# Machine Learning

Learning Model

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## A Formal Model (Statistical Learning)

We have a learner (us, or the machine) has access to:

- **1** Domain set  $\mathcal{X}$ : set of all possible objects to make predictions about
  - domain point  $x \in \mathcal{X} = instance$ , usually represented by a vector of *features*
  - $\chi$  is the *instance space*
- **2** Label set  $\mathcal{Y}$ : set of possible labels.
  - often two labels, e.g  $\{-1, +1\}$  or  $\{0, 1\}$
- **3 Training data**  $S = ((x_1, y_1), \dots, (x_m, y_m))$ : finite sequence of labeled domain points, i.e. pairs in  $\mathcal{X} \times \mathcal{Y}$ 
  - this is the learner's input
  - S: training example or training set

- **4 Learner's output** h: prediction rule  $h: \mathcal{X} \to \mathcal{Y}$ 
  - also called predictor, hypothesis, or classifier
  - *A(S)*: prediction rule produced by learning algorithm *A* when training set *S* is given to it
  - sometimes  $\hat{f}$  used instead of h
- **5 Data-generation model**: instances are generated by some probability distribution and labeled according to a function
  - D: probability distribution over X (NOT KNOWN TO THE LEARNER!)
  - labeling function  $f: \mathcal{X} \to \mathcal{Y}$  (NOT KNOWN TO THE LEARNER!)
  - label  $y_i$  of instance  $x_i$ :  $y_i = f(x_i)$ , for all i = 1, ..., m
  - each point in training set S: first sample  $x_i$  according to  $\mathcal{D}$ , then label it as  $y_i = f(x_i)$
- **6** Measures of success: error of a classifier = probability it does not predict the correct label on a random data point generate by distribution  $\mathcal{D}$

### Loss

Given domain subset  $A \subset \mathcal{X}$ ,  $\mathcal{D}(A) =$  probability of observing a point  $x \in A$ .

Let A be defined by a function  $\pi: \mathcal{X} \to \{0,1\}$ :

$$A = \{x \in \mathcal{X} : \pi(x) = 1\}$$

In this case we have  $\mathbb{P}_{x \sim \mathcal{D}}[\pi(x)] = \mathcal{D}(A)$ 

**Error of prediction rule**  $h: \mathcal{X} \to \mathcal{Y}$  is

$$L_{\mathcal{D},f}(h) \stackrel{\text{def}}{=} \mathbb{P}_{x \sim \mathcal{D}}[h(x) \neq f(x)] \stackrel{\text{def}}{=} \mathcal{D}(\{x : h(x) \neq f(x)\})$$

#### Notes:

- $L_{\mathcal{D},f}(h)$  has many different names: **generalization error**, *true* error, risk, **loss**, ...
- often f is obvious, so omitted:  $L_{\mathcal{D}}(h)$

Learner outputs  $h_{S}: \mathcal{X} \to \mathcal{Y}$ .

Lateral training set SGoal of the learner?

Goal: find  $h_S$  which minimizes the generalization error  $L_{\mathcal{D},f}(h)$ 

or LD,f(h)

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Training error:  $L_S(h) \stackrel{\text{def}}{=} \underbrace{\{i:h(x_i) \neq y_i, 1 \leq i \leq m\}}_{m}$  for which h predictions in  $S = \{(x_1, y_1), \dots, (x_m, y_m)\}$ 

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**Note**: the *training error* is also called *empirical error* or *empirical risk* 

## **Empirical Risk Minimization**

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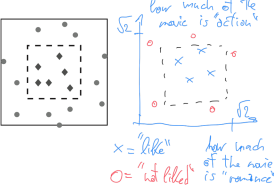
Empirical Risk Minimization (ERM): produce in output h minimizing  $L_S(h)$ 

### What can go wrong with ERM?

Consider our simplified movie ratings prediction problem. Assume

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Assume  $\mathcal{D}$  and f are such that:

- instance x is taken uniformly at random in the square  $(\mathcal{D})$
- label is 1 if x inside the inner square, 0 otherwise (f)
- area inner square = 1, area larger square = 2

Consider classifier given by
$$h_{S}(x) = \begin{cases} y_{i} & \text{if } \exists i \in \{1, \dots, m\} : x_{i} = x \\ 0 & \text{otherwise} \end{cases}$$

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Is it a good predictor?

$$L_S(h_S) = 0$$
 but  $L_{D,f}(h_S) = 1/2$ 

Good results on training data but poor generalization error ⇒ **overfitting** 

When does ERM lead to good performances in terms of generalization error?