "medfit", ...)
```

### Arguments

<code>x.fun</code>	Name of accessor method for spot intensity, usually <code>maA</code> .
<code>y.fun</code>	Name of accessor method for spot statistics, usually the log-ratio <code>maM</code> .
<code>z.fun</code>	Name of accessor method for spot statistic used to stratify the data, usually a layout parameter, e.g. <code>maPrintTip</code> or <code>maCompPlate</code> . If <code>z</code> is not a character, e.g. <code>NULL</code> , the data are not stratified.
<code>subset</code>	A "logical" or "numeric" vector indicating the subset of points used to compute the normalization values.
<code>fun</code>	Character string specifying the normalization procedure: <b>medfit</b> for global median location normalization <b>rlmfit</b> for global intensity or A-dependent location normalization using the <code>rlm</code> function <b>loessfit</b> for global intensity or A-dependent location normalization using the <code>loess</code> function
<code>...</code>	Miscs arguments to be passed in <code>fun</code>

### Details

Normalization is typically performed on the expression ratios of cDNA microarray data, using the function specified by argument `fun`. Currently, this function is to be chosen from: `medfit` (median), `rlmfit` (`rlm`) and `loessfit` (`loess`). When `z.fun` is provided as a character string, for example, `maPrintTip`, the normalization procedure is operated within each print-tip of the slide.

**Value**

The function `fitWithin` returns a function( $F$ ) with bindings for `x.fun`, `y.fun`, `z.fun`, `subset` and `fun`. When the function  $F$  is evaluated with an object of class `marrayNorm` or `marrayRaw`, it carries out normalization and returns an object of class `marrayFit` that contains the normalization information as a list with the following list components:

`varfun` : A character vector of names of predictor variables.  
`x` : A numeric matrix of predictor variables.  
`y` : A numeric matrix of responses.  
`residuals` : A numeric matrix of normalized values (typically log ratios ( $M$ )).  
`fitted` : A numeric matrix of the fitted values.  
`enp` : The equivalent number of parameters; see [loess](#).  
`df.residual` : The residual degrees of freedom.  
`fun` : A character string indicating the name of the function used for normalization.

Note that the `residuals` component stores the normalized ratios.

**Author(s)**

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
 Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

**References**

Y. H. Yang, S. Dudoit, P. Luu, and T. P. Speed (2001). Normalization for cDNA microarray data. In M. L. Bittner, Y. Chen, A. N. Dorsel, and E. R. Dougherty (eds), *Microarrays: Optical Technologies and Informatics*, Vol. 4266 of *Proceedings of SPIE*.

**See Also**

[fit2DWithin](#)

**Examples**

```
## using the swirl data as example
data(swirl)

## median normalization
med <- fitWithin(fun="medfit")
swirl1.med <- med(swirl[,1])
norm.M <- swirl1.med$residuals ## matrix of normalized ratios

## rlm normalization
rlmF <- fitWithin(fun="rlmfit")
swirl1.rlm <- rlmF(swirl[,1])

## loess normalization, default span=0.4
loessF <- fitWithin(fun="loessfit")
swirl1.loess <- loessF(swirl[,1])
```



```
## loess normalization, span=0.2
loessF.1 <- fitWithin(fun="loessfit", span=0.2)
swirl1.loess.1 <- loessF.1(swirl[,1])

## within-printtip loess normalization
loessP <- fitWithin(z.fun="maPrintTip", fun="loessfit")
swirl1.loessP <- loessP(swirl[,1])
```

---

`maCompPlate2`*Generate plate IDs*

---

## Description

This function is a modification of the `maCompPlate` function in the `marray` library. It generates plate IDs from the dimensions of the grid and spot matrices. Unlike the `maCompPlate` function, the number of spots is not necessarily a multiple of the number of wells on a plate, therefore this function allows empty spots on the slide.

## Usage

```
maCompPlate2(no.plates = NULL, n = 384)
```

## Arguments

<code>no.plates</code>	object of class "numeric", number of plates used specified by the user. If a number is not specified, then it is assumed that there are no empty spots on the slide.
<code>n</code>	object of class "numeric", number of wells in each plate, usually 384 or 96.

## Details

This function can be used to handle three cases: 1) the number of spots is a multiple of the number of wells on a plate (usually 96 or 384); 2) the number of spots is not a multiple of the number of wells on a plate, and several of spots on the slide are therefore left empty. In this case, the user needs to specify the number of total plates used; plate IDs of empty spots will be NAs; 3) the number of spots is not a multiple of the number of wells on a plate, but all spots on the slide are spotted, therefore there is one plate not fully used. In this case, the user does not need to specify the number of total plates (as this will not be an integer), the function assumes no empty spots on the slide automatically. See Examples below.

## Value

The function `maCompPlate2` returns a function with bindings for `no.plates` and `n`, which when receiving a object of `marrayRaw`, `marrayNorm` or `marrayLayout` class, it returns a vector of plate IDs (`factor`).

**Author(s)**

Yuanyuan Xiao

**See Also**

[maCompPlate](#), [marrayLayout](#).

**Examples**

```
##### case 1: no empty spots on the slide, full plates used
L<-new("marrayLayout", maNgr=4, maNgc=4, maNsr=22, maNsc=24)
### "compPlate" is a function
compPlate <- maCompPlate2(n=384)
plate <- compPlate(L)
table(plate)
### can also use:
plate<-maCompPlate(L,384)
table(plate)

##### case 2: with empty spots on the slide, full plates used
L<-new("marrayLayout", maNgr=4, maNgc=4, maNsr=22, maNsc=26)
### "compPlate" is a function
compPlate <- maCompPlate2(no.plates=22,n=384)
plate <- compPlate(L)
table(plate)
### empty spots are NAs
unique(plate)

##### case 3: no empty spots on the slide, one plate not full
L<-new("marrayLayout", maNgr=4, maNgc=4, maNsr=22, maNsc=26)
### argument no.plates not specified, the function assumes no empty spots
compPlate <- maCompPlate2(n=384)
plate <- compPlate(L)
### 23 full plates (384), the 24th not full (304)
table(plate)
### no NAs, no empty spots
unique(plate)
```

---

makeStepList

*Construction of a stepwise normalization list*

---

**Description**

This function provides a user friendly way to construct a list for input to the function [stepWithinNorm](#). The list indicates intended biases for correction and models for stepwise normalization.

**Usage**

```
makeStepList(A = c("median", "rlm", "loess"), PT = c("median", "rlm",
"loess"), PL = c("median", "rlm", "loess"), Spatial2D = c("rlm2D",
"loess2D", "aov2D", "spatialMedian"))
```

**Arguments**

- A** A character string specifying the normalization models for the adjustment of intensity or *A* bias:
- median:** global median location normalization
  - rlm:** global intensity or *A*-dependent robust linear normalization using the `rlm` function
  - loess:** global intensity or *A*-dependent robust nonlinear normalization using the `loess` function
- The user can specify any of these three choices and the selected model will be compared based the goodness fit and model parsimony; If the correction of the  $\$A\$$  bias is not desired, the user can set `A = NULL`.
- PT** A character string specifying the normalization models for the adjustment of print-tip or *PT* bias:
- median:** within-print-tip-group median normalization
  - rlm:** within-print-tip-group robust linear normalization using the `rlm` function
  - loess:** within-print-tip-group robust nonlinear normalization using the `loess` function
  - none:** no normalization for the *PT* bias
- If the correction of the  $\$PT\$$  bias is not desired, the user can set `PT = NULL`.
- PL** A character string specifying the normalization models for the adjustment of well-plate or *PL* bias:
- median:** within-well-plate median normalization
  - rlm:** within-well-plate robust linear normalization using the `rlm` function
  - loess:** within-well-plate robust nonlinear normalization using the `loess` function
  - none:** no normalization for the *PL* bias
- If the correction of the  $\$PL\$$  bias is not desired, the user can set `PL = NULL`.
- Spatial2D** A character string specifying the normalization models for the adjustment of spatial 2D bias:
- none:** no normalization for the spatial 2D bias
  - aov2D:** spatial bivariate location normalization using ANOVA
  - rlm2D:** spatial bivariate location normalization using the `rlm` function
  - loess2D:** spatial bivariate location normalization using the `loess` function
  - spatialMedian:** spatial location normalization using a spatial median approach (see Wilson et al. (2003) in reference)
- If the correction of the  $\$SPL\$$  bias is not desired, the user can set `Spatial2D = NULL`.

**Details**

This function provides a user friendly way to specify the parameter `wf.loc` for the main stepwise normalization function `stepWithinNorm`; see examples for details.

**Value**

An object of class "list" for input to the `stepWithinNorm` function.

**Author(s)**

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

**See Also**

[stepWithinNorm](#).

**Examples**

```
# Examples use swirl dataset, for description type ? swirl
data(swirl)

# To use the default parameters, which adjusts A, PT, PL and Spatial2D
# biases sequentially and compares all models available, simple type

wf.loc <- makeStepList()

# To apply loess for the A bias, and to omit the Spatial2D step
wf.loc <- makeStepList(A="loess", Spatial2D=NULL)

# To compare only rlm and loess in the A bias step, and other biases as default
wf.loc <- makeStepList(A=c("rlm","loess"))

# input to the stepWithinNorm function
## Not run:
step.swirl1 <- stepWithinNorm(swirl[,1],wf.loc=wf.loc)
## End(Not run)
```

---

marrayFit-class

*Class "marrayFit", storing parameters and results of post-normalization cDNA microarray data*

---

**Description**

A simple list-based class for the storage of parameters and results of normalization of cDNA microarray data.

## Creating Objects from the Class

Objects can be created by calls of the form `new('marrayFit', fit)` where `fit` is a list. Objects of `marrayFit` in the `StepNorm` package are typically created by functions `fitWithin` and `fit2DWithin`.

## List Components

This class contains no slots, but objects should contain the following list components:

**varfun** : A character vector of names of predictor variables.

**x** : A numeric matrix of predictor variables.

**y** : A numeric matrix of responses.

**residuals** : A numeric matrix of normalized values (typically log ratios ( $M$ )).

**fitted** : A numeric matrix of the fitted values.

**enp** : The equivalent number of parameters; see `loess`.

**df.residual** : The residual degrees of freedom.

**fun** : A character string indicating the name of the function used for normalization.

## Methods

This class inherits directly from class `list` so any operation appropriate for lists will work on objects of this class.

## Author(s)

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

## See Also

`fitWithin`, `fit2DWithin`.

## Examples

```
## load in swirl data
data(swirl)

## median normalization for the first slide of the swirl data
medWithin <- fitWithin(fun="medfit")
## medFit is an object of class marrayFit
medFit <- medWithin(swirl[,1])
## normalized ratios is stored in:
norm.M <- medFit$residuals
```

---

seqWithinNorm	<i>Sequential within-slide normalization function</i>
---------------	---

---

## Description

This function conducts cDNA microarray normalization in a sequential fashion. In a two-color cDNA array setting, within-slide normalization calibrates signals from the two channels to remove non-biological variation introduced by various processing steps.

## Usage

```
seqWithinNorm(marraySet, y = "maM", subset = TRUE, loss.fun = square,
A = c("loess", "rlm", "median", "none"),
PT = c("median", "rlm", "loess", "none"),
PL = c("median", "rlm", "loess", "none"),
Spatial2D = c("none", "aov2D", "rlm2D", "loess2D", "spatialMedian"),
criterion = c("BIC", "AIC"))
```

## Arguments

marraySet	Object of class <code>marrayRaw</code> or class <code>marrayNorm</code> , containing intensity data for the batch of arrays to be normalized.
y	Name of accessor method for spot statistics, usually the log-ratio <code>maM</code> .
subset	A "logical" or "numeric" vector indicating the subset of points used to compute the normalization values.
loss.fun	The loss function used in calculating deviance, the default uses squared sum of residuals; for absolute sum of residuals, use <code>abs</code>
A	A character string specifying the normalization model for the adjustment of intensity or <i>A</i> bias: <b>loess:</b> global intensity or <i>A</i> -dependent robust nonlinear normalization using the <code>loess</code> function <b>rlm:</b> global intensity or <i>A</i> -dependent robust linear normalization using the <code>rlm</code> function <b>median:</b> global median location normalization <b>none:</b> no normalization for the <i>A</i> bias If not specified, loess normalization will be applied.
PT	A character string specifying the normalization model for the adjustment of print-tip or <i>PT</i> bias: <b>median:</b> within-print-tip-group median normalization <b>rlm:</b> within-print-tip-group robust linear normalization using the <code>rlm</code> function <b>loess:</b> within-print-tip-group robust nonlinear normalization using the <code>loess</code> function <b>none:</b> no normalization for the <i>PT</i> bias

	If not specified, median normalization within print-tip will be applied.
PL	<p>A character string specifying the normalization model for the adjustment of well-plate or <i>PL</i> bias:</p> <p><b>median:</b> within-well-plate median normalization</p> <p><b>rlm:</b> within-well-plate robust linear normalization using the <code>rlm</code> function</p> <p><b>loess:</b> within-well-plate robust nonlinear normalization using the <code>loess</code> function</p> <p><b>none:</b> no normalization for the <i>PL</i> bias</p> <p>If not specified, median normalization within well-plate will be applied.</p>
Spatial2D	<p>A character string specifying the normalization model for the adjustment of spatial 2D bias:</p> <p><b>none:</b> no normalization for the spatial 2D bias</p> <p><b>aov2D:</b> spatial bivariate location normalization using ANOVA</p> <p><b>rlm2D:</b> spatial bivariate location normalization using the <code>rlm</code> function</p> <p><b>loess2D:</b> spatial bivariate location normalization using the <code>loess</code> function</p> <p><b>spatialMedian:</b> spatial location normalization using a spatial median approach (see Wilson et al. (2003) in reference)</p> <p>If not specified, no normalization will be carried out in this step.</p>
criterion	<p>Character string specifying the criterion:</p> <p><b>AIC:</b> the AIC criterion is used; see <code>calcAIC</code>.</p> <p><b>BIC:</b> the BIC criterion is used; see <code>calcBIC</code>.</p> <p>If no specification, BIC is used. Note that here we don't use the criterion to choose normalization model in each step. Criterion is calculated solely for information purpose.</p>

## Details

Typical systematic non-biological variations of a two-color cDNA microarray include the dependence of ratio measurements (*M*) on intensity (*A*), print-tip IDs (*PT*), plate IDs (*PL*) and spatial heterogeneity of the slide (Spatial 2D). The sequential normalization procedure in `seqWithinNorm` normalizes a slide in a sequential fashion: *A* -> *PT* -> *PL* -> Spatial2D. In each step one kind of variation is targeted for correction, and the user chooses the normalization method as desired. We calculate the AIC/BIC criterion along the normalization steps, but they are not used for selection of models.

## Value

An object of class "list":

<code>normdata</code>	an object of class <code>marrayNorm</code> , containing the normalized intensity data.
<code>res</code>	a list of the sequential normalization result for each slide within the <code>marray</code> dataset. Each list component is also a list containing the name of the biases, deviance, equivalent number of parameters, AIC/BIC value for a certain slide.

**Author(s)**

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
 Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

**References**

Y. H. Yang, S. Dudoit, P. Luu, and T. P. Speed (2001). Normalization for cDNA microarray data. In M. L. Bittner, Y. Chen, A. N. Dorsel, and E. R. Dougherty (eds), *Microarrays: Optical Technologies and Informatics*, Vol. 4266 of *Proceedings of SPIE*.

D. L. Wilson, M. J. Buckley, C. A. Helliwell and I. W. Wilson (2003). New normalization methods for cDNA microarray data. *Bioinformatics*, Vol. 19, pp. 1325-1332.

**See Also**

[stepWithinNorm](#), [withinNorm](#), [fitWithin](#), [fit2DWithin](#), [calcAIC](#), [calcBIC](#).

**Examples**

```
# Examples use swirl dataset, for description type ? swirl
data(swirl)

# Apply sequential normalization for the first slide
# default: loess(A)->median(PT)->median(PL)-> none (Spatial2D)
## Not run:
res.swirl1 <- seqWithinNorm(swirl[,1])

# normalized data
norm.swirl1 <- res.swirl1[[1]]

# sequential normalization information
step.swirl1 <- res.swirl1[[2]]

## End(Not run)
# median(A)->median(PT)->median(PL)->none(Spatial2D)
res.swirl <- seqWithinNorm(swirl[,1], A="median",PT="median",PL="median",Spatial2D="none")
```

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stepWithinNorm

*Stepwise within-slide normalization function*

---

**Description**

This function conducts cDNA microarray normalization in a stepwise fashion. In a two-color cDNA array setting, within-slide normalization calibrates signals from the two channels to remove non-biological variation introduced by various processing steps.



**Usage**

```
stepWithinNorm(marraySet, subset=TRUE, wf.loc, criterion = c("BIC", "AIC"), loss.fun = square)
```

**Arguments**

marraySet	Object of class <code>marrayRaw</code> or class <code>marrayNorm</code> , containing intensity data for the batch of arrays to be normalized.
subset	A "logical" or "numeric" vector indicating the subset of points used to compute the normalization values.
wf.loc	Object of class <code>list</code> , each component is a step for the removal of a particular systematic variation. Typically each step is also a list of several candidate models of different complexity, the best model will be chosen by the criterion specified. For a user friendly way of constructing such a list, consult the function <code>makeStepList</code> . If missing, the default procedure will be used, which we consider appropriate for most slides. See details for how to specify the list and how it is used.
criterion	Character string specifying the criterion used for the selection of the best normalization procedure in each step. This argument can be specified using the first letter of each method; if no specification is made, the default is BIC: <b>AIC:</b> the AIC criterion is used <b>BIC:</b> the BIC criterion is used.
loss.fun	loss function; default set at using residual sum of squares.

**Details**

Typical systematic non-biological variations of a two-color cDNA microarray include the dependence of ratio measurements (M) on intensity (A), print-tip IDs (PT), plate IDs (PL) and spatial heterogeneity of the slide (SP). The stepwise normalization procedure normalizes a slide in a stepwise fashion. In each step one kind of variation is targeted for correction. Within each step, various candidate models are assessed for their adequacy with respect to the observed data. The assessment is made based on a common model selection criterion, AIC (see `calcAIC`) or BIC (see `calcBIC`), and the best model is then chosen for the specified step.

The argument `wf.loc` is a list of steps. Each step is also a list of models. The user uses the function `fitWithin` or `fit2DWithin` to specify a model. Below is a table of how to do so:

systematic variation	model	function
intensity (A)	median	<code>fitWithin(fun="medfit")</code>
A	robust linear	<code>fitWithin(fun="rlmfit")</code>
A	robust nonlinear	<code>fitWithin(fun="loessfit")</code>
print-tip (PT)	median	<code>fitWithin(z.fun="maPrintTip", fun="medfit")</code>
PT	robust linear	<code>fitWithin(z.fun="maPrintTip", fun="rlmfit")</code>
PT	robust nonlinear	<code>fitWithin(z.fun="maPrintTip", fun="loessfit")</code>
plate (PL)	median	<code>fitWithin(z.fun="maCompPlate", fun="medfit")</code>
PL	robust linear	<code>fitWithin(z.fun="maCompPlate", fun="rlmfit")</code>
PL	robust nonlinear	<code>fitWithin(z.fun="maCompPlate", fun="loessfit")</code>
spatial (SP)	robust linear	<code>fit2DWithin(fun="rlm2Dfit")</code>
SP	robust nonlinear( <code>span=0.2</code> )	<code>fit2DWithin(fun="loess2Dfit", span=0.2)</code>

SP	anova	fit2DWithin(fun="aov2Dfit")
SP	spatial median (11X11)	fit2DWithin(fun="spatialMedfit", width=11)

If the `wf.loc` is not specified by the user, the default procedure conducts normalization in four steps: A -> PT -> PL -> SP and models are as described in the table above. The user can choose not to follow such a procedure by passing in a different list, however we advocate normalizing the intensity (A) variation first as it is usually the source of most variation in most slides. The list can be easier specified using the function `makeStepList` by inputting models as character strings, see `makeStepList` for details.

### Value

An object of class "list":

<code>normdata</code>	an object of class <code>marrayNorm</code> , containing the normalized intensity data.
<code>res</code>	a dataframe of the stepwise normalization result, containing the name of the model chosen for each step, deviance, equivalent number of parameters, AIC/BIC value.

### Author(s)

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

### References

- Y. H. Yang, S. Dudoit, P. Luu, and T. P. Speed (2001). Normalization for cDNA microarray data. In M. L. Bittner, Y. Chen, A. N. Dorsel, and E. R. Dougherty (eds), *Microarrays: Optical Technologies and Informatics*, Vol. 4266 of *Proceedings of SPIE*.
- D. L. Wilson, M. J. Buckley, C. A. Helliwell and I. W. Wilson (2003). New normalization methods for cDNA microarray data. *Bioinformatics*, Vol. 19, pp. 1325-1332.

### See Also

`seqWithinNorm`, `withinNorm`, `fitWithin`, `fit2DWithin`, `calcAIC`, `calcBIC`.

### Examples

```
# Examples use swirl dataset, for description type ? swirl
data(swirl)

# Apply stepwise normalization for the first slide
res.swirl1 <- stepWithinNorm(swirl[,1])

# normalized data
norm.swirl1 <- res.swirl1[[1]]

# stepwise procedure
step.swirl1 <- res.swirl1[[2]]
```

```

# using a stepwise procedure different than the default
# corrects intensity (A) and print-tip (PT), this can be
# carried out in two ways:
# 1)
steps <- list(
  wholeChipA = list(med = fitWithin(fun="medfit"),
                    rlm = fitWithin(fun="rlmfit"),
                    loess = fitWithin(fun="loessfit")),
  printTipA = list(med = fitWithin(z.fun="maPrintTip", fun="medfit"),
                   rlm = fitWithin(z.fun="maPrintTip", fun="rlmfit"),
                   loess = fitWithin(z.fun="maPrintTip", fun="loessfit")))

#2)
steps <- makeStepList(PL=NULL, Spatial2D=NULL)
## Not run:
res.swirl <- stepWithinNorm(swirl[,1], wf.loc=steps)
## End(Not run)

# using AIC criterion for the first slide
## Not run:
res.swirl <- stepWithinNorm(swirl[,1], criterion="A")
## End(Not run)

```

---

withinNorm

*Within-slide normalization function for cDNA spotted microarrays*


---

## Description

This function is a wrapper function around `fitWtihin` and `fit2DWithin`. It allows the user to choose from a set of thirteen basic location normalization procedures. The function operates on an object of class `marrayRaw` or `marrayNorm` and returns an object of class `marrayNorm`.

## Usage

```

withinNorm(marraySet, y = "maM", subset = TRUE, norm = c("none",
  "median", "rlm", "loess", "medianPrintTip", "rlmPrintTip",
  "loessPrintTip", "medianPlate", "rlmPlate", "loessPlate",
  "aov2D", "rlm2D", "loess2D", "spatialMedian"), ...)

```

## Arguments

<code>marraySet</code>	Object of class <code>marrayRaw</code> or class <code>marrayNorm</code> , containing intensity data for the batch of arrays to be normalized.
<code>y</code>	Name of accessor method for spot statistics, usually the log-ratio <code>maM</code> .
<code>subset</code>	A "logical" or "numeric" vector indicating the subset of points used to compute the normalization values.

norm            A character string specifying the normalization procedures:

- none:** no normalization
- median:** global median location normalization
- rlm:** global intensity or A-dependent robust linear normalization using the `rlm` function
- loess:** global intensity or A-dependent robust nonlinear normalization using the `loess` function
- medianPrintTip:** within-print-tip-group median normalization
- rlmPrintTip:** within-print-tip-group intensity or A-dependent robust linear normalization using the `rlm` function
- loessPrintTip:** within-print-tip-group intensity or A-dependent robust nonlinear normalization using the `loess` function
- medianPlate:** within-well-plate-group median normalization
- rlmPlate:** within-well-plate-group intensity or A-dependent robust linear normalization using the `rlm` function
- loessPlate:** within-well-plate-group intensity or A-dependent robust nonlinear normalization using the `loess` function
- aov2D:** spatial bivariate location normalization using ANOVA
- rlm2D:** spatial bivariate location normalization using the `rlm` function
- loess2D:** spatial bivariate location normalization using the `loess` function
- spatialMedian:** spatial location normalization using a spatial median approach (see Wilson et al. (2003) in reference)

...            Misc arguments for the specified norm function

### Details

The function `withinNorm` dispatches to the function `fitWithin` or `fit2DWithin` with specified arguments according to the choice of `norm`. For instance, when `norm="loess"` for global intensity dependent robust nonlinear normalization, `withinNorm` calls `fitWithin(fun="loess")` with the default `span` parameter set at 0.4. If a different `span` is preferred, it should be input by `span=0.2` through the argument `...` in the `withinNorm` function (see example below). For more details see `fitWithin`, `fit2DWithin` and individual fitting functions such as `loessfit`.

### Value

An object of class `marrayNorm`, containing the normalized intensity data.

### Author(s)

Yuanyuan Xiao, <yxiao@itsa.ucsf.edu>  
Jean Yee Hwa Yang, <jean@biostat.ucsf.edu>

### References

Y. H. Yang, S. Dudoit, P. Luu, and T. P. Speed (2001). Normalization for cDNA microarray data. In M. L. Bittner, Y. Chen, A. N. Dorsel, and E. R. Dougherty (eds), *Microarrays: Optical Technologies and Informatics*, Vol. 4266 of *Proceedings of SPIE*.

D. L. Wilson, M. J. Buckley, C. A. Helliwell and I. W. Wilson (2003). New normalization methods for cDNA microarray data. *Bioinformatics*, Vol. 19, pp. 1325-1332.

### See Also

[seqWithinNorm](#), [stepWithinNorm](#), [fitWithin](#), [fit2DWithin](#), [loessfit](#), [rlmfit](#).

### Examples

```
# Examples use swirl dataset, for description type ? swirl
data(swirl)

# Apply loess normalization for the first slide, span=0.4
## Not run:
res.swirl1 <- withinNorm(swirl[,1], norm="loess")
## End(Not run)

# Apply loess normalization for the first slide, span=0.2
## Not run:
res.swirl1 <- withinNorm(swirl[,1], norm="loess", span=0.2)
## End(Not run)
```

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