

# Neurocomputing

Basics in mathematics

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## **Outline**

- 1. Linear algebra
- 2. Calculus
- 3. Probability theory
- 4. Statistics
- 5. Information theory

# 1 - Linear algebra

## Mathematical objects

- Scalars x are 0-dimensional values. They can either take real values ( $x \in \Re$ , e.g. x = 1.4573, floats in CS) or natural values ( $x \in \mathbb{N}$ , e.g. x = 3, integers in CS).
- **Vectors**  ${\bf x}$  are 1-dimensional arrays of length d.
- The bold notation  $\mathbf x$  will be used in this course, but you may also be accustomed to the arrow notation  $\overrightarrow{x}$  used on the blackboard. When using real numbers, the **vector space** with d dimensions is noted  $\Re^d$ , so we can note  $\mathbf x \in \Re^d$ .
- Vectors are typically represented vertically to outline their d elements  $x_1, x_2, \ldots, x_d$ :

$$\mathbf{x} = egin{bmatrix} x_1 \ x_2 \ dots \ x_d \end{bmatrix}$$

## Mathematical objects

- Matrices A are 2-dimensional arrays of size (or shape)  $m \times n$  (m rows, n columns,  $A \in \Re^{m \times n}$ ).
- They are represented by a capital letter to distinguish them from scalars (classically also in bold  $\bf A$  but not here). The element  $a_{ij}$  of a matrix A is the element on the i-th row and j-th column.

$$A = egin{bmatrix} a_{11} & a_{12} & \cdots & a_{1n} \ a_{21} & a_{22} & \cdots & a_{2n} \ dots & dots & dots \ a_{m1} & a_{m2} & \cdots & a_{mn} \end{bmatrix}$$

• **Tensors** A are arrays with more than two dimensions. We will not really do math on these objects, but they are useful internally (hence the name of the tensorflow library).

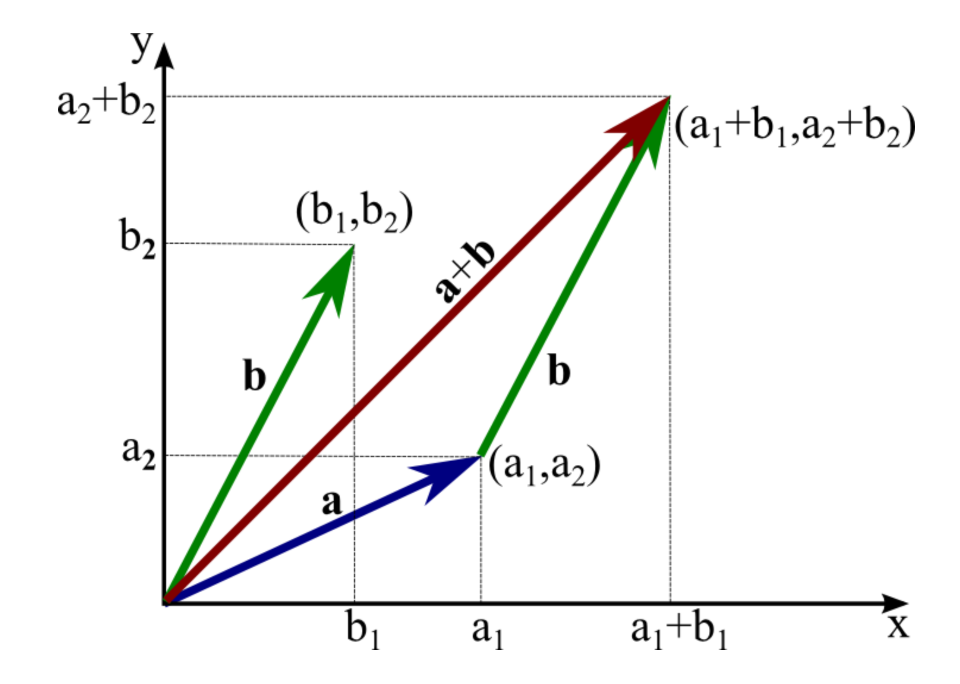
#### **Vectors**

- A vector can be thought of as the coordinates of a point in an Euclidean space (such the 2D space), relative to the origin.
- A vector space relies on two fundamental operations, which are that:
- Vectors can be added:

$$\mathbf{x}+\mathbf{y}=egin{bmatrix} x_1\ x_2\ dots\ x_d \end{bmatrix}+egin{bmatrix} y_1\ y_2\ dots\ y_d \end{bmatrix}=egin{bmatrix} x_1+y_1\ x_2+y_2\ dots\ x_d+y_d \end{bmatrix}$$

Vectors can be multiplied by a scalar:

$$egin{array}{c} egin{array}{c} egin{array}{c} x_1 \ x_2 \ dots \ x_d \end{bmatrix} &= egin{array}{c} a \ x_2 \ dots \ a \ x_d \end{bmatrix}$$



Source: https://mathinsight.org/image/vector\_2d\_add

## Properties of vector spaces

- These two operations generate a lot of nice properties (see https://en.wikipedia.org/wiki/Vector\_space for a full list), including:
  - associativity:

$$x + (y + z) = (x + y) + z$$

commutativity:

$$x + y = y + x$$

the existence of a zero vector

$$x + 0 = x$$

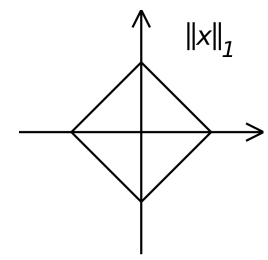
• inversion:

$$\mathbf{x} + (-\mathbf{x}) = \mathbf{0}$$

distributivity:

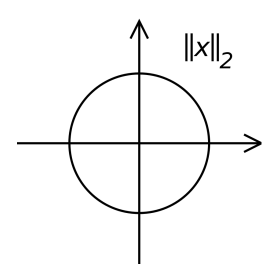
$$a\left(\mathbf{x}+\mathbf{y}\right)=a\,\mathbf{x}+a\,\mathbf{y}$$

#### Norm of a vector



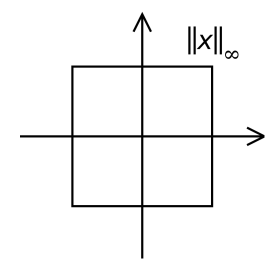
• Vectors have a **norm** (or length)  $||\mathbf{x}||$ . The most intuitive one (if you know the Pythagoras theorem) is the **Euclidean norm** or  $L^2$ -norm, which sums the square of each element:

$$||\mathbf{x}||_2 = \sqrt{x_1^2 + x_2^2 + \ldots + x_d^2}$$



ullet Other norms exist, distinguished by the subscript. The  $L^1$ -norm (also called the Manhattan norm) sums the absolute value of each element:

$$||\mathbf{x}||_1 = |x_1| + |x_2| + \ldots + |x_d|$$



• The **p-norm** generalizes the Euclidean norm to other powers p:

$$||\mathbf{x}||_p = (|x_1|^p + |x_2|^p + \ldots + |x_d|^p)^{rac{1}{p}}$$

ullet The **infinity norm** (or maximum norm)  $L^\infty$  returns the maximum element of the vector:

$$||\mathbf{x}||_{\infty} = \max(|x_1|,|x_2|,\ldots,|x_d|)$$

## **Dot product**

 One important operation for vectors is the dot product (also called scalar product or inner product) between two vectors:

$$egin{aligned} \langle \mathbf{x} \cdot \mathbf{y} 
angle = \langle egin{bmatrix} x_1 \ x_2 \ dots \ x_d \end{bmatrix} \cdot egin{bmatrix} y_1 \ y_2 \ dots \ y_d \end{bmatrix} 
angle = x_1 \, y_1 + x_2 \, y_2 + \ldots + x_d \, y_d \end{aligned}$$

- The dot product basically sums one by one the product of the elements of each vector. The angular brackets are sometimes omitted  $(\mathbf{x} \cdot \mathbf{y})$  but we will use them in this course for clarity.
- One can notice immediately that the dot product is symmetric:

$$\langle {f x} \cdot {f y} 
angle = \langle {f y} \cdot {f x} 
angle$$

and linear:

$$\langle (a\mathbf{x} + b\mathbf{y}) \cdot \mathbf{z} \rangle = a \langle \mathbf{x} \cdot \mathbf{z} \rangle + b \langle \mathbf{y} \cdot \mathbf{z} \rangle$$

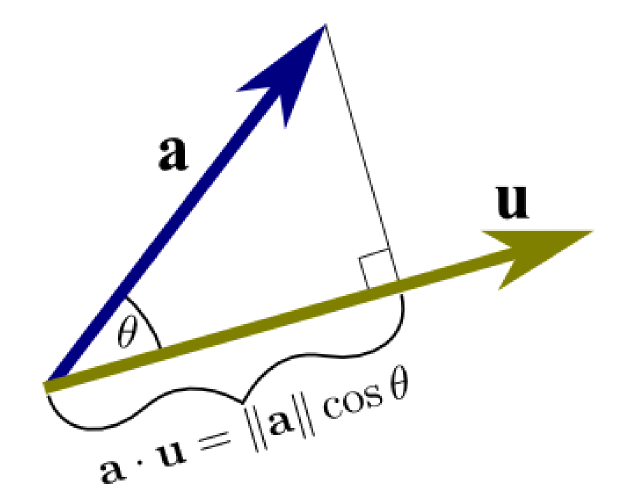
## **Dot product**

• The dot product is an indirect measurement of the angle  $\theta$  between two vectors:

$$\langle \mathbf{x} \cdot \mathbf{y} \rangle = ||\mathbf{x}||_2 \, ||\mathbf{y}||_2 \, \cos(\theta)$$

- If you normalize the two vectors by dividing them by their norm (which is a scalar), we indeed have the cosine of the angle between them
- The higher the normalized dot product, the more the two vectors point towards the same direction (cosine distance between two vectors).

$$\langle rac{\mathbf{x}}{||\mathbf{x}||_2} \cdot rac{\mathbf{y}}{||\mathbf{y}||_2} 
angle = \cos( heta)$$



Source: https://mathinsight.org/image/dot\_product\_projection\_unit\_vector

#### **Matrices**

• Matrices are derived from vectors, so most of the previous properties will be true. Let's consider this 4x3 matrix:

$$A = egin{bmatrix} a_{11} & a_{12} & a_{13} \ a_{21} & a_{22} & a_{23} \ a_{31} & a_{32} & a_{33} \ a_{41} & a_{42} & a_{43} \end{bmatrix}$$

• Each column of the matrix is a vector with 4 elements:

$$\mathbf{a}_1 = egin{bmatrix} a_{11} \ a_{21} \ a_{31} \ a_{41} \end{bmatrix} \qquad \mathbf{a}_2 = egin{bmatrix} a_{12} \ a_{22} \ a_{32} \ a_{42} \end{bmatrix} \qquad \mathbf{a}_3 = egin{bmatrix} a_{13} \ a_{23} \ a_{33} \ a_{43} \end{bmatrix}$$

ullet A m imes n matrix is therefore a collection of n vectors of size m put side by side column-wise:

$$A = egin{bmatrix} \mathbf{a}_1 & \mathbf{a}_2 & \mathbf{a}_3 \end{bmatrix}$$

## **Properties of matrix spaces**

• All properties of the vector spaces (associativity, commutativity, distributivity) also apply to matrices, as additions and multiplications with a scalar are defined.

$$lpha\,A + eta\,B = egin{bmatrix} lpha\,a_{11} + eta\,b_{11} & lpha\,a_{12} + eta\,b_{12} & lpha\,a_{13} + eta\,b_{13} \ lpha\,a_{21} + eta\,b_{21} & lpha\,a_{22} + eta\,b_{22} & lpha\,a_{23} + eta\,b_{23} \ lpha\,a_{31} + eta\,b_{31} & lpha\,a_{32} + eta\,b_{32} & lpha\,a_{33} + eta\,b_{33} \ lpha\,a_{41} + eta\,b_{41} & lpha\,a_{42} + eta\,b_{42} & lpha\,a_{43} + eta\,b_{43} \end{bmatrix}$$

**Note:** Beware, you can only add matrices of the same dimensions m imes n. You cannot add a 2 imes 3 matrix to a 5 imes 4 one.

## **Transposition**

• The **transpose**  $A^T$  of a  $m \times n$  matrix A is a  $n \times m$  matrix, where the row and column indices are swapped:

$$A = egin{bmatrix} a_{11} & a_{12} & \cdots & a_{1n} \ a_{21} & a_{22} & \cdots & a_{2n} \ dots & dots & \ddots & dots \ a_{m1} & a_{m2} & \cdots & a_{mn} \end{bmatrix}, \qquad A^T = egin{bmatrix} a_{11} & a_{21} & \cdots & a_{m1} \ a_{12} & a_{22} & \cdots & a_{m2} \ dots & dots & \ddots & dots \ a_{1n} & a_{2n} & \cdots & a_{mn} \end{bmatrix}$$

This is also true for vectors, which become horizontal after transposition:

$$\mathbf{x} = egin{bmatrix} x_1 \ x_2 \ dots \ x_d \end{bmatrix}, \qquad \mathbf{x}^T = egin{bmatrix} x_1 & x_2 & \dots & x_d \end{bmatrix}$$

## **Matrix multiplication**

ullet If A is a m imes n matrix and B a n imes p matrix:

$$A = egin{bmatrix} a_{11} & a_{12} & \cdots & a_{1n} \ a_{21} & a_{22} & \cdots & a_{2n} \ dots & dots & \ddots & dots \ a_{m1} & a_{m2} & \cdots & a_{mn} \end{bmatrix}, \quad B = egin{bmatrix} b_{11} & b_{12} & \cdots & b_{1p} \ b_{21} & b_{22} & \cdots & b_{2p} \ dots & dots & \ddots & dots \ b_{n1} & b_{n2} & \cdots & b_{np} \end{bmatrix}$$

we can multiply them to obtain a m imes p matrix:

$$C = A imes B = egin{bmatrix} c_{11} & c_{12} & \cdots & c_{1p} \ c_{21} & c_{22} & \cdots & c_{2p} \ dots & dots & \ddots & dots \ c_{m1} & c_{m2} & \cdots & c_{mp} \end{bmatrix}$$

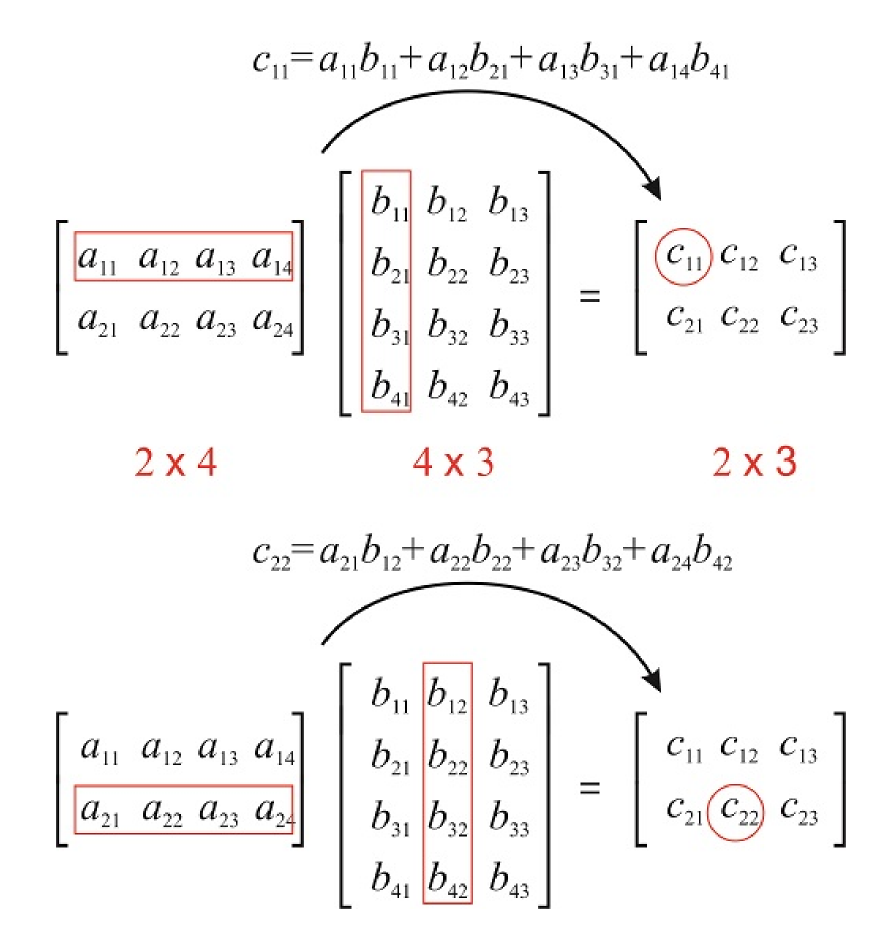
where each element  $c_{ij}$  is the dot product of the ith row of A and jth column of B:

$$c_{ij} = \langle A_{i,:} \cdot B_{:,j} 
angle = a_{i1}b_{1j} + a_{i2}b_{2j} + \dots + a_{in}b_{nj} = \sum_{k=1}^n a_{ik}b_{kj}$$

**Note:** n, the number of columns of A and rows of B, must be the same!

## **Matrix multiplication**

ullet The element  $c_{ij}$  of C=A imes B is the dot product between the ith row of A and the jth column of B .



Source:

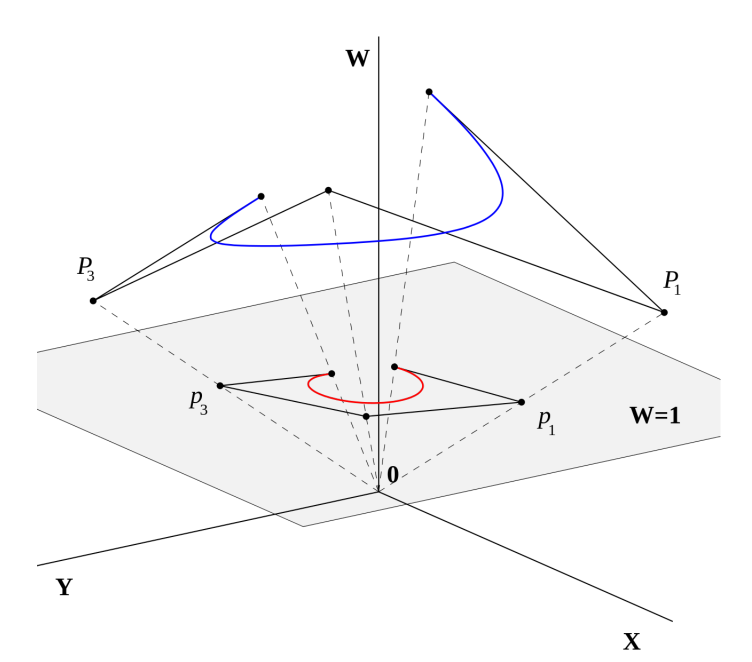
https://chem.libretexts.org/Bookshelves/Physical\_and\_Theoretical\_Chemistry\_Textbook\_Maps/Book%3A\_Mathematical\_Methods\_in\_Chemistry\_(Levitus)/15%3A\_CC BY-NC-SA; Marcia Levitus

## Matrix-vector multiplication

ullet Thinking of vectors as n imes 1 matrices, we can multiply a matrix m imes n with a vector:

$$\mathbf{y} = A imes \mathbf{x} = egin{bmatrix} a_{11} & a_{12} & \cdots & a_{1n} \ a_{21} & a_{22} & \cdots & a_{2n} \ dots & dots & \ddots & dots \ a_{m1} & a_{m2} & \cdots & a_{mn} \end{bmatrix} imes egin{bmatrix} x_1 \ x_2 \ dots \ x_n \end{bmatrix} = egin{bmatrix} y_1 \ y_2 \ dots \ y_m \end{bmatrix}$$

- The result  $\mathbf{y}$  is a vector of size m.
- In that sense, a matrix A can transform a vector of size n into a vector of size m:
  - A represents a **projection** from  $\Re^n$  to  $\Re^m$ .



#### Source:

https://en.wikipedia.org/wiki/Homogeneous\_coordinate

## **Dot product**

• Note that the **dot product** between two vectors of size n is the matrix multiplication between the transpose of the first vector and the second one:

$$\mathbf{x}^T imes \mathbf{y} = egin{bmatrix} x_1 & x_2 & \dots & x_n \end{bmatrix} imes egin{bmatrix} y_1 \ y_2 \ dots \ y_n \end{bmatrix} = x_1 \ y_1 + x_2 \ y_2 + \dots + x_n \ y_n = \langle \mathbf{x} \cdot \mathbf{y} 
angle$$

#### **Matrix inversion**

ullet Square matrices of size n imes n can be inverted. The **inverse**  $A^{-1}$  of a matrix A is defined by:

$$A imes A^{-1} = A^{-1} imes A = I$$

where I is the identity matrix (a matrix with ones on the diagonal and 0 otherwise).

• Matrix inversion allows to solve linear systems of equations. Given the problem:

$$egin{cases} a_{11}\,x_1+a_{12}\,x_2+\ldots+a_{1n}\,x_n=b_1\ a_{21}\,x_1+a_{22}\,x_2+\ldots+a_{2n}\,x_n=b_2\ \ldots\ a_{n1}\,x_1+a_{n2}\,x_2+\ldots+a_{nn}\,x_n=b_n \end{cases}$$

which is equivalent to:

$$A \times \mathbf{x} = \mathbf{b}$$

ullet We can multiply both sides to the left with  $A^{-1}$  (if it exists) and obtain:

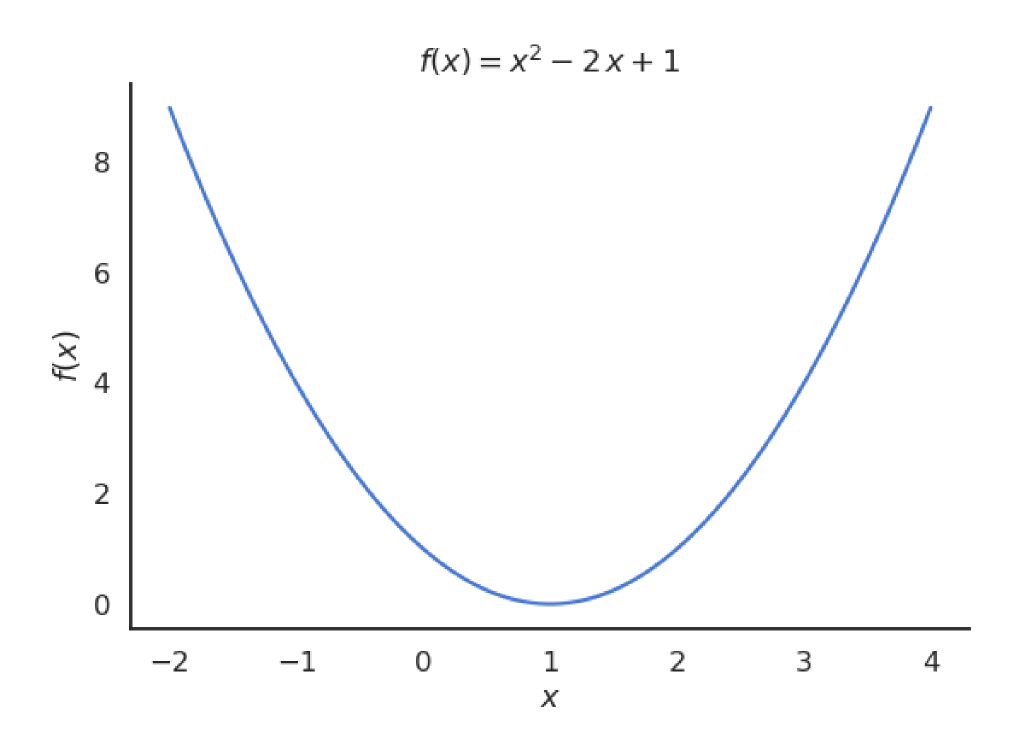
$$\mathbf{x} = A^{-1} \times \mathbf{b}$$

## 2 - Calculus

#### **Univariate functions**

ullet A **univariate function** f associates to any real number  $x\in\Re$  (or a subset of  $\Re$  called the support of the function) another (unique) real number f(x):

$$x\mapsto f(x),$$
 (2)



#### **Multivariate functions**

• A multivariate function f associates to any vector  $\mathbf{x} \in \Re^n$  (or a subset) a real number  $f(\mathbf{x})$ :

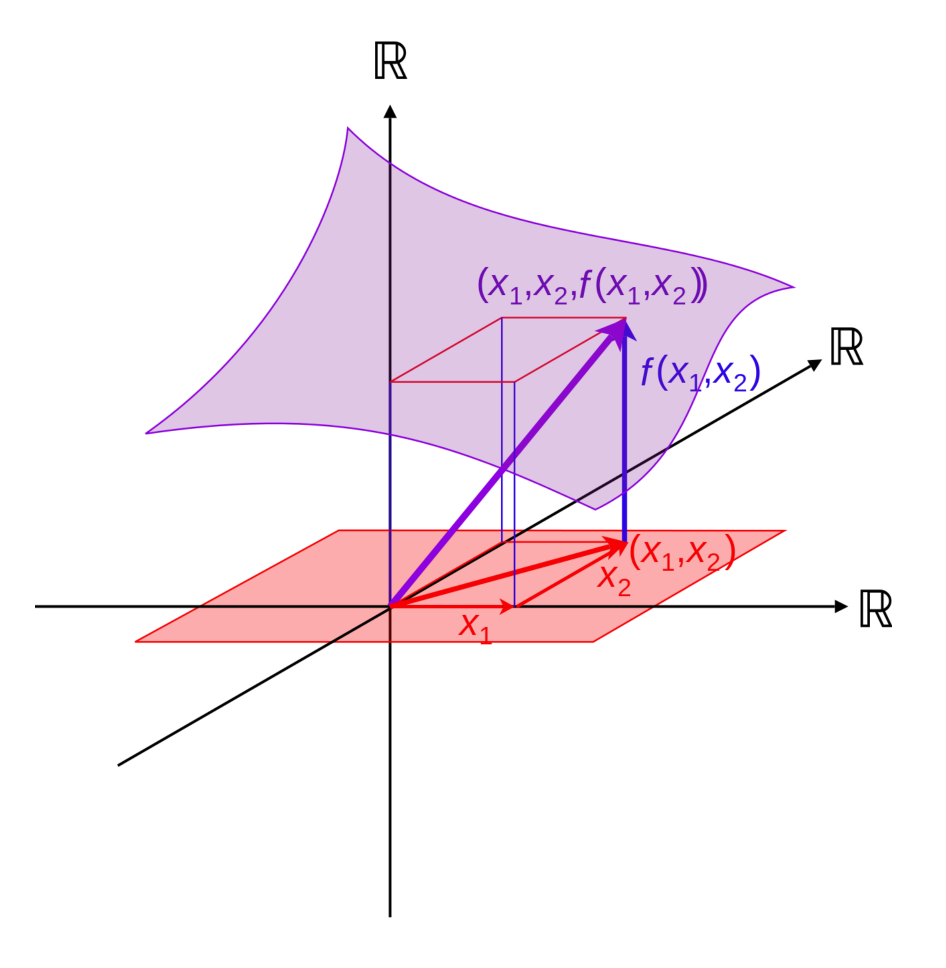
$$f: \quad \Re^n o \Re$$
 (3)

$$\mathbf{x}\mapsto f(\mathbf{x}), \qquad \qquad (4)$$

- The variables of the function are the elements of the vector.
- For low-dimensional vector spaces, it is possible to represent each element explicitly, for example:

$$f\colon \quad \Re^3 o \Re$$
 (5)

$$x, y, z \mapsto f(x, y, z),$$
 (6)



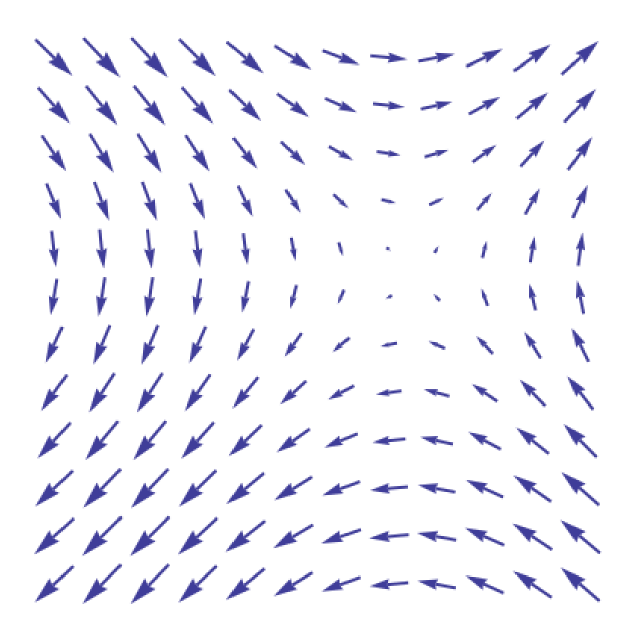
Source: https://en.wikipedia.org/wiki/Function\_of\_several\_real\_variables

#### **Vector fields**

• **Vector fields** associate to any vector  $\mathbf{x} \in \Re^n$  (or a subset) another vector (possibly of different size):

$$\overrightarrow{f} : \Re^n \to \Re^m$$
 (7)

$$\mathbf{x} \mapsto \overrightarrow{f}(\mathbf{x}),$$
 (8)



Source: https://en.wikipedia.org/wiki/Vector\_field

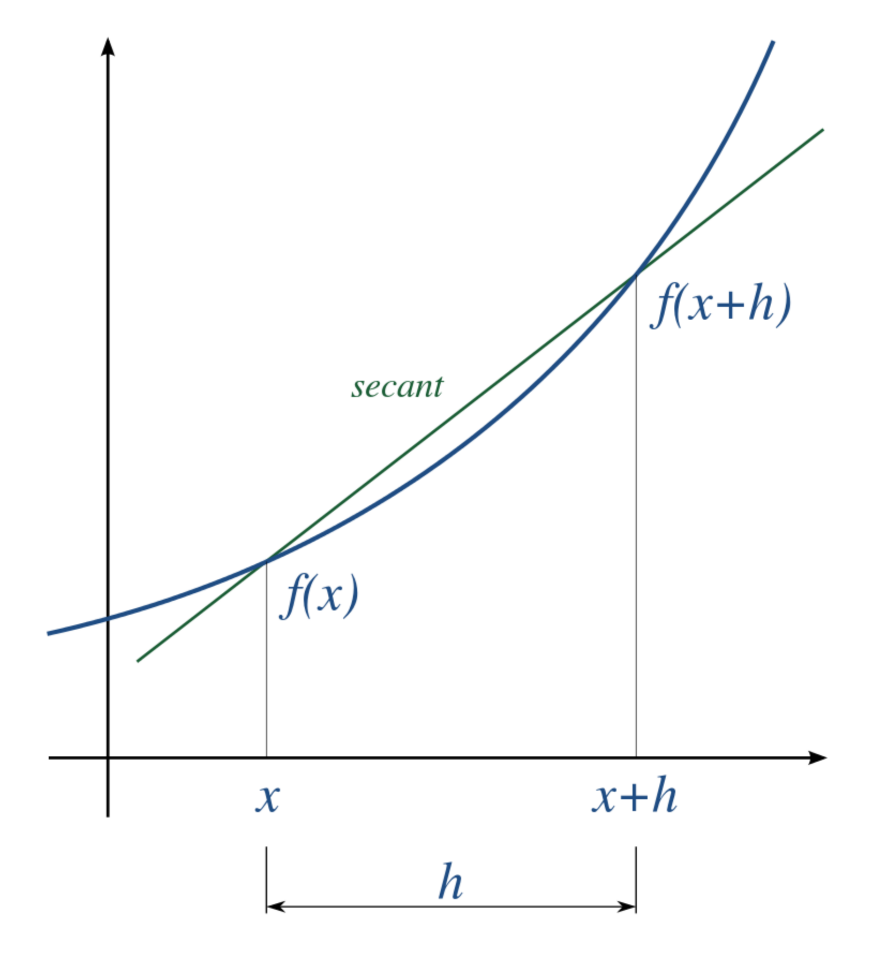
**Note:** The matrix-vector multiplication  $\mathbf{y} = A \times \mathbf{x}$  is a linear vector field, mapping any vector  $\mathbf{x}$  into another vector  $\mathbf{y}$ .

#### Differentiation

- Differential calculus deals with the **derivative** of a function, a process called differentiation.
- The derivative f'(x) or  $\frac{df(x)}{dx}$  of a univariate function f(x) is defined as the local slope of the tangent to the function for a given value of x:

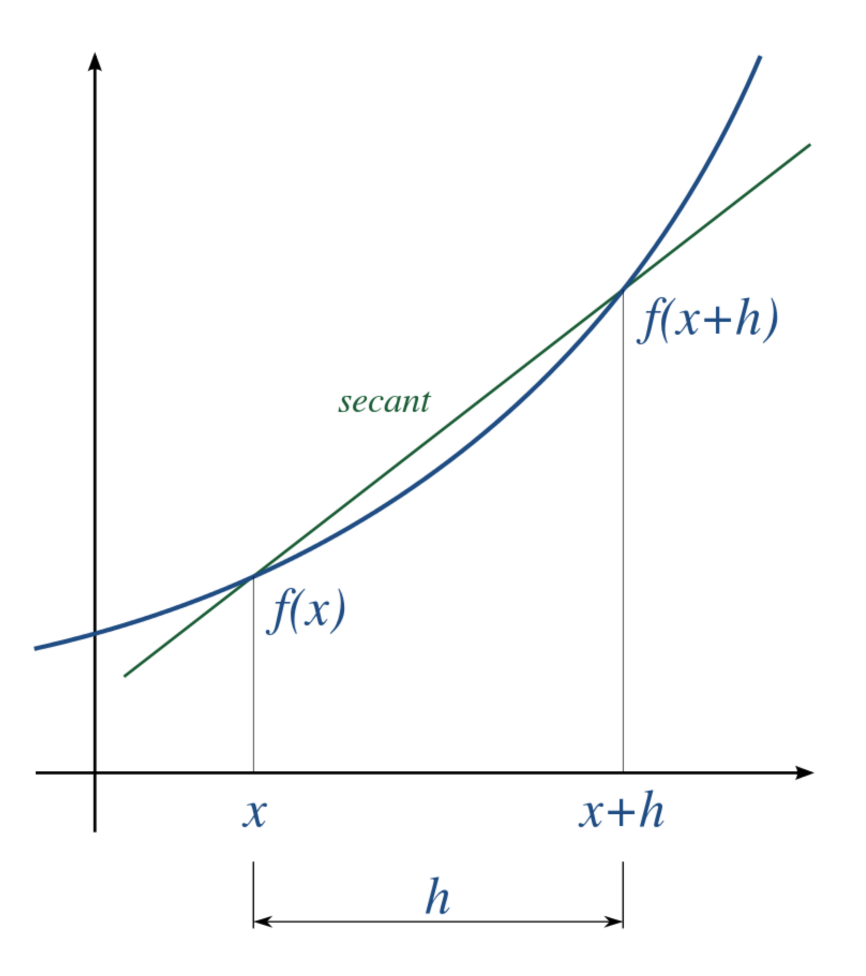
$$f'(x) = \lim_{h o 0} rac{f(x+h)-f(x)}{h}$$

• The line passing through the points (x,f(x)) and (x+h,f(x+h)) becomes tangent to the function when h becomes very small.



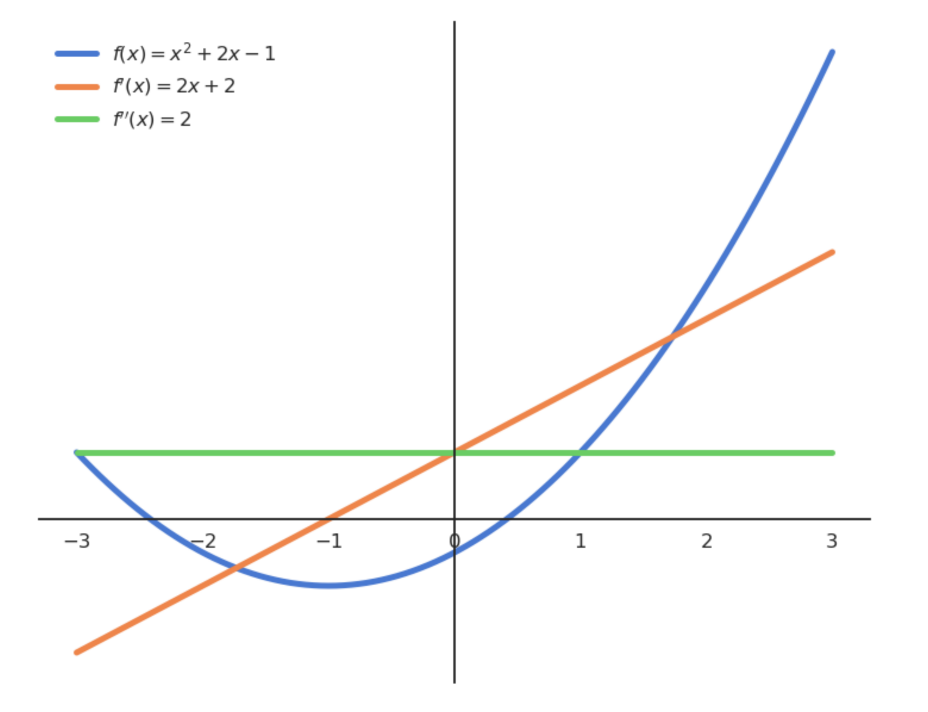
#### Differentiation

- The sign of the derivative tells you how the function behaves locally:
  - If the derivative is positive, increasing a little bit x increases the function f(x), so the function is **locally increasing**.
  - If the derivative is negative, increasing a little bit x decreases the function f(x), so the function is **locally decreasing**.
- It basically allows you to measure the local influence of x on f(x): if I change a little bit the value x, what happens to f(x)? This will be very useful in machine learning.



#### **Extrema**

- A special case is when the derivative is equal to 0 in x. x is then called an **extremum** (or optimum) of the function, i.e. it can be a maximum or minimum.
- You can tell whether an extremum is a maximum or a minimum by looking at its second-order derivative:
  - If f''(x) > 0, the extremum is a **minimum**.
  - If f''(x) < 0, the extremum is a **maximum**.
  - If f''(x) = 0, the extremum is a saddle point.



#### **Gradients**

• The derivative of a multivariate function  $f(\mathbf{x})$  is a vector of partial derivatives called the gradient of the function  $\nabla_{\mathbf{x}} f(\mathbf{x})$ :

$$abla_{\mathbf{x}} f(\mathbf{x}) = egin{bmatrix} rac{\partial f(\mathbf{x})}{\partial x_1} \ rac{\partial f(\mathbf{x})}{\partial x_2} \ rac{\partial f(\mathbf{x})}{\partial x_n} \end{bmatrix}$$

• The subscript to the  $\nabla$  operator denotes with respect to (w.r.t) which variable the differentiation is done.

#### **Partial derivatives**

A partial derivative w.r.t. to particular variable (or element of the vector) is simply achieved by
differentiating the function while considering all other variables to be constant. For example the function:

$$f(x,y) = x^2 + 3 x y + 4 x y^2 - 1$$

can be partially differentiated w.r.t. x and y as:

$$egin{cases} rac{\partial f(x,y)}{\partial x} = 2\,x + 3\,y + 4\,y^2 \ rac{\partial f(x,y)}{\partial y} = 3\,x + 8\,x\,y \end{cases}$$

#### Jacobian

• The gradient can be generalized to **vector fields**, where the **Jacobian** or **Jacobi matrix** is a matrix containing all partial derivatives.

$$J = egin{bmatrix} rac{\partial \mathbf{f}}{\partial x_1} & \cdots & rac{\partial \mathbf{f}}{\partial x_n} \end{bmatrix} = egin{bmatrix} rac{\partial f_1}{\partial x_1} & \cdots & rac{\partial f_1}{\partial x_n} \ dots & \ddots & dots \ rac{\partial f_m}{\partial x_1} & \cdots & rac{\partial f_m}{\partial x_n} \end{bmatrix}$$

## **Analytical properties**

• Differentiation is linear, which means that if we define the function:

$$h(x) = a f(x) + b g(x)$$

its derivative is:

$$h'(x) = a f'(x) + b g'(x)$$

• A product of functions can also be differentiated analytically (product rule):

$$(f(x) imes g(x))'=f'(x) imes g(x)+f(x) imes g'(x)$$

**Example:** 

$$f(x) = x^2 \, e^x$$

$$f'(x) = 2 x e^x + x^2 \cdot e^x$$

#### Chain rule

 A very important concept for neural networks is the chain rule, which tells how to differentiate function compositions (functions of a function) of the form:

$$(f\circ g)(x)=f(g(x))$$

• The derivative of  $f\circ g$  is:

$$(f\circ g)'(x)=(f'\circ g)(x) imes g'(x)$$

• The chain rule may be more understandable using Leibniz's notation:

$$rac{d(f\circ g)(x)}{dx} = rac{df(g(x))}{dg(x)} imes rac{dg(x)}{dx}$$

ullet By posing y=g(x) as an intermediary variable, it becomes:

$$rac{df(y)}{dx} = rac{df(y)}{dy} imes rac{dy}{dx}$$

#### Chain rule

• The function:

$$h(x) = \frac{1}{2\,x+1}$$

is the function composition of  $g(x)=2\,x+1$  and  $f(x)=rac{1}{x}$  , whose derivatives are known:

$$g'(x)=2$$

$$f'(x) = -rac{1}{x^2}$$

• Its derivative according to the **chain rule** is:

$$h'(x) = f'(g(x)) imes g'(x) = -rac{1}{(2\,x+1)^2} imes 2$$

### **Chain rule**

• The chain rule also applies to partial derivatives:

$$rac{\partial f \circ g(x,y)}{\partial x} = rac{\partial f \circ g(x,y)}{\partial g(x,y)} imes rac{\partial g(x,y)}{\partial x}$$

and gradients:

$$abla_{\mathbf{x}} \, f \circ g(\mathbf{x}) = 
abla_{g(\mathbf{x})} \, f \circ g(\mathbf{x}) imes 
abla_{\mathbf{x}} \, g(\mathbf{x})$$

• The opposite operation of differentiation is **integration**. Given a function f(x), we search a function F(x) whose *derivative* is f(x):

$$F'(x) = f(x)$$

• The **integral** of f is noted:

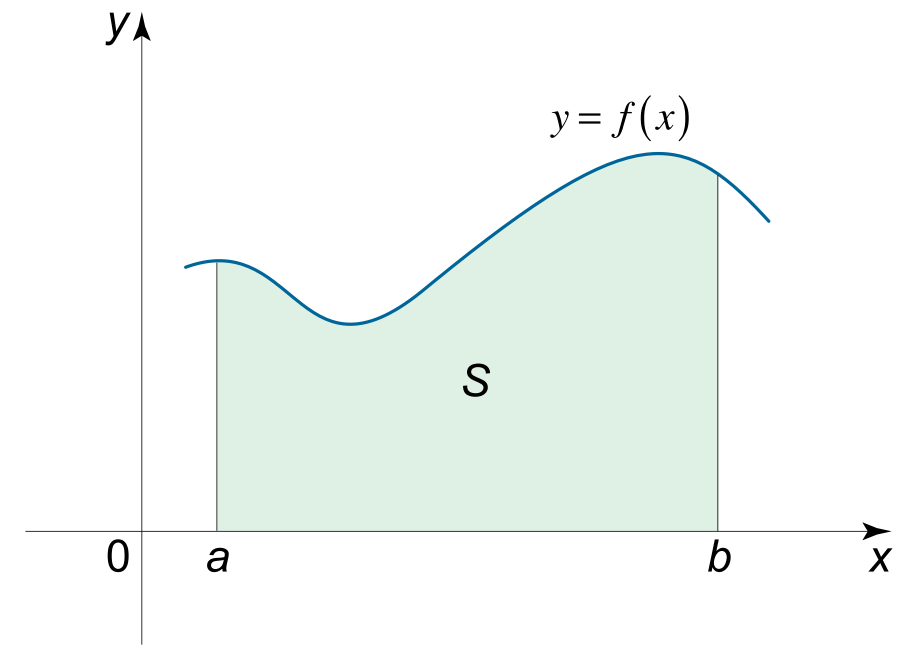
$$F(x) = \int f(x) \, dx$$

dx being an infinitesimal interval (similar to h in the definition of the derivative).

• There are tons of formal definitions of integrals (Riemann, Lebesgue, Darboux...) and we will not get into details here as we will not use integrals a lot.

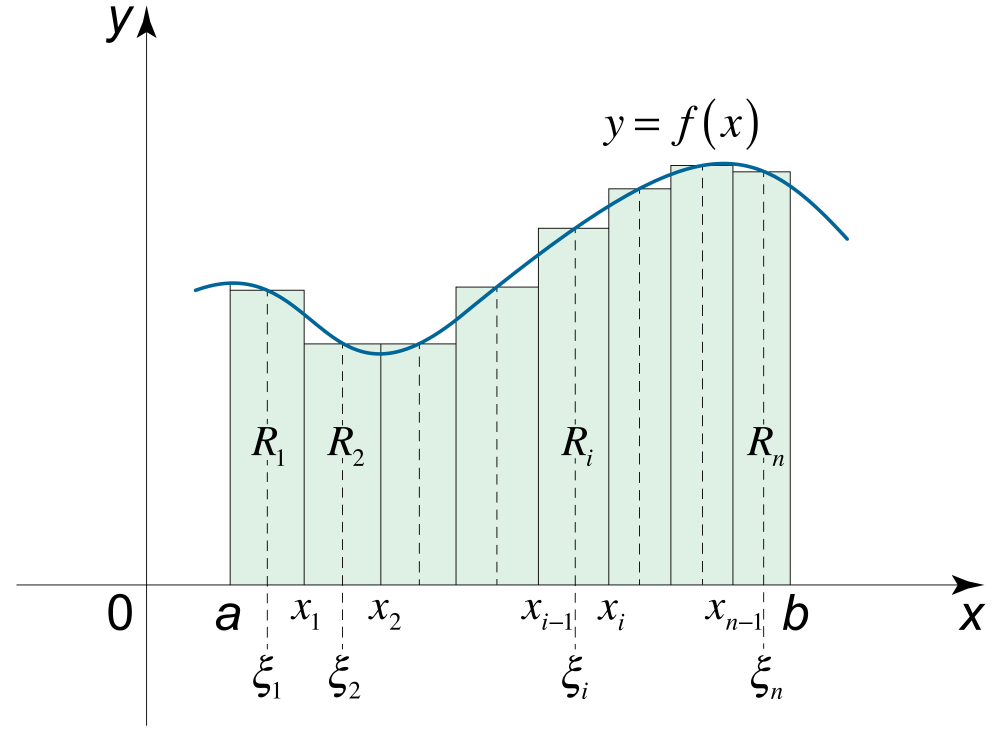
- The most important to understand for now is maybe that the integral of a function is the **area under the curve**.
- The area under the curve of a function f on the interval [a,b] is:

$$\mathcal{S} = \int_a^b f(x) \, dx$$



Source: https://www.math24.net/riemann-sums-definite-integral/

- One way to approximate this surface is to split the interval [a,b] into n intervals of width dx with the points  $x_1,x_2,\ldots,x_n$ .
- ullet This defines n rectangles of width dx and height  $f(x_i)$ , so their surface is  $f(x_i)\,dx$ .
- The area under the curve can then be approximated by the sum of the surfaces of all these rectangles.

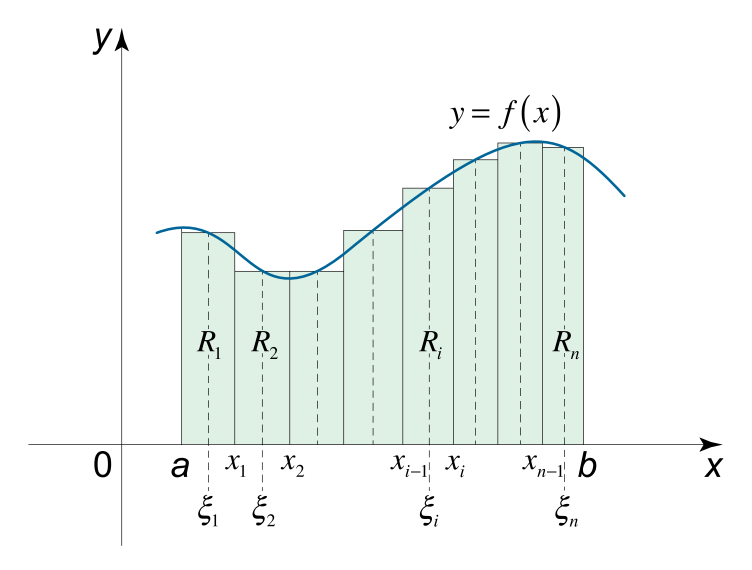


Source: https://www.math24.net/riemann-sums-definite-integral/

• When  $n \to \infty$ , or equivalently  $dx \to 0$ , the sum of these rectangular areas (called the Riemann sum) becomes exactly the area under the curve. This is the definition of the definite integral:

$$\int_a^b f(x)\,dx = \lim_{dx o 0} \sum_{i=1}^n f(x_i)\,dx$$

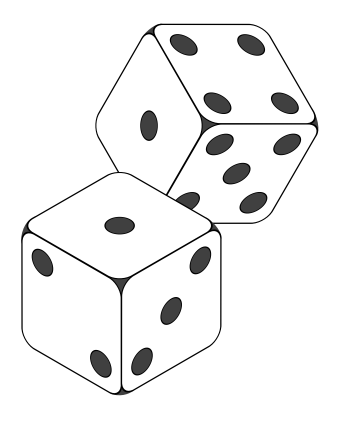
• Very roughly speaking, the integral can be considered as the equivalent of a sum for continuous functions.



Source: https://www.math24.net/riemann-sums-definite-integral/

# 3 - Probability theory

## Discrete probability distributions



Credit:
https://commons.wikimedia.org/wiki/File:2-Dice-Icon.svg

- Let's note X a **discrete random variable** with n realizations (or outcomes)  $x_1, \ldots, x_n$ .
- The **probability** that X takes the value  $x_i$  is defined by the *relative* frequency of occurrence, i.e. the proportion of samples having the value  $x_i$ , when the total number N of samples tends to infinity:

$$P(X = x_i) = rac{ ext{Number of favorable cases}}{ ext{Total number of samples}}$$

- The set of probabilities  $\{P(X=x_i)\}_{i=1}^n$  define the **probability distribution** for the random variable (or probability mass function, pmf).
- ullet By definition, we have  $0 \leq P(X=x_i) \leq 1$  and the probabilities **have** to respect:

$$\sum_{i=1}^n P(X=x_i)=1$$

## Mathematical expectation and variance

• An important metric for a random variable is its **mathematical expectation** or expected value, i.e. its "mean" realization weighted by the probabilities:

$$\mathbb{E}[X] = \sum_{i=1}^n P(X=x_i)\,x_i$$

The expectation does not even need to be a valid realization:

$$\mathbb{E}[ ext{Coin}] = rac{1}{2}\,0 + rac{1}{2}\,1 = 0.5$$
  $\mathbb{E}[ ext{Dice}] = rac{1}{6}\,(1+2+3+4+5+6) = 3.5$ 

• We can also compute the mathematical expectation of **functions of** a random variable:

$$\mathbb{E}[f(X)] = \sum_{i=1}^n P(X=x_i)\,f(x_i)$$

## Mathematical expectation and variance

• The **variance** of a random variable is the squared deviation around the mean:

$$\operatorname{Var}(X) = \mathbb{E}[(X - \mathbb{E}[X])^2] = \sum_{i=1}^n P(X = x_i) \left(x_i - \mathbb{E}[X]
ight)^2$$

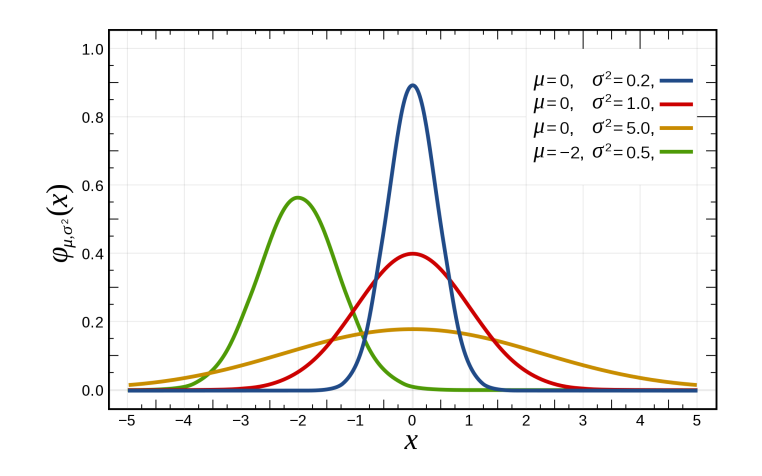
Variance of a coin:

$$\mathrm{Var}(\mathrm{Coin}) = rac{1}{2} \left( 0 - 0.5 \right)^2 + rac{1}{2} \left( 1 - 0.5 \right)^2 = 0.25$$

• Variance of a dice:

$$\mathrm{Var}(\mathrm{Dice}) = \frac{1}{6} \left( (1 - 3.5)^2 + (2 - 3.5)^2 + (3 - 3.5)^2 + (4 - 3.5)^2 + (5 - 3.5)^2 + (6 - 3.5)^2 \right) = \frac{105}{36}$$

## Continuous probability distributions



Source: https://en.wikipedia.org/wiki/Normal\_distribution

- Continuous random variables can take an infinity of continuous values, e.g.  $\Re$  or some subset.
- The closed set of values they can take is called the **support**  $\mathcal{D}_X$  of the probability distribution.
- The probability distribution is described by a **probability** density function (pdf) f(x).
- The pdf of a distribution must be positive ( $f(x) \geq 0 \ orall x \in \mathcal{D}_X$ ) and its integral must be equal to 1:

$$\int_{x\in\mathcal{D}_X}f(x)\,dx=1$$

• The pdf does not give the probability of taking a particular value x (it is 0), but allows to get the probability that a value lies in a specific interval:

$$P(a \leq X \leq b) = \int_a^b f(x) \, dx$$

ullet One can however think of the pdf as the **likelihood** that a value x comes from that distribution.

## Expectation and variance of continuous distributions

The mathematical expectation is now defined by an integral instead of a sum:

$$\mathbb{E}[X] = \int_{x \in \mathcal{D}_X} f(x) \, x \, dx$$

the variance:

$$\operatorname{Var}(X) = \int_{x \in \mathcal{D}_X} f(x) \, (x - \mathbb{E}[X])^2 \, dx$$

or a function of the random variable:

$$\mathbb{E}[g(X)] = \int_{x \in \mathcal{D}_X} f(x) \, g(x) \, dx$$

Note that the expectation operator is linear:

$$\mathbb{E}[a\,X+b\,Y]=a\,\mathbb{E}[X]+b\,\mathbb{E}[Y]$$

## Some parameterized probability distributions

- Probability distributions can in principle have any form: f(x) is unknown.
- However, specific parameterized distributions can be very useful: their pmf/pdf is fully determined by a couple of parameters.
- The **Bernouilli** distribution is a binary (discrete, 0 or 1) distribution with a parameter p specifying the probability to obtain the outcome 1:

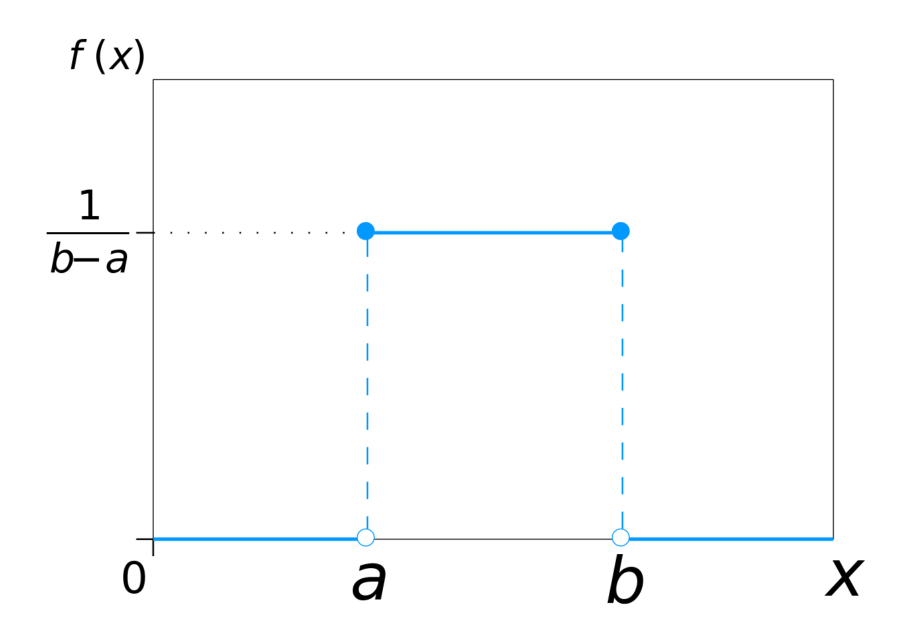
$$P(X=1)=p ext{ and } P(X=0)=1-p$$
  $P(X=x)=p^x \, (1-p)^{1-x}$   $\mathbb{E}[X]=p$ 

• The **Multinouilli** or **categorical** distribution is a discrete distribution with k realizations. Each realization  $x_i$  is associated with a parameter  $p_i>0$  representing its probability. We have  $\sum_i p_i=1$ .

$$P(X=x_i)=p_i$$

ullet Knowing p or the  $p_i$  tells us everything about the discrete distributions.

### The uniform distribution



- The **uniform distribution** has an equal and constant probability of returning values between a and b, never outside this range.
- It is parameterized by two parameters:
  - the start of the range a.
  - the end of the range b.
- Its support is [a,b].

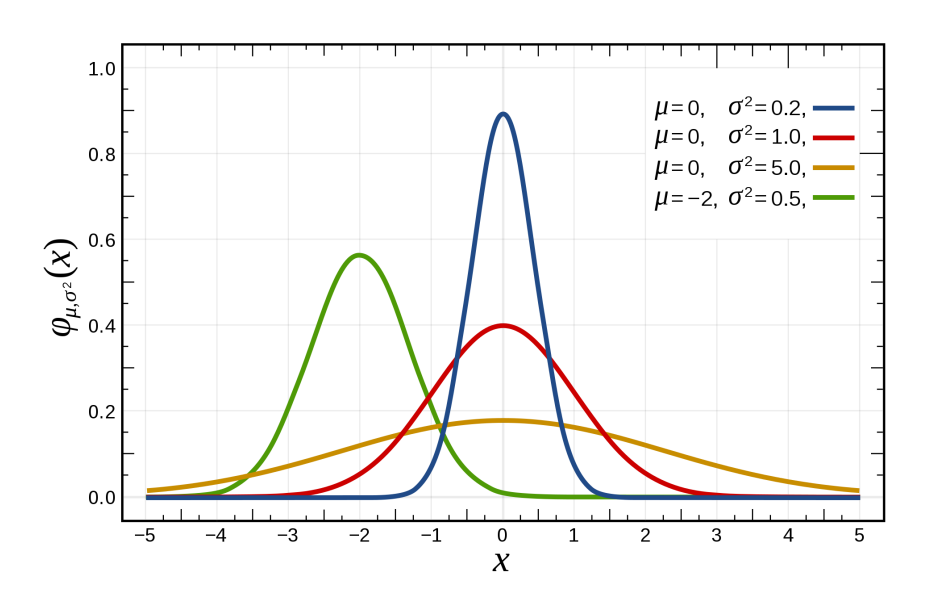
Credit: https://en.wikipedia.org/wiki/Uniform\_distribution\_(continuous)

ullet The pdf of the uniform distribution  $\mathcal{U}(a,b)$  is defined on [a,b] as:

$$f(x;a,b)=rac{1}{b-a}$$

ullet Knowing a and b completely defines the distribution.

### The normal or Gaussian distribution



- For continuous distributions, the normal distribution is the most frequently encountered one.
- It is parameterized by two parameters:
  - the mean  $\mu$ .
  - the variance  $\sigma^2$  (or standard deviation  $\sigma$ ).
- Its support is  $\Re$ .

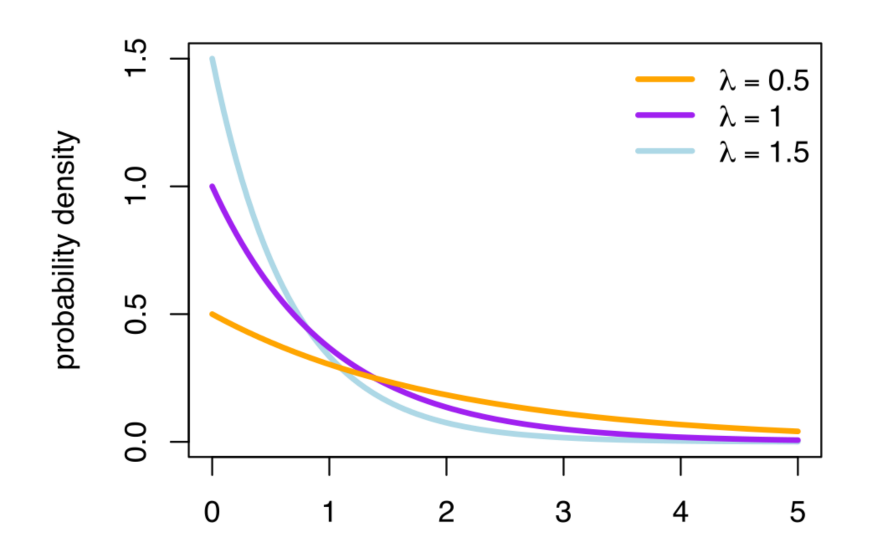
Credit: https://en.wikipedia.org/wiki/Normal\_distribution

• The pdf of the normal distribution  $\mathcal{N}(\mu,\sigma)$  is defined on  $\Re$  as:

$$f(x;\mu,\sigma) = rac{1}{\sqrt{2\,\pi\,\sigma^2}}\,e^{-rac{(x-\mu)^2}{2\,\sigma^2}}$$

• Knowing  $\mu$  and  $\sigma$  completely defines the distribution.

### The exponential distribution



- The **exponential distribution** is the probability distribution of the time between events in a Poisson point process, i.e., a process in which events occur continuously and independently at a constant average rate.
- It is parameterized by one parameter:
  - the rate  $\lambda$ .
- Its support is  $\Re^+$  (x>0).

Credit: https://en.wikipedia.org/wiki/Exponential\_distribution

ullet The pdf of the exponential distribution is defined on  $\Re^+$  as:

$$f(x;\lambda) = \lambda \, e^{-\lambda \, x}$$

ullet Knowing  $\lambda$  completely defines the distribution.

### Joint probabilities

- Let's now suppose that we have two random variables X and Y with different probability distributions P(X) and P(Y).
- The **joint probability** P(X,Y) denotes the probability of observing the realizations x **and** y at the same time:

$$P(X=x,Y=y)$$

• If the random variables are **independent**, we have:

$$P(X = x, Y = y) = P(X = x) P(Y = y)$$

• If you know the joint probability, you can compute the marginal probability distribution of each variable:

$$P(X = x) = \sum_{y} P(X = x, Y = y)$$

• The same is true for continuous probability distributions:

$$f(x) = \int f(x,y) \, dy$$

### **Conditional probabilities**

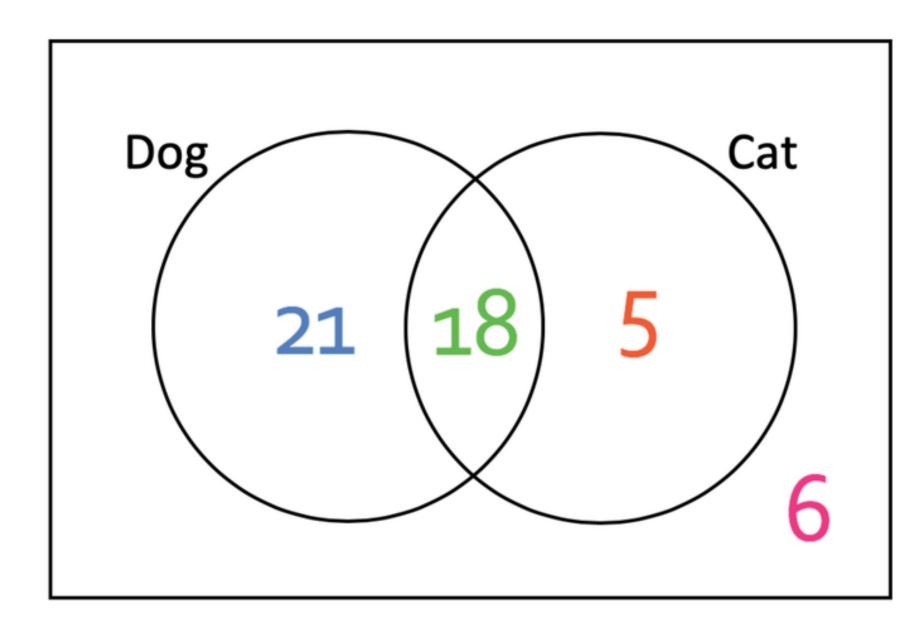
- Some useful information between two random variables is the conditional probability.
- ullet P(X=x|Y=y) is the conditional probability that X=x, **given** that Y=y is observed.
- Y=y is not random anymore: it is a **fact** (at least theoretically).
- ullet You wonder what happens to the probability distribution of X now that you know the value of Y .
- Conditional probabilities are linked to the joint probability by:

$$P(X=x|Y=y)=rac{P(X=x,Y=y)}{P(Y=y)}$$

- If X and Y are **independent**, we have P(X=x|Y=y)=P(X=x) (knowing Y does not change anything to the probability distribution of X).
- We can use the same notation for the complete probability distributions:

$$P(X|Y) = rac{P(X,Y)}{P(Y)}$$

## Joint and conditional probabilities: using a Venn diagram



Credit: https://www.elevise.co.uk/g-e-m-h-5-u.html

- You ask 50 people whether they like cats or dogs:
  - 18 like both cats and dogs.
  - 21 like only dogs.
  - 5 like only cats.
  - 6 like none of them.
- We consider loving cats and dogs as random variables (and that our sample size is big enough to use probabilities...)
- ullet We have  $P(\mathrm{dog})=rac{18+21}{50}$  and  $P(\mathrm{cat})=rac{18+5}{50}$ .
- Among the 23 who love cats, which proportion also loves dogs?
- The joint probability of loving both cats and dogs is  $P(\mathrm{cat},\mathrm{dog})=rac{18}{50}$ .
- The conditional probability of loving dogs given one loves cats is:

$$P( ext{dog}| ext{cat}) = rac{P( ext{cat}, ext{dog})}{P( ext{cat})} = rac{rac{18}{50}}{rac{23}{50}} = rac{18}{23}$$

## Bayes' rule

Noticing that the definition of conditional probabilities is symmetric:

$$P(X,Y) = P(X|Y) P(Y) = P(Y|X) P(X)$$

we can obtain the **Bayes' rule**:

$$P(Y|X) = rac{P(X|Y)P(Y)}{P(X)}$$

- ullet It is very useful when you already know P(X|Y) and want to obtain P(Y|X) (Bayesian inference).
  - P(Y|X) is called the **posterior probability**.
  - P(X|Y) is called the **likelihood**.
  - P(Y) is called the **prior probability** (belief).
  - ullet P(X) is called the **model evidence** or **marginal likelihood**.

### Bayes' rule: example

• Let's consider a disease D (binary random variable) and a medical test T (also binary). The disease affects 10% of the general population:

$$P(D=1) = 0.1$$
  $P(D=0) = 0.9$ 

• When a patient has the disease, the test is positive 80% of the time:

$$P(T=1|D=1)=0.8$$
  $P(T=0|D=1)=0.2$ 

• When a patient does not have the disease, the test is still positive 10% of the time:

$$P(T=1|D=0)=0.1$$
  $P(T=0|D=0)=0.9$ 

Given that the test is positive, what is the probability that the patient is ill?

### Bayes' rule: example

$$P(D = 1|T = 1) = \frac{P(T = 1|D = 1) P(D = 1)}{P(T = 1)}$$

$$= \frac{P(T = 1|D = 1) P(D = 1)}{P(T = 1|D = 1) P(D = 1) + P(T = 1|D = 0) P(D = 0)}$$

$$= \frac{0.8 \times 0.1}{0.8 \times 0.1 + 0.1 \times 0.9}$$

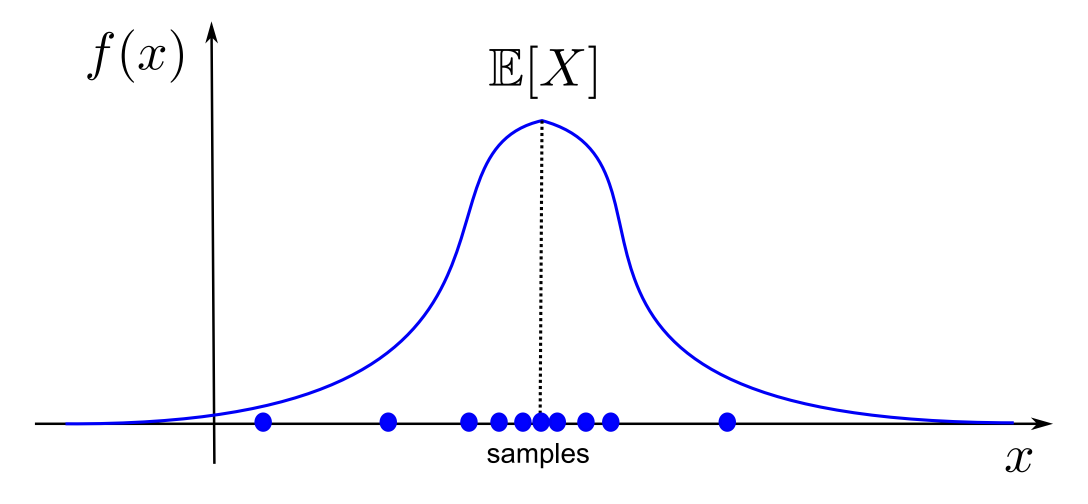
$$= 0.47$$

## 4 - Statistics

## Random sampling / Monte Carlo sampling

- In ML, we will deal with random variables whose exact probability distribution is unknown, but we are interested in their expectation or variance anyway.
- Random sampling or Monte Carlo sampling (MC) consists of taking N samples  $x_i$  out of the distribution X (discrete or continuous) and computing the sample average:

$$\mathbb{E}[X] = \mathbb{E}_{x \sim X}[x] pprox rac{1}{N} \, \sum_{i=1}^N x_i$$



• More samples will be obtained where f(x) is high (x is probable), so the average of the sampled data will be close to the expected value of the distribution.

### Random sampling / Monte Carlo sampling

#### Law of big numbers

As the number of identically distributed, randomly generated variables increases, their sample mean (average) approaches their theoretical mean.

MC estimates are only correct when:

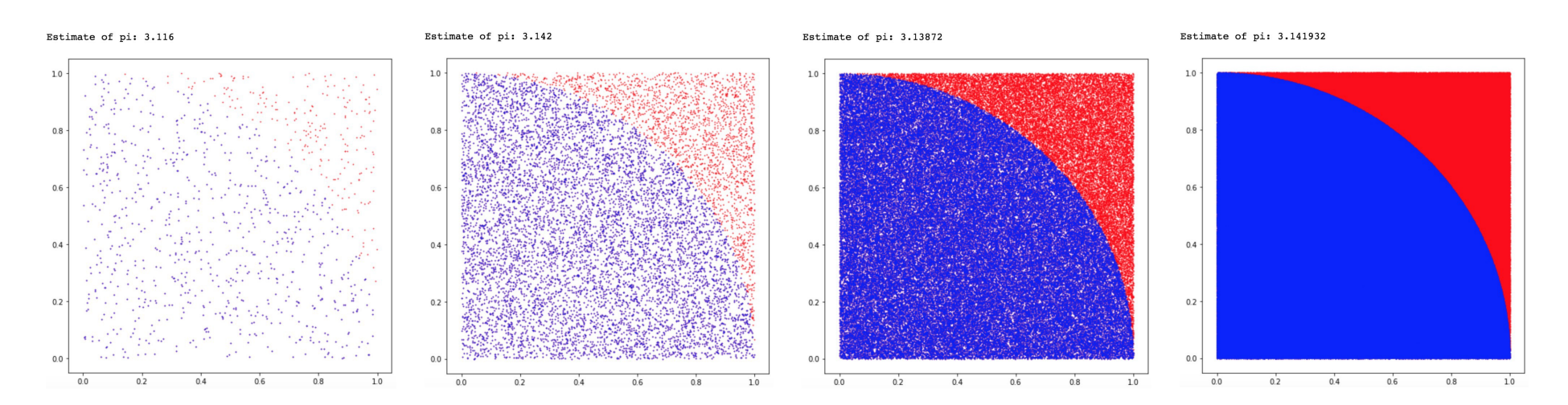
- the samples are i.i.d (independent and identically distributed):
  - independent: the samples must be unrelated with each other.
  - ullet identically distributed: the samples must come from the same distribution X.
- ullet the number of samples is large enough. Usually N>30 for simple distributions.

## Random sampling / Monte Carlo sampling

• One can estimate any function of the random variable with random sampling:

$$\mathbb{E}[f(X)] = \mathbb{E}_{x \sim X}[f(x)] pprox rac{1}{N} \, \sum_{i=1}^N f(x_i)$$

• Example of Monte Carlo sampling to estimate  $\pi/4$ :



### **Central limit theorem**

- ullet Suppose we have an unknown distribution X with expected value  $\mu=\mathbb{E}[X]$  and variance  $\sigma^2$  .
- ullet We can take randomly N samples from X to compute the sample average:

$$S_N = rac{1}{N} \, \sum_{i=1}^N x_i$$

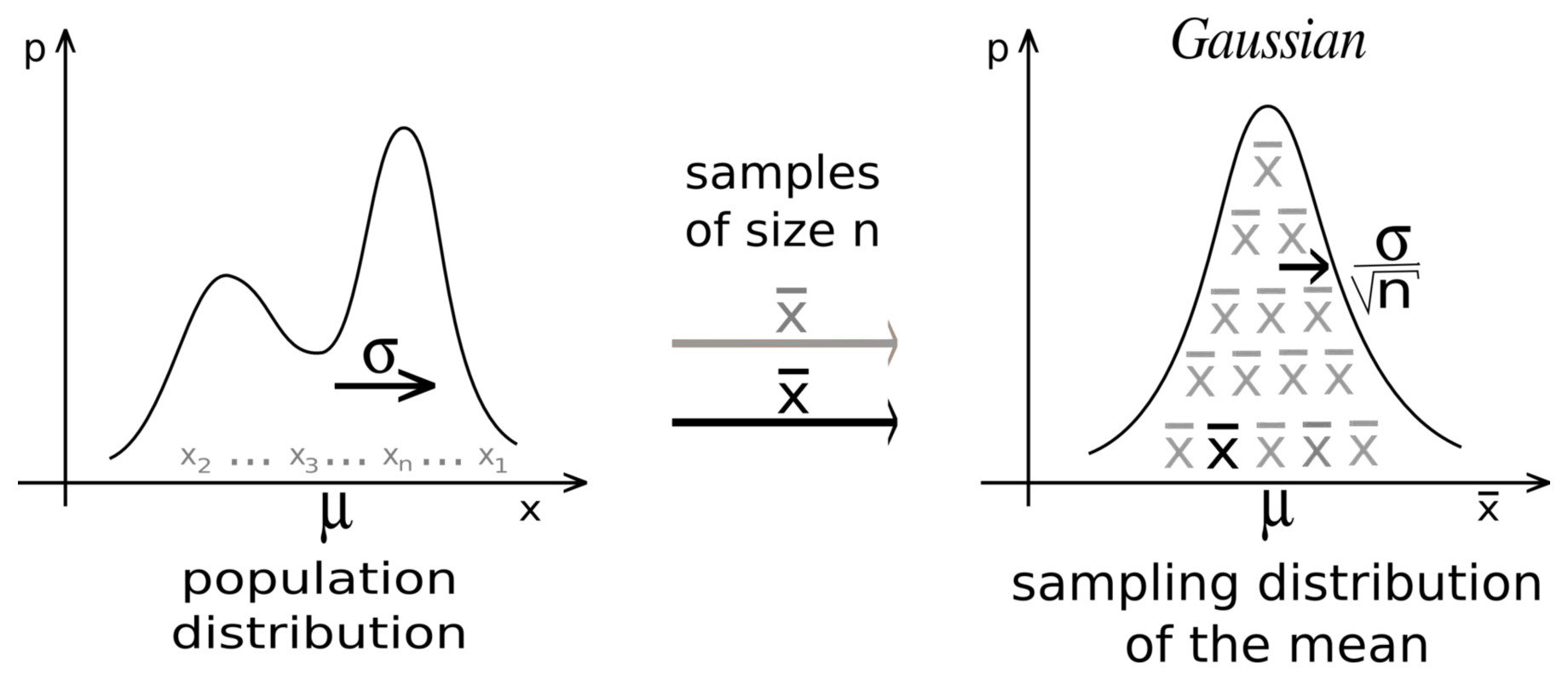
• The **Central Limit Theorem** (CLT) states that:

The distribution of sample averages is normally distributed with mean  $\mu$  and variance  $rac{\sigma^2}{N}$  .

$$S_N \sim \mathcal{N}(\mu, rac{\sigma}{\sqrt{N}})$$

### **Central limit theorem**

- If we perform the sampling multiple times, even with few samples, the average of the sampling averages will be very close to the expected value.
- The more samples we get, the smaller the variance of the estimates.
- ullet Although the distribution X can be anything, the sampling averages are normally distributed.



Credit: https://en.wikipedia.org/wiki/Central\_limit\_theorem

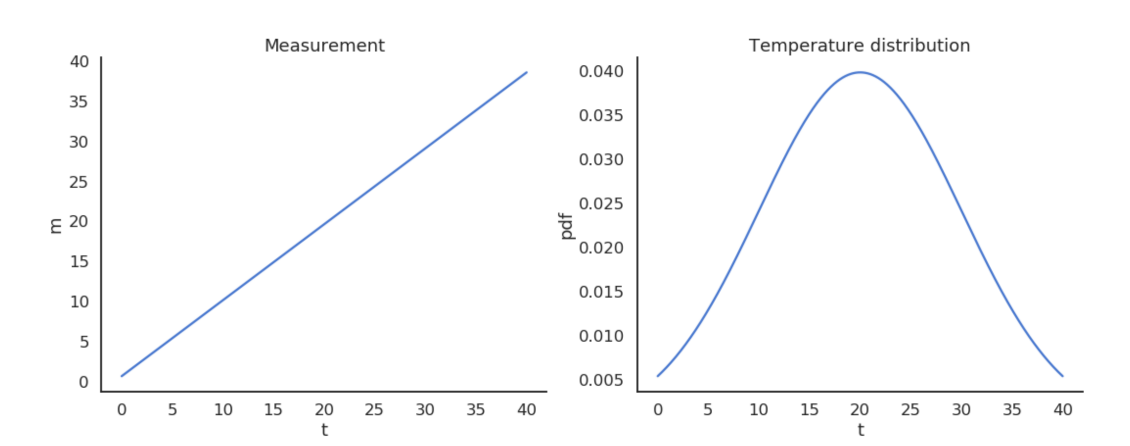
### **Estimators**

• CLT shows that the sampling average is an unbiased estimator of the expected value of a distribution:

$$\mathbb{E}(S_N) = \mathbb{E}(X)$$

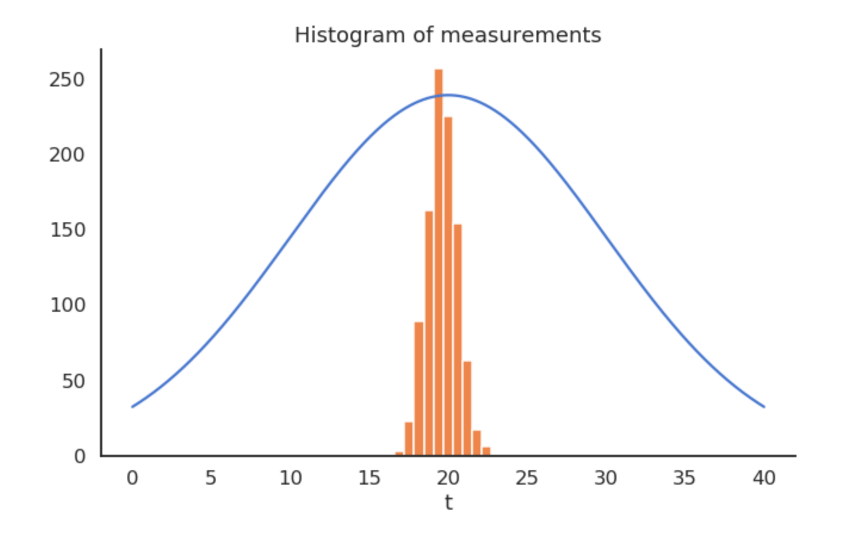
- An estimator is a random variable used to measure parameters of a distribution (e.g. its expectation). The problem is that estimators can generally be **biased**.
- Take the example of a thermometer M measuring the temperature T. T is a random variable (normally distributed with  $\mu=20$  and  $\sigma=10$ ) and the measurements M relate to the temperature with the relation:

$$M = 0.95 \, T + 0.65$$



### **Estimators**

- The thermometer is not perfect, but do random measurements allow us to estimate the expected value of the temperature?
- We could repeatedly take 100 random samples of the thermometer and see how the distribution of sample averages look like:



• But, as the expectation is linear, we actually have:

$$\mathbb{E}[M] = \mathbb{E}[0.95\,T + 0.65] = 0.95\,\mathbb{E}[T] + 0.65 = 19.65 
eq \mathbb{E}[T]$$

• The thermometer is a **biased estimator** of the temperature.

### **Estimators**

- Let's note  $\theta$  a parameter of a probability distribution X that we want to estimate (it does not have to be its mean).
- ullet An **estimator**  $\hat{ heta}$  is a random variable mapping the sample space of X to a set of sample estimates.
- The **bias** of an estimator is the mean error made by the estimator:

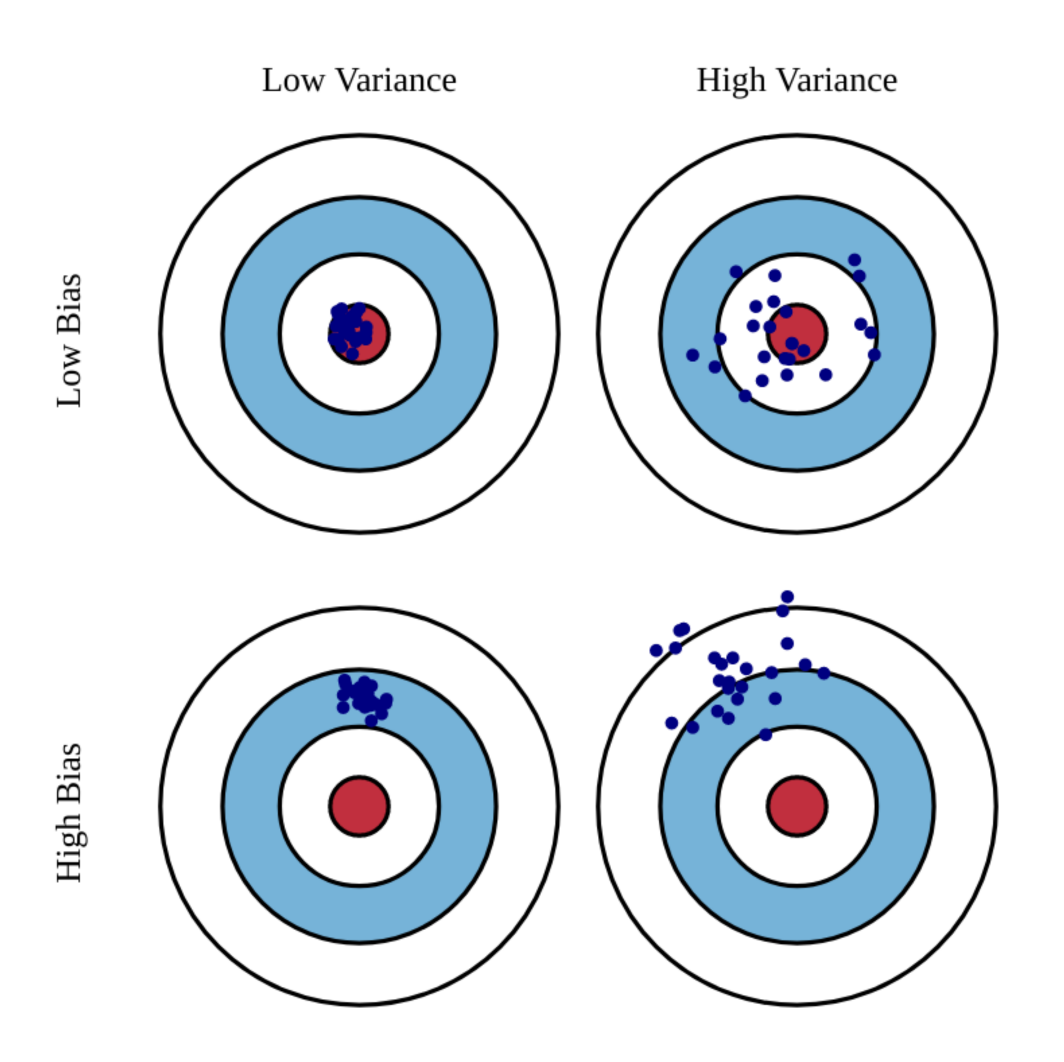
$$\mathcal{B}(\hat{ heta}) = \mathbb{E}[\hat{ heta} - heta] = \mathbb{E}[\hat{ heta}] - heta$$

• The variance of an estimator is the deviation of the samples around the expected value:

$$\operatorname{Var}(\hat{ heta}) = \mathbb{E}[(\hat{ heta} - \mathbb{E}[\hat{ heta}])^2]$$

- Ideally, we would like estimators with:
  - low bias: the estimations are correct on average (= equal to the true parameter).
  - low variance: we do not need many estimates to get a correct estimate (CLT:  $\frac{\sigma}{\sqrt{N}}$ )

### **Estimators: bias and variance**

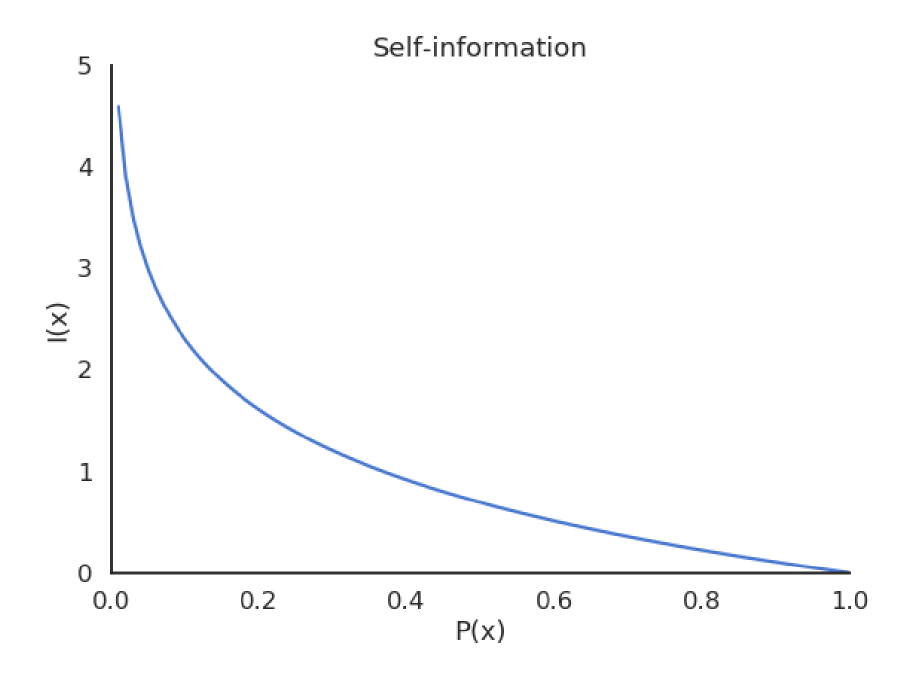


- Unfortunately, the perfect estimator does not exist.
- Estimators will have a bias and a variance:
  - **Bias**: the estimated values will be wrong, and the policy not optimal.
  - Variance: we will need a lot of samples (trial and error) to have correct estimates.
- One usually talks of a **bias/variance** trade-off: if you have a small bias, you will have a high variance, or vice versa.
- In machine learning, bias corresponds to underfitting, variance to overfitting.

# 5 - Information theory

### Information

- **Information theory** (Claude Shannon) asks how much information is contained in a probability distribution.
- Information is related to surprise or uncertainty: are the outcomes of a random variable surprising?
  - Almost certain outcomes ( $P\sim 1$ ) are not surprising because they happen all the time.
  - ullet Almost impossible outcomes ( $P\sim 0$ ) are very surprising because they are very rare.



• A useful measurement of how surprising is an outcome x is the **self-information**:

$$I(x) = -\log P(X = x)$$

- Depending on which log is used, self-information has different units:
  - $\log_2$ : bits or shannons.
  - $\log_e = \ln$ : nats.
- But it is just a rescaling, the base never matters.

### **Entropy**

• The **entropy** (or Shannon entropy) of a probability distribution is the expected value of the self-information of its outcomes:

$$H(X) = \mathbb{E}_{x \sim X}[I(x)] = \mathbb{E}_{x \sim X}[-\log P(X=x)]$$

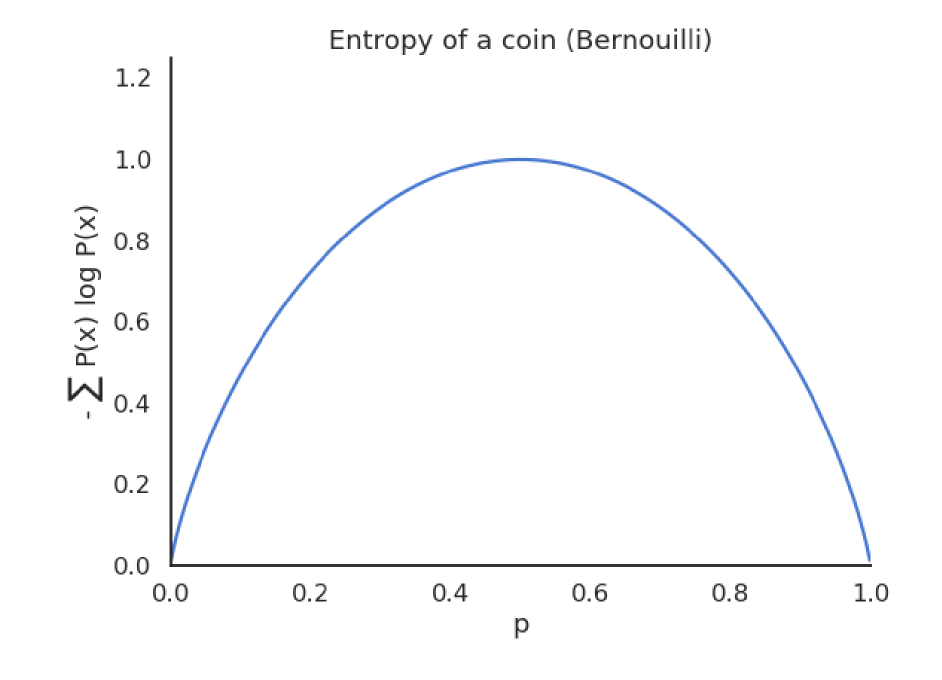
- It measures the uncertainty, randomness or information content of the random variable.
- In the discrete case:

$$H(X) = -\sum_x P(x) \, \log P(x)$$

• In the continuous case:

$$H(X) = -\int_x f(x) \, \log f(x) \, dx$$

- The entropy of a Bernouilli variable is maximal when both outcomes are **equiprobable**.
- If a variable is **deterministic**, its entropy is minimal and equal to zero.



## Joint and conditional entropies

ullet The **joint entropy** of two random variables X and Y is defined by:

$$H(X,Y) = \mathbb{E}_{x\sim X,y\sim Y}[-\log P(X=x,Y=y)]$$

ullet The **conditional entropy** of two random variables X and Y is defined by:

$$H(X|Y) = \mathbb{E}_{x\sim X,y\sim Y}[-\log P(X=x|Y=y)] = \mathbb{E}_{x\sim X,y\sim Y}[-\log rac{P(X=x,Y=y)}{P(Y=y)}]$$

If the variables are independent, we have:

$$H(X,Y) = H(X) + H(Y)$$
 or  $H(X|Y) = H(X)$ 

Both are related by:

$$H(X|Y) = H(X,Y) - H(Y)$$

The equivalent of Bayes' rule is:

$$H(Y|X) = H(X|Y) + H(Y) - H(X)$$

### **Mutual Information**

• The most important information measurement between two variables is the **mutual information** MI (or information gain):

$$I(X,Y) = H(X) - H(X|Y) = H(Y) - H(Y|X)$$

- ullet It measures how much information the variable X holds on Y:
  - ullet If the two variables are **independent**, the MI is 0 : X is as random, whether you know Y or not.

$$I(X,Y)=0$$

• If the two variables are **dependent**, knowing Y gives you information on X, which becomes less random, i.e. less uncertain / surprising.

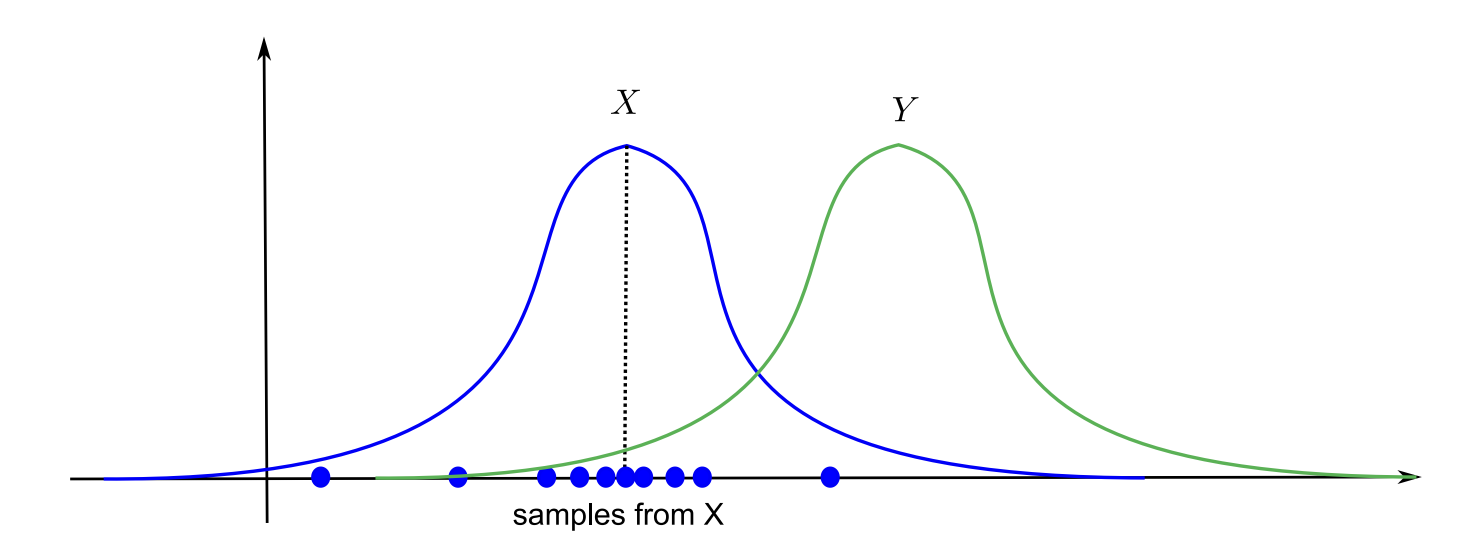
• If you can fully predict X when you know Y, it becomes deterministic (H(X|Y)=0) so the mutual information is maximal (I(X,Y)=H(X)).

### **Cross-entropy**

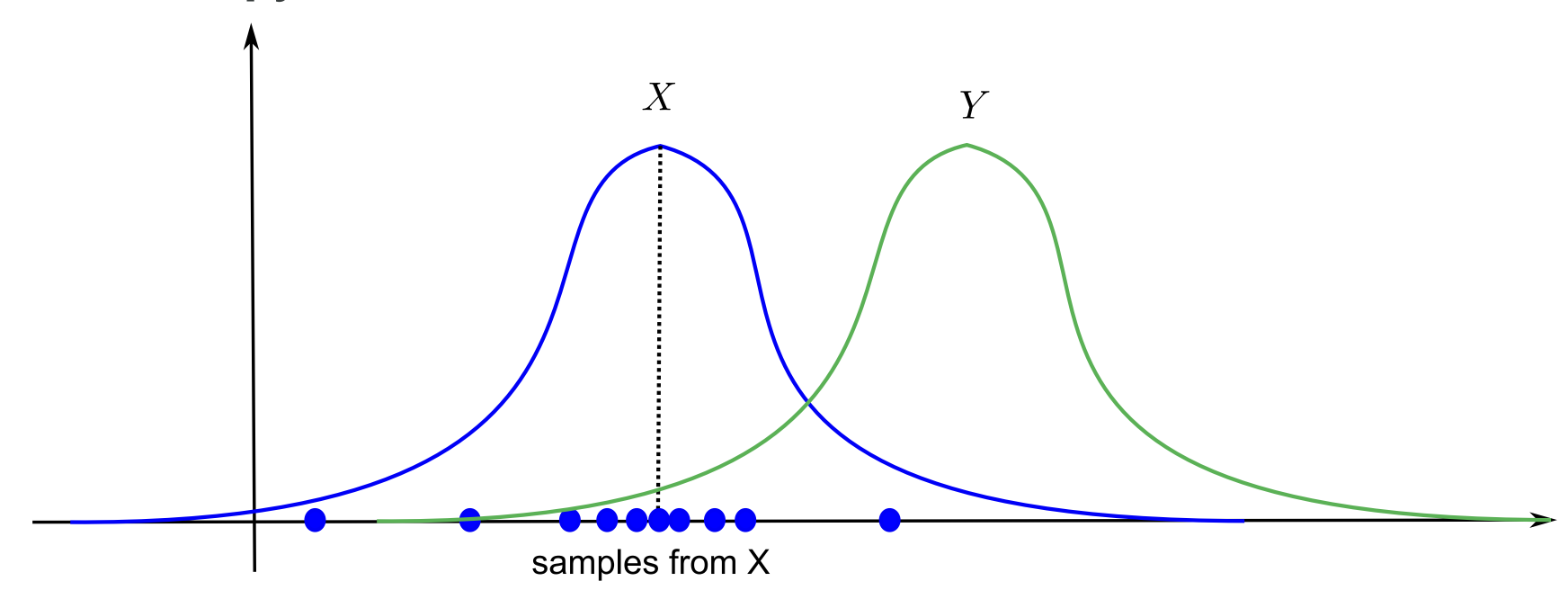
ullet The **cross-entropy** between two distributions X and Y is defined as:

$$H(X,Y) = \mathbb{E}_{x\sim X}[-\log P(Y=x)]$$

- ullet Beware that the notation H(X,Y) is the same as the joint entropy, but it is a different concept!
- The cross-entropy measures the **negative log-likelihood** that a sample x taken from the distribution X could also come from the distribution Y.
- ullet More exactly, it measures how many bits of information one would need to distinguish the two distributions X and Y.



### **Cross-entropy**



$$H(X,Y) = \mathbb{E}_{x\sim X}[-\log P(Y=x)]$$

- If the two distributions are the same *almost anywhere*, one cannot distinguish samples from the two distributions:
  - ullet The cross-entropy is the same as the entropy of X.
- ullet If the two distributions are completely different, one can tell whether a sample Z comes from X or Y:
  - ullet The cross-entropy is higher than the entropy of X.

## Kullback-Leibler divergence

• In practice, the **Kullback-Leibler divergence**  $\mathrm{KL}(X||Y)$  is a better measurement of the similarity (statistical distance) between two probability distributions:

$$ext{KL}(X||Y) = \mathbb{E}_{x\sim X}[-\lograc{P(Y=x)}{P(X=x)}]$$

It is linked to the cross-entropy by:

$$\mathrm{KL}(X||Y) = H(X,Y) - H(X)$$

- If the two distributions are the same *almost anywhere*:
  - The KL divergence is zero.
- If the two distributions are different:
  - The KL divergence is positive.
- Minimizing the KL between two distributions is the same as making the two distributions "equal".
- Again, the KL is not a metric, as it is not symmetric.