Classification

Machine Learning Course - CS-433 Oct 19, 2021 Nicolas Flammarion



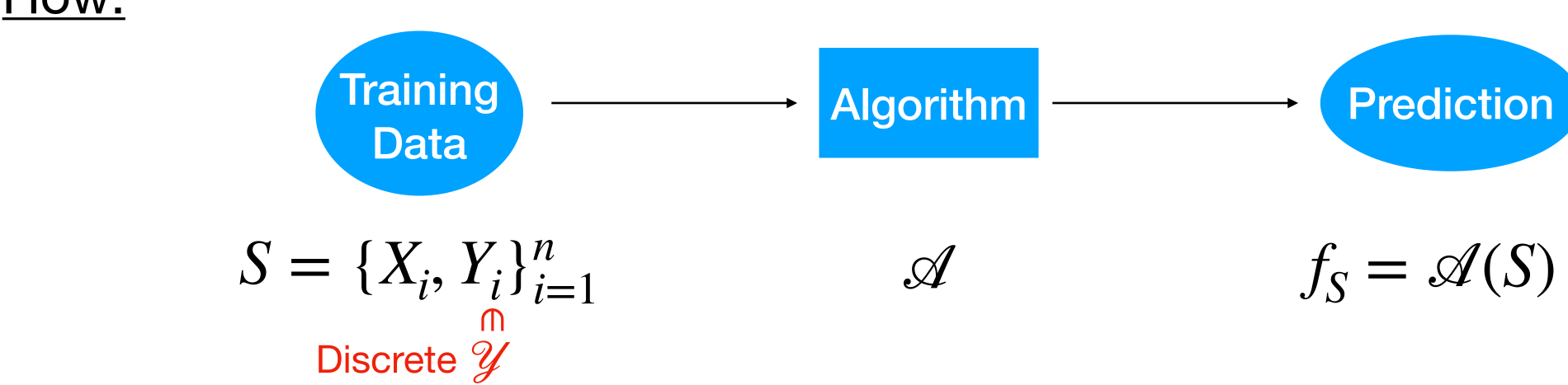
Definition of classification

We observe some data $S = \{X_i, Y_i\}_{i=1}^n \in \mathcal{X} \times \mathcal{Y}$

Discrete Set

Goal: given a new X, we want to predict its label Y

How:



Classification: relates input to a categorical variable

$$(X, Y) \in \mathcal{X} \times \mathcal{Y}$$
Discrete Set

Binary Classification: Y can take two values

 $Y \in \{c_1, c_2\}$ where c_i are the class labels . We often use $\{0,1\}$ or $\{-1,1\}$

Multi-class classification: Y can take more than two values

 $Y \in \{c_1, \dots, c_{K-1}\}$ for a K class problem. We often use $\{0, \dots, K-1\}$



no ordering between classes

Spam Detection



Patterns – Cell Press

Meet the newest members c Fri 10/8

The science of data Can't see this email p...

Volkan Cevher

▶ EPFL-CIS RIKEN-AIP Joint Talks on Thu 3:08 PM Dear All, I hope all is well. I am sending t...

sae.amenagements@...

Aménagements d'études pour ce Thu 2:36 PM English version below Madame, Monsieu...

westernunionrespo...

Western Union: Please verify 9/26/2021

Dear JULES ADAM, We noticed that you ...

Bachmann Jennifer

▶ coffee 10:35 AM

Hello, C'est Goodlife Coffee 😊 Ils m'ont...

Gestion de l'Ecole do...

Workshops for mentors 8:33 AM

Dear Thesis Directors, We hope our ema...

The Boston Globe

Parenting Unfiltered: Your W 2:00 AM

Beer. Blankets. Burgers. Dumplings! ...



Classifier



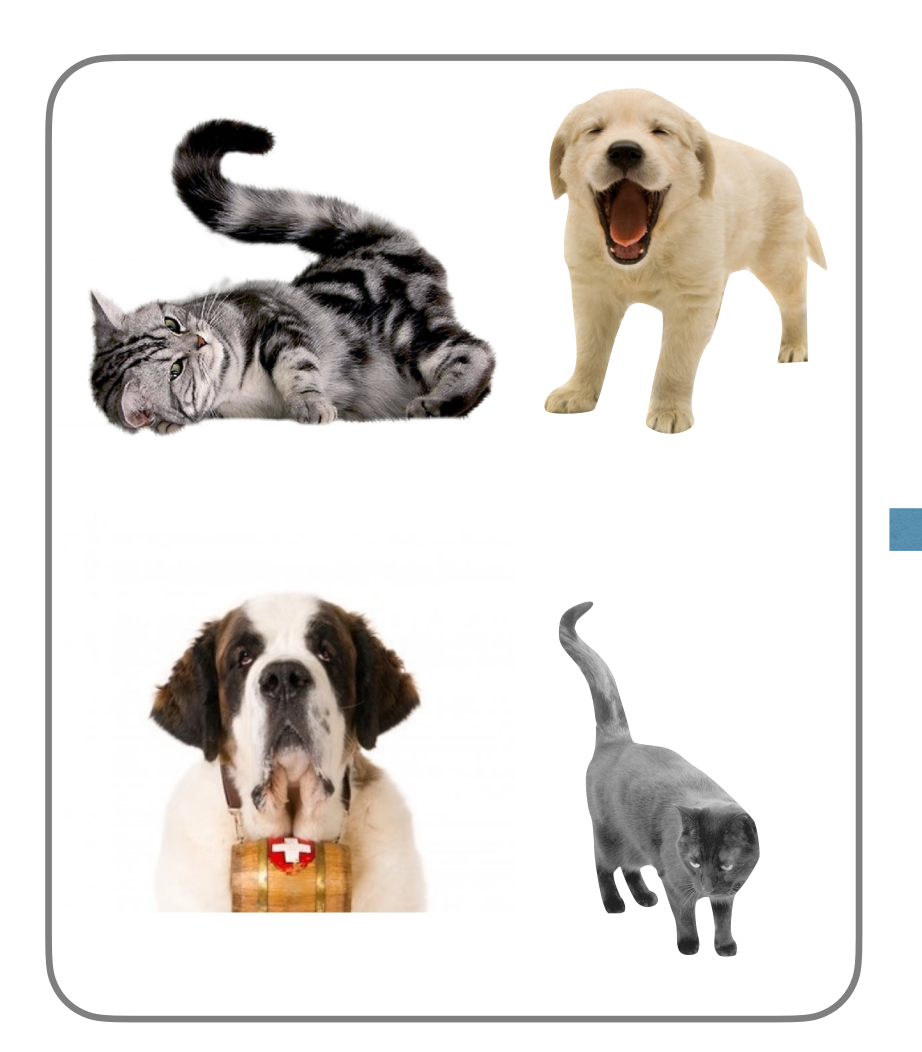


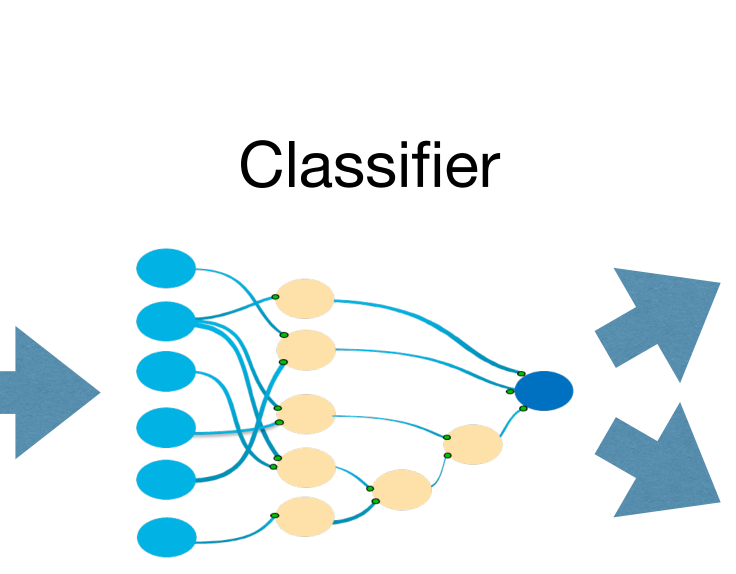
Inbox

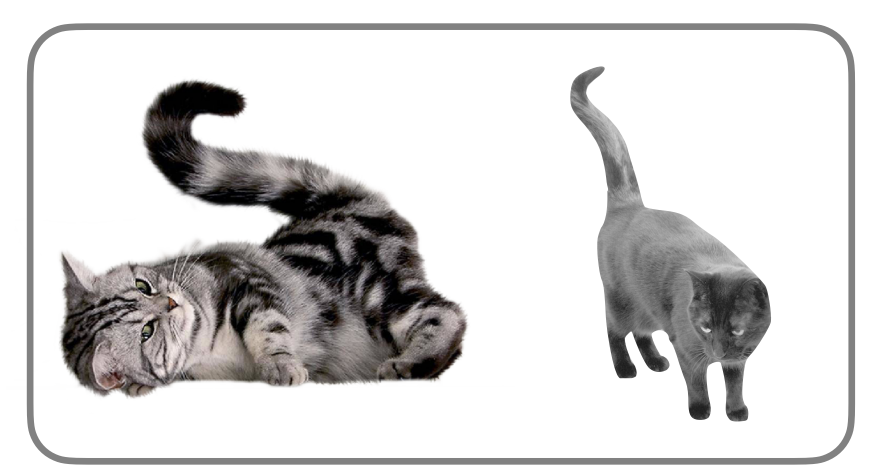


Spam Folder

Image classification

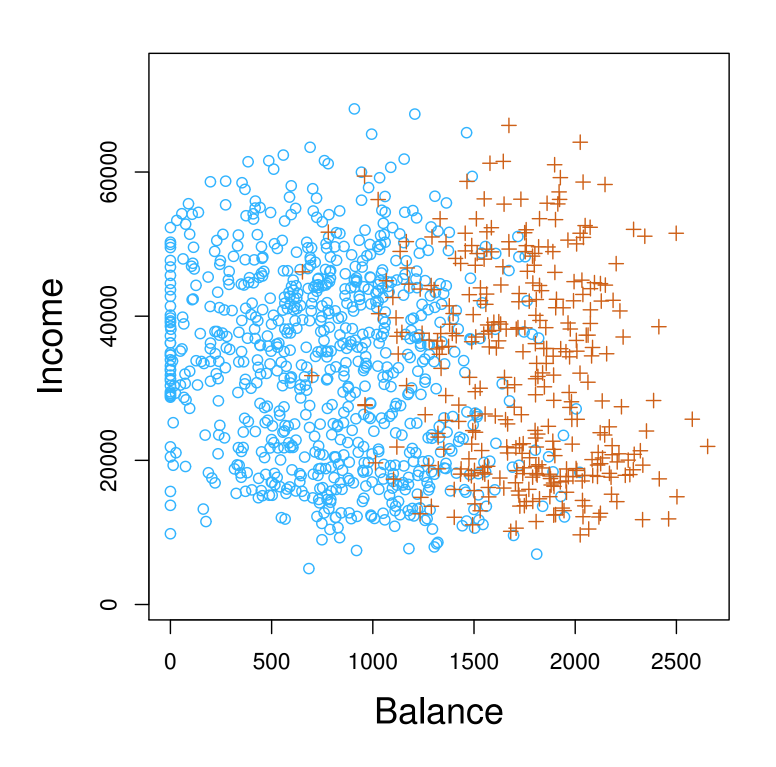








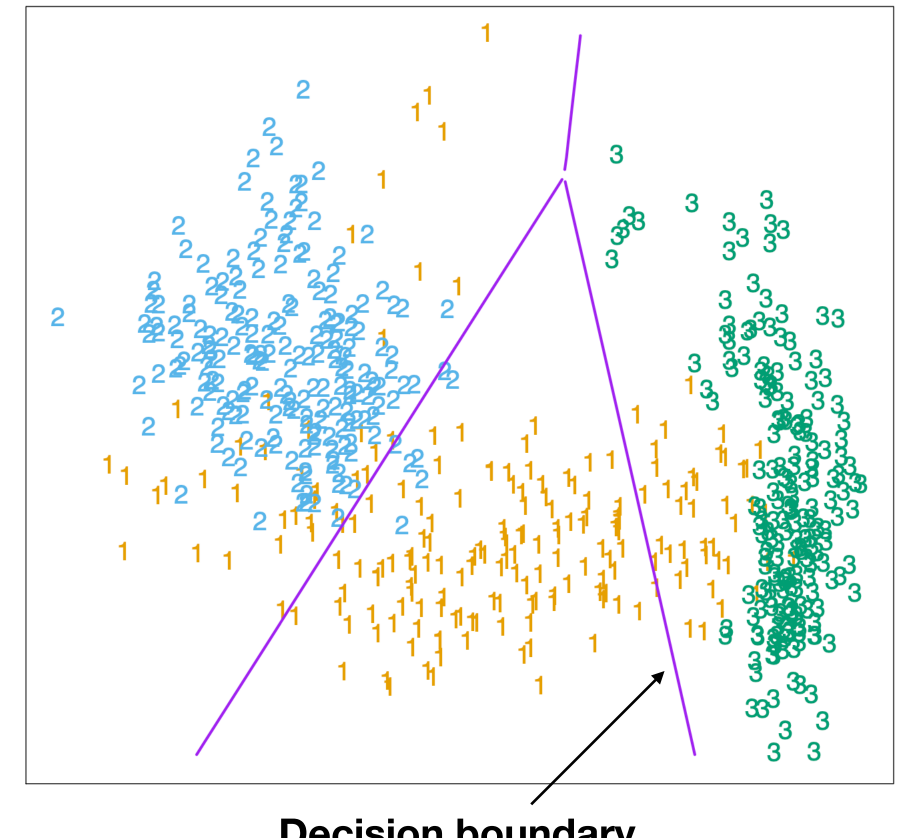
Credit Card Default



- + individual who defaulted on their credit card payments
- individual who did not

Classifier

A classifier $f:\mathcal{X} \to \mathcal{Y}$ divides the input space into a collection of region belonging to each class

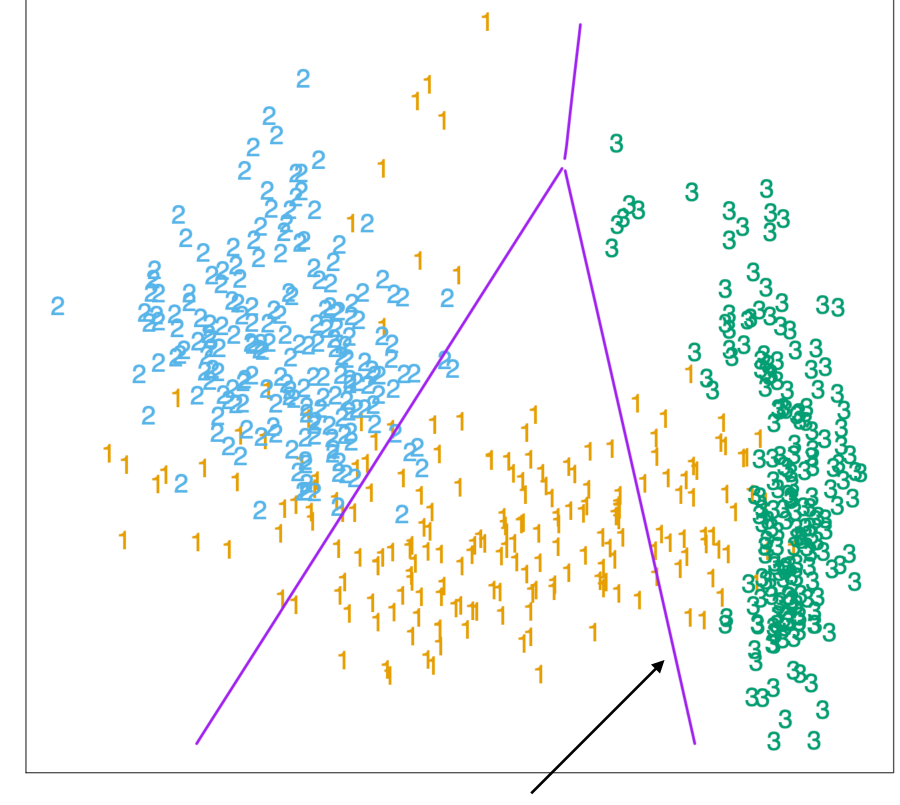


Decision boundary

Classifier

A classifier $f:\mathcal{X}\to\mathcal{Y}$ divides the input space into a collection of region belonging to each class

It can be linear

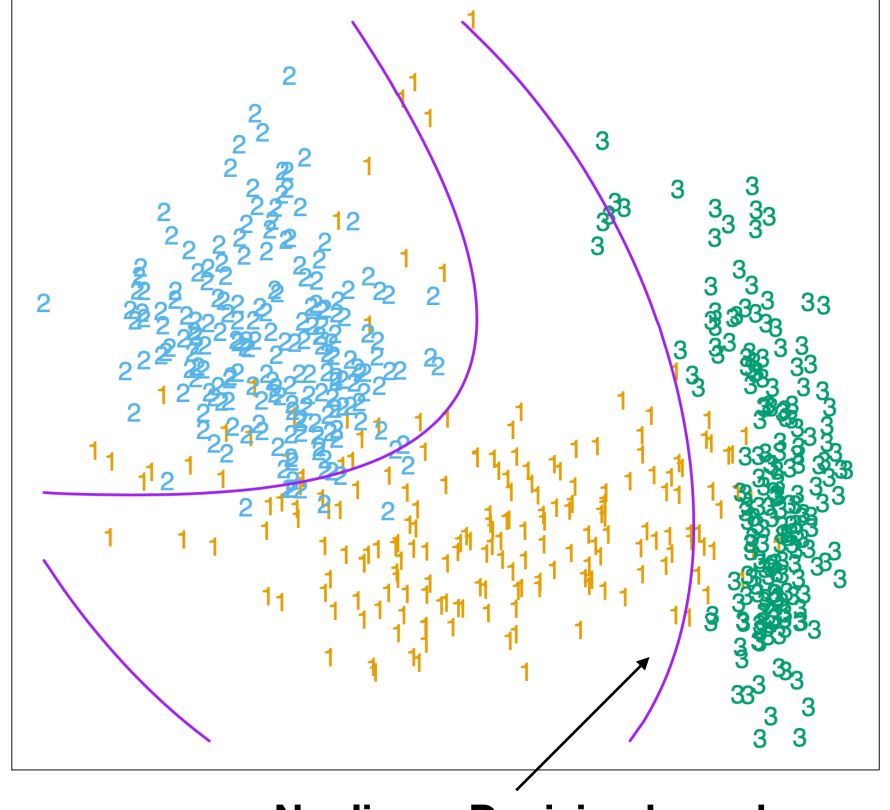


Linear Decision boundary

Classifier

A classifier $f:\mathcal{X}\to\mathcal{Y}$ divides the input space into a collection of region belonging to each class

It can also be nonlinear



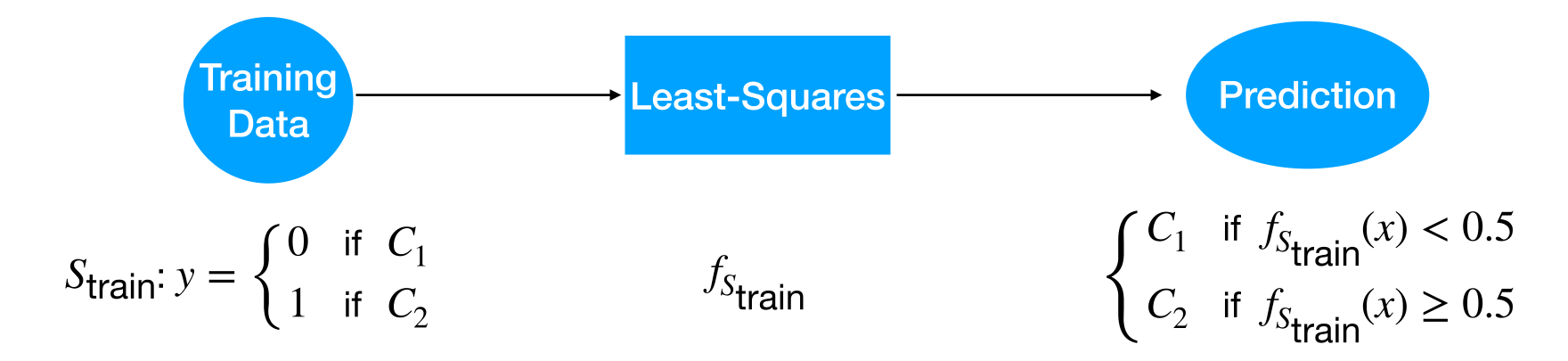
Nonlinear Decision boundary

Classification: a special case of regression?

Classification is a regression problem with discrete labels:

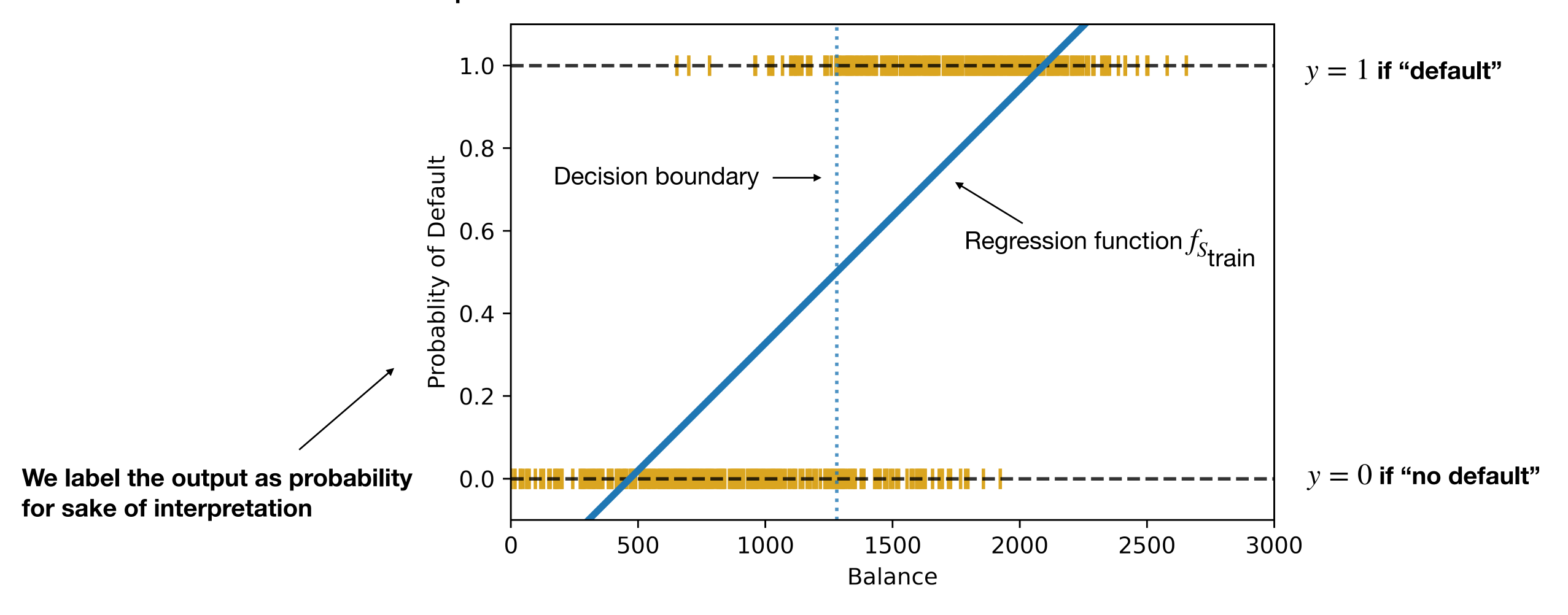
$$(x, y) \in \mathcal{X} \times \{0, 1\} \subset \mathcal{X} \times \mathbb{R}$$

Could we use previously seen regression methods to solve it?



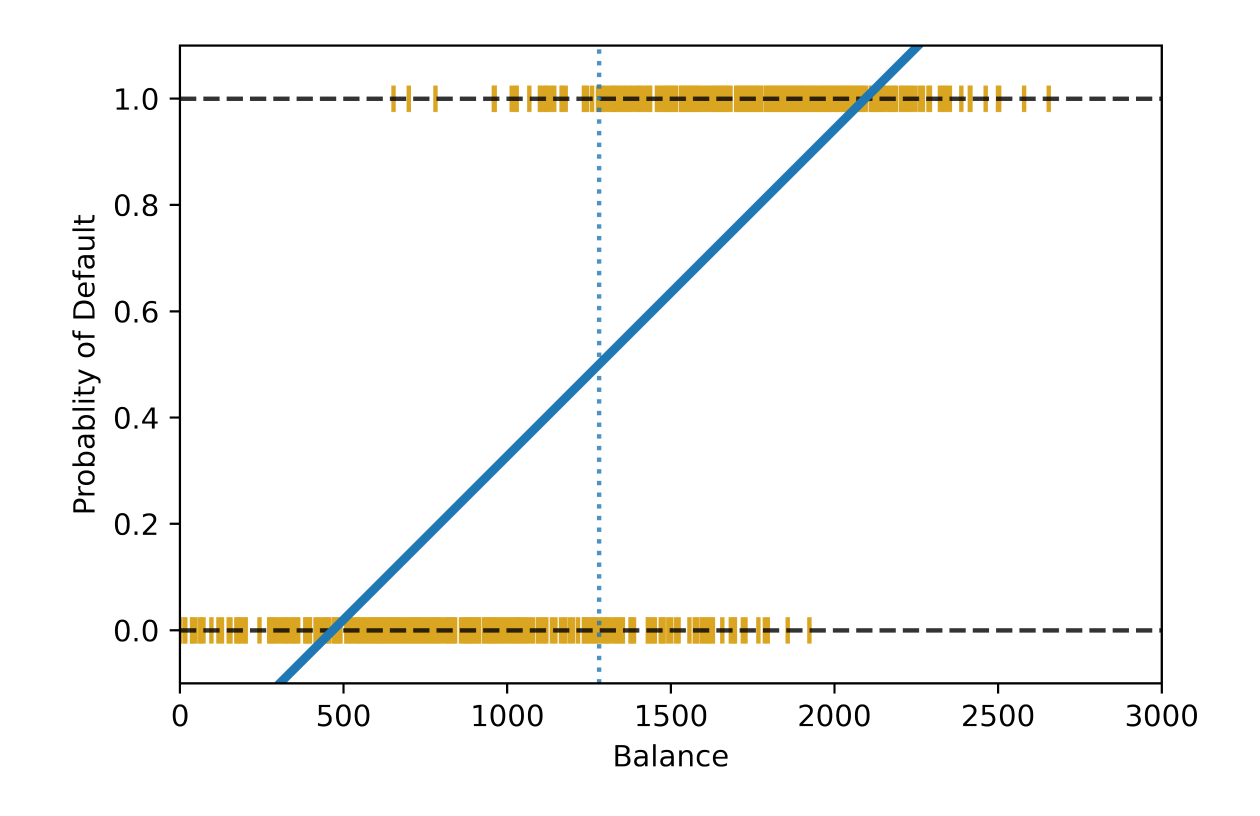
Is it a good idea?

Credit-card default problem:



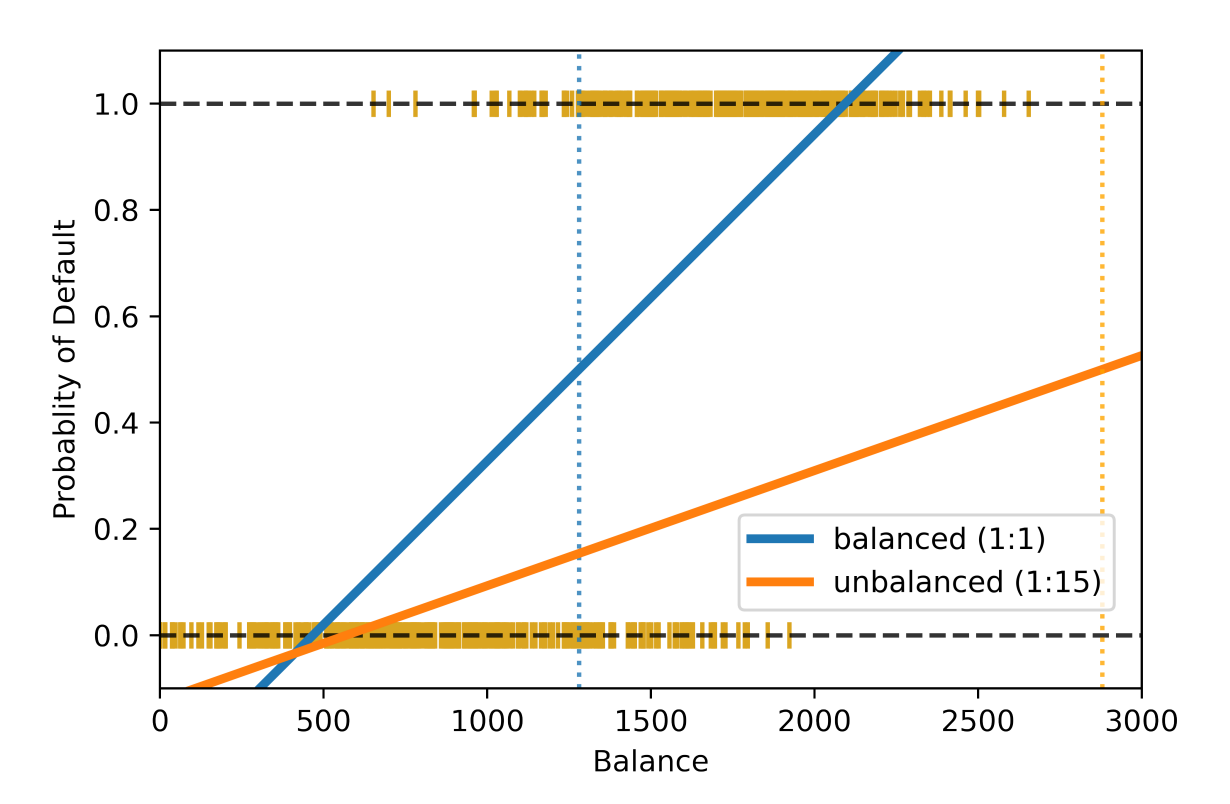
Classification is not just a special form of regression

A. The predicted values are not probabilities (not in [0,1])



Classification is not just a special form of regression

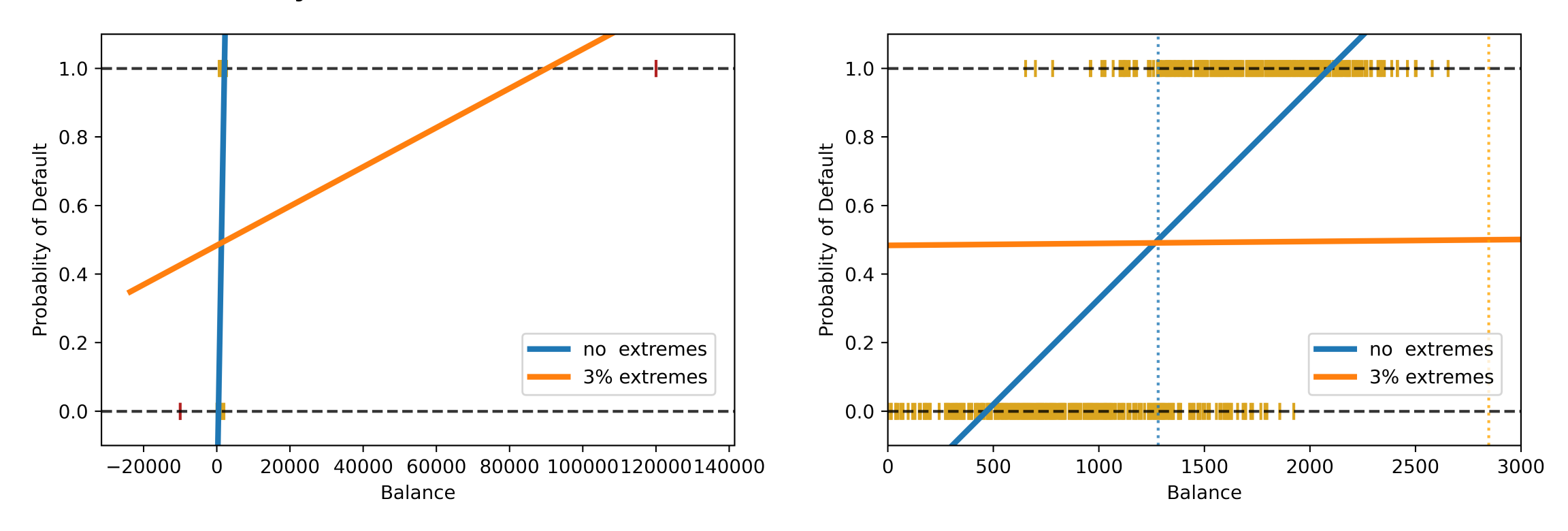
B. Sensitivity to unbalanced data



The position of the line depends crucially on how many points are in each class

Classification is not just a special form of regression

C. Sensitivity to extreme values:



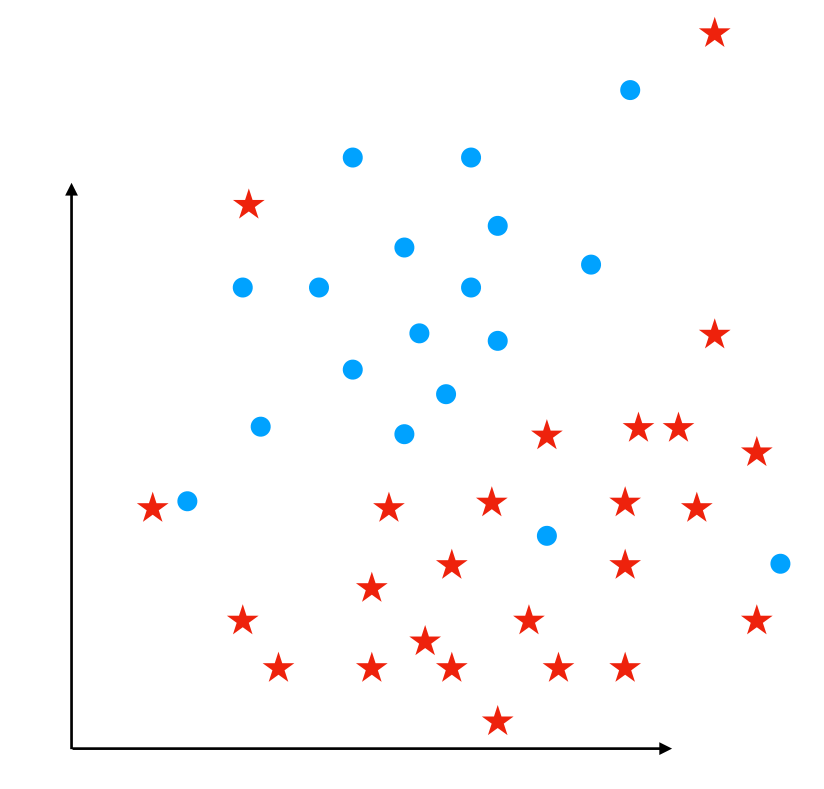
The position of the line depends crucially on where the points lie

Why: the square loss we used for regression is not suitable for classification

How to perform classification?

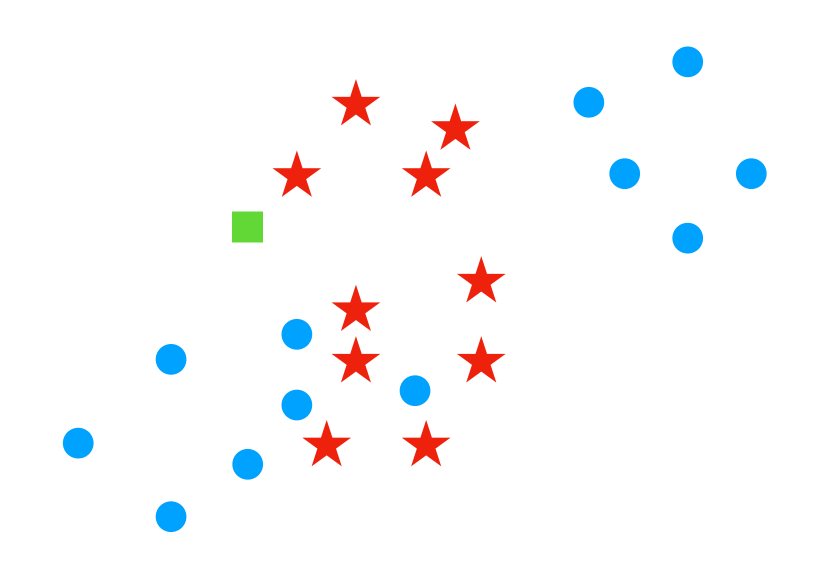
- A lot of different approaches have been developed
- We will not detail them exhaustively today
- Rather we will provide quick introductions

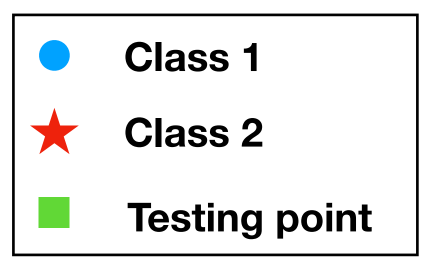




separate the space into various decision regions

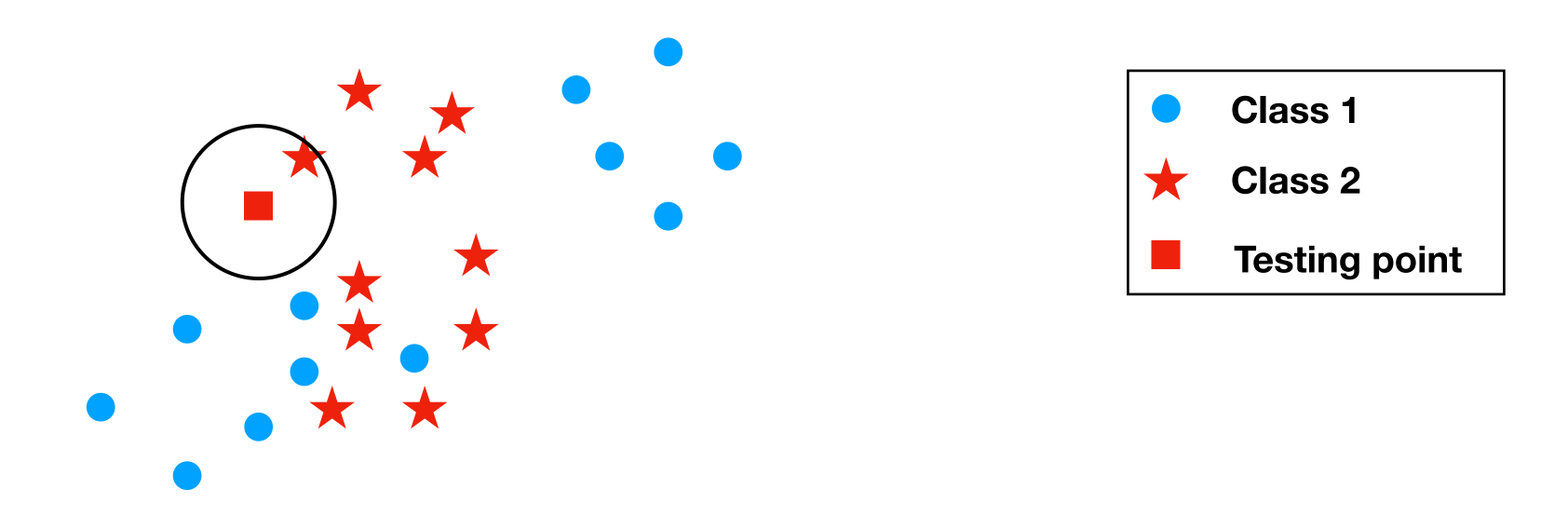
Nearest Neighbor





Nearest Neighbor

Assume that nearby points are likely to have similar label



You assign the label of the closest point in your training set.

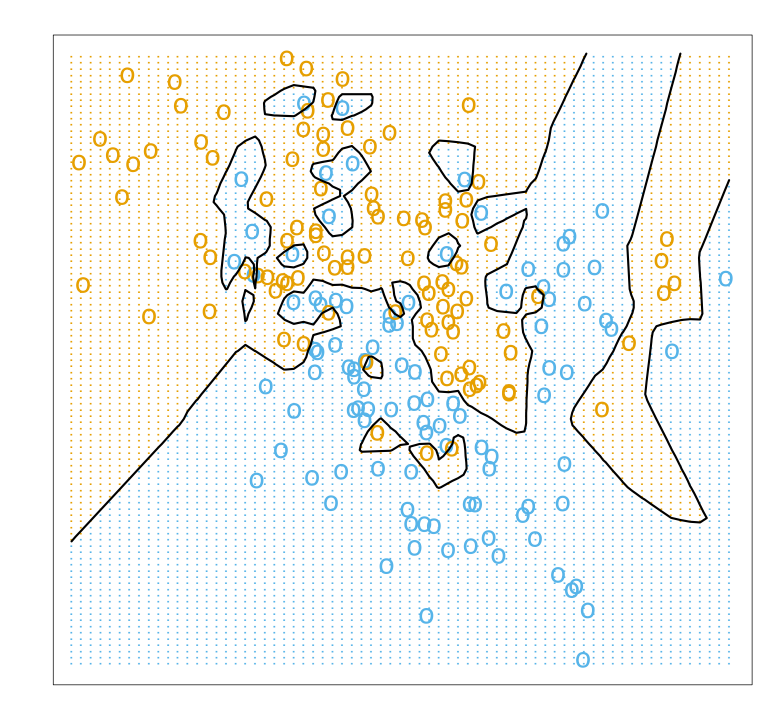
Nearest Neighbor

Pros:

- No optimization or training
- Easy to implement
- Works well in low dimension where you can get some very complex decision boundaries

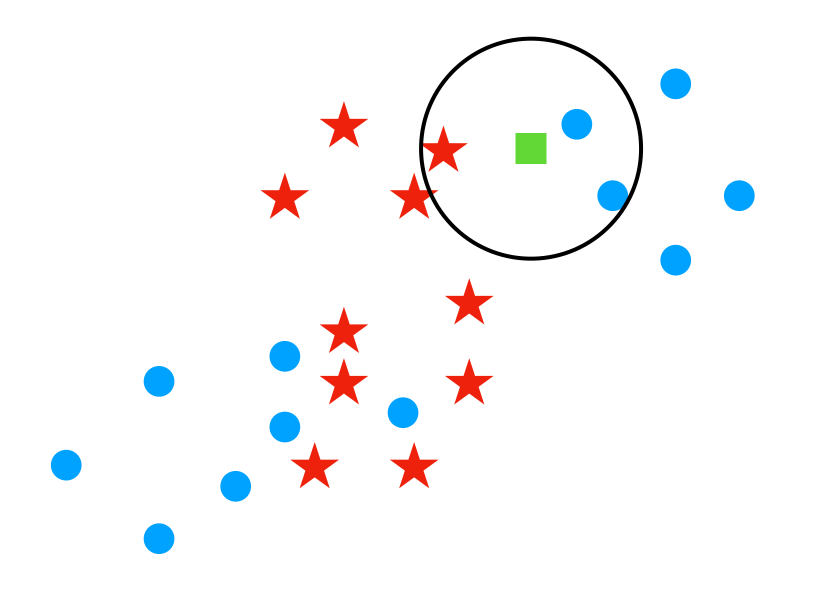
Cons:

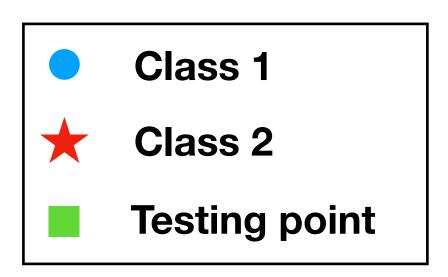
- Slow at query time
- Bad for high dimensional data
- Choice of local distance is crucial



k-Nearest Neighbor

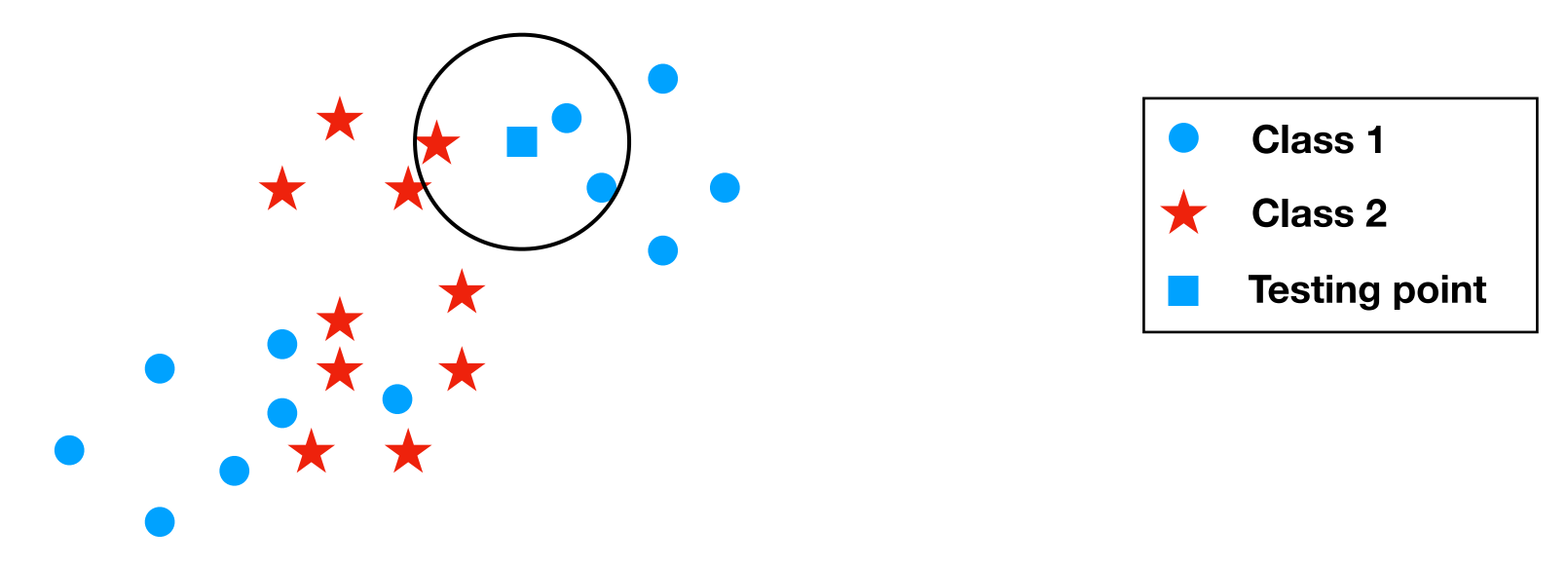
A new point X is classified by a majority vote among the k-nearest neighbor of X





k-Nearest Neighbor

A new point X is classified by a majority vote among the k-nearest neighbor of X

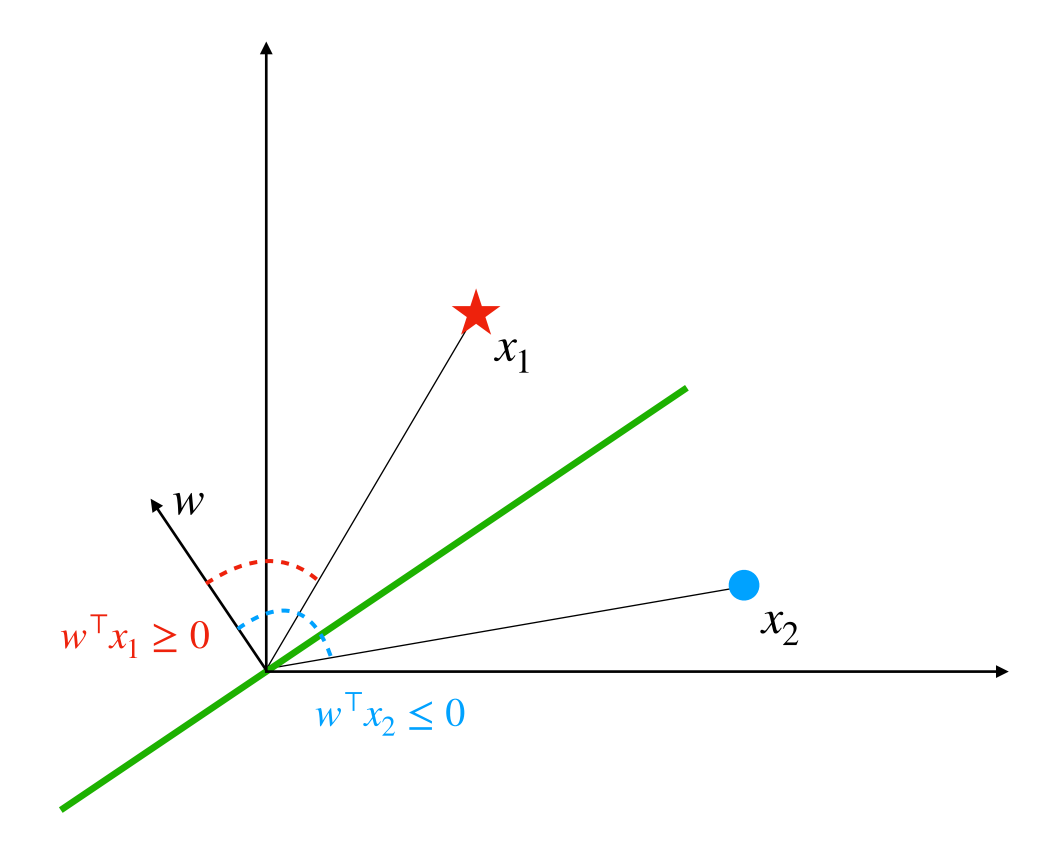


Generalization: smoothing kernels; weighted linear combination of elements

Linear Decision boundaries

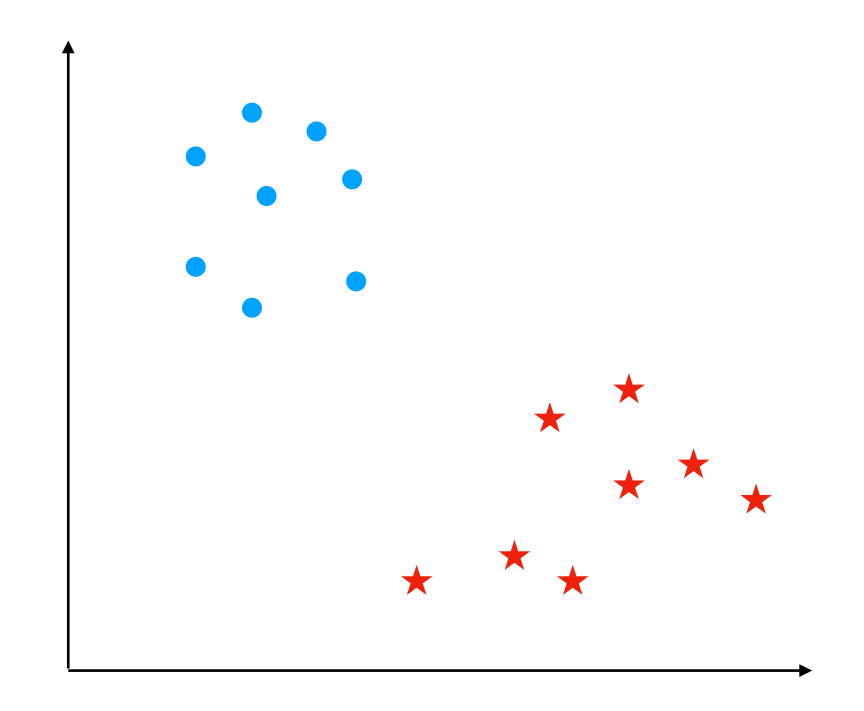
Assume we restrict ourself to linear decision boundaries (hyperplane):

 \rightarrow Prediction: $g(x) = \text{sign}(x^T w)$



Separating hyperplane

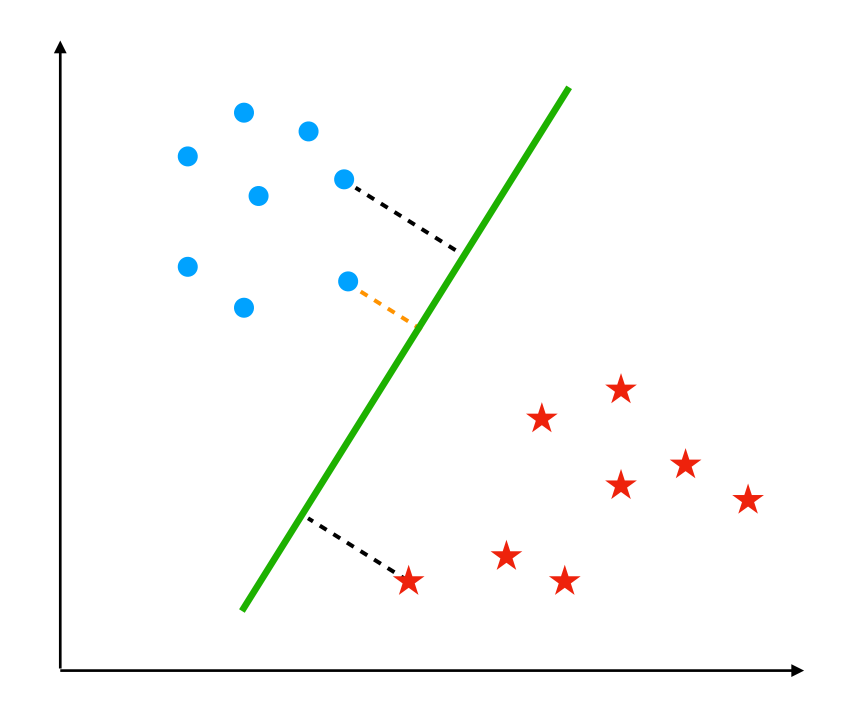
Assume we restrict ourself to linear decision boundaries (hyperplane): Assume the data are linearly separable, i.e, it exists a separating hyperplane



Which separating hyperplane would you pick?

Margin

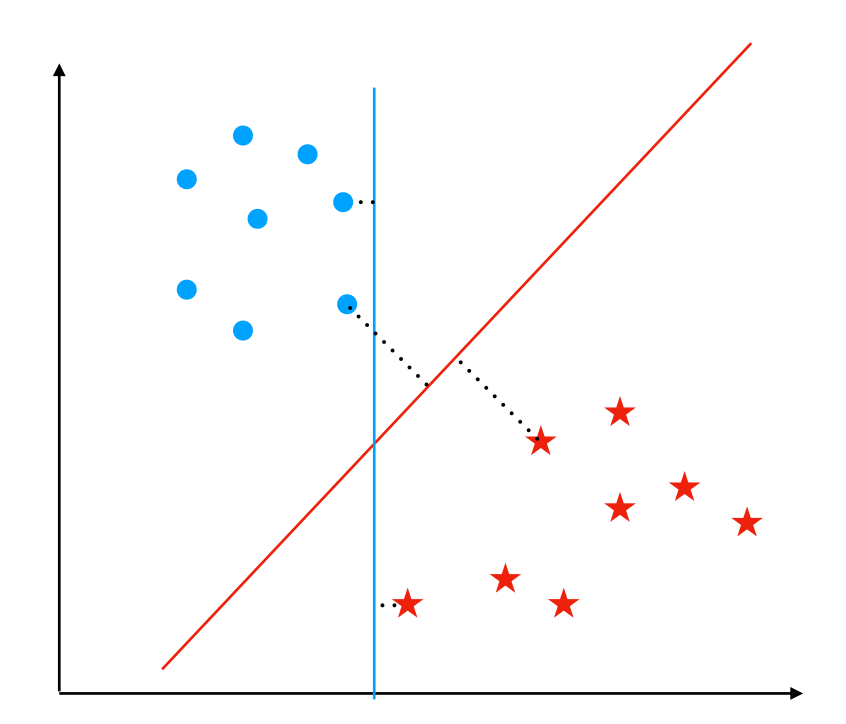
Key concept: The margin is the distance between the hyperplane and the closest point



→ Take the one with the largest margin!

Max-margin separating hyperplane

Pick the hyperplane which maximizes the margin

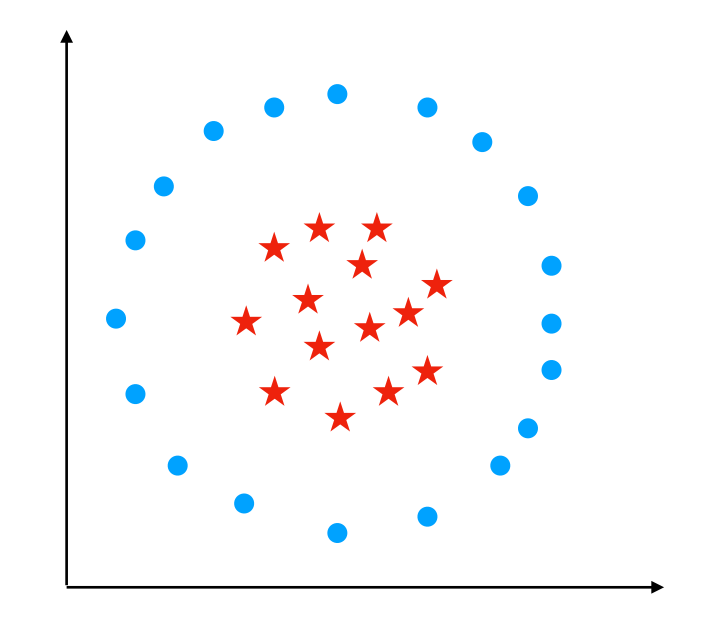


Why: If we slightly change the training set, the number of misclassification will stay low

→ It will lead us to support vector machine (SVM) and logistic regression

Non linear classifier

- Linear decision boundaries will not always works.
- Features augmentation (X, X^2, X^3, X^4)
- Kernel Method



Do we still have time?

A little bit of theory

$$(X, Y) \sim \mathcal{D}$$
 with $X \in \mathcal{X}, Y \in \mathcal{Y} = \{0,1\}$

Loss function:

$$\mathscr{E}(y, y') = 1_{y \neq y'} = \begin{cases} 1 & \text{if } y \neq y' \\ 0 & \text{if } y = y' \end{cases}$$

True risk for the classification:

$$L_{\mathcal{D}}(g) = \mathbb{E}_{\mathcal{D}}[1_{Y \neq g(X)}] = \mathbb{P}_{\mathcal{D}}[y \neq g(x)]$$
 classification error probability of making an error

Bayes classifier

What is the **optimal performance**, regardless of the finiteness of the training data?

Def: The classifier $g_* = \arg\min_{g} L_{\mathcal{D}}(g)$ is called the **Bayes classifier**

Claim:

$$g_*(x) = \arg \max_{y \in \{0,1\}} \mathbb{P}(Y = y | X = x)$$

Proof of the Bayes classifier

Claim 1: $\forall x \in \mathcal{X}, h_*(x) \in \arg\min_{y \in \mathcal{Y}} \mathbb{P}(Y \neq y \mid x) \implies h_* \in \arg\min_{h:\mathcal{X} \to \mathcal{Y}} L_{\mathcal{D}}(h)$

$$L_{\mathcal{D}}(h) = \mathbb{E}_{X,Y}[1_{Y \neq h(X)}] = \mathbb{E}_{X}[\mathbb{E}_{Y|X}[1_{Y \neq h(X)} | X = X]]$$

$$= \mathbb{E}_{X}[\mathbb{P}(Y \neq h(X) | X = X)]$$

$$\geq \mathbb{E}_{X}[\min_{y \in \mathcal{Y}} \mathbb{P}(Y \neq y | X = X)]$$

$$= \mathbb{E}_{X}[\mathbb{P}(Y \neq h_{*}(X) | X = X)] = \mathbb{E}_{X,Y}[1_{Y \neq h_{*}(X)}] = L_{\mathcal{D}}(h_{*})$$

Claim 2:
$$g_*(x) = \arg\min_{y \in \mathcal{Y}} \mathbb{P}(Y \neq y \mid x)$$

 $g_*(x) = \arg\max_{y \in \mathcal{Y}} \mathbb{P}(Y = y \mid x) = \arg\min_{y \in \mathcal{Y}} \mathbb{P}(Y \neq y \mid x)$

Bonus: a good regressor implies a good classifier

For all regression function $\eta:\mathcal{X} \to \mathbb{R}$ we can define a classifier as

$$\mathcal{X} \to \{0,1\}$$

$$g_{\eta}: x \mapsto 1_{\eta(x) \ge 1/2}$$

Claim:

$$L_{\mathcal{D}}^{\mathsf{classif}}(g_{\eta}) - L_{\mathcal{D}}^{\mathsf{classif}}(g^{*}) \leq 2\sqrt{L_{\mathcal{D}}^{\ell_{2}}(\eta) - L_{\mathcal{D}}^{\ell_{2}}(\eta^{*})}$$

$$\text{Where } L^{\text{classif}}_{\mathcal{D}}(g_{\eta}) = \mathbb{E}_{\mathcal{D}}[1_{g(X) \neq Y}], L^{\ell_2}_{\mathcal{D}}(f) = \mathbb{E}_{\mathcal{D}}[(Y - f(X))^2] \text{ and } \eta_* = \arg\min_{\eta} L^{\ell_2}_{\mathcal{D}}(\eta)$$

 \Rightarrow If η is good for regression then g_{η} is good for classification too (converse is not true)