

# Feature selection

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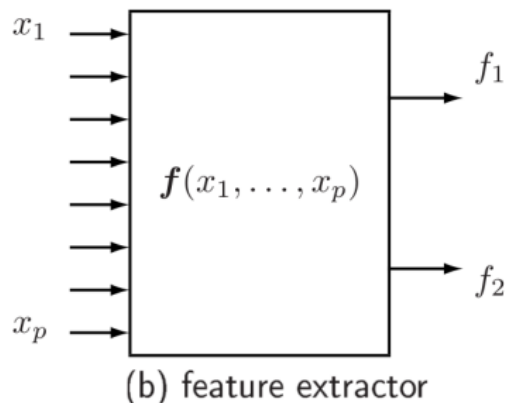
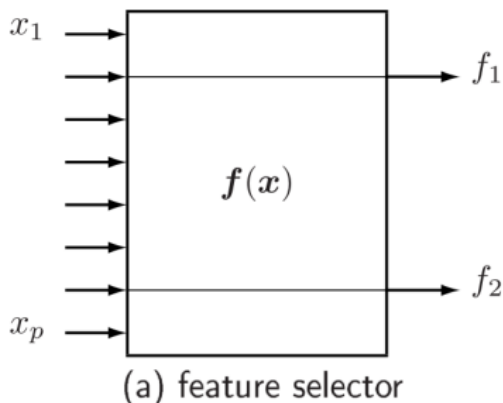
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# Feature selection

Feature selection is a process of selecting a subset of original features with minimum loss of information related to final task (classification, regression, etc.)



# Applications of feature selection

- Why feature selection?
  - increase predictive accuracy of classifier
  - improve optimization stability by removing multicollinearity
  - increase computational efficiency
  - reduce cost of future data collection
  - make classifier more interpretable
- Not always necessary step:
  - some methods have implicit feature selection:

# Applications of feature selection

- Why feature selection?
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- Not always necessary step:
  - some methods have implicit feature selection:
    - decision trees and tree-based (RF, ERT, boosting)
    - L1 regularization

## Types of features

Define  $f$  - the feature,  $F = \{f_1, f_2, \dots, f_D\}$  - full set of features,  
 $\tilde{F} = F \setminus \{f\}$ .

- **Strongly relevant feature:**

$$p(y|f, \tilde{F}) \neq p(y|\tilde{F})$$

- **Weakly relevant feature:**

$$p(y|f, \tilde{F}) = p(y|\tilde{F}), \text{ but } \exists S \subset \tilde{F} : p(y|f, S) \neq p(y|S)$$

- **Irrelevant feature:**

$$\forall S \subset \tilde{F} : p(y|f, S) = p(y|S)$$

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- **Irrelevant feature:**

$$\forall S \subset \tilde{F} : p(y|f, S) = p(y|S)$$

### Aim of feature selection

Find minimal features subset  $S \subset F$  such that  $P(y|S) \approx P(y|F)$ ,  
i.e. leave only *relevant* and *non-redundant* features.

# Categorization of feature selection algorithms

- Completeness of search:
  - Complete
    - exhaustive search complexity is  $2^D$ .
    - may be not exhaustive under certain conditions on  $J(S)$ <sup>1</sup>
  - Suboptimal
    - deterministic
    - random (deterministic with randomness / completely random)
- Integration with final predictor
  - independent (filter methods)
  - uses predictor quality (wrapper methods)
  - is embedded inside predictor (embedded methods)

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<sup>1</sup> $J(S)$  is a score of feature subset  $S$ .

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  - Feature subset generation
  - Feature importance estimation
- 2 Simultaneous feature selection specification

# Individual feature importances approach

- Estimate importances for individual features  $I(f_1), I(f_2), \dots, I(f_D)$ .
- Generate feature subset based on importances.

## 1 Individual feature importances approach

- Feature subset generation
- Feature importance estimation

## Incomplete search with suboptimal solution

- Order features with respect to feature importances  $I(f)$ :

$$I(f_1) \geq I(f_2) \geq \dots \geq I(f_D)$$

option 1: select top  $m$

$$\hat{F} = \{f_1, f_2, \dots, f_m\}$$

option 2: select best set from nested subsets:

$$S = \{\{f_1\}, \{f_1, f_2\}, \dots, \{f_1, f_2, \dots, f_D\}\}$$

$$\hat{F} = \arg \max_{F \in S} J(F)$$

- Comments:

- simple to implement
- when features are correlated, it will take many redundant features

## 1 Individual feature importances approach

- Feature subset generation
- Feature importance estimation

# Application of feature importances

- Feature importances can be used:
  - for feature selection
  - for rescaling features for adapting their impact on the model:
    - e.g.: in K-NN, in linear methods with regularization
  - for adapting feature sampling probability in random forest, extra random trees.

# Correlation

- two class:

$$\rho(f, y) = \frac{\sum_i (f_i - \bar{f})(y_i - \bar{y})}{[\sum_i (f_i - \bar{f})^2 \sum_i (y_i - \bar{y})^2]^{1/2}} = \frac{a}{b}$$

- multiclass  $\omega_1, \omega_2, \dots \omega_C$  (micro averaged  $\rho(f, y_c)$   $c = 1, 2, \dots C$ )

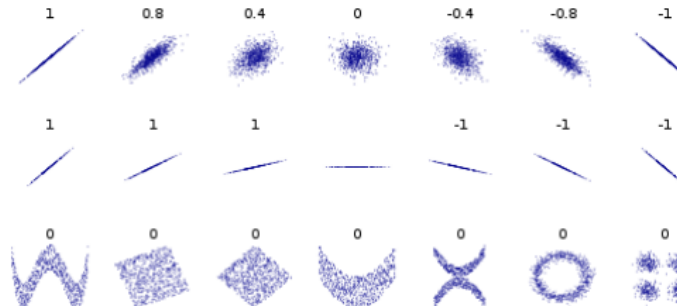
$$R^2 = \frac{\sum_{c=1}^C [\sum_i (f_i - \bar{f})(y_{ic} - \bar{y}_c)]^2}{\sum_{c=1}^C \sum_i (f_i - \bar{f})^2 \sum_i (y_{ic} - \bar{y}_c)^2} = \frac{\sum_c a_c^2}{\sum_c b_c^2}$$

- Benefits:

- simple to compute
- applicable both to continuous and discrete features/output.
- does not require calculation of probability density function.

# Correlation for non-linear relationship

- Correlation captures only linear relationship.
- Example: consider  $X$ -random variable, with  $\mathbb{E}X = 0$ ,  $\mathbb{E}X^3 = 0$  and random variable  $Z = X^2$ . Then  $X, Z$  are uncorrelated but dependent.
- Other examples of data and its correlation:



- Correlation between ranks.

# Defintitions

- Entropy<sup>2</sup> of random variable  $Y$ :

$$H(Y) := - \sum_y p(y) \ln p(y)$$

- Conditional entropy of  $Y$  after observing  $X$ :

$$H(Y|X) := - \sum_x p(x) \sum_y p(y|x) \ln p(y|x)$$

- Kullback-Leibler divergence for two p.d.f.  $P(x)$  and  $Q(x)$ :

$$KL(P||Q) := \sum_x P(x) \ln \frac{P(x)}{Q(x)}$$

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<sup>2</sup>measures level of uncertainty of r.v.  $Y$

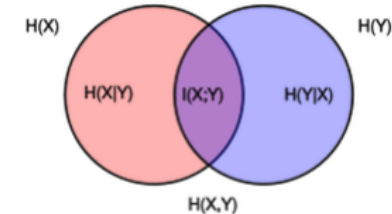
# Mutual information

Mutual information measures how much r.v.  $X$  and  $Y$  share information between each other:

$$MI(X, Y) := \sum_{x,y} p(x,y) \ln \left[ \frac{p(x,y)}{p(x)p(y)} \right] = KL(p(x,y)||p(x)p(y))$$

Properties:

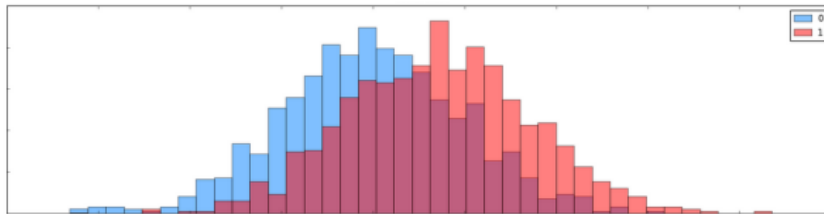
- $MI(X, Y) = MI(Y, X)$
- $MI(X, Y) = KL(p(x,y)||p(x)p(y)) \geq 0$
- $MI(X, Y) = H(Y) - H(Y|X)$
- $MI(X, Y) \leq \min \{H(X), H(Y)\}$
- $X, Y$ - independent  $\Leftrightarrow MI(X, Y) = 0$   
(for discrete r.v.)
- $X$  completely identifies  $Y$ , then  
 $MI(X, Y) = H(Y) \leq H(X)$



# Mutual information for feature selection

- Normalized variant  $NMI(X, Y) = \frac{MI(X, Y)}{H(Y)}$  equals
  - zero, when  $P(Y|X) = P(Y)$
  - one, when  $X$  completely identifies  $Y$ .
- Properties of  $MI$  and  $NMI$ :
  - identifies arbitrary non-linear dependencies
  - requires calculation of probability distributions
  - continuous variables need to be discretized

# importance based on probabilistic distance



Measure of feature  $f$  importance - distance between  $p(f|y = 0)$  and  $p(f|y = 1)$ , e.g. total variation:

$$\int |p(x|y = 1) - p(x|y = 0)| dx$$

# Relief criterion: 1-NN

**INPUT:**

Training set  $(x_1, y_1), (x_2, y_2), \dots (x_N, y_N)$

Number of neighbours  $K$

Distance metric  $\rho(x, x')$  # usually Euclidean

**for each** object  $x_n, y_n$ :

calculate nearest neighbour of the same class  $x_{s(n)}$

calculate  $K$  nearest neighbour of class  $x_{d(n)}$

**for each** feature  $f_i$  in  $f_1, f_2, \dots f_D$ :

calculate importance  $R(f_i) = \frac{1}{N} \sum_{n=1}^N \frac{|x_n^i - x_{d(n)}^i|}{|x_n^i - x_{s(n)}^i|}$

**OUTPUT:**

feature importances  $R$

# Relief criterion: K-NN

## INPUT:

Training set  $(x_1, y_1), (x_2, y_2), \dots (x_N, y_N)$

Number of neighbours  $K$

Distance metric  $\rho(x, x')$  # usually Euclidean

## **for each** object $x_n, y_n$ :

calculate  $K$  nearest neighbours of the same class  $y_n$ :

$x_{s(n,1)}, x_{s(n,2)}, \dots x_{s(n,K)}$

calculate  $K$  nearest neighbours of class other than  $y_n$ :

$x_{d(n,1)}, x_{d(n,2)}, \dots x_{d(n,K)}$

## **for each** feature $f_i$ in $f_1, f_2, \dots f_D$ :

calculate importance  $R(f_i) = \frac{1}{N} \sum_{n=1}^N \sum_{k=1}^K \frac{|x_n^i - x_{d(n,k)}^i|}{|x_n^i - x_{s(n,k)}^i|}$

## OUTPUT:

feature importances  $R$

# Tree feature importances

- Tree feature importances (`clf.feature_importances_` in `sklearn`).
  - Consider feature  $f$
  - Let  $T(f)$  be the set of all nodes, relying on feature  $f$  when making split.
  - efficiency of split at node  $t$ :  $\Delta I(t) = I(t) - \sum_{c \in \text{children}(t)} \frac{n_c}{n_t} I(c)$
  - feature importance of  $f$ :  $\sum_{t \in T(f)} n_t \Delta I(t)$
- Alternative: difference in decision tree prediction quality for
  - ① original validation set
  - ② validation set with  $j$ -th feature randomly shuffled

# Feature importances from linear model

- Feature importances from linear classification:
  - ① fit linear classifier with regularization to data
    - features should be normalized
  - ② retrieve  $w$  (`clf.coef_` in scikit-learn)
  - ③ importance of feature  $f_i$  is equal to  $|w_i|$ .

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  - Sequential search subset generation
  - Genetic search subset generation

# Simultaneous feature selection specification

- Need to specify:
  - quality criteria  $J(S)$  for any feature subset  $S$ 
    - typically: quality of model with these features (wrapper approach)
  - feature subset generation method  $S_1, S_2, S_3, \dots$

## 2 Simultaneous feature selection specification

- Sequential search subset generation
- Genetic search subset generation

# Sequential search

- Sequential forward selection algorithm:

- init:  $k = 0, F_0 = \emptyset$
- while  $k < \max\_features$ :
  - $f_{k+1} = \arg \max_{f \in F} J(F_k \cup \{f\})$
  - $F_{k+1} = F_k \cup \{f_{k+1}\}$
  - if  $J(F_{k+1}) < J(F_k)$ : break
  - $k=k+1$
- return  $F_k$

- Variants:

- sequential backward selection
- up-k forward search
- down-p backward search
- up-k down-p composite search
- up-k down-(variable step size) composite search

2 Simultaneous feature selection specification

- Sequential search subset generation
- Genetic search subset generation

# Genetic<sup>3</sup> algorithms

- Each feature set  $F = \{f_{i(1)}, f_{i(2)}, \dots, f_{i(K)}\}$  is represented using binary vector  $[b_1, b_2, \dots, b_D]$  where  $b_i = \mathbb{I}[f_i \in F]$
- Genetic operations:

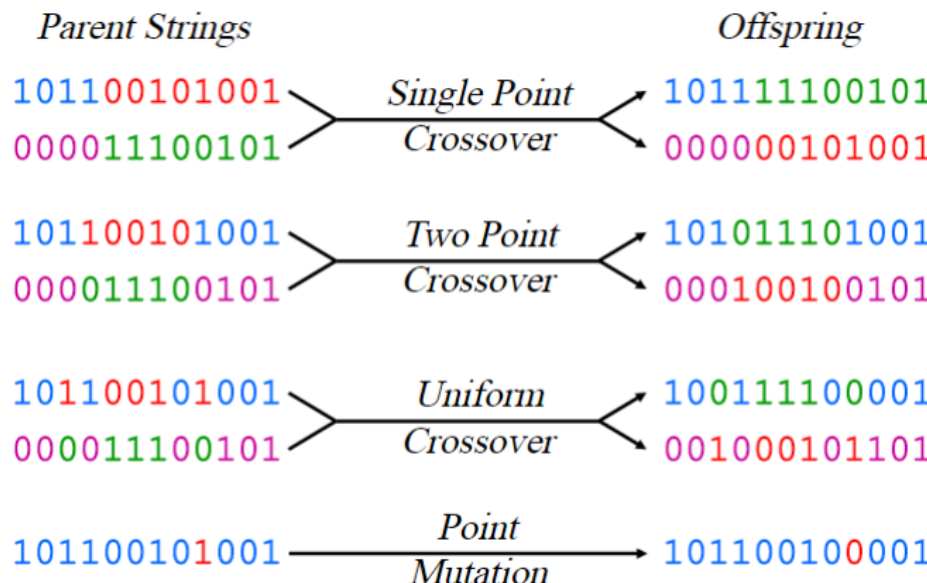
- $crossover(b^1, b^2) = b$ , where  $b_i = \begin{cases} b_i^1 & \text{with probability } \frac{1}{2} \\ b_i^2 & \text{otherwise} \end{cases}$

- $mutation(b^1) = b$ , where  $b_i = \begin{cases} b_i^1 & \text{with probability } 1 - \alpha \\ \neg b_i^1 & \text{with probability } \alpha \end{cases}$   
for some small  $\alpha$ .

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<sup>3</sup>Name inspired by genetic inheritance in biology.

# Genetic operations: demo



# Genetic algorithms

**INPUT:**

population size  $B$  and expanded population size  $B'$   
parameters of mutation and crossover  
maximum number of iterations  $T$ , minimum quality change  $\Delta J$

**ALGORITHM:**

generate  $B$  feature sets  $S_1, S_2, \dots, S_B$  randomly.

set  $t = 1$ ,  $P^0 = \{S_1, S_2, \dots, S_B\}$ ,  $J^0 = J(P^0)$

**while**  $t \leq T$  and  $|J^t - J^{t-1}| > \Delta J$ :

    modify  $P^{t-1}$  using crossover and mutation:

$S'_1, S'_2, \dots, S'_{B'} = \text{modify}(P^{t-1} | \theta)$

    order transformed sets by decreasing quality:

$J(S'^t_{i(1)}) \geq J(S'^t_{i(2)}) \geq \dots \geq J(S'^t_{i(B')})$

    set next population to consist of best representatives:

$P^t = \{S'_{i(1)}, S'_{i(2)}, \dots, S'_{i(B)}\}$

    set  $J^t = \max_{S \in P^t} J(S)$

$t = t + 1$

**OUTPUT:** suboptimal set of feature sets  $P^t$

# Modifications of genetic algorithm

- Preserve best features and best feature subsets:
  - Augment  $P'^t$  with  $K$  best representatives from  $P^{t-1}$ .
  - Make mutation probability lower for good features (that frequently appear in inside representatives).
- Increase breadth of search:
  - Crossover between more than two parents
- To prevent convergence to local optimum:
  - simultaneously modify several populations and allow rare random transitions between them.

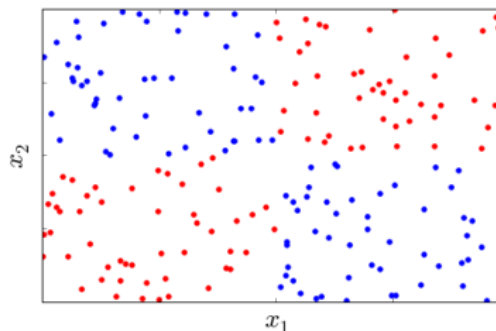
## Importance in context

Individually features do not affect  $y$ :

$$p(y|x^2) = p(y), \quad p(y|x^2) = p(y)$$

but may be relevant together:

$$p(y|x^1, x^2) \neq p(y)$$



Which methods will extract features relevant in context but irrelevant individually? [Decision tree \(?\)](#)