Package 'qMRI'

December 9, 2024

```
Type Package
Title Methods for Quantitative Magnetic Resonance Imaging ('qMRI')
Version 1.2.7.8
Date 2024-12-09
Maintainer Karsten Tabelow <karsten.tabelow@wias-berlin.de>
Depends R (>= 3.5), awsMethods (>= 1.0), methods, parallel
Imports or o.nifti (>= 0.9), stringr, aws (>= 2.4), adimpro (>= 0.9)
LazyData TRUE
Description Implementation of methods for estimation of quantitative maps
     from Multi-Parameter Mapping (MPM) acquisitions (Weiskopf et al. (2013)
     <a href="doi:10.3389/fnins.2013.00095">doi:10.3389/fnins.2013.00095</a>) and analysis of Inversion Recovery MRI data.
     Usage of the package is described in
     Polzehl and Tabelow (2023),
      ``Magnetic Resonance Brain Imaging", 2nd Edition, Chapter 6 and 7, Springer, Use R! Series.
     <doi:10.1007/978-3-031-38949-8>.
     J. Polzehl and K. Tabelow (2023), ``Magnetic Resonance Brain Imaging -
     Modeling and Data Analysis Using R: Code and Data."
     <doi:10.20347/WIAS.DATA.6> provides extensive example code and data.
License GPL (>= 2)
Copyright This package is Copyright (C) 2015-2024 Weierstrass
     Institute for Applied Analysis and Stochastics.
URL https://www.wias-berlin.de/research/ats/imaging/
Suggests covr, testthat, knitr, rmarkdown
VignetteBuilder knitr
RoxygenNote 6.1.1
NeedsCompilation yes
Author Joerg Polzehl [aut],
     Karsten Tabelow [aut, cre],
     WIAS Berlin [cph, fnd]
```

Repository CRAN

Date/Publication 2024-12-09 12:20:05 UTC

Contents

	qMRI-package					2
	awssigmc					6
	calculateQI					8
	colMT					10
	estimateESTATICS .					11
	estimateIR					13
	estimateIRfluid					16
	estimateIRsolid					18
	estimateIRsolidfixed					21
	extract-methods					23
	generateIRData					26
	MREdisplacement					27
	readMPMData					30
	smoothESTATICS					33
	smoothIRSolid					36
	writeESTATICS					37
	writeQI					39
Index						43
illuex						40
qMRI-	package <i>N</i>	Aethods fo	r Quantita	ative Magneti	c Resonance Imaging ('qMRI'))

Description

Implementation of methods for estimation of quantitative maps from Multi-Parameter Mapping (MPM) acquisitions (Weiskopf et al. (2013) <doi:10.3389/fnins.2013.00095>) and analysis of Inversion Recovery MRI data. Usage of the package is described in Polzehl and Tabelow (2023), "Magnetic Resonance Brain Imaging", 2nd Edition, Chapter 6 and 7, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8>. J. Polzehl and K. Tabelow (2023), "Magnetic Resonance Brain Imaging - Modeling and Data Analysis Using R: Code and Data." <doi:10.20347/WIAS.DATA.6> provides extensive example code and data.

Details

The DESCRIPTION file:

Package: qMRI Type: Package

Title: Methods for Quantitative Magnetic Resonance Imaging ('qMRI')

Version: 1.2.7.8 Date: 2024-12-09

Authors@R: c(person("Joerg", "Polzehl", role = c("aut"), email = "joerg.polzehl@wias-berlin.de"), person("Karsten", "

Maintainer: Karsten Tabelow <karsten.tabelow@wias-berlin.de> Depends: R (>= 3.5), awsMethods (>= 1.0), methods, parallel

Imports: oro.nifti (>= 0.9), stringr, aws (>= 2.4), adimpro (>= 0.9)

LazyData: TRUE

Description: Implementation of methods for estimation of quantitative maps from Multi-Parameter Mapping (MPM) ac

License: GPL (>= 2)

Copyright: This package is Copyright (C) 2015-2024 Weierstrass Institute for Applied Analysis and Stochastics.

URL: https://www.wias-berlin.de/research/ats/imaging/

Suggests: covr, testthat, knitr, rmarkdown

VignetteBuilder: knitr RoxygenNote: 6.1.1

Author: Joerg Polzehl [aut], Karsten Tabelow [aut, cre], WIAS Berlin [cph, fnd]

Index of help topics:

MREdisplacement Calculate the motion induced signal phase for

IR-MRE in biphasic material

awssigmc Estimate noise variance for multicoil MR

systems

calculateQI Obtain quantitative maps from estimated

 ${\tt ESTATICS} \ parameters.$

colMT MT map color scheme

estimateESTATICS Estimate parameters in the ESTATICS model.

estimateIR Estimate IRMRI parameters

estimateIRfluid Estimate parameters in Inversion Recovery MRI

experiments model for CSF voxel

estimateIRsolid Estimate parameters in Inversion Recovery MRI

experiments mixture model for non-fluid voxel

estimateIRsolidfixed Estimate mixture parameter in Inversion

Recovery MRI experiments mixture model for

non-fluid voxel

extract.ANY-method Methods to extract information from objects of

class '"MPMData"', '"ESTATICSModel"',
'"sESTATICSModel"', '"qMaps"', '"IRdata"',

'"IRfluid"' and '"IRmixed"'.

generateIRData generate IR MRI example data

qMRI-package Methods for Quantitative Magnetic Resonance

Imaging ('qMRI')

readIRData Prepare IRMRI dataset

readMPMData Read experimental Multi-Parameter Mapping (MPM)

data.

smoothESTATICS Adaptive smoothing of ESTATICS parameters and

MPM data

smoothIRSolid Smooth object generated by function

'estimateIRsolid'

writeESTATICS Write maps of ESTATICS parameters in

standardized form as NIfTI files.

writeQI Write estimated maps in standardized form as

NIfTI files.

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>

N"org Polzehl <polzehl@wias-berlin.de>

Maintainer: Karsten Tabelow <karsten.tabelow@wias-berlin.de>

References

Weiskopf, N.; Suckling, J.; Williams, G.; Correia, M. M.; Inkster, B.; Tait, R.; Ooi, C.; Bullmore, E. T. & Lutti, A. Quantitative multi-parameter mapping of R1, PD(*), MT, and R2(*) at 3T: a multi-center validation. Front Neurosci, Wellcome Trust Centre for Neuroimaging, UCL Institute of Neurology, University College London, UK., 2013, 7, 95

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6 and 7, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

aws

```
dataDir <- system.file("extdata",package="qMRI")</pre>
  set file names for T1w, MTw and PDw images
t1Names <- paste0("t1w_",1:8,".nii.gz")</pre>
mtNames <- paste0("mtw_",1:6,".nii.gz")</pre>
pdNames <- paste0("pdw_",1:8,".nii.gz")</pre>
t1Files <- file.path(dataDir, t1Names)</pre>
mtFiles <- file.path(dataDir, mtNames)</pre>
pdFiles <- file.path(dataDir, pdNames)</pre>
#
  file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")</pre>
maskFile <- file.path(dataDir, "mask.nii.gz")</pre>
  Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE \leftarrow c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
         2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR \leftarrow rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
#
    read MPM example data
library(qMRI)
```

```
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                    maskFile, TR = TR, TE = TE,
                    FA = FA, verbose = FALSE)
# Estimate Parameters in the ESTATICS model
#
modelMPM <- estimateESTATICS(mpm, method = "NLR")</pre>
#
  smooth maps of ESTATICS Parameters
setCores(2)
modelMPMsp1 <- smoothESTATICS(modelMPM,</pre>
                               kstar = 16,
                               alpha = 0.004,
                               patchsize=1,
                               verbose = TRUE)
#
   resulting ESTATICS parameter maps for central coronal slice
#
#
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
oldpar <- par(mfrow=c(2,4),mar=c(3,3,3,1),mgp=c(2,1,0))
on.exit(par(oldpar))
pnames <- c("T1","MT","PD","R2star")</pre>
modelCoeff <- extract(modelMPM, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(pnames[i])
   modelCoeff <- extract(modelMPMsp1, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(paste("smoothed",pnames[i]))
   }
}
#
#
   Compute quantitative maps (R1, R2star, PD, MT)
qMRIMaps <- calculateQI(modelMPM,
                         b1File = B1File,
                         TR2 = 3.4)
qMRISmoothedp1Maps <- calculateQI(modelMPMsp1,</pre>
                                      b1File = B1File,
                                      TR2 = 3.4)
   resulting quantitative maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
par(mfrow=c(2,4),mar=c(3,3,3,1),mgp=c(2,1,0))
nmaps <- c("R1","R2star","PD","MT")</pre>
qmap <- extract(qMRIMaps,nmaps)</pre>
for (i in 1:4) rimage(qmap[[i]][,11,],main=nmaps[i])
```

6 awssigme

```
qmap <- extract(qMRISmoothedp1Maps,nmaps)
for (i in 1:4) rimage(qmap[[i]][,11,],main=paste("Smoothed",nmaps[i]))
}
par(oldpar)</pre>
```

awssigmc

Estimate noise variance for multicoil MR systems

Description

The distribution of image intensity values S_i divided by the noise standard deviation in K-space σ in dMRI experiments is assumed to follow a non-central chi-distribution with 2L degrees of freedom and noncentrality parameter η , where L refers to the number of receiver coils in the system and $\sigma\eta$ is the signal of interest. This is an idealization in the sense that each coil is assumed to have the same contribution at each location. For realistic modeling L should be a locally smooth function in voxel space that reflects the varying local influence of the receiver coils in the the reconstruction algorithm used.

The functions assume L to be known and estimate either a local (function awslsigmc) or global (function awssigmc) σ employing an assumption of local homogeneity for the noncentrality parameter η .

Function afsigmc implements estimates from Aja-Fernandez (2009). Function aflsigmc implements the estimate from Aja-Fernandez (2013).

Usage

voxel extentions

Arguments

vext

У	3D array, usually obtained from an object of class dwi as obj@si[,,,i] for some i, i.e. one 3D image from an dMRI experiment. Alternatively a vector of length sum(mask) may be suppied together with a brain mask in mask.
steps	number of steps in adaptive weights smoothing, used to reveal the unerlying mean structure.
mask	restrict computations to voxel in mask, if is.null(mask) all voxel are used. In function afsigmc mask should refer to background for method %in% c("modem1chi", "bkm2chi", "bkm10 and to voxel within the head for method=="modevn".
ncoils	number of coils, or equivalently number of effective degrees of freedom of non- central chi distribution divided by 2.

awssigme 7

lambda scale parameter in adaptive weights smoothing

h0 initial bandwidth

verbose if verbose==TRUE density plots and quantiles of local estimates of sigma are

provided.

trace if trace==TRUE intermediate results for each step are returned in component

tergs for all voxel in mask.

sequence if sequence=TRUE a vector of estimates for the noise standard deviation sigma

for the individual steps is returned instead of the final value only.

hadj adjustment factor for bandwidth (chosen by bw.nrd) in mode estimation

q quantile to be used for interquantile-differences.

quantile of distribution of actual sum of weights $N_i = \sum_j w_{ij}$ in adaptive

smoothing. Only voxel i with $N_i > q_{qni}(N_i)$ are used for variance estimation.

Should be larger than 0.5.

method in case of function awssigme the method for variance estimation, either "VAR"

(variance) or "MAD" (mean absolute deviation). In function afsigmc see last

column in Table 2 in Aja-Fernandez (2009).

minni Minimum sum of weights for updating values of sigma.

hsig Bandwidth of the median filter.

sigma Initial estimate for sigma

family One of "Gauss" or "NCchi" (default) defining the probability distribution to

use.

u if verbose==TRUE an array of noncentrality paramters for comparisons. Internal

use for tests only

Value

a list with components

sigma either a scalar or a vector of estimated noise standard deviations.

theta the estimated mean structure

Author(s)

J\"org Polzehl <polzehl@wias-berlin.de>

References

K. Tabelow, H.U. Voss, J. Polzehl, Local estimation of the noise level in MRI using structural adaptation, Medical Image Analysis, 20 (2015), pp. 76–86.

See Also

aws

8 calculateQI

calculateQI	Obtain quantitative maps from estimated ESTATICS parameters.

Description

Quantitaive imaging parameters are calculated from the estimated parameters in the ESTATICS model. This involves a correction for magnetic field inhomogeneities if the information is provided in argument b1File and use of a second of a second recovery delay TR2 in case of Dual-Exitation FLASH measurements (Helms 2008).

Usage

```
calculateQI(mpmESTATICSModel, b1File = NULL, TR2 = 0, verbose = TRUE)
```

Arguments

mpmESTATICSModel

Object of class 'ESTATICSModel' as returned from function estimateESTATICS.

b1File (optional) Name of a file containing a B1-field inhomogeneity map (.nii)

TR2 second recovery delay TR2 in case of Dual-Exitation FLASH measurements.

verbose logical: Monitor process.

Value

List with components

b1Map

R1 Estimated map of R1

R2star Estimated map of R2star

PD Estimated map of PD

MT Estimated map of delta (if MT-series was used)

model Type of ESTATICS model used

t1Files filenames T1
mtFiles filenames MT
pdFiles filenames PD
mask brainmask

and class-attribute 'qMaps'.

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>

λ"org Polzehl <polzehl@wias-berlin.de>

calculateQI 9

References

Helms, G.; Dathe, H.; Kallenberg, K. & Dechent, P. High-Resolution Maps of Magnetization Transfer with Inherent Correction for RF Inhomogeneity and T1 Relaxation Obtained from 3D FLASH MRI Magn. Res. Med., 2008, 60, 1396-1407

Weiskopf, N.; Suckling, J.; Williams, G.; Correia, M. M.; Inkster, B.; Tait, R.; Ooi, C.; Bullmore, E. T. & Lutti, A. Quantitative multi-parameter mapping of R1, PD(*), MT, and R2(*) at 3T: a multi-center validation. Front Neurosci, Wellcome Trust Centre for Neuroimaging, UCL Institute of Neurology, University College London, UK., 2013, 7, 95

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_6>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

readMPMData, estimateESTATICS, smoothESTATICS, writeESTATICS, awsLocalSigma

```
dataDir <- system.file("extdata",package="qMRI")</pre>
#
   set file names for T1w, MTw and PDw images
t1Names <- paste0("t1w_",1:8,".nii.gz")
mtNames <- paste0("mtw_",1:6,".nii.gz")
pdNames <- paste0("pdw_",1:8,".nii.gz")
t1Files <- file.path(dataDir, t1Names)</pre>
mtFiles <- file.path(dataDir, mtNames)</pre>
pdFiles <- file.path(dataDir, pdNames)</pre>
#
   file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")</pre>
maskFile <- file.path(dataDir, "mask0.nii.gz")</pre>
   Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE \leftarrow c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR \leftarrow rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
#
    read MPM example data
library(qMRI)
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                    maskFile, TR = TR, TE = TE,
                    FA = FA, verbose = FALSE)
#
```

10 colMT

```
# limit calculations to voxel in the central coronal slice
# to reduce execution time of the example
#
# Estimate Parameters in the ESTATICS model
modelMPM <- estimateESTATICS(mpm, method = "NLR")</pre>
#
  resulting ESTATICS parameter maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
oldpar <- par(mfrow=c(2,2), mar=c(3,3,3,1), mgp=c(2,1,0))
on.exit(par(oldpar))
pnames <- c("T1","MT","PD","R2star")</pre>
modelCoeff <- extract(modelMPM, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(pnames[i])
   }
}
#
  Compute quantitative maps (R1, R2star, PD, MT)
#
qMRIMaps <- calculateQI(modelMPM,</pre>
                         b1File = B1File,
                         TR2 = 3.4)
#
  resulting quantitative maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
par(mfrow=c(2,2), mar=c(3,3,3,1), mgp=c(2,1,0))
nmaps <- c("R1","R2star","PD","MT")</pre>
qmap <- extract(qMRIMaps,nmaps)</pre>
for (i in 1:4){
   rimage(qmap[[i]][,11,],main=nmaps[i])
}
par(oldpar)
```

colMT

MT map color scheme

Description

Color map implementing the color scheme for MT maps. This is the plasma scale from Matplotlib (pyplot) generated by function plasma from package **viridisLite**.

estimateESTATICS 11

Usage

colMT

Format

A vector with 256 RGB color values.

estimateESTATICS

Estimate parameters in the ESTATICS model.

Description

Evaluation of the ESTATICS model (Weisskopf (2013) using nonlinear least squares regression and a quasi-likelihood approach assuming a noncentral chi- or a Rician distribuion for the data. The latter should be preferred in case of low SNR (high resolution) data to avoid biased parameter estimates. Quasi-likelihood estimation requires a specification of the scale parameter sigma of the data distribution.

Usage

```
estimateESTATICS(mpmdata, TEScale = 100, dataScale = 1000, method = c("NLR", "QL"),
                 sigma = NULL, L = 1, maxR2star = 50,
                 varest = c("RSS", "data"), verbose = TRUE)
```

Arguments

mpmdata	Object of class MPMData as created by readMPMData.
TEScale	scale factor for TE (used for improved numerical stability)
dataScale	scale factor for image intensities (used for improved numerical stability)
method	either "NLR" or "QL". Specifies non-linear regression or quasi-likelihood.
sigma	scale parameter sigma of signal distribution (either a scalar or a 3D array). (only needed in case of $method="QL"$.)
L	effective number of receiver coils ($2*L$ is degrees of freedom of the signal distribution). L=1 for Rician distribution. (only needed in case of method="QL".)
maxR2star	maximum value allowed for the R2star parameter in the ESTATICS model.
varest	For parameter covariance estimation use either residual sum of squares (RSS) or estimate variances for T1, MT (is available) and PD from higest intensity images using function awsLocalSigmafrom package aws .
verbose	logical: Monitor process.

12 estimateESTATICS

Value

list with components

modelCoeff Estimated parameter maps

invCov map of inverse covariance matrices rsigma map of residual standard deviations

isConv convergence indicator map

isThresh logical map indicating where R2star==maxR2star.

sdim image dimension
nFiles number of images
t1Files vector of T1 filenames
pdFiles vector of PD filenames
mtFiles vector of MT filenames

model model used (depends on specification of MT files)

maskFile filename of brain mask

mask brain mask sigma sigma

TR TR values
TE TE values

FA Flip angles (FA)

TEScale TEScale dataScale dataScale

and class-attribute 'ESTATICSModel'

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>

λ"org Polzehl <polzehl@wias-berlin.de>

References

Weiskopf, N.; Suckling, J.; Williams, G.; Correia, M. M.; Inkster, B.; Tait, R.; Ooi, C.; Bullmore, E. T. & Lutti, A. Quantitative multi-parameter mapping of R1, PD(*), MT, and R2(*) at 3T: a multi-center validation. Front Neurosci, Wellcome Trust Centre for Neuroimaging, UCL Institute of Neurology, University College London, UK., 2013, 7, 95

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_6>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

estimateIR 13

See Also

readMPMData, calculateQI, smoothESTATICS, writeESTATICS, awsLocalSigma

```
dataDir <- system.file("extdata",package="qMRI")</pre>
# set file names for T1w, MTw and PDw images
t1Names <- paste0("t1w_",1:8,".nii.gz")</pre>
mtNames <- paste0("mtw_",1:6,".nii.gz")</pre>
pdNames <- paste0("pdw_",1:8,".nii.gz")
t1Files <- file.path(dataDir, t1Names)</pre>
mtFiles <- file.path(dataDir, mtNames)</pre>
pdFiles <- file.path(dataDir, pdNames)</pre>
# file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")</pre>
maskFile <- file.path(dataDir, "mask0.nii.gz")</pre>
# Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE <- c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR <- rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
#
    read MPM example data
#
library(qMRI)
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                    maskFile, TR = TR, TE = TE,
                    FA = FA, verbose = FALSE)
   Estimate Parameters in the ESTATICS model
modelMPM <- estimateESTATICS(mpm, method = "NLR")</pre>
# Alternatively using Quasi-Likelihood
sigma <- 50
modelMPMQL <- estimateESTATICS(mpm, method = "QL",</pre>
                   sigma = array(sigma,mpm$sdim), L = 1)
```

14 estimateIR

Description

Parameter estimation (intensity, relaxation rate, proportion of fluid) in Inversion Recovery MRI data.

Usage

Arguments

8	
IRdataobj	4D array of IRMRI signals. First dimension corresponds to Inversion times (InvTime).
TEScale	Internal scale factor for Echo Times. This influences parameter scales in numerical calculations.
dataScale	Internal scale factor for MR signals. This influences parameter scales in numerical calculations.
method	Either "NLS" for nonlinear least squares (ignores Rician bias) or "QL" for Quasi-Likelihood. The second option is more accurate but requires additional information and is computationally more expensive.
varest	Method to, in case of method="QR", estimate sigmaif not provided. Either from residual sums of squares ("RSS") or MR signals ("data") using function varest from package aws. Only to be used in case that no image registration was needed as preprocessing.
fixed	Should adaptive smoothing performed for Sx and Rx maps and fx maps reestimated afterwards ?
smoothMethod	Either "PAWS" or "Depth". the second option is not yet implemented.
kstar	number of steps used in PAWS
alpha	significance level for decisions in aws algorithm (suggestion: between 1e-5 and 0.025)

TRUE: restrict smoothing to segments from segmentation, FALSE: restrict smooth-

Details

bysegment

verbose

This function implements the complete pipeline of IRMRI anlysis.

Logical. Provide some runtime diagnostics.

ing to solid mask.

Value

List of class "IRmixed" with components

IRdata 4D array containing the IRMRI data, first dimension refers to inversion times

InvTimes vector of inversion times

segm segmentation codes, 1 for CSF, 2 for GM, 3 for WM, 0 for out of brain

estimateIR 15

sigma	noise standard deviation, if not specified estimated fron CSF areas in image with largest inversion time
L	effective number of coils
fx	Array of fluid proportions
Sx	Array of maximal signals
Rx	Array of relaxation rates
Sf	Global estimate of maximal fluid signal
Rf	Global estimate of fluid relaxation rate
ICovx	Covariance matrix of estimates fx, Sx and Rx.
sigma	Array of provided or estimated noise standard deviations
Convx	Array of convergence indicators
rsdx	Residual standard deviations

The arrays contain entries for all voxel with segments%in%1:3.

Author(s)

```
Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>
```

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_7>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

estimateIRfluid, estimateIRsolid, estimateIRsolidfixed, smoothIRSolid

```
## runs about 30 seconds
dataDir0 <- system.file("extdataIR", package = "qMRI")
dataDir <- tempdir("IRdata")
library(oro.nifti)
library(qMRI)
segm <- readNIfTI(file.path(dataDir0,"Brainweb_segm"))
Sf <- 900
Rf <- 0.000285
Sgm <- 400
Rgm <- 0.00075
fgm <- .15
Swm <- 370
Rwm <- 0.0011
fwm <- .05
InvTimes0 <- c(100, 200, 400, 600, 800, 1200, 1600, 2000, 2500, 3000,</pre>
```

16 estimateIRfluid

```
3500, 4000, 4500, 5000, 6000, 15000)
nTimes <- length(InvTimes0)</pre>
sigma <- 40
## generate IR signal
IRdata <- generateIRData(segm, c(Sf,Rf), c(fgm,Rgm,Sgm), c(fwm,Rwm,Swm), InvTimes0, sigma)</pre>
for(i in 1:9) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR0",i)))
for(i in 10:nTimes) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR",i)))
## generate IRdata object
t1Files <- list.files(dataDir,"*.nii.gz",full.names=TRUE)</pre>
segmFile <- file.path(dataDir0,"Brainweb_segm")</pre>
IRdata <- readIRData(t1Files, InvTimes0, segmFile, sigma=sigma,</pre>
                      L=1, segmCodes=c("CSF","GM","WM"))
## estimate all
sIRmix <- estimateIR(IRdata, method="QL")</pre>
```

estimateIRfluid

Estimate parameters in Inversion Recovery MRI experiments model for CSF voxel

Description

The Inversion Recovery MRI signal in voxel containing only CSF follows is modeled as $S_{InvTime} = par[1] * abs(1 - 2 * exp(-InvTime*par[2])) dependings on two parameters. These parameters are assumed to be tissue (and scanner) dependent.$

Usage

```
estimateIRfluid(IRdataobj, TEScale = 100, dataScale = 1000, method = c("NLR", "QL"), varest = c("RSS", "data"), verbose = TRUE, lower = c(0, 0), upper = c(2, 2))
```

Arguments

IRdataobj	Object of class "IRdata" as generated by function readIRData.
TEScale	Internal scale factor for Echo Times. This influences parameter scales in numerical calculations.
dataScale	Internal scale factor for MR signals. This influences parameter scales in numerical calculations.
method	Either "NLS" for nonlinear least squares (ignores Rician bias) or "QL" for Quasi-Likelihood. The second option is more accurate but requires additional information and is computationally more expensive.

estimateIRfluid 17

varest Method to, in case of method="QR", estimate sigmaif not provided. Either from

residual sums of squares ("RSS") or MR signals ("data") using function varest from package aws. Only to be used in case that no image registration was needed

as preprocessing.

verbose Logical. Provide some runtime diagnostics.

lower Lower bounds for parameter values.

upper Upper bounds for parameter values.

Details

The Inversion Recovery MRI signal in voxel containing only CSF follows is modeled as $S_{InvTime} = par[1] * abs(1 - 2 * exp(-InvTime*par[2])) dependings on two parameters. These parameters are assumed to be tissue (and scanner) dependent.$

Value

List of class IRfluid with components

IRdata 4D array containing the IRMRI data, first dimension refers to inversion times

InvTimes vector of inversion times

segm segmentation codes, 1 for CSF, 2 for GM, 3 for WM, 0 for out of brain

sigma noise standard deviation, if not specified estimated fron CSF areas in image with

largest inversion time

L effective number of coils

Sf Global estimate of maximal fluid signal Rf Global estimate of fluid relaxation rate

Sx Array of maximal signals
Rx Array of relaxation rates

sigma Array of provided or estimated noise standard deviations

Convx Array of convergence indicators

method "NLS" for nonlinear regression or "QL" for quasi likelihood.

varest Method used for variance estimation

The arrays only contain entries for fluid voxel.

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. doi:10.1007/978-3-031-38949-8_7.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

18 estimateIRsolid

See Also

 ${\tt estimateIR}, {\tt estimateIR} solid, {\tt estimateIR} solid {\tt fixed}, {\tt smoothIR} Solid$

Examples

```
dataDir0 <- system.file("extdataIR", package = "qMRI")</pre>
dataDir <- tempdir("IRdata")</pre>
library(oro.nifti)
library(qMRI)
segm <- readNIfTI(file.path(dataDir0,"Brainweb_segm"))</pre>
Sf <- 900
Rf <- 0.000285
Sgm <- 400
Rgm <- 0.00075
fgm <- .15
Swm <- 370
Rwm <- 0.0011
fwm <- .05
InvTimes0 <- c(100, 200, 400, 600, 800, 1200, 1600, 2000, 2500, 3000,
              3500, 4000, 4500, 5000, 6000, 15000)
nTimes <- length(InvTimes0)</pre>
sigma <- 40
## generate IR signal
IRdata <- generateIRData(segm, c(Sf,Rf), c(fgm,Rgm,Sgm), c(fwm,Rwm,Swm), InvTimes0, sigma)</pre>
for(i in 1:9) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR0",i)))\\
for(i in 10:nTimes) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR",i)))
## generate IRdata object
t1Files <- list.files(dataDir,"*.nii.gz",full.names=TRUE)
segmFile <- file.path(dataDir0,"Brainweb_segm")</pre>
IRdata <- readIRData(t1Files, InvTimes0, segmFile, sigma=sigma,</pre>
                      L=1, segmCodes=c("CSF","GM","WM"))
## estimate fluid
setCores(2) # parallel mode using 2 threads
IRfluid <- estimateIRfluid(IRdata, method="NLR", verbose=FALSE)</pre>
cat("Estimated parameters Sf:", IRfluid$Sf,
                         " Rf:", IRfluid$Rf, "\n")
```

estimateIRsolid

Estimate parameters in Inversion Recovery MRI experiments mixture model for non-fluid voxel

Description

The Inversion Recovery MRI signal in non-fluid voxel follows is modeled as a mixture of a fluid and a solid compartment.

estimateIRsolid 19

Usage

```
estimateIRsolid(IRfluidobj, TEScale = 100, dataScale = 1000, verbose = TRUE, lower = c(0, 0, 0), upper = c(0.95, 2, 2))
```

Arguments

IRfluidobj Object of class "IRfluid" as generated by function estimateIRfluid.

TEScale Internal scale factor for Echo Times. This influences parameter scales in numer-

ical calculations.

dataScale Internal scale factor for MR signals. This influences parameter scales in numer-

ical calculations.

verbose Logical. Provide some runtime diagnostics.

lower Lower bounds for parameter values. upper Upper bounds for parameter values.

Details

The Inversion Recovery MRI signal in non-fluid voxel follows is modeled as a mixture of a fluid and a solid compartment. The function calculates estimates of the maximum signal and recovery rate for the solid compartment and a mixture coefficient (proportion of fluid) for all voxel with segment%in%2: 3using results from function estimateIRfluid.

Value

List of class IRmixed with components

IRdata 4D array containing the IRMRI data, first dimension refers to inversion time	IRdata	4D array containing the IRMRI d	data, first dimension refers to inversion time
---	--------	---------------------------------	--

InvTimes vector of inversion times

segm segmentation codes, 1 for CSF, 2 for GM, 3 for WM, 0 for out of brain

sigma noise standard deviation, if not specified estimated fron CSF areas in image with

largest inversion time

L effective number of coils
fx Array of fluid proportions
Sx Array of maximal signals
Rx Array of relaxation rates

Sf Global estimate of maximal fluid signal Rf Global estimate of fluid relaxation rate

ICovx Covariance matrix of estimates fx, Sx and Rx.

sigma Array of provided or estimated noise standard deviations

Convx Array of convergence indicators rsdx Residual standard deviations

method "NLS" for nonlinear regression or "QL" for quasi likelihood.

varest Method used for variance estimation

The arrays contain entries for all voxel with segments%in%1:3.

20 estimateIRsolid

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>

λ"org Polzehl <polzehl@wias-berlin.de>

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. doi:10.1007/978-3-031-38949-8_7.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

estimateIRfluid, estimateIR, estimateIRsolidfixed, smoothIRSolid

```
## runs about 7 seconds
dataDir0 <- system.file("extdataIR", package = "qMRI")</pre>
dataDir <- tempdir("IRdata")</pre>
library(oro.nifti)
library(qMRI)
segm <- readNIfTI(file.path(dataDir0,"Brainweb_segm"))</pre>
Sf <- 900
Rf <- 0.000285
Sgm <- 400
Rgm <- 0.00075
fgm <- .15
Swm <- 370
Rwm <- 0.0011
fwm <- .05
InvTimes0 <- c(100, 200, 400, 600, 800, 1200, 1600, 2000, 2500, 3000,
               3500, 4000, 4500, 5000, 6000, 15000)
nTimes <- length(InvTimes0)</pre>
sigma <- 40
## generate IR signal
IRdata <- generateIRData(segm, c(Sf,Rf), c(fgm,Rgm,Sgm), c(fwm,Rwm,Swm), InvTimes0, sigma)</pre>
for(i in 1:9) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR0",i)))
for(i in 10:nTimes) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR",i)))
## generate IRdata object
t1Files <- list.files(dataDir,"*.nii.gz",full.names=TRUE)</pre>
segmFile <- file.path(dataDir0,"Brainweb_segm")</pre>
IRdata <- readIRData(t1Files, InvTimes0, segmFile, sigma=sigma,</pre>
                      L=1, segmCodes=c("CSF","GM","WM"))
## estimate fluid
setCores(2) # parallel mode using 2 threads
IRfluid <- estimateIRfluid(IRdata, method="NLR", verbose=FALSE)</pre>
cat("Estimated parameters Sf:", IRfluid$Sf,
                         " Rf:", IRfluid$Rf, "\n")
```

estimateIRsolidfixed 21

```
## estimate solid
IRmix <- estimateIRsolid(IRfluid, verbose=FALSE)</pre>
```

mixture model for non-fluid voxel

Description

Reestimate proportion of fluid with Sx and Rx fixed after smoothing.

Usage

```
estimateIRsolidfixed(IRmixedobj, TEScale = 100, dataScale = 1000,
verbose = TRUE, lower = c(0), upper = c(0.95))
```

Arguments

IRmixedobj	Object of class "IRmixed" as generated by function smoothIRSolid or estimateIRsolid.
TEScale	Internal scale factor for Echo Times. This influences parameter scales in numerical calculations.
dataScale	Internal scale factor for MR signals. This influences parameter scales in numerical calculations.
verbose	Logical. Provide some runtime diagnostics.
lower	lower bound for fx (fluid proportion)
upper	upper bound for fx (fluid proportion)

Value

List of class "IRmixed" components

IRdata	4D array containing the IRMRI data, first dimension refers to inversion times
InvTimes	vector of inversion times
segm	segmentation codes, 1 for CSF, 2 for GM, 3 for WM, 0 for out of brain
sigma	noise standard deviation, if not specified estimated fron CSF areas in image with largest inversion time
L	effective number of coils
fx	Array of fluid proportions
Sx	Array of maximal signals
Rx	Array of relaxation rates
Sf	Global estimate of maximal fluid signal
Rf	Global estimate of fluid relaxation rate

22 estimateIRsolidfixed

ICovx	Covariance matrix of estimates fx, Sx and Rx.
sigma	Array of provided or estimated noise standard deviations
Convx	Array of convergence indicators
rsdx	Residual standard deviations
method	"NLS" for nonlinear regression or "QL" for quasi likelihood.
varest	Method used for variance estimation

The arrays contain entries for all voxel with segments%in%1:3.

Author(s)

```
Karsten Tabelow <tabelow@wias-berlin.de>

λ"org Polzehl <polzehl@wias-berlin.de>
```

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_7>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

estimateIRfluid, estimateIRsolid, estimateIR, smoothIRSolid, readIRData

```
## runs about 11 seconds
dataDir0 <- system.file("extdataIR", package = "qMRI")</pre>
dataDir <- tempdir("IRdata")</pre>
library(oro.nifti)
library(qMRI)
segm <- readNIfTI(file.path(dataDir0,"Brainweb_segm"))</pre>
Sf <- 900
Rf <- 0.000285
Sgm <- 400
Rgm <- 0.00075
fgm < - .15
Swm <- 370
Rwm <- 0.0011
fwm <- .05
InvTimes0 <- c(100, 200, 400, 600, 800, 1200, 1600, 2000, 2500, 3000,
              3500, 4000, 4500, 5000, 6000, 15000)
nTimes <- length(InvTimes0)</pre>
sigma <- 40
## generate IR signal
IRdata <- generateIRData(segm, c(Sf,Rf), c(fgm,Rgm,Sgm), c(fwm,Rwm,Swm), InvTimes0, sigma)</pre>
for(i in 1:9) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR0",i)))
for(i in 10:nTimes) writeNIfTI(as.nifti(IRdata[i,,,]),
```

extract-methods 23

```
file.path(dataDir,paste0("IR",i)))
## generate IRdata object
t1Files <- list.files(dataDir,"*.nii.gz",full.names=TRUE)</pre>
segmFile <- file.path(dataDir0,"Brainweb_segm")</pre>
IRdata <- readIRData(t1Files, InvTimes0, segmFile, sigma=sigma,</pre>
                      L=1, segmCodes=c("CSF","GM","WM"))
## estimate fluid
setCores(2) # parallel mode using 2 threads
IRfluid <- estimateIRfluid(IRdata, method="NLR", verbose=FALSE)</pre>
cat("Estimated parameters Sf:", IRfluid$Sf,
                         " Rf:", IRfluid$Rf, "\n")
## estimate solid
IRmix <- estimateIRsolid(IRfluid, verbose=FALSE)</pre>
## smoothing
sIRmix <- smoothIRSolid(IRmix, alpha=1e-4, partial=FALSE, verbose=FALSE)
## reestimate
sIRmix <- estimateIRsolidfixed(sIRmix, verbose=FALSE)</pre>
```

Description

The extract-methods extract and/or compute specified statistics from object of class "MPMData", "ESTATICSModel", "gMaps", "IRdata", "IRfluid" and "IRmixed". The [methods can be used to reduce objects of class "MPMData", "ESTATICSModel", "sESTATICSModel", "qMaps", "IRdata", "IRfluid" and "IRmixed" such that they contain a subcube of data and results.

Usage

```
## S3 method for class 'MPMData'
extract(x, what, ...)
## S3 method for class 'ESTATICSModel'
extract(x, what, ...)
## S3 method for class 'sESTATICSModel'
extract(x, what, ...)
## S3 method for class 'qMaps'
extract(x, what, ...)
## S3 method for class 'IRdata'
extract(x, what, ...)
## S3 method for class 'IRfluid'
extract(x, what, ...)
## S3 method for class 'IRmixed'
extract(x, what, ...)
## S3 method for class 'MPMData'
```

24 extract-methods

```
x[i, j, k, ...]
## S3 method for class 'ESTATICSModel'
x[i, j, k, ...]
## S3 method for class 'sESTATICSModel'
x[i, j, k, ...]
## S3 method for class 'qMaps'
x[i, j, k, ...]
## S3 method for class 'IRdata'
x[i, j, k, tind, ...]
## S3 method for class 'IRfluid'
x[i, j, k, ...]
## S3 method for class 'IRfluid'
x[i, j, k, ...]
```

Arguments

Х	object of class "MPMData", "ESTATICSModel", "sESTATICSModel" or "qMaps".
what	Character vector of of names of statistics to extract. See Methods Section for details.
i	index vector for first spatial dimension
j	index vector for second spatial dimension
k	index vector for third spatial dimension
tind	index vector for inversion times
	additional parameters, currently unused.

Value

A list with components carrying the names of the options specified in argument what.

Methods

```
class(x) = "ANY" Returns a warning for extract
class(x) = "MPMData" Depending the occurence of names in what a list with the specified components is returned
```

- ddata: mpm data
- sdim: dimension of image cube
- nFiles: number of images / image files
- t1Files: character filenames of t1Files
- pdFiles: character filenames of pdFiles
- mtFiles: character filenames of mtFiles
- model: Number of the ESTATICS model that can be used
- maskFile: character filenames of maskFile
- · mask: mask
- TR: vector of TR values
- TE: vector of TE values

extract-methods 25

• FA: vector of FA values

class(x) = "ESTATICSModel" Depending the occurrence of names in what a list with the specified
 components is returned

- modelCoeff: Estimated parameter maps
- invCov: map of inverse covariance matrices
- rsigma: map of residual standard deviations
- isConv: convergence indicator map
- isThresh: logical map indicating where R2star==maxR2star
- sdim: image dimension
- nFiles: number of images
- t1Files: vector of T1 filenames
- pdFiles: vector of PD filenames
- mtFiles: vector of MT filenames
- model: model used (depends on specification of MT files)
- · maskFile: filename of brain mask
- mask: brain mask
- sigma: standard deviation sigma
- L: effective number of receiver coils L
- TR: TR values
- TE: TE values
- FA: Flip angles (FA)
- TEScale: TEScale
- dataScale: dataScale

class(x) = "sESTATICSModel" Depending the occurrence of names in what a list with the specified components is returned

- modelCoeff: Estimated parameter maps
- invCov: map of inverse covariance matrices
- rsigma: map of residual standard deviations
- isConv: convergence indicator map
- bi: Sum of weights map from AWS/PAWS
- smoothPar: smooting parameters used in AWS/PAWS
- smoothedData: smoothed mpmData
- isThresh: logical map indicating where R2star==maxR2star
- sdim: image dimension
- nFiles: number of images
- t1Files: vector of T1 filenames
- pdFiles: vector of PD filenames
- mtFiles: vector of MT filenames
- model: model used (depends on specification of MT files)
- · maskFile: filename of brain mask
- mask: brain masksigma: sigma

26 generateIRData

- L: effective number of receiver coils L
- TR: TR values
- TE: TE values
- FA: Flip angles (FA)
- TEScale: TEScale
- dataScale: dataScale

class(x) = "qMaps" Depending the occurence of names in what a list with the specified components is returned

- b1Map: b1Map
- R1: Estimated map of R1
- R2star: Estimated map of R2star
- PD: Estimated map of PD
- MT: Estimated map of delta (if MT-series was used)
- model: Type of ESTATICS model used
- t1Files: filenames T1
- mtFiles: filenames MT
- pdFiles: filenames PD
- · mask: brainmask

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>

λ"org Polzehl <polzehl@wias-berlin.de>

References

J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_7>.

generateIRData

generate IR MRI example data

Description

The function generates IR MRI example data for specified parameters

Usage

```
generateIRData(segm, pCSF, pGM, pWM, InvTimes, sigma = 40)
```

MREdisplacement 27

Arguments

segm array containing segmentation resuts for an 3D MRI template. Contains 1 for

CSF, 2 for Gray Matter and 3 for White Matter

pCSF Parameters (S,R) for CSF

pGM Parameters (f,R,S) for Gray Matter pWM Parameters (f,R,S) for White Matter

InvTimes Vector of Inversion Times, length nTimes

sigma Noise standard variation

Value

array with dimension c(nTimes,dim(segm))

Note

used in examples for IR functions

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>

See Also

estimateIRfluid, estimateIRsolid, estimateIR, smoothIRSolid, readIRData

MREdisplacement	Calculate the motion induced signal phase for IR-MRE in biphasic
	material

Description

The function takes magnitude images and phase images (as NIfTI files) recordet with inversion IT1=Inf and a second inversion time IT2 that nulls the fluid signal. Tissue parameters (Relaxation rates) are extracted from an object of class "IRmixed" calculated from data of a related IRMRI experiment.

Usage

28 MREdisplacement

Arguments

MagnFiles1	Filenames of magnitude images recorded with inversion time IT=Inf.
PhaseFiles1	Filenames of phase images recorded with inversion time IT=Inf.
MagnFiles2	Filenames of magnitude images recorded with inversion time IT=IT2.
PhaseFiles2	Filenames of phase images recorded with inversion time IT=IT2.
TI2	Inversion time used for MagnFiles2 and PhaseFiles2. IT2 should be selected to extinguish the signal intendity for fluid.
IRmixobj	Object of class "IRmixed" obtained from a related IRMRI experiment.
method	Either "full" or "approx"
rescale	Logical, do we need to rescale phase images ?
verbose	Report scale range of phase images

Details

The first 4 arguments need to be vectors of filenames of identical length with files containing compatible 3D NIfTI images. Object IRmixobj needs to contain a components segm and Rx of compatible dimension that need to be registered to the MRE images.

Value

A list of class "IRMREbiphasic" with components

phisolid displacement solid phifluid displacement fluid

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>

References

J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. doi:10.1007/978-3-031-38949-8_7.

See Also

estimate IR fluid, estimate IR solid, estimate IR solid fixed, smooth IR Solid fixed, smo

readIRData 29

readIRData	Prepare IRMRI dataset	
------------	-----------------------	--

Description

The function reads IRMRI images given as NIfTI files in t1Files, inversion times and segmentation image(s) aund prepares an object class "IRdata"

Usage

Arguments

t1Files	Names of NIfTI files containing the recorded images.
InvTimes	Corresponding inversion times
segmFile	Either a NIfTI file containing a segmentation into GM, WM and CSF or three files containing probability maps for GM, WM and CSF
sigma	Noise standard deviation
L	Effective number of coils, L=1 assumes a Rician signal distribution
segmCodes	sequence of tissue code in segmFile

Value

A list of class "IRdata" with components

IRdata	4D array containing the IRMRI data, first dimension refers to inversion times
InvTimes	vector of inversion times
segm	segmentation codes, 1 for CSF, 2 for GM, 3 for WM, 0 for out of brain
sigma	noise standard deviation, if not specified estimated fron CSF areas in image with largest inversion time
L	effective number of coils

Author(s)

```
Karsten Tabelow < tabelow@wias-berlin.de>
J\"org Polzehl < polzehl@wias-berlin.de>
```

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_7>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

30 readMPMData

See Also

estimateIRfluid, estimateIRsolid, estimateIR, smoothIRSolid

Examples

```
dataDir0 <- system.file("extdataIR", package = "qMRI")</pre>
dataDir <- tempdir("IRdata")</pre>
library(oro.nifti)
library(qMRI)
segm <- readNIfTI(file.path(dataDir0,"Brainweb_segm"))</pre>
Sf <- 900
Rf <- 0.000285
Sgm <- 400
Rgm <- 0.00075
fgm <- .15
Swm <- 370
Rwm <- 0.0011
fwm <- .05
InvTimes0 <- c(100, 200, 400, 600, 800, 1200, 1600, 2000, 2500, 3000,
               3500, 4000, 4500, 5000, 6000, 15000)
nTimes <- length(InvTimes0)</pre>
sigma <- 40
## generate IR signal
IRdata <- generateIRData(segm, c(Sf,Rf), c(fgm,Rgm,Sgm), c(fwm,Rwm,Swm), InvTimes0, sigma)</pre>
for(i in 1:9) writeNIfTI(as.nifti(IRdata[i,,,]),
                           file.path(dataDir,paste0("IR0",i)))
for(i in 10:nTimes) writeNIfTI(as.nifti(IRdata[i,,,]),
                           file.path(dataDir,paste0("IR",i)))
## generate IRdata object
t1Files <- list.files(dataDir,"*.nii.gz",full.names=TRUE)</pre>
segmFile <- file.path(dataDir0,"Brainweb_segm")</pre>
IRdata <- readIRData(t1Files, InvTimes0, segmFile, sigma=sigma,</pre>
                      L=1, segmCodes=c("CSF", "GM", "WM"))
```

readMPMData

Read experimental Multi-Parameter Mapping (MPM) data.

Description

The function reads data generated in Multimodal Parameter Mapping (MPM) experiments.

Usage

```
readMPMData(t1Files = NULL, pdFiles = NULL, mtFiles = NULL, maskFile = NULL, TR = NULL, TE = NULL, FA = NULL, wghts = NULL, verbose = TRUE)
```

readMPMData 31

Arguments

t1Files	Vector of filenames corresponding to T1 weighted images (in Nifti-Format) with varying TE
pdFiles	Vector of filenames corresponding to PD weighted images (in Nifti-Format) with varying TE
mtFiles	optional Vector of filenames corresponding to MT weighted images (in Nifti-Format) with varying $\ensuremath{\mathrm{TE}}$
maskFile	optional filename for mask (in Nifti-Format)
TR	optional numeric TR vector, if omitted information is extracted from .nii files if possible
TE	optional numeric TE vector, if omitted information is extracted from .nii files if possible
FA	optional numeric FA (flip-angle) vector, if omitted information is extracted from .nii files if possible
wghts	optional weights for MPM data volumes. Only needed is volumes have different data variance, e.g., in case of averages of multiple acquisitions.
verbose	logical - provide information on progress

Value

List with components

ddata	mpm data	
sdim	dimension of image cube	
nFiles	number of images / image files	
t1Files	character - filenames of t1Files	
pdFiles	character - filenames of pdFiles	
mtFiles	character - filenames of mtFiles	
model	Number of the ESTATICS model that can be used	
maskFile	character - filenames of maskFile	
mask	mask	
TR	vector of TR values	
TE	vector of TE values	
FA	vector of FA values	
and class-attribute 'mpmData'		

Author(s)

Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>

32 readMPMData

References

Weiskopf, N.; Suckling, J.; Williams, G.; Correia, M. M.; Inkster, B.; Tait, R.; Ooi, C.; Bullmore, E. T. & Lutti, A. Quantitative multi-parameter mapping of R1, PD(*), MT, and R2(*) at 3T: a multi-center validation. Front Neurosci, Wellcome Trust Centre for Neuroimaging, UCL Institute of Neurology, University College London, UK., 2013, 7, 95

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6, Springer, Use R! Series. to-401:10.1007/978-3-031-38949-8_6.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

estimateESTATICS, calculateQI, smoothESTATICS, writeESTATICS, awsLocalSigma

```
dataDir <- system.file("extdata",package="qMRI")</pre>
  set file names for T1w, MTw and PDw images
t1Names <- paste0("t1w_",1:8,".nii.gz")</pre>
mtNames <- paste0("mtw_",1:6,".nii.gz")</pre>
pdNames <- paste0("pdw_",1:8,".nii.gz")</pre>
t1Files <- file.path(dataDir, t1Names)</pre>
mtFiles <- file.path(dataDir, mtNames)</pre>
pdFiles <- file.path(dataDir, pdNames)</pre>
# file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")</pre>
maskFile <- file.path(dataDir, "mask.nii.gz")</pre>
  Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE <- c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR \leftarrow rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
#
    read MPM example data
#
library(qMRI)
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                     maskFile, TR = TR, TE = TE,
                     FA = FA, verbose = FALSE)
```

smoothESTATICS 33

smoothESTATICS	Adaptive smoothing of ESTATICS parameters and MPM data	

Description

Performs adaptive smoothing of parameter maps in the ESTATICS model and if mpmData is specified these data. Implements both vectorized variants of the Adaptive Weights Smoothing (AWS, Polzehl and Spokoiny (2006)) and patchwise AWS (PAWS, Polzehl et al (2018)) algorithms with weighting schemes determined by the estimated parameter maps and their covariances.

Usage

Arguments

mpmESTATICSModel

Object of class 'ESTATICSModel' as returned from function estimateESTATICS.

mpmData (optional) Object of class MPMData as created by readMPMData from which the

parameter maps were obtained.

kstar Maximum number of steps.

alpha specifies the scale parameter for the adaptation criterion. smaller values are

more restrictive.

patchsize Patchsize in PAWS, 0 corresponds to AWS, alternative values are 1 and 2.

mscbw bandwidth for 3D median smoother used to stabilize the covariance estimates.

wghts (optional) voxel size if measurments are not isotropic.

verbose logical - provide information on progress

Value

list with components

modelCoeff Estimated parameter maps

invCov map of inverse covariance matrices

isConv convergence indicator map

bi Sum of weights map from AWS/PAWS smoothPar smooting parameters used in AWS/PAWS

smoothedData smoothed mpmData
sdim image dimension
nFiles number of images
t1Files vector of T1 filenames
pdFiles vector of PD filenames

34 smoothESTATICS

```
vector of MT filenames
mtFiles
mode1
                  model used (depends on specification of MT files)
maskFile
                  filename of brain mask
                  brain mask
mask
sigma
                  sigma
L
                 L
TR
                 TR values
                 TE values
ΤE
FΑ
                 Flip angles (FA)
TEScale
                  TEScale
dataScale
                  dataScale
and class-attribute 'sESTATICSModel'
```

Author(s)

```
Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>
```

References

- J. Polzehl, V. Spokoiny, Propagation-separation approach for local likelihood estimation, Probab. Theory Related Fields 135 (3), (2006), pp. 335–362.
- J. Polzehl, K. Papafitsorus, K. Tabelow (2018). Patch-wise adaptive weights smoothing. WIAS-Preprint 2520.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_6>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

```
readMPMData, estimateESTATICS
```

```
dataDir <- system.file("extdata",package="qMRI")
#
# set file names for T1w, MTw and PDw images
#
t1Names <- paste0("t1w_",1:8,".nii.gz")
mtNames <- paste0("mtw_",1:6,".nii.gz")
pdNames <- paste0("pdw_",1:8,".nii.gz")
t1Files <- file.path(dataDir, t1Names)
mtFiles <- file.path(dataDir, mtNames)
pdFiles <- file.path(dataDir, pdNames)
#</pre>
```

smoothESTATICS 35

```
# file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")</pre>
maskFile <- file.path(dataDir, "mask.nii.gz")</pre>
# Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE \leftarrow c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR \leftarrow rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
#
    read MPM example data
#
library(qMRI)
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                    maskFile, TR = TR, TE = TE,
                    FA = FA, verbose = FALSE)
  Estimate Parameters in the ESTATICS model
modelMPM <- estimateESTATICS(mpm, method = "NLR")</pre>
  smooth maps of ESTATICS Parameters
#
setCores(2)
modelMPMsp1 <- smoothESTATICS(modelMPM,</pre>
                                kstar = 16,
                                alpha = 0.004,
                                patchsize=1,
                                verbose = TRUE)
   resulting ESTATICS parameter maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
oldpar <- par(mfrow=c(2,4),mar=c(3,3,3,1),mgp=c(2,1,0))
on.exit(par(oldpar))
pnames <- c("T1","MT","PD","R2star")</pre>
modelCoeff <- extract(modelMPM, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(pnames[i])
modelCoeff <- extract(modelMPMsp1, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(paste("smoothed",pnames[i]))
par(oldpar)
```

36 smoothIRSolid

smoothIRSolid	Smooth object generated by function estimateIRsolid	

Description

Adaptive smoothing of Rx and Sx maps over WM and GM areas.

Usage

Arguments

IRmixedobj object of class IRmixed generated by function estimateIRsolid

kstar number of steps for AWS algorithm

patchsize patchsize in paws

alpha significance level for decisions in aws algorithm (suggestion: between 1e-5 and

0.025)

bandwidth for 3D median smoother used to stabilize the covariance estimates.

TRUE: restrict smoothing to segments from segmentation, FALSE: restrict smooth-

ing to solid mask.

partial TRUE: ignore information concerning parameter fx when smoothing.

verbose logical: Monitor process.

Details

This uses a vectorized version of the AWS algorithm that emloys inverse covariance estimates of the estimated parameters. Local smoothing is done for Rx and Sx maps in ergs which can be assumed to be locally smooth within tissue. No smoothing for fx maps since they may vary.

Value

an object of class "IRmixed", but with components Sx and Rx replaced. The object carries an additional component bi containing an array of sum of weights characterizing the amount of smoothing.

Author(s)

```
Karsten Tabelow <tabelow@wias-berlin.de>

J\"org Polzehl <polzehl@wias-berlin.de>
```

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 7, Springer, Use R! Series. doi:10.1007/978-3-031-38949-8_7.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

writeESTATICS 37

See Also

estimateIRfluid, estimateIRsolid, estimateIRsolidfixed, estimateIR

Examples

```
## runs about 10 seconds
dataDir0 <- system.file("extdataIR", package = "qMRI")</pre>
dataDir <- tempdir("IRdata")</pre>
library(oro.nifti)
library(qMRI)
segm <- readNIfTI(file.path(dataDir0,"Brainweb_segm"))</pre>
Sf <- 900
Rf <- 0.000285
Sgm <- 400
Rgm <- 0.00075
fgm <- .15
Swm <- 370
Rwm <- 0.0011
fwm <- .05
InvTimes0 <- c(100, 200, 400, 600, 800, 1200, 1600, 2000, 2500, 3000,
              3500, 4000, 4500, 5000, 6000, 15000)
nTimes <- length(InvTimes0)</pre>
sigma <- 40
## generate IR signal
IRdata <- generateIRData(segm, c(Sf,Rf), c(fgm,Rgm,Sgm), c(fwm,Rwm,Swm), InvTimes0, sigma)</pre>
for(i in 1:9) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR0",i)))
for(i in 10:nTimes) writeNIfTI(as.nifti(IRdata[i,,,]),
                          file.path(dataDir,paste0("IR",i)))
## generate IRdata object
t1Files <- list.files(dataDir,"*.nii.gz",full.names=TRUE)
segmFile <- file.path(dataDir0, "Brainweb_segm")</pre>
IRdata <- readIRData(t1Files, InvTimes0, segmFile, sigma=sigma,</pre>
                      L=1, segmCodes=c("CSF","GM","WM"))
## estimate fluid
setCores(2) # parallel mode using 2 threads
IRfluid <- estimateIRfluid(IRdata, method="NLR", verbose=FALSE)</pre>
cat("Estimated parameters Sf:", IRfluid$Sf,
                         " Rf:", IRfluid$Rf, "\n")
## estimate solid
IRmix <- estimateIRsolid(IRfluid, verbose=FALSE)</pre>
## smoothing
sIRmix <- smoothIRSolid(IRmix, alpha=1e-4, partial=FALSE, verbose=FALSE)</pre>
```

Write maps of ESTATICS parameters in standardized form as NIfTI files.

38 writeESTATICS

Description

R2, ST1, SPD and, if available, SMT-maps are written as compressed NIfTI files into directory the speecified directory. If class(mpmESTATICSModel) == "sESTATICSModel" and an smoothed data are stored in mpmESTATICSModel\$smoothedData the smoothed data are stored as ompressed NIfTI files in dir with filenames assembled using prefix and the names of the data source files.

Usage

```
writeESTATICS(mpmESTATICSModel, dir = NULL, prefix = "estatics", verbose = TRUE)
```

Arguments

mpmESTATICSModel

Object of class 'ESTATICSModel' or 'sESTATICSModel' as returned from func-

tion estimateESTATICS or smoothESTATICS.

dir Directory name (or path) for output.

prefix Prefix for file names

verbose logical - provide information on progress

Value

The function returns NULL

Author(s)

```
Karsten Tabelow <tabelow@wias-berlin.de>
J\"org Polzehl <polzehl@wias-berlin.de>
```

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6, Springer, Use R! Series. <doi:10.1007/978-3-031-38949-8_6>.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

```
readMPMData, estimateESTATICS, smoothESTATICS
```

```
dataDir <- system.file("extdata",package="qMRI")
outDir <- tempdir()
#
# set file names for T1w, MTw and PDw images
#
t1Names <- paste0("t1w_",1:8,".nii.gz")
mtNames <- paste0("mtw_",1:6,".nii.gz")
pdNames <- paste0("pdw_",1:8,".nii.gz")</pre>
```

```
t1Files <- file.path(dataDir, t1Names)
mtFiles <- file.path(dataDir, mtNames)</pre>
pdFiles <- file.path(dataDir, pdNames)</pre>
# file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")</pre>
maskFile <- file.path(dataDir, "mask0.nii.gz")</pre>
  Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE <- c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR \leftarrow rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
#
    read MPM example data
#
library(qMRI)
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                    maskFile, TR = TR, TE = TE,
                    FA = FA, verbose = FALSE)
   Estimate Parameters in the ESTATICS model
#
#
modelMPM <- estimateESTATICS(mpm, method = "NLR")</pre>
#
   resulting ESTATICS parameter maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
oldpar <- par(mfrow=c(2,2), mar=c(3,3,3,1), mgp=c(2,1,0))
on.exit(par(oldpar))
pnames <- c("T1","MT","PD","R2star")</pre>
modelCoeff <- extract(modelMPM, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(pnames[i])
   }
}
#
#
   write ESTATICS parameter maps
writeESTATICS(modelMPM, dir=outDir, prefix="estatics")
par(oldpar)
```

Description

Quantitative R2, R1, PD and, if available, MT-maps are written as compressed NIfTI files into directory the specified directory.

Usage

```
writeQI(qi, dir = NULL, prefix="qmap", verbose = TRUE)
```

Arguments

qi Object of class 'qMaps' as returned from function calculateQI

dir Directory name (or path) for output.

prefix Prefix for file names

verbose logical - provide information on progress

Value

The function returns NULL

Author(s)

```
Karsten Tabelow <tabelow@wias-berlin.de>

λ"org Polzehl <polzehl@wias-berlin.de>
```

References

- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging: Modeling and Data Analysis Using R, 2nd Edition, Chapter 6, Springer, Use R! Series. doi:10.1007/978-3-031-38949-8_6.
- J. Polzehl and K. Tabelow (2023), Magnetic Resonance Brain Imaging Modeling and Data Analysis Using R: Code and Data. <doi:10.20347/WIAS.DATA.6>.

See Also

```
readMPMData, estimateESTATICS, calculateQI
```

```
dataDir <- system.file("extdata",package="qMRI")
outDir <- tempdir()
#
# set file names for T1w, MTw and PDw images
#
t1Names <- paste0("t1w_",1:8,".nii.gz")
mtNames <- paste0("mtw_",1:6,".nii.gz")
pdNames <- paste0("pdw_",1:8,".nii.gz")
t1Files <- file.path(dataDir, t1Names)
mtFiles <- file.path(dataDir, mtNames)
pdFiles <- file.path(dataDir, pdNames)
#</pre>
```

```
# file names of mask and B1 field map
B1File <- file.path(dataDir, "B1map.nii.gz")
maskFile <- file.path(dataDir, "mask0.nii.gz")</pre>
# Acquisition parameters (TE, TR, Flip Angle) for T1w, MTw and PDw images
TE \leftarrow c(2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8,
        2.3, 4.6, 6.9, 9.2, 11.5, 13.8, 16.1, 18.4)
TR \leftarrow rep(25, 22)
FA \leftarrow c(rep(21, 8), rep(6, 6), rep(6, 8))
    read MPM example data
#
library(qMRI)
mpm <- readMPMData(t1Files, pdFiles, mtFiles,</pre>
                    maskFile, TR = TR, TE = TE,
                    FA = FA, verbose = FALSE)
  Estimate Parameters in the ESTATICS model
modelMPM <- estimateESTATICS(mpm, method = "NLR")</pre>
# resulting ESTATICS parameter maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
oldpar <- par(mfrow=c(2,2), mar=c(3,3,3,1), mgp=c(2,1,0))
on.exit(par(oldpar))
pnames <- c("T1","MT","PD","R2star")</pre>
modelCoeff <- extract(modelMPM, "modelCoeff")</pre>
for(i in 1:4){
   rimage(modelCoeff[i,,11,])
   title(pnames[i])
   }
}
#
   Compute quantitative maps (R1, R2star, PD, MT)
#
qMRIMaps <- calculateQI(modelMPM,</pre>
                         b1File = B1File,
                         TR2 = 3.4)
#
   resulting quantitative maps for central coronal slice
if(require(adimpro)){
rimage.options(zquantiles=c(.01,.99), ylab="z")
par(mfrow=c(2,2),mar=c(3,3,3,1),mgp=c(2,1,0))
nmaps <- c("R1","R2star","PD","MT")</pre>
qmap <- extract(qMRIMaps,nmaps)</pre>
for (i in 1:4) rimage(qmap[[i]][,11,],main=nmaps[i])
}
```

```
#
# write qmaps
#
writeQI(qMRIMaps, dir=outDir, prefix="qmap")
par(oldpar)
```

Index

* IO	smoothESTATICS, 33
readMPMData, 30	smoothIRSolid, 36
writeESTATICS, 37	* utililities
writeQI,39	generateIRData, 26
* IR-MRE	readIRData, 29
MREdisplacement, 27	* utilities
* IRMRI	smoothIRSolid, 36
estimateIR, 13	[.ANY-method(extract-methods), 23
estimateIRfluid, 16	[.ESTATICSModel (extract-methods), 23
estimateIRsolid, 18	[.IRdata(extract-methods), 23
estimateIRsolidfixed, 21	[.IRfluid(extract-methods), 23
generateIRData, 26	[.IRmixed(extract-methods), 23
readIRData, 29	[.MPMData(extract-methods), 23
smoothIRSolid, 36	[.qMaps(extract-methods), 23
* datasets	[.sESTATICSModel (extract-methods), 23
colMT, 10	
* manip	aws, 4, 7
extract-methods, 23	awsLocalSigma, 9, 13, 32
* methods	awslsigmc (awssigmc), 6
extract-methods, 23	awssigmc, 6
* models	coloulateOT 0 12 22 40
calculateQI, 8	calculateQI, 8, 13, 32, 40 colMT, 10
estimateESTATICS, 11	COIMI, IO
estimateIR, 13	estimateESTATICS, 8, 9, 11, 32–34, 38, 40
estimateIRfluid, 16	estimateIR, 13, 18, 20, 22, 27, 30, 37
estimateIRsolid, 18	estimateIRfluid, <i>15</i> , 16, <i>19</i> , 20, 22, 27, 28,
estimateIRsolidfixed, 21	30, 37
* model	estimateIRsolid, <i>15</i> , <i>18</i> , 18, <i>21</i> , 22, 27, 28,
smoothESTATICS, 33	30, 37
* package	estimateIRsolidfixed, 15, 18, 20, 21, 28, 37
qMRI-package, 2	extract-methods, 23
* regression	extract.ANY-method (extract-methods), 23
estimateESTATICS, 11	extract.ESTATICSModel
estimateIR, 13	(extract-methods), 23
estimateIRfluid, 16	extract.IRdata(extract-methods), 23
estimateIRsolid, 18	extract.IRfluid(extract-methods), 23
estimateIRsolidfixed, 21	extract.IRmixed(extract-methods), 23
* smooth	extract.MPMData(extract-methods), 23
awssigmc, 6	extract.qMaps(extract-methods), 23

INDEX INDEX

```
extract.sESTATICSModel (extract-methods), 23

generateIRData, 26

MREdisplacement, 27

qMRI (qMRI-package), 2

qMRI-package, 2

readIRData, 16, 22, 27, 29

readMPMData, 9, 11, 13, 30, 33, 34, 38, 40

smoothESTATICS, 9, 13, 32, 33, 38

smoothIRSolid, 15, 18, 20-22, 27, 28, 30, 36

writeESTATICS, 9, 13, 32, 37

writeQI, 39
```