In [2]: import numpy as np # For all our math needs n = 750# Number of data points X = np.random.uniform(-7.5, 7.5, n)# Training examples, in one dimension e = np.random.normal(0.0, 5.0, n)# Random Gaussian noise y = f true(X) + e# True labels with noise Now, we plot the raw data as well as the true function (without noise). In [3]: import matplotlib.pyplot as plt # For all our plotting needs plt.figure() # Plot the data plt.scatter(X, y, 12, marker='o') # Plot the true function, which is really "unknown" $x_{true} = np.arange(-7.5, 7.5, 0.05)$ y_true = f_true(x_true) plt.plot(x_true, y_true, marker='None', color='r') Out[3]: [<matplotlib.lines.Line2D at 0x7fde9a533460>] 20 10 0 -10-20Recall that we want to build a model to generalize well on future data, and in order to generalize well on future data, we need to pick a model that trade-off well between fit and complexity (that is, bias and variance). We randomly split the overall data set (D) into three • Training set: \mathcal{D}_{trn} consists of the actual training examples that will be used to train the model; • Validation set: \mathcal{D}_{val} consists of validation examples that will be used to tune model hyperparameters (such as $\lambda > 0$ in ridge regression) in order to find the best trade-off between fit and complexity (that is, the value of λ that produces the best model); • Test set: \mathcal{D}_{tst} consists of test examples to estimate how the model will perform on future data. Validation Train Test For this example, let us randomly partition the data into three non-intersecting sets: $D_{trn} = 60\%$ of D, $D_{val} = 10\%$ of D and $D_{tst} = 30\%$ In [4]: # scikit-learn has many tools and utilities for model selection from sklearn.model selection import train_test_split tst_frac = 0.3 # Fraction of examples to sample for the test set val frac = 0.1 # Fraction of examples to sample for the validation set # First, we use train_test_split to partition (X, y) into training and test sets X_trn, X_tst, y_trn, y_tst = train_test_split(X, y, test_size=tst_frac, random_state=42) # Next, we use train test split to further partition (X trn, y trn) into training and validation sets X_trn, X_val, y_trn, y_val = train_test_split(X_trn, y_trn, test_size=val_frac, random_state=42) # Plot the three subsets plt.figure() plt.scatter(X_trn, y_trn, 12, marker='o', color='orange') plt.scatter(X_val, y_val, 12, marker='o', color='green') plt.scatter(X_tst, y_tst, 12, marker='o', color='blue') Out[4]: <matplotlib.collections.PathCollection at 0x7fde9b6b43a0> 20 10 0 -10-201. **Regression with Polynomial Basis Functions**, 30 points. This problem extends ordinary least squares regression, which uses the hypothesis class of linear regression functions, to non-linear regression functions modeled using polynomial basis functions. In order to learn nonlinear models using linear regression, we have to explicitly transform the data into a higher-dimensional space. The nonlinear hypothesis class we will consider is the set of d-degree polynomials of the form $f(x) = w_0 + w_1 x + w_2 x^2 + ... + w_d x^d$ or a linear combination of polynomial basis function: $f(x) = [w_0, w_1, w_2 \dots, w_d]^T \begin{vmatrix} x \\ x^2 \end{vmatrix}$ The monomials $\{1, x, x^2, \dots, x^d\}$ are called **basis functions**, and each basis function x^k has a corresponding weight w_k associated with it, for all $k = 1, \ldots, d$. We transform each univariate data point x_i into into a multivariate (d-dimensional) data point via $\phi(x_i) \to [1, x_i, x_i^2, \dots, x_i^d]$. When this transformation is applied to every data point, it produces the **Vandermonde matrix**: $\Phi = \begin{bmatrix} 1 & x_1 & x_1^2 & \dots & x_1^d \\ 1 & x_2 & x_2^2 & \dots & x_2^d \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & x_n & x_n^2 & \dots & x_n^d \end{bmatrix}.$ a. (10 points) Complete the Python function below that takes univariate data as input and computes a Vandermonde matrix of dimension d. This transforms one-dimensional data into d-dimensional data in terms of the polynomial basis and allows us to model regression using a ddegree polynomial. In [15]: # X float(n,): univariate data # d int: degree of polynomial def polynomial transform(X, d): Phi = []for value in X: z = []for d in range (0, d+1): z.append(np.power(value, d)) Phi.append(z) Phi = np.asarray(Phi) return Phi **b.** (10 points) Complete the Python function below that takes a Vandermonde matrix Φ and the labels y as input and learns weights via **ordinary least squares regression**. Specifically, given a Vandermonde matrix Φ , implement the computation of $\mathbf{w} = (\Phi^T \Phi)^{-1} \Phi^T \mathbf{y}$. Remember that in Python, @ performs matrix multiplication, while * performs element-wise multiplication. Alternately, numpy.dot also performs matrix multiplication. In [6]: # Phi float(n, d): transformed data # y float(n,): labels def train model(Phi, y): w = np.linalg.inv(np.transpose(Phi) @ Phi) @ np.transpose(Phi) @ y return w

Complete the Python function below that takes a Vandermonde matrix Φ , corresponding labels y, and a linear regression model w as

We can explore the **effect of complexity** by varying $d=3,6,9,\cdots,24$ to steadily increase the non-linearity of the models. For each model, we train using the transformed training data (Φ , whose dimension increases) and evaluate its performance on the transformed

validationErr[d] = evaluate_model(Phi_val, y_val, w[d]) # Evaluate model on validation data

print ("The Validation and Test Error drop consistently until about a polynomial degree of 15 to 18, so

The Validation and Test Error drop consistently until about a polynomial degree of 15 to 18, so any d

Transform training data into d dimension

Transform validation data into d dimensi

Learn model on training data

Transform test data into d dimensions

input and evaluates the model using **mean squared error**. That is, $\epsilon_{MSE} = \frac{1}{n} \sum_{i=1}^{n} (y_i - \mathbf{w}^T \Phi_i)^2$.

Assignment I

Linear Regression

functions, to non-linear regression functions modeled using polynomial basis functions and radial basis functions. The function we want to fit is $y_{\text{true}} = f_{\text{true}}(x) = 6(\sin(x+2) + \sin(2x+4))$. This is a **univariate function** as it has only one input variable. First, we generate

This assignment shows how we can extend ordinary least squares regression, which uses the hypothesis class of linear regression

synthetic input (data) x_i by sampling n = 750 points from a uniform distribution on the interval [-7.5, 7.5].

Generating Synthetic Data

y = 6.0 * (np.sin(x + 2) + np.sin(2*x + 4))

We can generate a synthetic data set, with Gaussian noise.

The true function
def f true(x):

return y

c. (5 points)

In [7]: # Phi float(n, d): transformed data
y float(n,): labels

#ypred = Phi @ w

n = np.size(y)
for value in err:

#n = n + 1meansq = sum / n return meansq

d. (5 points, Discussion)

In $[8]: w = \{\}$

ons

testErr = {}

Plot all the models

plt.axis([2, 25, 15, 60])

plt.figure()

60

55

Validation/Test error

In [9]: plt.figure()

sum = 0

def evaluate_model(Phi, y, w):

err = ((Phi @ w) - y)**2#err = (y - (Phi @ w))**2

sum = sum + value

w float(d,): linear regression model

validation data and estimate what our future accuracy will be using the test data.

for d in range(3, 25, 3): # Iterate over polynomial degree

plt.legend(['Validation Error', 'Test Error'], fontsize=16)

Validation Error

Test Error

12

X_d = polynomial_transform(x_true, d)

plt.legend(['true'] + list(range(9, 25, 3)))

Finally, let's visualize each learned model.

for d in range(9, 25, 3):

plt.axis([-8, 8, -15, 15])

function**, which means that the model will be:

a. (15 points)

import math

Phi = []

return Phi

 $\mathbf{w} = (\Phi^T \Phi + \lambda I_n)^{-1} \Phi^T \mathbf{y}.$

In [11]: # Phi float(n, d): transformed data
y float(n,): labels

n = np.shape(Phi)[0]

lam float : regularization parameter

def train_ridge_model(Phi, y, lam):

validation error and (ii) λ vs. test error, as above.

In [12]: q = {} # Dictionary to store all the trained models
validationError = {} # Validation error of the models

Phi_train = radial_basis_transform(X_trn, X_trn)

Phi_value = radial_basis_transform(X_val, X_trn)

Phi_test = radial_basis_transform(X_tst, X_trn)

plt.legend(['Validation Error', 'Test Error'], fontsize=16)

plt.legend(['Validation Error', 'Test Error'], fontsize=16)

plt.ylabel('Validation/Test error', fontsize=16)
plt.xticks(list(validationError.keys()), fontsize=12)

plt.ylabel('Validation/Test error', fontsize=16)
plt.xticks(list(validationError.keys()), fontsize=12)

plt.xlabel('Regularization Parameter', fontsize=16)
plt.ylabel('Validation/Test error', fontsize=16)

he test and validation error are lowest then.")

Gamma

Gamma

Regularization Parameter

X b = radial basis transform(x true, X trn)

a = [0.001, 0.01, 0.1, 1, 10, 100, 1000]

-2

Ò

You have to submit a single .py file that contains all the code.

Out[14]: <matplotlib.legend.Legend at 0x7fde9c1dad60>

plt.plot(x_true, y b, marker='None', linewidth=2)

plt.plot(x true, y true, marker='None', linewidth=5, color='k')

Validation Error

Test Error

test and validation error are lowest then.

plt.xticks(list(validationError.keys()), fontsize=12)

plt.legend(['Validation Error', 'Test Error'], fontsize=16)

Validation Error

Validation Error

Test Error

Test Error

q[d] = train_ridge_model(Phi_train, y_trn, d)

Test error of all the models

What are some ideal values of λ ?

b. (15 points)

c. (30 points)

testError = {}

while d <= 10 ** 3:

Plot all the models

plt.xlabel('Gamma', fontsize=16)

plt.xlabel('Gamma', fontsize=16)

plt.axis([0, 0.01, 20, 65])

plt.axis([0, 1, 20, 65])

plt.axis([0, 1000, 20, 65])

d = 10 ** -3

mensions

dimensions

plt.figure()

plt.figure()

plt.figure()

60

50

20

00,00 0.1

65 60

25

20

with λ ?

20

15

10

5

0

-5

-10

In []:

plt.figure()

d = 10 ** -3

while d <= 10 ** 3:

y b = X b @ q[d]

plt.legend(['true'] + a)

true 0.001

0.01

1 10

1000

-6

In [14]:

0000100.0

d. (10 points, Discussion)

Validation/Test error

0.0

Validation/Test error

Validation/Test error

for value in X:
 z = []

Phi.append(z)
Phi = np.array(Phi)

In [10]:

nonlinear (kernel) regression.

X float(n,): univariate data
B float(n,): basis functions

def radial basis transform(X, B, gamma=0.1):

for a in B:#range(0, np.size(X)):

#Phi = Phi.reshape(np.size(X), np.size(B))

Out[9]: (-8.0, 8.0, -15.0, 15.0)

5

-5

-10

-15

15

Polynomial degree

18

plt.plot(x true, y true, marker='None', linewidth=5, color='k')

plt.plot(x true, y d, marker='None', linewidth=2)

21

2. **Regression with Radial Basis Functions**, 70 points

In the previous case, we considered a nonlinear extension to linear regression using a linear combination of polynomial basis functions, where each basis function was introduced as a feature $\phi(x) = x^k$. Now, we consider Gaussian radial basis functions of the form:

of trying to identify the number of radial basis functions or their centers, we can treat **each data point as the center of a radial basis

 $\phi(\mathbf{x}) = e^{-\gamma (x-\mu)^2}$, whose shape is defined by its center μ and its width $\gamma > 0$. In the case of polynomial basis regression, the user's choice of the dimension d determined the transformation and the model. For radial basis regression, we have to contend with deciding how many radial basis functions we should have, and what their center and width parameters should be. For simplicity, let's assume that $\gamma = 0.1$ is fixed. Instead

 $f(x) = [w_1, w_2, w_3 \dots, w_n]^T \begin{vmatrix} e^{-\gamma (x - x_1)^2} \\ e^{-\gamma (x - x_2)^2} \\ e^{-\gamma (x - x_2)^2} \end{vmatrix}.$

This transformation uses radial basis functions centered around data points $e^{-\gamma (x-x_i)^2}$ and each basis function has a corresponding weight w_i associated with it, for all $i=1,\ldots,n$. We transform each univariate data point x_i into into a multivariate (n-dimensional) data point

 $\Phi = \begin{bmatrix} 1 & e^{-\gamma (x_1 - x_2)^2} & e^{-\gamma (x_1 - x_3)^2} & \dots & e^{-\gamma (x_1 - x_n)^2} \\ e^{-\gamma (x_2 - x_1)^2} & 1 & e^{-\gamma (x_2 - x_3)^2} & \dots & e^{-\gamma (x_2 - x_n)^2} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ e^{-\gamma (x_n - x_1)^2} & e^{-\gamma (x_n - x_2)^2} & e^{-\gamma (x_n - x_3)^2} & \dots & 1 \end{bmatrix}.$

via $\phi(x_i) \to [\dots, e^{-\gamma (x_i - x_i)^2}, \dots]$. When this transformation is applied to every data point, it produces the **radial-basis kernel**:

Complete the Python function below that takes univariate data as input and computes a radial-basis kernel. This transforms one-

z.append(math.e ** (-1 * gamma * (value - a) ** 2)) # or neg gamma not sure

Complete the Python function below that takes a radial-basis kernel matrix Φ , the labels y, and a regularization parameter $\lambda > 0$ as input

and learns weights via **ridge regression**. Specifically, given a radial-basis kernel matrix Φ , implement the computation of

w = np.linalg.inv(np.transpose(Phi) @ Phi + lam * np.eye(n)) @ np.transpose(Phi) @ y

#for d in range(np.power(10, -3), np.power(10, 3), step): # Iterate over polynomial degree

testError[d] = evaluate_model(Phi_test, y_tst, q[d]) # Evaluate model on test data

plt.plot(testError.keys(), testError.values(), marker='s', linewidth=3, markersize=12)

plt.plot(testError.keys(), testError.values(), marker='s', linewidth=3, markersize=12)

plt.plot(testError.keys(), testError.values(), marker='s', linewidth=3, markersize=12)

plt.plot(validationError.keys(), validationError.values(), marker='o', linewidth=3, markersize=12)

plt.plot(validationError.keys(), validationError.values(), marker='o', linewidth=3, markersize=12)

plt.plot(validationError.keys(), validationError.values(), marker='o', linewidth=3, markersize=12)

print ("It seems that the validation and test error relative to the regularization paramater have a patt ern similar to a log function, thus the most optimal value for lambda would be 10^{-3} or 10^{-2} as both t

It seems that the validation and test error relative to the regularization paramater have a pattern s imilar to a log function, thus the most optimal value for lambda would be 10^-3 or 10^-2 as both the

0.0

1.0

1000.0

Plot the learned models as well as the true model similar to the polynomial basis case above. How does the linearity of the model change

validationError[d] = evaluate_model(Phi_value, y_val, q[d]) # Evaluate model on validation data

Transform training data into d di

Transform validation data into d

Transform test data into d dimensions

Learn model on training data

As before, we can explore the tradeoff between **fit and complexity** by varying $\lambda \in [10^{-3}, 10^{-2} \cdots, 1, \cdots 10^{3}]$. For each model, train using the transformed training data (Φ) and evaluate its performance on the transformed validation and test data. Plot two curves: (i) λ vs.

gamma float : standard deviation / scaling of radial basis kernel

dimensional data into n-dimensional data in terms of Gaussian radial-basis functions centered at each data point and allows us to model

24

validationErr = {} # Validation error of the models

Phi_trn = polynomial_transform(X_trn, d)

Phi_val = polynomial_transform(X_val, d)

Phi tst = polynomial transform(X tst, d)

plt.xlabel('Polynomial degree', fontsize=16)
plt.ylabel('Validation/Test error', fontsize=16)
plt.xticks(list(validationErr.keys()), fontsize=12)

w[d] = train_model(Phi_trn, y_trn)

From plot of d vs. validation error below, which choice of d do you expect will generalize best?

Test error of all the models

Dictionary to store all the trained models

testErr[d] = evaluate_model(Phi_tst, y_tst, w[d]) # Evaluate model on test data

plt.plot(testErr.keys(), testErr.values(), marker='s', linewidth=3, markersize=12)

any degree between these two numbers would yield the best accuracy in my opinion")

egree between these two numbers would yield the best accuracy in my opinion

plt.plot(validationErr.keys(), validationErr.values(), marker='o', linewidth=3, markersize=12)

In [1]: