

# Package ‘afmToolkit’

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**Title** Functions for Atomic Force Microscope Force-Distance Curves  
Analysis

**Version** 0.0.1

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**Description** Set of functions for analyzing Atomic Force Microscope (AFM) force-distance curves. It allows to obtain the contact and unbinding points, perform the baseline correction, estimate the Young's modulus, fit up to two exponential decay function to a stress-relaxation / creep experiment, obtain adhesion energies. These operations can be done either over a single F-d curve or over a set of F-d curves in batch mode.

**Depends** R (>= 3.2.2), ggplot2

**Imports** stats, utils, minpack.lm, grid, gridExtra, scales, dplyr, DBI,  
assertthat, tibble

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afmAdhesionEnergy	<i>Adhesion Energy</i>
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---

### Description

Finds the adhesion and the full detach energies from the retract segment of the AFM F-d curve.

### Usage

```
afmAdhesionEnergy(afmdata, width = 1, lagdiff = width, mul, mdj = NULL)
```

### Arguments

afmdata	An afmdata or afmexperiment class variables. Baseline correction should have been done already.
width	Width of the window for the local regression (in vector position units)
lagdiff	Lag for estimating the differences in Delta (or slopes) signal. By default it takes the same value as the window with.
mul	Multiplier for the calculating the threshold in the estimation of jumps and peaks in the Delta signal
mdj	Minimum distance between jumps. If none is given then it will be set equal to width

### Value

An afmdata class variable which will consist on the original input afmdata variable plus a new list named AdhEner with the following fields:

Points Array containing the indices of the retract segment where the adhesion begins, the unbinding event takes place and the adhesion ends.

Energies Data frame with three columns: E1adh, E2adh and Etotal, being the first one the energy from the beginning of the adhesion until the unbinding event, then second one the energy from the unbinding event until the full detachment of the tip, and the third one, the sum of them.

## Examples

```
path <- path.package("afmToolkit")
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path)
data <- afmContactPoint(data, width = 20, mul1 = 1, mul2 = 10)
data <- afmDetachPoint(data, width = 20, mul1 = 2, mul2 = 30)
data <- afmBaselineCorrection(data)
data <- afmAdhesionEnergy(data, width = 20, mul = 10)
str(data$AdhEner)
```

---

afmBaselineCorrection *Performs a baseline correction to an AFM F-z curve*

---

## Description

This function performs the baseline correction to an AFM F-z curve within an afmdata structure.

It subtracts a best fit line to the curve: for the approach and contact segments, it fits a line to the approach curve points where for which  $|z| > ZPointApp$  and for the retract segment, it fits a line to the retract curve where  $|z| > ZpointRet$ .

If no  $ZPointApp$  is given and the contact point has been already estimated (via `afmContactPoint()` function), then it is found as

$$ZPointApp = 0.7ContactPoint + 0.3max(Z)$$

## Usage

```
afmBaselineCorrection(afmdata, ZPointApp = NULL, ZPointRet = NULL,
  fitpause = c("approach", "retract", "none"), vsTime = FALSE)
```

## Arguments

afmdata	An afmdata structure.
ZPointApp	Point in the approach segment of the curve that defines the approach baseline
ZPointRet	Point in the retract segment of the curves that defines the retract baseline
fitpause	Behaviour for the baseline correction at the pause segment: if "approach" (default), the pause segment is corrected using the best line fit done on the approach segment, if "retract" the best line fit of the retract segment is used, if "none", no baseline correction is done on the pause segment.
vsTime	Logical. If TRUE then the baseline correction is performed following the Force vs time approach described by S. Moreno-Flores ( <i>Moreno Flores (2016)</i> ).

## Value

afmdata An afmdata structure identical to the one in the input, but with an additional ForceCorrected column in the data dataframe of the afmdata structure.

## References

Moreno Flores (2016). Baseline correction of AFM force curves in the force-time representation. *Microscopy Research and Technique*, 79, (11), pp. 1045-1049.

## Examples

```
AFMcurve <- afmReadJPK("force-save-JPK-2h.txt.gz", path = path.package("afmToolkit"))
ZPointApp <- 6.43e-6
ZPointRet <- 6.45e-6
AFMcurve <- afmBaselineCorrection(AFMcurve,ZPointApp = ZPointApp,ZPointRet = ZPointRet)
plot(AFMcurve)

# Without providing ZPointApp
AFMcurve <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path.package("afmToolkit"))
AFMcurve <- afmContactPoint(AFMcurve,width = 10,mul1 = 1,mul2 = 20,
                           loessSmooth = FALSE)
AFMcurve <- afmBaselineCorrection(AFMcurve)
plot(AFMcurve)
```

---

afmContactPoint	<i>Contact point</i>
-----------------	----------------------

---

## Description

Find the contact point in for the Force-Distance curve following the local regression and two thresholds methods described in *Microscopy Research and Technique* 2013 (see reference).

## Usage

```
afmContactPoint(afmdata, width = 1, mul1, mul2, lagdiff = width, Delta = TRUE,
               loessSmooth = FALSE)
```

## Arguments

afmdata	A Force-Distance curve with the afmdata structure. It should be a list with at least the 'data' field with a data frame of at least 4 columns.
width	Width of the window for the local regression (in vector position units)
mul1	First multiplier for the first alarm threshold
mul2	Second multiplier for the second alarm threshold
lagdiff	Lag for estimating the differences in Delta (or slopes) signal. By default it takes the same value as the window with.
Delta	Logical. If TRUE, then the statistic for determining the contact point is the differences between two consecutive values of the slope of the local regression line. If FALSE then the slope itself is used.
loessSmooth	Logical If TRUE, a loess smoothing (via loess.smooth()) is done prior to the determination of the contact point. The span of the smoothing is 0.05 (5 approach segment).

**Value**

An afmdata class variable which will consist on the original input afmdata variable plus a new list named CP with the following fields:

CP The contact point value.

iCP The position in the array for the contact point value.

delta The delta signal.

noise The noise of the delta signal

**References**

Benitez R., Moreno-Flores S., Bolos V. J. and Toca-Herrera J.L. (2013). "A new automatic contact point detection algorithm for AFM force curves". *Microscopy research and technique*, **76** (8), pp. 870-876.

**See Also**

[afmDetachPoint](#)

**Examples**

```
path <- path.package("afmToolkit")
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path)
width <- 20
mul1 <- 1
mul2 <- 10
data <- afmContactPoint(data, width = width, mul1 = mul1, mul2 = mul2)
## Not run:
plot(data, segment = "approach") + geom_vline(xintercept = data$CP$CP, lty = 2)

## End(Not run)
```

---

afmdata

*AFM data*

---

**Description**

This function creates an afmdata structure, which is as list with at least one field called data which is a data frame with a valid AFM data, that is, at least 3 variables called "Z", "Force", and "Segment".

**Usage**

```
afmdata(data, dstr = "Z", Fstr = "Force", Segstr = "Segment", tstr = "Time",
  params = list(SpringConstant = numeric(), curvename = NULL ))
```

**Arguments**

data	A data frame consisting in 3 or 4 columns. A minimum of "Z" (or "distance"), "Force" and "Segment". Optionally a fourth column with "Time" could be added.
dstr	Character string with the possible names for the distance variable.
Fstr	Character string with the possible names for the force variable.
Segstr	Character string with the possible names for the Segment variable.
tstr	Character string with the possible names for the time variable.
params	A list that may contain parameters describing the F-d curve. At least will contain the SpringConstant and the curvname, being the former the cantilever spring constant and the latter a F-d curve ID. Function afmReadJPK will try to obtain the spring constant from the file header and the curvname from the data file name.

**Value**

An object of class afmdata

**See Also**

[afmexperiment](#)

**Examples**

```
#Making some artificial data following a L-J 12-6 potential
n <- 1000
z <- seq(from = 9e-3, to = 1e-1, length.out = n )
u0 <- 1e-5
z0 <- 1e-2
Force <- -u0*(12*z0^6/z^7-12*z0^12/z^13)
Segment <- rep("approach",n)
AFMcurve <- afmdata(data.frame(Z = z, Force = Force, Segment = Segment))
plot(AFMcurve)
```

---

afmDetachPoint

*Detach point*


---

**Description**

Find the detach point (or unbinding point) for the Force-Distance curve following the local regression and two thresholds methods described in *Microscopy Research and Technique* 2013 (see reference).

The procedure is similar to the one used by the `afmContactPoint()` function for obtaining the contact point.

**Usage**

```
afmDetachPoint(afmdata,width=1,mul1,mul2, lagdiff = width, Delta=TRUE,
               loessSmooth = FALSE)
```

**Arguments**

afmdata	A Force-Distance curve with the afmdata structure. It should be a list with at least the 'data' field with a data frame of at least 4 columns.
width	Width of the window for the local regression (in vector position units)
mul1	First multiplier for the first alarm threshold
mul2	Second multiplier for the second alarm threshold
lagdiff	Lag for estimating the differences in Delta (or slopes) signal. By default it takes the same value as the window with.
Delta	Logical. If TRUE, then the statistic for determining the contact point is the differences between two consecutive values of the slope of the local regression line. If FALSE then the slope itself is used.
loessSmooth	Logical If TRUE, a loess smoothing (via loess.smooth()) is done prior to the determination of the contact point. The span of the smoothing is 0.05 (5 approach segment).

**Value**

An afmdata class variable which will consist on the original input afmdata variable plus a new list named DP with the following fields:

DP The detach point value.

iDP The position in the array for the detach point value.

delta The delta signal.

noise The noise of the delta signal

**References**

Benitez R., Moreno-Flores S., Bolos V. J. and Toca-Herrera J.L. (2013). "A new automatic contact point detection algorithm for AFM force curves". *Microscopy research and technique*, **76** (8), pp. 870-876.

**See Also**

[afmContactPoint](#)

**Examples**

```
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path.package("afmToolkit"))
width <- 10
mul1 <- 2
mul2 <- 40
data <- afmDetachPoint(data, width = width, mul1 = mul1, mul2 = mul2)
```

```
## Not run:
plot(data, segment = "retract") + geom_vline(xintercept = data$DP$DP, lty = 2)

## End(Not run)
```

---

afmExpDecay

*Exponential decay fit*


---

## Description

Fits a viscoelastic exponential decay in a Force-Relaxation or Creep experiments as described in Nanotechnology 2010 (see references).

## Usage

```
afmExpDecay(afmdata, nexpt = 2, tmax = NULL, type = c("CH", "CF"), plt = TRUE,
  ...)
```

## Arguments

afmdata	An object of afmdata class with a <b>pause</b> segment and a <b>Time</b> column in the data dataframe.
nexpt	Number of exponentials in the Prony series to be fitted. Currently only one or two exponentials are supported. Default is 2.
tmax	Maximum time considered in the relaxation curve. It defaults to Inf, meaning that the whole pause segment is considered.
type	Type of the experiment. Can be either "CH" (Constant Height) for a force-relaxation experiment or "CF" (Constant Force) for a creep experiment. Default is type = "CH".
plt	Logical. If TRUE (default) then a plot of the pause segment with the overlay of the fit is shown.
...	Options passed to the nlsM() function from the minpack.lm package. At least should contain the starting values (start = list(...)) for the Levenberg-Marquardt nonlinear least square method.

## Value

An afmdata class variable which will consist on the original input afmdata variable plus a new list named ExpFit with the following fields:

expdecayModel: A nls object returned from nlsM() function.

expdecayFit: The values predicted by the fit, returned from the predict() function.

## References

Susana Moreno-Flores, Rafael Benitez, Maria dM Vivanco and Jose Luis Toca-Herrera (2010). "Stress relaxation and creep on living cells with the atomic force microscope: a means to calculate elastic moduli and viscosities of cell components". Nanotechnology, **21** (44), pp. 445101.



## Examples

```
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path.package("afmToolkit"))
width <- 20
mul1 <- 1
mul2 <- 10
data <- afmContactPoint(data, width = width, mul1 = mul1, mul2 = mul2)
data <- afmDetachPoint(data, width = width, mul1 = mul1, mul2 = mul2)
data <- afmBaselineCorrection(data)
data <- afmExpDecay(data, nexp = 2, type = "CH")
```

---

afmexperiment

*AFM experiment*

---

## Description

This function creates an afmexperiment structure, which is a list (or an array) of elements of afmdata class.

## Usage

```
afmexperiment(data, ID=NULL)
```

## Arguments

data	A variable of afmdata class, or a list of elements of afmdata class.
ID	Character string with the identifier of the data variable or a string array in case data is a list of afmdata variables.

## Value

An object of class afmexp.

## See Also

[afmdata](#)

## Examples

```
dataFolder <- paste(path.package("afmToolkit"), "afmexperiment", sep = "/")
dataFiles <- list.files(dataFolder, pattern = "force", full.names = FALSE)
data <- lapply(dataFiles, afmReadJPK, path = dataFolder)
names(data) <- dataFiles
data <- afmexperiment(data)
plot(data[[1]])
```

---

afmExtract

*Extract computed parameters from an afmexperiment*


---

### Description

Extracts some parameters from an afmexperiment for an easy further analysis.

### Usage

```
afmExtract(afmexperiment, params = list("YM", "AE", "ED"), opt.param = NULL)
```

### Arguments

afmexperiment	Data of afmexperiment class.
params	List of parameters to extract from the data.
opt.param	Optional parameter or factor in the params field of the afmdata list to add to the data extraction.

### Value

A data frame with the name of the curve and the corresponding values of the parameters extracted.

### Examples

```
## Not run:
require(dplyr) # Not really necessary

# Load the data
data(batchExperiment)

# Process the afmexperiment
data <- afmContactPoint(batchExperiment, width = 50, mul1 = 1, mul2 = 10)
data <- afmDetachPoint(data, width = 50, mul1 = 1, mul2 = 10)
data <- afmBaselineCorrection(data)
data <- afmZeroPointSlope(data)
data <- afmIndentation(data)
data <- afmYoungModulus(data, thickness = 2e-7, params = list(alpha = 22))
data <- afmExpDecay(data, plt = FALSE)
data <- afmAdhesionEnergy(data, mul = 7)

# Extract the values of the parameters obtained in the analysis
afmExpParams <- afmExtract(data, opt.param = "type")

# Plotting the Young's Modulus
afmExpParams[[1]] %>% ggplot(aes(x = type, y = YM)) + geom_boxplot()
ylab("Young's Modulus (Pa)")

## End(Not run)
```

---

afmIndentation	<i>afmIndentation</i>
----------------	-----------------------

---

**Description**

This function computes the deformation of the sample from the calibrated Force-Distance curve, by subtracting Z to the Zero Force Point calculated with `afmZeroPointSlope` function.

**Usage**

```
afmIndentation(afmdata)
```

**Arguments**

afmdata	An afmdata object. It should be a valid afmdata object upon which the Contact Point, the baseline correction and the Zero Force Point must have been calculated first (using functions <code>afmContactPoint()</code> , <code>afmBaselineCorrection()</code> and <code>afmZeroPointSlope()</code> )
---------	---

**Value**

Returns a list with one field:

afmdata: An afmdata class in which a Indentation column is added in the data field.

**Examples**

```
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path.package("afmToolkit"))
data <- afmContactPoint(data, width = 20, mul1 = 1, mul2 = 20)
data <- afmDetachPoint(data, width = 40, mul1 = 3, mul2 = 40)
data <- afmBaselineCorrection(data)
data <- afmZeroPointSlope(data, segment = "approach")
data <- afmIndentation(data)
head(data$data)
```

---

afmReadJPK	<i>Read Nanowizard JPK ascii file</i>
------------	---------------------------------------

---

**Description**

Read an ascii JPK file.

Reads an ascii JPK file with one to three headers.

**Usage**

```
afmReadJPK(filename, path = "", FColStr = "Vertical",
ZColStr = "Height (measured & smoothed)", tColStr = "Segment Time")
```

**Arguments**

filename	String with the name of the jpk file.
path	Path to the folder where the file is.
FColStr	String with a pattern identifying the Force column.
ZColStr	String with a pattern identifying the Z column.
tColStr	String with a pattern identifying the Time column.

**Value**

A list containing a field 'data' which is a data frame

**Examples**

```
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path.package("afmToolkit"))
str(data)
```

---

afmReadJPKFolder	<i>Read all Nanowizard JPK ascii files in a folder</i>
------------------	--

---

**Description**

Read all JPK ascii files in a given folder. It searches for all files containing a given patter (".txt" by default) and uses the afmReadJPJ function.

**Usage**

```
afmReadJPKFolder(folder, pattern = ".txt", ...)
```

**Arguments**

folder	Name of the folder containing the jpk files.
pattern	Pattern that will identify the jok files (".txt" by default).
...	Other parameters passed to afmReadJPK function.

**Value**

An afmexperiment class data structure with all F-d curves.

**Examples**

```
folder <- paste(path.package("afmToolkit"), "afmexperiment", sep = "/")
data <- afmReadJPKFolder(folder = folder)
str(data)
```

---

afmReadVeeco	<i>Read Bruke Nanoscope Veeco ascii file</i>
--------------	--

---

**Description**

Read an ascii Veeco file.

Reads an ascii Veeco file with one or two segments.

**Usage**

```
afmReadVeeco(filename, path = "")
```

**Arguments**

filename       String with the name of the jpk file.

path            Path to the folder where the file is.

**Value**

A list containing a field 'data' which is a data frame

**Examples**

```
data <- afmReadVeeco("veeco_file.txt.gz", path = path.package("afmToolkit"))
str(data)
```

---

afmReadVeecoFolder	<i>Read all Bruke Nanoscope Veeco ascii files in a folder</i>
--------------------	---

---

**Description**

Read all Veeco ascii files in a given folder. It searches for all files containing a given patter (".txt" by default) and uses the afmReadVeeco function.

**Usage**

```
afmReadVeecoFolder(folder, pattern = ".txt")
```

**Arguments**

folder        Name of the folder containing the Veeco files.

pattern       Pattern that will identify the Veeco files (".txt" by default).

**Value**

An afmexperiment class data structure with all F-d curves.

**Examples**

```
folder <- paste(path.package("afmToolkit"), "veecoFolder", sep = "/")
data <- afmReadVeecoFolder(folder = folder)
str(data)
```

---

afmYoungModulus	<i>afmYoungModulus</i>
-----------------	------------------------

---

**Description**

This function computes the Young's Modulus of the sample from the approach curve using Hertz's contact model for a pyramidal tip.

**Usage**

```
afmYoungModulus(afmdata, thickness = NULL, model = "Hertz", geometry =
  c("pyramid", "paraboloid"), silent = TRUE, params)
```

**Arguments**

afmdata	An afmdata object. It should be a valid afmdata object upon which the Contact Point, the baseline correction and the Zero Force Point and the Indentation must have been calculated first (using functions <code>afmContactPoint()</code> , <code>afmBaselineCorrection()</code> , <code>afmZeroPointSlope()</code> , and <code>afmIndentation()</code> )
thickness	Thickness (in m) of the surface. The Force - Indentation fit will be done for values of the Indentation variable smaller than the thickness. If no value is given, it will be done for all values in the curve for which the Indentation is negative.
model	Contact mechanics model to be used. Currently only Hertz's pure elastic model is available.
geometry	Geometry of the tip. Currently only pyramidal (default) and paraboloid geometries are implemented.
silent	Logical value. If FALSE it prints the fit model summary (via <code>summary.lm()</code> ). Default value is TRUE
params	A list containing different parameters of the model: e.g. nu (Poisson's ratio) or alpha (internal angle, in degrees, of the pyramidal tip) or R (tip radius, in the paraboloid geometry)

**Value**

An afmdata class variable which will consist on the original input afmdata variable plus a new list named YoungModulus with the following fields:

`YoungModulus` The Young's modulus value (in Pa).

`fitYM` The Force vs Indentation<sup>2</sup> fit as an `lm` object.

`fitdata` The subset of the data used in the fit.

**Examples**

```

data <- afmReadJPK("force-save-JPK-2h.txt.gz", path = path.package("afmToolkit"))
data <- afmContactPoint(data, width = 20, mul1 = 1, mul2 = 20)
data <- afmDetachPoint(data, width = 40, mul1 = 3, mul2 = 40)
data <- afmBaselineCorrection(data)
data <- afmZeroPointSlope(data, segment = "approach")
data <- afmIndentation(data)
data <- afmYoungModulus(data, thickness = 1e-8, params = list(alpha = 22),
                        silent = TRUE)
print(data$YoungModulus$YoungModulus)

```

---

afmZeroPointSlope      *Zero Force Point and Slope*

---

**Description**

This function finds the point of zero force (real contact point) and the slope of the contact part of the Force-Distance curve.

**Usage**

```
afmZeroPointSlope(afmdata, fstar = 0, segment = c("approach", "retract"))
```

**Arguments**

afmdata	An afmdata object. It should be a valid afmdata object upon which the Contact Point and the baseline correction must have been calculated first (using functions afmContactPoint() and afmBaselineCorrection())
fstar	Value such that $fstar * sd$ is to be considered as zero Force, where $sd$ is the standard deviation of Force at the baseline. It takes $fstar = 0$ as default value, meaning that zero force is actually zero.
segment	The segment on which everything is calculated.

**Value**

An afmdata class variable which will consist on the original input afmdata variable plus a new list named Slopes with the following fields: Z0Point: Point of zero force. Slope: Slope of the best fit line in the contact part of the Force-Distance curve.

**Examples**

```

data <- afmReadJPK("force-save-JPK-2h.txt.gz", path = path.package("afmToolkit"))
data <- afmContactPoint(data, width = 20, mul1 = 1, mul2 = 20)
data <- afmDetachPoint(data, width = 40, mul1 = 3, mul2 = 40)
data <- afmBaselineCorrection(data)
data <- afmZeroPointSlope(data, segment = "approach")
## Not run:
plot(data, segment = "approach") + geom_vline(xintercept = data$Slopes$Z0Point, lty = 2)

```

```
## End(Not run)
```

---

append.afmdata	<i>Append to an afmdata list.</i>
----------------	-----------------------------------

---

### Description

This function appends a list to an existing afmdata structure. It is used internally by several afm\* functions when attaching the results to the input afmdata variable. This function should not be used directly unless by experienced users.

### Usage

```
append.afmdata(afmdata, x, name = NULL)
```

### Arguments

afmdata	The afmdata to which the new list is going to be joined.
x	A list to be appended.
name	The name of new field of the resulting afmdata object. If none is given, it is the same as x.

### Value

The new list of class afmdata

---

batchExperiment	<i>Example of an afmexperiment data class.</i>
-----------------	--

---

### Description

An afmexperiment list containing 14 afmdata Force-distance experiments. Each experiment has three segments ("approach", "pause" and "retract") and they are divided in two groups depending on the covering of the sample ("CHI" for Chitosan, and "PAH" for Polyallylamine hydrochloride).

### Usage

```
batchExperiment
```



**Format**

An afmexperiment class consisting on a list of 14 afmdata class elements each one having the following fields:

**data** Data frame with the data itself with a variable number of rows (between 4692 and 6142) and 4 variables:

**Z** Distance (in meters)

**Force** Force (in Newtons)

**Time** Time starting at the beginning of each segment (in seconds)

**Segment** Segment of the Force-distance curve (factor: "approach", "pause", "retract")

**params** List with the following fields describing the experiment:

**SpringConstant** Cantilever spring constant (in N/m)

**curvename** Name of the original AFM data file from which the data was obtained

**type** Type of sample covering: "CHI" for Chitosan, and "PAH" for Polyallylamine hydrochloride

---

is.afmdata

*Afmdata check.*

---

**Description**

Checks wether an R object is an afmdata or not.

**Usage**

```
is.afmdata(x)
```

**Arguments**

x                    Any **R** object.

**Value**

Returns TRUE if its argument is an afmdata (that is, has "afmdata" amongst its classes) and FALSE otherwise.

---

is.afmexperiment	<i>Afmexperiment check.</i>
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---

**Description**

Checks whether an R object is an afmexperiment or not.

**Usage**

```
is.afmexperiment(x)
```

**Arguments**

x	Any <b>R</b> object.
---	----------------------

**Value**

Returns TRUE if its argument is an afmdata (that is, has "afmexperiment" amongst its classes) and FALSE otherwise.

---

plot.afmdata	<i>Plot an afmdata object</i>
--------------	-------------------------------

---

**Description**

Plots an afmdata object.

**Usage**

```
## S3 method for class 'afmdata'
plot(x, y = NULL, vs = "Z", segment = "all", ...)
```

**Arguments**

x	An object of afmdata class.
y	Variable added for compatibility with plot.
vs	The variable for the x-axis. May take the values "Time" or "Z". It defaults to "Z", plotting thus a Force-Distance curve. If vs is set to "Time", then it plots a Force-Time curve.
segment	The segment of the curve to be plotted. If segment = "all" then all segments of the curve are plotted. Possible values are: "approach", "pause", "retract" and "all".
...	Additional parameters to be passed to the ggplot functions.

**Examples**

```

# Loading the data
path <- path.package("afmToolkit")
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path)
# Standard plot (out of the box)
plot(data)
# Computing the contact and detach points
data <- afmContactPoint(data, width = 20, mul1 = 1, mul2 = 10)
data <- afmDetachPoint(data, width = 40, mul1 = 3, mul2 = 20)
# Making the baseline correction
data <- afmBaselineCorrection(data)
# Plot once the baseline correction is done
plot(data)
# Plotting only retract segment
plot(data, segment = "retract")
# Plotting the pause segment: Force vs Time
plot(data, segment = "pause", vs = "Time")

```

---

summary.afmdata

*Summary of an afmdata class object.*


---

**Description**

This function summarizes the main features of an afmdata object and, optionnaly plots all segments available with all parameters estimated.

**Usage**

```

## S3 method for class 'afmdata'
summary(object, plt = TRUE, ...)

```

**Arguments**

object	An object of afmdata class.
plt	Logical variable. If TRUE plots all available segments with all available data.
...	Additional arguments (for compatibility with summary)

**Examples**

```

## Not run: path <- path.package("afmToolkit")
data <- afmReadJPK("force-save-JPK-3h.txt.gz", path = path)
data <- afmContactPoint(data, width = 20, mul1 = 1, mul2 = 10)
data <- afmDetachPoint(data, width = 20, mul1 = 2, mul2 = 30)
data <- afmBaselineCorrection(data)
data <- afmAdhesionEnergy(data, width = 20, mul = 10)
data <- afmZeroPointSlope(data, segment = "approach")
data <- afmIndentation(data)
data <- afmYoungModulus(data, thickness = 1e-7, params = list(alpha = 22),

```

```

                                silent = TRUE)
data <- afmExpDecay(data, nexp = 2, type = "CH")
summary(data)
## End(Not run)

```

---

windowedFit

*Linear fit in a running window*


---

### Description

This is an internal function used by the `afmContactPoint` and `afmDetachPoint` functions. It computes the slopes of a linear fit to the data in a window of a given radius. This function should not be used directly unless by experienced users.

### Usage

```
windowedFit(X, width)
```

### Arguments

X	Least squares matrix on the form [1 z Force], according to input parameters in function <code>lm.fit</code>
width	Width of the window for the local regression (in vector position units)

### Value

OUT A vector of length `nrow(X)-2*width`, containing with the slopes of the fits.

### Examples

```

n <- 100
x <- seq(0,2*pi,length.out = n)
y = sin(x)+0.1*rnorm(n)
X <- matrix(c(rep(1,n),x,y),nrow = n,ncol = 3)
width <- 5
b <- windowedFit(X,width)
plot(x[(width+1):(n-width)],b,xlab = "x",ylab = "y",type = "l")
lines(x,y,col = "red")
legend("bottomleft",c("Slopes","Signal"),col = c(1,2),lty = 1)

```

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