

ADRIAN BEVAN

INTRODUCTION TO MACHINE LEARNING PART 1

LECTURES GIVEN IN THE PARTICLE PHYSICS DIVISION AT THE RUTHERFORD APPLETON LABORATORY, MAY/JUNE 2020

OVERVIEW

- What is machine learning?
- Ethics
- Data wrangling
- Optimisation
- Decision Trees
- Neural Networks
 - Multilayer Perceptron (MLP)
 - Auto-encoders
 - Convolutional Neural Networks (CNN)
 - Generative Adversarial Networks
- Support Vector Machines
- **KNN**
- Explainability and Interpretability
- Appendix

Machine Learning is a huge field, and here I focus on a restricted set of topics, as an introduction into the subject.

Where algorithms or issues are explored in more depth this is generally done for pedagogical reasons, or the algorithm is widely used, or because the issue is general to the field.

OVERVIEW

- Examples are provided for the tutorials.
- These use ROOT or Python and have been tested using the following versions:
 - ▶ ROOT: 6.06/02
 - Python 3.7 (via an Anaconda install) with the following modules
 - TensorFlow 2.2 [Note: examples are not taking advantage of eager execution and are using the V1 backward compatibility mode]
 - matplotlib
 - numpy
 - sklearn



WHAT IS MACHINE LEARNING?



WHAT IS MACHINE LEARNING?

- Oxford English Dictionary:
 - ▶ "a type of artificial intelligence in which computers use huge amounts of data to learn how
 to do tasks rather than being programmed to do them"
- Collins Dictonary:
 - "a branch of artificial intelligence in which a computer generates rules underlying or based on raw data that has been fed into it"
- Google Developers glossary:
 - "A program or system that builds (trains) a predictive model from input data. The system uses the learned model to make useful predictions from new (never-before-seen) data drawn from the same distribution as the one used to train the model. Machine learning also refers to the field of study concerned with these programs or systems."
- There is no single definition agreed of machine learning, so I will use a working definition.
 - "The process of using an algorithm to approximate data using some underlying optimisation heuristic"



WHAT IS MACHINE LEARNING?

- "The process of using an algorithm to approximate data using some underlying optimisation heuristic"
- We are doing function approximation using some algorithm fit to some reference data using some heuristic.
- The fitting in this context is referred to as training or learning.
 - There are different learning paradigms, we will focus on supervised and unsupervised learning.
- The trained function is used as a model (i.e. for prediction).
- Dimensionality and parameters are implied in these slides when referring to general situations, i.e. $f(\underline{x},\underline{\theta}) \to f(x,\theta) \to f(x)$.

How does ethics fit into a lecture on machine learning?

Ethics plural in form but singular or plural in construction: the discipline dealing with what is good and bad and with moral duty and obligation.

Morals describes one's particular values concerning what is right and what is wrong.

- How does ethics fit into a lecture on machine learning?
 - Is it ethical to develop an algorithm to identify the political leaning of an individual with the intent of targeting advertising, to polarise the viewpoint of voters with positive reinforcement messages or fake news, to undermine or influence the outcome of an election?
 - Is it ethical to develop a pandemic model (e.g. using an SIR approach^[1]) using an AI, that could influence Government policy, without fully testing the robustness of predictions?

[1] e.g. see this article on wikipedia Compartmental models in epidemiology. A good illustration of a variant of this model can be found at https://upload.wikimedia.org/wikipedia/commons/3/31/SIRD_model_anim.gif



- But that is the "real world" why should I care about ethics?
 - Is it ethical to use an algorithm for science without checking that the result makes sense?

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- But that is the "real world" why should I care about ethics?
 - Is it ethical to use an algorithm for science without checking that the result makes sense?
 - Is it ethical to use an energy intensive algorithm when a computationally cheap alternative that performs just as well exists?
- What we think of as good scientific practice, is also ethical behaviour. It leads to robust results.
- Ethical behaviour in the wider world can have deeper ramifications than getting a robust scientific result or not.

- But that is the "real world" why should I care about ethics?
 - In reality most people who do a PhD in an STFC (or EPSRC) area of science will not end up in academia, and ethics will play a role beyond scientific correctness for those scientists.
 - For those of us who do stay in academia, then we have an obligation to help people understand the ramifications of algorithms and our rationale for using them (or not).
 - For all of us, we have transferable skills that can be applied to real world problems.



- ▶ The UK Government developed a data ethics framework^[1]:
 - "Public sector organisations should use the Data Ethics Framework to guide the appropriate use of data to inform policy and service design"



Other organisations such as the UN and IEEE (as well as the EU) are also concerned about ethical use of data, and ethical algorithms, from regulatory and the perspective of rights.





DATA WRANGLING (AKA FEATURE ENGINEERING)

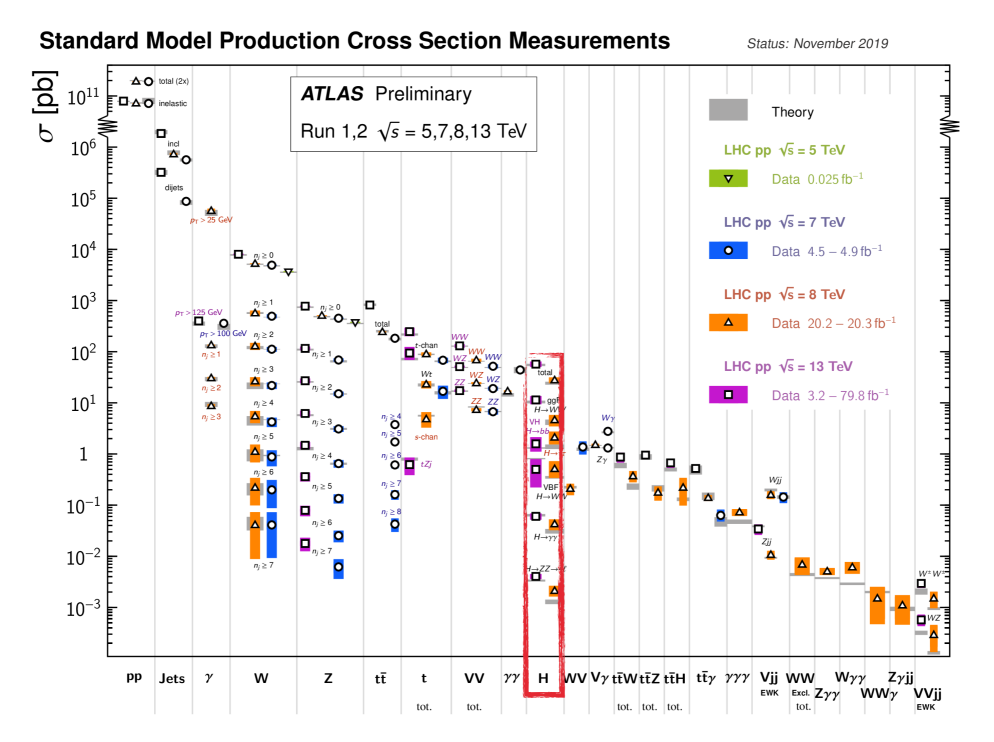
- Need to understand the data being analysed.
 - Domain context will provide most insight into getting information to highlight the solution to your problem.
 - Identify the features of interest in the data.
 - Allow you to reduce the dimensionality of the problem into as few a set of inputs as possible.
 - Most of the analysis can be done with this domain context background:
 - Deep Learning (DL) can replace the hard work of feature engineering for a resource cost and a lack of explainability and interpretability. [1]
 - "Smart Learning" (SL) is an alternative way of approaching the problem. [2]

^[2] I attribute the term Smart Learning to the use of domain knowledge and understandable AI, such as Bayesian Networks. A term that I first heard about from Norman Fenton who has a good book on the topic.

A. Bevan

- In HEP we use cut based selection to:
 - Remove pathological data (poorly calibrated or partially complete data, bad beam conditions, etc).
 - Remove obvious background examples from the data (well known SM processes that are just not interesting for study, or use as a calibration or control sample).
 - Prepare data for the "statistical analysis" that will include the use of multivariate analysis techniques and fitting.

• e.g. Higgs physics: Throw away $\sim 10^9$ recorded events for each interesting one.





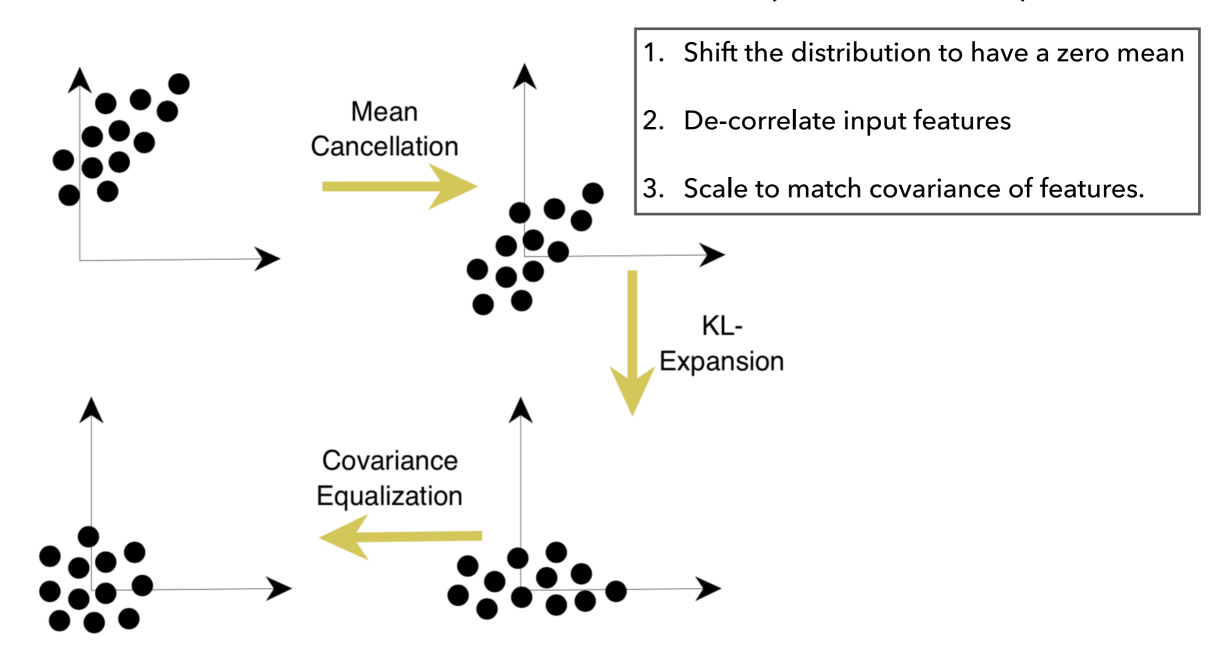
- Knowing what the task is allows one to identify the features of interest. This is the domain context knowledge.
 - If your signal is the process pp→HH+X, then:
 - Laws of physics provide insights for you to refine your list of features to train on.
 - e.g. the Higgs mass or p_T will play a role in identifying the signal, along with decay product properties (e.g. b-tag quality), etc.

- Reduce the number of dimensions of interest:
 - Trial and error with different inputs to identify what improves performance of the algorithm and what does not.
 - Linearly correlated features can be combined to reduce the number of features providing information for the algorithm to learn from.
 - Principal Component Analysis (PCA) to address this problem.
 - Some neural network configurations do that automatically for you (e.g. auto-encoders).
 - Let the algorithm learn what is important and what is not.
- Some algorithms (e.g. support vector machines) implicitly increase the dimensionality of the problem.

ACTIVATION FUNCTIONS: DATA PREPARATION

- Input features are arbitrary; whereas (for example) activation functions in neural networks work best for a standardised input domain of [-1, 1] or [0, 1].
 - We can map our input feature space onto a standardised domain that matches some range that matches that of the activation function.
 - Saves work for the optimiser in determining hyper-parameters.
 - > Standardises weights to avoid numerical inaccuracies; and set common starting weights.
 - e.g.
 - having an energy or momentum measured in units of 10^{12} eV, would require weights $O(10^{-12})$ to obtain an O(1) result for $w_i x_i$.
 - ▶ Mapping eV \mapsto TeV would translate 10^{12} eV \mapsto 1TeV, and allow for O(1) weights leading to an O(1) result for $w_i x_i$.
 - Comparing weights for features that are standardised allows the user to develop an intuition as to what the corresponding activation function will look like.

Variable transformations are also useful (can be vital).



- Data wrangling is an important part of ensuring you get the most out of applying machine learning; yet it is often not celebrated as the requirements can be very problem specific.
- If you are interested in this topic you may wish to review the data wrangling presentation and tutorials given by Dr Nick Barlow from the Alan Turing Institute at the GradNet meeting on Machine Learning and AI in January 2020:
 - Dr Barlow's talk and lecture can be found at: https://
 indico.ph.qmul.ac.uk/indico/conferenceOtherViews.py?
 view=standard&confld=543

TYPES OF LEARNING
GRID SEARCH
GRADIENT DESCENT
ADAM OPTIMISER
MISCELLANEOUS

OPTMISATION



OPTIMISATION TYPES OF LEARNING



TYPES OF LEARNING

- Consider a model y that depends on a parameter set θ , we can select a point in the parameter hyperspace $\hat{\theta}$ that has a corresponding estimator of the function \hat{y} .
- lacktriangle The heta are hyper-parameters (HPs) of the model.

Unsupervised learning:

- Infer the probability distribution $P(\hat{y})$ from the data without using labels.
- Widely used in many fields of research.

Supervised learning:

- Use training examples from labeled data sets with a known type or value, t, for each example.
- **>** Some loss function is used to compare t against model predictions \widehat{y} .
- ▶ Can be thought of as computing a conditional probability $P(t \mid \widehat{y})$.
- This is the most commonly used approach in particle physics today.



TYPES OF LEARNING: SUPERVISED LEARNING

- Define a heuristic based on some figure of merit (FOM) designed to improve the value of that metric through some iterative process.
- The FOM is called the objective function or loss function or cost function in machine learning.
- e.g. the L₂ norm loss function: this is like a χ^2 , but without the error term:

$$L_2 = \sum_{i=1}^{N_{examples}} \left[t_i - \hat{y}(\hat{\theta}) \right]^2$$

The θ are called HPs as they form a hyperspace; other variables in the optimisation process are often included in the set of HPs (e.g. learning rate for a neural network, cost for a support vector machine, tree depth for a decision tree etc.).

TYPES OF LEARNING

Batch learning

Use a sample (batch) of training data to evaluate an estimate of the error and update weights in the optimisation.

Advantages of Batch Learning

- 1. Conditions of convergence are well understood.
- 2. Many acceleration techniques (e.g. conjugate gradient) only operate in batch learning.
- 3. Theoretical analysis of the weight dynamics and convergence rates are simpler.

Stochastic learning

Use individual training examples to evaluate an estimate of the error and update weights in the optimisation.

Advantages of Stochastic Learning

- 1. Stochastic learning is usually *much* faster than batch learning.
- 2. Stochastic learning also often results in better solutions.
- 3. Stochastic learning can be used for tracking changes.



- •THIS IS A SIMPLE OPTIMISATION ALGORITHM THAT CAN BE USED WITH NO PROGRAMMING EXPERIENCE.
- •ONE CAN IMPLEMENT A 1 OR 2 DIMENSIONAL GRID SEARCH USING AN EXCEL SPREADSHEET, AND FROM INSPECTION OF THE TABULATED RESULTS YOU CAN OPTIMISE SIMPLE PROBLEMS.
- •THIS IS A BRUTE FORCE OPTIMISATION APPROACH, AND IT IS USED WIDELY IN CERTAIN APPLICATIONS ACROSS A NUMBER OF RESEARCH FIELDS.

OPTIMISATION GRID SEARCH

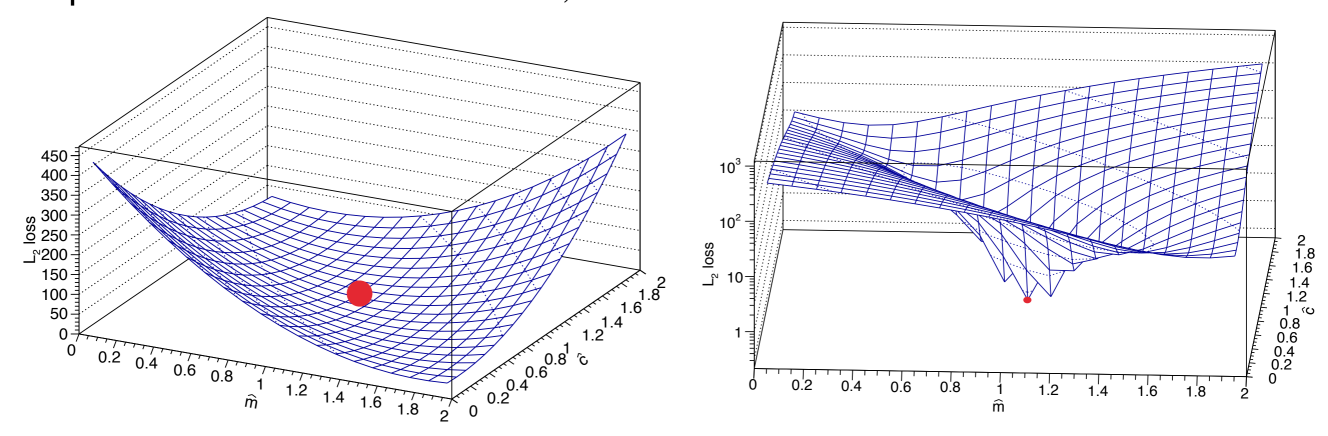


GRID SEARCH

- Scanning through the hyperspace for each $\hat{\theta}_i$ allows us to compute L₂, and the minimum value obtained is the "best fit" or optimal estimate of the parameters θ .
 - Simple heuristic to implement you can do this in Excel for 1 or 2D problems. ✓
 - ▶ Easy to understand. ✓
 - Expensive to compute: scanning n points in a dimension requires n^M computations for $M = dim(\theta)$.
- Heuristic suffers from the curse of dimensionality.

GRID SEARCH

• e.g. Consider an L2 loss function optimisation of the HPs for y = mx + c, i.e. optimise m and c. Here m = 1.0, c = 1.0



- ▶ The contours of the loss function show a minimum.
- The optimal value is selected from a discrete grid of points, only get an exact result if the grid maps onto the problem perfectly (as in this example).

GRID SEARCH

- e.g. The libsvm^[1] package uses a HP grid search for optimisation for the Support Vector Machine algorithm; where the cost C and Γ hyper-parameters need to be optimised.
 - The grid search is done efficiently by adapting to whatever step is sensible, in this case the algorithm has Γ as the parameter of an exponential (the radial basis function); and so a linear search in C and a logarithmic search in Γ is appropriate to sample the parameter hyperspace effectively.

[1] Chih-Chung Chang and Chih-Jen Lin, LIBSVM: a library for support vector machines. ACM Transactions on Intelligent Systems and Technology, 2:27:1--27:27, 2011. Software available at http://www.csie.ntu.edu.tw/~cjlin/libsvm.

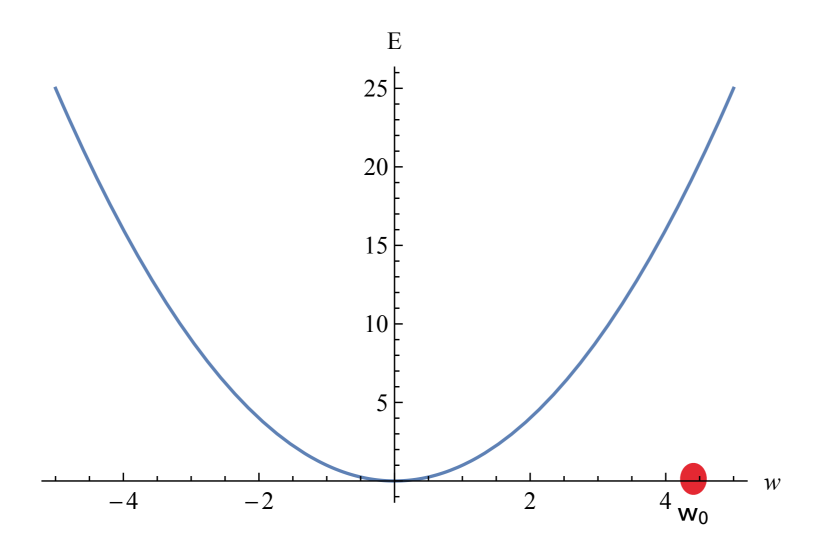


OPTIMISATION GRADIENT DESCENT



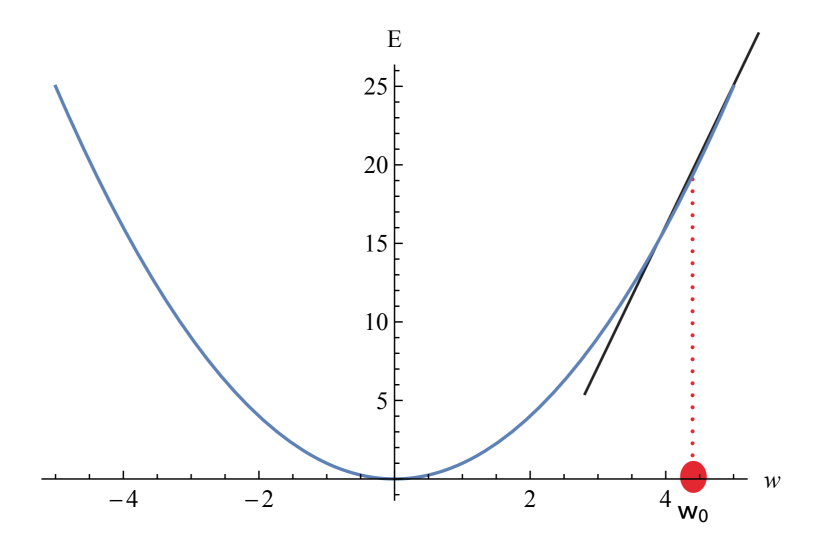
GRADIENT DESCENT

 \blacktriangleright Guess an initial value for the weight parameter: w_0 .



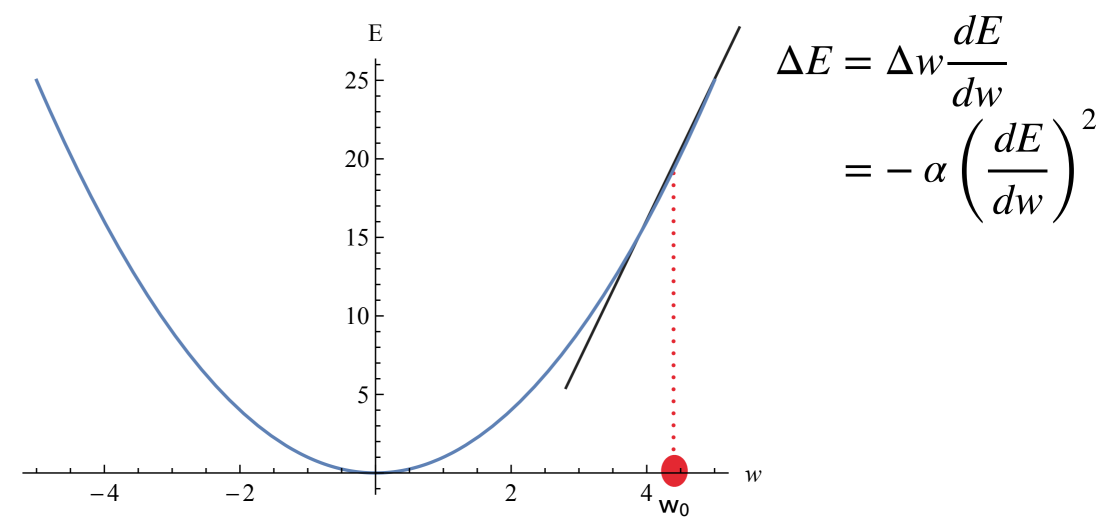
GRADIENT DESCENT

Estimate the gradient at that point (tangent to the curve)



GRADIENT DESCENT

• Compute Δw such that ΔE is negative (to move toward the minimum)



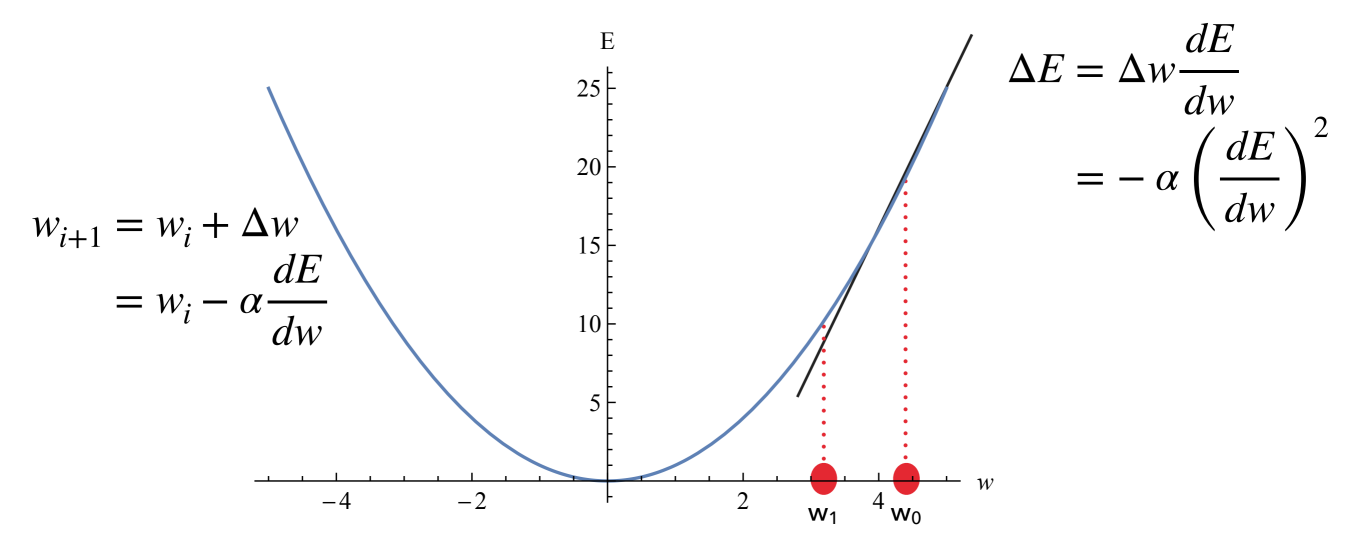
 α is the learning rate: a small positive number

Choose
$$\Delta w = -\, \alpha \frac{dE}{dw}$$
 to ensure $\Delta {\rm E}$ is always negative.



GRADIENT DESCENT

▶ Compute a new weight value: $w_1 = w_0 + \Delta w$



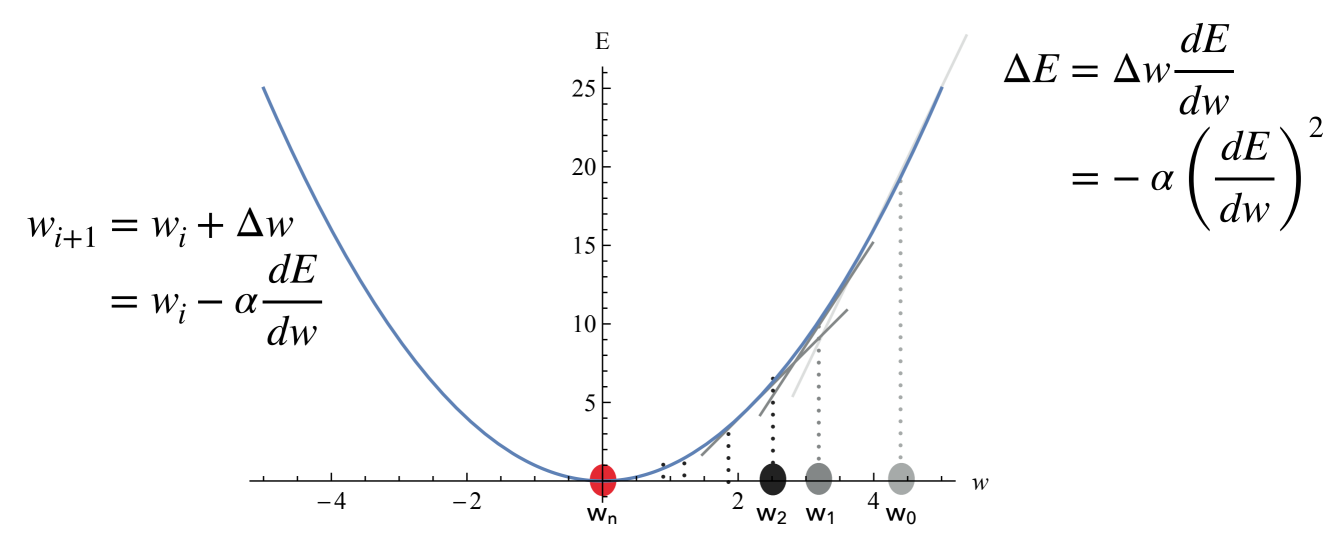
 α is the learning rate: a small positive number

Choose
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 to ensure $\Delta {\rm E}$ is always negative.



GRADIENT DESCENT

Repeat until some convergence criteria is satisfied.



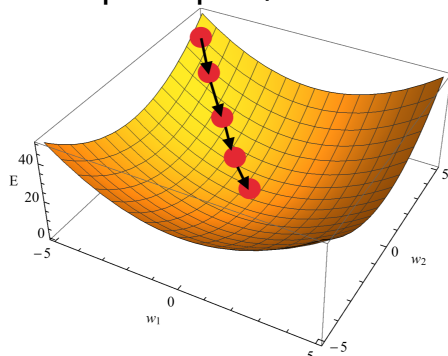
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$$\alpha$$
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GRADIENT DESCENT

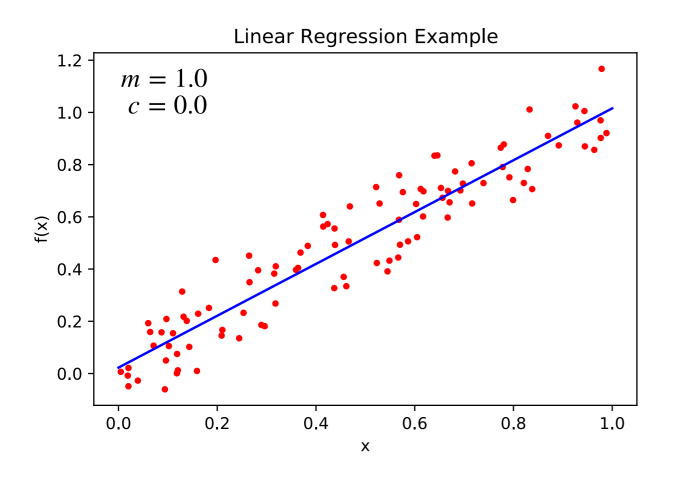
We can extend this from a one parameter optimisation to a 2 parameter one, and follow the same principles, now in 2D.

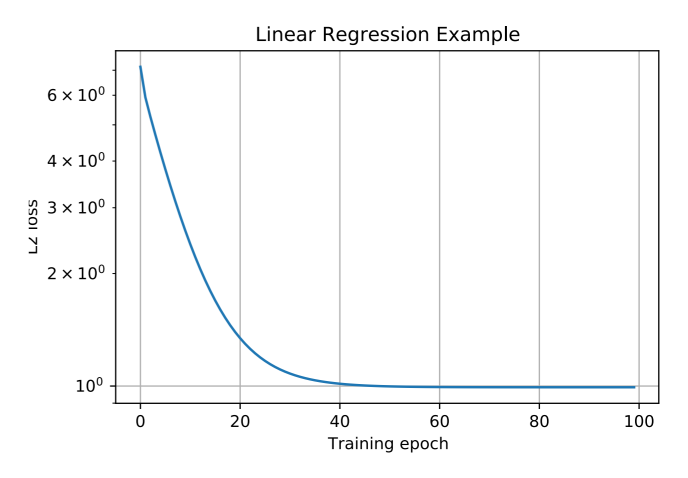


- The successive points w_i+1 can be visualised a bit like a ball rolling down a concave hill into the region of the minimum.
- In general update weights such that $\Delta E = \Delta w \nabla E = -\alpha \nabla^2 E$
- and $w_{i+1} = w_i \alpha \nabla E$.

GRADIENT DESCENT: EXAMPLE

Returning to the y = mx + c example, we can optimise this using the Gradient Descent algorithm.





Use N=100 and $\alpha=0.005$ with the TensorFlow GradientDescent optimiser to obtain:

$$\widehat{m} = 0.9928...$$

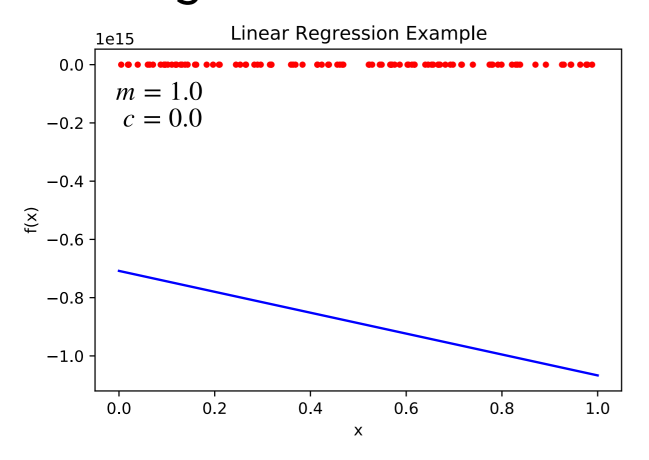
$$\hat{c} = 0.0226...$$

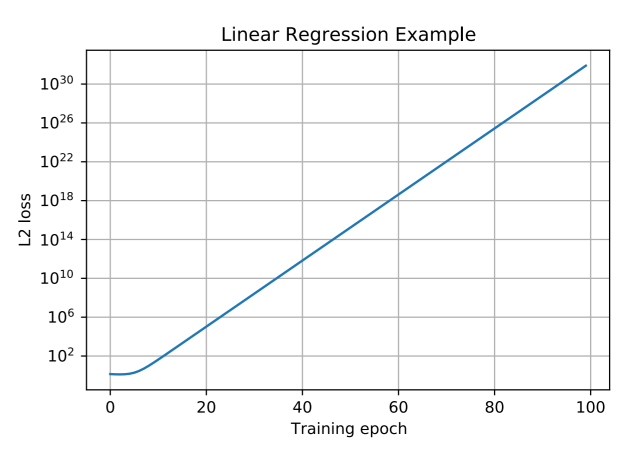
Implementation: TensorFlow 2.2 (using V1 compatibility) Jupyter Notebook



GRADIENT DESCENT: EXAMPLE

• Returning to the y = mx + c example, we can optimise this using the Gradient Descent algorithm.





Use N=100 and $\alpha = 0.01$ with the TensorFlow GradientDescent optimiser to obtain:

$$\widehat{m} = -3.5 \times 10^{14}$$

 $\widehat{c} = -7.08 \times 10^{14}$

The minimiser fails; too large a learning rate is being used.

Implementation: TensorFlow 2.2 (using V1 compatibility)
Jupyter Notebook



GRADIENT DESCENT: REFLECTION

- ▶ The examples shown illustrate problems with parabolic minima.
- With selection of an appropriate learning rate, α , to fix the step size, we can guarantee convergence to a sensible minimum in some number of steps.
- Translating the distribution to a fixed scale, then we can predict how many steps it will take to converge to the minimum from some distance away from it for a given α .
- If the problem hyperspace is not parabolic, this becomes more complicated, and there is no guarantee that we converge to the minimum.
- Modern machine learning algorithms use more refined variants on this method.

OPTIMISATION ADAM OPTIMISER



(ADAM: ADAptive Moment estimation based on gradient descent)

- This is a stochastic gradient descent algorithm.
- **Consider a model** $f(\theta)$ that is differentiable with respect to the HPs θ so that:
 - ightharpoonup t is the training epoch.
 - the gradient $g_t = \nabla f_t(\theta_{t-1})$ can be computed.
 - $m_t(v_t)$ are biased values of the first (second) moment.
 - $\widehat{m}_{t}(\widehat{v}_{t})$ are bias corrected estimator of the moments.
 - Some initial guess for the HP is taken: θ_0 , and the HPs for a given epoch are denoted by θ_r .
 - $m{\alpha}$ is the step size (i.e. learning rate).
 - β_1 and β_2 are exponential decay rates of moments.

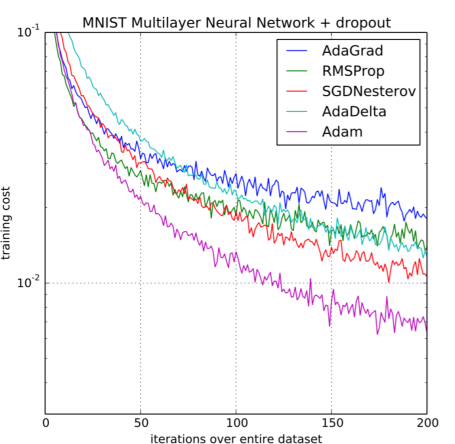
(ADAM: ADAptive Moment estimation based on gradient descent)

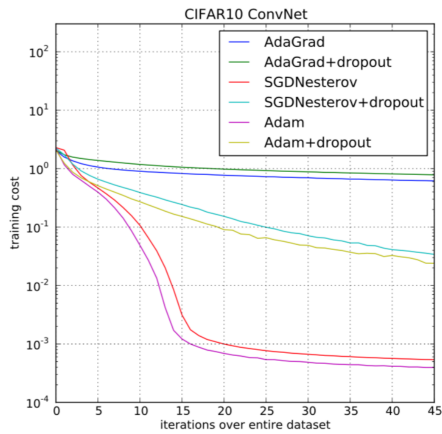
Algorithm 1: Adam, our proposed algorithm for stochastic optimization. See section 2 for details, and for a slightly more efficient (but less clear) order of computation. g_t^2 indicates the elementwise square $g_t \odot g_t$. Good default settings for the tested machine learning problems are $\alpha = 0.001$, $\beta_1 = 0.9$, $\beta_2 = 0.999$ and $\epsilon = 10^{-8}$. All operations on vectors are element-wise. With β_1^t and β_2^t we denote β_1 and β_2 to the power t.

```
Require: \alpha: Stepsize
Require: \beta_1, \beta_2 \in [0, 1): Exponential decay rates for the moment estimates
Require: f(\theta): Stochastic objective function with parameters \theta
Require: \theta_0: Initial parameter vector
   m_0 \leftarrow 0 (Initialize 1<sup>st</sup> moment vector)
   v_0 \leftarrow 0 (Initialize 2<sup>nd</sup> moment vector)
   t \leftarrow 0 (Initialize timestep)
   while \theta_t not converged do
       t \leftarrow t + 1
       g_t \leftarrow \nabla_{\theta} f_t(\theta_{t-1}) (Get gradients w.r.t. stochastic objective at timestep t)
       m_t \leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t (Update biased first moment estimate) v_t \leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2 (Update biased second raw moment estimate)
       \widehat{m}_t \leftarrow m_t/(1-\beta_1^t) (Compute bias-corrected first moment estimate)
       \widehat{v}_t \leftarrow v_t/(1-\beta_2^t) (Compute bias-corrected second raw moment estimate)
       \theta_t \leftarrow \theta_{t-1} - \alpha \cdot \widehat{m}_t / (\sqrt{\widehat{v}_t} + \epsilon) (Update parameters)
   end while
   return \theta_t (Resulting parameters)
```

(ADAM: ADAptive Moment estimation based on gradient descent)

Benchmarking performance using MNIST and CFAR10 data indicates that Adam with dropout minimises the loss function compared with other optimisers tested.





Faster grop oπ in loss, and lower overall loss obtained vs other algorithms benchmarked at the time.

(ADAM: ADAptive Moment estimation based on gradient descent)

Benchmarking performance using MNIST and CFAR10 data indicates that

MNIST and CFAR10 are standard benchmark data sets in CS (see appendix). MNIST is a set of handwritten numbers 0 through 9; CFAR10(0) is an image classification data set (cars, boats etc).



 Faster grop oπ in ioss, and iower overall loss obtained vs other algorithms benchmarked at the time.

MULTIPLE MINIMA MORE ON LOSS FUNCTIONS

OPTIMISATION MISCELLANEOUS



GRADIENT DESCENT: MULTIPLE MINIMA

 Often more complication hyperspace optimisation problems are encountered, where there are multiple

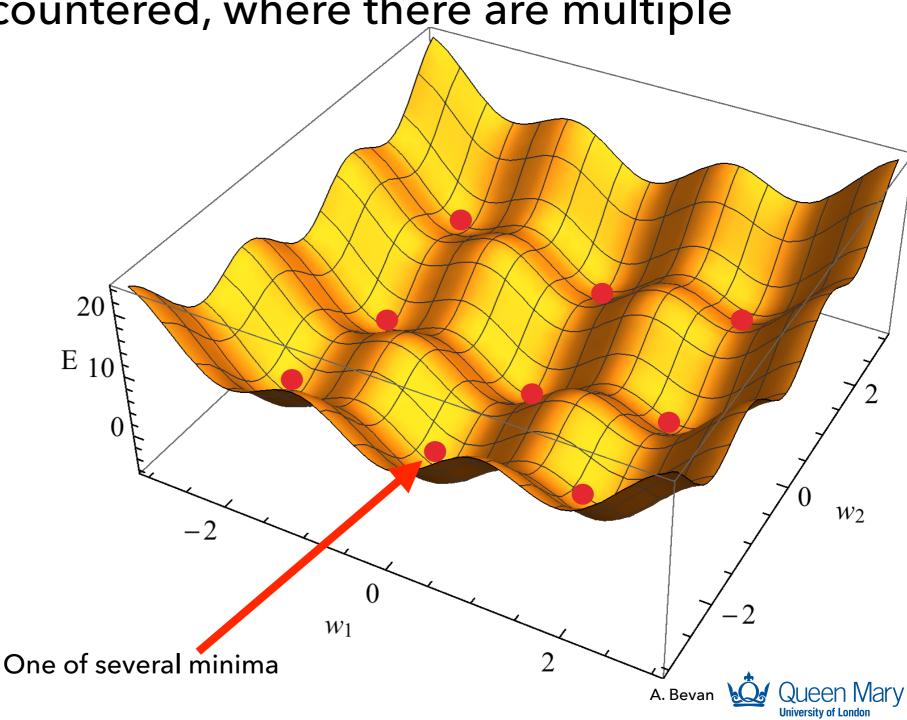
minima.

The gradient descent minimisation algorithm is based on the assumption that there is a single minimum to be found.

In reality there are often multiple minima.

Sometimes the minima are degenerate, or near degenerate.

How do we know we have converged on the global minimum?



GRADIENT DESCENT: MULTIPLE MINIMA

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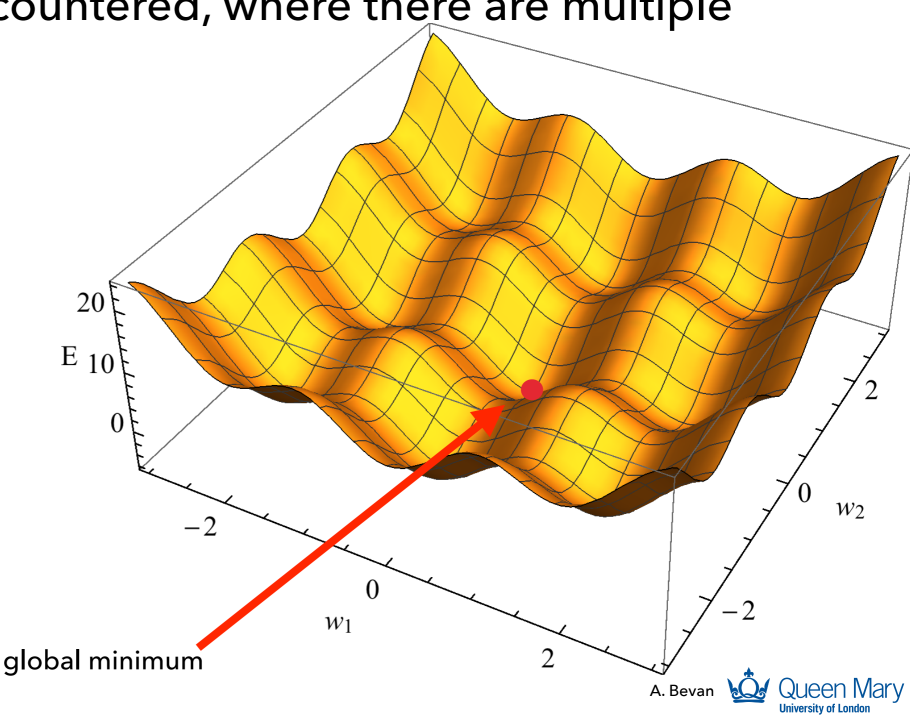
minima.

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How do we know we have converged on the global minimum?



OPTIMISATION: LOSS FUNCTIONS

- ▶ There are many types of loss function other than the L₂ loss
 - L1 norm loss:

$$L_1 \sum_{i=1}^{N_{examples}} \left| t_i - \hat{y}(\hat{\theta}) \right|$$

The mean square error (MSE) loss function:

$$L = \frac{1}{N}L_2 = \frac{1}{N}\sum_{i=1}^{N_{examples}} \left[t_i - \hat{y}(\hat{\theta})\right]^2$$

Cross entropy

$$L = -\sum_{i=1}^{N_{examples}} \widehat{y}(x_i) \ln t_i$$

The cross entropy can be thought of as the negative log likelihood function for the data t_i under the model \widehat{y}

SUPERVISED LEARNING OVERTRAINING

WEIGHT REGULARISATION CROSS VALIDATION

DROPOUT

TRAINING

TRAINING SUPERVISED LEARNING



SUPERVISED LEARNING

For supervised learning the loss function depends on both model parameters and a known target value t_i for a given example.

e.g. the L₂ loss:
$$L_2 = \sum_{i=1}^{N_{examples}} \left[t_i - \hat{y}(\hat{\theta}) \right]^2$$
.

- This requires (at a minimum) a sample of data with the input feature space of interest, and for each example in the data, the known target output value.
- ▶ The loss function is optimised using the data, to obtain a model.
- Unlike fitting however we don't care about the uncertainty on the model parameter estimates, only on the nominal values obtained, $\hat{\theta}$.
- Due to the complexity of models, it is generally not possible to understand if the optimisation converged to a sensible solution by inspecting marginalised projections of the model on the data.

SUPERVISED LEARNING

• e.g. the <u>Kaggle Higgs</u> data sample:

EventId, DER_mass_MMC, DER_mass_transverse_met_lep, ..., Weight Label, KaggleSet, KaggleWeight

Features (not all to be used for training)

Known example type

- The KaggleSet column in the CVS file for this data allows the user to understand if this is to be used for training or testing, or some other means (e.g. Kaggle Score Board evaluation).
- The Label column, is the known label; this defines the t_i used in the loss function.
- I use this challenge data as an assignment for undergraduate lectures (pdf, zip). N.B. the zip file is 189Mb.

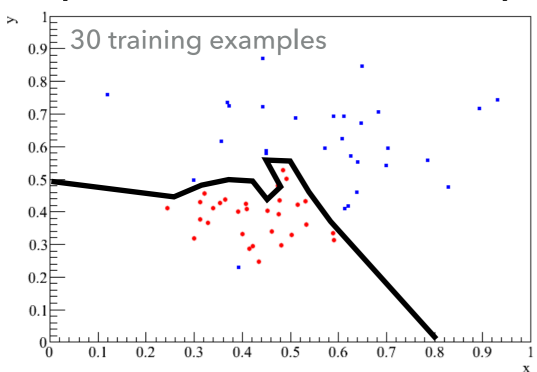
WEIGHT REGULARISATION CROSS VALIDATION

TRAINING OVERTRAINING



A model is over trained if the HPs that have been determined are tuned to the statistical fluctuations in the data set.

Simple illustration of the problem:



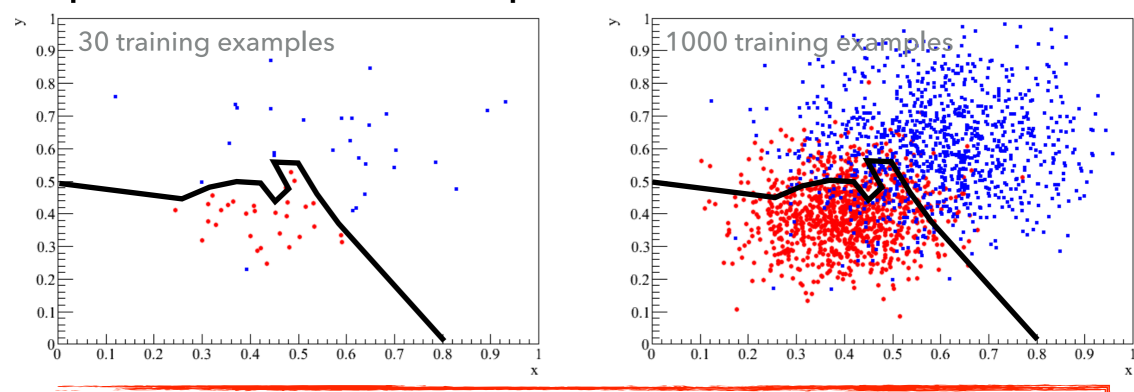
The decision boundary selected here does a good job of separating the red and blue dots.

Boundaries like this can be obtained by training models on limited data samples. The accuracies can be impressive.

But would the performance be as good with a new, or a larger data sample?

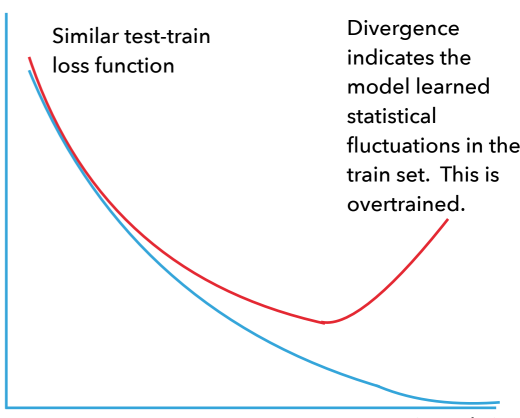
 A model is over trained if the HPs that have been determined are tuned to the statistical fluctuations in the data set.

Simple illustration of the problem:



Increasing to 1000 training examples we can see the boundary doesn't do as well. This illustrates the kind of problem encountered when we overfit HPs of a model.

- One way to avoid tuning to statistical fluctuations in the data is to impose a training convergence criteria based on a data sample independent from the training set: a test sample.
 - Use the loss evaluated for the training and test samples to check to see if the HPs are over trained.
 - If both samples have similar loss then the model response function is similar on two statistically independent samples.
 - Note: If the samples are large enough then one could reasonably assume that the response function would then be general when applied to an unseen data sample.
 - "large enough" is a model and problem dependent constraint.



Training iterations (epochs)

- Training convergence criteria that could be used:
 - Terminate training after N_{epochs}
 - Loss comparison:
 - Evaluate the performance on the training and validation sets.
 - Compare the two and place some threshold on the difference $\Delta_{LOSS} < \delta_{LOSS}$
 - Terminate the training when the gradient of the loss function with respect to the weights is below some threshold.
 - > Terminate the training when the Δ_{LOSS} starts to increase for the validation sample.

OVERTRAINING: WEIGHT REGULARISATION

- Weight regularisation involves adding a penalty term to the loss function used to optimise the HPs of a network.
- This term is based on the sum of the weights w_i in the network and takes the form:

$$\lambda \sum_{i=\forall weights} w_i$$

- The rationale is to add an additional cost term to the optimisation coming from the complexity of the network.
- \blacktriangleright The performance of the network will vary as a function of λ .
- To optimise a network using weight regularisation it will have to be trained a number of times in order to identify the value corresponding to the min(cost) from the set of trained solutions.

OVERTRAINING: WEIGHT REGULARISATION

For example we can consider extending an MSE loss function to allow for weight regularisation. The MSE loss is given by:

$$\varepsilon = \frac{1}{N} \sum_{i=1}^{N} (y_i - t_i)^2$$

To allow for regularisation we add the sum of weights term:

$$\varepsilon = \frac{1}{N} \sum_{i=1}^{N} (y_i - t_i)^2 + \lambda \sum_{i=\forall, weights} w_i$$

This is a simple modification to make to the NN training process.



OVERTRAINING: CROSS VALIDATION

- A well trained model will provide robust predictions, irrespective of the examples presented to it.
 - The variance of model predictions will be small.
 - The model predictions may be systematically biased independently of this.
- One can divide the training set up into k-folds, and then perform k trainings; each one leaving a single fold out.
- The amount of data used in a fold will depend (generally the more data the better.
- The ensemble of predictions indicates the variance on the model output.



OVERTRAINING: CROSS VALIDATION

- The use of cross validation to determine the spread of model predictions, and any systematic bias on prediction can be useful.
- e.g. in a physics context: Consider a new particle search at the Large Hadron Collider.
 - Using an overtrained model to suppress background and enhance signal will result in a lack of experimenter understanding as to how the expected and observed limits on the new particle relate to each other.
 - This could result in a false discovery of new physics.
 - It could result in missing out on a discovery of new physics.
 - Some people say it is not wrong to use an over trained model I argue strongly that it is not scientifically correct to use an over trained model, unless the variance on that model, and hence the implications are well understood.

TRAINING DROPOUT

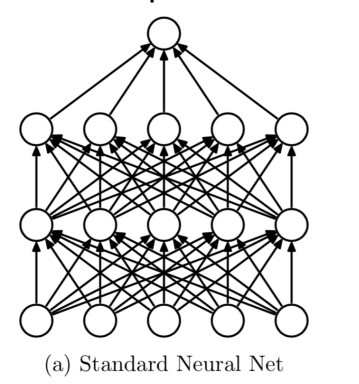


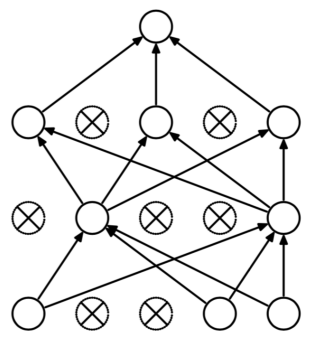
DROPOUT

- A problem with deep networks is the number of hyper-parameters that need to be determined.
- This leads to a requirement for very large data sets to avoid overfitting (or fine-tuning).
- Hyperparameters can also learn to "co-adapt" in the training process.
 - co-adapt means that as one parameter is changed, another in the network can be modified in a correlated way to offset that change.
 - This kind of behaviour intentionally exists in some algorithms (Sequential Minimal Optimisation for Support Vector machines, where pairs of HPs are changed to conserve an overall zero sum); but is generally unwelcome behaviour.

OVER FITTING: DROPOUT

A pragmatic way to mitigate these issues is to intentionally compromise the model randomly in different epochs of the training by removing units from the network.





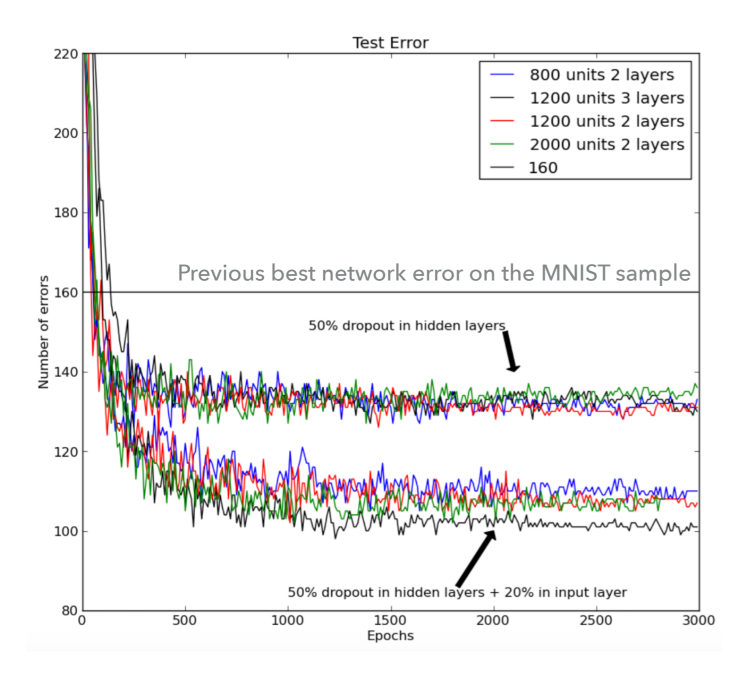
Dropout is used only during training.

The full model is used for making predictions.

(b) After applying dropout.

- That way the whole model will be effectively trained on a sub-sample of the data with the aim of limiting the ability to learn statistical fluctuations in the data, and mitigating the co-adaption issue.
- This does not remove the possibility that a model is overtrained, as with the previous discussion HP generalisation is promoted by using this method.

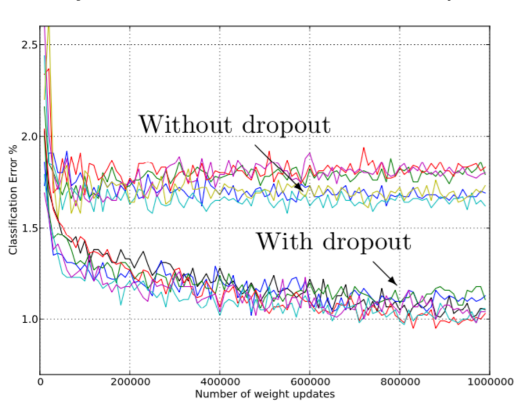
DROPOUT

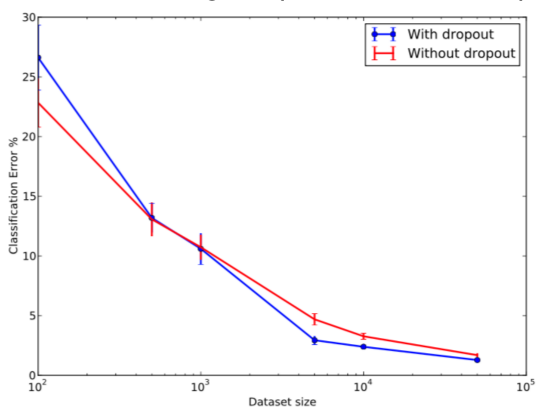


- Example from Hinton et al. for an MNIST sample.
- Using 50% drop out on the hidden layers gives an improvement over the previous best network architecture.
- Adding 20% drop out in the input layer provides further improvement in error.

OVER FITTING: DROPOUT

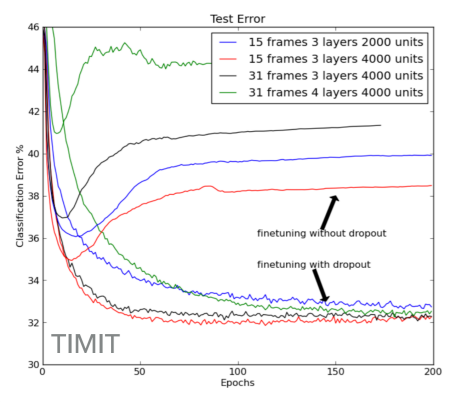
A variety of architectures has been explored with different training samples for this technique,

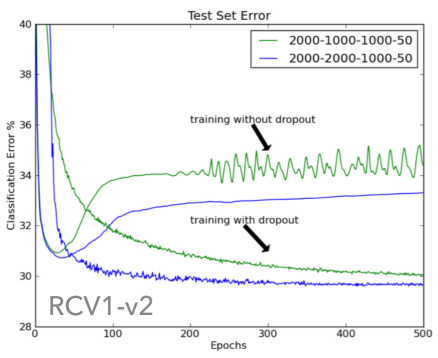




- Dropout can be detrimental for small training samples, however in general the results show that dropout is beneficial.
- For deep networks or typical training samples O(500) examples or more this technique is expected to be beneficial.
- For algorithms with very large training data sets, the benefits are less clear.

DROPOUT





- Example from Hinton et al. for a voice recognition sample of data (TIMIT, speech recognition data).
- Illustrates the classification error as a function of epoch with and without dropout.
- Similar results were obtained by Hinton and his team with the Reuters newsfeed data set (RCV1-v2).

DECISION TREES
BOOSTING

ADABOOST,M1

RANDOM FORESTS

DECISION TREES

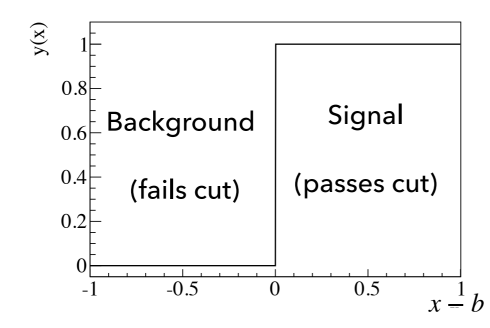


DECISION TREES

- The cut based and linear discriminant analysis methods are useful but limited [See Data Wrangling]
- The underlying concepts of applying Heaviside function constraints on data selection and on the use of a decision boundary definition (a plane in hyperspace) of the form of the dot product $\alpha^T x + \beta$ can be applied in more complicated algorithms.
- Here we consider extension to the concept of rectangular cuts to decision trees as a machine learning algorithm.
 - We will have to introduce the concepts of classification and regression; and methods to mitigate mis-classification of data.
 - The issue of overtraining is something we will come back to when discussing optimisation.



- Consider a data sample with an N dimensional input feature space X.
- X can be populated by examples from two or more different species of event (also called classes, categories or types).
- Consider the two types and call them signal and background, respectively*.
- We can use a Heaviside function to divide the data into two parts:
 - We can use this to distinguish between regions populated signal and background:



For an arbitrary cut position in x we can modify the Heaviside function

$$H(x) = \frac{1}{2}(1 + sign(x - b))$$

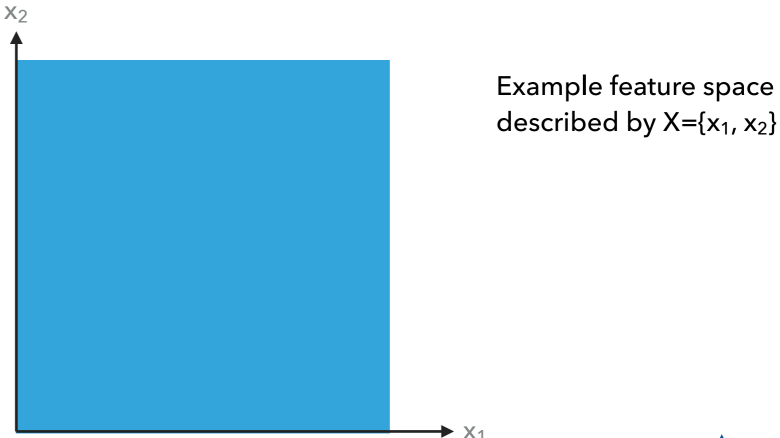
where b is the offset (bias) from zero.



- Decision trees divide the data feature space into a set of hypercubes that are classified as signal (+1) or background (-1) like.
 - Each region can be fitted with a constant to represent the data in that region.
 - We can recursively continue to sub-divide the data until some minimum number of examples are left in each sub-division.

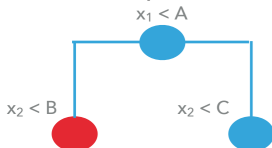
Can describe the data as the root node.

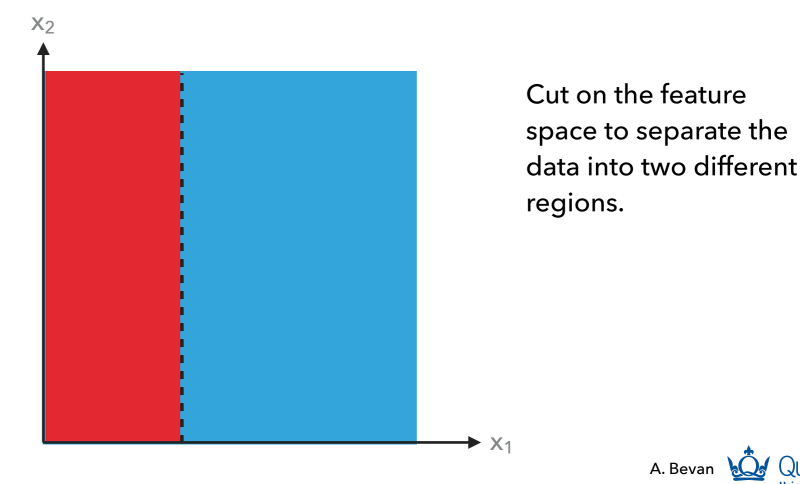




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The data get divided into two partitions.

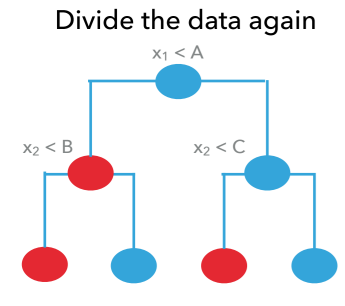


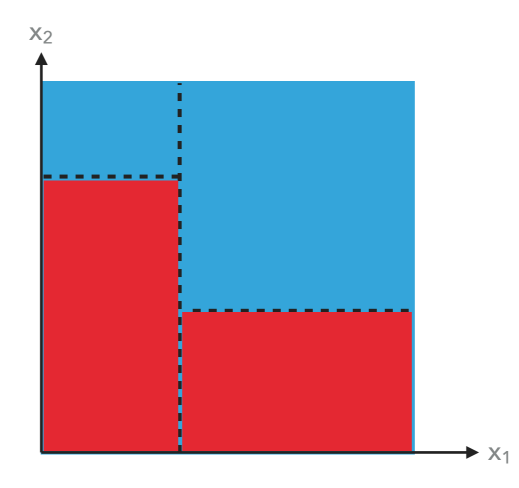




regions.

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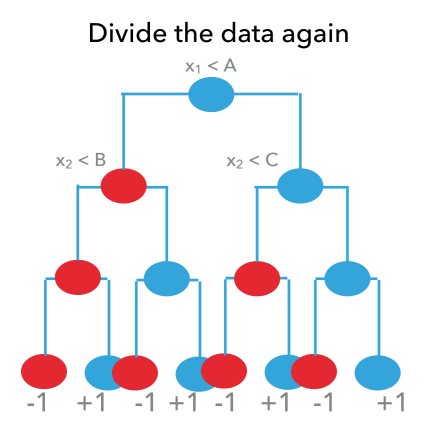


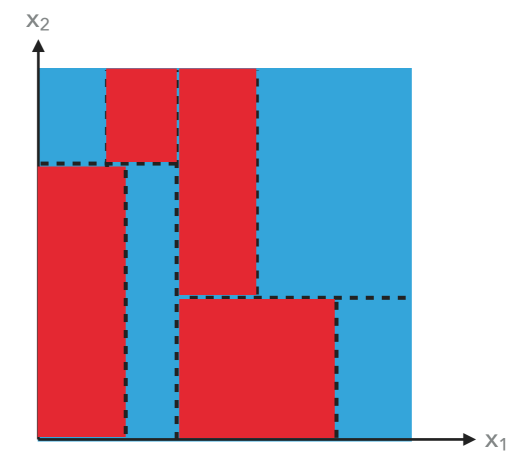


The feature space gets further sub-divided.



- Decision trees divide the data feature space into a set of hypercubes that are classified as signal (+1) or background (-1) like.
 - Each region can be fitted with a constant to represent the data in that region.
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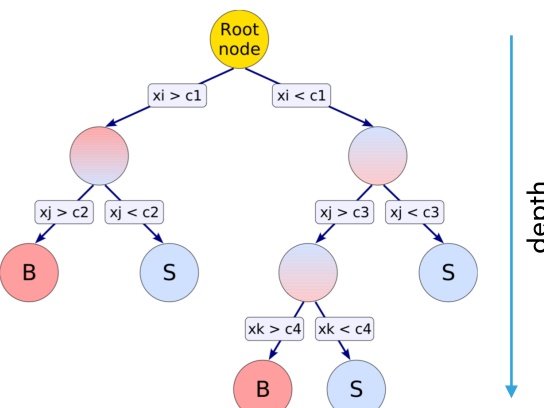
The feature space gets further sub-divided (again).



depth

DECISION TREES

- The set of rectangular cuts applied to the data allow us to build a tree from the root note.
- We can impose limits on:
 - Tree depth (how many divisions are performed).
 - Node size (how many examples per partition).
- Trees can be extended to more than 2 categories.
- They lend themselves to classifying examples or adapted to make a quantitative prediction (regression)



The decision tree output for a classification problem is

$$G(x) = +1 \text{ or } -1$$



- Decision trees are "weak learners", as they can take input features that only weakly separate types of example and combine those features to increase the separation.
- A single tree is susceptible to overtraining, and there are various methods of reducing this; including limiting the complexity of a tree, or the limiting the minimum number of examples in each node.
- The decision tree can be extended to an oblique decision tree, in which a linear combination of the features (instead of a single feature) is used to classify examples; so the Heaviside function cut becomes a hyperplane cut.

BOOSTING

- If a training example has been mis-classified in a training epoch, then the weight of that event can be increased for the next training epoch; so that the cost of mis-classification increases.
- The underlying aim is to take a weak learner and try and boost this into becoming a strong learner.
 - This example re-weighting technique is called boosting.
 - There are several re-weighting methods commonly used; here we discuss:
 - AdaBoost.M1 (popular variant of the Adaptive boosting method)
- Boosted Decision Trees are known as BDTs

- i is the ith example out of a data set with N examples.
- m is the mth training out of an ensemble of M learners to be trained.
- Step 1:
 - Assign event weights of $w_i = 1/N$ to all of the N examples.

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- Step 1:
 - Assign event weights of $w_i = 1/N$ to all of the N examples.
- Step 2: for m=1 through M
 - Train the weak learner (in our case this is a BDT): $G_m(x)$.
 - Compute the error rate \mathcal{E}_m .
 - Calculate the boost factor $\beta_m = \frac{\varepsilon_m}{1 \varepsilon_m}$.
 - Update weights for misclassified examples $w_i \mapsto w_i e^{\ln(1/\beta_m)}$.

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 - Update weights for misclassified examples $w_i \mapsto w_i e^{\ln(1/\beta_m)}$.
- Step 3:
 - Return the weighted committee: a combination of the M trees that have been

learned from the data:
$$G(x) = sign\left(\sum_{m=1}^{M} ln\left[\frac{1-\varepsilon_m}{\varepsilon_m}\right]G_m(x)\right)$$

▶ m=1

INITIAL TRAINING SAMPLE

▶ m=2

WEIGHTED SAMPLE

▶ m=3

WEIGHTED SAMPLE

 $-G_1(x)$

 $-G_3(x)$

 \bullet $G_M(x)$

The $G_m(x)$ are individual weak learners; each is derived from a training using the data.

The m=1 training uses the original data; all subsequent trainings use reweighted data.

m=M

WEIGHTED SAMPLE

$$G(x) = sign\left(\sum_{m=1}^{M} ln\left[\frac{1-\varepsilon_m}{\varepsilon_m}\right]G_m(x)\right)$$

The final classifier output is formed from a committee that is a weighted majority vote algorithm.

RANDOM FORESTS

- Random Forests are constructed from an ensemble of individual trees.
 - Each tree in the ensemble uses a randomly selected subset of the feature space, and the minimum node size is usually set to 1, so the classifier prediction is almost always accurate.
- The mode (classification) or mean (regression) of the ensemble is the output of the Random Forest.
- The probability that an example x_i is assigned to a given class c, is given by $P(c \mid x_i) = \frac{P(c \mid x_i)}{n}$

$$P(c | x_i) = \frac{P(c | x_i)}{\sum_{l=1}^{n} P(c_l | x_i)}$$

and the output score $g_c(x_i)$ is given by the aggregate over t trees in the forest:

$$g_c(x_i) = \frac{1}{t} \sum_{j=1}^t P_j(c \mid x_i)$$

The classification of x_i is simply the class c that maximises $g_c(x_i)$.



OVERVIEW

- Examples are provided for the tutorials.
- These use ROOT or Python and have been tested using the following versions:
 - ▶ ROOT: 6.06/02
 - > Python 3.7 (via an Anaconda install) with the following modules
 - TensorFlow 2.2 [Note: examples are not taking advantage of eager execution and are using the V1 backward compatibility mode]
 - matplotlib
 - numpy
 - sklearn



WHAT IS MACHINE LEARNING?



WHAT IS MACHINE LEARNING?

- Oxford English Dictionary:
 - ▶ "a type of artificial intelligence in which computers use huge amounts of data to learn how
 to do tasks rather than being programmed to do them"
- Collins Dictonary:
 - "a branch of artificial intelligence in which a computer generates rules underlying or based on raw data that has been fed into it"
- Google Developers glossary:
 - "A program or system that builds (trains) a predictive model from input data. The system uses the learned model to make useful predictions from new (never-before-seen) data drawn from the same distribution as the one used to train the model. Machine learning also refers to the field of study concerned with these programs or systems."
- There is no single definition agreed of machine learning, so I will use a working definition.
 - "The process of using an algorithm to approximate data using some underlying optimisation heuristic"



WHAT IS MACHINE LEARNING?

- "The process of using an algorithm to approximate data using some underlying optimisation heuristic"
- We are doing function approximation using some algorithm fit to some reference data using some heuristic.
- The fitting in this context is referred to as training or learning.
 - There are different learning paradigms, we will focus on supervised and unsupervised learning.
- The trained function is used as a model (i.e. for prediction).
- Dimensionality and parameters are implied in these slides when referring to general situations, i.e. $f(\underline{x},\underline{\theta}) \to f(x,\theta) \to f(x)$.



How does ethics fit into a lecture on machine learning?

Ethics plural in form but singular or plural in construction: the discipline dealing with what is good and bad and with moral duty and obligation.

Morals describes one's particular values concerning what is right and what is wrong.

- How does ethics fit into a lecture on machine learning?
 - Is it ethical to develop an algorithm to identify the political leaning of an individual with the intent of targeting advertising, to polarise the viewpoint of voters with positive reinforcement messages or fake news, to undermine or influence the outcome of an election?
 - Is it ethical to develop a pandemic model (e.g. using an SIR approach^[1]) using an AI, that could influence Government policy, without fully testing the robustness of predictions?

[1] e.g. see this article on wikipedia Compartmental models in epidemiology. A good illustration of a variant of this model can be found at https://upload.wikimedia.org/wikipedia/commons/3/31/SIRD_model_anim.gif



- But that is the "real world" why should I care about ethics?
 - Is it ethical to use an algorithm for science without checking that the result makes sense?

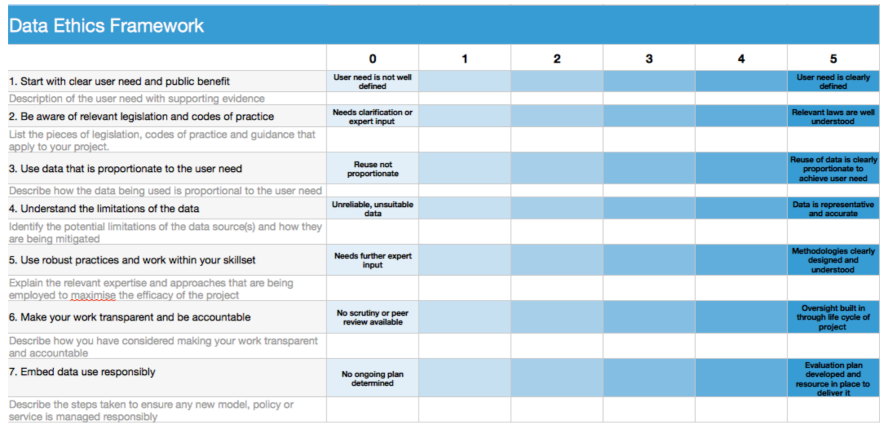
- But that is the "real world" why should I care about ethics?
 - Is it ethical to use an algorithm for science without checking that the result makes sense?
 - Is it ethical to use an energy intensive algorithm when a computationally cheap alternative that performs just as well exists?

- But that is the "real world" why should I care about ethics?
 - Is it ethical to use an algorithm for science without checking that the result makes sense?
 - Is it ethical to use an energy intensive algorithm when a computationally cheap alternative that performs just as well exists?
- What we think of as good scientific practice, is also ethical behaviour. It leads to robust results.
- Ethical behaviour in the wider world can have deeper ramifications than getting a robust scientific result or not.

- But that is the "real world" why should I care about ethics?
 - In reality most people who do a PhD in an STFC (or EPSRC) area of science will not end up in academia, and ethics will play a role beyond scientific correctness for those scientists.
 - For those of us who do stay in academia, then we have an obligation to help people understand the ramifications of algorithms and our rationale for using them (or not).
 - For all of us, we have transferable skills that can be applied to real world problems.



- ▶ The UK Government developed a data ethics framework^[1]:
 - "Public sector organisations should use the Data Ethics Framework to guide the appropriate use of data to inform policy and service design"



Other organisations such as the UN and IEEE (as well as the EU) are also concerned about ethical use of data, and ethical algorithms, from regulatory and the perspective of rights.

[1] See https://www.gov.uk/government/publications/data-ethics-framework



DATA WRANGLING (AKA FEATURE ENGINEERING)

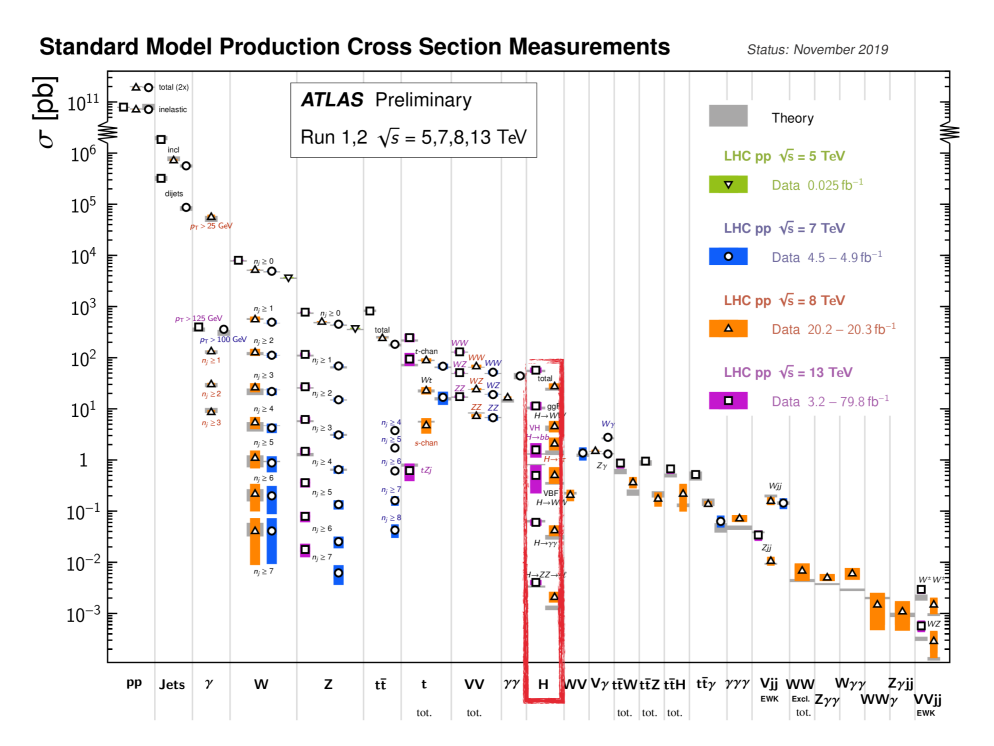
- Need to understand the data being analysed.
 - Domain context will provide most insight into getting information to highlight the solution to your problem.
 - Identify the features of interest in the data.
 - Allow you to reduce the dimensionality of the problem into as few a set of inputs as possible.
 - Most of the analysis can be done with this domain context background:
 - Deep Learning (DL) can replace the hard work of feature engineering for a resource cost and a lack of explainability and interpretability. [1]
 - "Smart Learning" (SL) is an alternative way of approaching the problem. [2]

^[2] I attribute the term Smart Learning to the use of domain knowledge and understandable AI, such as Bayesian Networks. A term that I first heard about from Norman Fenton who has a good book on the topic.

A. Bevan

- In HEP we use cut based selection to:
 - Remove pathological data (poorly calibrated or partially complete data, bad beam conditions, etc).
 - Remove obvious background examples from the data (well known SM processes that are just not interesting for study, or use as a calibration or control sample).
 - Prepare data for the "statistical analysis" that will include the use of multivariate analysis techniques and fitting.

• e.g. Higgs physics: Throw away $\sim 10^9$ recorded events for each interesting one.





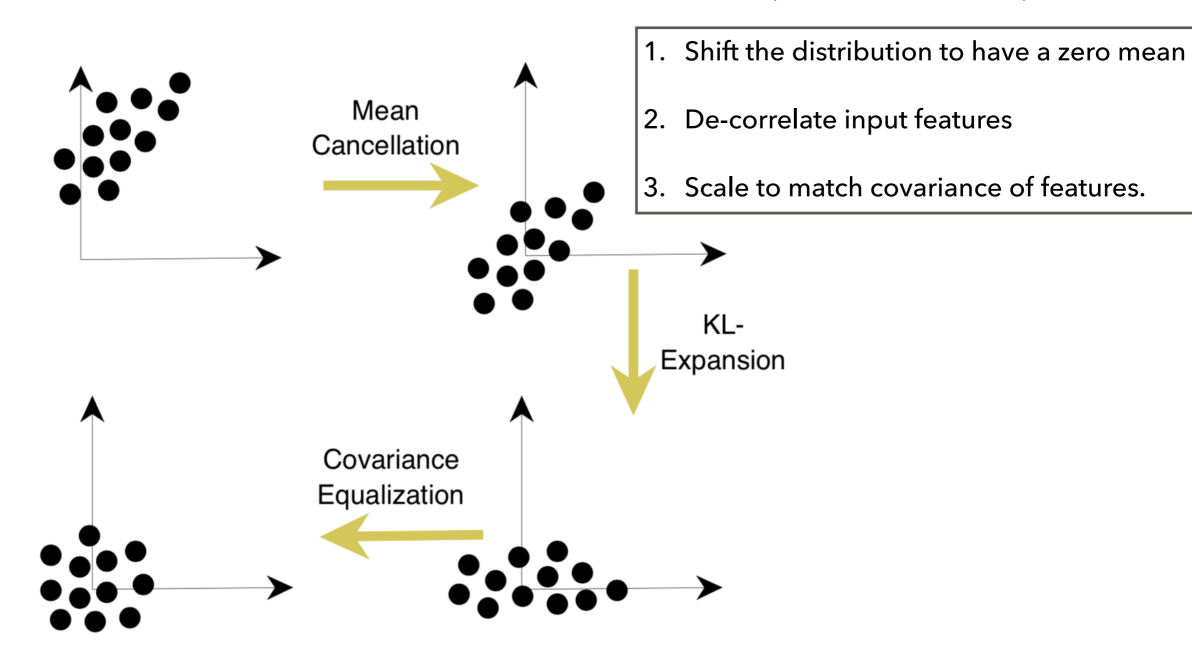
- Knowing what the task is allows one to identify the features of interest. This is the domain context knowledge.
 - If your signal is the process pp→HH+X, then:
 - Laws of physics provide insights for you to refine your list of features to train on.
 - e.g. the Higgs mass or p_T will play a role in identifying the signal, along with decay product properties (e.g. b-tag quality), etc.

- Reduce the number of dimensions of interest:
 - Trial and error with different inputs to identify what improves performance of the algorithm and what does not.
 - Linearly correlated features can be combined to reduce the number of features providing information for the algorithm to learn from.
 - Principal Component Analysis (PCA) to address this problem.
 - Some neural network configurations do that automatically for you (e.g. auto-encoders).
 - Let the algorithm learn what is important and what is not.
- Some algorithms (e.g. support vector machines) implicitly increase the dimensionality of the problem.

ACTIVATION FUNCTIONS: DATA PREPARATION

- Input features are arbitrary; whereas (for example) activation functions in neural networks work best for a standardised input domain of [-1, 1] or [0, 1].
 - We can map our input feature space onto a standardised domain that matches some range that matches that of the activation function.
 - Saves work for the optimiser in determining hyper-parameters.
 - > Standardises weights to avoid numerical inaccuracies; and set common starting weights.
 - e.g.
 - having an energy or momentum measured in units of 10¹² eV, would require weights O(10⁻¹²) to obtain an O(1) result for w_ix_i.
 - ▶ Mapping eV \mapsto TeV would translate 10^{12} eV \mapsto 1TeV, and allow for O(1) weights leading to an O(1) result for $w_i x_i$.
 - Comparing weights for features that are standardised allows the user to develop an intuition as to what the corresponding activation function will look like.

Variable transformations are also useful (can be vital).





- Data wrangling is an important part of ensuring you get the most out of applying machine learning; yet it is often not celebrated as the requirements can be very problem specific.
- If you are interested in this topic you may wish to review the data wrangling presentation and tutorials given by Dr Nick Barlow from the Alan Turing Institute at the GradNet meeting on Machine Learning and AI in January 2020:
 - Dr Barlow's talk and lecture can be found at: https://
 indico.ph.qmul.ac.uk/indico/conferenceOtherViews.py?
 view=standard&confld=543

TYPES OF LEARNING
GRID SEARCH
GRADIENT DESCENT
ADAM OPTIMISER
MISCELLANEOUS

OPTMISATION



OPTIMISATION TYPES OF LEARNING



TYPES OF LEARNING

- Consider a model y that depends on a parameter set θ , we can select a point in the parameter hyperspace $\hat{\theta}$ that has a corresponding estimator of the function \hat{y} .
- lacktriangle The heta are hyper-parameters (HPs) of the model.

Unsupervised learning:

- Infer the probability distribution $P(\hat{y})$ from the data without using labels.
- Widely used in many fields of research.

Supervised learning:

- Use training examples from labeled data sets with a known type or value, t, for each example.
- **>** Some loss function is used to compare t against model predictions \widehat{y} .
- ▶ Can be thought of as computing a conditional probability $P(t \mid \widehat{y})$.
- This is the most commonly used approach in particle physics today.



TYPES OF LEARNING: SUPERVISED LEARNING

- Define a heuristic based on some figure of merit (FOM) designed to improve the value of that metric through some iterative process.
- The FOM is called the objective function or loss function or cost function in machine learning.
- e.g. the L₂ norm loss function: this is like a χ^2 , but without the error term:

$$L_2 = \sum_{i=1}^{N_{examples}} \left[t_i - \hat{y}(\hat{\theta}) \right]^2$$

The θ are called HPs as they form a hyperspace; other variables in the optimisation process are often included in the set of HPs (e.g. learning rate for a neural network, cost for a support vector machine, tree depth for a decision tree etc.).

TYPES OF LEARNING

Batch learning

 Use a sample (batch) of training data to evaluate an estimate of the error and update weights in the optimisation.

Advantages of Batch Learning

- 1. Conditions of convergence are well understood.
- 2. Many acceleration techniques (e.g. conjugate gradient) only operate in batch learning.
- 3. Theoretical analysis of the weight dynamics and convergence rates are simpler.

Stochastic learning

Use individual training examples to evaluate an estimate of the error and update weights in the optimisation.

Advantages of Stochastic Learning

- 1. Stochastic learning is usually *much* faster than batch learning.
- 2. Stochastic learning also often results in better solutions.
- 3. Stochastic learning can be used for tracking changes.



- •THIS IS A SIMPLE OPTIMISATION ALGORITHM THAT CAN BE USED WITH NO PROGRAMMING EXPERIENCE.
- •ONE CAN IMPLEMENT A 1 OR 2 DIMENSIONAL GRID SEARCH USING AN EXCEL SPREADSHEET, AND FROM INSPECTION OF THE TABULATED RESULTS YOU CAN OPTIMISE SIMPLE PROBLEMS.
- •THIS IS A BRUTE FORCE OPTIMISATION APPROACH, AND IT IS USED WIDELY IN CERTAIN APPLICATIONS ACROSS A NUMBER OF RESEARCH FIELDS.

OPTIMISATION GRID SEARCH

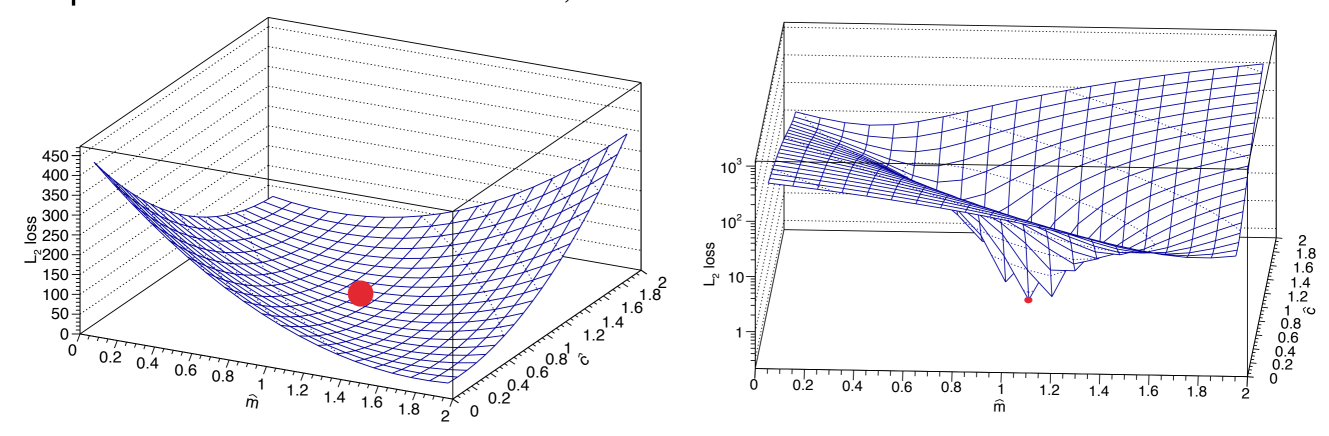


GRID SEARCH

- Scanning through the hyperspace for each $\hat{\theta}_i$ allows us to compute L₂, and the minimum value obtained is the "best fit" or optimal estimate of the parameters θ .
 - Simple heuristic to implement you can do this in Excel for 1 or 2D problems. ✓
 - ▶ Easy to understand. ✓
 - Expensive to compute: scanning n points in a dimension requires n^M computations for $M = dim(\theta)$.
- Heuristic suffers from the curse of dimensionality.

GRID SEARCH

• e.g. Consider an L2 loss function optimisation of the HPs for y = mx + c, i.e. optimise m and c. Here m = 1.0, c = 1.0



- ▶ The contours of the loss function show a minimum.
- The optimal value is selected from a discrete grid of points, only get an exact result if the grid maps onto the problem perfectly (as in this example).



GRID SEARCH

- e.g. The libsvm^[1] package uses a HP grid search for optimisation for the Support Vector Machine algorithm; where the cost C and Γ hyper-parameters need to be optimised.
 - The grid search is done efficiently by adapting to whatever step is sensible, in this case the algorithm has Γ as the parameter of an exponential (the radial basis function); and so a linear search in C and a logarithmic search in Γ is appropriate to sample the parameter hyperspace effectively.

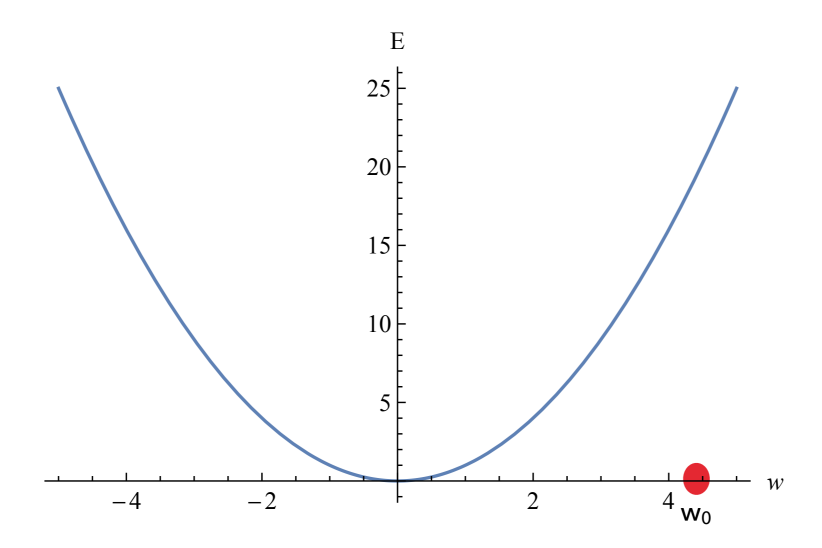
[1] Chih-Chung Chang and Chih-Jen Lin, LIBSVM: a library for support vector machines. ACM Transactions on Intelligent Systems and Technology, 2:27:1--27:27, 2011. Software available at http://www.csie.ntu.edu.tw/~cjlin/libsvm.



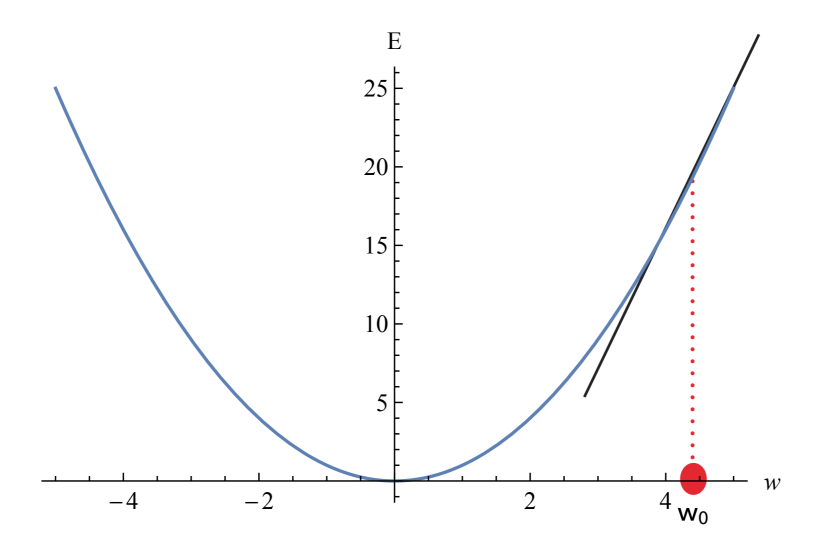
OPTIMISATION GRADIENT DESCENT



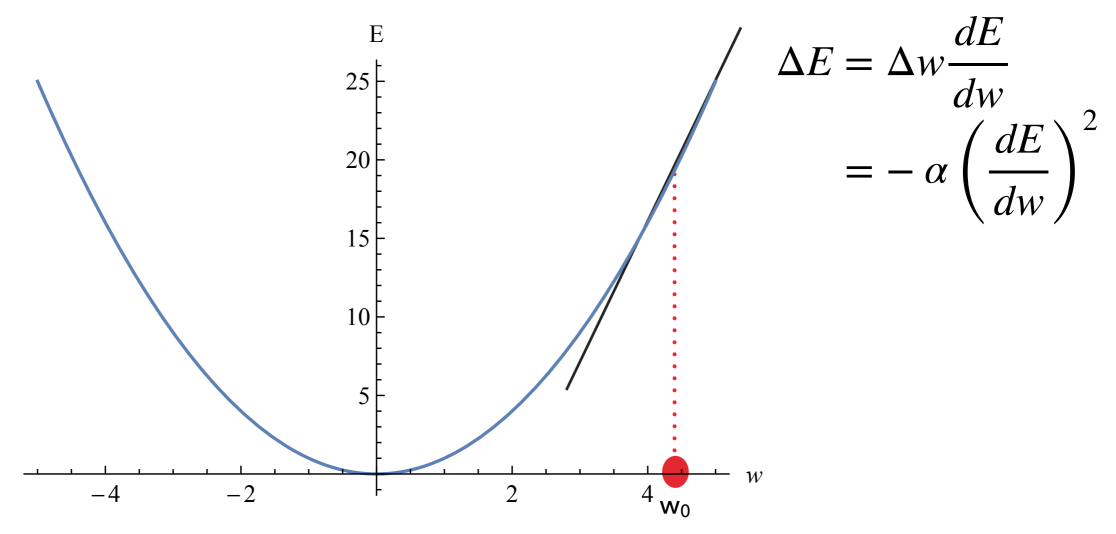
 \blacktriangleright Guess an initial value for the weight parameter: w_0 .



Estimate the gradient at that point (tangent to the curve)



• Compute Δw such that ΔE is negative (to move toward the minimum)

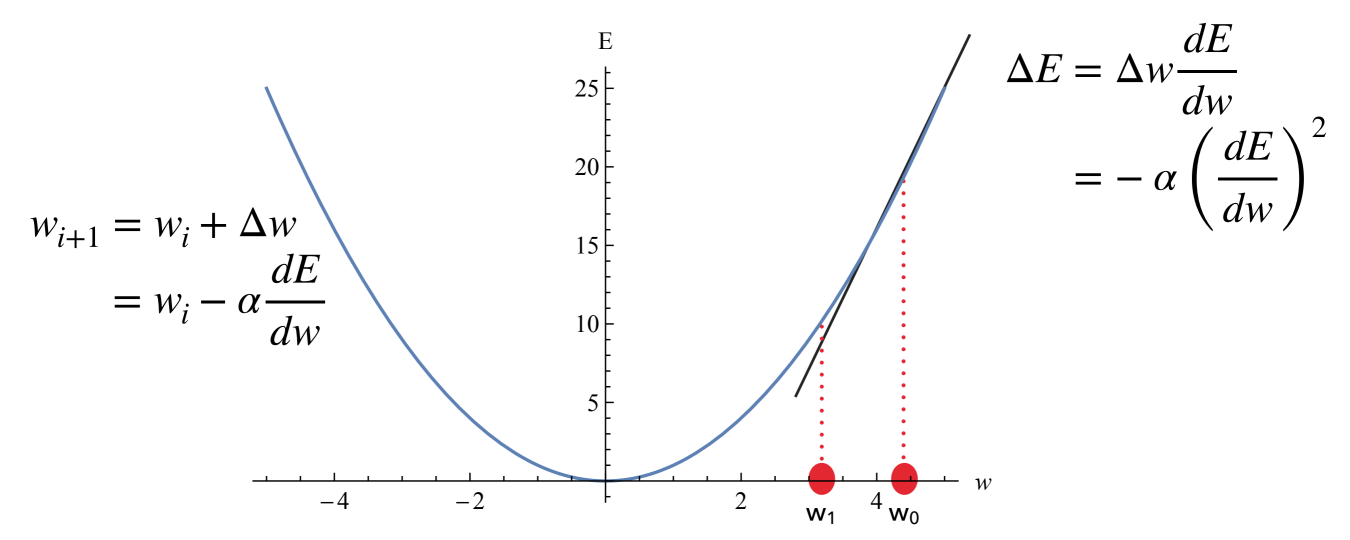


 α is the learning rate: a small positive number

Choose
$$\Delta w = -\, \alpha \frac{dE}{dw}$$
 to ensure $\Delta {\rm E}$ is always negative.



▶ Compute a new weight value: $w_1 = w_0 + \Delta w$

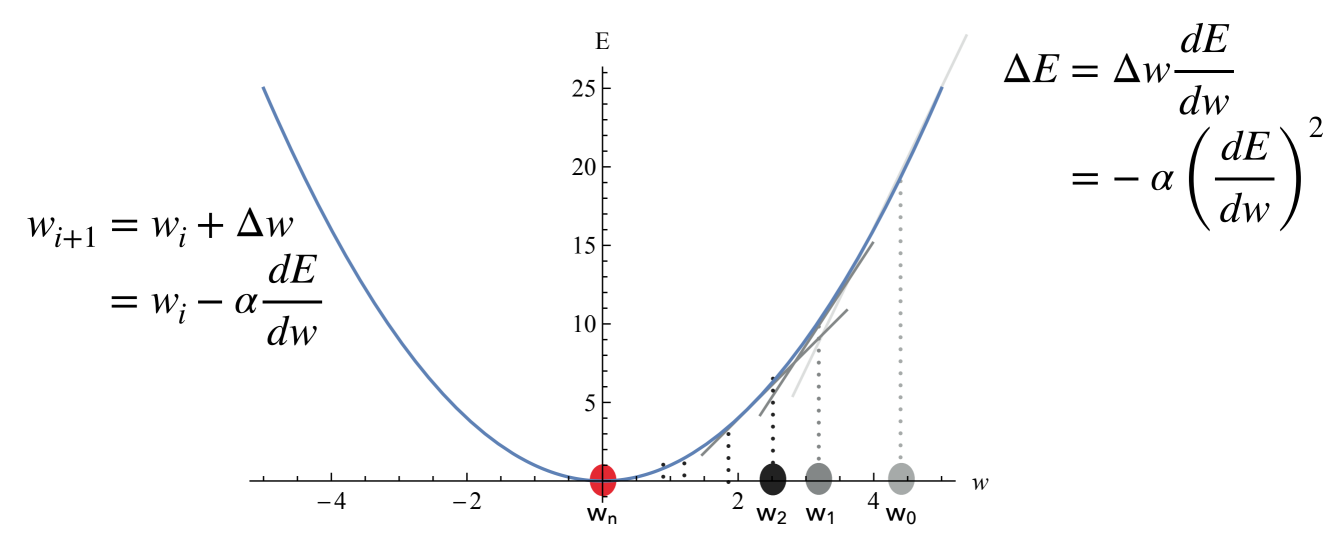


 α is the learning rate: a small positive number

Choose
$$\Delta w = -\, \alpha \frac{dE}{dw}$$
 to ensure $\Delta {\rm E}$ is always negative.



Repeat until some convergence criteria is satisfied.

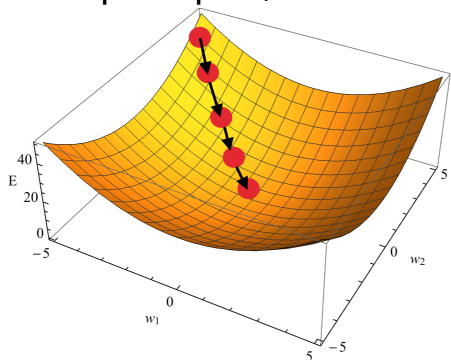


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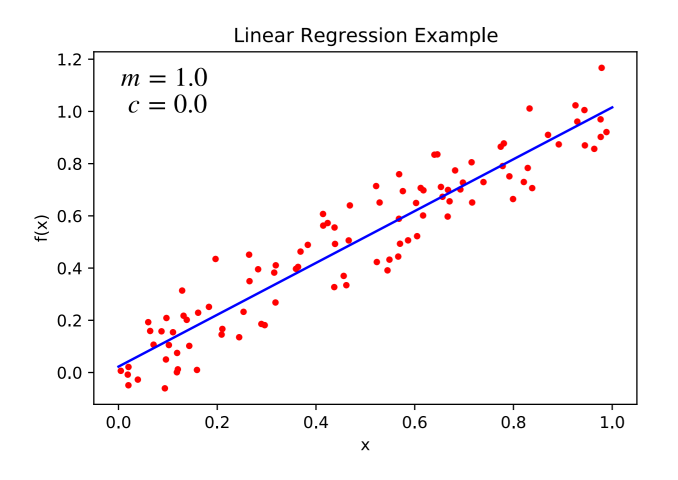
We can extend this from a one parameter optimisation to a 2 parameter one, and follow the same principles, now in 2D.

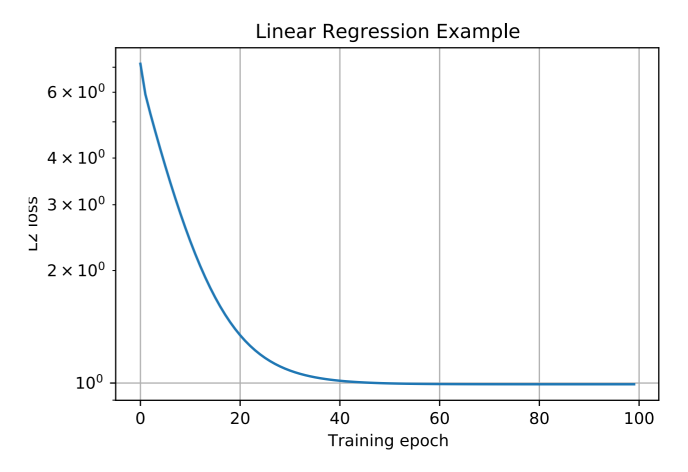


- The successive points w_i+1 can be visualised a bit like a ball rolling down a concave hill into the region of the minimum.
- In general update weights such that $\Delta E = \Delta w \nabla E = -\alpha \nabla^2 E$
- and $w_{i+1} = w_i \alpha \nabla E$.

GRADIENT DESCENT: EXAMPLE

• Returning to the y = mx + c example, we can optimise this using the Gradient Descent algorithm.





Use N=100 and $\alpha = 0.005$ with the TensorFlow GradientDescent optimiser to obtain:

$$\widehat{m} = 0.9928...$$

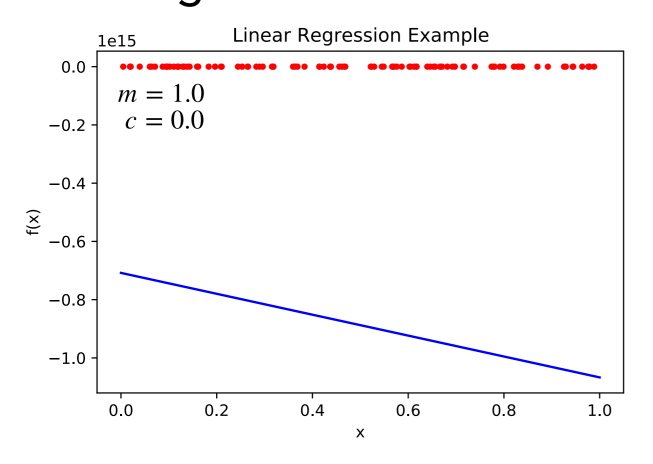
$$\hat{c} = 0.0226...$$

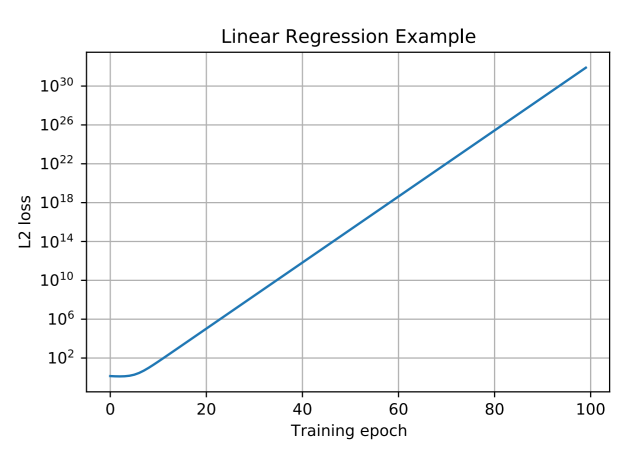
Implementation: TensorFlow 2.2 (using V1 compatibility) Jupyter Notebook



GRADIENT DESCENT: EXAMPLE

• Returning to the y = mx + c example, we can optimise this using the Gradient Descent algorithm.





Use N=100 and $\alpha = 0.01$ with the TensorFlow GradientDescent optimiser to obtain:

$$\widehat{m} = -3.5 \times 10^{14}$$

 $\widehat{c} = -7.08 \times 10^{14}$

The minimiser fails; too large a learning rate is being used.

Implementation: TensorFlow 2.2 (using V1 compatibility)
Jupyter Notebook



GRADIENT DESCENT: REFLECTION

- The examples shown illustrate problems with parabolic minima.
- With selection of an appropriate learning rate, α , to fix the step size, we can guarantee convergence to a sensible minimum in some number of steps.
- Translating the distribution to a fixed scale, then we can predict how many steps it will take to converge to the minimum from some distance away from it for a given α .
- If the problem hyperspace is not parabolic, this becomes more complicated, and there is no guarantee that we converge to the minimum.
- Modern machine learning algorithms use more refined variants on this method.

OPTIMISATION ADAM OPTIMISER



(ADAM: ADAptive Moment estimation based on gradient descent)

- This is a stochastic gradient descent algorithm.
- **Consider a model** $f(\theta)$ that is differentiable with respect to the HPs θ so that:
 - t is the training epoch.
 - the gradient $g_t = \nabla f_t(\theta_{t-1})$ can be computed.
 - $m_t(v_t)$ are biased values of the first (second) moment.
 - $\widehat{m}_{t}(\widehat{v}_{t})$ are bias corrected estimator of the moments.
 - Some initial guess for the HP is taken: θ_0 , and the HPs for a given epoch are denoted by θ_r .
 - ightharpoonup lpha is the step size (i.e. learning rate).
 - β_1 and β_2 are exponential decay rates of moments.

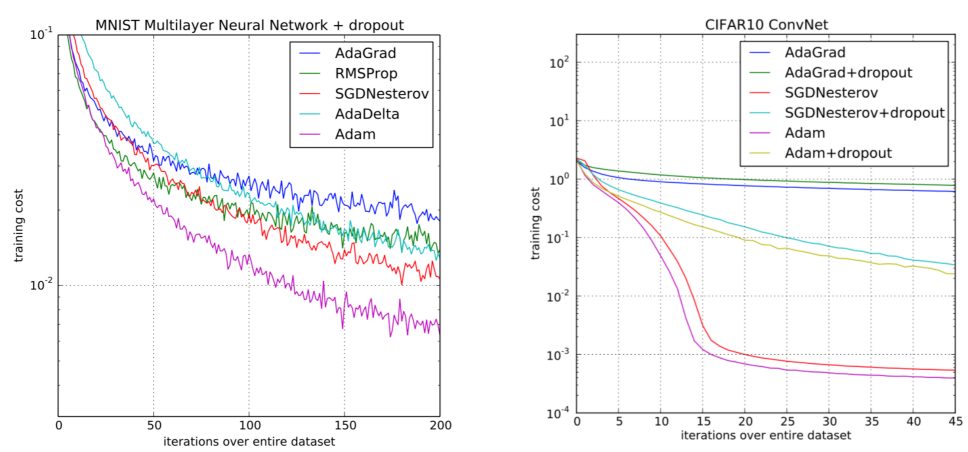
(ADAM: ADAptive Moment estimation based on gradient descent)

Algorithm 1: Adam, our proposed algorithm for stochastic optimization. See section 2 for details, and for a slightly more efficient (but less clear) order of computation. g_t^2 indicates the elementwise square $g_t \odot g_t$. Good default settings for the tested machine learning problems are $\alpha = 0.001$, $\beta_1 = 0.9$, $\beta_2 = 0.999$ and $\epsilon = 10^{-8}$. All operations on vectors are element-wise. With β_1^t and β_2^t we denote β_1 and β_2 to the power t.

```
Require: \alpha: Stepsize
Require: \beta_1, \beta_2 \in [0, 1): Exponential decay rates for the moment estimates
Require: f(\theta): Stochastic objective function with parameters \theta
Require: \theta_0: Initial parameter vector
   m_0 \leftarrow 0 (Initialize 1<sup>st</sup> moment vector)
   v_0 \leftarrow 0 (Initialize 2<sup>nd</sup> moment vector)
   t \leftarrow 0 (Initialize timestep)
   while \theta_t not converged do
       t \leftarrow t + 1
       g_t \leftarrow \nabla_{\theta} f_t(\theta_{t-1}) (Get gradients w.r.t. stochastic objective at timestep t)
       m_t \leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t (Update biased first moment estimate) v_t \leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2 (Update biased second raw moment estimate)
       \widehat{m}_t \leftarrow m_t/(1-\beta_1^t) (Compute bias-corrected first moment estimate)
       \widehat{v}_t \leftarrow v_t/(1-\beta_2^t) (Compute bias-corrected second raw moment estimate)
       \theta_t \leftarrow \theta_{t-1} - \alpha \cdot \widehat{m}_t / (\sqrt{\widehat{v}_t} + \epsilon) (Update parameters)
   end while
   return \theta_t (Resulting parameters)
```

(ADAM: ADAptive Moment estimation based on gradient descent)

Benchmarking performance using MNIST and CFAR10 data indicates that Adam with dropout minimises the loss function compared with other optimisers tested.



 Faster drop off in loss, and lower overall loss obtained vs other algorithms benchmarked at the time.

(ADAM: ADAptive Moment estimation based on gradient descent)

• Benchmarking performance using MNIST and CFAR10 data indicates that

benchmark data sets in CS (see appendix). MNIST is a set of handwritten numbers 0 through 9; CFAR10(0) is an image classification data set (cars, boats etc).



 Faster grop oπ in ioss, and iower overall loss obtained vs other algorithms benchmarked at the time.

MULTIPLE MINIMA MORE ON LOSS FUNCTIONS

OPTIMISATION MISCELLANEOUS



GRADIENT DESCENT: MULTIPLE MINIMA

 Often more complication hyperspace optimisation problems are encountered, where there are multiple

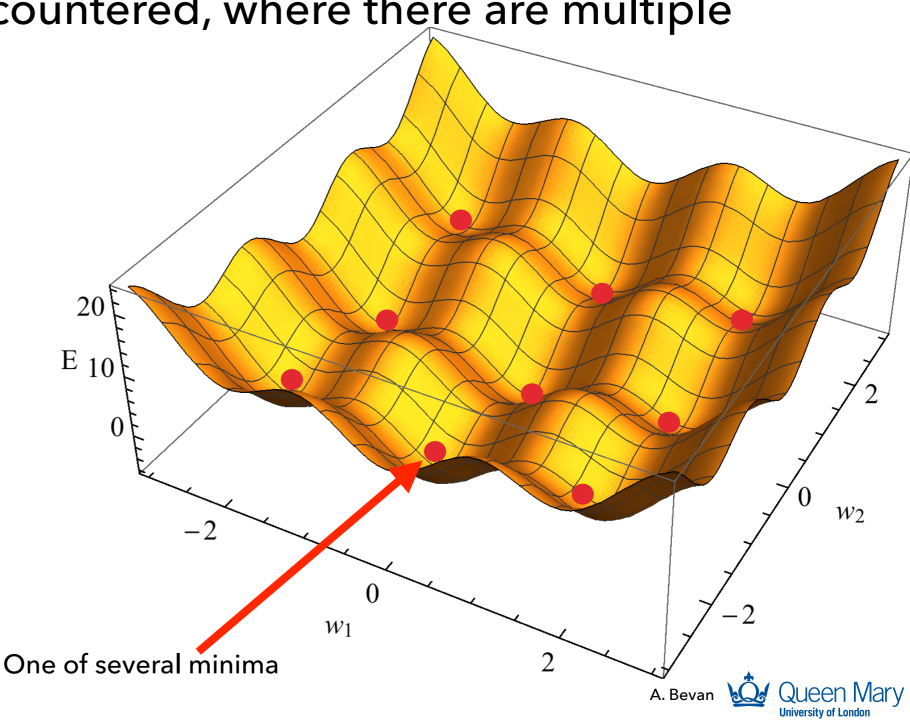
minima.

The gradient descent minimisation algorithm is based on the assumption that there is a single minimum to be found.

In reality there are often multiple minima.

Sometimes the minima are degenerate, or near degenerate.

How do we know we have converged on the global minimum?



GRADIENT DESCENT: MULTIPLE MINIMA

 Often more complication hyperspace optimisation problems are encountered, where there are multiple

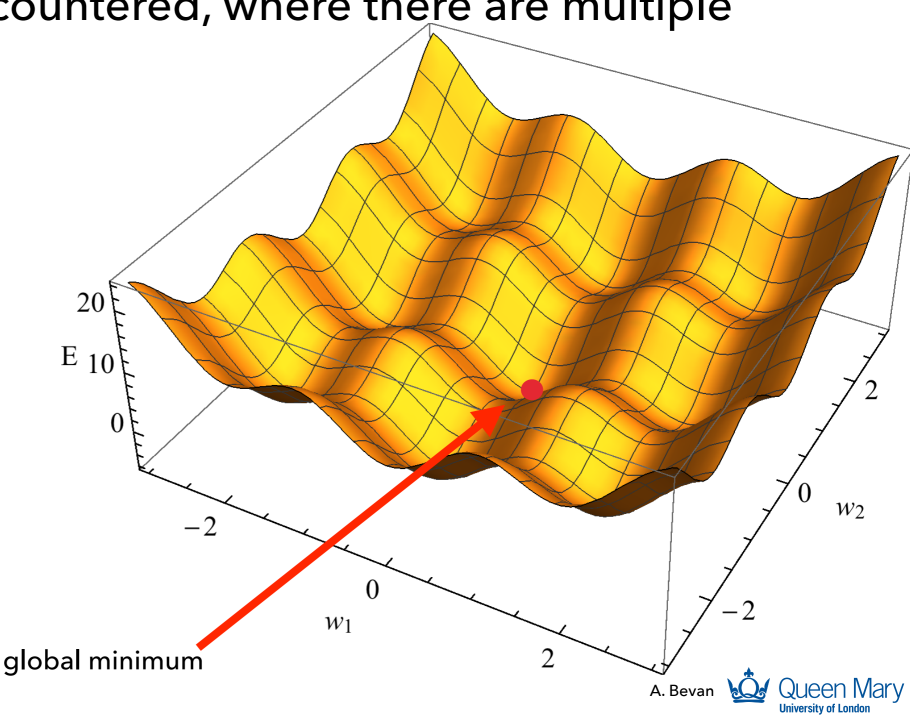
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How do we know we have converged on the global minimum?



OPTIMISATION: LOSS FUNCTIONS

- ▶ There are many types of loss function other than the L₂ loss
 - ▶ L₁ norm loss:

$$L_1 = \sum_{i=1}^{N_{examples}} \left| t_i - \hat{y}(\hat{\theta}) \right|$$

The mean square error (MSE) loss function:

$$L = \frac{1}{N_{examples}} L_2 = \frac{1}{N_{examples}} \sum_{i=1}^{N_{examples}} \left[t_i - \hat{y}(\hat{\theta}) \right]^2$$

Cross entropy

$$L = -\sum_{i=1}^{N_{examples}} \widehat{y}(x_i) \ln t_i$$

The cross entropy can be thought of as the negative log likelihood function for the data t_i under the model \hat{y}

SUPERVISED LEARNING OVERTRAINING

WEIGHT REGULARISATION CROSS VALIDATION

DROPOUT

TRAINING

TRAINING SUPERVISED LEARNING



SUPERVISED LEARNING

For supervised learning the loss function depends on both model parameters and a known target value t_i for a given example.

e.g. the L₂ loss:
$$L_2 = \sum_{i=1}^{N_{examples}} \left[t_i - \hat{y}(\hat{\theta}) \right]^2$$
.

- This requires (at a minimum) a sample of data with the input feature space of interest, and for each example in the data, the known target output value.
- ▶ The loss function is optimised using the data, to obtain a model.
- Unlike fitting however we don't care about the uncertainty on the model parameter estimates, only on the nominal values obtained, $\hat{\theta}$.
- Due to the complexity of models, it is generally not possible to understand if the optimisation converged to a sensible solution by inspecting marginalised projections of the model on the data.

SUPERVISED LEARNING

• e.g. the <u>Kaggle Higgs</u> data sample:

EventId, DER_mass_MMC, DER_mass_transverse_met_lep, ..., Weight Label, KaggleSet, KaggleWeight

Features (not all to be used for training)

Known example type

- The KaggleSet column in the CVS file for this data allows the user to understand if this is to be used for training or testing, or some other means (e.g. Kaggle Score Board evaluation).
- The Label column, is the known label; this defines the t_i used in the loss function.
- I use this challenge data as an assignment for undergraduate lectures (pdf, zip). N.B. the zip file is 189Mb.

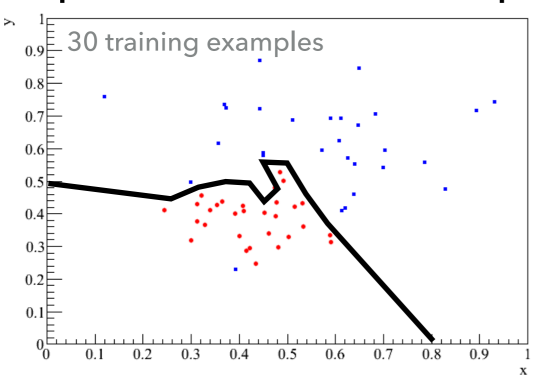
WEIGHT REGULARISATION CROSS VALIDATION

TRAINIG OVERTRAINIG



A model is over trained if the HPs that have been determined are tuned to the statistical fluctuations in the data set.

Simple illustration of the problem:



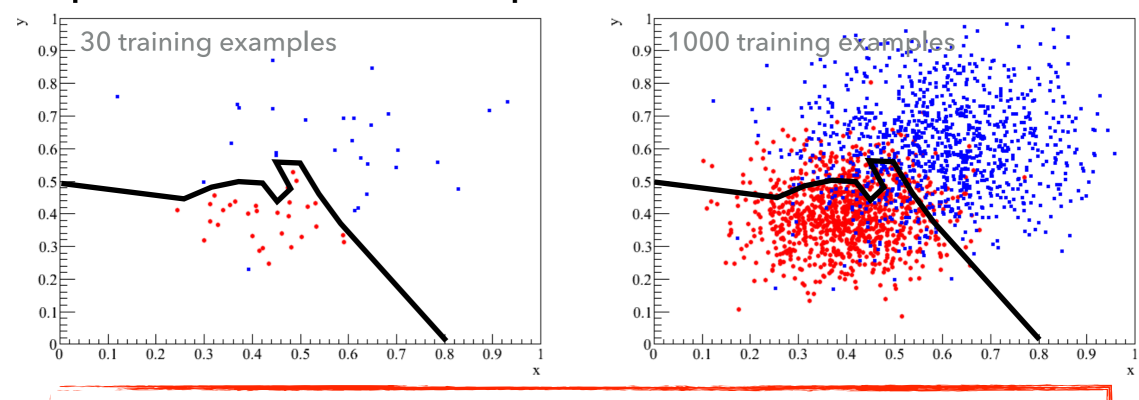
The decision boundary selected here does a good job of separating the red and blue dots.

Boundaries like this can be obtained by training models on limited data samples. The accuracies can be impressive.

But would the performance be as good with a new, or a larger data sample?

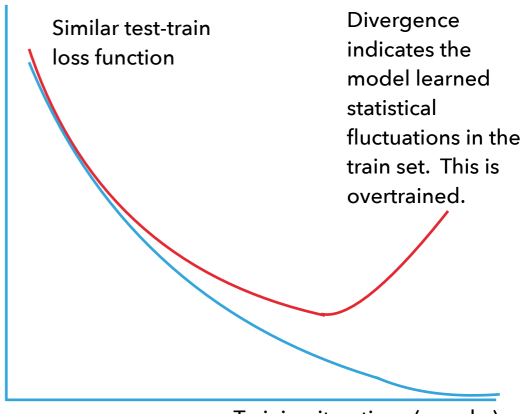
A model is over trained if the HPs that have been determined are tuned to the statistical fluctuations in the data set.

Simple illustration of the problem:



Increasing to 1000 training examples we can see the boundary doesn't do as well. This illustrates the kind of problem encountered when we overfit HPs of a model.

- One way to avoid tuning to statistical fluctuations in the data is to impose a training convergence criteria based on a data sample independent from the training set: a test sample.
 - Use the loss evaluated for the training and test samples to check to see if the HPs are over trained.
 - If both samples have similar loss then the model response function is similar on two statistically independent samples.
 - Note: If the samples are large enough then one could reasonably assume that the response function would then be general when applied to an unseen data sample.
 - "large enough" is a model and problem dependent constraint.
- Some also recommend a third validation set of data in order to provide a completely independent verification of algorithm performance.



Loss

Training iterations (epochs)

- Training convergence criteria that could be used:
 - ▶ Terminate training after N_{epochs}
 - Loss comparison:
 - Evaluate the performance on the training and test sets.
 - Compare the two and place some threshold on the difference $\Delta_{LOSS} < \delta_{LOSS}$
 - Terminate the training when the gradient of the loss function with respect to the weights is below some threshold.
 - > Terminate the training when the Δ_{LOSS} starts to increase for the test sample.

OVERTRAINING: WEIGHT REGULARISATION

- Weight regularisation involves adding a penalty term to the loss function used to optimise the HPs of a network.
- This term is based on the sum of the weights w_i in the network and takes the form:

$$\lambda \sum_{i=\forall weights} w_i$$

- The rationale is to add an additional cost term to the optimisation coming from the complexity of the network.
- The performance of the network will vary as a function of λ.
- To optimise a network using weight regularisation it will have to be trained a number of times in order to identify the value corresponding to the min(cost) from the set of trained solutions.

OVERTRAINING: WEIGHT REGULARISATION

For example we can consider extending an MSE loss function to allow for weight regularisation. The MSE loss is given by:

$$\varepsilon = \frac{1}{N} \sum_{i=1}^{N} (y_i - t_i)^2$$

To allow for regularisation we add the sum of weights term:

$$\varepsilon = \frac{1}{N} \sum_{i=1}^{N} (y_i - t_i)^2 + \lambda \sum_{i=\forall, weights} w_i$$

This is a simple modification to make to the NN training process.



OVERTRAINING: CROSS VALIDATION

- A well trained model will provide robust predictions, irrespective of the examples presented to it.
 - The variance of model predictions will be small.
 - The model predictions may be systematically biased independently of this.
- One can divide the training set up into k-folds, and then perform k trainings; each one leaving a single fold out.
- The amount of data used in a fold will depend (generally the more data the better.
- The ensemble of predictions indicates the variance on the model output.



OVERTRAINING: CROSS VALIDATION

- The use of cross validation to determine the spread of model predictions, and any systematic bias on prediction can be useful.
- e.g. in a physics context: Consider a new particle search at the Large Hadron Collider.
 - Using an overtrained model to suppress background and enhance signal will result in a lack of experimenter understanding as to how the expected and observed limits on the new particle relate to each other.
 - This could result in a false discovery of new physics.
 - It could result in missing out on a discovery of new physics.
 - Some people say it is not wrong to use an over trained model I argue strongly that it is not scientifically correct to use an over trained model, unless the variance on that model, and hence the implications are well understood.

TRAINING DROPOUT

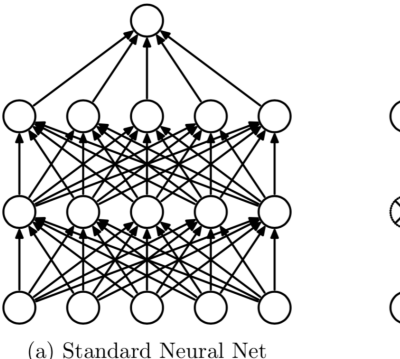


DROPOUT

- A problem with deep networks is the number of HPs that need to be determined.
- This leads to a requirement for very large data sets to avoid overfitting (or fine-tuning).
- HPs can also learn to "co-adapt" in the training process.
 - co-adapt means that as one parameter is changed, another in the network can be modified in a correlated way to offset that change.
 - This kind of behaviour intentionally exists in some algorithms (Sequential Minimal Optimisation for Support Vector machines, where pairs of HPs are changed to conserve an overall zero sum); but is generally unwelcome behaviour.

OVER FITTING: DROPOUT

A pragmatic way to mitigate these issues is to intentionally compromise the model randomly in different epochs of the training by removing units from the network.



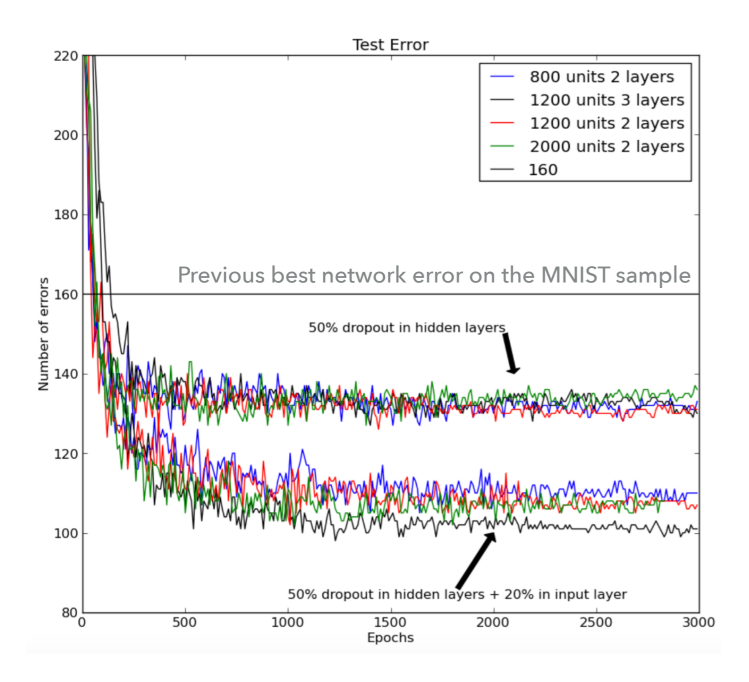
Dropout is used only during training.

The full model is used for making predictions.

ndard Neural Net (b) After applying dropout.

- That way the whole model will be effectively trained on a sub-sample of the data with the aim of limiting the ability to learn statistical fluctuations in the data, and mitigating the co-adaption issue.
- This does not remove the possibility that a model is overtrained, as with the previous discussion HP generalisation is promoted by using this method.

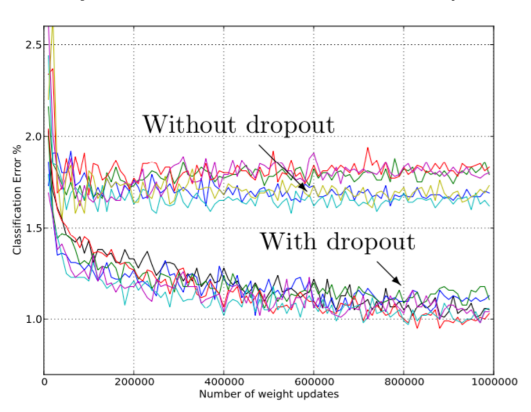
DROPOUT

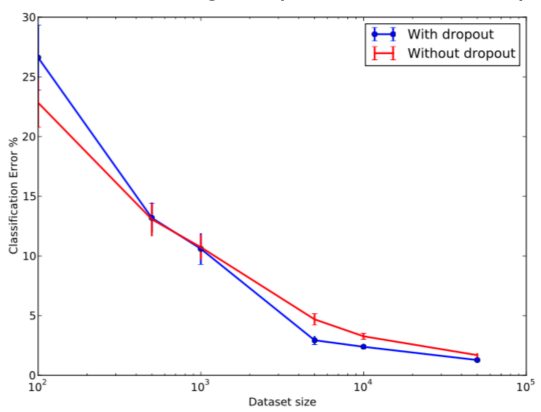


- Example from Hinton et al. for an MNIST sample.
- Using 50% drop out on the hidden layers gives an improvement over the previous best network architecture.
- Adding 20% drop out in the input layer provides further improvement in error.

OVER FITTING: DROPOUT

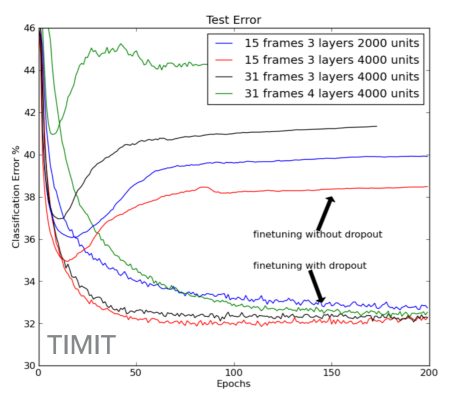
A variety of architectures has been explored with different training samples for this technique,

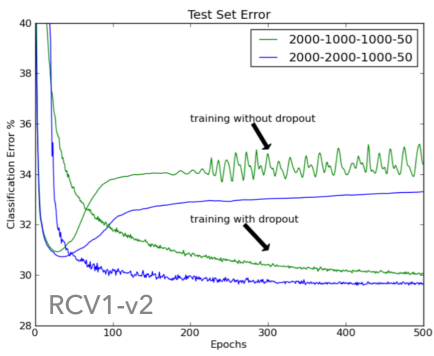




- Dropout can be detrimental for small training samples, however in general the results show that dropout is beneficial.
- For deep networks or typical training samples O(500) examples or more this technique is expected to be beneficial.
- For algorithms with very large training data sets, the benefits are less clear.

DROPOUT





- Example from Hinton et al. for a voice recognition sample of data (TIMIT, speech recognition data).
- Illustrates the classification error as a function of epoch with and without dropout.
- Similar results were obtained by Hinton and his team with the Reuters newsfeed data set (RCV1-v2).

DECISION TREES
BOOSTING

ADABOOST,M1

RANDOM FORESTS

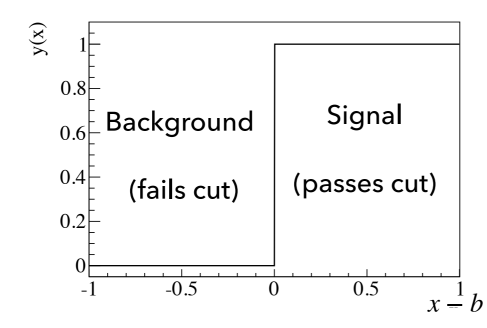
DECISION TREES



- The cut based [see Data Wrangling] and linear discriminant analysis methods are useful but limited.
- The underlying concepts of applying Heaviside function constraints on data selection and on the use of a decision boundary definition (a plane in hyperspace) of the form of the dot product $\alpha^T x + \beta$ can be applied in more complicated algorithms.
- Here we consider extension to the concept of rectangular cuts to decision trees as a machine learning algorithm.
 - We will have to introduce the concepts of classification and regression; and methods to mitigate mis-classification of data.
 - The issue of overtraining is something we discussed earlier regarding optimisation.



- Consider a data sample with an N dimensional input feature space X.
- X can be populated by examples from two or more different species of event (also called classes, categories or types).
- Consider the two types and call them signal and background, respectively*.
- We can use a Heaviside function to divide the data into two parts:
 - We can use this to distinguish between regions populated signal and background:



For an arbitrary cut position in x we can modify the Heaviside function

$$H(x) = \frac{1}{2}(1 + sign(x - b))$$

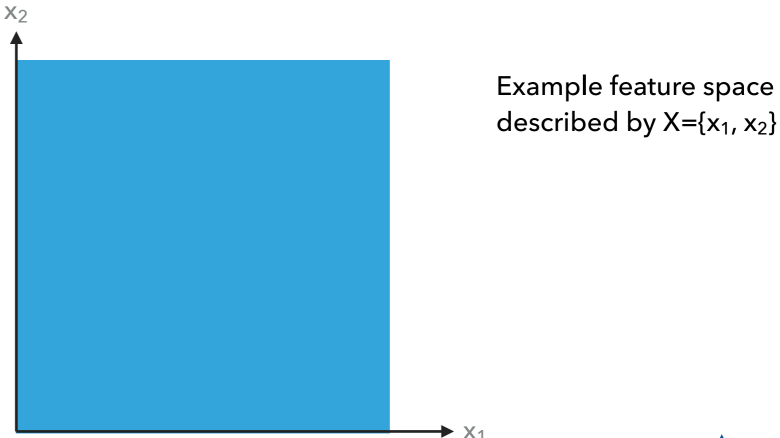
where b is the offset (bias) from zero.



- Decision trees divide the data feature space into a set of hypercubes that are classified as signal (+1) or background (-1) like.
 - Each region can be fitted with a constant to represent the data in that region.
 - We can recursively continue to sub-divide the data until some minimum number of examples are left in each sub-division.

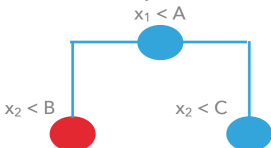
Can describe the data as the root node.

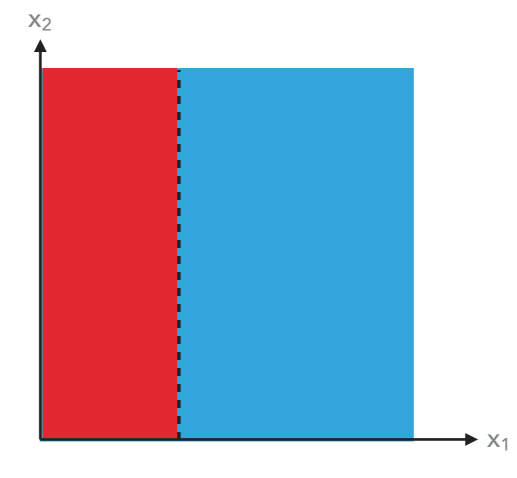




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The data get divided into two partitions.

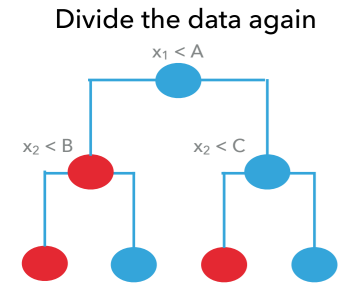


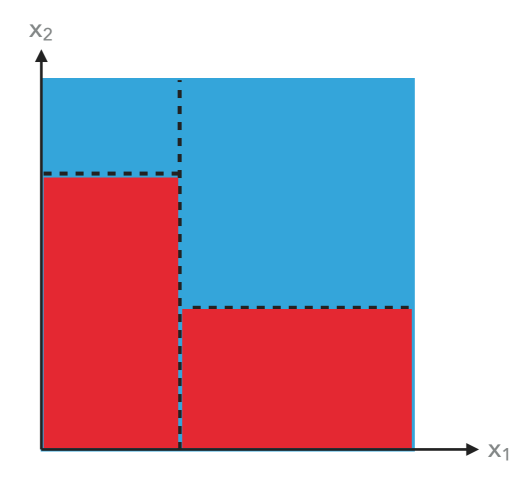


Cut on the feature space to separate the data into two different regions.



- Decision trees divide the data feature space into a set of hypercubes that are classified as signal (+1) or background (-1) like.
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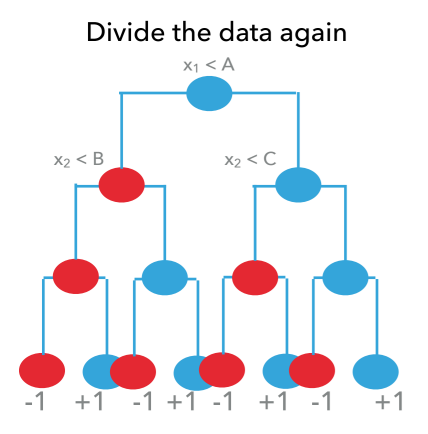


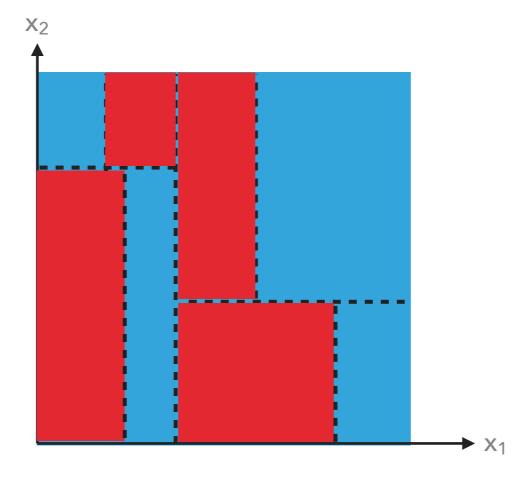


The feature space gets further sub-divided.



- Decision trees divide the data feature space into a set of hypercubes that are classified as signal (+1) or background (-1) like.
 - Each region can be fitted with a constant to represent the data in that region.
 - We can recursively continue to sub-divide the data until some minimum number of examples are left in each sub-division.

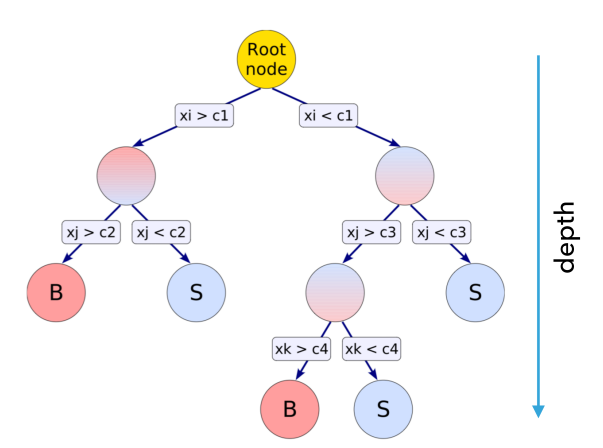




The feature space gets further sub-divided (again).



- The set of rectangular cuts applied to the data allow us to build a tree from the root note.
- We can impose limits on:
 - Tree depth (how many divisions are performed).
 - Node size (how many examples per partition).
- Trees can be extended to more than 2 categories.
- They lend themselves to classifying examples or adapted to make a quantitative prediction (regression)



The decision tree output for a classification problem is

$$G(x) = +1 \text{ or } -1$$



- Decision trees are "weak learners", as they can take input features that only weakly separate types of example and combine those features to increase the separation.
- A single tree is susceptible to overtraining, and there are various methods of reducing this; including limiting the complexity of a tree, or the limiting the minimum number of examples in each node.
- The decision tree can be extended to an oblique decision tree, in which a linear combination of the features (instead of a single feature) is used to classify examples; so the Heaviside function cut becomes a hyperplane cut.

BOOSTING

- If a training example has been mis-classified in a training epoch, then the weight of that event can be increased for the next training epoch; so that the cost of mis-classification increases.
- The underlying aim is to take a weak learner and try and boost this into becoming a strong learner.
 - This example re-weighting technique is called boosting.
 - There are several re-weighting methods commonly used; here we discuss:
 - AdaBoost.M1 (popular variant of the Adaptive boosting method)
- Boosted Decision Trees are known as BDTs

- i is the ith example out of a data set with N examples.
- m is the mth training out of an ensemble of M learners to be trained.
- Step 1:
 - Assign event weights of $w_i = 1/N$ to all of the N examples.

- i is the ith example out of a data set with N examples.
- m is the mth training out of an ensemble of M learners to be trained.
- Step 1:
 - Assign event weights of $w_i = 1/N$ to all of the N examples.
- Step 2: for m=1 through M
 - Train the weak learner (in our case this is a BDT): $G_m(x)$.
 - Compute the error rate \mathcal{E}_m .
 - Calculate the boost factor $\beta_m = \frac{\varepsilon_m}{1 \varepsilon_m}$.
 - Update weights for misclassified examples $w_i \mapsto w_i e^{\ln(1/\beta_m)}$.

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 - Calculate the boost factor $\beta_m = \frac{\varepsilon_m}{1 c}$.
 - Update weights for misclassified examples $w_i \mapsto w_i e^{\ln(1/\beta_m)}$.
- Step 3:
 - Return the weighted committee: a combination of the M trees that have been

learned from the data:
$$G(x) = sign\left(\sum_{m=1}^{M} ln\left[\frac{1-\varepsilon_m}{\varepsilon_m}\right]G_m(x)\right)$$

▶ m=1

INITIAL TRAINING SAMPLE

▶ m=2

WEIGHTED SAMPLE

▶ m=3

WEIGHTED SAMPLE

 $-G_1(x)$

 $-G_3(x)$

 \bullet $G_M(x)$

The $G_m(x)$ are individual weak learners; each is derived from a training using the data.

The m=1 training uses the original data; all subsequent trainings use reweighted data.

m=M

WEIGHTED SAMPLE

$$G(x) = sign\left(\sum_{m=1}^{M} ln\left[\frac{1-\varepsilon_m}{\varepsilon_m}\right]G_m(x)\right)$$

The final classifier output is formed from a committee that is a weighted majority vote algorithm.

RANDOM FORESTS

- Random Forests are constructed from an ensemble of individual trees.
 - Each tree in the ensemble uses a randomly selected subset of the feature space, and the minimum node size is usually set to 1, so the classifier prediction is almost always accurate.
- The mode (classification) or mean (regression) of the ensemble is the output of the Random Forest.
- The probability that an example x_i is assigned to a given class c, is given by

$$P(c | x_i) = \frac{P(c | x_i)}{\sum_{l=1}^{n} P(c_l | x_i)}$$

• and the output score $g_c(x_i)$ is given by the aggregate over t trees in the forest:

$$g_c(x_i) = \frac{1}{t} \sum_{j=1}^t P_j(c \mid x_i)$$

The classification of x_i is simply the class c that maximises $g_c(x_i)$.

