

Application News

Gas Chromatography Mass Spectrometry

No. **M282**

Discovery of Markers from Chromatogram Data by Machine Learning

Techniques for identifying or classifying samples from chromatogram data by using machine learning algorithms have attracted considerable attention. In machine learning from chromatogram data, it is necessary to create a data table from peak intensity information, but the accuracy of the discriminant model may be reduced if a correlation exists between the sizes of the peaks or the sizes of the peaks are extremely different. For this reason, pretreatment of the data after creation of the data table of peak information is generally necessary in machine learning of chromatogram data.

This article introduces an example of the workflow in discovery of discriminant markers from GC-MS scan data of food samples by using Python 3.7.

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Data Acquisition

The data used here was the dataset for beef introduced in Application News No. M276. As shown in Fig. 1, two types of samples were prepared using red meat from various cuts of beef: properly refrigerated samples (4 °C samples) and samples which were expected to display deterioration due to exposure to a 40 °C environment for 3 h (40 °C samples). 20 \pm 3 mg of each sample was taken and placed in individual measurement vials. A total of 116 samples was prepared, comprising 58 of the 4 °C samples and 58 of the 40 °C samples, and the composition of the gas generated when the samples were heated to 200 °C was analyzed by the Solid Phase Micro Extraction (SPME) method. An AOC-6000 auto injector, which enabled automatic SPME injection, was used in this analysis (Fig. 2).

Fig. 3 shows an example of a chromatogram obtained as a result. Classification of the sample types was not possible from the sample appearance or the appearance of the chromatogram analysis results.





Fig. 1 Left: Properly Refrigerated Sample (4 °C Sample), Right: Sample Exposed to 40 °C Environment for 3 H (40 °C Sample)

Fig. 2 Appearance of GCMS-QP™2020 + AOC-6000



Fig. 3 Example of Total Ion Chromatograms Black: 4 °C Sample, Blue: 40 °C Sample

Picking/alignment of the deconvolution/peaks of the chromatogram data were done using the mass spectrometry data analysis software MZmine 2 (Ver. 2.32). The peak heights were output as data, as this data is relatively unaffected by waveform processing.

Because the dataset was extremely wide, comprising 9,318 peaks $\times 116$ datafiles, we basically proceeded in the direction of reducing unnecessary peak data (features) for the data pretreatment.

Data Pretreatment and Model Creation

<u>1. Treatment of missing values</u>

Although imputation of missing values in chromatogram data is possible by adjusting the waveform processing parameters to some extent, it is almost always impossible to eliminate all missing values. In such cases, artificial substitution by a simple calculation, separate regression analysis, or the like is not considered advisable. Therefore, in this study, all features, including missing values, that occurred even once were deleted by using a commercial spreadsheet program. As a result, it was possible to narrow the features of the dataset to 200 peaks \times 116 datafiles.

2. Data reading by Python and division into training and test sets

After deletion of the missing values, the feature names, data names, and similar information were input in the same general spreadsheet program. The dataset was saved in a CSV file, and was then imported to the Python console.

In [1]: import pandas as pd data =pd.read_csv("Data.csv", header=0, index_col=0) data.head(5)						
Out [1]:	reshne	ss RT1 32	001	RT22	52 199	RT23.48 200
Data001	0	154497	_001	10100	9510	11125.10_200
Data002	0	154356		12702	10054	
Data003	0	179444		15774	11863	
Data004	0	211854		11123	9546	
Data005	0	129346		12236	10996	
[5 rows x 200 columns]						

Here, the data names in each line ("Data001", "Data002"...), the label data in the "freshness" column ("1" or "0", indicating 4 °C Sample or 40 °C Sample, respectively), and the other feature data in each column (where the peaks are denoted by RT followed by a serial number) were arranged. The values of the features are the height of the respective peaks.

As described in Application News No. M276, this dataset was split into a training set of 92 datafiles and a test set of 24 datafiles. The model was created using the training set, and predictions were made using the test set. Division was done using Stratified Shuffle Split so as to avoid skewing of the label data.



3. Correlation between features

If a strong correlation exists between two features, one of the features in which this correlation is recognized is deleted due to occurrence of the problem of multicollinearity. For this reason, the correlation coefficient between each pair of features is checked.



Fig. 4 Heatmap of Correlation Coefficient Matrix for Pairs of Features

Many parts are shown in blue and red, indicating correlation of multiple features.

In [6]: def drop_correlating_columns (train_X, test_X, corr_value=0.8): import numpy as np corr1 = train_X.corr().abs() corr2 = np.triu(corr1.values,1) corr3 = np.asarray(np.where(corr2>corr_value)) corr4 = np.unique(corr3[1]) del_list = train_X.columns[corr4] train_X_ncorr = train_X.drop(del_list, axis=1) test_X_ncorr = test_X.drop(del_list, axis=1) return train_X_ncorr, test_X_ncorr						
In [7]: train_X_ncorr, test_X_ncorr = drop_correlating_columns¥ (train_X, test_X, corr_value=0.8)						
In [8]: train_X_ncorr.head(5)						
Out[8] RT1.3 Data050 Data006 Data032 Data047 Data055	2_001 RT3 135257 129535 219201 121548 90842	.98_003 36010 34817 13716 42813 31363	 	RT19.56_155 14516 7891 14183 9761 13666	RT22.52_199 13188 10228 12402 11610 13254	
[5 rows x 30 columns]						

It can be understood that peaks with similar retention times include many combinations with high correlation coefficients. Here, when a combination of features had a correlation coefficient of 0.8 or higher, one of the features was deleted. This made it possible to narrow the number of features to 30.

4. Narrowing of features

After the number of features decreases, the content of each feature is checked.

4-1. Distribution of values

The histograms of the values are checked, and select the features that include less outliers and the distribution is simple.

	In [9]: def hist_features(data, num): import matplotlib import matplotlib.pyplot as plt plt.figure(figsize=(8,16)) plt.subplots_adjust(wspace=0.2, hspace=1) matplotlib.rc('xtick', labelsize=8) matplotlib.rc('ytick', labelsize=8) for i, col in enumerate(list(data.columns)[num:num + 60]): plt.subplot(10, 3, i + 1) plt.hist(data[col]) plt.title(col, fontsize=12) return					
	In [10]: hist_features (train_X_i	ncc	orr, 0)			
	RT1.32_001		RT3.98_003	RT4.64_004		
10 0		10 0				
	100000 150000 20000 25000 RT6 11 005	00	20000 40000 60000 RT6 56 011	7500 10000 12500 15000 RT7 09 016		
20		20		20		
0		0	50000 100000 150000 200000			
	RT7.36_021		RT7.39_022	RT7.49_023		
50		20	dia.			
0	25000 50000 75000 100000	0	10000 15000 20000 25000	20000 30000 40000 50000		
20	RT7.70_026		RT8.15_027	RT8.30_029		
0		0	din .			
1	0000 20000 30000 40000		100000 150000 200000	0 10000 20000 30000		
	RT9.21_035	20	RT9.40_039	20 RT10.09_049		
10 0	.dlllu.,	0	1			
	100000 150000 200000 RT10.94 057		20000 40000 60000 RT10.95.066	10000 15000 RT12 46 083		
		-		20		
10		20				
0	10000 15000 20000 25000	0	10000 15000 20000 25000 30000	25000 50000 75000 100000		
	RT12.88_111		RT13.19_113	RT14.15_116		
20 0	alles and	0		2		
	200000 400000 600000	-	10000 20000 30000	100000 200000 300000		
	RT14.55_122		RT14.78_125	RT15.54_135		
10		10		10		
0	20000 20000 40000 50000	0	10000 15000 20000	0 10000 20000 30000 40000		
	RT15.70_136		RT16.48_137	RT16.73_139		
20	and an a	20	the second			
0	20000 30000 40000	0	10000 15000 20000 25000	0 5000 10000 15000		
	RT18.33_141	20	RT19.56_155	RT22.52_199		
10			all.			
0		0		0		

Fig. 5 Histograms of Features

Features that are outliers or have skewed distributions are excluded from candidates because functioning as a marker is generally difficult.

4-2. Distribution of features by label

Features whose distribution differs by label have a high possibility as marker candidates. The candidates are normalized with z-score and validated by a boxplot or other appropriate method.

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rrom skiearn.preprocessing import standardscaler import pandas as pd ss = StandardScaler().fit(train_X) train_X_scaled = pd.DataFrame(ss.transform¥ (X=train_X), columns=train_X.columns, index=train_X.index) test_X_scaled = pd.DataFrame(ss.transform¥ (X=test_X), columns=test_X.columns, index=test_X.index) return train_X_scaled, test_X_scaled

In:[13] train_X_ncorr_scaled,test_X_ncorr_scaled=¥ standard_scaler_for_train_test (train_X_ncorr, test_X_ncorr) boxplot(train_X_ncorr_scaled, train_y)



Fig. 6 Boxplot of Peaks by Label

4-3. Contribution to model

A temporal model is created here using the current 30 features, and the contributions of each feature are calculated. Since this study concerns the discovery of marker compounds with a high contribution to target classification by a binary discriminant model for the above-mentioned 4 °C and 40 °C samples, a logistic regression type model was used, and the hyperparameters "C" and "tol" were optimized by grid search.



sin.set(font_scale=2) ax.set_xlabel("Predicted Label", fontsize=20) ax.set_ylabel("True Label", fontsize=20) print("accuracy score is {0}".format(ac_score)) return predict_y, model_01

In [15]: predict_y_01, model_01 = predict_with_logreg¥ (train_X_ncorr_scaled, test_X_ncorr_scaled, train_y, test_y)

accuracy score is 0.75



Fig. 7 Confusion Matrix of Predicted Label and True Label

Although the classification accuracy is 75% at this point, in further work, we improved the accuracy of the model by selecting features with higher contributions.

Although the coefficients in logistic regression can be considered as contributions of each features, here permutation importance algorithm is adopted in terms of its higher generality in various type of models.

<pre>def permutation_importances (train_X, train_y, iter=10): from sklearn.inear_model import LogisticRegression as logreg from sklearn.model_selection import GridSearchCV from sklearn.model_selection import train_test_split from eli5.sklearn import PermutationImportance import pandas as pd import matplotlib.pyplot as plt perm_imps = pd.DataFrame(columns = train_X.columns) for i in range(iter): tr_X, val_X, tr_y, val_y = train_test_split ¥ (train_X, train_y, test_size=0.2, random_state=i) parameters_logreg = ¥ {"CC":[100, 10, 1, 0.1, 0.01], "tol":[1e-3, 1e-4, 1e-5, 1e-6, 1e-7]} gscv = GridSearchCV(logreg(), parameters_logreg, cv=5) gscv.fit(tr_X, tr_y) best_params = gscv.best_params_ model_01 = logreg(C=best_params["C"], ¥ tol=best_params["tol"], solver="lbfgs", random_state=0) model_01.fit (tr_X, tr_y) perm_imps = perm_imps.append ¥ (pd.Series(perm.feature_importance(model_01, ¥ random_state=0).fit(val_X, val_y) perm_fingus.append ¥ (pd.Series(perm.feature_importances_, ¥ index=train_X.columns), ignore_index=True) plt.figure(figsize=(12,8)).subplots_adjust(bottom=0.3) plt.bar(x=train_X.columns, height=perm_imps.mean(), ¥ yerr=perm_imps.std()) plt.thines(y=0, xmin=-0.5, xmax=train_X.shape[1]-0.5) return perm_imps</pre>
In [17]: perm_imps = permutation_importances ¥ (train_X_ncorr_scaled, train_y)

Since the number of samples is relatively small, a single try of calculation may not reach true value. Therefore, several split patterns of training / test set were tried, and their average value was taken. Although the standard deviation was rather large, as expected, it was possible to grasp the overall trend.



Based on the study and trial-and-error process described above, 10 features were selected at this time.

In [17]: feature_list_01 = pd.Series(("RT1.32_001", "RT3.98_003",¥ "RT7.39_022", "RT9.21_035", "RT9.40_039", "RT12.46_083",¥ "RT12.88_111", "RT14.15_116", "RT16.73_139", "RT19.56_155"])	
In [18]: train_X_dropped = train_X_ncorr_scaled[feature_list_01] test_X_dropped = test_X_ncorr_scaled[feature_list_01]	

Because GC-MS scan data are used in this study, a qualitative analysis of the candidate marker compounds is possible by using library search and reference standards at this point in time (Table 1).

Table 1 Peaks Selected in Stuc	ly and Their Library	/ Search Results
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Peak	#1 hit compound in library search results	Variable
RT1.32_001	Trimethylamine	x1
RT3.98_003	2-hydroxypropanamide	х2
RT7.39_022	gamman-Amylbutyrolactone	х3
RT9.21_035	Uric acid	x4
RT9.40_039	Lauryl acetate	х5
RT12.46_083	9-Octadecen-1-ol	хб
RT12.88_111	Hexadecanamide	х7
RT14.15_116	Oleic Acid	x8
RT16.73_139	Benzyl icosanoate	x9
RT19.56_155	5-Cholestene	x10

5. Feature engineering

In the logistic regression method, the output probability is expressed by a first-order standard sigmoid function of the features. A higher-order model of the features may be more effective in some cases, depending on the dataset. However, there is currently a tendency to avoid extreme high-order models, which generally result in an increased calculation load and overfitting.

Here, the result of a single division operation for each case was used as a new feature "ratio of pairs of compound peaks," considering the fact that the data were chromatograms.

```
In [19]:

def features_ratio (train_X, test_X):

import pandas as pd

from sklearn.preprocessing import StandardScaler

train_X_r = train_X.copy()

for col1 in train_X.columns:

for col2 in train_X.columns:

train_X_r[col1+"_"+col2] = train_X[col1] / train_X[col2]
```

<pre>test_X_r = test_X.copy() for col1 in test_X.columns: for col2 in test_X.columns: test_X_r[col1+"_"+col2] = test_X[col1] / test_X[col2]</pre>
ss = StandardScaler().fit(train_X_r)
train_X_r = pd.DataFrame(ss.transform(train_X_r),¥ columns=train_X_r.columns, index=train_X_r.index) test_X_r = pd.DataFrame (ss.transform(test_X_r), ¥ columns=test_X_r.columns, index=test_X_r.index return train_X_r, test_X_r
In [20]: train_X_r, test_X_r = features_ratio (train_X_dropped, ¥ test_X_dropped)

Because the number of features became excessive, features values were selected arbitrarily based on the distribution and contribution of the features, in the same manner as above.

The final features were as follows:

In [21]: feature_list_final = pd.Series(["RT9.21_035",¥ "RT9.21_035_RT7.39_022", "RT3.98_003_RT9.21_035",¥ "RT3.98_003_RT16.73_139", "RT9.40_039_RT19.56_155",¥ "RT1.32_001_RT9.21_035", "RT16.73_139",¥ "RT14.15_116_RT1.32_001", "RT9.21_035_RT19.56_155",¥ "RT7.39_022_RT9.40_039"]) In [22]: train_X_final = train_X_r[feature_list_final] test_X_final = test_X_r[feature_list_final] In [23]: predict_y_final, model_final = predict_with_logreg¥ (train_X_final, test_X_final, train_y, test_y)	

accuracy score is 0.91666666666666666



Fig. 9 Confusion Matrix of Predicted Label and True Label

Finally, it was possible to classify unknown samples with precision of 91.67% by using this model. The equation for predicting the probability *P* of label 1 is as follows.

$$P = (1 + \exp(1.19 * x 4 + 1.22 * \frac{x4}{x3} - 0.21 * \frac{x2}{x4} - 1.32 * \frac{x2}{x9} + 1.72 * \frac{x5}{x10} - 0.50 * \frac{x1}{x4} + 0.18 * x9 + 1.25 * \frac{x8}{x1} + 0.16 * \frac{x4}{x10} - 0.07 * \frac{x3}{x5}))^{-1}$$

The classification process is based on $P \ge 0.5 \Rightarrow 1, P < 0.5 \Rightarrow 0$.

Table 2 Version Information

Python	3.7.3	seaborn	0.9.0
numpy	1.16.2	sklearn	0.20.3
pandas	0.24.2	eli5	0.8.2
matplotlib	3.0.3		

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