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# Internship report notes

Master 2 - Data Science

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# 1 Introduction

Given observations of random variables (X, Y), we suppose that there exist another set of random variables Z that we do not observe, yet that characterize (X, Y) conditionally to Z. Z is then called a latent variable, or a hidden variable.

For instance, if we observe the weights of a given population through X, and that we aim at inferring their height Y, knowing the sex of each individual through Z could improve our predictions on Y. Hence, assuming that there exist a latent variable to a given model adds structure to the model while improving the explainability, as Z characterizes the behavior of our dataset. Typically, clustering methods like KMeans or Gaussian Mixtures Models (GMM) provide a discrete Z given the observations, which are interesting in the sense that they provide a categorical representation of our data.

However, finding such Z given the observations is not straight forward, and not all dataset respond to a latent process, and may be not even a discrete one.

During this research internship, we aim at exploring latent models with discrete latent space in order to analyze the microbiota structure. Our first focus will be on various methods to conceive latent models like Expectation-Maximization and variational models.

# 2 Expectation-Maximization

# 2.1 Overview

The Expectation-Maximization (EM) algorithm, first introduced in [1], is vast class of latent models. It is based on the following smart decomposition of the data:

$$\log p_{\theta}(X) = \underbrace{\mathbb{E}_{p_{\widehat{\theta}}(Z|X)}[\log p_{\theta}(X,Z)|X]}_{Q(\widehat{\theta},\theta)} - \mathbb{E}_{p_{\widehat{\theta}}(Z|X)}[\log p_{\theta}(Z|X)|X]$$

The idea behind this decomposition is that  $\log p_{\theta}(X)$  is generally not tractable since it's an integral, while the complete likelihood  $p_{\theta}(X, Z)$  is generally manageable. Note that since  $\log p_{\theta}(X)$  can not be computed,  $\log p_{\theta}(Z|X)$  can't either by extension. Hence, we introduce  $\log p_{\widehat{\theta}}(Z|X)$  where  $\widehat{\theta}$  is the maximum likelihood estimator of  $\theta$ :

$$\widehat{\theta} = \arg\max_{\theta} \log p_{\theta}(X)$$

The main trick of the EM algorithm relies in the idea that  $Q(\widehat{\theta}, \theta)$  is sufficient to compute a maximum likelihood estimator of  $\theta$ . Indeed, consider the following algorithm:

# **Algorithm 1** Expectation-Maximization

Require:  $\widehat{\theta}$ 

Repeat until convergence

**Expectation**: compute  $p_{\widehat{\theta}}(Z|X)$  to compute  $Q(\widehat{\theta},.)$ 

**Maximization**:  $\widehat{\theta} = arg \max_{\theta} Q(\widehat{\theta}, \theta)$ 

**return**  $\widehat{\theta} = 0$ 

If we denote by  $\widehat{\theta}^h$  the iterates of this algorithm, one can show using Jensen's inequality that:

$$\log p_{\widehat{\theta}^{h+1}}(X) \ge \log p_{\widehat{\theta}^{h}}(X)$$

As a result, the EM algorithm maximizes the likelihood, producing an estimator  $\widehat{\theta}$  that is an MLE of  $\theta$ . Note that we don't have a convergence certainty towards the best maximizer of the likelihood, only to a local maxima. Hence, the EM algorithm is heavily sensitive to the initialization we pick.

All is required now is to choose a latent model so we can perform the EM algorithm, meaning that we have to define the distributions of the followings:

- $Z \sim \mathcal{B}(K,\pi)$ , conveniently set to a binomial of parameter  $\pi$  so that it's discrete and simple to manage.
- $X|Z=k\sim p_{\gamma(k)}(X|Z=k)$ , which is where we have the most choice to make.

In such models,  $\theta = (\pi, \gamma(0), ..., \gamma(K))$ . In the next section, we will study a specific fork architecture of the gaussian mixture case.

# 2.2 Gaussian Mixture Linear Classifier

# 2.2.1 Framework and computations

Consider the case for which we observe  $(X_i, Y_i)_{1 \leq i \leq n}$  i.i.d samples, where  $X_i$  denotes a feature vector and  $Y_i$  a label in a classification framework. We aim at introducing a linear classifier that exploits a latent structure over (X, Y), so that we have the following latent model:

- $Z_i \sim \mathcal{B}(K, \pi)$ , we note  $\pi_k = \mathbb{P}(Z_i = k)$
- $X_i|Z_i = k \sim \mathcal{N}(\mu_k, \sigma_k I)$ , we denote by  $f_k(X_i)$  its density.
- $\mathbb{P}(Y_i = 1 | X_i, Z_i = k) = \sigma(W_{e,k}^T e_k + W_{x,k}^T X_i) = p_k(X_i)$ , where  $e_k$  denotes a vector from the canonical basis of  $\mathbb{R}^K$ .

As we want to perform the EM algorithm find an MLE of

$$\theta = (\pi, \mu_1, ..., \mu_K, \sigma_1, ..., \sigma_K, W_{e,1}, ..., W_{e,K}, W_{x,1}, ..., W_{x,K})$$

we start by computing the **expectation** step by assessing  $p_{\widehat{\theta}}(Z_i = k|X_i, Y_i)$ , using Bayes rules  $(\hat{\pi}, \hat{p}, \hat{f} \text{ signify that we evaluate these quantities using the current estimate } \widehat{\theta})$ :

$$p_{\widehat{\theta}}(Z_i = k | X_i, Y_i) = \frac{\hat{\pi}_k \hat{f}_k(X_i) \left( Y_i \hat{p}_k(X_i) + (1 - Y_i)(1 - \hat{p}_k(X_i)) \right)}{\sum_{j=1}^K \hat{\pi}_j \hat{f}_j(X_i) \left( Y_i \hat{p}_j(X_i) + (1 - Y_i)(1 - \hat{p}_j(X_i)) \right)}$$
$$= \tau_{ik}$$

Now, we can safely evaluate  $Q(\widehat{\theta}, \theta)$  for any  $\theta$ :

$$Q(\widehat{\theta}, \theta) = \mathbb{E}_{p_{\widehat{\theta}}(Z|X)}[\log p_{\theta}(X, Y, Z)|X]$$

$$= \sum_{i=1}^{n} \sum_{k=0}^{K} \log p_{\theta}(X_{i}, Y_{i}, Z_{i} = k)\tau_{ik}$$

$$= \sum_{i=1}^{n} \sum_{k=0}^{K} (\log \pi_{k} + \log f_{k}(X_{i}) + Y_{i} \log p_{k}(X_{i}) + (1 - Y_{i}) \log(1 - p_{k}(X_{i})))\tau_{ik}$$

The **maximization** step now consists in deriving  $Q(\widehat{\theta}, \theta)$  regarding each parameters in  $\theta$  so that we obtain an either explicit value or iterative procedure to compute the next iterate of  $\widehat{\theta}$ .

• Maximization regarding  $\pi_k$  under constraint that  $\sum_{k=0}^K \pi_k = 1$  can be solved explicitly using Lagrange duality:

$$\pi_k^* = \frac{1}{n} \sum_{i=1}^n \tau_{ik}$$

• Maximization regarding  $(\mu_k, sigma_k)$  is given by the maximum of likelihood estimator on  $\sum_{i=1}^{n} \tau_{ik} \log f_k(X_i)$ :

$$\mu_k^* = \frac{1}{\sum_{i=1}^n \tau_{ik}} \sum_{i=1}^n \tau_{ik} X_i$$

$$\sigma_k^* = \frac{1}{\sum_{i=1}^n \tau_{ik}} \sum_{i=1}^n \tau_{ik} (X_i - \mu_k^*) (X_i - \mu_k^*)^\top$$

• Maximization regarding  $(W_{e,k}, W_{x,k})$  is not explicit, and requires a fixed point algorithm like gradient descent to determine an estimate of the optimal parameters. The iterations using full batch gradient descent are given below, with learning rate  $\alpha$ :

$$W_{e,k}^{l+1} \leftarrow W_{e,k}^{l} - \alpha \sum_{i=1}^{n} (p_k(X_i) - Y_i) \tau_{ik} e_k$$

$$W_{x,k}^{l+1} \leftarrow W_{x,k}^{l} - \alpha \sum_{i=1}^{n} (p_k(X_i) - Y_i) \tau_{ik} X_i$$

Each iteration should be confronted to the maximization criterion, so that each iterate improves  $Q(\widehat{\theta}, \theta)$ :

$$Q(\widehat{\theta}^{l+1}, \theta) \ge Q(\widehat{\theta}^{l}, \theta)$$

In the end, only the best improvement iterate is kept for  $W_{e,k}^*$  and  $W_{x,k}^*$ . Note that other methods could be used like SGD or CMAES as implemented during the internship.

After the maximization, we can update  $\widehat{\theta}$  with the previously computed parameters, and redo the (E) and (M) steps up to convergence. The convergence can be measured relatively to  $Q(\widehat{\theta}, \theta)$ , so that for a threshold  $\epsilon$ , we can use the following stopping criterion:

$$\frac{|Q(\widehat{\theta}^{h+1}, \widehat{\theta}^h) - Q(\widehat{\theta}^h, \widehat{\theta}^h)|}{|Q(\widehat{\theta}^h, \widehat{\theta}^h)|} \le \epsilon$$

We now have a ready-to-go Gaussian Mixture Linear Classifier that we can benchmark on an suited dataset against other common methods.

### 2.2.2 Dataset generation

Before we benchmark the Gaussian Mixture Linear Classifier, we need to generate a dataset that is well suited to its usage. Hence, we set random parameters for  $\theta$  and generate a new dataset following the latent framework we have set previously:

- For all  $k \leq K$ , generate n/K points following a gaussian parameterized by  $(\mu_k, \sigma_k I)$ . X is given by each point coordinate, Z by the gaussian from which the point was generated.
- For each sample, characterized by  $(X_i, Z_i)$ , draw the label of the sample as  $Y_i \sim \mathcal{B}(\sigma(W_{e,Z_i}^T e_{Z_i} + W_{x,Z_i}^T X_i))$ .

The following figure illustrates a generation with K=2:



Figure 1: Generated samples out of K=2 gaussians, labelized using a logistic model (label 1 or 0). The density of the hidden gaussians is represented in the background using shades of grey (black intense, white almost 0)

Note on the previous figure 1 that the limit between the labels in each subset gaussian is not sharp, as it is sorted out of a probabilistic modelisation (logistic). Consequently, an interesting observation can be made as we force the gaussians to have small values of mean and variance. Indeed, as shown on the next figure, if we take the same parameters as previously and divide them by a factor 10, the logistic model is ill conditioned.



Figure 2: Generated samples out of K=2 gaussians, using the same parameters as for figure 1 divided by a factor 10.

All points are close to the frontier in terms of norm on figure 2. Since we do not exploit any normalization term in the sigmoid modelization, this leads to a blurry area in which the linear separation model is not useful at all. This situation is heavily problematic as it prevents us from performing a general benchmark on that dataset. Indeed, the further larger the variance is, with sufficient samples, the better the linear approximation will be and therefore the better the modelization gets. On the contrary, with smaller variance the linear model isn't descriptive of the generated samples as they are all close to the frontier, therefore with heavily noisy labelization.

# 2.3 Dirichlet Mixture Linear Classifier

# 2.3.1 Framework and objectives

In the previous section, we have derived a very classical model (Gaussian Mixture Model) into a linear classifier using the EM algorithm. However, gaussian latent modelization is far from general, and may not be suited to our specific usage on the microbiota. Indeed, after performing the previous method onto our microbiota dataset, it turned out to be performing just as bad as the classical logistic regression, no matter the chosen latent space dimension. Therefore, we deduce that the gaussian latent modelization is not adapted to our practical settings.

As we analyze the data, we observe that each  $X_i$  belongs to the simplex. A natural distribution supported on the simplex is the Dirichlet distribution, parameterized by  $\alpha = (\alpha_1, \ldots, \alpha_p)$  where p is the dimension of  $X_i$ . We denote the dirichlet distribution by  $\mathcal{D}(\alpha)$ , for which the density is given by the following:

$$f(x|\alpha) = \frac{\Gamma\left(\sum_{j=1}^{p} \alpha_j\right)}{\prod_{j=1}^{p} \Gamma(\alpha_j)} \prod_{j=1}^{p} x_j^{\alpha_j - 1}$$

where  $\Gamma$  denotes the gamma function. For notation simplicity, we also introduce the digamma function that will play a key role in our model:

$$\psi(\alpha) = \frac{d}{d\alpha} \log \Gamma(\alpha) = \frac{\Gamma'(\alpha)}{\Gamma(\alpha)}$$

Since we are doing a Dirichlet mixture model, we have K Dirichlet distributions to handle. We introduce the notation  $\alpha^{(k)}$  to parameterize the k-th Dirichlet distribution.

As previously, we first perform the **expectation** step obtain the same result with a different conditional a priori distribution on X|Z:

$$p_{\widehat{\theta}}(Z_i = k | X_i, Y_i) = \frac{\hat{\pi}_k f(X_i | \hat{\alpha}^{(k)}) (Y_i \hat{p}_k(X_i) + (1 - Y_i) (1 - \hat{p}_k(X_i)))}{\sum_{j=1}^K \hat{\pi}_j f(X_i | \hat{\alpha}^{(j)}) (Y_i \hat{p}_j(X_i) + (1 - Y_i) (1 - \hat{p}_j(X_i)))}$$
$$= \tau_{ik}$$

We can now evaluate  $Q(\widehat{\theta}, \theta)$  for any  $\theta$ , which enables us to perform the **maximization** step:

• The maximization over  $pi_k$  under the simplex constraint  $\sum_{k=1}^k \pi_k = 1$  is again given by Lagrange duality as:

$$\pi_k^* = \frac{1}{n} \sum_{i=1}^n \tau_{ik}$$

• The maximization over  $\alpha^{(k)_j}$  is not straightforward on the other hand, and requires a fixed point algorithm. Indeed, deriving over  $\alpha^{(k)_j}$  we obtain:

$$\partial_{\alpha_j^{(k)}} Q(\widehat{\theta}, \theta) = \sum_{i=1}^n \left( \psi \left( \sum_{l=0}^K \alpha_l^{(k)} \right) - \psi(\alpha_j^{(k)}) + \log x_{ij} \right) \tau_{ik}$$
$$= \left( \psi \left( \sum_{l=0}^K \alpha_l^{(k)} \right) - \psi(\alpha_j^{(k)}) \right) \sum_{i=1}^n \tau_{ik} + \sum_{i=1}^n \tau_{ik} \log x_{ij}$$

Hence, as we look for  $\partial_{\alpha_i^{(k)}}Q(\widehat{\theta},\theta)=0$ , we obtain:

$$\psi(\alpha_j^{(k)}) - \psi\left(\sum_{l=0}^K \alpha_l^{(k)}\right) = \frac{\sum_{i=1}^n \tau_{ik} \log x_{ij}}{\sum_{i=1}^n \tau_{ik}}$$

Thankfully, [4] provides a few tricks to solve iteratively such equation, so that we can iterate as follows (5 steps are sufficient to obtain high-accuracy solution according to [4]):

$$\alpha_j^{(k)} \leftarrow \psi^{-1} \left( \frac{\sum_{i=1}^n \tau_{ik} \log x_{ij}}{\sum_{i=1}^n \tau_{ik}} + \psi \left( \sum_{l=0}^K \hat{\alpha}_l^{(k)} \right) \right)$$
$$\hat{\alpha}_j^{(k)} \leftarrow \alpha_j^{(k)}$$

However, this solution is a lower bound to the true objective, which makes our EM a generalized version of it.

• The maximization over  $(W_{e,k}, W_{x,k})$  is also given by a fixed point algorithm, which ends up being the same computation as previously for the Gaussian case:

$$W_{e,k}^{l+1} \leftarrow W_{e,k}^{l} - \alpha \sum_{i=1}^{n} (p_k(X_i) - Y_i) \tau_{ik} e_k$$
$$W_{x,k}^{l+1} \leftarrow W_{x,k}^{l} - \alpha \sum_{i=1}^{n} (p_k(X_i) - Y_i) \tau_{ik} X_i$$

### 2.3.2 Dataset generation

Now that we have defined the EM algorithm in the previous section, we aim at generating a dataset to benchmark the dirichlet mixture classifier. Following a similar procedure as for the gaussian case, we are able to generate a dataset that matches a dirichlet mixture and is labeled following the sigmoid modelisation. The next figure illustrates a given generation:

# 0.4 - 0.2 - 0.1 - 0.0 -

Figure 3: Generated samples out of K=2 dirichlet distributions, labelized using the sigmoid modelisation on  $\mathbb{P}(Y_i=1)$ .

0.6

0.8

0.4

0.2

0.0

As previously, since the data lives in the simplex, they are too close to the border of the next label set, which ends up creating a blurry dataset for which the linear model is not relevant anymore.

# 3 Variational methods

In this section, we aim at resourcing some variational methods that we are going to use in this study.

# 3.1 Variational Auto-Encoders

# 3.1.1 Framework and optimization objective

We are interested in another kind of latent models, this time based on variational inference results to achieve a new kind of deep latent structure: the Variational Auto-Encoder (VAE). These latent models were introduced in 2013 by Kingma, better described in a more in depth paper in 2019: see [3]

Once again, we assume the observations X to be modelizable by a given distribution parameterized by  $\theta$ :

$$X \sim p_{\theta}(x)$$

Determining  $\theta$  holds to find one  $\theta^*$  that would optimize a given objective, generally chosen as the maximum of likelihood. Indeed, if  $\theta^* \in arg \max_{\theta} p_{\theta}(x)$ , then such  $\theta^*$  maximizes the density around the dense areas of the observations, which makes them highly likely to happen under such distribution  $p_{\theta^*}$ . Hence, the maximum likelihood is a natural criterion:

$$\theta^* \in arg \max_{\theta} p_{\theta}(x)$$

However, such modelization does not include a latent structure. As a result, we try to enforce it by rewriting the objective as follows:

$$p_{\theta}(x) = \int_{\mathcal{Z}} p(x, z) dz$$

Using Bayes decomposition, we obtain the following objective:

$$p_{\theta}(x,z) = p_{\theta}(z)p_{\theta}(x|z)$$

Recall that the prior  $p_{\theta}(z)$  and the a priori  $p_{\theta}(x|z)$  are defined by the framework (ex: Bernoulli prior and Gaussian posterior gives the Gaussian mixture framework). However, the computation of the evidence  $p_{\theta}(x)$  is generally intractable in practice, which also leads to a non-tractable posterior distribution:  $p_{\theta}(z|x)$ . As a result, not being able to compute the evidence leads to not being able to provide a gradient regarding  $\theta$ , so we can not perform the backpropagation in a deep learning approach.

Note that there exist approximate inference techniques to compute the evidence and the posterior, but these are quite expensive and often yield poor convergence results.

To overcome this issue, we introduce a smart rewriting of the objective using variational

inference. Indeed, let  $q_{\Phi}(z|x) \approx p_{\theta}(z|x)$  to be learnt over  $\Phi$ , one can write:

$$\log p_{\theta}(x) = \mathbb{E}_{q_{\Phi}(z|x)} [\log p_{\theta}(x)]$$

$$= \mathbb{E}_{q_{\Phi}(z|x)} \left[ \log \frac{p_{\theta}(x)}{q_{\Phi}(z|x)} \frac{q_{\Phi}(z|x)}{p_{\theta}(z|x)} \right]$$

$$= \underbrace{\mathbb{E}_{q_{\Phi}(z|x)} \left[ \log \frac{p_{\theta}(x)}{q_{\Phi}(z|x)} \right]}_{ELBO(q_{\phi}(z|x), p_{\theta}(x, z))} + D_{KL} \left[ q_{\Phi}(z|x) || p_{\theta}(z|x) \right]$$

The first term of that decomposition is generally called the Evidence Lower BOund (ELBO), as it marks a lower bound to the evidence  $\log p_{\theta}(x)$  since the KL divergence is a positive quantity:

$$\log p_{\theta}(x) \ge ELBO(q_{\phi}(z|x), p_{\theta}(x, z))$$

 $q_{\Phi}(z|x)$  is an approximation of the true posterior  $p_{\theta}(z|x)$  that we aim at learning in a family of distributions. For instance,

$$q_{\Phi}(.|x) \sim \mathcal{N}(\mu(x), \Sigma(x))$$

would be an approximation of the true posterior by a Gaussian distribution. Notice that the true posterior may very not likely be Gaussian, which creates a first complexity error in our model.

Despite being a lower bound on the true maximum likelihood objective, the ELBO is actually tractable. Indeed, as we continue the computation:

$$ELBO(q_{\phi}(z|x), p_{\theta}(x, z)) = \mathbb{E}_{q_{\Phi}(z|x)}[\log p_{\theta}(x, z)] - \mathbb{E}_{q_{\Phi}(z|x)}[\log q_{\Phi}(z|x)]$$
$$= \mathbb{E}_{q_{\Phi}(z|x)}[\log p_{\theta}(x|z)] - D_{KL}[\log q_{\Phi}(z|x)||p_{\theta}(z)]$$

Another remarkable fact, is that when maximizing the ELBO, we are actually minimizing the KL divergence between the estimated and the true posterior. Hence, one can define the ELBO as a suboptimal objective to our problem that we get to maximize to obtain  $(\Phi^*, \theta^*)$ , the parameters of our model.

# 3.1.2 Reparameterization trick

Even though the gradient of the ELBO is well defined for  $\theta$ , it is not possible to compute the differential relatively to  $\Phi$  yet, as it requires samples from the approximation to the posterior  $q_{\Phi}(z|x)$  to compute  $\mathbb{E}_{q_{\Phi}(z|x)}[\log p_{\theta}(x|z)]$ .

Since, sampling is not a differentiable operation, we make use of the change of variable formula, so that for a bijective transformation  $z = \phi_x(\epsilon)$ , we get:

$$p(z) = p(\epsilon) \det \left| \frac{\partial \epsilon}{\partial z} \right|$$

Hence, if we take  $\epsilon$  a random variable of density  $p(\epsilon)$  that does not depend on  $\theta$ ,  $\Phi$  nor x, so that  $z = \phi_x(\epsilon)$ , for any  $L_1$  function f,

$$\mathbb{E}_{q_{\Phi}(z|x)}[f(z)] = \mathbb{E}_{p(\epsilon)}[f(z)]$$

As a result, the samples are not obtained through  $q_{\Phi}$  anymore but through  $p(\epsilon)$ , so that can safely perform derivation of the ELBO relatively to  $\Phi$  and backpropagate our gradient through the network.

### 3.1.3 Architecture

The vanilla architecture of the VAE is described by the following illustration:



Figure 4: Illustration of a VAE with Gaussian prior (wikipedia)

The first part is generally called the encoder, as it turns a sample x into its latent representation z by modelizing the posterior  $q_{\Phi}(z|x)$ . The second part is then called the decoder, as it throws a latent representation in the sample space. The latest can even serve as a generative architecture, as one can sample from the latent space through  $q_{\Phi}(z|x)$ , and decode it to obtain a new sample.

As we can see more clearly in that illustration, we can see that  $\Phi$  and  $\theta$  are trained jointly through the ELBO, both serving for one part of the VAE at a time.

The training procedure is straightforward: the entry is a sample x and the output objective is the same sample x. We aim at train the VAE for learning the data space and its latent representation by learning how to reconstruct the samples through it.

# 3.1.4 Limits for our problem

As we have seen through the reparameterization trick, training a VAE architecture requires to be able to backpropagate the gradient of the ELBO at each step. We namely had to perform the reparameterization trick to circumvent the randomness operation which is not differentiable. As a result, learning a discrete posterior is not possible with such architecture, since we would have to perform a projection of the output of the encoder on a discrete space, which is not a differentiable operation.

Yet, learning discrete representation of our data seems much more natural than continuous latent ones. As we tend to categorize things as much as we can, describing behaviors with words for instance. Furthermore, a discrete representation facilitates the interpretation of the latent space by ordering data distribution in simple bins.

The next architecture, called the VQ-VAE, stands as a first fork option to the VAE with discrete posterior.

# 3.2 VQ-VAE

# 3.2.1 Quick overview

Introduced in [7], VQ-VAE architecture provides a framework to compute discrete posterior distributions  $q_{\Phi}(z|x)$ . To compare that model with the VAE, we start by introducing the architecture of the model for which an illustration is given below:

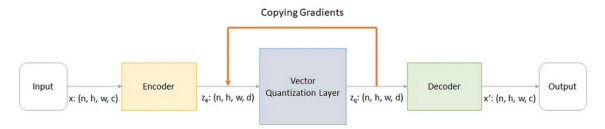


Figure 5: VQ-VAE architecture (source: Medium)

As we can see on figure 5, the major difference with the vanilla VAE architecture lies in the vector quantization step which enables to project the output of the encoder denoted by  $z_e(x)$  onto a discrete embedding dictionary  $(e_1, \ldots, e_K)$  by a simple distance argument:

$$k = arg \min_{j} ||z_q(x) - e_j||_2, \quad z_q(x) = e_k$$

The projection of  $z_e(x)$  on that discrete dictionary is denoted by  $z_q(x)$ , and serves as the input of the decoder. For further visual representation of the vector quantization layer, an illustration is given below.

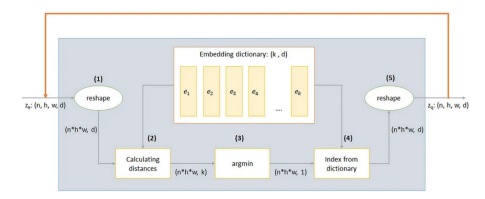


Figure 6: Architecture of the VQ-VAE: quantization layer (source: Medium)

Looking at the previous figures, we can grasp the challenge of backpropagation in such model with discrete prior and posterior. In the next section, we enter in the mathematical definition of the objective and how to train this architecture.

## 3.2.2 Framework and optimization objective

Contrary to the vanilla VAE, we have the following categorical distributions assumption:

- The prior  $p_{\theta}(z)$  is categorical. In the original paper, it is taken as uniform supported in  $\{1, \ldots, K\}$  during the training. When the training is over, it is fit to an autoregressive distribution through a PixelCNN (see [6]). It is left as an exploration research field to be able to learn the prior while training the model.
- The posterior  $q_{\Phi}(z|x)$  is categorical and set to the following:

$$q_{\Phi}(k|x) = \mathbb{1}_{\{k = arg \min_j \|z_q(x) - e_j\|_2\}}$$

This modelization of the posterior enables to obtain a discrete latent space, but it does not allow to differentiate regarding  $\Phi$ . As a result, the authors suggest two possible strategies:

- Straight-through: propagate the gradient through the discrete part (vector quantization layer) without changing it. The intuition is that the gradient propagated from the encoder contains sufficient information to update the encoder accordingly, but this is just intuition.
- Subgradient: compute the subgradient of the quantization layer (unexplored yet)

The optimization objective of the VQ-VAE is based on the ELBO, that one can compute as follow:

$$ELBO(q_{\Phi}(z|x), p_{\theta}(z|x)) = \mathbb{E}_{q_{\Phi}(z|x)}[\log p_{\theta}(x, z)] - \mathbb{E}_{q_{\Phi}(z|x)}[\log q_{\Phi}(z|x)]$$
$$= \mathbb{E}_{q_{\Phi}(z|x)}[\log p_{\theta}(x|z)] - D_{KL}[q_{\Phi}(z|X)||p_{\theta}(z)]$$

Notice then that:

• Since  $q_{\Phi}(k|x) = \mathbb{1}_{\{k = arg \min_j \|z_q(x) - e_j\|_2\}}$ , we have:

$$\mathbb{E}_{q_{\theta}(z|x)}[\log p_{\theta}(x|z)] = \log p_{\theta}(x|z_q(x))$$

• Since  $Z \sim \mathbb{U}(\{1,...,K\})$ ,  $\mathbb{P}(Z=k) = \frac{1}{K}$ . Also, notice that  $q_{\Phi}(z_q(x)|x) = 1$  by definition of  $q_{\Phi}(z|x)$  and  $z_q(x)$ . Combining those results, we obtain:

$$D_{KL}[q_{\Phi}(z|x)||p_{\theta}(z)] = \mathbb{E}_{q_{\Phi}(z|x)} \left[ \log \frac{q_{\Phi}(z|x)}{p_{\theta}(z)} \right]$$
$$= \log \frac{q_{\Phi}(z_q(x)|x)}{p_{\theta}(z_q(x))}$$
$$= \log K$$

Therefore, this KL divergence does not impact the optimization objective as it does not depend on  $(\theta, \Phi)$ .

Computing the previous quantities, we would obtain the following suboptimal objective for the VQ-VAE:

$$ELBO(\Phi,\theta) = log p_{\theta}(x|z_q(x))$$

However, such objective does not enable to learn the dictionary since we use a straightthrough approach over the quantization layer. To update the dictionary, the authors suggest to add the following term to the loss:

$$||sg[z_e(x)] - e||_2^2$$

Where sg denotes the stop-gradient operator, meaning we do not consider any gradient after the given operation.

Finally, to prevent embedding space over expansion, the authors suggest the addition of a commitment loss parameterized by  $\beta > 0$ :

$$\beta \|z_e(x) - sg[e]\|_2^2$$

Indeed, the embedding space was unconstrained so far, so its dimension could grow arbitrarily. Intuitively, adding that term forces the model to commit to a given embedding.

Overall, we obtain this final objective to maximize:

$$\mathcal{L}(\theta, \Phi) = \log p_{\theta}(x|z_{q}(x)) + ||sg[z_{e}(x)] - e||_{2}^{2} + \beta ||z_{e}(x) - sg[e]||_{2}^{2}$$

# 3.2.3 Discussion over some limiting aspects

Even though the previous objective seems natural, we only rigorously justified the first term thanks to the ELBO. Indeed, the dictionary update loss as well as the commitment loss keep dropping out from nowhere, while they seem necessary to ensure the training of our model.

Furthermore, we did not really deal with the non differentiability of the vector quantization loss, and the straight-through estimator can definitely be criticized about what information it actually provides to the encoder to update appropriately.

Finally, the usage of the PixelCNN after the training seems very unnatural, and could significantly boost the model as the PixelCNN did show amazing performances so far.

# 3.3 PixelCNN

### 3.3.1 Overview

Introducing the VQ-VAE, we have seen that an under-table tool that was being used is the Pixel CNN, first introduced in [6]. In the context of the VQ-VAE, it plays a major role after training by learning the prior p(z), that was set to a discrete uniform previously. This makes a drastic difference with the VAE, as we are not setting the prior ourselves as it's being learnt and modeled by the PixelCNN in this process.

Indeed, the PixelRNN/PixelCNN architectures are sequential deep neural networks that aims at modeling the distribution of a data space in an autoregressive fashion. Their main characteristic is that they take advantage of the structure of an image to learn the data space distribution.

- PixelRNN: Bi-directional recurrent networks with variable smart directions are used to model the spatial dependencies between pixels. (3 architectures are presented in the original paper, but our focus will be on the PixelCNN here).
- PixelCNN: the dependencies between the pixels is modeled through stacking of masked convolution layers (it's faster since the receptive field is bounded by the size of the convolution), no pooling layer is used. The masking ensures that we keep an autoregressive estimation of a new pixel, without seeing the future pixels.

The following figure illustrates the masked convolution technique.

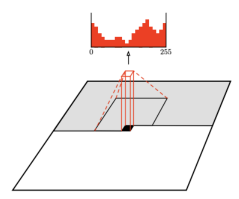


Figure 7: Masked convolution on an image (largest square), the big square represents the receptive field of the black pixel, the white area is masked

Hence, given x an image represented by its pixels  $x = (x_1, \ldots, x_n)$ , as usual for these distribution modeling problems, we aim at finding  $\theta^* = arg \max_{\theta} p_{\theta}(x)$  in a family of distributions parameterized by  $\theta$ : the maximum of likelihood. Contrary to the usual independent framework, we consider a local dependency between pixels given by the receptive field of our convolution. This local dependency is limited to the already seen pixels only as well, thanks to the masked convolution.

Furthermore, rather than using continuous outputs, these architecture use a softmax layer to determine the pixel of a given generation, leading to a discrete prior rather than a continuous

one, which is required for the VQ-VAE for instance. As a result, the distribution of a pixel conditionally to the ones in its receptive field is given by a multinomial in  $\{0, \ldots, 256\}$ .

# 3.3.2 Usage in the VQ-VAE

Once we have trained the VQ-VAE, the original paper states that we can replace the uniform prior on p(z) by a PixelCNN to model the prior. To perform the training of the PixelCNN, we turn the samples x in their latent representations z, and train the PixelCNN over the latent representations z. This way, we have created dependency between the z in the latent feature mapping, and we obtain a prior over their distribution modeled by the PixelCNN.

# 4 Microbiota analysis

This section aims at using the previous elements to design microbiota adapted latent methods.

# 4.1 Microbiota dataset

As we look into the microbiota data, we notice a major phylogenetic architecture to describe various levels of precision in the microbiota composition. Indeed, such phylogenetic structure can be represented as a tree as on the following figure:

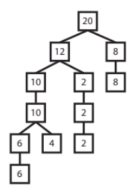


Figure 8: Phylogenetic tree example with abundance data (in the nodes) at each layer of the tree. Each node represents a bacterium specie at a given precision layer in the tree. From [2].

Such structure can not be used directly in a machine learning system since it's not a vectorizable representation. Hence, we first suggest to transform the tree in a matrix to image structure as in the following example:



Figure 9: Phylogenetic tree to image representation: opacity of the pixel relates to the abundance of the species at the given level of precision (normalized between 0 and 1). From [2].

To give some notations, we introduce the following framework:

- The maximum depth of a phylogenetic tree is D, the maximum number of nodes  $N_T$ , the maximum amount of unique species at any level is U.
- $X_i$  is the abundance matrix of the individual i (image of size  $D \times U$ ), which is observed.

- $T_i$  is the adjacent matrix of the phylogenetic tree of individual i, of size  $N_T \times N_T$ , which is observed. We could use any other encoding of a tree (contour function, depth function, ...).
- For a given tree, we will denote by  $U_l$  the number of species at a given precision layer  $\ell$  of the tree.
- We assume that we have n individuals observed through  $(X_i, T_i)_{1 \le i \le n}$  i.i.d samples.

Note that all tree follow the same global architecture, with branches and nodes not always being activated. The following figure illustrates the global architecture at precision layer 4:

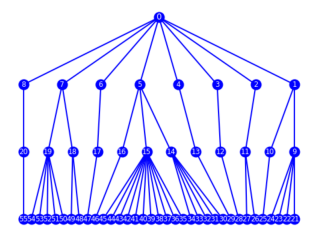


Figure 10: Taxonomy of the microbiota dataset (precision 4)

Notice that at this precision layer, we don't have any missing entries, which facilitates the current modelisation. The next figure illustrates a sample from the dataset as well, as taxa-abundance data represented through opacity.

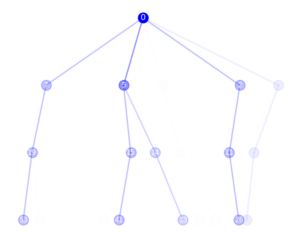


Figure 11: Taxa-abundance sample from the dataset (opacity relates the abundance value)

# 4.2 Mathematical context and objective without latent variables

In first approximation, we would like to define a generative model that does not exploit any latent structure. Such model, parameterized by  $\theta$ , aims at finding an optimal distribution in

the sense of the maximum of likelihood, within a family of distributions yet to be defined. The maximum likelihood objective is given below as:

$$\theta^* = \arg\max_{\theta} p_{\theta}(X, T)$$

One can rewrite the joint distribution as follow:

$$p_{\theta}(X,T) = \prod_{i=1}^{n} p_{\theta}(X_i, T_i)$$
$$= \prod_{i=1}^{n} p_{\theta}(T_i) p_{\theta}(X_i | T_i)$$

As a result, to compute this objective, we need to define a prior  $p_{\theta}(T_i)$  that generates trees, and a posterior distribution  $p_{\theta}(X_i|T_i)$  that generates abundance data from a sampled tree.

# 4.3 Markovian model without latent variables

# 4.3.1 Design of the prior

We aim at defining a parameterized distribution  $p_{\theta}(T)$  from which one can sample trees. We would also like this distribution to model the trees of the microbiota dataset, meaning that the generated trees should look like the ones from the dataset as well, and respect the phylogenetic constraints.

Consequently, we introduce a first simple generative process to characterize  $p_{\theta}(T)$  that we would call the markovian parenting tree generator. Before describing the generation method, let us introduce the framework of it:

- Describe T as a succession of L layers:  $T = (T^{(1)}, \ldots, T^{(L)})$ . We assume that we have no missing data, so to say, all the leaves the tree are reaching the precision layer L.
- Describe a given layer  $\ell$  as a discrete vector in  $\{0,1\}^U$ . To each possible node at layer  $\ell$  we can associate an index k so that we denote the nodes by  $u_k^{(\ell)} \in \{0,1\}$ .

$$T^{(\ell)} = (u_1^{(\ell)}, \dots, u_{K_\ell}^{(\ell)})$$

A node  $u_k^{(\ell)}$  is activated if it is valued as 1 in  $T^{(\ell)}$ , otherwise it is not.

• We introduce the function  $\mathcal{P}$  that takes a node  $u_k^{(\ell)}$  as input, and output the parent of the node if it is well defined in T, otherwise it outputs 0 as if the parent was not activated:

$$\mathcal{P}(u_k^{(\ell)}) = \begin{cases} 1 & \text{if parent of } u_k^{(\ell)} \text{ exists and is activated} \\ 0 & \text{otherwise} \end{cases}$$

Now that the framework is clear and defined, we describe the generative process:

• The root node of the tree is deterministic, since all trees begin to the same root ancestor. Hence, we have:

$$p(T^{(1)}) = \delta_{e_1}(T^{(1)})$$

• For all  $l \geq 2, k \in \{1, \dots, U\}$ , we assume that the activation of the node  $u_k^{(\ell)}$  is distributed as a Bernoulli conditionally to its parent activation, parameterized by  $\pi_k^{(\ell)}$ :

$$u_k^{(\ell)} | \left\{ \mathcal{P}(u_k^{(\ell)}) = 1 \right\} \sim \mathcal{B}(\pi_k^{(\ell)})$$

To respect the tree architecture, we assume that if  $\mathcal{P}(u_k^{(\ell)}) = 0$  then the probability for the children to be activated is deterministic and set to 0, as a child can not exist without his parent.

• For now, we make the major assumption that all nodes are independent within a given layer conditionally to their parents, and they only depend on their respective parent, so that:

$$p\left(u_1^{(\ell)}, \dots, u_{K_{\ell}}^{(\ell)} | \mathcal{P}(u_1^{(\ell)}), \dots, \mathcal{P}(u_{K_{\ell}}^{(\ell)})\right) = \prod_{k=1}^{K_{\ell}} p\left(u_k^{(\ell)} | \mathcal{P}(u_k^{(\ell)})\right)$$

• The dependency between the layers of the tree is markovian:

$$p_{\theta}(T^{(\ell+1)}|T^{(1:\ell)}) = p_{\theta}(T^{(\ell+1)}|T^{(\ell)})$$

Noting these framework properties, we can describe the log-likehood of such prior model on the trees using proposition 5.1:

$$p_{\theta}(T_1, \dots, T_n) = \sum_{i=1}^n \sum_{k=1}^{L-1} \sum_{k=1}^{K_{\ell}} \mathcal{P}(u_k^{(\ell+1)}) \left[ u_k^{(\ell+1)} \log \pi_k^{(\ell+1)} + (1 - u_k^{(\ell+1)}) \log(1 - \pi_k^{(\ell+1)}) \right]$$

Looking at the previous formula, we obtain that such prior is parameterized by the activation probabilities  $\pi_k^{(\ell)}$  of each node  $u_k^{(\ell)}$ . Furthermore, due to the indicator function expressed through  $\mathcal{P}(u_k^{(\ell+1)})$ , the update of a given activation probability will only be impacted by the trees which have the node  $u_k^{(\ell)}$  in any branch.

Now that the prior of the trees is well defined, we would like to compute an optimal value of  $\pi^{(\ell)} = \left(\pi_1^{(\ell)}, \dots, \pi_U^{(\ell)}\right)$  in the sense of the maximum of likelihood. Using proposition 5.2, we obtain

$$\left(\pi_k^{(\ell)}\right)^* = \frac{\sum_{i=1}^n \mathcal{P}(u_{k,i}^{(\ell)}) u_{k,i}^{(\ell)}}{\sum_{i=1}^n \mathcal{P}(u_{k,i}^{(\ell)})}$$

This estimator is actually the common MLE for a Bernoulli parameter estimation, except that it limits the computation of the estimation to all trees that respect the root constraint, and that could possess the node  $u_k^{(\ell)}$  since they must have the parent node  $\mathcal{P}(u_k^{(\ell)})$ .

# 4.3.2 Design of the posterior and maximum likelihood estimator

Now that we have a way to generate trees, we need to define an explicit stochastic relationship between the abundance data and the structure of the tree. Such inevitably exists, as when observing T, the abundance of an entity that isn't present in T is necessarily 0. Similarly, it is highly likely that when observing the presence of certain entities in T that induces a high abundance of another neighbour entity (interaction between bacteria).

For context, we recall that X is a matrix of shape (L, U), where L is the maximum precision level of the trees (assumed to be the same for all trees for now) and U the maximum number of entities at each depth of the trees. We denote by  $X^{(\ell)}$  the  $\ell$ -th line of the abundance matrix, for which up to  $U_{\ell}$  elements should be non-zero.

We assume the following framework:

- Let  $X = (X^1, \dots, X^{(L)})$  a given abundance matrix associated to a tree T.
- We denote  $X^{(\ell)} = (x_1^{(\ell)}, \dots, x_{K_\ell}^{(\ell)})$  the abundance vector at layer  $\ell$ .
- For an abundance node  $x_k^{(\ell)}$ , we denote by  $\mathcal{C}(x_k^{(\ell)})$  the set of abundance children associated to that node. Also, we introduce the notation  $T_k^{(\ell+1)}$ , the vector of activation states  $(u_j^{(\ell+1)})_j$  of the children of the node  $u_k^{(\ell)}$  in T (restriction of  $T^{(\ell)}$  to the children of the node  $u_k^{(\ell)}$ ).
- $X^{(1)} = [1, 0, \dots, 0]$ , since it's the root of the tree, only one entity gets the whole weight.
- Since we would like a simple explicit model at first, we design a posterior  $p_{\theta}(X|T)$  which is markovian relatively to the layers of the tree, so that the abundance at the next layer are only impacted by the previous layers abundance for now:

$$p_{\theta}(X^{(\ell)}|X^{(1:\ell-1)},T) = p_{\theta}(X^{(\ell)}|X^{(\ell-1)},T^{(\ell)})$$

• In addition to the markovianity, we assume that at a given layer, the abundance only depends of their respective parent and their siblings:

$$p(X^{(\ell+1)}|X^{(\ell)}, T^{(\ell+1)}) = \prod_{k=1}^{K_{\ell}} p(\mathcal{C}(x_k^{(\ell)})|x_k^{(\ell)}, T^{(\ell+1)})$$

- Each value in  $X^{(\ell)}$  is restricted by the following set of constraints due to the nature of the abundance in a tree structure:
  - If node k at layer  $\ell-1$  has one child, then its abundance value is the same for the child node
  - If node k at layer  $\ell-1$  has at least two children, the children abundance sums to the parent's abundance value.

• Since we deal with proportions in abundance vectors, it seems natural to use the Dirichlet distribution in first assumption. Hence, we assume that for all  $l \geq 2$ , if  $|\mathcal{C}(x_k^{(\ell)})| > 1$ ,

$$\mathcal{C}(x_k^{(\ell)})|x_k^{(\ell)}, T \sim x_k^{(\ell)} \mathcal{D}(\alpha_k^{(\ell)} \odot T_k^{(\ell+1)})$$

We denote by  $f_{\alpha_k^{(\ell)} \odot T^{(\ell+1)}}$  the density of this distribution, parameterized by  $\alpha_k^{(\ell)} \odot T^{(\ell+1)}$  which denotes a masked version of  $\alpha_k^{(\ell)}$  relatively to the activated nodes of the tree at  $T^{(\ell)}$ . Notice that setting this distribution framework enables us to verify the constraints given above by renormalizing the layer with the obtained weights. Naturally, if  $|\mathcal{C}(x_k^{(\ell)})| = 1$ , we have  $\mathcal{C}(x_k^{(\ell)}) \sim \delta_{x_k^{(\ell)}} \left(\mathcal{C}(x_k^{(\ell)})\right)$  to respect the constraints.

Noting the previous framework, the whole abundance distribution conditionally to a tree T can be written as (see proposition 5.4):

$$p(X|T) = \delta_{e_1}(X^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_{\ell}} \left( \delta_{x_k^{(\ell)}} \left( \mathcal{C}(x_k^{(\ell)}) \right)^{\mathbb{1}_{|\mathcal{C}(x_k^{(\ell)})|=1}} \frac{1}{x_k^{(\ell)}} f_{\alpha_k^{(\ell)} \odot T^{(\ell+1)}} \left( \frac{\mathcal{C}(x_k^{(l)})}{x_k^{(\ell)}} \right)^{\mathbb{1}_{|\mathcal{C}(x_k^{(\ell)})|>1}} \right)$$

One can then compute a maximum likelihood estimator of each  $\alpha_k^{(\ell)}$  using a fixed point iterative algorithm that is described in the appendix as proposition 5.5:

$$\boxed{\alpha_{k,v}^{(\ell)} \leftarrow \psi^{-1} \left( \frac{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[ \psi \left( \sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)} \right) + \log \frac{\left[ \mathcal{C}(x_{k,i}^{(\ell)}) \right]_{v}}{x_{k,i}^{(\ell)}} \right]}{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})}} \right)}$$

# 4.3.3 Optimization of the objective

Recall the optimization objective, written under the maximum of the log-likelihood:

$$\theta^* = \arg\max_{\theta} \log p_{\theta}(X, T)$$

$$= \arg\max_{\theta} \sum_{i=1}^{n} \log p_{\theta}(X_i, T_i)$$

$$= \arg\max_{\theta} \sum_{i=1}^{n} \log p_{\theta}(T_i) + \log p_{\theta}(X_i | T_i)$$

Notice that, in our context, this objective is separable since the prior and posterior do not share any common parameter in  $\theta$ . Hence, the optimal  $\theta^*$  is given by the concatenation of the MLE from the prior on the trees and the posterior of the abundance knowning the trees, which we both have computed in the previous sections.

### 4.3.4 Experiments

We implement the previous model to form a baseline to which we can compare the upcoming latent models to. Before we work with real data, we generate an artificial dataset based on the following structure.

We define the true parameters of the model:

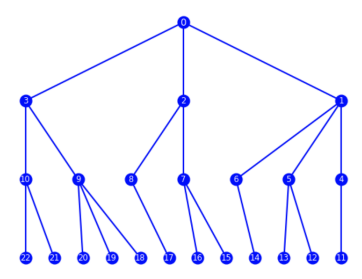


Figure 12: Global artificial tree structure

- The activation probabilities for each node:  $\pi_k^{(\ell)}$
- The abundance parameters for each parent node:  $\alpha_k^{(\ell)}$

Using the true parameters, we define the prior and posterior of the previous model to generate a dataset. The following figure illustrates a sample of that dataset.

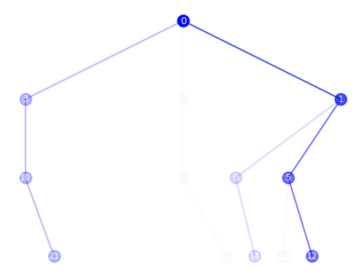


Figure 13: Artificial taxa-abundance data following the global structure of figure 12. Opacity accounts for the value of the abundance.

Now that we have a dataset, we train another set of 10 priors and posteriors onto the artifical data study the convergence of the model and the absolute error to the parameters. The following figure illustrates the convergence of the 10 models for the maximum likelihood objective, relatively to the the number of iterations done for the fixed point algorithm to compute the abundance parameters.

### Log-likelihood / iteration

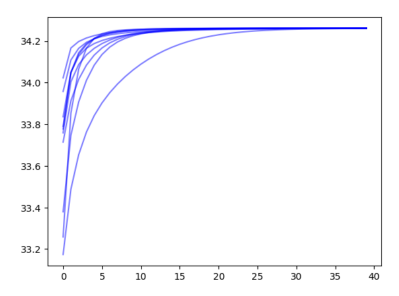


Figure 14: Log-likelihood per iteration of the optimization algorithm for 10 models being trained with random initializations.

Looking at figure 14, it seems that the optimization algorithm is converging to a given  $\widehat{\theta}^*$ . We would then like to compare such  $\widehat{\theta}^*$  to the true set of parameters  $\theta^*$ . The following figure illustrates the mean distance of the parameters to the original one:  $\|\theta^* - \widehat{\theta}^*\|_2$ .

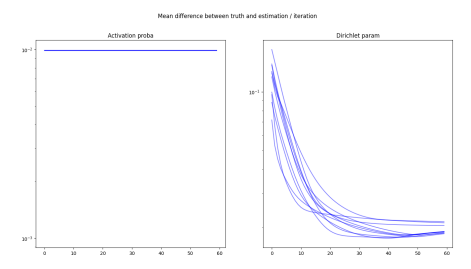


Figure 15: Average log optimization error to the real parameters for 10 models initialized with random parameters, per iteration.

Since the optimal value of  $\pi_k^{(\ell)}$  is explicit, the error does not change over the iterations of the fixed point algorithm. On the other hand, the dirichlet parameters  $\alpha_k^{(\ell)}$  are optimized coordinate per coordinate in the fixed point algorithm, leading to the convergence profile of figure 15, which shows that there is a convergence to a given  $\widehat{\theta}^*$  that is close to  $\theta^*$  with mean error rate of approximately  $10^{-2}$ .

The next graph illustrates the error per coordinate of the estimated parameter and the true one for the 10 trained models.

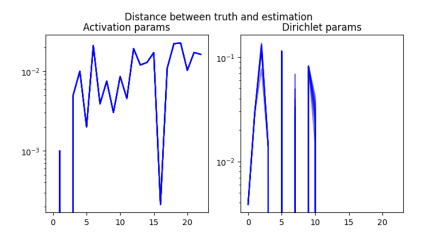


Figure 16: Coordinate-wise log optimization error to the real parameters for 10 models initialized with random parameters, per iteration.

As it seems, some coordinates are much more accurately determined than others, which highly relates to the structure of the tree (markovian) and the general presence of each node in the dataset. Indeed, the less likely a node is activated, the less samples we have to compute its activation probability. Likewise, the deeper a node is in the tree, the less likely it is to be present in a tree due to the conditional probability to the parent. That can then explain why deeper nodes are harder to grasp, and for which the parameter estimation is less good than the one of higher entities in the hierarchy of the tree.

Furthermore, note that we used an initialization of the  $\pi_k^{(\ell)}$  to 0, which artificially boosts the estimation quality of some node like the 16-th one, which has a probability of activation of 0.1 in our artificial model. Hence, while the estimation seems to be great for that node, this is just biased by the fact that this node rarely is present and that the initialization is quite close to the truth already.

Finally, we look into the average optimization error as we vary the amount of samples in figure 17.



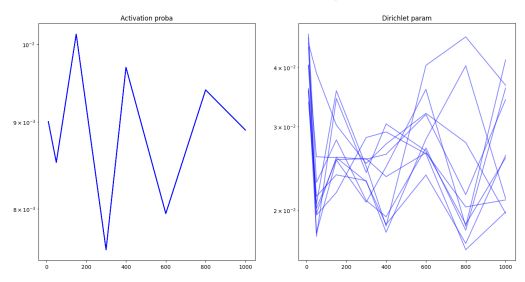


Figure 17: Average log optimization error to the real parameters for 10 models initialized with random parameters, per samples in the dataset.

As it seems, the number of samples is not impacting much the convergence towards the parameters of the models, which seems unnatural. We would like to observe that with more samples, the variance is reducing and the estimation getting better.

# 4.3.5 Conclusions

This first model is interesting as it provides a benchmark baseline for our upcoming tree-structured models. However, it clearly lacks of complexity:

- The nodes are modeled as independent, which prevents any modelisation of correlation between entities. As for many ecosystems, we would expect some bacteria to have symbiotic relationship, or domination roles, especially when it comes to critical systems like disease detection in which some bacteria may proliferate over others. One idea could be to model an interaction graph (see [5]) and use it as a correlation restriction between our bacteria.
- The abundance data generation takes the tree constraints into account, but the Dirichlet distribution is not quite expressive. Mixture models could enable us to provide more expressive priors and model multiple modes rather than one at the moment, to the cost of more parameters though.
- So far we have only considered trees without any missing entries at precision level L, which is not what we observe for microbiota datasets. Inference for missing data implementation could be interesting in the future.

# 4.4 Taxonomy clustering: mixture model

### 4.4.1 Context

A first natural extension to improve the model introduced in 4.3 would be to take into account a latent clustering of the microbiota. Indeed, before looking at functional areas, it is highly likely that some people may share a similar taxonomic architecture.

Introducing that latent clustering can be done by modelizing a hidden variable that we will generally denote as Z. We then define the following framework, which stands as a mixture of the previously introduced model 4.3 over the latent clusters:

- Denote  $Z \in \{1, ..., C\}$  the latent taxonomic clustering variable.
- Denote  $p(Z=c) = \gamma_c$  the probability of cluster c.
- Assume that T and X distributions parameters are now conditioned by the value of Z:

$$p(Z = c) = \gamma_c$$

$$p\left(u_k^{(\ell)} = 1 | \mathcal{P}(u_k^{(\ell)}) = 1, Z = c\right) = \pi_{c,k}^{(l)}$$

$$p\left(\mathcal{C}(x_k^{(\ell)}) | x_k^{(\ell)}, T^{(\ell+1)}, Z = c\right) \sim x_k^{(\ell)} \mathcal{D}(\alpha_{c,k}^{(l)} \odot T_k^{(\ell+1)}) \quad \text{if } |\mathcal{C}(x_k^{(\ell)})| > 1$$

# 4.4.2 Optimization

The usual optimization objective for this model would be the maximum likelihood regarding the observations, such that:

$$\theta^* = \arg\max_{\theta} p_{\theta}(T, X)$$

However, this does not implement the latent structure that we are aiming for, as we do not observe Z. Hence, using the Expectation-Maximization algorithm presented in 2, we introduce a suboptimal objective on the complete joint distribution:

$$\theta^* = \arg\max_{\theta} \underbrace{\mathbb{E}_{p_{\widehat{\theta}}(Z|T,X)} \left[\log p_{\theta}(T,X,Z)|T,X\right]}_{Q(\widehat{\theta},\theta)}$$

Such optimal  $\theta^*$  can be computed through a mix of explicit estimators and a fixed point algorithm detailled in proposition 5.6. Notice that the computed estimator of  $\theta^*$  is quite close to the one presented in section 4.3, as we simply introduced a global architecture clustering based on a mixture of the simple generative model.

## 4.4.3 Experiments

# 5 Appendix

# 5.1 Microbiota analysis

Proposition 5.1. (Bernoulli tree prior)

Let  $T = (T^{(1)}, \dots, T^{(L)})$  a random tree of depth L.

For all  $\ell \in \{1, \ldots, L\}$ , denote  $T^{(\ell)} = (u_1^{(\ell)}, \ldots, u_{K_\ell}^{(\ell)})$  the nodes at layer  $\ell$ , such that  $u_k^{(\ell)} \in \{0, 1\}$  denotes whether or not the node is activated (1 if activated).

Denote  $\mathcal{P}(u_k^{(\ell)})$  the parent of the node  $u_k^{(\ell)}$  that outputs 1 if the parent exists and is activated, 0 otherwise.

Assume that:

• 
$$p(T^{(1)}) = \delta_{e_1}(T^{(1)})$$

• 
$$\forall l \ge 2, p(T^{(\ell+1)}|T^{(1:\ell)}) = p(T^{(\ell+1)}|T^{(\ell)})$$

• 
$$\forall l \ge 2, k \in \{1, \dots, K_{\ell}\}, u_k^{(\ell)} | \left\{ \mathcal{P}(u_k^{(\ell)}) = 1 \right\} \sim \mathcal{B}(\pi_k^{(\ell)})$$

• 
$$p\left(u_1^{(\ell)}, \dots, u_{K_\ell}^{(\ell)} | \mathcal{P}(u_1^{(\ell)}), \mathcal{P}(u_{K_\ell}^{(\ell)})\right) = \prod_{k=1}^{K_\ell} p\left(u_k^{(\ell)} | \mathcal{P}(u_k^{(\ell)})\right)$$

Then, the distribution of T can be written as:

$$p(T) = \delta_{e_1}(T^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_\ell} \left( \left( \pi_k^{(\ell+1)} \right)^{u_k^{(\ell+1)}} \left( 1 - \pi_k^{(\ell+1)} \right)^{1 - u_k^{(\ell+1)}} \right)^{\mathcal{P}(u_k^{(\ell+1)})}$$

Proof.

$$\begin{split} p(T) &= p\left(T^{(1)}, \dots, T^{(L)}\right) \\ &= p\left(T^{(1)}\right) \prod_{l=1}^{L-1} p\left(T^{(\ell+1)} | T^{(\ell)}\right) \\ &= \delta_{e_1}(T^{(1)}) \prod_{l=1}^{L-1} p\left(u_1^{(\ell+1)}, \dots, u_{K_\ell}^{(\ell+1)} | \mathcal{P}(u_1^{(\ell+1)}), \dots, \mathcal{P}(u_{K_\ell}^{(\ell+1)})\right) \\ &= \delta_{e_1}(T^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_\ell} p\left(u_k^{(\ell+1)} | \mathcal{P}(u_k^{(\ell+1)})\right) \\ &= \delta_{e_1}(T^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_\ell} \left(\pi_k^{(\ell+1)} u_k^{(\ell+1)} + \left(1 - \pi_k^{(\ell+1)}\right) (1 - u_k^{(\ell+1)})\right)^{\mathcal{P}(u_k^{(\ell+1)})} \\ &= \delta_{e_1}(T^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_\ell} \left(\left(\pi_k^{(\ell+1)}\right)^{u_k^{(\ell+1)}} \left(1 - \pi_k^{(\ell+1)}\right)^{1 - u_k^{(\ell+1)}}\right)^{\mathcal{P}(u_k^{(\ell+1)})} \end{split}$$

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**Proposition 5.2** (MLE of the Bernoulli tree prior). Recall the context of 5.1. Let  $(T_1, \ldots, T_n)$  n trees i.i.d following the bernoulli tree prior. Denote the maximum likelihood objective by

$$\underset{\pi_{j}^{(m)}}{\arg\max} \quad \sum_{i=1}^{n} \sum_{l=1}^{L-1} \sum_{k=1}^{K_{\ell}} \mathcal{P}(u_{k,i}^{(\ell+1)}) \left[ u_{k,i}^{(\ell+1)} \log \pi_{k}^{(\ell+1)} + (1 - u_{k,i}^{(\ell+1)}) \log(1 - \pi_{k}^{(\ell+1)}) \right] \\
s.t. \quad \forall k, l, \pi_{k}^{(\ell)} \in [0, 1]$$
(1)

Then the maximum likelihood estimator is given by

$$\left(\pi_k^{(\ell)}\right)^* = \frac{\sum_{i=1}^n \mathcal{P}(u_{k,i}^{(\ell)}) u_{k,i}^{(\ell)}}{\sum_{i=1}^n \mathcal{P}(u_{k,i}^{(\ell)})}$$

*Proof.* Simply by deriving the objective we obtain:

$$\partial_{\pi_{j}^{(m)}} \log p(T_{1}, \dots, T_{n}) = \sum_{i=1}^{n} \mathcal{P}(u_{j,i}^{(m)}) \left[ u_{j,i}^{(m)} \frac{1}{\pi_{j}^{(m)}} + (1 - u_{j,i}^{(m)}) \frac{-1}{1 - \pi_{j}^{(m)}} \right]$$

$$= \frac{1}{\pi_{j}^{(m)}} \sum_{i=1}^{n} \mathcal{P}(u_{j,i}^{(m)}) u_{j,i}^{(m)} - \frac{1}{1 - \pi_{j}^{(m)}} \sum_{i=1}^{n} \mathcal{P}(u_{j,i}^{(m)}) (1 - u_{j,i}^{(m)})$$

Looking for 0 valued gradient, we end up with:

$$(1 - \pi_j^{(m)}) \sum_{i=1}^n \mathcal{P}(u_{j,i}^{(m)}) u_{j,i}^{(m)} - \pi_j^{(m)} \sum_{i=1}^n \mathcal{P}(u_{j,i}^{(m)}) (1 - u_{j,i}^{(m)}) = 0$$

$$\pi_j^{(m)} \sum_{i=1}^n \mathcal{P}(u_{j,i}^{(m)}) = \sum_{i=1}^n \mathcal{P}(u_{j,i}^{(m)}) u_{j,i}^{(m)}$$

$$\pi_j^{(m)} = \frac{\sum_{i=1}^n \mathcal{P}(u_{j,i}^{(m)}) u_{j,i}^{(m)}}{\sum_{i=1}^n \mathcal{P}(u_{j,i}^{(m)})}$$

Since the obtained value respects the constraint, that concludes the proof.

**Proposition 5.3** (Law of aX). Let  $X \sim f(x)$  a random variable with density f. Let  $a \in (0,1)$ .

The distribution of aX is given by:

$$p_{aX}(u) = \frac{1}{a} f\left(\frac{u}{a}\right)$$

*Proof.* We use the transfer theorem. Let g a continuous and measurable function.

$$\mathbb{E}[g(aX)] = \int g(ax)f(x)dx$$
$$= \int g(u)f\left(\frac{u}{a}\right)\frac{1}{a}du$$

Hence, one can identify the distribution of aX as  $p_{aX}(u) = \frac{1}{a}f\left(\frac{u}{a}\right)$ 

**Proposition 5.4** (Abundance distribution conditionally to the trees). Let  $X = (X^{(1)}, \dots, X^{(L)})$ 

the abundance matrix of a tree T with depth L. For all  $\ell \in \{1, \ldots, L\}$ , denote  $X^{(\ell)} = (x_1^{(\ell)}, \ldots, x_{K_\ell}^{(\ell)})$  the abundance value of each node of the

Denote  $\mathcal{C}(x_k^{(\ell)})$  the vector of children abundances related to  $x_k^{(\ell)}$ , which is empty if it has no

Denote  $T_k^{(\ell+1)}$  the vector of activation states  $(u_j^{(\ell+1)})_j$  of the children of the node  $u_k^{(l)}$  in T. Assume that:

• 
$$p(X^{(1)}) = \delta_{e_1}(X^{(1)})$$

• 
$$p(X^{(\ell+1)}|X^{(1:\ell)},T) = p(X^{(\ell+1)}|X^{(\ell)},T^{(\ell+1)})$$

• 
$$p(X^{(\ell+1)}|X^{(\ell)}, T^{(\ell+1)}) = \prod_{k=1}^{K_{\ell}} p(\mathcal{C}(x_k^{(\ell)})|x_k^{(\ell)}, T^{(\ell+1)})$$

• For all 
$$l \in \{2, ..., L\}, k \in \{1, ..., K_{\ell}\},$$

$$- |If |\mathcal{C}(x_k^{(\ell)})| > 1: |\mathcal{C}(x_k^{(\ell)})| x_k^{(\ell)}, T^{(\ell+1)} \sim x_k^{(\ell)} D(\alpha_k^{(\ell)} \odot T_k^{(\ell+1)})$$

$$- If |\mathcal{C}(x_k^{(\ell)})| = 1, \ \mathcal{C}(x_k^{(\ell)})|x_k^{(\ell)}, T^{(\ell+1)} \sim \delta_{x_k^{(\ell)}} \left( \mathcal{C}(x_k^{(\ell)}) \right)$$

Then, the distribution of X conditionally to T is given by:

$$p(X|T) = \delta_{e_1}(X^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_{\ell}} \left( \delta_{x_k^{(\ell)}} \left( \mathcal{C}(x_k^{(\ell)}) \right)^{\mathbb{1}_{|\mathcal{C}(x_k^{(\ell)})|=1}} \frac{1}{x_k^{(\ell)}} f_{\alpha_k^{(\ell)} \odot T_k^{(\ell+1)}} \left( \frac{\mathcal{C}(x_k^{(\ell)})}{x_k^{(\ell)}} \right)^{\mathbb{1}_{|\mathcal{C}(x_k^{(\ell)})|>1}} \right)$$

Proof.

$$p(X|T) = p(X^{(1)}, \dots, X^{(L)}|T)$$

$$= p(X^{(1)}|T^{(1)}) \prod_{l=1}^{L-1} p(X^{(\ell+1)}|X^{(\ell)}, T^{(\ell+1)})$$

$$= \delta_{e_1}(X^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_{\ell}} p(\mathcal{C}(x_k^{(\ell)})|x_k^{(\ell)}, T^{(\ell+1)})$$

$$= \delta_{e_1}(X^{(1)}) \prod_{l=1}^{L-1} \prod_{k=1}^{K_{\ell}} \left( \delta_{x_k^{(\ell)}} \left( \mathcal{C}(x_k^{(\ell)}) \right)^{\mathbb{1}_{|\mathcal{C}(x_k^{(\ell)})|=1}} \left[ \underbrace{\frac{1}{x_k^{(\ell)}} f_{\alpha_k^{(\ell)} \odot T_k^{(\ell+1)}} \left( \frac{\mathcal{C}(x_k^{(\ell)})}{x_k^{(\ell)}} \right)}_{\text{proposition 5.3}} \right]^{\mathbb{1}_{|\mathcal{C}(x_k^{(\ell)})|>1}}$$

**Proposition 5.5** (MLE of the abundance distribution a posteriori). Consider the context of proposition 5.4.

We consider a data set  $(X_i, T_i)_{1 \ge i \ge n}$  of abundance matrix and trees. Denote by  $\mathcal{V}(\alpha_k^{(\ell)}, T) = \left\{ v \in \mathbb{N} \middle| \left[ \alpha_k^{(\ell)} \odot T_k^{(\ell+1)} \right]_v \ne 0 \right\}$  the set of indexes so that the coordinate

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of  $\alpha_k^{(\ell)}$  is not masked by  $T^{(\ell+1)}$ .

The maximum likelihood estimator of the distribution characterising the abundance conditionally to the tree is then given by the following fixed point algorithm:

$$\forall l \in \{1, \dots, L\}, k \in \{1, \dots, K_{\ell}\}, v \in \{1, \dots, |\alpha_k^{(\ell)}|\},$$

$$\alpha_{k,v}^{(\ell)} \leftarrow \psi^{-1} \left( \frac{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[ \psi\left(\sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)}\right) + \log \frac{\left[\mathcal{C}(x_{k,i}^{(\ell)})\right]_{v}}{x_{k,i}^{(\ell)}} \right]}{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})}} \right)$$

*Proof.* The maximum likelihood objective can be written as:

$$\arg\max_{\alpha_{j,v}^{(m)}} \quad \sum_{i=1}^{n} \sum_{l=1}^{L-1} \sum_{k=1}^{K_{\ell}} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \log f_{\alpha_{k}^{(\ell)} \odot T_{k,i}^{(\ell+1)}} \left( \frac{C(x_{k,i}^{(\ell)})}{x_{k,i}^{(\ell)}} \right)$$

Using the Dirichlet distribution expression, we can write the following:

$$\log f_{\alpha_k^{(\ell)} \odot T_{k,i}^{(\ell+1)}} \left( \frac{C(x_{k,i}^{(\ell)})}{x_{k,i}^{(\ell)}} \right) = \log \Gamma \left( \sum_{v \in \mathcal{V}(\alpha_k^{(\ell)}, T_i)} \alpha_{k,v}^{(\ell)} \right) - \sum_{v \in \mathcal{V}(\alpha_k^{(\ell)}, T_i)} \Gamma \left( \alpha_{k,v}^{(\ell)} \right) + \sum_{v \in \mathcal{V}(\alpha_k^{(\ell)}, T_i)} \left( \alpha_{k,v}^{(\ell)} - 1 \right) \log \frac{\left[ \mathcal{C}(x_{k,i}^{(\ell)}) \right]_v}{x_{k,i}^{(\ell)}}$$

Writing  $\psi$  the digamma function, the derivative of the objective relatively to a fixed  $\alpha_{k,v}^{(\ell)}$  is given by:

$$\partial_{\alpha_{k,v}^{(\ell)}} \log p(X|T) = \sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[ \psi \left( \sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)} \right) - \psi \left( \alpha_{k,v}^{(\ell)} \right) + \log \frac{\left[ \mathcal{C}(x_{k,i}^{(\ell)}) \right]_{v}}{x_{k,i}^{(\ell)}} \right]$$

Looking for 0 valued gradient, we obtain the following equation:

$$\psi\left(\alpha_{k,v}^{(\ell)}\right) = \frac{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[\psi\left(\sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)}\right) + \log\frac{\left[\mathcal{C}(x_{k,i}^{(\ell)})\right]_{v}}{x_{k,i}^{(\ell)}}\right]}{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})}}$$

Using the reference trick from [4], one can compute an iterative fixed point algorithm to solve the previous equation, leading to:

$$\alpha_{k,v}^{(\ell)} \leftarrow \psi^{-1} \left( \frac{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[ \psi \left( \sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)} \right) + \log \frac{\left[ \mathcal{C}(x_{k,i}^{(\ell)}) \right]_{v}}{x_{k,i}^{(\ell)}} \right]}{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})}} \right)$$

**Proposition 5.6** (Expectation-Maximization for Taxonomy Clustering model). We use the same context as for proposition 5.4 of this section. Assume the following framework changes:

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- $(X_i, T_i)_{1 \le i \le n}$  samples of taxa-abundance data  $(X_i \text{ abundance}, T_i \text{ associated taxonomy})$
- Let  $Z_i$  a latent variable taking values in  $\{1, \ldots, C\}$ .
- We model the distributions as follow:

$$p(Z_{i} = c) = \gamma_{c}$$

$$p\left(u_{k,i}^{(\ell)} = 1 | \mathcal{P}(u_{k,i}^{(\ell)}) = 1, Z_{i} = c\right) = \pi_{c,k}^{(\ell)}$$

$$p\left(\mathcal{C}(x_{k,i}^{(\ell)}) | x_{k,i}^{(\ell)}, T^{(\ell+1)}, Z_{i} = c\right) \sim x_{k}^{(\ell)} \mathcal{D}(\alpha_{c,k}^{(\ell)} \odot T_{k}^{(\ell+1)}) \quad \text{if } |\mathcal{C}(x_{k}^{(\ell)})| > 1$$

Consider the optimization problem:

$$\theta^* = \arg\max_{\theta} \mathbb{E}_{p_{\widehat{\theta}}}[\log p_{\theta}(X, T, Z) | X, T]$$

Then, the Expectation-Maximization algorithm solving that problem is given by (for each sample i and cluster index c):

# Expectation step:

$$p_{\widehat{\theta}}(Z_i = c | X_i, T_i) = \tau_{ic} = \frac{\widehat{\gamma}_c p_{\widehat{\theta}}(T_i | Z_i = c) p_{\widehat{\theta}}(X_i | T_i, Z_i = c)}{\sum_{c'=1}^C \widehat{\gamma}_{c'} p_{\widehat{\theta}}(T_i | Z_i = c') p_{\widehat{\theta}}(X_i | T_i, Z_i = c')}$$

where:

$$p_{\widehat{\theta}}(T_i|Z_i = c) = \delta_{e_1}(T_i^{(1)}) \prod_{\ell=1}^{L-1} \prod_{k=1}^{K_\ell} \left[ \left( \widehat{\pi}_{c,k}^{(\ell+1)} \right)^{u_{k,i}^{(\ell+1)}} \left( 1 - \widehat{\pi}_{c,k}^{(\ell+1)} \right)^{1 - u_{k,i}^{(\ell+1)}} \right]^{\mathcal{P}(u_{k,i}^{(\ell+1)})}$$

$$p_{\widehat{\theta}}(X_i|T_i, Z_i = c) = \delta_{e_1}(X_i^{(1)}) \prod_{\ell=1}^{L-1} \prod_{k=1}^{K_\ell} \delta_{x_{k,i}^{(\ell)}} \left( \mathcal{C}(x_{k,i}^{(\ell)}) \right)^{1 |\mathcal{C}(x_{k,i}^{(\ell)})| = 1} \left( \frac{1}{x_{k,i}^{(\ell)}} f_{\widehat{\alpha}_{c,k}^{(\ell)} \odot T_{i,k}^{(\ell+1)}} \left( \frac{\mathcal{C}(x_{k,i}^{(\ell)})}{x_{k,i}^{(\ell)}} \right) \right)^{1 |\mathcal{C}(x_{k,i}^{(\ell)})| > 1}$$

# Maximization step:

$$\gamma_{c}^{*} = \frac{1}{n} \sum_{i=1}^{n} \tau_{ic} 
\left(\pi_{c,k}^{(\ell)}\right)^{*} = \frac{\sum_{i=1}^{n} \mathcal{P}(u_{k,i}^{(\ell)}) u_{k,i}^{(\ell)} \tau_{ic}}{\sum_{i=1}^{n} \mathcal{P}(u_{k,i}^{(\ell)}) \tau_{ic}} 
\alpha_{k,v}^{(\ell)} \leftarrow \psi^{-1} \left(\frac{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[\psi\left(\sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)}\right) + \log \frac{\left[\mathcal{C}(x_{k,i}^{(\ell)})\right]_{v}}{x_{k,i}^{(\ell)}}\right] \tau_{ic}}{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \tau_{ic}}\right)$$

*Proof.* We study the two parts of the EM algorithm, starting with the expectation step.

# Expectation step:

Applying Bayes formula, we obtain:

$$\begin{split} p_{\widehat{\theta}}(Z_{i} = c | X_{i}, T_{i}) &= \tau_{ic} \\ &= \frac{p_{\widehat{\theta}}(Z_{i} = c, X_{i}, T_{i})}{\sum_{c'=1}^{C} p_{\widehat{\theta}}(Z_{i} = c', X_{i}, T_{i})} \\ &= \frac{\widehat{\gamma}_{c} p_{\widehat{\theta}}(T_{i} | Z_{i} = c) p_{\widehat{\theta}}(X_{i} | T_{i}, Z_{i} = c)}{\sum_{c'=1}^{C} \widehat{\gamma}_{c'} p_{\widehat{\theta}}(T_{i} = Z_{i} = c') p_{\widehat{\theta}}(X_{i} | T_{i}, Z_{i} = c')} \end{split}$$

Exploiting the model framework, we obtain the following expression for the conditional distribution of the taxonomy:

$$p_{\theta}(T_{i}|Z_{i} = c) = p_{\theta}(T_{i}^{(1)}, \dots, T_{i}^{(L)}|Z_{i} = c)$$

$$= \delta_{e_{1}}(T_{i}^{(1)}) \prod_{\ell=1}^{L-1} p_{\theta}(T_{i}^{(\ell+1)}|T_{i}^{(\ell)}, Z_{i} = c)$$

$$= \delta_{e_{1}}(T_{i}^{(1)}) \prod_{\ell=1}^{L-1} \prod_{k=1}^{K_{\ell}} p_{\theta}(u_{k,i}^{(\ell+1)}|\mathcal{P}(u_{k,i}^{(\ell+1)}), Z_{i} = c)$$

$$= \delta_{e_{1}}(T_{i}^{(1)}) \prod_{\ell=1}^{L-1} \prod_{k=1}^{K_{\ell}} \left[ \left( \pi_{c,k}^{(\ell+1)} \right)^{u_{k,i}^{(\ell+1)}} \left( 1 - \pi_{c,k}^{(\ell+1)} \right)^{1 - u_{k,i}^{(\ell+1)}} \right]^{\mathcal{P}(u_{k,i}^{(\ell+1)})}$$

Similarly, the conditional abundance distribution is given by:

$$p_{\theta}(X_{i}|T_{i}, Z_{i} = c) = p_{\theta}(X_{i}^{(1)}, \dots, X_{i}^{(L)}|T_{i}, Z_{i} = c)$$

$$= \delta_{e_{1}}(X_{i}^{(1)}) \prod_{\ell=1}^{L-1} p_{\theta}(X_{i}^{(\ell+1)}|X_{i}^{(\ell)}, T_{i}^{(\ell+1)}, Z_{i} = c)$$

$$= \delta_{e_{1}}(X_{i}^{(1)}) \prod_{\ell=1}^{L-1} \prod_{k=1}^{K_{\ell}} p_{\theta}(\mathcal{C}(x_{k,i}^{(\ell)})|x_{k,i}^{(\ell)}, T_{i}^{(\ell+1)}, Z_{i} = c)$$

$$= \delta_{e_{1}}(X_{i}^{(1)}) \prod_{\ell=1}^{L-1} \prod_{k=1}^{K_{\ell}} \delta_{x_{k,i}^{(\ell)}} \left(\mathcal{C}(x_{k,i}^{(\ell)})\right)^{\mathbb{I}_{|\mathcal{C}(x_{k,i}^{(\ell)})|=1}} \left(\frac{1}{x_{k,i}^{(\ell)}} f_{\alpha_{c,k}^{(\ell)} \odot T_{i,k}^{(\ell+1)}} \left(\frac{\mathcal{C}(x_{k,i}^{(\ell)})}{x_{k,i}^{(\ell)}}\right)\right)^{\mathbb{I}_{|\mathcal{C}(x_{k,i}^{(\ell)})|>1}}$$

# Maximization step

Recall the general objective:

$$\theta^* = \arg\max_{\theta} Q(\widehat{\theta}, \theta)$$

$$= \arg\max_{\theta} \frac{1}{n} \sum_{i=1}^{n} \sum_{c=1}^{C} (\log \gamma_c + \log p(T_i | Z_i = c) + \log p(X_i | T_i, Z_i = c)) \tau_{ic}$$

We start by optimizing  $\gamma$ . The objective can be written as:

$$arg \max_{\gamma_j} \sum_{i=1}^n \sum_{c=1}^C \tau_{ic} \log \gamma_c$$
s.t. 
$$\sum_{c=1}^C \gamma_c = 1$$

$$\forall c, \gamma_c \ge 0$$
(2)

This optimization problem can be solved using Lagrange duality. KKT conditions are then written as (optimizing relatively to any  $\gamma_c$ ):

$$\frac{1}{\gamma_c} \sum_{i=1}^n \tau_{ic} + \phi - \psi = 0$$

$$\sum_{c=1}^C \gamma_c = 1$$

$$\psi \ge 0$$

$$\forall c, \gamma_c \ge 0$$

$$\psi \gamma_c = 0$$

Using the first identity, we get:

$$\gamma_c = \frac{1}{\psi - \phi} \sum_{i=1}^n \tau_{ic}$$

Let's set  $\phi = 0$ , which enables use to ensure the last identity. Using the second identity, we then have:

$$\sum_{c=1}^{C} \gamma_c = 1 = \sum_{c=1}^{C} \frac{1}{\psi} \sum_{i=1}^{n} \tau_{ic}$$

$$= \frac{1}{\psi} \sum_{i=1}^{n} \sum_{c=1}^{C} \tau_{ic}$$

$$= \frac{1}{\psi} \sum_{i=1}^{n} 1$$

$$= \frac{n}{\psi}$$

Hence, we get  $\psi = n$ , which leads us to the optimal  $\gamma_c$  that verifies the KKT constraints:

$$\gamma_c^* = \frac{1}{n} \sum_{i=1}^n \tau_{ic}$$

Looking at  $\pi_c$  now, the optimization objective can be written as:

$$\arg \max_{\substack{\pi_{c,k}^{(\ell)} \\ \text{s.t.}}} \sum_{i=1}^{n} \sum_{c=1}^{C} \tau_{ic} \log p_{\pi_c}(T_i | Z_i = c) \\
\text{s.t.} \quad \pi_{c,k}^{(\ell)} \in [0, 1]$$
(3)

This objective is very similar to the MLE computed for proposition 5.2, up to a multiplication by  $\tau_{ic}$ . Hence, we refer to the proposition 5.2 for the computation, which leads to the following optimal  $\pi_c$ :

$$\left(\pi_{c,k}^{(\ell)}\right)^* = \frac{\sum_{i=1}^n \mathcal{P}(u_{k,i}^{(\ell)}) u_{k,i}^{(\ell)} \tau_{ic}}{\sum_{i=1}^n \mathcal{P}(u_{k,i}^{(\ell)}) \tau_{ic}}$$

Similarly, the optimization objective of  $\alpha_c$  is close to the one of proposition 5.5, up to a multiplication by  $\tau_{ic}$ . Therefore, we refer to 5.5 for the original computation, that just requires a multiplication by  $\tau_{ic}$  to obtain the following result:

$$\alpha_{k,v}^{(\ell)} \leftarrow \psi^{-1} \left( \frac{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \left[ \psi\left(\sum_{u \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \alpha_{k,u}^{(\ell)}\right) + \log \frac{\left[\mathcal{C}(x_{k,i}^{(\ell)})\right]_{v}}{x_{k,i}^{(\ell)}} \right] \tau_{ic}}{\sum_{i=1}^{n} \mathbb{1}_{|\mathcal{C}(x_{k,i}^{(\ell)})| > 1} \mathbb{1}_{v \in \mathcal{V}(\alpha_{k}^{(\ell)}, T_{i})} \tau_{ic}} \right)$$

# 6 Bibliography

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