Lecture 9: Generalization Gap & Bias-variance decomposition

- We introduced Enew error on fresh unseen data
- Designing a method with small Enew is the central goal in supervised learning
- Cross-validation helps in estimating Enew (Unless one uses it for tuning hyperparameters)
- Enew can be further analyzed even more mathematically

$$E(\gamma, \hat{\gamma})$$

Recall that
$$E_{\text{train}} = \frac{1}{N} \sum_{i=1}^{N} E(y_i, \hat{y}(\underline{x}_i; T))$$

$$E_{\text{hew}} = \int E(y_*, \hat{y}(\underline{x}_*; T)) P(\underline{x}_*, y_*) d\underline{x}_* dy_*$$

$$multi-dimensional integral$$

- Note the values of E_{train} and E_{new} were calculated while keeping the training set T fixed

- We now emphasize the fact that Etrain and Enew are functions of the training set
 - · As the training set T changes, the values of Etrain and Enew changes too!

Etrain
$$(T) = \frac{1}{N} \sum_{i=1}^{N} E(y_i, \hat{y}(z_i; T))$$

of training data
$$E_{\text{new}}(T) = E(y_*, \hat{y}(\underline{x}_*, T)) P(\underline{x}_*, y_*) d\underline{x}_* dy_*$$

- We now emphasize the fact that Emain and Enew are functions of the braining set
 - · As the training set T changes, the values of Etrain and Enew changes too!

Function

Function

of training

$$E_{new}(T) = \frac{1}{N} \sum_{i=1}^{N} E(y^{(i)}, \hat{y}(z^{(i)}; T))$$
 $E_{new}(T) = E(y^*, \hat{y}(z^*; T)) P(z^*, y^*) dz^* dy^*$

- So we introduce another level of abstraction
 - · Training-data averaged versions of Enew and Ebain

$$E_{\text{new}} = E_{\tau} [E_{\text{new}}(\tau)]$$

$$\overline{E}_{\text{train}} = \underline{E}_{\tau} \left[E_{\text{train}}(\tau) \right]$$

- \mathbb{E}_{τ} denotes the expected value w.r.t. the training set $\mathcal{T} = \{\underline{x}^{(i)}, y^{(i)}\}_{i=1}^{N}$
- · We assume T consists of independent samples from p(x, y)

Enew is the average Enew if we were to train the model multiple times on on different training datasets of size N. (Same goes for Etrain)

$$E_{\text{new}} = E_{\mathcal{T}} [E_{\text{new}}(\mathcal{T})]$$

$$E_{\text{train}} = E_{\mathcal{T}} [E_{\text{train}}(\mathcal{T})]$$

- E_{τ} denotes the expected value w.r.t. the training set $T = \{\chi^{(i)}, \chi^{(i)}\}$
- We assume T consists of independent samples from p(x, y)

Enew is the average Enew if we were to train the model multiple times on on different training datasets of size N.

- Why introduce these average quantities Enew and Etrain?
 - It is easier to reason about the average behavior Enew and Errain than about the errors Enew and Etrain obtained when the model is trained on one specific training dataset T
 - Insights from \overline{E}_{new} are useful, even though we most often care about E_{new} in the end (as the training data is usually fixed)

- We have already seen that Enew & Etrain
 - Usually, $E_{train} < E_{new}$ and $E_{train} < E_{new}$

On an average, a method usually performs worse on new, unseen data than on training data

- A method generalizes well if it performs well on unseen data after training
- We call the difference between Enew and Etrain as the generalization gap

Generalization gap = \overline{E}_{new} - \overline{E}_{train} as the expected generalization gap)

is the difference between the expected performance on new unseen data and the expected performance on training data

- So Enew can be decomposed as:

Enew = Etrain + generalization gap

What affects the Generalization Gap?

- One can in some sense say, the more a method adapts to training data, the larger the generalization gap
- Vapnik-Chervonenkis (VC) dimension is a theoretical framework which assesses how much a method adapts to training data
 - · The framework provides probabilistic bounds on generalization gap
 - · However, the bounds are quite conservative
 - . Therefore, the method is not very useful in practice
- Instead, we will only use vague terms "Model Complexity" (or Model flexibility)

 ability of a method
 to adapt to patterns
 in training data
 - · High complexity models e.g. very deep decision trees, KNN with small 'K'
 - · Low complexity models e.g. linear regression, logistic regression

- Instead, we will only use vague terms "Model Complexity" (or Model flexibility)

 ability of a method

 to adapt to patterns
 in training data
 - · High complexity models e.g. very deep decision trees, KNN with small 'k'
 - Low complexity models e.g. linear regression, logistic regression basic parametric methods
 - · For parametric models, the model complexity is related to the number of parameters, but is also affected by regularization techniques

Model complexity 1

Etrain

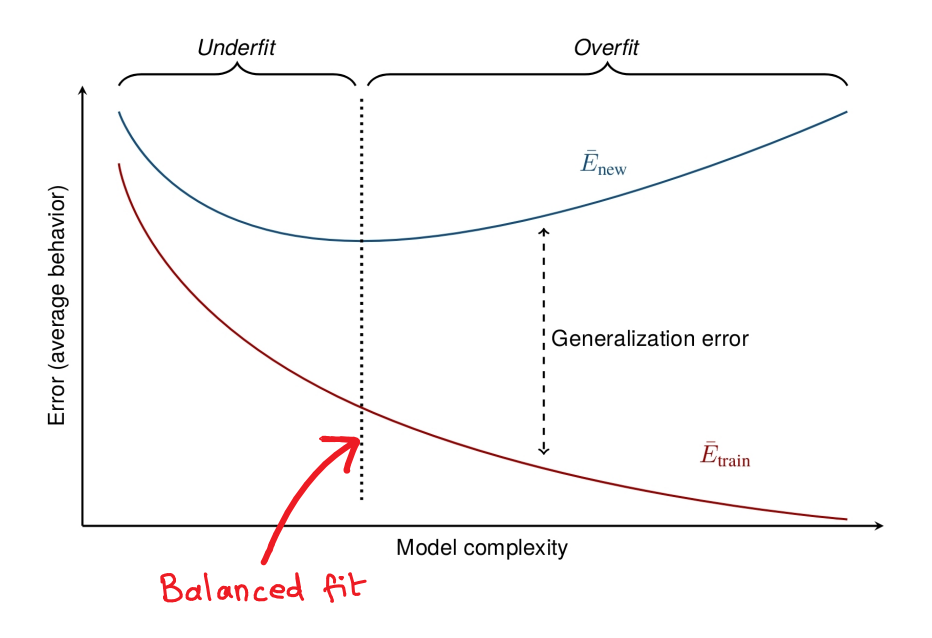
Generalization gap 1

Model complexity V

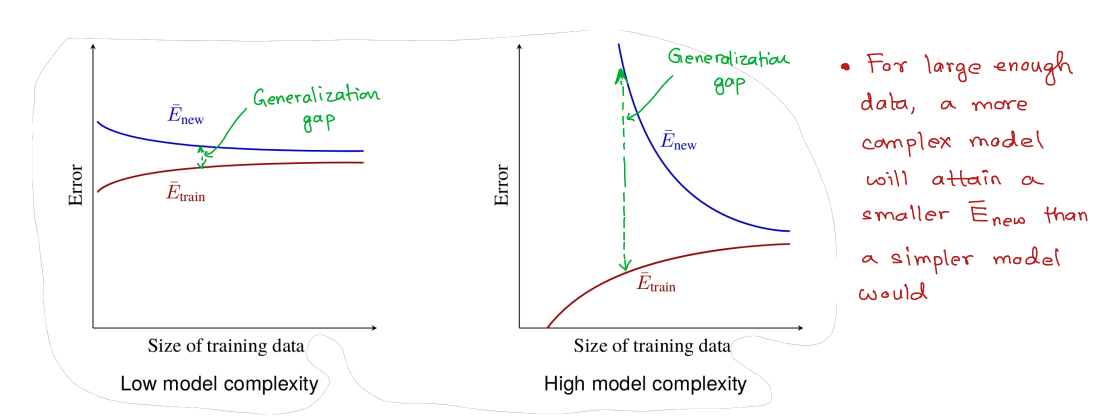
Etrain

Generalization gap 1

Enew usually attains a minimum at some intermediate complexity

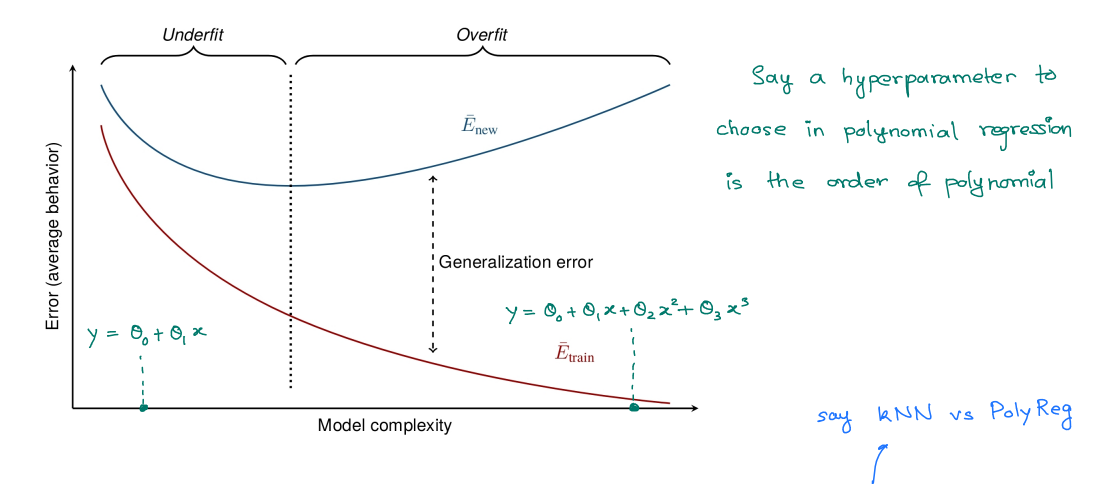


- Another important aspect is the size of training dataset N
- In general, the more the training data, the smaller the generalization gap
- Etrain typically increases as N increases
 - since most models are unable to fit all training data points well, more so if there are too many of them



Short comings of using Model Complexity

The figure below is relevant only when there is a single hyperparameter to choose



- However, when there are multiple hyperparameters (or multiple methods) to choose from, the above one-dimensional model complexity will not work well

- Lets take an example of jointly choosing the degree of polynomial regression and the regularization parameter
 - · higher degree of polynomial => more flexibility / complexity
 - · more regularization ⇒ less flexibility / complexity
- Example of a simulated problem: Sample data points from p(2,y)

Data

Generation

• N = 10 data points

- Input x ~ Uniform Dist (-5,10)
- $y = min(0.1x^2, 3) + \epsilon$
 - € ~ Normal Dist(0,1)

$$p(x,y) = p(y|x) p(x)$$

Then

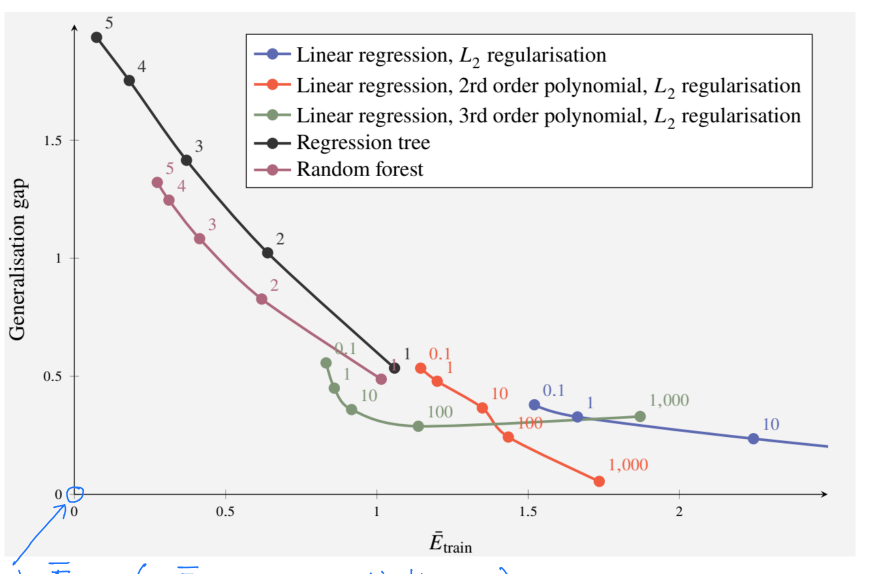
sample
First

y given x sample x

Now fit the input-output data { z(i), y(i)?10 using

- (a) Linear regression with Lz-regularization
- (b) Linear regression with a quadratic polynomial and Lz-regularization
- (c) Linear regression with a cubic polynomial and Lz-regularization
- (d) Regression Tree, (e) A random forest with 10 regression trees

- For each of the methods, we try different values of hyperparameters (regularization parameter λ and tree depth) and compute \overline{E}_{train} and generalization gap



In practice, we only have limited data and we cannot generate these plots!

Lowest Enew (= Errain + generalization gap)