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(https://stanford.edu/~shervine/teaching/cs-229/cheatsheet-supervised-learning#cs-229---machine-learning)CS 229 - Machine Learning (teaching/cs-229)

Supervised Learning	Unsupervised Learning	Deep Learning	Tips and tricks
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(https://stanford.edu/~shervine/teaching/cs-229/cheatsheet-supervisedlearning#cheatsheet)Supervised Learning cheatsheet

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(https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#introduction) Introduction to Supervised Learning

Given a set of data points $\{x^{(1)},...,x^{(m)}\}$ associated to a set of outcomes $\{y^{(1)},...,y^{(m)}\}$, we want to build a classifier that learns how to predict y from x.

 \Box **Type of prediction** — The different types of predictive models are summed up in the table below:

	Regression	Classification
Outcome	Continuous	Class
Examples	Linear regression	Logistic regression, SVM, Naive Bayes

\square Type of model $-$	- The different models	are summed up	in the table below:
,,,			

	Discriminative model	Generative model
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Goal	Directly estimate $P(y x)$	Estimate $P(x y)$ to then deduce $P(y x)$
What's learned	Decision boundary	Probability distributions of the data
Illustration		
Examples	Regressions, SVMs	GDA, Naive Bayes

[https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#notations) Notations and general concepts

 \Box **Hypothesis** — The hypothesis is noted h_{θ} and is the model that we choose. For a given input data $x^{(i)}$ the model prediction output is $h_{\theta}(x^{(i)})$.

 \square Loss function — A loss function is a function $L:(z,y)\in\mathbb{R}\times Y\longmapsto L(z,y)\in\mathbb{R}$ that takes as inputs the predicted value z corresponding to the real data value y and outputs how different they are. The common loss functions are summed up in the table below:

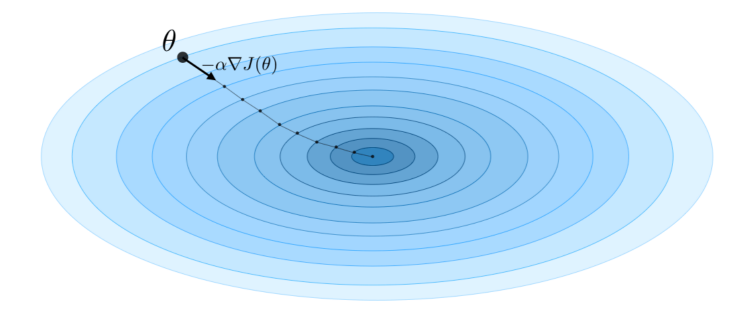
Least squared error	Logistic loss	Hinge loss	Cro
$\frac{1}{2}(y-z)^2$	$\log(1+\exp(-yz))$	$\max(0,1-yz)$	$-\Big[y \ y)$
$y\in\mathbb{R}$	y = -1 $y = 1$	y = -1 $y = 1$ $y = 1$	y = 0
Linear regression	Logistic regression	SVM	Neı

 \square **Cost function** — The cost function J is commonly used to assess the performance of a model, and is defined with the loss function L as follows:

$$J(heta) = \sum_{i=1}^m L(h_ heta(x^{(i)}), y^{(i)})$$

 \Box Gradient descent — By noting $\alpha \in \mathbb{R}$ the learning rate, the update rule for gradient descent is expressed with the learning rate and the cost function J as follows:

$$\theta \longleftarrow \theta - \alpha \nabla J(\theta)$$



Remark: Stochastic gradient descent (SGD) is updating the parameter based on each training example, and batch gradient descent is on a batch of training examples.

 \Box **Likelihood** — The likelihood of a model $L(\theta)$ given parameters θ is used to find the optimal parameters θ through likelihood maximization. We have:

$$heta^{ ext{opt}} = rg\max_{ heta} L(heta)$$

Remark: in practice, we use the log-likelihood $\ell(\theta) = \log(L(\theta))$ which is easier to optimize.

 \square **Newton's algorithm** — Newton's algorithm is a numerical method that finds θ such that $\ell'(\theta) = 0$. Its update rule is as follows:

$$heta \leftarrow heta - rac{\ell'(heta)}{\ell''(heta)}$$

Remark: the multidimensional generalization, also known as the Newton-Raphson method, has the following update rule:

$$heta \leftarrow heta - \left(
abla_{ heta}^2 \ell(heta)
ight)^{-1}
abla_{ heta} \ell(heta)$$

(https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#linear-models) Linear models

Linear regression

We assume here that $y|x; heta \sim \mathcal{N}(\mu, \sigma^2)$

 \square **Normal equations** — By noting X the design matrix, the value of θ that minimizes the cost function is a closed-form solution such that:

$$oxed{ heta = (X^TX)^{-1}X^Ty}$$

 \square LMS algorithm — By noting α the learning rate, the update rule of the Least Mean Squares (LMS) algorithm for a training set of m data points, which is also known as the Widrow-Hoff learning rule, is as follows:

$$oxed{ orall j, \quad heta_j \leftarrow heta_j + lpha \sum_{i=1}^m \left[y^{(i)} - h_ heta(x^{(i)})
ight] x_j^{(i)} }$$

Remark: the update rule is a particular case of the gradient ascent.

 \square LWR — Locally Weighted Regression, also known as LWR, is a variant of linear regression that weights each training example in its cost function by $w^{(i)}(x)$, which is defined with parameter $\tau \in \mathbb{R}$ as:

$$oxed{w^{(i)}(x)=\exp\left(-rac{(x^{(i)}-x)^2}{2 au^2}
ight)}$$

Classification and logistic regression

 \square **Sigmoid function** — The sigmoid function g, also known as the logistic function, is defined as follows:

$$orall z \in \mathbb{R}, \quad \boxed{g(z) = rac{1}{1 + e^{-z}} \in]0,1[}$$

 \Box Logistic regression — We assume here that $y|x; \theta \sim \mathrm{Bernoulli}(\phi)$. We have the following form:

$$\boxed{\phi = p(y=1|x; heta) = rac{1}{1+\exp(- heta^T x)} = g(heta^T x)}$$

Remark: logistic regressions do not have closed form solutions.

 \square Softmax regression — A softmax regression, also called a multiclass logistic regression, is used to generalize logistic regression when there are more than 2 outcome classes. By convention, we set $\theta_K=0$, which makes the Bernoulli parameter ϕ_i of each class i be such that:

$$\phi_i = rac{\exp(heta_i^T x)}{\displaystyle\sum_{j=1}^K \exp(heta_j^T x)}$$

Generalized Linear Models

 \square **Exponential family** — A class of distributions is said to be in the exponential family if it can be written in terms of a natural parameter, also called the canonical parameter or link function, η , a sufficient statistic T(y) and a log-partition function $a(\eta)$ as follows:

$$\boxed{p(y;\eta) = b(y) \exp(\eta T(y) - a(\eta))}$$

Remark: we will often have T(y) = y. Also, $\exp(-a(\eta))$ can be seen as a normalization parameter that will make sure that the probabilities sum to one.

The most common exponential distributions are summed up in the following table:

Distribution	η	T(y)	$a(\eta)$	b(y)
Bernoulli	$\log\left(rac{\phi}{1-\phi} ight)$	y	$\log(1+\exp(\eta))$	1
Gaussian	μ	y	$rac{\eta^2}{2}$	$\frac{1}{\sqrt{2\pi}}\exp\left(-\frac{y^2}{2}\right)$
Poisson	$\log(\lambda)$	y	e^{η}	$\frac{1}{y!}$
Geometric	$\log(1-\phi)$	y	$\log\left(rac{e^{\eta}}{1-e^{\eta}} ight)$	1

☐ **Assumptions of GLMs** — Generalized Linear Models (GLM) aim at predicting a random variable y as a function of $x \in \mathbb{R}^{n+1}$ and rely on the following 3 assumptions:

$$egin{aligned} (1) & egin{aligned} y|x; heta \sim ext{ExpFamily}(\eta) \ (2) & eta_{ heta}(x) = E[y|x; heta] \ (3) & egin{aligned} \eta = heta^T x \end{aligned} \end{aligned}$$

$$(2) \quad \boxed{h_{\theta}(x) = E[y|x;\theta]}$$

$$(3) \quad \Big| \, \eta = heta^T x \Big|$$

Remark: ordinary least squares and logistic regression are special cases of generalized linear models.

[https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#svm) **Support Vector Machines**

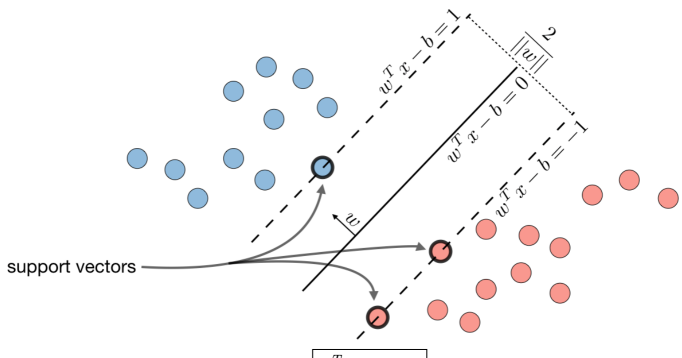
The goal of support vector machines is to find the line that maximizes the minimum distance to the line.

 \square **Optimal margin classifier** — The optimal margin classifier h is such that:

$$h(x) = \mathrm{sign}(w^Tx - b)$$

where $(w,b)\in\mathbb{R}^n imes\mathbb{R}$ is the solution of the following optimization problem:

$$oxed{\min rac{1}{2} ||w||^2} \qquad ext{such that} \quad oxed{y^{(i)}(w^T x^{(i)} - b) \geqslant 1}$$



Remark: the decision boundary is defined as $w^Tx - b = 0$

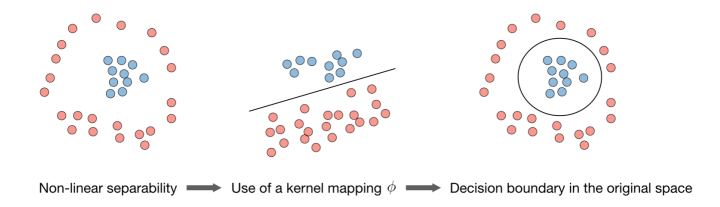
 \square **Hinge loss** — The hinge loss is used in the setting of SVMs and is defined as follows:

$$L(z,y)=[1-yz]_+=\max(0,1-yz)$$

 \square **Kernel** — Given a feature mapping ϕ , we define the kernel K as follows:

$$K(x,z) = \phi(x)^T \phi(z)$$

In practice, the kernel K defined by $K(x,z)=\exp\left(-\frac{||x-z||^2}{2\sigma^2}\right)$ is called the Gaussian kernel and is commonly used.



Remark: we say that we use the "kernel trick" to compute the cost function using the kernel because we actually don't need to know the explicit mapping ϕ , which is often very complicated. Instead, only the values K(x,z) are needed.

 \square **Lagrangian** — We define the Lagrangian $\mathcal{L}(w,b)$ as follows:

$$\mathcal{L}(w,b) = f(w) + \sum_{i=1}^l eta_i h_i(w)$$

Remark: the coefficients β_i are called the Lagrange multipliers.

[https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#generative-learning) **Generative Learning**

A generative model first tries to learn how the data is generated by estimating P(x|y), which we can then use to estimate P(y|x) by using Bayes' rule.

Gaussian Discriminant Analysis

 \square **Setting** — The Gaussian Discriminant Analysis assumes that y and x|y=0 and x|y=1are such that:

(1)
$$y \sim \text{Bernoulli}(\phi)$$

$$(2) \quad \boxed{x|y=0 \sim \mathcal{N}(\mu_0,\Sigma)}$$

$$egin{aligned} (1) & egin{aligned} y \sim \mathrm{Bernoulli}(\phi) \ \ (2) & x|y=0 \sim \mathcal{N}(\mu_0,\Sigma) \ \ \ (3) & x|y=1 \sim \mathcal{N}(\mu_1,\Sigma) \end{aligned}$$

☐ **Estimation** — The following table sums up the estimates that we find when maximizing the likelihood:

$\widehat{\phi}$	$\widehat{\mu_j} (j=0,1)$	$\widehat{\Sigma}$
$\frac{1}{m} \sum_{i=1}^m 1_{\{y^{(i)}=1\}}$	$rac{\sum_{i=1}^m 1_{\{y^{(i)}=j\}} x^{(i)}}{\sum_{i=1}^m 1_{\{y^{(i)}=j\}}}$	$rac{1}{m} \sum_{i=1}^m (x^{(i)} - \mu_{y^{(i)}}) (x^{(i)} - \mu_{y^{(i)}})^T$

Naive Bayes

☐ **Assumption** — The Naive Bayes model supposes that the features of each data point are all independent:

$$P(x|y) = P(x_1, x_2, ...|y) = P(x_1|y)P(x_2|y)... = \prod_{i=1}^n P(x_i|y)$$

☐ **Solutions** — Maximizing the log-likelihood gives the following solutions:

$$P(y=k)=rac{1}{m} imes\#\{j|y^{(j)}=k\}$$

$$P(y=k) = rac{1}{m} imes \#\{j|y^{(j)} = k\} \hspace{0.5cm} ext{and} \hspace{0.5cm} P(x_i = l|y = k) = rac{\#\{j|y^{(j)} = k ext{ and } x_i^{(j)} = k\}}{\#\{j|y^{(j)} = k\}}$$

with $k \in \{0,1\}$ and $l \in \llbracket 1,L
rbracket$

Remark: Naive Bayes is widely used for text classification and spam detection.

[https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#tree)

Tree-based and ensemble methods

These methods can be used for both regression and classification problems.

☐ **CART** — Classification and Regression Trees (CART), commonly known as decision trees, can be represented as binary trees. They have the advantage to be very interpretable.

☐ **Random forest** — It is a tree-based technique that uses a high number of decision trees built out of randomly selected sets of features. Contrary to the simple decision tree, it is highly uninterpretable but its generally good performance makes it a popular algorithm.

Remark: random forests are a type of ensemble methods.

☐ **Boosting** — The idea of boosting methods is to combine several weak learners to form a stronger one. The main ones are summed up in the table below:

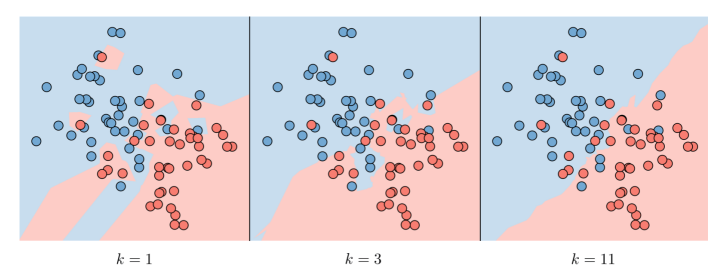
Adaptive boosting	Gradient boosting	
High weights are put on errors to improve at the next boosting stepKnown as Adaboost	Weak learners are trained on residualsExamples include XGBoost	

https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#other)

Other non-parametric approaches

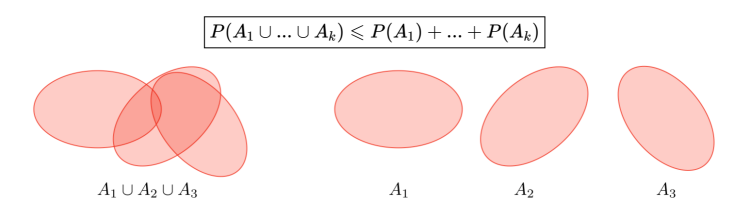
 \square k-nearest neighbors — The k-nearest neighbors algorithm, commonly known as k-NN, is a non-parametric approach where the response of a data point is determined by the nature of its kneighbors from the training set. It can be used in both classification and regression settings.

Remark: the higher the parameter k, the higher the bias, and the lower the parameter k, the higher the variance.



[https://stanford.edu/~shervine/teaching/cs-229/cheatsheetsupervised-learning#learning-theory) **Learning Theory**

 \square Union bound — Let $A_1,...,A_k$ be k events. We have:



 \square **Hoeffding inequality** — Let $Z_1,..,Z_m$ be m iid variables drawn from a Bernoulli distribution of parameter ϕ . Let $\widehat{\phi}$ be their sample mean and $\gamma > 0$ fixed. We have:

$$P(|\phi - \widehat{\phi}| > \gamma) \leqslant 2 \exp(-2\gamma^2 m)$$

Remark: this inequality is also known as the Chernoff bound.

 \square **Training error** — For a given classifier h, we define the training error $\hat{\epsilon}(h)$, also known as the empirical risk or empirical error, to be as follows:

$$\left|\widehat{\epsilon}(h) = rac{1}{m} \sum_{i=1}^m \mathbb{1}_{\{h(x^{(i)})
eq y^{(i)}\}}
ight|$$

- ☐ **Probably Approximately Correct (PAC)** PAC is a framework under which numerous results on learning theory were proved, and has the following set of assumptions:
 - the training and testing sets follow the same distribution
 - the training examples are drawn independently
- \square **Shattering** Given a set $S=\{x^{(1)},...,x^{(d)}\}$, and a set of classifiers $\mathcal H$, we say that $\mathcal H$ shatters S if for any set of labels $\{y^{(1)},...,y^{(d)}\}$, we have:

$$oxed{\exists h \in \mathcal{H}, \quad orall i \in \llbracket 1, d
rbracket}, \quad h(x^{(i)}) = y^{(i)}}$$

 \Box **Upper bound theorem** — Let $\mathcal H$ be a finite hypothesis class such that $|\mathcal H|=k$ and let δ and the sample size m be fixed. Then, with probability of at least $1-\delta$, we have:

$$\left| \epsilon(\widehat{h}) \leqslant \left(\min_{h \in \mathcal{H}} \epsilon(h)
ight) + 2 \sqrt{rac{1}{2m} \log \left(rac{2k}{\delta}
ight)}
ight.$$

☐ **VC dimension** — The Vapnik-Chervonenkis (VC) dimension of a given infinite hypothesis class \mathcal{H} , noted $VC(\mathcal{H})$ is the size of the largest set that is shattered by \mathcal{H} .

Remark: the VC dimension of $\mathcal{H} = \{\text{set of linear classifiers in 2 dimensions}\}\$ is 3.

















 \Box Theorem (Vapnik) — Let ${\mathcal H}$ be given, with ${
m VC}({\mathcal H})=d$ and m the number of training examples. With probability at least $1 - \delta$, we have:

$$\left| \epsilon(\widehat{h}) \leqslant \left(\min_{h \in \mathcal{H}} \epsilon(h)
ight) + O\left(\sqrt{rac{d}{m} \log\left(rac{m}{d}
ight) + rac{1}{m} \log\left(rac{1}{\delta}
ight)}
ight)
ight|$$





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