INT104 ARTIFICIAL INTELLIGENCE

Review II

Fang Kang
Fang.kang@xjtlu.edu.cn



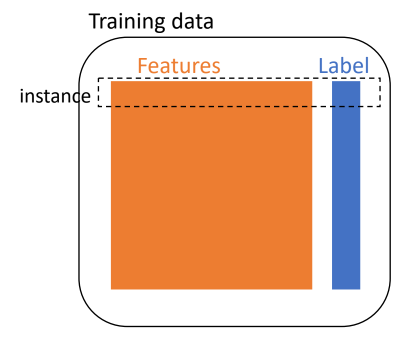


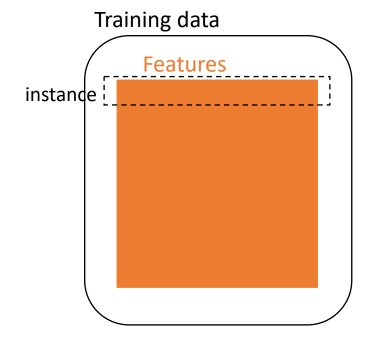
CONTENT

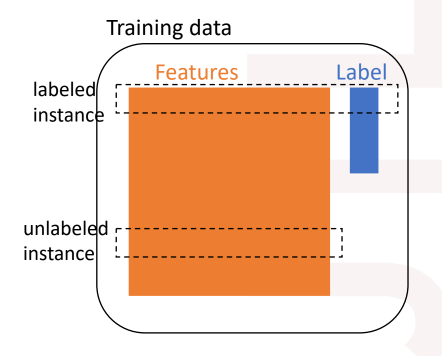
- Supervised methods
 - Classification and Regression
 - SVM
 - Decision Tree
 - Random Forest
- Unsupervised methods
 - K-means
 - Hierarchical clustering
 - GMM



Supervised vs. unsupervised







Supervised

Unsupervised

Semi-supervised

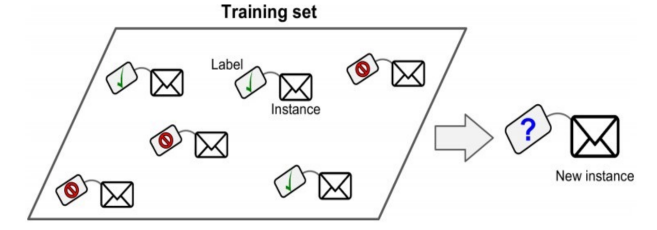




Classification

Classification: Classification algorithms find a function that determines which category the input data belongs to.

Binary Classification is a supervised learning algorithm that classifies new observations into one of two classes.

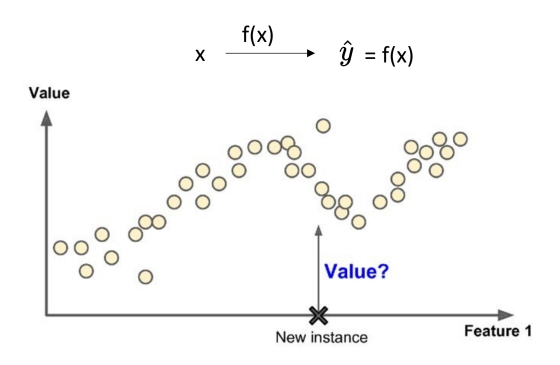


Multiclass/Multilabel Classification

- Multiclass classification refers to classification tasks that can distinguish between more than two classes.
- Multilabel classification refers to classification system that outputs multiple binary tags.

Regression

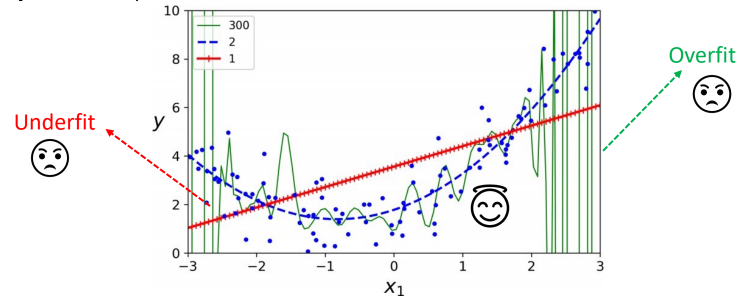
Regression attempts to determine the strength and character of the relationship between one dependent variable (usually denoted by Y) and a series of other variables (known as independent variables).





Learning Curves

If you perform high-degree Polynomial Regression, you will likely fit the training data much better than with plain Linear Regression. (Is high-degree polynomial always better?)



Bias: refers to the error from erroneous assumptions in the learning algorithm. (inability to capture the underlying patterns in the data).

Variance: refers an error from sensitivity to small fluctuations in the training data. (difference in fits between data sets)



Cross Validation

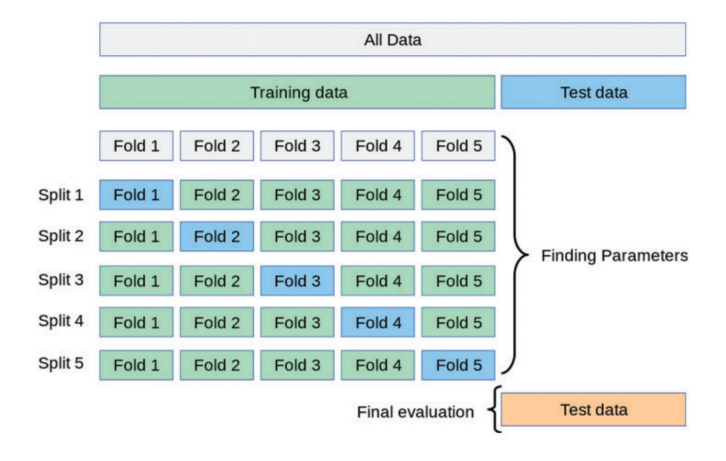
- Train/test/validation split
- To avoid selecting the parameters that perform best on the test data but maybe not the parameters that generalize best, we can further split the training set into training fold and validation fold
- Can maximize the accuracy on the training data



- Training fold: used to fit the model
- Validation fold: used to estimate prediction error for model selection
- Test set: used for assessment of the prediction error of the final chosen model

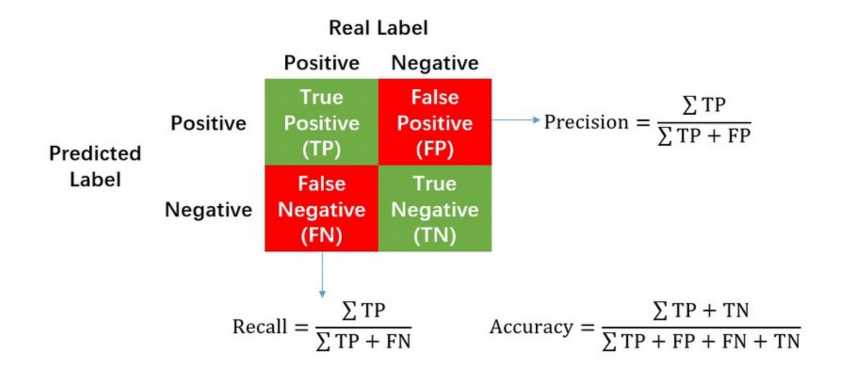


K-fold Cross-Validation





Confusion Matrix



 $F_1 = 2 * \frac{Precision * Recall}{Precision + Recall}$



Support Vector Machine (SVM)

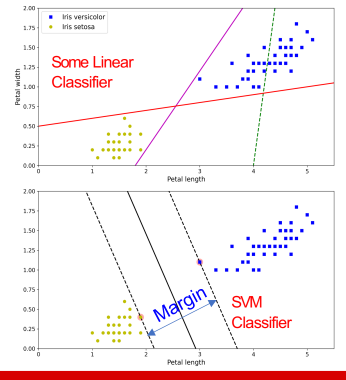
Linear SVM Classification

- Linear separability
- Fitting widest possible "street"

between classes

Performs better with new data

- Large Margin Classification
- Margin, Support Vectors



Hard Margin SVM

All instances being off the street and on the right side

Soft Margin SVM

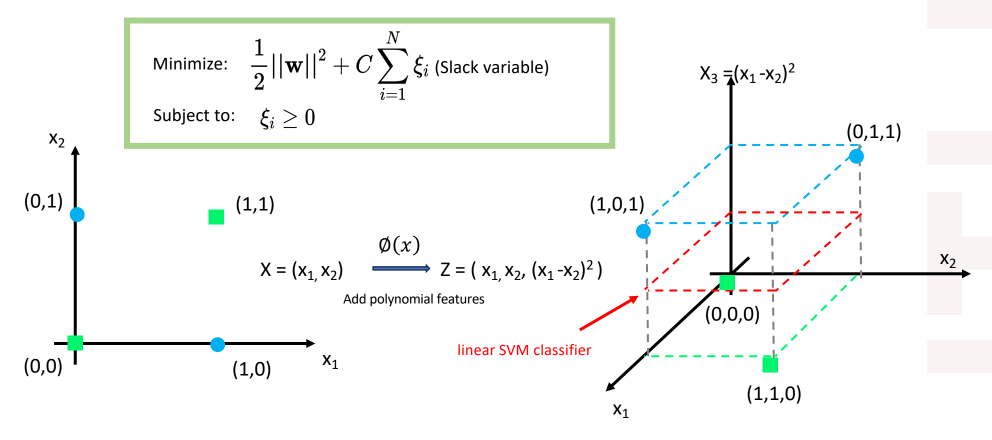
Allow margin violations

Support Vectors

- Decision boundary is not affected by more training instances
- It is determined by support vectors (instances located on the edge of street)



Nonlinear SVM Classification



Nonlinear transformation $\emptyset(x)$: not only one form

Cover's theorem: High-dimensional space is more likely to be linearly separable than in a low-dimensional space.



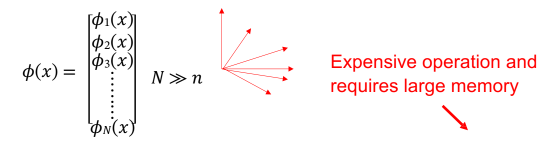
Nonlinear SVM: Kernel Trick

Input Space: dimension n

$$x = \begin{bmatrix} x_1 \\ x_2 \\ x_3 \\ \vdots \\ x_n \end{bmatrix}$$

$$K(oldsymbol{a},oldsymbol{b}) = oldsymbol{a}^Toldsymbol{b} = egin{bmatrix} a_1 & a_2 & a_3 & \cdots & a_n \end{bmatrix} egin{bmatrix} b_1 \ b_2 \ b_3 \ dots \ b_n \end{bmatrix}$$

High-dimensional Feature Space: dimension N >> n



$$K(\phi(oldsymbol{a}),\phi(oldsymbol{b})) = \phi(oldsymbol{a})^T\phi(oldsymbol{b}) = [\phi_1(a) \quad \phi_2(a) \quad \phi_3(a) \quad \dots \quad \phi_N(a)] egin{bmatrix} \phi_1(b) \ \phi_2(b) \ \phi_3(b) \ dots \ \phi_n(b) \end{bmatrix}$$

Kernel Trick

Common kernels:

 $\text{Linear:} K(\mathbf{a}, \mathbf{b}) = \mathbf{a}^{\top} \mathbf{b}$

 $\text{Polynomial:} K(\mathbf{a}, \mathbf{b}) = \left(\gamma \mathbf{a}^{\top} \mathbf{b} + r\right)^d$

Gaussian Radial Basis Function: $K(\mathbf{a}, \mathbf{b}) = \exp(-\gamma \|\mathbf{a} - \mathbf{b}\|^2)$

 $\text{Sigmoid: } K(\mathbf{a},\mathbf{b}) = \tanh \big(\gamma \mathbf{a}^{\top} \mathbf{b} + r \big)$

Universal approximator. Corresponding feature space $\phi(x)$ is infinite dimensional space

non-linearly separable data

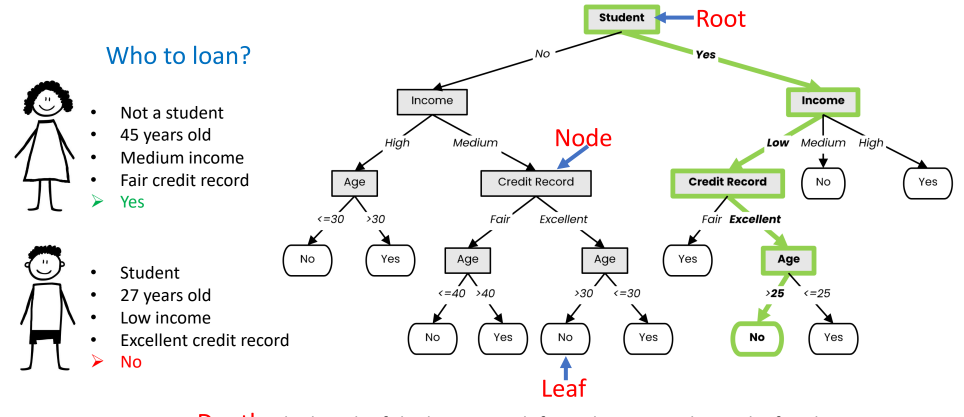


infinite-dimensional space



Decision Tree Definition

- A tree-like model that illustrates series of events leading to certain decisions
- Each node represents a test on an attribute and each branch is an outcome of that test



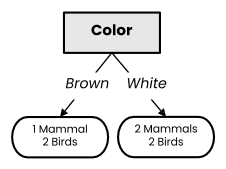
Depth: the length of the longest path from the root node to a leaf node

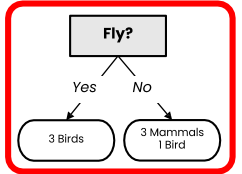


Best attribute = highest information gain

In practice, we compute entropy(X) only once!

Does it fly?	Color	Class
No	Brown	Mammal
No	White	Mammal
Yes	Brown	Bird
Yes	White	Bird
No	White	Mammal
No	Brown	Bird
Yes	White	Bird





$$entropy (X) = -p_{\text{mammal}} \log_2 p_{\text{mammal}} - p_{\text{bird}} \log_2 p_{\text{bird}} = -\frac{3}{7} \log_2 \frac{3}{7} - \frac{4}{7} \log_2 \frac{4}{7} \approx 0.985$$

$$entropy (X_{color=brown}) = -\frac{1}{3} \log_2 \frac{1}{3} - \frac{2}{3} \log_2 \frac{2}{3} \approx 0.918 \quad entropy (X_{color=white}) = 1$$

$$gain (X, color) = 0.985 - \frac{3}{7} \cdot 0.918 - \frac{4}{7} \cdot 1 \approx 0.020$$

$$entropy (X_{fly=yes}) = 0 \quad entropy (X_{fly=no}) = -\frac{3}{4} \log_2 \frac{3}{4} - \frac{1}{4} \log_2 \frac{1}{4} \approx 0.811$$

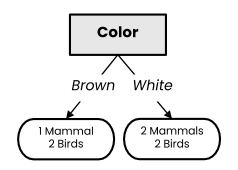
$$gain (X, fly) = 0.985 - \frac{3}{7} \cdot 0 - \frac{4}{7} \cdot 0.811 \approx 0.521$$

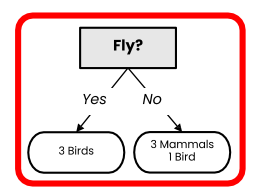


Best attribute = lowest Gini impurity

In practice, we compute gini(X) only once!

Does it fly?	Color	Class
No	Brown	Mammal
No	White	Mammal
Yes	Brown	Bird
Yes	White	Bird
No	White	Mammal
No	Brown	Bird
Yes	White	Bird





$$gini (X_{color=brown}) = 1 - \left(\frac{1}{3}\right)^{2} - \left(\frac{2}{3}\right)^{2} \approx 0.444 \qquad gini (X_{color=white}) = 0.5$$

$$gini (X, color) = \frac{3}{7} \cdot 0.444 + \frac{4}{7} \cdot 0.5 \approx 0.476$$

$$gini (X_{fly=yes}) = 0 \qquad gini (X_{fly=no}) = 1 - \left(\frac{3}{4}\right)^{2} - \left(\frac{1}{4}\right)^{2} \approx 0.375$$

$$gini (X, fly) = \frac{3}{7} \cdot 0 + \frac{4}{7} \cdot 0.375 \approx 0.214$$

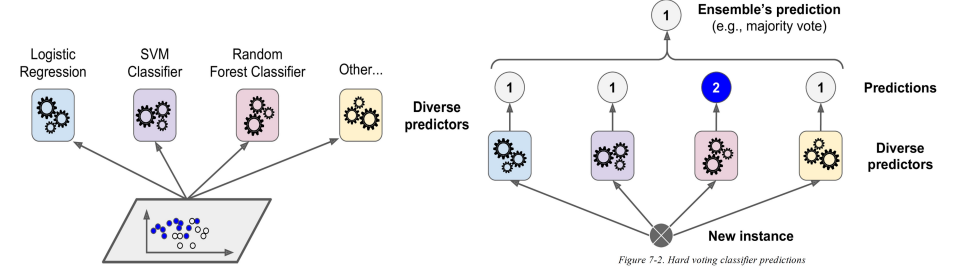


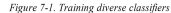
Ensemble Learning

Ensemble: A group of predictors

Voting Classifier

Hard Voting

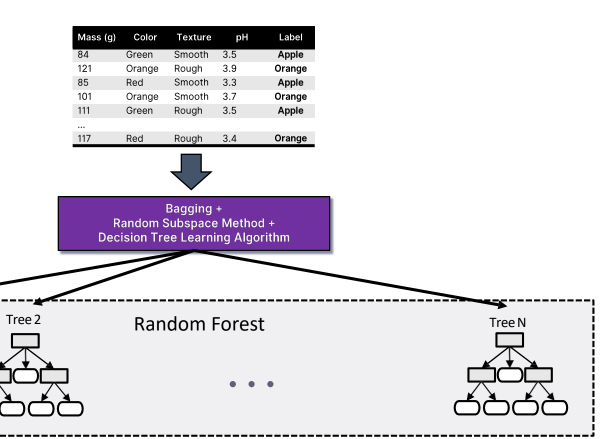






Random Forests

Tree 1





Ensemble method

- Random Forests are one of the most common examples of ensemble learning.
- Other commonly-used ensemble methods:
 - Bagging: multiple models on random subsets of data samples.
 - Random Subspace Method: multiple models on random subsets of features.
 - Boosting: train models iteratively, while making the current model focus on the mistakes of the previous ones by increasing the weight of misclassified samples.
 - Stacking: instead of using hard voting to aggregate the predictions of all predictors in an ensemble, train a model to perform this aggregation.



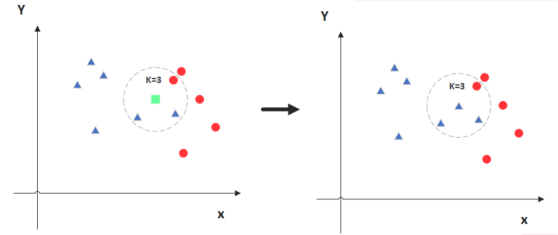
K Nearest Neighbors (KNN)

As in the general problem of classification, we have a set of data points for which we know the correct class labels

When we get a new data point, we compare it to each of our existing data points and find similarity

Take the most similar k data points (k nearest neighbours)

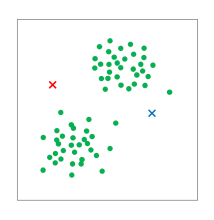
From these k data points, take the majority vote of their labels. The winning label is the label / class of the new data point

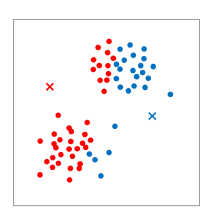


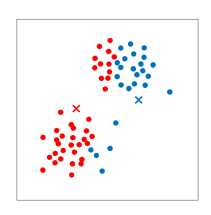


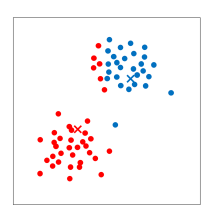


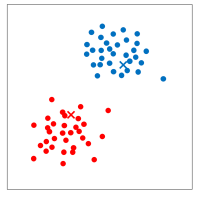
K-means clustering algorithm

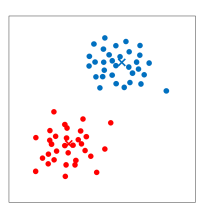












Euclidean distance $d(p,q)=\sqrt{(p_1-q_1)^2+(p_2-q_2)^2+\cdots+(p_n-q_n)^2}.$

Goal: Assign all data points to k clusters

Step 1: Pick k *random* initial cluster centroids

Step 2: Paint the data points that are closer to red centroid red, and those closer to blue centroid blue

Step 3: Update the positions of centroids

Red centroid := average of current red points

Blue centroid := average of current blue points

Repeat

Until no more pointes need to be repainted, i.e., the centroids no longer change

Clustering is done

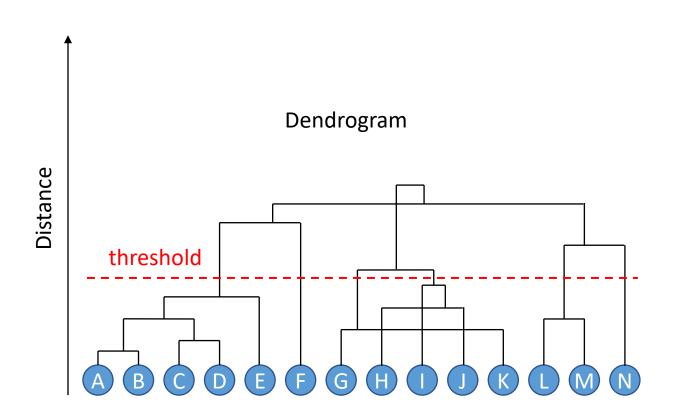


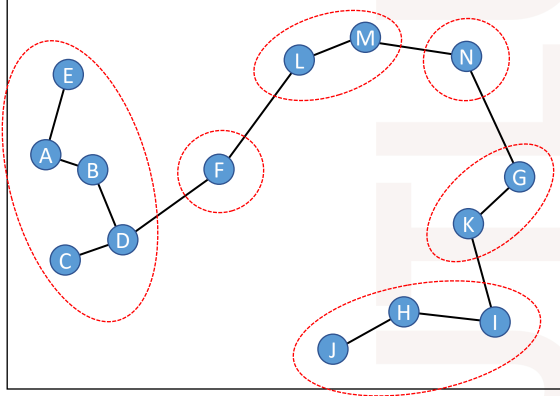
Hierarchical clustering

- Hierarchical Clustering is a set of clustering methods that aim at building a hierarchy of clusters
 - > A cluster is composed of smaller clusters
- There are two strategies for building the hierarchy of clusters:
 - Agglomerative (bottom-up): we start with each point in its own cluster and we merge pairs of clusters until only one cluster is formed.
 - > Divisive (top-down): we start with a single cluster containing the entire set of points and we recursively split until each point is in its own cluster.
- The most popular strategy in practical use is bottom-up (agglomerative)!



Agglomerative clustering example



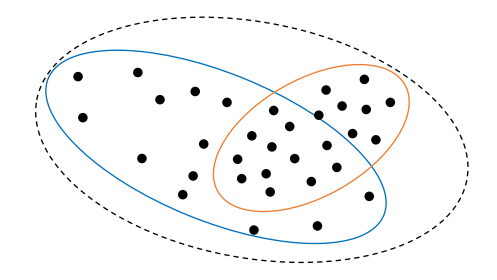




Gaussian mixture model (GMM)

K-means make <u>hard</u> assignments to data points: $x^{(i)}$ must belong to one of the clusters 1,2, \cdots , K

Sometimes, one data point can belong to multiple clusters



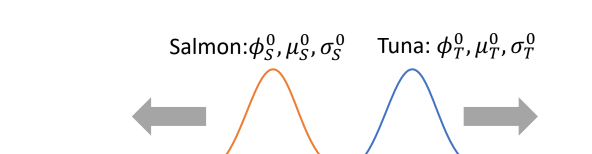
- Clusters may overlap
- Hard assignment may be simplistic
- Need a <u>soft</u> assignment:
 data points belong to clusters with different **probabilities**

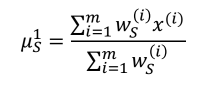


Demonstration with k = 2, 1-D Gaussian

Maximize likelihood of the whole data: $\mathcal{L}(\theta) = p(X|\Theta) = \prod_{i=1}^{m} p\big(x^{(i)}, z^{(i)}\big|\Theta\big) = \prod_{i=1}^{m} \sum_{z^{(i)}} Q_i(z^{(i)}) \frac{p(x^{(i)}, z^{(i)}|\Theta)}{Q_i(z^{(i)})}$

Random initialization





$$\mu_T^1 = \frac{\sum_{i=1}^m w_T^{(i)} x^{(i)}}{\sum_{i=1}^m w_S^{(i)}}$$



M-step



These points tend to have larger $w_S^{(i)}$

 $w_S^{(i)} = 0.6$

 $w_T^{(i)} = 0.4$

These points tend to have larger $w_T^{(i)}$

Weight (lbs)



