Cheat Sheet: Evaluating and Validating Machine Learning Models

Model evaluation metrics and methods

| Method Name | Description | Code Syntax |
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| classification_report | Generates a report with precision, recall, F1-score, and support for each class in classification problems. Useful for model evaluation. Hyperparameters: target_names: List of labels to include in the report. Pros: Provides a comprehensive evaluation of classification models. Limitations: May not provide enough insight for imbalanced datasets. | <pre>from sklearn.metrics import classification_report # y_true: True labels # y_pred: Predicted labels # target_names: List of target class names report = classification_report(y_true, y_pred, target_names=["class1", "class2"])</pre> |
| confusion_matrix | Computes a confusion matrix to evaluate the classification performance, showing counts of true positives, false positives, true negatives, and false negatives. Hyperparameters: labels: List of class labels to include. Pros: Essential for understanding classification errors. Limitations: Doesn't give insights into prediction probabilities. | <pre>from sklearn.metrics import confusion_matrix # y_true: True labels # y_pred: Predicted labels conf_matrix = confusion_matrix(y_true, y_pred)</pre> |
| mean_squared_error | Calculates the mean squared error (MSE), a common metric for regression models. Lower values indicate better performance. Hyperparameters: sample_weight: Weights to apply to each sample. Pros: Simple and widely used metric. Limitations: Sensitive to outliers, as large errors are squared. | <pre>from sklearn.metrics import mean_squared_error # y_true: True values # y_pred: Predicted values # sample_weight: Optional, array of sample weights mse = mean_squared_error(y_true, y_pred)</pre> |
| root_mean_squared_error | Calculates the root mean squared error (RMSE), which is the square root of the MSE. RMSE gives more interpretable results as it is in the same units as the target. Hyperparameters: sample_weight: Weights to apply to each sample. Pros: More interpretable than MSE. Limitations: Like MSE, it can be sensitive to large errors and outliers. | <pre>from sklearn.metrics import root_mean_squared_error # y_true: True values # y_pred: Predicted values # sample_weight: Optional, array of sample weights rmse = root_mean_squared_error(y_true, y_pred)</pre> |

| mean_absolute_error | Measures the average magnitude of errors in predictions, without considering their direction. Useful for understanding the average error size. Hyperparameters: sample_weight: Optional sample weights. Pros: Less sensitive to outliers compared to MSE. Limitations: Does not penalize large errors as much as MSE or RMSE. | <pre>from sklearn.metrics import mean_absolute_error # y_true: True values # y_pred: Predicted values mae = mean_absolute_error(y_true, y_pred)</pre> |
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| r2_score | Computes the coefficient of determination (R²), which represents the proportion of variance explained by the model. A higher value indicates a better fit. Pros: Provides a clear indication of model performance. Limitations: Doesn't always represent model quality, especially for non-linear models. | <pre>from sklearn.metrics import r2_score # y_true: True values # y_pred: Predicted values r2 = r2_score(y_true, y_pred)</pre> |
| silhouette_score | Measures the quality of clustering by assessing the cohesion within clusters and separation between clusters. Higher scores indicate better clustering. Hyperparameters: metric: Distance metric to use. Pros: Useful for validating clustering performance. Limitations: Sensitive to outliers and choice of distance metric. | <pre>from sklearn.metrics import silhouette_score # X: Data used in clustering # labels: Cluster labels for each sample score = silhouette_score(X, labels, metric='euclidean')</pre> |
| silhouette_samples | Provides silhouette scores for each individual sample, indicating how well it fits its assigned cluster. Hyperparameters: metric: Distance metric to use. Pros: Offers granular insight into each sample's clustering quality. Limitations: Same as silhouette_score; sensitive to outliers and distance metric. | <pre>from sklearn.metrics import silhouette_samples # X: Data used in clustering # labels: Cluster labels for each sample samples = silhouette_samples(X, labels, metric='euclidean')</pre> |
| davies_bouldin_score | Measures the average similarity ratio of each cluster with the most similar cluster. Lower values indicate better clustering. Pros: Provides a simple, effective clustering evaluation. Limitations: May not | <pre>from sklearn.metrics import davies_bouldin_score # X: Data used in clustering # labels: Cluster labels for each sample db_score = davies_bouldin_score(X, labels)</pre> |

| | work well with highly imbalanced clusters. | |
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| Voronoi | Computes the Voronoi diagram, which partitions space based on the nearest neighbor. Pros : Useful for spatial analysis and clustering. Limitations : Limited to use cases that involve spatial partitioning of data. | <pre>from scipy.spatial import Voronoi # points: Coordinates for Voronoi diagram vor = Voronoi(points)</pre> |
| voronoi_plot_2d | Plots the Voronoi diagram in 2D for visualizing clustering results. Hyperparameters: show_vertices: Whether to display the vertices. Pros: Great for visualizing spatial clustering. Limitations: Limited to 2D spaces and large datasets may cause performance issues. | <pre>from scipy.spatial import voronoi_plot_2d # vor: Voronoi diagram object voronoi_plot_2d(vor, show_vertices=True)</pre> |
| matplotlib.patches.Patch | Creates custom shapes such as rectangles, circles, or ellipses for adding to plots. Hyperparameters: color: Fills color of the shape. Pros: Versatile for visual customization. Limitations: May not support all shapes or complex customizations. | <pre>import matplotlib.patches as patches # Create a rectangle with specified width, height, and position rectangle = patches.Rectangle((0, 0), 1, 1, color='blue')</pre> |
| explained_variance_score | Measures the proportion of variance explained by the model's predictions. A higher score indicates better performance. Pros: Helps in assessing the fit of regression models. Limitations: Not suitable for classification tasks. | <pre>from sklearn.metrics import explained_variance_score # y_true: True values # y_pred: Predicted values ev_score = explained_variance_score(y_true, y_pred)</pre> |
| Ridge regression | Performs ridge regression (L2 regularization) to avoid overfitting by penalizing large coefficients. Hyperparameters: alpha: Regularization strength. Pros: Helps reduce overfitting in regression models. Limitations: May not work well with sparse data. | <pre>from sklearn.linear_model import Ridge # alpha: Regularization strength (larger values indicate stronger regularization) ridge = Ridge(alpha=1.0)</pre> |
| Lasso regression | Performs lasso regression (L1 regularization), which encourages sparsity by penalizing the absolute value of coefficients. | <pre>from sklearn.linear_model import Lasso # alpha: Regularization strength (larger values indicate stronger regularization) lasso = Lasso(alpha=0.1)</pre> |

| | Hyperparameters: alpha: Regularization strength. Pros: Encourages sparse solutions, useful for feature selection. Limitations: May struggle with multicollinearity. | |
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| Pipeline | Chains multiple steps of preprocessing and modeling into a single object, ensuring efficient workflow. Pros: Simplifies code, ensures reproducibility. Limitations: May not work well with complex pipelines requiring dynamic configurations. | <pre>from sklearn.pipeline import Pipeline # steps: List of tuples with name and estimator/transformer pipeline = Pipeline(steps=[('scaler', StandardScaler()), ('model', Ridge(alpha=1.0))])</pre> |
| GridSearchCV | Performs exhaustive search over a specified parameter grid to find the best model configuration. Hyperparameters: param_grid: Dictionary of parameter grids. Pros: Ensures optimal model parameters. Limitations: Computationally expensive for large grids. | <pre>from sklearn.model_selection import GridSearchCV # estimator: Model to be tuned # param_grid: Dictionary with parameters to search over grid_search = GridSearchCV(estimator=Ridge(), param_grid={'alpha': [0.1, 1.0, 10.0]})</pre> |

Visualization strategies for k-means evaluation

| Process Name | Brief Description | Code Snippet |
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| Multiple runs of k- means | Executes KMeans clustering multiple times with different random initializations to assess variability in cluster assignments. Advantage: Helps visualize consistency. Limitation: Computationally costly for large datasets. | <pre># Number of runs for KMeans with different random states n_runs = 4 inertia_values = [] plt.figure(figsize=(12, 12)) # Run K-Means multiple_times with different random states for i in range(n_runs): kmeans = KMeans(n_clusters=4, random_state=None) # Use the default `n_init` kmeans.fit(X) inertia_values.append(kmeans.inertia_) # Plot the clustering result plt.subplot(2, 2, i + 1) plt.scatter(X(:, 0), X[:, 1], c=kmeans.labels_, cmap='tab10', alpha=0.6, edgecolor='k') plt.scatter(Kmeans.cluster_centers_[:, 0], kmeans.cluster_centers_[:, 1], c='red', s=200, marker='x', plt.title(f'K-Means Clustering Run {i + 1}') plt.vlabel('Feature 1') plt.vlabel('Feature 2') plt.legend() plt.tight_layout() plt.show() # Print inertia values for i, inertia in enumerate(inertia_values, start=1): print(f'Run {i}: Inertia={inertia:.2f}')</pre> |
| Elbow method | Evaluates the optimal number of clusters by plotting inertia (within-cluster sum of squares) for different k values. | <pre># Range of k values to test k_values = range(2, 11) # Store performance metrics inertia_values = [] for k in k_values: kmeans = KMeans(n_clusters=k, random_state=42) y_kmeans = kmeans.fit_predict(X) # Calculate and store metrics inertia_values.append(kmeans.inertia_) # Plot the inertia values (Elbow Method)</pre> |

| | Advantage: Easy to interpret. Limitation: Subjective elbow point. | <pre>plt.figure(figsize=(18, 6)) plt.subplot(1, 3, 1) plt.plot(k_values, inertia_values, marker='o') plt.title('Elbow Method: Inertia vs. k') plt.xlabel('Number of Clusters (k)') plt.ylabel('Inertia')</pre> |
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| Silhouette method | Determines the optimal number of clusters by evaluating Silhouette Scores for different k values. Advantage: Considers both cohesion and separation. Limitation: High computation for large datasets. | <pre># Range of k values to test k_values = range(2, 11) # Store performance metrics silhouette_scores = [] for k in k_values: kmeans = KMeans(n_clusters=k, random_state=42) y_kmeans = kmeans.fit_predict(X) silhouette_scores.append(silhouette_score(X, y_kmeans)) # Plot the Silhouette Scores plt.figure(figsize=(18, 6)) plt.subplot(1, 3, 2) plt.plot(k_values, silhouette_scores, marker='o') plt.title('Silhouette Score vs. k') plt.xlabel('Number of Clusters (k)') plt.ylabel('Silhouette Score')</pre> |
| Davies- Bouldin Index | Evaluates clustering performance by calculating DBI for different k values. Advantage: Quantifies compactness and separation. Limitation: Sensitive to cluster shapes and density. | <pre># Range of k values to test k_values = range(2, 11) # Store performance metrics davies_bouldin_indices = [] for k in k_values: kmeans = KMeans(n_clusters=k, random_state=42) y_kmeans = kmeans.fit_predict(X) davies_bouldin_indices.append(davies_bouldin_score(X, y_kmeans)) # Plot the Davies=Bouldin Index plt.figure(figisize=(18, 6)) plt.subplot(1, 3, 3) plt.plot(k_values, davies_bouldin_indices, marker='o') plt.title('Davies=Bouldin Index vs. k') plt.xlabel('Number of Clusters (k)') plt.ylabel('Davies=Bouldin Index')</pre> |

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Skills Network