Introduction

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You have data from a stochastic system:

- from an airport camera,
- from a camera seeing an insect flying, a body suit on a human, ...
- weather observations, stock market data,
- a tactile sensor on a robot hand, etc.

and you want to use these data to

- detect suspicious behaviour,
- classify movement,
- predict future data,
- control a system

How?

The description is simple, in the form a model of your stochastic process:

$$y = f(x)$$

where y, x are vectors, and f() is the underlying model.

Consider the above also in the special form

$$x_{t+1} = f(x_t)$$

where we use the Markov assumption.

Key question: how do we get f()?

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approximating f() through system knowledge

If, e,g., we find all red objects in HSV space

- or we compute the Cartesian end position of a robot arm by inverse kinematics
- or we compute radiation of an exploding supernova
- or we want to determine a plant by analysing stem, leaves, flowers

in each of these cases, we formalise expert knowledge in terms of a set of rules or decisions.

This is known classical control, expert systems, etc.

It is your best solution if your approximation ('model') $\hat{f}_{\theta}()$ to f() can be described at your desired accuracy. The description of $\hat{f}()$ is obtained using innate knowledge of f. How do you know if your accuracy is good enough?

approximating f() when system knowledge fails

If, e.g., you want to recognise faces, or a plant from a picture

- or want to create a programme to play go
- or want to find good grip positions on a random object
- or want to recognise a composer by listening to their music

in each of these cases, the number of cases is too large to exhaustively describe accurately enough.

We have no detailed knowledge of f(), and instead we use a general form for $\hat{f}()$ that can represent any function.

But both approaches are in principle the same:

- 1. create a parameterised model $\hat{f}(x,\theta)$
- 2. find best values for the parameters θ

How do we do the second step? Two options:

- a. we make \hat{f} behave as close as possible to f (e.g., maximise number of go games won)
- b. we make \hat{f} be as close as possible to f through $\min_{\theta} |\hat{f}_{\theta}(x) f(x)|$. Failing f, we sample f in $\{x_i, y_i\}$ to $\min_{\theta} \sum_i |\hat{f}_{\theta}(x_i) y_i|$ as a proxy. We can write this as $\max_{\theta} \prod_i p_{\theta}(y_i \mid x_i)$.

The basic problem that we study in probability theory:

Given a data generating process, what are the properties of the outcomes?

The basic problem of statistics (or better statistical inference) is the inverse of probability theory:

Given the outcomes, what can we say about the process that generated the data?

Statistics uses the formal language of probability theory.

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discriminative vs. generative models

The above describes discriminative models, which map inputs to outputs and are described by p(y|x).

Often we are interested in *generative* models, represented by p(x,y) (if we have labels) or p(x) (if we have no labels).

We can still measure quality, through $\max_{\theta} \prod_{i} p_{\theta}(x_i)$.

Generative models can be used to generate new data, or to test the likelihood of a datum.

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examples of discriminative models

This is nothing new. Typically, such models use

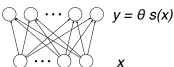
$$\hat{f}_{\theta}(x) = \sum_{k} \theta_k s_k(x)$$

which describes, for instance, Fourier transform, polynomials, etc.

This general form is known as *linear regression*, because the regression function is linear in the parameters θ .

This is important as we have a closed-form solution to find θ .

In the connectionism paradigm, the models can be drawn like this:



aka the perceptron

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curse of dimensionality

Even if x is only 100-dimensional

and you have a trillion data,

those data cover only 10^{-18} of the input space.

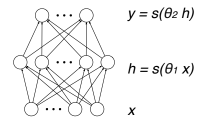
This can be very problematic for such models.

We need models that generalise further.

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extending the connectionist idea



did you see the small trick I did?

That's a great idea from the 1970s which suggests,

$$\hat{f}_{\theta}(x) = \sum_{k} \theta_{k} s \left(\sum_{i} \theta_{l} x \right)$$

and the neural network (= multi-layer perceptron) is born.

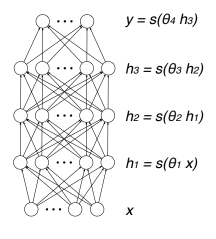
Problem: finding θ can no longer be done closed-form. Gradient-based optimisation is the standard approach.

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a deep neural network just has more layers



history of neural networks 1

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1960s: the linear perceptron learns from data (Rosenblatt et al)
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1969: the perceptron can't do XOR (Minsky & Papert)

1970–1980: nonlinear networks trained with back-propagation (Linnainmaa; Dreyfus; Werbos; Rumelhart)

1990s: one hidden layer can represent any (Borel-measurable) function

mid 90s: NN's can't do everything / do not generalise / ...

mid 90s: support vector machines (SVMs) are great!

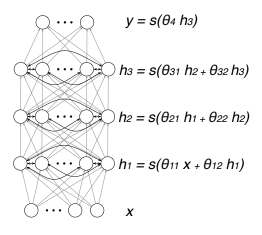
1995–2000: SVM too slow / too many SVs

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history of neural networks 2

- 2000-: probabilistic models for machine learning (ML)
 - 2006: **deep neural networks**, trained with *Restricted Boltzmann Machines* and backprop
 - 2009: deep NNs can be trained with just BP, **much compute power** (GPU) and **enough data**
- 2011-: recurrent neural networks resurrect for time-series modelling
- 2012–: convolutional neural networks (CNN) start winning most vision benchmarks; recurrent neural networks applied to speech recognition in Android
- 2013-: probabilistic NN (variance propagation; variational autoencoder)
- 2015-: show cases in robotics, sensory processing, ...
 - 2017: "Attention is all you need" . . .

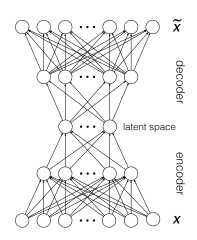
a recurrent neural network has an internal state



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Introduction

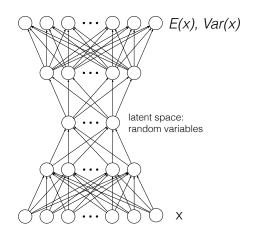
autoencoder: NN in a special form



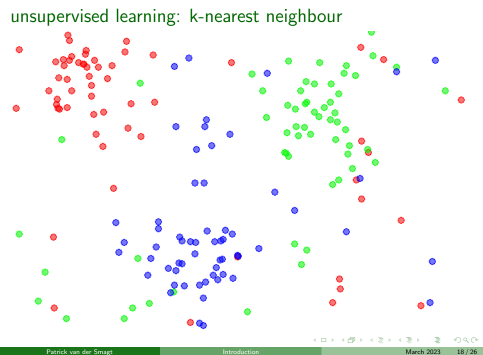
 $\mathsf{Dim}(\mathsf{latent\ space}) \ll \mathsf{Dim}(x)$

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VAE: probabilistic AE



'nonlinear PCA'



1-NN algorithm

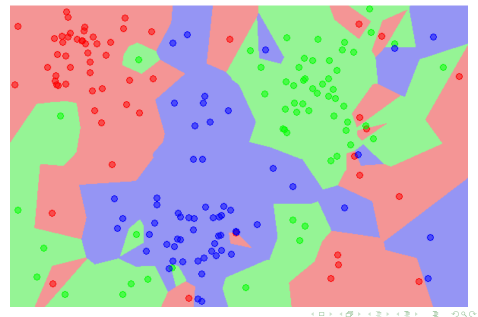
Example: let's take k = 1

- define a distance measure
- 2 determine the number of classes
- $\mathbf{3}$ for each new data point x:
 - 1 determine the distance to all other points
 - 2 find the nearest neighbours
- f 4 the class c of the new data point is determined

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1-NN algorithm: overfitting



k-NN equation

how can we describe this equation?

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k-NN equation

how can we describe this equation?

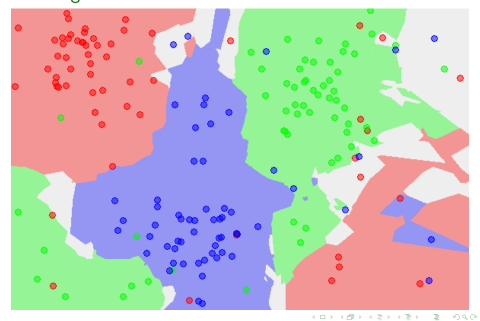
$$p(y = c \mid x) = \frac{1}{k} \sum_{i \in \text{neighbours}} \delta_{y_i c}$$

mit

$$\delta_{ij} = \begin{cases} 0 & \text{if } i \neq j \\ 1 & \text{if } i = j \end{cases}$$

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4-NN algorithm



k is a hyperparameter, the value of which deter	mines the outcome. How can we
optimise it?	
data set	

 \boldsymbol{k} is a hyperparameter, the value of which determines the outcome. How can we optimise it?

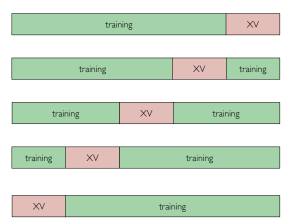
data set

by cross validation

training

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5-fold crossvalidation



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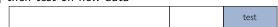
but a more honest method is:

first learn



but a more honest method is:

- 1 first learn
 - training XV
 - 2 then test on new data



problems with kNN

- finding the right distance measure is difficult and strongly influences the result
- in high-dimensional spaces, distances stop making sense

 https://stats.stackexchange.com/questions/99171/why-is-euclidean-distance-not-a-good-metric-in-high-dimensions
- finding the neighbours is very expensive in high-dimensional spaces
- finding the neighbours is very expensive if many data exist

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