

Package ‘fitnmr’

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Title Multidimensional Nuclear Magnetic Resonance Peak Fitting and Analysis

Description Tools for fitting and analyzing 1D-4D nuclear magnetic resonance spectra with analytical models of peak shapes and peak groups. The package reads spectra in 'NMRPipe' format, builds constrained parameter structures for chemical shifts, line widths, scalar couplings, volumes, and phases, and performs nonlinear least-squares optimization for iterative peak discovery or simultaneous fits across multiple spectra. It also provides methods for visualization, preprocessing, and kinetic analysis of 1D time-series data, including automated phase optimization, solvent suppression, time-domain correction for frequency shifts and line broadening, modeling spectra as linear combinations of two component spectra, and exponential rate fitting.

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URL <https://github.com/smith-group/fitnmr/>,
<https://smith-group.github.io/fitnmr/>

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Description

The functionality provided by the fitnmr package can be divided into several categories:

Spectrum reading, plotting, and analysis

Spectra in the NMRPipe file format can be read with [read_nmrpipe](#). Those spectra can be plotted with [contour_pipe](#). The noise level within a spectrum (or any numeric vector) can be calculated with [noise_estimate](#).

Low-level fitting and plotting

The core fitting procedure consists of a two-step process: First, the fit input is created with [make_fit_input](#). Second, the fit is executed with [perform_fit](#). Prior to running the fit, constraints can be added to the fit with either [update_fit_bounds](#) or [limit_omega0_by_r2](#).

Before or after a fit has been performed, you can extract the raw, starting, or fit spectral intensities with [get_spec_int](#). Furthermore, there are convenience plotting functions for plotting 1D ([plot_fit_1d](#)) or 2D ([plot_fit_2d](#)) fits.

The [make_fit_input](#) function takes many different parameters. To make a new fit from an existing fit, possibly with a different set of spectra or otherwise modified fitting parameters, [param_list_to_arg_list](#) can be helpful in generating a list of parameters for [make_fit_input](#).

Modifying parameter lists for fits

In fitnmr, a "parameter list" is a named list data structure that has all the information necessary to describe a set of peaks and interdependencies between the parameters for those peaks. It contains three elements: `start_list` (or also `fit_list` after a fit has been performed), `group_list`, and `comb_list`. Fitting constraints can also be stored in `lower_list` and `upper_list`. Each of those is itself a named list of arrays corresponding to different parameters, namely `omega0`, `r2`, `m0`, `p0`, `p1`, and `omega0_comb`. To help manage the values stored in those lists (particularly `omega0` and `omega0_comb`), there are several convenience functions to select particular subsets, including `omega0_param_idx` and `coupling_param_idx`. Once a subset is made, the parameters can be read or changed using `param_values`.

High-level fitting and plotting

`fit_peak_iter`
`param_list_to_peak_df`, `plot_peak_df`, `peak_df_to_param_list`, `peak_df_to_fit_input`
`fit_peak_cluster`, `fit_footprint`
`fit_peaks`, `make_param_list`
`spec_overlap_mat`

NMRPipe simulation and processing

`ppm_to_pts`, `whz_to_pts`, `write_nmrdraw_peak_tab`, `sim_time_nd`, `nmr_pipe`

Postprocessing and assignment

`height_assign`, `read_nmrdraw_peak_tab`

Deprecated

`extract_params`, `get_spec_peak_int`

Author(s)

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See Also

Useful links:

- <https://github.com/smith-group/fitnmr/>
- <https://smith-group.github.io/fitnmr/>

abind_list	<i>Combine multi-dimensional arrays with lists</i>
------------	--

Description

Combine multi-dimensional arrays with lists

Usage

```
abind_list(..., rev.along = NULL)
```

Arguments

...	any number of lists with dimensions or a single list of list arrays
rev.along	dimension to reverse before binding (passed to abind)

collapse_na	<i>Collapse strings of repeated NAs in a vector with numeric names</i>
-------------	--

Description

Collapse strings of repeated NAs in a vector with numeric names

Usage

```
collapse_na(x)
```

Arguments

x	1D array with dimnames having numeric positional values
---	---

collapse_na_array	<i>Collapse blocks of NAs in an array with numeric dimension names</i>
-------------------	--

Description

Collapse blocks of NAs in an array with numeric dimension names

Usage

```
collapse_na_array(x)
```

Arguments

x	array with names having numeric positional values
---	---

comb_vec_to_param_array

Determine array of destination parameters from a source vector

Description

Determine array of destination parameters from a source vector

Usage

```
comb_vec_to_param_array(
  comb_vec,
  comb_array,
  param_array = NULL,
  na_only = FALSE
)
```

Arguments

comb_vec	vector of source parameters
comb_array	array of 2xN data frames with mapping between vector and array
param_array	starting array of destination parameters
na_only	only overwrite values in param_array if they are NA

contour_pipe

Plot Spectra Contours

Description

Plot a two dimensional contour plot

Usage

```
contour_pipe(
  data_mat,
  nlevels = 10,
  zlim = range(data_mat, na.rm = TRUE),
  low_frac = 0.05,
  lwd = 0.25,
  main = NA,
  col_pos = "black",
  col_neg = grDevices::rgb(t(grDevices::col2rgb(col_pos))/255 * 0.25 + 0.75),
  add = FALSE,
  xlab = NULL,
  ylab = NULL,
  frame.plot = TRUE
)
```

Arguments

<code>data_mat</code>	matrix with spectral intensities.
<code>nlevels</code>	number of contour levels.
<code>zlim</code>	minimum and maximum values at which to show contours.
<code>low_frac</code>	minimum absolute value (as a fraction of <code>zlim</code>) at which to show contours.
<code>lwd</code>	width of contour lines.
<code>main</code>	title of the plot.
<code>col_pos</code>	color of positive contours.
<code>col_neg</code>	color of negative contours, defaults to lighter version of <code>col_pos</code> .
<code>add</code>	logical indicating whether to add to an existing plot (i.e. not start a new one).
<code>xlab</code>	label for x-axis, defaults to <code>names(dimnames(datamat))[1]</code> .
<code>ylab</code>	label for y-axis, defaults to <code>names(dimnames(datamat))[2]</code> .
<code>frame.plot</code>	a logical indicating whether a box should be drawn around the plot.

Details

The first dimension of `data_matrix` is drawn along the x-axis and the second dimension is drawn along the y-axis.

If `low_frac` is specified, then there can actually be up to `nlevels` total positive contour levels and/or `nlevels` total negative contour levels, whichever has the larger magnitude in `zlim`. The other dimension will have the mirror image of those up to the relevant limit in `zlim`. Note that the levels do not actually go up to `zlim`. There are `nlevels+1` log spaced levels calculated from the contour determined by `low_frac` up to the maximum absolute `zlim`, but the final level is not drawn because it is at the maximum absolute `zlim`.

If `low_frac` is set to `NA`, then the contour levels are calculated in the same way as `contour`, with there being approximately `nlevels` linearly spaced levels in a range close to `zlim`.

Note that this function does not directly take the data returned by `read_nmrpipe`. You must pass the int matrix from the value returned by that function.

Value

No return value, called for side effects (draws contour lines).

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
contour_pipe(spec$int)
```

coupling_param_idx	<i>Get a list of logical arrays indicating which parameters correspond to scalar couplings</i>
--------------------	--

Description

Get a list of logical arrays indicating which parameters correspond to scalar couplings

Usage

```
coupling_param_idx(param_list, comb_idx_offset = 0)
```

Arguments

param_list	parameter list
comb_idx_offset	offset for combination index mapping

Value

A named 'list' of logical index arrays/vectors (matching 'param_list\$group_list') that identify scalar-coupling parameters in 'omega0_comb'.

extract_params	<i>Extract parameters from fit object for use with make_fit_input</i>
----------------	---

Description

Extract parameters from fit object for use with make_fit_input

Usage

```
extract_params(fit, expand = 0)
```

Arguments

fit	fit_output structure
expand	number of empty peaks to append

fit_footprint	<i>Determine the region of a spectrum containing the majority of the fit peaks</i>
---------------	--

Description

Determine the region of a spectrum containing the majority of the fit peaks

Usage

```
fit_footprint(fit, frac = 0.12)
```

Arguments

fit	fit_output structure
frac	fraction of total intensity used to define the footprint

fit_peaks	<i>Fit peaks from a table of chemical shifts</i>
-----------	--

Description

Fit peaks from a table of chemical shifts

Usage

```
fit_peaks(  
  spec_list,  
  cs_mat,  
  fit_prev = NULL,  
  spec_ord = 1:2,  
  omega0_plus = c(0.075, 0.75),  
  r2_start = 5,  
  r2_bounds = c(0.5, 20),  
  sc_start = NULL,  
  sc_bounds = c(0, Inf),  
  positive_only = TRUE,  
  plot_fit_stages = TRUE  
)
```

Arguments

spec_list	list of spectra
cs_mat	matrix of chemical shifts
fit_prev	optional previous fit parameters
spec_ord	order of spectral dimensions
omega0_plus	omega0 bounds for each dimension
r2_start	starting r2 values
r2_bounds	numeric vector of length 2 with r2 bounds
sc_start	starting scalar coupling values
sc_bounds	numeric vector of length 2 with scalar coupling bounds
positive_only	logical indicating whether to constrain m0 to be positive
plot_fit_stages	logical indicating whether to plot intermediate fits

Value

A fitted parameter 'list' (fit-output structure) containing fitted values in 'fit_list' plus the original fit metadata and bounds.

fit_peak_cluster	<i>Fit a cluster of nearby peaks starting from a seed table of chemical shifts</i>
------------------	--

Description

Fit a cluster of nearby peaks starting from a seed table of chemical shifts

Usage

```
fit_peak_cluster(
  spec_list,
  cs_start,
  spec_ord,
  f_alpha_thresh = 0.001,
  omega0_plus = c(0.075, 0.75),
  r2_start = 5,
  r2_bounds = c(0.5, 20),
  sc_start = NULL,
  sc_bounds = c(0, Inf),
  plot_main_prefix = NULL,
  peak_num_offset = 0,
  plot_fit_stages = FALSE,
  verbose = TRUE
)
```

Arguments

spec_list	list of spectra
cs_start	matrix of starting chemical shifts
spec_ord	order of spectral dimensions
f_alpha_thresh	alpha threshold for F-test
omega0_plus	omega0 bounds for each dimension
r2_start	starting r2 values
r2_bounds	numeric vector of length 2 with r2 bounds
sc_start	starting scalar coupling values
sc_bounds	numeric vector of length 2 with scalar coupling bounds
plot_main_prefix	optional prefix for plot titles
peak_num_offset	offset added to peak numbers in plots
plot_fit_stages	logical indicating whether to plot intermediate fits
verbose	logical indicating whether to print progress updates

Value

Usually a fit-output 'list' (including 'fit_list') for the accepted peak cluster. In edge cases where no acceptable fit is retained, returns modeled intensity matrix/matrices used for subtraction.

fit_peak_iter	<i>Iterative Peak Fitting</i>
---------------	-------------------------------

Description

Iteratively fit peaks for whole spectra

Usage

```
fit_peak_iter(
  spectra,
  noise_sigma = NULL,
  noise_cutoff = 15,
  f_alpha = 0.001,
  iter_max = 100,
  omega0_plus = c(0.075, 0.75),
  r2_start = 5,
  r2_bounds = c(0.5, 20),
  sc_start = c(6, NA),
  sc_bounds = c(2, 12),
```

```

fit_list = list(),
plot_fit = FALSE,
plot_fit_stages = FALSE,
iter_callback = NULL,
verbose = TRUE
)

```

Arguments

spectra	list of spectrum objects read by read_nmrpipe .
noise_sigma	numeric vector of noise levels associated with each spectrum. If NULL, it is calculated with noise_estimate .
noise_cutoff	numeric value multiplied by noise_sigma to determine cutoffs for each spectrum. Peak fitting will terminate if the maximum residuals for all spectra fall below these cutoffs.
f_alpha	numeric value giving a F-test p-value threshold above which a peak will not be accepted.
iter_max	integer maximum number of iterations to apply.
omega0_plus	numeric vector giving the window size (ppm plus or minus the starting omega0 values) around which to use points from the spectra for fitting.
r2_start	numeric vector giving the starting r2 value(s) for the fit (in Hz).
r2_bounds	numeric vector of length two giving the lower and upper bounds for r2.
sc_start	numeric vector giving the starting scalar coupling values for doublets. It should be the same length as the number of dimensions in the spectrum. Set the value to NA for a given dimension to make it a singlet.
sc_bounds	numeric vector of length two giving the lower and upper bounds for scalar couplings.
fit_list	list of previous fits to which the new fits should be appended.
plot_fit	logical indicating whether produce a fit cluster plot for each iteration.
plot_fit_stages	logical indicating whether to plot each stage of fitting within the iterations.
iter_callback	function called after each iteration with two arguments: iter and iter_max
verbose	logical indicating whether to print progress updates

Details

This function uses an iterative algorithm to fit all the peaks in a given list of spectra, assuming identical peak positions and shapes across the spectra. Each iteration starts by identifying the maximum value across all spectra, using that position to fit a cluster of overlapping peaks with the [fit_peak_cluster](#) function. After the cluster of peaks is fit, the modeled intensity is subtracted from all spectra and another iteration is performed. Iterations are terminated if max_iter is reached or the residual intensity in all spectra falls below noise_sigma*noise_cutoff.

This function currently only supports fitting of 2D spectra, but will be generalized to work with spectra of any dimensionality in the near future. To reduce the number of false positives/negatives,

the most important parameters to adjust are `noise_cutoff`, `f_alpha`, and `iter_max`. If `iter_max` is reached before all peaks have been identified, then you can call this function again, setting the `fit_list` parameter to the return value of the previous invocation. In that case, `iter_max` new iterations will be performed and appended to `fit_list`.

For visualizing the iterative algorithm as it progresses, you can enable either the `plot_fit` or `plot_fit_stages` parameters.

Value

List of fit objects returned by `fit_peak_cluster`, one for each iteration. They are appended to `fit_list` if supplied.

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- list("1.ft2" = read_nmrpipe(spec_file, dim_order = "hx"))
peak_fits <- fit_peak_iter(spec, iter_max = 2)
```

get_spec_int	<i>Get arrays of spectral intensities for input, starting parameters, and fit peaks</i>
--------------	---

Description

Get arrays of spectral intensities for input, starting parameters, and fit peaks

Usage

```
get_spec_int(
  fit_data,
  spec_type = c("input", "start", "fit"),
  spec_idx = seq_along(fit_data$spec_data),
  peak_idx = seq_len(dim(fit_data$start_list$omega0)[2])
)
```

Arguments

<code>fit_data</code>	fit_input or fit_output structure
<code>spec_type</code>	character indicating which spectra to return
<code>spec_idx</code>	indices of spectra to return
<code>peak_idx</code>	indices of peaks to include (for start/fit)

Value

A ‘list’ of numeric arrays, one per selected spectrum. Each array is on contiguous ppm axes and contains either observed (‘input’) or modeled (‘start’/‘fit’) intensities.

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
fit_input <- make_fit_input(
  list(spec),
  omega0_start = matrix(c(8.5400, 119.76), nrow = 2),
  omega0_plus = c(0.075, 0.75),
  r2_start = 4,
  m0_start = 1e9
)
input_int <- get_spec_int(fit_input, "input")
start_int <- get_spec_int(fit_input, "start")
```

get_spec_peak_int *Get spectra for individual peaks*

Description

Get spectra for individual peaks

Usage

```
get_spec_peak_int(
  fit_data,
  spec_type = c("start", "fit"),
  spec_idx = seq_along(fit_data$spec_data),
  peak_idx = seq_len(dim(fit_data$start_list$omega0)[2])
)
```

Arguments

fit_data	fit_input or fit_output structure
spec_type	character indicating which spectra to return
spec_idx	indices of spectra to return
peak_idx	indices of peaks to include

height_assign	<i>Calculate mapping from assigned peak list onto an unknown peak list</i>
---------------	--

Description

This uses a greedy algorithm. It iterates over the unknown peaks in order of decreasing height and assigns each unknown to the closest assigned peak within a distance determined by the thresh parameter, as long as that peak wasn't already assigned.

Usage

```
height_assign(assigned, unknown, thresh = 0.01)
```

Arguments

assigned	two column matrix with assigned peak chemical shifts
unknown	three column matrix with unknown peak chemical shifts and heights
thresh	maximum distance (as a fraction of the two chemical shift ranges)

Value

An integer vector of length 'nrow(assigned)' giving matched row indices in 'unknown' (or 'NA' when unmatched). The 'thresh' attribute stores the effective distance thresholds in each dimension.

infer_acquisition_time	<i>Infer acquisition time for each dimension</i>
------------------------	--

Description

Infer acquisition time for each dimension

Usage

```
infer_acquisition_time(fheader, empirically_correct = TRUE)
```

Arguments

fheader	matrix of generalized ND parameters
empirically_correct	logical indicating whether to apply empirical correction

infer_aliasing	<i>Infer which dimension was directly acquired</i>
----------------	--

Description

Infer which dimension was directly acquired

Usage

```
infer_aliasing(fheader, phase_tolerance = 10)
```

Arguments

fheader	matrix of generalized ND parameters
phase_tolerance	tolerance in degrees for detecting half-dwell delay

infer_direct	<i>Infer which dimension was directly acquired</i>
--------------	--

Description

Infer which dimension was directly acquired

Usage

```
infer_direct(fheader)
```

Arguments

fheader	matrix of generalized ND parameters
---------	-------------------------------------

infer_sweep_width	<i>Infer original sweep width for each dimension</i>
-------------------	--

Description

Infer original sweep width for each dimension

Usage

```
infer_sweep_width(fheader)
```

Arguments

fheader	matrix of generalized ND parameters
---------	-------------------------------------

limit_omega0_by_r2 *Add upper/lower limits based on the r2 value*

Description

Add upper/lower limits based on the r2 value

Usage

```
limit_omega0_by_r2(fit_input, factor = 1.5)
```

Arguments

fit_input	fit_input structure
factor	multiplier applied to r2 to set omega0 bounds

Value

A modified 'fit_input' list with updated 'lower_list\$omega0' and 'upper_list\$omega0' bounds.

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnrm")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
fit_input <- make_fit_input(
  list(spec),
  omega0_start = matrix(c(8.5400, 119.76), nrow = 2),
  omega0_plus = c(0.075, 0.75),
  r2_start = 4,
  m0_start = 1e9
)
fit_input$start_list[1:2]
fit_input$lower_list[1]
fit_input$upper_list[1]
fit_input <- limit_omega0_by_r2(fit_input)
fit_input$lower_list[1]
fit_input$upper_list[1]
```

make_coupling_mat *Make a multiplet matrix with weights and scalar coupling coefficients*

Description

Make a multiplet matrix with weights and scalar coupling coefficients

Usage

```
make_coupling_mat(sc_names)
```

Arguments

sc_names character vector with names of scalar couplings

make_fit_input	<i>Prepare input data structure for peak fitting</i>
----------------	--

Description

Prepare input data structure for peak fitting

Usage

```
make_fit_input(  
  spectra,  
  omega0_start,  
  omega0_plus,  
  omega0_minus = omega0_plus,  
  omega0_trunc = NULL,  
  r2_start = NULL,  
  m0_start = NULL,  
  m0_region = (omega0_plus + omega0_minus)/2,  
  p0_start = 0,  
  p1_start = 0,  
  omega0_group = NULL,  
  r2_group = NULL,  
  m0_group = NULL,  
  p0_group = 0,  
  p1_group = 0,  
  omega0_comb = NULL,  
  omega0_comb_start = NULL,  
  omega0_comb_group = NULL,  
  coupling_comb = NULL,  
  resonance_names = NULL,  
  nucleus_names = NULL,  
  field_offsets = numeric(),  
  field_start = numeric(),  
  field_group = 0,  
  fheader = NULL  
)
```

Arguments

spectra	list of spectra or data frame with ppm and intensity columns
omega0_start	starting omega0 values
omega0_plus	positive omega0 bounds
omega0_minus	negative omega0 bounds
omega0_trunc	optional truncation bounds for omega0
r2_start	starting r2 values
m0_start	starting m0 values
m0_region	region for m0 estimation
p0_start	starting zero-order phase values
p1_start	starting first-order phase values
omega0_group	grouping for omega0 values
r2_group	grouping for r2 values
m0_group	grouping for m0 values
p0_group	grouping for p0 values
p1_group	grouping for p1 values
omega0_comb	list of omega0 combination matrices
omega0_comb_start	starting omega0 combination values
omega0_comb_group	grouping for omega0 combinations
coupling_comb	list of coupling combination matrices
resonance_names	optional resonance names
nucleus_names	optional nucleus names
field_offsets	matrix of field offsets
field_start	starting field values
field_group	grouping for field values
fheader	optional fheader matrix when spectra is a data frame

Value

A 'fit_input' 'list' containing 'spec_data', parameter lists ('start_list', 'group_list', 'comb_list'), parameter bounds ('lower_list', 'upper_list'), naming metadata, and 'num_points'.

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
fit_input <- make_fit_input(
  list(spec),
  omega0_start = matrix(c(8.5400, 119.76), nrow = 2),
```

```
    omega0_plus = c(0.075, 0.75),  
    r2_start = 4,  
    m0_start = 1e9  
  )  
  str(fit_input)
```

make_map	<i>Create a sparse axis map</i>
----------	---------------------------------

Description

Create a sparse axis map

Usage

```
make_map(  
  x,  
  max_spacing = 0.125,  
  new_spacing = 0.025,  
  starts = NULL,  
  ends = NULL  
)
```

Arguments

x	numeric vector or array used to define the sparse axis
max_spacing	maximum spacing between segments before inserting a gap
new_spacing	spacing used for the inserted gap
starts	optional numeric vector of segment start positions
ends	optional numeric vector of segment end positions

Value

A two-column numeric matrix mapping original coordinates (column 1) to sparse-axis coordinates (column 2), with 'starts' and 'ends' attributes giving segment boundaries.

make_param_list	<i>Make a parameter list for a set of spectra and chemical shifts</i>
-----------------	---

Description

Make a parameter list for a set of spectra and chemical shifts

Usage

```
make_param_list(  
  spec_list,  
  cs_mat,  
  fit_prev = NULL,  
  r2_start = 5,  
  m0_start = 1,  
  sc_start = NULL,  
  same_r2 = FALSE,  
  same_coupling = FALSE  
)
```

Arguments

spec_list	list of spectra
cs_mat	matrix of chemical shifts
fit_prev	optional previous fit parameters
r2_start	starting r2 values
m0_start	starting m0 values
sc_start	starting scalar coupling values
same_r2	logical indicating whether to tie r2 across dimensions
same_coupling	logical indicating whether to tie coupling across dimensions

Value

A parameter 'list' with 'start_list', 'group_list', and 'comb_list' components that can be passed to 'make_fit_input'.

nmrpipe_ft	<i>Fourier transform a 1D FID</i>
------------	-----------------------------------

Description

This function aims to numerically match the NMRPipe 'FT' function.

Usage

```
nmrpipe_ft(fid)
```

Arguments

fid list with 'int', 'header', and 'fheader' elements containing FID data

Details

See the NMRPipe documentation: <https://www.nmrscience.com/ref/nmrpipe/ft.html>

Value

A transformed spectrum 'list' with updated 'int', 'ppm', 'header', and 'fheader' fields.

Examples

```
fid_path <- system.file("extdata", "noesy1d", "11", "test.fid", package = "fitnmr")
fid <- read_nmrpipe(fid_path, complex_data = TRUE)
sp <- nmrpipe_sp(fid)
zf <- nmrpipe_zf(sp)
ft <- nmrpipe_ft(zf)
```

nmrpipe_fti	<i>Inverse Fourier transform a 1D spectrum</i>
-------------	--

Description

This function aims to numerically match the NMRPipe 'FT -inv' function.

Usage

```
nmrpipe_fti(ft)
```

Arguments

ft list with 'int', 'header', and 'fheader' elements containing spectrum data

Details

See the NMRPipe documentation: <https://www.nmrscience.com/ref/nmrpipe/ft.html>

Value

A modified FID-like 'list' with inverse-transformed 'int' values and updated 'header'/'fheader' metadata.

Examples

```
fid_path <- system.file("extdata", "noesy1d", "11", "test.fid", package = "fitnmr")
fid <- read_nmrpipe(fid_path, complex_data = TRUE)
sp <- nmrpipe_sp(fid)
zf <- nmrpipe_zf(sp)
ft <- nmrpipe_ft(zf)
fti <- nmrpipe_fti(ft)
```

nmrpipe_ps

Inverse Fourier transform a 1D spectrum

Description

This function aims to numerically match the NMRPipe 'PS' function.

Usage

```
nmrpipe_ps(ft, p0 = 0, p1 = 0)
```

Arguments

ft	list with 'int', 'header', and 'fheader' elements containing spectrum data
p0	equivalent to '-p0' flag
p1	equivalent to '-p1' flag

Details

See the NMRPipe documentation: <https://www.nmrscience.com/ref/nmrpipe/ps.html>

Value

A modified spectrum 'list' with phase-corrected 'int' values and updated phase metadata in 'header' and 'fheader'.

Examples

```
fid_path <- system.file("extdata", "noesy1d", "11", "test.fid", package = "fitnmr")
fid <- read_nmrpipe(fid_path, complex_data = TRUE)
sp <- nmrpipe_sp(fid)
zf <- nmrpipe_zf(sp)
ft <- nmrpipe_ft(zf)
ps <- nmrpipe_ps(ft, p0 = 129, p1 = 3)
```

nmrpipe_sp

Apply sine-based window function to a 1D FID

Description

This function aims to numerically match the NMRPipe ‘SP’ function.

Usage

```
nmrpipe_sp(fid, off = 0, end = 1, pow = 1, cval = 1, dmx = FALSE)
```

Arguments

fid	list with ‘int’, ‘header’, and ‘fheader’ elements containing FID data
off	equivalent to ‘-off’ flag
end	equivalent to ‘-end’ flag
pow	equivalent to ‘-pow’ flag
cval	equivalent to ‘-c’ flag
dmx	equivalent to ‘-dmx’ flag

Details

See the NMRPipe documentation: <https://www.nmrscience.com/ref/nmrpipe/sp.html>

Value

A modified FID ‘list’ with updated ‘int’, ‘header’, and ‘fheader’ values after sine-bell apodization.

Examples

```
fid_path <- system.file("extdata", "noesy1d", "11", "test.fid", package = "fitnmr")
fid <- read_nmrpipe(fid_path, complex_data = TRUE)
sp <- nmrpipe_sp(fid)
```

`nmrpipe_zf`*Apply zero filling to a 1D FID*

Description

This function aims to numerically match the NMRPipe ‘ZF’ function.

Usage

```
nmrpipe_zf(fid, zf = 1, pad = NULL, size = NULL, auto = FALSE)
```

Arguments

<code>fid</code>	list with ‘int’, ‘header’, and ‘fheader’ elements containing FID data
<code>zf</code>	equivalent to ‘-zf’ flag
<code>pad</code>	equivalent to ‘-pad’ flag
<code>size</code>	equivalent to ‘-size’ flag
<code>auto</code>	equivalent to ‘-auto’ flag

Details

See the NMRPipe documentation: <https://www.nmrscience.com/ref/nmrpipe/zf.html>

Value

A modified FID ‘list’ with zero-filled ‘int’ data and updated ‘header’/‘fheader’ metadata.

Examples

```
fid_path <- system.file("extdata", "noesy1d", "11", "test.fid", package = "fitnmr")
fid <- read_nmrpipe(fid_path, complex_data = TRUE)
sp <- nmrpipe_sp(fid)
zf <- nmrpipe_zf(sp)
```

`nmr_pipe`*Process an FID with NMRPipe*

Description

Process an FID with NMRPipe

Usage

```
nmr_pipe(
  in_path,
  out_path,
  ndim = 1,
  apod = NULL,
  sp = rbind(off = 0.5, end = 1, pow = 1, c = 0.5),
  zf = rbind(auto = ""),
  ps = rbind(p0 = 0, p1 = 0, di = ""),
  ext = NULL
)
```

Arguments

in_path	input file path
out_path	output file path
ndim	number of dimensions
apod	optional apodization argument matrix
sp	optional sine-bell argument matrix
zf	optional zero-fill argument matrix
ps	optional phase correction argument matrix
ext	optional extraction argument matrix

Value

The integer exit status from 'system(...)' (typically '0' on success).

noise_estimate	<i>Estimate Noise</i>
----------------	-----------------------

Description

Estimate properties of noise by fitting a Gaussian to a histogram of intensities

Usage

```
noise_estimate(
  x,
  height = TRUE,
  thresh = 10,
  plot_distributions = TRUE,
  peak_intensities = NULL
)
```

Arguments

<code>x</code>	numeric values for which to estimate noise.
<code>height</code>	logical indicating whether to use Gaussian function whose area is not fixed at one because of an additional height scaling factor.
<code>thresh</code>	numeric value specifying the factor by which to multiply the standard deviation to determine the threshold away from the mean value within which to include values for the next iteration and final histogram for fitting.
<code>plot_distributions</code>	logical indicating whether to plot the distribution and Gaussian fit used to estimate the noise.
<code>peak_intensities</code>	numeric values of peak intensities to determine the signal to noise.

Details

This function estimates noise using iterative calculation of the mean and standard deviation, followed by fitting a Gaussian function to a histogram of the values within a range determined at the end of the iterations.

The iterative algorithm first calculates the mean and standard deviation of the values in `x`. In the next iteration, only points within `thresh` times the standard deviation of the mean value are used for calculating a new mean and standard deviation. This is repeated 20 times. A histogram with 512 bins is then computed within a range determined from the final mean and standard deviation. The returned parameters are based on a Gaussian fit to this histogram.

Value

a named numeric vector with values:

mean mean value from the Gaussian fit

mu standard deviation from the Gaussian fit

h height of Gaussian fit (optional depending on value of `height` parameter)

max maximum value of `x`

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package="fitnmr")
spec <- read_nmrpipe(spec_file, dim_order="hx")
noise_estimate(spec$int)
```

omega0_comb_source_idx

Get the first index in the omega0 array corresponding to each TRUE value in omega0_idx

Description

Get the first index in the omega0 array corresponding to each TRUE value in omega0_idx

Usage

```
omega0_comb_source_idx(param_list, omega0_idx = omega0_param_idx(param_list))
```

Arguments

param_list	parameter list
omega0_idx	logical index list for omega0 parameters

omega0_param_idx

Get a list of logical arrays indicating which parameters correspond to peak positions

Description

Get a list of logical arrays indicating which parameters correspond to peak positions

Usage

```
omega0_param_idx(
  param_list,
  dims = seq_len(dim(param_list[["group_list"]][["omega0"]])[1]),
  peaks = seq_len(dim(param_list[["group_list"]][["omega0"]])[2]),
  specs = seq_len(dim(param_list[["group_list"]][["omega0"]])[3])
)
```

Arguments

param_list	parameter list
dims	dimensions to include
peaks	peaks to include
specs	spectra to include

Value

A named 'list' of logical index arrays/vectors (matching 'param_list\$group_list') that identify omega0-related parameters to extract or modify.

 param_array_to_comb_vec

Determine vector of source parameters from destination array via least squares

Description

Determine vector of source parameters from destination array via least squares

Usage

```
param_array_to_comb_vec(
  param_array,
  comb_array,
  group_vec = NULL,
  resid_thresh = 1e-08
)
```

Arguments

param_array	array of destination parameters
comb_array	array of 2xN data frames with mapping between vector and array
group_vec	optional vector of group numbers for source parameters
resid_thresh	all residuals from least squares fit must be less than this value

 param_list_to_arg_list

Convert a list of parameters for use with make_fit_input

Description

Convert a list of parameters for use with make_fit_input

Usage

```
param_list_to_arg_list(param_list)
```

Arguments

param_list	parameter list
------------	----------------

Value

A flattened, named 'list' of arguments suitable for 'do.call()' with 'make_fit_input()'.

param_list_to_peak_df *Convert Fit to Data Frame*

Description

Convert a parameter list into a peak data frame

Usage

```
param_list_to_peak_df(param_list, spec_names = NULL)
```

Arguments

param_list	list of fit parameters (or a list of such lists)
spec_names	character vector of spectrum names

Details

This function takes the input or (if present) output parameters from a fit and converts them into a data frame. A parameter list must contain three lists:

start_list **or** fit_list input or output values of the respective fit parameters

group_list group numbers for the fit parameters

comb_list coefficients for deriving fit parameters from a linear combination other auxiliary parameters

This function currently assumes the fit parameters were generated by [fit_peak_iter](#), [fit_peak_cluster](#), or [peak_df_to_param_list](#). These functions use a particular convention for group_list and comb_list to represent either singlets or doublets in each dimension of a 2D spectrum.

This function can take either a single parameter list or a list of parameter lists. If the latter is given, then the results from all the parameter lists will be combined into a single table.

Value

A data frame with the following columns:

peak peak number

fit fit cluster number, with all peaks in the same cluster having the same r2, scalar couplings, and m0

f_pvalue optional, p-value determined from F-test during iterative fitting

omega0_ppm_1 chemical shift of singlet/doublet center in the first dimension (ppm)

omega0_ppm_2 chemical shift of singlet/doublet center in the second dimension (ppm)

sc_hz_1 optional, scalar coupling of doublet in first dimension (Hz)

sc_hz_2 optional, scalar coupling of doublet in second dimension (Hz)

r2_hz_1 R2 in first dimension (Hz)

r2_hz_2 R2 in first dimension (Hz)

... m0 values for each spectrum

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- list("1.ft2" = read_nmrpipe(spec_file, dim_order = "hx"))
peak_fits <- fit_peak_iter(spec, iter_max = 2)
param_list_to_peak_df(peak_fits)
```

param_list_to_tables *Convert a parameter list into a set of tables with resonance/nuclei/couplings*

Description

Convert a parameter list into a set of tables with resonance/nuclei/couplings

Usage

```
param_list_to_tables(param_list, tables)
```

Arguments

param_list	param_list, fit_input, or fit_output structure
tables	list with resonance/nuclei/couplings tables to update

Value

A named 'list' with updated 'resonances', 'nuclei', and 'couplings' data frames.

Examples

```
spec_file <- system.file("extdata", "tyrosine", "proton.ft1", package = "fitnmr")
spec <- read_nmrpipe(spec_file)
```

```
start_resonances <- structure(
  list(
    x = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
    x_sc = c(
      "HA-HB3 HA-HB2",
      "HA-HB3 HB3-HB2",
      "HA-HB2 HB3-HB2",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5"
    )
  ),
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
  class = "data.frame"
)

start_nuclei <- structure(
```

```

list(
  omega0_ppm = c(3.364, 2.635, 2.799, 6.94, 6.538),
  r2_hz = c(0.7, 0.7, 0.7, 0.7, 0.7)
),
class = "data.frame",
row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2")
)

start_couplings <- structure(
  list(hz = c(7.153, 5.159, -13.941, 7.7, 2)),
  class = "data.frame",
  row.names = c("HA-HB3", "HA-HB2", "HB3-HB2", "HD1/2-HE1/2-3", "HD1/2-HE1/2-5")
)

start_tables <- list(
  resonances = start_resonances,
  nuclei = start_nuclei,
  couplings = start_couplings
)

param_list <- tables_to_param_list(list(spec), start_tables)
param_list$start_list$m0[] <- 1e9
arg_list <- param_list_to_arg_list(param_list)
fit_input <- do.call(make_fit_input, c(list(list(spec), omega0_plus = 0.075), arg_list))
fit_output <- perform_fit(fit_input)
fit_tables <- param_list_to_tables(fit_output, start_tables)

```

param_values	<i>Get/set a subset of fitting parameters specified by a list of logical vectors</i>
--------------	--

Description

Get/set a subset of fitting parameters specified by a list of logical vectors

Usage

```

param_values(params, idx_list)

param_values(params, idx_list) <- value

```

Arguments

params	a list of fitting parameters
idx_list	a list with the same structure as params but with logical vectors indicating which values should be return/set
value	replacement values (for setter)

Value

a vector of parameter values or the modified parameter list

peak_bind	<i>Combine parameter lists referring to different peaks</i>
-----------	---

Description

Combine parameter lists referring to different peaks

Usage

```
peak_bind(...)
```

Arguments

... any number of parameter lists or a single list of parameter lists

peak_df_to_fit_input	<i>Convert a peak data frame to fit input</i>
----------------------	---

Description

Convert a peak data frame to fit input

Usage

```
peak_df_to_fit_input(peak_df, spectra, ...)
```

Arguments

peak_df	data frame of peak parameters
spectra	list of spectra
...	additional arguments passed to make_fit_input

Value

A 'fit_input' list ready for optimization, as produced by 'make_fit_input'.

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spectra <- list("1.ft2" = read_nmrpipe(spec_file, dim_order = "hx"))

peak_df <- data.frame(
  peak = c(1, 2, 3, 4),
  fit = c(1, 1, 2, 2),
  f_pvalue = c(4.566421e-10, 1.118991e-05, 1.876528e-15, 5.817124e-04),
  omega0_ppm_1 = c(8.247602, 8.259565, 8.540030, 8.520232),
  omega0_ppm_2 = c(121.8666, 121.9299, 119.7611, 119.7266),
  sc_hz_1 = c(3.280589, 3.280589, 2.000000, 2.000000),
  r2_hz_1 = c(2.907218, 2.907218, 4.788566, 4.788566),
  r2_hz_2 = c(2.334497, 2.334497, 2.099646, 2.099646),
  `1.ft2` = c(824420657, 240560662, 1020008726, 89977216),
  check.names = FALSE
)

fit_input <- peak_df_to_fit_input(peak_df, spectra, omega0_plus=c(0.075, 0.75))
```

peak_df_to_param_list *Convert a peak data frame to a parameter list*

Description

Convert a peak data frame to a parameter list

Usage

```
peak_df_to_param_list(peak_df, spectra)
```

Arguments

peak_df	data frame of peak parameters
spectra	list of spectra

Value

A parameter 'list' with 'start_list', 'group_list', and 'comb_list' inferred from 'peak_df' and aligned to 'spectra'.

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spectra <- list("1.ft2" = read_nmrpipe(spec_file, dim_order = "hx"))

peak_df <- data.frame(
  peak = c(1, 2, 3, 4),
  fit = c(1, 1, 2, 2),
```

```

f_pvalue = c(4.566421e-10, 1.118991e-05, 1.876528e-15, 5.817124e-04),
omega0_ppm_1 = c(8.247602, 8.259565, 8.540030, 8.520232),
omega0_ppm_2 = c(121.8666, 121.9299, 119.7611, 119.7266),
sc_hz_1 = c(3.280589, 3.280589, 2.000000, 2.000000),
r2_hz_1 = c(2.907218, 2.907218, 4.788566, 4.788566),
r2_hz_2 = c(2.334497, 2.334497, 2.099646, 2.099646),
`1.ft2` = c(824420657, 240560662, 1020008726, 89977216),
check.names = FALSE
)

peak_df_to_param_list(peak_df, spectra)

```

perform_fit

*Perform a fit with an input data structure***Description**

Perform a fit with an input data structure

Usage

```

perform_fit(
  fit_input,
  method = c("minpack.lm", "gslnl", "sparseLM", "L-BFGS-B"),
  ...
)

```

Arguments

fit_input	fit_input structure
method	fitting method
...	additional arguments passed to the fitting routine

Value

The input 'fit_input' list, augmented with fitted parameters in 'fit_list' and fit diagnostics in 'fit_rsstrace', 'fit_counts', and 'fit_time'.

Examples

```

spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
fit_input <- make_fit_input(
  list(spec),
  omega0_start = matrix(c(8.5400, 119.76), nrow = 2),
  omega0_plus = c(0.075, 0.75),
  r2_start = 4,
  m0_start = 1e9
)

```

```

)
fit_output <- perform_fit(fit_input)
fit_output$start_list[1:3]
fit_output$fit_list[1:3]

```

plot_fit_1d

Plot a one dimensional peak fit

Description

Plot a one dimensional peak fit

Usage

```
plot_fit_1d(fit_data, always_show_start = FALSE)
```

Arguments

```
fit_data          fit_input or fit_output structure
always_show_start logical indicating whether to show starting model when fit is present
```

Value

No return value, called for side effects (draws plots).

Examples

```
spec_file <- system.file("extdata", "tyrosine", "proton.ft1", package = "fitnmr")
spec <- read_nmrpipe(spec_file)
```

```
start_resonances <- structure(
  list(
    x = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
    x_sc = c(
      "HA-HB3 HA-HB2",
      "HA-HB3 HB3-HB2",
      "HA-HB2 HB3-HB2",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5"
    )
  ),
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
  class = "data.frame"
)
```

```
start_nuclei <- structure(
  list(
```

```

    omega0_ppm = c(3.364, 2.635, 2.799, 6.94, 6.538),
    r2_hz = c(0.7, 0.7, 0.7, 0.7, 0.7)
  ),
  class = "data.frame",
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2")
)

start_couplings <- structure(
  list(hz = c(7.153, 5.159, -13.941, 7.7, 2)),
  class = "data.frame",
  row.names = c("HA-HB3", "HA-HB2", "HB3-HB2", "HD1/2-HE1/2-3", "HD1/2-HE1/2-5")
)

start_tables <- list(
  resonances = start_resonances,
  nuclei = start_nuclei,
  couplings = start_couplings
)
param_list <- tables_to_param_list(list(spec), start_tables)
param_list$start_list$omega0[] <- 1e9
arg_list <- param_list_to_arg_list(param_list)
fit_input <- do.call(make_fit_input, c(list(list(spec), omega0_plus = 0.075), arg_list))
fit_output <- perform_fit(fit_input)
plot_fit_1d(fit_output)

```

plot_fit_2d

Plot a two dimensional peak fit

Description

Plot a two dimensional peak fit

Usage

```

plot_fit_2d(
  fit_output,
  spec_ord = seq_len(dim(fit_output$start_list$omega0)[1]),
  always_show_start = FALSE,
  main = NULL
)

```

Arguments

fit_output	fit_output structure
spec_ord	order of spectral dimensions to plot
always_show_start	logical indicating whether to show starting model when fit is present
main	optional title for the plot

Value

No return value, called for side effects (draws plots).

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
fit_input <- make_fit_input(
  list(spec),
  omega0_start = matrix(c(8.5400, 119.76), nrow = 2),
  omega0_plus = c(0.075, 0.75),
  r2_start = 4,
  m0_start = 1e9
)
plot_fit_2d(fit_input)
```

plot_peak_df

Plot Peaks from a Peak Table

Description

Plot fits for series of spectra with parameters from a peak data frame

Usage

```
plot_peak_df(
  peak_df,
  spectra,
  noise_sigma = NULL,
  noise_cutoff = 4,
  omega0_plus = c(0.075, 0.75) * 2,
  cex = 0.2,
  lwd = 0.25,
  label = TRUE,
  label_col = "black",
  p0 = NULL,
  p1 = NULL,
  add = FALSE
)
```

Arguments

peak_df	data frame with peak data as produced by param_list_to_peak_df .
spectra	list of spectra corresponding to the volumes found in peak_df.
noise_sigma	numeric vector of noise levels associated with each spectrum. If NULL, it is calculated with noise_estimate .

noise_cutoff	numeric value used to calculate the lowest contour level according to noise_sigma*noise_cutoff.
omega0_plus	omega0 bounds for plotting
cex	numeric value by which to scale blue points and labels.
lwd	numeric value giving width of contour lines.
label	logical indicating whether to draw text labels and connecting lines.
label_col	color for text labels
p0	zero order phase for plotting modeled peaks.
p1	first order phase for plotting modeled peaks.
add	logical indicating whether to suppress generation of a new plot and add to an existing plot.

Details

The raw spectral data is shown in black contours and the modeled peak intensity is shown in red. The centers of peaks are shown with semi-transparent blue dots, with the area of the dot proportional to the volume of the peak (m_0). Blue lines connect peaks from modeled doublets. Singlets or doublets are labeled with the syntax <peak>:<fit>. If an F-test p-value column is present (f_pvalue), that will be given below the peak label.

Value

No return value, called for side effects (draws plots).

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")

peak_df <- data.frame(
  peak = 1:8,
  fit = c(1, 1, 2, 2, 3, 3, 3, 3),
  f_pvalue = c(4.5664e-10, 1.1190e-05, 1.8765e-15, 5.8171e-04,
              7.5270e-13, 2.5923e-27, 1.3424e-05, 1.4680e-13),
  omega0_ppm_1 = c(8.2476, 8.2596, 8.5400, 8.5202, 8.6124, 8.5853, 8.6449, 8.5370),
  omega0_ppm_2 = c(121.87, 121.93, 119.76, 119.73, 123.36, 123.03, 123.47, 122.96),
  sc_hz_1 = c(3.2806, 3.2806, 2.0000, 2.0000, 7.6481, 7.6481, 7.6481, 7.6481),
  r2_hz_1 = c(2.9072, 2.9072, 4.7886, 4.7886, 5.2849, 5.2849, 5.2849, 5.2849),
  r2_hz_2 = c(2.3345, 2.3345, 2.0996, 2.0996, 2.1801, 2.1801, 2.1801, 2.1801),
  `1.ft2` = c(824420657, 240560662, 1020008726, 89977216,
             848579189, 607904936, 147984411, 161971930),
  check.names = FALSE
)

plot_peak_df(peak_df, list("1.ft2" = spec), cex = 0.6)
```

plot_resonances_1d *Plot resonances from 1D fit*

Description

Plot resonances from 1D fit

Usage

```
plot_resonances_1d(fit_data, always_show_start = FALSE, omega0_plus = 0.05)
```

Arguments

fit_data fit_input or fit_output structure
 always_show_start show start parameters even if fit already done
 omega0_plus length 3 vector giving ppm range for each dimension

Value

No return value, called for side effects (draws one or more plots).

Examples

```
spec_file <- system.file("extdata", "tyrosine", "proton.ft1", package = "fitnmr")
spec <- read_nmrpipe(spec_file)
```

```
start_resonances <- structure(
  list(
    x = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
    x_sc = c(
      "HA-HB3 HA-HB2",
      "HA-HB3 HB3-HB2",
      "HA-HB2 HB3-HB2",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5"
    )
  ),
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
  class = "data.frame"
)
```

```
start_nuclei <- structure(
  list(
    omega0_ppm = c(3.364, 2.635, 2.799, 6.94, 6.538),
    r2_hz = c(0.7, 0.7, 0.7, 0.7, 0.7)
  ),
  class = "data.frame",
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2")
)
```

```

)

start_couplings <- structure(
  list(hz = c(7.153, 5.159, -13.941, 7.7, 2)),
  class = "data.frame",
  row.names = c("HA-HB3", "HA-HB2", "HB3-HB2", "HD1/2-HE1/2-3", "HD1/2-HE1/2-5")
)

start_tables <- list(
  resonances = start_resonances,
  nuclei = start_nuclei,
  couplings = start_couplings
)

param_list <- tables_to_param_list(list(spec), start_tables)
param_list$start_list$m0[] <- 1e9
arg_list <- param_list_to_arg_list(param_list)
fit_input <- do.call(make_fit_input, c(list(list(spec), omega0_plus = 0.075), arg_list))
fit_output <- perform_fit(fit_input)
plot_resonances_1d(fit_output, always_show_start = FALSE, omega0_plus = 0.075)

```

plot_resonances_2d *Plot resonances from 2D fit*

Description

Plot resonances from 2D fit

Usage

```

plot_resonances_2d(
  fit_data,
  omega0_plus,
  resonances = unique(fit_data$resonance_names),
  low_frac = 0.05,
  field = TRUE,
  proj_frac = 0.2
)

```

Arguments

fit_data	fit_input or fit_output structure
omega0_plus	length 2 vector giving ppm range for each dimension
resonances	character vector with resonances to plot
low_frac	minimum absolute value (as a fraction of maximum intensity) at which to show contours
field	logical indicating whether to show modeling of field inhomogeneity as separate peaks
proj_frac	fraction of plot area reserved for 1D projections

Value

No return value, called for side effects (draws one plot layout per resonance).

plot_resonances_3d *Plot resonances from 3D fit*

Description

Plot resonances from 3D fit

Usage

```
plot_resonances_3d(
  fit_data,
  omega0_plus,
  resonances = unique(fit_data$resonance_names)
)
```

Arguments

fit_data	fit_input or fit_output structure
omega0_plus	length 3 vector giving ppm range for each dimension
resonances	character vector with resonances to plot

Value

No return value, called for side effects (draws one set of projections per resonance).

plot_sparse_1d *Plot spectrum from 1D fit*

Description

Plot spectrum from 1D fit

Usage

```
plot_sparse_1d(
  fit_data,
  tables = NULL,
  spec_idx = 1,
  col_model = 2,
  col_resonance = NULL,
  lwd = 1,
  tick_spacing = 0.02,
```

```

coupling_spacing = 0.01,
coupling_marks = 0.009,
xaxs = "i",
yaxt = "n",
bty = "n",
always_show_start = FALSE,
add = FALSE,
ppm_map = make_map(get_spec_int(fit_data, "input", spec_idx)[[1]])
)

```

Arguments

fit_data	fit_input or fit_output structure
tables	list with resonances, nuclei, and couplings tables
spec_idx	index of spectrum to plot
col_model	color for modeled peak shapes
col_resonance	colors for resonances
lwd	line width
tick_spacing	spacing between axis ticks (ppm)
coupling_spacing	spacing between coupling annotations (fraction of plot)
coupling_marks	size of coupling tick marks (fraction of plot)
xaxs	x-axis style passed to plot
yaxt	y-axis style passed to plot
bty	box type passed to plot
always_show_start	logical indicating whether to show starting model when fit is present
add	logical indicating whether to add to existing plot
ppm_map	sparse axis map created by make_map

Value

No return value, called for side effects (draws a plot).

Examples

```

spec_file <- system.file("extdata", "tyrosine", "proton.ft1", package = "fitnmr")
spec <- read_nmrpipe(spec_file)

start_resonances <- structure(
  list(
    x = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
    x_sc = c(
      "HA-HB3 HA-HB2",
      "HA-HB3 HB3-HB2",
      "HA-HB2 HB3-HB2",
    )
  )
)

```

```

      "HD1/2-HE1/2-3 HD1/2-HE1/2-5",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5"
    )
  ),
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
  class = "data.frame"
)

start_nuclei <- structure(
  list(
    omega0_ppm = c(3.364, 2.635, 2.799, 6.94, 6.538),
    r2_hz = c(0.7, 0.7, 0.7, 0.7, 0.7)
  ),
  class = "data.frame",
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2")
)

start_couplings <- structure(
  list(hz = c(7.153, 5.159, -13.941, 7.7, 2)),
  class = "data.frame",
  row.names = c("HA-HB3", "HA-HB2", "HB3-HB2", "HD1/2-HE1/2-3", "HD1/2-HE1/2-5")
)

start_tables <- list(
  resonances = start_resonances,
  nuclei = start_nuclei,
  couplings = start_couplings
)

param_list <- tables_to_param_list(list(spec), start_tables)
param_list$start_list$m0[] <- 1e9
arg_list <- param_list_to_arg_list(param_list)
fit_input <- do.call(make_fit_input, c(list(list(spec), omega0_plus = 0.075), arg_list))
fit_output <- perform_fit(fit_input)
plot_sparse_1d(fit_output, start_tables)

```

plot_sparse_2d

Plot spectrum from 2D fit

Description

Plot spectrum from 2D fit

Usage

```

plot_sparse_2d(
  fit_data,
  tables = NULL,
  spec_idx = 1,
  spec_int = NULL,

```

```

    col_model = 2,
    col_nucleus = NULL,
    lwd = 1,
    tick_spacing = 0.02,
    coupling_spacing = 0.01,
    coupling_marks = 0.009,
    xaxs = "i",
    yaxs = "i",
    low_frac = 0.05,
    bty = "n",
    always_show_start = FALSE,
    add = FALSE,
    ppm_map_list = NULL
)

```

Arguments

fit_data	fit_input or fit_output structure
tables	list with resonances, nuclei, and couplings tables
spec_idx	index of spectrum to plot
spec_int	optional matrix to replace the input spectrum intensities
col_model	color for modeled peak shapes
col_nucleus	colors for nuclei
lwd	line width
tick_spacing	spacing between axis ticks (ppm)
coupling_spacing	spacing between coupling annotations (fraction of plot)
coupling_marks	size of coupling tick marks (fraction of plot)
xaxs	x-axis style passed to plot
yaxs	y-axis style passed to plot
low_frac	minimum absolute value (as a fraction of maximum intensity) at which to show contours
bty	box type passed to plot
always_show_start	logical indicating whether to show starting model when fit is present
add	logical indicating whether to add to existing plot
ppm_map_list	optional list of sparse axis maps created by make_map

Value

No return value, called for side effects (draws a plot).

ppm_to_pts	<i>Convert PPM values to points</i>
------------	-------------------------------------

Description

Convert PPM values to points

Usage

```
ppm_to_pts(ppm_mat, fheader)
```

Arguments

ppm_mat	matrix of ppm values
fheader	matrix of generalized ND parameters

Value

A numeric matrix of point indices with the same dimensions as 'ppm_mat', with column names converted from '*_PPM' to '*_AXIS'.

read_nmrdraw_peak_tab	<i>Read an NMRDraw formatted peak table</i>
-----------------------	---

Description

Read an NMRDraw formatted peak table

Usage

```
read_nmrdraw_peak_tab(file_path)
```

Arguments

file_path	path to the NMRDraw peak table file
-----------	-------------------------------------

Value

A 'data.frame' containing the peak table, with columns defined by the 'VARS' line in the NMR-Draw file.

read_nmrpipe	<i>Read NMRPipe spectrum</i>
--------------	------------------------------

Description

This function reads 1D-4D spectra stored in the NMRPipe format.

Usage

```
read_nmrpipe(inFormat, dim_order = NULL, complex_data = FALSE)
```

Arguments

<code>inFormat</code>	character with file name or format for multiple files
<code>dim_order</code>	integer vector used to reorder dimensions or character specifying
<code>complex_data</code>	logical value indicating whether complex data should be read

Details

For three and four dimensional datasets, the spectral data is often spread across multiple files. To read those, `inFormat` should be a `sprintf`-style string that describes how the files are named. For instance, if the files are named 001.ft3, 002.ft3, etc., then `inFormat` should be `"%03i.ft3"`. If there is no zero-padding, as in this case, 0 should be omitted from the format. If there are fewer digits, then the first 3 should be changed accordingly.

The default 2D NMRPipe scripts only have a single transpose ("TP") command, leaving the the indirect dimension as the first dimension in the resulting array. The 2D plotting functions in `fitnmr` usually plot this first dimension along the x-axis, which will make for generally non-standard contour plots. Furthermore, when peak fitting is employed, this will also be the first dimension. To fix this, you can change the spectral order with the `dim_order` parameter. The order of the dimensions should be specified in the same way that would be done for `aperm`. Alternatively, you can specify a character argument to have `fitnmr` attempt to automatically detect and correct the array order. The only currently supported type is "hx", which will put the dimension with the greatest observe frequency first.

This function is partly based on the `pipe2nmr` function from `rNMR`.

Value

a named list with four elements:

int multidimensional array with spectrum intensities (ppm values are given in the `dimnames`)

ppm list of numeric vectors giving the ppm values associated with each `int` array dimension

fheader matrix with dimension-specific header information

header numeric vector with the complete header contents

The fheader row definitions are as follows (taken from NMRPipe fdatp.h):

Row Name	Description
SIZE	Number of points in dimension
APOD	Current valid time-domain size
SW	Sweep Width, Hz
ORIG	Axis Origin (Last Point), Hz
OBS	Obs Freq, MHz
FTFLAG	1=Freq Domain 0=Time Domain
QUADFLAG	Data Type Code (See Below)
UNITS	Axis Units Code (See Below)
P0	Zero Order Phase, Degrees
P1	First Order Phase, Degrees
CAR	Carrier Position, PPM
CENTER	Point Location of Zero Freq
AQSIGN	Sign adjustment needed for FT
APODCODE	Window function used
APODQ1	Window parameter 1
APODQ2	Window parameter 2
APODQ3	Window parameter 3
C1	Add 1.0 to get First Point Scale
ZF	Negative of Zero Fill Size
X1	Extract region origin, if any, pts
XN	Extract region endpoint, if any, pts
OFFPPM	Additional PPM offset (for alignment)
FTSIZE	Size of data when FT performed
TDSIZE	Original valid time-domain size
LB	Extra Exponential Broadening, Hz
GB	Extra Gaussian Broadening, Hz
GOFF	Offset for Gaussian Broadening, 0 to 1
OBSMID	Original Obs Freq before 0.0ppm adjust

In addition several rows contain information inferred from the header data:

Row Name	Description
aq_s	Acquisition time, seconds
sw_ppm	Original Sweep Width, PPM
direct	Direct (1) or indirect (0) dimension
alias	Aliasing (0/1) with inversion (-1)
mag	Magnitude mode (direct from FDMCFLAG)

Examples

```
spec_file <- system.file("extdata", "t1", "1.ft2", package="fitnmr")
spec <- read_nmrpipe(spec_file, dim_order="hx")
str(spec)
```

 resonance_to_param_list

Convert data frame of resonances into a parameter list

Description

Convert data frame of resonances into a parameter list

Usage

```
resonance_to_param_list(spec, resonance, nuclei, couplings)
```

Arguments

spec	single spectrum
resonance	data frame with resonances
nuclei	data frame with nuclei chemical shifts and R2 rates
couplings	data frame with scalar couplings

 sim_time_nd

Simulate an FID using the NMRPipe SimTimeND function

Description

Simulate an FID using the NMRPipe SimTimeND function

Usage

```
sim_time_nd(
  peak_tab,
  fheader,
  rms = 0,
  iseed = stats::runif(1, max = .Machine$integer.max),
  file_path = NULL,
  verbose = FALSE
)
```

Arguments

peak_tab	NMRDraw peak table
fheader	matrix of generalized ND parameters
rms	RMS noise level
iseed	random seed
file_path	optional output path for the simulated FID
verbose	logical indicating whether to print the command

Value

The integer exit status from ‘system2("SimTimeND", ...)’ (typically ‘0’ on success).

spec_bind	<i>Combine parameter lists referring to different spectra</i>
-----------	---

Description

Combine parameter lists referring to different spectra

Usage

spec_bind(...)

Arguments

... any number of parameter lists or a single list of parameter lists

spec_overlap_mat	<i>Determine a matrix of fractional peak overlap</i>
------------------	--

Description

Determine a matrix of fractional peak overlap

Usage

spec_overlap_mat(peak_int_list)

Arguments

peak_int_list list of peak intensity arrays

split_coupling_names	<i>Split string of scalar coupling names</i>
----------------------	--

Description

The only currently implemented way of splitting string is using whitespace

Usage

split_coupling_names(name_char)

Arguments

name_char single string (character vector of length 1) with scalar coupling names

tables_to_param_list *Convert tables with resonance/nuclei/couplings to a parameter list*

Description

Convert tables with resonance/nuclei/couplings to a parameter list

Usage

```
tables_to_param_list(spec_list, tables)
```

Arguments

spec_list	list of spectra
tables	list with resonance/nuclei/couplings tables

Value

A parameter 'list' containing 'start_list', 'group_list', 'comb_list', and associated naming vectors used by fitting functions.

Examples

```
spec_file <- system.file("extdata", "tyrosine", "proton.ft1", package = "fitnmr")
spec <- read_nmrpipe(spec_file)
```

```
start_resonances <- structure(
  list(
    x = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
    x_sc = c(
      "HA-HB3 HA-HB2",
      "HA-HB3 HB3-HB2",
      "HA-HB2 HB3-HB2",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5",
      "HD1/2-HE1/2-3 HD1/2-HE1/2-5"
    )
  ),
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2"),
  class = "data.frame"
)
```

```
start_nuclei <- structure(
  list(
    omega0_ppm = c(3.364, 2.635, 2.799, 6.94, 6.538),
    r2_hz = c(0.7, 0.7, 0.7, 0.7, 0.7)
  ),
  class = "data.frame",
  row.names = c("HA", "HB3", "HB2", "HD1/2", "HE1/2")
)
```

```

start_couplings <- structure(
  list(hz = c(7.153, 5.159, -13.941, 7.7, 2)),
  class = "data.frame",
  row.names = c("HA-HB3", "HA-HB2", "HB3-HB2", "HD1/2-HE1/2-3", "HD1/2-HE1/2-5")
)

start_tables <- list(
  resonances = start_resonances,
  nuclei = start_nuclei,
  couplings = start_couplings
)

param_list <- tables_to_param_list(list(spec), start_tables)
param_list$start_list$m0[] <- 1e9
arg_list <- param_list_to_arg_list(param_list)
fit_input <- do.call(make_fit_input, c(list(list(spec), omega0_plus = 0.075), arg_list))
fit_output <- perform_fit(fit_input)

```

update_fit_bounds *Update bounds on fitting parameters*

Description

Update bounds on fitting parameters

Usage

```

update_fit_bounds(
  fit_input,
  omega0_r2_factor = NULL,
  r2_bounds = NULL,
  sc_bounds = NULL
)

```

Arguments

fit_input	fit_input structure
omega0_r2_factor	optional factor for omega0 bounds based on r2
r2_bounds	optional numeric vector of length 2
sc_bounds	optional numeric vector of length 2

Value

A modified 'fit_input' list with updated parameter bounds.

Examples

```

spec_file <- system.file("extdata", "t1", "1.ft2", package = "fitnmr")
spec <- read_nmrpipe(spec_file, dim_order = "hx")
fit_input <- make_fit_input(
  list(spec),
  omega0_start = matrix(c(8.5400, 119.76), nrow = 2),
  omega0_plus = c(0.075, 0.75),
  r2_start = 4,
  m0_start = 1e9
)
fit_input <- update_fit_bounds(fit_input, omega0_r2_factor=1.5, r2_bounds=c(0.5, 20))

```

whz_to_pts	<i>Convert widths in Hz into points</i>
------------	---

Description

Convert widths in Hz into points

Usage

```
whz_to_pts(whz_mat, fheader)
```

Arguments

whz_mat	matrix of widths in Hz
fheader	matrix of generalized ND parameters

write_nmrdraw_peak_tab	<i>Write an NMRDraw formatted peak table</i>
------------------------	--

Description

Write an NMRDraw formatted peak table

Usage

```
write_nmrdraw_peak_tab(peak_tab, file_path)
```

Arguments

peak_tab	data frame containing peak table data
file_path	path to write the NMRDraw peak table file

Value

No return value, called for side effects (writes 'file_path').

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