# Statistical Learning Common Core

**Understanding Machine Learning** 

## General Recipe

statistical learning algorithm by combining:

• model (e.g., linear function, Gaussian distribution)

objective function (e.g., squared residuals)

optimization algorithm (e.g., gradient descent)

## Consider Supervised Learning Scenario

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map input to output: y = f(x) (estimated: \hat{f}(x))
random variables Y and X = (X_1, X_2, \dots, X_p) \leftarrow usually high-dimensional
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curve fitting / parameter estimation: fit train data set of  $(y_i, x_i)$  pairs  $\rightarrow$  minimization of cost function

#### consider discriminative models:

- predict conditional density function  $p(y|x_i)$  (as opposed to generative models predicting p(y,x))
- often just conditional mean (depending on used loss)  $E[Y|X=x_i]$  of  $p(y|x_i)$

## Model

### Model in Linear Regression

fit: 
$$\hat{f}(x_i)$$

$$y_i = \hat{\alpha} + \sum_{j=1}^{p} \hat{\beta}_j x_{ij} + \varepsilon_i$$

to be estimated:

•  $\hat{\alpha}$ ,  $\hat{\beta}$ •  $\hat{\sigma}^2 = \frac{1}{n} \sum_{i=1}^n \left( y_i - \hat{f}(x_i) \right)^2$ 

(approximating assumed true  $\alpha$ ,  $\beta$ ,  $\sigma$ )

#### predict:

$$\hat{y}_i = E[Y|X = x_i] = \hat{f}(x_i)$$

$$p(y|\mathbf{x}_i) = \mathcal{N}(y; \hat{y}_i, \hat{\sigma}^2)$$

Gaussian mean variance (reflected by  $\varepsilon_i$  in fit)

## Model Ingredients

- underlying probability distribution of Y
- approximated functional form  $\hat{f}(x) \rightarrow$  depends on used ML algorithm

$$p(y|\mathbf{x}_i) = PDF(y; \hat{f}(\mathbf{x}_i), ...)$$

assumed Probability Density Function of target

 $\hat{f}(x)$  often estimation of location parameter

other parameters (e.g., scale parameter) often not predicted per sample (assume homoscedasticity)

## Principle of Maximum Entropy

How to choose the right model, i.e., the assumed underlying probability distribution of Y generating the observed data?

Occam's razor: pick distribution with fewest assumptions

 $\rightarrow$  distribution with maximum entropy under given range and moment constraints discrete:  $H(X) = -\sum p_k \log p_k$ , continuous:  $H(X) = -\int p(x) \log p(x) dx$  ( $\rightarrow$  bulk of entropy from tails of distributions)

examples for maximum entropy distributions:

- normal distribution: continuous on  $(-\infty, \infty)$  with particular mean and variance
- Poisson distribution: discrete on  $[0, \infty)$  with particular mean

## Objective Function

#### Loss Function

loss function L: expressing deviation between prediction and target

$$L(y_i, \hat{f}(\boldsymbol{x}_i); \widehat{\boldsymbol{\theta}})$$

with  $\widehat{\boldsymbol{\theta}}$  corresponding to parameters of model  $\widehat{f}(\boldsymbol{x})$ 

e.g.,  $\widehat{\alpha}$ ,  $\widehat{\beta}$  in linear regression

e.g., squared residuals (for regression problems):

$$L(y_i, \hat{f}(\boldsymbol{x}_i); \widehat{\boldsymbol{\theta}}) = (y_i - \hat{f}(\boldsymbol{x}_i; \widehat{\boldsymbol{\theta}}))^2$$

#### Cost Function

averaging losses over (empirical) training data set:

$$J(\widehat{\boldsymbol{\theta}}) = \frac{1}{n} \sum_{i=1}^{n} L(y_i, \widehat{f}(\boldsymbol{x}_i); \widehat{\boldsymbol{\theta}})$$

cost function to be minimized according to model parameters  $\widehat{\boldsymbol{\theta}}$   $\rightarrow$  objective function

## Optimization

#### Cost Minimization

minimize training costs  $J(\widehat{\boldsymbol{\theta}})$  according to model parameters  $\widehat{\boldsymbol{\theta}}$ :

$$\nabla_{\widehat{\boldsymbol{\theta}}} J(\widehat{\boldsymbol{\theta}}) = 0$$

for mean squared error (aka least squares method):

$$\nabla_{\widehat{\boldsymbol{\theta}}} \, \frac{1}{n} \sum_{i=1}^{n} \left( y_i - \hat{f}(\boldsymbol{x}_i; \widehat{\boldsymbol{\theta}}) \right)^2 = 0$$

## Ordinary Least Squares

linear regression:  $\hat{f}(x_i) = \hat{\alpha} + \sum_{j=1}^p \hat{\beta}_j x_{ij}$ matrix notation (drop intercept  $\alpha$  for simplicity here):  $\hat{y} = \mathbf{X} \hat{\beta}$ 

solve normal equations for training data set:

$$\nabla_{\widehat{\boldsymbol{\beta}}} (\mathbf{y} - \mathbf{X} \,\widehat{\boldsymbol{\beta}})^{\mathrm{T}} (\mathbf{y} - \mathbf{X} \,\widehat{\boldsymbol{\beta}}) = 0$$

$$\rightarrow \widehat{\boldsymbol{\beta}} = (\mathbf{X}^{\mathrm{T}} \mathbf{X})^{-1} \mathbf{X}^{\mathrm{T}} \mathbf{y}$$

but usually no closed-form solution  $\rightarrow$  need for numerical optimization

## Maximum Likelihood Estimation

#### Likelihood

special objective function: likelihood function  ${\cal L}$ 

- value of predicted probability density function (model) at observed target
- as function of model parameters  $\widehat{m{ heta}}$

$$\mathcal{L}(\widehat{\boldsymbol{\theta}}; \boldsymbol{y}|\boldsymbol{x}) = p(\boldsymbol{y}|\boldsymbol{x}; \widehat{\boldsymbol{\theta}})$$

a little bit confusing:

 $p(y|x; \hat{\theta})$  is conditional density function if function of data with  $\hat{\theta}$  fixed  $p(y|x; \hat{\theta})$  is conditional likelihood function if function of  $\hat{\theta}$  with data fixed

## Independence Assumption

consider training data set (of size n) as one sample from unknown joint probability distribution of n independent (i.i.d.) sets of random variables  $(Y_1, X_1), (Y_2, X_2), \dots, (Y_n, X_n)$  (same thing as random sampling n times from (Y, X))

→ likelihood as product of univariate probability density functions:

$$\mathcal{L}(\widehat{\boldsymbol{\theta}}; \boldsymbol{y}|\boldsymbol{x}) = \prod_{i=1}^{n} p(y_i|\boldsymbol{x}_i; \widehat{\boldsymbol{\theta}})$$

... to be maximized according to estimated model parameters  $\widehat{m{ heta}}$  (known as maximum likelihood estimation)

## Negative Log-Likelihood

logarithmic transformation of  $\mathcal{L}$ : log-likelihood function

model, e.g., for linear regression:  

$$p(y_i|x_i) = \mathcal{N}(y; \hat{\alpha} + x_i \hat{\beta}, \sigma^2)$$

- with y, X given in training
- $\widehat{\boldsymbol{\theta}} = (\widehat{\alpha}, \widehat{\boldsymbol{\beta}}, \widehat{\sigma}^2)$  to be estimated

$$\ell(\widehat{\boldsymbol{\theta}}; \boldsymbol{y}|\boldsymbol{x}) = \ln\left(\mathcal{L}(\widehat{\boldsymbol{\theta}}; \boldsymbol{y}|\boldsymbol{x})\right) = \sum_{i=1}^{n} \ln\left(p(y_i|\boldsymbol{x}_i; \widehat{\boldsymbol{\theta}})\right)$$

- logarithm monotonic function  $\rightarrow$  maximum of  $\ell$  and  $\mathcal L$  at same  $\widehat{m heta}$  values
- most probability distributions (e.g., exponential family) only logarithmically concave (important for optimization)
- sum computationally more convenient than product

negative log-likelihood: minimization instead of maximization

$$\widehat{\boldsymbol{\theta}} = \operatorname{argmax}_{\widehat{\boldsymbol{\theta}}} \ell(\widehat{\boldsymbol{\theta}}; \boldsymbol{y} | \boldsymbol{x}) = \operatorname{argmin}_{\widehat{\boldsymbol{\theta}}} \left( -\ell(\widehat{\boldsymbol{\theta}}; \boldsymbol{y} | \boldsymbol{x}) \right)$$
 objective function

#### Maximum Likelihood

estimate parameters  $\widehat{\boldsymbol{\theta}} = (\widehat{\theta}_1, \widehat{\theta}_2, \cdots, \widehat{\theta}_k)$  solving

$$\widehat{\boldsymbol{\theta}} = \operatorname{argmax}_{\widehat{\boldsymbol{\theta}}} \ell(\widehat{\boldsymbol{\theta}}; \boldsymbol{y} | \boldsymbol{x})$$
 mode of likelihood

mode of likelihood as point estimate

**least squares method** corresponds to maximum likelihood estimation with

Gaussian model:  $\sim e^{-\frac{1}{2}\left(\frac{x-\mu}{\sigma}\right)^2}$ 

 $\rightarrow$  likelihood equations (if  $\ell$  differentiable in  $\widehat{\boldsymbol{\theta}}$ ):

$$\frac{\partial \ell}{\partial \widehat{\theta}_1} = 0$$
,  $\frac{\partial \ell}{\partial \widehat{\theta}_2} = 0$ , ...,  $\frac{\partial \ell}{\partial \widehat{\theta}_k} = 0$  (in short:  $\nabla_{\widehat{\theta}} \ell = 0$ )

can be solved explicitly for some cases (e.g., ordinary least squares for linear regression)

but usually no closed-form solution

→ need for numerical optimization (e.g., gradient descent)

### Interpretation of Maximum Likelihood

quantification of difference between two probability distributions

maximum likelihood estimation corresponds to minimization of Kullback-Leibler divergence (as well as cross entropy:  $D_{KL}(p||q) = H(p,q) - H(p)$ ) between true data-generating probability distribution (manifested by empirical distribution of training data) and probability distribution of model:

$$\underset{\widehat{\boldsymbol{\theta}}}{\operatorname{argmin}}_{\widehat{\boldsymbol{\theta}}} D_{KL} \left( p_{\operatorname{model}}(\boldsymbol{y}|\boldsymbol{x}; \widehat{\boldsymbol{\theta}}) \mid\mid p_{\operatorname{data}}(\boldsymbol{y}|\boldsymbol{x}) \right) = \underset{\widehat{\boldsymbol{\theta}}}{\operatorname{argmin}}_{\widehat{\boldsymbol{\theta}}} \int p_{\operatorname{data}}(\boldsymbol{y}|\boldsymbol{x}) \log \frac{p_{\operatorname{data}}(\boldsymbol{y}|\boldsymbol{x})}{p_{\operatorname{model}}(\boldsymbol{y}|\boldsymbol{x}; \widehat{\boldsymbol{\theta}})} \, d\boldsymbol{y}$$

$$= \underset{\widehat{\boldsymbol{\theta}}}{\operatorname{argmin}}_{\widehat{\boldsymbol{\theta}}} \frac{1}{n} \sum_{i=1}^{n} \log \frac{p_{\operatorname{data}}(\boldsymbol{y}_{i}|\boldsymbol{x}_{i})}{p_{\operatorname{model}}(\boldsymbol{y}_{i}|\boldsymbol{x}_{i}; \widehat{\boldsymbol{\theta}})} = \underset{\widehat{\boldsymbol{\theta}}}{\operatorname{argmax}}_{\widehat{\boldsymbol{\theta}}} \sum_{i=1}^{n} \ln \left( p_{\operatorname{model}}(\boldsymbol{y}_{i}|\boldsymbol{x}_{i}; \widehat{\boldsymbol{\theta}}) \right) = \underset{no \text{ contribution from } p_{\operatorname{data}}}{\operatorname{pdata}} \ell \left( \widehat{\boldsymbol{\theta}}; \boldsymbol{y} \mid \boldsymbol{x} \right)$$

make the model distribution match the empirical distribution

(mean squared error corresponds to cross-entropy between empirical distribution and Gaussian model)

## Iterative Optimization

#### **Gradient Descent**

usually (except for special cases like ordinary least squares) no closed-form solution to ML optimization problems like minimization of a cost function or maximization of a likelihood function:

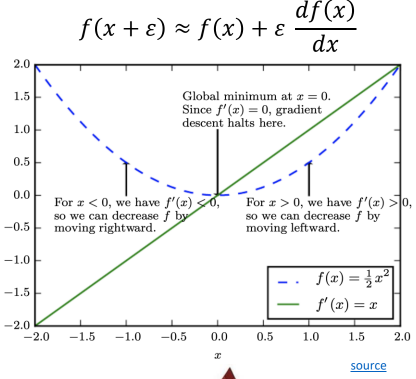
$$\nabla_{\widehat{\boldsymbol{\theta}}} J(\widehat{\boldsymbol{\theta}}) = 0$$

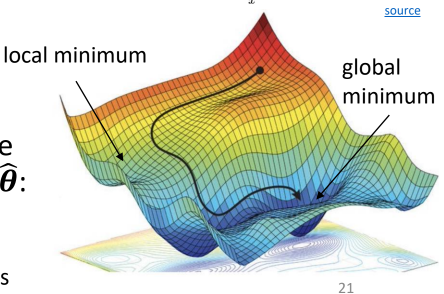
→ need for numerical methods

most popular choice: gradient descent

decreasing J by iteratively moving in direction of negative gradient (steepest descent) with respect to input vector  $\hat{\theta}$ :

$$\widehat{\boldsymbol{\theta}} \leftarrow \widehat{\boldsymbol{\theta}} - \eta \nabla_{\widehat{\boldsymbol{\theta}}} J(\widehat{\boldsymbol{\theta}})$$
 step size (learning rate) vector containing all partial derivatives

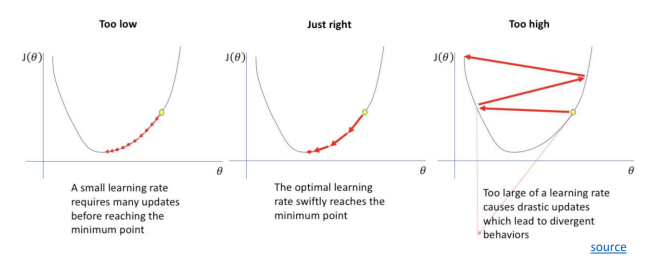




## Learning Rate

learning rate  $\eta$  corresponds to positive scalar value (hyperparameter)

- often set to small constant
- can be optimized via line search: evaluate  $J\left(\widehat{\boldsymbol{\theta}} \eta \nabla_{\widehat{\boldsymbol{\theta}}} J(\widehat{\boldsymbol{\theta}})\right)$  for several values of  $\eta$  and choose  $\eta$  resulting in smallest value of objective function
- can be varied from iteration to iteration (e.g., via some heuristic): smaller steps closer to the minimum (learning rate schedule/decay)



#### Batch and Stochastic Gradient Descent

in usual gradient descent: parameters  $\widehat{\theta}$  (and in turn objective function J) updated after full training epoch (one sweep through entire training data set)  $\rightarrow$  batch learning

consider: 
$$J(\widehat{\boldsymbol{\theta}}) = \frac{1}{n} \sum_{i=1}^{n} J_i(\widehat{\boldsymbol{\theta}})$$
 ( $J_i$  corresponds to loss function)

stochastic gradient descent updates after each training example (gradient of  $J(\widehat{\boldsymbol{\theta}})$  approximated with gradient of single loss):  $\widehat{\boldsymbol{\theta}} \leftarrow \widehat{\boldsymbol{\theta}} - \eta \nabla_{\widehat{\boldsymbol{\theta}}} J_i(\widehat{\boldsymbol{\theta}})$ 

- → online learning
- shuffling of samples after training epoch to prevent cycles
- adaptive learning rate to improve convergence

#### Mini Batches

Compared to its batch mode, stochastic gradient descent helps to avoid local minima (more robust) but is computationally expensive (less efficient).

→ mini-batch stochastic gradient descent as compromise:

splitting training data set into small batches used to calculate model error and update parameters

at the cost of one additional hyperparameter specifying mini-batch size

#### Gradient Descent with Momentum

goal: improve optimization process by avoiding gradients bouncing around the search space (dampening oscillations), accelerate in direction of minima

algorithm:

- estimate gradient g
- compute velocity update:  $\boldsymbol{v} \leftarrow \alpha \boldsymbol{v} \eta \boldsymbol{g}$
- apply parameter update:  $\widehat{m{ heta}} \leftarrow \widehat{m{ heta}} + m{ extit{v}}$

velocity: direction and speed at which the parameters move through parameter space, set to an exponentially decaying average of the past negative gradients (continues to move in its direction)

hyperparameter (0 <  $\alpha$  < 1) specifying exponential decay (like

learning rate  $\eta$ ,  $\alpha$  may be adapted over course of optimization)

## Bayesian Methods

### Bayesian Statistics

#### frequentist perspective:

- true model parameters  $\theta$  fixed (but unknown  $\rightarrow$  uncertain point estimates  $\widehat{\theta}$ )
- data set from random sampling

#### Bayesian perspective:

- true model parameters  $\theta$  random (estimation of full distributions over parameters)
- data not random (observed)

## Bayes Theorem

*H* : hypothesis (model)

*E* : evidence (data)

likelihood function (derived from statistical model)

$$P(H|E) = \frac{P(E|H) \cdot P(H)}{P(E)}$$

posterior probability (belief updated with observed evidence) new data (same for all hypotheses → uninteresting)

prior probability (belief in *H* before evidence)

## Bayes Theorem for Parameter Estimation

data likelihood:

model (function of  $\theta$  with data fixed), e.g., for linear regression:  $\mathcal{N}(y; \alpha + x \beta, \sigma^2)$ 

$$p(\boldsymbol{\theta}|y, \boldsymbol{x}) \propto p(y|\boldsymbol{x}, \boldsymbol{\theta}) \cdot p(\boldsymbol{\theta})$$

#### posterior:

probability distribution reflecting the effect of data on prior belief about parameters (increasing certainty with new evidence)

to be compared to frequentist point estimates  $\widehat{m{ heta}}$ 

prior (e.g., broad Gaussian): probability distribution reflecting initial belief about uncertain parameters (helps to reduce variance with few data)

## Bayesian Prediction

$$p\big(y_{n+1}|\boldsymbol{x}_{n+1},(y_1,\boldsymbol{x}_1),\cdots,(y_n,\boldsymbol{x}_n)\big) = \int p(y_{n+1}|\boldsymbol{x}_{n+1},\boldsymbol{\theta}) \, p\big(\boldsymbol{\theta}|(y_1,\boldsymbol{x}_1),\cdots,(y_n,\boldsymbol{x}_n)\big) \, d\boldsymbol{\theta}$$
 likelihood latest posterior

prediction of full probability distribution instead of frequentist point estimate(s) plugged into model distribution  $(p(y|x) = PDF(y; \hat{f}(x), ...))$ 

drawback: high computational costs due to integral over parameter distribution (especially for large training data sets)

#### Alternatives for PDF Predictions

#### quantile regression:

estimate quantile  $\tau$  of distribution instead of conditional mean by minimizing

$$(1-\tau)\sum_{y_i<\widehat{q}}(y_i-\widehat{q})+\tau\sum_{y_i\geq\widehat{q}}(y_i-\widehat{q})$$

instead of squared error loss (choice of loss function defines point estimate) possible with various ML methods, including neural networks and tree-based methods (like random forests or gradient boosting)

#### PDF assumption for model and estimation of its parameters:

assume, e.g., Gaussian or negative binomial  $\rightarrow$  estimate mean and variance (with different ML models or corresponding likelihood)

### Common Core of Statistical Learning

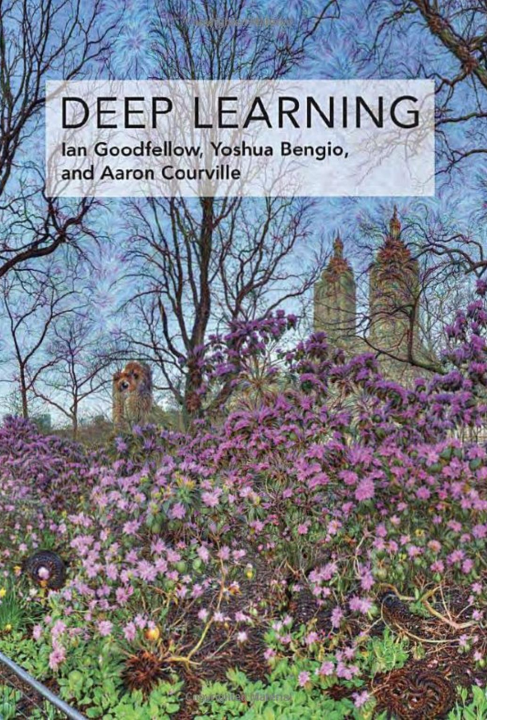
Most ML algorithms can be described by the general recipe of combining models, costs, and optimization methods.

#### including non-linear models:

- neural networks: backpropagation
- support-vector machines: minimizing hinge loss (soft-margin SVM)
- decision trees: minimizing impurity functions (mean squared error for regression, (kind of) Kullback-Leibler divergence for classification)

#### and even

- unsupervised learning: self-supervised, PCA by maximum variance directions
- reinforcement learning: express rewards in loss functions



#### Literature

not only deep learning, but also a nice ML introduction

https://www.deeplearningbook.org/

#### Overcome our Mathematical Limitations

evolution provided us with moderate math skills

#### AI/ML to the rescue:

- recognition of mathematical structures and patterns
- algorithm discovery: <u>AlphaTensor</u>
- symbolic regression (<u>e.g., for physics</u>)