In [1]:

```
# This line will add a button to toggle visibility of code blocks,
# for use with the HTML export version
from IPython.core.display import HTML
HTML('''<button style="margin:0 auto; display: block;" onclick="jQue
ry('.code_cell .input_area').toggle();
    jQuery('.prompt').toggle();">Toggle code</button>''')
```

Out[1]:

Toggle code



Week 8 - Support Vector Machines

Dr. David Elliott

- 1. Support Vector Classifier (SVC)
- 2. Support Vector Machine (SVM)

Common Notation

- X is a matrix containing all the feature values of all the observations
- *n* is the number of observations in the dataset
- x_i is a vector of all the feature values (except the label) of the ith instance in the dataset.
- y_i is the label (desired model output) of the *i*th instance in the dataset.
- p is the number of features in the dataset
- x_i is a vector of all the observations values of the jth feature in the dataset.

Notes

We can't always perfectly separate the data with a p-1 dimensional hyperplane. To overcome this problem we could either:

- Tweak the constraints on the hyperplane to allow some points to be misclassified (soft margin),
- Transform the data to be separable by a hyperplane in another space (kernel method).

In [2]:

```
%matplotlib inline
import os # locating directories
import numpy as np # Arrays
```

```
import pandas as pd # DataFrames
# Plotting
import matplotlib
import matplotlib.pyplot as plt
matplotlib.rcParams['animation.embed limit'] = 30000000.0
import seaborn as sns; sns.set()
from sklearn.datasets import load iris # for the Iris data
from IPython.display import Image
                                                  # displaying .png i
mages
                                                  # SVM
from sklearn.svm import SVC, LinearSVC
from mpl toolkits.mplot3d import Axes3D
                                            # 3d plots
from sklearn.preprocessing import StandardScaler # scaling features
from sklearn.preprocessing import LabelEncoder # binary encoding
from sklearn.pipeline import Pipeline
                                                  # combining classif
ier steps
from sklearn.preprocessing import PolynomialFeatures # make Polynomi
alFeatures
from sklearn.datasets import make classification, make moons # make
example data
import itertools
from time import time
import joblib # saving models
import warnings # prevent warnings
# colours for print()
class color:
   PURPLE = '\033[95m'
  CYAN = ' \setminus 033[96m']
  DARKCYAN = ' \setminus 033[36m']
  BLUE = ' \setminus 033[94m']
  GREEN = ' \ 033[92m']
  YELLOW = ' \setminus 033[93m']
  RED = ' \setminus 033[91m']
  BOLD = ' \setminus 033[1m']
  UNDERLINE = '\033[4m'
  END = '\033[0m'
image dir = os.path.join(os.getcwd(),"Images")
# Initial fig number
fig num=15
plt.rcParams['figure.dpi'] = 120
# golden ratio for figures ()
gr = 1.618
height pix = 500
width pix = height pix*gr
height inch = 4
width inch = height inch*gr
```

```
# if trying to make PDF
PDF=False
```

In [3]:

```
iris = load iris(as frame=True) # data stored in a `sklearn.utils.B
iris df = iris['data']
                                # get features DataFrame
target = iris['target']
                                # get target Series
# get the labels of flowers capitalised for visualisation
target names = list(map(lambda s: s.capitalize(), iris['target names
']))
# create a dictionary with the original labels decoded (inverse of L
abelEncoder)
decode label = dict(zip(range(3), target names))
# make a label encoder to use later if needed
le = LabelEncoder().fit(target names)
# add the target labels to df for visualisation purposes
iris vis = pd.concat([iris df, target],axis=1)
# turn the ints to labels
iris vis["target"] = iris vis["target"].replace(decode label)
# Capitalize column names for plotting
iris vis.columns = [x.capitalize() for x in list(iris vis.columns)]
# reduce the data for example
X AX LABEL = "Petal length (cm)"
Y AX LABEL = "Petal width (cm)"
REMOVE = "Virginica"
iris reduced = iris vis[[X AX LABEL, Y AX LABEL, "Target"]]
iris reduced = iris reduced[iris reduced.Target != REMOVE]
```

In [4]:

```
# Centered figures in the notebook and presentation
# ...was a real pain to find this:
# https://gist.githubusercontent.com/maxalbert/800b9f06c7b2dd365ea5

import matplotlib.pyplot as plt
import numpy as np
import urllib
import base64
from io import BytesIO, StringIO

def fig2str(fig, format='svg'):
    """
    Return a string containing the raw data of the matplotlib figure in the given format.
```

```
assert isinstance(fig, matplotlib.figure.Figure)
   imgdata = BytesIO()
   fig.savefig(imgdata, format=format, bbox inches='tight')
   imgdata.seek(0) # rewind the data
   output = imgdata.getvalue()
   if format == 'svq':
        return output
   else:
        return urllib.parse.quote(base64.b64encode(output))
class MatplotlibFigure(object):
    Thin wrapper around a matplotlib figure which provides a custom
   HTML representation that allows tweaking the appearance
    11 11 11
   def init (self, fig, centered=False):
        assert isinstance(fig, matplotlib.figure.Figure)
        self.centered = centered
   def repr html (self):
        img str png = fig2str(fig, format='png')
        uri = 'data:image/png;base64,' + img str png
        html repr = "<img src='{}'>".format(uri)
        if self.centered:
            html repr = "<center>" + html repr + "</center>"
        return html repr
```

1.4. Support Vector Classifier (SVC)

SVC's are a generalisation and extension of the maximal margin classifier so it can be applied to a broader range of cases¹.

In practice they are more robust to individual observations and better classify most training observations than the Maximal Margin Classifier. This is because they take the approach it is better to missclassify some training examples in order to do a better job classifying the rest.

This is called a *soft margin* as it allows some violations by the training data by a small subset of training observation, not only on the wrong side of the margin, but wrong side of the hyperplane.

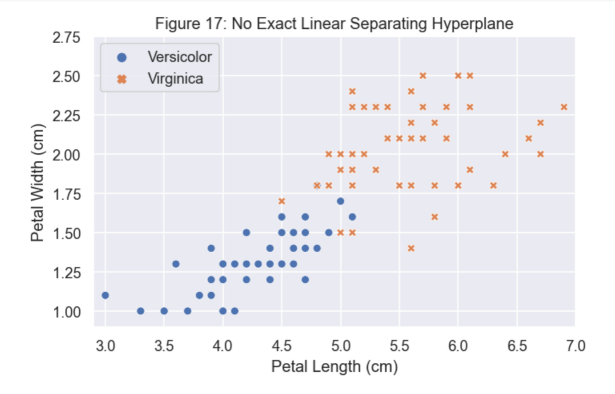
Notes

• "Developed in the computer science community in the 1990s"²

```
In [6]:
```

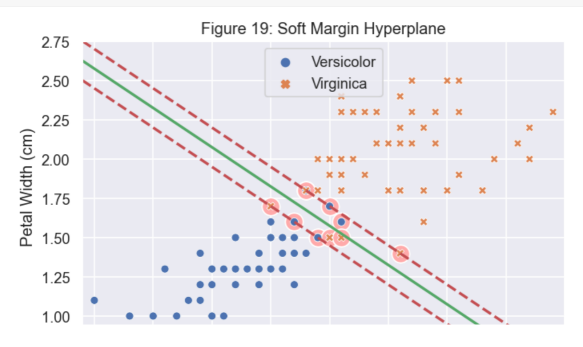
```
def svc_decision_boundary(clf, xmin=0, xmax=5.5, highlight=True, axe
s_limit = [0, 5.5, 0, 2]):
    w = clf.coef_[0]
    b = clf.intercept_[0]
```

```
# At the decision boundary, w0*x0 + w1*x1 + b = 0
    \# => x1 = -w0/w1 * x0 - b/w1
    x0 = np.linspace(xmin, xmax, 200)
    decision boundary = -w[0]/w[1] * x0 - b/w[1]
   margin = 1/w[1]
    gutter up = decision boundary + margin
    gutter down = decision boundary - margin
    svs = clf.support vectors
    if highlight:
        g = sns.scatterplot(x = svs[:, 0], y = svs[:, 1], s=180, fac
ecolors='#FFAAAA')
    plt.plot(x0, decision boundary, "g-", linewidth=2)
    plt.plot(x0, gutter up, "r--", linewidth=2)
    plt.plot(x0, gutter down, "r--", linewidth=2)
    plt.axis(axes limit)
def soft margin(title, hyperplane=False):
   virgin versi = iris vis[["Petal length (cm)", "Petal width (cm)"
, "Target"]]
    virgin versi = virgin versi[virgin versi.Target != "Setosa"]
    X = virgin versi[["Petal length (cm)", "Petal width (cm)"]].valu
es
    y = virgin versi[["Target"]].replace({'Versicolor':0, 'Virginica
':1}).values.ravel()
    if hyperplane:
        svm clf = SVC(kernel="linear", C=100)
        svm clf.fit(X, y)
        svc decision boundary (svm clf, 2.9, 7)
    labels = virgin versi[["Target"]].values.ravel()
    sns.scatterplot(x = X[:,0], y = X[:,1], hue=labels, style = labe
ls)
    plt.axis([2.9, 7, 0.9, 2.75])
    plt.title(title)
    plt.xlabel("Petal Length (cm)")
    plt.ylabel("Petal Width (cm)")
fig num+=1
fig = plt.figure(figsize=(width inch, height inch))
soft margin("Figure %d: No Exact Linear Separating Hyperplane"%fig n
um)
if PDF:
    plt.show()
   plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```



In [8]:

```
fig_num+=1
fig = plt.figure(figsize=(width_inch, height_inch))
soft_margin("Figure %d: Soft Margin Hyperplane"%fig_num, hyperplane=
True)
plt.savefig(os.path.join(image_dir, "Soft_Margin_Hyperplane.png"))
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```



We want to relax the following constraints when necessary:

$$\mathbf{w}^{\mathrm{T}}\mathbf{x}_{i} + b \geq 1 \text{ for } y_{i}$$

$$= 1,$$

$$\mathbf{w}^{\mathrm{T}}\mathbf{x}_{i} + b \leq$$

$$-1 \text{ for } y_{i} = -1$$

This can be done by introducing positive slack variables $\xi_i, i=1,\dots,n$ in the constraints 5,6,10.

$$egin{aligned} \mathbf{w}^{\mathrm{T}} \mathbf{x}_{i} + b &\geq 1 - \xi_{i} \\ & ext{if } y_{i} = 1, \\ \mathbf{w}^{\mathrm{T}} \mathbf{x}_{i} + b &\leq -1 \\ + \xi_{i} & ext{if } y_{i} = -1, \\ \xi_{i} &\geq 0 & \forall_{i}. \end{aligned}$$

Notes

- Slack variable ξ_1, \dots, ξ_n allow individual observations to be on the wrong side of the margin or hyperplane.
- $\sum_i \xi_i$ is an upper bound on the number of training errors.
- ullet ξ_i tells us where the ith observation is located relative to the hyperplane; $\xi_i=0$ being on the correct side of the margin, $\xi_i>0$ being on the wrong side of the margin, and $\xi_i>1$ on the wrong side of the hyperplane.
- $\xi_1=\ldots=\xi_n=0$ is the maximal margin hyperplane optimisation. Test observations are classified as before,

$$f(x^*) = eta_0 + eta_1 x_1^* + \dots + eta_p x_p^*$$

Tuning Parameter (C)

To ensure there is a penelty, C, for relaxing the constraint, we can change our objective function to be minimised from $\frac{1}{2}||\mathbf{w}||^2$ to,

minimise
$$\sum_{\substack{\mathbf{w},b,\xi}}^{n} \sum_{i=1}^{n} (1)$$

$$\mathbf{w},b,\xi = \frac{1}{2}||\mathbf{w}||^{2} + C = 1, (2)$$
subject to
$$y_{i}(\mathbf{w}^{T}\mathbf{x}_{i} + b) \geq 1$$

$$-\xi_{i}, \quad \xi_{i} \geq 0, \quad \forall_{i}.$$

C is a tuning parameter that controls the bias-variance trade-off¹.

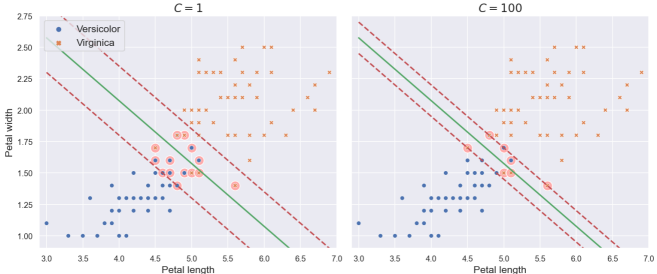
The strength of the regularization is inversely proportional to C, meaning a large C has a larger error penalty.

```
In [9]:
```

```
def soft margin(c list, main title):
   virgin versi = iris vis[["Petal length (cm)", "Petal width (cm)"
, "Target"]]
   virgin versi = virgin versi[virgin versi.Target != "Setosa"]
   X = virgin versi[["Petal length (cm)", "Petal width (cm)"]].valu
es
   y = virgin versi[["Target"]].replace({'Versicolor':0, 'Virginica
':1}).values.ravel()
    scaler = StandardScaler()
    svm clf1 = LinearSVC(C=c list[0], loss="hinge", random state=42)
    svm clf2 = LinearSVC(C=c list[1], loss="hinge", random state=42)
   scaled svm clf1 = Pipeline([
            ("scaler", scaler),
            ("linear svc", svm clf1),
        ])
    scaled svm clf2 = Pipeline([
            ("scaler", scaler),
            ("linear svc", svm clf2),
        ])
    scaled svm clf1.fit(X, y)
    scaled svm clf2.fit(X, y)
    # Convert to unscaled parameters
    b1 = svm clf1.decision function([-scaler.mean / scaler.scale ])
    b2 = svm clf2.decision function([-scaler.mean / scaler.scale ])
    w1 = svm clf1.coef [0] / scaler.scale
    w2 = svm clf2.coef [0] / scaler.scale
    svm_clf1.intercept_ = np.array([b1])
    svm clf2.intercept = np.array([b2])
    svm_clf1.coef_ = np.array([w1])
    svm clf2.coef = np.array([w2])
    # Find support vectors (LinearSVC does not do this automatically
    t = y * 2 - 1
    support vectors idx1 = (t * (X.dot(w1) + b1) < 1).ravel()
    support vectors idx2 = (t * (X.dot(w2) + b2) < 1).ravel()
    svm clf1.support vectors = X[support vectors idx1]
    svm clf2.support vectors = X[support vectors idx2]
    fig, axes = plt.subplots(ncols=2, figsize=(width inch*2, height
inch*1.5), sharey=True)
    plt.sca(axes[0])
    svc decision boundary (svm clf1, 3, 7)
    legend labels = virgin versi[["Target"]].values.ravel()
```

```
sns.scatterplot(x = X[:,0], y = X[:,1], hue=legend labels, style
= legend labels)
    plt.xlabel("Petal length", fontsize=14)
    plt.ylabel("Petal width", fontsize=14)
    plt.legend(loc="upper left", fontsize=14)
    plt.title("$C = {}$".format(svm clf1.C), fontsize=16)
    plt.axis([2.9, 7, 0.9, 2.75])
    plt.sca(axes[1])
    svc decision boundary (svm clf2, 3, 7)
    g = sns.scatterplot(x = X[:,0], y = X[:,1], hue=legend labels, s
tyle = legend labels, legend=False)
    plt.xlabel("Petal length", fontsize=14)
    plt.title("$C = {}$".format(svm clf2.C), fontsize=16)
    plt.axis([2.9, 7, 0.9, 2.75])
    plt.suptitle(main title)
    plt.tight layout()
    return fig
fig num+=1
fig = soft margin([1,100], "Figure %d: Tuning Parameter `C`"%fig num
if PDF:
    plt.show()
    plt.close()
else:
   plt.close()
    display(MatplotlibFigure(fig, centered=True))
```





Extra

Neither the ξ_i or their Lagrange multipliers appear in the Wolfe dual problem. This means we now have⁶:

$$egin{aligned} \max L_D &\equiv \sum_i \; lpha_i \ -rac{1}{2} \sum_{i,k}^n lpha_i lpha_k y_i y_k \mathbf{x}_i \ \cdot \mathbf{x}_k \quad ext{s.t.} \quad 0 \leq lpha_i \ \leq C, \quad \sum_i lpha_i y_i = 0 \end{aligned}$$

This also has the same solution as before:

$$\hat{\mathbf{w}} = i = 1 \frac{N_S}{\alpha_i y_i \mathbf{x}_i}.$$

Also, sometimes C is defined as $C = \frac{1}{\nu N}$ where $0 < \nu \le 1$ controls the fraction of misclasified points during the training phase⁷.

Notes

• Alike to maximal margin classifiers, SVC's only rely on a few observations, those on the margin or those that violate the margin (*Support Vectors*).

If they are on the correct side of the margin they dont change the classifier.

- This does mean that they are robust to observations far away from the hyperplane.
- When *C* is large we have narrow margins rarely violated, but highly fit to the training data (low bias-high variance).
- Coversely, when smaller it is less strict about missclassification errors, meaning the margin is wider (high bias-low variance).
- Like most hyper-parameters, it is often chosen using cross-validation.

1.5. Support Vector Machine (SVM)

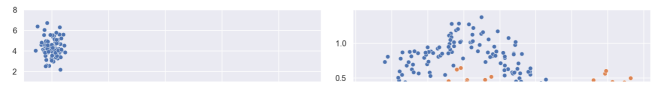
Aims to address the situation where the boundary between two classes is not linear.

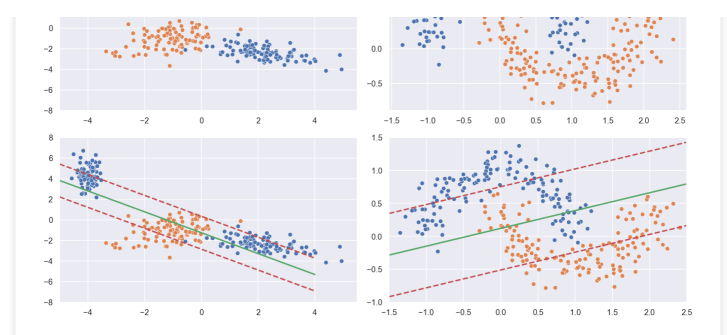
```
In [10]:
```

```
from sklearn.preprocessing import PolynomialFeatures
import warnings
nl1, y1 = make classification(n samples=300, n features=2,
                               n redundant=0, n informative= 2,
                              n classes=3, n clusters per class=1,
                               random state=14, class sep=30,
                               hypercube=False)
y1 = np.where(y1==1,0,y1)
y1 = np.where(y1==2,1,y1)
n12, y2 = make moons (n samples=300, noise=0.15, random state=42)
def non linear examples(main title):
    fig, axes = plt.subplots(ncols=2, nrows=2, figsize=(width inch*2
, height inch*2))
    axes = axes.flatten()
    plt.sca(axes[0])
    sns.scatterplot(nl1[:,0], nl1[:,1], y1, legend=False)
    plt.axis([-5, 5.5, -8, 8])
```

```
plt.sca(axes[1])
    sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
    svm clf1 = LinearSVC(random state=42).fit(nl1, y1)
    svm clf2 = LinearSVC(random state=42).fit(nl2, y2)
    # Convert to unscaled parameters
    b1 = svm clf1.intercept [0]
    b2 = svm clf2.intercept [0]
    w1 = svm clf1.coef [0]
    w2 = svm clf2.coef [0]
    # Find support vectors (LinearSVC does not do this automatically
    t1 = v1 * 2 - 1
    t2 = y2 * 2 - 1
    support vectors idx1 = (t1 * (nl1.dot(w1) + b1) < 1).ravel()
    support vectors idx2 = (t2 * (nl2.dot(w2) + b2) < 1).ravel()
    svm_clf1.support_vectors_ = nl1[support_vectors_idx1]
    svm clf2.support vectors = nl2[support vectors idx2]
    plt.sca(axes[2])
    svc decision boundary(svm clf1, -5, 4, highlight=False, axes lim
it = [-5, 5.5, -8, 8]
    sns.scatterplot(nl1[:,0], nl1[:,1], y1, legend=False)
    plt.sca(axes[3])
    svc decision boundary(svm clf2, -1.5, 2.5, highlight=False, axes
limit = [-1.5, 2.5, -1, 1.5])
    sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
    plt.suptitle(main title)
    plt.tight layout()
    return fig
with warnings.catch warnings():
    warnings.simplefilter("ignore")
    fig num+=1
    fig = non linear examples ("Figure %d: Linear SVM on Non-Linear D
ata"%fig num)
    if PDF:
        plt.show()
        plt.close()
    else:
        plt.close()
        display(MatplotlibFigure(fig, centered=True))
```

Figure 21: Linear SVM on Non-Linear Data





Feature Engineering

We could consider enlarging the feature space to make the dataset linearly separable.

Example: We can see below that our x_1 is not linearly separable but it is when we add in our second feature $x_2 = (x_1)^2$

In [11]:

```
# https://github.com/ageron/handson-m12/blob/master/05 support vecto
r machines.ipynb
def poly info(main title):
   X1D = np.linspace(-4, 4, 9).reshape(-1, 1)
    X2D = np.c [X1D, X1D**2]
    y = np.array([0, 0, 1, 1, 1, 1, 1, 0, 0])
    fig = plt.figure(figsize=(width inch*2, height inch))
    plt.subplot(121)
    plt.grid(True, which='both')
    plt.axhline(y=0, color='k')
    plt.plot(X1D[:, 0][y==0], np.zeros(4), "bs")
    plt.plot(X1D[:, 0][y==1], np.zeros(5), "g^{"}")
    plt.gca().get yaxis().set ticks([])
    plt.xlabel(r"$x 1$", fontsize=20)
    plt.axis([-4.5, 4.5, -0.2, 0.2])
    plt.subplot(122)
    plt.grid(True, which='both')
    plt.axhline(y=0, color='k')
    plt.axvline(x=0, color='k')
    plt.plot(X2D[:, 0][y==0], X2D[:, 1][y==0], "bs")
    plt.plot(X2D[:, 0][y==1], X2D[:, 1][y==1], "g^")
    plt.xlabel(r"$x_1$", fontsize=20)
    plt.ylabel(r"$x 2$ ", fontsize=20, rotation=0)
    plt.gca().get yaxis().set ticks([0, 4, 8, 12, 16])
```

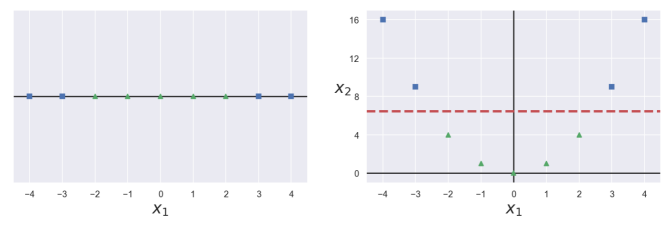
```
plt.plot([-4.5, 4.5], [6.5, 6.5], "r--", linewidth=3)
plt.axis([-4.5, 4.5, -1, 17])

plt.subplots_adjust(right=1)

plt.suptitle(main_title)
return fig

fig_num+=1
fig = poly_info("Figure %d: Adding Features to Linearly Separate Dat a"%fig_num)
if PDF:
   plt.show()
   plt.close()
else:
   plt.close()
display(MatplotlibFigure(fig, centered=True))
```

Figure 22: Adding Features to Linearly Separate Data



Using quadratic, cubic or higher-order polynomial functions we can project our data onto a higher-dimensional space via a mapping function ϕ where they are linearly separable (using a linear SVM model in this new feature space).

Example

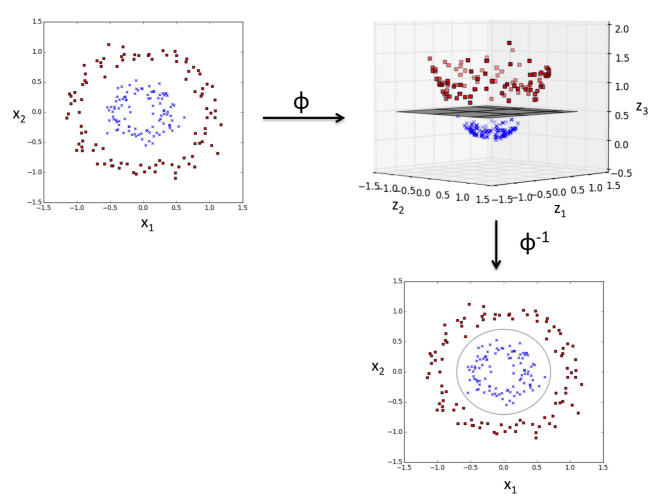
```
\phi(x_1, x_2) = (z_1, z_2, z_3)
\phi(x_1, x_2) = (x_1, x_2, x_1^2)
\phi(x_1, x_2) = (x_1, x_2, x_1^2)
```

In [12]:

```
# TODO - if time code something like this up
fig_num+=1
print(color.BOLD+color.UNDERLINE+"Figure %d: 2D Dataset into Separab
le 3D Feature Space"%fig_num+color.END)
# Image from https://github.com/rasbt/python-machine-learning-book-3
rd-edition/blob/master/ch03/ch03.ipynb
Image(filename='Images/03_13.png', width=width_pix)
```

Figure 23: 2D Dataset into Separable 3D Feature Space





Gaussian Radial Basis Function²

We could instead use a "similarity function", such as a Gaussian Radial Basis Function (RBF),

$$\phi_{\gamma}(\mathbf{x}, \ell) = \exp \left(-\gamma ||\mathbf{x} - \ell||^2\right)$$

This is a bell-shaped function which measures how much an instance resembles a *landmark*, with the function varying from 0 (far away) to 1 (at the landmark).

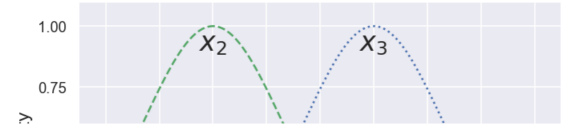
Example: Below we set our landmarks to $x_1 = -2$ and $x_1 = 1$.

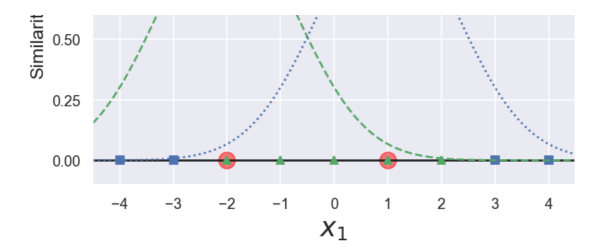
In [13]:

```
# copied from https://github.com/ageron/handson-m12/blob/master/05_s
upport_vector_machines.ipynb
def gaussian_rbf(x, landmark, gamma):
    return np.exp(-gamma * np.linalg.norm(x - landmark, axis=1) *
*2)
```

```
def rbk info(main title):
    qamma = 0.3
    X1D = np.linspace(-4, 4, 9).reshape(-1, 1)
    X2D = np.c [X1D, X1D**2]
    y = np.array([0, 0, 1, 1, 1, 1, 1, 0, 0])
    x1s = np.linspace(-4.5, 4.5, 200).reshape(-1, 1)
    x2s = gaussian rbf(x1s, -2, gamma)
    x3s = gaussian rbf(x1s, 1, gamma)
   XK = np.c_[gaussian_rbf(X1D, -2, gamma), gaussian_rbf(X1D, 1, gamma)]
mma)]
   yk = np.array([0, 0, 1, 1, 1, 1, 1, 0, 0])
    fig = plt.figure(figsize=(width inch, height inch))
    plt.grid(True, which='both')
    plt.axhline(y=0, color='k')
    plt.scatter(x=[-2, 1], y=[0, 0], s=150, alpha=0.5, c="red")
    plt.plot(X1D[:, 0][yk==0], np.zeros(4), "bs")
    plt.plot(X1D[:, 0][yk==1], np.zeros(5), "g^")
    plt.plot(x1s, x2s, "g--")
    plt.plot(x1s, x3s, "b:")
    plt.gca().get yaxis().set ticks([0, 0.25, 0.5, 0.75, 1])
    plt.xlabel(r"$x 1$", fontsize=20)
    plt.ylabel(r"Similarity", fontsize=14)
    plt.text(-2, 0.9, "$x 2$", ha="center", fontsize=20)
    plt.text(1, 0.9, "$x 3$", ha="center", fontsize=20)
    plt.axis([-4.5, 4.5, -0.1, 1.1])
    plt.suptitle(main title)
    return fig
fig num+=1
fig = rbk info("Figure %d: RBF Landmarks"%fig num)
if PDF:
   plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```

Figure 24: RBF Landmarks





Using the example of $x_1 = -1$ we can see it is a distance of 1 from the first landmark and 2 from the second.

If we set $\gamma = 0.3$ then our new features are:

```
x_2 = \exp \left( -0.3 \times 1^2 \right) \ pprox 0.74 \ x_3 = \exp \left( -0.3 \times 2^2 \right) \ pprox 0.30
```

In order to find the landmarks the simplist approach is just to create a landmark at each instance in the dataset.

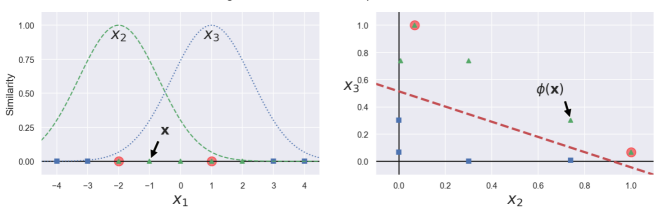
In [14]:

```
import math
# copied from https://github.com/ageron/handson-ml2/blob/master/05 s
upport vector machines.ipynb
def gaussian rbf(x, landmark, gamma):
        return np.exp(-gamma * np.linalg.norm(x - landmark, axis=1) *
*2)
def rbk info(main title):
    gamma = 0.3
    X1D = np.linspace(-4, 4, 9).reshape(-1, 1)
    X2D = np.c [X1D, X1D**2]
    y = np.array([0, 0, 1, 1, 1, 1, 1, 0, 0])
    x1s = np.linspace(-4.5, 4.5, 200).reshape(-1, 1)
    x2s = gaussian rbf(x1s, -2, gamma)
    x3s = gaussian rbf(x1s, 1, gamma)
    XK = np.c_[gaussian_rbf(X1D, -2, gamma), gaussian_rbf(X1D, 1, gamma)]
mma)]
    yk = np.array([0, 0, 1, 1, 1, 1, 1, 0, 0])
```

```
fig = plt.figure(figsize=(width inch*2, height inch))
    plt.subplot(121)
   plt.grid(True, which='both')
    plt.axhline(y=0, color='k')
   plt.scatter(x=[-2, 1], y=[0, 0], s=150, alpha=0.5, c="red")
   plt.plot(X1D[:, 0][yk==0], np.zeros(4), "bs")
   plt.plot(X1D[:, 0][yk==1], np.zeros(5), "g^")
   plt.plot(x1s, x2s, "g--")
   plt.plot(x1s, x3s, "b:")
   plt.gca().get yaxis().set ticks([0, 0.25, 0.5, 0.75, 1])
    plt.xlabel(r"$x 1$", fontsize=20)
   plt.ylabel(r"Similarity", fontsize=14)
    plt.annotate(r'$\mathbf{x}$',
                 xy = (X1D[3, 0], 0),
                 xytext = (-0.5, 0.20),
                 ha="center",
                 arrowprops=dict(facecolor='black', shrink=0.1),
                 fontsize=18,
    plt.text(-2, 0.9, "$x 2$", ha="center", fontsize=20)
    plt.text(1, 0.9, "$x 3$", ha="center", fontsize=20)
   plt.axis([-4.5, 4.5, -0.1, 1.1])
   plt.subplot(122)
   plt.grid(True, which='both')
   plt.axhline(y=0, color='k')
   plt.axvline(x=0, color='k')
   plt.plot(XK[:, 0][yk==0], XK[:, 1][yk==0], "bs")
   plt.plot(XK[:, 0][yk==1], XK[:, 1][yk==1], "g^")
   plt.scatter(x = [math.exp(-0.3*(0.**2.)), math.exp(-0.3*(3.**2.))]
y = [math.exp(-0.3*(3.**2.)), math.exp(-0.3*(0.**2.))], s=150, alpha
=0.5, c="red")
   plt.xlabel(r"$x 2$", fontsize=20)
   plt.ylabel(r"$x 3$ ", fontsize=20, rotation=0)
    plt.annotate(r'$\phi\left(\mathbf{x}\right)$',
                 xy = (XK[3, 0], XK[3, 1]),
                 xytext=(0.65, 0.50),
                 ha="center",
                 arrowprops=dict(facecolor='black', shrink=0.1),
                 fontsize=18,
   plt.plot([-0.1, 1.1], [0.57, -0.1], "r--", linewidth=3)
   plt.axis([-0.1, 1.1, -0.1, 1.1])
   plt.subplots adjust(right=1)
   plt.suptitle(main title)
   return fig
fig num+=1
fig = rbk info("Figure %d: RBF New Feature Space"%fig num)
if PDF:
```

```
plt.show()
  plt.close()
else:
  plt.close()
  display(MatplotlibFigure(fig, centered=True))
```

Figure 25: RBF New Feature Space



Kernels

However, by using feature engineering to enlarge our feature space, the larger the number of features, the higher computational burden.

Instead it is common to enlarge the feature space using an extension of a SVC termed a Support Vector Machine, which uses *kernels*.

The Kernel trick can relies on the fact we can define our SVM in the form of inner products.

$$L_D(lpha_i) = \sum_{i=lpha_i}^n lpha_i - rac{1}{2} \ \sum_{i,k}^n lpha_i lpha_k y_i y_k oldsymbol{\mathrm{x}}_i^{\mathrm{T}} oldsymbol{\mathrm{x}}_k \ \mathrm{s.t.} \quad orall_i lpha_i \geq 0, \ \sum_{i=lpha_i}^n y_i = 0.$$

Imagine we had a mapping ϕ which maps the data to some high dimensional Euclidean space

$$\phi: \mathbb{R}^d \mapsto H$$
,

then, we could do dot products between vectors after the mapping in H:

$$L_D(lpha_i) = \sum_{i}^n lpha_i - rac{1}{2} \ \sum_{i,k}^n lpha_i lpha_k y_i y_k \phi(\mathbf{x}_i^{\mathrm{T}}) \phi$$

$$(\mathbf{x}_k)$$

Instead we could use a kernel function,

$$K(\mathbf{x}_i, \mathbf{x}_k)$$

$$= \phi(\mathbf{x}_i^{\mathrm{T}})\Phi(\mathbf{x}_k).$$

Notes

- · Roughly speaking, a kernel can be interpreted as a similarity function betwen pairs of samples⁴.
- $\mathbf{a} \in \mathbb{R}^k$ just means it is a vector of length k. $\mathbf{A} \in \mathbb{R}^{r \times s}$ just means it is an $r \times s$ matrix. \mathbf{w} lives in H ("high dimensional") H is often refered to as a Hilbert space.

- "it is clear that the above implicit mapping trick will work for any algorithm in which the data only appear as dot products (for example, the nearest neighbor algorithm). This fact has been used to derive a nonlinear version of principal component analysis by (Scholkopf, Smola and Muller, 1998b); it seems likely that this trick will continue to find uses elsewhere."6

Second-Degree Polynomial Example²

Suppose we wanted to apply a second-degree polynomial transformation to two 2D vectors, a and b, and then compute the dot product. We could do this by:

$$\phi(\mathrm{a})^{\mathrm{T}}\phi(\mathrm{b}) \ \mathrm{T} \ = \left(rac{a_1^{\ 2}}{\sqrt{2a_1a_2}}
ight)^{\mathrm{T}} \left(rac{b_1^{\ 2}}{\sqrt{2b_1b_2}}
ight) \ = a_1^{\ 2}b_1^{\ 2} + 2a_1b_1a_2b_2 \ + a_2^{\ 2}b_2^{\ 2}.$$

Instead we could use the kernel approach:

$$K(\mathbf{a}, \mathbf{b}) = (\mathbf{a}^{\mathrm{T}} \mathbf{b})^{2}$$

$$= \left(\begin{pmatrix} a_{1} \\ a_{2} \end{pmatrix}^{\mathrm{T}} \begin{pmatrix} b_{1} \\ b_{2} \end{pmatrix}^{2} \right)^{2}$$

$$= (a_{1}b_{1} + a_{2}b_{2})^{2}$$

$$= a_{1}^{2}b_{1}^{2} + 2a_{1}b_{1}a_{2}b_{2}$$

$$+ a_{2}^{2}b_{2}^{2}.$$

This is useful as it means we didnt have to map our data using ϕ first. This saves us time!

Notes

• A second-degree polynomial for a training set with two features, $\mathbf{x} \in \mathbb{R}^2$, would be:

```
\begin{split} \Phi(x) &= \Phi((x_1, x_2)) \\ &= ({x_1}^2, \sqrt{2x_1x_2}, {x_2}^2) \end{split}
```

In [16]:

```
# TODO: Time allowing, look at if this increases as polynomial degre
# it doesnt always work - quess it depends how my computer is feelin
q : P
# https://github.com/ageron/handson-ml2/blob/master/05 support vecto
r machines.ipynb
from sklearn.datasets import make moons
from sklearn.pipeline import Pipeline
from sklearn.preprocessing import PolynomialFeatures
polynomial feat = Pipeline([
        ("poly features", PolynomialFeatures(degree=2)),
        ("scaler", StandardScaler()),
        ("svm clf", LinearSVC(C=10, loss="hinge", random state=42))
    ])
polynomial svm = Pipeline([
        ("scaler", StandardScaler()),
        ("svm clf", SVC(kernel='poly', degree=2, C=10, random state=
42))
   ])
with warnings.catch warnings():
    warnings.simplefilter("ignore")
    print(color.BOLD+color.UNDERLINE+"Polynomial Feature Engineering
(degree=2) "+color.END)
    %timeit polynomial feat.fit(nl2, y2)
    print(color.BOLD+color.UNDERLINE+"Polynomial Kernel (degree=2)"+
color.END)
    %timeit polynomial svm.fit(nl2, y2)
```

Polynomial Feature Engineering (degree=2)

2.17 ms \pm 66 μ s per loop (mean \pm std. dev. of 7 runs, 100 loops each) Polynomial Kernel (degree=2)

3.17 ms \pm 155 μ s per loop (mean \pm std. dev. of 7 runs, 100 loops eac h)

We still have all the same considerations, but replacing $\mathbf{x}_i^T \mathbf{x}_k$ with $K(\mathbf{x}_i, \mathbf{x}_k)$ allows us to produce a SVM in infinite dimensional space.

$$\sum_{i}^{n} \sum_{\alpha_{i} - \frac{1}{2}}^{n} \sum_{i}^{n}$$

$$\sum_{k}^{n} lpha_{i} lpha_{k} y_{i} y_{k} K(\mathbf{x}_{i}, \mathbf{x}_{k}) \ ext{s.t.} \quad orall_{i} lpha_{i} \geq 0, \ \sum_{i}^{n} lpha_{i} y_{i} = 0.$$

In the test phase, we can use the support vectors 6 :

$$f(\mathbf{x}^*) = \sum_{i \in s} \hat{\alpha}_i y_i \phi(\mathbf{x}_i) \cdot \phi(\mathbf{x}^*) + \hat{b}$$

$$= \sum_{i \in s} \hat{\alpha}_i y_i K(\mathbf{x}_i, \mathbf{x}^*) + \hat{b},$$
(3)

avoiding computing $\Phi(x^*)$.

Note

ullet Put another way, in the test phase 6 , we can use the support vectors, \mathbf{s}_i :

$$f(\mathbf{x}^*) = \sum_{i=1}^{n_s} \hat{\alpha}_i y_i \Phi(\mathbf{s}_i)$$
 (5)

Polynomial kernels

The polynomial kernel of degree d (a positive integer) can be defined as:

$$K(\mathbf{x}_i, \mathbf{x}_k) = (\gamma \langle \mathbf{x}_i, \mathbf{x}_k \rangle + r)^d$$

Hyperparameters

- γ (gamma) broadly defines the width or slope of kernel function 13 .
- r (coef0) roughly controls how much the model is influenced by high-degree polynomials².
- ullet \dot{d} (degree) is the degree of the polynomial.

Notes

• You will sometimes see a kernel defined in a general sense using K(x, x'), where x'means another point in the dataset.

```
egin{aligned} raket{x_i, x_j} &= x_i \cdot x_j \ &= \mathbf{x}_i^{\mathrm{T}} \mathbf{x}_k \end{aligned}
```

• γ is often 1.

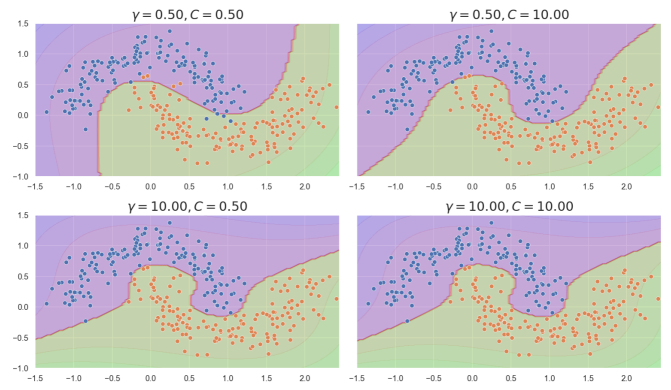
- "The 'curve' of the decision boundary becomes very low when gamma value is low making the decision region very broad. The 'curve' of the decision boundary becomes high when gamma is high, which creates islands of decision-boundaries around data points."
- *d* is a positive integer.
- If you start to overfit you should reduce the polynomial degree and underfitting try increasing it.

In [17]:

```
def plot predictions(clf, axes):
    x0s = np.linspace(axes[0], axes[1], 100)
    x1s = np.linspace(axes[2], axes[3], 100)
    x0, x1 = np.meshgrid(x0s, x1s)
    X = np.c [x0.ravel(), x1.ravel()]
    y pred = clf.predict(X).reshape(x0.shape)
    y decision = clf.decision function(X).reshape(x0.shape)
    plt.contourf(x0, x1, y pred, cmap=plt.cm.brg, alpha=0.2)
    plt.contourf(x0, x1, y decision, cmap=plt.cm.brg, alpha=0.1)
poly kernel svm clf = Pipeline([
        ("scaler", StandardScaler()),
        ("svm clf", SVC(kernel="poly", gamma=1, degree=3, coef0=1, C
=5))
poly kernel svm clf.fit(nl2, y2)
poly100 kernel svm clf = Pipeline([
        ("scaler", StandardScaler()),
        ("svm clf", SVC(kernel="poly", gamma=10, degree=3, coef0=1,
C = 5))
poly100 kernel svm clf.fit(nl2, y2)
fig, axes = plt.subplots(ncols=2, nrows =2, figsize=(width inch*2, h
eight inch*2), sharey=True)
axes = axes.flatten()
gamma list=[0.5, 10]
C list=[0.5, 10]
i = 0
for gamma in gamma list:
    for C in C list:
        poly kernel svm clf = Pipeline([
            ("scaler", StandardScaler()),
            ("svm clf", SVC(kernel="poly",
                             gamma=gamma,
                             C=C,
```

```
degree=3, coef0=1))
        ])
        poly kernel svm clf.fit(nl2, y2)
        warnings.simplefilter("ignore")
        plt.sca(axes[i])
        plot predictions (poly kernel svm clf, [-1.5, 2.45, -1, 1.5])
        with warnings.catch warnings():
            warnings.simplefilter("ignore")
            sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
        plt.title(r"\$\gamma = \$.2f, C=\$.2f$"\$(gamma, C), fontsize=18
        plt.ylabel("")
        i+=1
fig num+=1
plt.suptitle("Figure %d: Polynomial Kernel Decision Boundaries (degr
ee=3, coef0=1) "% fig num)
plt.tight layout()
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```





You shouldnt stick to the default settings, instead you want to search for optimal hyperparameters for the data. Good suggestions for search spaces are:

• "The authors of libsvm (Hsu et al. 2009) recommend using cv over a 2d grid with

```
C \in \{2^{-5}, 2^{-3} \dots, 2^{15} \\ \} \\ \textit{and} \\ \gamma \in \{2^{-15}, 2^{-13}, \dots, 2^{3} \\ \}_{\textit{A}}
```

In [18]:

```
from sklearn.model_selection import GridSearchCV, RandomizedSearchCV
, KFold
def hyper search(model, params, X, y, save path, n iter=60, metric="
accuracy",
                 cv = KFold(5), random state=42, refit=True,
                 overwrite=False):
    if os.path.exists(save path) and overwrite==False:
        #load the model
        models = joblib.load(save path)
    else:
        # check all param inputs are lists
        if all(type(x) == list for x in params.values()):
            search type = "Gridsearch"
            models = GridSearchCV(model, param grid=params, scoring=
metric, cv=cv,
                                   refit=refit, return train score=Tr
ue)
            n iter = len(list(itertools.product(*list(iter(params.va
lues())))))
        else:
            search type = "Randomsearch"
            models = RandomizedSearchCV(model, param distributions=p
arams,
                                         n iter=n iter, scoring=metri
c, cv=cv,
                                         refit=refit, random state=ra
ndom state,
                                         return train score=True)
        start = time()
        with warnings.catch warnings():
            warnings.simplefilter("ignore")
            models.fit(X, y)
        print(search type + " took %.2f seconds for %d candidates" %
((time() - start), n iter))
        joblib.dump(models, save path)
    return models
```

```
C list = []
pwr = -5
for i in range(11):
   C list.append(2**pwr)
   pwr+=2
gamma list = []
pwr = -15
for i in range (10):
   gamma list.append(2**pwr)
    pwr+=2
degree list = list(range(1,5))
coef0 list = []
pwr = 1
for i in range (7):
    coef0 list.append(2**pwr)
    pwr+=1
poly param grid = {'svm clf C':C list,
                   'svm clf gamma': gamma list,
                   'svm clf degree': degree list,
                   'svm clf coef0':coef0 list}
poly = SVC(random state=42, kernel = 'poly', max iter = 1e5)
poly svm = Pipeline([
        ("scaler", StandardScaler()),
        ("svm clf", poly)])
nl2 poly gs = hyper search(poly svm, poly param grid, nl2, y2,
                             os.path.join(os.getcwd(), "Models", "mo
ons poly qs.pkl"),
                             overwrite=False)
pd.DataFrame(nl2 poly gs.cv results ).sort values("rank test score")
[["param svm clf C",
"param svm clf gamma",
"param svm clf degree",
"param svm clf coef0",
"mean test score",
"std test_score"]].head()
```

Out[19]:

2829	param_svm_clf_6	param_svm_clfgamma	param_svm_clfdegree	paran
957	2	0.5	4	
1796	128	0.125	4	
1149	8	8	3	
757	0.5	0.5	4	
4				F

Radial Basis Function kernel

The most widely used kernel is the RBF kernel (also known as a Gaussian kernel) 4,8 :

$$K(\mathbf{x}_i, \mathbf{x}_k) = \exp \left(-\gamma ||\mathbf{x}_i - \mathbf{x}_k||^2\right),$$

where γ is a free parameter to be optimised and $||\mathbf{x}_i - \mathbf{x}_k||^2$ is the squared *Euclidean distance*.

 γ is often either a positive constant or $\frac{1}{2\sigma^2}$

When classifying a test observation $\mathbf{x}^* = (x_1^* \dots x_p^*)$, only training observations close to \mathbf{x}^* (in terms of Euclidean distance) will play a role in its class label. This is because $(x_j^* - x_{ij})^2$ will be large, so \exp

$$(-\gamma \sum_{j=1}^{P} (x_j^* - x_{ij})^2)$$
will be small 1

Notes

Euclidean distance⁸

$$egin{aligned} & \frac{d(\mathrm{x}_i,\mathrm{x}_k)}{\det} \ & \frac{\det}{1 - x_{k1}} \left(\frac{\left(x_{i1} - x_{k1} \right)^2 + \left(x_{i2} - x_{k2} \right)^2 + \ldots}{\left(x_{iN} - x_{kN} \right)^2} \right. \ & = \sqrt{\sum_{j=1}^{D} \left(x_{ij} - x_{kj} \right)^2} \end{aligned}$$

- _"if gamma='scale' (default) is passed then it uses 1 / (nfeatures * X.var()) as value of $gamma''^{12}$
- the minus sign inverts the the distance measure to a similarity score 10

Hyperparameters²

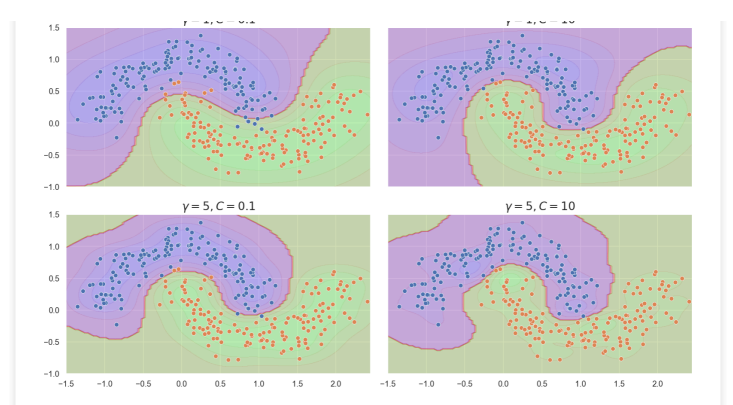
 γ is effectively acting like a regularization hyperparameter, so like C if your model is overfitting reduce it and underfitting then increase it.

- Increasing γ (gamma) makes the bell-shaped curve narrower.
 - Each instances range of influence is smaller.
 - The decision boundary becomes more irregular.
- Decreasing γ makes the bell-shaped curve wider.
 - Instances have a larger range of influence.
 - The decision boundary becomes smoother decision.

In [20]:

```
gamma1, gamma2 = 1, 5
C1, C2 = 0.1, 10
hyperparams = (gamma1, C1), (gamma1, C2), (gamma2, C1), (gamma2, C2)
svm clfs = []
for gamma, C in hyperparams:
    rbf kernel svm clf = Pipeline([
            ("scaler", StandardScaler()),
            ("svm clf", SVC(kernel="rbf", gamma=gamma, C=C))
        1)
    rbf kernel svm clf.fit(nl2, y2)
    svm clfs.append(rbf kernel svm clf)
fig, axes = plt.subplots(nrows=2, ncols=2, figsize=(width inch*2, he
ight inch*2), sharex=True, sharey=True)
with warnings.catch warnings():
    warnings.simplefilter("ignore")
    for i, svm clf in enumerate(svm clfs):
        plt.sca(axes[i // 2, i % 2])
        plot predictions (svm clf, [-1.5, 2.45, -1, 1.5])
        sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
        gamma, C = hyperparams[i]
        plt.title(r"$\qamma = {}, C = {}$".format(gamma, C), fontsiz
e = 16)
        if i in (0, 1):
            plt.xlabel("")
        if i in (1, 3):
            plt.ylabel("")
fig num+=1
plt.suptitle("Figure %d: Gamma Kernel Decision Boundaries"%fig num)
plt.tight layout()
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```

Figure 27: Gamma Kernel Decision Boundaries



In [21]:

```
rbf_param_grid = {'svm_clf__C':C_list,
                  'svm_clf__gamma':gamma_list
rbf = SVC(random state=42, gamma="auto")
rbf svm = Pipeline([
        ("scaler", StandardScaler()),
        ("svm_clf", rbf)])
nl2_rbf_gs = hyper_search(rbf_svm, rbf_param_grid, nl2, y2,
                            os.path.join(os.getcwd(), "Models", "moo
ns_rbf_gs.pkl"),
                            overwrite=False
pd.DataFrame(nl2_rbf_gs.cv_results_).sort_values("rank_test_score")[
["param_svm_clf__C",
                                                                   "pa
ram svm clf gamma",
                                                                   "me
an test score",
                                                                   "st
d_test_score"]].head()
```

Out[21]:

param_svm_clf__gamma mean_test_score std_test_score

58 32 2 0.996667 0.00666

78	param_svm_clf_51@	param_svm_clfgamma	mean_test9score	std_testoscor
107	32768	0.5	0.993333	0.00816
87	2048	0.5	0.993333	0.00816
88	2048	2	0.993333	0.00816
4				

C and γ are tightly coupled.

Generally if you have a larger/narrow gamma (e.g. $\gamma = 5$) you'll need more regularisation, so a larger C.

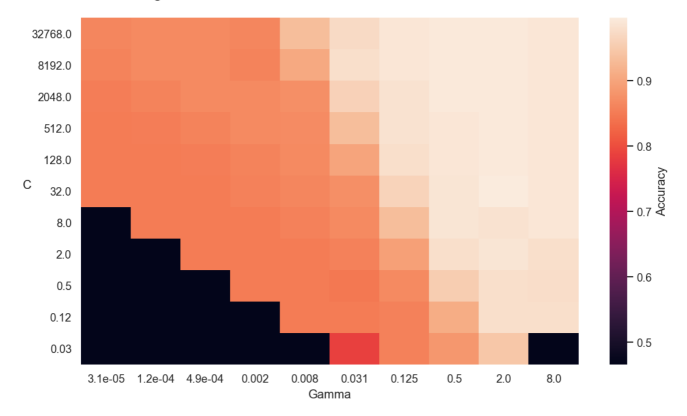
If you have a smaller/wider gamma (e.g. $\gamma = 1$), a smaller value of C should be used 7.

In [22]:

```
grid df = pd.DataFrame(nl2 rbf gs.cv results )[["param svm clf C",
                                                   "param svm clf ga
mma",
                                                   "mean test score"]
grid df = grid df.sort values(["param svm clf C", "param svm clf g
amma"])
grid np = np.array(grid df["mean test score"]).reshape(len(C list),
len(gamma list))
tidied gamma lst = list(pd.Series(gamma list).apply(np.format float
scientific, precision=1)[0:3].values)
tidied gamma lst = tidied gamma lst+list(np.array(gamma list[3:]).ro
und(3))
grid plt df = pd.DataFrame(grid np,
                           index = np.array(C list).round(2),
                           columns=tidied gamma lst)
fig = plt.figure(figsize = (width inch*1.5, height inch*1.5))
ax = sns.heatmap(grid plt df,
                 vmin=min(grid df["mean test score"]),
                 vmax=max(grid df["mean test score"]),
                 #cmap="YlGnBu r"
ax.collections[0].colorbar.set label("Accuracy")
ax.invert yaxis()
ax.set xlabel("Gamma")
ax.set ylabel("C", rotation = 0)
plt.yticks(rotation = 0)
plt.xticks(rotation = 0)
fig num+=1
plt.suptitle("Figure %d: GridSearch Results for Gamma and C on the M
oons Data"%fig num)
plt.tight layout()
```

```
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```

Figure 28: GridSearch Results for Gamma and C on the Moons Data



Extra Example

Lets have a look at the decision boundaries for our "optimal hyperparameters" as a comparison to those by the linear models we saw earlier.

In [23]:

```
"std_test_score"]].head()
```

Out[23]:

	param_svm_clfC	param_svm_clfgamma	param_svm_clfdegree	paran
387	0.125	0.5	3	
514	0.125	0.007812	4	
1545	32	0.03125	3	
906	2	0.125	3	
506	0.125	0.125	3	
4				•

In [24]:

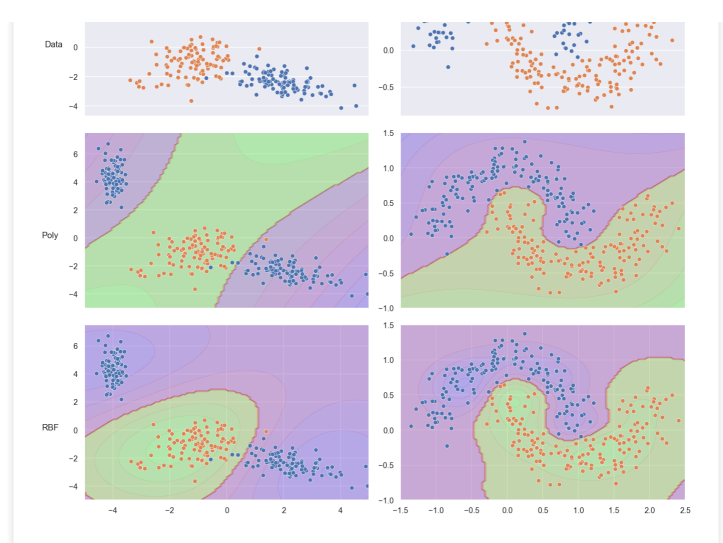
Out[24]:

	param_svm_clfC	param_svm_clfgamma	mean_test_score	std_test_score
29	0.5	8	0.983333	0.014907
27	0.5	0.5	0.983333	0.014907
28	0.5	2	0.983333	0.014907
37	2	0.5	0.983333	0.014907
18	0.125	2	0.983333	0.014907
4				[)

```
with warnings.catch warnings():
    warnings.simplefilter("ignore")
    fig, axes = plt.subplots(ncols=2, nrows=3, figsize=(width inch*2
, height inch*3))
    axes = axes.flatten()
   plt.sca(axes[0])
    sns.scatterplot(nl1[:,0], nl1[:,1], y1, legend=False)
    plt.sca(axes[1])
    sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
   plt.sca(axes[2])
   plot predictions (nl1 poly gs, [-5, 5, -5, 7.5])
    sns.scatterplot(nl1[:,0], nl1[:,1], y1, legend=False)
   plt.sca(axes[3])
   plot predictions (nl2 poly gs, [-1.5, 2.5, -1, 1.5])
    sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
   plt.sca(axes[4])
   plot predictions (nl1 rbf gs, [-5, 5, -5, 7.5])
    sns.scatterplot(nl1[:,0], nl1[:,1], y1, legend=False)
    plt.sca(axes[5])
    plot predictions (nl2 rbf gs, [-1.5, 2.5, -1, 1.5])
    sns.scatterplot(nl2[:,0], nl2[:,1], y2, legend=False)
    for i in range(4):
        axes[i].axes.xaxis.set visible(False)
    axes[0].text(-6.5, 0, 'Data')
    axes[2].text(-6.5, 0, 'Poly')
    axes[4].text(-6.5, 0, 'RBF')
    fiq num+=1
   plt.suptitle("Figure %d: Best Non-Linear SVM Model Decision Boun
dary Using GridSearch"%fig num)
   plt.tight layout()
    if PDF:
        plt.show()
       plt.close()
    else:
        plt.close()
        display(MatplotlibFigure(fig, centered=True))
```

Figure 29: Best Non-Linear SVM Model Decision Boundary Using GridSearch





Standardization¹¹

SVMs, along with many other machine learning estimators (e.g. I1 and I2 regularizers of linear models), are sensitive to feature scales.

If a feature has a variance orders of magnitude larger than others, it might dominate the objective function and make the estimator unable to learn from other features correctly as expected.

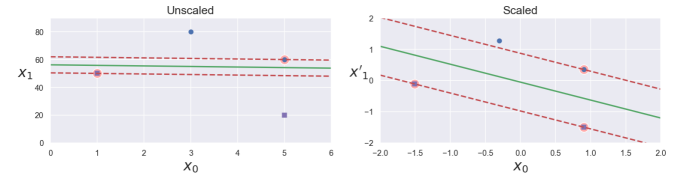
In [26]:

```
# copied from https://github.com/ageron/handson-m12/blob/master/05_s
upport_vector_machines.ipynb
Xs = np.array([[1, 50], [5, 20], [3, 80], [5, 60]]).astype(np.float6
4)
ys = np.array([0, 0, 1, 1])
svm_clf = SVC(kernel="linear", C=100)
svm_clf.fit(Xs, ys)

fig = plt.figure(figsize=(width_inch*2, height_inch))
plt.subplot(121)
plt.plot(Xs[:, 0][ys==1], Xs[:, 1][ys==1], "bo")
plt.plot(Xs[:, 0][ys==0], Xs[:, 1][ys==0], "ms")
svc_decision_boundary(svm_clf, 0, 6)
plt.xlabel("$x_0$", fontsize=20)
plt.ylabel("$x_1$  ", fontsize=20, rotation=0)
plt_title("Unscaled" fontsize=16)
```

```
bic. cicic/ and carea 1 rancoire in
plt.axis([0, 6, 0, 90])
from sklearn.preprocessing import StandardScaler
scaler = StandardScaler()
X scaled = scaler.fit transform(Xs)
svm clf.fit(X scaled, ys)
plt.subplot(122)
plt.plot(X scaled[:, 0][ys==1], X scaled[:, 1][ys==1], "bo")
plt.plot(X scaled[:, 0][ys==0], X scaled[:, 1][ys==0], "ms")
svc decision boundary(svm clf, -2, 2)
plt.xlabel("$x 0$", fontsize=20)
plt.ylabel("$x' 1$ ", fontsize=20, rotation=0)
plt.title("Scaled", fontsize=16)
plt.axis([-2, 2, -2, 2])
fig num+=1
plt.suptitle("Figure %d: Sensitivity to Feature Scales"%fig num)
plt.tight layout()
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display (MatplotlibFigure (fig, centered=True))
```

Figure 30: Sensitivity to Feature Scales



Notes

- We can see that in the unscaled plot that x_1 is much larger than x_0 so the widest street is very close to the horizontal.
- After scaling we have a much better decision boundary.

Standardization is especially important for RBF kernels.

These models assume that all features look like standard normally distributed data: Gaussian with zero mean and unit variance.

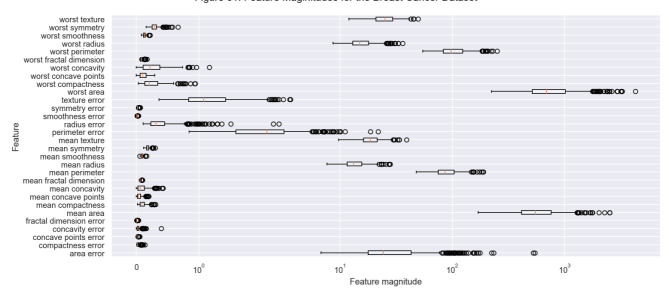
Dataset Example: Breast Cancer

We can see below for example that the features in the Breast Cancer dataset are of completely different orders of magnitude 16

In [27]:

```
from sklearn.datasets import load breast cancer
from sklearn.model selection import train test split
X, y = load breast cancer(return X y=True, as frame=True)
X = X.reindex(sorted(X.columns), axis=1)
X train, X test, y train, y test = train test split(X.values, y.valu
es,
                                                     test size = 0.2,
random state=42)
X train, X val, y train, y val = train test split(X train, y train,
                                                     test size = 0.2,
random state=42)
fig, ax = plt.subplots(figsize=(width inch*2, height inch*1.5))
plt.boxplot(X train, vert=False)
plt.xscale("symlog")
plt.ylabel("Feature")
plt.xlabel("Feature magnitude")
ax.set yticklabels(list(X.columns))
fig num+=1
plt.suptitle("Figure %d: Feature Maginitudes for the Breast Cancer D
ataset"%fig num)
plt.tight layout()
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```

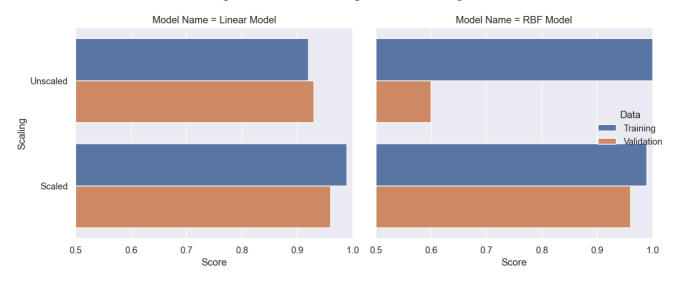
Figure 31: Feature Maginitudes for the Breast Cancer Dataset



```
from sklearn.metrics import accuracy score
lin = LinearSVC(random state=42)
rbf = SVC(random state=42, gamma="auto")
linear svm = Pipeline([
                     ("scaler", StandardScaler()),
                     ("svm clf", lin)
          ])
rbf svm = Pipeline([
                     ("scaler", StandardScaler()),
                     ("svm clf", rbf)])
model name = ["Linear Model", "Linear Model", "RBF M
1"]
for i, model in enumerate([lin, linear svm, rbf, rbf svm]):
          with warnings.catch warnings():
                     warnings.simplefilter("ignore")
                     trained model = model.fit(X train, y train)
                    pred train = trained model.predict(X train)
                    pred val = trained model.predict(X val)
                     if (i % 2) == 0:
                               scaled = "Unscaled"
                     else:
                               scaled = "Scaled"
                     score df = pd.DataFrame([round(accuracy score(y train, pred
train), 2),
                                                                                      round (accuracy score (y val, pred va
1),2)], columns = ["Score"])
                     score_df["Data"] = ['Training', 'Validation']
                     score df["Model Name"] = model name[i]
                     score df["Scaling"] = scaled
                     if i == 0:
                               scores df = score df
                     else:
                               scores df = pd.concat([scores df, score df], axis=0)
#fig = plt.figure(figsize=(width inch, height inch))
#fig, ax = plt.subplots(figsize=(width inch, height inch))
sns plot = sns.catplot(data = scores df, y="Scaling", x = "Score", h
ue="Data", kind="bar", col="Model Name", ax=ax)
fig = sns plot.fig
plt.xlim(0.5, 1.0)
fig num+=1
```

```
plt.suptitle("Figure %d: Effects of Scaling on SVM Overfitting"%fig_
num)
plt.tight_layout()
if PDF:
    plt.show()
    plt.close()
else:
    plt.close()
    display(MatplotlibFigure(fig, centered=True))
```

Figure 32: Effects of Scaling on SVM Overfitting



Notes

- In this example, the RBF model overfits quite substantially, with a perfect score on the training set and only 60% accuracy on the test set.
- Also you see the linear model also has a minor improvement in accuracy if scaled.

Recap¹

The kernel approach is an efficient computational approach to enlarge our feature space to accommodate a non-linear boundary.

Assume we have a new point x^* . If wanted to compute $f(x^*)$ using our linear classifier we would need to the inner product between x^* and each training point x_i :

$$f(x) = \sum_{oldsymbol{i} \in s} lpha_{oldsymbol{i}} \left\langle x^*, x_{oldsymbol{i}}
ight
angle + b.$$

Instead of actually calculating the inner product, we could instead use a *generalisation*, $K(x, x_{i'})$, where K is a *kernel*. We can now define the classifier as:

$$f(x) = \sum_{i \in s} lpha_i K(x^*, x_i) + b.$$

A kernel is a function that quantifies the similarity of two observations. For example, for a *linear kernel* we could use:

$$\sum_{K(x_i, x_{i'}) = j=1} \sum_{x_{ij} x_{i'j},$$

where we quantify the similarity of pairs of observations using Pearson (standard) correlation.

However, we could use other forms of kernel to fit the support vector classifier in a higher-dimensional space, such as a *polynomial kernel*:

$$K(x_i, x_{i'}) = \left(\gamma^{j=1} x_{ij} x_{i'j} + r \right)^d.$$

Another popular choice is the *radial kernel*:

$$K(x_i,x_{i'}) = \exp \left(\sum_{-\gamma \ j=1}^p (x_{ij}-x_{i'j})^2
ight).$$

Associated Exercises

Now might be a good time to try exercises 5-8.

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In [29]:

```
from shutil import copyfile
# where the HTML template is located
dst = os.path.join(sys.prefix, 'lib', 'site-packages', 'nbconvert',
'templates', "classic.tplx")
# If its not located where it should be
if not os.path.exists(dst):
    # uses a nb pdf template
    curr path = os.path.join(os.getcwd(),"..", "Extra", "classic.tpl
x")
    # copy where it is meant to be
    copyfile(curr path, dst)
if not PDF:
    # Create HTML notes document (preferred)
    !jupyter nbconvert 2 Support Vector Machines.ipynb \
        --to html \
        --output-dir . \
        --template classic
    !jupyter nbconvert 2 Support Vector Machines.ipynb \
        --to slides \
        --output-dir . \
        --TemplateExporter.exclude input=True \
        --TemplateExporter.exclude output prompt=True \
        --SlidesExporter.reveal scroll=True
else:
    # Create pdf notes document (issues)
    !jupyter nbconvert 2 Support Vector Machines.ipynb \
        --to pdf \
        --output-dir . \
        --TemplateExporter.exclude input=True \
        --TemplateExporter.exclude output prompt=True
[NbConvertApp] Converting notebook 2 Support Vector Machines.ipynb t
o html
[NbConvertApp] Writing 4536207 bytes to 2 Support Vector Machines.ht
[NbConvertApp] Converting notebook 2 Support Vector Machines.ipynb t
[NbConvertApp] Writing 3929286 bytes to 2 Support Vector Machines.sl
ides.html
```

In []: