# CPSC 340: Machine Learning and Data Mining

**Ensemble Methods** 

## Admin

- Welcome to the course!
  - Everyone on the waitlist made it in.
- Assignment 1 is due Sunday.
- Tutorial materials available on GitHub
  - See <a href="https://github.ubc.ca/cpsc340/home">https://github.ubc.ca/cpsc340/home</a>
- Office hours happen every day (see course website)
- (Persistent) feedback URL:
  - http://skaha.cs.ubc.ca:11616/cpsc340/
  - Link available from course website.
- Piazza Q&A guidelines added to course website

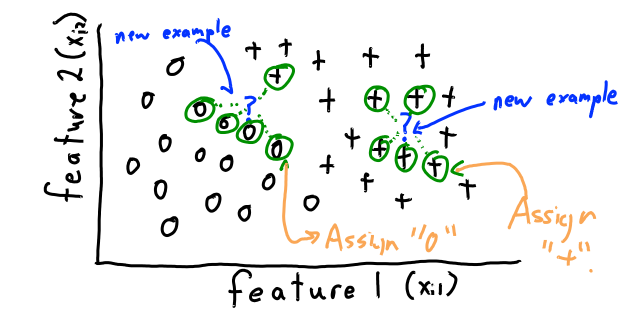
## Piazza guidelines (added to website)

- avoid including answers or partial answers in your question/post.
- do a quick search to make sure your question hasn't already been answered.
  - you can filter by tags like "hw1" or do a keyword search
- if you have a code problem, include relevant information like:
  - your version of Python and relevant libraries
  - your operating system
  - the command you used to execute the program
  - the entire error message
- (less important, but nice) when using equations, write them in LaTeX syntax and surround them with two dollar-signs. For example \$\$2^x\$\$ will render nicely in Piazza.

## K-Nearest Neighbours (KNN)

- K-nearest neighbours algorithm for classifying 'x':
  - Find 'k' values of  $x_i$  that are most similar to x.
  - Use mode of corresponding y<sub>i</sub>.

- Non-parametric:
  - Size of model grows with 'n'.



- Consistency:
  - Nearly-optimal test error with infinite data.
- But how many examples are needed?

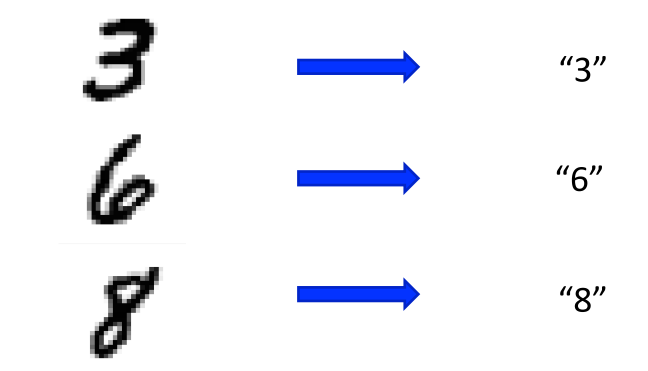
## **Curse of Dimensionality**

- "Curse of dimensionality": problems with high-dimensional spaces.
  - Volume of space grows exponentially with dimension.
  - Need exponentially more points to 'fill' a high-dimensional volume.
  - Distances become less meaningful:
    - All vectors may have similar distances.
  - Emergence of "hubs":
    - some datapoints are neighbours to many more points than average.
- KNN is also problematic if features have different scales.

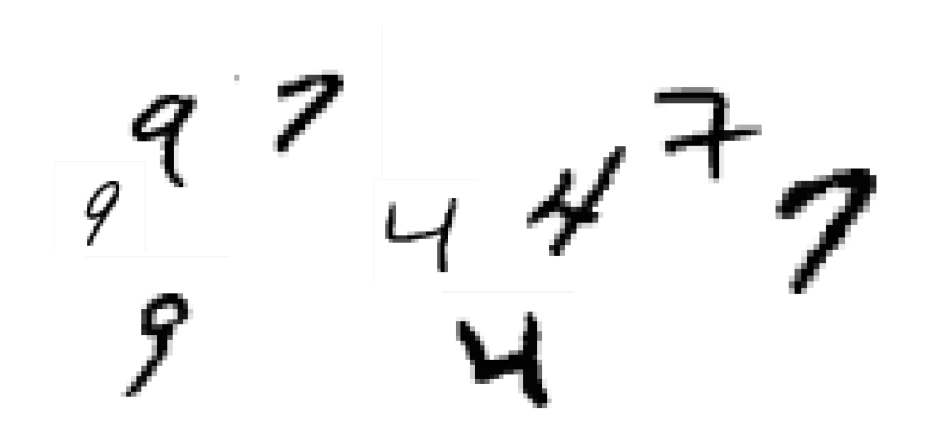
Nevertheless, KNN is really easy to use and often hard to beat!

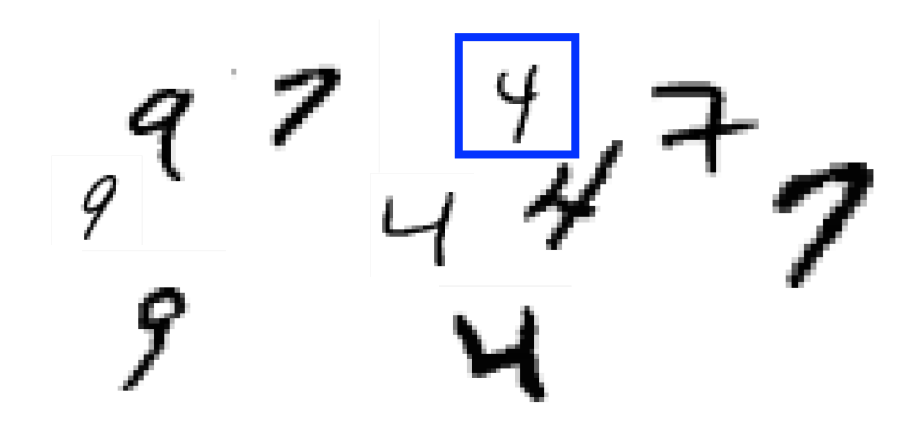
## Application: Optical Character Recognition

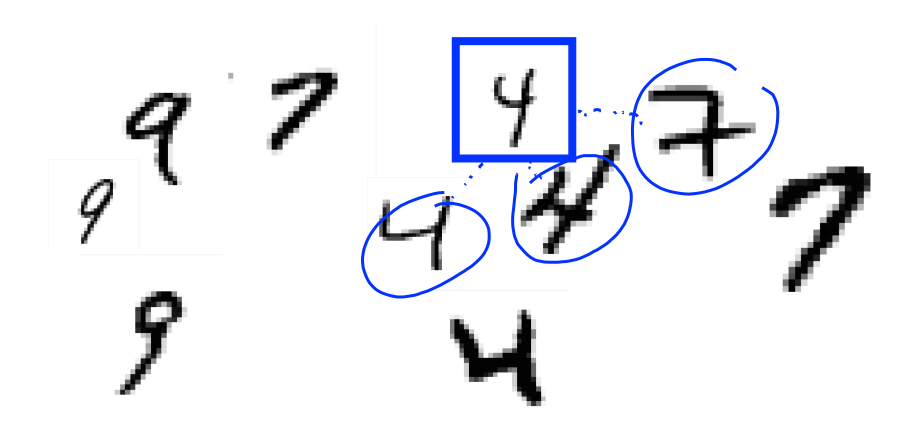
We have collection of letter/digit images, and corresponding labels:

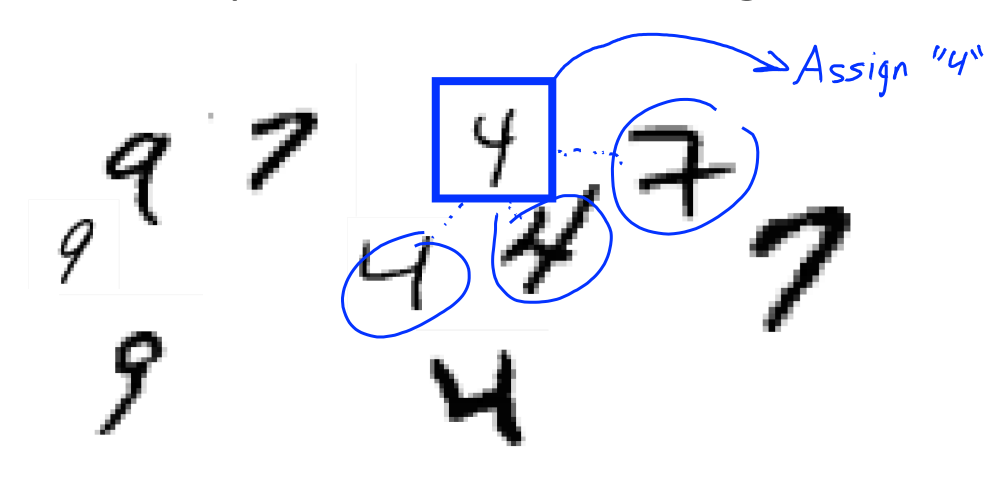


- Use supervised learning to automatically recognize letters/digits:
  - $-y_i$  could be the letter/digit,  $x_i$  could be the values of the pixels.









## Human vs. Machine Perception

There is huge difference between what we see and what KNN sees:

What we see:

3

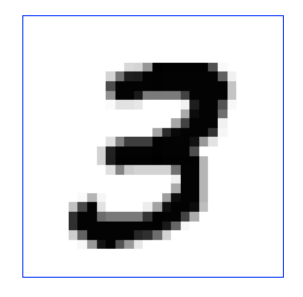
What the computer "sees":

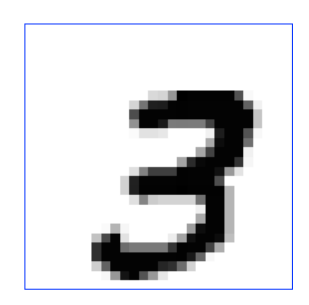


Actually, it's worse:

## What the Computer Sees

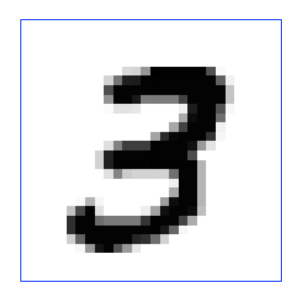
Are these two images 'similar'?

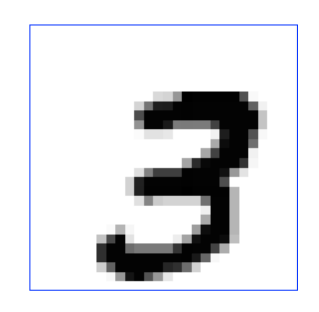




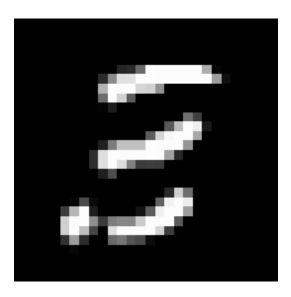
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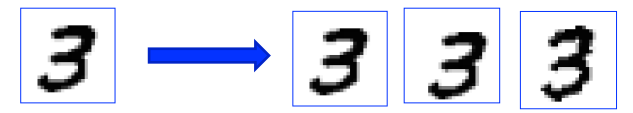
#### Difference:



KNN does not know that labels should be translation invariant.

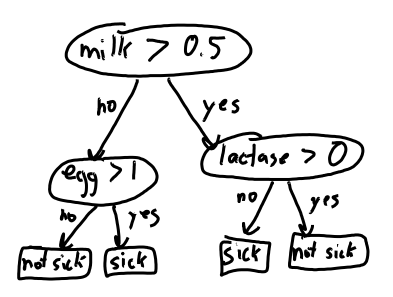
## **Encouraging Invariance**

- May want classifier to be invariant to certain feature transforms.
  - Digits: translations, small rotations, changes in size, mild warping,...
- The hard/slow way is to modify your distance function:
  - Find neighbours that require the 'smallest' transformation of image.
- The easy/fast way is to just add transformed data during training:
  - Add translated/rotate/resized/warped versions of training images.



Crucial part of many successful vision systems.

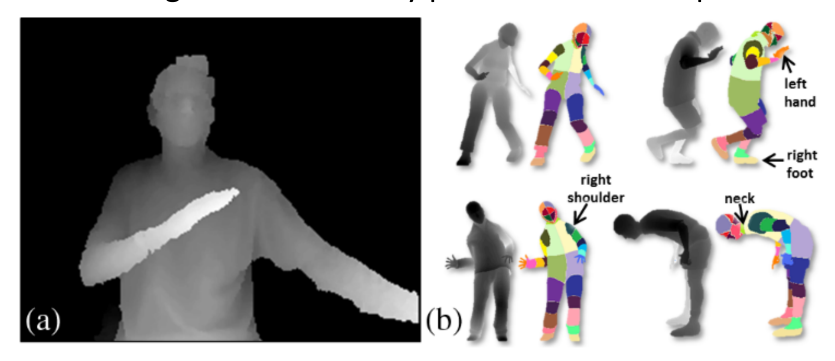
## Decision Trees vs. Naïve Bayes vs. KNN



$$(milk = 0.6, egg = 2, lactase = 0, ?)$$
 is close to  $(milk = 0.7, egg = 2, lactase = 0, sick)$  so predict sick.

## Application: Body-Part Recognition

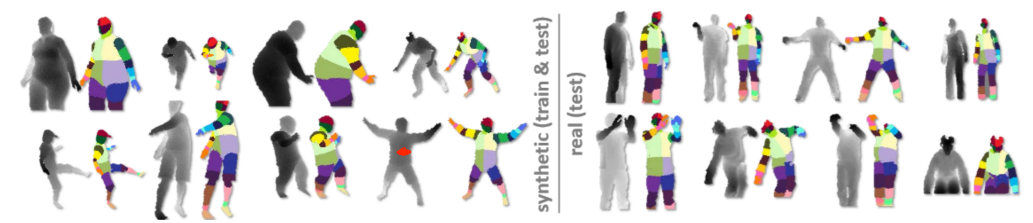
- Microsoft Kinect:
  - Real-time recognition of 31 body parts from laser depth data.



How could we write a program to do this?

## Some Ingredients of Kinect

- 1. Collect hundreds of thousands of labeled images (motion capture).
  - Variety of pose, age, shape, clothing, and crop.
- 2. Build a simulator that fills space of images by making even more images.



- Extract features of each location, that are cheap enough for real-time calculation (depth differences between pixel and pixels nearby.)
- 4. Treat classifying body part of a pixel as a supervised learning problem.
- 5. Run classifier in parallel on all pixels using graphical processing unit (GPU).

## Supervised Learning Step

ALL steps are important, but we'll focus on the learning step.

- Do we have any classifiers that are accurate and run in real time?
  - Decision trees and naïve Bayes are fast, but often not very accurate.
  - KNN is often accurate, but not very fast.

Deployed system uses an ensemble method called random forests.

## **Ensemble Methods**

- Ensemble methods are classifiers that have classifiers as input.
  - Also called "meta-learning".
- They have the best names:
  - Averaging.
  - Boosting.
  - Bootstrapping.
  - Bagging.
  - Cascading.
  - Random Forests.
  - Stacking.
- Meta-classifier often have higher accuracy than input classifiers.

## **Ensemble Methods**

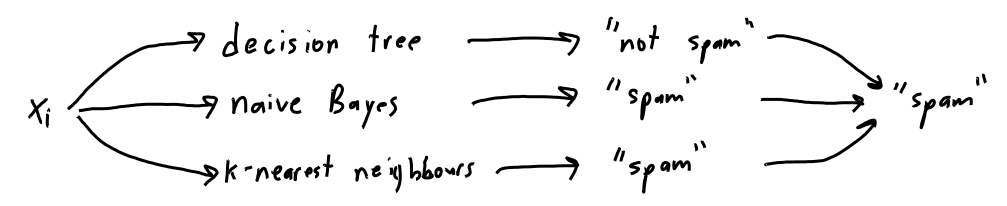
- Remember the fundamental trade-off:
  - 1. How small you can make the training error. vs.
  - 2. How well training error approximates the test error.
- Goal of ensemble methods is that meta-classifier:
  - Does much better on one of these than individual classifiers.
  - Doesn't do too much worse on the other.
- This (roughly) gives two types:
  - 1. Boosting: take simple classifier that underfits, improve its training error.
  - 2. Averaging: take complex classifier that overfits, improve its test error.

## Boosting (AdaBoost)

- Input to boosting is classifier that:
  - Is simple enough that it doesn't overfit much.
  - Can obtain >50% weighted training accuracy (for binary classification).
- Example: decision stumps or low-depth decision trees.
- Basic steps:
  - 1. Fit a classifier on the training data.
  - 2. Give a higher weight to examples that the classifier got wrong.
  - 3. Fit a classifier on the weighted training data.
  - 4. Go back to 2.
- Final prediction: weighted vote of individual classifier predictions.
- Boosted decision trees are very fast/accurate classifiers.

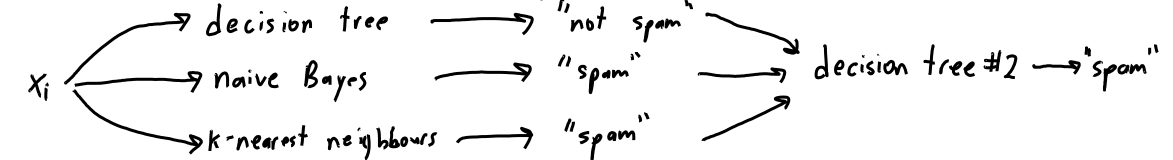
## Averaging

- Input to averaging is the predictions of a set of models:
  - Decision trees make one prediction.
  - Naïve Bayes makes another prediction.
  - KNN makes another prediction.
- Simple model averaging:
  - Take the mode of the predictions (or average if probabilistic).



## Averaging

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- Stacking:
  - Fit another classifier that uses the predictions as features.



## Averaging

- Averaging often performs better than individual models:
  - Averaging typically used by Kaggle winners.
  - E.g., Netflix \$1M user-rating competition winner was stacked classifier.
- Why does this work?
- Consider a set of classifiers that tend to overfit:
  - For example, deep decision trees.
- If they all overfit in exactly the same way, averaging does nothing.
- But if they make independent errors:
  - Probability of error of average can be lower than individual classifiers.
- Less attention on specific overfitting of each classifier.

## Random Forests

- Random forests average a set of deep decision trees.
  - Tend to be one of the best 'out of the box' classifiers.
    - Often close to the best performance of any method on the first run.
  - And predictions are very fast.
- Do deep decision trees make independent errors?
  - If just fit a decision tree repeatedly to same data, all trees will be the same.
- Two key ingredients in random forests:
  - Bootstrap aggregation (randomly sampling examples).
  - Random trees (randomly sampling features).

## Random Forest Ingredient 1: Bagging

- Bootstrap sample of a list of 'n' objects:
  - A set of 'n' objects, chosen independently with replacement.
  - Gives new dataset of 'n' objects, with some duplicated and some missing.
    - ~63% of original objects will be included.
  - Usually, it is used to estimate how sensitive a statistic is to the data.
- Bootstrap aggregation (bagging):

  - At test time, average the predictions.

Jootstrap aggregation (Dagging).

- Generate several bootstrap samples of the objects (x<sub>i</sub>,y<sub>i</sub>).

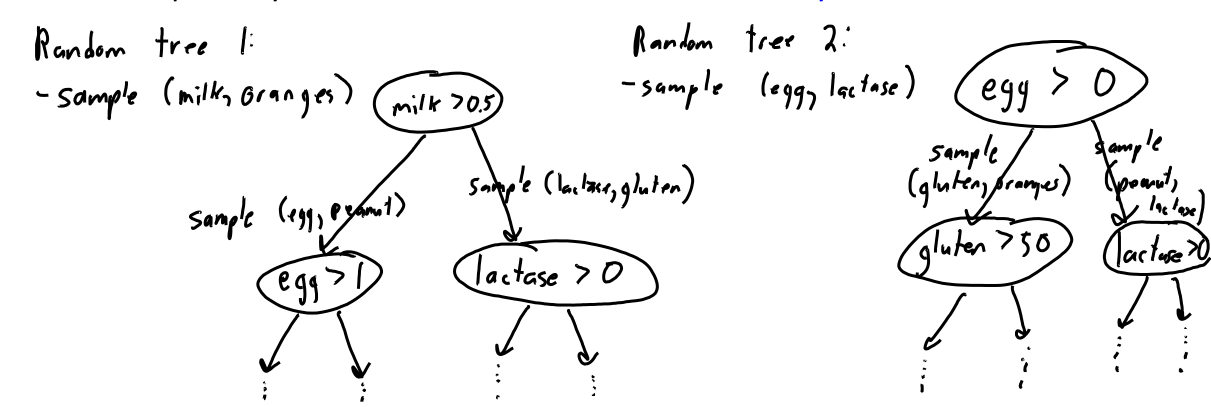
Decision trees will make different splits.

## Random Forest Ingredient 2: Random Trees

- When fitting each decision stump to construct deep decision tree:
  - Do not consider all features when searching for the optimal rule.
  - Each split only considers a small number of randomly-chosen features.

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  - Each split only considers a small number of randomly-chosen features.
- These random trees will tend to be very different from each other.
  - They will still overfit, but in \*different\* ways.
- The average tends to have a much lower test error.
- Empirically, random forests are one of the "best" classifiers.
- Fernandez-Delgado et al. [2014]:
  - Compared 179 classifiers on 121 datasets.
  - Random forests are most likely to be the best classifier.

## Summary

#### 1. Encouraging invariance:

- Add transformed data to be insensitive to the transformation.
- 2. Ensemble methods take classifiers as inputs.
- 3. Boosting turns 'weak' classifiers into 'strong' classifiers.
- 4. Averaging predictions often improves performance.
- 5. Random forests:
  - Averaging of deep randomized decision trees.
  - One of the best "out of the box" classifiers.

#### Next time:

We start unsupervised learning.



- All the remaining slides are "bonus".
- We may go through them briefly, if time permits.

## Bonus Slide: Why does Bootstrapping give 63%?

Probability of an arbitrary x<sub>i</sub> being selected in a bootstrap sample:

p(selected at least once in 'n' trials)

$$= |-p(not \text{ selected in any of 'n' trials})$$

$$= |-(p(not \text{ selected in one trial}))^n \qquad (trials \text{ are independent})$$

$$= |-(1 - 1/n)^n \qquad (prob = \frac{n-1}{n} \text{ for choosing any of the } n-1 \text{ other sample})$$

$$\approx |-1/e| \qquad (1-1/n)^n \rightarrow e^{-1} \text{ as } n \rightarrow \infty$$

## Bonus Slide: Why Random Forests Work

- Consider 'k' independent classifiers, whose errors have a variance of  $\sigma^2$ .
- If the errors are IID, the variance of the average is  $\sigma^2/k$ .
  - So the more classifiers you average, the more you decrease error variance.
     (And the more the training error approximates the test error.)
- Generalization to case where classifiers are not independent is:

$$co^2 + \underbrace{(1-c)o^2}_{k}$$

- Where 'c' is the correlation.
- So the decreasing correlation gets you closer to independent case.
- Randomization in random forests decreases correlation between trees.

## Bonus Slide: Bayesian Model Averaging

- Recall the key observation regarding ensemble methods:
  - If models overfit in "different" ways, averaging gives better performance.
- But should all models get equal weight?
  - E.g., decision trees of different depths, when lower depths have low training error.
  - E.g., a random forest where one tree does very well (on validation error) and others do horribly.
  - In science, research may be fraudulent or not based on evidence.
- In these cases, naïve averaging may do worse.

## Bonus Slide: Bayesian Model Averaging

- Suppose we have a set of 'm' probabilistic binary classifiers w<sub>i</sub>.
- If each one gets equal weight, then we predict using:

$$p(y_i|x_i) = \frac{1}{m}p(y_i|w_i,x_i) + \frac{1}{m}p(y_i|w_i,x_i) + \cdots + (\frac{1}{m})p(y_i|w_m,x_i)$$

• Bayesian model averaging treats model 'w<sub>j</sub>' as a random variable."

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$$P(y_{i}|x_{i}) = \underbrace{E}_{j=1}^{\infty} p(y_{i})w_{j}|x_{i}) = \underbrace{E}_{j=1}^{\infty} p(y_{i}|w_{j},x_{j})p(w_{j}|x_{i}) = \underbrace{E}_{j=1}^{\infty} p(y_{i}|x_{i})p(w_{j}|x_{i}) = \underbrace{E}_{j=1}^{\infty} p(y_{i}|x_{i})p(w_$$

- So we should weight by probability that  $w_i$  is the correct model:
  - Equal weights assume all models are equally probable.

# Bonus Slide: Bayesian Model Averaging

Can get better weights by conditioning on training set:

$$p(w_j|X,y) \propto p(y|w_j,X)p(w_j|X) = p(y|w_j,X)p(w_j)$$

- The 'likelihood' p(y | w<sub>i</sub>, X) makes sense:
  - We should give more weight to models that predict 'y' well.
  - Note that hidden denominator penalizes complex models.
- The 'prior' p(w<sub>i</sub>) is our 'belief' that w<sub>i</sub> is the correct model.
- This is how rules of probability say we should weigh models.
  - The 'correct' way to predict given what we know.
  - But it makes people uncomfortable because it is subjective.