Validation and Testing

COMPSCI 371D — Machine Learning

Outline

- 1 Training, Testing, and Model Selection
- 2 A Generative Data Model
- Model Selection: Validation
- 4 Model Selection: Cross-Validation
- 5 Model Selection: The Bootstrap

Training and Testing

Empirical risk is average loss over training set:

$$L_T(h) \stackrel{\text{def}}{=} \frac{1}{|T|} \sum_{(\mathbf{x}, y) \in T} \ell(y, h(\mathbf{x}))$$

Training is Empirical Risk Minimization:

$$\mathsf{ERM}_T(\mathcal{H}) \in \operatorname{arg\,min}_{h \in \mathcal{H}} L_T(h)$$
 (A fitting problem)

- Not enough for machine learning: Must generalize
- Small risk on "previously unseen data"
- How do we know? Evaluate on a separate test set S
- This is called testing the predictor
- How do we know that S and T are "related?"

Model Selection

- Hyper-parameters: Degree k for polynomials, number k of neighbors in k-NN
- How to choose? Why not just include with parameters, and train?
- Difficulty 0: k-NN has no training! No big deal
- Difficulty 1: $k \in \mathbb{N}$, while $\mathbf{v} \in \mathbb{R}^m$ for some predictors. Hybrid optimization. Medium deal, just technical difficulty
- Difficulty 2: Answer from training would be trivial!
- Can always achieve zero risk on T
- So k must be chosen separately from training. It tunes generalization
- This is what makes it a *hyper-parameter*
- Choosing hyper-parameters is called model selection
- Evaluate choices on a separate validation set V

Model Selection, Training, Testing

- Warning: We use "model" with two different meanings in the same slide deck! [Sorry, that's the literature]
- "Model" in "model selection" is H
- Given a (hyper-)parametric family of hypothesis spaces, model selection selects one particular member of the family
- Given a specific hypothesis space (hyper-parameter), training selects one particular predictor out of it
- Use V to select model, T to train, S to test
- Train on cats and test on horses?
- V, T, S are mutually disjoint but "related"
- What does "related" mean?



A Generative Data Model

- What does "related" mean?
- Every sample (\mathbf{x}, y) comes from a joint probability distribution $p(\mathbf{x}, y)$, the "data model" ("model" number 2)
- True for training, validation, and test data, and for data seen during deployment
- For the latter, y is "out there" but unknown
- The goal of machine learning:
 - Define the (statistical) risk $L_p(h) = \mathbb{E}_p[\ell(y, h(\mathbf{x}))] = \iint \ell(y, h(\mathbf{x})) p(\mathbf{x}, y) d\mathbf{x} dy$
 - Learning performs (Statistical) Risk Minimization: $RM_p(\mathcal{H}) \in arg \min_{h \in \mathcal{H}} L_p(h)$
- Lowest risk on \mathcal{H} : $L_p(\mathcal{H}) \stackrel{\text{def}}{=} \min_{h \in \mathcal{H}} L_p(h)$



p is Unknown

- $\mathsf{RM}_p(\mathcal{H}) \in \operatorname{arg\,min}_{h \in \mathcal{H}} \iint \ell(y, h(\mathbf{x})) p(\mathbf{x}, y) d\mathbf{x} dy$
- So, we don't need training data anymore?
- We typically do not know p(x, y)
- **x** = image? Or sentence?
- Can we not estimate p?
- The curse of dimensionality, again
- We typically cannot find $RM_{\rho}(\mathcal{H})$ or $L_{\rho}(\mathcal{H})$
- That's the goal all the same

So Why Talk About It?

- Why talk about $p(\mathbf{x}, y)$ if we cannot know it?
- $L_p(h)$ is a mean, and we can *estimate* means
- We can sandwich L_p(h) or L_p(H) between bounds over all possible choices of p
- What else would we do anyway?
- p is conceptually clean and simple
- The unattainable holy grail
- Think of p as an oracle that sells samples from X × Y
- She knows p, we don't
- Samples cost money and effort! [Example: MNIST Database]



Even More Importantly...

- We know what "related" means:
 T, V, S are all drawn independently from p(x, y)
- We know what "generalize" means: Find $RM_p(\mathcal{H}) \in \arg\min_{h \in \mathcal{H}} L_p(h)$
- We know the goal of machine learning

Validation

- Parametric family of hypothesis spaces $\mathcal{H} = \bigcup_{\pi \in \Pi} \mathcal{H}_{\pi}$
- Finding a good vector $\hat{\pi}$ of hyper-parameters is called model selection
- A popular method is called validation
- Use a validation set V separate from T
- Pick a hyper-parameter vector for which the predictor trained on the training set minimizes the validation risk

$$\hat{\pi} = \arg\min_{\pi \in \Pi} L_V(\mathsf{ERM}_T(\mathcal{H}_\pi))$$

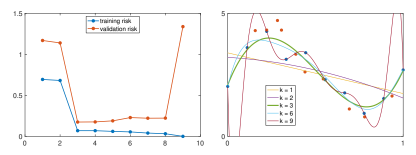
• When the set Π of hyper-parameters is finite, try them all

Validation Algorithm

```
procedure Validation(\mathcal{H}, \Pi, T, V, \ell)
       \hat{I} = \infty
                                                                                              Stores the best risk so far on V
      for \pi \in \Pi do
             h \in \operatorname{arg\,min}_{h' \in \mathcal{H}_{\pi}} L_{\mathcal{T}}(h')
                                                                  \triangleright Use loss \ell to compute best predictor ERM_{\tau}(\mathcal{H}_{\pi}) on T
             L = L_V(h)
                                                                           \triangleright Use loss \ell to evaluate the predictor's risk on V
              if l < \hat{l} then
                    (\hat{\pi},\hat{h},\hat{L})=(\pi,h,L) 
hinkspace Keep track of the best hyper-parameters, predictor, and risk
              end if
       end for
      return (\hat{\pi}, \hat{h}, \hat{L})
                                                                   Return best hyper-parameters, predictor, and risk estimate
end procedure
```

Validation for Infinite Sets

- When Π is not finite, scan and find a local minimum
- Example: Polynomial degree



 When Π is not countable, scan a grid and find a local minimum

Resampling Methods for Validation

- Validation is good but expensive: needs separate data
- A pity not to use V as part of T!
- Resampling methods split T into T_k and V_k for k = 1, ..., K
- (Nothing to do with number of classes or polynomial degree!)
- For each π , for each round k, train on T_k , test on V_k to measure performance
- Average performance over k taken as validation risk for π
- Let $\hat{\pi}$ be the best π
- When done, train the predictor in $\mathcal{H}_{\hat{\pi}}$ and on all of T
- Cross-validation and the bootstrap differ on how splits are made

K-Fold Cross-Validation

- V_1, \ldots, V_K are a partition of T into approximately equal-sized sets
- $T_k = T \setminus V_k$
- For $\pi \in \Pi$

For
$$k = 1, ..., K$$
:
train on T_k , measure performance on V_k
Average performance over k is validation risk for π

- Pick $\hat{\pi}$ as the π with best average performance
- When done, train the predictor in $\mathcal{H}_{\hat{\pi}}$ and on all of T
- Since performance is an average, we also get a variance!
- We don't have that for standard validation

How big should *K* be?

- T_k has |T|(K-1)/K samples, so the predictor in each fold is a bit worse than the final predictor
- Smaller K: More pessimistic risk estimate (upward bias b/c we train on smaller T_k)
- Bigger K decreases bias of risk estimate
- (training on bigger T_k)
- Why not *K* = *N*?
- LOOCV (Leave-One-Out Cross-Validation)
- Train on all but one data point, test on that data point, repeat
- Any issue?
- Nadeau and Bengio recommend K = 15

The Bootstrap

- Bag or multiset: A set that allows for multiple instances
- {*a*, *a*, *b*, *b*, *b*, *c*} has cardinality 6
- Multiplicities: 2 for a, 3 for b, and 1 for c
- A set is also a bag: {*a*, *b*, *c*}
- Bootstrap: Same as CV, except
 - T_k: N samples drawn uniformly at random from T, with replacement
 - $V_k = T \setminus T_k$
- T_k is a bag, V_k is a set

How Many Elements are Missing from T_k ?

- Fix attention on one sample s $\mathbb{P}[s \text{ is drawn in one draw}] = 1/N$ $\mathbb{P}[s \text{ is not drawn in one draw}] = 1 1/N$ $\mathbb{P}[s \text{ is not drawn ever}] = (1 1/N)^N$
- Average fraction of missing elements $(1 1/N)^N$
- For large *N*, this is about $\lim_{N\to\infty} \left(1-\frac{1}{N}\right)^N = \frac{1}{e} \approx 0.37$
- Good approximation: $(1 1/24)^{24} \approx 0.36$
- 37 % of elements are missing from T_k on average
- 63 % of elements end up in T_k on average

Cross-Validation vs Bootstrap

- Bootstrap estimates are good
- Sometimes somewhat more biased than CV
- CV is method of choice for model selection.
- Bootstrap leads to random decision forests