Text classification

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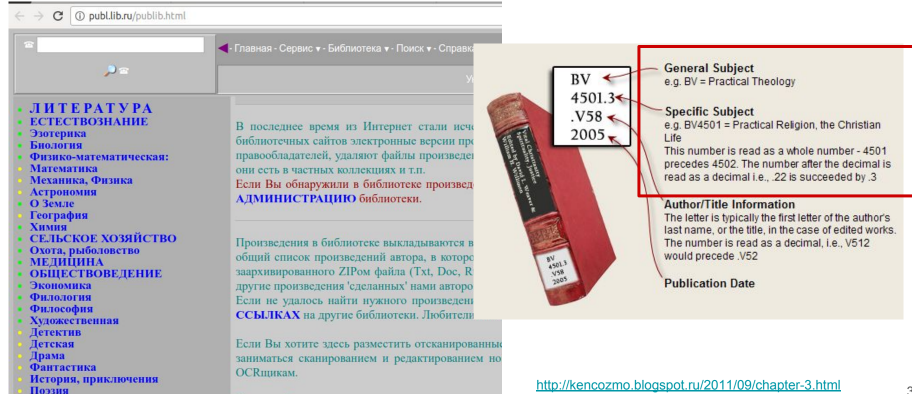
Plan

- Motivation
- 2. Classification task
- 3. Classifier example: Naive Bayes
- 4. Classification quality evaluation
- Classification methods review
 - a. Linear methods
 - b. Metric methods
 - c. Logical methods
 - d. Ensembles
- 6. Typical tasks and special cases

Motivation: roots bloody roots

Политика

Путешествия, природа



Прошу обратить внимание на то, что администрац

приславших. Кроме того, разрешается любое неког

http://kencozmo.blogspot.ru/2011/09/chapter-3.html

Motivation

- Sentiment analysis: track/check if the users are happy with the product or not (optional: + find out which particular feature user [dis]liked)
- 2. **Topic classification**: section the news article to be put in
- 3. **Spam detection**: predict if letters are unwanted by user based on those tagged by him/her as spam
- 4. **Incomplete data imputation**: predict user's gender based on text he/she publishes/likes/skips
- 5. **Many more**: authorship attribution, sociodemographic characteristics, etc...

Google Product Search



HP Officejet 6500A Plus e-All-in-One Color Ink-jet - Fax / copier / printer / s \$89 online, \$100 nearby ★★★★☆ 377 reviews

September 2010 - Printer - HP - Inkjet - Office - Copier - Color - Scanner - Fax



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Classification task

Supervised learning task. Given:

- a set of documents (texts) D = { d₁, d₂, ... d_n}
- a set of classes (categories) C = { c₁, c₂, ... c_k }
- usually there is a training set -- a subset of D x C, that is, document-class pairs

Task:

train a function f: D => C, matching each document with the correct class

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Classifier example: Naive Bayes

aka simple Bayes, independent Bayes is called so thanks to being a straightforward Bayes theorem application

Approach: learn **P(c|d)** and choose **c** with the largest conditional probability value for every **d**

$$p(C_k \mid x_1, \dots, \underbrace{x_n})$$
 class words in a document

Assumption: all words are conditionally independent

$$p(x_i \mid x_{i+1}, \ldots, x_n, C_k) = p(x_i \mid C_k)$$



$$P(A \mid B) = \frac{P(B \mid A)P(A)}{P(B)}$$

Naive Bayes: formulae

Class probability for the set of given words in a document

$$p(C_k \mid \mathbf{x}) = rac{p(C_k) \ p(\mathbf{x} \mid C_k)}{p(\mathbf{x})}$$

p(x) — constant!

...we care about the numerator only

$$p(C_k,x_1,\ldots,x_n)$$

Let's rewrite it using the chain rule

$$egin{aligned} p(C_k,x_1,\ldots,x_n) &= p(x_1,\ldots,x_n,C_k) \ &= p(x_1\mid x_2,\ldots,x_n,C_k) p(x_2,\ldots,x_n,C_k) \ &= p(x_1\mid x_2,\ldots,x_n,C_k) p(x_2\mid x_3,\ldots,x_n,C_k) p(x_3,\ldots,x_n,C_k) \ &= \ldots \ &= p(x_1\mid x_2,\ldots,x_n,C_k) p(x_2\mid x_3,\ldots,x_n,C_k) \ldots p(x_{n-1}\mid x_n,C_k) p(x_n\mid C_k) p(C_k) \end{aligned}$$

Naive Bayes: formulae

Using the conditional independence assumption, we get

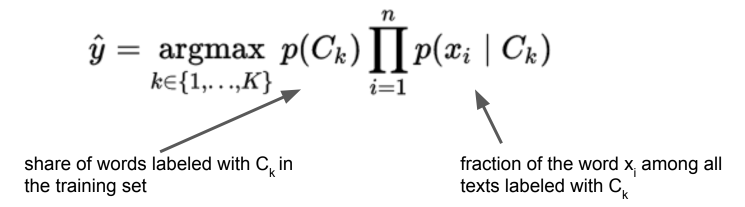
$$egin{split} p(C_k \mid x_1,\ldots,x_n) &\propto p(C_k,x_1,\ldots,x_n) \ &\propto p(C_k) \ p(x_1 \mid C_k) \ p(x_2 \mid C_k) \ p(x_3 \mid C_k) \ \cdots \ &\propto p(C_k) \prod_{i=1}^n p(x_i \mid C_k) \,. \end{split}$$

classifier is ready:

$$\hat{y} = rgmax_{k \in \{1,\ldots,K\}} p(C_k) \prod_{i=1}^n p(x_i \mid C_k).$$

the so-called MAP (maximum a posteriori) decision rule

Naive Bayes: how to compute this



- 1. Estimate probabilities using the formula above
- 2. Compute the value for every class for every new incoming document
- 3. Choose the class with the largest value

Yes, doing **smoothing** does make sense here; one can also consider taking

- a share of documents containing the word (binary Naive Bayes) instead of plain word frequencies
- log-frequencies instead of plain word frequencies

Naive Bayes: discussion

- independence assumption
 (natural language is not a bag of words)
- weights of long documents differ a great deal
- prone to the systematic error (bias) towards certain decisions
- robust to unknown words
- + simple and fast
- + is often used as a simple baseline

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 - d. Ensembles
- 6. Typical tasks and special cases

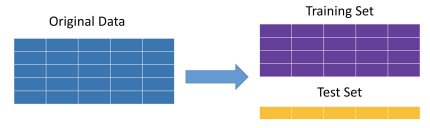
REMINDER! machine learning models quality evaluation

We have the data, we have the metric

Splitting into

- train set
- test set

Beleving these subsets are 'sampled from the same distribution' (otherwise training makes almost no sense)



REMINDER! machine learning models quality evaluation

Deadly Sin №1

Test data leaks into train set (this way we lose generalization capability and estimates validity)

Deadly Sin №2

Tuning hyperparameters on test set



But how do we tune the parameters? Ideas?

REMINDER! machine learning models quality evaluation

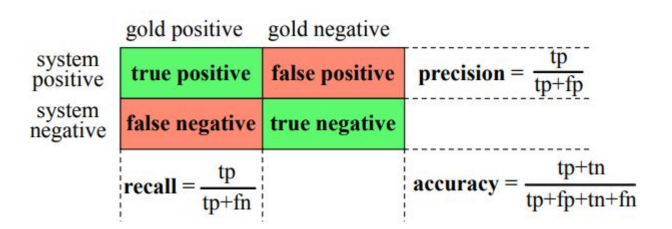
TRAIN DEV TEST

- 1. TRAIN training model
- 2. DEV evaluating quality + analyzing errors + tuning hyperparameters
- 3. TEST blind quality evaluation: looking at quality metric ONLY + not too often, so as not to overfit

Example: spam (positive) or not spam (negative) emails

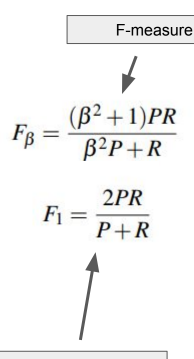
true labels

		spam!	not spam!			
predictions	spam!	TRUE POSITIVE we're happy	FALSE POSITIVE normal letter falling into a Spam folder a tragedy			
	not spam!	FALSE NEGATIVE spam in inbox not good	TRUE NEGATIVE we're happy			



Choose the target class and consider its prediction a **positive** case;

Correct prediction — **true positive**, incorrect — **false positive** + the same for the other class



F1-measure harmonic mean of recall and precision

Let's say 1 is a target class

ground_	_truth
prediction	on

1	1	0	0	0	1	0	1
1	0	0	1	0	1	1	1

Accuracy =
$$(3 + 2) / (3 + 2 + 2 + 1) = 0.625$$

Precision = $3 / (3 + 2) = 0.6$
Recall = $3 / (3 + 1) = 0.75$
F1 = $2 * 0.6 * 0.75 / (0.6 + 0.75) = 0.66(6)$

accuracy = share of correct hits, is in [0, 1]

 won't tell us much if samples counts of different classes shares are imbalanced

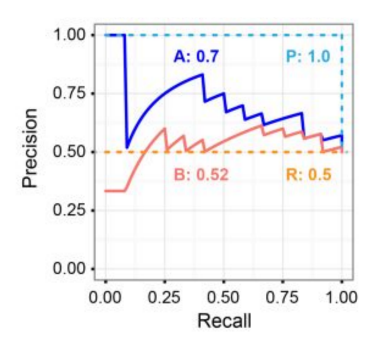
precision = a share of truly positive among predicted as positive ones
recall = a share of truly positive that were actually predicted as positive ones

Checking your understanding: if the classifier sets all labels as the target class (all samples are predicted as positive ones), what are precision and recall?

Precision-Recall Curve

we change the parameter that changes precision and recall and look at the behaviour of precision and recall values

(this parameter is usually a probability threshold in a decision rule)



https://classeval.wordpress.com/introduction/introduction-to-the-precision-recall-plot/

Classification quality evaluation: multi-class

= number of classes > 2

- Accuracy
 share of correctly predicted cases
- Micro-averaging: Precision, Recall, FScore first we compute TP, FP, ..., for every class and then we compute metrics values, summing all TPs, FPs, etc.
- 3. **Macro-averaging aka "all classes are equally important"**: Precision, Recall, FScore computing Precision, Recall,... for every class, then averaging (summing and dividing by the number of classes)

Classification quality evaluation: multi-class

ground_truth prediction

1	2	0	2	0	1	0	1
0	2	0	1	2	1	1	2

Label 0

TP = 1, FP = 1 FN = 2, TN = 4

Precision = 0.5

Recall = 0.33

Label 1

$$TP = 1, FP = 2$$

 $FN = 2, TN = 3$

Precision = 0.33

Recall = 0.33

Label 2

$$TP = 1, FP = 2$$

FN = 1, TN = 4

Precision = 0.33

Recall = 0.5

Macro-averaging

Pr = (0.5+0.33+0.33) / 3 = 0.387

R = (0.5+0.33+0.33) / 3 = 0.387

F1 = 2PrR / (Pr + R) = 0.387

Micro-averaging

Pr = (1+1+1) / (1+1+1 + 1+2+2) = 0.375

R = (1+1+1) / (1+1+1+1+2+2) = 0.375

F1 = 2PrR / (Pr + R) = 0.375

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Representing texts

Already discussed: ngrams, count/one-hot/tf-idf/normalized tf-idf Custom features may also help: POS counts, text length, weighted average word embeddings, RNN-based embeddings, etc.

Замечание о способах представления текстов

Способ #1, Bag-of-words: one hot (BOW; мешок слов)

~ one-hot-encoding / dummy coding; много интерпретируемых фич

"А не три, а не пять! Это надо знать!"

Bag-of-words: word counts (sklearn: CountVectorizer) вместо единиц — частоты / относительные частоты

8	не	три	шесть	пять	это	надо	семь	знать
2	2	1	0	1	1	1	0	1

Bag-of-words: weird numbers (sklearn: TfldfVectorizer) вместо единиц — TF-IDF или другие оценки "значимости"

BOW: специфика



много разреженных фич, можем столкнуться с проблемами больших размерностей, поэтому:

- надо уметь фильтровать и наказывать термы весами: частотные, редкие и т. д.- это есть из коробки есть в sklearn; кроме того — фильтрация по словарям (в т. ч. stopwords: словарь "вредных слов")
 - выбираем модели для работы с большим числом разреженных признаков, не всё можно загонять в Random Forest!
- обязательно экспериментируем с числом N в ngram-мах и вариациями one-hot/count/tf-idf/...

Замечание о способах представления текстов

Очевидно, с порядком слов в тексте мы теряем много информации, но есть простое средство для бедных!

Bag-of-ngrams (sklearn vectorizers поддерживают)

вместо отдельных термов наборы из n подряд идущих в тексте термов

"Нью Йорк"

"Нью Дели"

"не надо"

"catch up with"

Когда BoW плохо справляется?

- Мало обучающих данных
 - Закон Ципфа
 - Богатая морфология =>
 - слишком мало прецедентов для обучения
 - ...А если нормализуем => иногда теряем важную информацию
- Короткие тексты
 - Те же причины
 - + интуитивно: в большем тексте больше хороших слов-предикторов целевой переменной (or whatever)



Мешок пла мусора // Викиледиа

Conceptual stuff

In Naive Bayes we were training a **data model** that would allow us to **generate samples** given the class

$$\hat{y} = rgmax_{k \in \{1,\ldots,K\}} p(C_k) \prod_{i=1}^n p(x_i \mid C_k).$$

So we were **modeling the data**.

However, there is a family of models that are trained to predict this (exactly what we want classifier to do):

$$p(C_k \mid \mathbf{x})$$

They are focused on determining which features are the best to **separate the classes**

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- 6. Typical tasks and special cases

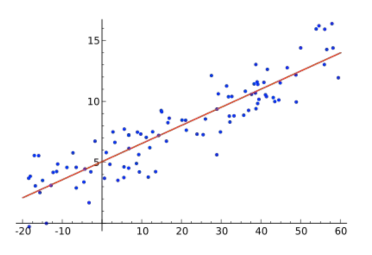
Linear models

Usually they look like this

$$?=\sum_{i=1}^{N} w_i f_i$$

where **f** are features (e.g., bag of ngrams), and **w** are the weights we are to find

Training the linear regression so that it would return the conditional **probability** of the class given the data is not really possible:)



Logistic regression

Let's try to fix it by making the outputs

- nonnegative

$$p(c|x) = \frac{1}{Z} \exp \left(\sum_{i} w_{i} f_{i}(c,x) \right)$$

- lying between 0 and 1

$$p(c|x) = \frac{\exp\left(\sum_{i=1}^{N} w_i f_i(c, x)\right)}{\sum_{c' \in C} \exp\left(\sum_{i=1}^{N} w_i f_i(c', x)\right)}$$

Logistic regression: predictions

Can be solved in a way similar to a linear regression

$$\hat{c} = \underset{c \in C}{\operatorname{argmax}} P(c|x)$$

$$= \underset{c \in C}{\operatorname{argmax}} \frac{\exp\left(\sum_{i=1}^{N} w_i f_i(c, x)\right)}{\sum_{c' \in C} \exp\left(\sum_{i=1}^{N} w_i f_i(c', x)\right)}$$

$$= \underset{c \in C}{\operatorname{argmax}} \exp \sum_{i=1}^{N} w_i f_i(c, x)$$

$$= \underset{c \in C}{\operatorname{argmax}} \sum_{i=1}^{N} w_i f_i(c, x)$$

Logistic regression: training

Maximizing conditional probability of the class given the data

$$\hat{w} = \underset{w}{\operatorname{argmax}} \log P(y^{(j)}|x^{(j)})$$

$$\hat{w} = \underset{w}{\operatorname{argmax}} \sum_{j} \log P(y^{(j)}|x^{(j)})$$

$$L(w) = \sum_{j} \log P(y^{(j)}|x^{(j)}) = \sum_{j} \log \exp \left(\sum_{i=1}^{N} w_{i} f_{i}(y^{(j)}, x^{(j)}) \right) - \sum_{j} \log \sum_{y' \in Y} \exp \left(\sum_{i=1}^{N} w_{i} f_{i}(y'^{(j)}, x^{(j)}) \right)$$

For gradient ascend we need a derivative

$$L'(w) = \sum_{j} f_k(y^{(j)}, x^{(j)}) - \sum_{j} \sum_{y' \in Y} P(y'|x^{(j)}) f_k(y'^{(j)}, x^{(j)})$$
 data-based feature counter predicted feature values

Important problem: overfitting

Machine learning models can fit the training set 'too well': features values that occur only with one class label are a strong signal for the classifier (even if the number of such cases is not large)!

E.g. logistic regression can assign a large weight to a particular feature $\mathbf{w}_{\mathbf{i}}$

However, such cases may be too specific and this may not be a good rule when using the model in the wild!

"Modell fitting too specific cases" usually fail to generalize. This is called **overfitting**.

Logistic regression: regularization

One way to fight overfitting is regularization: adding extra constraints to the task or restricting the possible solutions family

$$\hat{w} = \underset{w}{\operatorname{argmax}} \sum_{j} \log P(y^{(j)}|x^{(j)}) - \alpha R(w)$$

L2-regularizationaka **shrinkage**aka **Tikhonov's regularization**

$$R(W) = ||W||_2^2 = \sum_{j=1}^N w_j^2$$

...doesn't allow the weights to grow

Logistic regression: regularization

One way to fight overfitting is regularization: adding extra constraints to the task or restricting the possible solutions family

$$\hat{w} = \underset{w}{\operatorname{argmax}} \sum_{j} \log P(y^{(j)}|x^{(j)}) - \alpha R(w)$$

L1-regularization

aka **LASSO** (least absolute shrinkage and selection operator)

$$\hat{w} = \underset{w}{\operatorname{argmax}} \sum_{j} \log P(y^{(j)} | x^{(j)}) - \alpha \sum_{i=1}^{N} |w_{i}|$$

...doesn't just make the weights smaller but also allows to turn them into zero

Logistic regression: discussion

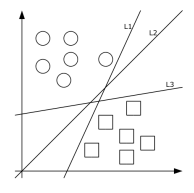
- + training and prediction is fast
- more robust then naive Bayes and works better with correlated features
- has to be done: tuning regularization, feature normalization, feature selection
- probabilities estimates may not reflect the data, see

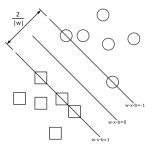
Linear models: SVM (Support Vector Machine)

Linear models usually build a separating hyperplane: different classes should be at different sides of it

Now let's try to build a hyperplace so that the objects with different labels are at max. distance from it

This should help to generalize and be more confident when predicting classes





Linear models: SVM

Points of classes **c** from the set { -1, 1 }:

$$\{(\mathbf{x}_1, c_1), (\mathbf{x}_2, c_2), \dots, (\mathbf{x}_n, c_n)\}$$

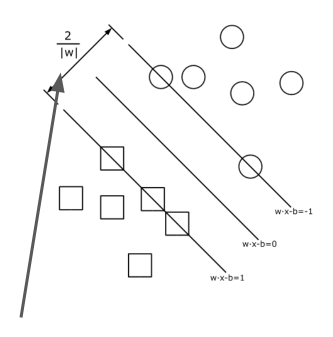
Separating hyperplane:

$$\mathbf{w} \cdot \mathbf{x} - b = 0.$$

two **parallel hyperplanes** that we can move without touching the samples in the case of linear separability:

$$\mathbf{w} \cdot \mathbf{x} - b = 1$$
, $\mathbf{w} \cdot \mathbf{x} - b = -1$.

So we minimize **|w|**, so that the distance between them was greater



Linear models: SVM

Quadratic programming task

$$egin{cases} \|\mathbf{w}\|^2 o \min \ c_i(\mathbf{w} \cdot \mathbf{x_i} - b) \geq 1, \quad 1 \leq i \leq n. \end{cases}$$

with a few transformations we can reformulate the task like this:

$$egin{cases} -\mathbf{L}(\lambda) = -\sum_{i=1}^n \lambda_{\mathbf{i}} + rac{1}{2} \sum_{i=1}^n \sum_{j=1}^n \lambda_{\mathbf{i}} \lambda_{\mathbf{j}} c_i c_j (\mathbf{x_i} \cdot \mathbf{x_j})
ightarrow \min_{\lambda} \ \lambda_{\mathbf{i}} \geq 0, \quad 1 \leq i \leq n \ \sum_{i=1}^n \lambda_{\mathbf{i}} c_i = 0 \end{cases}$$

this quadratic programming task has just one solution, which can be effectively found in the case if hundreds of thousands objects

Linear models: SVM

- there is an modification for multiple linearly inseparable classes
- take a look at the formulae at the previous slide: the features are used only in the scalar product

hence we can redefine it; this way we'll move objects into the space of higher dimensionality where they may be linearly separable

this is called the kernel trick

$$egin{aligned} k(\mathbf{x},\mathbf{x}') &= (\mathbf{x}\cdot\mathbf{x}')^d \ k(\mathbf{x},\mathbf{x}') &= (\mathbf{x}\cdot\mathbf{x}'+1)^d \ k(\mathbf{x},\mathbf{x}') &= \exp(-\gamma\|\mathbf{x}-\mathbf{x}'\|^2) \ k(\mathbf{x},\mathbf{x}') &= \exp\left(-\frac{\|\mathbf{x}-\mathbf{x}'\|^2}{2\sigma^2}\right) \ k(\mathbf{x},\mathbf{x}') &= anh(\kappa\mathbf{x}\cdot\mathbf{x}'+c) \end{aligned}$$

SVM, discussion

- + separating hyperplanes with margin usually deliver a more 'confident' solution
- + the optimization task has effective solution methods
- not robust to outliers (those that are close to the separation hyperplane)
- choosing the kernel is black magic; common sense doesn't always work
- when there is no prior belief in linear separability of the classes, one has to tune parameters

Plan

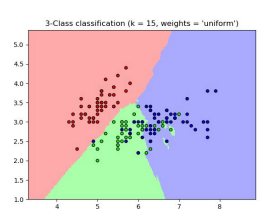
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 - a. Linear methods
 - b. Metric methods
 - c. Logical methods
 - d. Ensembles
- 6. Typical tasks and special cases

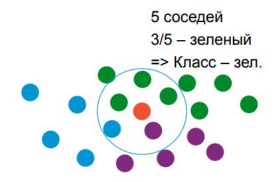
Metric classifiers: kNN

At the core -- **compactness hypothesis**: objects that are close to each other in a metric space should have the same label

k Nearest Neighbours method: no training, a classified object is given the most popular label among **k** closest objects in the train set

The larger the **k**, the more smooth are the borders between classes; however, if the k is too large, **underfitting** is possible





kNN: how to improve

One can

- use the order of the neighbours (when sorted by distance)
 as a 'vote weight' (the closer, the more important is the label vote)
- use neighbours distances to the classified object
 (vote weight is set by function, take a look at the Parzen window method)
- filter a set of representative objects in the training set
 (predictions are made faster + removing outliers helps)

kNN: discussion

- + non-linear, classes samples groups can be of arbitrary form and shape
- + a natural way to do the multiclass classification
- may be too expensive to store and use for predictions all/representative training set objects
- depends on the training set too much
- usually unsuitable for large dimensions

Plan

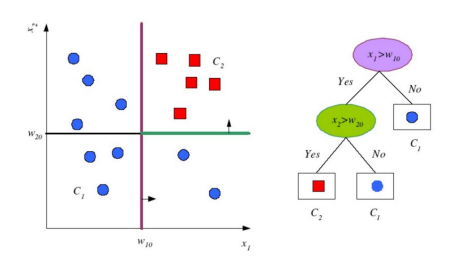
- 1. Motivation
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Logical classifiers: decision trees

We build a structure setting the conditions splitting the data

For every classified object we go through that structure (tree) checking the conditions on the way (e.g. "is there a word **genome** in the text?") from top to bottom, taking the label in the leaf as a result

The samples space is partitioned into the parallelograms, one label is set to each



https://www.slideshare.net/marinasantini1/lecture02-machine-learning

Logical classifiers: decision trees

ID3 algorithm

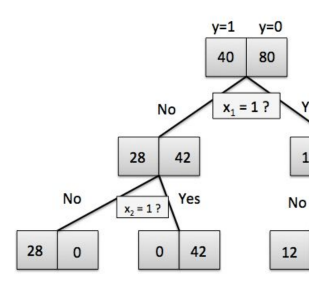
How "impure" the distribution of classes in S is is characterized by the entropy over shares of the samples of different labels

$$H(S) = \sum_{x \in X} -p(x) \log_2 p(x)$$

For every feature A and for every possible data partitioning
 T by it compute the information gain

$$IG(A,S) = H(S) - \sum_{t \in T} p(t)H(t)$$

- 2. Split the dataset using the feature and the partitioning with the MAX **IG**
- 3. Do 1-2 recursively with the subsets until there are no more samples or until **IG** stops to grow



Logical classifiers: decision trees

Classical algorithms: ID3, C4.5, CART, ...

A few heuristics to fight with overfitting

E.g. **pruning**: we replace the subtree with a leaf with the most frequent label in the former subtree if that doesn't hurt the quality of predictions on the **dev set**



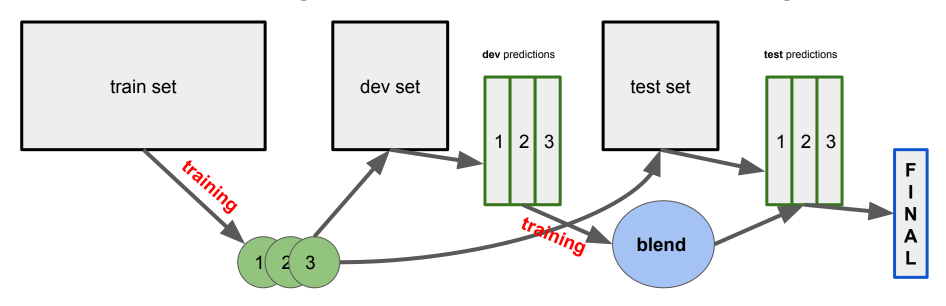
Decision trees: discussion

- + easy to interpret
- + don't have many assumptions on what the solution should look like
- overfit easily
- not that great for large dimensions

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Machine learning models ensembles: blending



We train at **train** set, we tune weights at **dev**, we check quality at **test**

- if you don't tune too hard, this helps to overcome overfitting
- main advantage: quick and dirty; the first thing to try

Machine learning models ensembles

Using multiple models for predictions may help

- not to ovefit
- to get a more rich solutions space than of any of the models the ensemble is composed of

Blending: joining the predictions of multiple models into one

- if we have probabilities, we can take the weighted sum
- weights for the linear combination may be trained + we can even train a model over predictions (BUT: overfitting alert!)
- if we predict classes, one can take a mode of the predicted labels



Machine learning models ensembles: bagging

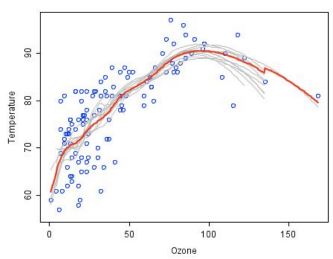
Bagging (bootstrap aggregating) —

sampling a few datasets from the training set, training classifiers on the independently

Feature bagging (attribute bagging, random subspace method) — sampling **subsets of features** and training classifiers on such sets independently

The resulting model is a consensus or a weighted vote

This allows for being more confident in predictions and helps overcome overfitting



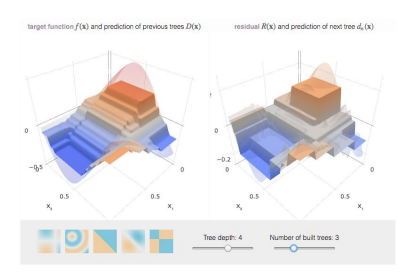
https://en.wikipedia.org/wiki/Bootstrap aggregating

Machine learning models ensembles: boosting

The core idea is to use a bunch of weak classifiers (non-random though) to build a strong one

Usually done like this:

- 1) incrementally training weak classifiers
- when adding each of them we increase the weight of previously wrongly classified samples
- 3) classifiers are added into the composition with the weight reflecting the quality they've shown



Cool demo

More stuff one needs to know

- other ways to measure quality, e.g., comparison with random predictions
- **feature selection** (PMI, DIA, Chi-square, ...)
- how to deal with label-imbalanced datasets
- how to deal with small training data
- tuning hyperparameters methods
 (grid search, random search, bayesian optimization, gradient-based optimization)

- ..

Important special cases

Sentiment analysis: building 'sentimental words' vocabularies, e.g.

- semi-automatic (given initial sentimental seed words)
- custom vocabulary building, e.g. for specific domains

Topic classification:

- topic hierarchy building; the less supervision there is, the better
- dealing with the case where there is no true topic in label list yet

Tools and instruments

Models zoos to give each a try:

- Weka (GUI)
- Scikit-Learn
- Mallet

Text classifiers can be implemented using: nltk, spaCy, H2O, mllib, Vowpal Wabbit, BigARTM, ...

Standard datasets for English:

- 20 Newsgroups (18k posts; 20 topics)
- Reuters Newswire Topic Classification (Reuters-21578; topical categories)
- IMDB Movie Review Sentiment Classification (stanford; sentiment)
- News Group Movie Review Sentiment Classification (cornell; sentiment)

Datasets are also many, any colour you like

Used/recommended literature

- Yandex Data School course on machine learning + similar lecture notes: this
- 2. The Elements of Statistical Learning and other classical books on machine learning (classification is everywhere)
- 3. Martin/Jurafsky, Chapters 6-7 in Ed. 3
- 4. Intro into IR (NB, kNN, Rocchio, SVM,...)
- 5. Wikipedia
- 6. CSC lectures, 2014 [Russian]

Text classification

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This time thanks go to Denis Kiryanov, Semyon Danilov, Viktor Evstratov

Machine learning models ensembles: boosting

AdaBoost

the final algorithm looks like this

$$F_T(x) = \sum_{t=1}^T f_t(x)$$

every classifier suggests a hypothesis per sample

$$h(x_i)$$

we are to find alpha parameter such that the error is minimized on the current iteration

$$E_t = \sum_i E[F_{t-1}(x_i) + lpha_t h(x_i)]$$