# **NBCT: Naive Bayes Clustering Toolkit**

(version 0.4.2)

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# **1** Introduction

NBCT (Naive Bayes Clustering Toolkit) is a C implementation of several probabilistic inference algorithms related to *naive Bayes clustering* — probabilistic clustering based on a *naive Bayes model* [2]. NBCT version 0.4.2 receives a dataset in a tabular format with discrete (nominal) and continuous (numeric) values, and performs the following tasks:<sup>1</sup>

- Clustering with EM learning
  - Avoiding undesired local maxima by:
    - \* Random restarts
    - \* Deterministic annealing EM algorithm
    - \* Split-merge EM algorithm (experimental)
  - Determining the number of clusters
- Other probabilistic inferences
  - Relevance analysis
  - Computing dissimilarities between objects, or between attribute values

NBCT is designed to be generic, and hence is expected to handle a variety of datasets including bag-of-words representation (with TF-IDF weighting) of documents [2].

Although NBCT was much influenced by *AutoClass* [4, 5], a popular probabilistic clustering tool in the data mining field, it has a couple of advantages over AutoClass. First, NBCT

<sup>&</sup>lt;sup>1</sup>This manual was unchanged from the one for version 0.4, except the descriptions related to the version numbers.

runs in three statistical frameworks — ML (maximum likelihood), MAP (maximum a posteriori) and VB (variational Bayes) [1, 3]. It is empirically shown in recent researches that the VB approach often shows a better performance than that by ML/MAP approaches in model selection (e.g. determining the optimal number of clusters) [3, 15], though ML/MAP is simpler and would be easier to work with. The second advantage is that NBCT is augmented with two EM algorithms aiming to escape from undesired local maxima, called the *deterministic annealing EM* (DAEM) algorithm [22] and the *split-merge EM* (SMEM) algorithm [23] (though the latter is now considered as experimental). Lastly, NBCT provides a flexible way of configurations and fully (i.e. even for multivariate Normal distributions) deals with missing values.

The rest of this document is comprised of three parts. First, in Section 2, we will see a typical usage of NBCT with an exemplar dataset included in the released package. Section 3 then gives a (rough) description of the algorithms used in NBCT. Section 4 is the third part which describes the detailed usage of NBCT. Lastly Section 5 makes some remarks on effective use of NBCT. We annotated '\*' to the title of the sections that relate to the advanced functions of NBCT, so the readers can skip these sections unless necessary.

# 2 Getting started

Let us use NBCT with an artificial dataset included in the released package. We suppose here that the installation of NBCT (see §4.2 for the procedure) has been successfully done. Also consider that we are working in the example directory of the unfolded package. In this section, we just focus on the look and feel of NBCT, and do not aim at listing all functions. We first describe the task of clustering.

## 2.1 What is clustering?

Given a dataset D of *objects*,<sup>2</sup> clustering [13] is a task to group similar objects in D into sets called *clusters*. For example, one may wish to obtain some groups of Web pages that seem to have the same topic, or one may group the patients who show similar medical states. Like associative rule mining, clustering requires no (or less) human annotations, and hence can be utilized as a first step in knowledge discovery.

Naive Bayes clustering can be seen as unsupervised classification of objects, based on a naive Bayes model. So we may use the term '*classes*' instead of clusters. Each object is classified into one of K anonymous classes. We also consider each object to be represented by a vector of values of one or more predefined *attributes*.

Table 1: The descriptions of attributes in ex1\_freq.csv.

			Formally
Index	Туре	Possible values	referred to
0	ID	P0, P1,, P999	—
1	Answer class	0,1,2,3	—
2	Discrete	a, b, o, ab	$A_1$
3	Continuous	(floating-point number)	$A_2$
4	Ignored	—	
5	Discrete	male, female	$A_3$
6	Continuous	(floating-point number)	$A_4$
7	Continuous	(floating-point number)	$A_5$
8	Discrete	high,mid,low	$A_6$
9	Ignored		

## 2.2 Target data

The dataset D is supposed to be in tabular form. To be concrete, let us open ex1\_freq.csv, an artificial dataset, in the directory by some text viewer:

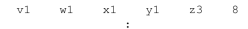
```
P0,0,b,4.74,6.48,female,0.36,1.51,mid,x0
P1,0,a,4.06,1.36,female,-0.38,0.99,mid,x0
P2,0,a,5.04,4.38,male,1.59,1.01,mid,x0
P3,0,b,5.13,5.33,female,0.84,0.69,mid,x0
:
```

Each of 1,000 lines in the file corresponds to an object, and each of comma-separated values in the line corresponds to a value of an attribute of the object. Each object has 10 attributes, which are described in Table 1. The attribute 0 (the indices of attributes are zero-based) indicates the ID of the corresponding object. The attribute 1 indicates the answer class to which the object is known to belong. Actually, in this dataset, all objects were sampled from a naive Bayes model (formally described in §2.3) with 4 classes, and the answer classes were determined in the process of sampling. The attributes 4 and 9 are noisy ones, and should be ignored. Only the rest of the attributes are used for clustering, and hereafter called descriptive attributes. In a formal context, we only consider these descriptive attributes, and T denotes the number of descriptive attributes. For instance, the first object in ex1\_freq.csv is written as x = (b, 4.74, c)female, 0.36, 1.51, mid), and T = 6.

On the other hand, there can be a dataset where several objects have the same attribute values. In NBCT, unless having continuous attributes or the ID attribute, these objects will be regarded as the same since there is no way to distinguish them. Consequently the dataset is a multiset of objects. In the dataset  $ex2\_freq.txt$ , objects who have the same vector  $\mathbf{x}$  of attribute values were suppressed, and instead their count  $N(\mathbf{x})$  is added to the right-most column, which is called the *count column*:

v1	w1	x1	y1	z1	16
v1	w1	x1	y1	z2	21

 $<sup>^2</sup> For$  a historical reason, in the terminal messages from the programs in NBCT, the term '*case*' is used instead of 'object.'



For example, we have  $N(\mathbf{x}) = 21$  for  $\mathbf{x} = (v1, w1, x1, y1, z2)$ . Please note here that the count column is not considered as an attribute. For the dataset where the count column is omitted, like  $ex1\_freq.csv$ ,  $N(\mathbf{x})$  is considered to be one for every object  $\mathbf{x}$ . Also note that NBCT accepts the tab character as the delimiter of attribute values for the input files whose file extension is 'txt'.

## 2.3 Naive Bayes models

To build clusters of objects in the dataset D, we attempt to use a probabilistic model called a naive Bayes model. In naive Bayes models, we consider that the objects in D were generated in a causal way depicted as a Bayesian network (Fig. 1), which has random variables C and  $A_i$  ( $1 \le i \le T$ ). C is called the *class variable*, and  $A_i$  are called the *attribute variables*. We then write the vector of  $A_1, A_2, \ldots, A_T$  as X. In the case of  $e \ge 1$ \_freq.csv, the descriptive attributes are referred to by  $A_i$  in the rightmost column of Table 1.

In a naive Bayes model, the class of each object is firstly determined as *k* under the class distribution p(C=k), and then attribute values **x** are conditionally determined under the attribute distribution p(X = x | C = k). On this conditional distribution, NBCT assumes that each attribute  $A_i$  is of three types:<sup>3</sup>

- Discrete (and single-dimensional) *A<sub>i</sub>* follows a categorical distribution.<sup>4</sup>
- Continuous and single-dimensional A<sub>i</sub> follows a univariate Normal distribution.
- Continuous and multidimensional some attribute variables including *A<sub>i</sub>* jointly follow a multivariate Normal distribution.

We let  $X_{\text{disc}}$  be a set of discrete attributes,  $X_{\text{cont-u}}$  be a set of continuous and single-dimensional attributes, and  $X_{\text{cont-m}}$  be a set of vectors each of which is comprised of the continuous attributes jointly following a certain multivariate Normal distribution. We define  $X \stackrel{\text{def}}{=} X_{\text{disc}} \cup X_{\text{cont-u}} \cup X_{\text{cont-m}}$  and let  $X = \{X_1, \ldots, X_j, \ldots, X_J\}$ . Note that  $J \leq T$  obviously holds, and that  $X_j \in X$  is a vector  $(A_{i_1}, A_{i_2}, \ldots, A_{i_n})$  of attributes. In the case of  $\text{exl_freq.csv}$ , we may have J = 4,  $X_1 = (A_1), X_2 = (A_2, A_4, A_5), X_3 = (A_3)$  and  $X_4 = (A_6)$ , where  $X_1, X_3, X_4 \in X_{\text{disc}}$  and  $X_2 \in X_{\text{cont-m}}$  (see also Table 1). Also it is possible to have  $J = 5, X_1 = (A_1), X_2 = (A_2, A_5)$ ,

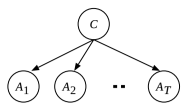


Figure 1: Bayesian network representation of a naive Bayes model, where all attributes are single-dimensional.

 $X_3 = (A_3), X_4 = (A_4) \text{ and } X_5 = (A_6), \text{ where } X_1, X_3, X_5 \in X_{\text{disc}}, X_2 \in X_{\text{cont-m}} \text{ and } X_4 \in X_{\text{cont-u}}.$ 

For simplicity, in the context where we know  $X_j$  (often) consists of a single attribute, i.e. a vector of length 1, we write  $X_j$  and its value as  $X_j$  and  $x_j$ , respectively. Furthermore, we refer to all nominal values by integers. That is, for an attribute  $X_j$  that can have  $V_j$  values, we number them from 1 to  $V_j$ . For instance, for  $A_1$  in Table 1, the values a, b, o and ab are referred to by 1, 2, 3 and 4, respectively.

Now, the *joint probability*, i.e. the probability that an object with attribute values  $\mathbf{x} = (a_1, a_2, ..., a_T)$  belongs to the class k, is simply computed as:

$$p(C=k, A_1 = a_1, A_2 = a_2, \dots, A_T = a_T)$$
  
=  $p(C=k) \prod_{j=1}^{J} p(X_j = x_j | C = k).$  (1)

According to the type of  $X_j$ ,  $p(X_j = x_j | C = k)$  is predefined as follows:

- When X<sub>j</sub> ∈ X<sub>disc</sub>, p(X<sub>j</sub> | C = k) follows a categorical distribution. We consider θ<sub>j,k,xj</sub> <sup>def</sup> = p(X<sub>j</sub> = x<sub>j</sub> | C = k) as a parameter of the distribution. V<sub>j</sub> stands for the number of its possible values, and we have 1 ≤ x<sub>j</sub> ≤ V<sub>j</sub>.
- When  $X_j \in X_{\text{cont-u}}$ ,  $p(X_j | C = k)$  follows a univariate Normal distribution. That is,  $p(X_j = x_j | C = k)$  is:

$$\mathcal{N}(x_{j} \mid \mu_{j,k}, \sigma_{j,k}^{2}) = \frac{1}{\sqrt{2\pi\sigma_{j,k}^{2}}} \exp\left(-\frac{(x_{j} - \mu_{j,k})^{2}}{2\sigma_{j,k}^{2}}\right),$$
(2)

where  $\mu_{j,k}$  and  $\sigma_{j,k}^2$  are the mean and the variance, respectively.

• When  $X_j \in X_{\text{cont-m}}$ ,  $p(X_j | C = k)$  follows a multivariate Normal distribution. That is,  $p(X_j = x_j | C = k)$  is:

$$\mathcal{N}(\boldsymbol{x}_{j} \mid \boldsymbol{\mu}_{j,k}, \boldsymbol{\Sigma}_{j,k}) = \frac{1}{\sqrt{(2\pi)^{n_{j}} |\boldsymbol{\Sigma}_{j,k}|}} \exp\left(-\frac{1}{2}(\boldsymbol{x}_{j} - \boldsymbol{\mu}_{j,k})^{\mathrm{T}} \boldsymbol{\Sigma}_{j,k}^{-1}(\boldsymbol{x}_{j} - \boldsymbol{\mu}_{j,k})\right),$$
(3)

where  $n_j$  is the dimension of  $X_j$ , and  $\mu_{j,k}$  and  $\Sigma_{j,k}$  are the mean vector and the covariance matrix, respectively.

<sup>&</sup>lt;sup>3</sup>AutoClass also introduces these three types. In a usual definition of naive Bayes models, on the other hand, all attributes are considered to be singledimensional. We may say that the naive Bayes models considered here are an extension of such usual ones.

<sup>&</sup>lt;sup>4</sup>The categorical distribution is defined as the generalization of the Bernoulli distribution to more than 2 categories.

The class variable *C*, on the other hand, is discrete and follows a categorical distribution, whose parameters are  $\theta_k \stackrel{\text{def}}{=} p(C = k)$  ( $1 \le k \le K$ ). We use  $\theta$  for a vector of all parameters above. Eq. 1 is then simplified and written in a parameterized form as:

$$p(k, \mathbf{x} \mid \boldsymbol{\theta}) = p(k \mid \boldsymbol{\theta}) \prod_{j=1}^{J} p(\mathbf{x}_j \mid k, \boldsymbol{\theta}),$$
(4)

where p(C = k,...) and  $p(X_j = x_j,...)$  are abbreviated as p(k,...) and  $p(x_j,...)$ , respectively.

## 2.4 Running NBCT

## 2.4.1 Basic usage

Now try NBCT with the dataset contained in ex1\_freq.csv. Let us assume here that the true number of classes is known as 4. Then, to get the clusters of objects in ex1\_freq.csv, we invoke an executable named nbc with several option flags:

```
nbc -f ex1 -k 4 -x 10 -I 0 -E 1 -y 4,9 -u 3,6,7
```

The option '-f ex1' specifies the names of input/output files including ex1\_freq.csv. Besides, the options '-k 4' and '-x 10' indicate that the number of clusters is 4 (K = 4), and that the number of attributes is 10, respectively. The rest of option flags specify the attribute types. The options '-I 0', '-E 1' and '-y 4, 9' respectively say that the attribute 0 is the ID attribute, and the attribute 1 is the answer class attribute, and the attributes 4 and 9 should be ignored (see also Table 1). The option '-u 3, 6, 7' says the attributes 3, 6 and 7 jointly follow a multivariate Normal distribution. The attributes that are not explicitly specified are considered as discrete attributes. After running the program, we may see the terminal messages below:

```
Reading cases from ex1_freq.csv...done
#classes = 4:
    #iters 0 (Converged: 19 iters)
    L=-5747.688373
    BIC = -5923.836132
```

In advance of clustering, the program runs the EM (expectationmaximization) algorithm (§3.1.1) to estimate the parameters of the naive Bayes model. The EM algorithm is an iterative, hill-climbing algorithm for parameter estimation. The messages above report the progress of the EM algorithm, i.e. we made 19 EM iterations, the log-likelihood was converged to -5747.688373, and so on.

One may notice it possible to have a different setting for continuous attributes. For example, we may specify<sup>5</sup> that attribute 3 and 7 jointly follow a bivariate Normal distribution and attribute 6 follows a univariate Normal distribution:<sup>6</sup>

As mentioned before, the count column is not considered as an attribute, so for a dataset having the count column, the number of attributes, which is specified by '-x' option, should be the number of columns minus one. Specifically, for  $ex2_freq.txt$ , we may run:

nbc -f ex2 -k 4 -x 5

## 2.4.2 Output files

After the run with ex1\_freq.csv, three files named ex1\_ {param,cluster,log}.csv should have been created. The first file, ex1\_param.csv, contains the parameters estimated by the EM algorithm:

```
#### ID:PARAM-CLASS
#### Class k, Class parameter P(k)
####
0,0.299999927172
1,0.20000072828
2,0.40000024986
3,0.0999999750144
#### ID:PARAM-CAT2
#### Attr. ID, Class k, Attr. value x2, Attr. par
ameter P(x2|k)
####
2,0,a,0.0933333559908
2,0,ab,0.196666862475
#### ID:PARAM-GAUSS-MEAN0
#### Gauss ID, Class k, Attr. ID, Mean
####
0,0,3,0.999999808665
0,0,6,5.00016666618
```

We see from above that the class parameters were estimated as 0.299, 0.200, ..., and so on. In general, each output file contains several data matrices, each of which has header lines beginning with '#'.

The second file, ex1\_cluster.csv, contains the data matrices on a mapping from objects to clusters:

```
#### ID:CLUSTER-CASE-BY-CASE
##### Case ID, Answer class, Cluster k, Attribute
values x, Membership prob. P(k|x)
####
P0,0,3,b,4.74,female,0.36,1.51,mid,1
P1,0,3,a,4.06,female,-0.38,0.99,mid,1
P2,0,3,a,5.04,male,1.59,1.01,mid,0.99999999999
:
```

We see that the object named P0, x = (b, 4.74, female, 0.36, 1.51, mid), belongs to the cluster 3, and so on. One may also notice that the rows in the data matrix named 'CLUSTER-CASE-BY-CASE' are ordered by the indices of objects. On the other hand, in the data matrix 'CLUSTER-CASE-BY-CLUSTER', the rows are ordered by the indices of clusters:

<sup>&</sup>lt;sup>5</sup> \\' means a continuation of the command line (as used in popular shell programs in UNIX/Linux), and is used here just for space limitation.

<sup>&</sup>lt;sup>6</sup>The question of which is better is a problem of *model selection*, and as will be mentioned later, we may decide it by a Bayesian score provided by NBCT.

```
% vnbc -f ex1 -k 2:10:1 -x 10 -I 0 -E 1 -y 4,9 -u 3,6,7
Reading cases from ex1_freq.csv...done
|C|asses| = 2:
  #iters 0 (Converged: 9 iters) F=-7485.127335
 Variational Free Energy = -7485.127335 (temporarily optimal)
|Classes| = 4:
  #iters 0 (Converged: 16 iters)
                                       F=-6014.731751
  Variational Free Energy = -6014.731751 (temporarily optimal)
|Classes| = 5:
  #iters 0. (Converged: 30 iters)
                                        F=-6017.392785
  Variational Free Energy = -6017.392785
|Classes| = 10:
  #iters 0. (Converged: 28 iters)
                                       F = -6027.032856
  Variational Free Energy = -6027.032856
Optimal |Classes| = 4
```

Figure 2: Terminal messages from vnbc in the 'model selection' mode.

```
#### ID:CLUSTER-CASE-BY-CLUSTER
##### Cluster k, Answer class, Case ID, Attribute
values x, Membership prob. P(k|x)
####
0,2,P300,a,-0.2,female,5.25,5.08,low,1
0,2,P301,ab,0.58,male,5.03,5.9,low,1
0,2,P302,a,1.74,male,4.12,4.45,low,1
:
```

The above tell us that the cluster 0 includes the objects P300, P301, and so on. Finally, the third file,  $ex1_log.csv$ , contains additional information on the (last) execution.

#### 2.4.3 Various execution options

NBCT provides many options for the EM algorithm. For example, since the EM algorithm is a hill-climbing algorithm, and is known to be often trapped in undesirable local maxima, we sometimes wish to restart the EM algorithm with several different initial settings. In NBCT, this method is enabled by giving the number of restarts to -n option:<sup>7</sup>

```
% nbc -f ex1 -k 4 -x 10 -I 0 -E 1 -y 4,9 ∖
                                   -u 3,6,7 -n 10
Reading cases from ex1_freq.csv...done
#classes = 4:
 [0] #iters 0 (Converged: 18 iters)
                                    L=-5747.688373
 [1] #iters 0 (Converged: 16 iters)
                                      L=-5747.688373
 [2] #iters 0 (Converged: 14 iters)
                                      L = -5747.688373
 [3] #iters 0. (Converged: 23 iters) L=-5747.688373
                                     L=-5747.688373
 [4] #iters 0. (Converged: 22 iters)
 [5] #iters 0 (Converged: 16 iters)
                                      L=-5747.688373
 [6] #iters 0. (Converged: 24 iters)
                                    L=-5747.688373
 [7] #iters 0. (Converged: 24 iters) L=-5747.688373
 [8] #iters 0 (Converged: 15 iters)
                                      L=-5747.688373
 [9] #iters 0 (Converged: 14 iters)
                                      L=-5747.688373
 <<Resumed best parameter set #5>>
                                      L=-5747.688373
 [5] #iters (Converged: 17 iters)
BIC = -5923.836132
```

We can find from above the sixth initial setting provides the best estimate of parameters (though the differences were subtle). To be precise, the parameter estimation method we have run is called 'maximum likelihood (ML) estimation.' On the other hand, maximum a posteriori (MAP) estimation is said to be more robust against the problem of data-sparseness, which often arises with a small data. In MAP estimation, we should tell the *pseudo counts* or *hyperparameters* to the program. In NBCT, it is possible to specify pseudo counts or hyperparameters in a flexible form, and we can make similar settings to AutoClass with –A option as well:

```
% nbc -f ex1 -k 4 -x 10 -I 0 -E 1 -y 4,9 -u 3,6,7 -A
Reading cases from ex1_freq.csv...done
#classes = 4:
    #iters 0. (Converged: 31 iters) L=-5907.149381
    Cheeseman-Stutz score = -6024.130102
```

We have assumed so far that the true number K of clusters is known in advance, but in practical situations, it is often unknown. So finding the optimal number of clusters (based on the dataset) is a key issue in clustering. Again, this can be seen a kind of model selection problem. NBCT provides a facility that computes the scores on K based on the marginal likelihood P(D | K), the plausibility of D given the number K of clusters.

Although the scores on *K* are provided for the frameworks of ML and MAP estimation, we here use variational free energy instead as a score on *K* in a variational Bayesian (VB) approach. In VB, we use another executable named vnbc. The terminal messages from vnbc are shown in Fig. 2, where the number of clusters to be examined ranges from 2 to 10. If we specify '-k *Kmin*: *Kmax*: *Kstep*', NBCT will be switched into the 'model selection' mode. In this example, we fortunately recovered the true number of clusters as 4.

#### 2.4.4 Auxiliary tool for post-processing

Lastly, we mention nbcsep, an auxiliary tool for postprocessing the outputs from nbc and vnbc. As seen before, each output file contains several data matrices, each having an

<sup>&</sup>lt;sup>7</sup>'%' is the prompt symbol.

identifier. Typically, nbcsep extracts these data matrices, and puts each of them into an individual file. The name of a new file includes the identifier of the corresponding data matrix. For example, let us apply nbcsep to ex1\_param.csv:

```
% nbcsep -f foo ex1_param.csv
Output: foo_PARAM-CLASS.csv
Output: foo_PARAM-CAT2.csv
Output: foo_PARAM-CAT5.csv
Output: foo_PARAM-CAT8.csv
Output: foo_PARAM-GAUSS-MEAN0.csv
Output: foo_PARAM-GAUSS-COVAR0.csv
```

Here, the common prefix of the names of newly created files was given by -f option.

# **3** Clustering algorithms

This section gives a detailed description on the clustering algorithms and the other related algorithms provided in NBCT.

## 3.1 ML/MAP based clustering

## 3.1.1 Parameter estimation based on ML

In advance of clustering and the other probabilistic inferences based on the joint distribution  $p(k, x | \theta)$ , we need to estimate the parameters  $\theta$  from the dataset D. Let us recall that we are given a set D of objects where N(x) is the number of occurrences of objects that have the attribute value x. We define N as the number of total occurrences of objects, that is,  $N = \sum_{x} N(x)$ . Since in clustering, we do not know the class to which each object belongs, the dataset D contains no information about k. In this sense, D is often called *incomplete data*. In *maximum likelihood (ML) estimation*, we try to find the parameters  $\theta$  that maximize the *likelihood*  $p(D | \theta)$ . That is, we have:

$$\hat{\boldsymbol{\theta}}_{ML} = \operatorname{argmax}_{\boldsymbol{\theta}} p(D \mid \boldsymbol{\theta}) = \operatorname{argmax}_{\boldsymbol{\theta}} \log p(D \mid \boldsymbol{\theta})$$

$$= \operatorname{argmax}_{\boldsymbol{\theta}} \sum_{\boldsymbol{x}} N(\boldsymbol{x}) \log p(\boldsymbol{x} \mid \boldsymbol{\theta})$$

$$= \operatorname{argmax}_{\boldsymbol{\theta}} \sum_{\boldsymbol{x}} N(\boldsymbol{x}) \log \sum_{k=1}^{K} p(k, \boldsymbol{x} \mid \boldsymbol{\theta}). \quad (5)$$

Due to the lack of the information on the cluster (*k*), it is not easy to analytically maximize  $p(D | \theta)$  or log  $p(D | \theta)$ . Instead, we use the *EM* (*expectation-maximization*) *algorithm* [9]. Fig. 3 is the EM algorithm derived for naive Bayes models.<sup>8</sup>

#### 3.1.2 Parameter estimation based on MAP

It is well-known in the machine learning literature that ML estimation often suffers from the problem of *data-sparseness* when

- 1. Initialize randomly the parameters  $\theta_k$   $(1 \le k \le K)$  and  $\theta_{j,k,x_j}$  $(1 \le k \le K, 1 \le j \le J, 1 \le x_j \le V_j).$
- 2. Repeat the following until the log-likelihood log  $p(D \mid \theta)$  converges:

$$p(k \mid \boldsymbol{x}, \boldsymbol{\theta}) \propto p(k, \boldsymbol{x} \mid \boldsymbol{\theta}) = p(k \mid \boldsymbol{\theta}) \prod_{j=1}^{J} p(x_j \mid k, \boldsymbol{\theta})$$
 (6)

$$E[k] := \sum_{\mathbf{x}} N(\mathbf{x}) p(k \mid \mathbf{x}, \boldsymbol{\theta})$$
(7)

$$\theta_k \propto E[k]$$

$$X_j \in \mathcal{X}_{\text{disc}} :$$
(8)

$$_{j,k,x_j} \propto \sum_{\mathbf{x}' \in D: x'_j = x_j} N(\mathbf{x}') p(k \mid \mathbf{x}', \boldsymbol{\theta})$$
 (9)

$$\mu_{j,k} := \frac{1}{E[k]} \sum_{x \in D} p(x_j \mid k, \theta) x_j$$
(10)

$$\sigma_{j,k}^{2} := \frac{1}{E[k]} \sum_{x \in D} p(x_{j} \mid k, \theta) (x_{j} - \mu_{j,k})^{2}$$
(11)

$$X_j \in \mathcal{X}_{\text{cont-m}}$$
:

ŀ

 $X_j \in \mathcal{X}_{\text{cont-u}}$ :

$$\boldsymbol{u}_{j,k} := \frac{1}{E[k]} \sum_{\boldsymbol{x} \in D} p(\boldsymbol{x}_j \mid \boldsymbol{k}, \boldsymbol{\theta}) \boldsymbol{x}_j$$
(12)

$$\boldsymbol{\Sigma}_{j,k} \quad := \quad \frac{1}{E[k]} \sum_{\boldsymbol{x} \in D} p(\boldsymbol{x}_j \mid k, \boldsymbol{\theta}) (\boldsymbol{x}_j - \boldsymbol{\mu}_{j,k}) (\boldsymbol{x}_j - \boldsymbol{\mu}_{j,k})^{\mathrm{T}}$$
(13)

Figure 3: The EM algorithm for ML estimation in naive Bayes models.

the data size *N* is not so large compared to the number of parameters. One way for avoiding this problem is to take a Bayesian approach, in which we consider a *prior distribution*  $p(\theta)$  on the parameter space  $\Theta$ . As is often done, in NBCT, we use *conjugate prior distributions* [8] for  $p(\theta)$ . To be more concrete:

• As a prior distribution of  $\theta_{class}$ , (a vector of) the parameters  $\theta_k$  ( $1 \le k \le K$ ) for the class variable *C*, we introduce a Dirichlet distribution:

$$p(\boldsymbol{\theta}_{\text{class}}) \stackrel{\text{def}}{=} \frac{\Gamma(\sum_{k=1}^{K} \alpha_k)}{\prod_{k=1}^{K} \Gamma(\alpha_k)} \prod_{k=1}^{K} \theta_k^{\alpha_k - 1} \quad . \tag{14}$$

Each  $\alpha_k$  corresponds to  $\theta_k$ , and is called a *hyperparameter*.

• As a prior distribution of  $\theta_{\text{disc}}$ , the parameters  $\theta_{j,k,x_j}$  for  $X_j \in X_{\text{disc}}$ , we also introduce a Dirichlet distribution:

$$p(\boldsymbol{\theta}_{\text{disc}}) \stackrel{\text{def}}{=} \prod_{k=1}^{K} \prod_{j \in \mathcal{J}_{d}} \left( \frac{\Gamma\left(\sum_{x_{j}=1}^{V_{j}} \alpha_{j,k,x_{j}}\right)}{\prod_{x_{j}=1}^{V_{j}} \Gamma\left(\alpha_{j,k,x_{j}}\right)} \prod_{x_{j}=1}^{V_{j}} \theta_{j,k,x_{j}}^{\alpha_{j,k,x_{j}}-1} \right)$$
(15)

<sup>&</sup>lt;sup>8</sup>In this document, the symbol ' $\propto$ ' means a substitution with normalization.

where  $\mathcal{J}_d$  is a set of the indices of discrete attributes, i.e.  $\mathcal{J}_d \stackrel{\text{def}}{=} \{j \mid X_j \in X_{\text{disc}}\}$ . Each  $\alpha_{j,k,x_j}$  is the hyperparameter corresponding to  $\theta_{j,k,x_j}$ .

• As a prior distribution of  $\theta_{\text{cont-u}}$ , the parameters  $\mu_{j,k}$  and  $\sigma_{j,k}^2$  for  $X_j \in \chi_{\text{cont-u}}$ , we introduce a Normal-Gamma distribution:

$$p(\boldsymbol{\theta}_{\text{cont-u}}) \stackrel{\text{def}}{=} \prod_{k=1}^{K} \prod_{j \in \mathcal{J}_{u}} p(\mu_{j,k}, \lambda_{j,k})$$
$$= \prod_{k=1}^{K} \prod_{j \in \mathcal{J}_{u}} \mathcal{N}(\mu_{j,k} \mid \mu_{0}, (\tau \lambda_{j,k})^{-1}) \cdot$$
$$\mathcal{G}(\lambda_{j,k} \mid a, b), \quad (16)$$

$$\mathcal{G}(\lambda_{j,k} \mid a, b) \stackrel{\text{def}}{=} \frac{1}{\Gamma(a)} b^a \lambda_{j,k}^{a-1} \exp(-b\lambda_{j,k}), \quad (17)$$

where  $\mathcal{J}_{u} \stackrel{\text{def}}{=} \{j \mid X_{j} \in \mathcal{X}_{\text{cont-u}}\}$ , and  $\lambda_{j,k} \stackrel{\text{def}}{=} \frac{1}{\sigma_{j,k}^{2}}$  is called the *precision* of the Normal distribution.  $\mu_{0}, \tau, a$  and *b* are the hyperparameters.

• As a prior distribution of  $\theta_{\text{cont-m}}$ , the parameters  $\mu_{j,k}$  and  $\Sigma_{j,k}$  for  $vX_j \in X_{\text{cont-m}}$ , we introduce a Normal-Wishart distribution:

$$p(\boldsymbol{\theta}_{\text{cont-m}}) \stackrel{\text{def}}{=} \prod_{k=1}^{K} \prod_{j \in \mathcal{J}_{m}} p(\boldsymbol{\mu}_{j,k}, \boldsymbol{\Lambda}_{j,k}) \cdot$$
$$= \prod_{k=1}^{K} \prod_{j \in \mathcal{J}_{m}} \mathcal{N}(\boldsymbol{\mu}_{j,k} \mid \boldsymbol{\mu}_{0}, (\tau \boldsymbol{\Lambda}_{j,k})^{-1})$$
$$\mathcal{W}(\boldsymbol{\Lambda}_{j,k} \mid \boldsymbol{\alpha}, \boldsymbol{\Sigma}_{0}), \quad (18)$$
$$\mathcal{U}(\boldsymbol{\Lambda}_{j,k} \mid \boldsymbol{\alpha}, \boldsymbol{\Sigma}_{0}) \stackrel{\text{def}}{=} c(n_{j}, \boldsymbol{\alpha}) |\boldsymbol{\Sigma}_{0}|^{\frac{\alpha}{2}} |\boldsymbol{\Lambda}_{j,k}|^{\frac{\alpha-n_{j}-1}{2}} \cdot$$

W

$$exp\left(-\frac{1}{2}tr(\boldsymbol{\Sigma}_{0}\boldsymbol{\Lambda}_{j,k})\right), \quad (19)$$

$$c(n_{j},\alpha) \stackrel{\text{def}}{=} \left(2^{\frac{\alpha n_{j}}{2}}\pi^{\frac{n_{j}(n_{j}-1)}{4}}\prod_{i=1}^{n_{j}}\Gamma\left(\frac{\alpha+1-i}{2}\right)\right)^{-1}, \quad (20)$$

where  $n_j$  is the dimension of  $X_j$ ,  $\mathcal{J}_m \stackrel{\text{def}}{=} \{j \mid X_j \in X_{\text{cont-m}}\}$ , and  $\Lambda_{j,k} \stackrel{\text{def}}{=} \Sigma_{j,k}^{-1}$  is also the precision.  $\mu_0$ ,  $\tau$ ,  $\alpha$  and  $\Sigma_0$  are the hyperparameters.

Finally, the whole prior distribution is the product of the above distributions:  $p(\theta) = p(\theta_{class})p(\theta_{disc})p(\theta_{cont-u})p(\theta_{cont-m})$ .

Now, instead of  $p(D \mid \theta)$ , we maximize  $p(\theta \mid D)$ , an *a posteriori probability* of  $\theta$  given the dataset *D*. That is, we have:

$$\hat{\theta}_{MAP} = \operatorname{argmax}_{\theta} p(\theta \mid D) = \operatorname{argmax}_{\theta} \log p(\theta \mid D)$$

$$= \operatorname{argmax}_{\theta} \log \frac{p(\theta)p(D \mid \theta)}{p(D)}$$

$$= \operatorname{argmax}_{\theta} \{\log p(\theta) + \log p(D \mid \theta)\}. \quad (29)$$

- 1. Initialize randomly the parameters  $\theta_k$   $(1 \le k \le K)$  and  $\theta_{j,k,x_j}$  $(1 \le k \le K, 1 \le j \le J, 1 \le x_j \le V_j).$
- 2. Repeat the following until the log-likelihood log  $p(D \mid \theta)$  converges:

$$p(k \mid \boldsymbol{x}, \boldsymbol{\theta}) \propto p(k, \boldsymbol{x} \mid \boldsymbol{\theta}) = p(k \mid \boldsymbol{\theta}) \prod_{j=1}^{J} p(x_j \mid k, \boldsymbol{\theta})$$
 (21)

$$E[k] := \sum_{\mathbf{x}} N(\mathbf{x}) p(k \mid \mathbf{x}, \boldsymbol{\theta})$$
(22)

$$\theta_k \propto E[k] + (\alpha_k - 1)$$
 (23)  
 $X_j \in X_{\text{disc}}$ :

$$\theta_{j,k,x_j} \propto \sum_{\boldsymbol{x}' \in D: x'_j = x_j} N(\boldsymbol{x}') p(k \mid \boldsymbol{x}', \boldsymbol{\theta}) + (\alpha_{j,k,x_j} - 1)$$
(24)

$$X_j \in \mathcal{X}_{\text{cont-u}}$$
:

$$_{i,k} := \frac{\tau \mu_0 + \sum_{x \in D} p(x_j \mid k, \theta) x_j}{\tau + E[k]}$$
(25)

$$\sum_{j,k}^{2} := \frac{\tau(\mu_{j,k} - \mu_{0})^{2} + \sum_{x \in D} p(x_{j} \mid k, \theta)(x_{j} - \mu_{j,k})^{2} + 2b}{2a - 1 + E[k]}$$
(26)

$$X_j \in \mathcal{X}_{\text{cont-m}}$$
:

μ

 $\sigma$ 

$$\boldsymbol{\mu}_{j,k} := \frac{\tau \boldsymbol{\mu}_0 + \sum_{\boldsymbol{x} \in D} p(\boldsymbol{x}_j \mid \boldsymbol{k}, \boldsymbol{\theta}) \boldsymbol{x}_j}{\tau + E[k]}$$
(27)

$$\Sigma_{j,k} := \frac{1}{\alpha - n_j + E[k]} \cdot \left( \sum_{\mathbf{x} \in D} p(\mathbf{x}_j \mid k, \boldsymbol{\theta}) (\mathbf{x}_j - \boldsymbol{\mu}_{j,k}) (\mathbf{x}_j - \boldsymbol{\mu}_{j,k})^{\mathrm{T}} \right)$$

$$+ \tau(\boldsymbol{\mu}_{j,k} - \boldsymbol{\mu}_0)(\boldsymbol{\mu}_{j,k} - \boldsymbol{\mu}_0)^{\mathrm{T}} + \boldsymbol{\Sigma}_0 \Big)$$
(28)

Figure 4: The EM algorithm for MAP estimation in naive Bayes models.

This procedure is usually called *MAP* (maximum a posteriori) estimation.

The EM algorithm for MAP estimation is shown in Fig. 4. The algorithm is obtained by modifying the convergence condition and the procedure in M-step of the ML version (Fig. 3) [10]. Let us introduce  $\delta_k \stackrel{\text{def}}{=} \alpha_k - 1$  and  $\delta_{j,k,x_j} \stackrel{\text{def}}{=} \alpha_{j,k,x_j} - 1$  as the *pseudo counts* or the *smoothing constants* in estimating the corresponding parameters.<sup>9</sup> If we let  $\delta_k$  and  $\delta_{j,k,x_j}$  be positive, the estimated parameters will also be positive, so we would be able to avoid the problem of data-sparseness.

<sup>&</sup>lt;sup>9</sup>In NBCT, we use the term 'pseudo counts' in the sense of ones used in the MAP estimator, and for compatibility, it is designed that the users are expected to configure the hyperparameters  $\alpha_k$  and  $\alpha_{j,k,x_j}$  through the corresponding pseudo counts  $\delta_k$  and  $\delta_{j,k,x_j}$  (even in VB learning). In the Bayesian estimator, on the other hand, hyperparameters  $\alpha_k$  and  $\alpha_{j,k,x_j}$  themselves can be considered as pseudo counts. Another confusing issue in NBCT is that the current version does not allow  $\delta_{j,k,x_j}$  to be negative. Of course this restriction is reasonable for MAP estimation, but theoretically it is important to note that the prior distribution  $p(\theta)$  is defined for  $\alpha_k \ge 0$  and  $\alpha_{j,k,x_j} \ge 0$ .

For simplicity, these pseudo counts are set to be uniform. When specifying all  $\delta_k$  and  $\delta_{j,k,x_j}$  to be 1.0, the estimation procedure is sometimes called *Laplace's estimation*. In Auto-Class [4], all  $\delta_k$  are fixed at 1/K, and all  $\delta_{j,k,x_j}$  are fixed at  $1/V_j$ .

A similar discussion is possible for continuous attributes. To see this, let us introduce  $\bar{x}_{j,k}$  defined as:

$$\bar{x}_{j,k} \stackrel{\text{def}}{=} \frac{1}{E[k]} \sum_{\boldsymbol{x} \in D} p(x_j \mid \boldsymbol{k}, \boldsymbol{\theta}) x_j.$$
(30)

Then, we can rewrite Eqs. 10 and 25 as:

$$\mu_{j,k} := \bar{x}_{j,k}$$
  
$$\mu_{j,k} := \frac{\tau \mu_0 + E[k] \bar{x}_{j,k}}{\tau + E[k]}$$

where the former (resp. the latter) is the ML (resp. MAP) estimate of the mean of  $X_j$  given the class k. It is easily seen that the latter is an weighted average of the former and  $\mu_0$ , where the weights are E[k] and  $\tau$ , respectively. So if we set  $\tau$  and  $\mu_0$ appropriately, the latter will show a stable behavior even for a small dataset. Also in Eq. 28 with an appropriate  $\Sigma_0$ , we can prevent the updated  $\Sigma_{j,k}$  from being singular.

#### 3.1.3 Membership distribution

Given the estimated parameters  $\hat{\theta}$  ( $\hat{\theta}_{ML}$  or  $\hat{\theta}_{MAP}$ ) and an attribute vector  $\boldsymbol{x}$  of some object, we can obtain a *membership distribution*  $p(k \mid \boldsymbol{x}, \boldsymbol{\theta})$  under which the object belongs to the class k. The probabilities are computed by:

$$p(k \mid \boldsymbol{x}, \hat{\boldsymbol{\theta}}) = \frac{p(k, \boldsymbol{x} \mid \hat{\boldsymbol{\theta}})}{p(\boldsymbol{x} \mid \hat{\boldsymbol{\theta}})} \propto p(k, \boldsymbol{x} \mid \hat{\boldsymbol{\theta}}).$$
(31)

Since we often consider that a class summarizes some (hidden) characteristic of an object, membership distributions play an important role in probabilistic inferences based on naive Bayes models.

In addition, for an attribute (vector)  $X_j$ , we may consider the *attribute membership distribution*  $p(k | x_j)$ , which is computed as follows:

$$p(k \mid x_j, \hat{\theta}) \propto p(k \mid \hat{\theta}) p(x_j \mid k, \hat{\theta}).$$
(32)

In inferences using attribute membership distributions (described in §3.1.4 – §3.1.6), NBCT makes different treatments between discrete and continuous attributes. That is, for a *discrete* attribute  $X_j$ ,  $p(k | x_j, \hat{\theta})$  is taken into account for each class k and each possible value  $1 \le x_j \le V_j$ . For each *continuous* attribute (vector)  $X_{j'}$ , on the other hand, we cannot enumerate all possible values, and thus  $p(k | x_{j'}, \hat{\theta})$  is taken into account for each class k and each value  $x_{j'}$  appeared in the dataset D.

#### 3.1.4 Clustering

As is mentioned above, clustering is a task to partition the objects in the dataset D into clusters of similar ones. One way

is to classify each object, whose attribute values are x, into its most probable cluster (class)  $k^*$ . More specifically, based on the estimated parameters  $\hat{\theta}$ , we compute  $k^*$  by using a membership probability as a score for clustering:

$$k^* = \underset{k:1 \le k \le K}{\operatorname{argmax}} p(k \mid \boldsymbol{x}, \hat{\boldsymbol{\theta}}) = \underset{k:1 \le k \le K}{\operatorname{argmax}} p(k, \boldsymbol{x} \mid \hat{\boldsymbol{\theta}}).$$
(33)

In this sense, naive Bayes clustering can be seen as an *unsupervised* classification task based on a naive Bayes model.

Some may be interested in clustering of attribute values for each attribute. Given an attribute value  $x_j$  of the *j*-th attribute (vector), the most probable cluster is predicted as follows:

$$k^* = \underset{k:1 \le k \le K}{\operatorname{argmax}} p(k \mid x_j, \hat{\theta}) = \underset{k:1 \le k \le K}{\operatorname{argmax}} p(k, x_j \mid \hat{\theta})$$
(34)

#### 3.1.5 Relevance analysis

Using the estimated parameters  $\hat{\theta}$ , we may want to know the most relevant objects to the class *k* of interest. One promising way is to rank the objects  $\mathbf{x}^{(1)}, \mathbf{x}^{(2)}, \ldots$  according to the magnitude of  $R(k, \mathbf{x}) \stackrel{\text{def}}{=} p(k \mid \mathbf{x})$ , i.e. the membership probability [21]. To understand this way of ranking, let us transform the probability as follows:

$$R(k, \mathbf{x}) = p(k \mid \mathbf{x}, \hat{\boldsymbol{\theta}})$$
  
=  $\frac{p(k \mid \hat{\boldsymbol{\theta}})p(\mathbf{x} \mid k, \hat{\boldsymbol{\theta}})}{p(\mathbf{x} \mid \hat{\boldsymbol{\theta}})} \propto \frac{p(\mathbf{x} \mid k, \hat{\boldsymbol{\theta}})}{p(\mathbf{x} \mid \hat{\boldsymbol{\theta}})}$  (35)

(note that *k* is fixed here). Now  $R(k, \mathbf{x})$  can be seen as a relevance score, because it indicates the significance of  $p(\mathbf{x} | k, \hat{\theta})$  compared to  $p(\mathbf{x} | \hat{\theta}) = \sum_{k=1}^{K} p(k | \hat{\theta}) p(\mathbf{x} | k, \hat{\theta})$ , the unconditional (or averaged) probability of  $\mathbf{x}$  being occurred. In some applications, the highly relevant objects would be a help for characterizing the cluster [12, 21]. One may find here that, for a fixed *k*, the relevance score  $R(k, \mathbf{x})$  yields the same ranking (among different  $\mathbf{x}$ 's) as that via the *pointwise* (or *pairwise*) *mutual information* [6] which is defined as follows:

$$I(k, \mathbf{x}) \stackrel{\text{def}}{=} \log \frac{p(\mathbf{x}, k \mid \hat{\boldsymbol{\theta}})}{p(\mathbf{x} \mid \hat{\boldsymbol{\theta}})p(k \mid \hat{\boldsymbol{\theta}})} = \log \frac{p(\mathbf{x} \mid k, \hat{\boldsymbol{\theta}})}{p(\mathbf{x} \mid \hat{\boldsymbol{\theta}})}.$$
(36)

One (subtle) practical advantage of  $R(k, \mathbf{x})$  over  $I(k, \mathbf{x})$  is that it is normalized (i.e.  $0 \le R(k, \mathbf{x}) \le 1$  by definition). For example, we may collect  $\{\mathbf{x} \mid R(k, \mathbf{x}) \ge \rho\}$  as the relevant objects to the class k, where  $\rho$  is a threshold *common* to all classes ( $0 < \rho < 1$ ).

Furthermore, we can consider the attribute-wise version of  $R(k, \mathbf{x})$ . That is,  $R^j(k, x_j)$  indicates the relevance between the class k and the value  $x_j$  of the attribute (vector)  $X_j$ , and is defined as  $R^j(k, x_j) = p(k | x_j, \hat{\theta})$ . This score measures the degree of relevance between a particular class and a particular attribute value, and so would be useful in the case where we are interested in the behavior of each attribute (e.g. in distributional clustering [18]). It should be addressed that AutoClass uses the attribute-wise version of  $I(k, \mathbf{x})$  to measure the 'influence' of  $x_j$  to the class k, so essentially NBCT yields the same rankings of attribute values as those by AutoClass.

#### 3.1.6 Dissimilarity

Since the membership distributions (§3.1.3)  $p(k \mid x, \hat{\theta})$  can be considered to characterize the object represented by x, we measure the dissimilarity  $\Delta$  between two objects x and x' by the Kullback-Liebler divergence between the corresponding membership distributions:

$$\Delta(\mathbf{x}, \mathbf{x}') \stackrel{\text{def}}{=} \operatorname{KL}(p(k \mid \mathbf{x}, \hat{\boldsymbol{\theta}}) \parallel p(k \mid \mathbf{x}', \hat{\boldsymbol{\theta}}))$$
$$= \sum_{k=1}^{K} p(k \mid \mathbf{x}, \hat{\boldsymbol{\theta}}) \log \frac{p(k \mid \mathbf{x}, \hat{\boldsymbol{\theta}})}{p(k \mid \mathbf{x}', \hat{\boldsymbol{\theta}})}.$$
(37)

In a similar way, the attribute-wise version of the dissimilarity is given as  $\Delta_j(x_j, x'_j) \stackrel{\text{def}}{=} \text{KL}(p(k \mid x_j, \hat{\theta}) \parallel p(k \mid x'_j, \hat{\theta})).$ 

## 3.1.7 Model selection

In clustering, we are often in question of how to determine the number of clusters. This is a problem of model selection, and in NBCT, we attempt to find a solution in a Bayesian approach. To be specific, we first consider the joint distribution  $p(D, M, \theta)$  of complete data D, a probabilistic model M, and its parameters  $\theta$ .  $p(D, M, \theta)$  is factored as  $p(D | M, \theta)p(\theta | M)p(M)$  by the chain rule, where p(M) is the prior distribution of the model M,  $p(\theta | M)$  is the prior distribution of the model M, and  $p(D | M, \theta)$  is the likelihood of the data D based on the model M with the parameters  $\theta$ . In naive Bayes models, for instance, each  $d \in D$  corresponds to the attribute vector x of an object. Here, M corresponds to K, the number of classes (clusters). From the settings above, our goal is to find the most probable model  $M^*$  based on the data D at hand, that is, we aim to find  $M^*$  such that:

$$M^{*} = \operatorname{argmax}_{M} p(M \mid D)$$
  
=  $\operatorname{argmax}_{M} \frac{p(D \mid M)p(M)}{p(D)}$   
=  $\operatorname{argmax}_{M} p(D \mid M),$  (38)

where we assume p(M) to be uniform for simplicity. Now the goal is reduced to finding  $M (= M^*)$  that maximizes p(D | M). p(D | M) is commonly called the *marginal likelihood* of D given M, and is used as a score for model selection. The marginal likelihood can be interpreted as the expectation (or the average) of the likelihood  $p(D | M, \theta)$  with respect to the prior distribution  $p(\theta | M)$ :

$$p(D \mid M) = \int_{\Theta} p(D, \theta \mid M) d\theta$$
  
= 
$$\int_{\Theta} p(D \mid M, \theta) p(\theta \mid M) d\theta$$
  
= 
$$\langle p(D \mid M, \theta) \rangle_{p(\theta \mid M)} .$$
(39)

If the dataset were complete data  $D_c$ , where each  $d \in D_c$  is a pair  $(k, \mathbf{x})$  of the attribute vector  $\mathbf{x}$  of an object and the class k to which the object belongs, then  $p(D_c \mid M)$  is obtained in closed form (see [7, 11] for the case with Bayesian networks).

On the other hand, when the data is incomplete, as in the case of probabilistic clustering, the integral in Eq. 39 is difficult to compute. So, including MCMC (Markov chain Monte Carlo) sampling, several approximation methods of log of the marginal likelihood are proposed so far [5]. *Bayesian information criterion* (BIC) [20] should be the most popular 'deterministic' approximation method, in which Laplace approximation is introduced based on the large-data assumption. The *Cheeseman-Stutz score* [4, 5] is used in AutoClass. The general forms of these two scores are respectively written as follows:

$$Score_{\rm BIC}(M) \stackrel{\text{def}}{=} p(D \mid M, \hat{\theta}_{\rm MAP}) - \frac{|\theta|}{2} \log N \qquad (40)$$

$$Score_{CS}(M) \stackrel{\text{def}}{=} p(\tilde{D}_c \mid M) - p(\tilde{D}_c \mid M, \hat{\theta}_{MAP}) + p(D \mid M, \hat{\theta}_{MAP}), \quad (41)$$

where N is the total size of dataset,  $|\theta|$  denotes the number of free parameters, and  $\tilde{D}_c$  is pseudo complete data whose sufficient statistics are the expected statistics obtained in the E-step of the EM algorithm.

## 3.1.8 Random restarts

Since the EM algorithm is a hill-climbing algorithm, being trapped in undesirable *local maxima* is known as one of practical problems in the EM algorithm. NBCT provides three facilities for avoiding such local maxima — random restarts [19], the deterministic annealing EM (DAEM) algorithm [22], and the split-merge EM (SMEM) algorithm [23].

In random restarts, we first prepare *n* different initial parameter sets. Then, from each initial parameter set, we run a series of EM iterations, and record the converged likelihood  $p(D \mid \theta)$  or the a posteriori probability  $p(\theta \mid D)$ . Finally we pick up the estimated parameters that bring the highest likelihood or the highest a posteriori probability.

#### 3.1.9 Deterministic annealing EM algorithm\*

Since the final estimate of the parameters depends on the choice of initial parameters, in the DAEM algorithm, we attempt to reduce an undesirable influence from the initial parameters in the early stage of EM iterations. To achieve this, by an analogy to statistical mechanics, the free energy is introduced first as:

$$\mathcal{F}_{\beta} = -\frac{1}{\beta} \log \sum_{k=1}^{K} p(k, \boldsymbol{x} \mid \boldsymbol{\theta})^{\beta}, \qquad (42)$$

where  $\beta$  is the *inverse temperature* which controls the influence from the initial parameters. Then we can obtain the DAEM algorithm, which tries to minimize the free energy  $\mathcal{F}_{\beta}$  at each temperature  $\frac{1}{\beta}$ . Fig. 5 shows an expected behavior of the DAEM algorithm, where  $L_{\beta}$  is introduced as  $-\mathcal{F}_{\beta}$  (then we will try to maximize  $L_{\beta}$ ). In the DAEM algorithm, we start from the small  $\beta$ , under which the free energy is expected to have a smooth shape, and hopefully has only one local maximum (i.e. the

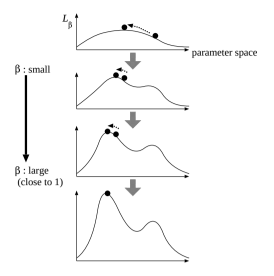


Figure 5: Image of the deterministic annealing EM algorithm.

- 1. Initialize randomly the parameters  $\theta_k$   $(1 \le k \le K)$  and  $\theta_{j,k,x_j}$  $(1 \le k \le K, 1 \le j \le J, 1 \le x_j \le V_j).$
- 2. Repeat Step 3 for each  $\beta = \beta_0, \beta_1, \beta_2, \dots, 1.0$
- 3. Repeat the following until the log-likelihood log  $p(D \mid \theta)$  converges:

$$p(k \mid \boldsymbol{x}, \boldsymbol{\theta}) \propto p(k \mid \boldsymbol{\theta})^{\beta} \prod_{j=1}^{J} p(x_j \mid k, \boldsymbol{\theta})^{\beta}.$$
 (43)

(The rest are the same as those in Fig. 3, the original EM algorithm.)

Figure 6: The DAEM algorithm for ML estimation in naive Bayes models.

global maximum). So under the smaller  $\beta$ , we may be able to find the global maximum or a good local maximum. When  $\beta$  increases, on the other hand, the shape of the free energy changes (becomes sharper), and hence we should continue to update the parameters by EM iterations. However please note that the starting point of these EM iterations is expected to be more promising than the initial parameters. Finally we perform EM iterations at  $\beta = 1$ , which is equivalent to the usual EM iterations.

To be more concrete, we show the DAEM algorithm in naive Bayes models in Fig. 6. For an effective use of DAEM algorithm, the temperature schedule is important. In NBCT, following [22], we start from  $\beta_0 = \beta_{init}$  and then update  $\beta$  by the update rule  $\beta_{t+1} \leftarrow \beta_t \cdot \beta_{rate}$ , where  $\beta_{init}$  and  $\beta_{rate}$  are given by the user (the default values are 0.1 and 1.2, respectively).

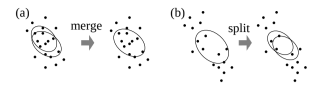


Figure 7: Image of a merge operation and a split operation, where the dots and the ovals stand for objects and clusters, respectively.

#### 3.1.10 Split-merge EM algorithm\*

The SMEM algorithm is applicable to mixture models, and with this algorithm, we attempt to escape from local maxima by forcedly applying the *split operation* and the *merge operation* to unpromising clusters.

To be more specific, we merge two clusters, say  $k_1$  and  $k_2$ , that closely overlap with each other (Fig. 7 (a)), and split a cluster, say  $k_3$ , that excessively covers objects (Fig. 7 (b)). In the paper that firstly proposed the SMEM algorithm [23], the split operation and the merge operation are always paired, so the number of resulting clusters will not change. After a splitmerge operation pair executed, the EM algorithm is conducted until the convergence of likelihood  $p(D | \theta)$  or the a posteriori probability  $p(\theta | D)$ .

The possible triplets of clusters  $(k_1, k_2, k_3)$  are kept as prioritized candidates for the next split-merge operation. If the converged likelihood or a posteriori probability is improved with the first candidate (i.e. the candidate is said to be *accepted*), we will proceed to further split-merge operations. If there is no (significant) improvement with the first candidate, we will discard the result of the EM algorithm (i.e. the candidate is said to be *rejected*) and try the next candidate. In the sequel, for making simpler descriptions, we will concentrate on the case that the attributes are all discrete.

#### Prioritizing the split-merge candidates

To prioritize the split-merge candidates, we first get the pairs  $\{(k_1, k_2) \mid 1 \le k_1 < k_2 \le K\}$  of classes to be merged, in the descending order of heuristic scores  $J_{\text{merge}}(k_1, k_2 \mid \theta)$ . The score is defined as follows:

$$J_{\text{merge}}(k_1, k_2 \mid \boldsymbol{\theta}) \stackrel{\text{def}}{=} \mathbf{p}_{k_1}(\boldsymbol{\theta})^T \mathbf{p}_{k_2}(\boldsymbol{\theta})$$
$$= \sum_{\boldsymbol{x}} N(\boldsymbol{x}) p(k_1 \mid \boldsymbol{x}, \boldsymbol{\theta}) p(k_2 \mid \boldsymbol{x}, \boldsymbol{\theta}),$$
(44)

where we have a multiset of objects  $D = \{x^{(1)}, x^{(2)}, \dots, x^{(N)}\}$  as the observed data, and  $\mathbf{p}_k(\boldsymbol{\theta})$  is the vector of the membership probabilities to the class k:

$$\mathbf{p}_k(\boldsymbol{\theta}) = (p(k \mid \boldsymbol{x}^{(1)}, \boldsymbol{\theta}), p(k \mid \boldsymbol{x}^{(2)}, \boldsymbol{\theta}), \dots, P(k \mid \boldsymbol{x}^{(N)}, \boldsymbol{\theta}))^T.$$
(45)

Intuitively,  $J_{\text{merge}}(k_1, k_2 | \theta)$  measures a (partially empirical) similarity between the classes  $k_1$  and  $k_2$  based on the data D.

Besides, we may use the normalized version:

$$\tilde{J}_{\text{merge}}(k_1, k_2 \mid \boldsymbol{\theta}) \stackrel{\text{def}}{=} \frac{\mathbf{p}_{k_1}(\boldsymbol{\theta})^T \mathbf{p}_{k_2}(\boldsymbol{\theta})}{\|\mathbf{p}_{k_1}(\boldsymbol{\theta})\| \cdot \|\mathbf{p}_{k_2}(\boldsymbol{\theta})\|}.$$
(46)

Then, for each  $(k_1, k_2)$  pair to be merged, we also get the classes  $\{k_3 \mid 1 \le k_3 \le K, k_3 \ne k_1, k_3 \ne k_2\}$  to be split, in the descending order of another heuristic score  $J_{\text{split}}(k_3 \mid \theta)$ :

$$J_{\text{split}}(k_3 \mid \boldsymbol{\theta}) \stackrel{\text{def}}{=} \text{KL}(\tilde{p}(\boldsymbol{x} \mid k_3, \boldsymbol{\theta}) \parallel p(\boldsymbol{x} \mid k_3, \boldsymbol{\theta}))$$
$$= \sum_{\boldsymbol{x}} \tilde{p}(\boldsymbol{x} \mid k_3, \boldsymbol{\theta}) \log \frac{\tilde{p}(\boldsymbol{x} \mid k_3, \boldsymbol{\theta})}{p(\boldsymbol{x} \mid k_3, \boldsymbol{\theta})}, \quad (47)$$

where  $\tilde{p}(\mathbf{x} | k_3, \boldsymbol{\theta})$  is a local empirical probability computed by:

$$\tilde{p}(\boldsymbol{x} \mid k_3, \boldsymbol{\theta}) \propto \tilde{p}(\boldsymbol{x}) p(k_3 \mid \boldsymbol{x}, \boldsymbol{\theta}) \propto N(\boldsymbol{x}) p(k_3 \mid \boldsymbol{x}, \boldsymbol{\theta}).$$
(48)

In the above,  $\tilde{p}(\mathbf{x})$  denotes  $N(\mathbf{x})/N$ , the empirical unconditional probability of  $\mathbf{x}$ . So using  $J_{\text{split}}(k_3 | \boldsymbol{\theta})$ , the class that does not fit to the data will be split earlier.

#### ♦ Partial EM iterations

After a split-merge operation, we should re-initialize the parameters of the modified classes. Let  $k'_1$  be the new class obtained by merging  $k_1$  and  $k_2$ . Also we consider two classes  $k'_2$  and  $k'_3$ which are obtained by splitting  $k_3$ . Then, for the merged class  $k'_1$ , we re-initialize the parameters related to  $k'_1$  as follows:

$$\theta_{k_1'} := \theta_{k_1} + \theta_{k_2} \tag{49}$$

$$\theta_{j,k'_1,x_j} := \frac{E[k_1]\theta_{j,k_1,x_j} + E[k_2]\theta_{j,k_2,x_j}}{E[k_1] + E[k_2]}$$
(50)

On the other hand, for the split classes  $k'_2$  and  $k'_3$ , we re-initialize  $\theta_{k'_2} := \frac{1}{2}\theta_{k_3}, \ \theta_{k'_3} := \frac{1}{2}\theta_{k_3}, \ \theta_{j,k'_2,x_j} := \theta_{j,k_3,x_j} + \epsilon$  and  $\theta_{j,k'_3,x_j} := \theta_{j,k_3,x_j} + \epsilon'$ , where  $\epsilon$  and  $\epsilon'$  are some random noises.

In partial EM iterations, to make it reasonable to compute Estep and M-step only for the modified classes  $(k'_1, k'_2 \text{ and } k'_3)$ , M-step is modified as follows  $(k' = k'_1, k'_2, k'_3)$ :

$$\theta_{j,k',x_j} = \frac{E[k']}{\sum_{k''=k'_1,k'_2,k'_3} E[k'']} \cdot \sum_{k''=k'_1,k'_2,k'_3} \theta_{j,k'',x_j}$$
(51)

#### **\diamond** Behavior of the SMEM algorithm

Fig. 8 shows a typical pattern on the changes in log-likelihood by the SMEM algorithm, where we apply five split-merge operations. Note here that we have only plotted a sequence with the accepted split-merge operations. It is seen from Fig. 8 that the log-likelihood decreases at the moment of applying a splitmerge operation, but as a whole, the log-likelihood steadily increases, and finally we can obtain the estimates with a higher log-likelihood than the one obtained by the usual EM algorithm.

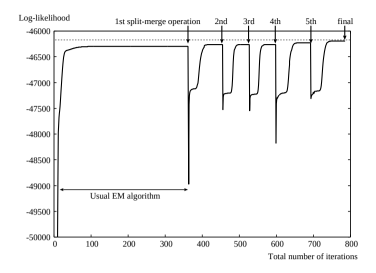


Figure 8: Changes in log-likelihood by the SMEM algorithm.

## 3.2 VB based clustering

## 3.2.1 Model selection based on VB

As described in §3.1.7, to obtain the model  $M^*$  that explains best the data D at hand, we consider  $M = M^*$  is the model that maximizes the marginal likelihood p(D | M). In naive Bayes models, the optimal number  $K^*$  of classes is the number of classes in  $M^*$ . It has been known that if D is complete data  $D_c$ , p(D | M) can be obtained in closed form. However, when D is incomplete, i.e. there is some hidden data z such that  $D_c = (D, z)$  (for instance, in naive Bayes clustering, the classes of the objects are hidden in D), some approximation method is required. In this section, we briefly describe the approximation via the VB approach.

First, let us consider log of the marginal likelihood  $L(D) \stackrel{\text{def}}{=} \log p(D \mid M)$ , and then we have:

$$L(D) = \log \sum_{z} \int_{\Theta} p(D, z, \theta \mid M) d\theta$$
  
= 
$$\log \sum_{z} \int_{\Theta} q(z, \theta \mid D, M) \frac{p(D, z, \theta \mid M)}{q(z, \theta \mid D, M)} d\theta$$
  
$$\geq \sum_{z} \int_{\Theta} q(z, \theta \mid D, M) \log \frac{p(D, z, \theta \mid M)}{q(z, \theta \mid D, M)} d\theta.$$
  
(from Jensen's inequality) (52)

For the space limitation, we fix the model *M* for the moment, and simply write  $p(\cdot | M) = p(\cdot)$  and  $q(\cdot | D, M) = q(\cdot | D)$ , and then obtain:

$$L(D) \geq F[q]$$

$$\stackrel{\text{def}}{=} \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log \frac{p(D, z, \theta)}{q(z, \theta \mid D)} d\theta, \quad (53)$$

where F[q] can be seen as a lower limit of L(D), and is called

the *variational* (or *negative*) *free energy*. So to get a good approximation of L(D), we find a distribution function  $q = q^*$  that maximizes a functional F[q]. In model selection, we use the free energy F[q] as a model score.

Besides, to get another view, we have the following by considering  $L(D) = \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log p(D) d\theta$ :

$$L(D) - F[q]$$

$$= \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log \left\{ p(D) \cdot \frac{q(z, \theta \mid D)}{p(D, z, \theta)} \right\} d\theta$$

$$= \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log \frac{q(z, \theta \mid D)}{p(z, \theta \mid D)} d\theta$$

$$= \operatorname{KL}(q(z, \theta \mid D) \parallel p(z, \theta \mid D)). \quad (54)$$

Here, maximizing F[q] implies minimizing the Kullback-Liebler divergence between  $q(z, \theta \mid D)$  and  $p(z, \theta \mid D)$ . So finding  $q^*$  is to make a good approximation of  $p(z, \theta \mid D)$ , the conditional distribution of hidden variables and the parameters.

In VB learning, we further assume  $q(z, \theta \mid D) \approx q(z \mid D)q(\theta \mid D)$ , and obtain a generic form of *variational Bayesian EM (VB-EM) algorithm* as an iterative procedure consisting of the following two updating rules:

. .

$$q(z \mid D) \propto \exp\left(\int_{\Theta} q(\theta \mid D) \log p(D, z \mid \theta) d\theta\right), \quad (55)$$
  
$$q(\theta \mid D) \propto p(\theta) \exp\left(\sum_{z} q(z \mid D) \log p(D, z \mid \theta)\right). \quad (56)$$

Now we can derive a VB-EM algorithm specific to naive Bayes clustering, shown in Fig. 9, by substituting the distribution form of a naive Bayes model (Eq. 1 and the predefined distributions for attributes) to the generic VB-EM procedure above. For simplicity, we assume in Fig. 9 that the attributes are all discrete. For the descriptions of the VB-EM algorithm for Gaussian mixture models, please see [17, 24].<sup>10</sup> In Fig. 9,  $\pi_k$  and  $\pi_{i,k,x_i}$  are defined as follows:

$$\pi_k \stackrel{\text{def}}{=} \exp\left(\Psi\left(\alpha_k\right) - \Psi\left(\sum_{k'=1}^K \alpha_{k'}\right)\right),\tag{57}$$

$$\pi_{j,k,x_j} \stackrel{\text{def}}{=} \exp\left(\Psi\left(\alpha_{j,k,x_j}\right) - \Psi\left(\sum_{x_j'=1}^{V_j} \alpha_{j,k,x_j'}\right)\right), \quad (58)$$

where  $\Psi(\cdot)$  is the digamma function. One may find that the resulting VB-EM algorithm receives the data *D* as input, and outputs the learned hyperparameters  $\alpha^*$  (or the learned pseudo counts  $\delta^*$ ). In NBCT, the initial hyperparameters such as  $\alpha_k^{(0)}$  are obtained by adding small random noises. For instance, we obtain  $\alpha_k^{(0)}$  by:

$$\alpha_k^{(0)} := \alpha_k + \epsilon_k \quad , \tag{59}$$

where  $\alpha_k$  is the quantity specified by the user, and  $\epsilon_k$  is a small random noise.

- 1. Initialize randomly the hyperparameters  $\alpha_k^{(0)}$   $(1 \le k \le K)$  and  $\alpha_{j,k,x_i}^{(0)}$   $(1 \le k \le K, 1 \le j \le J, 1 \le x_j \le V_j)$ .
- 2. Repeat the following steps until the free energy F[q] converges:

$$q(k \mid \mathbf{x}) \quad \propto \quad \pi_k \prod_{j=1}^J \pi_{j,k,x_j} \tag{60}$$

$$E[k] := \sum_{\mathbf{x}} N(\mathbf{x})q(k \mid \mathbf{x})$$
(61)

$$E_j[k, x_j] \quad := \quad \sum_{\boldsymbol{x}' \in D: \boldsymbol{x}'_j = x_j} N(\boldsymbol{x}') q(k \mid \boldsymbol{x}') \tag{62}$$

$$\alpha_k := \alpha_k^{(0)} + E[k] \tag{63}$$

$$\alpha_{j,k,x_j} := \alpha_{j,k,x_j}^{(0)} + E_j[k,x_j]$$
(64)

Figure 9: The VB-EM algorithm in naive Bayes models.

## 3.2.2 Clustering

As in the ML/MAP case (§3.1.4), we may regard the cluster  $k^*$  which has the highest membership probability for an object x as the cluster to which the object belongs. The difference is that we use the distribution  $p(k, x) = \int_{\Theta} q^*(\theta \mid D)p(k, x \mid \theta)d\theta$ , an averaged distribution on the prior distribution  $q^*(\theta \mid D)$  with learned hyperparameters  $\alpha^*$ , instead of  $p(k, x \mid \hat{\theta})$  which relies on the point-estimated value  $\hat{\theta}$ . That is, we have:

$$k^{*} = \operatorname{argmax}_{k:1 \le k \le K} p(k \mid \mathbf{x}) = \operatorname{argmax}_{k:1 \le k \le K} p(k, \mathbf{x})$$
$$= \operatorname{argmax}_{k:1 \le k \le K} \int_{\Theta} q^{*}(\boldsymbol{\theta} \mid D) p(k, \mathbf{x} \mid \boldsymbol{\theta}) d\boldsymbol{\theta}$$
$$= \operatorname{argmax}_{k:1 \le k \le K} p^{*}(k) \prod_{j=1}^{J} p^{*}(x_{j} \mid k) , \qquad (65)$$

where  $p^*$  is the *predictive distribution*. That is, for the class *C* and each discrete attribute  $X_j$ , the predictive distributions are obtained in closed form:

$$p^{*}(k) = \alpha_{k}^{*} / \sum_{k'} \alpha_{k'}^{*}$$
 (66)

$$p^{*}(x_{j} \mid k) = \alpha^{*}_{j,k,x_{j}} / \sum_{x'_{j}} \alpha^{*}_{j,k,x'_{j}} .$$
(67)

On the other hand, for continuous attributes, each  $p^*(x_j | k)$  is analytically obtained as the Student's *t*-distribution.

#### 3.2.3 Other inference tasks

In VB learning, it seems not easy in straightforward ways to conduct the probabilistic inferences other than clustering based on the membership distribution (§3.2.2) — relevance analysis (§3.1.5) or computing dissimilarities (§3.1.6). One compromise is to use the probabilities  $p^*$  obtained by Eqs. 66 and 67 instead

<sup>&</sup>lt;sup>10</sup>Precisely speaking, the VB-EM algorithm implemented in NBCT is based on the author's derivation, and is a bit different from the ones in these papers (e.g. some coefficients of terms in the update rules).

of  $\hat{\theta}$ , the point-estimated parameters in ML/MAP [3], and actually NBCT adopts this way. It is important to note however that the resulting probabilistic inferences might *not* be good approximations of the inferences based on a posteriori quantities.

## 3.2.4 Split-merge EM algorithm\*

Basically, the SMEM algorithm for VB-EM (say, VB-SMEM) follow almost the same procedures as the ones for the ML/MAP-EM algorithms (§3.1.8 and §3.1.10). The difference is that we need to use the probabilities  $p^*$  obtained by Eqs. 66 and 67 to compute two heuristic scores  $J_{\text{merge}}$  and  $J_{\text{split}}$  (Eqs. 44–47).

#### 3.2.5 Deterministic annealing EM algorithm\*

Since version 0.3, the deterministic-annealing version of VB-EM has been introduced. To be specific, following [14], let us revisit the definition of the variational free energy (Eq. 53):

$$F[q] \stackrel{\text{def}}{=} \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log \frac{p(D, z, \theta)}{q(z, \theta \mid D)} d\theta$$
$$= \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log p(D, z, \theta) d\theta$$
$$- \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log q(z, \theta \mid D) d\theta \quad (68)$$

Again, by an analogy to statistical mechanics, we correspond F[q] with  $-\mathcal{F}$  ( $\mathcal{F}$ : the free energy), the first term in the right hand side of Eq. 68 with  $-\mathcal{U}$  ( $\mathcal{U}$ : the internal energy) and the second term with S (S: the entropy). Then we newly introduce the variational free energy that takes into account the (inverse) temperature:

$$F_{\beta}[q] \stackrel{\text{def}}{=} \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log p(D, z, \theta) d\theta$$
$$-\frac{1}{\beta} \sum_{z} \int_{\Theta} q(z, \theta \mid D) \log q(z, \theta \mid D) d\theta.$$
(69)

The VB-EM algorithm that tries to maximize  $F_{\beta}[q]$  (i.e. the DAversion of the VB-EM algorithm) has a similar procedure to that of the DAEM algorithm for ML/MAP estimation.

# 4 How to use NBCT

## 4.1 Overall organization of NBCT

NBCT 0.4.2 provides three executables named nbc, vnbc and nbcsep:

• nbc covers probabilistic inferences for ML/MAP-based clustering (§3.1).

- vnbc covers probabilistic inferences for VB-based clustering (§3.2).
- nbcsep is an auxiliary tool for post-processing of the output files from nbc or vnbc.

See §4.2 for the installation procedure. The usages of nbc, vnbc and nbcsep are described in §4.3, §4.4 and §4.5, respectively.

#### 4.2 Installation

#### 4.2.1 Preliminary installation (GNU libiconv)

To handle the dataset with non-ASCII-safe character encodings, NBCT provides a way to utilize GNU libiconv (http://www.gnu.org/software/libiconv/). If you are sure that the target dataset only contains ASCII characters or is ASCII-safe, GNU libiconv will *not* be required. Otherwise and if GNU libiconv is not installed in your environment, you need to install GNU libiconv in advance of the installation of NBCT.

On Windows, a DLL file iconv.dll should be placed in the folder specified by the PATH environment variable, or in one of the system folders such as c:\WINDOWS\system32. To obtain iconv.dll, we may download http://ftp.gnu.org/pub/gnu/ libiconv/libiconv-1.9.1.bin.woe32.zip, which includes a couple of executables and DLL files already built.<sup>11</sup> On the other hand, the latest version can be built from the source files.<sup>12</sup>

#### 4.2.2 Contents of the package

The original package of NBCT is distributed as a source tarball (nbct-0.4.2.tar.gz) or a zip archive (nbct-0.4.2\_win.zip) including Windows executables, compiled by MSVC++ 9.0. After the package unfolded, you may find the following subdirectories (or sub-folders):

- src/ contains C source code.
- doc/ contains the document files including this manual.
- bin/ contains executables or script files (but might be empty just after unfolding).
- example/ contains data examples.

For the users who use the executables in nbct-0.4.2\_win.zip on Windows, the installation is quite easy — we only need to place these executables in the

<sup>&</sup>lt;sup>11</sup>It is confirmed that the NBCT binaries built using MSVC++ 9.0 can run with iconv.dll from libiconv-1.9.1.bin.woe32.

 $<sup>^{12}</sup>$ According to README.woe32 in the released package of GNU libiconv 1.12 (the latest version as of Dec. 2008), building requires MinGW, and MSVC++ is no longer supported.

folder that appears in the PATH environment variable. However it should be noted that the executables in nbct-0.4.2\_win. zip do not link with GNU libiconv, and so are only applicable to the datasets with ASCII-safe characters.

## 4.2.3 Using configure

For Linux, Mac OS X, and Win32 with Cygwin or MinGW, we can use the configure script in the package. That is, after moving to the src directory, just type:

```
./configure Options make
```

where *Options* are some appropriate setting to your system. For the details on *Options*, please consult the INSTALL file in the src directory. Also,

```
./configure --help
```

will show the details of the options for the configure script.

## 4.2.4 Using Makefile.msvc

To build NBCT on Win32 with MSVC++, we attempt to compile the source code by the cl command of MSVC++. Please follow the steps below:

- 1. Edit src/Makefile.msvc as suitable for your environment.
- Invoke the Command Prompt window prepared for MSVC++. For instance, if you are using MS Visual Studio 2008 on Windows XP, please follow the menus: [Start] → [All Programs] → [Microsoft Visual Studio 2008] → [Visual Studio Tools] → [Visual Studio 2008 Command Prompt].
- 3. At the Command Prompt invoked, visit the src folder.
- 4. Type the following command to compile NBCT:

```
nmake -f Makefile.msvc
```

5. Type the following command to install NBCT:

nmake -f Makefile.msvc install

By default (i.e. without modifying Makefile.msvc), all executables will be copied into the bin directory of the unfolded package.

## 4.3 ML/MAP based clustering

## 4.3.1 Overview

As is mentioned above, we use the executable nbc for ML/MAP based clustering. Roughly speaking, nbc conducts the EM algorithm first to estimate the parameters, and then, based on the estimated parameters, makes a couple of requested

Table 2: Input/Output files for nbc. *Base* is the base name, and *Ext* is txt (for the file in the tab-separated value format) or csv (for the file in the comma-separated value format).

	Filename	Content
Input	Base_freq.Ext	Occurrences of
		objects
Input/Output	Base_param.Ext	Parameters
Input	Base_hparam.Ext	Hyperparameters
Output	Base_cluster.Ext	Clustering results
Output	Base_memp.Ext	Membership probs.
Output	Base_rank.Ext	Result of
		relevance analysis
Output	Base_dissim.Ext	Dissimilarities
Output	Base_log.Ext	Logs
Output	Base_msg.Ext	Terminal messages

probabilistic inferences including clustering. We can pass our settings and tasks to nbc through the command line arguments:  $^{13}$ 

nbc -f Base -x NumAttr \ -k NumClass Options...

Here '-f Base' indicates that we have a file named Base \_freq.Ext containing the target dataset D. Base will work as the base name of the input/output files listed in Table 2. Ext is the file extension, which can be txt or csv according to the format (See §4.3.3). NumAttr, given with -x, indicates the number of attributes in Base\_freq.Ext (Note that the count column is not counted as an attribute). Typically we also specify the number of classes by -k option. The details of optional flags will be described in §4.3.5.

#### 4.3.2 Attribute types

By default, in nbc, all attribute values in the dataset D are considered to be discrete. So we need to specify (the indices of) continuous attributes explicitly, using -gauss (-u) option. Also we may have to specify the following attributes which can be included in real datasets:

- The *ID attribute*, which only identifies the object, but is irrelevant to the characteristics of the object.
- The *answer class attribute*, which is annotated by human for the purpose of cluster evaluation.
- *Ignored attributes*, whose values are recorded in the dataset, but are not used in clustering.

Since version 0.4, the answer class attribute and the ignored attributes are similarly treated — the only difference is that the

<sup>&</sup>lt;sup>13</sup>The symbol '\' just means a continuation of the command line, and is used here just for space limitation.

Figure 10: Common file format in NBCT.

answer classes are recorded into the output files, while the values of the ignored attributes are not. The information about the answer classes can be used for evaluating the obtained clusters, which is planned to be implemented in future.

The indices of these attributes are specified by -id(-I), -eval(-E) and -ignore(-y) options. See §4.3.5 for the details of these options.

## 4.3.3 File format

Fig. 10 illustrates the file format that is common to all NBCT's input/output files. In this format, a file contains several data matrices each having a header part which consists of the lines starting with '#'. A header part includes the identifier to the corresponding data matrix and gives a brief description on columns. Each data matrix follows the *tab-separated value format* or the *CSV format*, where a row in the matrix is separated into data cells by tab characters or commas, respectively. Note that the CSV format used here is just a restricted one — e.g. the current version of NBCT does not understand the double quote characters. Missing values, denoted by '?' or 'NA', are allowed to exist in the data matrix.

The input file format is automatically determined based on the file extension, and the files in the tab-separated value (resp. CSV) format must have the file extension txt (resp. csv). By default, NBCT first tries to read *Base\_freq.txt*,<sup>14</sup> and if the file does not exist, NBCT then tries to read *Base\_freq.csv* (or *Base\_param.csv*). So we only need to put the input files with the valid file extension in the current working directory. On the other hand, when '--input *Ext*' is given, NBCT reads *Base\_freq.Ext* (or *Base\_param.Ext*) only. The output file format is the same as the input file format by default, while we can change it by giving '--output *Ext*' or '-j *Ext*'.

#### 4.3.4 Compression of the indistinguishable objects

As described in §2.1, the objects having the same attribute values will be compressed into a single object unless they have the ID attribute or continuous attributes. So the number of distinct objects can be less than the number of lines in the data file *Base\_freq.Ext*. The number of occurrences of the resulting object is of course the sum of the numbers of occurrences of the distinct objects. On the other hand, we can inhibit such a compression by giving --uncompress (-U) option to the command.

## 4.3.5 Command line options

## Model specification (general)

--n-class NumClass

-k NumClass

By this option, we set *NumClass* to *K*, the number of classes. Given '-k *Kmin:Kmax:Kstep*', nbc will enter into the 'model scoring' mode (§3.1.7). That is, with varying the number *K* of classes from *Kmin* to *Kmax* with step *Kstep*, we evaluate the plausibility of *K* based on the score specified by --score(-g) option. This option is mandatory except when --read-param(-R) or --read-hparam(-H) is given.

-x NumAttr

This option tells nbc that the total number of attributes, *including* the ID attribute, the answer class attribute, ignored attributes (§4.3.2), is *NumAttr*.

--gauss ContAttrs

## -u ContAttrs

This option specifies the zero-based indices of continuous attributes. Basically we specify  $A_i$ , the *i*-th attribute, as continuous and single-dimensional by '-u *i*', whereas the *n* continuous attributes that jointly follow a multivariate Normal distribution is specified by '-u  $i_1, i_2, \ldots, i_n$ '. For two or more Normal distributions, we use this option for each of them. For example, '-u 1, 3 -u 2' declares that we have two Normal distributions, where the first corresponds to a bivariate Normal distribution the attributes  $(A_1, A_3)$  follow, and the second corresponds to a univariate Normal distribution followed by  $A_2$ . For convenience, nbc allows a variety of abbreviated forms:

- '-u 0-3' is the same as '-u 0, 1, 2, 3'.
- '-u -3' is the same as '-u 0-3'.
- '-u 2-' is the same as '-u 2, 3, ..., (T 1)', where T is the number of attributes specified by -x option.
- '-u 2-4, 8' is the same as '-u 2, 3, 4, 8'.

<sup>&</sup>lt;sup>14</sup>When --read-param (-R) option specified, NBCT first tries to read *Base\_param.txt*, and NBCT then tries to read *Base\_param.csv*.

- '-u 1#3, 4' is the same as '-u 1 -u 3, 4'.
- '-u 2~4' is the same as '-u 2#3#4', and hence as '-u 2 -u 3 -u 4'.

## --id *IdAttr*

-I IdAttr

This option specifies the zero-based index of the ID attribute.

## --ignore IgnoreAttrs

-y IgnoreAttrs

This option specifies the zero-based index (indices) of ignored attribute(s). If there are more than one ignored attribute, we can specify them using hyphen and comma. The meanings of hyphen and comma are the same as those for -gauss(-u) option, i.e. '-y 2–4, 8' will be interpreted as '-y 2, 3, 4, 8', and hence as '-y 2 –y 3 –y 4 –y 8'.

--eval AnsAttr

```
-E AnsAttr
```

This option specifies the zero-based index of the answer class attribute. The answer classes will be recorded into the output files. To introduce a facility for cluster evaluation is future work.

--read-param (no argument)

-R (no argument)

With this option, nbc will skip the EM algorithm and read the parameters from  $Base\_param.Ext$ , which may have been created by hand or by a previous run of nbc. With this option, the number of classes is then determined according to the content of  $Base\_param.Ext$  (so -n-class(-k) option will be ignored).

#### Model specification (hyperparameters)

On contrary to AutoClass, NBCT allows us to specify raw hyperparameters. In addition, for the cases where it is tedious to specify them, --autoclass (-A) option is provided to make a similar<sup>15</sup> configuration to AutoClass.

--smooth-c ClassPseudoCount

-c ClassPseudoCount

This option specifies a uniform pseudo count for the class variable. That is, for each class k, the pseudo count  $\delta_k$  is equally set to *ClassPseudoCount*. When 'auto' is specified for *ClassPseudoCount*, the pseudo counts are set as the ones used in AutoClass (§3.1.2). The default is 0.

--smooth-a AttrPseudoCount

```
-w AttrPseudoCount
```

This option specifies uniform pseudo counts for discrete attributes. If *AttrPseudoCount* is just a non-negative number  $\zeta$ , all pseudo counts  $\delta_{j,k,x_j}$  for discrete attributes are equally set as  $\zeta$ . If *AttrPseudoCount* takes a commaseparated form ' $\zeta_{j_1}$ ,  $\zeta_{j_2}$ ,  $\cdots$ ,  $\zeta_{j_M}$ ', where  $X_{j_m}$  is a discrete attribute and M is the number of discrete attributes, the pseudo counts  $\delta_{j_m,k,x_{j_m}}$  ( $1 \le x_{j_m} \le V_{j_m}$ ) are equally set as  $\zeta_{j_m}$ . When 'auto' is specified for  $\zeta$  or for some  $\zeta_{j_m}$ , the corresponding pseudo counts are set as the ones used in AutoClass (§3.1.2). The default values are all 0.

--tau *Tau* 

-т Таи

Basically this option sets *Tau* to  $\tau$ , a hyperparameter of the Normal-Gamma or the Normal-Wishart distribution (see §3.1.2). If *Tau* is given as 'auto', similarly to the pseudo counts for discrete attributes,  $\tau$  will be set as 1/K, where *K* is the number of classes. If this option is not given, ML estimation will be conducted for the continuous attributes.

--alpha Alpha

## -0 Alpha

This option sets *Alpha*/2 (resp. *Alpha*) to *a* (resp.  $\alpha$ ), a hyperparameter of the Normal-Gamma (resp. the Normal-Wishart) distribution (see §3.1.2). If this option is not given, *a* will be set as  $n_j/2 + \epsilon$  (resp.  $\alpha$  will be set as  $n_j + \epsilon$ ), where  $n_j$  is the dimension of an attribute (resp. attributes)  $X_j$ , and  $\epsilon$  is a small positive number.

--smooth-mean Mu



This option sets Mu to  $\mu_0$ , a hyperparameter of the Normal-Gamma or the Normal-Wishart distribution (see §3.1.2). If this option is not given, we use  $\hat{\mu}_j$  instead for attribute(s)  $X_j$ , where  $\hat{\mu}_j$  is the sample mean of  $X_j$  in the dataset D as if there is no latent class variable.

--smooth-var DiagElem

-L *DiagElem* 

This option sets *DiagElem* to  $\kappa$ , and then, *b* (resp.  $\Sigma_0$ ) will be set as  $\kappa/2$  (resp.  $\kappa I$ , where *I* is the identity matrix). Here *b* (resp.  $\Sigma_0$ ) is a hyperparameter of the Normal-Gamma (resp. the Normal-Wishart) distribution (see §3.1.2). If this option is not given, we use  $\hat{\sigma}_j^2/2$  (resp.  $\hat{\Sigma}_j$ ) instead for attribute (resp. attributes)  $X_j$ , where  $\hat{\sigma}_j^2$  (resp.  $\hat{\Sigma}_j$ ) is the sample variance (resp. covariances) of  $X_j$  in the dataset *D* as if there is no latent class variable.

--autoclass (no argument)

-A (no argument)

With this option, we can make a similar configuration to AutoClass. This option is equivalent to '--smooth-c auto --smooth-a auto --tau auto --score cs'.

 $<sup>^{15}</sup>$ To be precise, the configuration with --autoclass (-A) for continuous attributes is different from Autoclass, because AutoClass uses different prior distributions for continuous attributes.

--read-hparam (no argument) -H (no argument)

> With this option, nbc will run the MAP-based EM algorithm (§3.1.2) under the pseudo counts (the smoothing constants) read from a file named Base\_hparam. Ext, which may have been created by hand or by a previous run of vnbc (§4.4). The number of classes is determined according to the content of Base hparam. Ext (so --n-class (-k) option will be ignored).

## **\diamond EM algorithm**

The ML/MAP-based EM algorithms for naive Bayes models are described in §3.1.1 and §3.1.2.

--seed RandomSeed

-r RandomSeed

nbc will use RandomSeed as a random seed for initialization of parameters in the EM algorithm (Step 1 in Fig. 3 and Fig. 4).

--epsilon Threshold

```
-e Threshold
```

nbc will use *Threshold* as the threshold  $\xi$  for judging the convergence of the likelihood or the a posteriori probability (Step 2 in Fig. 3 and Fig. 4). That is, if the difference between the values of the log-likelihood (or log of the a posteriori probability) before the update and the one after the update becomes less than  $\xi$ , we will consider that the parameters have been converged. Since version 0.4, the default value has been changed to  $10^{-3}$ .

```
--restart NumInit
```

-n NumInit

With this option, random restarts will be enabled in the EM algorithm (§3.1.8), where the number of trials is NumInit. By default, NumInit is set as 1, that is, nbc will not perform random restarts.

```
--max-iter MaxIter
-m MaxIter
```

This option indicates the maximum number of iterations to be performed is MaxIter. That is, nbc will stop the EM algorithm when the number of iterations exceeds MaxIter. If this option is omitted or MaxIter = 0, the EM iterations will be continued until the convergence.

--max-iter-restart MaxInitIter

```
-1 MaxInitIter
```

With this option, the maximum number of preliminary EM iterations in random restarts (§3.1.8) will be set as MaxInitIter.

--init-c InitClassMethod

This option specifies the initialization method for the class parameters (Step 1 in Fig. 3 and Fig. 4). There are two alternatives — 'noisy\_u' initializes the parameters based on a uniform distribution with small noises, and 'random' initializes the parameters more randomly. The default method is 'noisy u'.

--init.-a InitAttrMethod

This option specifies the initialization method for the parameters of discrete attributes (Step 1 in Fig. 3 and Fig. 4). Similarly to --init-c (-I) option, 'noisy\_u' initializes the parameters based on a uniform distribution with small noises, and 'random' initialize the parameters more randomly. In addition, 'v\_freq' initializes the parameters based on the empirical frequencies with small noises. That is, each parameter  $\theta_{j,k,x_i}$  will be initialized to  $\frac{1}{N} \sum_{\mathbf{x}' \in D: \mathbf{x}'_i = \mathbf{x}_i} N(\mathbf{x}')$  with a small noise. The default method is 'noisy\_u'.

#### **\diamond** Membership distribution

```
--mem-prob (no argument)
```

-D (no argument)

With this option, the membership probabilities  $p(k \mid x)$ (§3.1.3) will be written into a file 'Base memp.txt'.

### ◊ Dissimilarity

--dissim (no argument)

This option enables the computation of the dissimilarities between objects, and the dissimilarities between attribute values (§3.1.6). The computed dissimilarities are written into 'Base\_dissim. Ext'. Since nbc will try to compute the dissimilarities for all possible pairs of objects in the dataset D, it may take a quite long time and may create a very large file for large D.

## 

--rank RankArg

## -z RankArg

This option enables relevance analysis (§3.1.5), in which the most relevant objects or attribute values to each class (each cluster) are recorded into 'Base\_rank.Ext'. If *RankArg* is a floating-point number  $\rho$  between 0 and 1, nbc only records the objects (resp. attribute values) whose relevance score  $R(k, x) = p(k \mid x, \theta)$  (resp.  $R^{j}(k, x_{i}) = p(k \mid x_{i})$  $(x_i, \theta)$ ) is greater than or equal to  $\rho$ . If *RankArg* is a positive integer  $N \ge 1$ , only N most relevant objects or attribute values will be recorded. If RankArg is set as 0, the ranking over all objects or attribute values will be recorded.

--hide-rank (no argument)

With this option, the objects (resp. the attribute values) whose relevance score  $R(k, x) = p(k \mid x, \theta)$  (resp.  $R^{j}(k, x_{j}) = p(k \mid x_{j}, \theta)$ ) is smaller than 1/K, where K is the number of classes, will not be written into 'Base\_rank.Ext'.

## Model selection

If we give an option '-k *Kmin:Kmax:Kstep*', nbc will enter into the 'model scoring' mode (§3.1.7) — that is, if we specify '-k 3:10:2', nbc will compute the scores of the models with the numbers of classes K = 3, 5, 7, 9.

--score ModelScore

-g ModelScore

This option specifies the model score as *ModelScore*. The available model scores are 'bic' (BIC, Eq. 40) and 'cs' (the Cheeseman-Stutz score, Eq. 41). The default score is BIC. The Cheeseman-Stutz score is also enabled by --autoclass (-A) option.

## **\diamond DAEM algorithm**

- --daem (no argument)
- -b (no argument)

This option enables the DAEM algorithm.

--itemp InvTempInit:InvTempRate

#### -B InvTempInit: InvTempRate

This option sets the initial value  $\beta_{init}$  of the inverse temperature as *InvTempInit*, and the updating coefficient  $\beta_{rate}$  as *InvTempRate*. It is required that  $0 < \beta_{init} \le 1$  and  $\beta_{rate} > 1$ . If '*InvTempInit*:' is given as the argument, the default value 1.2 is used for  $\beta_{rate}$ . If '*InvTempRate*' is given, the default value 0.1 is used for  $\beta_{init}$ . From our experience, the appropriate values of  $\beta_{init}$  and  $\beta_{rate}$  seem quite different according to the target data.

When we enable the DAEM algorithm, the terminal messages like below will be displayed ( $\beta_{init} = 0.4$  and  $\beta_{rate} = 1.2$ ). In these messages, each '\*' indicates that the inverse temperature has just been updated.

% nbc -f ex1 -k 4 -x 10 -I 0 -E 1 -y 4,9 -u 3,6,7 -b -B 0.4: -n 3 Reading cases from ex1\_freq.csv...done #classes = 4: [0] #iters \*0\*...100.\*.\*\*\*\*\* (Converged: 163 iters) L=-5747.688373 [1] #iters \*0\*...100.\*.\*\*\*\*\* (Converged: 167 iters) L=-5747.688373 [2] #iters \*0\*\*...\*\*\*\* (Converged: 75 iters) L=-574 7.688373 <<Resumed best parameter set #0>> [0] #iters \* (Converged: 165 iters) L=-5747.688373 BIC = -5923.836132

#### **\diamond SMEM algorithm**

Since the number of split-merge candidates to be examined is K(K-1)(K-2) at maximum, where *K* is the number of classes, the SMEM algorithm could take a long time to find a better set of parameters. nbc provides several (ad-hoc) control flags not to explore unpromising search space. In the current version, the SMEM algorithm is still considered as experimental.

--n-sm NumOp

 $-\circ$  NumOp

This option enables the SMEM algorithm, where the number of operations is set as *NumOp*.

--n-cand NumCand

-a NumCand

This option specifies the maximum number of the splitmerge candidates to be examined.

```
--n-split NumSplitCand
```

## -G NumSplitCand

This option limits the number of the split candidates to be taken into account for each merge candidate.

#### --min-c-param MinClassPar

This option specifies the lower limit of the class probability of a class to be split. That is, if the class probability of a split candidate is lower than *MinClassPar*, nbc will skip the candidate. If this option is omitted, no limit will be set.

#### --min-new-c-param MinNewClassPar

This option specifies the lower limit of the class probability of a new class obtained by a split-merge operation. That is, if one of the probabilities of three newly obtained classes is lower than *MinNewClassPar*, the result of the split-merge operation will be discarded. If this option is omitted, no limit will be set.

--min-sm-imp MinImprove

This option specifies the lower limit of the improvement of a split-merge operation. That is, the improvement of such a operation does not exceed *MinImprove*, the result of the operation will be discarded. The default value is  $10 \times \xi$ , where  $\xi$  is the threshold for judging convergence of the likelihood or the a posteriori probability, which is specified with --epsilon(-e) option.

This option enables the normalized version of  $J_{\text{merge}}$  (Eq. 46), instead of the original one (Eq. 44).

<sup>--</sup>norm-jmerge (no argument)

## Input/Output files

With GNU libiconv, we can handle datasets with non-ASCIIsafe character codes. The default codeset can be specified at the installation time, by the configure script (§4.2.3) or Makefile.msvc (§4.2.4). Without such a configuration, the default codeset will be set as UTF-8. Running 'iconv -1' would show a list of the available codesets. As described in §4.3.3, each input/output file is either in the CSV format or in the tab-separated value format.

```
--target Base
```

```
-f Base
```

This option indicates that we have a file named *Base* \_freq.*Ext* containing the target dataset *D*. *Base* will also be used as the base name of the input/output files as listed in Table 2.

## --suffix Suffix

## -s Suffix

nbc adds a suffix *Suffix* to the base name of the output files. For example, '*Base\_param.Ext*' will be changed as '*Base\_Suffix\_param.Ext*'.

#### --input Ext

As described in §4.3.3, nbc tries to determine the input file format automatically. On the other hand, with this option, nbc only reads the input files that have the file extension *Ext.* The input files should be in the tab-separated value (resp. CSV) format when *Ext* is txt (resp. csv).

## --output Ext

-j Ext

As described in §4.3.3, the output file format is the same as the input file format by default. On the other hand, with this option, nbc outputs the files whose file extension is *Ext*, where the output files are in the tab-separated value (resp. CSV) format when *Ext* is txt (resp. csv).

## --code-in Charcode

-p *Charcode* 

(Available only with GNU libiconv)

nbc assumes the character codeset of the input files is *Charcode*. If this option is omitted, the default codeset (which can be specified at compilation time) will be used.

## --code-out Charcode

-q Charcode

(Available only with GNU libiconv)

nbc outputs files with the character codeset *Charcode*. If this option is omitted, the default codeset (which can be specified at compilation time) will be used.

## ◊ Miscellaneous

- --log-valued (no argument)
- -J (no argument)

With this option, most of probability values are stored as log-valued. This would remedy the problem of underflow of very small probabilities, but on the other hand, the probability computations will take a bit longer time. Also it is highly recommended to combine this option with the settings for MAP estimation (e.g. with --autoclass (-A) option).

## --save-msg (no argument)

## -S (no argument)

By default, nbc outputs the terminal messages to the standard error output. This option switches the output to a file named '*Base\_msg.Ext*'. Then the buffer will not be flushed for each dot symbol, so for a small model/dataset, this option would shorten the runtime.

--uncompress (no argument)

-U (no argument)

This option inhibits the compression of indistinguishable objects, which is described in §2.1 and §4.3.4.

## --hide-attr (no argument)

#### -V (no argument)

By default, the attribute values appearing in the dataset D will always be recorded into files. With this option, on the other hand, they will not be recorded.

--prec-out NumDigits

This option sets *NumDigits* to the number of significant digits in each floating-point number to be written into files. At the implementation level, *NumDigits* will be passed as N in "\$. Ng" to fprintf() in the standard C library.

## --prec-msg NumDigits

This option sets *NumDigits* to the number of digits after the decimal point in each floating-point number to be displayed on the terminal. At the implementation level, *NumDigits* will be passed as N in "\$.Nf" or "\$.Ne" to fprintf() in the standard C library.

--verbose (no argument)

-v (no argument)

With this option, several temporary results are recorded into '*Base\_log.Ext*'. It should be noted that the resulting log file may become quite large.

--occ-order (no argument)

By default, the discrete attribute values output from nbc are ordered according to their character codes. With this option, on the other hand, nbc will write them in the order of appearances in '*Base\_freq.Ext*'.

--help (no argument)

-h (no argument)

This option displays a short description on major optional flags.

Table 3: Input/Output files for vnbc. *Base* is the base name, and *Ext* is txt (for the file in the tab-separated value format) or csv (for the file in the comma-separated value format).

	Filename	Content	
Input	Base_freq.Ext	Occurrences of	
		objects	
Input/Output	Base_hparam.Ext	Hyperparameters	
Output	Base_cluster.Ext	Clustering results	
Output	Base_memp.Ext	Membership probs	
Output	Base_rank.Ext	Result of	
		relevance analysis	
Output	Base_dissim.Ext	Dissimilarities	
Output	Base_log.Ext	Logs	
Output	Base_msg.Ext	Terminal messages	

--helpall (no argument)

-X (no argument)

This option displays a short description on all optional flags.

## 4.4 VB based clustering

Generally speaking, the usage of vnbc is quite similar to that of nbc. The names of input/output files are shown in Table 3. Also the file format for vnbc is the same as that of nbc except the descriptions in the header parts (see §4.3.3). The differences are as follows:

- The options --read-param (-R), --autoclass (-A), --score (-g), --init-c, --init-a and --log-valued (-J)<sup>16</sup> are ignored in vnbc.
- The meaning of --read-hparam (-H) is changed as follows:

--read-hparam (no argument)

-H (no argument)

If this option is given, vnbc will skip the VB-EM algorithm and read the hyperparameters from a file named *Base\_hparam.Ext*. The number of classes is then determined according to the content of *Base\_hparam.Ext*.

• The meanings of -smooth-c(-c), -smooth-a(-w), -tau(-T), -alpha(-0), -smooth-mean(-K) and -smooth-var(-L) are changed to specify the initial hyperparameters such as  $\alpha_k^{(0)}$  and  $\alpha_{j,k,x_j}^{(0)}$  in Step 1 of Fig. 9. For instance, if we give '--smooth-c *ClassPseudoCount*', the initial hyperparameters  $\alpha_k^{(0)}$  are set as  $(1 + ClassPseudoCount) + \epsilon_k$ , where  $\epsilon_k$  is a small random noise.

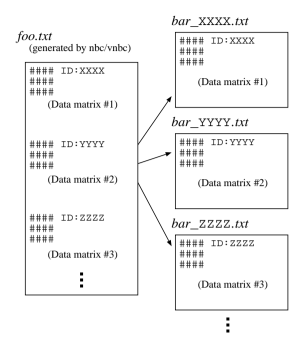


Figure 11: Output file split by nbcsep with '-f bar'.

## 4.5 Auxiliary tool

NBCT provides an auxiliary tool named nbcsep for postprocessing of the outputs from the executables nbc and vnbc. Hopefully, using nbcsep combined with some UNIX commands such as grep, cut, sort, sed, and so on, we may not have to use extra text processing programs (written in Perl, Python, Ruby, etc.).

As described above, an output file contains several data matrices each of which has a header part with a predefined identifier. In a typical case, we use nbcsep to split the output file into the files whose names have the corresponding identifiers (Fig. 11). The synopsis of nbcsep is as follows:

```
nbcsep [Options] InputFile [InputFile ...]
```

If '-' is given as an *InputFile*, nbcsep assumes that one input is passed from the standard input. If no options are given, nbcsep displays all identifiers of the data matrices in *Input-Files*. The below is a list of available options:

--target Base

-f Base

nbcsep writes each data matrix in *InputFile* into the file named '*Base\_ID.Ext*', where *ID* is the identifier of the data matrix, and *Ext* is the file extension of *InputFile*. The default value of *Base* is out.

--view ID -v ID

nbcsep outputs only the data matrices whose identifiers match with *ID*, ignoring the case of letters, to the standard

<sup>&</sup>lt;sup>16</sup>Most of probability(-like) values are unconditionally stored as log-valued.

output. The matching criterion can be configured by -m option.

--output ID

-0 *ID* 

This is the same as ' $\neg v$  *ID*' except that the output will be saved into the file(s) whose names have a prefix specified with -f option.

```
--match Criterion
```

```
-m Criterion
```

This option specifies the matching criterion as *Criterion* for -v and -o options. *Criterion* is one from exact, infix, prefix and suffix. The default value is infix.

--suffix Suffix

-s Suffix

nbcsep adds *Suffix* to the name of the output file of nbcsep. The resulting filename will be *'Base\_Suffix\_ID.Ext'*.

```
--ext Ext
```

−e *Ext* 

nbcsep uses *Ext* as a new file extension for the output files. If this option is omitted, the file extension will be set as the one commonly appearing in *InputFiles*. If different extensions are used in *InputFiles*, 'out' will be used as the default file extension.

```
--del-head (no argument)
```

```
-d (no argument)
```

nbcsep outputs only the data matrices.

```
--del-body (no argument)
```

```
-D (no argument)
```

nbcsep outputs only the header parts.

--code-in Charcode

```
-p Charcode
```

(Available only with GNU libiconv)

nbcsep assumes the character codeset of the input files is *Charcode*.

- --code-out Charcode
- -q *Charcode* (Available only with GNU libiconv)

nbcsep outputs the files with the character codeset *Charcode*.

```
--help (no argument)
```

```
-h (no argument)
```

nbcsep displays a simple help message.

# **5** Miscellaneous remarks

## 5.1 Settings for EM algorithms

The computation time and space required in an EM iteration is almost linear in each of the number of classes, the number of single-dimensional attributes, and the number of distinct objects in the dataset. Exceptionally, however, it is cubic in the largest dimension of the multivariate Normal distributions included in the model.

For some applications, the default value of the threshold for judging convergence of the likelihood or the a posteriori probability, which is specified by --epsilon(-e) option (§4.3.5), might be small. Early researches on statistical natural language processing reported that, at least in ML estimation, a large number of EM iterations do not necessarily lead to a good performance in probabilistic inferences after EM learning (e.g. Section 10.3.2 of [16] discusses the case of hidden Markov models). So there should be cases where it is reasonable to give a larger value to -epsilon(-e) option, or set the maximum number of iterations with -max-iterate(-m) option (§4.3.5). In addition, it is empirically observed that the EM algorithm run with larger pseudo counts tends to converge with a smaller number of iterations.

## 5.2 Precision of floating-point numbers

For large-scale datasets, NBCT might show an unexpected behavior due to the problems with arithmetic precision such as underflow or precision errors. --log-valued option provides a way to avoid underflow. Also, NBCT uses the long double type for floating-point numbers on Linux, Mac OS X, and Win32 with Cygwin. Of course long double can be memory consuming than double,<sup>17</sup> and thus, to use double instead of long double, we add the '--disable-long-double' flag to the configure script. Also it should be addressed that, in the current implementation of the digamma function  $\Psi(\cdot)$ , floating-point numbers are treated as double.

# 5.3 Numerical problems in ML/MAP-based clustering

As described in §3.1.2, ML/MAP-based clustering with multivariate Gaussian distributions tends to show an unstable behavior for a small dataset (e.g. the updated covariances may be singular). For such cases, the MAP estimation is more preferred than the ML estimation, and --log-valued (-J) option would make the probability computation more stable.

<sup>&</sup>lt;sup>17</sup>The physical size of long double is system-dependent, and is only guaranteed not to be smaller than that of double. If you are using long double, the physical size will be recorded into the log file.

## 5.4 Parallelization via OpenMP\*

Nowadays a couple of popular C compilers support OpenMP for shared-memory parallel computing. In NBCT's source code, several 'pragma' declarations for OpenMP are added to the 'for' loops which require massive numerical computation. An interested user may enable these 'pragma' declarations by giving compiler-dependent flags to the C compiler. With the GNU C compiler (version 4.2 or later), for instance, we may add CFLAGS='-fopenmp -O3' and LIBS='-lgomp' to the arguments of the configure script, and specify the number of threads at runtime by the OMP\_NUM\_THREADS environment variable.

# **Contact information**

NBCT is still under development, and bug reports, questions, suggestions, or any other feedbacks are highly welcome. To make a contact, please send an e-mail to nbct[AT]mi.cs. titech.ac.jp (please replace [AT] with @).

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<sup>&</sup>lt;sup>18</sup>SPECFUN was developed by W. J. Cody et al. at Argonne National Laboratory, and is now available in public domain at http://www.netlib. org/specfun/.

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