

Bayesian Methods

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1.

1.1.

The log likelihood function of the samples $\{k_1, k_2, \dots, k_n\}$ is

$$L(\{k_i\}|\lambda) = \ln\left(\prod_{i=1}^n \frac{\lambda^{k_i} \exp(-\lambda)}{k_i!}\right) = \sum_{i=1}^n (k_i \ln(\lambda) - \lambda - \ln(k_i!))$$

Calculate the derivative with respect to λ , we get:

$$\frac{\partial L}{\partial \lambda} = \sum_{i=1}^n \frac{k_i}{\lambda} - 1 = \frac{\sum_{i=1}^n k_i}{\lambda} - n$$

And we can see that the second derivative of the log likelihood function is always negative as $\lambda > 0$, so this function is a concave function, and the maximum of this function is achieved at its unique extreme point, so at the extreme point we have:

$$\frac{\sum_{i=1}^n k_i}{\lambda} - n = 0 \rightsquigarrow \lambda_{mle} = \frac{\sum_{i=1}^n k_i}{n}$$

1.2.

$$\begin{aligned} E[K] &= \sum_{k=0}^{\infty} \frac{\lambda^k \exp(-\lambda)}{k!} k = \sum_{k=1}^{\infty} \lambda \frac{\lambda^{k-1} \exp(-\lambda)}{(k-1)!} = \lambda \sum_{k'=0}^{\infty} \frac{\lambda^{k'} \exp(-\lambda)}{k'!} = \lambda \\ \text{Var}[K] &= E[K^2] - E[K]^2 = \sum_{k=0}^{\infty} \frac{\lambda^k \exp(-\lambda)}{k!} k((k-1) + 1) - \lambda^2 \\ &= \sum_{k=0}^{\infty} \frac{\lambda^k \exp(-\lambda)}{k!} k(k-1) + E[K] - E[K]^2 \\ &= \lambda^2 \sum_{k'=0}^{\infty} \frac{\lambda^{k'} \exp(-\lambda)}{k'!} + E[K] - E[K]^2 \\ &= \lambda^2 + \lambda - \lambda^2 = \lambda \end{aligned}$$

The expectation and variance of λ_{mle} is:

$$E[\lambda_{mle}] = \frac{1}{n} \sum_{i=1}^n E(k_i) = E(k_i) = \lambda$$

$$Var[\lambda_{mle}] = E(\lambda_{mle}^2) - E(\lambda_{mle})^2 = \frac{nE(K^2) + (n^2 - n)E(K)^2}{n^2} - \lambda^2 = \frac{\lambda}{n}$$

From the two results above, we can see the λ_{mle} is the unbiased estimation of λ . And as the number of samples increase, the variance of the estimation decreases, the estimation is more accurate, so this estimation is a effective estimation of λ .

1.3.

The posterior probability of λ is:

$$P(\lambda|\{k_i\}) = \frac{L(\{k_i\}|\lambda)p(\lambda)}{P(\{k_i\})}$$

In which $P(\{k_i\})$ is the evidence of the observed samples $\{k_i\}$ intergrated on all the possible value of the parameter λ , the parameter λ is marginalized out, so it is unrelated to λ . We take the logarithm of of the above equation, get:

$$\begin{aligned} \log(P(\lambda|\{k_i\})) &= \sum_{i=1}^N (k_i \ln(\lambda) - \lambda - \ln(k_i!)) + \ln(p(\lambda)) - \ln(P(\{k_i\})) \\ &= \sum_{i=1}^N (k_i \ln(\lambda) - \lambda - \ln(k_i!)) + (a-1) \ln(\lambda) - \frac{\lambda}{\beta} - \ln(\Gamma(a)) - a \ln(\beta) \end{aligned}$$

Take the derivative and make it equal to 0:

$$\begin{aligned} \frac{\partial P(\lambda|\{k_i\})}{\lambda} &= \frac{\sum_{i=1}^N + a - 1}{\lambda} - (1 + \frac{1}{\beta}) = 0 \\ \rightsquigarrow \lambda_{map} &= \frac{a - 1 + \sum_{i=1}^n k_i}{n + \frac{1}{\beta}} \end{aligned}$$

1.4.

- When $n \rightarrow 0$, $\lambda_{map} \rightarrow (a-1)\beta$, this is the mode value of the prior gamma distribution (the value with the maximum prior).
- When $n \rightarrow \infty$, $\lambda_{map} \rightarrow \frac{\sum_{i=1}^n k_i}{n}$, this is the maximum-likelihood estimation of λ : λ_{mle} .

We can see from the two limiting case: when there are few samples, the prior distribution contributes much to the MAP estimation; while as the number of samples increase, the effect of the prior distribution fades out, and the actual samples contributes more to the MAP estimation, which will drive the MAP estimation to λ_{mle} .

Also, we can notice that the gamma distribution $\text{Gamma}(a, \beta)$ is the *conjugate prior* of the poison likelihood function: the posterior distribution of λ is of the same distribution family as the prior (both Gamma distribution). So the sample-adding process can be regard as an progressively parameter adjusting process of the gamma distribution (adding a new sample just adjust a and β a little). This perspective will make the MAP estimation easy to calculate progressively (at every step, the last posterior distribution can be regard as a new prior), and unrelated to the exact number of samples n that have been observed.

2.

2.1.

The log likelihood function is:

$$\begin{aligned}
L(X) &= P(\{x_1, \dots, x_n\} | \mu, \Lambda^{-1}) = \prod_{i=1}^n P(x_i | \mu, \Lambda) \\
&= \prod_{i=1}^n \frac{1}{\sqrt{(2\pi)^p |\Lambda^{-1}|}} \exp\left(-\frac{1}{2}(x_i - \mu)^T \Lambda (x_i - \mu)\right) \\
&= \frac{1}{(2\pi)^{\frac{pn}{2}}} \Lambda^{\frac{n}{2}} \exp\left(-\frac{1}{2} \sum_{i=1}^n (x_i - \mu)^T \Lambda (x_i - \mu)\right) \\
\log L(X) &= \log(L(X)) = \frac{n}{2} \log(|\Lambda|) - \frac{pn}{2} \log(2\pi) - \frac{1}{2} \sum_{i=1}^n (x_i - \mu)^T \Lambda (x_i - \mu)
\end{aligned}$$

To do the MLE estimation of μ , we take the partial derivative:

$$\frac{\partial \log L(X)}{\partial \mu} = \sum_{i=1}^n \Lambda (x_i - \mu) = \Lambda \sum_{i=1}^n (x_i - \mu) = 0$$

As Λ^{-1} is non-singular, its null space is empty set. So, we must have:

$$\sum_{i=1}^n (x_i - \mu) = 0 \rightsquigarrow \mu_{mle} = \frac{1}{n} \sum_{i=1}^n x_i$$

To do the MLE estimation of Λ , we take the partial derivative:

$$\frac{\partial \log L(X)}{\partial \Lambda} = \frac{n}{2} \Lambda^{-T} - \frac{1}{2} \sum_{i=1}^n (x_i - \mu)(x_i - \mu)^T = 0$$

As Λ is non-singular and symmetric, we get:

$$\Lambda_{mle} = \left(\frac{1}{n} \sum_{i=1}^n (x_i - \mu_{mle})(x_i - \mu_{mle})^T \right)^{-1}$$

2.2.

The log posterior function is:

$$\begin{aligned}
&\log(L(X)) + \log(gw(\mu, \Lambda)) - \log(P(X)) \\
&= C + \frac{v-p+n}{2} \ln(|\Lambda|) - \frac{1}{2} \sum_{i=1}^n (x_i - \mu)^T \Lambda (x_i - \mu) - \frac{1}{2} s(\mu - \mu_0)^T \Lambda (\mu - \mu_0) - \frac{\text{tr}(V^{-1} \Lambda)}{2}
\end{aligned}$$

C is a constant that do not influence the derivative. The MAP estimation of μ satisfy:

$$\frac{\partial \text{posterior}}{\partial \mu} = \sum_{i=1}^n \Lambda (x_i - \mu) - s \Lambda (\mu - \mu_0) = \Lambda \left(\sum_{i=1}^n x_i + s \mu_0 - (n+s) \mu \right) = 0$$

As Λ is non-singular, we have $\mu_{map} = \frac{\sum_{i=1}^n x_i + s \mu_0}{n+s}$. The MAP estimation of Λ satisfy:

$$\frac{\partial \text{posterior}}{\partial \Lambda} = \frac{v-p+n}{2} \Lambda^{-T} - \frac{1}{2} \sum_{i=1}^n (x_i - \mu)(x_i - \mu)^T - \frac{1}{2} s(\mu - \mu_0)(\mu - \mu_0)^T - \frac{V^{-T}}{2} = 0$$

Substitute in μ_{map} , we get:

$$\Lambda_{map} = \left(\frac{\sum_{i=1}^n (x_i - \mu_{map})(x_i - \mu_{map})^T + s(\mu_{map} - \mu_0)(\mu_{map} - \mu_0)^T + V^{-T}}{n+v-p} \right)^{-1}$$

2.3.

- When $n \rightarrow 0$, $\mu_{map} \rightarrow \mu_0$; $\Lambda_{map} \rightarrow \frac{V}{v-p}$.
- When $n \rightarrow \infty$, $n \gg s$, $\mu_{map} \rightarrow \frac{\sum_{i=1}^n x_i}{n} = \mu_{mle}$; substitute μ_{map} into Λ_{map} , also use $n \gg s$, we have $\Lambda_{map} \rightarrow \Lambda_{mle}$.

Very intuitively, when there are few samples, the prior distribution of μ and Λ contributes much to the posterior distribution, so when $n = 0$, the MAP estimation is just the mode value of the prior distribution of the parameters. However, as n increases, the samples $\{x_i\}$ contribute more and more to the posterior distribution, and when $n \rightarrow \infty$, the influence of the prior distribution fades out, so the MAP estimation when $n \rightarrow \infty$ will approach the MLE estimation where parameter prior is not used.

Also, the Gaussian-Wishart prior is the conjugate prior distribution of the Gaussian likelihood function, so we can see the sample-adding process as a progressively adjustment process of the hyper-parameters V, v, s, μ_0 to get the posterior distribution. This property leads to quicker and simpler posterior calculation and hence MAP estimation.

3.

3.1.

Suppose we have a failure model of a certain type of device, we use exponential distribution to model the failure process of this type of device ($F_{\text{failure}}(t) = P(\text{failure_time} \leq t) = 1 - \exp(-\lambda t)$), in which the failure rate of this type of device remains unchanged during the lifetime of the device:

$$\lambda(t) = \frac{f(t)}{1 - F(t)} = \lambda$$

Suppose we observe the behavior of N such devices for a period: T . We assume in this time period, there are n device failure observed, we assume the n devices failed at t_1, t_2, \dots, t_n . We can write the likelihood function of these observation as:

$$\log L = \log\left(\prod_{i=1}^n \lambda \exp(-\lambda t_i) (1 - F(T))^{N-n}\right) = n \ln(\lambda) - \lambda \sum_{i=1}^n t_i - (N - n)\lambda T$$

When $n = 0$, the log likelihood function is:

$$\log L(\lambda) = -N\lambda T$$

The maximum is achieved at $\lambda = 0$, however, $\lambda = 0$ does not correspond to a legal exponential distribution, because every exact point have a probability of 0. So, in this situation, the MLE estimation do not exist.

3.2.

Consider a mixture of two fixed-mean fixed-variance gaussian distribution with the same Σ , the parameter $\theta, 0 \leq \theta \leq 1$ is the combination factor of the two gaussian probabilistic function:

$$p(x|\theta) = \theta \frac{1}{(2\pi)^{n/2} |\Sigma^{-1}|} \exp\left(-\frac{(x - \mu_1)^T \Sigma^{-1} (x - \mu_1)}{2}\right) + (1 - \theta) \frac{1}{(2\pi)^{n/2} |\Sigma^{-1}|} \exp\left(-\frac{(x - \mu_2)^T \Sigma^{-1} (x - \mu_2)}{2}\right)$$

When the observed sample x_s is at the middle point of μ_1 and μ_2 , that is $x_s = \frac{\mu_1 + \mu_2}{2}$, the MLE of the parameter θ is not unique, because θ can be every value in $[0, 1]$, and the likelihood of this special sample won't change.

3.3.

To guarantee the MLE of parameter always exists and its uniqueness, the log likelihood function should have the only maximum point no matter what samples are observed.

4.

4.1.

The log likelihood of the observation $T = \{t_i\}$ is:

$$\log(p(T|X, \omega, \beta)) = -N \ln\left(\frac{2\pi}{\sqrt{\beta}}\right) - \frac{\beta}{2} \sum_{i=1}^N (t - y(x, \omega))^2$$

In maximum likelihood estimation, we maximize the log likelihood function, which is to minimize $\sum_{i=1}^N (t - y(x, \omega))^2$ according to the equation above. So this is equivalent to minimizing the sum of square error.

4.2.

$$\log(p(\omega)) \propto \log(p(T|X, \omega, \beta)) + \log(p(\omega|\alpha)) = C - \frac{\beta}{2} \sum_{i=1}^N (t - y(x, \omega))^2 - \alpha \frac{\omega^T \omega}{2}$$

Doing the MAP estimation of the posterior distribution of ω , is equivalent to minimize $\frac{\beta}{2} \sum_{i=1}^N (t - y(x, \omega))^2 + \alpha \frac{\|\omega\|^2}{2}$, this is the form of the square error loss with a weight decay regularizing term (when $\alpha = \lambda\beta$).

Naive Bayes

1.

When $Y = T$, we need a parameter $P(X_1 = T|Y = T)$ for X_1 , and two parameter μ_i, σ_i for every other continuous variable X_i . And another set of parameters are needed when $Y = F$. So we need $2(1 + 2(n - 1)) = 2(2n - 1)$ parameters to build the Naive Bayes classifier.

$$P(Y = y|X) = \frac{P(X|Y)P(Y)}{P(X)} = \frac{P(X_1|Y = y) \prod_{i=2}^n P(X_i|Y)P(Y)}{P(X_1|Y = T) \prod_{i=2}^n P(X_i|Y = T) + P(X_1|Y = F) \prod_{i=2}^n P(X_i|Y = F)}$$

In the above equation, $P(Y)$ prior is estimated using the occurring times of $Y = T$ and $Y = F$ in the training data.

2.

2.1.

We use S for the abbreviation of the random variable *Sunny*, W for *Windy* and H for *Hike*. The posterior distribution of H is:

$$P(H|S, W) = \frac{P(S, W|H)P(H)}{P(S, W, H)} = \frac{P(S|H)P(W|H)P(H)}{P(S|H = T)P(W|H = T)P(H = T) + P(S|H = F)P(W|H = F)P(H = F)}$$

The decision rule is:

$$\begin{cases} \hat{H} = T : P(H = T|S, W) > P(H = F|S, W) \rightsquigarrow P(S|H = T)P(W|H = T) > P(S|H = F)P(W|H = F) \\ \hat{H} = F : \text{otherwise} \end{cases}$$

The four cases is listed below:

- $S = T, W = T$, $0.27 = P(S = T|H = T)P(W = T|H = T) < P(S = T|H = F)P(W = T|H = F) = 0.32$, so the decision is $\hat{H} = F$.
- $S = T, W = F$, $0.63 = P(S = T|H = T)P(W = F|H = T) > P(S = T|H = F)P(W = F|H = F) = 0.08$, so the decision is $\hat{H} = T$.
- $S = F, W = T$, $0.03 = P(S = F|H = T)P(W = T|H = T) < P(S = F|H = F)P(W = T|H = F) = 0.48$, so the decision is $\hat{H} = F$.
- $S = F, W = F$, $0.07 = P(S = F|H = T)P(W = F|H = T) > P(S = F|H = F)P(W = F|H = F) = 0.12$, so the decision is $\hat{H} = F$.

2.2.

$$\begin{aligned}
 P(\text{error}) &= \sum_{s,w \text{ s.t. } \hat{H}=T} P(H = F|S = s, W = w)P(S = s, W = w) \\
 &\quad + \sum_{s,w \text{ s.t. } \hat{H}=F} P(H = T|S = s, W = w)P(S = s, W = w) \\
 &= \sum_{s,w \text{ s.t. } \hat{H}=T} P(S = s, W = w|H = F)P(H = F) \\
 &\quad + \sum_{s,w \text{ s.t. } \hat{H}=F} P(S = s, W = w|H = T)P(H = T) \\
 &= 0.08 * 0.5 + (0.27 + 0.03 + 0.07) * 0.5 = 0.225
 \end{aligned}$$

2.3.

The joint probability is:

$$P(S = T, W = T, H = T) = P(H = T)p(S = T|H = T)P(W = T|H = T) = 0.135$$

2.4.

The naive bayes assumption is violated, because the Sunny random variable is dependent to Rainy variable even if the decision is given. The joint probability remains unchange here:

$$P(S = T, W = T, H = T) = P(H = T)p(S = T|H = T)P(W = T|H = T) = 0.135$$

2.5.

The error rate remains unchanged as 0.225, and the performance of Naive Bayes does not improve by observing the new attribute Rainy, because $\text{Rainy} = \neg \text{Sunny}$, and it brought no additional information.

Programming

In this problem you will implement Naive Bayes and Logistic Regression, then compare their performance on a document classification task. The data for this task is taken from the 20 Newsgroups data set, and is available from the attached zip file. The included README.txt describes the data set and file format.

Our Naive Bayes model will use the bag-of-words assumption. This model assumes that each word in a document is drawn independently from a multinomial distribution over possible words. (A multinomial distribution is a generalization of a Bernoulli distribution to multiple values.) Although this model ignores the ordering of words in a document, it works surprisingly well for a number of tasks. We number the words in our vocabulary from 1 to m , where m is the total number of distinct words in all of the documents. Documents from class y are drawn from a class-specific multinomial distribution parameterized by θ_y . θ_y is a vector, where $\theta_{y,i}$ is the probability of drawing word i and $\sum_{i=1}^m \theta_{y,i} = 1$. Therefore, the class-conditional probability of drawing document x from our Naive Bayes model is $P(X = x|Y = y) = \prod_{i=1}^m (\theta_{y,i})^{count_i(x)}$, where $count_i(x)$ is the number of times word i appears in x .

1. Provide high-level descriptions of the Naive Bayes and Logistic Regression algorithms. Be sure to describe how to estimate the model parameters and how to classify a new example.

Naive Bayes Classifier For Naive Bayes Classifier, we learn the generative model of the documents by learning $P(Y)$ and $P(X|Y)$. The likelihood function $P(X|Y)$ can be factorized as $P(X = x|Y = y) = \prod_{i=1}^m (\theta_{y,i})^{count_i(x)}$ from the bag-of-word assumption; The prior distribution of document labels Y can be easily modelled using the normalized count of the occurrence of $Y = y$ in the training data.

Assume there are M_y samples for every document label y :

$$P(Y = y) = \frac{M_y}{\sum_i M_i}$$

In the learning process, we want to obtain the MLE estimation of the parameter $\{\theta_y, i = 1, \dots, m\}$ set for every document label y . Assume there are N samples that are labeled y in the training data, we can write the log likelihood function:

$$\ln\left(\prod_{n=1}^N P(X|Y = y)\right) = \sum_{n=1}^N \sum_{i=1}^m count_i(x^{(n)}) \ln(\theta_{y,i})$$

Also, there is a constraint for $\theta_{y,i}$ that $\sum_{i=1}^m \theta_{y,i} = 1$. We can obtain the Lagrangian of this problem:

$$\sum_{n=1}^N \left(\sum_{i=1}^m count_i(x^{(n)}) \ln(\theta_{y,i}) \right) - a \left(\sum_{i=1}^m \theta_{y,i} - 1 \right)$$

Take the derivative with respect to $\theta_{y,i}$, we have:

$$\begin{aligned} \frac{\sum_{n=1}^N count_i(x^{(n)})}{\theta_{y,i}} - a &= 0 \\ \rightsquigarrow \theta_{y,i} &\propto \sum_{n=1}^N count_i(x^{(n)}) \\ \rightsquigarrow \theta_{y,i} &= \frac{\sum_{n=1}^N count_i(x^{(n)})}{\sum_{i=1}^m \sum_{n=1}^N count_i(x^{(n)})} \end{aligned}$$

From the above estimation equation, we can see that we just need to count the occurrence of words i in samples grouped by the label y , after which the parameters can be obtained by doing some normalization.

In the test phase, we get a test sample x , we should calculate $P(X = x|Y = y)P(Y = y)$ for every y (The evidence term $P(X = x)$ is ignored in the posterior as it's the same for every y). And take the y that maximise $P(X = x|Y = y)P(Y = y)$ as the classification result.

Logistic Regression Here, we are using logsitic regression for multi-class classification, in which the logistic regression can also be called Softmax Regression. It is a discriminative model that directly model the conditional/posterior distribution $P(Y = y|X = x)$ instead of the joint distribution. The model is parameterized as follow:

$$P(Y = y|X = x; \theta) = \frac{\exp(\theta_y^\top x)}{\sum_{j=1}^K \exp(\theta_j^\top x)}$$

We take the logarithm of the likelihood of all the training data, and we can get:

$$J = \ln\left(\prod_{n=1}^N P(y^{(n)}|x^{(n)}; \theta)\right) = \sum_{n=1}^N (\theta_{y^{(n)}}^\top x^{(n)}) - \ln\left(\sum_{j=1}^K \exp(\theta_j^\top x)\right)$$

Take the derivative with respect to the parameter vector θ_k , we get:

$$\nabla_{\theta_k} J = \sum_{n=1}^N (I(y = k) - \frac{\exp(\theta_k^\top x)}{\sum_{j=1}^K \exp(\theta_j^\top x)})x$$

In the above equation, $I(\bullet)$ is the indicator function that is 1 when the condition is true otherwise 0. However, a closed-form solution cannot be found that make this derviative equals to 0. So we can use a local optimization method such as gradient descent or its variants to solve this problem.

In GD, the update of parameters follows:

$$\theta_k \leftarrow \theta_k - a \nabla_{\theta_k} J$$

Also, we can consider using SGD or adding regularization, momentum.

In the test phase, we get a test sample x , we calculate $\exp(\theta_y^\top x)$ for every y , and take the y that miximise that value as the classification result.

- Imagine that a certain word is never observed in the training data, but occurs in a test instance. What will happen when our Naive Bayes classifier predicts the probability of the this test instance? Explain why this situation is undesirable. How to avoid this problem? Will logistic regression have a similar problem? Why or why not?

In Naive Bayse method, if we just make the likelihood of words given document label y that did not occur in the training data labeled y to be 0, once such a word occurs in one training data, the posterior probability of the document label being y will be 0. This situation is certainly undesirable, because there are so many words that may be not included in a relatively small training set. Also, we can considier that when there is a test sample with a word with id i that never occur in any training samples, then the posterior probability for all y given this sample is 0.

So, we can use a method that is analogous to ridge estimation in which a bias of estimation is introduced to guarantee numerical stability (to smooth the problem) – we can add one to the count of every words so that no likelihood of one word will be 0. We do not need to do the add-one smoothing explicitly, this procedure should be done lazily when testing for efficiency, because this problem has a very sparse but large input space.

As for logistic regression, there might also be some word that do not occur in any of the training data – for example, in the test data, there is a word id ‘61188’ that is bigger than any other word ids in the training data. So in this situation, we should ignore the word count of this word j (it’s equivalent as the weight of this word j satisfy: $\theta_{ij} = 0$ for all document labels i).

- Implement Logistic Regression and Naive Bayes. Use add-one smoothing when estimating the parameters of your Naive Bayes classifier. For logistic regression, we found that a step size around 0.0001 worked well. Train both models on the provided training data and predict the labels of the test data. Report the training and test error of both models. Submit your code along with your homework.


```

$ ./main.py train -h
usage: main.py train [-h] [-v {None,test}] [-c CONFIG] [-s SAVE]
                    {Logistic,NaiveBayes} train_path test_path

positional arguments:
  {Logistic,NaiveBayes}  classifier type
  train_path             the path of the train data files
  test_path             the path of the test data files

optional arguments:
  -h, --help            show this help message and exit
  -v {None,test}, --validation {None,test}
                        use what method for validation. TODO: k-fold
  -c CONFIG, --config CONFIG
                        a string of space-separated configurations for the
                        classifier. wrong configuration will be discarded. eg.
                        `max_epoch=5 momentum=0.9
  -s SAVE, --save SAVE  the model name prefix, if not specified, the model
                        will not be saved.

```

Figure 1: View The Help Message

```

$ ./main.py train NaiveBayes ../PRHW_bayes_data/20news-bydate/matlab/ ../PRHW_bayes_data/20news-bydate/matlab/ -s "naivebayes"
[PROFILE] training: 0.667083978653 s elapsed.
2017-03-08 03:44:17,239 [doc-classify] Finished training NaiveBayes classifier.
[PROFILE] test the training set: 14.8545451164 s elapsed.
2017-03-08 03:44:32,094 [doc-classify] The error rate on **training set** of NaiveBayes classifier is 0.0567042328512 (639/11269)
[PROFILE] test the test set: 9.55620408058 s elapsed.
2017-03-08 03:44:41,650 [doc-classify] The error rate on **test set** of NaiveBayes classifier is 0.214790139907 (1612/7505)
2017-03-08 03:44:41,650 [doc-classify] Saving model to naivebayes-03-08-03-44-41.dat...
2017-03-08 03:44:41,973 [doc-classify] Saving model finished.

```

Figure 2: Naive Bayes Training

Model	Batch Size	Optimizer	Epoch	Train Error Rate	Test Error Rate
Naive Bayes				5.67%	21.48%
Logistic Regression-exp1	no	GD	40	35.22%	46.60%
Logistic Regression-exp2	250	SGD	50	19.42%	35.95%
Logistic Regression-exp3	125	SGD	50	16.72%	34.24%

Some screenshots is shown in help_message, NB_training, LR_training.

In the experiment, I find that randomized parameter initialization will slightly outperforms 0-initialization in Logistic Regression when the learning rate is small (If learning rate is sufficiently large, the randomized parameter initialization contribute very little after a few steps). 0-initialization is used here as the initial learning rate is large enough in the experiment. And the experiment results of non-stochastic GD showed that there would still be improvement with more training epochs, as no plateau of error rate is achieved in the two experiment: they all ended because the number of epochs exceeds the pre-defined limit (20/40), however, the convergence is too slow so I did not carry on GD training any more.

The training of Naive Bayes is much quicker than Logistic Regression. Logistic Regression needs multiple epochs to converge. Because in our Naive Bayes model, we assume the conditional independence of different words given y , and the feature space is really large, so the parameters that Naive Bayes need to learn is much less than Logistic regression. Assume there are d different words (input feature space size), the convergence time of NB is $O(\log(d))$, and the convergence time of LR is $O(d)$.

The hyper-parameters of non-stochastic GD are hard to tune... I tried for a while and give up trying any more. The "0.0001" learning rate does not work well in my case. SGD outperforms GD, the error rate of SGD training decreases faster. Note that there are randomness in each run of SGD training process

```

foxfi@foxfi-eva6:~/homework/pattern_recognition/homework1/code$ ./main.py train Logistic .
./data/matlab ../data/matlab -c "base_learning_rate=1 batch_size=125 weight_decay=0" -s lo
gistic 2>&1|tee logs/batch_125_wd0.log

x_dim: 53976; y_dim: 20
Configuration: {'max_word_id': 53975, 'decay_learning_rate': 0.5, 'learn_controller': 'pl
ateau', 'batch_size': '125', 'base_learning_rate': '1', 'epoch': 50, 'sparse': True, 'weig
ht_decay': '0', 'momentum': 0}

2017-03-08 14:37:16,171 [doc-classify] Finished training Logistic classifier.
[PROFILE] val: 12.7909350395 s elapsed.
[PROFILE] training: 2063.98967505 s elapsed.
2017-03-08 14:37:36,016 [doc-classify] The error rate on **training set** of Logistic clas
sifier is 0.167184310942 (1884/11269)
[PROFILE] test the training set: 19.8445670605 s elapsed.
2017-03-08 14:37:48,979 [doc-classify] The error rate on **test set** of Logistic classifi
er is 0.342438374417 (2570/7505)
2017-03-08 14:37:48,979 [doc-classify] Saving model to logistic-03-08-14-37-48.dat...
2017-03-08 14:37:49,790 [doc-classify] Saving model finished.
[PROFILE] test the test set: 12.9625918865 s elapsed.

```

Figure 3: Logistic Regression Training of exp3. SGD (batch size = 125)

because of the random shuffling step, but the overall convergence speed remains similar for the same configuration.

Future exploration: I think momentum will help the convergence in SGD case as it keep a running average of gradient which will make every gradient more accurate. Also, due to the large sparse input feature space, use some regularization, especially those that can lead to weight sparsity (eg. L1 Lasso) can help reducing overfit and accelerate the convergence.

The entrance of this program is “main.py”, and the usage of this scripts can be viewed using “-help” option. There are two subcommand “train” and “test” of which the detailed help message can also be viewd. This program has a dependency on the “numpy” lib for just using some basic matrix multiply and element-wise operation (When I’m coding the Naive Bayes classifier, I tried to keep the compatibility of this project for the situation that “numpy” is not installed. However, I get tired to write a counterpart and adapter for “numpy.ndarray”, so the logistic part of program assumes the installation of “numpy”). On my Mac OS X, the “numpy” version is “1.10.0”, but any version with compatible basic matrix operation will do.

NOTE: Although the tool will print out several configurations like weight_decay, I just tried some of them while debugging, but in the submitting code, code for some of the configurations are untested, and might not work well. Anyway, the default configuration (only specify “batch_size=blabla” with “-c” option) will work. And there are many “FIXME” and “TODO” in the code as this code is finished in a rush. Apologize

4. Which model performs better on this task? Why do you think this is the case?

In my experiements, the Naive Bayes classifier performs better on this task. I think the reasons might come from several aspects including problem properties and model properties:

- Independence Assumption(Generative vs. Discriminative): In the Naive Bayes classifier, we make the (bag-of-words) independence assumption between the occurrence of different words when the document label y is given.
 - Although this assumption seems quite naive and unreal, the actual mutual information of most pair of words is small. So a generative model that *utilize the useful prior knowledge in this domain (modeling of the world)* has the potential to perform better.
 - The imperfection of the assumption might not be so important for this specific classification task. This simple assumption and feature selection can lead to simple learning process.
- Sparse input space: This problem have a large but sparse input space, Logistic Regression will spend more effort learning unimportant weight than Naive Bayes. Naive Bayes model exploits the sparsity better, and hence have less parameters than Logistic Regression, this helps avoiding overfit. I think some regularization methods with feature selection property (learn sparse weights) will improve the Logistic Regression model.
- Tuning difficulty: There are quite a few hyper-parameters to tune in Logistic Regression, it's hard and time-consuming to find a good hyper-parameters set. Conversely, the Naive Bayes barely need tuning to achieve a fairly good result.