Deep-Learning Do-It-Yourself
What is machine learning? What is data science?

The process of learning from data?
What is machine learning? What is data science?

Not so clear...
Simpler question: what do machine learning people do?
Supervised Classification & Regression

Given a dataset with label, find a function that can assign labels to new unlabelled data

Labeled data: \[
\{ \vec{x}_1, \vec{x}_2, \vec{x}_3, \ldots, \vec{x}_n \} \quad \vec{x} \in \mathbb{R}^d \\
\{ y_1, y_2, y_3, \ldots, y_n \} \quad y \in \mathbb{R} \quad \text{or} \quad y \in \mathbb{N}
\]

Goal: Find a function \( f_W(\vec{x}) \) that outputs the right class/value for an object \( \vec{x} \)
Cats vs dogs classification

Sample of cats & dogs images from Kaggle Dataset

0 = dog
1 = cat
MNIST dataset classification
Regression: predicting house price

[Graph showing the relationship between house size (m²) and house price (£). The graph includes a line of best fit and historic data points.]

- House Price (£) on the y-axis.
- House Size (m²) on the x-axis.
- Predicted House Price at the intersection of the line of best fit and the orange line from the x-axis.
- Historic Data points scattered around the line of best fit.
Medical applications for regression

Breast Cancer Wisconsin (Prognostic) Data Set

Abstract: Prognostic Wisconsin Breast Cancer Database

<table>
<thead>
<tr>
<th>Data Set Characteristics:</th>
<th>Multivariate</th>
<th>Number of Instances:</th>
<th>198</th>
<th>Area:</th>
<th>Life</th>
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<tr>
<td>Attribute Characteristics:</td>
<td>Real</td>
<td>Number of Attributes:</td>
<td>34</td>
<td>Date Donated:</td>
<td>1995-12-01</td>
</tr>
<tr>
<td>Associated Tasks:</td>
<td>Classification, Regression</td>
<td>Missing Values?:</td>
<td>Yes</td>
<td>Number of Web Hits:</td>
<td>124725</td>
</tr>
</tbody>
</table>

4-33) Ten real-valued features are computed for each cell nucleus:

a) radius (mean of distances from center to points on the perimeter)
b) texture (standard deviation of gray-scale values)
c) perimeter
d) area
e) smoothness (local variation in radius lengths)
f) compactness (perimeter^2 / area - 1.0)
g) concavity (severity of concave portions of the contour)
h) concave points (number of concave portions of the contour)
i) symmetry
j) fractal dimension ("coastline approximation" - 1)

Several of the papers listed above contain detailed descriptions of how these features are computed.

The mean, standard error, and "worst" or largest (mean of the three largest values) of these features were computed for each image, resulting in 30 features. For instance, field 4 is Mean Radius, field 14 is Radius SE, field 24 is Worst Radius.

Values for features 4-33 are recoded with four significant digits.

34) Tumor size - diameter of the excised tumor in centimeters
35) Lymph node status - number of positive axillary lymph nodes observed at time of surgery

8. Missing attribute values:
   Lymph node status is missing in 4 cases.

2) Predicting Time To Recur (field 3 in recurrent records)
   - Estimated mean error 13.9 months using Recurrence Surface Approximation. (See references (i) and (ii) above)
Medical applications for classification

Dermatologist-level classification of skin cancer

An artificial intelligence trained to classify images of skin lesions as benign lesions or malignant skin cancers achieves the accuracy of board-certified dermatologists.

In this work, we pretrain a deep neural network at general object recognition, then fine-tune it on a dataset of ~130,000 skin lesion images comprised of over 2000 diseases.
A new tool in the box!

MACHINE LEARNING

New tool in the box

A recent burst of activity in applying machine learning to tackle fundamental questions in physics suggests that associated techniques may soon become as common in physics as numerical simulations or calculus.

Lenka Zdeborová

The goal of machine learning, broadly speaking, is to design a computer code — the eponymous machine — capable of discovering meaningful structure in data. The last decade saw a game-changing revolution unfold in this field: with the development of deep neural networks, tasks that were considered inaccessible to automated learning became possible. This prompted fierce competition in the artificial intelligence market, but it also brought promise to many areas of data-intensive fundamental science — with physics being no exception. Current machine-learning systems are not yet able to divine the laws of general relativity from planetary data, but they are able to reliably recognize human faces, detect objects in photographs and even beat world champions of Go. And now, writing in Nature Physics, two groups have used artificial neural networks to recognize different phases of matter and localize associated phase transitions.

Juan Carrasquilla and Roger Melko

It should be stressed that saying one applied machine learning to a given problem is about as generic as saying that one used numerical simulations. It is clear to every researcher in physics that there are many kinds of numerical learning. Another group recently showed that a support vector machine can be used to classify which particles in a glassy system are susceptible to rearrangement. Decision forests have been used to classify metals from insulators based on the hybridization function, combined with kernel ridge regression to predict correlation functions in many-body physics. For the quantum systems
3.1. The Mathematical Problem

The data for the Challenge consisted of 800,000 fully simulated events provided by the ATLAS Collaboration corresponding to the signal process with Higgs decaying to $\tau^+\tau^-$ and background events from top-antitop and $Z \rightarrow \tau^+\tau^-$. Details of the ATLAS Experiment can be found in Ref. [13]. For each event, 30 numbers were recorded, including “primitive” quantities such as the jet and lepton momenta as well as derived quantities like the missing transverse energy and visible mass.

A subsample of 250,000 of these events were labeled accordingly as signal or background. This training sample was used by participants to design an algorithm that could be used to search for the signal process. The remaining 550,000 events were used for testing the performance of the algorithm.

Abstract. The Higgs Machine Learning Challenge was an open data analysis competition that took place between May and September 2014. Samples of simulated data from the ATLAS Experiment at the LHC corresponding to signal events with Higgs bosons decaying to $\tau^+\tau^-$ together with background events were made available to the public through the website of the data science organization Kaggle (kaggle.com). Participants attempted to identify the search region in a space of 30 kinematic variables that would maximize the expected discovery significance of the signal process. One of the primary goals of the Challenge was to promote communication of new ideas between the Machine Learning (ML) and HEP communities. In this regard it was a resounding success, with almost 2,000 participants from HEP, ML and other areas. The process of understanding and integrating the new ideas, particularly from ML into HEP, is currently underway.
A structural approach to relaxation in glassy liquids


In contrast with crystallization, there is no noticeable structural change at the glass transition. Characteristic features of glassy dynamics that appear below an onset temperature, $T_0$ (refs 1–3), are qualitatively captured by mean field theory$^{4-6}$, which assumes uniform local structure. Studies of more realistic systems have found only weak correlations between structure and dynamics$^{7-11}$. This raises the question: is structure important to glassy dynamics in three dimensions? We answer this question affirmatively, using machine learning to identify a new field, ‘softness’ which characterizes local structure and is strongly correlated with dynamics. We find that the onset of glassy dynamics at $T_0$ corresponds to the onset of correlations between softness (that is, structure) and dynamics. Moreover, we construct a simple model of relaxation that agrees well with our simulation results, showing that a theory of the evolution of softness in time would constitute a theory of glassy dynamics.

Figure 1 | The characteristics of the softness field. a, A snapshot of the system at $T = 0.47$ and $\rho = 1.20$ with particles coloured according to their softness from red (soft) to blue (hard). b, The distribution of softness of all particles in the system (black) and of those particles that are about to rearrange (red). 90% of the particles that are about to rearrange have $S > 0$ (shaded region). None of the data included in this plot were in the training set.
What are the tasks solved by machine learning people?

- **Supervised**
  - Classification
  - Regression

- **Un-Supervised**
  - Clustering
  - Generative models

Most data are actually unlabelled!
Un-Supervised Clustering

Find « groups » in unlabelled data
Un-Supervised Clustering

https://challengedata.ens.fr/en/home

21 Challenge providers
21 Projects
2428 Participants

Supported by the CFM chair on Data Sciences at ENS

Announcement: New Season in 2018

Cluster actor faces from TV show
In this challenge, we provide you with faces extracted from 20 episodes of a TV show. The goal is to gather, for each movie, all the faces that belong to the
Hierarchical Block Structures and High-Resolution Communities in Large Networks

Tiago P. Peixoto
Phys. Rev. X 4, 011047 – Published 24 March 2014

IMDb Datasets

Subsets of IMDb data are available for access to customers for personal use. You can hold local copies of this data, and it is subject to our terms. Please refer to the Non-Commercial Licensing and copyright/license and verification.
| Un-Supervised | Generative models |
Spent a weekend training a Generative Adversarial Network on 25k 19th century portraits; results seem good. These people have never existed:
Many applications to generative models

Cleaning data: image completion

https://medium.com/@Synced/generative-face-completion-cfa85cc4e835
Many applications to generative models

Solving the quantum many-body problem with artificial neural networks

Giuseppe Carleo¹,*, Matthias Troyer¹,²

¹ See all authors and affiliations

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Supervised problems

Let's introduce some of the machine learning vernacular...
Supervised machine learning

Labeled data: \[ \{ \vec{x}_1, \vec{x}_2, \vec{x}_3, \ldots, \vec{x}_n \} \quad \vec{x} \in \mathbb{R}^d \]
\[ \{ y_1, y_2, y_3, \ldots, y_n \} \quad y \in \mathbb{R} \quad \text{or} \quad y \in \mathbb{N} \]

Goal: Find a function \( f_W(\vec{x}) \) that outputs the right class/value for an object \( \vec{x} \)

\[ f_W(\vec{x}) \] has many parameters, denoted \( W \in \mathbb{R}^p \)

Ideal: Find \( f_W(\vec{x}) \) such that the prediction error is minimal among all unseen vectors
Supervised machine learning

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Ideal: Find \( f_W(\vec{x}) \) such that the prediction error is minimal among all unseen vectors

Instead: Find \( f_W(\vec{x}) \) such that the prediction error is minimal among all vectors in the dataset
Example for cats and dog classification

Ideally: find a function

$$f_W(x)$$

That minimise the error on all possible images of cats and dogs!

Define the loss as

$$\text{loss}(F_W(.), \tilde{x}_i, y_i) = (F_W(\tilde{x}_i) - y_i)^2$$

We want to minimise the population risk defined as

$$\mathcal{R}_{\text{pop}}(F_W(.)) = \mathbb{E}_{\text{population}}(F_W(\tilde{x}) - y)^2$$

Cannot do this: I do not have access to ALL images in the universe, and most of them are not labeled anyway.
Example for cats and dog classification

Define the loss as

$$\text{loss}(F_W(.), \vec{x}_i, y_i) = (F_W(\vec{x}_i) - y_i)^2$$

We want to minimise the empirical risk defined as

$$R_{\text{empirical}}(F_W(.)) = \frac{1}{N} \sum_{i}^{\text{dataset}} (F_W(\vec{x}_i) - y_i)^2$$

We are actually minimizing the wrong function (but we have no choice)

Instead: find a function

$$f_W(\vec{x})$$

That minimise the error on the images on the dataset
The workhorse: Empirical Risk Minimisation

Minimize

$$R_{\text{empirical}}(W) = \frac{1}{N} \sum_{i}^{\text{dataset}} \ell(W, (\vec{x}_i), y_i)$$

Rationale: it should be close to

$$R_{\text{population}}(W) = \mathbb{E} \ell(W, (\vec{x}), y)$$
Statistics: How close is the empirical risk to the population risk?

Computational: How do we minimise the empirical risk?
Statistics: How close is the empirical risk to the population risk?

**Generalization bound**

Theorem (Vapnik, Chervonenkis, 1968; . . . )

Under conditions [omitted], with high probability

\[
\sup_{W \in \theta} |R_{emp}^{n}(W) - R_{pop}(W)| \leq Cst \sqrt{\frac{VC \log n}{n}}
\]

VCd dimension = capacity of your classifier

Error on unseen data = Error on training data + Additional term

Decay with number of training sample
Increase with complexity of function f
Statistics: How close is the empirical risk to the population risk?

Those results are cool and beautiful, but they have no practical consequence. No one uses generalization bounds.

Yann LeCun, Facebook IA

Error on unseen data = Error on training data + Additional term

Generalisation error Training error

Additional term
Decay with number of training sample
Increase with complexity of function f
In practice

Divide the labelled set into training, validation and testing sets

* **Training set**: used to train the classifier
* **Validation set (optional)**: choose between different methods, fine-tune parameters, 
* **Testing set**: predict the generalization error

No cheat: do not use the test set to train your algorithm!
No cheat: do not use the test set to train your algorithm!

A View from Tom Simonite

Why and How Baidu Cheated an Artificial Intelligence Test

Machine learning gets its first cheating scandal.

How much reliable is the paper "Stacked Approximated Regression Machine"? Are we going to rewrite DL frameworks?

1 Answer

Zhaojun Zhang, Trained as a Bayesian for two years
Answered Sep 10, 2016

From Arxiv, the paper has been withdrawn: A Simple Deep Learning Approach. So, it is unlikely that we are going to rewrite DL frameworks based on this paper.

326 Views

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Questions?

**Statistics**: How close is the empirical risk to the population risk?

**Computational**: How do we minimise the empirical risk?
Computational: How do we minimise the empirical risk?

**Example 1**: binary classification with the perceptron (Rosenblath 1958)

\[ z_i = (y_i, x_i), \quad y_i \in \{0, 1\}, \quad x_i \in \mathbb{R}^d \]

\[ \hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^{n} (y_i - \sigma(w, x_i))^2, \]

\[ \sigma(u) = \frac{1}{1 + e^{-u}}. \]
Computational: How do we minimise the empirical risk?

**Example 2: Multi-layered networks**

1 layer neural network: \( w \in \mathbb{R}^d \)

\[
\ell(w; y, x) = (y - \sigma(\langle w, x \rangle))^2
\]

2 layers neural network: \( w_1 \in \mathbb{R}^d, \ W_2 \in \mathbb{R}^{d \times d} \)

\[
\ell(\theta; y, x) = (y - \sigma(w_1^T \sigma(W_2x)))^2
\]

3 layers neural network: \( w_1 \in \mathbb{R}^d, \ W_2 \in \mathbb{R}^{d \times d}, \ W_3 \in \mathbb{R}^{d \times d} \)

\[
\ell(\theta; y, x) = (y - \sigma(w_1^T \sigma(W_2 \sigma(W_3x))))^2
\]
Minimising the cost function by gradients descent

\[ \theta^{t+1} = \theta^t - \gamma \nabla R(\theta^t) \]

If \( \gamma \) small enough, converge to a (possible local) minima
Minimising the cost function by gradients descent

\[ \theta^{t+1} = \theta^t - \gamma \nabla R(\theta^t) \]

If \( \gamma \) small enough, converge to a (possible local) minima

Standard (or "batch") gradient descent

Compute the gradient by averaging the derivative of the loss is the entire training set

\[ \theta^{t+1} = \theta^t - \gamma \sum_i \frac{1}{N} \nabla l(\theta^t; \bar{x}_i, y_i) \]
Minimising the cost function by gradients descent

\[ \theta^{t+1} = \theta^t - \gamma \nabla R(\theta^t) \]

If \( \gamma \) small enough, converge to a (possible local) minima

Stochastic (or « mini-batch") gradient descent

Compute the gradient by averaging the derivative of the loss in a mini-batch

1) Divide the training set into \( P \) batch of size \( B \)
2) For each batch, do

\[ \frac{\theta^{t+1}}{P} = \theta^t - \gamma \sum_{i \text{ in mini batch}} \frac{1}{B} \nabla l(\theta^t; x_i, y_i) \]

3) One « epoch » (t->t+1) means running the algorithm through all mini-batches
Why Mini-batch gradient descent?

\[
\hat{\theta}^{t+1} = \hat{\theta}^{t} - \gamma \sum_{i \text{ in mini batch}} \frac{1}{B} \nabla l(\hat{\theta}^{t}; \tilde{x}_i, y_i)
\]

- The model update frequency is higher than batch gradient descent: faster and memory efficient (often nothing else is actually possible)

- Effective noise in the dynamics helps optimization/regularization: works better than full batch minimisation in practice
Many mini-batch algorithms
(but we shall discuss them later)
That’s all for today

Supervised learning

Empirical risk minimisation

Training, Validation, and Test sets

Gradient descent

Mini-batch Gradient descent