skbold Documentation

Release 0.1

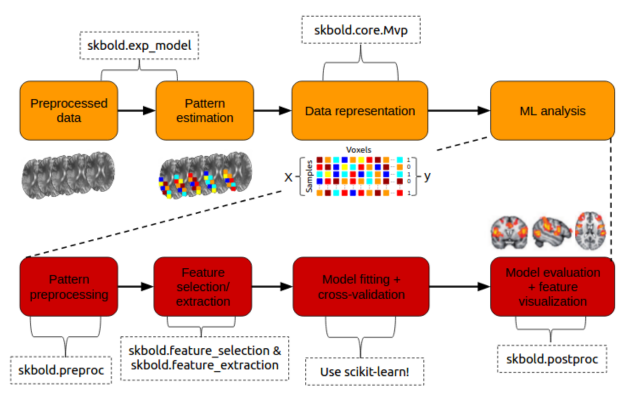
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Getting started

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The Python package skbold offers a set of tools and utilities for machine learning analyses of functional MRI (BOLD-fMRI) data. Instead of (largely) reinventing the wheel, this package builds upon an existing machine learning framework in Python: scikit-learn. The modules of skbold are applicable in several 'stages' of typical pattern analyses (see image below), including pattern estimation, data representation, pattern preprocessing, feature selection/extraction, and model evaluation/feature visualization.



The documentation of skbold is split up into three sections:

- Getting started
- Examples
- API

Installation & dependencies

Although the package is very much in development, it can be installed using *pip*:

\$ pip install skbold

However, the pip-version is likely behind compared to the code on Github, so to get the most up to date version, use git:

\$ pip install git+https://github.com/lukassnoek/skbold.git@master

Skbold is largely Python-only (both Python2.7 and Python3) and is built around the "PyData" stack, including:

- Numpy
- Scipy
- Pandas
- · Scikit-learn

And it uses the awesome nibabel package for reading/writing nifti-files. Also, skbold uses FSL (primarily the FLIRT and applywarp functions) to transform files from functional (native) to standard (here: MNI152 2mm) space. These FSL-calls are embedded in the convert2epi and convert2mni functions, so avoid this functionality if you don't have a working FSL installation.

Data organization

Skbold is quite flexible in how to work with data organized in different ways. However, to optimally use the *Mvp* classes (especially the *MvpBetween* class), it's best to organize your data hierarchically. For example, suppose you have a *.feat*-directory with first-level pattern estimates of two two runs ('run-1', 'run-2') for 3 subjects ('sub-01', 'sub-02', 'sub-03'). A nice organization would be as follows:

```
project_dir
  — sub-01
      - run-1
        L____ some_task.feat
       run-2
        sub-02
     — run-1
        run-2
        L____ some_task.feat
    sub-03
       run-1
        _____ some_task.feat
        run-2
        L_____ some_task.feat
```

This organization makes it easy for *Mvp* objects to find, load in, and organize patterns from these *.feat*-directories, as will become more clear in the examples-section on *Mvp* objects.

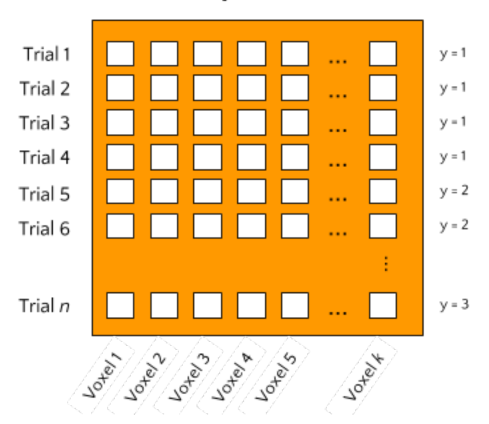
Mvp-objects

One of skbold's main features is the data-structure Mvp (an abbreviation of MultiVoxel Pattern). This custom object allows for an efficient way to store and access data and metadata necessary for multivoxel analyses of fMRI data. A nice feature of this Mvp objects is that they can easily load data (i.e., sets of nifti-files) from disk and automatically organize it in a format that is used in ML-analyses (i.e., a sample-by-feature matrix).

So, at the core, an Mvp-object is simply a collection of data - a 2D array of samples by features - and fMRI-specific metadata necessary to perform customized preprocessing and feature engineering. However, machine learning analyses, or more generally any type of multivoxel-type analysis (i.e. MVPA), can be done in two basic ways, which provide the basis of the two 'flavors' of Mvp-objects: MvpWithin and MvpBetween, as explained in more detail below.

3.1 MvpWithin

One way is to perform analyses *within subjects*. This means that a model is fit on each subjects' data separately. Data, in this context, often refers to single-trial data, in which each trial comprises a sample in our data-matrix and the values per voxel constitute our features. This type of analysis is alternatively called *single-trial decoding*, and is often performed as an alternative to (whole-brain) univariate analysis.



MvpWithin

Ultimately, this type of analysis aims to predict some kind of attribute of the trials (for example condition/class membership in classification analyses or some continuous feature in regression analyses), which skbold calls y, based on a model trained on the samples-by-features matrix, which skbold calls X. After obtaining model performance scores (such as accuracy, F1-score, or R-squared) for each subject, a group-level random effects (RFX) analysis can be done on these scores. Skbold does not offer any functionality in terms of group-level analyses; we advise researchers to look into the prevalance inference method of Allefeld and colleagues.

3.2 MvpBetween

With the increase in large-sample neuroimaging datasets, another type of MVPA starts to become feasible, which we'll call *between subject* analyses. In this type of analysis, single subjects constitute the data's samples and a corresponding single multivoxel pattern constitutes the data's features. The type of multivoxel pattern, or 'feature-set', can be any set of voxel values. For example, features from a single first-level contrast (note: this should be a condition average contrast, as opposed to single-trial contrasts in MvpWithin!) can be used. But voxel patterns from VBM, TBSS (DTI), and dual-regression maps can equally well be used. Crucially, this package allows for the possibility to stack feature-sets such that models can be fit on features from multiple data-types simultaneously.

	Feature-set 1 (e.g. fMRI)	Feature-set 2 (e.g. VBM)	Feature-set 3 (e.g. fMRI)			
Subject 1 Subject 2				y = 1 y = 1		
Subject 2				y = 1		
Subject 4				y = 1		
Subject 5				y = 2		
Subject 6				y = 2		
	: :		:			
Subject n				y = 3		
Jot						

MvpBetween

MvpResults: model evaluation and feature visualization

Given that an appropriate Mvp-object exists, it is really easy to implement a machine learning analysis using standard *scikit-learn* modules. However, as fMRI datasets are often relatively small, K-fold cross-validation is often performed to keep the training-set as large as possible. Additionally, it might be informative to visualize which features are used and are most important in your model. (But, note that feature mapping should not be the main objective of decoding analyses!) Doing this - model evaluation and feature visualization across multiple folds - complicates the process of implementing machine learning pipelines on fMRI data.

The MvpResults object offers a solution to the above complications. Simply pass your scikit-learn pipeline to MvpResults after every fold and it automatically calculates a set of model evaluation metrics (accuracy, precision, recall, etc.) and keeps track of which features are used and how 'important' these features are (in terms of the value of their weights).

Feature selection/extraction

The feature_selection and feature_extraction modules in skbold contain a set of scikit-learn type transformers that can perform various types of feature selection and extraction specific to multivoxel fMRI-data. For example, the RoiIndexer-transformer takes a (partially masked) whole-brain pattern and indexes it with a specific region-of-interest defined in a nifti-file. The transformer API conforms to scikit-learn transformers, and as such, (almost all of them) can be used in scikit-learn pipelines.

To get a better idea of the package's functionality - including the use of Mvp-objects, transformers, and MvpResults - a typical analysis workflow using skbold is described below.

For some example usages of the Mvp-objects and how to incorporate them in a scikit-learn-based ML-pipeline, check the examples below:

An example workflow: MvpWithin

Suppose you have data from an fMRI-experiment for a set of subjects who were presented with images which were either emotional or neutral in terms of their content. You've modelled them using a single-trial GLM (i.e. each trial is modelled as a separate event/regressor) and calculated their corresponding contrasts against baseline. The resulting FEAT-directory then contains a directory ('stats') with contrast-estimates (COPEs) for each trial. Now, using MvpWithin, it is easy to extract a sample by features matrix and some meta-data associated with it, as shown below.

Now, we have an Mvp-object on which machine learning pipeline can be applied:

```
import joblib
from sklearn.preprocessing import StandardScaler
from sklearn.svm import SVC
from sklearn.pipeline import Pipeline
from sklearn.model_selection import StratifiedKFold
from sklearn.metrics import accuracy_score, f1_score
from skbold.feature_selection import fisher_criterion_score, SelectAboveCutoff
from skbold.feature_extraction import RoiIndexer
```

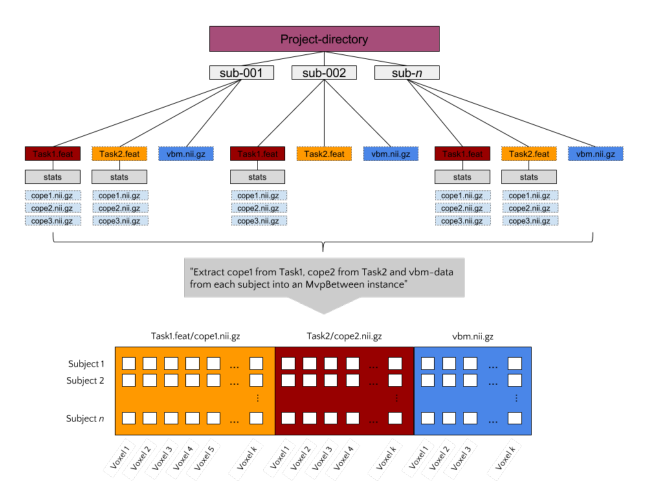
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```
from skbold.postproc import MvpResults
mvp = joblib.load('~/mvp_sub001.jl')
roiindex = RoiIndexer(mvp=mvp,
                      mask='Amygdala',
                      atlas_name='HarvardOxford-Subcortical',
                      lateralized=False) # loads in bilateral mask
# Extract amygdala patterns from whole-brain
mvp.X = roiindex.fit().transform(mvp.X)
# Define pipeline
pipe = Pipeline([
    ('scaler', StandardScaler()),
    ('anova', SelectAboveCutoff(fisher_criterion_score, cutoff=5)),
    ('svm', SVC(kernel='linear'))
1)
cv = StratifiedKFold(y=mvp.y, n_splits=5)
# Initialization of MvpResults; 'forward' indicates that it keeps track of
# the forward model corresponding to the weights of the backward model
# (see Haufe et al., 2014, Neuroimage)
mvp_results = MvpResults(mvp=mvp, n_iter=len(cv), feature_scoring='forward',
                         f1=f1_score, accuracy=accuracy_score)
for train_idx, test_idx in cv.split(mvp.X, mvp.y):
   train, test = mvp.X[train_idx, :], mvp.X[test_idx, :]
   train_y, test_y = mvp.y[train_idx], mvp.y[train_idx]
   pipe.fit(train, train_y)
   pred = pipe.predict(test)
   mvp_results.update(test_idx, pred, pipeline=pipe) # update after each fold!
mvp_results.compute_scores() # compute!
mvp_results.write(out_path) # write file with metrics and niftis with feature-scores!
```

An example workflow: MvpBetween

Suppose you have MRI data from a large set of subjects (let's say >50), including (task-based) functional MRI, structural MRI (T1-weighted images, DTI), and behavioral data (e.g. questionnaires, behavioral tasks). Such a dataset would qualify for a *between subject* decoding analysis using the MvpBetween object. To use the MvpBetween functionality effectively, it is important that the data is organized sensibly. An example is given below.



In this example, each subject has three different data-sources: two FEAT- directories (with functional contrasts) and one VBM-file. Let's say that we'd like to use all of these sources of information together to predict some behavioral variable, neuroticism for example (as measured with e.g. the NEO-FFI). The most important argument passed to MvpBetween is source. This variable, a dictionary, should contain the data-types you want to extract and their corresponding paths (with wildcards at the place of subject-specific parts):

```
import os
from skbold import roidata_path
gm_mask = os.path.join(roidata_path, 'GrayMatter.nii.gz')
source = dict(
    Contrast_tlcope1={'path': '~/Project_dir/sub*/Task1.feat/cope1.nii.gz'},
    Contrast_t2cope2={'path': '~/Project_dir/sub*/Task2.feat/cope2.nii.gz'},
    VBM={'path': '~/Project_dir/sub*/vbm.nii.gz', 'mask': gm_mask}
```

Now, to initialize the MvpBetween object, we need some more info:

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This is basically all you need to create a MvpBetween object! It is very similar to MvpWithin in terms of attributes (including X, y, and various meta-data attributes). In fact, MvpResults works exactly in the same way for MvpWithin and MvpBetween! The major difference is that MvpResults keeps track of the feature-information for each feature-set separately and writes out a summarizing nifti-file for each feature-set. Transformers also work the same for MvpBetween objects/data, with the exception of the cluster-threshold transformer.

skbold.core package

The core subpackage contains skbold's most important data-structure: the Mvp. This class forms the basis of the 'multivoxel-patterns' (i.e. mvp) that are used throughout the package. Subclasses of Mvp (MvpWithin and MvpBetween) are also defined in this core module.

The MvpWithin object is meant as a data-structure that contains a set of multivoxel fMRI patterns of *single trials, for a single subject*, hence the 'within' part (i.e. within-subjects). Currently, it has a single public method, create(), loading a set of contrasts from a FSL-firstlevel directory (i.e. a .feat-directory). Thus, importantly, it assumes that the single-trial patterns are already modelled, on a single-trial basis, using some kind of GLM. These trialwise patterns are then horizontally stacked to create a 2D samples by features matrix, which is set to the X attribute of MvpWithin.

The MvpBetween object is meant as a data-structure that contains a set of multivoxel fMRI patterns of *single conditions, for a set of subjects*. It is, so to say, a 'between-subjects' multivoxel pattern, in which subjects are 'samples'. In contrast to MvpWithin, contrasts that will be loaded are less restricted in terms of their format; the only requisite is that they are nifti files. Notably, the MvpBetween format allows to vertically stack different kind of 'feature-sets' in a single MvpBetween object. For example, it is possible to, for a given set of subjects, stack a functional contrast (e.g. a high-load minus low-load functional contrast) with another functional contrast (e.g. a conflict minus no-conflict functional contrast) in order to use features from both sets to predict a certain psychometric or behavioral variable of the corresponding subjects (such as, e.g., intelligence). Also, the MvpBetween format allows to load (and stack!) VBM, TBSS, resting-state (to extract connectivity measures), and dual-regression data. More information can be found below in the API. A use case can be found on the main page of ReadTheDocs.

Also, functional-to-standard (i.e. convert2mni) and standard-to-functional (i.e. convert2epi) warp-functions for niftis are defined here, because they have caused circular import errors in the past.

class Mvp (X=None, y=None, mask=None, mask_thres=0)

Bases: object

Mvp (multiVoxel Pattern) class. Creates an object, specialized for storing fMRI data that will be analyzed using machine learning or RSA-like analyses, that stores both the data (X: an array of samples by features, y: numeric labels corresponding to X's classes/conditions) and the corresponding meta-data (e.g. nifti header, mask info, etc.).

Parameters

• **X** (*ndarray*) – A 2D numpy-array with rows indicating samples and columns indicating features.

- y (list or ndarray) Array/list with labels/targets corresponding to samples in X.
- **mask** (*str*) Absolute path to nifti-file that will mask (index) the patterns.
- mask_thres (int or float) Minimum value for mask (in cases of probabilistic masks).

Variables

- **mask_shape** (*tuple*) Shape of mask that patterns will be indexed with.
- **nifti_header** (*Nifti1Header object*) **Nifti-header** from corresponding mask.
- **affine** (*ndarray*) Affine corresponding to nifti-mask.
- **voxel_idx** (*ndarray*) Array with integer-indices indicating which voxels are used in the patterns relative to whole-brain space. In other words, it allows to map back the patterns to a whole-brain orientation.
- X (ndarray) The actual patterns (2D: samples X features)
- **y** (list or ndarray) Array/list with labels/targets corresponding to samples in X.

Notes

This class is mainly meant to serve as a parent-class for MvpWithin and MvpBetween, but it can alternatively be used as an object to store a 'custom' multivariate-pattern set with meta-data.

```
update_mask (mask, threshold=0)
```

```
write (path=None, name='mvp', backend='joblib')
Writes the Mvp-object to disk.
```

Parameters

- **path** (*str*) Absolute path where the file will be written to.
- **name** (*str*) Name of to-be-written file.
- **backend** (*str*) Which format will be used to save the files. Default is 'joblib', which conveniently saves the Mvp-object as one file. Alternatively, and if the Mvp-object is too large to be save with joblib, a data-header format will be used, in which the data (X) will be saved using Numpy and the meta-data (everythin except X) will be saved using joblib.

convert2epi (file2transform, reg_dir, out_dir=None, interpolation='trilinear', suffix='epi', overwrite=False)

Transforms a nifti from mni152 (2mm) to EPI (native) format. Assuming that reg_dir is a directory with transformation-files (warps) including standard2example_func warps, this function uses nipype's fsl interface to flirt a nifti to EPI format.

- **file2transform** (*str or list*) Absolute path(s) to nifti file(s) that needs to be transformed
- **reg_dir** (*str*) Absolute path to registration directory with warps
- **out_dir** (*str*) Absolute path to desired out directory. Default is same directory as the to-be transformed file.
- interpolation (*str*) Interpolation used by flirt. Default is 'trilinear'.
- **suffix** (*str*) What to suffix the transformed file with (default : 'epi')
- **overwrite** (bool) Whether to overwrite existing transformed files

Returns out_all – Absolute path(s) to newly transformed file(s).

Return type list

convert2mni (file2transform, reg_dir, out_dir=None, interpolation='trilinear', suffix=None, overwrite=False, apply_warp=True)

Transforms a nifti to mni152 (2mm) format. Assuming that reg_dir is a directory with transformation-files (warps) including example_func2standard warps, this function uses nipype's fsl interface to flirt a nifti to mni format.

Parameters

- **file2transform** (*str or list*) Absolute path to nifti file(s) that needs to be transformed
- **reg_dir** (*str*) Absolute path to registration directory with warps
- **out_dir** (*str*) Absolute path to desired out directory. Default is same directory as the to-be transformed file.
- interpolation (*str*) Interpolation used by flirt. Default is 'trilinear'.
- **suffix** (*str*) What to append to name when converted (default : basename file2transform).
- **overwrite** (bool) Whether to overwrite already existing transformed file(s)
- apply_warp (bool) Whether to use the non-linear warp transform (if available).

Returns out_all – Absolute path(s) to newly transformed file(s).

Return type list

class MvpBetween (source, subject_idf='sub0???', remove_zeros=True, X=None, y=None, mask=None, mask thres=0, subject list=None)

Bases: skbold.core.mvp.Mvp

Extracts and stores multivoxel pattern information across subjects. The MvpBetween class allows for the extraction and storage of multivoxel (MRI) pattern information across subjects. The MvpBetween class can handle various types of information, including functional contrasts, 3D (subject-specific) and 4D (subjects stacked) VBM and TBSS data, dual-regression data, and functional-connectivity data from resting-state scans (experimental).

Parameters

• **source** (*dict*) – Dictionary with types of data as keys and data-specific dictionaries as values. Keys can be 'Contrast_*' (indicating a 3D functional contrast), '4D_anat' (for 4D anatomical - VBM/TBSS - files), 'VBM', 'TBSS', and 'dual_reg' (a subject-spedific 4D file with components as fourth dimension).

The dictionary passed as values must include, for each data-type, a path with wildcards to the corresponding (subject-specific) data-file. Other, optional, key-value pairs per data-type can be assigned, including 'mask': 'path', to use data-type-specific masks.

An example:

```
>>> source = {}
>>> path_emo = '~/data/sub0*/*.feat/stats/tstat1.nii.gz'
>>> source['Contrast_emo'] = {'path': path_emo}
>>> vbm_mask = '~/vbm_mask.nii.gz'
>>> path_vbm = '~/data/sub0*/*vbm.nii.gz'
>>> source['VBM'] = {'path': path_vbm, 'mask': vbm_mask}
```

- **subject_idf** (*str*) Subject-identifier. This identifier is used to extract subject-names from the globbed directories in the 'path' keys in source, so that it is known which pattern belongs to which subject. This way, MvpBetween can check which subjects contain complete data!
- **X** (*ndarray*) Not necessary to pass MvpWithin, but needs to be defined as it is needed in the super-constructor.
- **y** (*ndarray or list*) Labels or targets corresponding to the samples in X.
- **mask** (str) Absolute path to nifti-file that will be used as a common mask. Note: this will only be applied if its shape corresponds to the to-be-indexed data. Otherwise, no mask is applied. Also, this mask is 'overridden' if source[data_type] contains a 'mask' key, which implies that this particular data-type has a custom mask.
- **mask_threshold** (*int or float*) Minimum value to binarize the mask when it's probabilistic.

Variables

- **mask_shape** (*tuple*) Shape of mask that patterns will be indexed with.
- **nifti_header**(*list of Nifti1Header objects*) Nifti-headers from original data-types.
- affine (list of ndarray) Affines corresponding to nifti-masks of each data-type.
- X (ndarray) The actual patterns (2D: samples X features)
- **y** (list or ndarray) Array/list with labels/targets corresponding to samples in X.
- **common_subjects** (*list*) List of subject-names that have complete data specified in source.
- **featureset_id** (*ndarray*) Array with integers of size X.shape[1] (i.e. the amount of features in X). Each unique integer, starting at 0, refers to a different feature-set.
- voxel_idx (ndarray) Array with integers of size X.shape[1]. Per feature-set, these
 voxel- indices allow the features to be mapped back to whole-brain space. For example, to
 map back the features in X from feature set 1 to MNI152 (2mm) space, do:

```
>>> mni_vol = np.zeros((91, 109, 91))
>>> tmp_idx = mvp.featureset_id == 0
>>> mni_vol[mvp.featureset_id[tmp_idx]] = mvp.X[0, tmp_idx]
```

- **data_shape** (*list of tuples*) Original (whole-brain) shape of the loaded data, per data-type.
- **data_name** (*list of str*) List of names of data-types.

add_y (file_path, col_name, sep=`\t', index_col=0, normalize=False, remove=None, ensure_balanced=False, nan_strategy='remove', **kwargs) Sets y attribute to an outcome-variable (target).

- **file_path** (*str*) Absolute path to spreadsheet-like file including the outcome var.
- col_name (str) Column name in spreadsheet containing the outcome variable
- **sep** (*str*) Separator to parse the spreadsheet-like file.
- index_col (*int*) Which column to use as index (should correspond to subject-name).
- **normalize** (bool) Whether to normalize (0 mean, unit std) the outcome variable.

- **remove** (*int* or *float* or *str*) Removes instances in which y == remove from MvpBetween object.
- **ensure_balanced** (bool) Whether to ensure balanced classes (if True, done by undersampling the majority class).
- **nan_strategy** (*str*) Strategy on how to deal with NaNs. Default: 'remove'. Also, a specific string, int, or float can be specified when you want to impute a specific value. Other options, see: sklearn.preprocessing.Imputer.
- ****kwargs** Arbitrary keyword arguments passed to pandas read_csv.

```
apply_binarization_params (param_file, ensure_balanced=False)
Applies binarization-parameters to y.
```

binarize_y (*params*, *save_path=None*, *ensure_balanced=False*) Binarizes mvp's y-attribute using a specified method.

Parameters

• **params** (*dict*) – The outcome variable (y) will be binarized along the key-value pairs in the params-argument. Options:

```
>>> params = {'type': 'percentile', 'high': 75, 'low': 25}
>>> params = {'type': 'zscore', 'std': 1}
>>> params = {'type': 'constant', 'cutoff': 10}
>>> params = {'type': 'median'}
```

- **save_path** (*str*) If not None (default), this should be an absolute path referring to where the binarization-params should be saved.
- **ensure_balanced** (*bool*) Whether to ensure balanced classes (if True, done by undersampling the majority class).

create()

Extracts and stores data as specified in source.

Raises ValueError – If data-type is not one of ['VBM', 'TBSS', '4D_anat*', 'dual_reg', 'Contrast*']

Runs a searchlight on the mvp object.

Parameters

- **out_dir** (*str*) Path to which to save the searchlight results
- **name** (*str*) Name for the searchlight-results-file (nifti)
- **n_folds** (*int*) The amount of folds in sklearn's StratifiedKFold.
- radius (int/list) Radius for the searchlight. If list, it iterates over radii.
- **mask** (*str*) Path to mask to apply to mvp. If nothing is listed, it will use the masks applied when the mvp was created.
- **estimator** (*sklearn estimator or pipeline*) Estimator to use in the classification process.
- **kwargs Other keyword arguments for initializing nilearn's searchlight object (see nilearn.github.io/decoding/searchlight.html).

split (*file_path, col_name, target, sep='\t', index_col=0, nan_strategy='train', **kwargs*) Splits an MypBetween object based on some external index.

Parameters

- **file_path** (*str*) Absolute path to spreadsheet-like file including the outcome var.
- col_name (str) Column name in spreadsheet containing the outcome variable
- **target** (*str* or *int* or *float*) Target to which the data in col_name needs to be compared to, in order to create an index.
- **sep** (*str*) Separator to parse the spreadsheet-like file.
- index_col (*int*) Which column to use as index (should correspond to subject-name).
- **nan_strategy** (*str*) Which value to impute if the labeling is absent. Default: 'train'.
- ****kwargs** Arbitrary keyword arguments passed to pandas read_csv.

update_sample(*idx*)

Updates the data matrix and associated attributes.

write_4D (path=None, return_nimg=False)
Writes a 4D nifti (subs = 4th dimension) of X.

Parameters

- **path** (*str*) Absolute path to save nifti to.
- **return_nimg** (*bool*) Whether to actually return the Nifti1-image object.

Extracts and stores subject-specific single-trial multivoxel-patterns The MvpWithin class allows for the extraction of subject-specific single-trial, multivoxel fMRI patterns from a FSL feat-directory.

- **source** (*str*) An absolute path to a subject-specific first-level FEAT directory.
- **read_labels** (*bool*) Whether to read the labels/targets (i.e. y) from the contrast names defined in the design.con file.
- **remove_contrast** (*list*) Given that all contrasts (COPEs) are loaded from the FEAT-directory, this argument can be used to remove irrelevant contrasts (e.g. contrasts of nuisance predictors). Entries in remove_contrast do not have to literal; they may be a substring of the full name of the contrast.
- **invert_selection** (bool) Sometimes, instead of loading in all contrasts and excluding some, you might want to load only a single or a couple contrasts, and exclude all other. By setting invert_selection to True, it treats the remove_contrast variable as a list of contrasts to include.
- **ref_space** (*str*) Indicates in which 'space' the patterns will be stored. The default is 'epi', indicating that the patterns will be loaded and stored in subject-specific (native) functional space. The other option is 'mni', which indicates that MvpWithin will first transform contrasts to MNI152 (2mm) space before it loads them. This option assumes that a 'reg' directory is present in the .feat-directory, including warp-files from functional to mni space (i.e. example_func2standara.nii.gz).
- **statistic** (*str*) Which statistic (beta = (CO)PE, tstat, zstat, etc.) from FEAT directories to use as patterns.

- **remove_zeros** (*bool*) Whether to remove features (i.e. voxels) which are 0 across all trials (due to, e.g., being located outside the brain).
- **X** (*ndarray*) Not necessary to pass MvpWithin, but needs to be defined as it is needed in the super-constructor.
- **y** (*ndarray* or *list*) Labels or targets corresponding to the samples in X. This can be used when read_labels is set to False.
- **mask** (*str*) Absolute path to nifti-file that will be used as mask.
- mask_threshold (*int or float*) Minimum value to binarize the mask when it's probabilistic.

Variables

- **mask_shape** (*tuple*) Shape of mask that patterns will be indexed with.
- **nifti_header** (*Nifti1Header object*) **Nifti-header** from corresponding mask.
- **affine** (*ndarray*) Affine corresponding to nifti-mask.
- **voxel_idx** (*ndarray*) Array with integer-indices indicating which voxels are used in the patterns relative to whole-brain space. In other words, it allows to map back the patterns to a whole-brain orientation.
- X (*ndarray*) The actual patterns (2D: samples X features)
- y (list or ndarray) Array/list with labels/targets corresponding to samples in X.
- **contrast_labels** (*list*) List of names corresponding to the y-values.

create()

Extracts (meta-)data from FEAT-directory given appropriate settings during initialization.

Raises

- ValueError If the 'source'-directory doesn't exist.
- ValueError If the number of loaded contrasts does not equal the number of extracted labels.

skbold.exp_model package

The exp_model subpackage contains some (pre)processing functions and classes that help in preparing to fit a first-level GLM on fMRI data across multiple subjects.

The PresentationLogfileCrawler (and its function-equivalent 'parse_presentation_logfile') can be used to parse Presentation-logfile, which are often used at the University of Amsterdam.

Also, there is an experimental Eprime-logfile converter, which converts the Eprime .txt-file to a tsv-file format.

Function-interface for PresentationLogfileCrawler. Can be used to create a Nipype node.

Parameters

- **in_file** (*str or list*) Absolute path to logfile (can be a list of paths).
- **con_names** (*list*) List with names for each condition
- **con_codes** (*list*) List with codes for conditions. Can be a single integer or string (in the latter case, it may be a substring) or a list with possible values.
- **con_design** (*list* or *str*) Which 'design' to assume for events (if 'multivar', all events regardless of condition are treated as a separate condition/regressor; if 'univar', all events from a single condition are treated as a single condition). Default: 'univar' for all conditions.
- **con_duration** (*list*) If the duration cannot be parsed from the logfile, you can specify them here manually (per condition).
- **pulsecode** (*int*) Code with which the first (or any) pulse is logged.

class PresentationLogfileCrawler (*in_file*, *con_names*, *con_codes*, *con_duration=None*, *pulsec-ode=30*, *write_tsv=True*, *verbose=True*, *write_code=False*)

Bases: object

Logfile crawler for Presentation (Neurobs) files; cleans logfile, calculates event onsets and durations, and (optionally) writes out .bfsl files per condition.

- in_file (*str or list*) Absolute path to logfile (can be a list of paths).
- con_names (list) List with names for each condition
- **con_codes** (*list*) List with codes for conditions. Can be a single integer or string (in the latter case, it may be a substring) or a list with possible values.
- **con_design** (*list* or *str*) Which 'design' to assume for events (if 'multivar', all events regardless of condition are treated as a separate condition/regressor; if 'univar', all events from a single condition are treated as a single condition). Default: 'univar' for all conditions.
- **con_duration** (*list*) If the duration cannot be parsed from the logfile, you can specify them here manually (per condition).
- **pulsecode** (*int*) Code with which the first (or any) pulse is logged.
- write_bfsl (bool) Whether to write out a .bfsl file per condition.
- **verbose** (*bool*) Print out intermediary output.

Variables df (Dataframe) – Dataframe with cleaned and parsed logfile.

parse()

Parses logfile, writes bfsl (optional), and return subject-info.

Returns subject_info_list – Bunch object to be used in Nipype pipelines.

Return type Nilearn bunch object

class Eprime2tsv(*in_file*)

Bases: object

Converts Eprime txt-files to tsv.

Parameters in_file (*str*) – Absolute path to Eprime txt-file.

Variables df (Dataframe) – Pandas dataframe with parsed and cleaned txt-file

convert (*out_dir=None*) Converts txt-file to tsv.

Parameters out_dir(*str*) – Absolute path to desired directory to save tsv to (default: current directory).

Bases: object

Given an fsf-template, this crawler creates subject-specific fsf-FEAT files assuming that appropriate .bfsl files exist.

- **data_dir** (*str*) Absolute path to directory with BIDS-formatted data.
- **run_idf** (*str*) Identifier for run to apply template fsf to.
- **template** (*str*) Absolute path to template fsf-file. Default is 'mvpa', which models each event as a separate regressor (and contrast against baseline).
- **preprocess** (*bool*) Whether to apply preprocessing (as specified in template) or whether to only run statistics/GLM.
- **register** (bool) Whether to calculate registration (func -> highres, highres -> standard)

- **mvpa_type** (*str*) Whether to estimate patterns per trial (mvpa_type='trial_wise') or to estimate patterns per condition (or per run, mvpa_type='run_wise')
- **output_dir** (*str*) Path to desired output dir of first-levels.
- **subject_idf** (*str*) Identifier for subject-directories.
- event_file_ext (*str*) Extension for event-file; if 'bfsl/txt' (default, for legacy reasons), then assumes single event-file per predictor. If 'tsv' (cf. BIDS), then assumes a single tsv-file with all predictors.
- **sort_by_onset** (*bool*) Whether to sort predictors by onset (first trial = first predictor), or, when False, sort by condition (all trials condition A, all trials condition B, etc.).
- n_cores (int) How many CPU cores should be used for the batch-analysis.
- **feat_options** (*key-word arguments*) Which preprocessing options to set (only relevant if template='mvpa' or if you want to deviate from template). Examples:

mc='1' (apply motion correction), st='1' (apply regular-up slice-time correction), bet_yn='1' (do brain extraction of func-file), smooth='5.0' (smooth with 5 mm FWHM), temphp_yn='1' (do HP-filtering), paradigm_hp='100' (set HP-filter to 100 seconds), prewhiten_yn='1' (do prewhitening), motionevs='1' (add motion-params as nuisance regressors)

crawl()

Crawls subject-directories and spits out subject-specific fsf.

skbold.feature_extraction package

This module contains some feature-extraction methods/transformers.

```
class PatternAverager(method='mean')
```

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Reduces the set of features to its average.

Parameters method (*str*) – method of averaging (either 'mean' or 'median')

fit (X=None, y=None)

Does nothing, but included to be used in sklearn's Pipeline.

 $\texttt{transform}\left(X\right)$

Transforms patterns to its average.

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X_new – Transformed ndarray of shape = [n_samples, 1]

Return type ndarray

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Transforms a whole-brain voxel pattern into a region-average pattern Computes the average from different regions from a given parcellation and returns those as features for X.

- **atlas** (*str*) Atlas to extract ROIs from. Available: 'HarvardOxford-Cortical', 'HarvardOxford-Subcortical', 'HarvardOxford-All' (combination of cortical/subcortical), 'Talairach' (not tested), 'JHU-labels', 'JHU-tracts', 'Yeo2011'.
- mvp (Mvp-object (see core.mvp)) Mvp object that provides some metadata about previous masks

- mask_threshold (*int* (*default:* 0)) Minimum threshold for probabilistic masks (such as Harvard-Oxford)
- **reg_dir** (*str*) Path to directory with registration info (warps/transforms).
- ****kwargs** (key-word arguments) Other arguments that can be passed to *skbold.utils.load_roi_mask.*

```
fit (X=None, y=None)
```

Does nothing, but included to be used in sklearn's Pipeline.

transform(X, y=None)

Transforms features from X (voxels) to region-average features.

Parameters

- X (*ndarray*) Numeric (float) array of shape = [n_samples, n_features]
- **y** (Optional[List[str] or numpy ndarray[str]]) List of ndarray with strings indicating label-names
- **Returns X_new** array with transformed data of shape = [n_samples, n_features] in which features are region-average values.

Return type ndarray

class PCAfilter (*n_components=5*, *reject=None*)

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Filters out a (set of) PCA component(s) and transforms it back to original representation.

Parameters

- n_components (int) number of components to retain.
- **reject** (*list*) Indices of components which should be additionally removed.

Variables pca (scikit-learn PCA object) - Fitted PCA object.

fit (X, y=None, *args)

Fits PcaFilter.

Parameters

- X (*ndarray*) Numeric (float) array of shape = [n_samples, n_features]
- **y** (*List of str*) List or ndarray with floats corresponding to labels

transform(X)

Transforms a pattern (X) by the inverse PCA transform with removed components.

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X – Transformed array of shape = [n_samples, n_features] given the PCA calculated during fit().

Return type ndarray

class ClusterThreshold (*mvp*, *min_score*, *selector=<function* <u>f_classif></u>, *min_cluster_size=20*) **Bases:** sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Implements a cluster-based feature selection method. This feature selection method performs a univariate feature selection method to yield a set of voxels which are then cluster-thresholded using a minimum (contiguous) cluster size. These clusters are then averaged to yield a set of cluster-average features. This method is described in detail in my master's thesis: github.com/lukassnoek/MSc_thesis.

- **transformer** (*scikit-learn style transformer class*) transformer class used to perform some kind of univariate feature selection.
- **mvp** (*Mvp*-object (see core.*mvp*)) Necessary to provide mask metadata (index, shape).
- **min_cluster_size** (*int*) minimum cluster size to be set for cluster-thresholding

fit (X, y, *args)

Fits ClusterThreshold transformer.

Parameters

- **X** (*ndarray*) Numeric (float) array of shape = [n_samples, n_features]
- **y** (*List[str]* or *numpy ndarray[str]*) List of ndarray with floats corresponding to labels

transform(X)

Transforms a pattern (X) given the indices calculated during fit().

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X_cl – Transformed array of shape = [n_samples, n_clusters] given the indices calculated during fit().

Return type ndarray

skbold.feature_selection package

The transformer subpackage provides several scikit-learn style transformers that perform feature selection and/or extraction of multivoxel fMRI patterns. Most of them are specifically constructed with fMRI data in mind, and thus often need an Mvp object during initialization to extract necessary metadata. All comply with the scikit-learn API, using fit() and transform() methods.

class GenericUnivariateSelect (*score_func=<function f_classif>*, *mode='percentile'*, *param=le-*

05) Bases: sklearn.feature_selection.univariate_selection._BaseFilter

Univariate feature selector with configurable strategy.

Updated version from scikit-learn: http://scikit-learn.org/6.

Parameters

- **score_func** (*callable*) Function taking two arrays X and y, and returning a pair of arrays (scores, pvalues). For modes 'percentile' or 'kbest' it can return a single array scores.
- mode ({'percentile', 'k_best', 'fpr', 'fdr', 'fwe', 'cutoff'})
 - Feature selection mode.
- **param** (float or int depending on the feature selection mode) Parameter of the corresponding mode.

Variables

- **scores** (*array-like*, *shape=*(*n_features*,)) **Scores** of features.
- **pvalues** (*array-like*, *shape=(n_features,)*) p-values of feature scores, None if *score_func* returned scores only.

class SelectAboveCutoff (*cutoff*, *score_func=<function f_classif>*)

Bases: sklearn.feature_selection.univariate_selection._BaseFilter

Filter: Select features with a score above some cutoff.

Parameters

• **cutoff** (*int/float*) – Cutoff for feature-scores to be selected.

• **score_func** (*callable*) – Function that takes a 2D array X (samples x features) and returns a score reflecting a univariate difference (higher is better).

fisher_criterion_score (X, y, norm='ll', balance=False)

Calculates fisher score.

See [1]_ for more info.

References

[1] P. E. H. R. O. Duda and D. G. Stork. Pattern Classification. Wiley-Interscience Publication, 2001.

Parameters

- X ({array-like, sparse matrix} shape = (n_samples, n_features)) The set of regressors that will be tested sequentially.
- **y**(array of shape(n_samples)) The data matrix
- **norm** (*str*) Whether to use the 11-norm or 12-norm.

Returns scores_ – Fisher criterion scores for each feature.

Return type array, shape=(n_features,)

class IncrementalFeatureCombiner(scores, cutoff)

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Indexes a set of features with a number of (sorted) features.

Parameters

- **scores** (*ndarray*) Array of shape = n_features, or [n_features, n_class] in case of soft/hard voting in, e.g., a roi_stacking_classifier (see classifiers.roi_stacking_classifier).
- **cutoff** (*int* or *float*) If int, it refers the absolute number of features included, sorted from high to low (w.r.t. scores). If float, it selects a proportion of features.

fit (X, y=None)

Fits IncrementalFeatureCombiner transformer.

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

transform (X, y=None)

Transforms a pattern (X) given the indices calculated during fit().

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X – Transformed array of shape = [n_samples, n_features] given the indices calculated during fit().

Return type ndarray

skbold.pipelines package

The pipelines module contains some standard MVPA pipelines using the scikit-learn style Pipeline objects.

```
create_ftest_kbest_svm (kernel='linear', k=100, **kwargs)
Creates an svm-pipeline with f-test feature selection.
```

Uses SelectKBest from scikit-learn.feature_selection.

Parameters

- kernel (str) Kernel for SVM (default: 'linear')
- **k** (*int*) How many voxels to select (from the k best)
- ****kwargs** Arbitrary keyword arguments for SVC() initialization.

Returns ftest_svm – Pipeline with f-test feature selection and svm.

Return type scikit-learn Pipeline object

create_ftest_percentile_svm(kernel='linear', perc=10, **kwargs)

Creates an svm-pipeline with f-test feature selection.

Uses SelectPercentile from scikit-learn.feature_selection.

Parameters

- kernel (str) Kernel for SVM (default: 'linear')
- perc (int or float) Percentage of voxels to select
- ****kwargs** Arbitrary keyword arguments for SVC() initialization.

Returns ftest_svm - Pipeline with f-test feature selection and svm.

Return type scikit-learn Pipeline object

create_pca_svm (*kernel='linear'*, *n_comp=10*, *whiten=False*, ***kwargs*) Creates an svm-pipeline with f-test feature selection.

Parameters

• kernel (str) – Kernel for SVM (default: 'linear')

- **n_comp** (*int*) How many PCA-components to select
- whiten (bool) Whether to use whitening in PCA
- ****kwargs** Arbitrary keyword arguments for SVC() initialization.

Returns pca_svm – Pipeline with PCA feature extraction and svm.

Return type scikit-learn Pipeline object

skbold.postproc package

The postproc subpackage contains all off skbold's 'postprocessing' tools. Most prominently, it contains the MvpResults objects (both MvpResultsClassification and MvpResultsRegression) which can be used in analyses to keep track of model performance across iterations/folds (in cross-validation). Additionally, it allows for keeping track of featurescores (e.g. f-values from the univariate feature selection procedure) or model weights (e.g. SVM-coefficients). These coefficients can kept track of as raw weights [1]_ or as 'forward-transformed' weights [2]_.

The postproc subpackage additionally contains the function 'extract_roi_info', which allows to calculate the amount of voxels (and other statistics) per ROI in a single statistical brain map and output a csv-file.

The cluster_size_threshold function allows you to set voxels to zero which do not belong to a cluster of a given extent/size. This is NOT a statistical procedure (like GRF thresholding), but merely a tool for visualization purposes.

References

R., and Turner, R. (2014). Prioritizing spatial accuracy in high-resolution fMRI data using multivariate feature weight mapping. Front. Neurosci., http://dx.doi.org/10.3389/fnins.2014.00066.

Blankertz, B., and Biessmann, F. et al. (2014). On the interpretation of weight vectors of linear models in multivariate neuroimaging. Neuroimage, 87, 96-110.

extract_roi_info(statfile, stat_name=None, roi_type='unilateral', per_cluster=True, cluster_engine='scipy', min_clust_size=20, stat_threshold=None, mask_threshold=20, save indices=True, verbose=True)

Extracts information per ROI for a given statistics-file. Reads in a thresholded (!) statistics-file (such as a thresholded z- or t-stat from a FSL first-level directory) and calculates for a set of ROIs the number of significant voxels included and its maximum value (+ coordinates). Saves a csv-file in the same directory as the statistics-file. Assumes that the statistics file is in MNI152 2mm space.

- **statfile** (*str*) Absolute path to statistics-file (nifti) that needs to be evaluated.
- **stat_name** (*str*) Name for the contrast/stat-file that is being analyzed.

- **roi_type** (*str*) Whether to use unilateral or bilateral masks (thus far, only Harvard-Oxford atlas masks are supported.)
- **per_cluster** (bool) Whether to evaluate the statistics-file as a whole (per_cluster=False) or per cluster separately (per_cluster=True).
- **cluster_engine** (*str*) Which 'engine' to use for clustering; can be 'scipy' (default), using scipy.ndimage.measurements.label, or 'fsl' (using FSL's cluster command).
- **min_clust_size** (*int*) Minimum cluster size (i.e. clusters with fewer voxels than this number are discarded; also, ROIs containing fewer voxels than this will not be listed on the CSV.
- **stat_threshold** (*int or float*) If the stat-file contains uncorrected data, stat_threshold can be used to set a lower bound.
- mask_threshold (bool) Threshold for probabilistics masks, such as the Harvard-Oxford masks. Default of 25 is chosen as this minimizes overlap between adjacent masks while still covering most of the entire brain.
- **save_indices** (bool) Whether to save the indices (coordinates) of peaks of clusters.
- **verbose** (bool) Whether to print some output regarding the parsing process.

Returns df – Dataframe corresponding to the written csv-file.

Return type Dataframe

class MvpResults (*mvp*, *n_iter*, *type_model='classification'*, *feature_scoring=None*, *confmat=False*, *verbose=False*, ***metrics*)

Bases: object

Class to keep track of model evaluation metrics and feature scores. See the ReadTheDocs homepage for more information on its API and use.

Parameters

- **mvp** (*mvp-object*) Necessary to extract some metadata from.
- n_iter (*int*) Number of folds that will be kept track of.
- type_model (*str*) Either 'classification' or 'regression'
- feature_scoring (str) Which method to use to calculate feature-scores with. Can be: 1) 'fwm': feature weight mapping [1] keep track of raw voxel-weights (coefficients) 2) 'forward': transform raw voxel-weights to corresponding forward- model [2].
- confmat (bool) Whether to keep track of the confusion-matrix across folds (only relevant for type_model='classification')
- **verbose** (*bool*) Whether to print extra output.
- ****metrics** (keyword-arguments) Keyword arguments of the form: *name_metric: metric_function*; any metric from scikit-learn works (or other metrics, as long as they have two input args, *y_true* and *y_pred*).

References

```
compute_scores (multiclass='ovr', maps_to_tstat=True)
Computes scores across folds.
```

```
load_model (path, param=None)
Load model or pipeline from disk.
```

Parameters

- **path** (*str*) Absolute path to model.
- **param** (*str*) Which, if any, specific param needs to be loaded.

save_model (model, out_path)

Method to serialize model(s) to disk.

Parameters model (pipeline or scikit-learn object.) - Model to be saved.

update (*test_idx*, *y_pred*, *pipeline=None*) Updates with information from current fold.

Parameters

- test_idx (ndarray) Indices of current test-trials.
- **y_pred** (*ndarray*) Predictions of current test-trials.
- **pipeline** (*scikit-learn Pipeline object*) **pipeline** from which relevant scores/coefficients will be extracted.

write (out_path, confmat=True, to_tstat=True, multiclass='ovr')
Writes results to disk.

Parameters

- **out_path** (*str*) Where to save the results to
- **feature_viz** (bool) Whether to write out (and optionally return) feature-visualization information
- **confmat** (*bool*) Whether to write out (and optionally return) the confusion-matrix (across folds).
- to_tstat (bool) Whether to convert averaged feature-scores to t-tstats (by dividing them by sqrt(score.std(axis=0)).

class MvpAverageResults (mvp_results_list, identifiers=None)

Bases: object

Averages results from MVPA analyses on, for example, different subjects or different ROIs.

Parameters

- mvp_results_list (list) List with MvpResults objects (after updating across folds)
- identifiers (*list of str*) List of identifiers (e.g. subject-name) that correspond to the different MvpResults objects

compute_statistics (*metric='accuracy'*, *h0=0.5*) Computes statistics across MvpResults objects

Parameters

- metric (str) Which metric should be used in the MvpResults dataframes
- h0 (float) The null-hypothesis in terms of model performance (e.g. accuracy equals 1/n_classes)

write (path, name='average_results')

cluster_size_threshold (*data*, thresh=None, min_size=20, save=False)

Removes clusters smaller than a prespecified number in a stat-file.

Parameters

- **data** (*numpy-array or str*) 3D Numpy-array with statistic-value or a string to a path pointing to a nifti-file with statistic values.
- thresh (*int*, *float*) Initial threshold to binarize the image and extract clusters.
- min_size (*int*) Minimum size (i.e. amount of voxels) of cluster. Any cluster with fewer voxels than this amount is set to zero ('removed').
- **save** (bool) If data is a file-path, this parameter determines whether the cluster- corrected file is saved to disk again.

class PrevalenceInference (obs, perms, P2=100000, gamma0=0.5, alpha=0.05)

Bases: object

Class that performs PrevalenceInference based on the paper by Allefeld, Gorgen, & Haynes (2016), NeuroImage.

Parameters

- **obs** (*numpy ndarray*) A 2D array of shape [N (subjects) x K (voxels)], or a 1D array of shape [N, 1].
- **perms** (*numpy ndarray*) A 3D array of shape [N (subjects) x K (voxels) x P1 (first level permutations)], or a 2D array of shape [N x P1]
- P2 (int) Number of second level permutations to run
- gamma0 (float) What prevalence inference null (gamma < gamma0) to test
- **alpha** (*float*) Significance level for hypothesis testing

Examples

___init___(obs, perms, P2=100000, gamma0=0.5, alpha=0.05) Initializes PrevalenceInference object.

run()

Runs actual prevalence inference algorithm.

write(path)

Writes results from Prevalence Inference procedure to disk.

Parameters path (str) – Where to write the results to disk

skbold.preproc package

class LabelFactorizer(grouping)

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Transforms labels according to a given factorial grouping.

Factorizes/encodes labels based on part of the string label. For example, the label-vector ['A_1', 'A_2', 'B_1', 'B_2'] can be grouped based on letter (A/B) or number (1/2).

Parameters grouping (List of str) – List with identifiers for condition names as strings

Variables new_labels (*list*) – List with new labels.

```
fit (y=None, X=None)
```

Does nothing, but included to be used in sklearn's Pipeline.

```
get_new_labels()
```

Returns new labels based on factorization.

transform(*y*, *X*=*None*)

Transforms label-vector given a grouping.

Parameters

- **y** (List/ndarray of str) List of ndarray with strings indicating label-names
- X (*ndarray*) Numeric (float) array of shape = [n_samples, n_features]

Returns

- y_new (ndarray) array with transformed y-labels
- **X_new** (*ndarray*) array with transformed data of shape = [n_samples, n_features] given new factorial grouping/design.

class MajorityUndersampler(verbose=False)

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Undersamples the majority-class(es) by selecting random samples.

Parameters verbose (bool) – Whether to print downsamples number of samples.

___init___(verbose=False) Initializes MajorityUndersampler object.

fit (X=None, y=None)

Does nothing, but included for scikit-learn pipelines.

transform (X, y)

Downsamples majority-class(es).

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X – Transformed array of shape = [n_samples, n_features] given the indices calculated during fit().

Return type ndarray

class LabelBinarizer(params)

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

```
__init__ (params)
```

Initializes LabelBinarizer object.

fit (X=None, y=None)
 Does nothing, but included for scikit-learn pipelines.

transform(X, y)

Binarizes y-attribute.

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X – Transformed array of shape = [n_samples, n_features] given the indices calculated during fit().

Return type ndarray

class ConfoundRegressor (*confound*, *X*, *cross_validate=True*, *precise=False*, *stack_intercept=True*) Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Fits a confound onto each feature in X and returns their residuals.

___init___(confound, X, cross_validate=True, precise=False, stack_intercept=True) Regresses out a variable (confound) from each feature in X.

- **confound** (*numpy array*) Array of length (n_samples, n_confounds) to regress out of each feature; May have multiple columns for multiple confounds.
- **X** (*numpy array*) Array of length (n_samples, n_features), from which the confound will be regressed. This is used to determine how the confound-models should be cross-validated (which is necessary to use in in scikit-learn Pipelines).
- **cross_validate** (*bool*) Whether to cross-validate the confound-parameters (*y*~confound) estimated from the train-set to the test set (cross_validate=True) or whether to fit the confound regressor separately on the test-set (cross_validate=False). Setting this parameter to True is equivalent to "foldwise confound regression" (FwCR) as described in our paper (https://www.biorxiv.org/content/early/2018/03/28/290684). Setting this parameter to False, however, is NOT equivalent to "whole dataset confound regression" (WDCR) as it does not apply confound regression to the *full* dataset, but simply refits the confound model on the test-set. We recommend setting this parameter to True.
- **precise** (bool) Transformer-objects in scikit-learn only allow to pass the data (X) and optionally the target (y) to the fit and transform methods. However, we need to index the confound accordingly as well. To do so, we compare the X during initialization (self.X)

with the X passed to fit/transform. As such, we can infer which samples are passed to the methods and index the confound accordingly. When setting precise to True, the arrays are compared feature-wise, which is accurate, but relatively slow. When setting precise to False, it will infer the index by looking at the sum of all the features, which is less accurate, but much faster. For dense data, this should work just fine. Also, to aid the accuracy, we remove the features which are constant (0) across samples.

• **stack_intercept** (*bool*) – Whether to stack an intercept to the confound (default is True)

Variables weights (*numpy array*) – Array with weights for the confound(s).

fit (X, y=None)

Fits the confound-regressor to X.

Parameters

- X (*numpy array*) An array of shape (n_samples, n_features), which should correspond to your train-set only!
- **y** (*None*) Included for compatibility; does nothing.

transform(X)

Regresses out confound from X.

Parameters X (*numpy array*) – An array of shape (n_samples, n_features), which should correspond to your train-set only!

Returns X_new – ndarray with confound-regressed features

Return type ndarray

skbold.utils package

The utils subpackage contains some extra utilities for machine learning pipelines on fMRI data. For example, the *CrossvalSplitter* creates balanced train/test-sets given a (set of) confound(s). Also, the *load_roi_mask* function allows for loading ROIs from the Harvard-Oxford (sub)cortical atlas. This function is also integrated in the *RoiIndexer* transformer.from

Lastly, the ArrayPermuter, RowIndexer, and SelectFeatureset transformers can be used in, for example. permutation analyses.

sort_numbered_list (stat_list)

Sorts a list containing numbers.

Sorts list with paths to statistic files (e.g. COPEs, VARCOPES), which are often sorted wrong (due to single and double digits). This function extracts the numbers from the stat files and sorts the original list accordingly.

Parameters stat_list (list or str) – list with absolute paths to files

Returns sorted_list - sorted stat_list

Return type list of str

class CrossvalSplitter(data, train_size, vars, cb_between_splits=False, binarize=None, include=None, exclude=None, interactions=True, sep='t', index_col=0, ignore=None, iterations=1000)

Bases: object

plot_results (out_dir)

save (out_dir, save_plots=True)

split (verbose=False)

parse_roi_labels (atlas_type='Talairach', lateralized=False, debug=False)
Parses xml-files belonging to FSL atlases.

Parameters

• **atlas_type** (*str*) – String identifying which atlas needs to be parsed.

• **lateralized** (*bool*) – Whether to use the lateralized version of the atlas (only applicable to HarvardOxford masks)

Returns info_dict – Dictionary with indices and coordinates (values) per ROI (keys).

Return type dict

print_mask_options (atlas_name='HarvardOxford-Cortical')

Prints the options for ROIs given a certain atlas.

Parameters atlas_name (*str*) – Name of the atlas. Availabel: 'HarvardOxford-Cortical', 'HarvardOxford-Subcortical', 'Yeo2011'.

class ArrayPermuter

Bases: sklearn.base.BaseEstimator, sklearn.base.TransformerMixin

Permutes (shuffles) rows of matrix.

___init__()

Initializes ArrayPermuter object.

fit (X=None, y=None)
Does nothing, but included to be used in sklearn's Pipeline.

transform(X)

Permutes rows of data input.

Parameters X (*ndarray*) – Numeric (float) array of shape = [n_samples, n_features]

Returns X_new – ndarray with permuted rows

Return type ndarray

class RowIndexer(mvp, train_idx)

Bases: object

Selects a subset of rows from an Mvp object.

Notes

NOT a scikit-learn style transformer.

Parameters

- **idx** (*ndarray*) Array with indices.
- **mvp** (*mvp*-object) **Mvp**-object to drawn metadata from.

transform()

Returns

- **mvp** (*mvp-object*) Indexed mvp-object.
- X_not_selected (*ndarray*) Data which has not been selected.
- **y_not_selected** (*ndarray*) Labels which have not been selected.

class SelectFeatureset (mvp, featureset_idx)

 $Bases: \verb+sklearn.base.BaseEstimator, \verb+sklearn.base.TransformerMixin+$

Selects only columns of a certain featureset. CANNOT be used in a scikit-learn pipeline!

Parameters

• **mvp** (*mvp*-*object*) – Used to extract meta-data.

• **featureset_idx** (*ndarray*) – Array with indices which map to unique feature-set voxels.

fit()

Does nothing, but included due to scikit-learn API.

transform(X=None)

Transforms mvp.

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