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Cheat Sheet: Building Unsupervised Learning Models

Unsupervised learning models

| Model Name | Brief Description | Code Syntax |
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| UMAP | UMAP (Uniform Manifold Approximation and Projection) is used for dimensionality reduction. Pros: High performance, preserves global structure. Cons: Sensitive to parameters. Applications: Data visualization, feature extraction. Key hyperparameters: • n_neighbors: Controls the local neighborhood size (default = 15). • min_dist: Controls the minimum distance between points in the embedded space (default = 0.1). • n_components: The dimensionality of the embedding (default = 2). | <pre>from umap.umap_ import UMAP umap = UMAP(n_neighbors=15, min_dist=0.1, n_components=2)</pre> |
| t-SNE | t-SNE (t-Distributed Stochastic Neighbor Embedding) is a nonlinear dimensionality reduction technique. Pros: Good for visualizing high-dimensional data. Cons: Computationally expensive, prone to overfitting. Applications: Data visualization, anomaly detection. Key hyperparameters: • n_components: The number of dimensions for the output (default = 2). • perplexity: Balances attention between local and global aspects of the data (default = 30). • learning_rate: Controls the step size during optimization (default = 200). | <pre>from sklearn.manifold import TSNE tsne = TSNE(n_components=2, perplexity=30, learning_rate=200)</pre> |
| PCA | PCA (principal component analysis) is used for linear dimensionality reduction. Pros: Easy to interpret, reduces noise. Cons: Linear, may lose information in nonlinear data. Applications: Feature extraction, compression. Key hyperparameters: • n_components: Number of principal components to retain (default = 2). • whiten: Whether to scale the components (default = False). • svd_solver: The algorithm to compute the components (default = 'auto'). | <pre>from sklearn.decomposition import PCA pca = PCA(n_components=2)</pre> |
| DBSCAN | DBSCAN (Density-Based Spatial Clustering of Applications with Noise) is a density-based clustering algorithm. Pros: Identifies outliers, does not require the number of clusters. Cons: Difficult with varying density clusters. Applications: Anomaly detection, spatial data clustering. Key hyperparameters: • eps: The maximum distance between two points to be considered neighbors (default = 0.5). • min_samples: Minimum number of samples in a neighborhood to form a cluster (default = 5). | from sklearn.cluster import DBSCAN dbscan = DBSCAN(eps=0.5, min_samples=5) |
| HDBSCAN | HDBSCAN (Hierarchical DBSCAN) improves on DBSCAN by handling varying density clusters. Pros: Better handling of varying densities. Cons: Can be slower than DBSCAN. Applications: Large datasets, complex clustering problems. Key hyperparameters: • min_cluster_size: The minimum size of clusters (default = 5). • min_samples: Minimum number of samples to form a cluster (default = 10). | import hdbscan clusterer = hdbscan.HDBSCAN(min_cluster_size=5) |
| K-Means clustering | K-Means is a centroid-based clustering algorithm that groups data into k clusters. Pros: Efficient, simple to implement. Cons: Sensitive to initial cluster centroids. Applications: Customer segmentation, pattern recognition. Key hyperparameters: • n_clusters: Number of clusters (default = 8). • init: Method for initializing the centroids ('k-means++' or | <pre>from sklearn.cluster import KMeans kmeans = KMeans(n_clusters=3)</pre> |

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| Model Name | Brief Description | Code Syntax |
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| | • n_init: Number of times the algorithm will run with different centroid seeds (default = 10). | |

Associated fuctions used

| Method | Brief Description | Code Syntax |
|-------------------------------|---------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------|
| make_blobs | Generates isotropic Gaussian blobs for clustering. | from sklearn.datasets import make_blobs X, y = make_blobs(n_samples=100, centers=2, random_state=42) |
| multivariate_normal | Generates samples from a multivariate normal distribution. | from numpy.random import multivariate_normal samples = multivariate_normal(mean=[0, 0], cov=[[1, 0], [0, 1]], size=100) |
| plotly.express.scatter_3d | Creates a 3D scatter plot using Plotly Express. | <pre>import plotly.express as px fig = px.scatter_3d(df, x='x', y='y', z='z') fig.show()</pre> |
| geopandas.GeoDataFrame | Creates a GeoDataFrame from a Pandas DataFrame. | <pre>import geopandas as gpd gdf = gpd.GeoDataFrame(df, geometry='geometry')</pre> |
| geopandas.to_crs | Transforms the coordinate reference system of a GeoDataFrame. | gdf = gdf.to_crs(epsg=3857) |
| contextily.add_basemap | Adds a basemap to a GeoDataFrame plot for context. | <pre>import contextily as ctx ax = gdf.plot(figsize=(10, 10)) ctx.add_basemap(ax)</pre> |
| pca.explained_variance_ratio_ | Returns the proportion of variance explained by each principal component. | <pre>from sklearn.decomposition import PCA pca = PCA(n_components=2) pca.fit(X) variance_ratio = pca.explained_variance_ratio_</pre> |

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