Unit 10 Ensemble Methods and Decision Trees

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ECE 5307: Introduction to Machine Learning, Sp23

Learning objectives

- Understand intuition behind ensemble methods: "the wisdom of the crowd"
- Understand parallel ensemble methods
 - bagging, pasting
 - random feature selection
- Understand decision trees
 - feature thresholding and decision regions
 - training via top-down tree induction
 - homogeneity metrics: variance reduction, gini impurity
 - ensemble extension: random forests
- Understand boosting, or sequentially trained ensemble methods
 - Adaboost
 - gradient boosting
 - XGBoost

Outline

Parallel Ensemble Methods: Bagging and Pasting

Decision Trees and Random Forests

Boosting: Sequentially Trained Ensemble Methods

The wisdom of the crowd

- Suppose that you want to determine the heavy side of a biased coin
 - Suppose "heads" occurs 70% and "tails" occurs 30%
- To predict the heavy side, you flip it N times and choose the majority vote
 - You'll be correct with probability $\frac{N}{\text{prob}}$ $\frac{1}{70\%}$ $\frac{11}{92\%}$ $\frac{21}{97\%}$ $\frac{31}{99\%}$
 - These values follow from the cdf of the binomial distribution
 - Interpretation: As you use more independent trials, the variance decreases
- Implications for classification:
 - Say you have 31 binary classifiers, each correct only 70% of the time
 - Individual classifiers ("weak learners") do only a bit better than random guessing
 - A majority vote ("strong learner") will classify with 99% accuracy!
 - Caveat: We've assumed the classifiers generate statistically independent outcomes
 - As they become more dependent, there will be less improvement from ensembling

Voting classifiers

- Given an ensemble (i.e., a collection) of base classifiers . . .
 - the hard voting classifier takes the majority vote of hard decisions
 - the soft voting classifier averages their soft outputs (i.e., pmfs) and then chooses the maximizing class . . . which usually works better
 - Both are implemented in sklearn. VotingClassifier
- The voting classifier will be better than the base classifiers if the base classifiers are sufficiently diverse (i.e., sufficiently independent)
- There are different ways to generate diverse ensembles of classifiers:
 - 1 Train a common prediction model using different datasets
 - 2 Train different prediction models using a common dataset:
 - Use different types of model (e.g., logistic regression, SVM, neural net)
 - Use same type but inject randomness (e.g., use random subsets of features)
- Similar ideas can be used to generate an ensemble of base regressors

Bagging and pasting

- Two main ways to introduce data diversity:
 - 1 pasting: draw training samples i without replacement
 - each sample is used exactly once by one predictor
 - f 2 bootstrap aggregating (or bagging): draw training samples i with replacement
 - each sample may be used several times, and some samples never used
 - bagging usually works better than pasting
- Bagging:
 - Usually draw n samples per predictor (where n is total # training samples)
 - When n is large, this implies $e^{-1} \approx 37\%$ samples go unused (per predictor)
 - Can use these "out of bag" samples for cross-validation!
- Implemented in sklearn.BaggingClassifier & sklearn.BaggingRegressor:
 - Choose bootstrap=True for bagging (or =False for pasting)
 - Choose oob_score=True to report out-of-bag cross-validation

Random patches and random subspaces

- Recall that using random subsets of features is another way to add diversity
 - Useful when features are high-dimensional and redundant
 - Can sample features j with replacement (i.e., "bootstrap") or without
- Terminology:
 - Using both random data i and features j is known as "random patches"
 - Using random features j but full data is known as "random subspaces"
- Implemented in sklearn.BaggingClassifier & sklearn.BaggingRegressor:
 - Set max_features < d for feature randomization
 - Set bootstrap_features=True for bootstrapping

Outline

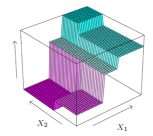
Parallel Ensemble Methods: Bagging and Pasting

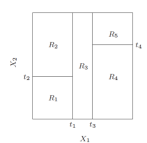
Decision Trees and Random Forests

Boosting: Sequentially Trained Ensemble Methods

Decision trees

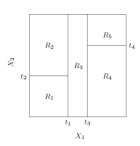
- Consider a supervised learning task (e.g., regression or classification)
 - Suppose data is $\{(\boldsymbol{x}_i,y_i)\}_{i=1}^n$ with $\boldsymbol{x}_i \in \mathbb{R}^d$ and $y_i \in \mathbb{R}$
- Approach:
 - 1 Partition feature domain \mathbb{R}^d into L regions $\{R_\ell\}_{\ell=1}^L$
 - 2 Output $z_\ell \in \mathbb{R}$ whenever ${\boldsymbol x} \in R_\ell$
 - For regression, could set z_ℓ at sample mean of $\{y_i\}_{i\in S_\ell}$, where $S_\ell=\{i: \ x_i\in R_\ell\}$
 - lacksquare For classification, could set z_ℓ at sample mode, i.e., most common value in $\{y_i\}_{i\in S_\ell}$
- Regression example with dimension d = 2:

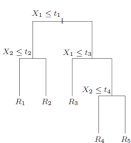




Why are they "trees"?

- We normally construct the regions by thresholding one feature at a time!
 - Thus, the decision boundaries are always parallel to the coordinate axes
- Then can view prediction as a decision tree:
 - The domain of x (i.e., \mathbb{R}^d) forms the root of the tree
 - The decision regions $\{R_\ell\}$ are the L leaves of the tree
- **Example** with dimension d=2 and L=5 leaves:





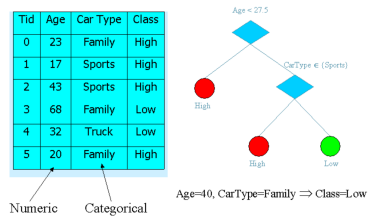
Top-down induction of decision trees

Decision trees are trained in a top-down manner:

- Start with entire data set: $\{(\boldsymbol{x}_i,y_i)\}_{i=1}^n$ and define index set $S \triangleq \{1,\ldots,n\}$
- First split S into subsets (S_1, S_2) , so that labels $\{y_i\}$ are most homogeneous (i.e., similar) within each subset
 - Splitting is performed by thresholding some feature j:
 - Ordinal feature: $S_1 = \{i: x_{ij} \le t\}$ and $S_2 = \{i: x_{ij} > t\}$
 - Categorical feature: $S_1 = \{i: x_{ij} \in \{A,C\}\}$ and $S_2 = \{i: x_{ij} \in \{B,D\}\}$
 - lacktriangle To find "best" split, must search jointly over feature j and threshold t
 - Many ways to quantify homogeneity (will discuss later)
- lacktriangle Then split each subset S_1 and S_2 further, using the same procedure
- Repeat until . . .
 - \blacksquare labels are perfectly homogeneous within a subset (e.g., all y_i are same), or
 - stopping condition: e.g., subsets have min # samples, tree has max depth, etc

Decision tree example

Example: Risk prediction with n=6 samples, d=2 features, L=3 leaves. Feature j=1 (age) is ordinal and feature j=2 (car type) is categorical:



https://web.fhnw.ch/personenseiten/taoufik.nouri/Data%20Mining/Course/Course3/DM-Part%203.htm

Homogeneity metrics for regression

- lacktriangle Recall that the goal is to maximize homogeneity within S_1 and S_2
 - lacksquare Equivalently, we want to minimize inhomogeneity within S_1 and S_2
- Let us first consider regression, where labels $y_i \in \mathbb{R}$
- lacksquare Inhomogeneity could be measured by the variance after splitting $S o (S_1,S_2)$:
 - The mean in subset S_{ℓ} is $\mu_{\ell} \triangleq \frac{1}{|S_{\ell}|} \sum_{i \in S_{\ell}} y_i$, for $\ell \in \{1, 2\}$
 - The variance in subset S_ℓ is $v_\ell \triangleq \frac{1}{|S_\ell|} \sum_{i \in S_\ell} (y_i \mu_\ell)^2$, for $\ell \in \{1, 2\}$
 - Thus the (average) variance after splitting is $v \triangleq \frac{|S_1|}{|S|} v_1 + \frac{|S_2|}{|S|} v_2$
- Another option is to use absolute error $\frac{1}{|S_{\ell}|} \sum_{i \in S_{\ell}} |y_i \mu_{\ell}|$ instead of v_{ℓ} , since it is more robust to outliers
- We'll see a more sophisticated metric when discussing XGBoost on page 31

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Homogeneity metrics for classification

- Now consider K-ary classification, where $y_i \in \{1, \dots, K\} \ \forall i$
 - For subset S_ℓ , the empirical label pmf is $\{p_{\ell k}\}_{k=1}^K$, where $p_{\ell k} = \frac{\sum_{i \in S_\ell} \mathbb{1}_{y_i = k}}{|S_\ell|}$
- Again, want to minimize inhomogeneity within S_1 and S_2
- One popular measure of inhomogeneity is Gini impurity:

$$I_{\mathsf{G}} = \frac{|S_1|}{|S|} I_{\mathsf{G},1} + \frac{|S_2|}{|S|} I_{\mathsf{G},2}, \quad \text{where} \quad I_{\mathsf{G},\ell} \triangleq 1 - \sum_{k=1}^K p_{\ell k}^2$$

- Note that $I_{G,\ell} \in \left[0, 1 \frac{1}{L}\right]$
- $I_{G,\ell} = 0$ means perfectly pure/homogeneous, i.e., constant y_i for $i \in S_\ell$
- Other criteria include entropy, misclassification error, and chi-square
 - See Hastie, Tibshirani, Friedman, The Elements of Statistical Learning, 2nd Ed., 2009

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Top-down induction for generic loss

- lacksquare Say we want a tree $f(\cdot)$ that minimizes a generic loss $\sum_{i=1}^n L(y_i,f(m{x}_i))$
- lacksquare To design the tree, it helps to write $f(m{x})=z_{q(m{x})}$, where
 - lacksquare z_ℓ is the value assigned to leaf ℓ
 - $lacksquare q(\cdot)$ is the decision function that maps $oldsymbol{x}$ to ℓ
- Thus tree design becomes " $\arg\min_{{m{z}},q}\sum_{\ell}\sum_{i:q({m{x}}_i)=\ell}L(y_i,z_\ell)$ "
- Remember that we design the tree one split at a time. Consider splitting $S \to (S_1, S_2)$. This would cause the loss to go from

$$\min_{z_1} \sum_{i \in S} L(y_i, z_1) \quad \rightarrow \quad \min_{z_1} \sum_{i \in S_1} L(y_i, z_1) + \min_{z_2} \sum_{i \in S_2} L(y_i, z_2)$$

- lacksquare So, we try all feature/threshold pairs (j,t), each of which leads to a split (S_1,S_2) . We evaluate the loss for each split and choose the best one.
- If the best split does not improve the loss, we do no splitting.

Advantages and disadvantages of decision trees

Advantages:

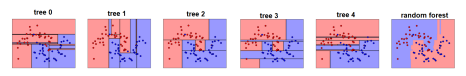
- Very general
 - Very few assumptions made on data; can work with any dataset
 - Don't need to standardize the features.
- Very interpretable: "white box model"
 - Easy to understand how a tree arrives at its prediction
- lacksquare Prediction is fast: $O(\log_2 n)$ decisions to process a test sample $oldsymbol{x}$

Disadvantages:

- lacktriangle Training can be expensive if all features j are considered at every split
 - $lue{}$ Solution: Restrict to a few randomly chosen features j at each split
- Prone to overfitting
 - Highly dependent on training data: changing one sample can change entire tree!
 - Can regularize by enforcing min # samples per subset, max depth, etc
- → There is a good overview of decision trees in the sklearn documentation!

Random forests

- Decision trees tend to overfit: low bias & high variance
- Idea: Use a random forest: An ensemble of trees generated by . . .
 - using a random subset of data to train each tree (i.e., bagging or pasting)
 - considering a random subset of features j at each split
- As with other ensemble methods, results are averaged to make a prediction
- Implemented in sklearn.RandomForestClassifier & Regressor



from machine-learning-algorithms-ensemble-methods-bagging-boosting-and-random-forests

Outline

Parallel Ensemble Methods: Bagging and Pasting

Decision Trees and Random Forests

Boosting: Sequentially Trained Ensemble Methods

Boosting

- Previously we described ensemble methods that train in parallel
- Now we discuss ensemble methods that train sequentially
 - This is called "boosting"
- Some of the best known boosting methods are:
 - adaptive boosting, or Adaboost (1996)
 - gradient boosting (2001)
 - extreme gradient boosting or XGBoost (2016)



Adaboost

Adaboost sequentially trains an ensemble of predictors $\{f_m(\cdot)\}_{m=1}^M$ as follows:

For
$$m = 1 \dots M$$
,

- **1** Train predictor $f_m(\cdot)$ to minimize some weighted loss $\sum_{i=1}^n w_{mi} L(y_i, f_m(\boldsymbol{x}_i))$
- 2 For next round, assign larger weights $w_{m+1,i}$ to samples i with higher loss
- The final prediction is done using a weighted average

$$F_M(m{x}) = \sum_{m=1}^M lpha_m f_m(m{x}), \;\; ext{for some learned weights } \{lpha_m\}$$

- For simplicity, we will focus on the design of binary classifiers $f_m(\cdot) \in \{-1,1\}$
 - Other versions of Adaboost exist for regression and non-binary classification

Adaboost: Derivation

- Consider binary classification, with training data $\{(x_i, y_i)\}_{i=1}^n$ for $y_i = \pm 1$
- Adaboost's base classifiers $f_m(\cdot)=\pm 1$, for $m=1\dots M$, are trained as follows:
 - Define the step-m boosted classifier $F_m(x) \triangleq F_{m-1}(x) + \alpha_m f_m(x)$
 - Initialize $F_0(x) = 0$. For m = 1, ..., M, design $\alpha_m > 0$ and f_m so that F_m minimizes the exponential loss $\mathcal{L}_m = \sum_{i=1}^n e^{-y_i F_m(x_i)}$
- Plugging $F_m(\cdot)$ into \mathcal{L}_m , we find that

$$\mathcal{L}_m = \sum_{i=1}^n e^{-y_i(F_{m-1}(\boldsymbol{x}_i) + \alpha_m f_m(\boldsymbol{x}_i))} = \sum_{i=1}^n e^{-y_i F_{m-1}(\boldsymbol{x}_i)} e^{-y_i \alpha_m f_m(\boldsymbol{x}_i)}$$

$$= e^{-\alpha_m} \sum_{i:y_i = f_m(\boldsymbol{x}_i)} w_{mi} + e^{\alpha_m} \sum_{i:y_i \neq f_m(\boldsymbol{x}_i)} w_{mi} \text{ since } f_m \text{ and } y_i \text{ are } \pm 1$$

$$= e^{-\alpha_m} \sum_{i=1}^n w_{mi} + (e^{\alpha_m} - e^{-\alpha_m}) \sum_{i:y_i \neq f_m(\boldsymbol{x}_i)} w_{mi}$$

Adaboost: Derivation (cont.)

■ From the previous expression, we see that the \mathcal{L}_m -minimizing classifier $f_m(\cdot)$ is that which minimizes the weighted misclassification loss

$$\sum_{i:y_i \neq f_m(\boldsymbol{x}_i)} w_{mi} = \sum_{i=1}^n w_{mi} \mathbb{1}_{y_i \neq f_m(\boldsymbol{x}_i)}$$

- At step m=1, use $w_{1i}=1$ $\forall i$, and thus train $f_1(\cdot)$ on $\{(\boldsymbol{x}_i,y_i)\}_{i=1}^n$ as usual
- At step m>1, the weight w_{mi} is large if F_{m-1} made a wrong decision on y_i
- The \mathcal{L}_m -minimizing α_m can be found by solving $\partial \mathcal{L}_m/\partial \alpha_m=0$, which gives

$$\alpha_m = \frac{1}{2} \bigg(\frac{1 - \epsilon_m}{\epsilon_m} \bigg) \quad \text{for weighted error rate } \epsilon_m = \frac{\sum_{i: y_i \neq f_m(\boldsymbol{x}_i)} w_{mi}}{\sum_i w_{mi}}$$

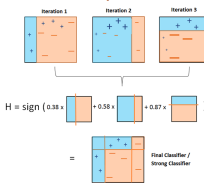
- In summary, to train Adaboost, initialize $w_{1i}=1 \ \forall i$ and then, for $m=1\ldots M$, compute $f_m(\cdot)$, ϵ_m , α_m , $F_m(\cdot)$, and $\{w_{m+1,i}\}_{i=1}^n$
- lacksquare Once trained, predict the label of a test sample $m{x}$ using $F_M(m{x})$

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Adaboost: Implementation

- Adaboost classification implemented via sklearn.AdaBoostClassifier
 - By default, it uses 50 decision tree classifiers of depth 1 ("stumps")
- Adaboost regression implemented via sklearn.AdaBoostRegressor
 - By default, it uses 50 decision tree regressors of depth 3

AdaBoost Classifier Working Principle with Decision Stump as a Base Classifier



Gradient boosting

■ Goal: Design predictor $F_M(\cdot)$ to minimize a training loss of the form

$$\mathcal{L}(F_M) = \sum_{i=1}^n Lig(y_i, F_M(oldsymbol{x}_i)ig)$$
 for some $L(\cdot, \cdot)$

where $F_M(\cdot) = \sum_{m=1}^M \alpha_m f_m(\cdot)$ with $f_m(\cdot) \in \mathcal{F}$ is sequentially learned

- \blacksquare Adaboost did this with exponential loss $L(y,F)=e^{-yF}$ and binary $f_m=\pm 1$
- lacksquare Gradient boosting tries to do this for general L and ${\mathcal F}$
- Ideally, we would like to do the following:
 - lacksquare Train $f_1 \in \mathcal{F}$ to minimize $\sum_{i=1}^n Lig(y_i, f_1(m{x}_i)ig)$, set $F_1 = f_1$
 - For $m = 2 \dots M$: $F_m = F_{m-1} + \operatorname*{arg\,min}_{\alpha_m, f_m \in \mathcal{F}} \sum_{i=1}^n L(y_i, F_{m-1}(\boldsymbol{x}_i) + \alpha_m f_m(\boldsymbol{x}_i))$
 - lacksquare But for many choices of loss L, this optimization is too difficult
- Idea: Instead of exact minimization, settle for a gradient step, i.e.,

For
$$m = 2 \dots M$$
: $F_m = F_{m-1} - \alpha_m \nabla \mathcal{L}(F_{m-1})$

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Gradient boosting: Details

- Problem:
 - We are limited to updates of the form

$$F_m = F_{m-1} + \alpha_m f_m$$
, for $f_m \in \mathcal{F}$, where \mathcal{F} is set of base predictors

• Gradient descent does not constrain the update to \mathcal{F} :

$$F_m = F_{m-1} + \alpha_m \big[-\nabla \mathcal{L}(F_{m-1}) \big], \text{ with } \nabla \mathcal{L}(F_{m-1}) = \sum_{i=1}^n \underbrace{\frac{\partial L(y_i, F_{m-1}(\boldsymbol{x}_i))}{\partial F}}_{\triangleq -r_{m,i}}$$

- Solution: Train $f_m \in \mathcal{F}$ to be close to $-\nabla \mathcal{L}(F_{m-1})$
- In particular, for m = 2, ..., M, we do the following:
 - lacksquare First train $f_m \in \mathcal{F}$ to minimize $\sum_{i=1}^n ig(r_{mi} f_m(m{x}_i)ig)^2$
 - Then choose α_m via line-search: $\alpha_m = \arg\min_{\alpha} \sum_{i=1}^n L(y_i, F_{m-1}(\boldsymbol{x}_i) + \alpha f_m(\boldsymbol{x}_i))$
 - lacksquare Finally, update boosted predictor: $F_m(oldsymbol{x}) = F_{m-1}(oldsymbol{x}) + lpha_m f_m(oldsymbol{x})$

Gradient boosting: Implementation

■ In practice, it helps to slow down the updates using "learning rate" $\mu \in (0,1]$:

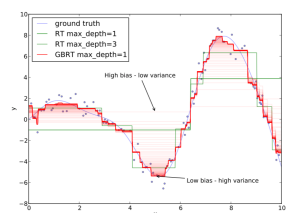
$$F_m(\mathbf{x}) = F_{m-1}(\mathbf{x}) + \mu \alpha_m f_m(\mathbf{x})$$

- Classification implemented via sklearn.GradientBoostingClassifier
 - By default, it uses 100 decision tree classifiers of depth 3
 - By default, uses $\mu = 0.1$
- Regression implemented via sklearn.GradientBoostingRegressor
 - By default, it uses 100 decision tree regressors of depth 3
 - By default, uses $\mu = 0.1$

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Gradient boosting: Example

```
from sklearn.ensemble import GradientBoostingRegressor
est = GradientBoostingRegressor(n_estimators=2000, max_depth=1).fit(X, y)
for pred in est.staged_predict(X):
   plt.plot(X[:, 0], pred, color='r', alpha=0.1)
```



from Gradient Boosted Regression Trees by Prettenhofer & Louppe

Extreme gradient boosting (XGBoost)

- XGBoost is an evolution of gradient boosting that has won many recent machine-learning contests
 - On low-dimensional tasks, XGBoost often performs as well as neural nets, but it is much easier to design/train!
- Compared to gradient boosting, XGBoost uses...
 - second-order Taylor series approximation
 - regularization
 - sophisticated tree-search
 - many parallelization and hardware acceleration tricks
- The XGBoost package is not part of sklearn, but compatible with it
 - Once you download the XGBoost package, you can use it just like sklearn

XGBoost: Derivation

- \blacksquare XGBoost sequentially designs a boosted predictor $F_M(m{x}) = \sum_{m=1}^M f_m(m{x})$
- Similar to gradient boosting, we'd ideally like to do the following:
 - Train $f_1 \in \mathcal{F}$ to minimize $\sum_{i=1}^n L(y_i, f_1(\boldsymbol{x}_i))$, set $F_1 = f_1$
 - For

$$m = 2...M$$
: $F_m = F_{m-1} + \underset{f_m \in \mathcal{F}}{\operatorname{arg min}} \left[\sum_{i=1}^n L(y_i, F_{m-1}(\boldsymbol{x}_i) + f_m(\boldsymbol{x}_i)) + \phi(f_m) \right]$

- lacksquare Note that we added regularization ϕ to the cost function
- Unfortunately, this optimization problem is too difficult to solve exactly
- Idea: Simplify using a 2nd-order Taylor series approximation
 - Use $L(y_i, F_{m-1}(\boldsymbol{x}_i) + f_m(\boldsymbol{x}_i)) \approx L(y_i, F_{m-1}(\boldsymbol{x}_i)) + g_{mi}f_m(\boldsymbol{x}_i) + \frac{1}{2}h_{mi}f_m^2(\boldsymbol{x}_i)$ where $g_{mi} \triangleq \frac{\partial L(y_i, F_{m-1}(\boldsymbol{x}_i))}{\partial F}$ and $h_{mi} \triangleq \frac{\partial^2 L(y_i, F_{m-1}(\boldsymbol{x}_i))}{\partial^2 F}$
 - Then the optimization problem becomes

$$\underset{f_m \in \mathcal{F}}{\operatorname{arg \, min}} \left[\sum_{i=1}^{n} \left(g_{mi} f_m(\boldsymbol{x}_i) + \frac{1}{2} h_{mi} f_m^2(\boldsymbol{x}_i) \right) + \phi(f_m) \right] \tag{1}$$

XGBoost: Derivation 2

- Now suppose that $f(\cdot)$ is a decision tree with max L leaves
 - As on page 15, write $f(\boldsymbol{x}) = z_{q(\boldsymbol{x})}$ where . . .
 - $z_{\ell} \in \mathbb{R}$ is the output assigned to leaf ℓ
 - $ullet q(\cdot)$ is the function that assigns $oldsymbol{x} \in \mathbb{R}^d$ to a leaf $\ell \in \{1,\dots,L\}$
- XGBoost uses the regularization

$$\phi(f) = \frac{\lambda}{2} \sum_{\ell=1}^L z_\ell^2 + \gamma L, \quad \text{for tunable } \lambda > 0 \text{ and } \gamma > 0$$

lacktriangle Omitting the "m" notation, the XGBoost optimization problem (1) becomes

$$\arg\min_{q, \mathbf{z}} \sum_{\ell=1}^{L} \sum_{i: q(\mathbf{x}_i) = \ell} \left(g_i z_\ell + \frac{1}{2} h_i z_\ell^2 \right) + \frac{\lambda}{2} \sum_{\ell=1}^{L} z_\ell^2 + \gamma L$$

$$= \arg\min_{q, \mathbf{z}} \sum_{\ell=1}^{L} \left[\left(\sum_{i: q(\mathbf{x}_i) = \ell} g_i \right) z_\ell + \frac{1}{2} \left(\lambda + \sum_{i: q(\mathbf{x}_i) = \ell} h_i \right) z_\ell^2 + \gamma \right]$$
(2)

XGBoost: Derivation 3

lacksquare We can optimize over $oldsymbol{z} \in \mathbb{R}^L$ to yield

$$z_{\ell} = -\frac{\sum_{i:q(\boldsymbol{x}_i)=\ell} g_i}{\lambda + \sum_{i:q(\boldsymbol{x}_i)=\ell} h_i}, \quad \ell = 1, \dots, L$$

and plug this back into (2) to obtain an optimization problem over $q(\cdot)$:

$$\arg\min_{q} \sum_{\ell=1}^{L} \left[2\gamma - \frac{\left(\sum_{i:q(\boldsymbol{x}_{i})=\ell} g_{i}\right)^{2}}{\lambda + \sum_{i:q(\boldsymbol{x}_{i})=\ell} h_{i}} \right]$$

- \blacksquare Using the above loss, the tree $q(\cdot)$ can be similarly to page 15: Choose a feature j and search for the threshold t that maximally reduces loss after a split
- For example, consider the split $S o (S_1, S_2)$. The loss would change as

$$2\gamma - \frac{\left(\sum_{i \in S} g_i\right)^2}{\lambda + \sum_{i \in S} h_i} \rightarrow \left[2\gamma - \frac{\left(\sum_{i \in S_1} g_i\right)^2}{\lambda + \sum_{i \in S_1} h_i} + 2\gamma - \frac{\left(\sum_{i \in S_2} g_i\right)^2}{\lambda + \sum_{i \in S_2} h_i}\right]$$

If the loss doesn't decrease, then it's better *not* to split!

■ Further details can be found in the original paper

XGBoost: Example

```
# First XGBoost model for Pima Indians dataset
2 from numpy import loadtxt
3 from xaboost import XGBClassifier
4 from sklearn.model_selection import train_test_split
5 from sklearn.metrics import accuracy_score
6 # load data
7 dataset = loadtxt('pima-indians-diabetes.csv', delimiter=",")
8 # split data into X and y
9 X = dataset[:,0:8]
10 Y = dataset\Gamma:.87
11 # split data into train and test sets
12 \text{ seed} = 7
13 test size = 0.33
14 X_train, X_test, v_train, v_test = train_test_split(X, Y, test_size=test_size, ran
15 # fit model no training data
16 model = XGBClassifier()
17 model.fit(X_train, y_train)
18 # make predictions for test data
19 v pred = model.predict(X test)
20 predictions = [round(value) for value in v_pred]
21 # evaluate predictions
22 accuracy = accuracy_score(y_test, predictions)
23 print("Accuracy: %.2f%%" % (accuracy * 100.0))
```

Running this example produces the following output.

```
1 Accuracy: 77.95%
```

This is a good accuracy score on this problem, which we would expect, given the capabilities of the model and the modest complexity of the problem.

from develop-first-xgboost-model-python-scikit-learn

Tuning trees, random forests, and XGBoost

To get the best performance on test data, several hyperparameters must be tuned. Here are some good resources on how to do that:

- Importance of decision tree hyperparameters on generalization
- A beginner's guide to random forest hyperparameter tuning
- XGBoost: Notes on parameter tuning
- Complete guide to parameter tuning in XGBoost with code
- Hyperparameter tuning for hyperaccurate XGBoost model

Learning objectives

- Understand intuition behind ensemble methods: "the wisdom of the crowd"
- Understand parallel ensemble methods
 - bagging, pasting
 - random feature selection
- Understand decision trees
 - feature thresholding and decision regions
 - training via top-down tree induction
 - homogeneity metrics: variance reduction, gini impurity
 - ensemble extension: random forests
- Understand boosting, or sequentially trained ensemble methods
 - Adaboost
 - gradient boosting
 - XGBoost