

# Package ‘FTICRMS’

February 14, 2012

**Type** Package

**Title** Programs for Analyzing Fourier Transform-Ion Cyclotron Resonance Mass Spectrometry Data

**Version** 0.8

**Date** 2009-08-20

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**Depends** Matrix,lattice,splines

**Description** This package was developed partially with funding from the  
NIH Training Program in Biomolecular Technology (2-T32-GM08799).

**License** GPL-2

**Repository** CRAN

**Date/Publication** 2009-12-01 17:18:55

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FTICRMS-package      *Fourier Transform-Ion Cyclotron Resonance Mass Spectrometry (FT-ICR MS) Analysis*

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**Description**

Contains programs for identifying baseline curves and peaks and for statistical analysis of FT-ICR MS data.

**Details**

Package: FTICRMS  
Type: Package  
Version: 0.8  
Date: 2009-08-20  
License: GPL-2

This package was developed partially with funding from the NIH Training Program in Biomolecular Technology (2-T32-GM08799).

**Author(s)**

Don Barkauskas

Maintainer: Don Barkauskas (<barkda@wald.ucdavis.edu>)

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baseline      *Calculate Baselines for Spectroscopic Data*

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**Description**

Computes an estimated baseline curve for a spectrum using the “BXR algorithm,” a method of Xi and Rocke generalized by Barkauskas and Rocke.

**Usage**

```
baseline(spect, init.bd, sm.par = 1e-11, sm.ord = 2, max.iter = 20, tol = 5e-8,  
sm.div = NA, sm.norm.by = c("baseline", "overestimate", "constant"),  
neg.div = NA, neg.norm.by = c("baseline", "overestimate", "constant"),  
rel.conv.crit = TRUE, zero.rm = TRUE, halve.search = FALSE)
```

**Arguments**

spect	vector containing the intensities of the spectrum
init.bd	initial value for baseline; default is flat baseline at median height
sm.par	smoothing parameter for baseline calculation
sm.ord	order of derivative to penalize in baseline analysis
max.iter	convergence criterion in baseline calculation
tol	convergence criterion; see below
sm.div	smoothness divisor in baseline calculation
sm.norm.by	method for smoothness penalty in baseline analysis
neg.div	negativity divisor in baseline calculation
neg.norm.by	method for negativity penalty in baseline analysis
rel.conv.crit	logical; whether convergence criterion should be relative to size of current baseline estimate
zero.rm	logical; whether to replace zeros with average of surrounding values
halve.search	logical; whether to use a halving-line search if step leads to smaller value of function

**Details**

If the spectrum is given by  $y_i$ , then the algorithm works by maximizing the objective function

$$F(\{b_i\}) = \sum_{i=1}^n b_i - \sum_{i=2}^{n-1} A_{1,i}(b_{i-1} - 2b_i + b_{i+1})^2 - \sum_{i=1}^n A_{2,i}[\max\{b_i - y_i, 0\}]^2$$

using Newton's method (with embedded halving line search if `halve.search == TRUE`) using starting value  $b[i] = \text{init.bd}[i]$  for all  $i$ . The middle term controls the smoothness of the baseline and the last term applies a "negativity penalty" when the baseline is above the spectrum.

The smoothing factor `sm.par` corresponds to  $A_1^*$  in Barkauskas (2009) and controls how large the estimated  $n$ th derivative of the baseline is allowed to be (for `sm.ord = n`). From a practical standpoint, values of `sm.ord` larger than two do not seem to adequately smooth the baseline because the Hessian becomes computationally singular for any reasonable value of `sm.par`.

The parameters `sm.div`, `sm.norm.by`, `neg.div`, and `neg.norm.by` determine the methods used to normalize the smoothness and negativity terms. The general forms are  $A_{1,i} = n^4 A_1^* / M_i / p$  and  $A_{2,i} = 1 / M_i / p$ . Here,  $n = \text{length}(\text{spect})$ ;  $p$  is `sm.div` or `neg.div`, as appropriate; and  $M_i$  is determined by `sm.norm.by` or `neg.norm.by`, as appropriate. Values of "baseline" make  $M_i = b'_i$ , where  $b'_i$  is the currently estimated value of the baseline; values of "overestimate" make  $M_i = b'_i - y_i$ ; and values of "constant" make  $M_i = \sigma$ , where  $\sigma$  is an estimate of the noise standard deviation.

The values of `sm.norm.by` and `neg.norm.by` can be abbreviated and both have default value "baseline". The default values of NA for `sm.div` and `neg.div` are translated by default to `sm.div = 0.5223145` and `neg.div = 0.4210109`, which are the appropriate parameters for the FT-ICR mass spectrometry machine that generated the spectra which were used to develop this package. It is distinctly possible that other machines will require different parameters, and almost certain that

other spectroscopic technologies will require different parameters; see Barkauskas (2009a) for a description for how these parameters were obtained.

If `zero.rm == TRUE` and  $y_a, \dots, y_{a+k} = 0$ , then these values of the spectrum are set to be  $(y_{a-1} + y_{a+k+1})/2$ . (For typical MALDI FT-ICR spectra, a spectrum value of zero indicates an erased harmonic and should not be considered a real data point.)

### Value

A list containing the following items:

<code>baseline</code>	The computed baseline
<code>iter</code>	The number of iterations for convergence
<code>changed</code>	Numeric vector of length <code>iter</code> containing the number of indicator variables that switched value on each iteration
<code>hs</code>	Numeric vector of length <code>iter</code> containing the number of halving line-searches done on each iteration

### Note

The original algorithm was developed by Yuanxin Xi and David Rocke. The code in this package was first adapted from a Matlab program by Yuanxin Xi, then modified to account for the new methodology in Barkauskas (2009a).

`halve.search = FALSE` is recommended unless both `sm.norm.by == "constant"` and `neg.norm.by == "constant"`.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

### References

- Barkauskas, D.A. and D.M. Rocke. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043
- Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.
- Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.
- Xi, Y. and Rocke, D.M. (2008) "Baseline Correction for NMR Spectroscopic Metabolomics Data Analysis". *BMC Bioinformatics*, **9**:324.

### See Also

[run.baselines](#)

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display.tests	<i>Display Full Test Information for Peaks</i>
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### Description

Displays full test information (not just  $p$ -values) for peaks generated by `run.analysis`.

### Usage

```
display.tests(sig.rows = "all", summ = "anova", tests,
             form = parameter.list$form,
             use.model = parameter.list$use.model, ...)
```

### Arguments

sig.rows	numeric or character vector used to select rows of sigs; default value returns all significant tests
summ	either a function or string representing a function which can be applied to the output of use.model or "none"
tests	numeric or character vector used to select rows of clust.mat; default value returns the rows in clust.mat corresponding to the rows in sigs[sig.rows,]
form	formula for use in lm; default is the one that was used to generate the significant peaks
use.model	function or string representing a function; what test to apply to data
...	arguments to be passed to use.model

### Details

If use.model in `run.analysis` evaluates to anything other than `t.test`, then the only thing reported on each peak by `run.analysis` is the  $p$ -value. This program takes a specified subset of the significant peaks and returns a list consisting of the models generated by use.model (if summ = "none") or summ applied to those models. Typical values for summ include `anova` and `summary`.

Although the program is designed to be used on significant peaks, by defining tests directly in the function call, you can access any of the peaks in clust.mat. If tests is defined in the function call, its value overrides anything specified by sig.rows.

### Value

A list with components equal to the models or summaries for the requested peaks.

### Note

clust.mat and sig.mat must be defined in the workspace for this program to work—for example, in the results file output by `run.analysis`.

**Author(s)**

Don Barkauskas (<barkda@wald.ucdavis.edu>)

**See Also**

[run.analysis](#), [anova](#), [lm](#), [t.test](#)

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extract.pars

*Extract Parameters from File*

---

**Description**

Extracts the parameters in the file specified by `par.file` and returns them in list form.

**Usage**

```
extract.pars(par.file = "parameters.RData", root.dir = ".")
```

**Arguments**

<code>par.file</code>	string containing name of parameters file
<code>root.dir</code>	string containing directory of parameters file to be extracted from

**Details**

Used by [run.analysis](#) to record all the parameter choices in an analysis for future reference.

**Value**

A list with the following components:

<code>add.norm</code>	logical; whether to normalize additively or multiplicatively on the log scale
<code>add.par</code>	additive parameter for "shiftedlog" or "glog" options for <code>trans.method</code>
<code>align.fcn</code>	function (and inverse) to apply to masses before (and after) applying <code>align.method</code>
<code>align.method</code>	alignment algorithm for peaks
<code>base.dir</code>	directory for baseline files
<code>bhbysubj</code>	logical; whether to look for number of large peaks by subject (i.e., combining replicates) or by spectrum
<code>calc.all.peaks</code>	whether to calculate all possible peaks or only sufficiently large ones
<code>cluster.constant</code>	parameter used in running <code>cluster.method</code>
<code>cluster.method</code>	method for determining when two peaks from different spectra are the same
<code>cor.thresh</code>	threshold correlation for declaring isotopes
<code>covariates</code>	data frame containing covariates used in analysis

FDR	False Discovery Rate in Benjamini-Hochberg test
FTICRMS.version	Version of <a href="#">FTICRMS</a> that created file
form	formula used in use.model
gengamma.quantiles	whether to use generalized gamma quantiles when calculating large peaks
halve.search	whether to use a halving-line search if step leads to smaller value of function
isotope.dist	maximum distance for declaring isotopes
lrg.dir	directory for significant peaks file
lrg.file	name of file for storing large peaks
lrg.only	whether to consider only peaks that have at least one “large” peak; i.e., identified by <a href="#">run.lrg.peaks</a>
masses	specific masses to test
max.iter	convergence criterion in baseline calculation
min.spect	minimum number of spectra necessary for peak to be used in <a href="#">run.analysis</a>
neg.div	negativity divisor in baseline calculation
neg.norm.by	method for negativity penalty in baseline analysis
norm.peaks	which peaks to use in normalization
norm.post.repl	logical; whether to normalize after combining replicates
normalization	type of normalization to use on spectra before statistical analysis
num.pts	number of points needed for peak fitting
oneside.min	minimum number of points on each side of local maximum for peak fitting
overwrite	whether to replace existing files with new ones
par.file	string containing name of parameters file
peak.dir	directory for peak location files
peak.method	method for locating peaks
peak.thresh	threshold for declaring large peak
pre.align	shifts to apply before running <a href="#">run.strong.peaks</a>
pval.fcn	function to calculate $p$ -values
R2.thresh	$R^2$ value needed for peak fitting
raw.dir	directory for raw data files
rel.conv.crit	whether convergence criterion should be relative to size of current baseline estimate
repl.method	how to deal with replicates
res.dir	directory for result file
res.file	name for results file
root.dir	directory for parameters file and raw data directory
sm.div	smoothness divisor in baseline calculation

sm.norm.by	method for smoothness penalty in baseline analysis
sm.ord	order of derivative to penalize in baseline analysis
sm.par	smoothing parameter for baseline calculation
subs	subset of spectra to use for analysis
subtract.base	whether to subtract calculated baseline from spectrum
tol	convergence criterion in baseline calculation
trans.method	data transformation method
use.model	what model to apply to data
zero.rm	whether to replace zeros in spectra with average of surrounding values

### Note

`do.call(make.par.file, extract.pars())` recreates the original parameter file

`align.method`, `cluster.method`, `neg.norm.by`, `normalization`, `peak.method`, `sm.norm.by`, and `trans.method` can be abbreviated.

See [make.par.file](#) for a summary of which programs use each of the parameters in the list.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

### References

- Barkauskas, D.A. and D.M. Rocke. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043
- Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.
- Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.
- Benjamini, Y. and Hochberg, Y. (1995) "Controlling the false discovery rate: a practical and powerful approach to multiple testing." *J. Roy. Statist. Soc. Ser. B*, **57**:1, 289–300.
- Xi, Y. and Rocke, D.M. (2008) "Baseline Correction for NMR Spectroscopic Metabolomics Data Analysis". *BMC Bioinformatics*, **9**:324.

### See Also

[make.par.file](#), [run.analysis](#)

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locate.peaks	<i>Locate Peaks in a FT-ICR MS Spectrum</i>
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### Description

Locates peaks in FT-ICR MS spectra assuming that the peaks are roughly parabolic on the log scale.

### Usage

```
locate.peaks(peak.base, num.pts = 5, R2.thresh = 0.98,
             onside.min = 1, peak.method = c("parabola", "locmaxes"),
             thresh = -Inf)
```

### Arguments

peak.base	numeric matrix with two columns containing the masses and the transformed spectrum intensities
num.pts	minimum number of points needed to have a peak
R2.thresh	minimum $R^2$ needed to have a peak
onside.min	minimum number of points needed on each side of the local maximum
peak.method	how to locate peaks
thresh	only local maxes that are larger than this will be checked to see if they are peaks

### Details

If `peak.method == "parabola"`, the algorithm works by locating local maxima in the spectrum, then seeing if any `num.pts` consecutive points with at least `onside.min` point(s) on each side of the local maximum have a coefficient of determination ( $R^2$ ) of at least `R2.thresh` when fitted with a quadratic. If, in addition, the coefficient of the squared term is negative, then this is declared a peak and the vertex of the corresponding parabola is located. The coordinates of the vertex give the components `Center_hat` and `Max_hat` in the return value. The `Width_hat` component is the negative reciprocal of the coefficient of the squared term.

If `peak.method == "locmax"`, then the algorithm merely returns the set of local maxima larger than `thresh`, and the `Width_hat` component of the return value is NA.

### Value

A data frame with columns

<code>Center_hat</code>	estimated mass of peak
<code>Max_hat</code>	estimated intensity of peak
<code>Width_hat</code>	estimated width of peak

**Note**

An extremely large value for `Width_hat` most likely indicates a bad fit.

`peak.method` can be abbreviated. Using `peak.method = "locmax"` will vastly speed up the runtime, but may affect the quality of the analysis.

As noted in both papers in the References, a typical FT-ICR MS spectrum has far more peaks than can be accounted for by actual compounds. Thus, defining a good value of `thresh` will vastly speed up the computation without materially affecting the analysis.

**Author(s)**

Don Barkauskas (<barkda@wald.ucdavis.edu>)

**References**

Barkauskas, D.A. and D.M. Rocke. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043

Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.

Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.

**See Also**

[run.peaks](#)

---

make.par.file

*Create Parameter File for FT-ICR MS Analysis*

---

**Description**

Creates a file of parameters that can be read by the functions in the **FTICRMS** package

**Usage**

```
make.par.file(covariates, form, par.file = "parameters.RData", root.dir = ".", ...)
```

**Arguments**

<code>covariates</code>	data frame with rownames given by raw data files with extensions (e.g., ".txt") stripped
<code>form</code>	object of class " <b>formula</b> " to be used for testing using <code>covariates</code>
<code>par.file</code>	string containing name of file
<code>root.dir</code>	string containing location for file
<code>...</code>	parameters whose default values are to be overwritten (see below)

## Details

Creates a file with name given by `par.file` in directory given by `root.dir` which contains values for all of the parameters used in the programs in the **FTICRMS** package. The possible parameters that can be included in `...`, their default values, their descriptions, and the program(s) in which they are used are as follows:

<code>add.norm = TRUE</code>	logical; whether to normalize additively or multiplicatively
<code>add.par = 0</code>	additive parameter for "shiftedlog" or "glog" options for t
<code>align.fcn = NA</code>	function (and inverse) to apply to masses before (and after) ap
<code>align.method = "spline"</code>	alignment algorithm for peaks
<code>base.dir = paste(root.dir, "/Baselines", sep="")</code>	directory for baseline files
<code>bhbysubj = FALSE</code>	logical; whether to look for number of large peaks by subject
<code>calc.all.peaks = FALSE</code>	logical; whether to calculate all possible peaks or only sufficie
<code>cluster.constant = 10</code>	parameter used in running <code>cluster.method</code>
<code>cluster.method = "ppm"</code>	method for determining when two peaks from different spectra
<code>cor.thresh = 0.8</code>	threshold correlation for declaring isotopes
<code>FDR = 0.1</code>	False Discovery Rate in Benjamini-Hochberg test
<code>FTICRMS.version = "0.8"</code>	Version of <b>FTICRMS</b> that created file
<code>gengamma.quantiles = TRUE</code>	logical; whether to use generalized gamma quantiles when cal
<code>halve.search = FALSE</code>	logical; whether to use a halving-line search if step leads to sn
<code>isotope.dist = 7</code>	maximum distance for declaring isotopes
<code>lrg.dir = paste(root.dir, "/Large_Peaks", sep="")</code>	directory for large peaks file
<code>lrg.file = "lrg_peaks.RData"</code>	name of file for storing large peaks
<code>lrg.only = TRUE</code>	logical; whether to consider only peaks that have at least one 's
<code>masses = NA</code>	specific masses to test
<code>max.iter = 20</code>	convergence criterion in baseline calculation
<code>min.spect = 1</code>	minimum number of spectra necessary for peak to be used in
<code>neg.div = NA</code>	negativity divisor in baseline calculation
<code>neg.norm.by = "baseline"</code>	method for negativity penalty in baseline analysis
<code>norm.peaks = "common"</code>	which peaks to use in normalization
<code>norm.post.repl = FALSE</code>	logical; whether to normalize after combining replicates
<code>num.pts = 5</code>	number of consecutive points needed for peak fitting
<code>oneside.min = 1</code>	minimum number of points on each side of local maximum fo
<code>overwrite = FALSE</code>	logical; whether to replace existing files with new ones
<code>par.file = "parameters.RData"</code>	string containing name of parameters file
<code>peak.dir = paste(root.dir, "/All_Peaks", sep="")</code>	directory for peak location files
<code>peak.method = "parabola"</code>	method for locating peaks
<code>peak.thresh = 3.798194</code>	threshold for declaring large peak
<code>pre.align = FALSE</code>	shifts to apply before running <code>run.strong.peaks</code>
<code>pval.fcn = "default"</code>	function to calculate $p$ -values; default is overall $p$ -value of test
<code>R2.thresh = 0.98</code>	$R^2$ value needed for peak fitting
<code>raw.dir = paste(root.dir, "/Raw_Data", sep="")</code>	directory for raw data files
<code>rel.conv.crit = TRUE</code>	whether convergence criterion should be relative to size of cur
<code>repl.method = "max"</code>	how to deal with replicates
<code>res.dir = paste(root.dir, "/Results", sep="")</code>	directory for results file
<code>res.file = "analyzed.RData"</code>	name for results file
<code>root.dir = "."</code>	directory for parameters file and raw data

sm.div = NA	smoothness divisor in baseline calculation
sm.norm.by = "baseline"	method for smoothness penalty in baseline analysis
sm.ord = 2	order of derivative to penalize in baseline analysis
sm.par = 1e-11	smoothing parameter for baseline calculation
subs	subset of spectra to use for analysis
subtract.base = FALSE	logical; whether to subtract calculated baseline from spectrum
tol = 5e-8	convergence criterion in baseline calculation
trans.method = "shiftedlog"	data transformation method
use.model = "lm"	what model to apply to data
zero.rm = TRUE	whether to replace zeros in spectra with average of surrounding

### Value

No value returned; the file `par.file` is simply created in `root.dir`.

### Note

`do.call(make.par.file, extract.pars())` recreates the original parameter file.

See the individual function help pages for each function for more detailed descriptions of the above parameters.

`align.method`, `cluster.method`, `neg.norm.by`, `normalization`, `peak.method`, `sm.norm.by`, and `trans.method` can be abbreviated.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

### References

- Barkauskas, D.A. and D.M. Rocke. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043
- Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.
- Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.
- Xi, Y. and Rocke, D.M. (2008) "Baseline Correction for NMR Spectroscopic Metabolomics Data Analysis". *BMC Bioinformatics*, **9**:324.

### See Also

[extract.pars](#)

---

`run.all`*Complete Analysis of FT-ICR MS Data*

---

**Description**

A wrapper that calls all six functions needed for a full analysis.

**Usage**

```
run.all(par.file = "parameters.RData", root.dir = ".")
```

**Arguments**

<code>par.file</code>	string containing the name of the parameters file
<code>root.dir</code>	string containing location of raw data directory and parameters file

**Details**

Requires `par.file` to be in place before starting—for example by creating it with `make.par.file`.

Calls (in order) `run.baselines`, `run.peaks`, `run.lrg.peaks`, `run.strong.peaks`, `run.cluster.matrix`, and `run.analysis`.

**Note**

The analysis described in Barkauskas *et al.* (2008) can be (approximately) reproduced using the following parameter values instead of the defaults:

```
add.par = 10
calc.all.peaks = TRUE
gengamma.quantiles = FALSE
max.iter = 30
neg.norm.by = "constant"
peak.thresh = 4
pval.fcn = function(x){anova(x)[2,5]}
rel.conv.crit = FALSE
sm.norm.by = "constant"
subtract.base = TRUE
zero.rm = FALSE
```

(It is only an approximate reproduction because the stopping criterion for baseline calculation used in the article turned out to be a poor one and is no longer supported in the package. This shouldn't make a very large difference, however.)

**Author(s)**

Don Barkauskas (<barkda@wald.ucdavis.edu>)

## References

- Barkauskas, D.A. and D.M. Roche. (2009a) “A general-purpose baseline estimation algorithm for spectroscopic data”. to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043
- Barkauskas, D.A. *et al.* (2009b) “Analysis of MALDI FT-ICR mass spectrometry data: A time series approach”. *Analytica Chimica Acta*, **648**:2, 207–214.
- Barkauskas, D.A. *et al.* (2009c) “Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data”. *Bioinformatics*, **25**:2, 251–257.
- Benjamini, Y. and Hochberg, Y. (1995) “Controlling the false discovery rate: a practical and powerful approach to multiple testing.” *J. Roy. Statist. Soc. Ser. B*, **57**:1, 289–300.
- Xi, Y. and Roche, D.M. (2008) “Baseline Correction for NMR Spectroscopic Metabolomics Data Analysis”. *BMC Bioinformatics*, **9**:324.

## See Also

[make.par.file](#), [run.baselines](#), [run.peaks](#), [run.lrg.peaks](#), [run.strong.peaks](#), [run.cluster.matrix](#), [run.analysis](#)

---

run.analysis

*Test for Significant Peaks in FT-ICR MS by Controlling FDR*

---

## Description

Takes the file generated by [run.cluster.matrix](#) and tests the peaks using Benjamini-Hochberg to control the False Discovery Rate.

## Usage

```
run.analysis(form, covariates, FDR = 0.1, norm.post.repl = FALSE,
             norm.peaks = c("common", "all", "none"), normalization,
             add.norm = TRUE, repl.method = "max", use.model = "lm",
             pval.fcn = "default", lrg.only = TRUE, masses = NA,
             isotope.dist = 7, root.dir = ".", lrg.dir,
             lrg.file = lrg_peaks.RData, res.dir,
             res.file = "analyzed.RData", overwrite = FALSE,
             use.par.file = FALSE, par.file = "parameters.RData",
             bhbysubj = TRUE, subs, ...)
```

## Arguments

form	object of class “ <a href="#">formula</a> ” to be used by <code>use.model</code> for testing using <code>covariates</code>
covariates	data frame containing covariates used in analysis
FDR	False Discovery Rate in Benjamini-Hochberg test
norm.post.repl	logical; whether to normalize after combining replicates
norm.peaks	which peaks to use in normalization

normalization	type of normalization to use on spectra before statistical analysis; kept for compatibility (see below)
add.norm	logical; whether to normalize additively or multiplicatively on the log scale
repl.method	function or string representing the name of a function; how to deal with replicates
use.model	function or string representing the name of a function; what test to apply to data
pval.fcn	function to extract $p$ -values; default is overall $p$ -value of test
lrg.only	logical; whether to consider only peaks that have at least one “large” peak; i.e., identified by <code>run.lrg.peaks</code>
masses	specific masses to test
isotope.dist	maximum distance for declaring isotopes
root.dir	directory for parameters file and raw data
lrg.dir	directory for large peaks file; default is <code>paste(root.dir, "/Large_Peaks", sep = "")</code>
lrg.file	name of file to store large peaks in
res.dir	directory for results file; default is <code>paste(root.dir, "/Results", sep = "")</code>
res.file	name for results file
overwrite	logical; whether to replace existing files with new ones
use.par.file	logical; if TRUE, then parameters are read from <code>par.file</code> in directory <code>root.dir</code>
par.file	string containing name of parameters file
bhbysubj	logical; whether to look for number of large peaks by subject (i.e., combining replicates) or by spectrum
subs	subset of spectra to use for analysis; see below
...	additional parameters to be passed to <code>use.model</code>

## Details

Reads in information from file created by `run.cluster.matrix` and creates a file named `res.file` in directory `res.dir` which contains the following variables:

amps	matrix of transformed amplitudes of alignment peaks
bysubjvar	a vector which tells which rows of <code>covariates</code> are identified as the same subject
centers	matrix of calculated masses of alignment peaks
clust.mat	matrix of transformed amplitudes of peaks used in statistical testing
min.FDR	FDR level required to get at least one significant test given the starting set of peaks
sigs	matrix containing all tests which are significant under at least one scenario
which.sig	matrix containing all peaks tested
parameter.list	if <code>use.par.file = TRUE</code> , a list generated by <code>extract.pars</code> ; otherwise not defined

**Value**

No value returned; the file is simply created.

**Note**

If `use.par.file == TRUE` and other parameters are entered into the function call, then the parameters entered in the function call overwrite those read in from the file. Note that this is opposite from the behavior for **FTICRMS** versions 0.7 and earlier.

`norm.peaks` determines the peaks used for normalization: "common" normalizes each spectrum using the average peak height of the alignment peaks from that spectrum in amps; "all" normalizes each spectrum using the average peak height of all peaks in that spectrum.

`normalization` is obsolete but is included for compatibility with previous versions of the package. The valid normalization schemes translate to the new scheme as follows: "common" is `norm.post.repl = FALSE` and `norm.peaks = "common"`; "postbase" is `norm.post.repl = FALSE` and `norm.peaks = "all"`; "postrepl" is `norm.post.repl = TRUE` and `norm.peaks = "all"`; and "none" is `norm.peaks = "none"` (and `norm.post.repl = FALSE`, although this value is irrelevant).

Replicates for the same subject are assumed to be determined by the unique values of `covariates$subj`. (Future implementations will allow for other methods of defining this.) To analyze replicates as independent samples, use `repl.method = "none"`. This will also speed up the run time if there are no replicates in the data set.

The argument `subs` can be logical or numeric or character; if it is defined, then `covariates` is modified to `covariates[subs, , drop=F]`.

If `masses` is not `NULL`, then the listed masses plus anything that could be in the first `isotope.dist - 1` isotope peaks of each mass are tested.

If something other than the  $p$ -value for the overall test statistic is needed, then the user-defined function for `pval.fcn` should have the form `pval.fcn = function(x){...}`, where `x` is a model object of the type returned by `use.model`; and should have a return value of the desired  $p$ -value.

If `use.model` evaluates to `t.test`, then the difference between the two groups for each peak is recorded in `which.sig$Delta` and `sigs$Delta`; otherwise, these columns consist entirely of NA entries.

Each rowname of `sigs` and `which.sig` represents the range of masses that were used to form that peak. The columns of those objects give the  $p$ -value of the peaks in each row, the number of samples that had large peaks for each row, and the significance of each test, coded as

NA	peak not eligible for B-H
0	peak eligible for B-H but not declared significant
1	peak declared significant

The "S" labels refer to the number of large peaks that were necessary for a row to be eligible. For example, the column labeled S5 in `sigs` used as its starting set of  $p$ -values all rows which had `which.sig$num.lrg >= 5`. If `bhbysubj == TRUE`, then the entries of `num.lrg` are obtained by going subject-by-subject and for each mass counting the number of subjects who had at least one spectrum with a large peak at that mass; otherwise, `num.lrg` for each mass is simply the total

number of spectra that had a large peak at that mass.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

### References

Barkauskas, D.A. and D.M. Rocke. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043

Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.

Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.

Benjamini, Y. and Hochberg, Y. (1995) "Controlling the false discovery rate: a practical and powerful approach to multiple testing." *J. Roy. Statist. Soc. Ser. B*, **57**:1, 289–300.

### See Also

[run.strong.peaks](#)

---

run.baselines

*Calculate and Store Baselines for Spectroscopic Data*

---

### Description

Takes the spectra from files in `raw.dir`, calculates the baselines from them, and writes the results in the directory `base.dir`.

### Usage

```
run.baselines(root.dir = ".", raw.dir, base.dir, overwrite = FALSE,
              use.par.file = FALSE, par.file = "parameters.RData",
              sm.par = 1e-11, sm.ord = 2, max.iter = 20, tol = 5e-8,
              sm.div = NA, sm.norm.by = c("baseline", "overestimate", "constant"),
              neg.div = NA, neg.norm.by = c("baseline", "overestimate", "constant"),
              rel.conv.crit = TRUE, zero.rm = TRUE, halve.search = FALSE)
```

### Arguments

<code>root.dir</code>	directory for parameters file and raw data
<code>raw.dir</code>	directory for raw data files; default is <code>paste(root.dir, "/Raw_Data", sep = "")</code>
<code>base.dir</code>	directory for baseline files; default is <code>paste(root.dir, "/Baselines", sep = "")</code>
<code>overwrite</code>	logical; whether to replace existing files with new ones

use.par.file	logical; if TRUE, then parameters are read from par.file in directory root.dir
par.file	string containing name of parameters file
sm.par	smoothing parameter for baseline calculation
sm.ord	order of derivative to penalize in baseline analysis
max.iter	convergence criterion in baseline calculation
tol	convergence criterion
sm.div	smoothness divisor in baseline calculation
sm.norm.by	method for smoothness penalty in baseline analysis
neg.div	negativity divisor in baseline calculation
neg.norm.by	method for negativity penalty in baseline analysis
rel.conv.crit	logical; whether convergence criterion should be relative to size of current baseline estimate
zero.rm	logical; whether to replace zeros with average of surrounding values
halve.search	logical; whether to use a halving-line search if step leads to smaller value of function

### Details

Goes through the entire directory `raw.dir` file-by-file and computes each baseline using [baseline](#), then writes the spectrum and the baseline to a file in directory `base.dir`. The name of the new file is the same as the name of the old file with “.txt” replaced by “.RData”, and the new file is ready to be used by [run.peaks](#).

The files in `raw.dir` must be in a specific format (future versions of the package will allow for more flexibility). The files should be two-column text files with mass in the first column and spectrum intensity in the second column. There should be no header row (just start the file with the first data point). The columns can be either comma-separated or whitespace-separated and the program will automatically detect which each file is. The decimal separator should be “.”, as using “,” will cause errors in reading the files.

See [baseline](#) for details of all the parameters after `par.file`.

### Value

No value returned; the files are simply created.

### Note

If `use.par.file == TRUE` and other parameters are entered into the function call, then the parameters entered in the function call overwrite those read in from the file. Note that this is opposite from the behavior for [FTICRMS](#) versions 0.7 and earlier.

The values of `sm.norm.by` and `neg.norm.by` can be abbreviated and both have default value “baseline”.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

## References

- Barkauskas, D.A. (2009) “Statistical Analysis of Matrix-Assisted Laser Desorption/Ionization Fourier Transform Ion Cyclotron Resonance Mass Spectrometry Data with Applications to Cancer Biomarker Detection”. Ph.D. dissertation, University of California at Davis.
- Barkauskas, D.A. *et al.* (2009) “Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data”. *Bioinformatics*, **25**:2, 251–257.
- Xi, Y. and Rocke, D.M. (2008) “Baseline Correction for NMR Spectroscopic Metabolomics Data Analysis”. *BMC Bioinformatics*, **9**:324.

## See Also

[baseline](#), [run.peaks](#)

---

run.cluster.matrix      *Identify Equivalent Peaks from Different Subjects*

---

## Description

Takes the file generated by [run.lrg.peaks](#), identifies equivalent peaks in each spectrum, and fills in missing values.

## Usage

```
run.cluster.matrix(pre.align = FALSE, align.method = c("PL",
  "spline", "affine", "none"), align.fcn = NA,
  trans.method = c("shiftedlog", "glog", "none"),
  add.par = 0, subtract.base = FALSE,
  lrg.only = TRUE, calc.all.peaks = FALSE,
  masses = NA, isotope.dist = 7,
  cluster.method = c("ppm", "constant", "usewidth"),
  cluster.constant = 10, num.pts = 5,
  R2.thresh = 0.98, onside.min = 1, min.spect = 1,
  peak.method = c("parabola", "locmaxes"),
  bhbysubj = TRUE, covariates, root.dir = ".",
  base.dir, peak.dir, lrg.dir,
  lrg.file = "lrg_peaks.RData", overwrite = FALSE,
  use.par.file = FALSE, par.file = "parameters.RData")
```

## Arguments

- |              |  |
|--------------|--|
| pre.align    | either FALSE, or a numeric vector of shifts to apply to spectra, or a four-component list (of the form described in the Note section below) to be used before identifying peaks from different spectra |
| align.method | alignment algorithm for peaks  |
| align.fcn    | function (and inverse) to apply to masses before (and after) applying align.method; see below  |

trans.method	type of transformation to use on spectra before statistical analysis
add.par	additive parameter for "shiftedlog" or "glog" options for trans.method
subtract.base	logical; whether to subtract calculated baseline from spectrum
lrg.only	logical; whether to consider only peaks that have at least one "large" peak; i.e., identified by run.lrg.peaks
calc.all.peaks	logical; whether to calculate all possible peaks or only sufficiently large ones
masses	specific masses to test
isotope.dist	maximum distance for declaring isotopes
cluster.method	method for determining when two peaks from different spectra are the same
cluster.constant	parameter used in running cluster.method
num.pts	number of consecutive points needed for peak fitting
R2.thresh	$R^2$ value needed for peak fitting
oneside.min	minimum number of points on each side of local maximum for peak fitting
min.spect	minimum number of spectra necessary for peak to be used in <a href="#">run.analysis</a>
peak.method	method for locating peaks
bhbysubj	logical; whether to look for number of large peaks by subject (i.e., combining replicates) or by spectrum
covariates	data frame with rownames given by raw data files with extensions (e.g., ".txt") stripped; only needed if bhbysubj == TRUE
root.dir	directory for parameters file and raw data
base.dir	directory for baseline files; default is paste(root.dir, "/Baselines", sep = "")
peak.dir	directory for peak location files; default is paste(root.dir, "/All_Peaks", sep = "")
lrg.dir	directory for large peaks file; default is paste(root.dir, "/Large_Peaks", sep = "")
lrg.file	name of file to store large peaks in
overwrite	logical; whether to replace existing files with new ones
use.par.file	logical; if TRUE, then parameters are read from par.file in directory root.dir
par.file	string containing name of parameters file

## Details

Reads in information from file created by [run.strong.peaks](#), calculates the cluster matrix, fills in missing values, and overwrites the file named lrg.file in lrg.dir. The resulting file contains variables

amps	data frame of amplitudes created by <a href="#">run.strong.peaks</a>
centers	data frame of centers created by <a href="#">run.strong.peaks</a>
clust.mat	data frame with columns given by samples and rows given by the distinct peaks in the samples
lrg.mat	data frame of same size as clust.mat with entries given by TRUE if the peak was large in that spectrum and FALSE otherwise
lrg.peaks	the data frame of significant peaks created by <a href="#">run.lrg.peaks</a>
num.lrg	number of subjects (or spectra if bhbysubj == TRUE) with a large peak at the corresponding mass

and is ready to be used by [run.analysis](#).

### Value

No value returned; the file is simply created.

### Note

If `use.par.file == TRUE` and other parameters are entered into the function call, then the parameters entered in the function call overwrite those read in from the file. Note that this is opposite from the behavior for **FTICRMS** versions 0.7 and earlier.

`align.method`, `cluster.method`, `peak.method`, and `trans.method` can be abbreviated.

If `align.fcn` is not NA, then it should consist of a list with components `fcn` and `inv`, each of class function. `align.fcn$fcn` should take a vector of masses as its argument and return a vector of transformed masses. (Typically, this will be transforming masses to frequencies; see Zhang (2005).) `align.fcn$inv` should be the inverse function of `align.fcn$fcn`.

If `align.method == "spline"`, then alignment consists of making the transformed masses of the strong peaks all agree exactly with their means, then shifting the rest of the transformed masses via an interpolation spline generated using [interpSpline](#). If `align.method == "PL"`, then the same is done but interpolation is done piecewise linearly between the strong peaks. If `align.method == "leastsq"`, then the transformed masses of the strong peaks are aligned to their means using a least-squares affine fit for each spectrum. In any of these cases, if there are no strong peaks, `align.method` is changed to "none" with a warning. If there is exactly one strong peak, then alignment is by a simple shift in each spectrum on the transformed masses. If there are exactly two strong peaks, then the alignment is by a simple affine transformation on the transformed masses in each spectrum. If `align.method = "spline"` and there are exactly three strong peaks, then alignment is piecewise affine on the transformed masses (i.e., identical to `align.method = "PL"`).

If `align.method = "leastsq"`, it is strongly recommended that you supply a value for `align.fcn` that makes the data points (approximately) equally-spaced.

Defining a value for `min.spect` can vastly speed up the run time at the (small) cost of a little flexibility in doing the statistical analysis in [run.analysis](#). For exploratory data analysis, this should probably be left alone, but once the peak criterion has been established, further analyses will go much more quickly with `min.spect` re-defined. The value can either be an integer, which is interpreted as the number of spectra; or a number between 0 and 1, in which case it is interpreted as a fraction of the total number of spectra. In either case, the values of `clust.mat`, `lrg.mat`, and `num.lrg` saved in `lrg.file` are only those masses which have at least `min.spect` large peaks among the spectra.

`pre.align = FALSE` is used if the spectra have already been aligned by the mass spectroscopists. If it is not FALSE, it can either be a vector of additive shifts to be applied to the spectra, or a list with components `targets`, `actual`, and `align.method`. In the last case, `targets` is a vector of target masses, and `actual` is a matrix with `length(targets)` columns and a row for each spectrum, `actual[i,j]` being the mass in spectrum `i` that should be matched exactly to `target[j]`, with NA being a valid entry in `actual`. The alignment is then done as in the description in the above paragraph, depending on the number of non-missing values in row `i`.

Suppose `cluster.constant = K` and we have two peaks in different spectra with masses  $m_1 < m_2$ . If `cluster.method == "constant"`, then the peaks are considered to be the same peak if we have  $m_2 - m_1 < K$ . If `cluster.method == "ppm"`, then the peaks are considered to be the same peak if

we have  $m_2 - m_1 < Km_2/10^6$ . If `cluster.method == "usewidth"`, then the algorithm uses the observation that  $\log(\text{Width\_hat})$  and  $\log(\text{Center\_hat})$  appear to be linearly related. Tolerances are computed using this relationship.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

### References

Barkauskas, D.A. and D.M. Rocke. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043

Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.

Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.

Zhang, L.-K. *et al.* (2005) "Accurate mass measurements by Fourier transform mass spectrometry". *Mass Spectrom Rev*, **24**:2, 286–309.

### See Also

[run.lrg.peaks](#), [run.strong.peaks](#), [interpSpline](#)

---

run.lrg.peaks                      *Extract "Large" Peaks from Files*

---

### Description

Takes the files output by [run.peaks](#), extracts "large" peaks, combines them into a single data frame, and writes the data frame to a file.

### Usage

```
run.lrg.peaks(trans.method = c("shiftedlog", "glog", "none"),
             add.par = 0, subtract.base = FALSE,
             root.dir = ".", peak.dir, base.dir, lrg.dir,
             lrg.file = lrg_peaks.RData, overwrite = FALSE,
             use.par.file = FALSE, par.file = "parameters.RData",
             calc.all.peaks = FALSE, gengamma.quantiles = TRUE,
             peak.thresh = 3.798194, subs)
```

**Arguments**

trans.method	type of transformation to use on spectra before statistical analysis
add.par	additive parameter for "shiftedlog" or "glog" options for trans.method
subtract.base	logical; whether to subtract calculated baseline from spectrum
root.dir	directory for parameters file and raw data
peak.dir	directory for peak location files; default is paste(root.dir, "/All_Peaks", sep = "")
base.dir	directory for baseline files; default is paste(root.dir, "/Baselines", sep = "")
lrg.dir	directory for large peaks file; default is paste(root.dir, "/Large_Peaks", sep = "")
lrg.file	name of file to store large peaks in
overwrite	logical; whether to replace existing files with new ones
use.par.file	logical; if TRUE, then parameters are read from par.file in directory root.dir
par.file	string containing name of parameters file
calc.all.peaks	logical; whether to calculate all possible peaks or only sufficiently large ones
gengamma.quantiles	logical; whether to use generalized gamma quantiles when calculating large peaks
peak.thresh	threshold for declaring large peak; see below
subs	subset of spectra to use for analysis; see below

**Details**

Reads in information from each file created by [run.peaks](#), extracts peaks which are "large" (see below), and creates the file lrg.file in lrg.dir. The resulting file contains the data frame lrg.peaks, which has columns

Center_hat	estimated mass of peak
Max_hat	estimated intensity of peak
Width_hat	estimated width of peak
File	name of file the peak was extracted from, with "\_peaks.RData" deleted

and is ready to be used by [run.strong.peaks](#).

**Value**

No value returned; the file is simply created.

**Note**

If `use.par.file == TRUE` and other parameters are entered into the function call, then the parameters entered in the function call overwrite those read in from the file. This is opposite from the behavior for **FTICRMS** versions 0.7 and earlier.

`trans.method` can be abbreviated.

If `gengamma.quantiles == TRUE`, then a peak is “large” if it is at least `peak.thresh` times as large as the estimated baseline at that point.

If `gengamma.quantiles == FALSE`, then a peak is “large” if it has zero weight in the data generated by `run.peaks` for the spectrum it comes from when using Tukey’s biweight with parameter  $K = 1.5 * \text{peak.thresh}$  to estimate center and scale.

If `subs` is not defined, the algorithm finds large peaks for all files in `peak.dir`. If it is defined, `subs` can be logical or numeric or character; if it is defined, then the algorithm finds large peaks for all entries in `subs` (character) or `list.files(peak.dir)[subs]` (logical or numeric).

**Author(s)**

Don Barkauskas (<barkda@wald.ucdavis.edu>)

**References**

Barkauskas, D.A. and D.M. Rocke. (2009a) “A general-purpose baseline estimation algorithm for spectroscopic data”. to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043

Barkauskas, D.A. *et al.* (2009b) “Analysis of MALDI FT-ICR mass spectrometry data: A time series approach”. *Analytica Chimica Acta*, **648**:2, 207–214.

Barkauskas, D.A. *et al.* (2009c) “Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data”. *Bioinformatics*, **25**:2, 251–257.

**See Also**

[run.peaks](#), [run.cluster.matrix](#)

---

run.peaks

*Locate Potential Peaks in FT-ICR MS Spectra*

---

**Description**

Takes baseline-corrected data and locates potential peaks in the spectra.

**Usage**

```
run.peaks(trans.method = c("shiftedlog", "glog", "none"),
          add.par = 0, subtract.base = FALSE, root.dir = ".",
          base.dir, peak.dir, overwrite = FALSE,
          use.par.file = FALSE, par.file = "parameters.RData",
          num.pts = 5, R2.thresh = 0.98, onside.min = 1,
```

```

peak.method = c("parabola", "locmaxes"),
calc.all.peaks = FALSE, gengamma.quantiles = TRUE,
peak.thresh = 3.798194)

```

### Arguments

trans.method	type of transformation to use on spectra before statistical analysis
add.par	additive parameter for "shiftedlog" or "glog" options for trans.method
subtract.base	logical; whether to subtract calculated baseline from spectrum
root.dir	directory for parameters file and raw data
base.dir	directory for baseline files; default is paste(root.dir, "/Baselines", sep = "")
peak.dir	directory for peak location files; default is paste(root.dir, "/All_Peaks", sep = "")
overwrite	logical; whether to replace existing files with new ones
use.par.file	logical; if TRUE, then parameters are read from par.file in directory root.dir
par.file	string containing name of parameters file
num.pts	number of consecutive points needed for peak fitting
R2.thresh	$R^2$ value needed for peak fitting
oneside.min	minimum number of points on each side of local maximum for peak fitting
peak.method	method for locating peaks
calc.all.peaks	logical; whether to calculate all possible peaks or only sufficiently large ones
gengamma.quantiles	logical; whether to use generalized gamma quantiles when calculating large peaks
peak.thresh	threshold for declaring large peak; see below

### Details

Reads in information from each file created by [run.baselines](#), calls [locate.peaks](#) to find potential peaks, and writes the output to a file in directory `peak.dir`. The name of each new file is the same as the name of the old file with ".RData" replaced by "\\_peaks.RData". The resulting file contains the data frame `all.peaks`, which has columns

Center_hat	estimated mass of peak
Max_hat	estimated intensity of peak
Width_hat	estimated width of peak

and is ready to be used by [run.lrg.peaks](#).

The parameters `gengamma.quantiles` and `peak.thresh` are relevant only if `calc.all.peaks = FALSE`. In that case, if `gengamma.quantiles == TRUE`, then `peak.thresh` is interpreted as a multiplier for the baseline. Anything larger than `peak.thresh` times the estimated baseline is declared to

be a real peak. If `gengamma.quantiles == FALSE`, then `peak.thresh` is interpreted as two-thirds of the value of  $K$  used in a Tukey's biweight estimation of center and scale (so roughly equal to the number of standard deviations above the mean for iid normal data). Anything with weight zero in the calculation is then declared to be a real peak.

### Value

No value returned; the files are simply created.

### Note

If `use.par.file == TRUE` and other parameters are entered into the function call, then the parameters entered in the function call overwrite those read in from the file. Note that this is opposite from the behavior for **FTICRMS** versions 0.7 and earlier.

`peak.method` and `trans.method` can be abbreviated.

Using `calc.all.peaks == FALSE` can speed up computation time immensely, but will affect the final result. It probably won't affect it much, but *caveat emptor*.

### Author(s)

Don Barkauskas (<barkda@wald.ucdavis.edu>)

### References

Barkauskas, D.A. and D.M. Roche. (2009a) "A general-purpose baseline estimation algorithm for spectroscopic data". to appear in *Analytica Chimica Acta*. doi:10.1016/j.aca.2009.10.043

Barkauskas, D.A. *et al.* (2009b) "Analysis of MALDI FT-ICR mass spectrometry data: A time series approach". *Analytica Chimica Acta*, **648**:2, 207–214.

Barkauskas, D.A. *et al.* (2009c) "Detecting glycan cancer biomarkers in serum samples using MALDI FT-ICR mass spectrometry data". *Bioinformatics*, **25**:2, 251–257.

### See Also

[run.baselines](#), [run.lrg.peaks](#), [locate.peaks](#)

---

run.strong.peaks

*Locate Peaks that are "Large" in All Samples*

---

### Description

Takes the file generated by [run.peaks](#), extracts all peaks that are "large" in all samples, and writes the results to a file.

**Usage**

```
run.strong.peaks(cor.thresh = 0.8, isotope.dist = 7, pre.align = FALSE,
  align.method = c("PL", "spline", "affine", "none"),
  align.fcn = NA, root.dir = ".", lrg.dir,
  lrg.file = "lrg_peaks.RData", overwrite = FALSE,
  use.par.file = FALSE, par.file = "parameters.RData")
```

**Arguments**

<code>cor.thresh</code>	threshold correlation for declaring isotopes
<code>isotope.dist</code>	maximum distance for declaring isotopes
<code>pre.align</code>	either FALSE, or a numeric vector of shifts to apply to spectra, or a three-component list (of the form described in the Note section below) to be used before identifying peaks from different spectra
<code>align.method</code>	alignment algorithm for peaks
<code>align.fcn</code>	function (and inverse) to apply to masses before (and after) applying <code>align.method</code> ; see below
<code>root.dir</code>	directory for parameters file and raw data
<code>lrg.dir</code>	directory for large peaks file; default is <code>paste(root.dir, "/Large_Peaks", sep = "")</code>
<code>lrg.file</code>	name of file to store large peaks in
<code>overwrite</code>	logical; whether to replace existing files with new ones
<code>use.par.file</code>	logical; if TRUE, then parameters are read from <code>par.file</code> in directory <code>root.dir</code>
<code>par.file</code>	string containing name of parameters file

**Details**

Reads in information from file created by `run.lrg.peaks`, locates peaks which appear in all samples, and overwrites the file `lrg.file` in `lrg.dir`. The resulting file contains variables

<code>amps</code>	data frame of amplitudes of non-isotope peaks that occur in all samples
<code>centers</code>	data frame of centers of non-isotope peaks that occur in all samples
<code>lrg.peaks</code>	the data frame of significant peaks created by <code>run.lrg.peaks</code>

and is ready to be used by `run.cluster.matrix`.

**Value**

No value returned; the file is simply created.

**Note**

If `use.par.file == TRUE` and other parameters are entered into the function call, then the parameters entered in the function call overwrite those read in from the file. Note that this is opposite from

the behavior for **FTICRMS** versions 0.7 and earlier.

If `align.fcn` is not NA, then it should consist of a list with components `fcn` and `inv`, each of class function. `align.fcn$fcn` should take a vector of masses as its argument and return a vector of transformed masses. (Typically, this will be transforming to the frequency domain; see Zhang (2005).) `align.fcn$inv` should be the inverse function of `align.fcn$fcn`. If `align.method == "leastsq"`, it is strongly recommended that you supply a value for `align.fcn` that makes the masses (approximately) equally-spaced.

`align.method` can be abbreviated. If `align.method == "spline"`, then alignment consists of making the transformed masses of the strong peaks all agree exactly with their means, then shifting the rest of the transformed masses via a cubic interpolation spline generated using `interpSpline`. If `align.method == "PL"`, then the same is done but interpolation is piecewise linear between the strong peaks. If `align.method == "leastsq"`, then the transformed masses of the strong peaks are aligned to their means using a least-squares affine fit for each spectrum. In any of these cases, if there are no strong peaks, `align.method` is changed to "none" with a warning. If there is exactly one strong peak, then alignment is by a simple shift in each spectrum on the transformed masses. If there are exactly two strong peaks, then the alignment is by a simple affine transformation on the transformed masses in each spectrum. If `align.method == "spline"` and there are exactly three strong peaks, then alignment is piecewise affine on the transformed masses (i.e., identical to using `align.method = "PL"`).

`pre.align = FALSE` is used if the spectra have already been aligned by the mass spectroscopists. If it is not FALSE, it can either be a vector of additive shifts to be applied to the spectra, or a list with components `targets`, `actual`, and `align.method`. In the last case, `targets` is a vector of target masses, and `actual` is a matrix with `length(targets)` columns and a row for each spectrum, `actual[i, j]` being the mass in spectrum `i` that should be matched exactly to `target[j]`, with NA being a valid entry in `actual`. The alignment is then done row-by-row as in the description in the above paragraph, depending on the number of non-missing values in row `i`).

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### See Also

[run.lrg.peaks](#), [run.cluster.matrix](#), [interpSpline](#)

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