Regularization and Model Selection

Prof. Alessandro Lucantonio

Aarhus University

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Underfitting and overfitting - intuition

Imagine that you have to study for a written exam consisting of exercises.

Doing only a few exercises while studying leads to poor perfomance both on homework and at the exam. This is called *underfitting*: the performance at the exam will be bad because of poor training.

Moreover, memorize all the solutions to homework leads to the maximum score on homework (trivially) but probably a bad score on the exam exercises. This is called *overfitting*: you have a bad performance on the exam because you did not capture the true essence of the homework, rather you have memorized their peculiarities.

Underfitting and overfitting - analytical example

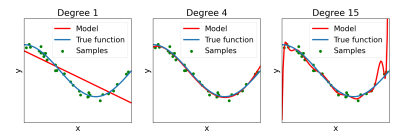


Figure: Underfitting (polynomial degree 1), good fitting (polynomial degree 4), overfitting (polynomial degree 15).

Overfitting can be countered through:

- validation;
- regularization.

Regularization

Overfitting is correlated with "complex" models: to avoid them, we penalize models with large weights. One way to do this is to consider the following cost function

$$E_r(\mathbf{w}) = E(\mathbf{w}) + R_{\lambda}(\mathbf{w})$$

where R_{λ} is the *regularization term*.

 $\lambda > 0$ is a hyperparameter that must be chosen in the **model** selection phase (see later).

Most common regularization techniques:

- ► Tikhonov (or L^2 or Ridge): $R_{\lambda}(\mathbf{w}) = \lambda ||\mathbf{w}||_2^2 = \lambda \sum_i w_i^2$. Tends to make *all* weights small.
- ▶ LASSO (or L^1): $R_{\lambda}(\mathbf{w}) = \lambda ||\mathbf{w}||_1 = \lambda \sum_i |w_i|$. Tends to make some weights 0 (feature selection).

LR with Tikhonov regularization

► Cost function (without regularization):

$$E(\mathbf{w}) = \frac{1}{N} ||\mathbf{X}\mathbf{w} - \mathbf{y}||^2$$

Regularized cost function:

$$E_r(\mathbf{w}) = E(\mathbf{w}) + \lambda ||\mathbf{w}||^2$$

Gradient of the regularized cost function:

$$\nabla E_r(\mathbf{w}) = \nabla E(\mathbf{w}) + 2\lambda \mathbf{w} = 2\left(\frac{1}{N}X^T(X\mathbf{w} - \mathbf{y}) + \lambda \mathbf{w}\right)$$

► Normal equation:

$$\mathbf{w} = (\mathsf{X}^\mathsf{T} \mathsf{X} + \lambda \mathsf{I})^{-1} \mathsf{X}^\mathsf{T} \mathbf{y}$$

Note that in this case $X^TX + \lambda I$ is *always* invertible (why?).

Bias-Variance trade-off

Bias: expected discrepancy between targets and predictions given by the model.

Variance: measure of the deviation from the expected value given by the model that any particular sampling of the data (from the same underlying data-generating distribution) is likely to cause.

Some intuitions on the effect of regularization:

- ▶ High $\lambda \rightsquigarrow$ simple model \rightsquigarrow high bias (underfitting).
- ▶ Low $\lambda \leadsto$ complex model \leadsto high variance (overfitting).
- Intermediate λ : optimal solution (good bias-variance trade-off).

Generalization

Central challenge in Machine Learning: **generalization**! Our algorithm must perform well on *new, previously unseen* inputs.

Recall that we have to find a balance between bias and variance. Even if we limit the complexity of our model using regularization, the training set does not provide a good estimate of the test (generalization) error.

In other words, generalization is compromised if we choose hyperparameters (including the regularization factor) only according to the training error.

Generalization - Training vs test error

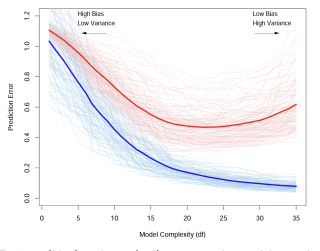


Figure: Training (blue) and test (red) errors as the model complexity varies.

Model selection and model assessment

Do **not** use the test set to tune hyperparameters: introduce the **validation set**.

In general, we should distinguish between:

- 1. Model selection: estimate the performance of different models trained with different hyperparameters.
- 2. Model assessment: after choosing a final model we evaluate its performance on *new*, *previously unseen* (test) data.

Once we have selected a model using the validation set, we can assess its generalization error on the test set. Splitting the dataset into training and validation sets is called *hold-out*.

Double Hold-out



- ▶ We split the entire dataset in three sets: training, validation and test (usually 60%-20%-20% or 70%-20%-10% of the total dataset).
- Training set is used to fit the model. Then, we evaluate its validation error and adjust its hyperparameters to reduce it. We re-fit the model and keep adjusting the hyperparameters until we find a model with a good trade-off between training and validation errors.
- ▶ We assess the generalization capability of the best model by computing the error on the test set.

Cross-Validation

	Fold 2	Fold 3	Fold 4	Fold 5
Fold 1	Fold 2	Fold 3	Fold 4	Fold 5
Fold 1	Fold 2	Fold 3	Fold 4	Fold 5
Fold 1	Fold 2	Fold 3	Fold 4	Fold 5
Fold 1	Fold 2	Fold 3	Fold 4	Fold 5

Figure: 5-fold CV

- ▶ Split the data in *k* disjoint *folds*.
- Use k-1 folds as the training set and the remaining fold as the validation set. Repeat k times (see figure).
- ► The performance will be the mean ± standard deviation computed across the k runs.

Pro: Not sensitive to a particular partition of the data. Mean filter the error.

Cons: Computationally expensive (but parallellizable).

CV for selection and hold-out for assessment

- 1. Split dataset into two sets (hold-out): development D (model selection) and test T (model assessment).
- Use the cross validation method on D to select the best model. Optional: retrain the best model using the entire dataset D.
- 3. Evaluate the generalization error on the test set T.

Grid search

How to choose the best set of hyperparameters?

Example: two hyperparameters (learning rate α and regularization parameter λ). Consider sets of three values for each parameter, e.g. $\alpha_{\rm vals} = \{0.001, 0.01, 0.1\}, \lambda_{\rm vals} = \{0.0001, 0.001, 0.01\}.$ Evaluate the model with $(\alpha, \lambda) = (i, j)$ for $i \in \alpha_{\rm vals}, j \in \lambda_{\rm vals}$.

λ

	(0.001,0.0001)	(0.001, 0.001)	(0.001, 0.01)
α	(0.01, 0.0001)	(0.01, 0.001)	(0.01, 0.01)
	(0.1, 0.0001)	(0.1, 0.001)	(0.1, 0.01)

Choose the pair of value corresponding to the *best performance on* the validation set.

Refinement: Suppose that the best is (0.1,0.001). Then we can "zoom" near the best set doing another grid search with e.g. $\alpha_{\text{vals}} = \{0.075, 0.1, 0.125\}, \lambda_{\text{vals}} = \{0.00075, 0.001, 0.00125\}.$