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Python与机器学习 ——机器学习算法

华算科技 黄老师 2022年1月18日



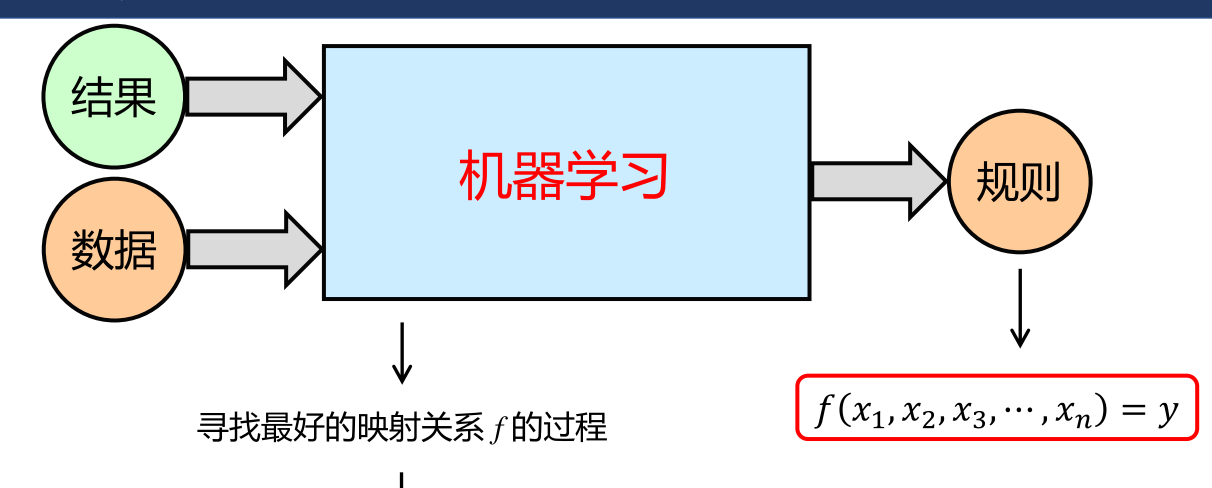
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- 2. 回归算法
- 3. 双金属吸附中的回归
- 4. 非线性回归求速率常数
- 5. 模型评价
- 6. 决策树分类算法
- 7. 支持向量机

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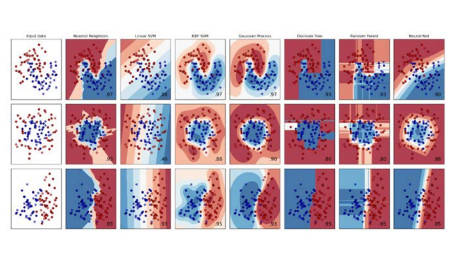
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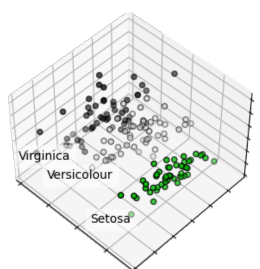
机器学习算法



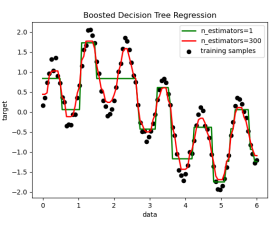
算法

机器学习算法





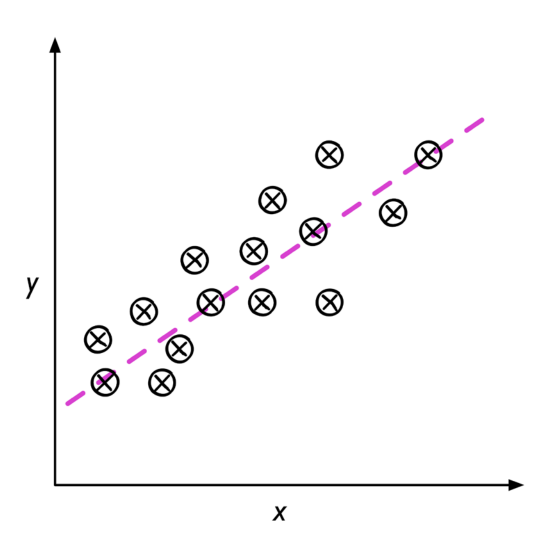


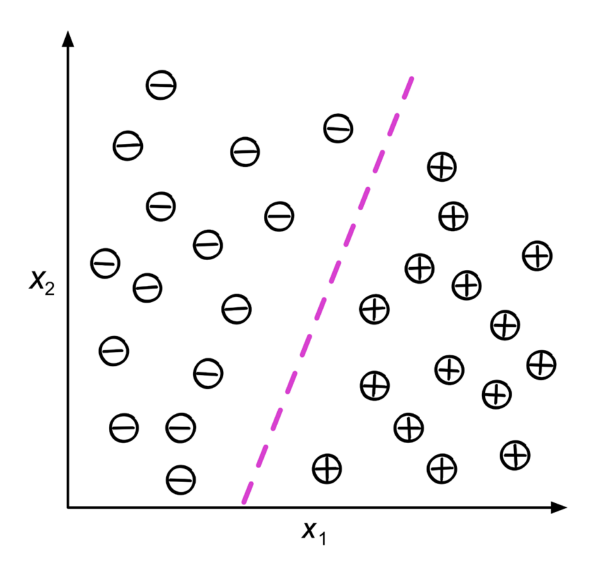


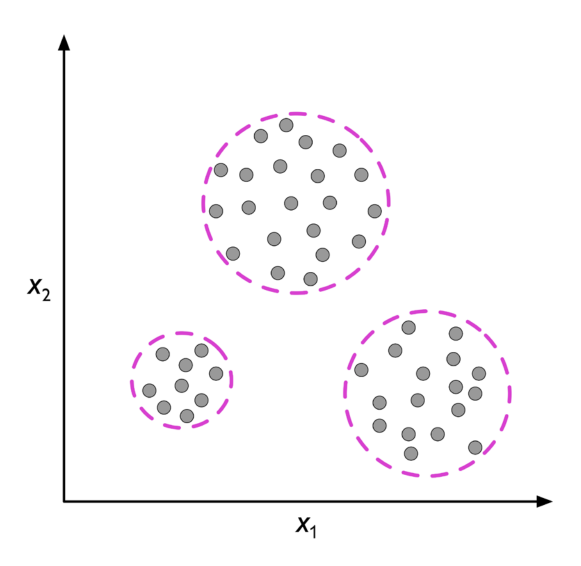
K-means clustering on the digits dataset (PCA-reduced data) Centroids are marked with white cross



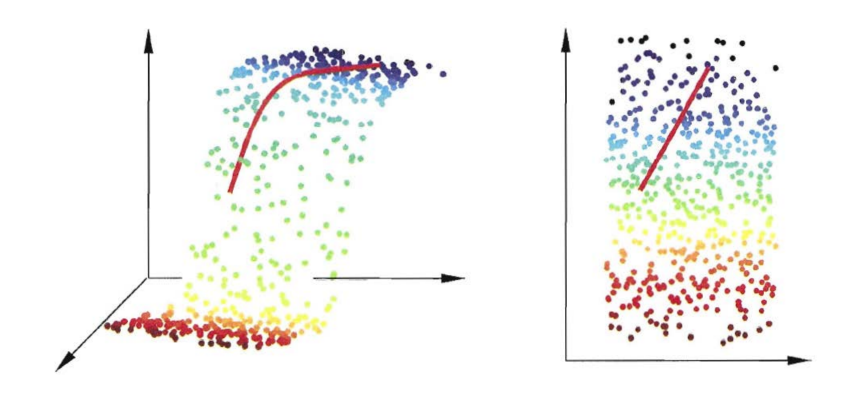
回归算法







维度约减



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回归算法

寻找目标值(通常是连续值)与输入值之间关系的算法。

线性回归

Linear Regression

逻辑回归

Logistic Regression

岭回归

Ridge Regression

多项式回归

Polynomial Regression

套索回归

Lasso Regression

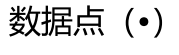
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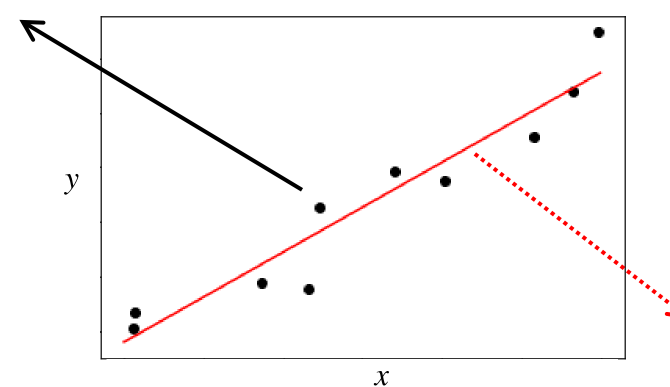
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回归算法原理

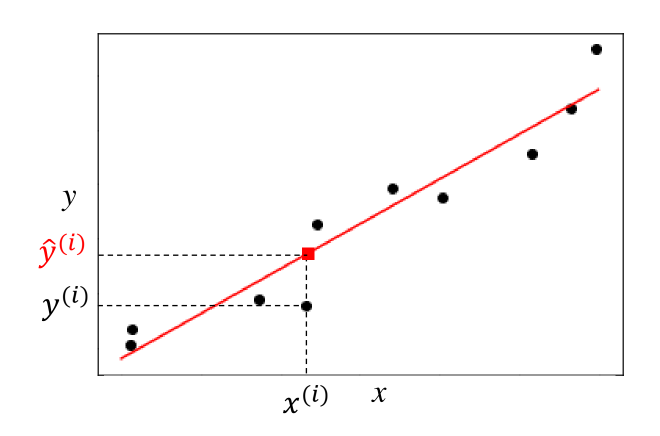
$$\{(x^{(1)}, y^{(1)}), (x^{(2)}, y^{(2)}), \cdots, (x^{(m)}, y^{(m)})\}$$





映射关系 (f)

$$\hat{y} = f(x) = w_0 + w_1 x$$



$$\hat{y}^{(i)} = f(x^{(i)}) = w_0 + w_1 x^{(i)}$$

残差

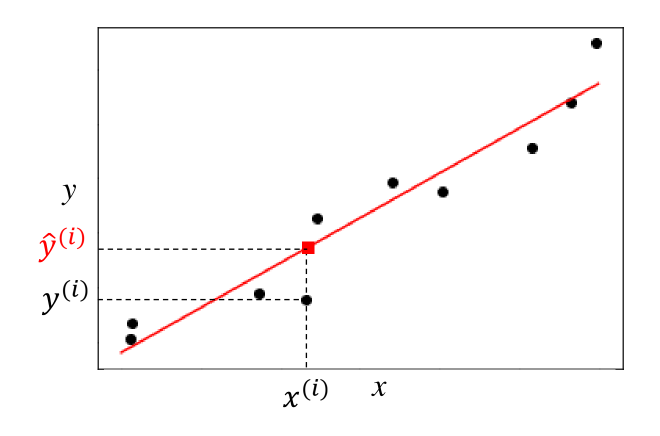
$$\varepsilon_i = \hat{y}^{(i)} - y^{(i)}$$

残差平方和

$$\sum \varepsilon_i^2 = \sum (\hat{y}^{(i)} - y^{(i)})^2$$

最小二乘法原理

$$\min \sum (\hat{y}^{(i)} - y^{(i)})^2$$



$$\hat{y}^{(i)} = f(x^{(i)}) = w_0 + w_1 x^{(i)}$$

$$\min \sum (\hat{y}^{(i)} - y^{(i)})^2$$

$$S = \sum (w_0 + w_1 x^{(i)} - y^{(i)})^2$$

最小值处 S 对 w_0 与 w_1 的偏导数为0

$$\int \frac{\partial S}{\partial w_0} = 2 \times \sum (w_0 + w_1 x^{(i)} - y^{(i)}) = 0$$
$$\frac{\partial S}{\partial w_1} = 2 \times \sum (w_0 + w_1 x^{(i)} - y^{(i)}) x^{(i)} = 0$$

$$\int \frac{\partial S}{\partial w_0} = 2 \times \sum (w_0 + w_1 x^{(i)} - y^{(i)}) = 0$$
$$\frac{\partial S}{\partial w_1} = 2 \times \sum (w_0 + w_1 x^{(i)} - y^{(i)}) x^{(i)} = 0$$

一元线性回归

解简单的二元一次方程组

得到
$$w_0$$
与 w_1 $f(x) = w_0 + w_1 x$

$$w_0 = \bar{y} - w_1 \bar{x}$$

$$w_1 = \frac{\sum (x^{(i)} - \bar{x})(y^{(i)} - \bar{y})}{\sum (x^{(i)} - \bar{x})^2}$$

$$\bar{x} = \frac{\sum x^{(i)}}{m}$$
 $\bar{y} = \frac{\sum y^{(i)}}{m}$

n元情形

数据点

$$\left\{ \left(x_1^{(1)}, x_2^{(1)}, \dots, x_n^{(1)}, y^{(1)} \right), \left(x_1^{(2)}, x_2^{(2)}, \dots, x_n^{(2)}, y^{(2)} \right), \cdots, \left(x_1^{(m)}, x_2^{(m)}, \dots, x_n^{(m)}, y^{(m)} \right) \right\}$$

映射关系

$$f(x) = w_0 + w_1 x_1 + w_2 x_2 + \dots + w_n x_n$$

目标: 寻找到一组 $w(w_0, w_1, w_2, \cdots, w_n)$,

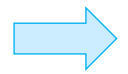
使得 $S(w) = \sum (\hat{y}^{(i)} - y^{(i)})^2$ 最小。

$$\hat{y}^{(1)} = w_0 + w_1 x_1^{(1)} + w_2 x_2^{(1)} + \dots + w_n x_n^{(1)}$$

$$\hat{y}^{(2)} = w_0 + w_1 x_1^{(2)} + w_2 x_2^{(2)} + \dots + w_n x_n^{(2)}$$
.....

$$\hat{y}^{(m)} = w_0 + w_1 x_1^{(m)} + w_2 x_2^{(m)} + \dots + w_n x_n^{(m)}$$

$$\mathbf{X} = \begin{pmatrix} x_1^{(1)} & \cdots & x_n^{(1)} \\ \vdots & \ddots & \vdots \\ x_1^{(m)} & \cdots & x_n^{(m)} \end{pmatrix} \qquad \mathbf{w} = (w_1, w_2, \cdots, w_n)$$



$$\widehat{\boldsymbol{y}} = \mathbf{X} \boldsymbol{w}^{\mathrm{T}} + \boldsymbol{w}_{0} \qquad \widehat{\boldsymbol{y}} = \mathbf{X} \boldsymbol{w}^{\mathrm{T}}$$

$$\widehat{\mathbf{y}} = \mathbf{X} \mathbf{w}^{\mathrm{T}}$$

n元情形
$$S(w) = \sum (\hat{y}^{(i)} - y^{(i)})^2 = \sum (w_0 + w_1 x_1^{(i)} + w_2 x_2^{(i)} + \dots + w_n x_n^{(i)} - y^{(i)})^2$$
$$= ||\mathbf{X} \mathbf{w}^T - \mathbf{y}||^2$$

同样的, 求导
$$\frac{\partial S(\mathbf{w})}{\partial \mathbf{w}^T} = 2 \sum_{i=1}^{T} \sum_{j=1}^{T} x_j^{(i)} (w_j x_j^{(i)} - y^{(i)})$$
$$= 2\mathbf{X}^T (\mathbf{X} \mathbf{w}^T - \mathbf{y})$$

令偏导数为0
$$\mathbf{w}^T = (\mathbf{X}^T \mathbf{X})^{-1} \mathbf{X}^T \mathbf{y}$$

正规方程 The Normal Equation

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文献中的线性回归

Journal of Cluster Science https://doi.org/10.1007/s10876-018-1346-x

ORIGINAL PAPER



Mechanism of the Reverse Water-Gas Shift Reaction Catalyzed by Cu₁₂TM Bimetallic Nanocluster: A Density Functional Theory Study

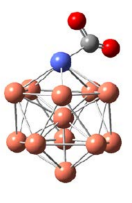
Qian Zhang^{1,2} · Ling Guo^{1,2}

Received: 20 July 2017 © Springer Science+Business Media, LLC, part of Springer Nature 2018

Abstract

Density functional theory calculations were carried out to investigate $Cu_{12}TM$ (TM = Co, Rh, Ir, Ni, Pd, Pt, Ag, Au) bimetallic metal catalysts for the mechanism of reverse water–gas shift (RWGS) reaction. The three possible reaction pathways relevant to the RWGS reaction are explored, including the CO_2 dissociation, carboxyl, and formate mechanisms. Our results indicate that the RWGS reaction prefers to follow the CO_2 dissociation mechanism on $Cu_{12}TM$ surfaces. A detailed potential energy diagram of the kinetically favored mechanism is presented that shows that the RDS of reaction are the formation of H_2O and carboxyl (HOCO), formate (HCOO) dissociation, respectively. And, $Cu_{12}TM$ (TM = CO, Pt) are lower than other catalysts from the energy barrier of elementary step. Moreover, the catalytic behavior of a $Cu_{12}TM$ cluster is changed significantly due to the modifiers, via the electron transfer from TM to Cu-based cluster, and the activation barrier decreases with doped TM. The turnover frequency of the $Cu_{12}CO$ is the highest value, which thus is more efficiency catalyst to RWGS reaction. To gain insights into the synergistic effect in catalytic activity of the $Cu_{12}TM$ bimetallic cluster, a projected density of states analysis has been performed. Our works will be important for predicting the energetic trends and designing a better catalyst of RWGS reaction.

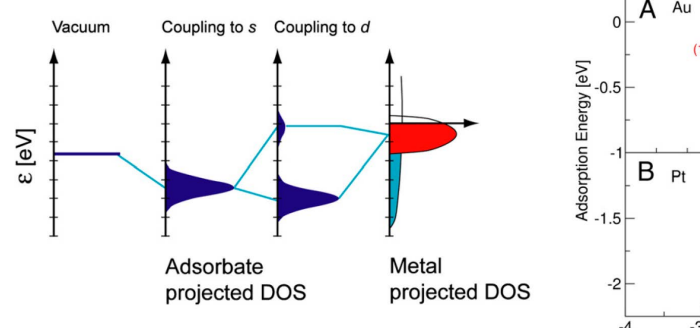
Keywords Cu₁₂TM bimetallic nanocluster · RWGS reaction · Mechanism · TOF · d-Band center

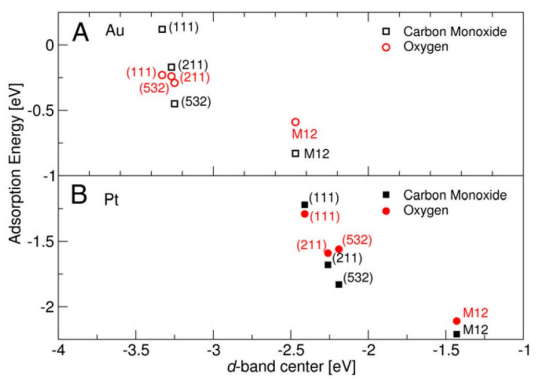


Cu₁₂TM-CO2*

d带中心理论

CO、O₂在Au与 Pt上的吸附能





J. K. Nørskov, T. Bligaardc, et. al. Proc. Natl. Acad. Sci 2011, 108, 937.

d带中心与吸附能

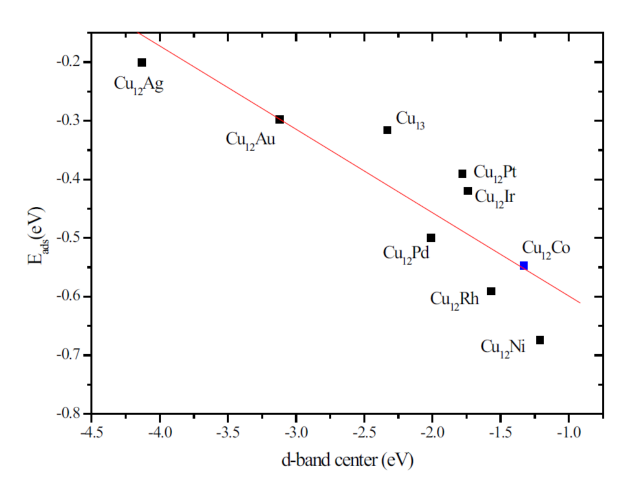
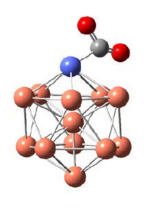


Fig. 6 Relationship between d-band center of TM metal atom and adsorption energy of CO₂ molecular



Cu₁₂TM-CO2*

d带中心 \longleftrightarrow CO₂吸附能

Cu ₁₂ Ir	Cu ₁₂ Ni	Cu ₁₂ Pd	Cu ₁₂ Pt	Cu ₁₃	Cu ₁₂ Ag	Cu ₁₂ Au
- 1.74	- 1.21	- 2.01	- 1.78	- 2.33	- 4.13	- 3.12
-0.420	-0.674	-0.500	- 0.391	- 0.316	- 0.201	- 0.298

```
import numpy as np
import pandas as pd

columns_names = ['formula', 'db_center', 'ad_energy']

data = pd.read_csv('db_center.csv', names = columns_names)

t

文取文件
```

	formula	db_center	ad_energy
0	Cu12Co	-1.33	-0.547
1	Cu12Rh	-1.57	-0.591
2	Cu12lr	-1.74	-0.420
3	Cu12Ni	-1.21	-0.674
4	Cu12Pd	-2.01	-0.500

4	А	В	С	
1	Cu12Co	-1.33	-0.547	
2	Cu12Rh	-1.57	-0.591	
3	Cu12lr	-1.74	-0.42	
4	Cu12Ni	-1.21	-0.674	
5	Cu12Pd	-2.01	-0.5	
6	Cu12Pt	-1.78	-0.391	
7	Cu13	-2.33	-0.316	
8	Cu12Ag	-4.13	-0.201	
9	Cu12Au	-3.12	-0.298	
4.0				

```
1 \mid X = data['db\_center']. values. reshape (-1, 1)
 2 | y = data['ad_energy']. values
  3 print(X)
  4 print(y)
[[-1.33]
 [-1.57]
 [-1.74]
                                                                            查看文件
 [-1.21]
 [-2.01]
 [-1.78]
 [-2.33]
 [-4.13]
 [-3.12]
[-0.547 -0.591 -0.42 -0.674 -0.5 -0.391 -0.316 -0.201 -0.298]
```

```
from sklearn.linear_model import LinearRegression
lr = LinearRegression()

1r.fit(X,y)
R2 = 1r.score(X,y)
print("R2 = " + str(R2))
```

R2 = 0.7646612597264623

```
1 predicts = 1r. predict([[-2.2]])
2 print("precdicts = " + str(predicts))

precdicts = [-0.42839089]
```

输出斜率与截距的方法

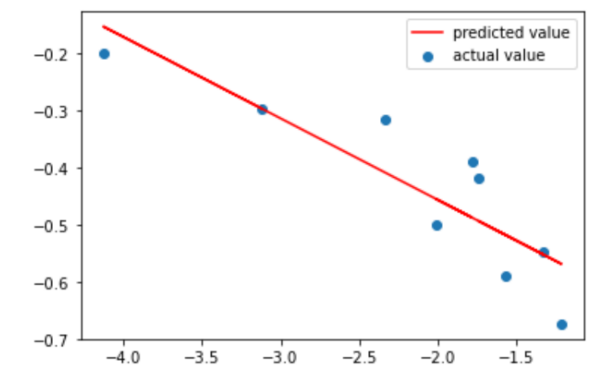
```
斜率 coef = lr.coef_
截距 intercept = lr.intercept
```

```
In [4]: coef = lr.coef_
intercept = lr.intercept_
print(lr.coef_, lr.intercept_)

[-0.14221031] -0.7412535628655477
```

```
import matplotlib.pyplot as plt

x = data['db_center'].values
plt.scatter(x, y, label='actual value')
plt.plot(x, lr.predict(X), color='red', label='predicted value')
plt.legend()
plt.show()
```



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非线性回归背景: 连续反应动力学

$$A \xrightarrow{k_1} B \xrightarrow{k_2} C$$

$$\frac{dc_A}{dt} = -k_1 c_A$$

$$A - k_2 c_B$$

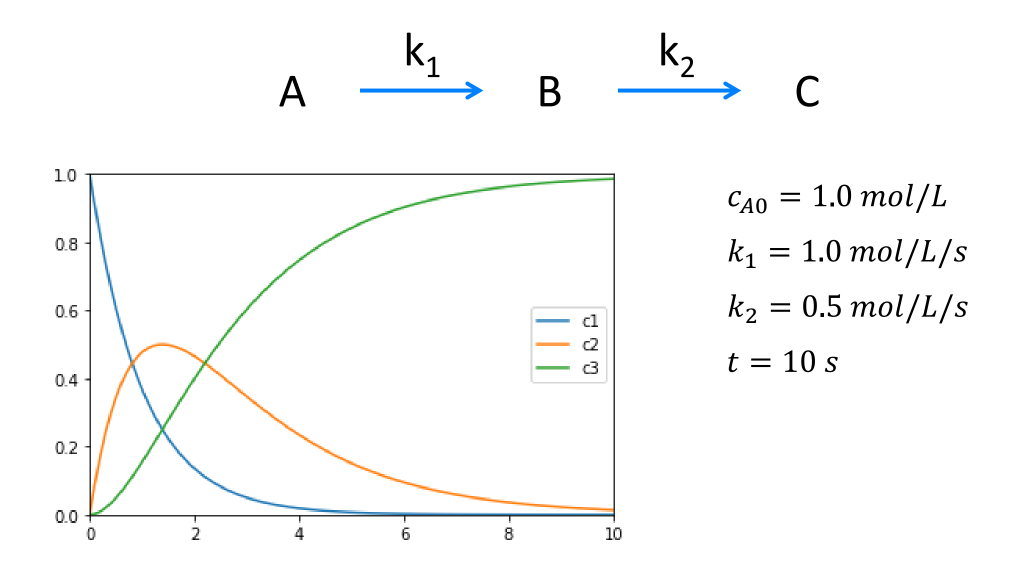
$$\frac{dc_c}{dt} = k_2 c_B$$

$$c_A = c_{A0} e^{k_1 t}$$

$$\frac{dc_B}{dt} = k_1 c_A - k_2 c_B \qquad \qquad c_B = \frac{k_1 c_{A0}}{k_2 - k_1} (e^{-k_1 t} - e^{-k_2 t})$$

$$c_C = c_{A0}(1 - \frac{k_2}{k_2 - k_1}e^{-k_1t} - \frac{k_1}{k_2 - k_1}e^{-k_2t})$$

连续反应动力学



实操: 反应速率的确定

```
In [4]: def func(t, k1, k2):
             c10 = 1.0
             c1 = c10 * np. exp(- k1 * t)
             c2 = k1 * c10 / (k2 - k1) * (np. exp(-k1 * t) - np. exp(-k2 * t))
             c3 = c10 - c1 - c2
             return c3
In [5]: from scipy.optimize import curve_fit
         pfit, pcov = curve_fit(func, df['t'], df['c3'], p0 = [1, 0])
         print(pfit)
         [0.93621629 0.51942599]
In [6]: plt. scatter(df['t'], df['c3'], s = 50, c = 'k')
         t_{cuv} = np. linspace(0, 10, 100)
         plt.plot(t_cuv, func(t_cuv, 0.936, 0.519), c = 'r')
         plt. show()
          1.0
          0.8
          0.6
          0.4
          0.2
          0.0
```

非线性回归

Python提供了自定义 函数回归的方式

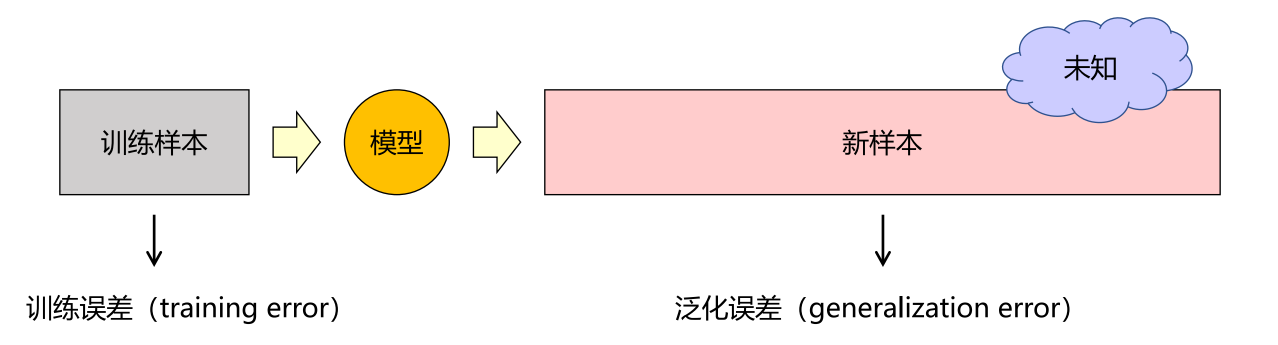
curve_fit()

def func(x, p1, p2): pfit, pcov = curve fit(func, x val, y val) $\sigma(x,y) = \frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})(y_i - \bar{y})$ np数组,拟合参数

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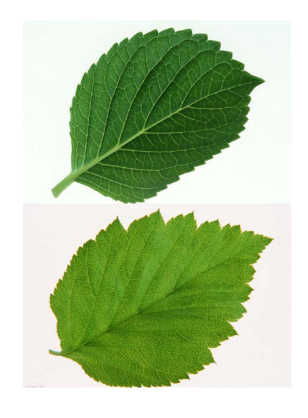
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误差



欠拟合与过拟合

判断图片中是否是树叶



树叶训练集



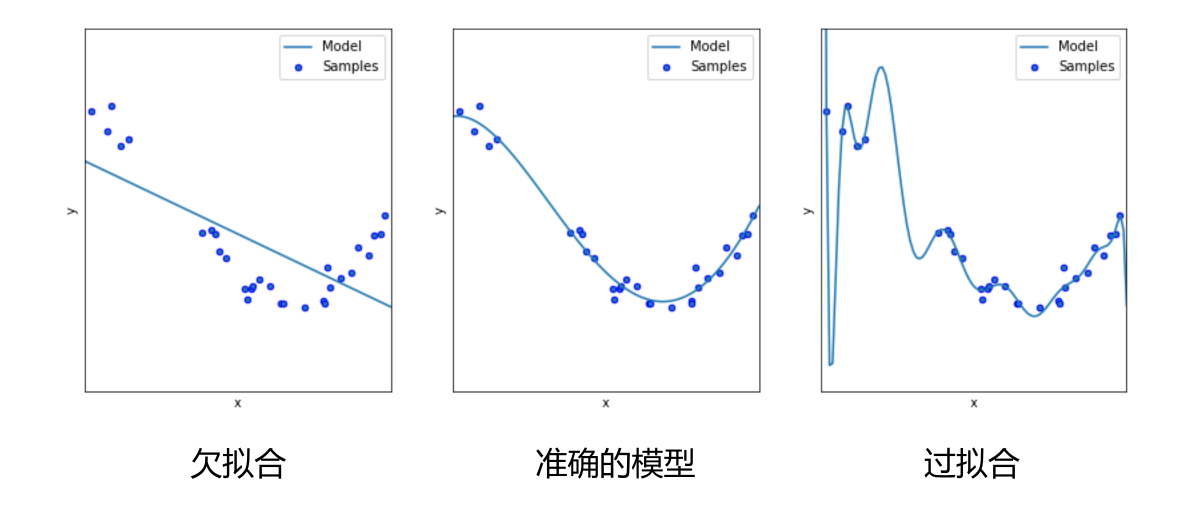
欠拟合(Underfitting) 结果:是树叶 绿色的就是树叶



过拟合(Overfitting) 结果:不是树叶 有锯齿的菜才是树叶

测试集

欠拟合与过拟合



偏差与方差

低偏差

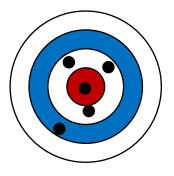
高偏差

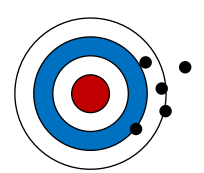
低方差



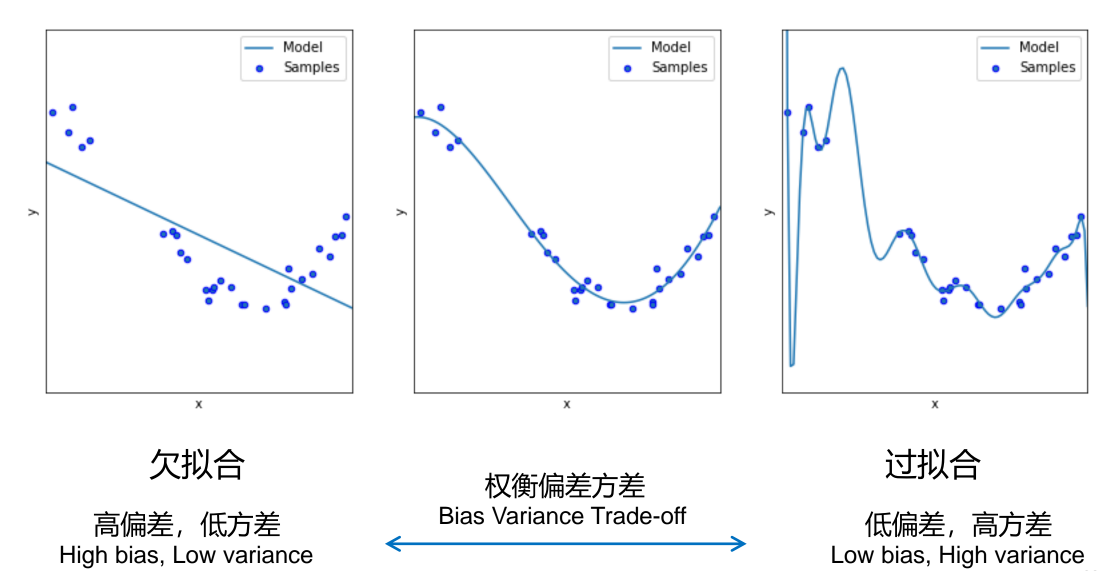


高方差

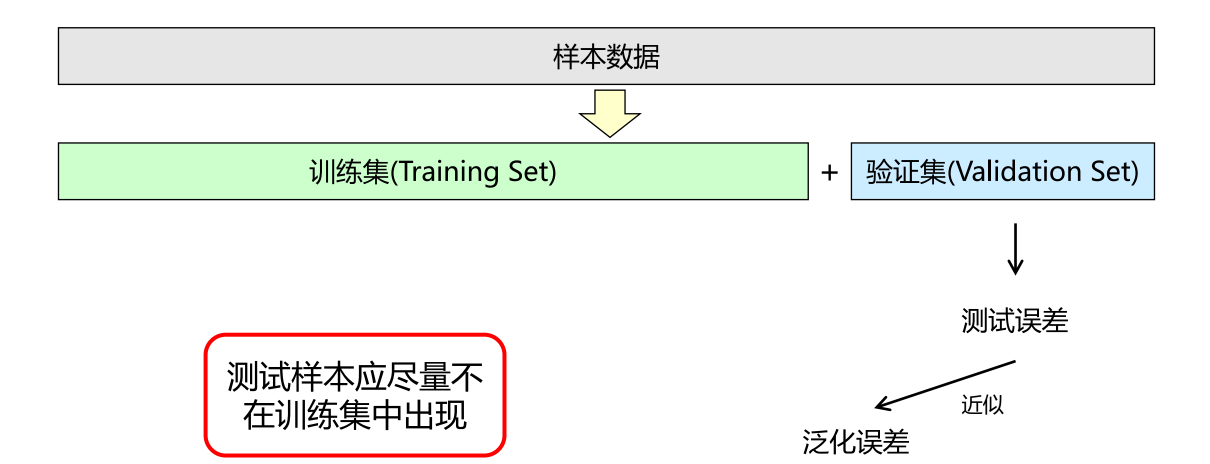




准确的概念

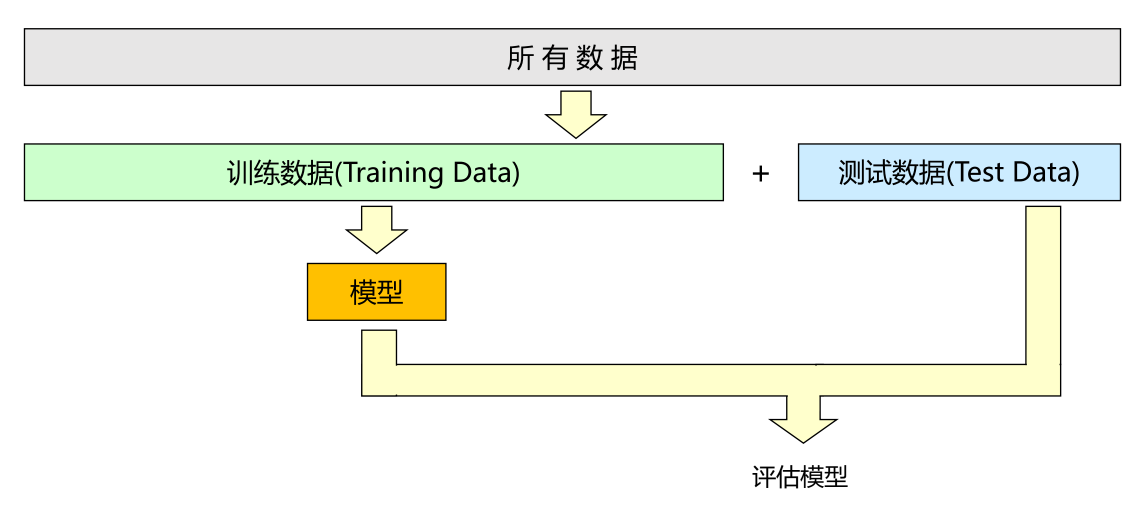


训练集与测试集



留出法

留出法 (hold-out)





DFT+ML预测晶格热导率

www.acsami.org Research Article

Lattice Thermal Conductivity: An Accelerated Discovery Guided by Machine Learning

Russlan Jaafreh, Yoo Seong Kang, and Kotiba Hamad*



Cite This: ACS Appl. Mater. Interfaces 2021, 13, 57204–57213



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ABSTRACT: In the present work, we used machine learning (ML) techniques to build a crystal-based model that can predict the lattice thermal conductivity (LTC) of crystalline materials. To achieve this, first, LTCs of 119 compounds at various temperatures (100–1000 K) were obtained based on density functional theory (DFT) and phonon calculations, and then, these data were employed in the next learning process to build a predictive model using various ML algorithms. The ML results showed that the model built based on the random forest (RF) algorithm with an R² score of 0.957 was the most accurate compared with the models built using other algorithms. Additionally, the accuracy of this model was validated using new cases of four compounds, which was not seen for the model before, where a good matching between calculated and predicted LTCs of the new compounds was found. To find candidates with ultralow LTCs (<1 W m⁻¹ K⁻¹) at room temperature, the model was used to screen compounds (32116) in the Inorganic Crystal Structure Database. From the screened compounds, Cs₂SnI₆ and SrS were selected to validate the ML prediction

示例

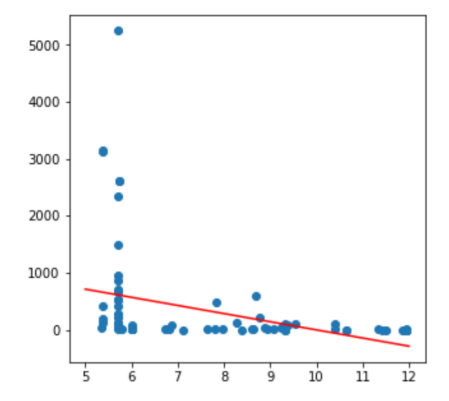
整个模型的拟合结果

```
In [1]:
            1 import pandas as pd
            2 import matplotlib.pyplot as plt
            3 import numpy as np
              from sklearn.linear_model import LinearRegression
            6 df = pd. read csv('ltc. csv')
            7 df
 Out[1]:
                Compound Temperature(K) mean_EffectiveCoordination var_EffectiveCoordination min_EffectiveCoordination m
                      AlAs
                                152.91209
                                                            5.720212
                                                                                 3.110000e-15
                                                                                                              5.720212
             0
             1
                      AIP
                                140.72993
                                                            5.368858
                                                                                 3.150000e-10
                                                                                                              5.368858
                    AIP (F-
             2
                                162.26119
                                                            5.720212
                                                                                 1.330000e-15
                                                                                                              5.720212
                     43m)
             3
                     AISb
                                148.62687
                                                            5.720212
                                                                                 0.000000e+00
                                                                                                              5.720212
                    B2AsP
                                142.34014
                                                            5.731569
                                                                                 1.021271e-01
                                                                                                              5.541343
             4
                                                                                 4.440000e-16
           114
                     TePb
                                147.79412
                                                            6.000000
                                                                                                              6.000000
           115
                   Ti2SnC
                                100.70368
                                                            9.383733
                                                                                 1.691867e+00
                                                                                                              6.000000
                      TIBr
                                160.52725
                                                           11.956922
                                                                                 8.880000e-16
                                                                                                             11.956922
           116
           117
                      TICI
                                160.26508
                                                           11.956922
                                                                                 8.880000e-16
                                                                                                             11.956922
           118
                       TII
                                148.12925
                                                           11.956922
                                                                                 8.880000e-16
                                                                                                             11.956922
```

示例

```
In [2]:

1    X = df['mean_EffectiveCoordination'].values.reshape(-1,1)
2    y = df['LTC'].values
3    lr = LinearRegression()
4    lr.fit(X, y)
5    plt.figure(figsize = (5, 5))
6    plt.scatter(df['mean_EffectiveCoordination'], df['LTC'])
7    plot_x = np.linspace(5, 12)
8    plot_y = plot_x * lr.coef_ + lr.intercept_
9    plt.plot(plot_x, plot_y, c = 'r')
10    plt.show()
```



示例

留出法评价

```
In [4]:
         1 X lo train = X[:-35]
           2 | y 1o train = y[:-35]
           3 X 1o test = X[-35:]
           4 | y 1o test = y[-35:]
   [5]:
          1 | 1r 1o = LinearRegression()
           2 lr_lo.fit(X_lo_train, y_lo_train)
           3 print(lr_lo.coef_, lr_lo.intercept_)
          [-136. 98658328] 1375. 0261185904187
    [6]:
              from sklearn.metrics import mean_squared_error
             R2 lo = lr lo. score(X lo test, y lo test)
           4 print('RMSE = %.3f'% np. sqrt(mean squared error(y true = y lo test, y pred = lr lo.predict(X lo test))))
           5 | print("R2 = " + str(R2_{10})) |
         RMSE = 831.068
         R2 = 0.1063898139220959
```

交叉验证法 (Cross Validation)

所有数据								
数据1	数据2	数据3	•••	数据k				
数据1	数据2	数据3	•••	数据k				
数据1	数据2	数据3	•••	数据k				
数据1	数据2	数据3	•••	数据k				
数据1	数据2	数据3	•••	数据k				

样例:交叉验证

Folds: 5, mean RMSE: 646.926

留一法 (Leave-One-Out)

所有数据 (m个样本)



数据1

数据2

数据3

•••

数据k

k = m

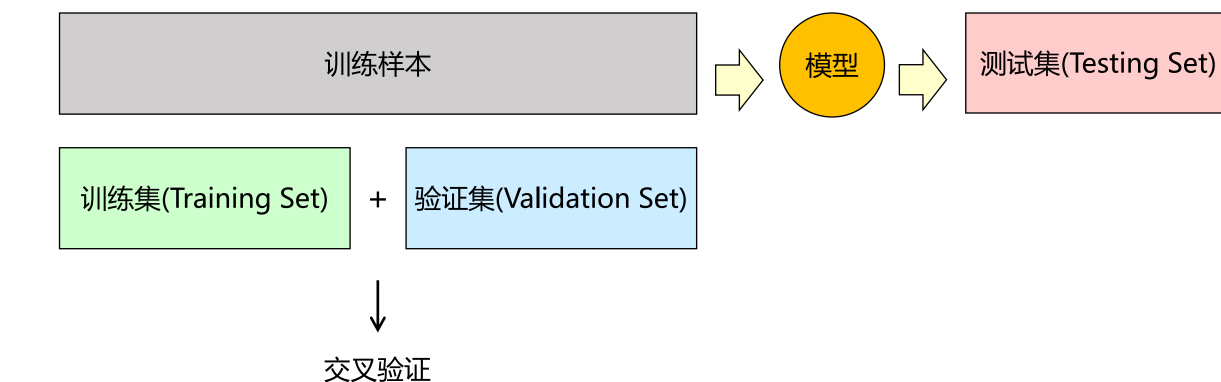
每次只拿一个 样本做测试

样例:留一法的使用

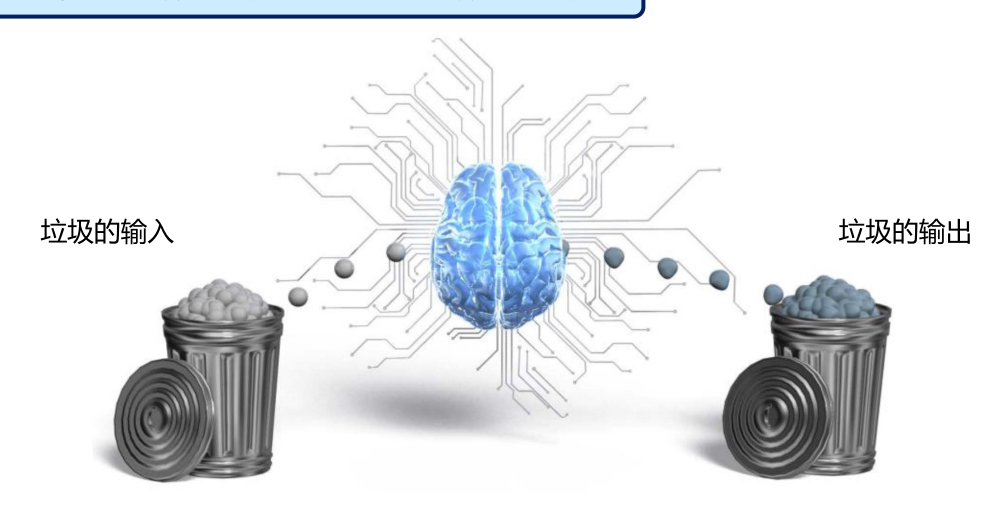
Folds: 119, mean RMSE: 358.903

```
[11]:
            from sklearn.model selection import LeaveOneOut
            sample x = range(5)
           loo = LeaveOneOut()
           for train, test in loo.split(sample_x):
                print(train, test)
       [1 \ 2 \ 3 \ 4] \ [0]
       [0\ 2\ 3\ 4] [1]
       [0\ 1\ 3\ 4] [2]
       [0 \ 1 \ 2 \ 4] \ [3]
       [0 \ 1 \ 2 \ 3] \ [4]
         1 rmse_scores = cross_val_score(1r, X, y, scoring = 'neg_root_mean_squared_error', cv=len(X))
[12]:
         2 print ('Folds: %i, mean RMSE: %.3f' % (len(rmse_scores), -np. mean(rmse_scores)))
```

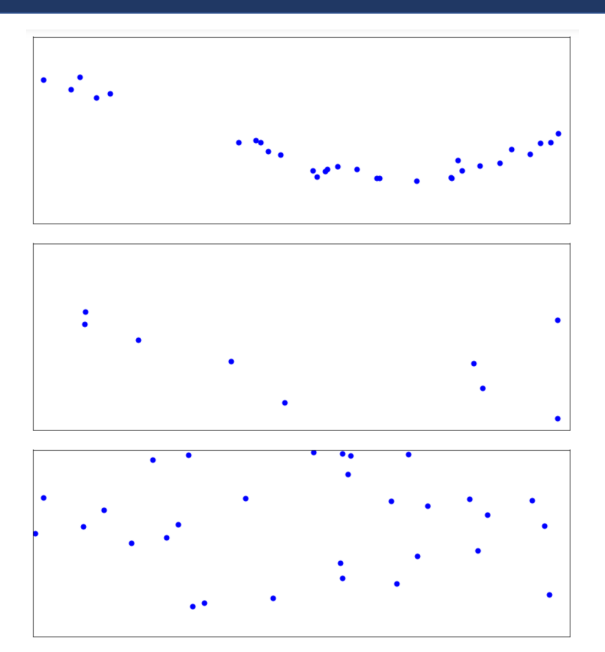
交叉验证法 (Cross Validation)



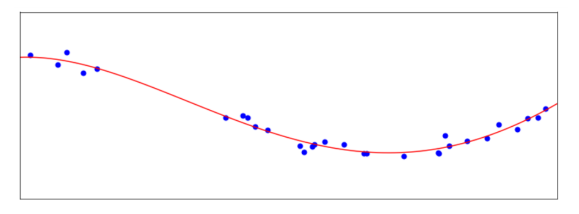
永远记住,你<mark>输入</mark>数据的质量决定了你<mark>输出</mark>数据集的质量

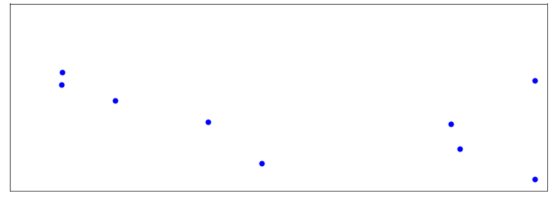


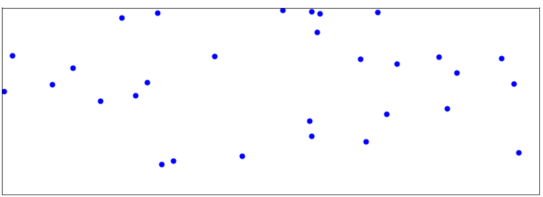
T. Toyao, K. Shimizu, et. al. ACS Catal. 2020, 10, 2260.



对这三组数据分别进行回归



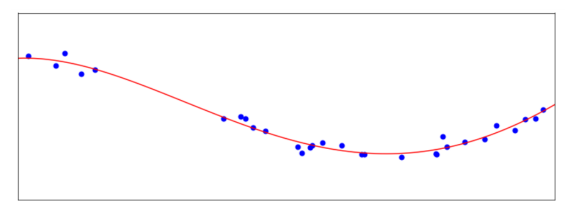


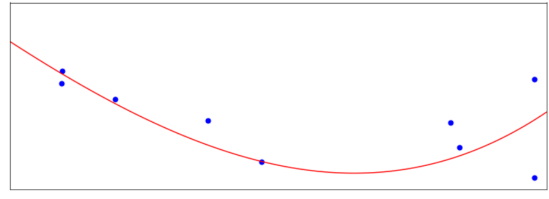


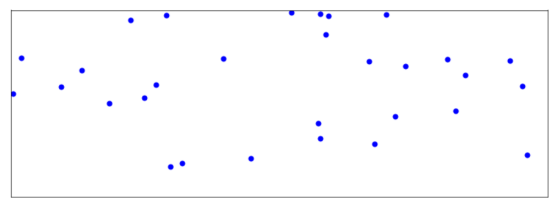
回归曲线基本符合数据点趋势

效果很棒

充足的有规律的数据







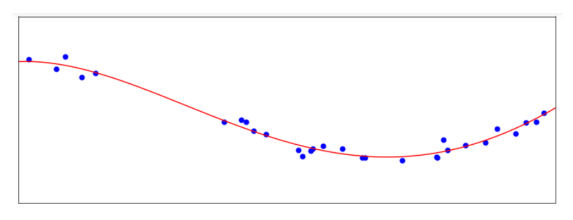
回归曲线基本符合数据点趋势

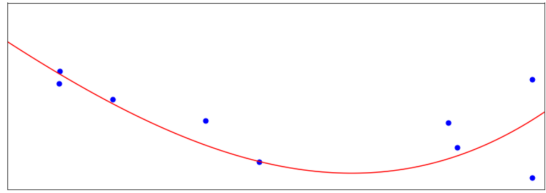
效果很棒

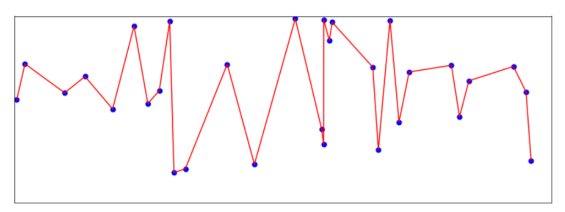
充足的有规律的数据

效果还不好说

数据不充足







回归曲线基本符合数据点趋势

效果很棒

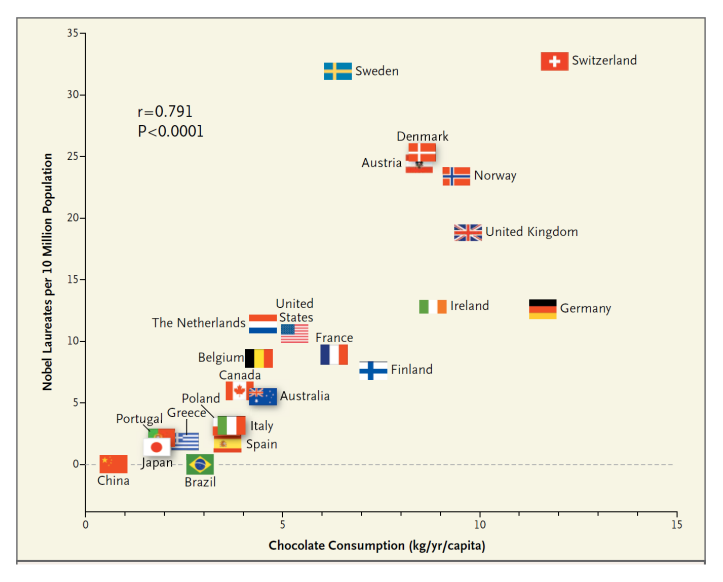
充足的有规律的数据

效果还不好说

数据不充足

回归曲线完全不可用 输入与输出之间完全没有相关性 充足的"垃圾"数据

无关数据



巧克力消费量与获得诺贝尔奖人数的关系

N. Engl. J. Med. 可不是"野鸡"杂志, 而是医学界声望非常高的杂志。



74.699

~ 5.7%

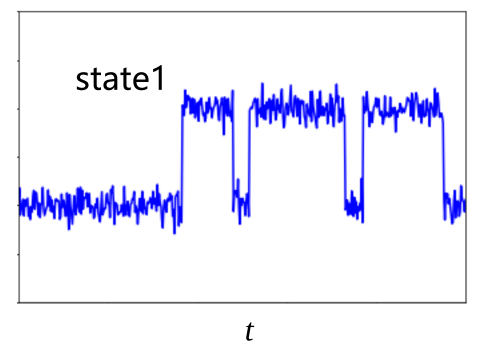
H. Franz and M. D. Messerli, *N. Engl. J. Med.* 2012, 367, 1562.

目录

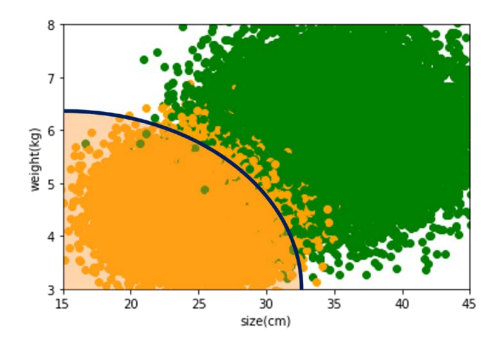
- 1. 算法简介
- 2. 回归算法
- 3. 双金属吸附中的回归
- 4. 非线性回归求速率常数
- 5. 模型评价
- 6. 决策树分类算法
- 7. 支持向量机

分类算法

在一个已知标号的样本中找到一种分类器,使其能对未知样品进行分类。

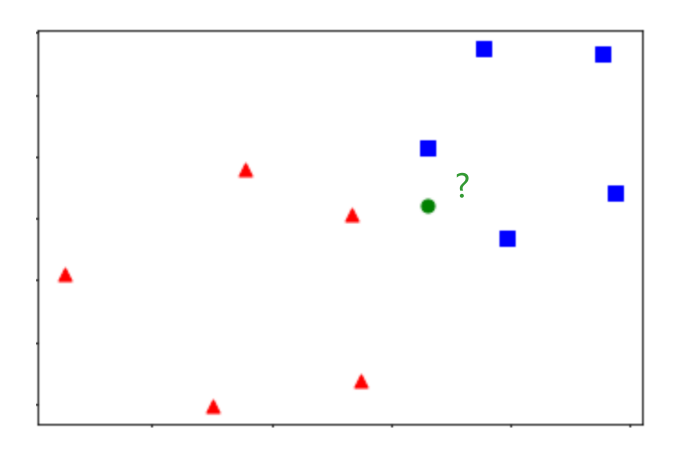


state0



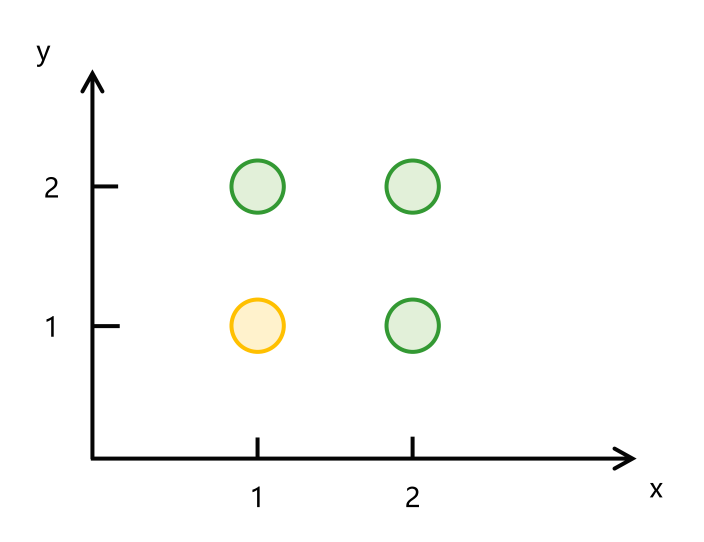
分类问题

分类问题:已知红色点为A类,蓝色点为B类,绿色点应该属于哪一类?



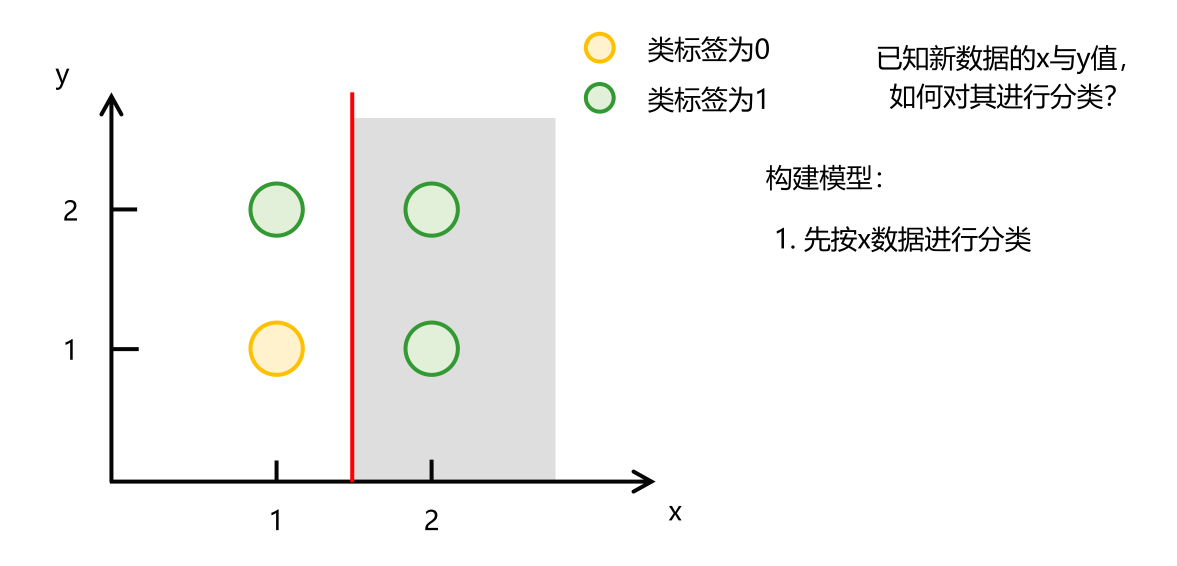
十大经典算法

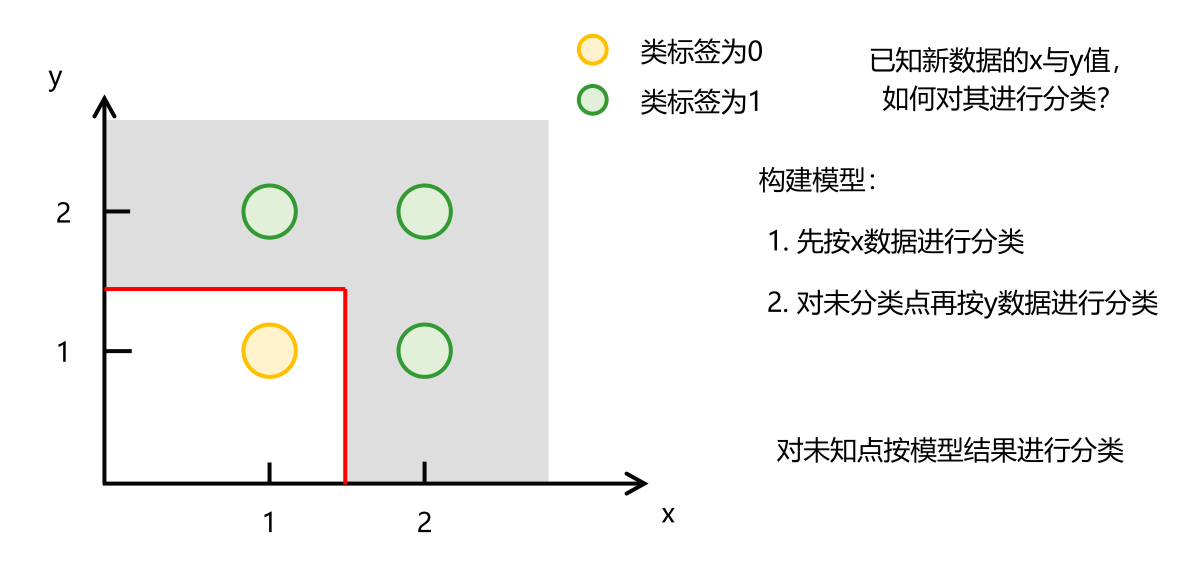
排名	算法	作者	发表年份	
1	C4.5	Quinlan, J. R.	1993	
2	k-Means	MacQueen, J. B.	1967	
3	SVM	Vapnik, V. N.	1995	
4	Apriori	Agrawal, R.	1994	
5	EM	McLachlan, G.	2000	
6	PageRank	Brin, S.	1998	
7	AdaBoost	Freund, Y.	1997	
8	kNN	Hastie, T.	1996	
9	Naïve Bayes	Hand, D. J.	2001	
10	CART	Breiman, L.	1984	

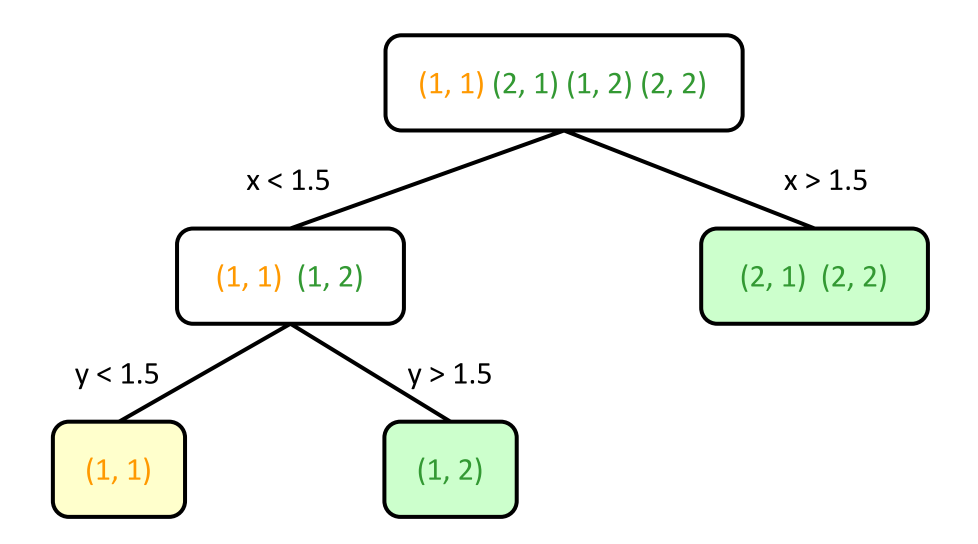


- 类标签为0
- 类标签为1

已知新数据的x与y值, 如何对其进行分类?







```
In [1]: from sklearn import tree

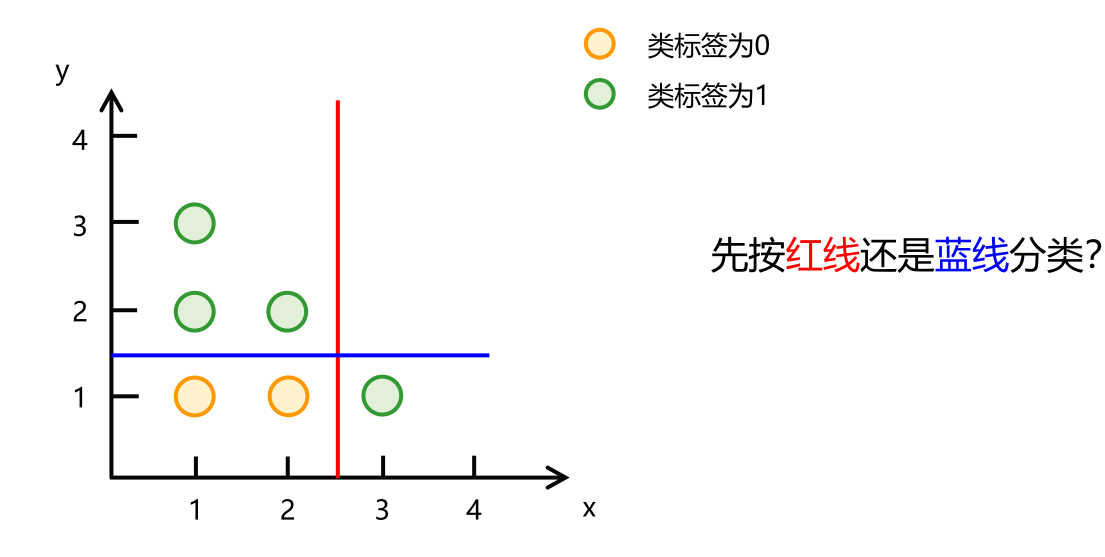
X = [[1, 1], [2, 1], [1, 2], [2, 2]]
y = [0, 1, 1, 1]
clf = tree.DecisionTreeClassifier()
clf = clf.fit(X, y)

In [2]: clf.predict([[3, 3]])
Out[2]: array([1])

In [3]: clf.predict_proba([[3, 3]])
Out[3]: array([[0., 1.]])
```

```
In [4]: import graphviz
         dot_data = tree.export_graphviz(clf, out_file = None, filled=True)
         graph = graphviz.Source(dot data)
         graph
Out[4]:
                               X[1] \le 1.5
                               gini = 0.375
                               samples = 4
                               value = [1, 3]
                            True
                                            False
                     X[0] \le 1.5
                                           gini = 0.0
                       gini = 0.5
                                          samples = 2
                     samples = 2
                                         value = [0, 2]
                     value = [1, 1]
             gini = 0.0
                                 gini = 0.0
            samples = 1
                                samples = 1
           value = [1, 0]
                               value = [0, 1]
```

划分选择



信息熵

信息熵 (information entropy):

假定当前样本集合D中第k类样本所占的比例为pk,则D的信息熵为

$$Ent(D) = -\sum p_k log_2 p_k$$

Ent(D)的值越小,D的纯度越高

假定共含有n种样本,最大熵对应的分布为

$$p_k = \frac{1}{n}$$

ID3算法

信息增益 (information gain):

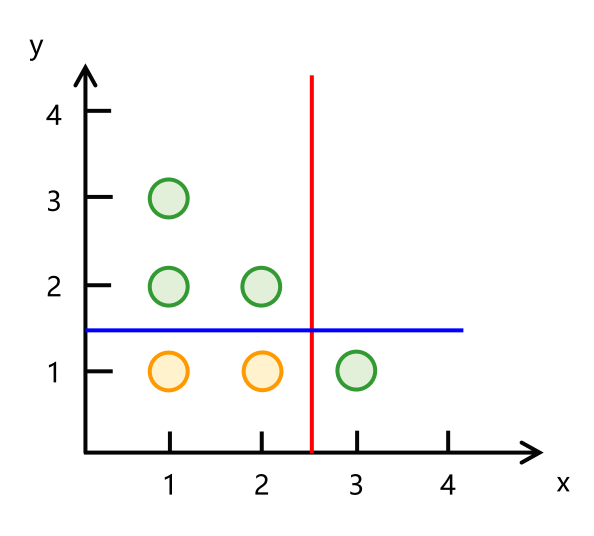
划分后样本的纯度提升

$$Gain(D,a) = Ent(D) - \sum \frac{|D^{v}|}{|D|} Ent(D^{v})$$

ID3学习算法(Quinlan, 1975):

$$a_* = \underset{a \in A}{argmax} \ Gain(D, a)$$

划分选择



划分前:

$$Ent(D) = -\frac{1}{3}log_2\frac{1}{3} - \frac{2}{3}log_2\frac{2}{3} = 0.9183$$

红线划分后:

$$Ent(D^{1}) = -\frac{2}{5}log_{2}\frac{2}{5} - \frac{3}{5}log_{2}\frac{3}{5} = 0.9709$$

$$Ent(D^2) = 0$$

$$Gain(D, a) = 0.9183 - \frac{5}{6} \times 0.9709 = 0.1092$$

蓝线划分后:

$$Ent(D^1) = -\frac{1}{3}log_2\frac{1}{3} - \frac{2}{3}log_2\frac{2}{3} = 0.9183$$

$$Ent(D^2) = 0$$

$$Gain(D, a) = 0.9183 - \frac{1}{2} \times 0.9183 = 0.4592$$

划分选择

```
In [5]: from sklearn import tree
         X = [[1, 1], [2, 1], [3, 1], [1, 2], [2, 2], [1, 3]]
         y = [0, 0, 1, 1, 1, 1]
         clf = tree. DecisionTreeClassifier(criterion = 'entropy')
         clf = clf. fit(X, y)
In [6]: dot_data = tree.export_graphviz(clf, out_file = None, filled=True)
         graph = graphviz.Source(dot_data)
         graph
Out[6]:
                                X[1] \le 1.5
                              entropy = 0.918
                                samples = 6
                               value = [2, 4]
                            True
                                            False
                      X[0] \le 2.5
                                          entropy = 0.0
                    entropy = 0.918
                                           samples = 3
                      samples = 3
                                          value = [0, 3]
                     value = [2, 1]
           entropy = 0.0
                               entropy = 0.0
            samples = 2
                                samples = 1
           value = [2, 0]
                               value = [0, 1]
```

C4.5算法

C4.5算法(Quinlan, 1993)

使用增益率进行划分

$$Gain_ratio(D, a) = \frac{Gain(D, a)}{IV(a)}$$

其中

$$IV(a) = -\sum \frac{|D^v|}{|D|} log_2 \frac{|D^v|}{|D|}$$

CART算法

基尼值

$$Gini(D) = \sum_{k} \sum_{k' \neq k} p_k p_{k'}$$
$$= 1 - \sum_{k} p_k^2$$

属性a的基尼指数

$$Gini_index(D, a) = \sum \frac{|D^v|}{|D|} Gini(D^v)$$

CART算法(Breiman, 1984)

$$a_* = \underset{a \in A}{argmin \ Gini_index(D, a)}$$

分类背景: 有机小分子

定量构效关系(quantitative structure-activity relationships, QSAR)

药物、农药、化学毒剂等研究

麻醉化合物小分子通常被分为两类,他们的毒性机理有较大的区别

第一类: 非极性麻醉剂

第二类:极性麻醉剂

Table 7 Chemical Compounds, Theoretical Descriptors (E_{HOMO} , E_{LUMO} and Q⁻), and Mechanism of Toxic Action (nonpolar, class +1; polar, class -1)

No	Compound	Еномо	E_{LUMO}	Q^-	MOA	Class
1	tetrachloroethene	-9.902	-0.4367	-0.0372	1	+1
2	1,2-dichloroethane	-11.417	0.6838	-0.1151	1	+1
3	1,3-dichloropropane	-11.372	1.0193	-0.1625	1	+1
4	dichloromethane	-11.390	0.5946	-0.1854	1	+1
5	1,2,4-trimethylbenzene	-8.972	0.5030	-0.2105	1	+1
6	1,1,2,2-tetrachloroethane	-11.655	-0.0738	-0.2785	1	+1
7	2,4-dichloroacetophenone	-9.890	-0.5146	-0.4423	1	+1
8	4-methyl-2-pentanone	-10.493	0.8962	-0.4713	1	+1
•	.1 1	44 000	4.4250	0.5045	4	. 4

决策树分类

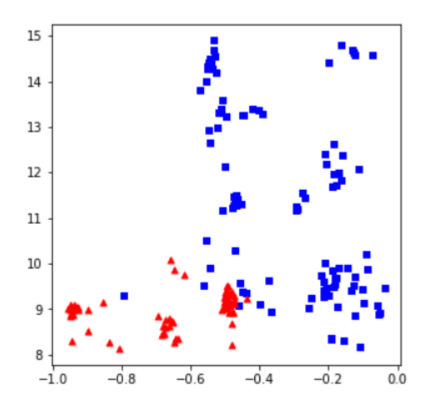
Out[1]:

	Compound	log(Kow)	EHOMO	Q-	ELUMO	Q+	Class
0	methanol	-0.77	-11.135	-0.5353	3.7775	0.3182	1
1	ethanol	-0.31	-11.050	-0.5360	3.6513	0.3107	1
2	1-propanol	0.25	-10.940	-0.5317	3.6324	0.3122	1
3	2-propanol	0.05	-10.895	-0.5469	3.4925	0.3166	1
4	1-butanol	0.88	-10.940	-0.5422	3.5041	0.3141	1
185	2-nitroaniline	1.85	-9.068	-0.6488	-0.7937	0.3510	2
186	3-nitroaniline	1.37	-9.254	-0.9468	-0.9503	0.3922	2
187	4-nitroaniline	1.39	-9.160	-0.6493	0.7050	0.3134	2
188	2-chloro-4-nitroaniline	2.06	-9.256	-0.6434	-0.9066	0.3183	2
189	4-ethoxy-2-nitroaniline	2.38	-8.994	-0.8070	-0.8747	0.3969	2

190 rows × 7 columns

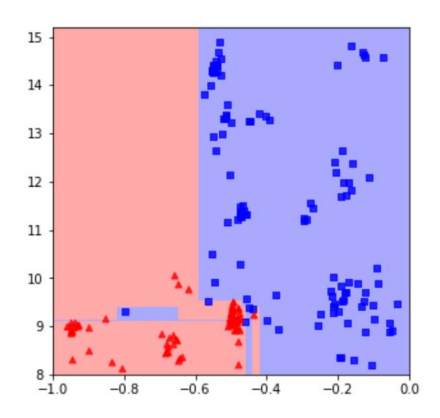
决策树分类

```
In [2]: df['Gap'] = df['ELUMO'] - df['EHOMO']
In [3]: X = df[['Q-','Gap']]
         y = df['Class']. values
In [4]: from sklearn import tree
         clf_tree = tree. DecisionTreeClassifier()
         clf_tree.fit(X, y)
   [5]: import matplotlib.pyplot as plt
         plt. figure (figsize=(5, 5))
         x1 = df. iloc[:114, 3]
         y1 = df. iloc[:114, 7]
         x2 = df. iloc[114:, 3]
         y2 = df. iloc[114:, 7]
         plt. scatter(x1, y1, c='b', marker='s', s=25)
         plt. scatter(x2, y2, c='r', marker='^', s=25)
```



决策树分类

```
[6]: import numpy as np
         xx, yy = np. meshgrid (np. arange (-1.0, 0.1, 0.01), np. arange (8, 15.2, 0.01))
         coords = np. stack((xx. reshape(-1), yy. reshape(-1)), axis = 1)
In [7]: Z = clf tree. predict (coords)
         Z = Z. reshape (xx. shape)
    [8]: from matplotlib.colors import ListedColormap
         light_rgb = ListedColormap(['#AAAAFF', '#FFAAAA'])
         plt.figure(figsize=(5,5))
         plt.pcolormesh(xx, yy, Z, cmap = light_rgb)
         plt. scatter(x1, y1, c='b', marker='s', s=25, alpha=0.8)
         plt. scatter(x2, y2, c='r', marker='^', s=25, alpha=0.8)
         plt. axis ((-1.0, 0.0, 8, 15.2))
```



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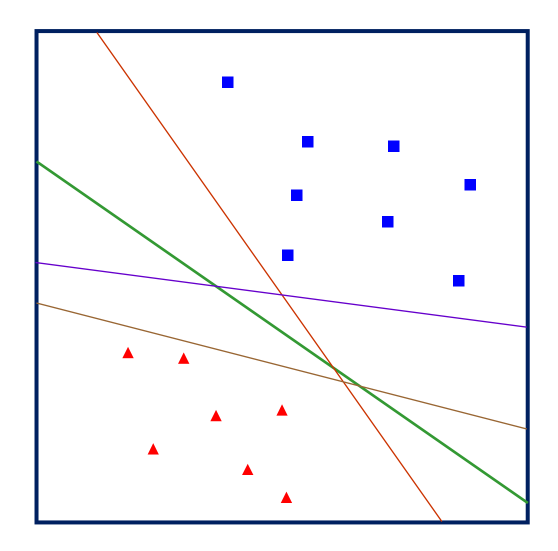
支持向量机(Support Vector Machines)

问题: 最佳分类方式?

1963 V. N. Vapnik and A. Y. Chervonenkis 原始支持向量机

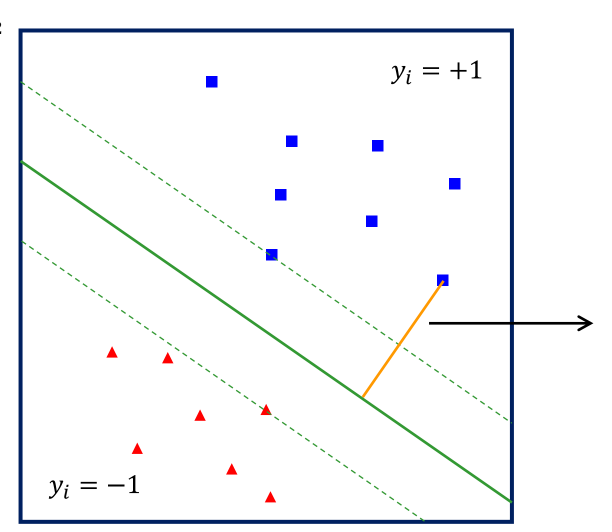
1992 B. E. Boser, I. M. Guyon and V. N. Vapnik 核技巧

1995 C. Cortes and V. N. Vapnik 软间隔



划分超平面

 x_2 划分超平面 $\mathbf{w}^T \mathbf{x} + b = 0$ $\boldsymbol{w} = (w_1, w_2)$ $\boldsymbol{x} = (x_1, x_2)$



点到超平 面的距离

$$r = \frac{|\boldsymbol{w}^T \boldsymbol{x} + b|}{\|\boldsymbol{w}\|}$$

支持向量

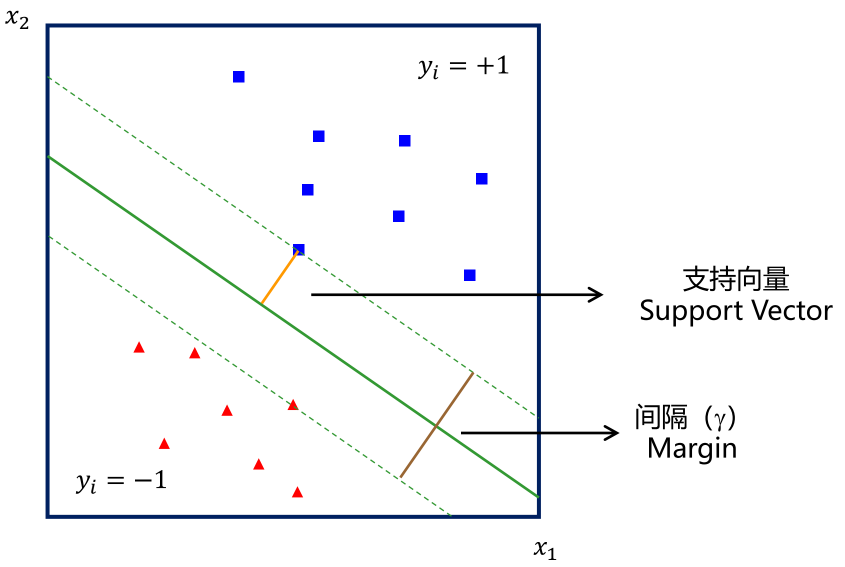
\$

$$\begin{cases} \mathbf{w}^T \mathbf{x} + b \ge 1, & y_i = +1 \\ \mathbf{w}^T \mathbf{x} + b \le -1, & y_i = -1 \end{cases}$$

等号在距离最近的点满足。

支持向量到超平面的距离 之和

$$\gamma = \frac{2}{\|\boldsymbol{w}\|}$$



支持向量机(Support Vector Machines)

寻找具有最大间隔的划分超平面

$$\max_{\mathbf{w}, b} \frac{2}{\|\mathbf{w}\|} \quad s.t. \ y_i(\mathbf{w}^T \mathbf{x} + b) \ge 1, \quad i = 1, 2, 3, \dots, m.$$

支持向量机

$$\min_{\mathbf{w},b} \frac{1}{2} \|\mathbf{w}\|^2 \quad s.t. \quad y_i(\mathbf{w}^T \mathbf{x} + b) \ge 1, \qquad i = 1,2,3,\cdots, m.$$

对偶问题

添加拉格朗日乘子 $\alpha_i \geq 0$

拉格朗日函数为

$$L(\mathbf{w}, b, \alpha) = \frac{1}{2} \|\mathbf{w}\|^2 + \sum_{i=1}^{n} \alpha_i (1 - y_i(\mathbf{w}^T \mathbf{x}_i + b))$$

L对w, b偏导为0

$$\mathbf{w} = \sum \alpha_i y_i \mathbf{x}_i$$

$$0 = \sum \alpha_i y_i$$

对偶问题为

$$\max_{\alpha} (\sum \alpha_i - \frac{1}{2} \sum \sum \alpha_i \alpha_j y_i y_j \mathbf{x}_i^T \mathbf{x}_j) \qquad s.t. \sum \alpha_i y_i = 0,$$

$$\alpha_i \geq 0, \qquad i = 1, 2, \cdots, m.$$

KKT条件

模型变为

$$f(\mathbf{x}) = \mathbf{w}^T \mathbf{x} + b$$
$$= \sum_{i=1}^{n} \alpha_i y_i \mathbf{x}_i^T \mathbf{x} + b$$

KKT (Karush-Kuhn-Tucker) 条件

$$\begin{cases} \alpha_i \ge 0; \\ y_i f(x_i) - 1 \ge 0; \\ \alpha_i (y_i f(x_i) - 1) = 0. \end{cases}$$

SMO算法

SMO (Sequential Minimal Optimization)

选取一对需更新的变量 α_i 和 α_j

固定 α_i 和 α_i 以外的参数,求解对偶问题更新 α_i 和 α_i

仅考虑 α_i 和 α_i 时,对偶问题的约束条件为

$$a_i y_i + a_j y_j = c, \qquad a_i \ge 0, a_j \ge 0$$

其中

$$c = -\sum_{k \neq i,j} a_k y_k$$

代入对偶问题,得到单变量二次规划问题,可以轻易解出

b的确定

对于任意支持向量 (x_s, y_s)

$$y_{S}\left(\sum_{i\in S}\alpha_{i}y_{i}\boldsymbol{x}_{i}^{T}\boldsymbol{x}_{S}+b\right)=1, \qquad S=\left\{i|\alpha_{i}>0, i=1,2,\cdots,m\right\}$$

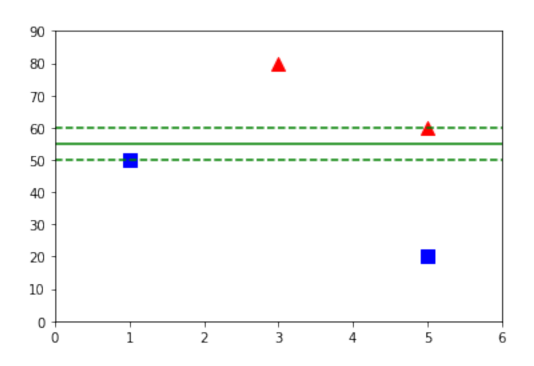
取任意支持向量 (x_s, y_s) 求解即可得到b

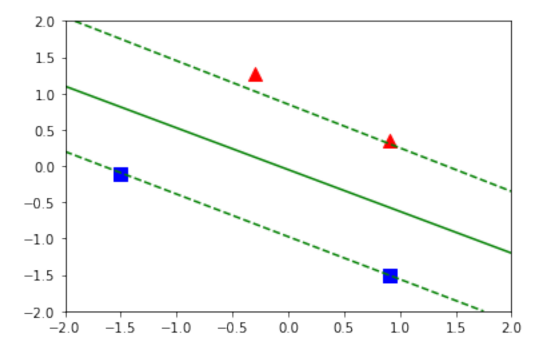
还可通过下式求解

$$b = \frac{1}{|S|} \sum_{S \in S} (y_S - \sum_{i \in S} \alpha_i y_i \mathbf{x}_i^T \mathbf{x}_S)$$

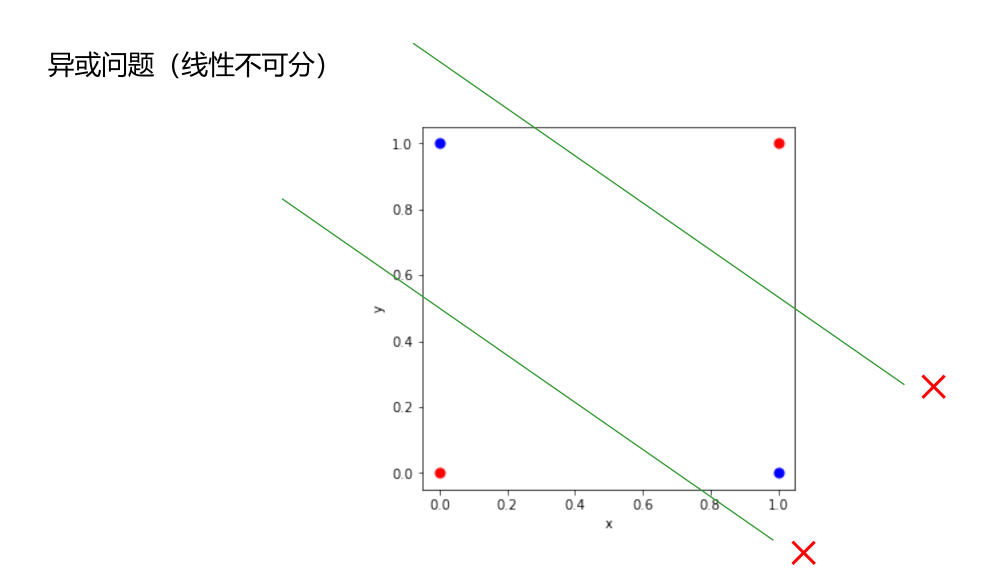
特征缩放

SVM对特征缩放敏感

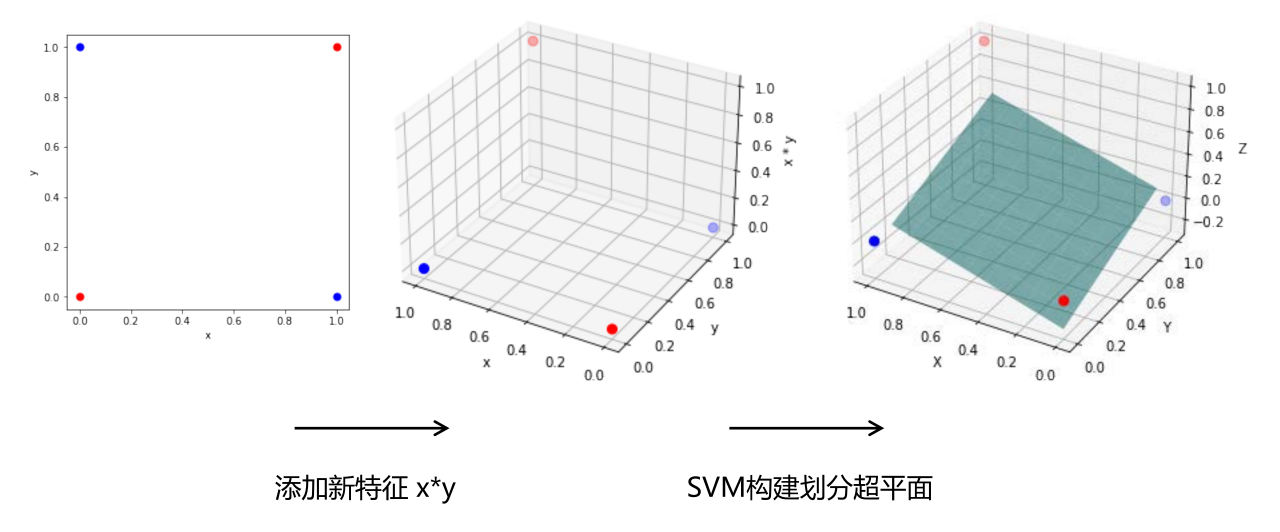




非线性问题



高维特征空间



高维空间对偶问题

 $\phi(x)$ 表示映射函数,则划分超平面对应的模型可表示为

$$f(\mathbf{x}) = \mathbf{w}^T \phi(\mathbf{x}) + b$$

类似的,我们需要

$$\min_{\mathbf{w},b} \frac{1}{2} \|\mathbf{w}\|^2 \quad s.t. \quad y_i(\mathbf{w}^T \mathbf{x} + b) \ge 1, \qquad i = 1,2,3,\cdots, m.$$

对偶问题是

$$\max_{\alpha} (\sum \alpha_i - \frac{1}{2} \sum \sum \alpha_i \alpha_j y_i y_j \phi(\mathbf{x}_i)^T \phi(\mathbf{x}_j)) \qquad s.t. \sum \alpha_i y_i = 0,$$

$$\alpha_i \geq 0, \qquad i = 1, 2, \cdots, m.$$
 高维特征空间内积

核技巧

构建核函数

$$\kappa(\mathbf{x}_i, \mathbf{x}_j) = \langle \phi(\mathbf{x}_i), \phi(\mathbf{x}_j) \rangle = \phi(\mathbf{x}_i)^T \phi(\mathbf{x}_j)$$

对偶问题

$$\max_{\alpha} \sum \alpha_i - \frac{1}{2} \sum \sum \alpha_i \alpha_j y_i y_j \kappa(\mathbf{x}_i, \mathbf{x}_j) \qquad s. t. \sum \alpha_i y_i = 0,$$

$$\alpha_i \ge 0, \qquad i = 1, 2, \dots, m.$$

求解后即可得到

$$f(\mathbf{x}) = \sum \alpha_i y_i \kappa(\mathbf{x}_i, \mathbf{x}_j) + b$$

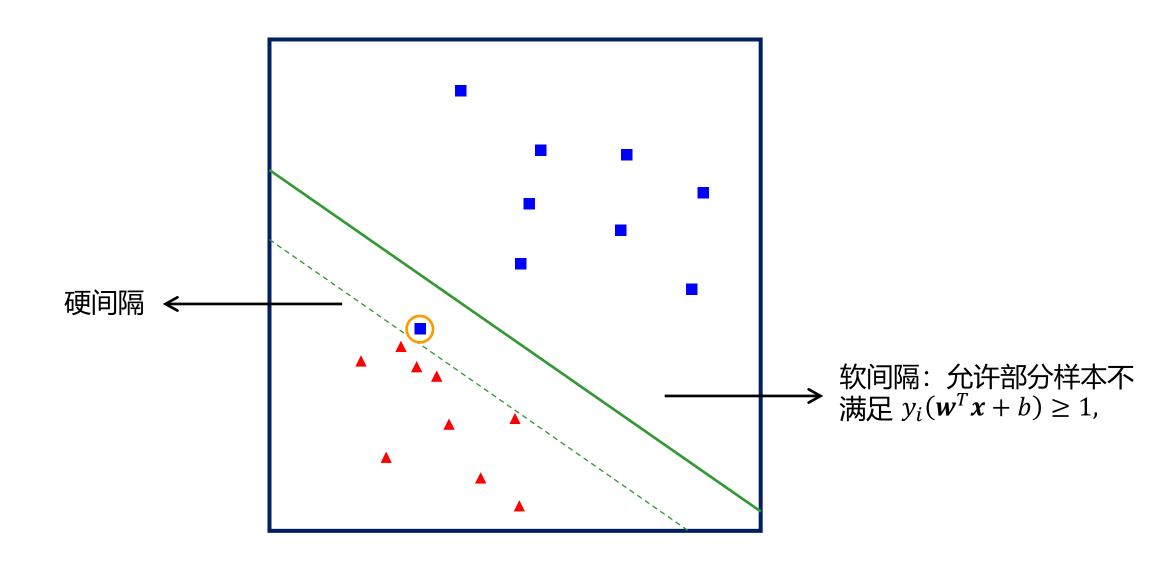
常用核函数

scikit-learn中的核函数

The *kernel function* can be any of the following:

- linear: $\langle x, x' \rangle$.
- polynomial: $(\gamma\langle x,x'\rangle+r)^d$, where d is specified by parameter degree, r by coef0.
- rbf: $\exp(-\gamma ||x-x'||^2)$, where γ is specified by parameter gamma, must be greater than 0.
- sigmoid $\tanh(\gamma\langle x,x'\rangle+r)$, where r is specified by coef0.

软间隔



软间隔支持向量机

优化目标改为

$$\min \frac{1}{2} \|\mathbf{w}\|^2 + C \sum \iota_{0/1} (1 - y_i(\mathbf{w}^T \mathbf{x}_i + b))$$

C为惩罚系数, $\iota_{0/1}$ 为0/1损失函数

$$\iota_{0/1}(z) = \begin{cases} 1, & if \ y_i(\mathbf{w}^T \mathbf{x}_i + b) < 0; \\ 0, & otherwise. \end{cases}$$

hinge损失函数

$$\iota_{hinge}(z) = \max(0, 1 - y_i(\mathbf{w}^T \mathbf{x}_i + b))$$

优化目标

$$\min \frac{1}{2} ||\mathbf{w}||^2 + C \sum \max(0, 1 - y_i(\mathbf{w}^T \mathbf{x}_i + b))$$

松弛变量 $\xi_i \geq 0$

$$\min_{\mathbf{w}, b, \xi_i} \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i} \xi_i \qquad s.t. \quad y_i(\mathbf{w}^T \mathbf{x}_i + b) \ge 1 - \xi_i,$$

$$\xi_i \ge 0, \qquad i = 1, 2, \cdots, m.$$

支持向量机(Support Vector Machines)

```
In [9]: from sklearn. pipeline import make_pipeline
          from sklearn.preprocessing import StandardScaler
          from sklearn. svm import SVC
          clf_svm = make_pipeline(StandardScaler(), SVC(kernel = 'linear'))
          clf_svm.fit(X, y)
 Out[9]: Pipeline(steps=[('standardscaler', StandardScaler()),
                           ('svc', SVC(kernel='linear'))])
In [10]: Z = clf_svm. predict(coords)
          Z = Z. reshape (xx. shape)
   [11]: | light_rgb = ListedColormap(['#AAAAFF', '#FFAAAA'])
          plt. figure (figsize=(5, 5))
          plt.pcolormesh(xx, yy, Z, cmap = light_rgb)
          plt. scatter(x1, y1, c='b', marker='s', s=25, alpha=0.8)
          plt. scatter(x2, y2, c='r', marker='^', s=25, alpha=0.8)
          plt. axis ((-1.0, 0.0, 8, 15.2))
```

