



# *Machine Learning*

CS 485



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# Preface

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Since the course is online, we are watching recordings from a previous offering. Videos are available on <https://www.newworldai.com/understanding-machine-learning-course/>. The textbook for this course is [Understanding Machine Learning: From Theory to Algorithms](#).

Some notations:

- $D[A]$  denotes the probability hitting the set  $A$ .

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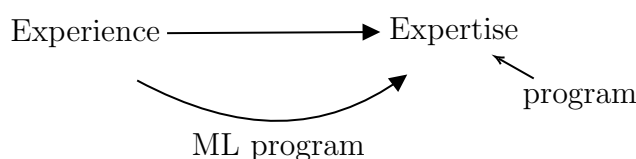
# Introduction

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## Reading

Up to page 41 of the textbook.

What is learning?



Process takes us from experience and leads us to expertise. Expertise would be another program that can do something you need expertise to do. For example, develop a spam filter. The outcome program is the spam filter.

## 1.1 Learning in Nature

**Bait Shyness:** It's difficult to poison the rats with the bait food. The rats will find that the shape might be different. If they take a bite and feel sick, they will immediately associate the sickness with the food and then never touch it again. It's a clear example of learning from a single experience.

**Spam filters:** Inputs are emails which are labeled.

$(\text{email1}, \text{spam}), (\text{email2}, \text{not spam}), \dots$

Then we have to come up with the program which filters the spam. The simplest way is to **memorize** all the emails that are spam. So what's wrong with such a program?

It does not generalize. We want **generalization**. Memorization is not enough. Generalization is sometimes called **inductive reasoning**: take previous cases and try to extend it to something new.

**Pigeon Superstition:** discovered by Skinner in 1947. He took a collection of pigeons and put them in the cage. Also he put different kinds of toys. Above the cage, there is

some mechanism that can spread grains. Something interesting happens. When the birds get hungry, they pick around for worms. Suddenly there's a spread of food. The birds start to learn: maybe the toys the bird is picking at that particular moment had some influence on getting food. So the next time the bird is hungry, the bird is more likely to pick on this toy than others. Then the next time food spreads, it reinforces what the bird did. After several times, the birds are completely devoted to some specific toys.

This is silly generalization. For rats, it's important generalization making them survive.

Garcia 1996, looks at the rats again. He gave the rats the poisoned bait which smelled and looked exactly like the usual food they get. Then the question: does the rat learn the connection between sickness and the poisoned food? Rats fail to associate the bell ringing with the poison effect. Here note that unlike the Pavlov's dog experiment which did repeatedly many times, the rat only has one chance to learn.

The key point here is prior knowledge: the rat already knows the shape and smell of the food through generation. Why have this limitation, why not paying attention to everything? In terms of rats, if they feel sick, every experience/feed is special, then the rat don't know what to associate to. Therefore, the prior knowledge is very important.

If we have little prior knowledge, we need a lot of training. If we have much prior knowledge, maybe we can do without much experiences. ML is living somewhere between these two.

Why do we need Machine Learning?

1. Some tasks that we (animals) can carry out may be too complex to program. E.g., Spam filter, driving, speech recognition.
2. Tasks that require experience with amounts of data that are beyond human capabilities. E.g., ads placement, genetic data.
3. Adaptivity.

## 1.2 Many types of machine learning

1. Supervised vs. Unsupervised. Supervised: spam filter. Unsupervised: outlier detection, clustering.

There's also an intermediate scenario called reinforcement learning.

2. Batch vs. Online. Batch: get all training data in advance. Online: need to response as you learn.
3. Cooperative  $\rightarrow$  indifferent  $\rightarrow$  adversarial. Teacher.
4. Passive vs. Active learner.

## 1.3 Relationships to other fields

**AI:** two important differences: We are going beyond what human/animals can do, not try to imitate; This area is rigorous, mathematical, nothing like "happens to be".

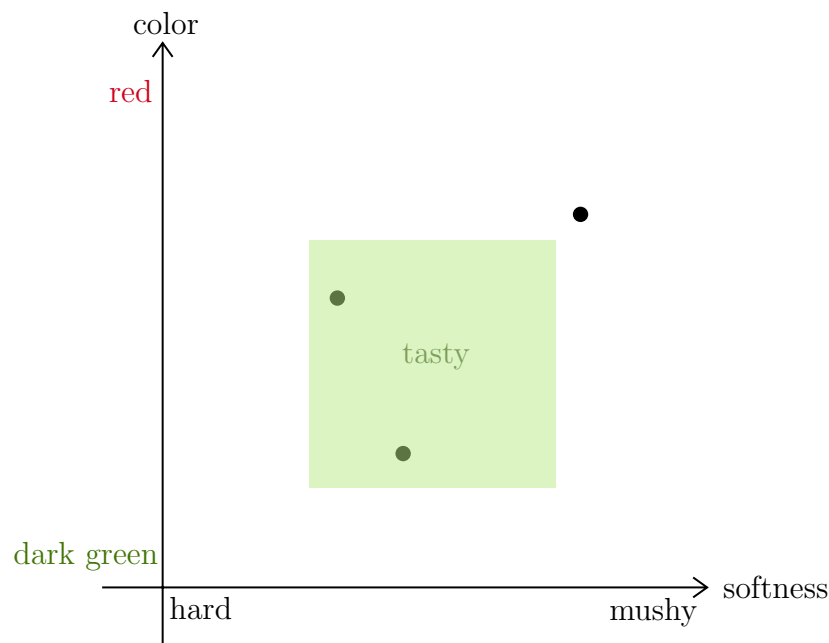
Also we need: Algorithms & complexity, statistics, linear algebra, combinatorics, optimization. However, there's something different with statistics in several ways.

1. algorithmic statistics
2. Distribution free. No clue on how spam is generated.
3. finite samples.

## Outline of the course

- Principles: supervised, batch, ...
- Algorithmic paradigms.
- Other types of learning.

### 1.4 Papaya Tasting



Each papaya corresponds to a coordinate  $(c, s)$ .

Training data:  $(x_1, y_1), \dots, (x_m, y_m) = S$ , where  $x_i \in \mathbb{R}^2$  and  $y_i \in \{T, N\}$

Domain set:  $[0, 1]^2$

Label set:  $\{T, N\}$

Output:  $f : [0, 1]^2 \rightarrow \{T, N\}$ . Prediction rule.

Assumption about data generation:

1. Training data are randomly generated.
2. Reliability by rectangles. See the picture above.

Measure of success: probability of my predictor  $f$  to err on randomly generated papaya.

# A Gentle Start

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## 2.1 Formal model for learning

In the context of papaya example last time,

Domain set $X$	$[0, 1]^2$
Label set $y$	$\{T, N\}$
Training input (sample) $S = ((x_1, y_1), \dots, (x_m, y_m))$	set of already tasted papayas
Learners output $h : X \rightarrow Y$	Prediction rule for tasteness
The quality of such $h$ is determined with respect to some data generating distribution and labeling rule	$L_{D,f}(h) = D[\{x : h(x) \neq f(x)\}]$ where $L$ stands for loss, $D$ is the distribution of papaya generated in the world, $f$ is the function to determine the true tastefulness of papaya. So it determines the probability that our hypothesis $h$ fails

The goal of the learner is given  $S$  to come up with  $h$  with small loss.

## 2.2 Empirical Risk Minimization

Basic learning strategy: **empirical risk minimization** (ERM) which minimizes the empirical loss.

We define **empirical loss (risk)** over a sample  $S$ :

$$L_S(h) = \frac{|\{i : h(x_i) \neq y_i\}|}{|S|}$$

A very simple rule for finding  $h$  with small empirical risk (ER)

$$h_S(x) \triangleq \begin{cases} y_i & \text{if } x = x_i \text{ for some } (x_i, y_i) \in S \\ N & \text{otherwise} \end{cases}$$

Then  $L_S(h_S) = 0$ .

Although the ERM rule seems very natural, without being careful, this approach may fail miserably. It **overfits** our sample.

To guard against overfitting we introduce some prior knowledge (Inductive Bias).

There exists a good prediction rule that is some axis aligned rectangle. Let  $H$  denote a fixed collection of potential (candidate) prediction rules, i.e.,

$$H \subseteq \{f : X \rightarrow Y\} = X^Y,$$

and we call it **hypothesis class**. Then we have a revised learning rule:  $\text{ERM}_H$  - “pick  $h \in H$  that minimizes  $L_S(h)$ ”, i.e.,

$$\text{ERM}_H(s) \in \underset{h \in H}{\text{argmin}} \{L_S(h)\}$$

### Theorem 2.1

Let  $X$  be any set,  $Y = \{0, 1\}$ , and let  $H$  be a finite set of functions from  $X$  to  $Y$ . Assume:

1. The training sample  $S$  is generated by some probability distribution  $D$  over  $X$  and labeled some  $f \in H$ , and elements of  $S$  are picked i.i.d.<sup>a</sup>
2. **Realizability assumption:**  $\exists h \in H$  such that  $L_{D,f}(h) = 0$

Then  $\text{ERM}_H$  is guaranteed to come up with an  $h$  that has small true loss, given sufficiently large sample  $S$ .

<sup>a</sup>identically and independently distributed

### Remark:

This is quite different from hypothesis testing. Unlike hypothesis testing, here we have assumptions after seeing the data. We are developing theories based on the data, and here  $H$  is a finite set.

### Proof:

Confusing samples are those on which  $\text{ERM}_H$  may come up with a bad  $h$ .

Fix some success parameter  $\varepsilon > 0$ , and the set of confusing  $S$ 's is

$$\{S : L_{D,f}(h_S) > \varepsilon\}$$

We wish to upper bound the probability of getting such a bad sample.

$$D^m[\{S|_X : L_{D,f}(h_S) > \varepsilon\}]$$

where  $S|_X = x_1, \dots, x_m$ .

Consider  $H_B = \{h \in H : L_{D,f}(h) > \varepsilon\}$  which is the set we want to avoid.

The misleading samples is the set of samples that may lead to an out come in  $H_B$ , formally:

$$M = \{S|_X : \exists h \in H_B \text{ such that } L_S(h) = 0\}$$



We claim that  $\{S|_X : L_{D,f}(h_S) > \varepsilon\} \subseteq M$ . The former set is the cases that we select bad hypothesis, and  $M$  is the cases there exist bad hypothesis. So it is a subset. We might not have selected a bad hypothesis from  $M$ , then we cannot put an equal between these two sets. We want to upper bound a probability of the set we defined. Therefore, it suffices to upper bound  $D^m(M)$ .

$$D^m(M) = D^m \left[ \bigcup_{h \in H_B} : \{S|_X : L_S(h) = 0\} \right]$$

Now we need two basic probability rules:

1. The union bound: For any two events  $A, B$  and any probability distribution  $P$ ,  $P(A \cup B) \leq P(A) + P(B)$ .
2. If  $A$  and  $B$  are independent events then  $P(A \cap B) = P(A) \cdot P(B)$ .

For any fixed  $h \in H_B$ . Let us upper bound  $D^m[\{S|_X : L_S(h) = 0\}]$ . For a single one, the probability of  $h$  is doing wrong on  $X$  is at least  $\varepsilon$ , then

$$\begin{aligned} D^m[\{S|_X : L_S(h) = 0\}] &= D^m[\{S_X : h(x_1) = f(x_1) \wedge \dots \wedge h(x_m) = f(x_m)\}] \\ &\leq (1 - \varepsilon)^m \end{aligned}$$

Then we conclude

$$D^m[\text{Bad } S] \leq D^m(M) = D^m[\bigcup_{h \in H_B} (L_S(h) = 0)] \leq |H_B| \cdot (1 - \varepsilon)^m \leq |H| \cdot (1 - \varepsilon)^m$$

Then  $\text{ERM}_H$  has small probability of failure as  $m \rightarrow \infty$ .  $\square$

Note that here we call it paradigm not algorithm since it doesn't tell you which  $H$  to pick.

This time we use a different notation:

$$\Pr_{S \sim D^m} [L_{D,f}(\text{ERM}_H(s)) > \varepsilon] \leq |H| \cdot (1 - \varepsilon)^m$$

for every  $\varepsilon \geq 0$  and for every  $m$ , where

$$L_{D,f}(h) = \Pr_{x \sim D} [h(x) \neq f(x)].$$

Trust that for every  $1 > \varepsilon > 0$ ,  $m$

$$(1 - \varepsilon)^m \leq e^{-\varepsilon m}$$

Thus the probability of making an error is going down exponentially fast in the sample size.

Also recall the proof idea:

- Step 1: For any given  $h : X \rightarrow \{0, 1\}$ ,  $\Pr_{S \sim D^m} [L_S(h) = 0]$  is small and getting smaller with  $m$ , provided that  $L_{D,f}(h) > \varepsilon$ . We are looking at samples that make  $h$  look good in spite of  $h$  being bad.
- Step 2: Take union over all  $h \in H$ .

# A Formal Learning Model

## 3.1 A formal notion of learnability

### PAC learnability

We say that a class of predictors  $H$  is **PAC learnable** (Probably Approximately Correct) if there exists a function  $m_H : (0, 1) \times (0, 1) \rightarrow \mathbb{N}$  such that there exists a learner

$$A : \bigcup_{m=1}^{\infty} (X \times \{0, 1\})^m \rightarrow \{f \mid f : X \rightarrow \{0, 1\}\}$$

that for every  $D$  probability distribution over  $X$  and every  $f \in H$  and every  $\varepsilon, \delta > 0$

$$\Pr_{S \sim D^m, f} [L_{D, f}(A(S)) > \varepsilon] < \delta$$

for every  $m \geq m_H(\varepsilon, \delta)$ .

Here we can think of  $\varepsilon$  as an accuracy parameter and  $\delta$  as confidence parameter.

If we rephrase Theorem 2.1, we get

### Theorem

Every finite  $H$  is PAC learnable with  $m_H(\varepsilon, \delta) \leq \frac{\ln |H| + \ln(1/\delta)}{\varepsilon}$ . Furthermore, any  $\text{ERM}_H$  learner will be successful.

Last time we have showed

$$\Pr_{S \sim D^m, f} [L_{D, f}(A(s)) > \varepsilon] \leq |H| \cdot e^{-\varepsilon m} \leq \delta$$

Then take  $\ln$ ,

$$\ln |H| - \varepsilon m \leq \ln(\delta)$$

Then we get results as desired.

**Strength** of the PAC definition is that we can guarantee the number of needed examples (for training) regardless of the data distribution  $D$  and of which  $f \in H$  is used for labeling. We call this is a “**Distribution free guarantee**”.

**Weakness:** It only works if the labeling rule  $f$  comes from  $H$ .

**Relaxation** The data is generated by some probability distribution  $D$  over  $X \times Y$ .

We still wish to output a labeling rule  $h : X \rightarrow Y$ .

Assume  $Y = \{0, 1\}$ , we claim the best predictor  $h$  should be

$$h^*(x) = \begin{cases} 1 & \text{if } D((x, 1)|x) \geq \frac{1}{2} \\ 0 & \text{otherwise} \end{cases}$$

which is called the Bayes rule. The problem is we *do not* know  $D$ . We only see a sample (generated by  $D$ ).

Now redefine successful learning to have only a relative error guarantee.

#### Agnostic PAC learnability

A class of predictors  $H$  is **agnostic PAC learnable** if there exist some function  $m_H(\varepsilon, \delta) : (0, 1) \times (0, 1) \rightarrow \mathbb{N}$ , and a learner  $A$  (taking samples, outputting predictors) such that for every  $D$  over  $X \times Y$  and every  $\varepsilon, \delta > 0$

$$\Pr_{s \sim D^m} [L_D(A(s)) > \min_{h \in H} (L_D(h)) + \varepsilon] < \delta$$

whenever  $m \geq m_H(\varepsilon, \delta)$ .

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