6.7900 Fall 2024: Lecture Notes 1

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There was a lot of logistical information which you can find at gradml.mit.edu.

1 What is machine learning?

Broadly, machine learning is a set of methods for *using data* to *become better at a task.*

Problem formalization

- Input space : X is often (but not necessarily) a vector space, often \mathbb{R}^d
- Output space : Y is a fixed finite discrete set, for *classification* problems; it could be a continuous set, such as R *d* in a *regression* problem
- Training set : $\{(x^{(n)}, y^{(n)})\}_{n=1}^N$: set of *N* pairs of values, where $x^{(n)} \in \mathcal{X}$ and $y^{(n)} \in \mathcal{Y}$

• Loss function : $L : \mathcal{Y} \times \mathcal{Y} \rightarrow \mathbb{R}$. This function is potentially *asymmetric*, so that $L(a, g)$ is the loss (badness) of predicting or guessing *g* when the actual label is *a*.

Example problem: spam detection Our *data* is a set of pairs, of an email message and a binary value describing whether that message is spam. Our *task* is to predict whether future email messages are or are not spam. We represent each email message as a vector of *features*, $x^{(n)}$, for example with each $x^{(n)}_i$ $i^{(n)}$ having value 1 if some word i occurs in the n th message and 0 otherwise. The associated *label* $y^{(n)} \in \{0,1\}$ indicates not-spam or spam.

What should our loss function be? Maybe something like:

$$
L(a, g) = \begin{cases} 0 & \text{if } a = g \\ 1 & \text{if } a = 1 \text{ and } g = 0 \\ 15 & \text{if } a = 0 \text{ and } g = 1 \end{cases}
$$

This encodes the idea that it's worse to say a message is spam when it is not (because we might miss an important message), than to say it is not spam when it is (because it is a mild annoyance).

Supervised learning problem: Given training set $\{(x^{(n)}, y^{(n)})\}_{n=1}^N$, find a decision rule $h: \mathcal{X} \rightarrow \mathcal{Y}$ that *performs well*.

To "perform well" is to make predictions on as-yet-unseen inputs that have low loss. That is, that $L(y^{(N+1)}, h(x^{(N+1)}))$ is low. But we don't know what these inputs will be! Probability to the rescue!
Assume training data and future data are drawn from some distribution $p(x, y)$ guage of uncertainty.

Assume training data and future data are drawn from some distribution $p(x, y)$. Now we can define *risk* of *h* to be the expected loss of using *h* to make a prediction on a new example:

$$
risk(h) = \mathbb{E}[L(Y, h(X))]
$$

Exercise: Show that, if $\{(X^{(N+m)}, Y^{(N+m)})\}_{m=1}^M$ are independent and identically distributed, the average expected loss of using *h* to make predictions on the next *M* points is equal to risk(*h*).

Optimal *h* for classification We'd love to pick *h* to minimize risk $\mathbb{E}[L(Y, h(X))]$. If we know $p(X, Y)$, then we can!

Proposition: Consider *K*-class classification, in which $\mathcal{Y} = \{1, \dots K\}$, and suppose *X* is a discrete random variable, *Y* is a random variable with domain \mathcal{Y} , and we know $p(x, y)$. Then the rule

$$
h(x) = \arg\min_{k} \sum_{j=1}^{K} L(j,k)p(y=j | x)
$$

minimizes $\mathbb{E}[L(Y, h(X))].$

Proof. We start by expanding and rearranging terms in the risk definition:

$$
\mathbb{E}[L(Y, h(X))] = \sum_{x, y \in \mathcal{X} \times \mathcal{Y}} L(y, h(x))p(x, y)
$$

$$
= \sum_{j=1}^{K} \sum_{x \in \mathcal{X}} L(j, h(x))p(x, y = j)
$$

$$
= \sum_{j=1}^{K} \sum_{x \in \mathcal{X}} L(j, h(x))p(y = j | x)p(x)
$$

$$
= \sum_{x \in \mathcal{X}} p(x) \sum_{j=1}^{K} L(j, h(x))p(y = j | x)
$$

Now observe that this sum over *x* can be minimized by independently minimizing the *j* for each *x*, and that $h(x) = \arg \min_k \sum_{j=1}^K L(j,k)p(y=j \mid x)$ is exactly the *h* that does so. \Box

Exercise: State and prove a similar result for continuous χ .

Two-class decision rule So, if we have two classes 0 and 1, and known $p(x, y)$, what is the form of the optimal h ? (Remember that the loss function is $L(a, g)$ where *a* is the true value of *Y* and *g* is the "guess" we are proposing to make.)

$$
h(x) = \begin{cases} 0 & \text{if } L(1,0)p(Y=1 \mid X=x) < L(0,1)p(Y=0 \mid X=x) \\ 1 & \text{otherwise} \end{cases}
$$

Exercise: What happened to $L(0,0)$ and $L(1,1)$? Where would they go in the expression above? Why was it okay to omit them?

Optimal *h* for regression Now let's consider a regression problem where $\mathcal{Y} = \mathbb{R}$. If we know $p(x, y)$ we can also determine the optimal (minimum risk) $h!$

Proposition: Assume $\mathcal{X} = \mathbb{R}^d$ and $\mathcal{Y} = \mathbb{R}$, and we know $p(x, y)$. If we are using *squared loss* $L(a, g) = (a - g)^2$, then the decision rule

$$
h(x) = \mathbb{E}[Y \mid X = x]
$$

minimizes $\mathbb{E}[L(Y, h(X))].$

Exercise: Complete the proof! It may be useful to make this (common, but counter-intuitive) move at some point:

$$
y - h(x) = y - \mathbb{E}[Y | X = x] + \mathbb{E}[Y | X = x] - h(x)
$$

Pretty cool! It says that the best prediction, under squared loss, is the conditional mean of *Y* given $X = x$.

Major obstacle We don't know $p(x, y)$!

But we do have training data! So, let's use it in the process of picking *h*. It can only be helpful if it is related, somehow, to the data that we are going to have to make predictions about in the future. A typical, strong assumption is that *all data*, including training data and future data, are independent and identically distributed.

Given this relationship between past and future data, we might think about approximating the *risk* by the *empirical risk* (average loss on the training data), observing that

$$
\mathbb{E}[L(Y, h(X))] \approx \frac{1}{N} \sum_{n=1}^{N} L(y^{(n)}, h(x^{(n)}))
$$

Exercise: Will this be the key to solving all our problems in machine learning?