STAT5003

Week 5: Introduction to classification techniques

Dr. Justin Wishart Semester 2, 2020





Readings



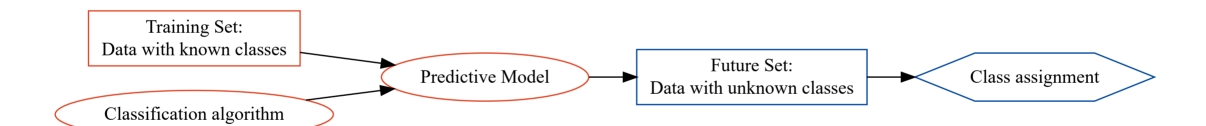
- Classification covered in Chapter 4 in James, Witten, Hastie, and Tibshirani (2013)
- Support Vector Machines covered in Chapter 9 in James, Witten, Hastie, et al. (2013)
- Optional for SVMs
 - Section 4.5.2 and Sections 14.1-14.3 in Hastie, Tibshirani, and Friedman (2017)

Classification



Basic principles of classification

- Each observation has two properties
 - A class label or response, y
 - $\circ~$ A feature vector (vector of predictor variables), $oldsymbol{x}=(x_1,x_2,\ldots,x_p)$
- Goal is to classify y using x



Classification vs Clustering

Clustering: classes are unknown, want to discover them from the data (unsupervised)

Classification: classes are predefined, want to use a (training or learning) set of labeled objects to form a classifier for classification of future observations (supervised)

Classification vs Regression

Regression: no class definition, the response variable is a continuous value. Model the relationship between explanatory variables and the response variable.

Classification: samples are predefined to be from a given class. Classification models produce a continuous valued prediction, which is usually in the form of a probability (i.e. the predicted values of class membership for any individual sample are between 0 and 1 and sum to 1). A predicted class is required in order to make a decision.

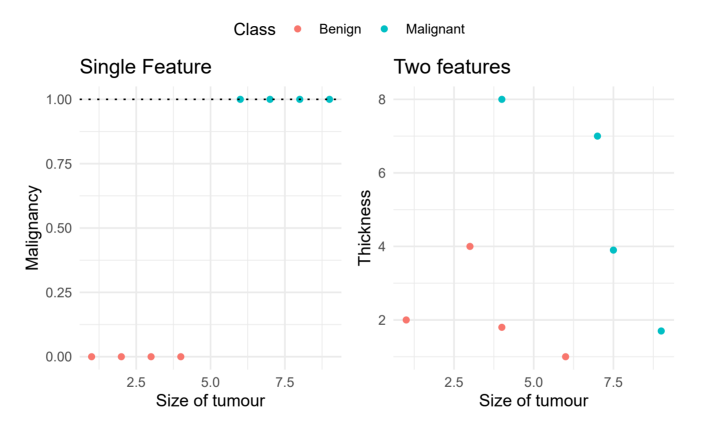
Classification algorithms to discuss

- Logistic Regression
- Linear discriminant analysis (LDA)
- *k*-nearest neighbours
- Supper vector machines (SVM)

Binary or Two class classification

- Binary in there are two possible values (0 or 1, TRUE or FALSE)
- Examples of binary classification:
 - Email: Spam / Not Spam
 - Tumour: Malignant /Benign
- ullet Labels are similarly described, $y \in \{0,1\}$
 - 0: "negative class"
 - 1: "positive class"

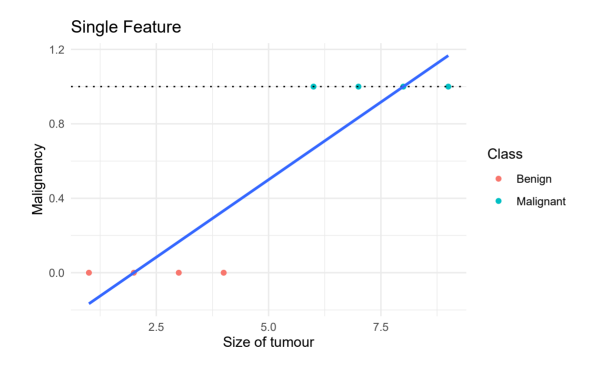
Problem setup



- Threshold classifier output $h_{\theta}(x)$ at 0.5:
 - $\circ \:$ if $h_{ heta}(x) > 0.5$, predict y = 1
 - $\circ \;$ if $h_{ heta}(x) < 0.5$, predict y = 0

Why not use simple linear regression?

• Y is the target value is a *binary* outcome.



- Linear regression is not constrained to 0 < y < 1 for all x
 - \circ What is the interpretation when $\hat{y}>1$ (or <0)

Linear regression misspecifications here

- The regression line $\beta_0 + \beta_1 x$ can span the entire real line
 - \circ all values between $-\infty$ to ∞
- In the tumour diagnosis problem, the target variable y only takes two values: 0 or 1.
- The linear regression model is not well specified for this purpose.

Logistic regression



Logistic regression

Previously we had the multiple regression

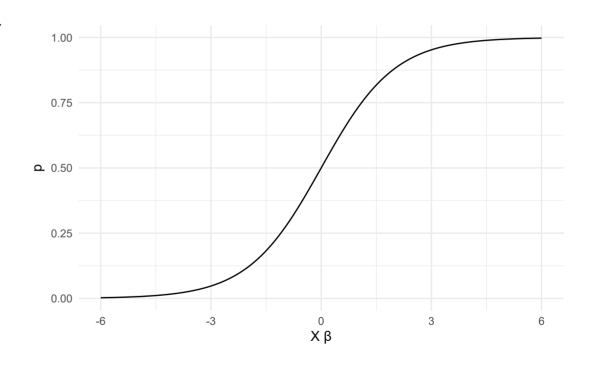
$$oldsymbol{Y} = oldsymbol{X}oldsymbol{eta} + oldsymbol{arepsilon} \ y = eta_0 + eta_1x_1 + eta_2x_2 + \dots + eta_px_p + arepsilon \ \mathbf{ heta}^Toldsymbol{x} = eta_0 + eta_1x_1 + eta_2x_2 + \dots + eta_px_p \$$

- \circ Could write this as $\mathbb{E}Y = heta^T oldsymbol{x} = \mu$
- ullet Can **generalise** this to $g(\mathbb{E}Y) = heta^T oldsymbol{x} = g(\mu)$
- Logistic regression is a special case of one of these generalised linear models.

$$ullet \log \left(rac{p}{1-p}
ight) = oldsymbol{ heta}^T oldsymbol{x}$$

• Solve for *p* gives

$$\stackrel{\circ}{p} = P(Y=1|oldsymbol{x}) = g^{-1}(oldsymbol{ heta}^Toldsymbol{x}) = rac{1}{1+\exp(-oldsymbol{ heta}^Toldsymbol{x})}$$



Logistic regression terminology

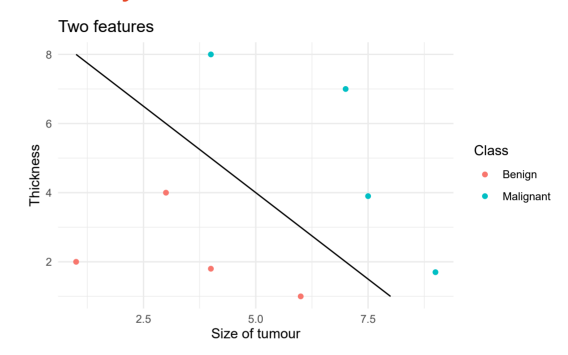
- Logistic function $\frac{1}{1+e^{-\theta^T x}}$
 - \circ Responsible from mapping the features from $(-\infty,\infty)=\mathbb{R}$ to (0, 1)
- Odds ratio: $\frac{p}{1-p}$
 - \circ Maps the probability from (0, 1) to $(0, \infty)$
- Log-odds or logit: $\log \left(\frac{p}{1-p} \right)$
- ullet In logistic regression we want the values in the logit space to be linear in X

Logistic regression: decision boundary

• Decision boundary

$$P(Y=1|oldsymbol{x}) = rac{1}{1+\exp(-oldsymbol{ heta}^Toldsymbol{x})}$$

• Predict Y = 1 if $\boldsymbol{\theta}^T \boldsymbol{x} \geq 0$



Linear Discriminant Analysis (LDA)



Linear Discriminant Analysis (LDA)

LDA undertakes the same task as Logistic Regression. It classifies data based on categorical variables

- Malignant or benign
- Making profit or not
- Buy a product or not
- Satisfied customer or not

Bayes' Theorem in the classification context

$$p_k(x) = extbf{ extit{P}(Y=k|X=x)} = rac{\pi_k f_k(x)}{\sum_{\ell=1}^K \pi_\ell f_\ell(x)}$$

Posterior: The probability of classifying observation to group k given it has features x

Prior: The prior probability of an observation in general belonging to group k

• $f_k(x) = P(X = x | Y = k)$ is the density function for feature x given it's in group k

Logistic Regression vs LDA formulations

ullet With Logistic Regression we modeled the probability of Y being from the $k^{
m th}$ class as

$$p_k(x)=P(Y=k|X=x)=rac{e^{eta_0+eta_1X}}{1+e^{eta_0+eta_1X}}$$

• Bayes' Theorem states

$$p_k(x) = P(Y=k|X=x) = rac{\pi_k f_k(x)}{\sum_{\ell=1}^K \pi_\ell f_\ell(x)}$$

 π_k : Probability of coming from class k (prior probability)

 $f_k(x)$: Density function for X given that X is an observation from class k.

LDA estimates of π_k and $f_k(x)$

- We can estimate π_k and $f_k(x)$ to compute $p_k(x)$
- The most common model for $f_k(x)$ is the Normal Density (LDA)

$$f_k(x) = rac{1}{\sigma\sqrt{2\pi}} \mathrm{exp}igg(-rac{(x-\mu_k)^2}{2\sigma^2}igg).$$

- Using the above density, we only need to estimate three quantities to compute $p_k(x)$
 - \circ That is, μ_k , σ_k^2 and π_k
- For simplicity, assume common variance.

Use training data set for estimation

- The mean $\widehat{\mu_k}$ could be estimated by the average of all training observations from the $k^{ ext{th}}$ class.
- The variance σ^2 could be estimated as the weighted average of variances of all k classes.
- The proportion π_k is estimated as the proportion of the training observations that belong to the $k^{\rm th}$ class.

$$egin{align} \widehat{\mu_k} &= rac{1}{n_k} \sum_{i:y_i = k} x_i \ & \ \sigma^2 &= rac{1}{n-K} \sum_{k=1}^K \sum_{i:y_i = k} (x_i - \widehat{\mu}_k)^2 \ & \ \end{pmatrix}$$

Simple example with one predictor

- Suppose we have only one predictor
- Two normal density function $f_1(x)$ and $f_2(x)$, represent two distinct classes
- The two density functions overlap, so there is some uncertainty about the class to which an observation with an unknown class belongs
- The dashed vertical line represents Bayes' decision boundary

Deriving LDA for one predictor

Assuming one predictor (and common variance)

$$f_k(x) = rac{1}{\sigma_k \sqrt{2\pi}} \mathrm{exp}igg(-rac{1}{2\sigma_k^2}(x-\mu_k)^2igg)$$

$$p_k(x) = P(Y=k|x) = rac{rac{\pi_k}{\sigma_k\sqrt{2\pi}} ext{exp}\Big(-rac{1}{2\sigma_k^2}(x-\mu_k)^2\Big)}{\sum_{l=0}^Krac{\pi_\ell}{\sigma_k\sqrt{2\pi}} ext{exp}\Big(-rac{1}{2\sigma_k^2}(x-\mu_k)^2\Big)}$$

• Find the class k which we maximize:

$$xrac{\mu_k}{\sigma^2} - rac{\mu_k^2}{2\sigma^2} + \log(\pi_k)$$

LDA Decision boundary

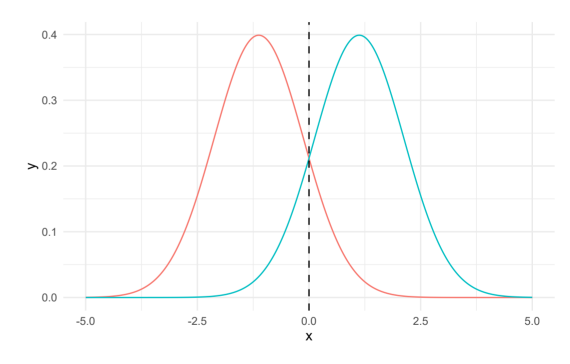
If K=2 and $\pi_1=\pi_2$, then assigns an observation to class 1 if $\log p_1(x)>\log p_2(x)\leadsto\log\Bigl(rac{p_1(x)}{p_2(x)}\Bigr)>0$

Substituting in the previous equation (assuming $\sigma_i = \sigma$) we have,

$$\log \left(rac{p_1(x)}{p_2(x)}
ight) > 0 \ \log(\pi_1) - \log(\pi_2) + rac{x\mu_1}{2\sigma^2} - rac{x\mu_2}{2\sigma^2} - rac{\mu_1^2}{2\sigma^2} + -rac{\mu_1^2}{2\sigma^2} > 0 \ 2x(\mu_1 - \mu_2) > \mu_1^2 - \mu_2^2$$

Decision boundary at

$$x=rac{\mu_1^2-\mu_2^2}{2(\mu_1-\mu_2)}=rac{\mu_1+\mu_2}{2}$$



Why not logistic regression?

- In the case where n is small, and the distribution of predictors X is approximately normal, then LDA is more stable than Logistic Regression
- LDA is more popular when we have more than two response classes. More intuitive to predict class assignment.
- When the classes are well separated, the parameter estimates for logistic regression are unstable. However, LDA doesn't suffer any stability issues in this case.

Logistic Regression vs LDA

Similarity:

Both Logistic Regression and LDA produce linear boundaries

Differences:

- LDA assumes that the observations are drawn from the normal distribution with common variance in each class, while logistic regression does not have this assumption.
- LDA would do better than Logistic Regression if the assumption of normality hold, otherwise logistic regression may outperform LDA

k-Nearest Neighbours (kNN)

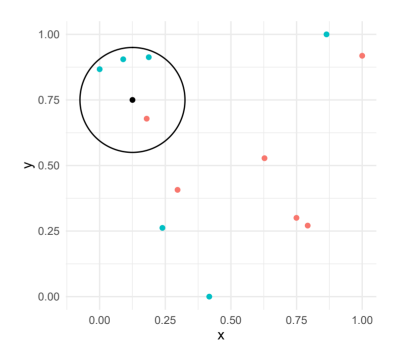


k-Nearest Neighbours

• kNN model is probability of an observation with features $m{x}$ belonging to group ℓ depends on the membership of the nearest points to $m{x}$

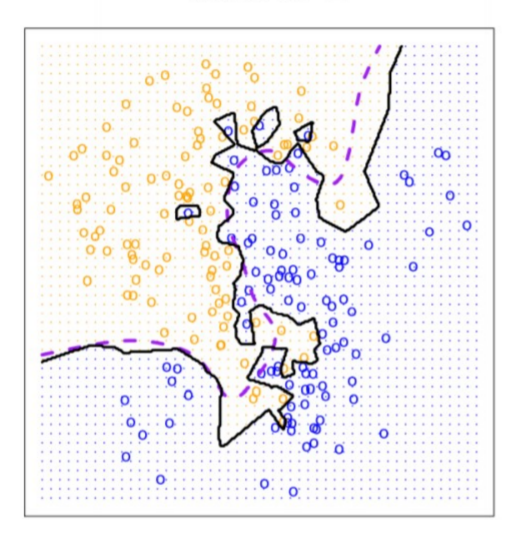
$$P(Y=\ell|m{x})=rac{1}{k}\sum_{N_x^k}1_{\{y=\ell\}}=rac{1}{k} imes ext{Count of the closest }k ext{ points that belong to group }\ell$$

 Suppose k = 4 is chosen. At the candidate black point. The four nearest neighbours are inspected. There is probability 3/4 of being in the green group and 1/4 for being in the orange group.

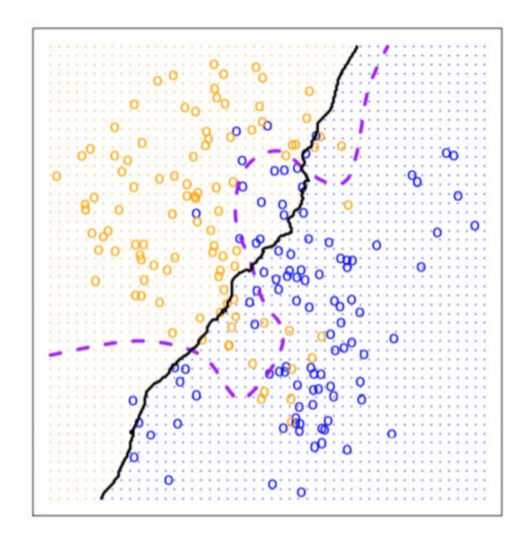


k-Nearest Neighbours

KNN: K=1



KNN: K=100



kNN vs (LDA and Logistic Regression)

- kNN takes a completely different approach
- kNN is completely non-parametric: No assumptions are made about the shape of the decision boundary
- Advantage of kNN: We can expect kNN to dominate both LDA and Logistic Regression when the decision boundary is highly non-linear
- Disadvantage of kNN: kNN does not tell us which predictors are important (no table of coefficients)

Support Vector Machines (SVM)



Support Vector Machines (SVM)

Basic idea behind SVM

Find a plane that separates the classes in the feature space.

- If a basic mathematical plane is not possible due to overlap
 - Relax the idea of complete separation into
 - Enrich and enlarge the feature space so that separation is possible
 - Think dimension expansion

What is a hyperplane?

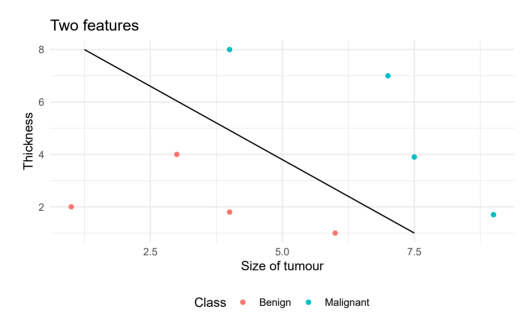
- In p dimensions it is a flat affine subspace of dimension p-1
- General equation has the form

$$\beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_p X_p = 0$$

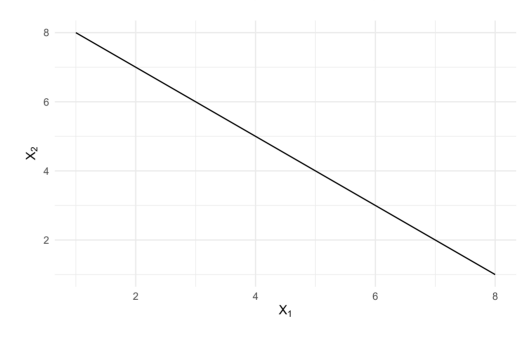
- In p=2 dimensions, the hyperplane is a line.
- If $\beta_0 = 0$, the hyperplane passes through the origin, otherwise it does not.
- The vector $(\beta_1, \beta_2, \dots, \beta_p)$ is called the normal vector
 - It points in a direction orthogonal to the surfance of the hyperplane

Hyperplane example

• Earlier hypothetical example

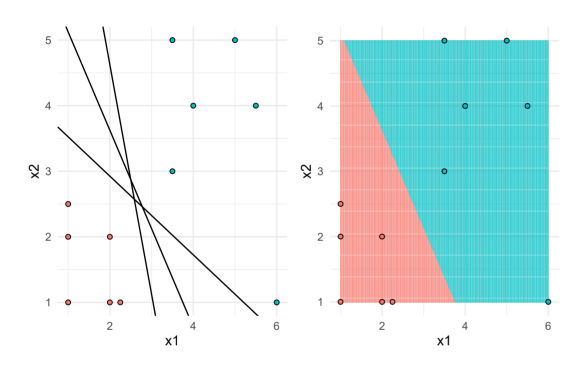


• Consider just the line (hyperplane)



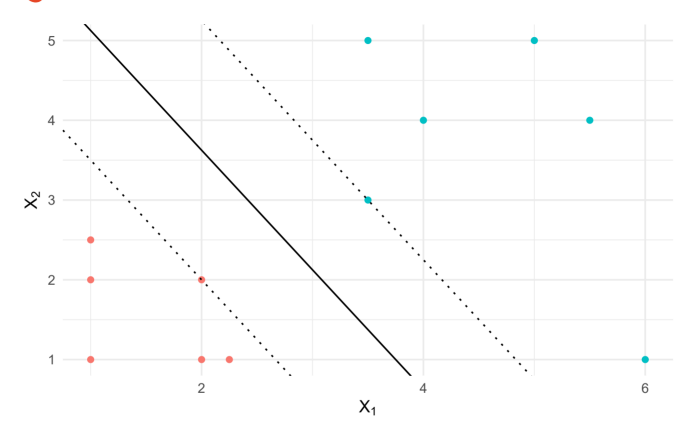
• Equation of the hyperplane here is $-9 + X_1 + X_2 = 0$

Separating hyperplanes



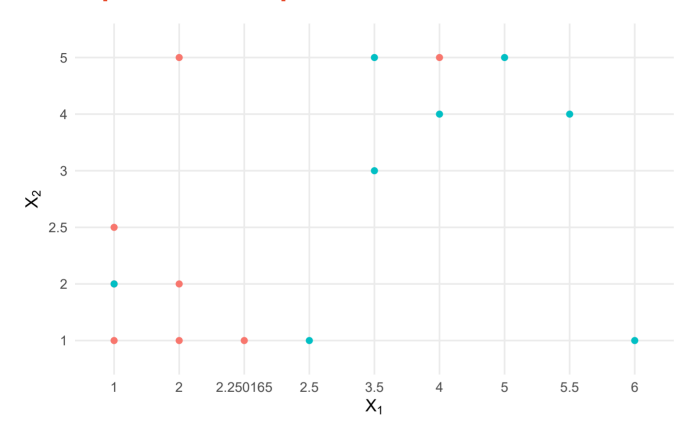
- ullet Consider coding Benign (red?) as $y_i=-1$ and malignant (blue?) as $y_i=1$
- Then $y_i f(x_i) > 0$ for all i, $f(x_i)$ defines a separating hyperplane.
- ullet If $f(x_i)=eta_0+eta_1x_1+\cdots+eta_px_p$ defines a hyperplane
 - $\circ f(x) > 0$ defines a region on one side of the hyperplane
 - $\circ \ f(x) < 0$ defines a region on one other side of the hyperplane

Maximal Margin Classifier



$$egin{aligned} \max_{eta_0,eta_1,eta_2,\dots,eta_p} & ext{such that } \sum_{j=1}^p eta_j^2 = 1 \ y_i(eta_0 + eta_1 x_{i1} + eta_2 x_{i2} + \dots eta_p x_{ip}) \geq M \end{aligned}$$

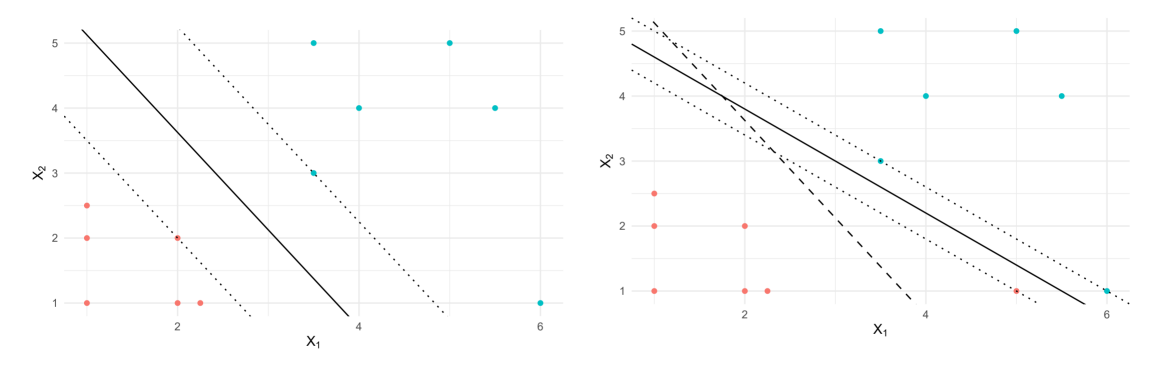
Non-perfect separation



- There is no linear boundary (hyperplane) that perfectly separates the classes.
- This is typically the case that observations don't have a perfect boundary of separation.
 - \circ Except in the case when n < p (more features than observations)

Effect of noisy data

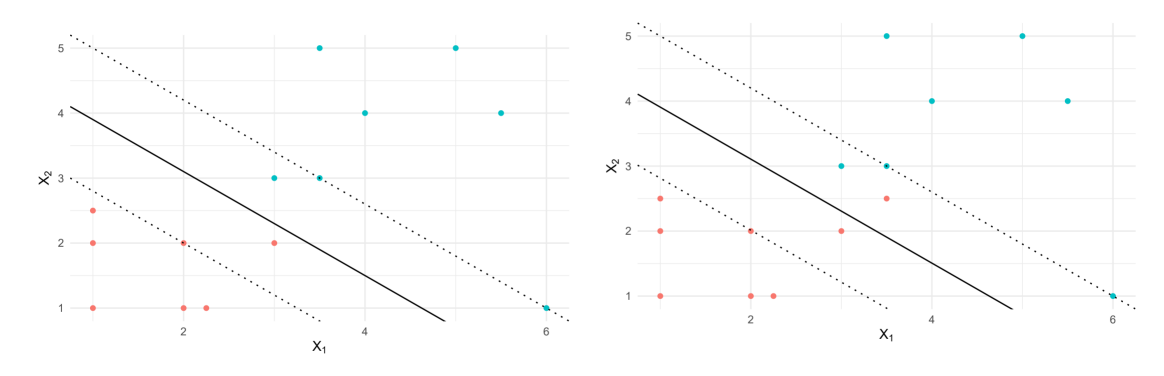
Consider the impact of one extra observation



- Data could be separable, but noisy → unstable solution for the maximal margin classifier.
- The support vector classifier maximizes a soft margin.
 - relaxes requirement for all observations to be on the correct side of the margin

Soft margin examples

• Observations allowed in the margin



- Observations on the correct side of hyperplane
- Allow observations on incorrect side of hyperplane

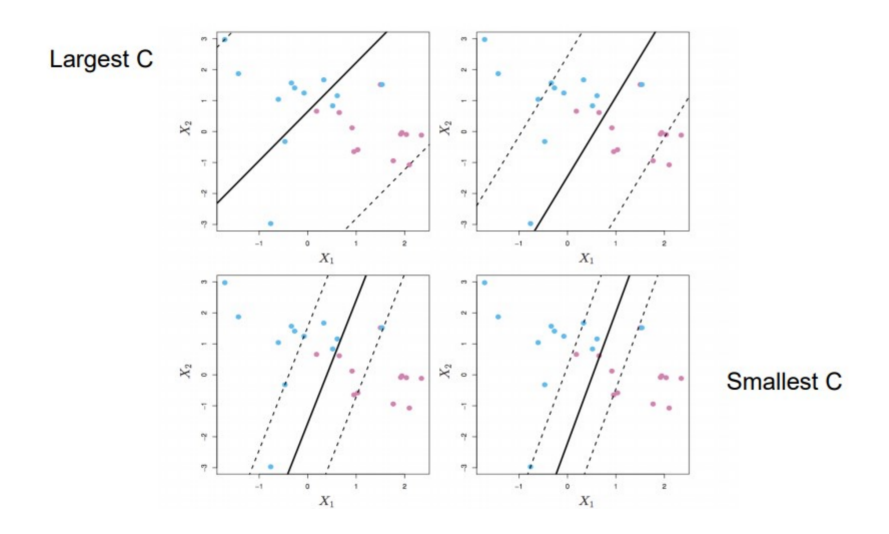
Support Vector Classifier

Support Vector Classifier solves the following optimization problem:

$$egin{aligned} \max_{eta_0,eta_1,eta_2,\ldots,eta_p,\epsilon_1,\ldots,\epsilon_n} M & ext{ such that } \sum_{j=1}^p eta_j^2 = 1 \ y_i(eta_0+eta_1x_{i1}+eta_2x_{i2}+\ldotseta_px_{ip}) \geq M(1-\epsilon_i) \ \epsilon_i \geq 0, \sum_{i=1}^n \epsilon_i \leq C \end{aligned}$$

- C is a non-negative tuning parameter,
- M is the width of the margin,
- ullet ϵ_i are slack variables that allow observations to be on the wrong side of the margin,
 - \circ if $\epsilon_i > 1$, then observation i is on the wrong side of the hyperplane
 - \circ if $0 < \epsilon_i \le 1$, then observation *i* is on the correct side but inside margin
 - \circ if $\epsilon_i = 0$, then observation i is on the correct side and past the margin.

Impact of cost parameter C



Limitations of support vector classifier

• Single linear boundary can be insufficient

Feature space expansion

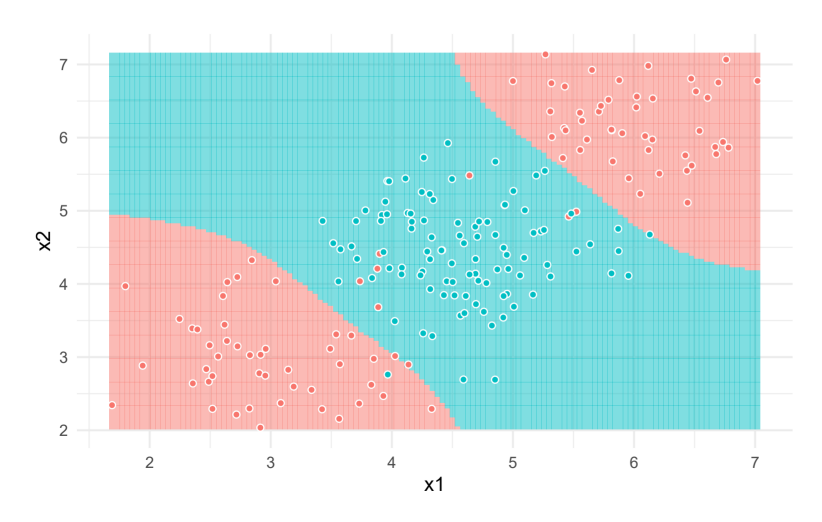
- Enlarge the space of features by including transformations:
 - $\circ \,$ e.g. new features that are powers and products $X_1^2, X_1^3, X_1 X_2$
 - \circ Hence go from p-dimensional space to P>p dimensional
- Fit (linear) support vector classifier in the expanded feature space.
 - o Impact is a **non-linear** decision boundary in original feature space.
- Example: Suppose we start off in 2-dimensional feature space (X_1, X_2) .
 - Make new feature space $(X_1, X_2, X_1^2, X_2^2, X_1X_2)$.
 - Then the decision boundary would be of the form:

$$eta_0 + eta_1 X_1 + eta_2 X_2 + eta_3 X_1^2 + eta_4 X_2^2 + eta_5 X_1 X_2 = 0$$

This leads to non-linear decision boundary in the original space (quadratic conic sections)

Sixth order polynomial

• Using degree six polynomial expansion



Non-linearity and kernels

- Polynomials get complicated and a burden very quickly as dimension increases.
- More elegant solution is to induce non-linear structure in Support vector classifier with kernels
- The elegance comes from the role of the **inner product** in the support vector classifier definition

Inner products and support vectors

- ullet Recall $oldsymbol{x}_i = (x_{i1}, x_{i2}, \dots, x_{ip}) \in \mathbb{R}^p$
- Inner product between vectors given by,

$$\langle oldsymbol{x}_i, oldsymbol{x}_j
angle = \sum_{k=1}^p x_{ik} x_{jk}$$

Recall the hyperplane equation

$$f(oldsymbol{x}) = eta_0 + eta_1 x_1 + eta_2 x_2 + \dots + eta_p x_p$$

• The linear support vector classifier can be represented as

$$f(oldsymbol{x}) = eta_0 + \sum_{j=1}^n lpha_j \langle oldsymbol{x}, oldsymbol{x}_j
angle$$

Support set

- Estimation of the parameters $\alpha_1, \alpha_2, \ldots, \alpha_n$ and β_0 required
 - All that is required are the inner products between pairs of training observations.
- Usually $\widehat{\alpha}_i = 0$ with the non-zero values occurring on the support vectors
 - I.e. ones that lie on the margin.

$$f(oldsymbol{x}) = eta_0 + \sum_{j \in S} lpha_j \langle oldsymbol{x}, oldsymbol{x}_j
angle$$

ullet Here, S is the support set of indices such that $lpha_j>0$

Kernel functions

Suppose now we replace the inner product with a generalized function of the form

$$K(oldsymbol{x}_i,oldsymbol{x}_j)$$

This function is called a kernel.

In this context is quantifies the similarity of two observations.

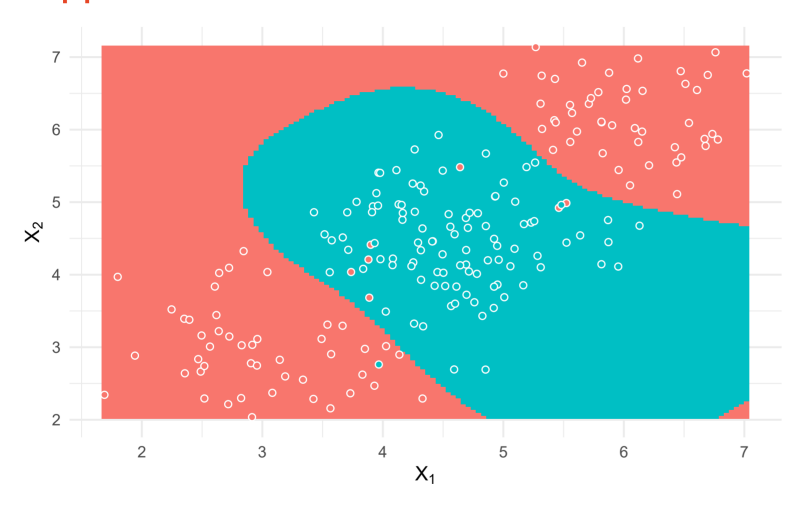
- Examples
 - Polynomial kernel

$$K(oldsymbol{x}_i,oldsymbol{x}_j) = ig(1+\langleoldsymbol{x}_i,oldsymbol{x}_j
angleig)^d$$

Gaussian radial kernel

$$K(oldsymbol{x}_i,oldsymbol{x}_j) = \exp\left(-\gamma + \sum_{k=1}^p (x_{ik} - x_{jk})^2
ight)^d$$

Support vector machines with the radial kernel



SVM with more than two classes

ullet SVM covered previously are design for binary classification. To expand this to K classes there are two options

1. One vs all:

- \circ Fit K different binary classifiers, $f_k(x)$ for $k=1,2,\ldots,K$ where each boundary attempts to separate class k vs the rest.
- Then x_i is classified to k^* where $f_{k^*}(x_i) > f_j(x_i)$ for all $j \neq k^*$. (i.e. the largest distance from the boundary).

2. One vs one:

- Fit all $\binom{K}{2}$ pairwise classifiers
- \circ Fit \boldsymbol{x}_i to the class that wins the most pairwise comparisons.
- Which to use?
 - \circ If K is small, do one vs one. Otherwise recommended One vs all.

References

Hastie, T, R. Tibshirani, and J. Friedman (2017). *The elements of statistical learning: data mining, inference, and prediction.* Second Edition, 12th printing. Springer Science & Business Media.

James, G, D. Witten, T. Hastie, et al. (2013). *An introduction to statistical learning*. Vol. 112. Springer.