

# 2.5. Decomposing signals in components (matrix factorization problems)

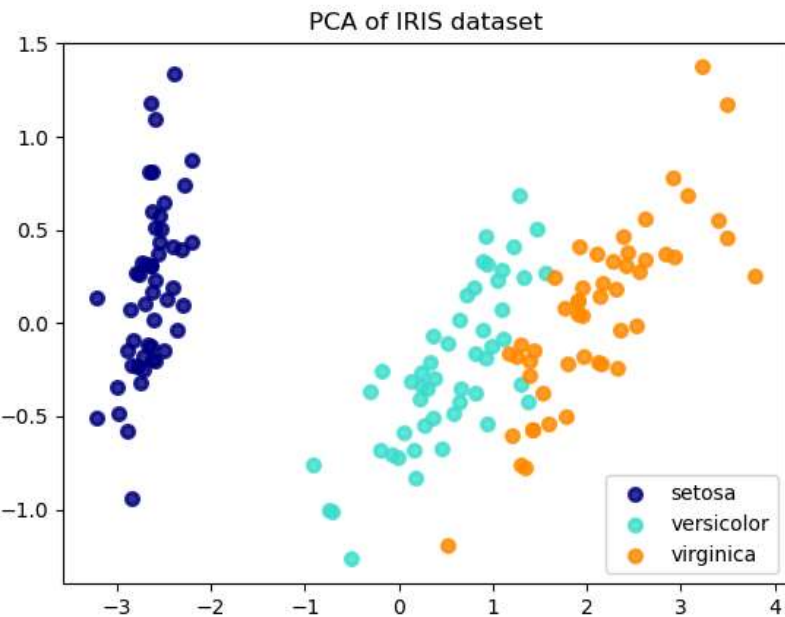
## 2.5.1. Principal component analysis (PCA)

### 2.5.1.1. Exact PCA and probabilistic interpretation #

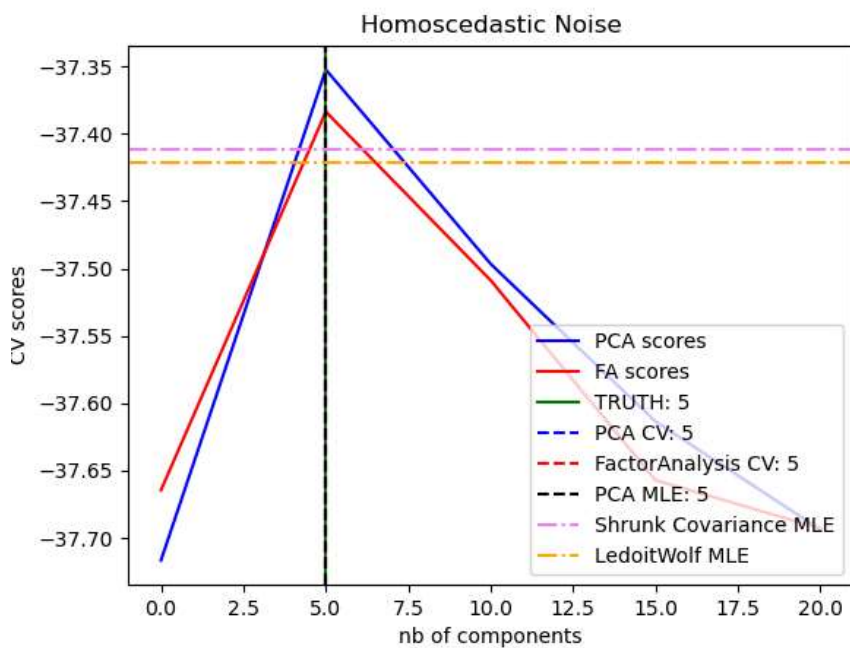
PCA is used to decompose a multivariate dataset in a set of successive orthogonal components that explain a maximum amount of the variance. In scikit-learn, [PCA](#) is implemented as a *transformer* object that learns  $n$  components in its `fit` method, and can be used on new data to project it on these components.

PCA centers but does not scale the input data for each feature before applying the SVD. The optional parameter `whiten=True` makes it possible to project the data onto the singular space without scaling each component to unit variance. This is often useful if the models down-stream make strong assumptions on the isotropy of the signal: this is for example the case for Support Vector Machines with the RBF kernel and the K-Means clustering algorithm.

Below is an example of the iris dataset, which is comprised of 4 features, projected on the 2 dimensions that explain most variance:



The [PCA](#) object also provides a probabilistic interpretation of the PCA that can give a likelihood of data based on the amount of variance it explains. As such it implements a [score](#) method that can be used in cross-validation:



## Examples

- [PCA example with Iris Data-set](#)

- [Comparison of LDA and PCA 2D projection of Iris dataset](#)
- [Model selection with Probabilistic PCA and Factor Analysis \(FA\)](#)

## 2.5.1.2. Incremental PCA

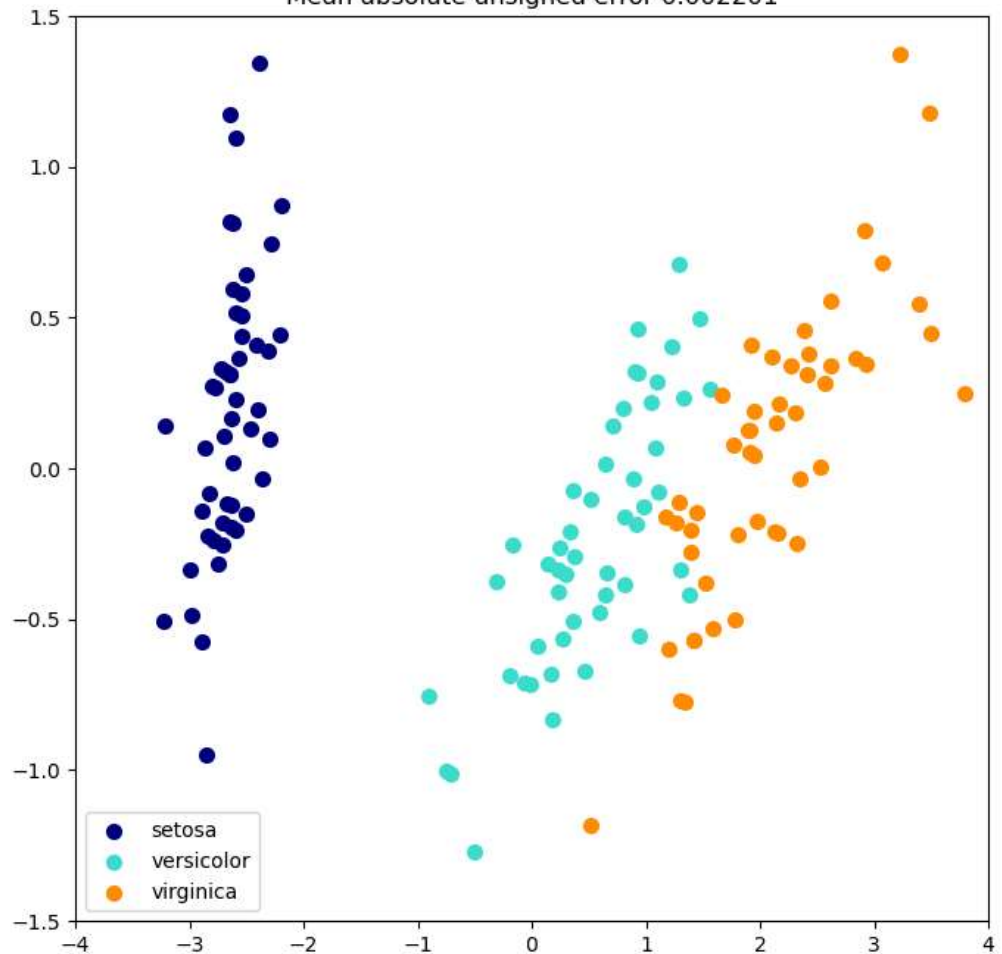
The `PCA` object is very useful, but has certain limitations for large datasets. The biggest limitation that `PCA` only supports batch processing, which means all of the data to be processed must fit in main memory. The `IncrementalPCA` object uses a different form of processing and allows for partial computations which almost exactly match the results of `PCA` while processing the data in a minibatch fashion. `IncrementalPCA` makes it possible to implement out-of-core Principal Component Analysis either by:

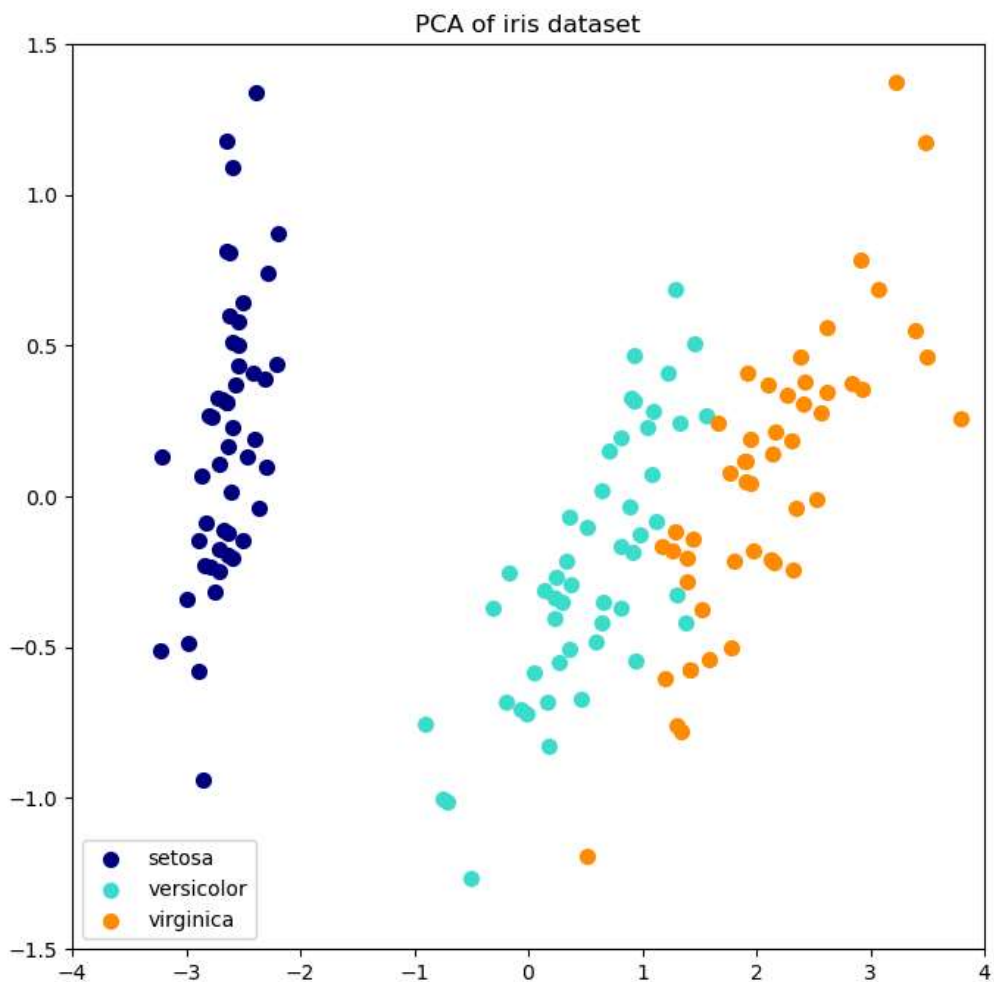
- Using its `partial_fit` method on chunks of data fetched sequentially from the local hard drive or a network database.
- Calling its `fit` method on a memory mapped file using `numpy.memmap`.

`IncrementalPCA` only stores estimates of component and noise variances, in order to update `explained_variance_ratio_` incrementally. This is why memory usage depends on the number of samples per batch, rather than the number of samples to be processed in the dataset.

As in `PCA`, `IncrementalPCA` centers but does not scale the input data for each feature before applying the SVD.

Incremental PCA of iris dataset  
Mean absolute unsigned error 0.002201





## Examples

- [Incremental PCA](#)

### 2.5.1.3. PCA using randomized SVD

It is often interesting to project data to a lower-dimensional space that preserves most of the variance, by dropping the singular vector of components associated with lower singular values.

For instance, if we work with 64x64 pixel gray-level pictures for face recognition, the dimensionality of the data is 4096 and it is slow to train an RBF support vector machine on such wide data. Furthermore we know that the intrinsic dimensionality of the data is much lower than 4096 since a

pictures of human faces look somewhat alike. The samples lie on a manifold of much lower dimension (say around 200 for instance). The PCA algorithm can be used to linearly transform the data while both reducing the dimensionality and preserve most of the explained variance at the same time.

The class `PCA` used with the optional parameter `svd_solver='randomized'` is very useful in that case: since we are going to drop most of the singular vectors it is much more efficient to limit the computation to an approximated estimate of the singular vectors we will keep to actually perform the transform.

For instance, the following shows 16 sample portraits (centered around 0.0) from the Olivetti dataset. On the right hand side are the first 16 singular vectors reshaped as portraits. Since we only require the top 16 singular vectors of a dataset with size  $n_{samples} = 400$  and  $n_{features} = 64 \times 64 = 4096$ , the computation time is less than 1s:



If we note  $n_{\max} = \max(n_{\text{samples}}, n_{\text{features}})$  and  $n_{\min} = \min(n_{\text{samples}}, n_{\text{features}})$ , the time complexity of the randomized [PCA](#) is  $O(n_{\max}^2 \cdot n_{\text{components}})$  instead of  $O(n_{\max}^2 \cdot n_{\min})$  for the exact method implemented in [PCA](#).

The memory footprint of randomized [PCA](#) is also proportional to  $2 \cdot n_{\max} \cdot n_{\text{components}}$  instead of  $n_{\max} \cdot n_{\min}$  for the exact method.

Note: the implementation of `inverse_transform` in [PCA](#) with `svd_solver='randomized'` is not the exact inverse transform of `transform` even when `whiten=False` (default).

## Examples

---

- [Faces recognition example using eigenfaces and SVMs](#)
- [Faces dataset decompositions](#)

## References

---

- Algorithm 4.3 in "[Finding structure with randomness: Stochastic algorithms for constructing approximate matrix decompositions](#)" Halko, et al., 2009
- "[An implementation of a randomized algorithm for principal component analysis](#)" A. Szlam et al., 2014

## 2.5.1.4. Sparse principal components analysis (SparsePCA and MiniBatchSparsePCA)

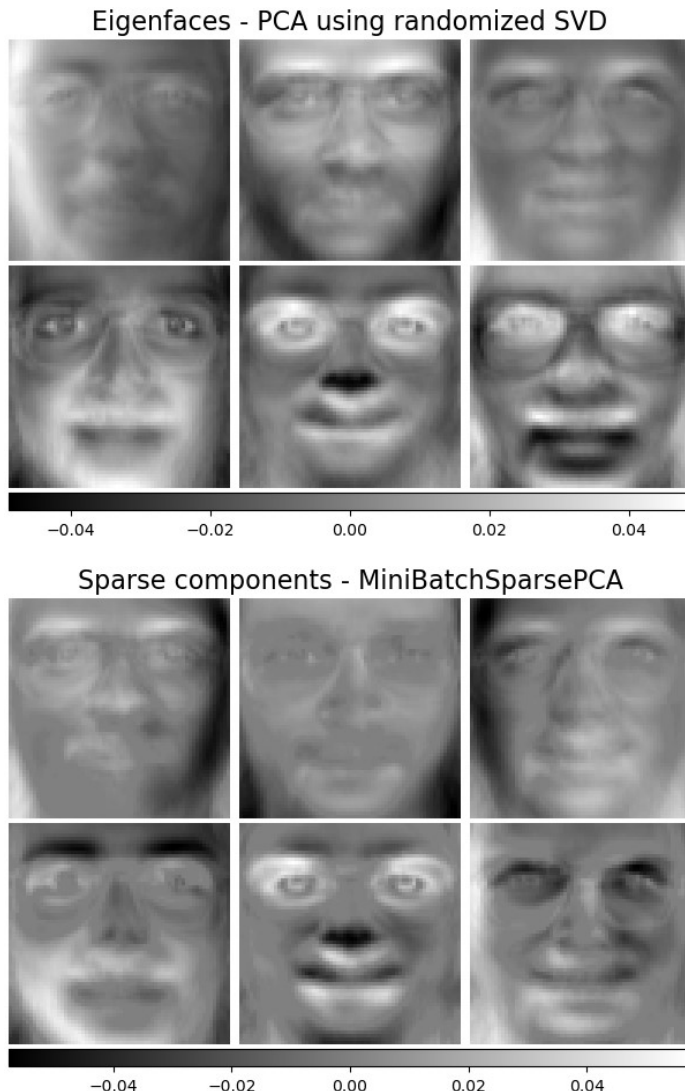
[SparsePCA](#) is a variant of PCA, with the goal of extracting the set of sparse components that best reconstruct the data.

Mini-batch sparse PCA ([MiniBatchSparsePCA](#)) is a variant of [SparsePCA](#) that is faster but less accurate. The increased speed is reached by iterating over small chunks of the set of features, for a given number of iterations.

Principal component analysis ([PCA](#)) has the disadvantage that the components extracted by this method have exclusively dense expressions, i.e. they have non-zero coefficients when expressed as linear combinations of the original variables. This can make interpretation difficult. In many cases, the real underlying components can be more naturally imagined as sparse vectors; for example in face recognition, components might naturally map to parts of faces.

Sparse principal components yields a more parsimonious, interpretable representation, clearly emphasizing which of the original features contribute to the differences between samples.

The following example illustrates 16 components extracted using sparse PCA from the Olivetti face dataset. It can be seen how the regularization term induces many zeros. Furthermore, the natural structure of the data causes the non-zero coefficients to be vertically adjacent. The model does not enforce this mathematically: each component is a vector  $h \in \mathbf{R}^{4096}$ , and there is no notion of vertical adjacency except during the human-friendly visualization as 64x64 pixel images. The fact that the components shown below appear local is the effect of the inherent structure of the data, which makes such local patterns minimize reconstruction error. There exist sparsity-inducing norms that take into account adjacency and different kinds of structure; see [\[Jen09\]](#) for a review of such methods. For more details on how to use Sparse PCA, see the Examples section, below.



Note that there are many different formulations for the Sparse PCA problem. The one implemented here is based on [\[Mrl09\]](#). The optimization problem solved is a PCA problem (dictionary learning)

with an  $\ell_1$  penalty on the components:

$$(U^*, V^*) = \arg \min_{U, V} \frac{1}{2} \|X - UV\|_{\text{Fro}}^2 + \alpha \|V\|_{1,1}$$

subject to  $\|U_k\|_2 \leq 1$  for all  $0 \leq k < n_{\text{components}}$

$\|\cdot\|_{\text{Fro}}$  stands for the Frobenius norm and  $\|\cdot\|_{1,1}$  stands for the entry-wise matrix norm which is the sum of the absolute values of all the entries in the matrix. The sparsity-inducing  $\|\cdot\|_{1,1}$  matrix norm also prevents learning components from noise when few training samples are available. The degree of penalization (and thus sparsity) can be adjusted through the hyperparameter `alpha`. Small values lead to a gently regularized factorization, while larger values shrink many coefficients to zero.

### Note

While in the spirit of an online algorithm, the class `MiniBatchSparsePCA` does not implement `partial_fit` because the algorithm is online along the features direction, not the samples direction.

## Examples

- [Faces dataset decompositions](#)

## References

- [Mrl09] [“Online Dictionary Learning for Sparse Coding”](#) J. Mairal, F. Bach, J. Ponce, G. Sapiro, 2009  
[Jen09] [“Structured Sparse Principal Component Analysis”](#) R. Jenatton, G. Obozinski, F. Bach, 2009

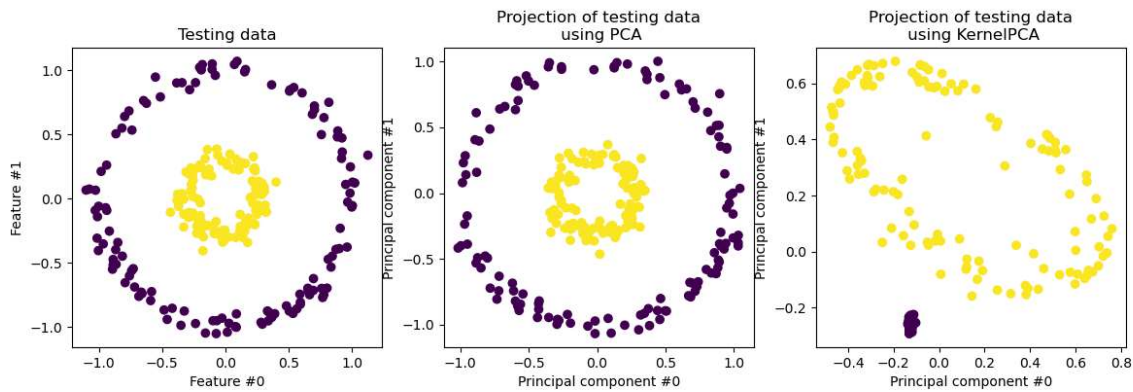
## 2.5.2. Kernel Principal Component Analysis (kPCA)

### 2.5.2.1. Exact Kernel PCA

[KernelPCA](#) is an extension of PCA which achieves non-linear dimensionality reduction through the use of kernels (see [Pairwise metrics, Affinities and Kernels](#)) [\[Scholkopf1997\]](#). It has many applicatio

including denoising, compression and structured prediction (kernel dependency estimation).

`KernelPCA` supports both `transform` and `inverse_transform`.



### Note

`KernelPCA.inverse_transform` relies on a kernel ridge to learn the function mapping samples from the PCA basis into the original feature space [Bakir2003]. Thus, the reconstruction obtained with `KernelPCA.inverse_transform` is an approximation. See the example linked below for more details.

### Examples

- [Kernel PCA](#)
- [Image denoising using kernel PCA](#)

### References

[Schölkopf1997] Schölkopf, Bernhard, Alexander Smola, and Klaus-Robert Müller. "[Kernel principal component analysis.](#)" International conference on artificial neural networks. Springer, Berlin Heidelberg, 1997.

[Bakir2003] Bakır, Gökhan H., Jason Weston, and Bernhard Schölkopf. "[Learning to find pre-image](#)" Advances in neural information processing systems 16 (2003): 449-456.

## 2.5.2.2. Choice of solver for Kernel PCA

While in `PCA` the number of components is bounded by the number of features, in `KernelPCA` the number of components is bounded by the number of samples. Many real-world datasets have large

number of samples! In these cases finding *all* the components with a full kPCA is a waste of computation time, as data is mostly described by the first few components (e.g. `n_components<=100`). In other words, the centered Gram matrix that is eigendecomposed in the Kernel PCA fitting process has an effective rank that is much smaller than its size. This is a situation where approximate eigensolvers can provide speedup with very low precision loss.

## Eigensolvers

### 2.5.3. Truncated singular value decomposition and latent semantic analysis

[TruncatedSVD](#) implements a variant of singular value decomposition (SVD) that only computes the largest singular values, where  $k$  is a user-specified parameter.

[TruncatedSVD](#) is very similar to [PCA](#), but differs in that the matrix  $X$  does not need to be centered. When the columnwise (per-feature) means of  $X$  are subtracted from the feature values, truncated SVD on the resulting matrix is equivalent to PCA.

## About truncated SVD and latent semantic analysis (LSA)

### Examples

- [Clustering text documents using k-means](#)

### References

- Christopher D. Manning, Prabhakar Raghavan and Hinrich Schütze (2008), *Introduction to Information Retrieval*, Cambridge University Press, chapter 18: [Matrix decompositions & latent semantic indexing](#)

### 2.5.4. Dictionary Learning

#### 2.5.4.1. Sparse coding with a precomputed dictionary

The [SparseCoder](#) object is an estimator that can be used to transform signals into sparse linear combination of atoms from a fixed, precomputed dictionary such as a discrete wavelet basis. This object therefore does not implement a `fit` method. The transformation amounts to a sparse coding problem: finding a representation of the data as a linear combination of as few dictionary atoms as possible. All variations of dictionary learning implement the following transform method controllable via the `transform_method` initialization parameter:

- Orthogonal matching pursuit ([Orthogonal Matching Pursuit \(OMP\)](#))
- Least-angle regression ([Least Angle Regression](#))
- Lasso computed by least-angle regression
- Lasso using coordinate descent ([Lasso](#))
- Thresholding

Thresholding is very fast but it does not yield accurate reconstructions. They have been shown useful in literature for classification tasks. For image reconstruction tasks, orthogonal matching pursuit yields the most accurate, unbiased reconstruction.

The dictionary learning objects offer, via the `split_code` parameter, the possibility to separate the positive and negative values in the results of sparse coding. This is useful when dictionary learning is used for extracting features that will be used for supervised learning, because it allows the learning algorithm to assign different weights to negative loadings of a particular atom, from to the corresponding positive loading.

The split code for a single sample has length `2 * n_components` and is constructed using the following rule: First, the regular code of length `n_components` is computed. Then, the first `n_components` entries of the `split_code` are filled with the positive part of the regular code vector. The second half of the split code is filled with the negative part of the code vector, only with a positive sign. Therefore, the `split_code` is non-negative.

## Examples

---

- [Sparse coding with a precomputed dictionary](#)

### 2.5.4.2. Generic dictionary learning

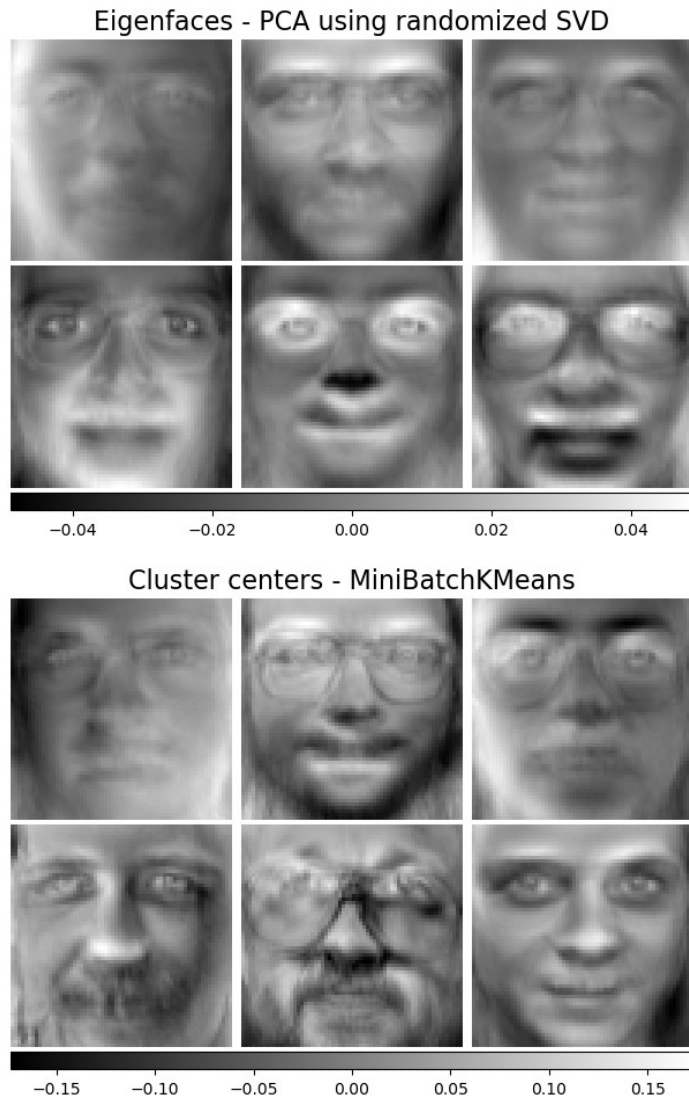
Dictionary learning ([DictionaryLearning](#)) is a matrix factorization problem that amounts to finding a (usually overcomplete) dictionary that will perform well at sparsely encoding the fitted data.

Representing data as sparse combinations of atoms from an overcomplete dictionary is suggested to be the way the mammalian primary visual cortex works. Consequently, dictionary learning application on image patches has been shown to give good results in image processing tasks such as image completion, inpainting and denoising, as well as for supervised recognition tasks.

Dictionary learning is an optimization problem solved by alternatively updating the sparse code, a solution to multiple Lasso problems, considering the dictionary fixed, and then updating the dictionary to best fit the sparse code.

$$(U^*, V^*) = \arg \min_{U, V} \frac{1}{2} \|X - UV\|_{\text{Fro}}^2 + \alpha \|U\|_{1,1}$$

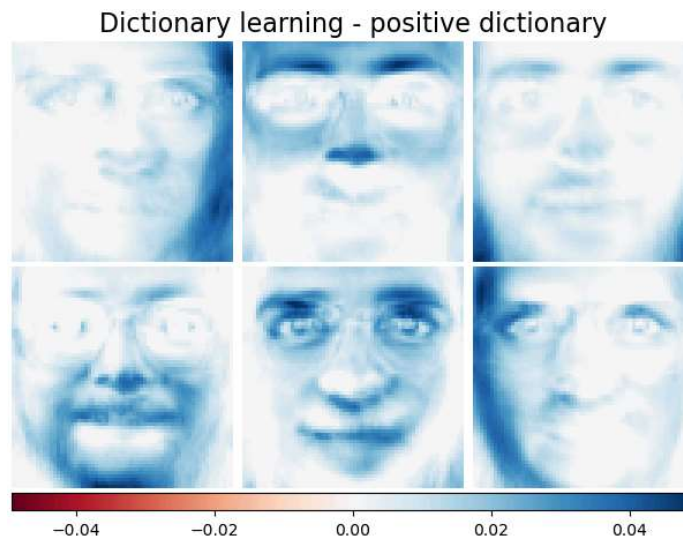
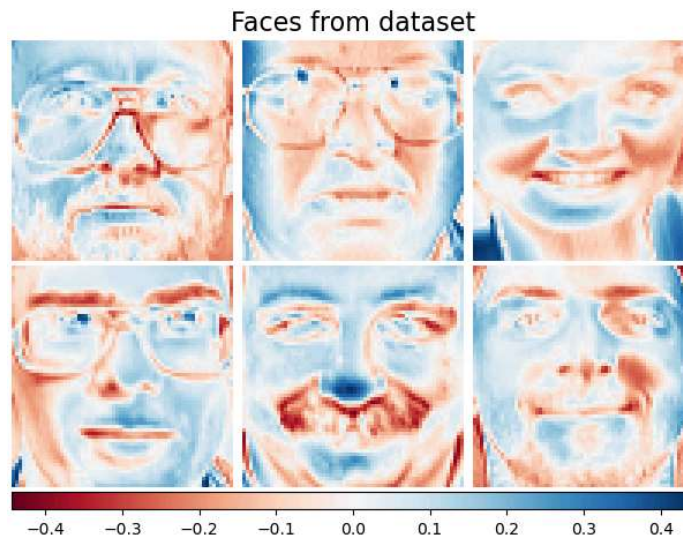
subject to  $\|V_k\|_2 \leq 1$  for all  $0 \leq k < n_{\text{atoms}}$



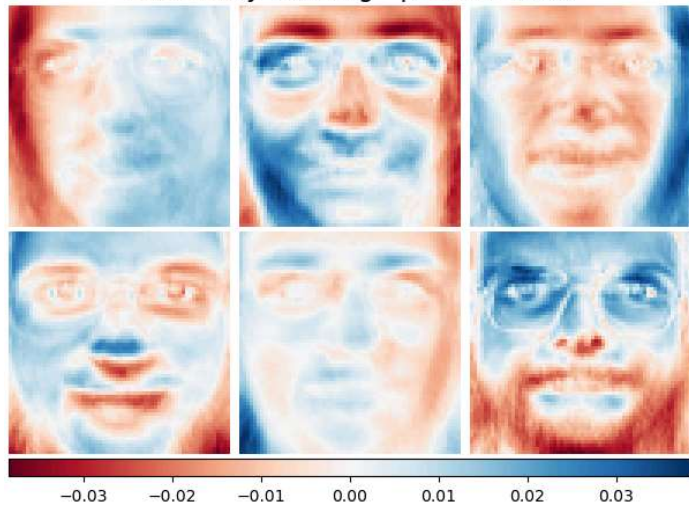
$\|\cdot\|_{\text{Fro}}$  stands for the Frobenius norm and  $\|\cdot\|_{1,1}$  stands for the entry-wise matrix norm which is the sum of the absolute values of all the entries in the matrix. After using such a procedure to fit the

dictionary, the transform is simply a sparse coding step that shares the same implementation with dictionary learning objects (see [Sparse coding with a precomputed dictionary](#)).

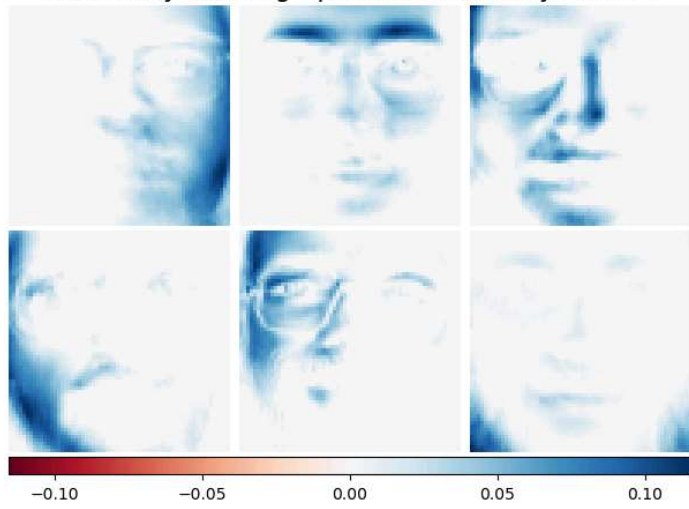
It is also possible to constrain the dictionary and/or code to be positive to match constraints that may be present in the data. Below are the faces with different positivity constraints applied. Red indicates negative values, blue indicates positive values, and white represents zeros.



Dictionary learning - positive code

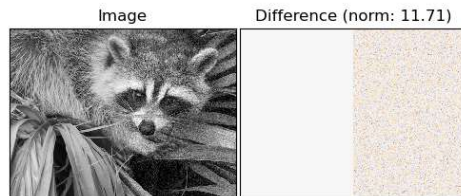


Dictionary learning - positive dictionary & code



The following image shows how a dictionary learned from 4x4 pixel image patches extracted from part of the image of a raccoon face looks like.

Distorted image



## Examples

- [Image denoising using dictionary learning](#)

## References

- “[Online dictionary learning for sparse coding](#)” J. Mairal, F. Bach, J. Ponce, G. Sapiro, 2009

### 2.5.4.3. Mini-batch dictionary learning

[\*\*MiniBatchDictionaryLearning\*\*](#) implements a faster, but less accurate version of the dictionary learning algorithm that is better suited for large datasets.

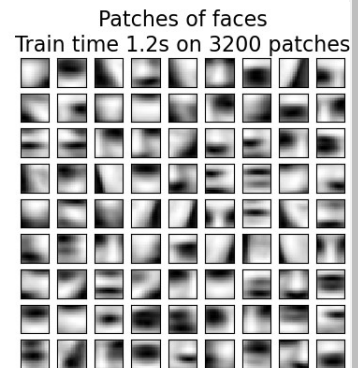
By default, [\*\*MiniBatchDictionaryLearning\*\*](#) divides the data into mini-batches and optimizes in an online manner by cycling over the mini-batches for the specified number of iterations. However, at the moment it does not implement a stopping condition.

The estimator also implements `partial_fit`, which updates the dictionary by iterating only once over a mini-batch. This can be used for online learning when the data is not readily available from the start, or for when the data does not fit into the memory.

#### Clustering for dictionary learning

Note that when using dictionary learning to extract a representation (e.g. for sparse coding) clustering can be a good proxy to learn the dictionary. For instance the [\*\*MiniBatchKMeans\*\*](#) estimator is computationally efficient and implements on-line learning with a `partial_fit` method.

Example: [Online learning of a dictionary of parts of faces](#)



### 2.5.5. Factor Analysis

In unsupervised learning we only have a dataset  $X = \{x_1, x_2, \dots, x_n\}$ . How can this dataset be described mathematically? A very simple `continuous latent variable` model for  $X$  is

$$x_i = Wh_i + \mu + \epsilon$$

The vector  $h_i$  is called “latent” because it is unobserved.  $\epsilon$  is considered a noise term distributed according to a Gaussian with mean 0 and covariance  $\Psi$  (i.e.  $\epsilon \sim \mathcal{N}(0, \Psi)$ ),  $\mu$  is some arbitrary offset.

vector. Such a model is called “generative” as it describes how  $x_i$  is generated from  $h_i$ . If we use all the  $x_i$ ’s as columns to form a matrix  $\mathbf{X}$  and all the  $h_i$ ’s as columns of a matrix  $\mathbf{H}$  then we can write (with suitably defined  $\mathbf{M}$  and  $\mathbf{E}$ ):

$$\mathbf{X} = W\mathbf{H} + \mathbf{M} + \mathbf{E}$$

In other words, we *decomposed* matrix  $\mathbf{X}$ .

If  $h_i$  is given, the above equation automatically implies the following probabilistic interpretation:

$$p(x_i|h_i) = \mathcal{N}(Wh_i + \mu, \Psi)$$

For a complete probabilistic model we also need a prior distribution for the latent variable  $h$ . The most straightforward assumption (based on the nice properties of the Gaussian distribution) is  $h \sim \mathcal{N}(0, \mathbf{I})$ . This yields a Gaussian as the marginal distribution of  $x$ :

$$p(x) = \mathcal{N}(\mu, WW^T + \Psi)$$

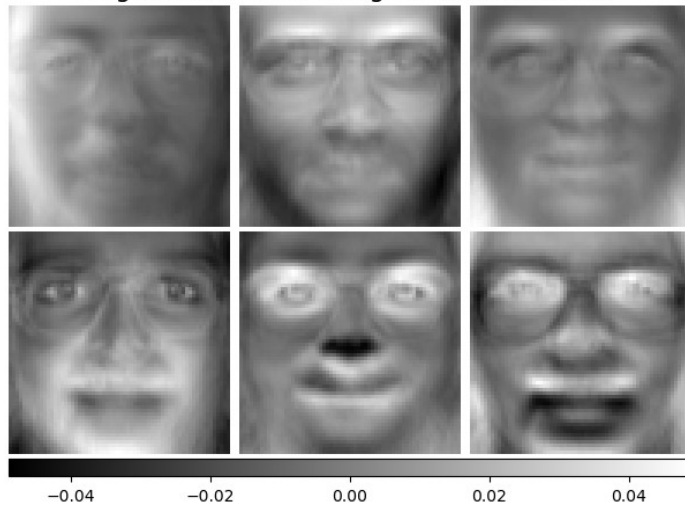
Now, without any further assumptions the idea of having a latent variable  $h$  would be superfluous:  $x$  can be completely modelled with a mean and a covariance. We need to impose some more specific structure on one of these two parameters. A simple additional assumption regards the structure of the error covariance  $\Psi$ :

- $\Psi = \sigma^2 \mathbf{I}$ : This assumption leads to the probabilistic model of [PCA](#).
- $\Psi = \text{diag}(\psi_1, \psi_2, \dots, \psi_n)$ : This model is called [FactorAnalysis](#), a classical statistical model. The matrix  $W$  is sometimes called the “factor loading matrix”.

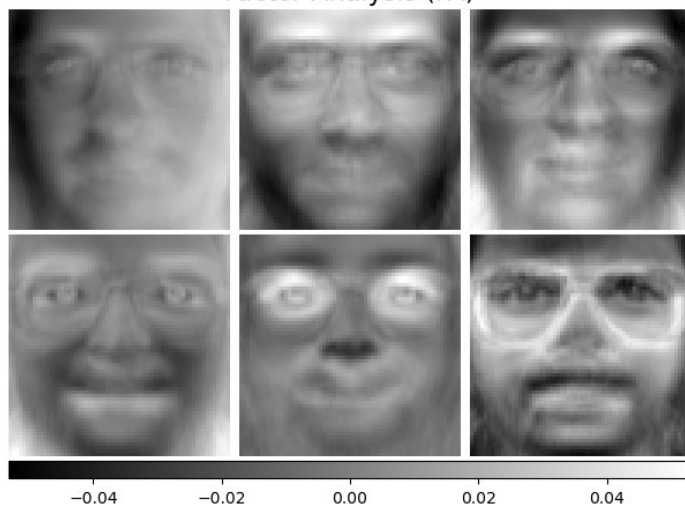
Both models essentially estimate a Gaussian with a low-rank covariance matrix. Because both models are probabilistic they can be integrated in more complex models, e.g. Mixture of Factor Analysers. One gets very different models (e.g. [FastICA](#)) if non-Gaussian priors on the latent variables are assumed.

Factor analysis *can* produce similar components (the columns of its loading matrix) to [PCA](#). However, one can not make any general statements about these components (e.g. whether they are orthogonal):

Eigenfaces - PCA using randomized SVD

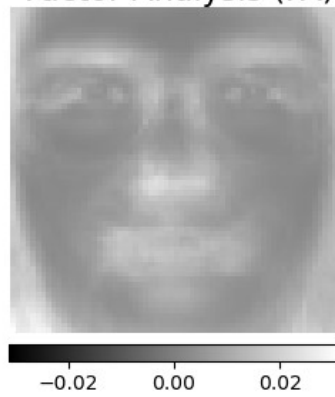


Factor Analysis (FA)

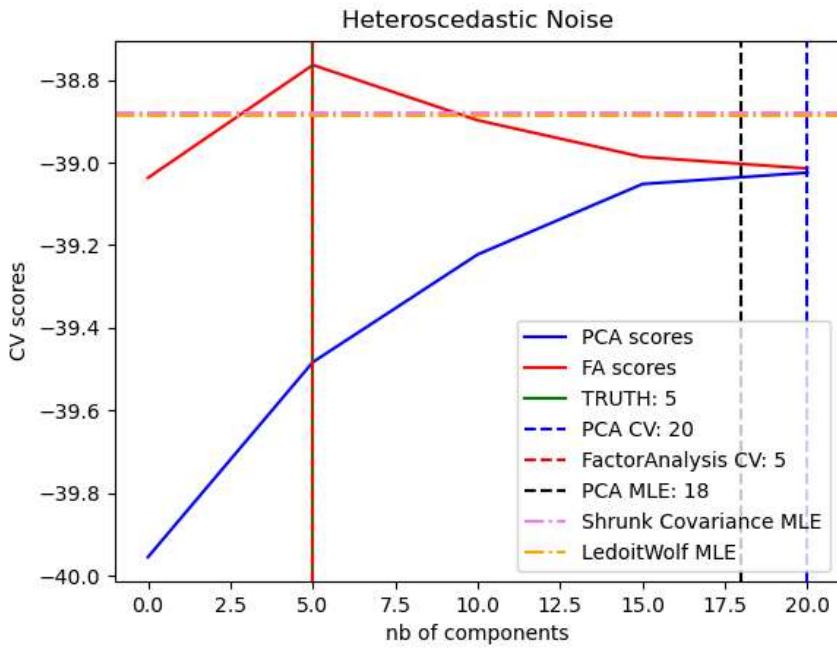


The main advantage for Factor Analysis over [PCA](#) is that it can model the variance in every direction of the input space independently (heteroscedastic noise):

Pixelwise variance from Factor Analysis (FA)



This allows better model selection than probabilistic PCA in the presence of heteroscedastic noise



Factor Analysis is often followed by a rotation of the factors (with the parameter `rotation`), usual to improve interpretability. For example, Varimax rotation maximizes the sum of the variances of the squared loadings, i.e., it tends to produce sparser factors, which are influenced by only a few features each (the “simple structure”). See e.g., the first example below.

## Examples

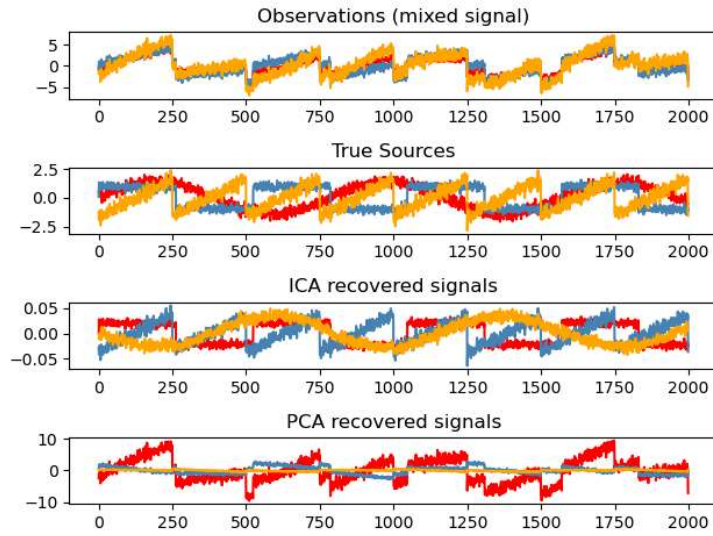
---

- [Factor Analysis \(with rotation\) to visualize patterns](#)
- [Model selection with Probabilistic PCA and Factor Analysis \(FA\)](#)

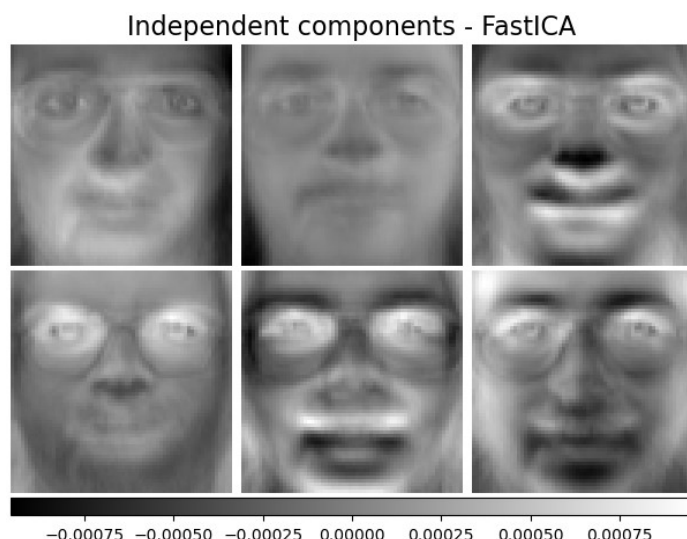
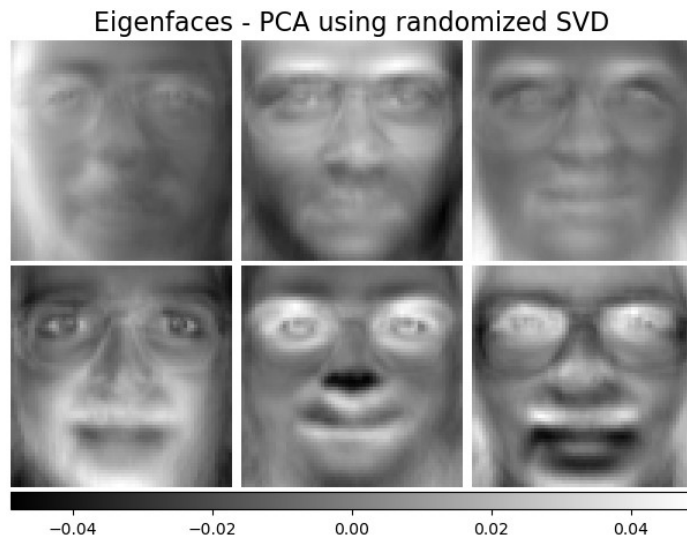
## 2.5.6. Independent component analysis (ICA)

Independent component analysis separates a multivariate signal into additive subcomponents that are maximally independent. It is implemented in scikit-learn using the `Fast ICA` algorithm. Typically ICA is not used for reducing dimensionality but for separating superimposed signals. Since the ICA model does not include a noise term, for the model to be correct, whitening must be applied. This can be done internally using the `whiten` argument or manually using one of the PCA variants.

It is classically used to separate mixed signals (a problem known as *blind source separation*), as in the example below:



ICA can also be used as yet another non linear decomposition that finds components with some sparsity:



## Examples

- [Blind source separation using FastICA](#)
- [FastICA on 2D point clouds](#)
- [Faces dataset decompositions](#)

## 2.5.7. Non-negative matrix factorization (NMF and NNMF)

### 2.5.7.1. NMF with the Frobenius norm

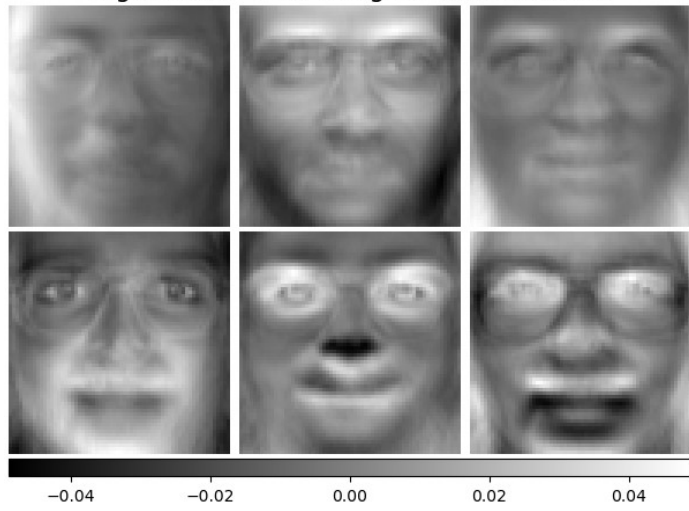
[NMF](#) [1] is an alternative approach to decomposition that assumes that the data and the components are non-negative. [NMF](#) can be plugged in instead of [PCA](#) or its variants, in the cases where the data matrix does not contain negative values. It finds a decomposition of samples  $X$  into two matrices  $W$  and  $H$  of non-negative elements, by optimizing the distance  $d$  between  $X$  and the matrix product  $WH$ . The most widely used distance function is the squared Frobenius norm, which is an obvious extension of the Euclidean norm to matrices:

$$d_{\text{Fro}}(X, Y) = \frac{1}{2} \|X - Y\|_{\text{Fro}}^2 = \frac{1}{2} \sum_{i,j} (X_{ij} - Y_{ij})^2$$

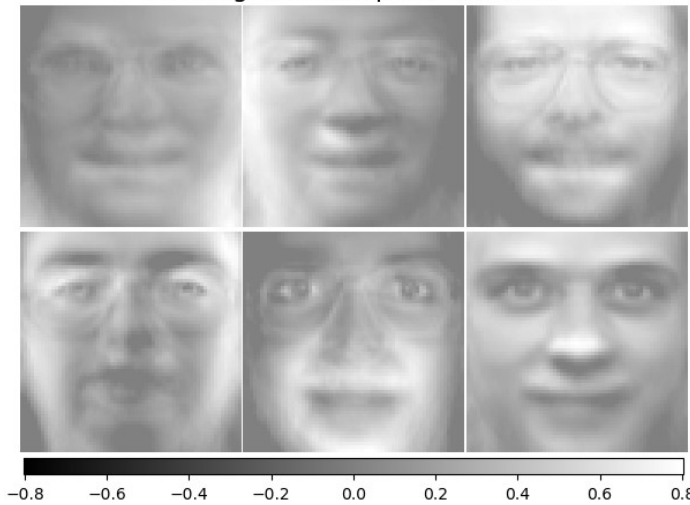
Unlike [PCA](#), the representation of a vector is obtained in an additive fashion, by superimposing the components, without subtracting. Such additive models are efficient for representing images and text.

It has been observed in [Hoyer, 2004] [2] that, when carefully constrained, [NMF](#) can produce a part based representation of the dataset, resulting in interpretable models. The following example displays 16 sparse components found by [NMF](#) from the images in the Olivetti faces dataset, in comparison with the PCA eigenfaces.

Eigenfaces - PCA using randomized SVD



Non-negative components - NMF



The `init` attribute determines the initialization method applied, which has a great impact on the performance of the method. [NMF](#) implements the method Nonnegative Double Singular Value Decomposition. NNDSVD [4] is based on two SVD processes, one approximating the data matrix, the other approximating positive sections of the resulting partial SVD factors utilizing an algebraic property of unit rank matrices. The basic NNDSVD algorithm is better fit for sparse factorization. Its variants NNDSVDA (in which all zeros are set equal to the mean of all elements of the data), and NNDSVDAR (in which the zeros are set to random perturbations less than the mean of the data divided by 100) are recommended in the dense case.

Note that the Multiplicative Update ('mu') solver cannot update zeros present in the initialization, it leads to poorer results when used jointly with the basic NNDSVD algorithm which introduces a lot of zeros; in this case, NNDSVDA or NNDSVDAR should be preferred.

[NMF](#) can also be initialized with correctly scaled random non-negative matrices by setting `init="random"`. An integer seed or a `RandomState` can also be passed to `random_state` to control

reproducibility.

In [NMF](#), L1 and L2 priors can be added to the loss function in order to regularize the model. The L2 prior uses the Frobenius norm, while the L1 prior uses an elementwise L1 norm. As in [ElasticNet](#), we control the combination of L1 and L2 with the `l1_ratio` ( $\rho$ ) parameter, and the intensity of the regularization with the `alpha_W` and `alpha_H` ( $\alpha_W$  and  $\alpha_H$ ) parameters. The priors are scaled by the number of samples (`n_samples`) for `H` and the number of features (`n_features`) for `W` to keep their impact balanced with respect to one another and to the data fit term as independent as possible of the size of the training set. Then the priors terms are:

$$(\alpha_W \rho \|W\|_1 + \frac{\alpha_W(1 - \rho)}{2} \|W\|_{\text{Fro}}^2) * n\_features + (\alpha_H \rho \|H\|_1 + \frac{\alpha_H(1 - \rho)}{2} \|H\|_{\text{Fro}}^2) * n\_samples$$

and the regularized objective function is:

$$d_{\text{Fro}}(X, WH) + (\alpha_W \rho \|W\|_1 + \frac{\alpha_W(1 - \rho)}{2} \|W\|_{\text{Fro}}^2) * n\_features + (\alpha_H \rho \|H\|_1 + \frac{\alpha_H(1 - \rho)}{2} \|H\|_{\text{Fro}}^2) * n\_samples$$

### 2.5.7.2. NMF with a beta-divergence

As described previously, the most widely used distance function is the squared Frobenius norm, which is an obvious extension of the Euclidean norm to matrices:

$$d_{\text{Fro}}(X, Y) = \frac{1}{2} \|X - Y\|_{\text{Fro}}^2 = \frac{1}{2} \sum_{i,j} (X_{ij} - Y_{ij})^2$$

Other distance functions can be used in NMF as, for example, the (generalized) Kullback-Leibler (KL) divergence, also referred as I-divergence:

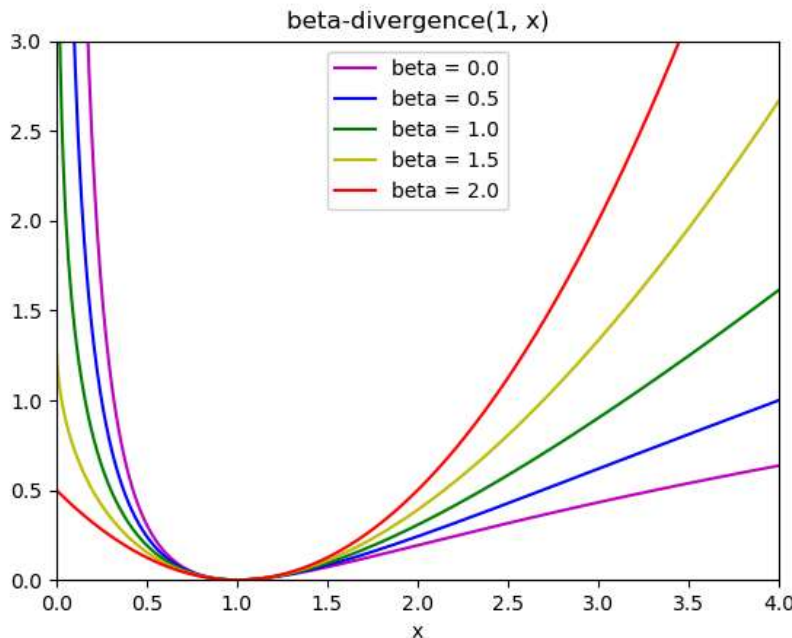
$$d_{KL}(X, Y) = \sum_{i,j} (X_{ij} \log(\frac{X_{ij}}{Y_{ij}}) - X_{ij} + Y_{ij})$$

Or, the Itakura-Saito (IS) divergence:

$$d_{IS}(X, Y) = \sum_{i,j} \left( \frac{X_{ij}}{Y_{ij}} - \log\left(\frac{X_{ij}}{Y_{ij}}\right) - 1 \right)$$

These three distances are special cases of the beta-divergence family, with  $\beta = 2, 1, 0$  respectively [6]. The beta-divergence are defined by :

$$d_\beta(X, Y) = \sum_{i,j} \frac{1}{\beta(\beta-1)} (X_{ij}^\beta + (\beta-1)Y_{ij}^\beta - \beta X_{ij} Y_{ij}^{\beta-1})$$



Note that this definition is not valid if  $\beta \in (0; 1)$ , yet it can be continuously extended to the definitions of  $d_{KL}$  and  $d_{IS}$  respectively.

## NMF implemented solvers

NMF is best used with the `fit_transform` method, which returns the matrix W. The matrix H is stored into the fitted model in the `components_` attribute; the method `transform` will decompose new matrix X\_new based on these stored components:

```
>>> import numpy as np
>>> X = np.array([[1, 1], [2, 1], [3, 1.2], [4, 1], [5, 0.8], [6, 1]])
>>> from sklearn.decomposition import NMF
>>> model = NMF(n_components=2, init='random', random_state=0)
>>> W = model.fit_transform(X)
>>> H = model.components_
>>> X_new = np.array([[1, 0], [1, 6.1], [1, 0], [1, 4], [3.2, 1], [0, 4]])
>>> W_new = model.transform(X_new)
```

## Examples

- [Faces dataset decompositions](#)
- [Topic extraction with Non-negative Matrix Factorization and Latent Dirichlet Allocation](#)

### 2.5.7.3. Mini-batch Non Negative Matrix Factorization

[`MiniBatchNMF`](#) [7] implements a faster, but less accurate version of the non negative matrix factorization (i.e. [`NMF`](#)), better suited for large datasets.

By default, [`MiniBatchNMF`](#) divides the data into mini-batches and optimizes the NMF model in an online manner by cycling over the mini-batches for the specified number of iterations. The `batch_size` parameter controls the size of the batches.

In order to speed up the mini-batch algorithm it is also possible to scale past batches, giving them less importance than newer batches. This is done introducing a so-called forgetting factor control by the `forget_factor` parameter.

The estimator also implements `partial_fit`, which updates `H` by iterating only once over a mini-batch. This can be used for online learning when the data is not readily available from the start, or when the data does not fit into memory.

## References

- [1] ["Learning the parts of objects by non-negative matrix factorization"](#) D. Lee, S. Seung, 1999
- [2] ["Non-negative Matrix Factorization with Sparseness Constraints"](#) P. Hoyer, 2004
- [4] ["SVD based initialization: A head start for nonnegative matrix factorization"](#) C. Boutsidis, E. Gallopolous, 2008

