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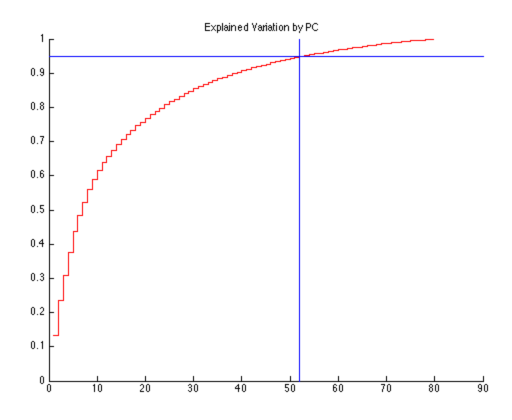
Question 1

```
load('CancerMicroarray.mat')
numTumors = length(G)
numGenes = length(X)
% There are 2308 different genes sequenced from 83 different tumors.
[coeffPC, scorePC, latentPC] = pca(X);
cumulativeLatent = cumsum(latentPC) / sum(latentPC);
numPC = max(find(cumulativeLatent < 0.95));</pre>
% You need 52 or 53 PCs to explain 95% of the data
figure;
hold on;
stairs(cumulativeLatent, 'Color', 'r');
title('Explained Variation by PC');
refline(0, 0.95);
line([52 52],[0 1]);
hold off;
linCValAcc = zeros(400,1);
quadCValAcc = zeros(400,1);
for i = 1:400
trainingindicies = randsample(1:83, 60);
testindicies = 1:83;
testindicies(ismember(testindicies,trainingindicies)) = [];
trainingdata = X(trainingindicies, :);
testdata = X(testindicies, :);
trainingclass = G(trainingindicies);
testclass = G(testindicies);
```

```
linClass = fitcdiscr(trainingscorePC(:, 1:10), trainingclass);
quadClass = fitcdiscr(trainingscorePC(:, 1:10), trainingclass, ...
    'DiscrimType', 'pseudoQuadratic');
predictedLinClass = predict(linClass, testdata(:,1:10));
predictedQuadClass = predict(quadClass, testdata(:,1:10));
linCValAcc(i) = mean(predictedLinClass == testclass');
quadCValAcc(i) = mean(predictedQuadClass == testclass');
end;
avgLinearCrossVal = mean(linCValAcc)
avgQuadraticCrossVal = mean(quadCValAcc)
% My mean linear and quadratic cross validated accuracies are 22.9% and
% 26.0%. This seems low, but I can't identify anything incorrect in my
% implementation of the PCA/fitcdiscr functionality so I assume there's
% some sort of logic error in my test/training data set creation in the
% loop, or the test data actually passed to predict. I was a little fuzzy
% Not sure, but it at least compiles and produces a number!
        numTumors =
            83
        numGenes =
                2308
        avgLinearCrossVal =
            0.2349
        avgQuadraticCrossVal =
            0.2667
```

[trainingcoeffPC, trainingscorePC, traininglatentPC] = pca(trainingdata);

2



```
% Decision trees are super cool. They seek to partition data space into a
% dichotomous outcome, is a, or is not a for example that minimizes
% "impurity". It does this recursively, producing some sort of tree that
% will have multiple branches looking like is a / is not a with is not a
% subdivided further. Classification trees deal with classifications alone,
% and then there are regression trees that can deal with continuous data.
% There are then CART trees, which can deal with both. (I have played with
% these a little bit in analyzing large ecological data sets where we were
% trying to determine disease load based upon multiple terrain and
% oceanographic types)
treeCVal = zeros(400,1);
for i = 1:400
trainingindicies = randsample(1:83, 60);
testindicies = 1:83;
testindicies(ismember(testindicies,trainingindicies)) = [];
trainingdata = X(trainingindicies, :);
testdata = X(testindicies, :);
trainingclass = G(trainingindicies);
testclass = G(testindicies);
[trainingcoeffPC, trainingscorePC, traininglatentPC] = pca(trainingdata);
```

```
treeDiscr = fitctree(trainingscorePC(:, 1:10), trainingclass);

treePredict = predict(treeDiscr, testdata(:, 1:10));

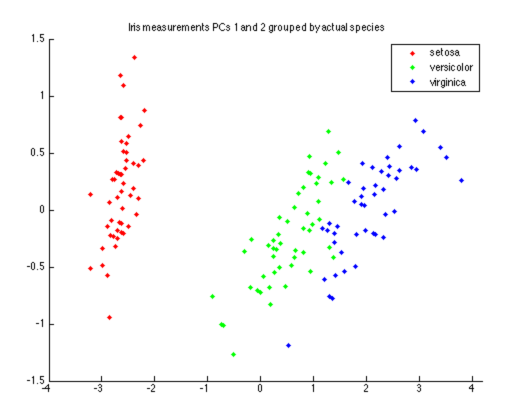
treeCVal(i) = mean(treePredict == testclass');

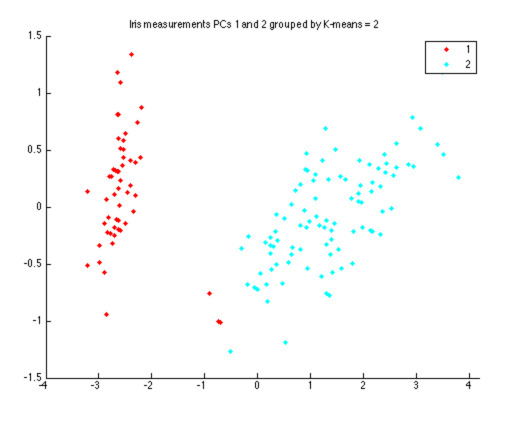
end
% the Decision tree has ~ 25% cross validated accuracy. Still somewhat % sucky. I'm pretty sure it's a problem with the predict function, but I % don't know how to fix it. I'm still unsure of what exactly the first 10 % PCs represents in my data, like how do I access the test data that % corresponds to those PCs, or is there any way to represent that data?
```

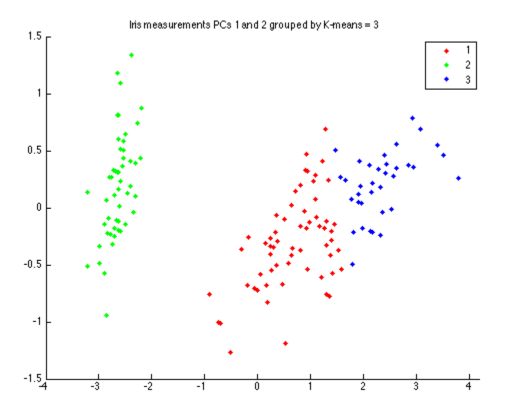
Question 2

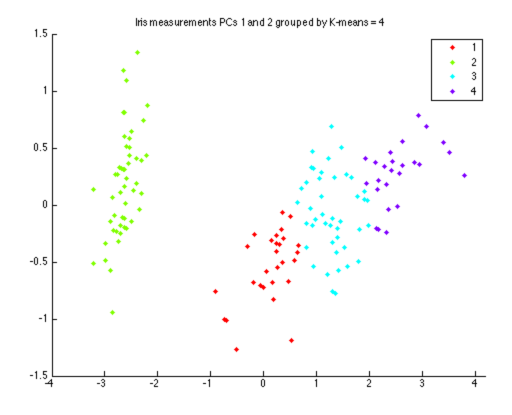
```
load('fisheriris');
numSpecies = length(unique(species));
numIris = length(species);
numMeas = size(meas,2);
% The measruements Ronald Fisher took were Petal Width, Petal Length, Sepal
% Width, and Sepal Length, measured in mm There were three different
% species, setsoa, versicolor, and virginica. There were 150 different
% flowers sampled.
[coeffPC2, scorePC2, latentPC2] = pca(meas);
figure;
gscatter(scorePC2(:, 1), scorePC2(:, 2), species);
title('Iris measurements PCs 1 and 2 grouped by actual species');
kmeans2 = kmeans(meas, 2);
figure;
gscatter(scorePC2(:, 1), scorePC2(:, 2), kmeans2);
title('Iris measurements PCs 1 and 2 grouped by K-means = 2');
kmeans3 = kmeans(meas, 3);
figure;
gscatter(scorePC2(:, 1), scorePC2(:, 2), kmeans3);
title('Iris measurements PCs 1 and 2 grouped by K-means = 3');
kmeans4 = kmeans(meas, 4);
figure;
gscatter(scorePC2(:, 1), scorePC2(:, 2), kmeans4);
title('Iris measurements PCs 1 and 2 grouped by K-means = 4');
% Determining the number of groups, k, is hard algorithmically. Some guys
% out of Stanford apparently have developed a Gap statistic that determines
% the k such that the Gap(k) >= Gap(k + 1) - sim error(k+1), It seems to
% work well. Some other guys ahve figured out an algorithm based on a
% statistical test that determines a K in which the data about each k
% centroid closest approximates a Gaussian distribution. There's also a way
```

% to do it graphically via silhouette graphs or information theory % critirions, but i'm well out of sentence now so lets just say those % exist.









Question 3

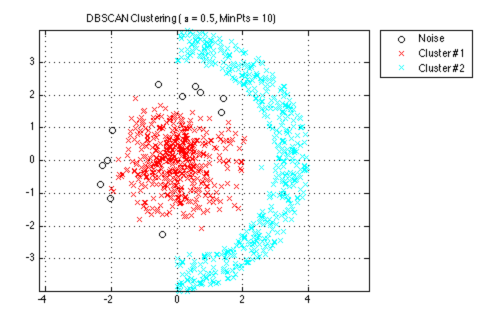
```
% Pt a.
% DBSCAN uses three different types of points, core points, border points,
% and noise points to determine clusters based upon the comparison of high
% density and low density regions. Core points have some minimum number of
% points within some radius epsilon, border points fall within a
% neighborhood of a core point, but do not have the minimum number of points
% to qualify as a core point, and noise points are any points which are not
% noise or border points. The general algoritmn is the classify points as
% one of the three, remove noise points, and then cluster based upon
% remaining points.
% Pt b.
load('mydata');
epsilon=0.5;
MinPts=10;
IDX=DBSCAN(X,epsilon,MinPts);
figure;
PlotClusterinResult(X, IDX);
title(['DBSCAN Clustering (\epsilon = ' num2str(epsilon) ...
```

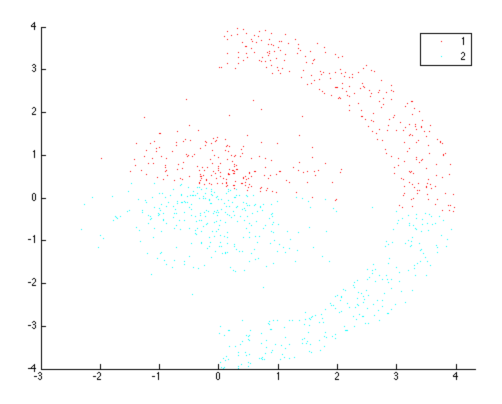
```
', MinPts = ' num2str(MinPts) ')']);

Q3kmeans = kmeans(X, 2);
figure;
gscatter(X(:, 1), X(:, 2), Q3kmeans);

% pt c.

% The DBSCAN algorithm is density based and has the ability to deal with
% non-linear groupings. Visible in the comparison between the two graphs is
% k-means needs to draw a straight line between the two groups, while
% DBSCAN can draw a non-straight/curved line. I found a thing that
% describes it as k-means can't deal with "non-globular" shapes.
```





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