

Package ‘spant’

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Type Package

Title MR Spectroscopy Analysis Tools

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Description Tools for reading, visualising and processing Magnetic Resonance Spectroscopy data. The package includes methods for spectral fitting: Wilson (2021) <[DOI:10.1002/mrm.28385](https://doi.org/10.1002/mrm.28385)> and spectral alignment: Wilson (2018) <[DOI:10.1002/mrm.27605](https://doi.org/10.1002/mrm.27605)>.

BugReports <https://github.com/martin3141/spant/issues/>

License GPL-3

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R topics documented:

spnt-package	8
abfit_opts	9
abfit_opts_v1_9_0	12
acquire	12
add_noise	13
align	13
apodise_xy	14
append_basis	15
append_coils	15
append_dyns	16
apply_axes	16
apply_mrs	17
apply_pvc	17
Arg.mrs_data	18
array2mrs_data	18
auto_phase	19
back_extrap_ar	20
basis2mrs_data	20
bbase	21
bc_als	22
bc_constant	22
beta2lw	23
bin_spec	23
calc_coil_noise_cor	24
calc_coil_noise_sd	24
calc_ed_from_lambda	25
calc_peak_info_vec	25
calc_sd_poly	26
calc_spec_diff	26
calc_spec_snr	27
check_lcm	28
check_tqn	28
circ_mask	28
collapse_to_dyns	29
comb_coils	29
comb_fit_list_fit_tables	30
comb_fit_list_result_tables	31
comb_fit_tables	32
comb_metab_ref	32
Conj.mrs_data	33
conv_mrs	33
crop_spec	34
crop_td_pts	34
crop_td_pts_pot	35
crop_xy	35
crossprod_3d	36

decimate_mrs_fd	36
decimate_mrs_td	37
def_acq_paras	37
def_fs	38
def_ft	38
def_N	39
def_nuc	39
def_ref	39
dicom_reader	40
diff_mrs	40
downsample_mrs_fd	41
downsample_mrs_td	41
ecc	42
elliptical_mask	42
est_noise_sd	43
fd2td	43
fd_conv_filt	44
fitamps	44
fit_diags	45
fit_mrs	45
fit_res2csv	47
fp_phase	47
fp_phase_correct	48
fp_scale	48
fs	49
ft_dyns	49
ft_shift	50
ft_shift_mat	50
gausswin_2d	51
gen_F	51
gen_F_xy	52
get_1h_brain_basis_paras	52
get_1h_brain_basis_paras_v1	53
get_1h_brain_basis_paras_v2	53
get_1h_brain_basis_paras_v3	54
get_2d_psf	54
get_acq_paras	55
get_dyns	55
get_even_dyns	56
get_fh_dyns	56
get_fit_map	57
get_fp	57
get_gaussian_pulse	58
get_head_dyns	58
get_lcm_cmd	59
get_metab	59
get_mol_names	59
get_mol_paras	60

get_mrsi2d_seg	60
get_mrsi_voi	61
get_mrsi_voxel	61
get_mrsi_voxel_xy_psf	62
get_mrs_affine	62
get_odd_dyns	63
get_ref	63
get_seg_ind	64
get_sh_dyns	64
get_slice	65
get_subset	65
get_svs_voi	66
get_tail_dyns	67
get_td_amp	67
get_tqn_cmd	68
get_uncoupled_mol	68
get_voi_cog	69
get_voi_seg	69
get_voi_seg_psf	70
get_voxel	70
gridplot	71
gridplot.mrs_data	71
grid_shift_xy	72
hsvd	72
hsvd_filt	73
hsvd_vec	74
hz	74
ift_shift	75
ift_shift_mat	75
Im.mrs_data	76
image.mrs_data	76
img2kspace_xy	77
interleave_dyns	78
int_spec	78
inv_even_dyns	79
inv_odd_dyns	79
is.def	80
is_fd	80
kspace2img_xy	81
l2_reg	81
lb	82
lw2alpha	83
lw2beta	83
mask_dyns	84
mask_fit_res	84
mask_xy	85
mask_xy_mat	85
mat2mrs_data	86

max_mrs	86
max_mrs_interp	87
mean.list	87
mean.mrs_data	88
mean_dyns	88
mean_dyn_blocks	89
mean_dyn_pairs	89
mean_mrs_list	90
median_dyns	90
Mod.mrs_data	91
mod_td	91
mrs_data2basis	92
mrs_data2mat	92
mrs_data2vec	93
mvfftshift	93
mvifftshift	94
n2coord	94
Ncoils	95
Ndyns	95
nifti_flip_lr	95
Npts	96
Nspec	96
Nx	97
Ny	97
Nz	97
ortho3	98
ortho3_inter	99
peak_info	99
pg_extrap_xy	100
phase	101
plot.fit_result	101
plot.mrs_data	103
plot_bc	105
plot_slice_fit	105
plot_slice_fit_inter	106
plot_slice_map	106
plot_slice_map_inter	107
plot_voi_overlay	109
plot_voi_overlay_seg	109
ppm	110
precomp	110
print.fit_result	111
print.mrs_data	111
qn_states	112
rats	112
Re.mrs_data	113
read_basis	114
read_basis_ac	114

read_ima_coil_dir	115
read_ima_dyn_dir	115
read_lcm_coord	116
read_mrs	116
read_mrs_tqn	117
read_siemens_txt_hdr	118
read_tqn_fit	119
read_tqn_result	119
recon_twix_2d_mrsi	120
rectangular_mask	120
rep_array_dim	121
rep_dyn	121
rep_mrs	122
resample_img	122
resample_voi	123
reslice_to_mrs	123
reson_table2mrs_data	124
re_weighting	124
rm_dyns	125
scale_amp_molal_pvc	125
scale_amp_molar	126
scale_amp_ratio	126
scale_amp_water_ratio	127
scale_mrs_amp	127
scale_spec	128
sd	128
sd.mrs_data	129
seconds	129
seq_cpmg_ideal	130
seq_mega_press_ideal	130
seq_press_ideal	131
seq_pulse_acquire	132
seq_pulse_acquire_31p	132
seq_laser_ideal	133
seq_spin_echo_ideal	133
seq_spin_echo_ideal_31p	134
seq_steam_ideal	134
set_def_acq_paras	135
set_lcm_cmd	135
set_lw	136
set_mask_xy_mat	136
set_precomp_mode	137
set_precomp_verbose	137
set_ref	137
set_td_pts	138
set_tqn_cmd	138
shift	139
shift_basis	139

sim_basis	140
sim_basis_1h_brain	140
sim_basis_1h_brain_press	141
sim_basis_tqn	142
sim_brain_1h	142
sim_mol	143
sim_noise	144
sim_resonances	145
sim_zero	146
sort_basis	146
spant_abfit_benchmark	147
spant_mpress_drift	147
spant_simulation_benchmark	148
spec_decomp	148
spec_op	149
spin_sys	149
spm_pve2categorical	150
ssp	150
stackplot	151
stackplot.fit_result	151
stackplot.mrs_data	153
sub_mean_dyns	154
sum_coils	155
sum_dyns	155
sum_mrs	156
sum_mrs_list	156
svs_1h_brain_analysis	157
svs_1h_brain_batch_analysis	158
td2fd	159
tdsr	159
td_conv_filt	160
varpro_3_para_opts	160
varpro_basic_opts	161
varpro_opts	162
vec2mrs_data	163
write_basis	163
write_basis_tqn	164
write_mrs	164
write_mrs_nifti	165
zero_fade_spec	165
zero_nzoc	166
zf	166
zf_xy	167

spant-package

spant: spectroscopy analysis tools.

Description

spant provides a set of tools for reading, visualising and processing Magnetic Resonance Spectroscopy (MRS) data.

Details

To get started with spant, take a look at the introduction vignette:

```
vignette("spant-intro", package="spant")
```

Full list of vignettes:

```
browseVignettes(package = "spant")
```

Full list of functions:

```
help(package = spant, help_type = "html")
```

An online version of the documentation is available from:

<https://martin3141.github.io/spant/>

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See Also

Useful links:

- <https://martin3141.github.io/spant/>
- <https://github.com/martin3141/spant/>
- Report bugs at <https://github.com/martin3141/spant/issues/>

abfit_opts	<i>Return a list of options for an ABfit analysis.</i>
------------	--

Description

Return a list of options for an ABfit analysis.

Usage

```
abfit_opts(  
  init_damping = 5,  
  maxiters = 1024,  
  max_shift = 0.078,  
  max_damping = 15,  
  max_phase = 360,  
  lambda = NULL,  
  ppm_left = 4,  
  ppm_right = 0.2,  
  zp = TRUE,  
  bl_ed_pppm = 2,  
  auto_bl_flex = TRUE,  
  bl_comps_pppm = 15,  
  export_sp_fit = FALSE,  
  max_asym = 0.25,  
  max_basis_shift = 0.0078,  
  max_basis_damping = 2,  
  maxiters_pre = 1000,  
  algo_pre = "NLOPT_LN_NELDERMEAD",  
  min_bl_ed_pppm = NULL,  
  max_bl_ed_pppm = 7,  
  auto_bl_flex_n = 20,  
  pre_fit_bl_ed_pppm = 1,  
  remove_lip_mm_prefit = FALSE,  
  pre_align = TRUE,  
  max_pre_align_shift = 0.1,  
  pre_align_ref_freqs = c(2.01, 3.03, 3.22),  
  noise_region = c(-0.5, -2.5),  
  optimal_smooth_criterion = "maic",  
  aic_smoothing_factor = 5,  
  anal_jac = TRUE,  
  pre_fit_ppm_left = 4,  
  pre_fit_ppm_right = 1.8,  
  phi1_optim = FALSE,  
  phi1_init = 0,  
  max_dphi1 = 0.2,  
  max_basis_shift_broad = 0.0078,  
  max_basis_damping_broad = 2,
```

```

ahat_calc_method = "lh_pnnls",
prefit_phase_search = TRUE,
freq_reg = NULL,
output_all_paras = FALSE
)

```

Arguments

<code>init_damping</code>	initial value of the Gaussian global damping parameter (Hz). Very poorly shimmed or high field data may benefit from a larger value.
<code>maxiters</code>	The maximum number of iterations to run for the detailed fit.
<code>max_shift</code>	The maximum allowable shift to be applied in the optimisation phase of fitting (ppm).
<code>max_damping</code>	maximum permitted value of the global damping parameter (Hz).
<code>max_phase</code>	the maximum absolute permitted value of the global zero-order phase term (degrees). Note, the <code>prefit_phase_search</code> option is not constrained by this term.
<code>lambda</code>	manually set the the baseline smoothness parameter.
<code>ppm_left</code>	downfield frequency limit for the fitting range (ppm).
<code>ppm_right</code>	upfield frequency limit for the fitting range (ppm).
<code>zp</code>	zero pad the data to twice the original length before fitting.
<code>b1_ed_pppm</code>	manually set the the baseline smoothness parameter (ED per ppm).
<code>auto_b1_flex</code>	automatically determine the level of baseline smoothness.
<code>b1_comps_pppm</code>	spline basis density (signals per ppm).
<code>export_sp_fit</code>	add the fitted spline functions to the fit result.
<code>max_asym</code>	maximum allowable value of the asymmetry parameter.
<code>max_basis_shift</code>	maximum allowable frequency shift for individual basis signals (ppm).
<code>max_basis_damping</code>	maximum allowable Lorentzian damping factor for individual basis signals (Hz).
<code>maxiters_pre</code>	maximum iterations for the coarse (pre-)fit.
<code>algo_pre</code>	optimisation method for the coarse (pre-)fit.
<code>min_b1_ed_pppm</code>	minimum value for the candidate baseline flexibility analyses (ED per ppm).
<code>max_b1_ed_pppm</code>	minimum value for the candidate baseline flexibility analyses (ED per ppm).
<code>auto_b1_flex_n</code>	number of candidate baseline analyses to perform.
<code>pre_fit_b1_ed_pppm</code>	level of baseline flexibility to use in the coarse fitting stage of the algorithm (ED per ppm).
<code>remove_lip_mm_prefit</code>	remove broad signals in the coarse fitting stage of the algorithm.
<code>pre_align</code>	perform a pre-alignment step before coarse fitting.
<code>max_pre_align_shift</code>	maximum allowable shift in the pre-alignment step (ppm).

```

pre_align_ref_freqs
    a vector of prominent spectral frequencies used in the pre-alignment step (ppm).

noise_region    spectral region to estimate the noise level (ppm).

optimal_smooth_criterion
    method to determine the optimal smoothness.

aic_smoothing_factor
    modification factor for the AIC calculation.

anal_jac        use a analytical approximation to the jacobian in the detailed fitting stage.

pre_fit_ppm_left
    downfield frequency limit for the fitting range in the coarse fitting stage of the
    algorithm (ppm).

pre_fit_ppm_right
    upfield frequency limit for the fitting range in the coarse fitting stage of the
    algorithm (ppm).

phi1_optim      apply and optimise a frequency dependant phase term.

phi1_init        initial value for the frequency dependant phase term (ms).

max_dphi1       maximum allowable change from the initial frequency dependant phase term
                (ms).

max_basis_shift_broad
    maximum allowable shift for broad signals in the basis (ppm). Determined based
    on their name beginning with Lip or MM.

max_basis_damping_broad
    maximum allowable Lorentzian damping for broad signals in the basis (Hz).
    Determined based on their name beginning with Lip or MM.

ahat_calc_method
    method to calculate the metabolite amplitudes. May be one of: "lh_pnnls" or
    "ls".

prefit_phase_search
    perform a 1D search for the optimal phase in the prefit stage of the algorithm.

freq_reg        frequency shift parameter.

output_all_paras
    include more fitting parameters in the fit table, e.g. individual shift and damping
    factors for each basis set element.

```

Value

full list of options.

Examples

```
opts <- abfit_opts(ppm_left = 4.2, noise_region = c(-1, -3))
```

`abfit_opts_v1_9_0` *Return a list of options for an ABfit analysis to maintain comparability with analyses performed with version 1.9.0 (and earlier) of spant.*

Description

Return a list of options for an ABfit analysis to maintain comparability with analyses performed with version 1.9.0 (and earlier) of spant.

Usage

```
abfit_opts_v1_9_0(...)
```

Arguments

...	arguments passed to <code>abfit_opts</code> .
-----	---

Value

full list of options.

`acquire` *Simulate pulse sequence acquisition.*

Description

Simulate pulse sequence acquisition.

Usage

```
acquire(sys, rec_phase = 180, tol = 1e-04, detect = NULL)
```

Arguments

<code>sys</code>	spin system object.
<code>rec_phase</code>	receiver phase in degrees.
<code>tol</code>	ignore resonance amplitudes below this threshold.
<code>detect</code>	detection nuclei.

Value

a list of resonance amplitudes and frequencies.

add_noise	<i>Add noise to an mrs_data object.</i>
-----------	---

Description

Add noise to an mrs_data object.

Usage

```
add_noise(mrs_data, sd = 0.1, fd = TRUE)
```

Arguments

mrs_data	data to add noise to.
sd	standard deviation of the noise.
fd	generate the noise samples in the frequency-domain (TRUE) or time-domain (FALSE). This is required since the absolute value of the standard deviation of noise samples changes when data is Fourier transformed.

Value

mrs_data object with additive normally distributed noise.

align	<i>Align spectra to a reference frequency using a convolution based method.</i>
-------	---

Description

Align spectra to a reference frequency using a convolution based method.

Usage

```
align(  
  mrs_data,  
  ref_freq = 4.65,  
  zf_factor = 2,  
  lb = 2,  
  max_shift = 20,  
  ret_df = FALSE,  
  mean_dyncs = FALSE  
)
```

Arguments

<code>mrs_data</code>	data to be aligned.
<code>ref_freq</code>	reference frequency in ppm units. More than one frequency may be specified.
<code>zf_factor</code>	zero filling factor to increase alignment resolution.
<code>lb</code>	line broadening to apply to the reference signal.
<code>max_shift</code>	maximum allowable shift in Hz.
<code>ret_df</code>	return frequency shifts in addition to aligned data (logical).
<code>mean_dync</code>	align the mean spectrum and apply the same shift to each dynamic.

Value

aligned data object.

apodise_xy

Apodise MRSI data in the x-y direction with a k-space filter.

Description

Apodise MRSI data in the x-y direction with a k-space filter.

Usage

```
apodise_xy(mrs_data, func = "hamming", w = 2.5)
```

Arguments

<code>mrs_data</code>	MRSI data.
<code>func</code>	must be "hamming" or "gaussian".
<code>w</code>	the reciprocal of the standard deviation for the Gaussian function.

Value

apodised data.

append_basis*Combine a pair of basis set objects.*

Description

Combine a pair of basis set objects.

Usage

```
append_basis(basis_a, basis_b)
```

Arguments

basis_a first basis.

basis_b second basis.

Value

combined basis set object.

append_coils*Append MRS data across the coil dimension, assumes they matched across the other dimensions.*

Description

Append MRS data across the coil dimension, assumes they matched across the other dimensions.

Usage

```
append_coils(...)
```

Arguments

... MRS data objects as arguments, or a list of MRS data objects.

Value

a single MRS data object with the input objects concatenated together.

`append_dyncs`

Append MRS data across the dynamic dimension, assumes they matched across the other dimensions.

Description

Append MRS data across the dynamic dimension, assumes they matched across the other dimensions.

Usage

```
append_dyncs(...)
```

Arguments

... MRS data objects as arguments, or a list of MRS data objects.

Value

a single MRS data object with the input objects concatenated together.

`apply_axes`

Apply a function over specified array axes.

Description

Apply a function over specified array axes.

Usage

```
apply_axes(x, axes, fun, ...)
```

Arguments

x	an array.
axes	a vector of axes to apply fun over.
fun	function to be applied.
...	optional arguments to fun.

Value

array.

Examples

```
z <- array(1:1000, dim = c(10, 10, 10))
a <- apply_axes(z, 3, fft)
a[1,1,] == fft(z[1,1,])
a <- apply_axes(z, 3, sum)
a[1,1,] == sum(z[1,1,])
```

apply_mrs

Apply a function across given dimensions of a MRS data object.

Description

Apply a function across given dimensions of a MRS data object.

Usage

```
apply_mrs(mrs_data, dims, fun, ..., data_only = FALSE)
```

Arguments

mrs_data	MRS data.
dims	dimensions to apply the function.
fun	name of the function.
...	arguments to the function.
data_only	return an array rather than an MRS data object.

apply_pvc

Convert default LCM/TARQUIN concentration scaling to molal units with partial volume correction.

Description

Convert default LCM/TARQUIN concentration scaling to molal units with partial volume correction.

Usage

```
apply_pvc(fit_result, p_vols, te, tr)
```

Arguments

fit_result	a fit_result object to apply partial volume correction.
p_vols	a numeric vector of partial volumes.
te	the MRS TE.
tr	the MRS TR.

Value

a fit_result object with a rescaled results table.

`Arg.mrs_data`

Apply Arg operator to an MRS dataset.

Description

Apply Arg operator to an MRS dataset.

Usage

```
## S3 method for class 'mrs_data'
Arg(z)
```

Arguments

`z` MRS data.

Value

MRS data following Arg operator.

`array2mrs_data`

Convert a 7 dimensional array in into a mrs_data object. The array dimensions should be ordered as : dummy, X, Y, Z, dynamic, coil, FID.

Description

Convert a 7 dimensional array in into a mrs_data object. The array dimensions should be ordered as : dummy, X, Y, Z, dynamic, coil, FID.

Usage

```
array2mrs_data(
  data_array,
  fs = def_fs(),
  ft = def_ft(),
  ref = def_ref(),
  nuc = def_nuc(),
  fd = FALSE
)
```

Arguments

data_array	7d data array.
fs	sampling frequency in Hz.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
nuc	nucleus that is resonant at the transmitter frequency.
fd	flag to indicate if the matrix is in the frequency domain (logical).

Value

mrs_data object.

auto_phase	<i>Perform zeroth-order phase correction based on the minimisation of the squared difference between the real and magnitude components of the spectrum.</i>
------------	---

Description

Perform zeroth-order phase correction based on the minimisation of the squared difference between the real and magnitude components of the spectrum.

Usage

```
auto_phase(mrs_data, xlim = NULL, ret_phase = FALSE)
```

Arguments

mrs_data	an object of class mrs_data.
xlim	frequency range (default units of PPM) to including in the phase.
ret_phase	return phase values (logical).

Value

MRS data object and phase values (optional).

<code>back_extrap_ar</code>	<i>Back extrapolate time-domain data points using an autoregressive model.</i>
-----------------------------	--

Description

Back extrapolate time-domain data points using an autoregressive model.

Usage

```
back_extrap_ar(
  mrs_data,
  extrap_pts,
  pred_pts = NULL,
  method = "burg",
  rem_add = TRUE,
  ...
)
```

Arguments

<code>mrs_data</code>	mrs_data object.
<code>extrap_pts</code>	number of points to extrapolate.
<code>pred_pts</code>	number of points to base the extrapolation on.
<code>method</code>	character string specifying the method to fit the model. Must be one of the strings in the default argument (the first few characters are sufficient). Defaults to "burg".
<code>rem_add</code>	remove additional points from the end of the FID to maintain the original length of the dataset. Default to TRUE.
...	additional arguments to specific methods, see ?ar.

Value

back extrapolated data.

<code>basis2mrs_data</code>	<i>Convert a basis object to an mrs_data object - where basis signals are spread across the dynamic dimension.</i>
-----------------------------	--

Description

Convert a basis object to an mrs_data object - where basis signals are spread across the dynamic dimension.

Usage

```
basis2mrs_data(basis, sum_elements = FALSE, amps = NULL, shifts = NULL)
```

Arguments

- | | |
|--------------|---|
| basis | basis set object. |
| sum_elements | return the sum of basis elements (logical) |
| amps | a vector of scaling factors to apply to each basis element. |
| shifts | a vector of frequency shifts (in ppm) to apply to each basis element. |

Value

an mrs_data object with basis signals spread across the dynamic dimension or summed.

bbase	<i>Generate a spline basis, slightly adapted from : "Splines, knots, and penalties", Eilers 2010.</i>
-------	---

Description

Generate a spline basis, slightly adapted from : "Splines, knots, and penalties", Eilers 2010.

Usage

```
bbase(N, number, deg = 3)
```

Arguments

- | | |
|--------|---|
| N | number of data points. |
| number | number of spline functions. |
| deg | spline degree : deg = 1 linear, deg = 2 quadratic, deg = 3 cubic. |

Value

spline basis as a matrix.

<code>bc_als</code>	<i>Baseline correction using the ALS method.</i>
---------------------	--

Description

Eilers P. H. C. and Boelens H. F. M. (2005) Baseline correction with asymmetric least squares smoothing. Leiden Univ. Medical Centre Report.

Usage

```
bc_als(mrs_data, lambda = 10000, p = 0.001, ret_bc_only = TRUE)
```

Arguments

<code>mrs_data</code>	mrs_data object.
<code>lambda</code>	controls the baseline flexibility.
<code>p</code>	controls the penalty for negative data points.
<code>ret_bc_only</code>	return the baseline corrected data only. When FALSE the baseline estimate and input data will be returned.

Value

baseline corrected data.

<code>bc_constant</code>	<i>Remove a constant baseline offset based on a reference spectral region.</i>
--------------------------	--

Description

Remove a constant baseline offset based on a reference spectral region.

Usage

```
bc_constant(mrs_data, xlim)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>xlim</code>	spectral range containing a flat baseline region to measure the offset.

Value

baseline corrected data.

beta2lw	<i>Covert a beta value in the time-domain to an equivalent linewidth in Hz: $x * \exp(-i * t * t * \text{beta})$.</i>
---------	--

Description

Covert a beta value in the time-domain to an equivalent linewidth in Hz: $x * \exp(-i * t * t * \text{beta})$.

Usage

```
beta2lw(beta)
```

Arguments

beta beta damping value.

Value

linewidth value in Hz.

bin_spec	<i>Bin equally spaced spectral regions.</i>
----------	---

Description

Bin equally spaced spectral regions.

Usage

```
bin_spec(mrs_data, width = 0.05, unit = "ppm")
```

Arguments

mrs_data data to be "binned".
width bin width.
unit bin width unit, can be "ppm" (default) or "pts".

Value

binned mrs_data object.

calc_coil_noise_cor *Calculate the noise correlation between coil elements.*

Description

Calculate the noise correlation between coil elements.

Usage

```
calc_coil_noise_cor(noise_data)
```

Arguments

noise_data *mrs_data object with one FID for each coil element.*

Value

correlation matrix.

calc_coil_noise_sd *Calculate the noise standard deviation for each coil element.*

Description

Calculate the noise standard deviation for each coil element.

Usage

```
calc_coil_noise_sd(noise_data)
```

Arguments

noise_data *mrs_data object with one FID for each coil element.*

Value

array of standard deviations.

calc_ed_from_lambda *Calculate the effective dimensions of a spline smoother from lambda.*

Description

Calculate the effective dimensions of a spline smoother from lambda.

Usage

```
calc_ed_from_lambda(spline_basis, deriv_mat, lambda)
```

Arguments

spline_basis spline basis.
deriv_mat derivative matrix.
lambda smoothing parameter.

Value

the effective dimension value.

calc_peak_info_vec *Calculate the FWHM of a peak from a vector of intensity values.*

Description

Calculate the FWHM of a peak from a vector of intensity values.

Usage

```
calc_peak_info_vec(data_pts, interp_f)
```

Arguments

data_pts input vector.
interp_f interpolation factor to improve the FWHM estimate.

Value

a vector of: x position of the highest data point, maximum peak value in the y axis, FWHM in the units of data points.

<code>calc_sd_poly</code>	<i>Perform a polynomial fit, subtract and return the standard deviation of the residuals.</i>
---------------------------	---

Description

Perform a polynomial fit, subtract and return the standard deviation of the residuals.

Usage

```
calc_sd_poly(y, degree = 1)
```

Arguments

<code>y</code>	array.
<code>degree</code>	polynomial degree.

Value

standard deviation of the fit residuals.

<code>calc_spec_diff</code>	<i>Calculate the sum of squares differences between two mrs_data objects.</i>
-----------------------------	---

Description

Calculate the sum of squares differences between two mrs_data objects.

Usage

```
calc_spec_diff(mrs_data, ref = NULL, xlim = c(4, 0.5))
```

Arguments

<code>mrs_data</code>	mrs_data object.
<code>ref</code>	reference mrs_data object to calculate differences.
<code>xlim</code>	spectral limits to perform calculation.

Value

an array of the sum of squared difference values.

calc_spec_snr	<i>Calculate the spectral SNR.</i>
---------------	------------------------------------

Description

SNR is defined as the maximum signal value divided by the standard deviation of the noise.

Usage

```
calc_spec_snr(  
  mrs_data,  
  sig_region = c(4, 0.5),  
  noise_region = c(-0.5, -2.5),  
  p_order = 2,  
  interp_f = 4,  
  full_output = FALSE  
)
```

Arguments

mrs_data	an object of class <code>mrs_data</code> .
sig_region	a ppm region to define where the maximum signal value should be estimated.
noise_region	a ppm region to defined where the noise level should be estimated.
p_order	polynomial order to fit to the noise region before estimating the standard deviation.
interp_f	interpolation factor to improve detection of the highest signal value.
full_output	output signal, noise and SNR values separately.

Details

The mean noise value is subtracted from the maximum signal value to reduce DC offset bias. A polynomial detrending fit (second order by default) is applied to the noise region before the noise standard deviation is estimated.

Value

an array of SNR values.

`check_lcm`*Check LCModel can be run***Description**

Check LCModel can be run

Usage

```
check_lcm()
```

`check_tqn`*Check the TARQUIN binary can be run***Description**

Check the TARQUIN binary can be run

Usage

```
check_tqn()
```

`circ_mask`*Create a logical circular mask spanning the full extent of an n x n matrix.***Description**

Create a logical circular mask spanning the full extent of an n x n matrix.

Usage

```
circ_mask(d, n, offset = 1)
```

Arguments

- `d` diameter of the mask.
- `n` number of matrix rows and columns.
- `offset` offset the mask centre in matrix dimension units.

Value

logical n x n mask matrix.

collapse_to_dync	<i>Collapse MRS data by concatenating spectra along the dynamic dimension.</i>
------------------	--

Description

Collapse MRS data by concatenating spectra along the dynamic dimension.

Usage

```
collapse_to_dync(x, rm_masked = FALSE)

## S3 method for class 'mrs_data'
collapse_to_dync(x, rm_masked = FALSE)

## S3 method for class 'fit_result'
collapse_to_dync(x, rm_masked = FALSE)
```

Arguments

x	data object to be collapsed (mrs_data or fit_result object).
rm_masked	remove masked dynamics from the output.

Value

collapsed data with spectra or fits concatenated along the dynamic dimension.

comb_coils	<i>Combine coil data based on the first data point of a reference signal.</i>
------------	---

Description

By default, elements are phased and scaled prior to summation. Where a reference signal is not given, the mean dynamic signal will be used instead.

Usage

```
comb_coils(
  metab,
  ref = NULL,
  noise = NULL,
  scale = TRUE,
  scale_method = "sig_noise_sq",
  sum_coils = TRUE,
  noise_region = c(-0.5, -2.5),
```

```

    average_ref_dyns = TRUE,
    ref_pt_index = 1,
    ret_metab_only = FALSE
)

```

Arguments

metab	MRS data containing metabolite data.
ref	MRS data containing reference data (optional).
noise	MRS data from a noise scan (optional).
scale	option to rescale coil elements based on the first data point (logical).
scale_method	one of "sig_noise_sq", "sig_noise" or "sig".
sum_coils	sum the coil elements as a final step (logical).
noise_region	the spectral region (in ppm) to estimate the noise.
average_ref_dyns	take the mean of the reference scans in the dynamic dimension before use.
ref_pt_index	time-domain point to use for estimating phase and scaling values.
ret_metab_only	return the metabolite data only, even if reference data has been specified.

Value

MRS data.

comb_fit_list_fit_tables

Combine all fitting data points from a list of fits into a single data frame.

Description

Combine all fitting data points from a list of fits into a single data frame.

Usage

```

comb_fit_list_fit_tables(
  fit_list,
  add_extra = TRUE,
  harmonise_ppm = TRUE,
  inc_basis_sigs = FALSE,
  inc_indices = TRUE,
  add_res_id = TRUE
)

```

Arguments

fit_list	list of fit_result objects.
add_extra	add variables in the extra data frame to the output (TRUE).
harmonise_ppm	ensure the ppm scale for each fit is identical to the first.
inc_basis_sigs	include the individual fitting basis signals in the output table, defaults to FALSE.
inc_indices	include indices such as X, Y and coil in the output, defaults to TRUE. These are generally not useful for SVS analysis.
add_res_id	add a res_id column to the output to distinguish between datasets.

Value

a data frame containing the fit data points.

comb_fit_list_result_tables

Combine the fit result tables from a list of fit results.

Description

Combine the fit result tables from a list of fit results.

Usage

```
comb_fit_list_result_tables(fit_list, add_extra = TRUE, add_res_id = TRUE)
```

Arguments

fit_list	a list of fit_result objects.
add_extra	add variables in the extra data frame to the output (TRUE).
add_res_id	add a res_id column to the output to distinguish between datasets.

Value

a data frame combine all fit result tables with an additional id column to differentiate between data sets. Any variables in the extra data frame may be optionally added to the result.

<code>comb_fit_tables</code>	<i>Combine all fitting data points into a single data frame.</i>
------------------------------	--

Description

Combine all fitting data points into a single data frame.

Usage

```
comb_fit_tables(fit_res, inc_basis_sigs = FALSE, inc_indices = TRUE)
```

Arguments

- | | |
|-----------------------------|---|
| <code>fit_res</code> | a single fit_result object. |
| <code>inc_basis_sigs</code> | include the individual fitting basis signals in the output table, defaults to FALSE. |
| <code>inc_indices</code> | include indices such as X, Y and coil in the output, defaults to TRUE. These are generally not useful for SVS analysis. |

Value

a data frame containing the fit data points.

<code>comb_metab_ref</code>	<i>Combine a reference and metabolite mrs_data object.</i>
-----------------------------	--

Description

Combine a reference and metabolite mrs_data object.

Usage

```
comb_metab_ref(metab, ref)
```

Arguments

- | | |
|--------------------|-----------------------------|
| <code>metab</code> | metabolite mrs_data object. |
| <code>ref</code> | reference mrs_data object. |

Value

combined metabolite and reference mrs_data object.

Conj.mrs_data *Apply Conj operator to an MRS dataset.*

Description

Apply Conj operator to an MRS dataset.

Usage

```
## S3 method for class 'mrs_data'  
Conj(z)
```

Arguments

z MRS data.

Value

MRS data following Conj operator.

conv_mrs *Convolve two MRS data objects.*

Description

Convolve two MRS data objects.

Usage

```
conv_mrs(mrs_data, conv)
```

Arguments

mrs_data MRS data to be convolved.
conv convolution data stored as an mrs_data object.

Value

convolved data.

crop_spec*Crop mrs_data object based on a frequency range.***Description**

Crop mrs_data object based on a frequency range.

Usage

```
crop_spec(mrs_data, xlim = c(4, 0.2), scale = "ppm")
```

Arguments

<code>mrs_data</code>	MRS data.
<code>xlim</code>	range of values to crop in the spectral dimension eg <code>xlim = c(4, 0.2)</code> .
<code>scale</code>	the units to use for the frequency scale, can be one of: "ppm", "hz" or "points".

Value

cropped mrs_data object.

crop_td_pts*Crop mrs_data object data points in the time-domain.***Description**

Crop mrs_data object data points in the time-domain.

Usage

```
crop_td_pts(mrs_data, start = NULL, end = NULL)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>start</code>	starting data point (defaults to 1).
<code>end</code>	ending data point (defaults to the last saved point).

Value

cropped mrs_data object.

crop_td_pts_pot	<i>Crop mrs_data object data points in the time-domain rounding down to the next smallest power of two (pot). Data that already has a pot length will not be changed.</i>
-----------------	---

Description

Crop mrs_data object data points in the time-domain rounding down to the next smallest power of two (pot). Data that already has a pot length will not be changed.

Usage

```
crop_td_pts_pot(mrs_data)
```

Arguments

mrs_data MRS data.

Value

cropped mrs_data object.

crop_xy	<i>Crop an MRSI dataset in the x-y direction</i>
---------	--

Description

Crop an MRSI dataset in the x-y direction

Usage

```
crop_xy(mrs_data, x_dim, y_dim)
```

Arguments

mrs_data MRS data object.
x_dim x dimension output length.
y_dim y dimension output length.

Value

selected subset of MRS data.

<code>crossprod_3d</code>	<i>Compute the vector cross product between vectors x and y. Adapted from http://stackoverflow.com/questions/15162741/what-is-rs-crossproduct-function</i>
---------------------------	---

Description

Compute the vector cross product between vectors x and y. Adapted from <http://stackoverflow.com/questions/15162741/what-is-rs-crossproduct-function>

Usage

```
crossprod_3d(x, y)
```

Arguments

- | | |
|----------------|---------------------|
| <code>x</code> | vector of length 3. |
| <code>y</code> | vector of length 3. |

Value

vector cross product of x and y.

<code>decimate_mrs_fd</code>	<i>Decimate an MRS signal to half the original sampling frequency by filtering in the frequency domain before down sampling.</i>
------------------------------	--

Description

Decimate an MRS signal to half the original sampling frequency by filtering in the frequency domain before down sampling.

Usage

```
decimate_mrs_fd(mrs_data)
```

Arguments

- | | |
|-----------------------|------------------|
| <code>mrs_data</code> | MRS data object. |
|-----------------------|------------------|

Value

decimated data at half the original sampling frequency.

decimate_mrs_td	<i>Decimate an MRS signal by filtering in the time domain before down-sampling.</i>
-----------------	---

Description

Decimate an MRS signal by filtering in the time domain before downsampling.

Usage

```
decimate_mrs_td(mrs_data, q = 2, n = 4, ftype = "iir")
```

Arguments

mrs_data	MRS data object.
q	integer factor to downsample by (default = 2).
n	filter order used in the downsampling.
ftype	filter type, "iir" or "fir".

Value

decimated data.

def_acq_paras	<i>Return (and optionally modify using the input arguments) a list of the default acquisition parameters.</i>
---------------	---

Description

Return (and optionally modify using the input arguments) a list of the default acquisition parameters.

Usage

```
def_acq_paras(  
    ft = getopt("spant.def_ft"),  
    fs = getopt("spant.def_fs"),  
    N = getopt("spant.def_N"),  
    ref = getopt("spant.def_ref"),  
    nuc = getopt("spant.def_nuc")  
)
```

Arguments

<code>ft</code>	specify the transmitter frequency in Hz.
<code>fs</code>	specify the sampling frequency in Hz.
<code>N</code>	specify the number of data points in the spectral dimension.
<code>ref</code>	specify the reference value for ppm scale.
<code>nuc</code>	specify the resonant nucleus.

Value

A list containing the following elements:

- `ft` transmitter frequency in Hz.
- `fs` sampling frequency in Hz.
- `N` number of data points in the spectral dimension.
- `ref` reference value for ppm scale.
- `nuc` resonant nucleus.

`def_fs`

Return the default sampling frequency in Hz.

Description

Return the default sampling frequency in Hz.

Usage

```
def_fs()
```

Value

sampling frequency in Hz.

`def_ft`

Return the default transmitter frequency in Hz.

Description

Return the default transmitter frequency in Hz.

Usage

```
def_ft()
```

Value

transmitter frequency in Hz.

def_N

Return the default number of data points in the spectral dimension.

Description

Return the default number of data points in the spectral dimension.

Usage

def_N()

Value

number of data points in the spectral dimension.

def_nuc

Return the default nucleus.

Description

Return the default nucleus.

Usage

def_nuc()

Value

number of data points in the spectral dimension.

def_ref

Return the default reference value for ppm scale.

Description

Return the default reference value for ppm scale.

Usage

def_ref()

Value

reference value for ppm scale.

dicom_reader *A very simple DICOM reader.*

Description

Note this reader is very basic and does not use a DICOM dictionary or try to convert the data to the correct datatype. For a more robust and sophisticated reader use the *oro.dicom* package.

Usage

```
dicom_reader(
    input,
    tags = list(sop_class_uid = "0008,0016"),
    endian = "little",
    debug = FALSE
)
```

Arguments

input	either a file path or raw binary object.
tags	a named list of tags to be extracted from the file. eg tags <- list(spec_data = "7FE1,1010", pat_name = "0010,0010")
endian	can be "little" or "big".
debug	print out some debugging information, can be "little" or "big".

Value

a list with the same structure as the input, but with tag codes replaced with the corresponding data in a raw format.

diff_mrs *Apply the diff operator to an MRS dataset in the FID/spectral dimension.*

Description

Apply the diff operator to an MRS dataset in the FID/spectral dimension.

Usage

```
diff_mrs(mrs_data, ...)
```

Arguments

mrs_data	MRS data.
...	additional arguments to the diff function.

Value

MRS data following diff operator.

downsample_mrs_fd	<i>Downsample an MRS signal by a factor of 2 using an FFT "brick-wall" filter.</i>
-------------------	--

Description

Downsample an MRS signal by a factor of 2 using an FFT "brick-wall" filter.

Usage

```
downsample_mrs_fd(mrs_data)
```

Arguments

mrs_data MRS data object.

Value

downsampled data.

downsample_mrs_td	<i>Downsample an MRS signal by a factor of 2 by removing every other data point in the time-domain. Note, signals outside the new sampling frequency will be aliased.</i>
-------------------	---

Description

Downsample an MRS signal by a factor of 2 by removing every other data point in the time-domain. Note, signals outside the new sampling frequency will be aliased.

Usage

```
downsample_mrs_td(mrs_data)
```

Arguments

mrs_data MRS data object.

Value

downsampled data.

<code>ecc</code>	<i>Eddy current correction.</i>
------------------	---------------------------------

Description

Apply eddy current correction using the Klose method.

Usage

```
ecc(metab, ref, rev = FALSE)
```

Arguments

<code>metab</code>	MRS data to be corrected.
<code>ref</code>	reference dataset.
<code>rev</code>	reverse the correction.

Details

In vivo proton spectroscopy in presence of eddy currents. Klose U. Magn Reson Med. 1990 Apr;14(1):26-30.

Value

corrected data in the time domain.

<code>elliptical_mask</code>	<i>Create an elliptical mask stored as a matrix of logical values.</i>
------------------------------	--

Description

Create an elliptical mask stored as a matrix of logical values.

Usage

```
elliptical_mask(xN, yN, x0, y0, xr, yr, angle)
```

Arguments

<code>xN</code>	number of pixels in the x dimension.
<code>yN</code>	number of pixels in the y dimension.
<code>x0</code>	centre of ellipse in the x direction in units of pixels.
<code>y0</code>	centre of ellipse in the y direction in units of pixels.
<code>xr</code>	radius in the x direction in units of pixels.
<code>yr</code>	radius in the y direction in units of pixels.
<code>angle</code>	angle of rotation in degrees.

Value

logical mask matrix with dimensions `fov_yN` x `fov_xN`.

<code>est_noise_sd</code>	<i>Estimate the standard deviation of the noise from a segment of an mrs_data object.</i>
---------------------------	---

Description

Estimate the standard deviation of the noise from a segment of an `mrs_data` object.

Usage

```
est_noise_sd(mrs_data, n = 100, offset = 100, p_order = 2)
```

Arguments

<code>mrs_data</code>	MRS data object.
<code>n</code>	number of data points (taken from the end of array) to use in the estimation.
<code>offset</code>	number of final points to exclude from the calculation.
<code>p_order</code>	polynomial order to fit to the data before estimating the standard deviation.

Value

standard deviation array.

<code>fd2td</code>	<i>Transform frequency-domain data to the time-domain.</i>
--------------------	--

Description

Transform frequency-domain data to the time-domain.

Usage

```
fd2td(mrs_data)
```

Arguments

<code>mrs_data</code>	MRS data in frequency-domain representation.
-----------------------	--

Value

MRS data in time-domain representation.

fd_conv_filt *Frequency-domain convolution based filter.*

Description

Frequency-domain convolution based filter.

Usage

```
fd_conv_filt(mrs_data, K = 25, ext = 1)
```

Arguments

mrs_data	MRS data to be filtered.
K	window width in data points.
ext	point separation for linear extrapolation.

fit_amps *Extract the fit amplitudes from an object of class fit_result.*

Description

Extract the fit amplitudes from an object of class `fit_result`.

Usage

```
fit_amps(
  x,
  inc_index = FALSE,
  sort_names = FALSE,
  append_common_1h_comb = TRUE
)
```

Arguments

x	<code>fit_result</code> object.
inc_index	include columns for the voxel index.
sort_names	sort the basis set names alphabetically.
append_common_1h_comb	append commonly used 1H metabolite combinations eg tNAA = NAA + NAAG.

Value

a dataframe of amplitudes.

fit_diags*Calculate diagnostic information for object of class fit_result.*

Description

Calculate diagnostic information for object of class `fit_result`.

Usage

```
fit_diags(x, amps = NULL)
```

Arguments

x	fit_result object.
amps	known metabolite amplitudes.

Value

a dataframe of diagnostic information.

fit_mrs*Perform a fit based analysis of MRS data.*

Description

Note that TARQUIN and LCModel require these packages to be installed, and the functions `set_tqn_cmd` and `set_lcm_cmd` (respectively) need to be used to specify the location of these software packages.

Usage

```
fit_mrs(  
  metab,  
  basis = NULL,  
  method = "ABFIT",  
  w_ref = NULL,  
  opts = NULL,  
  parallel = FALSE,  
  time = TRUE,  
  progress = "text",  
  extra = NULL  
)
```

Arguments

<code>metab</code>	metabolite data.
<code>basis</code>	basis class object or character vector to basis file in LCModel .basis format.
<code>method</code>	'ABFIT' (default), 'VARPRO', 'VARPRO_3P', 'TARQUIN' or 'LCMODEL'.
<code>w_ref</code>	water reference data for concentration scaling (optional).
<code>opts</code>	options to pass to the analysis method.
<code>parallel</code>	perform analyses in parallel (TRUE or FALSE).
<code>time</code>	measure the time taken for the analysis to complete (TRUE or FALSE).
<code>progress</code>	option is passed to plyr::alply function to display a progress bar during fitting. Default value is "text", set to "none" to disable.
<code>extra</code>	an optional data frame to provide additional variables for use in subsequent analysis steps, eg id or grouping variables.

Details

Fitting approaches described in the following references: ABfit Wilson, M. Adaptive baseline fitting for ^1H MR spectroscopy analysis. Magn Reson Med 2012;85:13-29.

VARPRO van der Veen JW, de Beer R, Luyten PR, van Ormondt D. Accurate quantification of in vivo ^{31}P NMR signals using the variable projection method and prior knowledge. Magn Reson Med 1988;6:92-98.

TARQUIN Wilson, M., Reynolds, G., Kauppinen, R. A., Arvanitis, T. N. & Peet, A. C. A constrained least-squares approach to the automated quantitation of in vivo ^1H magnetic resonance spectroscopy data. Magn Reson Med 2011;65:1-12.

LCModel Provencher SW. Estimation of metabolite concentrations from localized in vivo proton NMR spectra. Magn Reson Med 1993;30:672-679.

Value

MRS analysis object.

Examples

```
fname <- system.file("extdata", "philips_spar_sdat_WS.SDAT", package =
"spant")
svs <- read_mrs(fname)
## Not run:
basis <- sim_basis_1h_brain_press(svs)
fit_result <- fit_mrs(svs, basis)

## End(Not run)
```

fit_res2csv	<i>Write fit results table to a csv file.</i>
-------------	---

Description

Write fit results table to a csv file.

Usage

```
fit_res2csv(fit_res, fname, unscaled = FALSE)
```

Arguments

fit_res	fit result object.
fname	filename of csv file.
unscaled	output the unscaled result table (default = FALSE).

fp_phase	<i>Return the phase of the first data point in the time-domain.</i>
----------	---

Description

Return the phase of the first data point in the time-domain.

Usage

```
fp_phase(mrs_data)
```

Arguments

mrs_data	MRS data.
----------	-----------

Value

phase values in degrees.

<code>fp_phase_correct</code>	<i>Perform a zeroth order phase correction based on the phase of the first data point in the time-domain.</i>
-------------------------------	---

Description

Perform a zeroth order phase correction based on the phase of the first data point in the time-domain.

Usage

```
fp_phase_correct(mrs_data, ret_phase = FALSE)
```

Arguments

<code>mrs_data</code>	MRS data to be corrected.
<code>ret_phase</code>	return phase values (logical).

Value

corrected data or a list with corrected data and optional phase values.

<code>fp_scale</code>	<i>Scale the first time-domain data point in an mrs_data object.</i>
-----------------------	--

Description

Scale the first time-domain data point in an mrs_data object.

Usage

```
fp_scale(mrs_data, scale = 0.5)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>scale</code>	scaling value, defaults to 0.5.

Value

scaled mrs_data object.

fs

Return the sampling frequency in Hz of an MRS dataset.

Description

Return the sampling frequency in Hz of an MRS dataset.

Usage

```
fs(mrs_data)
```

Arguments

mrs_data MRS data.

Value

sampling frequency in Hz.

ft_dync

Apply the Fourier transform over the dynamic dimension.

Description

Apply the Fourier transform over the dynamic dimension.

Usage

```
ft_dync(mrs_data, ft_shift = FALSE, ret_mod = FALSE, fd = TRUE)
```

Arguments

mrs_data MRS data where the dynamic dimension is in the time-domain.
ft_shift apply FT shift to the output, default is FALSE.
ret_mod return the modulus out the transform, default is FALSE.
fd transform the chemical shift axis to the frequency domain first, default is TRUE.

Value

transformed MRS data.

ft_shift

Perform a fft and ffshift on a vector.

Description

Perform a fft and ffshift on a vector.

Usage

```
ft_shift(vec_in)
```

Arguments

vec_in vector input.

Value

output vector.

ft_shift_mat

Perform a fft and fftshift on a matrix with each column replaced by its shifted fft.

Description

Perform a fft and fftshift on a matrix with each column replaced by its shifted fft.

Usage

```
ft_shift_mat(mat_in)
```

Arguments

mat_in matrix input.

Value

output matrix.

gausswin_2d	<i>Create a two dimensional Gaussian window function stored as a matrix.</i>
-------------	--

Description

Create a two dimensional Gaussian window function stored as a matrix.

Usage

```
gausswin_2d(xN, yN, x0, y0, xw, yw)
```

Arguments

xN	number of pixels in the x dimension.
yN	number of pixels in the y dimension.
x0	centre of window function in the x direction in units of pixels. Note, only integer values are applied.
y0	centre of window function in the y direction in units of pixels. Note, only integer values are applied.
xw	the reciprocal of the standard deviation of the Gaussian window in x direction.
yw	the reciprocal of the standard deviation of the Gaussian window in y direction.

Value

matrix with dimensions `fov_yN x fov_xN`.

gen_F	<i>Generate the F product operator.</i>
-------	---

Description

Generate the F product operator.

Usage

```
gen_F(sys, op, detect = NULL)
```

Arguments

sys	spin system object.
op	operator, one of "x", "y", "z", "p", "m".
detect	detection nuclei.

Value

F product operator matrix.

`gen_F_xy`*Generate the Fxy product operator with a specified phase.***Description**

Generate the Fxy product operator with a specified phase.

Usage

```
gen_F_xy(sys, phase, detect = NULL)
```

Arguments

- | | |
|---------------------|-------------------------|
| <code>sys</code> | spin system object. |
| <code>phase</code> | phase angle in degrees. |
| <code>detect</code> | detection nuclei. |

Value

product operator matrix.

`get_1h_brain_basis_paras`*Return a list of mol_parameter objects suitable for 1H brain MRS analyses.***Description**

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Usage

```
get_1h_brain_basis_paras(ft, metab_lw = NULL, lcm_compat = FALSE)
```

Arguments

- | | |
|-------------------------|--|
| <code>ft</code> | transmitter frequency in Hz. |
| <code>metab_lw</code> | linewidth of metabolite signals (Hz). |
| <code>lcm_compat</code> | when TRUE, lipid, MM and -CrCH molecules will be excluded from the output. |

Value

list of mol_parameter objects.

```
get_1h_brain_basis_paras_v1
```

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Description

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Usage

```
get_1h_brain_basis_paras_v1(ft, metab_lw = NULL, lcm_compat = FALSE)
```

Arguments

ft	transmitter frequency in Hz.
metab_lw	linewidth of metabolite signals (Hz).
lcm_compat	when TRUE, lipid, MM and -CrCH molecules will be excluded from the output.

Value

list of mol_parameter objects.

```
get_1h_brain_basis_paras_v2
```

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Description

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Usage

```
get_1h_brain_basis_paras_v2(ft, metab_lw = NULL, lcm_compat = FALSE)
```

Arguments

ft	transmitter frequency in Hz.
metab_lw	linewidth of metabolite signals (Hz).
lcm_compat	when TRUE, lipid, MM and -CrCH molecules will be excluded from the output.

Value

list of mol_parameter objects.

get_1h_brain_basis_paras_v3

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Description

Return a list of mol_parameter objects suitable for 1H brain MRS analyses.

Usage

```
get_1h_brain_basis_paras_v3(ft, metab_lw = NULL, lcm_compat = FALSE)
```

Arguments

- | | |
|------------|--|
| ft | transmitter frequency in Hz. |
| metab_lw | linewidth of metabolite signals (Hz). |
| lcm_compat | when TRUE, lipid, MM and -CrCH molecules will be excluded from the output. |

Value

list of mol_parameter objects.

get_2d_psf

Get the point spread function (PSF) for a 2D phase encoded MRSI scan.

Description

Get the point spread function (PSF) for a 2D phase encoded MRSI scan.

Usage

```
get_2d_psf(
  FOV = 160,
  mat_size = 16,
  sampling = "circ",
  hamming = FALSE,
  ensure_odd = TRUE
)
```

Arguments

FOV	field of view in mm.
mat_size	acquisition matrix size (not interpolated).
sampling	can be either "circ" for circular or "rect" for rectangular.
hamming	should Hamming k-space weighting be applied (default FALSE).
ensure_odd	add 1mm to the FOV when required to ensure the output pdf has odd dimensions. Required when using get_mrsi2d_seg.

Value

A matrix of the PSF with 1mm resolution.

`get_acq_paras`

Return acquisition parameters from a MRS data object.

Description

Return acquisition parameters from a MRS data object.

Usage

```
get_acq_paras(mrs_data)
```

Arguments

mrs_data	MRS data.
----------	-----------

Value

list of acquisition parameters.

`get_dyncs`

Extract a subset of dynamic scans.

Description

Extract a subset of dynamic scans.

Usage

```
get_dyncs(mrs_data, subset)
```

Arguments

mrs_data	dynamic MRS data.
subset	vector containing indices to the dynamic scans to be returned.

Value

MRS data containing the subset of requested dynamics.

get_even_dyns	<i>Return even numbered dynamic scans starting from 1 (2,4,6...).</i>
---------------	---

Description

Return even numbered dynamic scans starting from 1 (2,4,6...).

Usage

```
get_even_dyns(mrs_data)
```

Arguments

mrs_data	dynamic MRS data.
----------	-------------------

Value

dynamic MRS data containing even numbered scans.

get_fh_dyns	<i>Return the first half of a dynamic series.</i>
-------------	---

Description

Return the first half of a dynamic series.

Usage

```
get_fh_dyns(mrs_data)
```

Arguments

mrs_data	dynamic MRS data.
----------	-------------------

Value

first half of the dynamic series.

get_fit_map	<i>Get a data array from a fit result.</i>
-------------	--

Description

Get a data array from a fit result.

Usage

```
get_fit_map(fit_res, name)
```

Arguments

fit_res	fit_result object.
name	name of the quantity to plot, eg "tNAA".

get_fp	<i>Return the first time-domain data point.</i>
--------	---

Description

Return the first time-domain data point.

Usage

```
get_fp(mrs_data)
```

Arguments

mrs_data	MRS data.
----------	-----------

Value

first time-domain data point.

get_gaussian_pulse *Generate a gaussian pulse shape.*

Description

Generate a gaussian pulse shape.

Usage

```
get_gaussian_pulse(angle, n, trunc = 1)
```

Arguments

angle	pulse angle in degrees.
n	number of points to generate.
trunc	percentage truncation factor.

get_head_dyns *Return the first scans of a dynamic series.*

Description

Return the first scans of a dynamic series.

Usage

```
get_head_dyns(mrs_data, n = 1)
```

Arguments

mrs_data	dynamic MRS data.
n	the number of dynamic scans to return.

Value

first scans of a dynamic series.

get_lcm_cmd	<i>Print the command to run the LCModel command-line program.</i>
-------------	---

Description

Print the command to run the LCModel command-line program.

Usage

```
get_lcm_cmd()
```

get_metab	<i>Extract the metabolite component from an mrs_data object.</i>
-----------	--

Description

Extract the metabolite component from an mrs_data object.

Usage

```
get_metab(mrs_data)
```

Arguments

mrs_data MRS data.

Value

metabolite component.

get_mol_names	<i>Return a character array of names that may be used with the get_mol_paras function.</i>
---------------	--

Description

Return a character array of names that may be used with the get_mol_paras function.

Usage

```
get_mol_names()
```

Value

a character array of names.

`get_mol_paras`*Get a mol_parameters object for a named molecule.***Description**

Get a `mol_parameters` object for a named molecule.

Usage

```
get_mol_paras(name, ...)
```

Arguments

- | | |
|-------------------|--|
| <code>name</code> | the name of the molecule. |
| <code>...</code> | arguments to pass to molecule definition function. |

`get_mrsi2d_seg`*Calculate the partial volume estimates for each voxel in a 2D MRSI dataset.***Description**

Localisation is assumed to be perfect in the z direction and determined by the `ker` input in the x-y direction.

Usage

```
get_mrsi2d_seg(mrs_data, mri_seg, ker)
```

Arguments

- | | |
|-----------------------|---|
| <code>mrs_data</code> | 2D MRSI data with multiple voxels in the x-y dimension. |
| <code>mri_seg</code> | MRI data with values corresponding to the segmentation class. Must be 1mm isotropic resolution. |
| <code>ker</code> | MRSI PSF kernel in the x-y direction compatible with the <code>mmnd</code> package, eg: <code>mmnd::shapeKernel(c(10, 10), type = "box")</code> . |

Value

a data frame of partial volume estimates and individual segmentation maps.

get_mrsi_voi	<i>Generate a MRSI VOI from an mrs_data object.</i>
--------------	---

Description

Generate a MRSI VOI from an mrs_data object.

Usage

```
get_mrsi_voi(mrs_data, target_mri = NULL, map = NULL, ker = mmand::boxKernel())
```

Arguments

mrs_data	MRS data.
target_mri	optional image data to match the intended volume space.
map	optional voi intensity map.
ker	kernel to rescale the map data to the target_mri. Default value is mmand::boxKernel(), use mmand::mnKernel() for a smoothed map.

Value

volume data as a nifti object.

get_mrsi_voxel	<i>Generate a MRSI voxel from an mrs_data object.</i>
----------------	---

Description

Generate a MRSI voxel from an mrs_data object.

Usage

```
get_mrsi_voxel(mrs_data, target_mri, x_pos, y_pos, z_pos)
```

Arguments

mrs_data	MRS data.
target_mri	optional image data to match the intended volume space.
x_pos	x voxel coordinate.
y_pos	y voxel coordinate.
z_pos	z voxel coordinate.

Value

volume data as a nifti object.

`get_mrsi_voxel_xy_psf` *Generate a MRSI voxel PSF from an mrs_data object.*

Description

Generate a MRSI voxel PSF from an `mrs_data` object.

Usage

```
get_mrsi_voxel_xy_psf(mrs_data, target_mri, x_pos, y_pos, z_pos)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>target_mri</code>	optional image data to match the intended volume space.
<code>x_pos</code>	x voxel coordinate.
<code>y_pos</code>	y voxel coordinate.
<code>z_pos</code>	z voxel coordinate.

Value

volume data as a nifti object.

`get_mrs_affine` *Generate an affine for nifti generation.*

Description

Generate an affine for nifti generation.

Usage

```
get_mrs_affine(mrs_data, x_pos = 1, y_pos = 1, z_pos = 1)
```

Arguments

<code>mrs_data</code>	input data.
<code>x_pos</code>	x_position coordinate.
<code>y_pos</code>	y_position coordinate.
<code>z_pos</code>	z_position coordinate.

Value

affine matrix.

get_odd_dync

Return odd numbered dynamic scans starting from 1 (1,3,5...).

Description

Return odd numbered dynamic scans starting from 1 (1,3,5...).

Usage

```
get_odd_dync(mrs_data)
```

Arguments

mrs_data dynamic MRS data.

Value

dynamic MRS data containing odd numbered scans.

get_ref

Extract the reference component from an mrs_data object.

Description

Extract the reference component from an mrs_data object.

Usage

```
get_ref(mrs_data)
```

Arguments

mrs_data MRS data.

Value

reference component.

`get_seg_ind`

Get the indices of data points lying between two values (end > x > start).

Description

Get the indices of data points lying between two values (end > x > start).

Usage

```
get_seg_ind(scale, start, end)
```

Arguments

`scale` full list of values.

`start` smallest value in the subset.

`end` largest value in the subset.

Value

set of indices.

`get_sh_dyns`

Return the second half of a dynamic series.

Description

Return the second half of a dynamic series.

Usage

```
get_sh_dyns(mrs_data)
```

Arguments

`mrs_data` dynamic MRS data.

Value

second half of the dynamic series.

get_slice*Return a single slice from a larger MRSI dataset.*

Description

Return a single slice from a larger MRSI dataset.

Usage

```
get_slice(mrs_data, z_pos)
```

Arguments

mrs_data	MRSI data.
z_pos	the z index to extract.

Value

MRS data.

get_subset*Extract a subset of MRS data.*

Description

Extract a subset of MRS data.

Usage

```
get_subset(  
    mrs_data,  
    x_set = NULL,  
    y_set = NULL,  
    z_set = NULL,  
    dyn_set = NULL,  
    coil_set = NULL,  
    fd_set = NULL,  
    td_set = NULL  
)
```

Arguments

<code>mrs_data</code>	MRS data object.
<code>x_set</code>	x indices to include in the output (default all).
<code>y_set</code>	y indices to include in the output (default all).
<code>z_set</code>	z indices to include in the output (default all).
<code>dyn_set</code>	dynamic indices to include in the output (default all).
<code>coil_set</code>	coil indices to include in the output (default all).
<code>fd_set</code>	frequency domain data indices to include in the output (default all).
<code>td_set</code>	time-domain indices to include in the output (default all).

Value

selected subset of MRS data.

`get_svs_voi`

Generate a SVS acquisition volume from an `mrs_data` object.

Description

Generate a SVS acquisition volume from an `mrs_data` object.

Usage

```
get_svs_voi(mrs_data, target_mri)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>target_mri</code>	optional image data to match the intended volume space.

Value

volume data as a nifti object.

get_tail_dyns	<i>Return the last scans of a dynamic series.</i>
---------------	---

Description

Return the last scans of a dynamic series.

Usage

```
get_tail_dyns(mrs_data, n = 1)
```

Arguments

mrs_data	dynamic MRS data.
n	the number of dynamic scans to return.

Value

last scans of a dynamic series.

get_td_amp	<i>Return an array of amplitudes derived from fitting the initial points in the time domain and extrapolating back to t=0.</i>
------------	--

Description

Return an array of amplitudes derived from fitting the initial points in the time domain and extrapolating back to t=0.

Usage

```
get_td_amp(mrs_data, nstart = 10, nend = 50, method = "spline")
```

Arguments

mrs_data	MRS data.
nstart	first data point to fit.
nend	last data point to fit.
method	method for measuring the amplitude, one of "spline" or "exp".

Value

array of amplitudes.

`get_tqn_cmd`*Print the command to run the TARQUIN command-line program.***Description**

Print the command to run the TARQUIN command-line program.

Usage

```
get_tqn_cmd()
```

`get_uncoupled_mol`*Generate a mol_parameters object for a simple spin system with one resonance.***Description**

Generate a `mol_parameters` object for a simple spin system with one resonance.

Usage

```
get_uncoupled_mol(  
    name,  
    chem_shift,  
    nucleus,  
    scale_factor,  
    lw,  
    lg,  
    full_name = NULL  
)
```

Arguments

<code>name</code>	abbreviated name of the molecule.
<code>chem_shift</code>	chemical shift of the resonance (PPM).
<code>nucleus</code>	nucleus (1H, 31P...).
<code>scale_factor</code>	multiplicative scaling factor.
<code>lw</code>	linewidth in Hz.
<code>lg</code>	Lorentz-Gauss lineshape parameter (between 0 and 1).
<code>full_name</code>	long name of the molecule (optional).

Value

`mol_parameters` object.

get_voi_cog

Calculate the centre of gravity for an image containing 0 and 1's.

Description

Calculate the centre of gravity for an image containing 0 and 1's.

Usage

```
get_voi_cog(voi)
```

Arguments

voi nifti object.

Value

triplet of x,y,z coordinates.

get_voi_seg

Return the white matter, gray matter and CSF composition of a volume.

Description

Return the white matter, gray matter and CSF composition of a volume.

Usage

```
get_voi_seg(voi, mri_seg)
```

Arguments

voi volume data as a nifti object.
mri_seg segmented brain volume as a nifti object.

Value

a vector of partial volumes expressed as percentages.

get_voi_seg_psf	<i>Return the white matter, gray matter and CSF composition of a volume.</i>
-----------------	--

Description

Return the white matter, gray matter and CSF composition of a volume.

Usage

```
get_voi_seg_psf(psf, mri_seg)
```

Arguments

psf	volume data as a nifti object.
mri_seg	segmented brain volume as a nifti object.

Value

a vector of partial volumes expressed as percentages.

get_voxel	<i>Return a single voxel from a larger mrs dataset.</i>
-----------	---

Description

Return a single voxel from a larger mrs dataset.

Usage

```
get_voxel(mrs_data, x_pos = 1, y_pos = 1, z_pos = 1, dyn = 1, coil = 1)
```

Arguments

mrs_data	MRS data.
x_pos	the x index to plot.
y_pos	the y index to plot.
z_pos	the z index to plot.
dyn	the dynamic index to plot.
coil	the coil element number to plot.

Value

MRS data.

gridplot	<i>Arrange spectral plots in a grid.</i>
----------	--

Description

Arrange spectral plots in a grid.

Usage

```
gridplot(x, ...)
```

Arguments

x	object for plotting.
...	arguments to be passed to methods.

gridplot.mrs_data	<i>Arrange spectral plots in a grid.</i>
-------------------	--

Description

Arrange spectral plots in a grid.

Usage

```
## S3 method for class 'mrs_data'  
gridplot(  
  x,  
  rows = NA,  
  cols = NA,  
  mar = c(0, 0, 0, 0),  
  oma = c(3.5, 1, 1, 1),  
  bty = "o",  
  restore_def_par = TRUE,  
  ...  
)
```

Arguments

x	object of class mrs_data.
rows	number of grid rows.
cols	number of grid columns.
mar	option to adjust the plot margins. See ?par.
oma	outer margin area.

bty	option to draw a box around the plot. See ?par.
restore_def_par	restore default plotting par values after the plot has been made.
...	other arguments to pass to the plot method.

grid_shift_xy	<i>Grid shift MRSI data in the x/y dimension.</i>
---------------	---

Description

Grid shift MRSI data in the x/y dimension.

Usage

```
grid_shift_xy(mrs_data, x_shift, y_shift)
```

Arguments

mrs_data	MRSI data in the spatial domain.
x_shift	shift to apply in the x-direction in units of voxels.
y_shift	shift to apply in the y-direction in units of voxels.

Value

shifted data.

hsvd	<i>HSVD of an mrs_data object.</i>
------	------------------------------------

Description

HSVD method as described in: Barkhuijsen H, de Beer R, van Ormondt D. Improved algorithm for noniterative and timedomain model fitting to exponentially damped magnetic resonance signals. J Magn Reson 1987;73:553-557.

Usage

```
hsvd(mrs_data, comps = 40, irlba = TRUE, max_damp = 10)
```

Arguments

mrs_data	mrs_data object to be decomposed.
comps	number of Lorentzian components to use for modelling.
irlba	option to use irlba SVD (logical).
max_damp	maximum allowable damping factor.

Value

basis matrix and signal table.

hsvd_filt	<i>HSVD based signal filter.</i>
-----------	----------------------------------

Description

HSVD based signal filter described in: Barkhuijsen H, de Beer R, van Ormondt D. Improved algorithm for noniterative and timedomain model fitting to exponentially damped magnetic resonance signals. J Magn Reson 1987;73:553-557.

Usage

```
hsvd_filt(  
  mrs_data,  
  xlim = c(-30, 30),  
  comps = 40,  
  irlba = TRUE,  
  max_damp = 10,  
  scale = "hz",  
  return_model = FALSE  
)
```

Arguments

mrs_data	MRS data to be filtered.
xlim	frequency range to filter, default units are Hz which can be changed to ppm using the "scale" argument.
comps	number of Lorentzian components to use for modelling.
irlba	option to use irlba SVD (logical).
max_damp	maximum allowable damping factor.
scale	either "hz" or "ppm" to set the frequency units of xlim.
return_model	by default the filtered spectrum is returned. Set return_model to TRUE to return the HSVD model of the data.

Value

filtered data or model depending on the return_model argument.

<code>hsvd_vec</code>	<i>HSVD of a complex vector.</i>
-----------------------	----------------------------------

Description

HSVD method as described in: Barkhuijsen H, de Beer R, van Ormondt D. Improved algorithm for noniterative and timedomain model fitting to exponentially damped magnetic resonance signals. J Magn Reson 1987;73:553-557.

Usage

```
hsvd_vec(y, fs, comps = 40, irlba = TRUE, max_damp = 0)
```

Arguments

<code>y</code>	time domain signal to be filtered as a vector.
<code>fs</code>	sampling frequency of <code>y</code> .
<code>comps</code>	number of Lorentzian components to use for modelling.
<code>irlba</code>	option to use irlba SVD (logical).
<code>max_damp</code>	maximum allowable damping factor. Default value of 0 ensures resultant model is damped.

Value

basis matrix and signal table.

<code>hz</code>	<i>Return the frequency scale of an MRS dataset in Hz.</i>
-----------------	--

Description

Return the frequency scale of an MRS dataset in Hz.

Usage

```
hz(mrs_data, fs = NULL, N = NULL)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>fs</code>	sampling frequency in Hz.
<code>N</code>	number of data points in the spectral dimension.

Value

frequency scale.

ift_shift	<i>Perform an ifffshift and ifft on a vector.</i>
-----------	---

Description

Perform an ifffshift and ifft on a vector.

Usage

```
ift_shift(vec_in)
```

Arguments

vec_in vector input.

Value

output vector.

ift_shift_mat	<i>Perform an ifft and ifftshift on a matrix with each column replaced by its shifted ifft.</i>
---------------	---

Description

Perform an ifft and ifftshift on a matrix with each column replaced by its shifted ifft.

Usage

```
ift_shift_mat(mat_in)
```

Arguments

mat_in matrix input.

Value

output matrix.

Im.mrs_data*Apply Im operator to an MRS dataset.***Description**

Apply Im operator to an MRS dataset.

Usage

```
## S3 method for class 'mrs_data'
Im(z)
```

Arguments

z MRS data.

Value

MRS data following Im operator.

image.mrs_data*Image plot method for objects of class mrs_data.***Description**

Image plot method for objects of class mrs_data.

Usage

```
## S3 method for class 'mrs_data'
image(
  x,
  xlim = NULL,
  mode = "re",
  col = NULL,
  plot_dim = NULL,
  x_pos = NULL,
  y_pos = NULL,
  z_pos = NULL,
  dyn = 1,
  coil = 1,
  restore_def_par = TRUE,
  y_ticks = NULL,
  vline = NULL,
  hline = NULL,
  ...
)
```

Arguments

x	object of class mrs_data.
xlim	the range of values to display on the x-axis, eg xlim = c(4,1).
mode	representation of the complex numbers to be plotted, can be one of: "re", "im", "mod" or "arg".
col	Colour map to use, defaults to viridis.
plot_dim	the dimension to display on the y-axis, can be one of: "dyn", "x", "y", "z", "coil" or NULL. If NULL (the default) all spectra will be collapsed into the dynamic dimension and displayed.
x_pos	the x index to plot.
y_pos	the y index to plot.
z_pos	the z index to plot.
dyn	the dynamic index to plot.
coil	the coil element number to plot.
restore_def_par	restore default plotting par values after the plot has been made.
y_ticks	a vector of indices specifying where to place tick marks.
vline	draw a vertical line at the value of vline.
hline	draw a horizontal line at the value of hline.
...	other arguments to pass to the plot method.

img2kspace_xy

*Transform 2D MRSI data to k-space in the x-y direction.***Description**

Transform 2D MRSI data to k-space in the x-y direction.

Usage

```
img2kspace_xy(mrs_data)
```

Arguments

mrs_data	2D MRSI data.
----------	---------------

Value

k-space data.

interleave_dync *Interleave the first and second half of a dynamic series.*

Description

Interleave the first and second half of a dynamic series.

Usage

```
interleave_dync(mrs_data)
```

Arguments

mrs_data dynamic MRS data.

Value

interleaved data.

int_spec *Integrate a spectral region.*

Description

See spec_op function for a more complete set of spectral operations.

Usage

```
int_spec(mrs_data, xlim = NULL, freq_scale = "ppm", mode = "re")
```

Arguments

mrs_data MRS data.

xlim spectral range to be integrated (defaults to full range).

freq_scale units of xlim, can be : "ppm", "hz" or "points".

mode spectral mode, can be : "re", "im", "mod" or "cplx".

Value

an array of integral values.

inv_even_dyns *Invert even numbered dynamic scans starting from 1 (2,4,6...).*

Description

Invert even numbered dynamic scans starting from 1 (2,4,6...).

Usage

```
inv_even_dyns(mrs_data)
```

Arguments

mrs_data dynamic MRS data.

Value

dynamic MRS data with inverted even numbered scans.

inv_odd_dyns *Invert odd numbered dynamic scans starting from 1 (1,3,5...).*

Description

Invert odd numbered dynamic scans starting from 1 (1,3,5...).

Usage

```
inv_odd_dyns(mrs_data)
```

Arguments

mrs_data dynamic MRS data.

Value

dynamic MRS data with inverted odd numbered scans.

is.def

Check if an object is defined, which is the same as being not NULL.

Description

Check if an object is defined, which is the same as being not NULL.

Usage

```
is.def(x)
```

Arguments

x object to test for being NULL.

Value

logical value.

is_fd

Check if the chemical shift dimension of an MRS data object is in the frequency domain.

Description

Check if the chemical shift dimension of an MRS data object is in the frequency domain.

Usage

```
is_fd(mrs_data)
```

Arguments

mrs_data MRS data.

Value

logical value.

kspace2img_xy	<i>Transform 2D MRSI data from k-space to image space in the x-y direction.</i>
---------------	---

Description

Transform 2D MRSI data from k-space to image space in the x-y direction.

Usage

```
kspace2img_xy(mrs_data)
```

Arguments

mrs_data 2D MRSI data.

Value

MRSI data in image space.

l2_reg	<i>Perform l2 regularisation artefact suppression.</i>
--------	--

Description

Perform l2 regularisation artefact suppression using the method proposed by Bilgic et al. JMRI 40(1):181-91 2014.

Usage

```
l2_reg(  
  mrs_data,  
  thresh = 0.05,  
  b = 1e-11,  
  A = NA,  
  xlim = NA,  
  thresh_xlim = NULL,  
  A_append = NULL,  
  ret_norms = FALSE  
)
```

Arguments

<code>mrs_data</code>	input data for artefact suppression.
<code>thresh</code>	threshold parameter to extract lipid signals from <code>mrs_data</code> based on the spectral integration of the <code>thresh_xlim</code> region in magnitude mode.
<code>b</code>	regularisation parameter.
<code>A</code>	set of spectra containing the artefact basis signals. The <code>thresh</code> parameter is ignored when <code>A</code> is specified.
<code>xlim</code>	spectral limits in ppm to restrict the reconstruction range. Defaults to the full spectral width.
<code>thresh_xlim</code>	spectral limits in ppm to integrate for the threshold map.
<code>A_append</code>	additional spectra to append to the <code>A</code> basis.
<code>ret_norms</code>	return the residual norm and solution norms.

Value

`l2` reconstructed `mrs_data` object.

`lb`

Apply line-broadening (apodisation) to MRS data or basis object.

Description

Apply line-broadening (apodisation) to MRS data or basis object.

Usage

```
lb(x, lb, lg = 1)

## S3 method for class 'list'
lb(x, lb, lg = 1)

## S3 method for class 'mrs_data'
lb(x, lb, lg = 1)

## S3 method for class 'basis_set'
lb(x, lb, lg = 1)
```

Arguments

<code>x</code>	input <code>mrs_data</code> or <code>basis_set</code> object.
<code>lb</code>	amount of line-broadening in Hz.
<code>lg</code>	Lorentz-Gauss lineshape parameter (between 0 and 1).

Value

line-broadened data.

lw2alpha	<i>Convert a linewidth in Hz to an equivalent alpha value in the time-domain ie: $x * \exp(-t * \alpha)$.</i>
----------	--

Description

Covert a linewidth in Hz to an equivalent alpha value in the time-domain ie: $x * \exp(-t * \alpha)$.

Usage

```
lw2alpha(lw)
```

Arguments

lw linewidth in Hz.

Value

beta damping value.

lw2beta	<i>Convert a linewidth in Hz to an equivalent beta value in the time-domain ie: $x * \exp(-t * t * \beta)$.</i>
---------	--

Description

Covert a linewidth in Hz to an equivalent beta value in the time-domain ie: $x * \exp(-t * t * \beta)$.

Usage

```
lw2beta(lw)
```

Arguments

lw linewidth in Hz.

Value

beta damping value.

mask_dyncs*Mask an MRS dataset in the dynamic dimension.***Description**

Mask an MRS dataset in the dynamic dimension.

Usage

```
mask_dyncs(mrs_data, mask)
```

Arguments

- | | |
|-----------------------|---|
| <code>mrs_data</code> | MRS data object. |
| <code>mask</code> | vector of boolean values specifying the dynamics to mask, where a value of TRUE indicates the spectrum should be removed. |

Value

masked dataset.

mask_fit_res*Mask fit result spectra depending on a vector of bool values.***Description**

Mask fit result spectra depending on a vector of bool values.

Usage

```
mask_fit_res(fit_result, mask_vec, amps_only = FALSE)
```

Arguments

- | | |
|-------------------------|---|
| <code>fit_result</code> | fit result object to be masked. |
| <code>mask_vec</code> | a Boolean vector with the same number of rows as there are rows in the results table. |
| <code>amps_only</code> | only mask the amplitude and associated error estimate columns. |

Value

a masked fit result object.

mask_xy	<i>Mask an MRSI dataset in the x-y direction</i>
---------	--

Description

Mask an MRSI dataset in the x-y direction

Usage

```
mask_xy(mrs_data, x_dim, y_dim)
```

Arguments

mrs_data	MRS data object.
x_dim	x dimension output length.
y_dim	y dimension output length.

Value

masked MRS data.

mask_xy_mat	<i>Mask a 2D MRSI dataset in the x-y dimension.</i>
-------------	---

Description

Mask a 2D MRSI dataset in the x-y dimension.

Usage

```
mask_xy_mat(mrs_data, mask, value = NA)
```

Arguments

mrs_data	MRS data object.
mask	matrix of boolean values specifying the voxels to mask, where a value of TRUE indicates the voxel should be removed.
value	the value to set masked data to (usually NA or 0).

Value

masked dataset.

<code>mat2mrs_data</code>	<i>Convert a matrix (with spectral points in the column dimension and dynamics in the row dimensions) into a mrs_data object.</i>
---------------------------	---

Description

Convert a matrix (with spectral points in the column dimension and dynamics in the row dimensions) into a mrs_data object.

Usage

```
mat2mrs_data(
  mat,
  fs = def_fs(),
  ft = def_ft(),
  ref = def_ref(),
  nuc = def_nuc(),
  fd = FALSE
)
```

Arguments

<code>mat</code>	data matrix.
<code>fs</code>	sampling frequency in Hz.
<code>ft</code>	transmitter frequency in Hz.
<code>ref</code>	reference value for ppm scale.
<code>nuc</code>	resonant nucleus.
<code>fd</code>	flag to indicate if the matrix is in the frequency domain (logical).

Value

mrs_data object.

<code>max_mrs</code>	<i>Apply the max operator to an MRS dataset.</i>
----------------------	--

Description

Apply the max operator to an MRS dataset.

Usage

```
max_mrs(mrs_data)
```

Arguments

mrs_data MRS data.

Value

MRS data following max operator.

max_mrs_interp *Apply the max operator to an interpolated MRS dataset.*

Description

Apply the max operator to an interpolated MRS dataset.

Usage

max_mrs_interp(mrs_data, interp_f = 4)

Arguments

mrs_data MRS data.
interp_f interpolation factor.

Value

Array of maximum values (real only).

mean.list *Calculate the mean spectrum from an mrs_data object.*

Description

Calculate the mean spectrum from an mrs_data object.

Usage

```
## S3 method for class 'list'  
mean(x, ...)
```

Arguments

x object of class mrs_data.
... other arguments to pass to the colMeans function.

Value

mean mrs_data object.

`mean.mrs_data` *Calculate the mean spectrum from an mrs_data object.*

Description

Calculate the mean spectrum from an mrs_data object.

Usage

```
## S3 method for class 'mrs_data'  
mean(x, ...)
```

Arguments

`x` object of class mrs_data.
`...` other arguments to pass to the colMeans function.

Value

mean mrs_data object.

`mean_dync` *Calculate the mean dynamic data.*

Description

Calculate the mean dynamic data.

Usage

```
mean_dync(mrs_data)
```

Arguments

`mrs_data` dynamic MRS data.

Value

mean dynamic data.

mean_dyn_blocks	<i>Calculate the mean of adjacent dynamic scans.</i>
-----------------	--

Description

Calculate the mean of adjacent dynamic scans.

Usage

```
mean_dyn_blocks(mrs_data, block_size)
```

Arguments

mrs_data	dynamic MRS data.
block_size	number of adjacent dynamics scans to average over.

Value

dynamic data averaged in blocks.

mean_dyn_pairs	<i>Calculate the pairwise means across a dynamic data set.</i>
----------------	--

Description

Calculate the pairwise means across a dynamic data set.

Usage

```
mean_dyn_pairs(mrs_data)
```

Arguments

mrs_data	dynamic MRS data.
----------	-------------------

Value

mean dynamic data of adjacent dynamic pairs.

`mean_mrs_list` *Return the mean of a list of mrs_data objects.*

Description

Return the mean of a list of mrs_data objects.

Usage

```
mean_mrs_list(mrs_list)
```

Arguments

`mrs_list` list of mrs_data objects.

Value

mean mrs_data object.

`median_dyns` *Calculate the median dynamic data.*

Description

Calculate the median dynamic data.

Usage

```
median_dyns(mrs_data)
```

Arguments

`mrs_data` dynamic MRS data.

Value

median dynamic data.

Mod.mrs_data	<i>Apply Mod operator to an MRS dataset.</i>
--------------	--

Description

Apply Mod operator to an MRS dataset.

Usage

```
## S3 method for class 'mrs_data'  
Mod(z)
```

Arguments

z MRS data.

Value

MRS data following Mod operator.

mod_td	<i>Apply the Modulus operator to the time-domain MRS signal.</i>
--------	--

Description

Apply the Modulus operator to the time-domain MRS signal.

Usage

```
mod_td(mrs_data)
```

Arguments

mrs_data MRS data input.

Value

time-domain modulus of input.

<code>mrs_data2basis</code>	<i>Convert an mrs_data object to basis object - where basis signals are spread across the dynamic dimension in the MRS data.</i>
-----------------------------	--

Description

Convert an mrs_data object to basis object - where basis signals are spread across the dynamic dimension in the MRS data.

Usage

```
mrs_data2basis(mrs_data, names)
```

Arguments

<code>mrs_data</code>	mrs_data object with basis signals spread across the dynamic dimension.
<code>names</code>	list of names corresponding to basis signals.

Value

basis set object.

<code>mrs_data2mat</code>	<i>Convert mrs_data object to a matrix, with spectral points in the column dimension and dynamics in the row dimension.</i>
---------------------------	---

Description

Convert mrs_data object to a matrix, with spectral points in the column dimension and dynamics in the row dimension.

Usage

```
mrs_data2mat(mrs_data, collapse = TRUE)
```

Arguments

<code>mrs_data</code>	MRS data object or list of MRS data objects.
<code>collapse</code>	collapse all other dimensions along the dynamic dimension, eg a 16x16 MRSI grid would be first collapsed across 256 dynamic scans.

Value

MRS data matrix.

mrs_data2vec*Convert mrs_data object to a vector.*

Description

Convert mrs_data object to a vector.

Usage

```
mrs_data2vec(mrs_data, dyn = 1, x_pos = 1, y_pos = 1, z_pos = 1, coil = 1)
```

Arguments

mrs_data	MRS data object.
dyn	dynamic index.
x_pos	x index.
y_pos	y index.
z_pos	z index.
coil	coil element index.

Value

MRS data vector.

mvfftshift*Perform a fftshift on a matrix, with each column replaced by its shifted result.*

Description

Perform a fftshift on a matrix, with each column replaced by its shifted result.

Usage

```
mvfftshift(x)
```

Arguments

x	matrix input.
---	---------------

Value

output matrix.

<code>mvifftshift</code>	<i>Perform an ifftshift on a matrix, with each column replaced by its shifted result.</i>
--------------------------	---

Description

Perform an ifftshift on a matrix, with each column replaced by its shifted result.

Usage

```
mvifftshift(x)
```

Arguments

`x` matrix input.

Value

output matrix.

<code>n2coord</code>	<i>Print fit coordinates from a single index.</i>
----------------------	---

Description

Print fit coordinates from a single index.

Usage

```
n2coord(n, fit_res)
```

Arguments

`n` fit index.
`fit_res` fit_result object.

Ncoils

Return the total number of coil elements in an MRS dataset.

Description

Return the total number of coil elements in an MRS dataset.

Usage

```
Ncoils(mrs_data)
```

Arguments

mrs_data MRS data.

Ndyncs

Return the total number of dynamic scans in an MRS dataset.

Description

Return the total number of dynamic scans in an MRS dataset.

Usage

```
Ndyncs(mrs_data)
```

Arguments

mrs_data MRS data.

nifti_flip_lr

Flip the x data dimension order of a nifti image. This corresponds to flipping MRI data in the left-right direction, assuming the data is save in neurological format (can check with fslorient program).

Description

Flip the x data dimension order of a nifti image. This corresponds to flipping MRI data in the left-right direction, assuming the data is save in neurological format (can check with fslorient program).

Usage

```
nifti_flip_lr(x)
```

Arguments

- x nifti object to be processed.

Value

nifti object with reversed x data direction.

Npts

Return the number of data points in an MRS dataset.

Description

Return the number of data points in an MRS dataset.

Usage

```
Npts(mrs_data)
```

Arguments

- | | |
|----------|-----------|
| mrs_data | MRS data. |
|----------|-----------|

Value

number of data points.

Nspec

Return the total number of spectra in an MRS dataset.

Description

Return the total number of spectra in an MRS dataset.

Usage

```
Nspec(mrs_data)
```

Arguments

- | | |
|----------|-----------|
| mrs_data | MRS data. |
|----------|-----------|

Nx

Return the total number of x locations in an MRS dataset.

Description

Return the total number of x locations in an MRS dataset.

Usage

`Nx(mrs_data)`

Arguments

`mrs_data` MRS data.

Ny

Return the total number of y locations in an MRS dataset.

Description

Return the total number of y locations in an MRS dataset.

Usage

`Ny(mrs_data)`

Arguments

`mrs_data` MRS data.

Nz

Return the total number of z locations in an MRS dataset.

Description

Return the total number of z locations in an MRS dataset.

Usage

`Nz(mrs_data)`

Arguments

`mrs_data` MRS data.

ortho3*Display an orthographic projection plot of a nifti object.*

Description

Display an orthographic projection plot of a nifti object.

Usage

```
ortho3(
  underlay,
  overlay = NULL,
  xyz = NULL,
  zlim = NULL,
  zlim_ol = NULL,
  alpha = 0.7,
  col_ol = viridisLite::viridis(64),
  orient_lab = TRUE,
  rescale = 1,
  crosshairs = TRUE,
  ch_lwd = 1,
  colourbar = TRUE,
  bg = "black",
  mar = c(0, 0, 0, 0),
  smallplot = c(0.63, 0.65, 0.07, 0.42)
)
```

Arguments

underlay	underlay image to be shown in grayscale.
overlay	optional overlay image.
xyz	x, y, z slice coordinates to display.
zlim	underlay intensity limits.
zlim_ol	overlay intensity limits.
alpha	transparency of overlay.
col_ol	colour palette of overlay.
orient_lab	display orientation labels (default TRUE).
rescale	rescale factor for the underlay and overlay images.
crosshairs	display the crosshairs (default TRUE).
ch_lwd	crosshair linewidth.
colourbar	display a colourbar for the overlay (default TRUE).
bg	plot background colour.
mar	plot margins.
smallplot	smallplot option for positioning the colourbar.

ortho3_inter	<i>Display an interactive orthographic projection plot of a nifti object.</i>
--------------	---

Description

Display an interactive orthographic projection plot of a nifti object.

Usage

```
ortho3_inter(  
    underlay,  
    overlay = NULL,  
    xyz = NULL,  
    zlim = NULL,  
    zlim_ol = NULL,  
    alpha = 0.7,  
    ...  
)
```

Arguments

underlay	underlay image to be shown in grayscale.
overlay	optional overlay image.
xyz	x, y, z slice coordinates to display.
zlim	underlay intensity limits.
zlim_ol	overlay intensity limits.
alpha	transparency of overlay.
...	other options to be passed to the ortho3 function.

peak_info	<i>Search for the highest peak in a spectral region and return the frequency, height and FWHM.</i>
-----------	--

Description

Search for the highest peak in a spectral region and return the frequency, height and FWHM.

Usage

```
peak_info(  
    mrs_data,  
    xlim = c(4, 0.5),  
    interp_f = 4,  
    scale = "ppm",  
    mode = "real"  
)
```

Arguments

<code>mrs_data</code>	an object of class <code>mrs_data</code> .
<code>xlim</code>	frequency range (default units of PPM) to search for the highest peak.
<code>interp_f</code>	interpolation factor, defaults to 4x.
<code>scale</code>	the units to use for the frequency scale, can be one of: "ppm", "hz" or "points".
<code>mode</code>	spectral mode, can be : "real", "imag" or "mod".

Value

list of arrays containing the highest peak frequency, height and FWHM in units of PPM and Hz.

`pg_extrap_xy`

Papoulis-Gerchberg (PG) algorithm method for k-space extrapolation.

Description

PG method as described in: Haupt CI, Schuff N, Weiner MW, Maudsley AA. Removal of lipid artifacts in 1H spectroscopic imaging by data extrapolation. Magn Reson Med. 1996 May;35(5):678-87. Extrapolation is performed to expand k-space coverage by a factor of 2, with the aim to reduce Gibbs ringing.

Usage

```
pg_extrap_xy(
  mrs_data,
  img_mask = NULL,
  kspace_mask = NULL,
  intensity_thresh = 0.15,
  iters = 50
)
```

Arguments

<code>mrs_data</code>	MRS data object.
<code>img_mask</code>	a boolean matrix of voxels with strong signals to be extrapolated. Must be twice the dimensions of the input data.
<code>kspace_mask</code>	a boolean matrix of kspace points that have been sampled. Typically a circle for MRSI, but defaults to the full rectangular area of k-space covered by the input data. Must match the x-y dimensions of the input data.
<code>intensity_thresh</code>	used to define <code>img_mask</code> based on the strength of the signal in each voxel. Defaults to intensities greater than 15% of the maximum. Ignored if <code>img_mask</code> is specified as argument.
<code>iters</code>	number of iterations to perform.

Value

extrapolated mrs_data object.

phase	<i>Apply phasing parameters to MRS data.</i>
-------	--

Description

Apply phasing parameters to MRS data.

Usage

```
phase(mrs_data, zero_order, first_order = 0)
```

Arguments

mrs_data	MRS data.
zero_order	zero'th order phase term in degrees.
first_order	first order (frequency dependent) phase term in ms.

Value

MRS data with applied phase parameters.

plot.fit_result	<i>Plot the fitting results of an object of class fit_result.</i>
-----------------	---

Description

Plot the fitting results of an object of class fit_result.

Usage

```
## S3 method for class 'fit_result'  
plot(  
  x,  
  dyn = 1,  
  x_pos = 1,  
  y_pos = 1,  
  z_pos = 1,  
  coil = 1,  
  xlim = NULL,  
  data_only = FALSE,  
  label = NULL,  
  plot_sigs = NULL,
```

```

n = NULL,
sub_b1 = FALSE,
mar = NULL,
restore_def_par = TRUE,
ylim = NULL,
y_scale = FALSE,
show_grid = TRUE,
grid_nx = NULL,
grid_ny = NA,
...
)

```

Arguments

x	fit_result object.
dyn	the dynamic index to plot.
x_pos	the x index to plot.
y_pos	the y index to plot.
z_pos	the z index to plot.
coil	the coil element number to plot.
xlim	the range of values to display on the x-axis, eg xlim = c(4,1).
data_only	display only the processed data (logical).
label	character string to add to the top left of the plot window.
plot_sigs	a character vector of signal names to add to the plot.
n	single index element to plot (overrides other indices when given).
sub_b1	subtract the baseline from the data and fit (logical).
mar	option to adjust the plot margins. See ?par.
restore_def_par	restore default plotting par values after the plot has been made.
ylim	range of values to display on the y-axis, eg ylim = c(0,10).
y_scale	option to display the y-axis values (logical).
show_grid	plot gridlines behind the data (logical). Defaults to TRUE.
grid_nx	number of cells of the grid in x and y direction. When NULL the grid aligns with the tick marks on the corresponding default axis (i.e., tickmarks as computed by axTicks). When NA, no grid lines are drawn in the corresponding direction.
grid_ny	as above.
...	further arguments to plot method.

plot.mrs_data *Plotting method for objects of class mrs_data.*

Description

Plotting method for objects of class mrs_data.

Usage

```
## S3 method for class 'mrs_data'  
plot(  
  x,  
  dyn = 1,  
  x_pos = 1,  
  y_pos = 1,  
  z_pos = 1,  
  coil = 1,  
  fd = TRUE,  
  x_units = NULL,  
  xlim = NULL,  
  y_scale = FALSE,  
  x_ax = TRUE,  
  mode = "re",  
  lwd = NULL,  
  bty = NULL,  
  label = "",  
  restore_def_par = TRUE,  
  mar = NULL,  
  xaxis_lab = NULL,  
  yaxis_lab = NULL,  
  xat = NULL,  
  xlabs = TRUE,  
  yat = NULL,  
  ylabs = TRUE,  
  show_grid = TRUE,  
  grid_nx = NULL,  
  grid_ny = NA,  
  col = NULL,  
  alpha = NULL,  
  bl_lty = NULL,  
  ...  
)
```

Arguments

x	object of class mrs_data.
dyn	the dynamic index to plot.

x_pos	the x index to plot.
y_pos	the y index to plot.
z_pos	the z index to plot.
coil	the coil element number to plot.
fd	display data in the frequency-domain (default), or time-domain (logical).
x_units	the units to use for the x-axis, can be one of: "ppm", "hz", "points" or "seconds".
xlim	the range of values to display on the x-axis, eg <code>xlim = c(4,1)</code> .
y_scale	option to display the y-axis values (logical).
x_ax	option to display the x-axis values (logical).
mode	representation of the complex numbers to be plotted, can be one of: "re", "im", "mod" or "arg".
lwd	plot linewidth.
bty	option to draw a box around the plot. See <code>?par</code> .
label	character string to add to the top left of the plot window.
restore_def_par	restore default plotting par values after the plot has been made.
mar	option to adjust the plot margins. See <code>?par</code> .
xaxis_lab	x-axis label.
yaxis_lab	y-axis label.
xat	x-axis tick label values.
xlabs	x-axis tick labels.
yat	y-axis tick label values.
ylabs	y-axis tick labels.
show_grid	plot gridlines behind the data (logical). Defaults to TRUE.
grid_nx	number of cells of the grid in x and y direction. When NULL the grid aligns with the tick marks on the corresponding default axis (i.e., tickmarks as computed by <code>axTicks</code>). When NA, no grid lines are drawn in the corresponding direction.
grid_ny	as above.
col	set the line colour, eg <code>col = rgb(0.5, 0.5, 0.5)</code> .
alpha	set the line transparency, eg <code>alpha = 0.5</code> is 50% transparency. Overrides any transparency levels set by col.
bl_lty	linetype for the $y = 0$ baseline trace. A default value NULL results in no baseline being plotted.
...	other arguments to pass to the plot method.

plot_bc	<i>Convenience function to plot a baseline estimate with the original data.</i>
---------	---

Description

Convenience function to plot a baseline estimate with the original data.

Usage

```
plot_bc(orig_data, bc_data, ...)
```

Arguments

orig_data	the original data.
bc_data	the baseline corrected data.
...	other arguments to pass to the stackplot function.

plot_slice_fit	<i>Plot a 2D slice from an MRSI fit result object.</i>
----------------	--

Description

Plot a 2D slice from an MRSI fit result object.

Usage

```
plot_slice_fit(  
  fit_res,  
  map,  
  map_denom = NULL,  
  slice = 1,  
  zlim = NULL,  
  interp = 1  
)
```

Arguments

fit_res	fit_result object.
map	fit result values to display as a colour map. Can be specified as a character string or array of numeric values. Defaults to "tNAA".
map_denom	fit result values to divide the map argument by. Can be specified as a character string (eg "tCr") or array of numeric values.
slice	slice to plot in the z direction.
zlim	range of values to plot.
interp	interpolation factor.

`plot_slice_fit_inter` *Plot a 2D slice from an MRSI fit result object.*

Description

Plot a 2D slice from an MRSI fit result object.

Usage

```
plot_slice_fit_inter(
  fit_res,
  map = NULL,
  map_denom = NULL,
  slice = 1,
  zlim = NULL,
  interp = 1,
  xlim = NULL
)
```

Arguments

<code>fit_res</code>	fit_result object.
<code>map</code>	fit result values to display as a colour map. Can be specified as a character string or array of numeric values. Defaults to "tNAA".
<code>map_denom</code>	fit result values to divide the map argument by. Can be specified as a character string (eg "tCr") or array of numeric values.
<code>slice</code>	slice to plot in the z direction.
<code>zlim</code>	range of values to plot.
<code>interp</code>	interpolation factor.
<code>xlim</code>	spectral plot limits for the x axis.

`plot_slice_map` *Plot a slice from a 7 dimensional array.*

Description

Plot a slice from a 7 dimensional array.

Usage

```
plot_slice_map(  
    data,  
    zlim = NULL,  
    mask_map = NULL,  
    mask_cutoff = 20,  
    interp = 1,  
    slice = 1,  
    dyn = 1,  
    coil = 1,  
    ref = 1,  
    denom = NULL,  
    horizontal = FALSE  
)
```

Arguments

data	7d array of values to be plotted.
zlim	smallest and largest values to be plotted.
mask_map	matching map with logical values to indicate if the corresponding values should be plotted.
mask_cutoff	minimum values to plot (as a percentage of the maximum).
interp	map interpolation factor.
slice	the slice index to plot.
dyn	the dynamic index to plot.
coil	the coil element number to plot.
ref	reference index to plot.
denom	map to use as a denominator.
horizontal	display the colourbar horizontally (logical).

plot_slice_map_inter *Plot an interactive slice map from a data array where voxels can be selected to display a corresponding spectrum.*

Description

Plot an interactive slice map from a data array where voxels can be selected to display a corresponding spectrum.

Usage

```
plot_slice_map_inter(  
    mrs_data,  
    map = NULL,  
    xlim = NULL,  
    slice = 1,  
    zlim = NULL,  
    mask_map = NULL,  
    denom = NULL,  
    mask_cutoff = 20,  
    interp = 1,  
    mode = "re",  
    y_scale = FALSE,  
    ylim = NULL,  
    coil = 1,  
    fd = TRUE  
)
```

Arguments

<i>mrs_data</i>	spectral data.
<i>map</i>	array of values to be plotted, defaults to the integration of the modulus of the full spectral width.
<i>xlim</i>	spectral region to plot.
<i>slice</i>	the slice index to plot.
<i>zlim</i>	smallest and largest values to be plotted.
<i>mask_map</i>	matching map with logical values to indicate if the corresponding values should be plotted.
<i>denom</i>	map to use as a denominator.
<i>mask_cutoff</i>	minimum values to plot (as a percentage of the maximum).
<i>interp</i>	map interpolation factor.
<i>mode</i>	representation of the complex spectrum to be plotted, can be one of: "re", "im", "mod" or "arg".
<i>y_scale</i>	option to display the y-axis values (logical).
<i>ylim</i>	intensity range to plot.
<i>coil</i>	coil element to plot.
<i>fd</i>	display data in the frequency-domain (default), or time-domain (logical).

plot_voi_overlay *Plot a volume as an image overlay.*

Description

Plot a volume as an image overlay.

Usage

```
plot_voi_overlay(mri, voi, export_path = NULL, zlim = NULL, ...)
```

Arguments

mri	image data as a nifti object or path to data file.
voi	volume data as a nifti object or path to data file.
export_path	optional path to save the image in png format.
zlim	underlay intensity limits.
...	additional arguments to the ortho3 function.

plot_voi_overlay_seg *Plot a volume as an overlay on a segmented brain volume.*

Description

Plot a volume as an overlay on a segmented brain volume.

Usage

```
plot_voi_overlay_seg(mri_seg, voi, export_path = NULL, ...)
```

Arguments

mri_seg	segmented brain volume as a nifti object.
voi	volume data as a nifti object.
export_path	optional path to save the image in png format.
...	additional arguments to the ortho3 function.

ppm*Return the ppm scale of an MRS dataset or fit result.***Description**

Return the ppm scale of an MRS dataset or fit result.

Usage

```
ppm(x, ft = NULL, ref = NULL, fs = NULL, N = NULL)

## S3 method for class 'mrs_data'
ppm(x, ft = NULL, ref = NULL, fs = NULL, N = NULL)

## S3 method for class 'fit_result'
ppm(x, ft = NULL, ref = NULL, fs = NULL, N = NULL)
```

Arguments

x	MRS dataset or fit result.
ft	transmitter frequency in Hz, does not apply when the object is a fit result.
ref	reference value for ppm scale, does not apply when the object is a fit result.
fs	sampling frequency in Hz, does not apply when the object is a fit result.
N	number of data points in the spectral dimension, does not apply when the object is a fit result.

Value

ppm scale.

precomp*Save function results to file and load on subsequent calls to avoid repeat computation.***Description**

Save function results to file and load on subsequent calls to avoid repeat computation.

Usage

```
precomp(file, fun, ...)
```

Arguments

- | | |
|------|---------------------------------|
| file | file name to write the results. |
| fun | function to run. |
| ... | arguments to be passed to fun. |

print.fit_result *Print a summary of an object of class fit_result.*

Description

Print a summary of an object of class fit_result.

Usage

```
## S3 method for class 'fit_result'  
print(x, ...)
```

Arguments

- | | |
|-----|--------------------|
| x | fit_result object. |
| ... | further arguments. |

print.mrs_data *Print a summary of mrs_data parameters.*

Description

Print a summary of mrs_data parameters.

Usage

```
## S3 method for class 'mrs_data'  
print(x, full = FALSE, ...)
```

Arguments

- | | |
|------|---------------------------------------|
| x | mrs_data object. |
| full | print all parameters (default FALSE). |
| ... | further arguments. |

qn_states*Get the quantum coherence matrix for a spin system.***Description**

Get the quantum coherence matrix for a spin system.

Usage

```
qn_states(sys)
```

Arguments

sys	spin system object.
-----	---------------------

Value

quantum coherence number matrix.

rats*Robust Alignment to a Target Spectrum (RATS).***Description**

Robust Alignment to a Target Spectrum (RATS).

Usage

```
rats(
  mrs_data,
  ref = NULL,
  xlim = c(4, 0.5),
  max_shift = 20,
  p_deg = 2,
  sp_N = 2,
  sp_deg = 3,
  max_t = 0.2,
  basis_type = "poly",
  rescale_output = TRUE,
  phase_corr = TRUE,
  ret_corr_only = TRUE
)
```

Arguments

<code>mrs_data</code>	MRS data to be corrected.
<code>ref</code>	optional MRS data to use as a reference, the mean of all dynamics is used if this argument is not supplied.
<code>xlim</code>	optional frequency range to perform optimisation, set to NULL to use the full range.
<code>max_shift</code>	maximum allowable frequency shift in Hz.
<code>p_deg</code>	polynomial degree used for baseline modelling. Negative values disable baseline modelling.
<code>sp_N</code>	number of spline functions, note the true number will be <code>sp_N + sp_deg</code> .
<code>sp_deg</code>	degree of spline functions.
<code>max_t</code>	truncate the FID when longer than <code>max_t</code> to reduce time taken, set to NULL to use the entire FID.
<code>basis_type</code>	may be one of "poly" or "spline".
<code>rescale_output</code>	rescale the <code>bl_matched_spec</code> and <code>bl</code> output to improve consistency between dynamic scans.
<code>phase_corr</code>	apply phase correction (in addition to frequency). TRUE by default.
<code>ret_corr_only</code>	return the corrected <code>mrs_data</code> object only.

Value

a list containing the corrected data; phase and shift values in units of degrees and Hz respectively.

`Re.mrs_data`

Apply Re operator to an MRS dataset.

Description

Apply Re operator to an MRS dataset.

Usage

```
## S3 method for class 'mrs_data'
Re(z)
```

Arguments

<code>z</code>	MRS data.
----------------	-----------

Value

MRS data following Re operator.

read_basis*Read a basis file in LCModel .basis format.*

Description

Read a basis file in LCModel .basis format.

Usage

```
read_basis(basis_file, ref = def_ref(), sort_basis = TRUE)
```

Arguments

basis_file	path to basis file.
ref	assumed ppm reference value.
sort_basis	sort the basis set based on signal names.

Value

basis object.

read_basis_ac*Read a basis file in LCModel .basis format (for testing only).*

Description

Read a basis file in LCModel .basis format (for testing only).

Usage

```
read_basis_ac(basis_file, ref = def_ref(), sort_basis = TRUE)
```

Arguments

basis_file	path to basis file.
ref	assumed ppm reference value.
sort_basis	sort the basis set based on signal names.

Value

basis object.

`read_ima_coil_dir`

Read a directory containing Siemens MRS IMA files and combine along the coil dimension. Note that the coil ID is inferred from the sorted file name and should be checked when consistency is required between two directories.

Description

Read a directory containing Siemens MRS IMA files and combine along the coil dimension. Note that the coil ID is inferred from the sorted file name and should be checked when consistency is required between two directories.

Usage

```
read_ima_coil_dir(dir, extra = NULL)
```

Arguments

dir	data directory path.
extra	an optional data frame to provide additional variables for use in subsequent analysis steps, eg id or grouping variables.

Value

mrs_data object.

`read_ima_dyn_dir`

Read a directory containing Siemens MRS IMA files and combine along the dynamic dimension. Note that the coil ID is inferred from the sorted file name and should be checked when consistency is required.

Description

Read a directory containing Siemens MRS IMA files and combine along the dynamic dimension. Note that the coil ID is inferred from the sorted file name and should be checked when consistency is required.

Usage

```
read_ima_dyn_dir(dir, extra = NULL)
```

Arguments

dir	data directory path.
extra	an optional data frame to provide additional variables for use in subsequent analysis steps, eg id or grouping variables.

Value

mrs_data object.

read_lcm_coord

Read an LCModel formatted coord file containing fit information.

Description

Read an LCModel formatted coord file containing fit information.

Usage

```
read_lcm_coord(coord_f)
```

Arguments

coord_f	path to the coord file.
---------	-------------------------

Value

list containing a table of fit point and results structure containing signal amplitudes, errors and fitting diagnostics.

read_mrs

Read MRS data from a file.

Description

Read MRS data from a file.

Usage

```
read_mrs(
  fname,
  format = NULL,
  ft = NULL,
  fs = NULL,
  ref = NULL,
  n_ref_scans = NULL,
  full_fid = FALSE,
  omit_svs_ref_scans = TRUE,
  verbose = FALSE,
  extra = NULL
)
```

Arguments

fname	filename of the dpt format MRS data.
format	string describing the data format. Must be one of the following : "spar_sdat", "rda", "dicom", "twix", "pfile", "list_data", "paravis", "dpt", "lcm_raw", "rds", "nifti", "varian". If not specified, the format will be guessed from the filename extension.
ft	transmitter frequency in Hz (required for list_data format).
fs	sampling frequency in Hz (required for list_data format).
ref	reference value for ppm scale (required for list_data format).
n_ref_scans	override the number of water reference scans detected in the file header (GE p-file only).
full_fid	export all data points, including those before the start of the FID (default = FALSE), TWIX format only.
omit_svs_ref_scans	remove any reference scans sometimes saved in SVS twix data (default = TRUE).
verbose	print data file information (default = FALSE).
extra	an optional data frame to provide additional variables for use in subsequent analysis steps, eg id or grouping variables.

Value

MRS data object.

Examples

```
fname <- system.file("extdata", "philips_spar_sdat_WS.SDAT", package = "spant")
mrs_data <- read_mrs(fname)
print(mrs_data)
```

read_mrs_tqn

Read MRS data using the TARQUIN software package.

Description

Read MRS data using the TARQUIN software package.

Usage

```
read_mrs_tqn(fname, fname_ref = NA, format, id = NA, group = NA)
```

Arguments

<code>fname</code>	the filename containing the MRS data.
<code>fname_ref</code>	a second filename containing reference MRS data.
<code>format</code>	format of the MRS data. Can be one of the following: siemens, philips, ge, dcm, dpt, rda, lcm, varian, bruker, jmrui_txt.
<code>id</code>	optional ID string.
<code>group</code>	optional group string.

Value

MRS data object.

Examples

```
fname <- system.file("extdata","philips_spar_sdat_WS.SDAT",package="spant")
## Not run:
mrs_data <- read_mrs_tqn(fname, format="philips")

## End(Not run)
```

`read_siemens_txt_hdr` *Read the text format header found in Siemens IMA and TWIX data files.*

Description

Read the text format header found in Siemens IMA and TWIX data files.

Usage

```
read_siemens_txt_hdr(input, version = "vd", verbose)
```

Arguments

<code>input</code>	file name to read or raw data.
<code>version</code>	software version, can be "vb" or "vd".
<code>verbose</code>	print information to the console.

Value

a list of parameter values

read_tqn_fit *Reader for csv fit results generated by TARQUIN.*

Description

Reader for csv fit results generated by TARQUIN.

Usage

```
read_tqn_fit(fit_f)
```

Arguments

fit_f TARQUIN fit file.

Value

A data frame of the fit data points.

Examples

```
## Not run:  
fit <- read_tqn_fit(system.file("extdata","fit.csv",package="spant"))  
  
## End(Not run)
```

read_tqn_result *Reader for csv results generated by TARQUIN.*

Description

Reader for csv results generated by TARQUIN.

Usage

```
read_tqn_result(result_f, remove_rcs = TRUE)
```

Arguments

result_f TARQUIN result file.
remove_rcs omit row, column and slice ids from output.

Value

list of amplitudes, crlbs and diagnostics.

Examples

```
## Not run:
result <- read_tqn_result(system.file("extdata", "result.csv", package="spant"))

## End(Not run)
```

recon_twix_2d_mrsi *Reconstruct 2D MRSI data from a twix file loaded with read_mrs.*

Description

Reconstruct 2D MRSI data from a twix file loaded with read_mrs.

Usage

```
recon_twix_2d_mrsi(twix_mrs)
```

Arguments

twix_mrs raw dynamic data.

Value

reconstructed data.

rectangular_mask *Create a rectangular mask stored as a matrix of logical values.*

Description

Create a rectangular mask stored as a matrix of logical values.

Usage

```
rectangular_mask(xN, yN, x0, y0, xw, yw, angle)
```

Arguments

xN	number of pixels in the x dimension.
yN	number of pixels in the y dimension.
x0	centre of rectangle in the x direction in units of pixels.
y0	centre of rectangle in the y direction in units of pixels.
xw	width in the x direction in units of pixels.
yw	width in the y direction in units of pixels.
angle	angle of rotation in degrees.

Value

logical mask matrix with dimensions `fov_yN` x `fov_xN`.

`rep_array_dim`

Repeat an array over a given dimension.

Description

Repeat an array over a given dimension.

Usage

`rep_array_dim(x, rep_dim, n)`

Arguments

<code>x</code>	array.
<code>rep_dim</code>	dimension to extend.
<code>n</code>	number of times to repeat.

Value

extended array.

`rep_dyn`

Replicate a scan in the dynamic dimension.

Description

Replicate a scan in the dynamic dimension.

Usage

`rep_dyn(mrs_data, times)`

Arguments

<code>mrs_data</code>	MRS data to be replicated.
<code>times</code>	number of times to replicate.

Value

replicated data object.

<code>rep_mrs</code>	<i>Replicate a scan over a given dimension.</i>
----------------------	---

Description

Replicate a scan over a given dimension.

Usage

```
rep_mrs(
  mrs_data,
  x_rep = 1,
  y_rep = 1,
  z_rep = 1,
  dyn_rep = 1,
  coil_rep = 1,
  warn = TRUE
)
```

Arguments

<code>mrs_data</code>	MRS data to be replicated.
<code>x_rep</code>	number of x replications.
<code>y_rep</code>	number of y replications.
<code>z_rep</code>	number of z replications.
<code>dyn_rep</code>	number of dynamic replications.
<code>coil_rep</code>	number of coil replications.
<code>warn</code>	print a warning when the data dimensions do not change.

Value

replicated data object.

<code>resample_img</code>	<i>Resample an image to match a target image space.</i>
---------------------------	---

Description

Resample an image to match a target image space.

Usage

```
resample_img(source, target, interp = 3L)
```

Arguments

source	image data as a nifti object.
target	image data as a nifti object.
interp	interpolation parameter, see nifyreg.linear definition.

Value

resampled image data as a nifti object.

resample_voi	<i>Resample a VOI to match a target image space using nearest-neighbour interpolation.</i>
--------------	--

Description

Resample a VOI to match a target image space using nearest-neighbour interpolation.

Usage

```
resample_voi(voi, mri)
```

Arguments

voi	volume data as a nifti object.
mri	image data as a nifti object.

Value

volume data as a nifti object.

reslice_to_mrs	<i>Reslice a nifti object to match the orientation of mrs data.</i>
----------------	---

Description

Reslice a nifti object to match the orientation of mrs data.

Usage

```
reslice_to_mrs(mri, mrs, interp = 3L)
```

Arguments

mri	nifti object to be resliced.
mrs	mrs_data object for the target orientation.
interp	interpolation parameter, see nifyreg.linear definition.

Value

resliced imaging data.

reson_table2mrs_data *Generate mrs_data from a table of single Lorentzian resonances.*

Description

Generate mrs_data from a table of single Lorentzian resonances.

Usage

```
reson_table2mrs_data(
    reson_table,
    acq_paras = def_acq_paras(),
    back_extrap_pts = 0
)
```

Arguments

reson_table	as produced by the hsvgd function.
acq_paras	list of acquisition parameters. See
back_extrap_pts	number of data points to back extrapolate def_acq_paras

Value

mrs_data object.

re_weighting *Apply a weighting to the FID to enhance spectral resolution.*

Description

Apply a weighting to the FID to enhance spectral resolution.

Usage

```
re_weighting(mrs_data, re, alpha)
```

Arguments

mrs_data	data to be enhanced.
re	resolution enhancement factor (rising exponential factor).
alpha	alpha factor (Gaussian decay)

Value

resolution enhanced mrs_data.

`rm_dync`

Remove a subset of dynamic scans.

Description

Remove a subset of dynamic scans.

Usage

`rm_dync(mrs_data, subset)`

Arguments

<code>mrs_data</code>	dynamic MRS data.
<code>subset</code>	vector containing indices to the dynamic scans to be removed.

Value

MRS data without the specified dynamic scans.

`scale_amp_molal_pvc`

Apply partial volume correction to a fitting result object.

Description

Apply partial volume correction to a fitting result object.

Usage

`scale_amp_molal_pvc(fit_result, ref_data, p_vols, te, tr, ...)`

Arguments

<code>fit_result</code>	result object generated from fitting.
<code>ref_data</code>	water reference MRS data object.
<code>p_vols</code>	a numeric vector of partial volumes.
<code>te</code>	the MRS TE in seconds.
<code>tr</code>	the MRS TR in seconds.
<code>...</code>	additional arguments to get_td_amp function.

Value

A `fit_result` object with a rescaled results table.

scale_amp_molar *Apply water reference scaling to a fitting results object to yield metabolite quantities in millimolar (mM) units (mol/litre).*

Description

See the LCModel manual section on water-scaling for details on the assumptions and relevant references.

Usage

```
scale_amp_molar(fit_result, ref_data, w_att = 0.7, w_conc = 35880, ...)
```

Arguments

fit_result	a result object generated from fitting.
ref_data	water reference MRS data object.
w_att	water attenuation factor (default = 0.7).
w_conc	assumed water concentration (default = 35880). Default value corresponds to typical white matter. Set to 43300 for gray matter, and 55556 for phantom measurements.
...	additional arguments to get_td_amp function.

Value

a fit_result object with a rescaled results table.

scale_amp_ratio *Scale fitted amplitudes to a ratio of signal amplitude.*

Description

Scale fitted amplitudes to a ratio of signal amplitude.

Usage

```
scale_amp_ratio(fit_result, name)
```

Arguments

fit_result	a result object generated from fitting.
name	the signal name to use as a denominator (usually, "tCr" or "tNAA").

Value

a fit_result object with a rescaled results table.

scale_amp_water_ratio *Scale metabolite amplitudes as a ratio to the unsuppressed water amplitude.*

Description

Scale metabolite amplitudes as a ratio to the unsuppressed water amplitude.

Usage

```
scale_amp_water_ratio(fit_result, ref_data, ...)
```

Arguments

fit_result	a result object generated from fitting.
ref_data	a water reference MRS data object.
...	additional arguments to get_td_amp function.

Value

a fit_result object with a rescaled results table.

scale_mrs_amp *Scale an mrs_data object by a scalar or vector or amplitudes.*

Description

Scale an mrs_data object by a scalar or vector or amplitudes.

Usage

```
scale_mrs_amp(mrs_data, amp)
```

Arguments

mrs_data	data to be scaled.
amp	multiplicative factor, must have length equal to 1 or Nspec(mrs_data).

Value

mrs_data object multiplied by the amplitude scale factor.

scale_spec*Scale mrs_data to a spectral region.***Description**

Scale mrs_data to a spectral region.

Usage

```
scale_spec(
  mrs_data,
  xlim = NULL,
  operator = "sum",
  freq_scale = "ppm",
  mode = "re",
  mean_dyncs = TRUE,
  ret_scale_factor = FALSE
)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>xlim</code>	spectral range to be integrated (defaults to full range).
<code>operator</code>	can be "sum" (default), "mean", "l2", "max", "min" or "max-min".
<code>freq_scale</code>	units of xlim, can be : "ppm", "Hz" or "points".
<code>mode</code>	spectral mode, can be : "re", "im", "mod" or "cplx".
<code>mean_dyncs</code>	mean the dynamic scans before applying the operator. The same scaling value will be applied to each individual dynamic.
<code>ret_scale_factor</code>	option to return the scaling factor in addition to the scaled data.

Value

normalised data.

sd*Calculate the standard deviation spectrum from an mrs_data object.***Description**

Calculate the standard deviation spectrum from an mrs_data object.

Usage

```
sd(x, na.rm)
```

Arguments

- x object of class mrs_data.
na.rm remove NA values.

Value

sd mrs_data object.

sd.mrs_data *Calculate the standard deviation spectrum from an mrs_data object.*

Description

Calculate the standard deviation spectrum from an mrs_data object.

Usage

```
## S3 method for class 'mrs_data'  
sd(x, na.rm = FALSE)
```

Arguments

- x object of class mrs_data.
na.rm remove NA values.

Value

sd mrs_data object.

seconds *Return a time scale vector to match the FID of an MRS data object.*

Description

Return a time scale vector to match the FID of an MRS data object.

Usage

```
seconds(mrs_data)
```

Arguments

- mrs_data MRS data.

Value

time scale vector in units of seconds.

<code>seq_cpmg_ideal</code>	<i>CPMG style sequence with ideal pulses.</i>
-----------------------------	---

Description

CPMG style sequence with ideal pulses.

Usage

```
seq_cpmg_ideal(spin_params, ft, ref, TE = 0.03, echoes = 4)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
TE	echo time in seconds.
echoes	number of echoes.

Value

list of resonance amplitudes and frequencies.

<code>seq_mega_press_ideal</code>	<i>MEGA-PRESS sequence with ideal localisation pulses and Gaussian shaped editing pulse.</i>
-----------------------------------	--

Description

MEGA-PRESS sequence with ideal localisation pulses and Gaussian shaped editing pulse.

Usage

```
seq_mega_press_ideal(
    spin_params,
    ft,
    ref,
    ed_freq = 1.89,
    TE1 = 0.015,
    TE2 = 0.053,
    BW = 110,
    steps = 50
)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
ed_freq	editing pulse frequency in ppm.
TE1	TE1 sequence parameter in seconds (TE=TE1+TE2).
TE2	TE2 sequence parameter in seconds.
BW	editing pulse bandwidth in Hz.
steps	number of hard pulses used to approximate the editing pulse.

Value

list of resonance amplitudes and frequencies.

seq_press_ideal *PRESS sequence with ideal pulses.*

Description

PRESS sequence with ideal pulses.

Usage

```
seq_press_ideal(spin_params, ft, ref, TE1 = 0.01, TE2 = 0.02)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
TE1	TE1 sequence parameter in seconds (TE=TE1+TE2).
TE2	TE2 sequence parameter in seconds.

Value

list of resonance amplitudes and frequencies.

seq_pulse_acquire *Simple pulse and acquire sequence with ideal pulses.*

Description

Simple pulse and acquire sequence with ideal pulses.

Usage

```
seq_pulse_acquire(spin_params, ft, ref)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.

Value

list of resonance amplitudes and frequencies.

seq_pulse_acquire_31p *Simple pulse and acquire sequence with ideal pulses.*

Description

Simple pulse and acquire sequence with ideal pulses.

Usage

```
seq_pulse_acquire_31p(spin_params, ft, ref)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.

Value

list of resonance amplitudes and frequencies.

seq_slaser_ideal *sLASER sequence with ideal pulses.*

Description

sLASER sequence with ideal pulses.

Usage

```
seq_slaser_ideal(spin_params, ft, ref, TE1 = 0.008, TE2 = 0.011, TE3 = 0.009)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
TE1	first echo time (between exc. and 1st echo) in seconds.
TE2	second echo time (between 2nd echo and 4th echo) in seconds.
TE3	third echo time (between 4th echo and 5th echo) in seconds.

Value

list of resonance amplitudes and frequencies.

seq_spin_echo_ideal *Spin echo sequence with ideal pulses.*

Description

Spin echo sequence with ideal pulses.

Usage

```
seq_spin_echo_ideal(spin_params, ft, ref, TE = 0.03)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
TE	echo time in seconds.

Value

list of resonance amplitudes and frequencies.

seq_spin_echo_ideal_31p

Spin echo sequence with ideal pulses.

Description

Spin echo sequence with ideal pulses.

Usage

```
seq_spin_echo_ideal_31p(spin_params, ft, ref, TE = 0.03)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
TE	echo time in seconds.

Value

list of resonance amplitudes and frequencies.

seq_steam_ideal

STEAM sequence with ideal pulses.

Description

STEAM sequence with ideal pulses.

Usage

```
seq_steam_ideal(spin_params, ft, ref, TE = 0.03, TM = 0.02)
```

Arguments

spin_params	spin system definition.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
TE	sequence parameter in seconds.
TM	sequence parameter in seconds.

Value

list of resonance amplitudes and frequencies.

set_def_acq_paras *Set the default acquisition parameters.*

Description

Set the default acquisition parameters.

Usage

```
set_def_acq_paras(  
    ft = getopt("spant.def_ft"),  
    fs = getopt("spant.def_fs"),  
    N = getopt("spant.def_N"),  
    ref = getopt("spant.def_ref"),  
    nuc = getopt("spant.nuc")  
)
```

Arguments

ft	transmitter frequency in Hz.
fs	sampling frequency in Hz.
N	number of data points in the spectral dimension.
ref	reference value for ppm scale.
nuc	resonant nucleus.

set_lcm_cmd *Set the command to run the LCModel command-line program.*

Description

Set the command to run the LCModel command-line program.

Usage

```
set_lcm_cmd(cmd)
```

Arguments

cmd	path to binary.
-----	-----------------

`set_lw`

Apply line-broadening to an mrs_data object to achieve a specified linewidth.

Description

Apply line-broadening to an mrs_data object to achieve a specified linewidth.

Usage

```
set_lw(mrs_data, lw, xlim = c(4, 0.5), lg = 1)
```

Arguments

<code>mrs_data</code>	data in.
<code>lw</code>	target linewidth in units of ppm.
<code>xlim</code>	region to search for peaks to obtain a linewidth estimate.
<code>lg</code>	Lorentz-Gauss lineshape parameter.

Value

line-broadened data.

`set_mask_xy_mat`

Set the masked voxels in a 2D MRSI dataset to given spectrum.

Description

Set the masked voxels in a 2D MRSI dataset to given spectrum.

Usage

```
set_mask_xy_mat(mrs_data, mask, mask_mrs_data)
```

Arguments

<code>mrs_data</code>	MRSI data object.
<code>mask</code>	matrix of boolean values specifying the voxels to set, where a value of TRUE indicates the voxel should be set to <code>mask_mrs_data</code> .
<code>mask_mrs_data</code>	the spectral data to be assigned to the masked voxels.

Value

updated dataset.

set_precomp_mode *Set the precompute mode.*

Description

Set the precompute mode.

Usage

```
set_precomp_mode(mode = NA)
```

Arguments

mode can be one of: "default", "overwrite", "clean" or "disabled".

set_precomp_verbose *Set the verbosity of the precompute function.*

Description

Set the verbosity of the precompute function.

Usage

```
set_precomp_verbose(verbose = NA)
```

Arguments

verbose can be TRUE or FALSE.

set_ref *Set the ppm reference value (eg ppm value at 0Hz).*

Description

Set the ppm reference value (eg ppm value at 0Hz).

Usage

```
set_ref(mrs_data, ref)
```

Arguments

mrs_data MRS data.

ref reference value for ppm scale.

<code>set_td_pts</code>	<i>Set the number of time-domain data points, truncating or zero-filling as appropriate.</i>
-------------------------	--

Description

Set the number of time-domain data points, truncating or zero-filling as appropriate.

Usage

```
set_td_pts(mrs_data, pts)
```

Arguments

<code>mrs_data</code>	MRS data.
<code>pts</code>	number of data points.

Value

MRS data with pts data points.

<code>set_tqn_cmd</code>	<i>Set the command to run the TARQUIN command-line program.</i>
--------------------------	---

Description

Set the command to run the TARQUIN command-line program.

Usage

```
set_tqn_cmd(cmd)
```

Arguments

<code>cmd</code>	path to binary.
------------------	-----------------

shift	<i>Apply a frequency shift to MRS data.</i>
-------	---

Description

Apply a frequency shift to MRS data.

Usage

```
shift(mrs_data, shift, units = "ppm")
```

Arguments

mrs_data	MRS data.
shift	frequency shift (in ppm by default).
units	of the shift ("ppm" or "hz").

Value

frequency shifted MRS data.

shift_basis	<i>Apply frequency shifts to basis set signals.</i>
-------------	---

Description

Apply frequency shifts to basis set signals.

Usage

```
shift_basis(basis, shifts)
```

Arguments

basis	the basis to apply the shift to.
shifts	a vector of frequency shifts to apply in ppm units. Must be the same length as there are basis elements.

Value

modified basis set object.

sim_basis *Simulate a basis set object.*

Description

Simulate a basis set object.

Usage

```
sim_basis(
  mol_list,
  pul_seq = seq_pulse_acquire,
  acq_paras = def_acq_paras(),
  xlim = NULL,
  ...
)
```

Arguments

<code>mol_list</code>	list of <code>mol_parameter</code> objects.
<code>pul_seq</code>	pulse sequence function to use.
<code>acq_paras</code>	list of acquisition parameters or an <code>mrs_data</code> object. See <code>def_acq_paras</code>
<code>xlim</code>	ppm range limiting signals to be simulated.
<code>...</code>	extra parameters to pass to the pulse sequence function.

Value

basis object.

sim_basis_1h_brain *Simulate a basis-set suitable for 1H brain MRS analysis acquired with a PRESS sequence. Note, ideal pulses are assumed.*

Description

Simulate a basis-set suitable for 1H brain MRS analysis acquired with a PRESS sequence. Note, ideal pulses are assumed.

Usage

```
sim_basis_1h_brain(
  pul_seq = seq_press_ideal,
  acq_paras = def_acq_paras(),
  xlim = c(0.5, 4.2),
  lcm_compat = FALSE,
  ...
)
```

Arguments

pul_seq	pulse sequence function to use.
acq_paras	list of acquisition parameters or an mrs_data object. See def_acq_paras .
xlim	range of frequencies to simulate in ppm.
lcm_compat	exclude lipid and MM signals for use with default LCModel options.
...	extra parameters to pass to the pulse sequence function.

Value

basis object.

sim_basis_1h_brain_press

Simulate a basis-set suitable for 1H brain MRS analysis acquired with a PRESS sequence. Note, ideal pulses are assumed.

Description

Simulate a basis-set suitable for 1H brain MRS analysis acquired with a PRESS sequence. Note, ideal pulses are assumed.

Usage

```
sim_basis_1h_brain_press(  
  acq_paras = def_acq_paras(),  
  xlim = c(0.5, 4.2),  
  lcm_compat = FALSE,  
  TE1 = 0.01,  
  TE2 = 0.02  
)
```

Arguments

acq_paras	list of acquisition parameters or an mrs_data object. See def_acq_paras
xlim	range of frequencies to simulate in ppm.
lcm_compat	exclude lipid and MM signals for use with default LCModel options.
TE1	TE1 of PRESS sequence (TE = TE1 + TE2).
TE2	TE2 of PRESS sequence.

Value

basis object.

sim_basis_tqn	<i>Simulate a basis file using TARQUIN.</i>
---------------	---

Description

Simulate a basis file using TARQUIN.

Usage

```
sim_basis_tqn(
  fs = def_fs(),
  ft = def_ft(),
  N = def_N(),
  ref = def_ref(),
  opts = NULL
)
```

Arguments

fs	sampling frequency
ft	transmitter frequency
N	number of data points
ref	chemical shift reference
opts	list of options to pass to TARQUIN.

Examples

```
## Not run:
write_basis_tqn('test.basis',mrs_data,c("--echo","0.04"))

## End(Not run)
```

sim_brain_1h	<i>Simulate MRS data with a similar appearance to normal brain (by default).</i>
--------------	--

Description

Simulate MRS data with a similar appearance to normal brain (by default).

Usage

```
sim_brain_1h(
  acq_paras = def_acq_paras(),
  type = "normal_v1",
  pul_seq = seq_press_ideal,
  xlim = c(0.5, 4.2),
  full_output = FALSE,
  amps = NULL,
  ...
)
```

Arguments

acq_paras	list of acquisition parameters or an mrs_data object. See def_acq_paras .
type	type of spectrum, only "normal" is implemented currently.
pul_seq	pulse sequence function to use.
xlim	range of frequencies to simulate in ppm.
full_output	when FALSE (default) only output the simulated MRS data. When TRUE output a list containing the MRS data, basis set object and corresponding amplitudes.
amps	a vector of basis amplitudes may be specified to modify the output spectrum.
...	extra parameters to pass to the pulse sequence function.

Value

see full_output option.

sim_mol

Simulate a mol_parameter object.

Description

Simulate a mol_parameter object.

Usage

```
sim_mol(
  mol,
  pul_seq = seq_pulse_acquire,
  ft = def_ft(),
  ref = def_ref(),
  fs = def_fs(),
  N = def_N(),
  xlim = NULL,
  ...
)
```

Arguments

<code>mol</code>	<code>mol_parameter</code> object.
<code>pul_seq</code>	pulse sequence function to use.
<code>ft</code>	transmitter frequency in Hz.
<code>ref</code>	reference value for ppm scale.
<code>fs</code>	sampling frequency in Hz.
<code>N</code>	number of data points in the spectral dimension.
<code>xlim</code>	ppm range limiting signals to be simulated.
<code>...</code>	extra parameters to pass to the pulse sequence function.

Value

`mrs_data` object.

`sim_noise`

Simulate an mrs_data object containing simulated Gaussian noise.

Description

Simulate an `mrs_data` object containing simulated Gaussian noise.

Usage

```
sim_noise(
  sd = 0.1,
  fs = def_fs(),
  ft = def_ft(),
  N = def_N(),
  ref = def_ref(),
  dyns = 1,
  fd = TRUE
)
```

Arguments

<code>sd</code>	standard deviation of the noise.
<code>fs</code>	sampling frequency in Hz.
<code>ft</code>	transmitter frequency in Hz.
<code>N</code>	number of data points in the spectral dimension.
<code>ref</code>	reference value for ppm scale.
<code>dyns</code>	number of dynamic scans to generate.
<code>fd</code>	return data in the frequency-domain (TRUE) or time-domain (FALSE)

Value

`mrs_data` object.

sim_resonances	<i>Simulate a MRS data object containing a set of simulated resonances.</i>
----------------	---

Description

Simulate a MRS data object containing a set of simulated resonances.

Usage

```
sim_resonances(  
    freq = 0,  
    amp = 1,  
    lw = 0,  
    lg = 0,  
    phase = 0,  
    freq_ppm = TRUE,  
    acq_paras = def_acq_paras(),  
    fp_scale = TRUE,  
    back_extrap_pts = 0,  
    sum_resonances = TRUE  
)
```

Arguments

freq	resonance frequency.
amp	resonance amplitude.
lw	line width in Hz.
lg	Lorentz-Gauss lineshape parameter (between 0 and 1).
phase	phase in degrees.
freq_ppm	frequencies are given in ppm units if set to TRUE, otherwise Hz are assumed.
acq_paras	list of acquisition parameters. See def_acq_paras
fp_scale	multiply the first data point by 0.5.
back_extrap_pts	number of data points to back extrapolate.
sum_resonances	sum all resonances (default is TRUE), otherwise return a dynamic mrs_data object.

Value

MRS data object.

Examples

```
sim_data <- sim_resonances(freq = 2, lw = 5)
```

sim_zero*Simulate an mrs_data object containing complex zero valued samples.***Description**

Simulate an mrs_data object containing complex zero valued samples.

Usage

```
sim_zero(fs = def_fs(), ft = def_ft(), N = def_N(), ref = def_ref(), dyns = 1)
```

Arguments

<code>fs</code>	sampling frequency in Hz.
<code>ft</code>	transmitter frequency in Hz.
<code>N</code>	number of data points in the spectral dimension.
<code>ref</code>	reference value for ppm scale.
<code>dyns</code>	number of dynamic scans to generate.

Value

mrs_data object.

sort_basis*Sort the basis-set elements alphabetically.***Description**

Sort the basis-set elements alphabetically.

Usage

```
sort_basis(basis)
```

Arguments

<code>basis</code>	input basis.
--------------------	--------------

Value

sorted basis.

`spant_abfit_benchmark` *Simulate and fit some spectra with ABfit for benchmarking purposes. Basic timing and performance metrics will be printed.*

Description

Simulate and fit some spectra with ABfit for benchmarking purposes. Basic timing and performance metrics will be printed.

Usage

```
spant_abfit_benchmark(noise_reps = 10, return_res = FALSE, opts = abfit_opts())
```

Arguments

<code>noise_reps</code>	number of spectra to fit with differing noise samples.
<code>return_res</code>	return a list of <code>fit_result</code> objects.
<code>opts</code>	ABfit options structure.

`spant_mpress_drift` *Example MEGA-PRESS data with significant B0 drift.*

Description

Example MEGA-PRESS data with significant B0 drift.

Usage

```
spant_mpress_drift
```

Format

An object of class `mrs_data` of length 13.

spant_simulation_benchmark

Simulate a typical metabolite basis set for benchmarking. Timing metrics will be printed on completion.

Description

Simulate a typical metabolite basis set for benchmarking. Timing metrics will be printed on completion.

Usage

```
spant_simulation_benchmark(sim_reps = 10, N = 1024)
```

Arguments

sim_reps	number of times to simulate the basis set.
N	number of FID data points to simulate.

spec_decomp

Decompose an mrs_data object into white and gray matter spectra.

Description

An implementation of the method published by Goryawala et al MRM 79(6) 2886-2895 (2018). "Spectral decomposition for resolving partial volume effects in MRSI".

Usage

```
spec_decomp(mrs_data, wm, gm, norm_fractions = TRUE)
```

Arguments

mrs_data	data to be decomposed into white and gray matter spectra.
wm	vector of white matter contributions to each voxel.
gm	vector of gray matter contributions to each voxel.
norm_fractions	option to normalise the wm, gm vectors for each voxel.

Value

a list of two mrs_data objects corresponding to the two tissue types.

spec_op	<i>Perform a mathematical operation on a spectral region.</i>
---------	---

Description

Perform a mathematical operation on a spectral region.

Usage

```
spec_op(  
    mrs_data,  
    xlim = NULL,  
    operator = "sum",  
    freq_scale = "ppm",  
    mode = "re"  
)
```

Arguments

mrs_data	MRS data.
xlim	spectral range to be integrated (defaults to full range).
operator	can be "sum" (default), "mean", "l2", "max", "min" or "max-min".
freq_scale	units of xlim, can be : "ppm", "hz" or "points".
mode	spectral mode, can be : "re", "im", "mod" or "cplx".

Value

an array of integral values.

spin_sys	<i>Create a spin system object for pulse sequence simulation.</i>
----------	---

Description

Create a spin system object for pulse sequence simulation.

Usage

```
spin_sys(spin_params, ft, ref)
```

Arguments

spin_params	an object describing the spin system properties.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.

Value

spin system object.

spm_pve2categorical	<i>Convert SPM style segmentation files to a single categorical image where the numerical values map as: 0) Other, 1) CSF, 2) GM and 3) WM.</i>
---------------------	---

Description

Convert SPM style segmentation files to a single categorical image where the numerical values map as: 0) Other, 1) CSF, 2) GM and 3) WM.

Usage

```
spm_pve2categorical(fname)
```

Arguments

fname any of the segmentation files (eg c1_MY_T1.nii).

Value

nifti object.

ssp	<i>Signal space projection method for lipid suppression.</i>
-----	--

Description

Signal space projection method as described in: Tsai SY, Lin YR, Lin HY, Lin FH. Reduction of lipid contamination in MR spectroscopy imaging using signal space projection. Magn Reson Med 2019 Mar;81(3):1486-1498.

Usage

```
ssp(mrs_data, comps = 5, xlim = c(1.5, 0.8))
```

Arguments

mrs_data	MRS data object.
comps	the number of spatial components to use.
xlim	spectral range (in ppm) covering the lipid signals.

Value

lipid suppressed mrs_data object.

stackplot*Produce a plot with multiple traces.*

Description

Produce a plot with multiple traces.

Usage

```
stackplot(x, ...)
```

Arguments

x	object for plotting.
...	arguments to be passed to methods.

stackplot.fit_result *Plot the fitting results of an object of class fit_result with individual basis set components shown.*

Description

Plot the fitting results of an object of class fit_result with individual basis set components shown.

Usage

```
## S3 method for class 'fit_result'  
stackplot(  
  x,  
  xlim = NULL,  
  y_offset = 0,  
  dyn = 1,  
  x_pos = 1,  
  y_pos = 1,  
  z_pos = 1,  
  coil = 1,  
  n = NULL,  
  sub_bl = FALSE,  
  labels = FALSE,  
  label_names = NULL,  
  sig_col = "black",  
  restore_def_par = TRUE,  
  omit_signals = NULL,  
  combine_lipmm = FALSE,  
  combine_metab = FALSE,
```

```

  mar = NULL,
  show_grid = TRUE,
  grid_nx = NULL,
  grid_ny = NA,
  ...
)

```

Arguments

<code>x</code>	fit_result object.
<code>xlim</code>	the range of values to display on the x-axis, eg <code>xlim = c(4,1)</code> .
<code>y_offset</code>	separate basis signals in the y-axis direction by this value.
<code>dyn</code>	the dynamic index to plot.
<code>x_pos</code>	the x index to plot.
<code>y_pos</code>	the y index to plot.
<code>z_pos</code>	the z index to plot.
<code>coil</code>	the coil element number to plot.
<code>n</code>	single index element to plot (overrides other indices when given).
<code>sub_bl</code>	subtract the baseline from the data and fit (logical).
<code>labels</code>	print signal labels at the right side of the plot.
<code>label_names</code>	provide a character vector of signal names to replace the defaults determined from the basis set.
<code>sig_col</code>	colour of individual signal components.
<code>restore_def_par</code>	restore default plotting par values after the plot has been made.
<code>omit_signals</code>	a character vector of basis signal names to be removed from the plot.
<code>combine_lipmm</code>	combine all basis signals with names starting with "Lip" or "MM".
<code>combine_metab</code>	combine all basis signals with names not starting with "Lip" or "MM".
<code>mar</code>	option to adjust the plot margins. See <code>?par</code> .
<code>show_grid</code>	plot gridlines behind the data (logical). Defaults to TRUE.
<code>grid_nx</code>	number of cells of the grid in x and y direction. When NULL the grid aligns with the tick marks on the corresponding default axis (i.e., tickmarks as computed by <code>axTicks</code>). When NA, no grid lines are drawn in the corresponding direction.
<code>grid_ny</code>	as above.
<code>...</code>	further arguments to plot method.

stackplot.mrs_data *Stackplot plotting method for objects of class mrs_data.*

Description

Stackplot plotting method for objects of class mrs_data.

Usage

```
## S3 method for class 'mrs_data'  
stackplot(  
  x,  
  xlim = NULL,  
  mode = "re",  
  x_units = NULL,  
  fd = TRUE,  
  col = NULL,  
  alpha = NULL,  
  x_offset = 0,  
  y_offset = 0,  
  plot_dim = NULL,  
  x_pos = NULL,  
  y_pos = NULL,  
  z_pos = NULL,  
  dyn = 1,  
  coil = 1,  
  bty = NULL,  
  labels = NULL,  
  lab_cex = 1,  
  right_marg = NULL,  
  bl_lty = NULL,  
  restore_def_par = TRUE,  
  show_grid = NULL,  
  grid_nx = NULL,  
  grid_ny = NA,  
  lwd = NULL,  
  ...  
)
```

Arguments

- x object of class mrs_data.
- xlim the range of values to display on the x-axis, eg xlim = c(4,1).
- mode representation of the complex numbers to be plotted, can be one of: "re", "im", "mod" or "arg".
- x_units the units to use for the x-axis, can be one of: "ppm", "hz", "points" or "seconds".

fd	display data in the frequency-domain (default), or time-domain (logical).
col	set the colour of the line, eg col = rgb(1, 0, 0, 0.5).
alpha	set the line transparency, eg alpha = 0.5 is 50% transparency. Overrides any transparency levels set by col.
x_offset	separate plots in the x-axis direction by this value. Default value is 0.
y_offset	separate plots in the y-axis direction by this value.
plot_dim	the dimension to display on the y-axis, can be one of: "dyn", "x", "y", "z", "coil" or NULL. If NULL (the default) all spectra will be collapsed into the dynamic dimension and displayed.
x_pos	the x index to plot.
y_pos	the y index to plot.
z_pos	the z index to plot.
dyn	the dynamic index to plot.
coil	the coil element number to plot.
bty	option to draw a box around the plot. See ?par.
labels	add labels to each data item.
lab_cex	label size.
right_marg	change the size of the right plot margin.
bl_lty	linetype for the y = 0 baseline trace. A default value NULL results in no baseline being plotted.
restore_def_par	restore default plotting par values after the plot has been made.
show_grid	plot gridlines behind the data (logical). Defaults to TRUE.
grid_nx	number of cells of the grid in x and y direction. When NULL the grid aligns with the tick marks on the corresponding default axis (i.e., tickmarks as computed by axTicks). When NA, no grid lines are drawn in the corresponding direction.
grid_ny	as above.
lwd	plot linewidth.
...	other arguments to pass to the matplot method.

sub_mean_dyncs*Subtract the mean dynamic spectrum from a dynamic series.***Description**

Subtract the mean dynamic spectrum from a dynamic series.

Usage

```
sub_mean_dyncs(mrs_data)
```

Arguments

mrs_data dynamic MRS data.

Value

subtracted data.

sum_coils

Calculate the sum across receiver coil elements.

Description

Calculate the sum across receiver coil elements.

Usage

sum_coils(mrs_data)

Arguments

mrs_data MRS data split across receiver coil elements.

Value

sum across coil elements.

sum_dyncs

Calculate the sum of data dynamics.

Description

Calculate the sum of data dynamics.

Usage

sum_dyncs(mrs_data)

Arguments

mrs_data dynamic MRS data.

Value

sum of data dynamics.

<code>sum_mrs</code>	<i>Sum two mrs_data objects.</i>
----------------------	----------------------------------

Description

Sum two mrs_data objects.

Usage

```
sum_mrs(a, b, force = FALSE)
```

Arguments

- | | |
|--------------------|---|
| <code>a</code> | first mrs_data object to be summed. |
| <code>b</code> | second mrs_data object to be summed. |
| <code>force</code> | set to TRUE to force mrs_data objects to be summed, even if they are in different time/frequency domains. |

Value

`a + b`

<code>sum_mrs_list</code>	<i>Return the sum of a list of mrs_data objects.</i>
---------------------------	--

Description

Return the sum of a list of mrs_data objects.

Usage

```
sum_mrs_list(mrs_list)
```

Arguments

- | | |
|-----------------------|---------------------------|
| <code>mrs_list</code> | list of mrs_data objects. |
|-----------------------|---------------------------|

Value

sum mrs_data object.

`svs_1h_brain_analysis` Standard SVS 1H brain analysis pipeline.

Description

Standard SVS 1H brain analysis pipeline.

Usage

```
svs_1h_brain_analysis(  
  metab,  
  basis = NULL,  
  w_ref = NULL,  
  mri_seg = NULL,  
  mri = NULL,  
  output_dir = NULL,  
  extra = NULL,  
  decimate = NULL,  
  rats_corr = TRUE,  
  ecc = FALSE,  
  comb_dyns = TRUE,  
  hsvd_filt = FALSE,  
  scaleamps = TRUE,  
  te = NULL,  
  tr = NULL,  
  preproc_only = FALSE,  
  method = "ABFIT",  
  opts = NULL  
)
```

Arguments

<code>metab</code>	filepath or mrs_data object containing MRS metabolite data.
<code>basis</code>	basis set object to use for analysis.
<code>w_ref</code>	filepath or mrs_data object containing MRS water reference data.
<code>mri_seg</code>	filepath or nifti object containing segmented MRI data.
<code>mri</code>	filepath or nifti object containing anatomical MRI data.
<code>output_dir</code>	directory path to output fitting results.
<code>extra</code>	data.frame with one row containing additional information to be attached to the fit results table.
<code>decimate</code>	option to decimate the input data by a factor of two. The default value of NULL does not perform decimation unless the spectral width is greater than 20 PPM.
<code>rats_corr</code>	option to perform rats correction, defaults to TRUE.
<code>ecc</code>	option to perform water reference based eddy current correction, defaults to FALSE.

comb_dyncs	option to combine dynamic scans, defaults to TRUE.
hsvd_filt	option to apply hsvc water removal, defaults to FALSE.
scale_amps	option to scale metabolite amplitude estimates, defaults to TRUE.
te	metabolite mrs data echo time in seconds.
tr	metabolite mrs data repetition time in seconds.
preproc_only	only perform the preprocessing steps and omit fitting. The preprocessed metabolite data will be returned in this case.
method	analysis method to use, see fit_mrs help.
opts	options to pass to the analysis method.

Value

a fit_result or mrs_data object depending on the preproc_only option.

svs_1h_brain_batch_analysis

Batch interface to the standard SVS 1H brain analysis pipeline.

Description

Batch interface to the standard SVS 1H brain analysis pipeline.

Usage

```
svs_1h_brain_batch_analysis(
  metab_list,
  w_ref_list = NULL,
  mri_seg_list = NULL,
  mri_list = NULL,
  output_dir_list = NULL,
  extra = NULL,
  ...
)
```

Arguments

metab_list	list of file paths or mrs_data objects containing MRS metabolite data.
w_ref_list	list of file paths or mrs_data objects containing MRS water reference data.
mri_seg_list	list of file paths or nifti objects containing segmented MRI data.
mri_list	list of file paths or nifti objects containing anatomical MRI data.
output_dir_list	list of directory paths to output fitting results.
extra	a data frame with the same number of rows as metab_list, containing additional information to be attached to the fit results table.
...	additional options to be passed to the svs_1h_brain_analysis function.

Value

a list of fit_result objects.

td2fd

Transform time-domain data to the frequency-domain.

Description

Transform time-domain data to the frequency-domain.

Usage

```
td2fd(mrs_data)
```

Arguments

mrs_data MRS data in time-domain representation.

Value

MRS data in frequency-domain representation.

tdsr

Time-domain spectral registration.

Description

An implementation of the method published by Near et al MRM 73:44-50 (2015).

Usage

```
tdsr(mrs_data, ref = NULL, xlim = c(4, 0.5), max_t = 0.2)
```

Arguments

mrs_data MRS data to be corrected.
ref optional MRS data to use as a reference, the mean of all dynamics is used if this argument is not supplied.
xlim optional frequency range to perform optimisation, set to NULL to use the full range.
max_t truncate the FID when longer than max_t to reduce time taken.

Value

a list containing the corrected data; phase and shift values in units of degrees and Hz respectively.

td_conv_filt*Time-domain convolution based filter.*

Description

Time-domain convolution based filter described by: Marion D, Ikura M, Bax A. Improved solvent suppression in one-dimensional and twodimensional NMR spectra by convolution of time-domain data. J Magn Reson 1989;84:425-430.

Usage

```
td_conv_filt(mrs_data, K = 25, ext = 1)
```

Arguments

mrs_data	MRS data to be filtered.
K	window width in data points.
ext	point separation for linear extrapolation.

varpro_3_para_opts*Return a list of options for VARPRO based fitting with 3 free parameters.*

Description

Return a list of options for VARPRO based fitting with 3 free parameters.

Usage

```
varpro_3_para_opts(
  nstart = 20,
  init_damping = 2,
  maxiters = 200,
  max_shift = 5,
  max_damping = 5,
  anal_jac = FALSE,
  bl_smth_pts = 80
)
```

Arguments

nstart	position in the time-domain to start fitting, units of data points.
init_damping	starting value for the global Gaussian line-broadening term - measured in Hz.
maxiters	maximum number of levmar iterations to perform.
max_shift	maximum global shift allowed, measured in Hz.
max_damping	maximum damping allowed, FWHM measured in Hz.
anal_jac	option to use the analytic or numerical Jacobian (logical).
bl_smth_pts	number of data points to use in the baseline smoothing calculation.

Value

list of options.

varpro_basic_opts *Return a list of options for a basic VARPRO analysis.*

Description

Return a list of options for a basic VARPRO analysis.

Usage

```
varpro_basic_opts(method = "fd_re", nnls = TRUE)
```

Arguments

method	one of "td", "fd", "fd_re".
nnls	restrict basis amplitudes to non-negative values.

Value

full list of options.

varpro_opts*Return a list of options for VARPRO based fitting.***Description**

Return a list of options for VARPRO based fitting.

Usage

```
varpro_opts(
    nstart = 20,
    init_g_damping = 2,
    maxiters = 200,
    max_shift = 5,
    max_g_damping = 5,
    max_ind_damping = 5,
    anal_jac = TRUE,
    bl_smth_pts = 80
)
```

Arguments

nstart	position in the time-domain to start fitting, units of data points.
init_g_damping	starting value for the global Gaussian line-broadening term - measured in Hz.
maxiters	maximum number of levmar iterations to perform.
max_shift	maximum shift allowed to each element in the basis set, measured in Hz.
max_g_damping	maximum permitted global Gaussian line-broadening.
max_ind_damping	maximum permitted Lorentzian line-broadening for each element in the basis set, measured in Hz.
anal_jac	option to use the analytic or numerical Jacobian (logical).
bl_smth_pts	number of data points to use in the baseline smoothing calculation.

Value

list of options.

Examples

```
varpro_opts(nstart = 10)
```

vec2mrs_data	<i>Convert a vector into a mrs_data object.</i>
--------------	---

Description

Convert a vector into a mrs_data object.

Usage

```
vec2mrs_data(  
  vec,  
  fs = def_fs(),  
  ft = def_ft(),  
  ref = def_ref(),  
  nuc = def_nuc(),  
  dyns = 1,  
  fd = FALSE  
)
```

Arguments

vec	the data vector.
fs	sampling frequency in Hz.
ft	transmitter frequency in Hz.
ref	reference value for ppm scale.
nuc	resonant nucleus.
dyns	replicate the data across the dynamic dimension.
fd	flag to indicate if the matrix is in the frequency domain (logical).

Value

mrs_data object.

write_basis	<i>Write a basis object to an LCModel .basis formatted file.</i>
-------------	--

Description

Write a basis object to an LCModel .basis formatted file.

Usage

```
write_basis(basis, basis_file, fwhmba = 0.1)
```

Arguments

- basis basis object to be exported.
- basis_file path to basis file to be generated.
- fwhmba parameter used by LCModel.

<code>write_basis_tqn</code>	<i>Generate a basis file using TARQUIN.</i>
------------------------------	---

Description

Generate a basis file using TARQUIN.

Usage

```
write_basis_tqn(basis_file, metab_data, opts = NULL)
```

Arguments

- basis_file filename of the basis file to be generated.
- metab_data MRS data object to match the generated basis parameters.
- opts list of options to pass to TARQUIN.

Examples

```
## Not run:  
write_basis_tqn('test.basis',mrs_data,c("--echo","0.04"))  
  
## End(Not run)
```

<code>write_mrs</code>	<i>Write MRS data object to file.</i>
------------------------	---------------------------------------

Description

Write MRS data object to file.

Usage

```
write_mrs(mrs_data, fname, format = NULL, force = FALSE)
```

Arguments

mrs_data	object to be written to file, or list of mrs_data objects.
fname	one or more filenames to output.
format	string describing the data format. Must be one of the following : "nifti", "dpt", "lcm_raw", "rds". If not specified, the format will be guessed from the filename extension.
force	set to TRUE to overwrite any existing files.

write_mrs_nifti *Write MRS data object to file in NIFTI format.*

Description

Write MRS data object to file in NIFTI format.

Usage

```
write_mrs_nifti(mrs_data, fname)
```

Arguments

mrs_data	object to be written to file.
fname	the filename of the output NIFTI MRS data.

zero_fade_spec *Fade a spectrum to zero by frequency domain multiplication with a tanh function. Note this operation distorts data points at the end of the FID.*

Description

Fade a spectrum to zero by frequency domain multiplication with a tanh function. Note this operation distorts data points at the end of the FID.

Usage

```
zero_fade_spec(mrs_data, start_ppm, end_ppm)
```

Arguments

mrs_data	data to be faded.
start_ppm	start point of the fade in ppm units.
end_ppm	end point of the fade in ppm units.

Value

modified mrs_data object.

zero_nzoc*Zero all non-zero-order coherences.*

Description

Zero all non-zero-order coherences.

Usage

```
zero_nzoc(sys, rho)
```

Arguments

sys	spin system object.
rho	density matrix.

Value

density matrix.

zf*Zero-fill MRS data in the time domain.*

Description

Zero-fill MRS data in the time domain.

Usage

```
zf(x, factor = 2)

## S3 method for class 'list'
zf(x, factor = 2)

## S3 method for class 'mrs_data'
zf(x, factor = 2)

## S3 method for class 'basis_set'
zf(x, factor = 2)
```

Arguments

x	input mrs_data or basis_set object.
factor	zero-filling factor, factor of 2 returns a dataset with twice the original data points.

Value

zero-filled data.

zf_xy

Zero-fill MRSI data in the k-space x-y direction.

Description

Zero-fill MRSI data in the k-space x-y direction.

Usage

```
zf_xy(mrs_data, factor = 2)
```

Arguments

mrs_data	MRSI data.
factor	zero-filling factor, a factor of 2 returns a dataset with twice the original points in the x-y directions. Factors smaller than one are permitted, such that a factor of 0.5 returns half the k-space points in the x-y directions.

Value

zero-filled data.

Index

* datasets

spant_mpress_drift, 147

abfit_opts, 9, 12
abfit_opts_v1_9_0, 12
acquire, 12
add_noise, 13
align, 13
apodise_xy, 14
append_basis, 15
append_coils, 15
append_dyns, 16
apply_axes, 16
apply_mrs, 17
apply_pvc, 17
Arg.mrs_data, 18
array2mrs_data, 18
auto_phase, 19

back_extrap_ar, 20
basis2mrs_data, 20
bbase, 21
bc_als, 22
bc_constant, 22
beta2lw, 23
bin_spec, 23

calc_coil_noise_cor, 24
calc_coil_noise_sd, 24
calc_ed_from_lambda, 25
calc_peak_info_vec, 25
calc_sd_poly, 26
calc_spec_diff, 26
calc_spec_snr, 27
check_lcm, 28
check_tqn, 28
circ_mask, 28
collapse_to_dyns, 29
comb_coils, 29
comb_fit_list_fit_tables, 30

comb_fit_list_result_tables, 31
comb_fit_tables, 32
comb_metab_ref, 32
Conj.mrs_data, 33
conv_mrs, 33
crop_spec, 34
crop_td_pts, 34
crop_td_pts_pot, 35
crop_xy, 35
crossprod_3d, 36

decimate_mrs_fd, 36
decimate_mrs_td, 37
def_acq_paras, 37, 124, 140, 141, 143, 145
def_fs, 38
def_ft, 38
def_N, 39
def_nuc, 39
def_ref, 39
dicom_reader, 40
diff_mrs, 40
downsample_mrs_fd, 41
downsample_mrs_td, 41

ecc, 42
elliptical_mask, 42
est_noise_sd, 43

fd2td, 43
fd_conv_filt, 44
fitamps, 44
fit_diags, 45
fit_mrs, 45
fit_res2csv, 47
fp_phase, 47
fp_phase_correct, 48
fp_scale, 48
fs, 49
ft_dyns, 49
ft_shift, 50

ft_shift_mat, 50
gausswin_2d, 51
gen_F, 51
gen_F_xy, 52
get_1h_brain_basis_paras, 52
get_1h_brain_basis_paras_v1, 53
get_1h_brain_basis_paras_v2, 53
get_1h_brain_basis_paras_v3, 54
get_2d_psf, 54
get_acq_paras, 55
get_dyns, 55
get_even_dyns, 56
get_fh_dyns, 56
get_fit_map, 57
get_fp, 57
get_gaussian_pulse, 58
get_head_dyns, 58
get_lcm_cmd, 59
get_metab, 59
get_mol_names, 59
get_mol_paras, 60
get_mrs_affine, 62
get_mrsi2d_seg, 60
get_mrsi_voi, 61
get_mrsi_voxel, 61
get_mrsi_voxel_xy_psf, 62
get_odd_dyns, 63
get_ref, 63
get_seg_ind, 64
get_sh_dyns, 64
get_slice, 65
get_subset, 65
get_svs_voi, 66
get_tail_dyns, 67
get_td_amp, 67
get_tqn_cmd, 68
get_uncoupled_mol, 68
get_voi_cog, 69
get_voi_seg, 69
get_voi_seg_psf, 70
get_voxel, 70
grid_shift_xy, 72
gridplot, 71
gridplot.mrs_data, 71
hsvd, 72
hsvd_filt, 73
hsvd_vec, 74
hz, 74
ift_shift, 75
ift_shift_mat, 75
Im.mrs_data, 76
image.mrs_data, 76
img2kspace_xy, 77
int_spec, 78
interleave_dyns, 78
inv_even_dyns, 79
inv_odd_dyns, 79
is.def, 80
is_fd, 80
kspace2img_xy, 81
l2_reg, 81
lb, 82
lw2alpha, 83
lw2beta, 83
mask_dyns, 84
mask_fit_res, 84
mask_xy, 85
mask_xy_mat, 85
mat2mrs_data, 86
max_mrs, 86
max_mrs_interp, 87
mean.list, 87
mean.mrs_data, 88
mean_dyn_blocks, 89
mean_dyn_pairs, 89
mean_dyns, 88
mean_mrs_list, 90
median_dyns, 90
Mod.mrs_data, 91
mod_td, 91
mrs_data2basis, 92
mrs_data2mat, 92
mrs_data2vec, 93
mvfftshift, 93
mvifftshift, 94
n2coord, 94
Ncoils, 95
Ndyns, 95
nifti_flip_lr, 95
Npts, 96
Nspec, 96

Nx, 97
 Ny, 97
 Nz, 97
 ortho3, 98
 ortho3_inter, 99
 peak_info, 99
 pg_extrap_xy, 100
 phase, 101
 plot.fit_result, 101
 plot.mrs_data, 103
 plot_bc, 105
 plot_slice_fit, 105
 plot_slice_fit_inter, 106
 plot_slice_map, 106
 plot_slice_map_inter, 107
 plot_voi_overlay, 109
 plot_voi_overlay_seg, 109
 ppm, 110
 precomp, 110
 print.fit_result, 111
 print.mrs_data, 111
 qn_states, 112
 rats, 112
 Re.mrs_data, 113
 re_weighting, 124
 read_basis, 114
 read_basis_ac, 114
 read_ima_coil_dir, 115
 read_ima_dyn_dir, 115
 read_lcm_coord, 116
 read_mrs, 116
 read_mrs_tqn, 117
 read_siemens_txt_hdr, 118
 read_tqn_fit, 119
 read_tqn_result, 119
 recon_twix_2d_mrsi, 120
 rectangular_mask, 120
 rep_array_dim, 121
 rep_dyn, 121
 rep_mrs, 122
 resample_img, 122
 resample_voi, 123
 reslice_to_mrs, 123
 reson_table2mrs_data, 124
 rm_dyns, 125
 scale_amp_molal_pvc, 125
 scale_amp_molar, 126
 scale_amp_ratio, 126
 scale_amp_water_ratio, 127
 scale_mrs_amp, 127
 scale_spec, 128
 sd, 128
 sd.mrs_data, 129
 seconds, 129
 seq_cpmg_ideal, 130
 seq_mega_press_ideal, 130
 seq_press_ideal, 131
 seq_pulse_acquire, 132
 seq_pulse_acquire_31p, 132
 seq_slaser_ideal, 133
 seq_spin_echo_ideal, 133
 seq_spin_echo_ideal_31p, 134
 seq_steam_ideal, 134
 set_def_acq_paras, 135
 set_lcm_cmd, 135
 set_lw, 136
 set_mask_xy_mat, 136
 set_precomp_mode, 137
 set_precomp_verbose, 137
 set_ref, 137
 set_td_pts, 138
 set_tqn_cmd, 138
 shift, 139
 shift_basis, 139
 sim_basis, 140
 sim_basis_1h_brain, 140
 sim_basis_1h_brain_press, 141
 sim_basis_tqn, 142
 sim_brain_1h, 142
 sim_mol, 143
 sim_noise, 144
 sim_resonances, 145
 sim_zero, 146
 sort_basis, 146
 spant (spant-package), 8
 spant-package, 8
 spant_abfit_benchmark, 147
 spant_mpress_drift, 147
 spant_simulation_benchmark, 148
 spec_decomp, 148
 spec_op, 149
 spin_sys, 149
 spm_pve2categorical, 150

ssp, 150
stackplot, 151
stackplot.fit_result, 151
stackplot.mrs_data, 153
sub_mean_dyns, 154
sum_coils, 155
sum_dyns, 155
sum_mrs, 156
sum_mrs_list, 156
svs_1h_brain_analysis, 157
svs_1h_brain_batch_analysis, 158

td2fd, 159
td_conv_filt, 160
tdsr, 159

varpro_3_para_opts, 160
varpro_basic_opts, 161
varpro_opts, 162
vec2mrs_data, 163

write_basis, 163
write_basis_tqn, 164
write_mrs, 164
write_mrs_nifti, 165

zero_fade_spec, 165
zero_nzoc, 166
zf, 166
zf_xy, 167