## Introduction and Overview

Traditional Algorithms vs ML

**Understanding Machine Learning** 

#### goals of this course

 understanding of foundational ML concepts and commonalities between different methods

 ability to properly use ML for scientific or business problems

#### schedule of lectures

- 1. introduction and overview
- 2. statistical learning
- 3. non-linear models
- 4. generalization
- 5. deep learning
- 6. transformers
- 7. generative models
- 8. causality
- 9. reinforcement learning

# AI/ML Overview

## Main Areas of Artificial Intelligence

#### computer vision

(spatial structures, state-of-the-art: Convolutional Neural Networks)

natural language processing

(sequential structures, state-of-the-art: transformers)

• automated decision making, robotics (reinforcement learning)

All of these are enabled by one key ingredient:

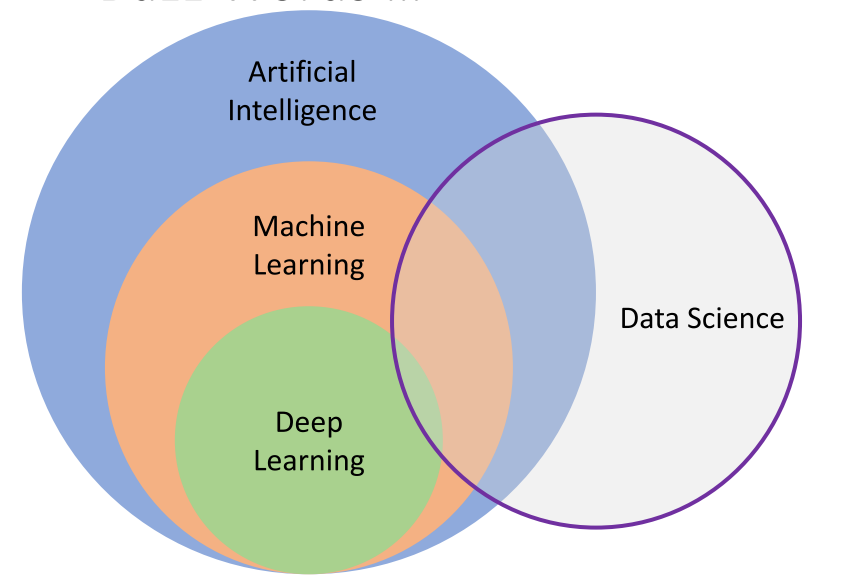
- learning from experience (Machine Learning)
- knowledge representation, automated reasoning



from wikipedia

agency:perception – thought – action

## Buzz Words ...



#### Deep Learning:

special kind of ML algorithms using (deep) neural networks

#### Data Science:

extract knowledge from data (by means of ML, among other things)

## Traditional Algorithms and GOFAI

traditional algorithms:

explicit (handcrafted) instructions for each situation



from wikipedia

symbolic AI (aka GOFAI):

use knowledge by means of symbols (as representations), logic, search (e.g., expert systems like Deep Blue)

Public perception is changing over time: A modern chess program, nowadays disparaged as brute computing, would have been considered intelligent in the 50s.

## ML: Learning from Experience/Data

mainly exploiting statistical dependencies with the aim of generalization to new (e.g., future) data

training (usually offline optimization):

ML algorithm + data = explicit algorithm (to be used at inference time)

> reduction of complexity and much better generalizability compared to handcrafted algorithms

analogy: Humans do not hit the ground running, but have learning capabilities. 

develop ML algorithm and let it learn from data

## Hot Debate: Connectionism vs Symbolic Al

#### connectionists:

learn from (big) data without prior knowledge

#### symbolists:

use knowledge with only modest input data

(crude) analogy: learning and evolution philosophical: empiricist and rationalist schools of mind

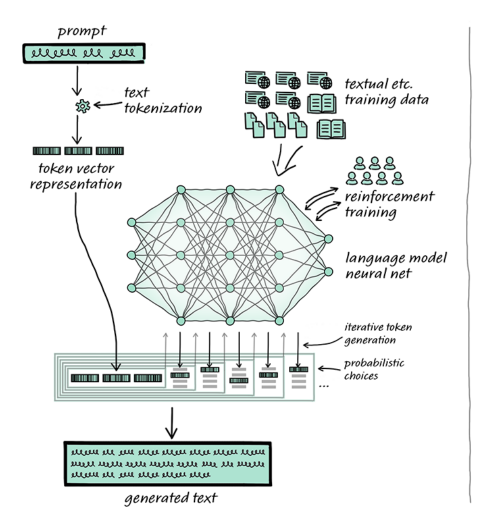
#### famous move 37 from Max Tegmark

hybrid approaches often most successful:

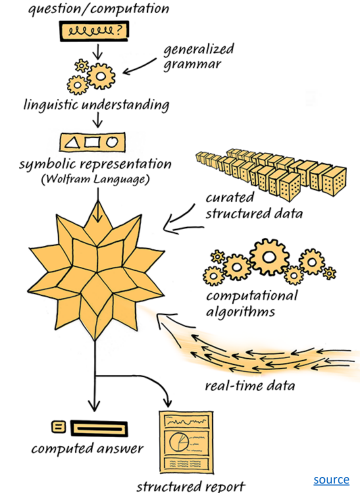
- e.g., AlphaGo with Deep Reinforcement Learning and Monte Carlo Tree Search
- feature engineering for ML models also kind of symbolic knowledge representation

## Hybrid Approach for Language Models?

#### **ChatGPT**



#### Wolfram Alpha



tool usage:

LangChain

## Supercharging the Scientific Method

use ML and data to replace or enhance explicit methods relying on detailed domain knowledge (Software 2.0)

- → overcome our evolutionary limitations in math with clever learning algorithms and collecting data
- → immediate impact on many aspects of industry, business, and science, formulated as narrow tasks with strictly defined inputs (aka weak AI)

more imminent than (still philosophical) long-term quest for human-level AI (aka strong AI, AGI), i.e., general-purpose intelligence

(although recent language models show multi-purpose capabilities)

## When to apply ML?

#### complexity

- decisions under uncertainty, many influencing factors
- e.g., demand forecasting, DNA sequencing
- difficult for humans, direct model inexpressible

#### automation

- e.g., face and speech recognition, autonomous driving
- goal to reach human-level performance

... and of course you need data to learn from

# and more recently: generative tasks

- rather than predictive (or discriminative) ones
- e.g., image generation, conversational AI, new proteins or materials

# Learning Paradigms

## Supervised Learning

#### **learning by teacher** $\rightarrow$ usually rather narrow tasks

#### **Target Quantity**

- **known in training**: labeled samples or observations from past
- to be **predicted** for unknown cases (e.g., future values)

#### **Features**

input information that is

- correlated to target quantity
- known at prediction time

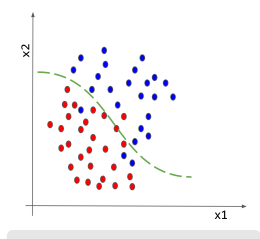


#### **Example: Spam Filtering**

Classify emails as spam or no spam

use accordingly **labeled** emails as training set

use information like occurrence of specific words or email length as features



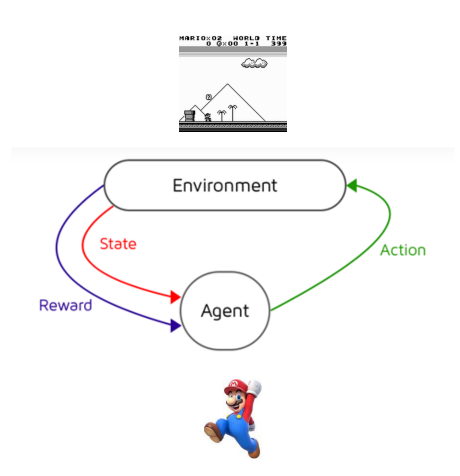
features x1 and x2 spam, no spam

## Reinforcement Learning

#### learning by trial-and-error

goal-based approach → more generic than supervised learning (but sparse reward signals)

- receiving feedback from the environment
- no supervision
- sequential decision making (delayed rewards)
- exploration and exploitation



## ML needs lots of training data

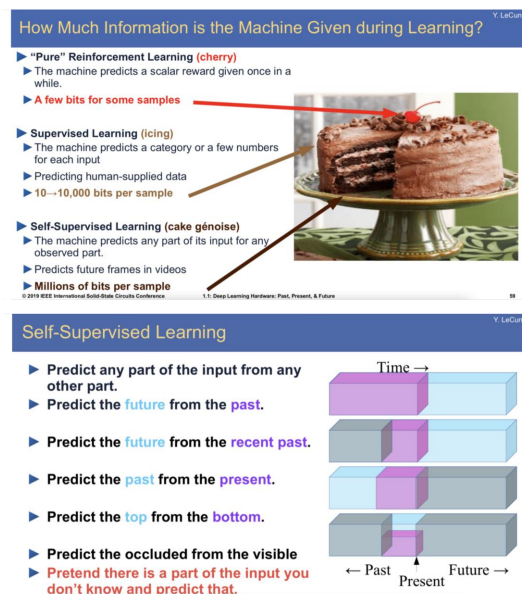
## Unsupervised Learning

#### learning by observation

no target information  $\rightarrow$  kind of "vague" pattern recognition (but plenty of data)

#### self-supervised:

- input-output mapping like supervised learning
- but generating labels itself from input information
- learning of semantic feature representations
- e.g., word2vec, BERT, GPT

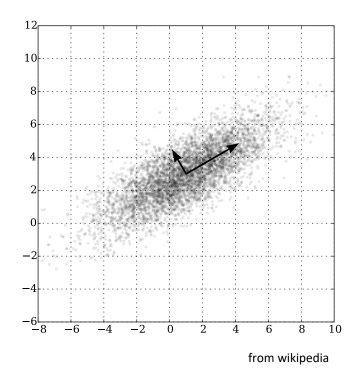


## Example for Unsupervised Learning

dimensionality reduction by principal component analysis (PCA) using only first few principal components (eigenvectors of data's covariance matrix)

intuition: fitting p-dimensional ellipsoid to data

- axes representing principal components
- large axis → high variance, small axis → low variance
- successively choose directions of maximum variance
- → account for as much variability (uniqueness) of data set as possible



often used as lower-dimensional features in other (supervised) methods

# Fitting / Statistical Learning

### Notation

- random variable: *X*
- vector of p random variables  $X_i$  (features): X
- observation of random variable X: x
- matrix of n observations of p features  $x_{ij}$ :  $\mathbf{X}$
- vector of observations: x
  - vector of n observations  $x_i$ : x
  - vector of observation of p features  $x_i$ : x
- column vector: *x*
- row vector:  $x^T$

- parameter:  $\beta$
- vector of parameters  $\beta_k$ :  $\beta$
- probability that X takes on value  $x_0$ :  $P(X = x_0)$
- probability distribution: p(x) = P(X = x)

design matrix

## Supervised Learning Scenario

map inputs to output: y = f(x) (estimated:  $\hat{f}(x)$ ) random variables Y and  $X = (X_1, X_2, \dots, X_p)$ 



... ML ...



## ML domain:

no deterministic dependencies between input and output

#### classification

- categorical target (e.g., image of cat or not  $\rightarrow y = 0$  or y = 1)
- predict probability to belong to specific class

#### regression

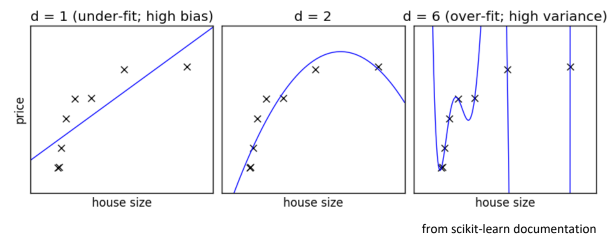
- real-valued target
- $Y \in [0, \infty)$  (e.g., demand forecasting) or  $Y \in (-\infty, \infty)$

## Curve Fitting / Parameter Estimation

fit train data set of  $(y_i, x_i)$  pairs

- minimization of cost function
- e.g., least squares method, maximium likelihood estimation
- usually many dimensions (features  $X_j$ )

d: degree of fitted polynomial  $\rightarrow$  number of parameters



apply learned statistical dependencies from training to new test data set

different  $(y_i, x_i)$  pairs considered as random samples of underlying data-generating process (i.i.d. assumption), for both train and test data sets

## Generalization

#### core of ML:

**empirical risk minimization** (training error) as proxy for minimizing unknown population risk (test error, aka generalization error or out-of-sample error)

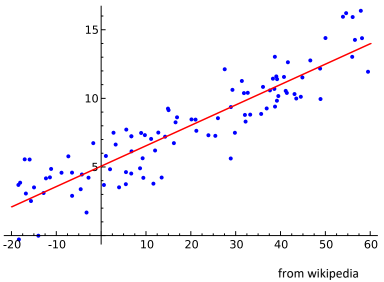
generalization gap: difference between test and training error

- interpolation: to unencountered samples from training environment
- **extrapolation**: to testing conditions differing from training environment curse of dimensionality: "learning in high dimensions always amounts to extrapolation"  $\rightarrow$  need for appropriate inductive bias

## Generalized Linear Models

## Linear Regression

$$y_i = \alpha + \sum_{j=1}^p \beta_j x_{ij} + \varepsilon_i$$
 (model)



y: dependent variable / target

x: p independent variables / features

 $\alpha$ ,  $\beta$ : p+1 parameters

ε: error term / statistical noise

vector (y) or matrix (X) of given data

 $\rightarrow$  to be fitted reflects assumed data distribution (here: Gaussian with same variance  $\sigma^2$  for all samples)

- X and Y jointly distributed random variables
- $\hat{f}(x)$ : predict, e.g., conditional mean of conditional density function p(y|x)

depending on used loss function (conditional mean for squared loss of least squares method)

## Linear Regression

fit: 
$$f(x_i)$$

$$y_i = \hat{\alpha} + \sum_{j=1}^p \hat{\beta}_j x_{ij} + \varepsilon_i$$
predict: 
$$\hat{y}_i = E[Y|X = x_i] = \hat{f}(x_i) = \hat{\alpha} + \sum_{j=1}^p \hat{\beta}_j x_{ij}$$

$$p(y|x_i) = \mathcal{N}(y; \hat{y}_i, \hat{\sigma}^2)$$
Gaussian mean variance

to be estimated:

• 
$$\hat{\alpha}$$
,  $\hat{\beta}$ 

$$\Rightarrow \hat{\sigma}^2 = \frac{1}{n} \sum_{i=1}^n \left( y_i - \hat{f}(\mathbf{x}_i) \right)^2$$

(approximating assumed true  $\alpha$ ,  $\beta$ ,  $\sigma$ )

## Multiplicative Model

- count data:  $Y \in [0, \infty)$
- Y follows Poisson (or negative binomial / Poisson-gamma) distribution

log-linear model (Gaussian errors in fit, Poisson with mean  $\hat{y}_i$  predicted):

$$\log(E[Y|X=x_i]) = \hat{\alpha} + \sum_{j=1}^{p} \hat{\beta}_j x_{ij}$$
 single parameter

• further advantage: usually multiplicative effects for count data, i.e., proportional (small effects for small counts, large effects for large counts)

## Scheme of Generalized Linear Models

$$g(E[Y|X = x_i]) = \hat{\alpha} + \sum_{j=1}^{p} \hat{\beta}_j x_{ij}$$

#### link function *g*:

- linking range of Y to linear predictor
- canonical forms for different
   Y distributions (e.g., log for
   Poisson, identity for Gaussian
   → linear regression)

Y following probability distribution from exponential family (e.g., Poisson or Gaussian)

## Classification: Logistic Regression

- predict probability  $p_i$  for y=1 respectively y=0 for each sample
- link function: logit (log-odds)
- *Y* following Bernoulli distribution

$$logit(E[Y|X = x_i]) = ln\left(\frac{p_i}{1 - p_i}\right)$$
$$= \hat{\alpha} + \sum_{i=1}^{p} \hat{\beta}_i x_{ij}$$

## Toward Non-Linear Models

## Generalized Additive Models

blending of Generalized Linear Models and additive models

$$g(E[Y|X = x_i]) = \hat{\alpha} + \sum_{j=1}^p \hat{h}_j(x_{ij})$$

smooth functions

- potentially non-parametric form
- describe non-linear effects
- estimated, e.g., via backfitting algorithm

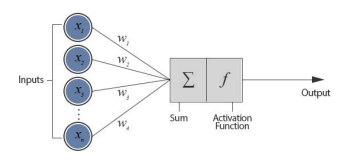
extension: add interaction terms between different features, e.g.,  $X_3$  and  $X_4$ 

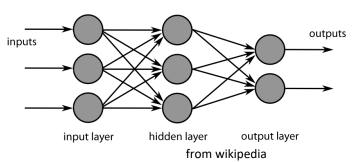
## Algorithmic Families and Linear Building Blocks

#### linear (parametric) models

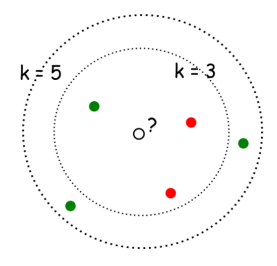
- linear regression
- Generalized Linear Models
- Generalized Additive Models

**neural networks**: non-linear just by means of activation functions





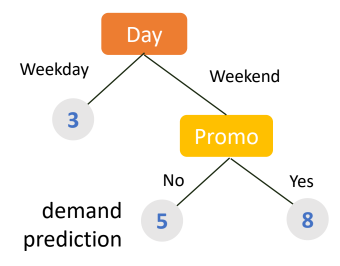
nearest neighbors (local methods, instance-based learning) – non-parametric models



with k = 3, • with k = 5, •

**kernel/support-vector machines**: linear model (maximum-margin hyperplane) with kernel trick

#### decision trees



#### often used in ensemble methods

- bagging: random forests
- boosting: gradient boosting

At its heart, all the diverse statistical learning methods are reflections of the **same underlying concept**, and just differ in their applicability for different use cases.

(need to find method with best inductive bias for the task at hand → generalization capability)

# ML Workflow

## Modeling

#### **extract features**

- help the ML algorithm to better understand the data
- impose assumptions hard to discover in the raw data

#### choose ML algorithm

- from open-source libraries like scikit-learn or pytorch, rarely write an own one
- many different algorithms available, differently suited for given task

## execute hyperparameter tuning

- variety of different forms
- model settings not all automatically adjusted by the machine

## Evaluation

# cross-validation Train on k-1 splits time Trest ... Train Test ... Train Test

decide on acceptance of model changes by means of A/B tests: improved model vs baseline (current best)

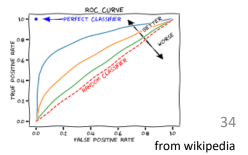
#### measure accuracy of predictions

#### regression

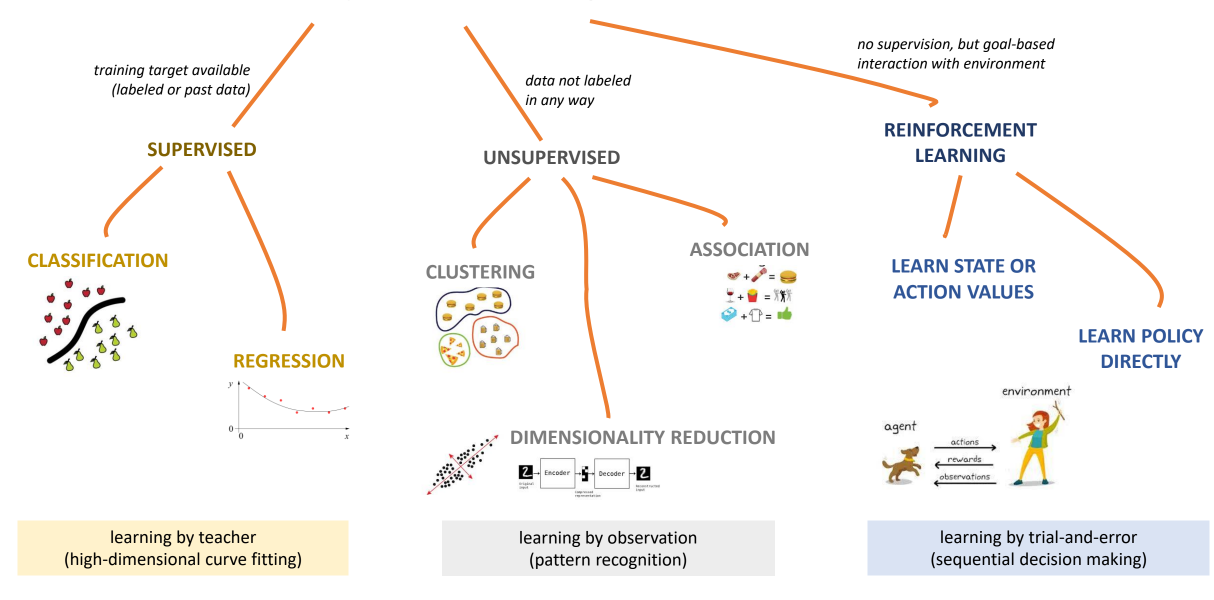
- point estimate: absolute (MAD, MSE, ...) or relative (MAPE, ...) metrics
- full probability distribution: tricky;)

#### classification

ROC curve (true and false positve rates)

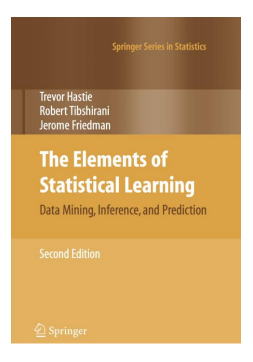


#### **MACHINE LEARNING**



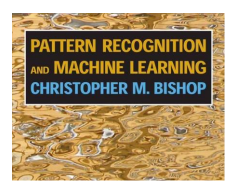
### Literature

nice book on the foundations of ML (relevant for the whole course):

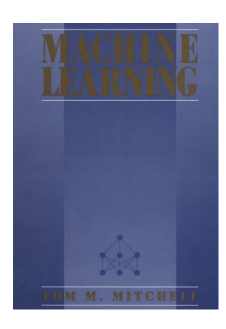


<u>Hastie</u>

other general overviews:



**Bishop** 



**Mitchell** 

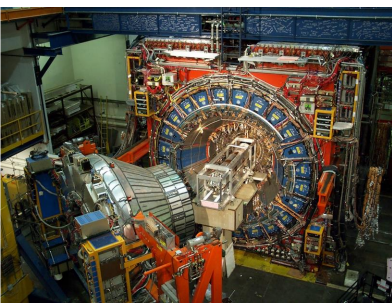
## Scientific Application: ML in Particle Physics

example: classification of decay signatures in particle colliders

Tevatron accelerator at Fermilab



CDF detector at Tevatron



## charmed baryon signals filtered out of background

