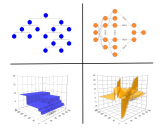
Important Learning Algorithms in ML





Learning goals

- General idea of important ML algorithms
- Overview of strengths and weaknesses

CONTENTS

- k-Nearest Neighbors (k-NN)
- Generalized Linear Models (GLM)
- Generalized Additive Models (GAM)
- Classification & Regression Trees (CART)
- Random Forests
- Gradient Boosting
- Linear Support Vector Machines (SVM)
- Nonlinear Support Vector Machines
- Neural Networks (NN)



K-NN - METHOD SUMMARY

REGRESSION

CLASSIFICATION

NONPARAMETRIC

WHITE-BOX

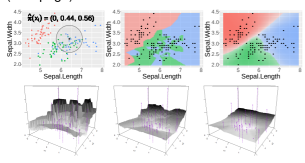
General idea

- similarity in feature space (w.r.t. certain distance metric
 d(x⁽ⁱ⁾, x)) → similarity in target space
- **Prediction** for **x**: construct k-neighborhood $N_k(\mathbf{x})$ from k points closest to **x** in \mathcal{X} , then predict
 - (weighted) mean target for **regression**: $\hat{y} = \frac{1}{\sum\limits_{i:\mathbf{x}^{(i)} \in N_k(\mathbf{x})} w_i} \sum\limits_{i:\mathbf{x}^{(i)} \in N_k(\mathbf{x})} w_i y^{(i)}$ with $w_i = \frac{1}{d(\mathbf{x}^{(i)},\mathbf{x})}$
 - \rightarrow optional: higher weights w_i for close neighbors
 - most frequent class for classification: $\hat{y} = \underset{\ell \in \{1, ..., g\}}{\operatorname{arg max}} \sum_{i: \mathbf{x}^{(i)} \in N_k(\mathbf{x})} \mathbb{I}(y^{(i)} = \ell)$
 - \Rightarrow Estimating posterior probabilities as $\hat{\pi}_{\ell}(\mathbf{x}^{(i)}) = \frac{1}{k} \sum_{i:\mathbf{x}^{(i)} \in N_k(\mathbf{x})} \mathbb{I}(\mathbf{y}^{(i)} = \ell)$
- Nonparametric behavior: no compression of information
- Not immediately interpretable



K-NN - METHOD SUMMARY

Hyperparameters Neighborhood **size** k (locality), **distance** metric (next page)

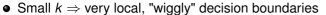


Classification

Left: Neighborhood for exemplary observation in iris, k=50 Middle: Prediction surface for k=1 Right: Prediction surface for k=50

Regression

Left: Prediction surface for k=3 Middle: Prediction surface for k=7 Right: Prediction surface for k=15



• Large $k \Rightarrow$ rather global, smooth decision boundaries



K-NN – METHOD SUMMARY

Popular distance metrics

• Numerical feature space: Typically, Minkowski distances

$$d(\mathbf{x}, \tilde{\mathbf{x}}) = \|\mathbf{x} - \tilde{\mathbf{x}}\|_q = \left(\sum_j |x_j - \tilde{x}_j|^q\right)^{\frac{1}{q}}$$

- q = 1: **Manhattan** distance $\rightarrow d(\mathbf{x}, \tilde{\mathbf{x}}) = \sum_{j} |x_j \tilde{x}_j|$
- q=2: **Euclidean** distance $\rightarrow d(\mathbf{x}, \tilde{\mathbf{x}}) = \sqrt{\sum_{j} (x_{j} \tilde{x}_{j})^{2}}$





Manhattan vs. Euclidean (green)

https://es.m.wikipedia.org/wiki/Archivo:Manhattan_distance.svg

- Mixed feature space:
 - Gower distance for numerical, categorical and missing data:

- numerical:
$$d(x_i, x_j) = \frac{|x_i - x_j|}{\max(x) - \min(x)}$$

- categorical:
$$d(x_i, x_j) = \begin{cases} 1, & \text{if } x_i \neq x_j \\ 0, & \text{if } x_i = x_j \end{cases}$$

- Gower distance as average over individual scores
- Optional weighting for beliefs about varying feature importance

K-NN – IMPLEMENTATION & PRACTICAL HINTS

Preprocessing Features should be standardized or normalized

Implementation

- R: mlr3 learners (calling kknn::kknn())
 - Classification:
 - LearnerClassifKKNN
 - -fnn::knn()
 - Regression:
 - LearnerRegrKKNN
 - -fnn::knn.reg()
 - Nearest Neighbour Search in O(N log N): RANN::nn2()



K-NN – IMPLEMENTATION & PRACTICAL HINTS

X O O X

- **Python:** From package sklearn.neighbors
 - Classification:
 - KNeighborsClassifier()
 - RadiusNeighborsClassifier() as alternative if data not uniformly sampled
 - Regression:
 - KNeighborsRegressor()
 - RadiusNeighborsRegressor() as alternative if data not uniformly sampled

K-NN - PROS & CONS

Advantages

- + Algorithm **easy** to explain and implement
- assumptions

 → able to model data of arbitrary complexity

No distributional or functional

- + No training or optimization required
- + **local model** \rightarrow **nonlinear** decision boundaries
- + Easy to tune (few hyperparameters)
 → number of neighbors k, distance metric
- Custom distance metrics can often be easily designed to incorporate domain knowledge

Disadvantages

- Sensitivity w.r.t. noisy or irrelevant features and outliers due to dependency on distance measure
- Heavily affected by curse of dimensionality
- Bad performance when feature scales are not consistent with feature relevance
- Poor handling of data **imbalances** (worse for more global model, i.e., large k)



REGRESSION

CLASSIFICATION

PARAMETRIC

WHITE-BOX

FEATURE SELECTION

General idea Represent target as function of linear predictor $\boldsymbol{\theta}^{\top} \mathbf{x}$ (weighted sum of features)

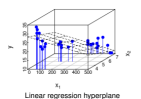
 \rightarrow **Interpretation:** if feature x_j increases by 1 unit, the linear predictor changes by θ_j units

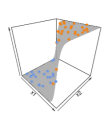
Hypothesis space $\mathcal{H} = \{ f : \mathcal{X} \to \mathbb{R} \mid f(\mathbf{x}) = \phi(\boldsymbol{\theta}^{\top}\mathbf{x}) \}$, with suitable transformation $\phi(\cdot)$, e.g.,

- ullet Linear Regression: $\mathcal{Y} = \mathbb{R}$, ϕ identity
- Logistic Regression: $\mathcal{Y} = \{0, 1\}$, logistic sigmoid $\phi(\boldsymbol{\theta}^{\top}\mathbf{x}) = \frac{1}{1 + \exp(-\boldsymbol{\theta}^{\top}\mathbf{x})} =: \pi(\mathbf{x} \mid \boldsymbol{\theta})$
 - ⇒ Decision rule: Linear hyperplane

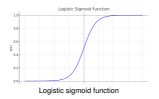


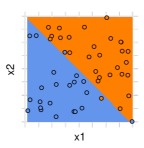
Loss functions





Logistic function for bivariate input and loss-minimal heta





Corresponding separating hyperplane



- Lin. Regr.:
 - Typically, based on quadratic loss (OLS estimation):

$$L(y, f) = (y - f)^2$$

- Log. Regr.: Based on bernoulli / log / cross-entropy loss
 - Loss based on scores

$$L(y, f) = \ln(1 + \exp(-y \cdot f)) \text{ for } y \in \{-1, +1\}$$

 $L(y, f) = -y \cdot f + \log(1 + \exp(f)) \text{ for } y \in \{0, 1\}$

Loss based on probabilities:

$$L(y,\pi) = \ln(1 + \exp(-y \cdot \log(\pi))) \text{ for } y \in \{-1, +1\}$$

$$L(y,\pi) = -y \log(\pi) - (1-y) \log(1-\pi) \text{ for } y \in \{0, 1\}$$



Optimization

- Minimization of the empirical risk
- For **OLS**: analytical solution $\hat{\theta} = (\mathbf{X}^{\top}\mathbf{X})^{-1}\mathbf{X}^{\top}\mathbf{y}$
- For other loss functions:
 - Log. Regr.: Convex problem, solvable via second-order optimization methods (e.g. BFGS)
 - Else: Numerical optimization

Multi-class extension of logistic regression

• Estimate **class-wise** scoring functions:

$$\Rightarrow \pi: \mathcal{X} \rightarrow [0,1]^g, \ \pi(\mathbf{x}) = (\pi_1(\mathbf{x}), \dots, \pi_g(\mathbf{x})), \ \sum_{k=1}^g \pi_k(\mathbf{x}) = 1$$

• Achieved through **softmax** transformation:

$$\pi_k(\mathbf{x} \mid \boldsymbol{\theta}) = \exp(\boldsymbol{\theta}_k^{\top} \mathbf{x}) / \sum_{j=1}^g \exp(\boldsymbol{\theta}_j^{\top} \mathbf{x})$$

- Multi-class log-loss: $L(y, \pi(\mathbf{x})) = -\sum_{k=1}^{g} \mathbb{I}_{\{y=k\}} \log(\pi_k(\mathbf{x}))$
- Predict class with maximum score (or use thresholding variant)



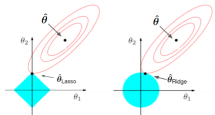
GENERALIZED LINEAR MODELS – REGULARIZATION

General idea

- Unregularized LM: risk of overfitting in high-dimensional space with only few observations
- Goal: avoidance of overfitting by adding penalty term

Regularized empirical risk

- Empirical risk function **plus complexity penalty** $J(\theta)$, controlled by shrinkage parameter $\lambda > 0$: $\mathcal{R}_{reg}(\theta) := \mathcal{R}_{emp}(\theta) + \lambda \cdot J(\theta)$
- Ridge regression: L2 penalty $J(\theta) = \|\theta\|_2^2$
- LASSO regression: L1 penalty $J(\theta) = \|\theta\|_1$





GENERALIZED LINEAR MODELS – REGULARIZATION

Optimization under regularization

- Ridge: analytically with $\hat{\theta}_{\text{Ridge}} = (\mathbf{X}^{\top}\mathbf{X} + \lambda \mathbf{I})^{-1}\mathbf{X}^{\top}\mathbf{y}$
- LASSO: numerically with, e.g., (sub-)gradient descent

Choice of regularization parameter

- Standard hyperparameter optimization problem
- ullet E.g., choose λ with minimum mean cross-validated error

Ridge vs. LASSO

Ridge

- ullet Global shrinkage \Rightarrow overall smaller but still dense heta
- Applicable with large number of influential features, correlated variables' coefficients are shrinked by equal amount

LASSO

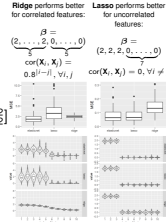
- Actual variable selection by shrinking coefficients to zero
- Suitable for sparse problems, ineffective with correlated features (randomly selecting one)



GENERALIZED LINEAR MODELS – REGULARIZATION

- Neither overall better ⇒ elastic net
- Weighted combination of Ridge and LASSO
- Introducing additional penalization coefficient:

$$\mathcal{R}_{\text{reg}}(\theta) = \mathcal{R}_{\text{emp}}(\theta) + \lambda \cdot P_{\alpha}(\theta), \text{ with } P_{\alpha}(\theta) = [\alpha \cdot \|\theta\|_{1} + (1 - \alpha) \cdot \frac{1}{2} \cdot \|\theta\|_{2}^{2}]$$





GENERALIZED LINEAR MODELS – IMPLEMENTATION

Implementation

- R:
 - Unregularized: mlr3 learner LearnerRegrLM, calling stats::lm()/mlr3 learner LearnerClassifLogReg, calling stats::glm()
 - Regularized / ElasticNet: mlr3 learners
 LearnerClassifGlmnet / LearnerRegrGlmnet, calling
 glmnet::glmnet()
 - For large classification data: mlr3 learner
 LearnerClassifLiblineaR, calling
 LiblineaR::LiblineaR() uses fast coordinate descent



GENERALIZED LINEAR MODELS -IMPLEMENTATION

- Python: From package sklearn.linear_model
 - Unregularized:
 - LinearRegression()
 - LogisticRegression(penalty = None)
 - Regularized:
 - Linear regression: Lasso(), Ridge(), ElasticNet()
 - Logistic regression: LogisticRegression(penalty = {'11', '12', 'elasticnet'})
 - Package for advanced statistical models: statsmodels.api



GENERALIZED LINEAR MODELS - PROS & CONS

Advantages

- + Simple and fast implementation
- + **Analytical** solution for L2 loss
- + Applicable for any dataset size, as long as number of observations ≫ number of features
- Flexibility beyond linearity with polynomials, trigonometric transformations, interaction terms etc.
- + Intuitive **interpretability** via feature effects
- + Statistical hypothesis **tests** for effects available

Disadvantages

- Nonlinearity of many real-world problems
- Further restrictive
 assumptions: linearly
 independent features,
 homoskedastic residuals,
 normality of conditional
 response
- Sensitivity: outliers, noise
- LM can overfit (e.g., many features and few observations)
- Feature interactions must be handcrafted
 - ightarrow practically infeasible for higher orders



REGRESSION

CLASSIFICATION

NON)PARAMETRIC

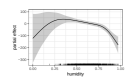
WHITE-BOX

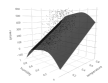
FEATURE SELECTION

General idea

- Same as GLM, but introduce flexibility through nonlinear (smooth) effects f_i(x_i)
- Typically, combination of linear & smooth effects
- Smooth effects also conceivable for feature interactions

Hypothesis space
$$\mathcal{H} = \left\{ f : \mathcal{X} \to \mathbb{R} \mid f(\mathbf{x}) = \phi \left(\theta_0 + \sum_{j=1}^p f_j(x_j) \right) \right\}$$
, suitable transformation $\phi(\cdot)$, intercept θ_0 , smooth functions $f_i(\cdot)$



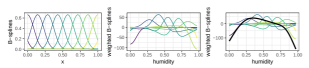


Prediction of bike rentals from smooth term of humidity (left: partial effect) and linear term of temperature (right: bivariate



Smooth functions

- Non-/semi-/parametric approaches conceivable
- Frequently: express f_j as weighted sum of basis functions ~> model linear in weight coefficients again
 - Use fixed basis of functions b₁,..., b_K and estimate associated coefficients γ₁,..., γ_K
 ∴ f_i(x_i) = ∑_{k=1}^{K_i} γ_{i,k}b_k(x_i)
 - Popular types of basis functions
 - Polynomial → smoothing/TP-/B-splines
 - Radial → Kriging
 - Trigonometric → Fourier/wavelet forms
- Alternatives: local regression (LOESS), kernel-smoothing, ...

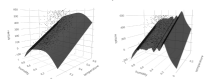


Left: B-spline basis with 9 basis functions. Middle: BFs weighted with coefficients estimated for humidity. Right: sum of weighted BFs in black (= partial effect).



Regularization

- Smooth functions possibly very flexible → regularization vital to prevent overfitting
- Control smoothness
 - Basis-function approaches: control number; impose penalty on coefficients (e.g., magnitude or differences between coefficients of neighboring components) & control associated hyperparameter
 - Local smoothers: control width of smoothing window (larger
 → smoother)



Prediction surfaces for bike rentals with 9 (left) and 500 (right) basis functions in smooth humidity term. Higher number of basis functions yields more local, less smooth model.





Loss functions Same as in GLM → essentially: use **negative** log-likelihood Optimization

- Coefficients (of smooth + linear terms): penalized MLE, Bayesian inference
- Smoothing hyperparameters: typically, generalized cross-validation

GENERALIZED ADDITIVE MODELS – IMPLEMENTATION



Implementation

- R: mlr3 learner LearnerRegrGam, calling mgcv::gam()
 - Smooth terms: s(..., bs="<basis>") or te(...) for multivariate (tensorproduct) effects
 - Link functions: family={Gamma, Binomial, ...}
- Python: GLMGam from package statsmodels; package pygam

GENERALIZED ADDITIVE MODELS – PROS & CONS

Advantages

- + Simple and fast
- + Applicable for any dataset size, as long as number of observations ≫ number of features
- + High **flexibility** via smooth effects
- Easy to combine linear & nonlinear effects
- + Rather intuitive interpretability via feature effects
- Statistical hypothesis **tests** for effects available

Disadvantages

- Sensitivity w.r.t. outliers and noisy data
- Feature interactions must be handcrafted
 - → practically infeasible for higher orders
- Harder to optimize than GLM
- Additional hyperparameters (type of smooth functions, smoothness degree, ...)



CART – METHOD SUMMARY

REGRESSION CLASSIFICATION

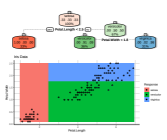
NONPARAMETRIC

WHITE-BOX

FEATURE SELECTION

General idea (CART - Classification and Regression Trees)

- Start at root node containing all data
- Perform repeated axis-parallel binary splits in feature space to obtain rectangular partitions at terminal nodes Q_1, \ldots, Q_M
- Splits based on reduction of node impurity → empirical risk minimization (**ERM**)
- In each step:
 - Find **optimal split** (feature-threshold combination)
 - \rightarrow greedy search
 - Assign constant prediction c_m to all obs. in Q_m
 - \rightarrow Regression: c_m is average of v
 - \rightarrow Classif.: c_m is majority class (or class proportions)
 - Stop when a pre-defined criterion is reached
 - → See Complexity control





CART – METHOD SUMMARY

Hypothesis space
$$\mathcal{H} = \left\{ f(\mathbf{x}) : f(\mathbf{x}) = \sum_{m=1}^{M} c_m \mathbb{I}(\mathbf{x} \in Q_m) \right\}$$

 Splitting feature x_j at split point t divides a parent node N_p into two child nodes:

$$N_l = \{(\mathbf{x}, y) \in N_p : x_i < t\} \text{ and } N_r = \{(\mathbf{x}, y) \in N_p : x_i > t\}$$

 Compute empirical risks in child nodes and minimize their sum to find best split (impurity reduction):

$$\underset{j,t}{\arg\min} \mathcal{R}(N_p, j, t) = \underset{j,t}{\arg\min} \mathcal{R}(N_l) + \mathcal{R}(N_r)$$
 (1)

Note: If \mathcal{R} is the average instead of the sum of loss functions, we need to reweight: $\frac{|N_{pt}|}{|N_{cl}|} \mathcal{R}(N_{pt})$



CART – METHOD SUMMARY

- In general, compatible with arbitrary losses typical choices:
 - *g*-way classification:

g many construction		
	Brier score → Gini impurity	Bernoulli loss $ o$ entropy impurity
	$\mathcal{R}(N_p) = \sum_{(\mathbf{x}, \mathbf{y}) \in N_p} \sum_{k=1}^g \hat{\pi_k}^{(N_p)} (1 - \hat{\pi_k}^{(N_p)})$	$\mathcal{R}(N_p) = -\sum_{(\mathbf{x}, y) \in N_p} \sum_{k=1}^g \hat{\pi_k}^{(N_p)} \log \hat{\pi_k}^{(N_p)}$
•	Regression (quadratic loss): $\mathcal{R}(N_p) =$	
	$c = \frac{1}{ N_p } \sum_{(\mathbf{x}, \mathbf{y}) \in N_p} \mathbf{y} \tag{S}$	$\mathbf{x}, \mathbf{y}) \in N_{\mathcal{D}}$



- Exhaustive search over all split candidates, choice of risk-minimal split
- In practice: reduce number of split candidates (e.g., using quantiles instead of all observed values)



CART – IMPLEMENTATION & PRACTICAL HINTS

Hyperparameters and complexity control

- Unless interrupted, splitting continues until we have pure leaf nodes (costly + overfitting)
- Hyperparameters: Complexity (i.e., number of terminal nodes) controlled via tree depth, minimum number of observations per node, maximum number of leaves, minimum risk reduction per split, ...
- Limit tree growth / complexity via
 - Early stopping: stop growth prematurely
 - \rightarrow hard to determine good stopping point before actually trying all combinations
 - Pruning: grow deep trees and cut back in risk-optimal manner afterwards



CART – IMPLEMENTATION & PRACTICAL HINTS

Implementations

- R:
 - CART: mlr3 learners LearnerClassifRpart / LearnerRegrRpart, calling rpart::rpart()
 - Conditional inference trees: partykit::ctree()
 mitigates overfitting by controlling tree size via p-value-based splitting
 - Model-based recursive partitioning: partykit::mob()
 fits a linear model within each terminal node of the decision tree
 - Rule-based models: Cubist::cubist() for regression and C50::C5.0() for classification; more flexible frameworks for fitting various types of models (e.g., GLMs) within a tree's terminal nodes
- Python: DecisionTreeClassifier / DecisionTreeRegressor from package scikit-learn



CART – PROS & CONS



Dual purpose of CART

- **Exploration purpose** to obtain interpretable decision rules (here: performance/tuning is secondary)
- Prediction model: CART as base learner in ensembles (bagging, random forest, boosting) can improve stability and performance (if tuned properly), but becomes less interpretable

CART - PROS & CONS

Advantages

- + **Easy** to understand & visualize (interpretable)
- + Built-in feature selection
 → e.g., when features are not used for splitting
- Applicable to categorical features

 → e.g., 2^m possible binary splits for m categories
 → trick for regr. with L2-loss and binary classif.: categories can be sorted ⇒ m − 1 binary splits
- + Handling of **missings** possible via surrogate splits
- Models interactions,
- + Fast well scalable
- + High **flexibility** with custom split criteria or leaf-node prediction rules

Disadvantages

- Rather poor generalization
- High variance/instability: model can change a lot when training data is minimally changed
- Can overfit if tree is grown too deep
- Not well-suited to model linear relationships
- Bias toward features with many unique values or categories



RANDOM FORESTS – METHOD SUMMARY

REGRESSION

CLASSIFICATION

NONPARAMETRIC

BLACK-BO

FEATURE SELECTION

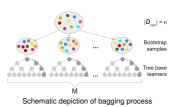
General idea

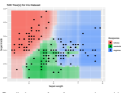
- Bagging ensemble of M tree base learners fitted on bootstrap data samples
 - \Rightarrow Reduce **variance** by ensembling while slightly increasing **bias** by bootstrapping
 - Use unstable, high-variance base learners by letting trees grow to full size
 - Promoting decorrelation by random subset of candidate features for each split
- Predict via averaging (regression) or majority vote (classification) of base learners

Hypothesis space
$$\mathcal{H} = \left\{ f(\mathbf{x}) : f(\mathbf{x}) = \frac{1}{M} \sum_{m=1}^{M} \sum_{t=1}^{T^{[m]}} c_t^{[m]} \mathbb{I}(\mathbf{x} \in Q_t^{[m]}) \right\}$$



RANDOM FORESTS – METHOD SUMMARY





Prediction surface for iris data with 500-tree ensemble base learners

Empirical risk & Optimization Just like tree base learners
Out-of-bag (OOB) error

- Ensemble prediction for obs. outside individual trees' bootstrap training sample ⇒ unseen test sample
- Use resulting loss as unbiased estimate of generalization error
- Mainly useful for tuning and less for model comparison as we usually compare all models uniformly by CV



RANDOM FORESTS – METHOD SUMMARY

Feature importance

- Based on improvement in split criterion: aggregate improvements by all splits using j-th feature
- Based on **permutation:** permute *j*-th feature in OOB observations and compute impact on OOB error

Hyperparameters

- Ensemble size, i.e., number of trees
- Complexity of base learners, e.g., tree depth, min-split, min-leaf-size
- Number of split candidates, i.e., number of features to be considered at each split
 - \Rightarrow frequently used heuristics with total of p features: $\lfloor \sqrt{p} \rfloor$ for classification, $\lfloor p/3 \rfloor$ for regression



RANDOM FORESTS – IMPLEMENTATION & PRACTICAL HINTS

Extremely Randomized Trees

- Variance of trees can be further increased by randomizing split points instead of using the optimal one
- Alternatively consider k random splits and pick the best one according to impurity

Tuning

- While default values for number of split points is often good, tuning it can still improve performance
- Tuning the minimum samples in leafs and minimum samples for splitting can be benificial but no huge performance increases are to be expected



RANDOM FORESTS – IMPLEMENTATION & PRACTICAL HINTS



Implementation

- R: mlr3 learners LearnerClassifRanger /
 LearnerRegrRanger, calling ranger::ranger() as a highly efficient and flexible implementation
- Python: RandomForestClassifier / RandomForestRegressor from package scikit-learn

RANDOM FORESTS – PROS & CONS

Advantages

- Retains most of trees' advantages (e.g., feature selection, feature interactions)
- Fairly good predictor: mitigating base learners' variance through bagging
- + Quite **robust** w.r.t. small changes in data
- Good with high-dimensional data, even in presence of noisy features
- + Easy to parallelize
- Robust to its hyperparameter configuration
- Intuitive measures of feature importance

Disadvantages

- Loss of individual trees' interpretability
- Can be suboptimal for regression when extrapolation is needed
- Bias toward selecting features with many categories (same as CART)
- Rather large model size and slow inference time for large ensembles
- Typically inferior in performance to tuned gradient tree boosting.



GRADIENT BOOSTING – METHOD SUMMARY

REGRESSION

CLASSIFICATION

(NON)PARAMETRIC

BLACK-BOX

FEATURE SELECTION

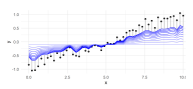
General idea

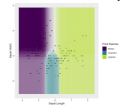
- Sequential ensemble of M base learners by greedy forward stagewise additive modeling
 - In each iteration a base learner is fitted to current pseudo residuals ⇒ one boosting iteration is one approximate gradient step in function space
 - Base learners are typically trees, linear regressions or splines
- Predict via (weighted) sum of base learners



GRADIENT BOOSTING – METHOD SUMMARY

Hypothesis space
$$\mathcal{H} = \left\{ f(\mathbf{x}) : f(\mathbf{x}) = \sum_{m=1}^{M} \beta m b(\mathbf{x}, \theta m) \right\}$$





Boosting prediction function with GAM base learners for univariate regression problem after 10 iterations

Boosting prediction surface with tree base learners for iris data after 100 iterations (right: contour lines of discriminant functions)

Empirical risk

- In general, compatible with any differentiable loss
- Base learner in iteration *m* is fitted on **Pseudo residuals**:

Base learner in iteration
$$m$$
 is fitted on **Pseudo residuals**:
$$\tilde{r}^{(i)} = -\frac{\partial L(y^{(i)}, f(\mathbf{x}^{(i)}))}{\partial f(\mathbf{x}^{(i)})} \text{ by minimizing the } \mathbf{L2\text{-loss}} : \sum_{i=1}^{n} (\tilde{r}^{(i)} - b(\mathbf{x}^{(i)}, \boldsymbol{\theta}))^2$$



GRADIENT BOOSTING – METHOD SUMMARY

Optimization

- Same optimization procedure as base learner, while keeping the current ensemble fmdh fixed
 - ⇒ Efficient and generally applicable since *inner* loss is always L2
- βm is found via **line search** or fixed to a **small constant value** and combined with the leaf values ctm for tree base learners: $ctmt = \beta m \cdot ctm$

Hyperparameters

- Ensemble size, i.e., number of base learners
- Complexity of base learners (depending on type used)
- Learning rate β , i.e., impact of next base learner



GRADIENT BOOSTING – PRACTICAL HINTS

Scalable Gradient Boosting

- Feature and data subsampling for each base learner fit
- Parallelization and approximate split finding for tree base learners
- GPU accelaration

Explainable / Componentwise Gradient Boosting

- Base learners of simple linear regression models or splines, selecting a single feature in each iteration
- Allows feature selection and creates an interpretable model since uni- and bivariate effects can be visualized directly.
- Feature interactions can be learned via ranking techniques (e.g., GA²M FAST)

Tuning

- Use early-stopping to determine ensemble size
- Various regularization parameters, e.g., L1/L2, number of leaves, ... that need to be carefully tuned
- Tune learning rate and base learner complexity hyperparameters on log-scale



GRADIENT BOOSTING – IMPLEMENTATION

Gradient Tree Boosting

- R:mlr3 learners LearnerClassifXgboost / LearnerRegrXgboost, LearnerClassifLightGBM / LearnerRegrLightGBM
- Python: GradientBoostingClassifier / GradientBoostingRegressor from package scikit-learn, XGBClassifier / XGBRegressor from package xgboost, lgb.train from package lightgbm
- $\Rightarrow {\tt LightGBM}$ current state-of-the-art but slightly more complicated to use than ${\tt xgboost}$

Componentwise Gradient Boosting

- R: mboost from package mboost, boostLinear / boostSplines from package compboost
- Python: /
- \Rightarrow mboost very flexible but slow while compboost is much faster with limited features



GRADIENT BOOSTING - PROS & CONS

Advantages

- Retains of most of base learners' advantages
- Very good predictor due to aggressive loss minimization, typically only outperformed by heterogenous stacking ensembles
- High flexibility via custom loss functions and choice of base learner
- Highly efficient implementations exist (lightgbm / xgboost) that work well on large (distributed) data sets
- Componentwise boosting: Good combination of (a) high performance (b) interpretable model and (c) feature selection

Disadvantages

- Loss of base learners' potential interpretability
- Many hyperparameters to be carefully tuned
- Hard to parallelize (→ solved by efficient implementation)



LINEAR SVM - METHOD SUMMARY

CLASSIFICATION

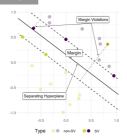
EGRESSION

PARAMETRIC

WHITE-BOX

highlightGeneral idea (Soft-margin SVM)

- Find linear decision boundary (separating hyperplane) that
 - maximizes distance (margin γ) to closest points (support vectors, SVs) on each side of decision boundary
 - while minimizing margin violations (points either on wrong side of hyperplane or between dashed margin line and hyperplane)





- 3 types of training points
 - non-SVs with no impact on decision boundary
 - SVs that are margin violators and affect decision boundary
 - SVs located exactly on dashed margin lines and affect decision boundary



LINEAR SVM – METHOD SUMMARY

Hypothesis space (primal) $\mathcal{H} = \{f(\mathbf{x}) : f(\mathbf{x}) = \boldsymbol{\theta}^{\top} \mathbf{x} + \theta_0\}$ Empirical risk Soft-margin

SVM as **L2-regularized ERM**:

$$\frac{1}{2} \|\boldsymbol{\theta}\|_{2}^{2} + C \sum_{i=1}^{n} L(y^{(i)}, f(\mathbf{x}^{(i)}))$$

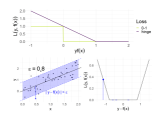
- $\|\theta\| = 1/\gamma$ ($\hat{=}$ maximizing margin)
- C > 0: penalization for margin violations
- Loss aims at minimizing margin violations

$$\rightarrow$$
 Classif. (**hinge** loss):

$$L(y, f) = \max(1 - yf, 0)$$

 \rightarrow Regr. (ϵ -insensitive loss):

$$L(y, f) = \max(|y - f| - \epsilon, 0)$$





LINEAR SVM - METHOD SUMMARY

Dual problem SVMs as a constraint optimization (primal) problem (maximize margin s.t. constraints on obs. to limit margin violations) can be formulated as a Lagrangian dual problem with Lagrange multipliers $\alpha_i \geq 0$:

$$\max_{\alpha v \in \mathbb{R}^n} \Leftrightarrow \text{ s.t. } 0 \le \alpha_i \le C \ \forall i \in \{1, \dots, n\} \text{ and } \sum_{i=1}^n \alpha_i y^{(i)} = 0$$

Solution Non-SVs have $\alpha_i = 0$ as they do not affect the hyperplane

$$f(\mathbf{x}) = \sum_{i=1}^{n} \alpha_i y^{(i)} \langle \mathbf{x}^{(i)}, \mathbf{x} \rangle + \theta_0$$

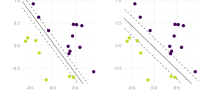
Optimization

- Typically, tackling dual problem (though feasible in corresponding primal) via quadratic programming
- Popular: sequential minimal optimization ⇒ iterative algorithm based on breaking down objective into bivariate quadratic problems with analytical solutions

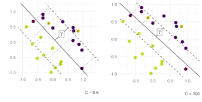


LINEAR SVM - METHOD SUMMARY

Hyperparameters Cost parameter *C* to control maximization of the margin vs. minimizing margin violations



Hard-margin SVM: margin is maximized by boundary on the right

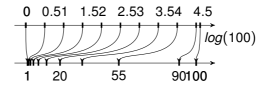


Soft-margin SVM: large margin and few margin violations on the right (best trade-off)



Preprocessing Features should be scaled before applying SVMs (applies generally to regularized models)
Tuning

- Tuning of cost parameter C advisable
 ⇒ strong influence on resulting hyperplane
- C it is often tuned on a log-scale grid for optimal and space-filling search space





LINEAR SVM - IMPLEMENTATION

Implementation

- R: mlr3 learners LearnerClassifSVM / LearnerRegrSVM, calling e1071::svm() with linear kernel (libSVM interface). Further implementations in mlr3extralearners based on
 - kernlab::ksvm() allowing custom kernels
 - LiblineaR::LiblineaR() for a fast implementation with linear kernel
- Python: sklearn.svm.SVC from package scikit-learn/ package libSVM



NONLINEAR SVM – METHOD SUMMARY

CLASSIFICATION

REGRESSION

NONPARAMETRIC

BLACK-BOX

General idea

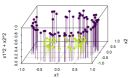
- Move beyond linearity by mapping data to transformed space where they are linearly separable
- Kernel trick
 - No need for explicit construction of feature maps
 - Replace inner product of feature map $\phi: \mathcal{X} \to \Phi$ by **kernel**: $\langle \phi \mathbf{x}, \phi \mathbf{x} \mathbf{t} \rangle = \mathbf{k} \mathbf{x} \mathbf{x} \mathbf{t}$

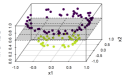
Hypothesis space

$$\mathcal{H} = \{ f(\mathbf{x}) : f(\mathbf{x}) = \text{sign} (\boldsymbol{\theta}^{\top} \phi x + \theta_0) \} \text{ (primal)}$$

$$\mathcal{H} = \left\{ f(\mathbf{x}) : f(\mathbf{x}) = \operatorname{sign}\left(\sum_{i=1}^{n} \alpha_i y^{(i)} k(\mathbf{x}^{(i)}, \mathbf{x}) + \theta_0\right) \mid \alpha_i \ge 0, \sum_{i=1}^{n} \alpha_i y^{(i)} = 0 \right\} \text{ (dual)}$$







Nonlinear problem in original

Mapping to 3D space and subsequent linear separation - implicitly handled by kernel in nonlinear

Kernelize dual (soft-margin) SVM problem, replacing all inner



NONLINEAR SVM – IMPLEMENTATION & PRACTICAL HINTS

Common kernels

- Linear kernel: dot product of given observations $\Rightarrow kxxt = \mathbf{x}^{\top}\tilde{\mathbf{x}} \Rightarrow \text{linear SVM}$
- Polynomial kernel of degree $d \in \mathbb{N}$: monomials (i.e., feature interactions) up to d-th order $\Rightarrow kxxt = (\mathbf{x}^{\top}\tilde{\mathbf{x}} + b)^d$, $b \ge 0$
- Radial basis function (RBF) kernel: infinite-dimensional feature space, allowing for perfect separation of all finite datasets $\Rightarrow kxxt = \exp\left(-\gamma \|\mathbf{x} \tilde{\mathbf{x}}\|_2^2\right)$ with bandwidth parameter $\gamma > 0$

Tuning

- ◆ High sensitivity w.r.t. hyperparameters, especially those of kernel ⇒ tuning very important
- For RBF kernels, use RBF sigma heuristic to determine bandwidth

Implementation

- R: mlr3 learners LearnerClassifSVM / LearnerRegrSVM, calling e1071::svm() with nonlinear kernel (libSVM interface), kernlab::ksvm() allowing custom kernels
- Python: sklearn.svm.SVC from package scikit-learn / package libSVM



SVM - PRO'S & CON'S

Advantages

- Often **sparse** solution (w.r.t. observations)
- Robust against overfitting (regularized); especially in high-dimensional space
- Stable solutions (w.r.t. changes in train data) → Non-SV do not affect decision boundary
- Convex optimization problem
- → local minimum $\hat{=}$ global minimum Advantages (nonlinear SVM)

Can learn nonlinear decision Disadvantages (nonlinear SVM) boundaries

- Very flexible due to custom kernels
 - → RBF kernel yields local

model

Disadvantages

- **Long** training times $\rightarrow O(n^2p + n^3)$
- Confined to linear model
- Restricted to continuous features
- Optimization can also fail or get stuck



- Poor **interpretability** due to complex kernel
- Not easy tunable as it is highly important to choose the

right kernel (which also

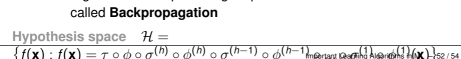
NEURAL NETWORKS – METHOD SUMMARY

CLASSIFICATION

BLACK-BOX

General idea

- Learn **composite function** through series of nonlinear feature transformations, represented as **neurons**, organized hierarchically in layers
 - Basic neuron operation: 1) affine **transformation** ϕ (weighted sum of inputs), 2) nonlinear activation σ
 - Combinations of simple building blocks to create a complex model
- Optimize via mini-batch stochastic gradient descent (SGD) variants:
 - Gradient of each weight can be infered from the computational graph of the network
 - → **Automatic Differentiation** (AutoDiff)
 - Algorithm to compute weight updates based on the loss is





NEURAL NETWORKS – IMPLEMENTATION & PRACTICAL HINTS

General hints

- Instead of NAS, use a standard architecture and tune training hyperparameters
- Training pipeline (data-augmentation, training schedules, ...) is more crucial than the specific architecture
- While NNets are state-of-the-art for computer vision (CV) and natural language processing (NLP), we recommend not to use them for tabular data because alternatives perform better
- Computational efforts for training (and inference) can be very high, requiring specific hardware.
 - \rightarrow Using a service (esp. for foundation models) can be more cost efficient

Implementation

- R: Use python libraries (below) via reticulate, but not really recommended except for toy applications.
- Python libraries:
 - keras for simple high level API



NEURAL NETWORKS - PROS & CONS

Advantages

- Applicable to complex, nonlinear problems
- Very versatile w.r.t. architectures
- State-of-the-art for CV and NLP
- + Strong **performance** if done right
- Built-in feature extraction, obtained by intermediate representations
- + Easy handling of **high-dimensional** data
- + Parallelizable training

Disadvantages

- Typically, high computational cost
- High demand for training data
- Strong tendency to overfit
- Requiring lots of tuning expertise
- Black-box model hard to interpret or explain

