Advanced Machine Learning

Over-Fitting





Overfitting, regularisation, feature selection

4. Regularisation

1. Over-fitting?

2. Controlling Complexity 3. Hidden structure



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Over-fitting

- Complex machine can over-fit over-fitting: fitting the training data well at the cost of $getting\ poorer\ generalisation\ performance \blacksquare$
- Three red cars. . . I
- If we used an infinitely flexible machine we can fit our training data perfectly, but would have no generalisation ability.

Binary Classification Task for You

Outline



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Which Category?

• Which category does the following image belong to?



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Spurious Rules

- You ask a learning machine to solve a task based on datal
- It will find a rule that does this, but not necessary the rule you had in mind─machine learning isn't magic, it can't read your mind
- Infinitely flexible machines have an infinity of spurious rules they can learn

 —they are useless

 ■
- What should we do?

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All Binary Functions

 $x_1 = 100 \quad y_1 = \begin{cases} 0 \\ 1 \end{cases}$ $\mathcal{D} = \{(000, 0), (010, 1), (110, 1), (001, 0), (101, 0)\}$ $x_3 = 110 \ y_3 = \begin{cases} 1 \\ 1 \\ 1 \end{cases}$ $x_4 = 001 \quad y_4 = \begin{cases} 0 \\ 1 \end{cases}$ $x_6 = 011 \quad y_6 = \begin{cases} 0 \\ 1 \end{cases}$

Are MLPs Universal Approximators?

- Yesland Nol
- Yes: If you give me any function, I can find an MLP that approximates that function to any desired accuracy
- No: If you give me an MLP, I can find a function with an arbitrary high approximation error
- Would an MLP that could approximate any function be useful?
- Absolutely not!

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Training Examples

- As we increase the number of training examples, we make it hard to find a spurious rule
- Bigger data sets allow us to use more complicated machines
- Part of the success of deep learning is because they use huge training sets—but this is only a part of their success
- (Labelled) data is often expensive to collect so we sometimes have no choice but to use a small training set
- One of the limitations of using deep learning comes because we often have limited data

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Preprocessing

- Structure might often be obscure to the learning machine
- If we are trying to predict the spread of disease then a list of place names might be a lot less useful than their coordinates
- Imposing an ordering on an unordered set might not be useful

 $\big\{\,\text{``blue''}:0,\,\,\text{``brown''}:1,\,\,\text{``green''}:2,\,\,\text{``black''}:3\big\}\blacksquare$

• Choosing an encoding that reflect meaningful structure is essential to successful machine learning

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Controlling Complexity

- Infinitely flexible machine don't generalise (because any unseen data could have any value)
- Machine learning only works because there is some structure in the data
- A successful machine should capture this structure
- Even deep learning machines with millions of parameters only work because they successfully capture the structure of images or
- Different learning machines have different performance on different problems because the data has different structure

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Identifying Structure

- ullet In some cases we know $a\ priori$ some of the structure in the datall
- In images we believe the identity of an object is invariant to translation and scaling
- The success of convolutional neural networks (CNNs) in deep learning is in large part because the convolutions respect translational invariance

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Automatic Preprocessing

- One view of deep learning is that each layer (particularly in CNNs) acts as a preprocessor
- That is, it finds filters that captures features salient to the problem being tackled
- For both images and texts we expect salient features to be spatially localised (CNN finds localised filter)
- The deep structure allows ever more complicated features to be captured—that is, we can find spatially localised features on different scales
- Having very large datasets and simple filters (the number of weights in the CNN layers tends to be small) stops overfitting

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Hidden Structure

- Often the structure of data is invisible to us
- A very successful strategy is to try many different machine learning techniques and choose the best (stupid but effective)
- Often learning machines have adjustable parameters (hyper-parameters) that we have to set (they are the same for all input data, but change with the problem)
- We need to choose the hyper-parameters to fit the data in our problem
- Fine tuning hyper-parameter is important and almost always required (true in SVMs, MLP, deep learning)

Measuring Generalisation Performance

- Recall, we want to predict unseen data
- You cannot use data that you have trained on! you will overfit
- Need to split your data up into training and validation set
- Use the validation set to choose the hyper-parameters
- You need a separate testing set if you want to measure your generalisation performance

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Cross Validation

- If you want to use more data for training then you can use cross
- ullet K-fold cross validation splits the data into K groups

$$\mathcal{D} = \{D_i\}_{i=1}^P \quad D_i = (\mathbf{x}_i, y_i)$$

$$\frac{77}{10} + \frac{5.5.8 \cdot 5.5 \cdot 1.8.2 \cdot 1.8.2 \cdot 1.8.8 \cdot 3.7 \cdot 3.6}{10} + \frac{17.4 \cdot 4.6 \cdot 0.99 + 4.5 \cdot 4.6 \cdot 5.4}{10} + 4.6.2 \cdot 3.3 \cdot 2.7}{10} = 4.3$$

Leave-one-out cross-validation is extreme case

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Dimensionality Reduction

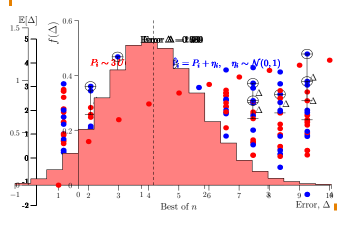
- We can sometimes simplify our machines by using less features
- We can project our data onto a lower dimensional sub-space (e.g. one with the maximum variation in the data: PCA)
- We can use clustering to find exemplars and recode our data in terms of distances from the exemplars (radial basis functions)
- Whether this helps depends on whether the information we discard is pertinent to the task we are trying to perform!

Normalising Features

- Measuring a feature in millimeters or kilometers is going to make a lot of difference to the size of that feature
- Many learning algorithms are sensitive to the size of a feature (larger features are more important)
- If we don't know how important different features are then it makes sense to normalise our features. E.g.

$$x_i^\alpha \leftarrow \frac{x_i^\alpha - \hat{\mu}_i}{\hat{\sigma}_i}, \quad \hat{\mu} = \frac{1}{m} \sum_{\beta=1}^m x_i^\beta, \quad \hat{\sigma}_i^2 = \frac{1}{m-1} \sum_{\beta=1}^m (x_i^\beta - \hat{\mu}_i)^2 \mathbf{I}$$

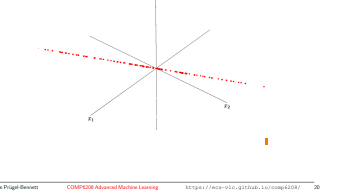
The Overfitting Game



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Hidden Structure

Can't spot low dimensional data by looking at numbers



Feature Selection

- Spurious features will allow us to find spurious rules (over-fitting)
- We can try different combinations of features to find the best set, although it rapidly becomes intractable to do this in all ways
- We can use various heuristics to decide which features to keep, but no heuristic is fail-safe method to find the best set of features
- Feature selection however can be powerful, often we can get very good results by keeping only a few variables
- As well as possibly improving generalisation we also get a more interpretable rule

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Explicit Regularisation

 As you've seen in the foundations of ML course, we can modify our error function to choose smoother functions

$$L = \sum_{k=1}^{m} \left(\boldsymbol{w}^{\mathsf{T}} \boldsymbol{x}_{k} - y_{k} \right)^{2} + \nu \| \boldsymbol{w} \|^{2}$$

(Good to normalise features)

- Second term is minimised when $w_i = 0$
- If w_i is large then

$$f(\boldsymbol{x}|\boldsymbol{w}) = \boldsymbol{w}^{\mathsf{T}} \boldsymbol{x} = \sum_{i=1}^{p} w_i x_i$$

varies rapidly as we change x_i

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Implicit Regularisation

- In the last two examples we added an explicit regularisation term that made the function we learnt simpler
- Some learning machines do this less explicitly
- Some deep learning architectures do subtle averaging
- Sometimes the architecture biases the machine to find a simple solution

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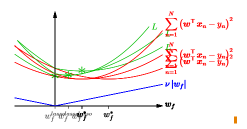
Success of SVMs

- SVMs regularise themselves by choosing the machine with the largest margin
- This ensures maximum stability to noise on the datal
- It leads to very good generalisation on small datasets—usually beats everything else
- But you still need to normalise the features
- ullet You also need to tune its hyper-parameters (C and sometimes $\gamma)$

Lasso

• We can use other regularisers

• Spurious features (e.g. colour of flag on energy consumption) will give us a small improvement in training error



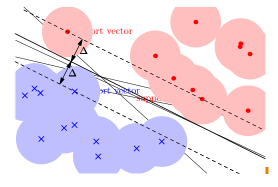
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Maximum Margin Machines

Perceptrons have many options to separate data



• SVMs choose the machine with the biggest margins

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Lessons

- Machine learning isn't magic
- It works when the learning machine is well attuned to the problem
- Sometimes you can help by preprocessing your data
- Sometimes there is a regularisation term that helps select a simpler machine
- Most machines have hyper-parameter that you tune to match the machine to the datal
- Really clever machines try to do this matching automatically

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