

---

# PyPairs Documentation

*Release v3.1.1+10.g2dcd414*

**R. Fechtner, A. Scialdone**

**Jul 11, 2019**

---

## Contents

---

<b>1 PyPairs - A python scRNA-Seq classifier</b>	<b>2</b>
1.1 Core Dependencies . . . . .	2
1.2 Authors . . . . .	2
1.3 Release notes . . . . .	2
1.3.1 Announcement . . . . .	2
1.3.2 Versions . . . . .	3
<b>2 Getting Started</b>	<b>5</b>
2.1 Installation . . . . .	5
2.2 Minimal Example . . . . .	5
<b>3 Documentation</b>	<b>6</b>
3.1 Sandbag . . . . .	6
3.1.1 pypairs.pairs.sandbag . . . . .	6
3.2 Cyclone . . . . .	8
3.2.1 pypairs.pairs.cyclone . . . . .	8
3.3 Datasets . . . . .	9
3.3.1 pypairs.datasets.leng15 . . . . .	9
3.3.2 pypairs.datasets.default_cc_marker . . . . .	10
3.4 Quality Assesment . . . . .	10
3.4.1 pypairs.utils.evaluate_prediction . . . . .	10
3.5 Utils . . . . .	11
3.5.1 pypairs.utils.export_marker . . . . .	11
3.5.2 pypairs.utils.load_marker . . . . .	11
3.6 Settings . . . . .	11
3.6.1 pypairs.log.print_versions . . . . .	12
<b>4 References</b>	<b>13</b>
<b>Bibliography</b>	<b>14</b>
<b>Python Module Index</b>	<b>15</b>
<b>Index</b>	<b>16</b>



# CHAPTER 1

---

## PyPairs - A python scRNA-Seq classifier

---

This is a python-reimplementation of the Pairs algorithm as described by A. Scialdone et. al. (2015). Original Paper available under: <<https://doi.org/10.1016/j.ymeth.2015.06.021>>

A supervised machine learning algorithm aiming to classify single cells based on their transcriptomic signal. Initially created to predict cell cycle phase from scRNA-Seq data, this algorithm can be used for various applications.

Build to be fully compatible with [Scanpy](#) [Wolf18].

Code available on [GitHub](#).

### 1.1 Core Dependencies

- [Numpy](#)
- [Numba](#)
- [Pandas](#)
- [AnnData](#)

### 1.2 Authors

- **Antonio Scialdone** - *original algorithm*
- **Ron Fechtner** - *implementation and extension in Python*

### 1.3 Release notes

#### 1.3.1 Announcement

---

**Note:** Please only use `pypairs` >= 3.1.0

---

### 1.3.2 Versions

#### Version 3.1.0, Apr 4, 2019

- **New feature:**
  - Multithreading now available for `pais.cyclone()`
- **Minor changes and fixes:**
  - `pais.sandbag()` now significantly faster
  - `pais.sandbag()` more stable in terms of memory access

#### Version 3.0.1 - 3.0.13, Mar 13, 2019

- **Various bug fixes, including:**
  - Bioconda compatibility
  - Dataset loading
  - Cache file required
  - Cell Cycle specific scoring

#### Version 3.0.0, Jan 18, 2019 - Jan 31, 2019

- Complete restructuring of the package. Now fully compatible with `scanpy`.
- **Added:**
  - This documentation
  - Default (oscope) dataset & marker pairs [Leng15]
- **Changed:**
  - `sandbag()` and `cyclone()` now accept `AnnData`, `DataFrame` and `ndarray`
  - Multiprocessing now completely handled by `numba`.

#### Version 2.0.1 - 2.0.6, Nov 22, 2018

- Minor bug fixes and improvements.

#### Version 2.0.0, Aug 14, 2018

- Major restructuring of the package
- Improved parallel processing
- **New features:**
  - `sandbag()` and `cyclone()` can now deal with any number of classes to predict

**Version 1.0.1 - 1.0.3, Jul 29, 2018**

- Bug fixes and improvements. (Mostly bugs though)
- Added multi-core processing

**Version 1.0.0, Mar 4, 2018**

- Speed and performance improvements.

**Version 0.1, Feb 22, 2018**

- Simple python reimplementation of the Pairs algorithm.
- Included sandbag() and cyclone() algorithms

# CHAPTER 2

---

## Getting Started

---

### 2.1 Installation

This package is hosted at PyPi (<https://pypi.org/project/pypairs/>) and can be installed on any system running Python3 via pip with:

```
pip install pypairs
```

Alternatively, pypairs can be installed using Conda (most easily obtained via the Miniconda Python distribution):

```
conda install -c bioconda pypairs
```

### 2.2 Minimal Example

*Datasets* provide a example scRNA dataset and default marker pairs for cell cycle prediction:

```
from pypairs import pairs, datasets

# Load samples from the oscope scRNA-Seq dataset with known cell cycle
training_data = datasets.leng15(mode='sorted')

# Run sandbag() to identify marker pairs
marker_pairs = pairs.sandbag(training_data, fraction=0.6)

# Load samples from the oscope scRNA-Seq dataset without known cell cycle
testing_data = datasets.leng15(mode='unsorted')

# Run cyclone() score and predict cell cycle classes
result = pairs.cyclone(testing_data, marker_pairs)

# Further downstream analysis
print(result)
```

# CHAPTER 3

---

## Documentation

---

To use PyPairs import the package as i.e. follows:

```
from pypairs import pairs, datasets, settings, utils
```

### 3.1 Sandbag

This function implements the classification step of the pair-based prediction method described by Scialdone et al. (2015) [Scialdone15].

To illustrate, consider classification of cells into G1 phase. Pairs of marker genes are identified with `sandbag()`, where the expression of the first gene in the training data is greater than the second in G1 phase but less than the second in all other phases.

---

`pairs.sandbag(data[, annotation, ...])`

Calculate âĂŹmarker pairsâĂŹ from a genecount matrix.

---

#### 3.1.1 pypairs.pairs.sandbag

```
pypairs.pairs.sandbag(data, annotation=None, gene_names=None, sample_names=None, fraction=0.65, filter_genes=None, filter_samples=None)
```

Calculate âĂŹmarker pairsâĂŹ from a genecount matrix. Cells x Genes.

A Pair of genes ( $g1, g2$ ) is considered a marker for a category if its expression changes from  $g1 > g2$  in one category to  $g1 < g2$  in all other categories, for at least a fraction of cells in this category.

data can be of type `AnnData`, `DataFrame` or `ndarray` and should contain the raw or normalized gene counts of shape `n_obs * n_vars`. Rows correspond to cells and columns to genes.

- If data is `AnnData` object, the category for each sample should be in `in data.vars['category']`, gene names in `data.var_names` and sample names in `data.obs_names`.

- If data is `DataFrame` object, gene names can be in `df.columns` or passed via `gene_names` and sample names in `df.index` or passed via `sample_names`. The category for each sample must be passed via annotation.
  - annotation must be in form of `{category_1: [sample_1, sample_2], ...}`. List of samples for indexing can be integer, str or a boolean mask of `len(sample_names)`.
- If data `ndarray`, all information must be passed via annotation, `gene_names` and `sample_names` parameters.

Marker pairs are returned as a mapping from category to list of 2-tuple Genes: `{category: [(Gene_1, Gene_2), ...]}`

#### Parameters

`data: Union[AnnData, DataFrame, ndarray, Collection[Collection[float]]]`

The (annotated) data matrix of shape `n_obs * n_vars`. Rows correspond to cells and columns to genes.

`annotation: Optional[Mapping[str, Collection[Union[str, int, bool]]]]`

Mapping from category to genes. If data is not `AnnData`, this is required. List of genes can be index, names or logical mask.

`gene_names: Optional[Collection[str]]` Names for genes, must be same length as `n_vars`. If data is not `AnnData`, this is required.

`sample_names: Optional[Collection[str]]` Names for samples, must be same length as `n_obs`. If data is not `AnnData`, this is required.

`fraction: float` Fraction of cells per category where marker criteria must be satisfied. Default: 0.65

`filter_genes: Optional[Collection[Union[str, int, bool]]]` A list of genes to keep. If not `None` all genes not in this list will be removed. List can be index, names or logical mask.

`filter_samples: Optional[Collection[Union[str, int, bool]]]` A list of samples to keep. If not `None` all samples not in this list will be removed. List can be index, names or logical mask.

**Return type** `Mapping[str, Collection[Tuple[str, str]]]`

**Returns** `marker_pairs_dict` – A dict mapping from str to a list of 2-tuple, where the key is the category and the list contains the marker pairs: `{category_1: [(Gene_1, Gene_2), ...]}`.

#### Examples

To generate marker pairs for a different fraction than the default (0.65) based on the bundled `oscope-dataset` [`Leng15`] run:

```
from pypairs import pairs, datasets

adata = datasets.leng15()
marker_pairs = pairs.sandbag(adata, fraction=0.5)
```

## 3.2 Cyclone

For each cell, `cyclone()` calculates the proportion of all marker pairs where the expression of the first gene is greater than the second in the new data (pairs with the same expression are ignored). A high proportion suggests that the cell is likely to belong to this category, as the expression ranking in the new data is consistent with that in the training data. Proportions are not directly comparable between phases due to the use of different sets of gene pairs for each phase. Instead, proportions are converted into scores that account for the size and precision of the proportion estimate. The same process is repeated for all phases, using the corresponding set of marker pairs in pairs.

<code>pairs.cyclone(data[, marker_pairs, ...])</code>	Score samples for each category based on marker pairs.
---	--

### 3.2.1 pypairs.pairs.cyclone

`pypairs.pairs.cyclone(data, marker_pairs=None, gene_names=None, sample_names=None, iterations=1000, min_iter=100, min_pairs=50)`

Score samples for each category based on marker pairs.

`data` can be of type `AnnData`, `DataFrame` or `ndarray` and should contain the raw or normalized gene counts of shape `n_obs * n_vars`. Rows correspond to cells and columns to genes.

- If a `AnnData` object is passed, the category scores and the final prediction will be added to `data.obs` with key `pypairs_{category}_score` and `pypairs_max_class`.
  - If marker pairs contain only the cell cycle categories G1, S and G2M an additional column `pypairs_cc_prediction` will be added. Where category S is assigned to samples where G1 and G2M score are below 0.5, as described in [Scialdone15].

`marker_pairs`, i.e. output from `sandbag()`, must be a mapping from category to list of 2-tuple Genes: `{category: [(Gene_1, Gene_2), ...]}`.

- If no `marker_pairs` are passed the default are used from `default_marker()` based on [Leng15] (marker pairs for cell cycle prediction).

#### Parameters

`data : Union[AnnData, DataFrame, ndarray, Collection[Collection[float]]]`

The (annotated) data matrix of shape `n_obs * n_vars`. Rows correspond to cells and columns to genes.

`marker_pairs : Optional[Mapping[str, Collection[Tuple[str, str]]]]` A dict mapping from str to a list of 2-tuple, where the key is the category and the list contains the marker pairs: `{Category_1: [(Gene_1, Gene_2), ...]}`. If not provided default marker pairs are used

`gene_names : Optional[Collection[str]]` Names for genes, must be same length as `n_vars`.

`sample_names : Optional[Collection[str]]` Names for samples, must be same length as `n_obs`.

`iterations : Optional[int]` An integer specifying the number of iterations for random sampling to obtain a cycle score. Default: 1000

`min_iter : Optional[int]` An integer specifying the minimum number of iterations for score estimation. Default: 100

`min_pairs : Optional[int]` An integer specifying the minimum number of pairs for cycle estimation. Default: 50

**Return type** DataFrame

**Returns**

- A DataFrame with samples as index and categories as columns with scores for each category for each
- sample and a additional column with the name of the max scoring category for each sample.
- If marker pairs contain only the cell cycle categories G1, S and G2M an additional column pypairs\_cc\_prediction will be added. Where category S is assigned to samples where G1 and G2M score are below 0.5, as described in [Scialdone15].

## Examples

To predict the cell cycle phase of the unsorted cell from the [Leng15] dataset run:

```
import pypairs import pairs, datasets
adata = datasets.leng15('unsorted')
marker_pairs = datasets.default_cc_marker()
scores = pairs.cyclone(adata, marker_pairs)
print(scores)
```

While this method is described for cell cycle phase classification, any biological groupings can be used here. However, for non-cell cycle phase groupings users should manually apply their own score thresholds for assigning cells into specific groups.

## 3.3 Datasets

<code>datasets.leng15([mode, gene_sub, sample_sub])</code>	Single cell RNA-seq data of human hESCs to evaluate Oscope [Leng15]
<code>datasets.default_cc_marker([dataset])</code>	Cell cycle marker pairs derived from [Leng15] with the default <code>sandbag()</code> settings.

### 3.3.1 pypairs.datasets.leng15

`pypairs.datasets.leng15(mode='all', gene_sub=None, sample_sub=None)`

Single cell RNA-seq data of human hESCs to evaluate Oscope [Leng15]

Total 213 H1 single cells and 247 H1-Fucci single cells were sequenced. The 213 H1 cells were used to evaluate Oscope in identifying oscillatory genes. The H1-Fucci cells were used to confirm the cell cycle gene cluster identified by Oscope in the H1 hESCs. Normalized expected counts are provided in GSE64016\_H1andFUCCI\_normalized\_EC.csv.gz

GEO-Dataset: <https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE64016>

#### Parameters

`mode : Optional[str]`

**sample selection mode:**

- ‘all’ for all samples, default

- sorted for all samples with known cell cycle (G2, S or G1)
- unsorted for all samples with unknown cell cycle (H1)

**gene\_sub** : `Optional[Iterable[int]]` Index based array of subsetted genes

**sample\_sub** : `Optional[Iterable[int]]` Index based array of subsetted samples

**Return type** AnnData

**Returns** `adata` (`AnnData`) – Annotated data matrix containing the normalized gene counts

### 3.3.2 pypairs.datasets.default\_cc\_marker

`pypairs.datasets.default_cc_marker(dataset='leng15')`

Cell cycle marker pairs derived from [Leng15] with the default `sandbag()` settings.

For description of the dataset see `leng15()`.

**Parameters**

**dataset** : `Optional[str]` placeholder. only options currently 'leng15'. Everything else with raise a `NotImplementedError`

**Return type** `Mapping[str, Collection[Tuple[str, str]]]`

**Returns** Mapping from category [`G1`, `G2M`, `S`] to list of gene marker pairs [(`Gene1`, `Gene2`), ...].

## 3.4 Quality Assesment

---

<code>utils.evaluate_prediction(prediction, reference)</code>	Calculates F1 Score, Recall and Precision of a <code>cyclone()</code> prediction.
---	---

---

### 3.4.1 pypairs.utils.evaluate\_prediction

`pypairs.utils.evaluate_prediction(prediction, reference)`

Calculates F1 Score, Recall and Precision of a `cyclone()` prediction.

**Parameters**

**prediction** : `Iterable[str]` List of predicted classes.

**reference** : `Iterable[str]` List of actual classes

**Return type** DataFrame

**Returns**

- A `DataFrame` with columns f1, recall, precision and average
- for all categories and a overall average containing the respective score.

#### Example

To get the prediction quality for the example usecase of `cyclone()` run:

```

from pypairs import pairs, datasets, utils, plotting
import numpy as np

adata = datasets.leng15('sorted')
marker_pairs = datasets.default_cc_marker()
scores = pairs.cyclone(adata, marker_pairs)

ref_labels = list(np.repeat("G2M", 76)) + list(np.repeat("S", 80)) + list(np.
    ↪repeat("G1", 91))

prediction_quality = utils.evaluate_prediction(scores['max_class'], ref_labels)

print(prediction_quality)

```

## 3.5 Utils

---

`utils.export_marker(marker, fname[, defaultpath])` Export marker pairs to json-File.

---

`utils.load_marker(fname[, defaultpath])` Export marker pairs to json-File.

---

### 3.5.1 pypairs.utils.export\_marker

`pypairs.utils.export_marker(marker, fname, defaultpath=True)`

Export marker pairs to json-File.

#### Parameters

`marker : Mapping[str, Iterable[Tuple[str, str]]]` Marker pairs as from `sandbag()`

`fname : str` Name of the json-File in the writedir (see settings)

`defaultpath : Optional[bool]` Use settings.writedir as root. Default: True

### 3.5.2 pypairs.utils.load\_marker

`pypairs.utils.load_marker(fname, defaultpath=True)`

Export marker pairs to json-File.

#### Parameters

`fname : str` Name of the json-File to write to

`defaultpath : Optional[bool]` Use settings.writedir as root. Default: True

## 3.6 Settings

The default directories for saving figures and caching files.

<code>settings.figdir</code>	Directory for saving figures (default: <code>'./figures/'</code> ).
<code>settings.cachedir</code>	Directory for cache files (default: <code>'./cache/'</code> ).

The verbosity of logging output, where verbosity levels have the following meaning: 0=error, 1=warning, 2=info, 3=hint

settings.verbosity	Verbosity level (default: 1).
--------------------	-------------------------------

Print versions of packages that might influence numerical results.

---

<code>log.print_versions()</code>	Versions that might influence the numerical results.
-----------------------------------	--

---

### 3.6.1 pypairs.log.print\_versions

`pypairs.log.print_versions()`

Versions that might influence the numerical results. Matplotlib and Seaborn are excluded from this.

## CHAPTER 4

---

### References

---

---

## Bibliography

---

- [Leng15] Leng *et al.* (2015) *Oscope identifies oscillatory genes in unsynchronized single-cell RNA-seq experiments.*, *Nat Methods*.
- [Scialdone15] Scialdone *et al.* (2015), *Computational assignment of cell-cycle stage from single-cell transcriptome data*, *Methods*.
- [Wolf18] Wolf *et al.* (2018) *SCANPY: large-scale single-cell gene expression data analysis*, *Genome Biology*.

---

## Python Module Index

---

p

[pypairs](#), 5

---

## Index

---

### C

`clone()` (*in module pypairs.pairs*), 8

### D

`default_cc_marker()` (*in module pypairs.datasets*), 10

### E

`evaluate_prediction()` (*in module pypairs.utils*), 10

`export_marker()` (*in module pypairs.utils*), 11

### L

`length()` (*in module pypairs.datasets*), 9

`load_marker()` (*in module pypairs.utils*), 11

### P

`print_versions()` (*in module pypairs.log*), 12

`pypairs(module)`, 5

### S

`sandbag()` (*in module pypairs.pairs*), 6