Learning via Probabilistic Modeling

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Introduction to Machine Learning (CS771A)

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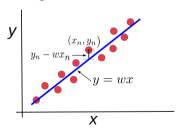


Recap: Linear Models and Linear Regression

• Linear model: Each output is a linearly weighted combination of the inputs

$$y_n = \sum_{d=1}^{D} w_d x_{nd} = \boldsymbol{w}^{\top} \boldsymbol{x}_n, \forall n \quad \Rightarrow \quad \boldsymbol{y} = \boldsymbol{X} \boldsymbol{w}$$

- The weights are the parameters of the model
- Can use linear models for doing linear regression. Amounts to fitting a line/plane to the data.



- Finding the best line/plane = finding w that minimizes the total error/loss of the fit
- This requires optimizing the loss w.r.t. w



Recap: Least Squares and Ridge Regression

- Least squares and ridge regression are both linear regression models based on squared loss
- Least squares regression minimizes the simple sum of squared errors

$$\hat{\boldsymbol{w}} = \arg\min_{\boldsymbol{w}} \sum_{n=1}^{N} (y_n - \boldsymbol{w}^{\top} \boldsymbol{x}_n)^2 = (\sum_{n=1}^{N} \boldsymbol{x}_n \boldsymbol{x}_n^{\top})^{-1} \sum_{n=1}^{N} y_n \boldsymbol{x}_n = (\boldsymbol{\mathsf{X}}^{\top} \boldsymbol{\mathsf{X}})^{-1} \boldsymbol{\mathsf{X}}^{\top} \boldsymbol{y}$$

• Ridge regression minimizes the ℓ_2 regularized sum of squared errors

$$\hat{\boldsymbol{w}} = \arg\min_{\boldsymbol{w}} \left[\sum_{n=1}^{N} (y_n - \boldsymbol{w}^{\top} \boldsymbol{x}_n)^2 + \underbrace{\frac{\lambda}{2} \boldsymbol{w}^{\top} \boldsymbol{w}}_{\text{regularizer}} \right] = (\mathbf{X}^{\top} \mathbf{X} + \lambda \mathbf{I}_D)^{-1} \mathbf{X}^{\top} \boldsymbol{y}$$

- Regularization helps prevent overfitting the training data
- The ℓ_2 regularization $\mathbf{w}^{\top}\mathbf{w} = \sum_{d=1}^{D} w_d^2$ promotes small individual weights



Recap: Learning as Optimization

- Supervised learning is essentially a function approximation problem
- Given training data $\{(\boldsymbol{x}_n, y_n)_{n=1}^N\}$, find a function f s.t. $f(\boldsymbol{x}_n) \approx y_n, \forall n$
- In addition, we want f to be simple (i.e., want to regularize it)
- Can learn such a function f by solving the following optimization problem

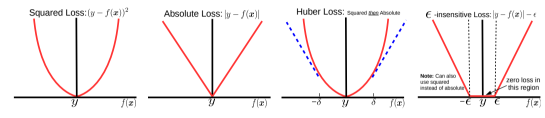
$$\hat{f} = \arg\min_{f} \mathcal{L}_{reg}(f) = \arg\min_{f} \underbrace{\sum_{n=1}^{N} \ell(y_n, f(\mathbf{x}_n))}_{\text{training loss}} + \underbrace{\lambda R(f)}_{\text{regularization}}$$

- Different supervised learning problems differ in the choice of f, $\ell(.,.)$ and R(.)
 - f depends on the model (e.g., for linear models $f(x) = w^{\top}x$)
 - $\ell(.,.)$: loss function that measures the error of model's prediction (e.g., squared loss)
 - R(.) denotes the regularizer chosen to make f simple (e.g., ℓ_2 regularization)



A Brief Detour: Some Loss Functions

• Some popular loss functions for regression problems¹



- Absolute/Huber loss preferred if there are outliers in the data
 - Less affected by large errors |y f(x)| as compared to the squared loss
- Overall objective function = loss func + some regularizer (e.g., ℓ_2 , ℓ_1), as we saw for ridge reg.
- Some objectives easy to optimize (convex and differentiable), some not so (e.g., non-differentiable)
- Will revisit many of these aspects when we talk about optimization techniques for ML

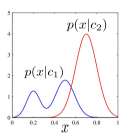
¹will look at loss functions for classification later when discussing classification in detail

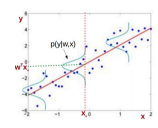
Brief Detour: Inductive Bias of ML Algorithms

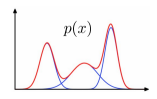
- No ML algorithm is "universally good"
- Should not expect it to work well on all datasets
- Each algorithm makes some assumption about data ("no free lunch")
 - Work best when assumptions are correct. May fail in other case.
- Inductive Bias: Set of assumptions made about outputs of previously unseen inputs
- Learning is impossible without making assumptions!
- Some common examples of such assumptions
 - Classes are separable by a large margin
 - The function is "smooth"
 - Only a few features are relevant for the prediction



Learning via Probabilistic Modeling







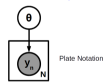


Probabilistic Modeling of Data

• Assume the data $\mathbf{y} = \{y_1, y_2, \dots, y_N\}$ as generated from a probability model

$$y_n \sim p(y|\theta) \quad \forall n$$

- Each y_n assumed drawn from distribution $p(y|\theta)$, with unknown parameters θ
- We usually assume data to be independently & identically distributed (i.i.d.)



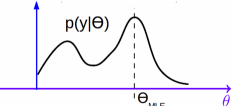
- Some of the things we may be interested in
 - ullet Parameter Estimation: Estimate heta given the observed data $oldsymbol{y}$
 - Prediction: Compute predictive distribution $p(y_*|y)$ for new data (or mean/variance of $p(y_*|y)$)
- Important: Pretty much any ML problem (sup/unsup) can be formulated like this

Parameter Estimation in Probabilistic Models

• Since data is i.i.d., the probability (or probability density) of observed data $\mathbf{y} = \{y_1, y_2, \dots, y_N\}$

$$p(\mathbf{y}|\theta) = p(y_1, y_2, \dots, y_N|\theta) = \prod_{n=1}^N p(y_n|\theta)$$

- $p(y|\theta)$ also called the model's likelihood, $p(y_n|\theta)$ is likelihood w.r.t. a single data point
- ullet The likelihood will be a function of the parameters heta

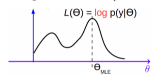


- How do we estimate the "best" model parameters θ ?
- One option: Find value of θ that makes observed data most probable (i.e., most *likely*)
 - Maximize the likelihood $p(y|\theta)$ w.r.t. θ : Maximum Likelihood Estimation (MLE)



Maximum Likelihood Estimation (MLE)

• We doing MLE, we typically maximize log-likelihood instead of the likelihood, which is easier (doesn't affect the estimation because log is monotonic)



Log-likelihood:

$$\mathcal{L}(\theta) = \log p(\mathbf{y} \mid \theta) = \log \prod_{n=1}^{N} p(y_n \mid \theta) = \sum_{n=1}^{N} \log p(y_n \mid \theta)$$

• Maximum Likelihood Estimation (MLE)

$$egin{aligned} \hat{ heta}_{MLE} = rg \max_{ heta} \mathcal{L}(heta) = rg \max_{ heta} \sum_{n=1}^{N} \log p(y_n \mid heta) \end{aligned}$$

Now this becomes an optimization problem w.r.t. θ



Maximum Likelihood Estimation (MLE)

• Maximum Likelihood parameter estimation

$$\hat{\theta}_{MLE} = \arg\max_{\theta} \sum_{n=1}^{N} \log p(y_n \mid \theta) = \arg\min_{\theta} - \sum_{n=1}^{N} \log p(y_n \mid \theta)$$

• We thus also think of it as minimizing the negative log-likelihood (NLL)

$$\hat{ heta}_{ extit{MLE}} = rg\min_{ heta} extit{NLL}(heta)$$

where
$$NLL(\theta) = -\sum_{n=1}^{N} \log p(y_n \mid \theta)$$

- We can think of the negative log-likelihood as a loss function
- Thus MLE is equivalent to doing empirical risk (training data loss) minimization
- Important: This view relates/unifies the optimization and probabilistic modeling approaches
- Something is still missing (we will look at that shortly)

MLE: An Example

- Consider a sequence of *N* coin toss outcomes (observations)
- Each observation y_n is a binary random variable. Head = 1, Tail = 0
- Since each y_n is binary, let's use a **Bernoulli distribution** to model it

$$p(y_n \mid \theta) = \theta^{y_n} (1 - \theta)^{1 - y_n}$$

- ullet Here heta is the unknown parameter (probability of head). Want to learn heta using MLE
- Log-likelihood: $\sum_{n=1}^{N} \log p(y_n \mid \theta) = \sum_{n=1}^{N} y_n \log \theta + (1 y_n) \log(1 \theta)$
- ullet Taking derivative of the log-likelihood w.r.t. heta, and setting it to zero gives

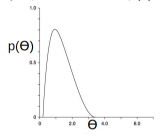
$$\hat{\theta}_{MLE} = \frac{\sum_{n=1}^{N} y_n}{N}$$

- $\hat{\theta}_{MLE}$ in this example is simply the fraction of heads!
- What can go wrong with this approach (or MLE in general)?
 - We haven't "regularized" θ . Can do badly (i.e., overfit), e.g., if we don't have enough data



Prior Distributions

• In probabilistic models, we can specify a prior distribution $p(\theta)$ on parameters



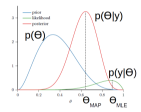
- The prior distribution expresses our a priori belief about the unknown θ . Plays two key roles
 - ullet The prior helps us specify that some values of heta are more likely than others
 - ullet The prior also works as a regularizer for heta (we will see this soon)
- Note: A uniform prior distribution is the same as using no prior!



Using a Prior in Parameter Estimation

• We can **combine** the prior $p(\theta)$ with the likelihood $p(y|\theta)$ using Bayes rule and define the posterior distribution over the parameters θ

$$p(heta|oldsymbol{y}) = rac{p(oldsymbol{y}| heta)p(heta)}{p(oldsymbol{y})}$$



• Now, instead of doing MLE which maximizes the likelihood, we can find the θ that is most likely given the data, i.e., which maximizes the posterior probability $p(\theta|\mathbf{y})$

$$\hat{\theta}_{\textit{MAP}} = \arg\max_{\boldsymbol{\theta}} p(\boldsymbol{\theta}|\boldsymbol{y})$$

• Note that the prior sort of "pulls" θ_{MLE} toward's the prior distribution's mean/mode



Maximum-a-Posteriori (MAP) Estimation

• We will work with the **log** posterior probability (it is easier)

$$\begin{split} \hat{\theta}_{MAP} &= \arg\max_{\theta} p(\theta|\mathbf{y}) &= \arg\max_{\theta} \log p(\theta|\mathbf{y}) \\ &= \arg\max_{\theta} \log \frac{p(\mathbf{y}|\theta)p(\theta)}{p(\mathbf{y})} \\ &= \arg\max_{\theta} \log p(\mathbf{y}|\theta) + \log p(\theta) \\ \\ &\hat{\theta}_{MAP} &= \arg\max_{\theta} \sum_{n=1}^{N} \log p(y_n|\theta) + \log p(\theta) \end{split}$$

- Same as MLE with an extra log-prior-distribution term (acts as a regularizer)
- Can also write the same as the following (equivalent) minimization problem

$$\hat{\theta}_{MAP} = \arg\min_{\theta} NLL(\theta) - \log p(\theta)$$

• When $p(\theta)$ is a uniform prior, MAP reduces to MLE

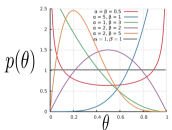


MAP: An Example

- Let's again consider the coin-toss problem (estimating the bias of the coin)
- Each likelihood term is Bernoulli: $p(y_n|\theta) = \theta^{y_n}(1-\theta)^{1-y_n}$
- Since $\theta \in (0,1)$, we assume a Beta prior: $\theta \sim \text{Beta}(\alpha,\beta)$

$$\rho(\theta) = \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha)\Gamma(\beta)} \theta^{\alpha - 1} (1 - \theta)^{\beta - 1}$$

• Note: Γ is the gamma function. α, β are called hyperparameters of the prior



For Beta, using $\alpha = \beta = 1$ corresponds to using a uniform prior distribution



MAP: An Example

• The log posterior probability for the coin-toss model

$$\sum_{n=1}^{N} \log p(y_n|\theta) + \log p(\theta)$$

• Ignoring the constants w.r.t. θ , the log posterior probability simplifies to

$$\sum_{n=1}^{N} \{y_n \log \theta + (1-y_n) \log (1-\theta)\} + (\alpha-1) \log \theta + (\beta-1) \log (1-\theta)$$

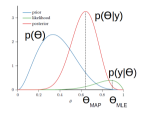
ullet Taking derivative w.r.t. heta and setting to zero gives

$$\hat{\theta}_{MAP} = \frac{\sum_{n=1}^{N} y_n + \alpha - 1}{N + \alpha + \beta - 2}$$

- **Note:** For $\alpha = 1, \beta = 1$, i.e., $p(\theta) = \text{Beta}(1,1)$ (which is equivalent to a uniform prior, hence no regularization). Thus, for $\alpha = 1, \beta = 1$, we get the same solution as $\hat{\theta}_{MLE}$
- Note: Hyperparameters of a prior distribution usually have intuitive meaning. E.g., in the coin-toss example, $\alpha-1$, $\beta-1$ are like "pseudo-observations" expected numbers of heads and tails, respectively, before tossing the coin

Inferring the Full Posterior (a.k.a. Fully Bayesian Inference)

• MLE/MAP only give us a **point estimate** of θ . Doesn't capture the uncertainty in θ



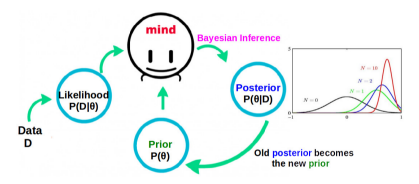
• The Bayes rule (at least in theory) also allows us to compute the full posterior

$$p(\theta|\mathbf{y}) = \frac{p(\mathbf{y}|\theta)p(\theta)}{p(\mathbf{y})} = \frac{p(\mathbf{y}|\theta)p(\theta)}{\int p(\mathbf{y}|\theta)p(\theta)d\theta}$$

- In general, much harder problem than MLE/MAP! Easy if the prior and likelihood are "conjugate" to each other (then the posterior will then have the same "form" as the prior)
- Many pairs of distributions are conjugate to each other (e.g., Beta-Bernoulli, Gaussian is conjugate to itself, etc.). May refer to Wikipedia for a list of conjugate pairs of distributions

Fully Bayesian Inference

• Fully Bayesian inference fits naturally into an "online" learning setting



ullet Our belief about heta keeps getting updated as we see more and more data



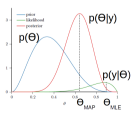
Fully Bayesian Inference: An Example

- Let's again consider the coin-toss example
- With Bernoulli likelihood and Beta prior (a conjugate pair), the posterior is also Beta (exercise)

$$\mathsf{Beta}(\alpha + \mathsf{N}_1, \beta + \mathsf{N}_0)$$

where N_1 is the number of heads and $N_0 = N - N_1$ is the number of tails

• Can verify the above by simply plugging in the expressions of likelihood and prior into the Bayes rule and identifying the form of resulting posterior (note: this may not always be easy)





Making Predictions: MLE/MAP/Fully Bayesian

- \bullet Once θ is learned, we can use it to make predictions about the future observations
- E.g., for the coin-toss example, we can predict the probability of next toss being head
- This can be done using the MLE/MAP estimate, or using the full posterior (harder)
- In the coin-toss example, $\theta_{MLE} = \frac{N_1}{N}$, $\theta_{MAP} = \frac{N_1 + \alpha 1}{N + \alpha + \beta 2}$, and $p(\theta|\mathbf{y}) = \text{Beta}(\theta|\alpha + N_1, \beta + N_0)$
- Thus for this example (where observations are assumed to come from a Bernoulli)

MLE prediction:
$$p(y_{N+1} = 1|\mathbf{y}) = \int p(y_{N+1} = 1|\theta)p(\theta|\mathbf{y})d\theta \approx p(y_{N+1} = 1|\theta_{MLE}) = \theta_{MLE} = \frac{N_1}{N}$$

MAP prediction: $p(y_{N+1} = 1|\mathbf{y}) = \int p(y_{N+1} = 1|\theta)p(\theta|\mathbf{y})d\theta \approx p(y_{N+1} = 1|\theta_{MAP}) = \theta_{MAP} = \frac{N_1 + \alpha - 1}{N + \alpha + \beta - 2}$

Fully Bayesian: $p(y_{N+1} = 1|\mathbf{y}) = \int p(y_{N+1} = 1|\theta)p(\theta|\mathbf{y})d\theta = \int \theta p(\theta|\mathbf{y})d\theta = \int \theta \operatorname{Beta}(\theta|\alpha + N_1, \beta + N_0)d\theta = \frac{N_1 + \alpha}{N + \alpha + \beta}$

• Note that the fully Bayesian approach to prediction averages over all possible values of θ , weighted by their respective posterior probabilities (easy in this example, but a hard problem in general)

Probabilistic Modeling: Summary

- A flexible way to model data by specifying a proper probabilistic model
- Likelihood corresponds to a loss function; prior corresponds to a regularizer
- Can choose likelihoods and priors based on the nature/property of data/parameters
- MLE estimation = unregularized loss function minimization
- MAP estimation = regularized loss function minimization
- Allows us to do fully Bayesian learning
 - Allows learning the full distribution of the parameters (note that MLE/MAP only give a "single best" answer as a point estimate of the parameters)
 - Makes more robust predictions by posterior averaging (rather than using a single point estimate)
 - Many other benefits, such as
 - Estimate of confidence in the model's prediction (useful for doing Active Learning)
 - Can do automatic model selection, hyperparameter estimation, handle missing data, etc.
 - .. and many other benefits (a proper treatment deserves a separate course :))
- MLE/MAP estimation is also related to the optimization view of ML

