

# Package ‘PERFect’

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**Type** Package

**Title** Permutation filtration for microbiome data

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**Description** PERFect is a novel permutation filtering approach designed to address two unsolved problems in microbiome data processing: (i) define and quantify loss due to filtering by implementing thresholds, and (ii) introduce and evaluate a permutation test for filtering loss to provide a measure of excessive filtering.

**Depends** R (>= 3.6.0), sn (>= 1.5.2)

**Imports** ggplot2 (>= 3.0.0), phyloseq (>= 1.28.0), zoo (>= 1.8.3),  
psych (>= 1.8.4), stats (>= 3.6.0), Matrix (>= 1.2.14),  
fitdistrplus (>= 1.0.12), parallel (>= 3.6.0)

**License** Artistic-2.0

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**LazyData** false

**BugReports** <https://github.com/cxquy91/PERFect/issues>

**URL** <https://github.com/cxquy91/PERFect>

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DiffFiltLoss	<i>Difference in filtering loss</i>
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---

### Description

This function calculates differences in filtering loss due to removing a set of J taxa sequentially.

### Usage

```
DiffFiltLoss(X, Order_Ind, Plot = TRUE, Taxa_Names = NULL)
```

### Arguments

X	OTU table, where taxa are columns and samples are rows of the table. It should be a in dataframe format with columns corresponding to taxa names.
Order_Ind	Numeric column order corresponding to taxa importance arrangement.
Plot	A binary TRUE/FALSE value. If TRUE, the function returns plot of sequential differences in filtering loss.
Taxa_Names	Optional taxa labels corresponding to the columns ordering given by Order_Ind.

### Details

This function calculates and plots (if Plot = TRUE) differences in filtering loss sequentially for removing the first j taxa as  $DFL(j+1) = FL(J_{j+1}) - FL(J_j)$  for taxa  $j=1, \dots, p$ .

**Value**

DFL                   Differences in filtering loss values.  
p\_FL                   Plot of the differences in filtering loss.

**Author(s)**

Ekaterina Smirnova

**References**

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

**See Also**

[FiltLoss](#)

**Examples**

```
data(mock2)

# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

#arrange counts in order of increasing number of samples taxa are present in
NP <- NP_Order(Counts)

#obtain numeric column order corresponding to taxa importance arrangement
Order_Ind <- match(NP, names(Prop))
DFL <- DiffFiltLoss(X=Prop, Order_Ind = Order_Ind, Plot = TRUE, Taxa_Names = NP)

#Differences in filtering loss values
DFL$DFL

#Plot of the differences in filtering loss
DFL$p_FL
```

---

FiltLoss

*Filtering Loss*

---

**Description**

Sequential filtering loss calculation for removing a set of  $J_j$  taxa for  $J= 1, \dots, p$ .

**Usage**

```
FiltLoss(X, Order = "NP", Order.user = NULL, type = "Cumu", Plot = TRUE)
```

**Arguments**

X	OTU table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
Order	Taxa ordering. The default ordering is the number of occurrences (NP) of the taxa in all samples. Other types of order are number of connected taxa and weighted number of connected taxa, denoted as "NC", "NCw" respectively. More details about taxa ordering are described in Smirnova et al. User can also specify their preference order with Order.user.
Order.user	User's taxa ordering. This argument takes a character vector of ordered taxa names.
type	Type of filtering loss calculation. "Ind" Individual taxon's filtering loss $FL_u(j)$ "Cumu" Cumulative filtering loss $FL(J)$ due to removing a set of taxa J
Plot	Binary TRUE/FALSE value. If TRUE, the function returns plot of sequential differences in filtering loss.

**Details**

The individual filtering loss due to removing one taxon  $j$  is defined as:

$$FL_u(j) = 1 - (\|X_{-j}\|_F^2 / \|X\|_F^2),$$

where  $X_{-j}$  is the matrix  $X$  without column corresponding to  $j$ th taxon and  $\|Z\|_F$  is the Frobenious norm of a matrix  $Z$ .

The cumulative filtering loss due to removing a set of taxa is defined as:

$$FL(J) = 1 - (\|X_{-J}\|_F^2 / \|X\|_F^2),$$

where  $X_{-J}$  is the  $n \times (p - |J|)$  dimensional matrix obtained by removing the columns indexed by the set  $J$  from the data matrix  $X$ .

The cumulative filtering loss is calculated sequentially for each set of taxa  $J_j, j=1, \dots, p$ .

**Value**

FL	Filtering loss values.
p_FL	Plot of filtering loss values.

**Author(s)**

Ekaterina Smirnova

**References**

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

**See Also**[DiffFiltLoss](#)**Examples**

```
data(mock2)

# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

#Calculate cumulative filtering loss
FL <- FiltLoss(X=Prop, Order = "NP", type = "Cumu", Plot = TRUE)

#Differences in filtering loss values
FL$FL

#Plot of the differences in filtering loss
FL$p_FL
```

---

**FL\_J***Filtering Loss for a set of filtered taxa J*

---

**Description**

This function calculates filtering loss due to removing a group of J taxa.

**Usage**

```
FL_J(X, J)
```

**Arguments**

X	OTU table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
J	A vector of J taxa to be removed. It must be subset of column names of X.

**Value**

FL Filtering loss value.

**Author(s)**

Ekaterina Smirnova

## References

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

## See Also

[FiltLoss](#)

## Examples

```
data(mock2)

# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

#arrange counts in order of increasing number of samples taxa are present in
NP <- NP_Order(Counts)
Counts <- Counts[,NP]

# Extract the taxa names to be removed
J <- colnames(Counts)[1:30]

#Calculate filtering loss due to removing these taxa
FL_J(Counts,J)
```

---

mock2

*Bias experiment data*

---

## Description

These publicly available data (Brooks et al., 2015) were generated as a part of a study designed to evaluate the bias at each step of the VCU sequencing protocol, namely, DNA extraction, PCR amplification, sequencing and taxonomic classification. Mock community samples were created out of 7 vaginally relevant bacteria by mixing prescribed quantities of cells, with quantities varying across samples according to an experimental design described in Brooks et al, 2015. As opposed to the positive controls data, bacteria appear in different proportions across samples. The number of taxa identified by the sequencing and bioinformatics pipeline was 46.

## Usage

```
data(mock2)
```

**Format**

This file contains a count OTU table and a proportion OTU table, each with 240 samples and 46 taxa. A list of true taxa is also given.

---

NCw\_Order                      *Taxa importance ordering by the weighted number of connected taxa*

---

**Description**

Taxa importance ordering by the weighted number of connected taxa

**Usage**

```
NCw_Order(Counts)
```

**Arguments**

Counts                      OTU COUNTS table, where taxa are columns and samples are rows of the table. It should be in data frame format with columns corresponding to taxa names.

**Value**

NCW Taxa names in increasing order of the weighted number of connected taxa.

**Author(s)**

Ekaterina Smirnova

**References**

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

**Examples**

```
data(mock2)
# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

#arrange counts in order of increasing number of samples taxa are present in
NCw <- NCw_Order(Counts)
```

---

NC\_Order

*Taxa importance ordering by the number of connected taxa*

---

### **Description**

Taxa importance ordering by the number of connected taxa

### **Usage**

```
NC_Order(Counts)
```

### **Arguments**

Counts            OTU COUNTS table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.

### **Value**

NC Taxa names in increasing order of the number of connected taxa.

### **Author(s)**

Ekaterina Smirnova

### **References**

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

### **Examples**

```
data(mock2)
# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

#arrange counts in order of increasing number of samples taxa are present in
NC <- NC_Order(Counts)
```



---

NP_Order	<i>Taxa importance ordering by the number of occurrences of the taxa in the n samples</i>
----------	---

---

**Description**

Taxa importance ordering by the number of occurrences of the taxa in the n samples

**Usage**

```
NP_Order(Counts)
```

**Arguments**

Counts            OTU COUNTS table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.

**Value**

NP Taxa names in increasing order of the number of samples taxa are present in.

**Author(s)**

Ekaterina Smirnova

**References**

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

**Examples**

```
data(mock2)
# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

#arrange counts in order of increasing number of samples taxa are present in
NP <- NP_Order(Counts)
```

---

PERFect\_perm                      *Permutation PERFect filtering for microbiome data*

---

### Description

Permutation filtering of the provided OTU table  $X$  at a test level  $\alpha$ . Each set of  $j$  taxa significance is evaluated by fitting the Skew-Normal, Normal,  $t$  or Cauchy distribution to the sampling distribution obtained by permuted taxa labels.

### Usage

```
PERFect_perm(X, infocol = NULL, Order = "NP", Order.user = NULL, normalize = "counts",
  algorithm = "fast", center = FALSE, quant = c(0.1, 0.25, 0.5),
  distr = "sn", alpha = 0.1, rollmean = TRUE, direction = "left", pvals_sim = NULL,
  k = 10000, nbins = 30, hist = TRUE, col = "red", fill = "green",
  hist_fill = 0.2, linecol = "blue")
```

### Arguments

<code>X</code>	OTU table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
<code>infocol</code>	Index vector of the metadata. We assume user only gives a taxa table, but if the metadata of the samples are included in the columns of the input, this option needs to be specified.
<code>Order</code>	Taxa ordering. The default ordering is the number of occurrences (NP) of the taxa in all samples. Other types of order are p-value ordering, number of connected taxa and weighted number of connected taxa, denoted as "pvals", "NC", "NCw" respectively. More details about taxa ordering are described in Smirnova et al. User can also specify their preference order with <code>Order.user</code> .
<code>Order.user</code>	User's taxa ordering. This argument takes a character vector of ordered taxa names.
<code>normalize</code>	Normalizing taxa count. The default option does not normalize taxa count, but user can convert the OTU table into a proportion table using the option "prop" or convert it into a presence/absence table using "pres".
<code>algorithm</code>	Algorithm speed. The default is speed is "fast", which allows the program to efficiently search for significant taxa without computing all the p-values. User must use the default option "hist = FALSE" for the fast algorithm. The alternative setting is "full", which computes all the taxa's p-values.
<code>center</code>	Centering OTU table. The default option does not center the OTU table.
<code>quant</code>	Quantile values used to fit the distribution to log DFL values. The number of quantile values corresponds to the number of parameters in the distribution the data is fitted to. Assuming that at least 50% of taxa are not informative, we suggest fitting the log Skew-Normal distribution by matching the 10%, 25% and 50% percentiles of the log-transformed samples to the Skew-Normal distribution.

distr	The type of distribution to fit log DFL values to. While we suggest using Skew-Normal distribution, and set as the default distribution, other choices are available. "sn" Skew-Normal distribution with 3 parameters: location xi, scale omega^2 and shape alpha "norm" Normal distribution with 2 parameters: mean and standard deviation sd
alpha	Test level alpha, set to 0.1 by default.
rollmean	Binary TRUE/FALSE value. If TRUE, rolling average (moving mean) of p-values will be calculated, with the lag window set to 3 by default.
direction	Character specifying whether the index of the result should be left- or right-aligned or centered compared to the rolling window of observations, set to "left" by default.
pvals_sim	Object resulting from simultaneous PERFect with taxa abundance ordering, allowing user to perform Simultaneous PERFect with p-values ordering. Be aware that the choice of distribution for both methods must be the same.
k	The number of permutations, set to 10000 by default.
nbins	Number of bins used to visualize the histogram of log DFL values, set to 30 by default.
hist	Binary TRUE/FALSE value. If TRUE, the function builds histograms for each taxon.
col	Graphical parameter for color of histogram bars border, set to "red" by default.
fill	Graphical parameter for color of histogram fill, set to "green" by default.
hist_fill	Graphical parameter for intensity of histogram fill, set to 0.2 by default.
linecol	Graphical parameter for the color of the fitted distribution density, set to "blue" by default.

## Details

Filtering is the process of identifying and removing a subset of taxa according to a particular criterion. As opposed to the simultaneous filtering approach, we do not assume that all distributions for each set of taxa are identical and equal to the distribution of simultaneous filtering. Function `PERFect_perm()` filters the provided OTU table `X` and outputs a filtered table that contains signal taxa. `PERFect_perm()` calculates differences in filtering loss DFL for each taxon according to the given taxa order. By default, the function fits Skew-Normal distribution to the log-differences in filtering loss but Normal, t, or Cauchy distributions can be also used.

## Value

If `"algorithm = full"` is chosen, a list is returned containing:

<code>filtX</code>	Filtered OTU table.
<code>info</code>	The metadata information.
<code>pvals</code>	P-values of the test.
<code>DFL</code>	Differences in filtering loss values.

<code>fit</code>	Fitted values and further goodness of fit details passed from the <code>fitdistr()</code> function.
<code>hist</code>	Histogram of log differences in filtering loss.
<code>est</code>	Estimated distribution parameters.
<code>dfl_distr</code>	Plot of differences in filtering loss values.

If "algorithm = fast" is chosen, `fit`, `hist`, `est`, `dfl_distr` will not be returned.

### Author(s)

Ekaterina Smirnova

### References

Azzalini, A. (2005). The skew-normal distribution and related multivariate families. *Scandinavian Journal of Statistics*, 32(2), 159-188.

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutationfiltration of microbiome data", to be submitted.

### See Also

[PERFect\\_sim](#)

### Examples

```
data(mock2)

# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

# Perform simultaenous filtering of the data
res_sim <- PERFect_sim(X=Counts)

#order according to p-values
pvals_sim <- pvals_Order(Counts, res_sim)

## Not run:
# obtain permutation PERFECT results using NP taxa ordering
res_perm <- PERFect_perm(X = Prop, Order.user = pvals_sim, algorithm = "fast")

# permutation perfect colored by FLu values
pvals_Plots(PERFect = res_perm, X = Counts, quantiles = c(0.25, 0.5, 0.8, 0.9), alpha=0.05)

## End(Not run)
```

---

PERFect\_perm\_reorder *Permutation PERFect filtering for microbiome data*

---

### Description

This function filters the provided OTU table  $X$  at a test level  $\alpha$  given a fitted object `perfect_perm` obtained by running `PERFect_perm()` function. `PERFect_perm_reorder()` reevaluates taxa significance p-values for a different taxa ordering.

### Usage

```
PERFect_perm_reorder(X, Order = "NP", Order.user = NULL, res_perm, normalize = "counts",
  center = FALSE, alpha = 0.1, distr = "sn", rollmean = TRUE, direction = "left",
  pvals_sim = NULL)
```

### Arguments

<code>X</code>	OTU table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
<code>Order</code>	Taxa ordering. The default ordering is the number of occurrences (NP) of the taxa in all samples. Other types of order are p-value ordering, number of connected taxa and weighted number of connected taxa, denoted as "pvals", "NC", "NCw" respectively. More details about taxa ordering are described in Smirnova et al. User can also specify their preference order with <code>Order.user</code> .
<code>Order.user</code>	User's taxa ordering. This argument takes a character vector of ordered taxa names.
<code>res_perm</code>	Output of <code>PERFect_perm()</code> function.
<code>normalize</code>	Normalizing taxa count. The default option does not normalize taxa count, but user can convert the OTU table into a proportion table using the option "prop" or convert it into a presence/absence table using "pres".
<code>center</code>	Centering OTU table. The default option does not center the OTU table.
<code>alpha</code>	Test level $\alpha$ , set to 0.1 by default.
<code>distr</code>	The type of distribution used in <code>PERFect_perm()</code> function to obtain <code>res_perm</code> object. "sn" Skew-Normal distribution with 3 parameters: location $\xi$ , scale $\omega^2$ and shape $\alpha$ "norm" Normal distribution with 2 parameters: mean and standard deviation $sd$ "t" Student t-distribution with 2 parameters: $n$ degrees of freedom and noncentrality $ncp$ "cauchy" Cauchy distribution with 2 parameters: location and scale
<code>rollmean</code>	Binary TRUE/FALSE value. If TRUE, rolling average (moving mean) of p-values will be calculated, with the lag window set to 3 by default.

direction	Character specifying whether the index of the result should be left- or right-aligned or centered compared to the rolling window of observations, set to "left" by default.
pvals_sim	Object resulting from simultaneous PERFect with taxa abundance ordering, allowing user to perform Simultaneous PERFect with p-values ordering. Be aware that the choice of distribution for both methods must be the same.

### Details

This function is designed to save computational time needed to obtain and fit the sampling distribution for each taxon if taxa ordering different from the one used in PERFect\_perm() is used. Note, the distribution and OTU table X should match the distribution used in PERFect\_perm().

### Value

res_perm	The perfect_perm object updated according to the alternative taxa ordering. All elements in this list are same as in perfect_perm object given by PERFect() function.
----------	---

### Author(s)

Ekaterina Smirnova

### References

- Azzalini, A. (2005). The skew-normal distribution and related multivariate families. *Scandinavian Journal of Statistics*, 32(2), 159-188.
- Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutationfiltration of microbiome data", to be submitted.

### See Also

[PERFect\\_perm](#)

### Examples

```
data(mock2)

# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

## Not run:
# obtain permutation PERFect results using NP taxa ordering
system.time(res_perm <- PERFect_perm(X=Prop, k = 1000, algorithm = "fast"))

# run PERFect_sim() function and obtain p-values ordering
res_sim <- PERFect_sim(X=Prop)
```

```

# order according to p-values
pvals_sim <- pvals_Order(Counts, res_sim)

# update perfect_perm object according to p-values ordering
res_reorder <- PERFect_perm_reorder(X=Prop, Order.user = pvals_sim, res_perm = res_perm)

# permutation perfect colored by FLu values
pvals_Plots(PERFect = res_perm, X = Counts, quantiles = c(0.25, 0.5, 0.8, 0.9), alpha=0.05)

## End(Not run)

```

---

 PERFect\_sim

*Simulation PERFect filtering for microbiome data*


---

### Description

Simultaneous filtering of the provided OTU table X at a test level alpha. One distribution is fit to taxa simultaneously.

### Usage

```

PERFect_sim(X, infocol = NULL, Order = "NP", Order.user = NULL, normalize = "counts",
            center = FALSE, quant = c(0.1, 0.25, 0.5), distr = "sn",
            alpha = 0.1, rollmean = TRUE, direction = "left", pvals_sim = NULL,
            nbins = 30, col = "red", fill = "green", hist_fill = 0.2,
            linecol = "blue")

```

### Arguments

X	OTU table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names. It could contains columns of metadata.
infocol	Index vector of the metadata. We assume user only gives a taxa table, but if the metadata of the samples are included in the columns of the input, this option needs to be specified.
Order	Taxa ordering. The default ordering is the number of occurrences (NP) of the taxa in all samples. Other types of order are p-value ordering, number of connected taxa and weighted number of connected taxa, denoted as "pvals", "NC", "NCw" respectively. More details about taxa ordering are described in Smirnova et al. User can also specify their preference order with Order.user.
Order.user	User's taxa ordering. This argument takes a character vector of ordered taxa names.
normalize	Normalizing taxa count. The default option does not normalize taxa count, but user can convert the OTU table into a proportion table using the option "prop" or convert it into a presence/absence table using "pres".
center	Centering OTU table. The default option does not center the OTU table.

quant	Quantile values used to fit the distribution to log DFL values. The number of quantile values corresponds to the number of parameters in the distribution the data is fitted to. Assuming that at least 50% of taxa are not informative, we suggest fitting the log Skew-Normal distribution by matching the 10%, 25% and 50% percentiles of the log-transformed samples to the Skew-Normal distribution.
distr	The type of distribution to fit log DFL values to. While we suggest using Skew-Normal distribution, and set as the default distribution, other choices are available. "sn" Skew-Normal distribution with 3 parameters: location xi, scale omega^2 and shape alpha "norm" Normal distribution with 2 parameters: mean and standard deviation sd "t" Student t-distribution with 2 parameters: n degrees of freedom and noncentrality ncp "cauchy" Cauchy distribution with 2 parameters: location and scale
alpha	Test level alpha, set to 0.1 by default.
rollmean	Binary TRUE/FALSE value. If TRUE, rolling average (moving mean) of p-values will be calculated, with the lag window set to 3 by default.
direction	Character specifying whether the index of the result should be left- or right-aligned or centered compared to the rolling window of observations, set to "left" by default.
pvals_sim	Object resulting from simultaneous PERFect with taxa abundance ordering, allowing user to perform Simultaneous PERFect with p-values ordering. Be aware that the choice of distribution for both methods must be the same.
nbins	Number of bins used to visualize the histogram of log DFL values, set to 30 by default.
col	Graphical parameter for color of histogram bars border, set to "red" by default.
fill	Graphical parameter for color of histogram fill, set to "green" by default.
hist_fill	Graphical parameter for intensity of histogram fill, set to 0.2 by default.
linecol	Graphical parameter for the color of the fitted distribution density, set to "blue" by default.

### Details

Filtering is the process of identifying and removing a subset of taxa according to a particular criterion. Function `PERFect_sim()` filters the provided OTU table *X* and outputs a filtered table that contains signal taxa. `PERFect_sim()` calculates differences in filtering loss DFL for each taxon according to the given taxa order. By default, the function fits Skew-Normal distribution to the log-differences in filtering loss but Normal, t, or Cauchy distributions can be also used. This is implementation of Algorithm 1 described in Smirnova et al.

### Value

A list is returned containing:

`filtX` Filtered OTU table.



info	The metadata information.
pvals	P-values of the test.
DFL	Differences in filtering loss values.
fit	Fitted values and further goodness of fit details passed from the fitdistr() function.
hist	Histogram of log differences in filtering loss.
est	Estimated distribution parameters.
pDFL	Plot of differences in filtering loss values.

### Author(s)

Ekaterina Smirnova

### References

Azzalini, A. (2005). The skew-normal distribution and related multivariate families. *Scandinavian Journal of Statistics*, 32(2), 159-188.

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutationfiltration of microbiome data", to be submitted.

### See Also

[PERFect\\_perm](#)

### Examples

```
data(mock2)
# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts
dim(Counts) # 240x46

# Perform simultaenous filtering of the data
res_sim <- PERFect_sim(X=Counts)
dim(res_sim$filtX) # 240x10, removing 36 taxa
colnames(res_sim$filtX) # signal taxa

#permutation perfect colored by FLu values
pvals_Plots(PERFect = res_sim, X = Counts, quantiles = c(0.25, 0.5, 0.8, 0.9), alpha=0.05)
```

---

pvals\_Order                      *Taxa importance ordering by PERFect p-values*

---

**Description**

This function orders taxa by increasing significance of simultaneous PERFect p-values.

**Usage**

```
pvals_Order(Counts, res_sim)
```

**Arguments**

Counts	OTU COUNTS table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
res_sim	Output of PERFect_sim() function.

**Value**

Order\_pvals Taxa names in increasing order of p-values significance.

**Author(s)**

Ekaterina Smirnova

**References**

Smirnova, E., Huzurbazar, H., Jafari, F. "PERFect: permutation filtration of microbiome data", to be submitted.

**See Also**

[PERFect\\_sim](#), [PERFect\\_perm](#)

**Examples**

```
data(mock2)

# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts

# Perform simultaenous filtering of the data
res_sim <- PERFect_sim(X=Counts)

#order according to p-values
pvals_sim <- pvals_Order(Counts, res_sim)
```

---

pvals\_Plots

*Plots of PERFect p-values*

---

### Description

Graphical representation of p-values obtained by running `PERFect_sim()` or `PERFect_perm()` for *j*th taxon colored by quantile values of individual filtering loss.

### Usage

```
pvals_Plots(PERFect, X, quantiles = c(0.25, 0.5, 0.8, 0.9), alpha = 0.1)
```

### Arguments

<code>PERFect</code>	Output of <code>PERFect_sim()</code> or <code>PERFect_perm()</code> function.
<code>X</code>	OTU table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
<code>quantiles</code>	Quantile values for coloring, these are set to 25%, 50%, 80% and 90% percentiles of the individual filtering loss values.
<code>alpha</code>	Alpha level of the test, set to 0.1 by default.

### Value

A list is returned containing:

<code>df</code>	Dataframe of taxa names, p-values, Flu values and quantiles.
<code>p_vals</code>	Plot of p-values.

### Author(s)

Ekaterina Smirnova

### See Also

[PERFect\\_sim](#), [PERFect\\_perm](#)

### Examples

```
data(mock2)
# Proportion data matrix
Prop <- mock2$Prop

# Counts data matrix
Counts <- mock2$Counts
dim(Counts) # 240x46

# Perform simultaenous filtering of the data
res_sim <- PERFect_sim(X=Counts)
```

```

dim(res_sim$filtX)      # 240x10, removing 36 taxa
colnames(res_sim$filtX) # signal taxa

#permutation perfect colored by FLu values
pvals_Plots(PERfect = res_sim, X = Counts, quantiles = c(0.25, 0.5, 0.8, 0.9), alpha=0.05)

```

---

TraditR1

*Traditional Filtering Rule 1*


---

### Description

This rule suggests to remove taxa that are mostly absent in all samples.

### Usage

```
TraditR1(X, thresh =5)
```

### Arguments

X	OTU COUNTS table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
thresh	Numerical value, set to 5 by default. Throughout all samples, taxa that are present for less than this threshold with be removed.

### Value

filtX	Filtered OTU table
-------	--------------------

### Author(s)

Ekaterina Smirnova

### References

Smirnova, E., Huzurbazar, H., Jafari, F. “PERFect: permutation filtration of microbiome data.

### Examples

```

data(mock2)

# Counts data matrix
Counts <- mock2$Counts

# Filtering
Filtered_X <- TraditR1(Counts)

```

**Description**

This rule is adopted from Milici et al. (2016) that removes taxa with low abundance level. Specifically, it keeps taxa with abundance level higher than 0.001%. Then it further selects taxa that satisfy at least one of the following conditions: Present in at least one sample at a relative abundance higher than 1% of the reads of that sample, present in at least 2% of samples at a relative abundance higher than 0.1% for a given sample, present in at least 5% of samples at any abundance level.

**Usage**

```
TraditR2(X, Ab_min = 0.001)
```

**Arguments**

X	OTU COUNTS table, where taxa are columns and samples are rows of the table. It should be a in data frame format with columns corresponding to taxa names.
Ab_min	Numerical value, set to 0.001 by default. Throughout all samples, taxa with abundance less than this threshold with be removed.

**Value**

filtX	Filtered OTU table
-------	--------------------

**Author(s)**

Ekaterina Smirnova

**References**

Smirnova, E., Huzurbazar, H., Jafari, F. “PERFect: permutation filtration of microbiome data.

**Examples**

```
data(mock2)

# Counts data matrix
Counts <- mock2$Counts

# Filtering
Filtered_X <- TraditR2(Counts)
```

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