API Documentation

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1 Package meet

This is the Modular EEg Toolkit (MEET) for Python 2.

Only Python 2 is supported at the moment, however modifications should be easy to do.

Dependencies:

- -Python 2
- -Numpy
- -Scipy

Version Compatibility:

I try to avoid incompatibilities when updating functions, this however cannot be totaly avoided from time to time. However functions are thoroughly tested.

Citation:

If you use this software for scientific publications please give proper citation.

In the moment please cite as (or similar)

G. Waterstraat, 2014. Modular EEg toolkit (MEET) for Python.

http://<code.com>. Retrieved on <date>

There is a properly citable publication on the way as well which may be cited additionally.

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Modules Package meet

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Author & Contact

Written by Gunnar Waterstraat
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1.1 Modules

- _dot_new: A drop in replacement for numpy.dot (Section 2, p. 5)
- _interp: hidden functions for interpolating EEG (Section 3, p. 6)
- basic: Basic functions for reading binaries and EEG manipulation (Section 4, p. 7)
- **eeg_viewer**: Simple interactive EEG viewer (Section 5, p. 13)
- elm: Extreme Learning Machine Classification (Section 6, p. 15)
- iir: IIR Filtering (Section 7, p. 23)
- spatfilt: Spatial Filters (Section 8, p. 25)
- **sphere**: Spherical spline interpolation and CSD (Section 9, p. 30)
- tf: S transform (time-frequency transformation) (Section 10, p. 38)

Variables Module meet._dot_new

2 Module meet._dot_new

A drop in replacement for numpy.dot

Avoid temporary copies of non C-contiguous arrays

The code is available here: http://pastebin.com/raw.php?i=M8TfbURi In future this will be available in numpy directly: https://github.com/numpy/numpy/pull/2730

2.1 Functions

dot(A, B, out=None)
A drop in replaement for numpy.dot Computes A.B optimized using fblas call
$test_dot()$
$test_to_fix()$
1d array, complex and 3d

2.2 Variables

Name	Description
package	Value: 'meet'

3 Module meet._interp

hidden functions for interpolating EEG

 $\mbox{{\it Hidden}}$ submodule of the Modular EGg Toolkit - $\mbox{{\it MEET}}$ for Python.

Author:

Gunnar Waterstraat
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3.1 Functions

 $\mathbf{mchi}(x, y)$

3.2 Variables

Name	Description
_package	Value: 'meet'

4 Module meet.basic

Basic functions for reading binaries and $\ensuremath{\mathsf{EEG}}$ manipulation

Submodule of the Modular EEg Toolkit - MEET for Python.

Author:

Gunnar Waterstraat
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4.1 Functions

```
readBinary(fname, num_channels, channels='all', readnum_dp='all',
data_type='float4', buffermem=512)
Read EEG from a binary file and output as numpy array.
The binary of a signal with k channels and n datapoints must be
of the type:
       t0 t1 ... tn-1
ch_0 0
         k ... (n−1)*k
          k+1 ... (n-1)*k+1
ch_1 | 1
... | ... ...
ch_{k-1} k-1 2*k-1 ... n*k-1
The endianess of the runtime system is used.
Input:
-- fname - (str) - input file name
-- num_channels - int - total number of channels in the file
-- channels - numpy array OR 'all' - iterable of channels to read
             (starting with 0) if 'all', all channels are read
-- readnum_dp - int OR 'all' - number of datapints to read
-- data_type - str - any of 'int2', 'int4', 'int8', 'float4',
                    'float8', 'float16' where the digit determins
                    the number of bytes (= 8 bits) for each element
-- buffermem - float - number of buffer to us in MegaBytes
Output:
-- data - numpy array - data shaped k x n where k is number of
                       channels and n is number of datapoints
Example:
>>> readBinary(_path.join(_path.join(_packdir, 'test_data'),
                                                                       'sample.dat'), 2, data_ty
array([[0, 2, 4, 6, 8],
      [1, 3, 5, 7, 9]])
```

```
interpolateEEG(data, markers, win, interpolate_type='mchs')
Interpolates segemnets in the data
Input:
-- data - one or two dimensional array
         1st dimension: channels (can be ommitted if single channel)
         2nd dimension: datapoints
-- markers - marker positions arranged in 1d array
-- win - iterable of len 2 - determining the window in datapoints to
        be interpolated (win[0] is in, win[1] is out of the window)
-- interpolate_type: ['linear', 'mchs', 'akima'] - linear or
                   Monotone Cubic Hermite Spline
  or Akima interpolation
Output:
interpolated dataset
Examples:
>>> data = _np.arange(20, dtype=float).reshape(2,-1)
>>> interpolateEEG(data, [5], [-1,2], 'linear')
array([[ 0., 1.,
                   2., 3., 4., 5., 6.,
                                               7., 8.,
                                                          9.],
      [ 10., 11., 12., 13., 14., 15., 16., 17., 18., 19.]])
>>> interpolateEEG(data, [5], [-1,2], 'mchs')
array([[ 0., 1., 2.,
                                               7., 8.,
                                                          9.],
                        3., 4., 5., 6.,
      [ 10., 11., 12., 13., 14., 15., 16., 17., 18., 19.]])
>>> interpolateEEG(data, [5], [-1,2], 'akima')
                          2. , 3. , 3.625, 5. ,
array([[ 0.
              , 1. ,
                                                          6.375,
                        9.
         7.
                8.
                              ],
                        12.
                              , 13. , 13.625, 15. , 16.375,
      [ 10.
              , 11.
        17.
                18.
                         19.
                              ]])
```

```
epochEEG(data, marker, win)
Arange the dataset into trials (=epochs) according to the marker and
window.
markers and the window borders are sorted in ascending order.
Input:
-- data - numpy array - 1st dim channels (can be ommitted if single
                                         channel)
                       2nd dim datapoints
-- marker - iterable - the marker
-- win - iterable of len 2 - determing the start and end of epchos
        in dp (win[0] is in, win[1] is out of the window)
Output:
-- epochs - numpy array - dimension one more then data input
                       - 1st dim: channel (might be ommited - see
                                           above)
                       - 2nd dim: epoch length = win[1] - win[0]
                       - 3rd dim: number of epochs
Example:
-----
>>> data = _np.arange(20, dtype=float).reshape(2,-1)
>>> epochEEG(data, [3,5,7], [-2,2])
array([[[ 1.,
               3., 5.],
        [ 2.,
                4., 6.],
                     7.],
        [ 3.,
               5.,
        [ 4.,
               6.,
                     8.]],
<BLANKLINE>
      [[ 11., 13., 15.],
       [ 12., 14., 16.],
        [ 13., 15., 17.],
        [ 14., 16., 18.]])
```

```
calculateRMS(data, axis=-1)
Calculate rms value of the input data along the indicated axis
Input:
-- data - numpy array - input data
-- axis - int - axis along which the rms is calculated; if None, the
               flattened array is used
Output:
-----
-- rms value along the indicated axis
Example:
>>> data = _np.arange(20, dtype=float).reshape(2,-1)
>>> calculateRMS(data, None)
11.113055385446435
>>> calculateRMS(data, 0)
array([ 7.07106781, 7.81024968, 8.60232527, 9.43398113,
       10.29563014, 11.18033989, 12.08304597, 13.
       13.92838828, 14.86606875])
>>> calculateRMS(data, 1)
array([ 5.33853913, 14.7817455])
```

```
getMarker(marker, width=50, mindist=100)
Gets position of markers from the trigger channel
GetMarkerPosFromData(marker)
input:
-- marker - one-dimensional array with trigger channel - each
            impulse or zero crossing is treated a marker
--width - int - calculates the local mean in window of size width
               - defaults to 50
--mindist - int - minimal distance between triggers in dp
                 - defaults to 100
output:
-- marker - one-dimensional array containing trigger positions
Example:
_____
>>> x = np.ones(1000)
>>> x[200:400] = -1
>>> x[600:800] = -1
>>> getMarker(x)
array([ 400, 600, 800, 1000])
```

Variables Module meet.basic

4.2 Variables

Name	Description
package	Value: 'meet'

5 Module meet.eeg_viewer

Simple interactive EEG viewer

Submodule of the Modular EEg Toolkit - MEET for Python.

Author:

Gunnar Waterstraat
gunnar[dot]waterstraat[at]charite.de

5.1 Variables

Name	Description
package	Value: 'meet'

5.2 Class plotEEG

5.2.1 Methods

```
\_init\_(self, signals, ylabels, t, t_res=30, title=False)
Function to Plot EEG-Signals in of several channels
Def: PlotEEG(signals, ylabels, t, t_res, title = False)
Interaction:
PageUp -> Go backward a big step
PageDown -> Go forward a big step
Up -> Go backward a small step
Down -> go forward a small step
i -> Zoom in (show smaller temporal window)
o -> Zoom out (show larger temporal window)
+ -> increase gain
- -> decrease gain
Pos1 -> Go to start
End \rightarrow Go to end
LeftMouseClick -> saves the x coordinate in self.clicks
Input:
    -signals: 2d array of input data
    -ylabels: list of channel names in same order as in
              'signals' (unit is presumed to be mikro V)
    -t: array of time values in s
    -t_res: mm per second (standard: 30 mm / s)
    -title: title string
Output:
    -EEG_viewer class: plot with EEG.show()
     clicks are saved in self.clicks
```

```
change_gain(self, new_offset)
```

```
\mathbf{change\_t}(self,\ new\_t0,\ new\_t\_show)
```

```
\mathbf{show}(\mathit{self})
```

6 Module meet.elm

Extreme Learning Machine Classification

Submodule of the Modular EEg Toolkit - MEET for Python.

This module implements regularized Extreme Learning Machine Classification and Weighted Extreme Learning Machine Classification.

Classification is implemented in the ClassELM class

For faster execution of dot product the module dot_new is imported since it avoids the need of temporary copy and calls fblas directly. The code is available here: http://pastebin.com/raw.php?i=M8TfbURi In future this will be available in numpy directly: https://github.com/numpy/numpy/pull/2730

- 1. Extreme Learning Machine for Regression and Multiclass Classification Guang-Bin Huang, Hongming Zhou, Xiaojian Diang, Rui Zhang IEEE Transactions of Systems, Man and Cybernetics Pat B: Cybernetics, Vol. 42, No. 2. April 2012
- 2. Weighted extreme learning machine for imbalance learning. Weiwei Zong, Guang-Bin Huang, Yiqiang Chen Neurocomputing 101 (2013) 229-242

Author & Contact

Written by Gunnar Waterstraat

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Functions Module meet.elm

6.1 Functions

accuracy(conf_matrix) Measure of the performance of the classifier. The Accuracy is the proportion of correctly classified items in relation to the total number of items. You should be aware that this is very sensitive to imbalanced data (data with very unequal sizes of each class): Imagine a sample with 99% of the items belonging to class 0 and 1% of items belonging to class 1. A classifier might have an accuracy of 99% by just assigning all items to class 0. However, the sensitivity for class 1 is 0% in that case. It depends on your needs if this is acceptable or not. Input: conf_matrix - shape ny x ny, where ny is the number of classes the rows belong to the actual, the columns to the predicted class: item ij is hence predicted as class j, while it would have belonged to class i Output: float - the accuracy

$G_{-mean}(conf_{-matrix})$

The G-mean is the geometric mean of the per-class-sensitivities. It is much more stable to imbalance of the dataset than the global accuray. However it depends on your needs, which measure of performance of the classifier to use.

Input:

Output:

the geometric mean of per-class sensitivities

Functions Module meet.elm

Matthews(conf_matrix)

The Matthews correlation coefficient is used in machine learning as a measure of the quality of binary (two-class) classifications. It takes into account true and false positives and negatives and is generally regarded as a balanced measure which can be used even if the classes are of very different sizes. The MCC is in essence a correlation coefficient between the observed and predicted binary classifications; it returns a value between -1 and +1. A coefficient of +1 represents a perfect prediction, O no better than random prediction and -1 indicates total disagreement between prediction and observation.

Source: Wikipedia (2013-09-25)

Input:

conf_matrix - shape 2 x 2, where 2 is the number of classes
 the rows belong to the actual, the columns to the
 predicted class: item ij is hence predicted as class j,
 while it would have belonged to class i
 AN ERROR IS THROWN IF THE SHAPE OF THE MATRIX IS NOT
 CORRECT

Output:

float - the the Matthews Correlation Coefficient

$PPV(conf_matrix)$

Calculate the Positive Predictive Value

Input:

conf_matrix - shape 2 x 2, where 2 is the number of classes
 the rows belong to the actual, the columns to the
 predicted class: item ij is hence predicted as class j,
 while it would have belonged to class i
 AN ERROR IS THROWN IF THE SHAPE OF THE MATRIX IS NOT
 CORRECT

Output:

float - the PPV

```
Ssk_cv(data, labels, folds=3)

Cut data into folds with method:
shuffled, stratified, k-folds cross-validation

Input:
-----
data - numpy array - shape n x p, with n items anf p features

Output:
-----
returns a list, with each list-element including the indices of one fold
```

6.2 Variables

Name	Description
package	Value: 'meet'

6.3 Class ClassELM

Class for Extreme Learning Machine Classification

Input:

L - (int) - dimensionality of the feature space (defaults to 1000) change_alg - (int) - number of samples to change from implementation

6.3.1 Methods

 $_$ init $_$ (self, L=1000, kernel='sigmoid')

```
cv(self, data, labels, method='ssk_cv', C_array=None, folds=3,
precision_func='accuracy', scale=True, weights=True, mem_size=512, verbose=True)
Perform Cross-Validation of Extreme Learning Machine parameter C
Input:
data - numpy array - shape (n x p) with n being sample number
       and p being number of features
labels - numpy array - shape (n) with the class labels
         0,1,\ldots,ny-2,ny-1, where ny is the number of classes
method - string - cross-validation method
                - 'ssk_cv' - shuffled stratified k-folds
                  cross-validation
C_{-}array - numpy array - default is None - the C's which are
                        cross-validated
                      - if None from 2**(-25), 2**(-24), ...,
                                      2**(24), 2**(25)
folds - integer - default 3 - number of folds
precision_func - string or function - standard is 'accuray' -
                 Measure of performance
               - as string implemented: 'accuracy' -
                                           proportion of
                                           correctly classified
                                           to total number of
                                           samples
                                         'G_mean' - geometric
                                           mean of per-class
                                           accuracies
                                         'Matthews' - Matthews
                                           Correlation
                                           Coefficient - Only
                                           for binary
                                           classification
                                         'PPV' - Positive
                                           Predictive Value -
                                           Only for binary
                                           classification
               - if function: with confusion matrix as single
                 input and float (0,1) as single output
scale - bool (True | False) - whether data should be scaled to
                              range (-1,1)
weights - can be: - bool (True | False): - standard is True
                                            if True, data is
                                            re-weighted to a
                                            class ratio of 1.0
                                            if False, data is not
                                            re-weighted
                  - float in half-open interval [0,1)
                        - data is re-weighted such that the
                          minority / majority ratio is this
                          float
                        - minority clas@es are the ones having
                          less members than on average, majority
                          classes have more than average
                        - Zong et al. proposed to use the golden
                          ratio (approx. 0.618 -
```

```
train(self, data, labels, C, scale=True, weights=True, mem_size=512)
Train the ELM Classifier
Input:
data - (numpy array) - shape n x p
                       n - number of observations
                       p - number of dimensions
       if data.ndim > 2, the array is reshaped as (n,-1)
labels - array with integer labels
C - regularization parameter
scale - bool (True | False) - standard is True
                            - switch, if the features of the
                              dataset should be scaled
                              to the interval (-1,1)
weights - can be: - bool (True | False): - standard is True
                                           if True, data is re
                                           weighted to a class
                                           ratio of 1.0 if
                                           False, data is not
                                           re-weighted
                  - float in half-open interval [0,1)
                        - data is re-weighted such that the
                          minority / majority ratio is this
                          float
                        - minority classes are the ones having
                          less members than on average, majority
                          classes have more than average
                        - Zong et al. proposed to use the golden
                        ratio (approx. 0.618 -
                        scipy.Constants.golden) as a good value
                  - numpy array with weights for each sorted
                  unique class in labels, each class weight is
                  expected to be in half-open interval [0,1)
mem_size - number - memory size of temporary array in Mb -
                    defaulte to 512
Output:
No user ouput (Weights are generated and stored in the Class as
self._beta) self.istrained is set to True
```

7 Module meet.iir

```
IIR Filtering
Submodule of the Modular EEg Toolkit - MEET for Python.
So far only butterworth filtering is implemented
Author:
-----
Gunnar Waterstraat
gunnar[dot]waterstraat[at]charite.de
```

Variables Module meet.iir

7.1 Functions

```
butterworth(data, fp, fs, s_rate, gpass=3, gstop=8, axis=-1, zero_phase=True,
return_param=False)
Apply a butterworth filter to the data.
fp is the passband frequency in Hz
fs is the stopband frequency in Hz
Example of Notations:
For fs = 10, fp = 15 a high-pass filter is applied with the
  transition band between 10-15 Hz
For fs = 15, fp = 10 a low-pass filter is applied with the
  transition band between 10-15 Hz
For fs = [10, 40], fp = [20,30] a bandpass-filter with the passband
  20-30 Hz and transitions between 10-20 Hz and 30-40 Hz is applied
For fs = [20, 30], fp = [10,40] a bandstop-filter with the stopband
   20-30 Hz and transitions between 10-20 Hz and 30-40 Hz is applied
Input:
_____
-- data - numpy array
-- fp - float - pass-band frequencies
-- fs - float - stop-band frequencies
-- s_rate - float - the sampling rate in Hz
-- gpass - float - maximal attenuation in the passband (in dB)
-- gstop - float - minimal attenuation in the passband (in dB)
-- axis - int -along which axis the filter is applied
-- zero_phase - bool - if zero phase filter is applied by filtering
                in both directions
-- return_param - bool - if the order, b, and a shoud be returned
Output:
-- data - numpy array - the filtered data array
if return_param:
   -- data
   -- ord - int - filter order (this must be doubled for the
                  zero_phase implementation)
   -- (b, a) - Numerator ('b') and denominator ('a') polynomials of
               the IIR filter.
```

7.2 Variables

Name	Description
package	Value: 'meet'

8 Module meet.spatfilt

Spatial Filters

Submodule of the Modular EEg Toolkit - MEET for Python.

This module implements some spatial filters such as CSP, CCA, CCAvReg, bCSTP and QCA.

Author & Contact

Written by Gunnar Waterstraat
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Functions Module meet.spatfilt

8.1 Functions

```
CSP(data1, data2, center=True)
Common Spatial Pattern (CSP)
This takes the multivariate data in two conditions and finds the
spatial filters which minimize the variance in condition 2, while
simultaneously maximizing the variance in condition one.
The algorithm uses Singular Value Decomposition since this is more
stable than the Eigenvalue Decomposition of the Covariance Matrices
The filters are scaled such, that Var(cond1) + Var(cond2) = 1. The
eigenvalues give the variance in condition 1.
The filters are in the columns of the filter matrix. Patterns can be
obtained by inverting the filter matrix and are located in the rows
of the inverted filter matrix.
filter, eigval = CSP(data1, data2)
filtered_data = filter.T.dot(data1)
Input:
-- data1 - 2d array - data in condition 1 (variables in rows,
     observations in columns)
-- data2 - 2d array - data in condition 2 (variables in rows,
     observations in columns)
  The number of variables in data1 and data2 must be equal
-- center - bool - if data should be centered, defaults to True
Output:
-- filter - where the individual filters are in the columns of the
     matrix (in order of decreasing eigenvalues)
-- eigvals - eigenvalues in decreasing order
```

$\mathbf{CCA_data}(X, Y)$

Cannonical Correlation Analysis – by a combination of ${\tt QR}$ decomposition and Singular Value Decomposition

The filters are scaled to give unit variance in the components and are given in the order of decreasing canonical correlations.

Patterns can be obtained by inverting the filter matrices.

Inputs:

- ${\tt X}$ shaped p1 x ${\tt N}$ variables in rows, obervations in columns
- Y shaped p2 x N variables in rows, obervations in columns

Outputs:

- a filters for X (shape p1 x d), where d is min(rank(X), rank(Y)), each filter is in one column
- b filters for Y (shape p2 x d), where d is min(rank(X), rank(Y)), each filter is in one column
- s canonical correlations in non-increasing order, each corresponding to the respective colum in a and b

$\mathbf{CCAvReg}(trials)$ Canonical Correlation Average Regression Calculate the CCA between trials and their average across trials The filters are scaled to give unit variance in the components and are given in the order of decreasing canonical correlations. Patterns can be obtained by inverting the filter matrices. notably, all single trials can be filtered at once by: _np.tensordot(a, trials, axes=(0,0)) the filtered average can be obtained as: _np.dot(b.T, trials.mean(-1)) Input: -- trials - 3d numpy array - 1st dim: channels - 2nd dim: epoch length - 3rd dim: number of epochs Output: -- a - 2d numpy array (channel x channel) - spatial filters for single trials, each filter is in one column -- b - 2d numpy array (channel x channel) - spatial filters for average, each filter is in one column

-- s - the canonical correlations between single trials and averages

```
bCSTP(trials1, trials2, num_iter=30, s_keep=2, t_keep=2, verbose=True)
bilinear Common Spatial-Temporal Patterns
In each iteration the number of kept patterns is reduced by one,
in the last 5 iterations the finally desired number of patterns is
The minimal number of iterations must therefor be > 10
Filter matrices won't be square if the data was rank deficient,
patterns can then be obtained by using the pseudo-inverse
(numpy.linalg.pinv)
Input:
-- trials1 - 3d numpy array - 1st dim: channels
                            - 2nd dim: epoch length
                            - 3rd dim: number of epochs
  variance in condition 1 is maximized
-- trials2 - 3d numpy array - 1st dim: channels
                            - 2nd dim: epoch length
                            - 3rd dim: number of epochs
  variance in condition 2 is minimized
-- num_iter - int - number of iterations to do, defaults to 30
                    (minimum 10)
-- s_keep - int - number of spatial patterns to keep, defaults to 2
-- t_keep - int - number of temporal patterns to keep, defaults to 2
-- verbose - bool - if number of iterations should be printed during
                    execution
Output:
-- W - list of 2d arrays - spatial filter matrix for each iteration,
      the final spatial filters are in W[-1]
-- V - list of 2d arrays - temporal filter matrix for eachiteration,
                           the final temporal filters are in V[-1]
-- s_eigvals - list of 1d array - spatial eigenvalues,
               the final spatial eigenvalues are in s_eigvals[-1]
-- t_eigvals - list of 1d array - temporal eigenvalues, the final
               temporal eigenvalues are in s_eigvals[-1]
```

8.2 Variables

Name	Description
package	Value: 'meet'

output is in data-unit/m**

9 Module meet.sphere

Author:

Gunnar Waterstraat

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Spherical spline interpolation and CSD

72:184-187, Corrigenda in 1990, 76: 565-566

Submodule of the Modular EEg Toolkit - MEET for Python.

Algorithm from Perrin et al., Electroenceph Clin Neurophysiol 1989,

>>> CSD = calcCSD(coords, data) # get CSD using spherical splines -

```
While the implementation was done independently, the code was tested
using the sample data of the CSD Toolbox for Matlab
(http://psychophysiology.cpmc.columbia.edu/Software/CSDtoolbox/)
and results are the same. Thanks a lot th Juergen Kayer for sharing his
work there.
Example:
For Plotting a scalp map:
>>> import numpy as _np
>>> coords = getStandardCoordinates(['Fp1', 'Fp2', 'C1', 'C2', 'C3',
                                                                             'C4', 'P2', 'P4'])
>>> coords = projectCoordsOnSphere(coords) # be sure that these coords
                                                                               are on the surface of a
>>> data = _np.random.random(coords.shape[0]) # just create a random
                                                                             vector for testing
>>> X, Y, Z = potMap(coords, data) # interpolate using spherical splines
For Calculating current source densities:
>>> import numpy as _np
>>> coords = getStandardCoordinates(['Fp1', 'Fp2', 'C1', 'C2', 'C3',
                                                                            'C4', 'P2', 'P4'])
>>> coords = projectCoordsOnSphere(coords) # be sure that these coords
                                                                               are on the surface of a
>>> data = _np.random.random([coords.shape[0], 1000]) # just create a
                                                                              random vector for testin
```

Functions Module meet.sphere

9.1 Functions

```
getStandardCoordinates(elecnames, fname='standard')
Read (cartesian) Electrode Coordinates from tab-seperated text file
The standard is plotting_1005.txt obtained from:
http://robertoostenveld.nl/electrode/plotting_1005.txt
Thanks to Robert Oostenveld for sharing these files!!!
Input:
-- elecnames - iterable - list of Electrode names
-- fname - str - filename from which positions should be read
              - this is a tab delimeed file with the UPPERCASE
                electrode names in first column
                x,y,z in subsequent columns
Output:
-- coords - 2D array containing the cartesian coordinates in rows
               - 1st column: x
               - 2nd column: y
               - 3rd column: z
Example:
>>> getStandardCoordinates(['fc3', 'FC1', 'xx']) # 'xx' is not a
                                                                            valid electrode
array([[-0.6638 , 0.3610691 , 0.6545
                                           ],
      [-0.3581 , 0.37682936, 0.8532
                                           ],
                            nan, nan]])
              nan,
```

Module meet.sphere

```
projectCoordsOnSphere(coords)
Get the Coords with same azimuth and altitude on on a spherical
surface with center 0 and radius 1
Input:
----
--coords - 3D array containing the points in rows
         - 1st column: x
         - 2nd column: y
         - 3rd column: z
Output:
-- out_coords - 3D array containing the coordinates in rows
Examples:
>>> coords = _np.array([[0,0,1], [0,1,0], [1,1,1]]) # the last one
                                                                         is not on a spheri
>>> projectCoordsOnSphere(coords)
],
                                        ],
      [0.57735027, 0.57735027, 0.57735027]
```

```
Calculate a 3D meshgrid on a sphere from a 2D meshgrid on a circle with center (0,0) and radius 1

Input:
-----
-- coords - 2D array containing the coordinates in rows
- 1st column: x
- 2nd column: y

Output:
-----
-- sphere_coords - masked 3D array containing the coordinates in rows
- 1st column: x
- 2nd column: x
- 2nd column: y
- 3rd column: z

(all coordinates outside a unit sphere are masked)
```

Module meet.sphere

```
meshSphereOnCircle(sphere_coords)
Calculate a 2D meshgrid on a circle from a 3D meshgrid on a sphere
Input:
-- sphere_coords - 2D array containing the coordinates in rows
         - 1st column: x
          - 2nd column: v
         - 3rd column: z
Output:
-- coords - 2D array containing the coordinates in rows
                - 1st column: x
                - 2nd column: y
Example:
_____
>>> coords_2d = meshCircle(4) # mesh a circle with 4 points on the
>>> coords_3d = meshCircleOnSphere(coords_2d) # project that mesh
>>> _np.all(_np.abs((meshSphereOnCircle(coords_3d) - coords_2d) /
True
```

diagonal
on a sphere
coords_2d) < 1E-10)

```
potMap(RealCoords, data, diameter\_samples=200, n=(7, 7), m=4, smooth=0)
Get a scalp map of potentials interpolated on a sphere.
Input:
-- RealCoords - array of shape channels x \ 3 - (x,y,z) cartesian
                coordinates of physical electrodes on a sphere with
                center (0,0,0) and radius 1
-- data - array of shape channels x datapoints - containing values
          at electrodes in same order as in 'RealCoords'
-- diameter_samples - int - number of points along the diameter of
                      the scalp to interpolate
-- n - tuple of ints - how many terms of the Legendre-Polynomial
       should be calculated, defaults to (7,7)
-- m - int - order of the spherical spline interpolation, defaults
      to 4
-- smooth - float - amount of smoothing, defaults to 0
Output:
-----
-- X, Y, Z - grid containing the X and Y coordinates and the
             interpolated values where everything outside the unit
             circle is masked.
```

```
csdMap(RealCoords, data, diameter\_samples=200, n=(7, 20), m=4, smooth=1e-05)
Get a scalp map of CSDs calculated from spherical splines.
Input:
-- RealCoords - array of shape channels x 3 - (x,y,z) cartesian
               coordinates physical electrodes on a sphere with
                center (0,0,0) and radius 1
-- data - array of shape channels x datapoints - containing values
         at electrodes in same order as in 'RealCoords'
-- diameter_samples - int - number of points along the diameter of
                     the scalp to interpolate
-- n - tuple of ints - how many terms of the Legendre-Polynomial
      should be calculated, defaults to (7,20)
-- m - int - order of the spherical spline interpolation,
      defaults to 4
-- smooth - float - amount of smoothing, defaults to 1E-5
Output:
-- X, Y, Z - grid containing the X and Y coordinates and the
             interpolated values where everything outside the unit
             circle is masked. the unit of z is data_unit / m**2
```

```
calcCSD(Coords, data, n=(7, 20), m=4, smooth=1e-05, buffersize=64)
Calculate current source densities at the same electrode posisionts
as in the original data set
Input:
-- Coords - 2d array - channels x (x,y,z) cartesian coordinates of
           physical electrodes
-- data - array containing values at electrodes in same order as in
         'Coords'
   -- 1st dim: channels
   -- 2nd dim: datapoints
-- n - tuple of ints - How many terms of the Legendre-Polynomial
      should be calculated, defaults to (7, 20) - 7 for the first
       and 20 for the 2nd polynomial
-- m - order of the spherical spline interpolation, defaults to 4
-- smooth - float - amount of smoothing, defaults to 1E-5
-- buffersize - float - determine the size of a buffer in MB to do
               the calculations
Output:
-- pot - array containing CSD-values in data-unit / m**2
Example:
>>> Coords = getStandardCoordinates(['C1', 'C3'])
>>> Coords = projectCoordsOnSphere(Coords) # be sure that the
                                                                           points are on a sphere
>>> data = _np.arange(6, dtype=float).reshape(2,-1)
>>> calcCSD(Coords, data)
array([[-3.5474569 , -3.5474569 , -3.5474569 ],
       [ 3.60884286, 3.60884286, 3.60884286]])
```

Module meet.sphere

```
smoothSP(Coords, data, n=(7, 7), m=4, smooth=1e-05, buffersize=64)
Calculate spatially smoothed potential values at the same electrode
positions as in the original data set
Input:
-- Coords - 2d array - channels x (x,y,z) cartesian coordinates of
           physical electrodes
-- data - array containing values at electrodes in same order as in
         'Coords'
   -- 1st dim: channels
    -- 2nd dim: datapoints
-- n - tuple of ints - How many terms of the Legendre-Polynomial
       should be calculated, defaults to (7, 7) - 7 for the first
       and 7 for the 2nd polynomial
-- m - order of the spherical spline interpolation, defaults to 4
-- smooth - float - amount of smoothing, defaults to 1E-5
-- buffersize - float - determine the size of a buffer in MB to do
               the calculations
Output:
-- pot - array containing the spatially smoothed potentials
Example:
>>> Coords = getStandardCoordinates(['C1', 'C3'])
>>> Coords = projectCoordsOnSphere(Coords) # be sure that the
                                                                        points are on a sphere
>>> data = _np.arange(6, dtype=float).reshape(2,-1)
>>> smoothSP(Coords, data)
array([[ 0.01220571, 1.01220571, 2.01220571],
       [ 2.98778429, 3.98778429, 4.98778429]])
```

9.2 Variables

Name	Description
package	Value: 'meet'

10 Module meet.tf

```
S transform (time-frequency transformation)
Submodule of the Modular EEg Toolkit - MEET for Python.
the 'standard' S transform:
Stockwell, Robert Glenn, Lalu Mansinha, and R. P. Lowe. "Localization of
the complex spectrum: the S transform." Signal Processing,
IEEE Transactions on 44.4 (1996): 998-1001.
as well as
the fast dyadic S transform:
-----
Brown, Robert A., M. Louis Lauzon, and Richard Frayne. "A general
description of linear time-frequency transforms and formulation of a
fast, invertible transform that samples the continuous S-transform
spectrum nonredundantly." Signal Processing, IEEE Transactions on 58.1
(2010): 281-290.
Author:
_____
Gunnar Waterstraat
gunnar[dot]waterstraat[at]charite.de
Example for a standard S transform:
_____
>>> import matplotlib.pyplot as plt
>>> import numpy as np
>>> data = np.random.random(1000) # generate a random 1000 dp signal
>>> tf_coords, S = gft(data, sampling='full')
>>> S = S.reshape(-1, data.shape[0]) # for the full transform no
                                                                      interpolation is needed -> j
>>> plt.imshow(_np.abs(S), aspect='equal', origin='lower') #plot the
Example for a dyadic S transform:
-----
>>> import matplotlib.pyplot as plt
>>> import numpy as np
>>> data = np.random.random(1000) # generate a random 1000 dp signal
>>> tf_coords, S = gft(data, sampling='dyadic')
>>> t, f, S_interp = interpolate_gft(tf_coords, S, (data.shape[0]//2,
                                                                          data.shape[0]), data.shap
                                                                      extent=[t[0], t[-1], f[0], f
>>> plt.imshow(_np.abs(S_interp), aspect='equal', origin='lower',
```

Functions Module meet.tf

10.1 Functions

```
gft(sig, window='gaussian', axis=-1, sampling='full', full=False, hanning=True)
Calculate the discrete general fourier family transform
Inputs:
-----
-- sig - signal of interest - 1 or 2 dimensional
-- window - window function, standard is gaussian
-- axis - axis along which transform should be performed
-- sampling - 'full' | 'dyadic' - should the S-domain be sampled
              completely, i.e. redundant or dyadic
-- full - True False - return also negative frequencies, standard
         is False
-- hanning - True False - Apply a hanning window to 5% first and
             last datapoints
Outputs:
-- Coords - (frequency, time) Coordinates of each point in S
-- S - complex S transform array
  if sampling is dyadic interpolate_gft should be used to
   interpolate the result onto a regular grid
   if sampling is full, this can be easily done by reshaping each {\tt S}
  transform to the shape (-1, sig.shape[axis])
```

Variables Module meet.tf

```
interpolate_gft(Coords, S, IM_shape, data_len, kindf='nearest', kindt='nearest')
Interpolate the result of a gft-transform - standard is nearest
neighbor interpolation
Input:
-- Coords, Coords as output by the gft command
-- S - A gft result list
-- IM\_shape - requested shape of the interpolated matrix
      1st axis frequency
       2nd axis time
-- data_len - length of inital data
-- kindf - interpolation method along frequency axis - any of
           ['nearest', 'linear']
-- kindt - interpolation method along time axis - any of
           ['nearest', 'linear']
Output:
-- wanted times - time point array
-- wanted freqs - frequency array
-- IM - interpolated matrix
```

10.2 Variables

Name	Description
package	Value: 'meet'

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