Clustering Patients with Tensor Decomposition

Matteo Ruffini

Department of Computer Science Universitat Politècnica de Catalunya Barcelona, Spain.

Ricard Gavaldà

Department of Computer Science Universitat Politècnica de Catalunya Barcelona, Spain.

Esther Limón

Institut Català de la Salut EAP Mataró-7, Barcelona, Spain MRUFFINI@CS.UPC.EDU

GAVALDA@CS.UPC.EDU

ELIMON.MN.ICS@GENCAT.CAT

Abstract

In this paper we present a method for the unsupervised clustering of high-dimensional binary data, with a special focus on electronic healthcare records. We present a robust and efficient heuristic to face this problem using tensor decomposition. We present the reasons why this approach is preferable for tasks such as clustering patient records, to more commonly used distance-based methods. We run the algorithm on two datasets of healthcare records, obtaining clinically meaningful results.

1. Introduction

Clustering patients in groups with similar clinical profiles is a strategic activity for modern healthcare systems. First, similar patients share the need for similar cares, so the system can design specific guidelines to treat and prescribe diagnostic patterns rather than single diagnostics. Also, clear and tested clusters based on comorbidities help clinicians to decide treatments on specific patients. Third, characterizing patient patterns helps the system in planning resources and in performing fair comparisons between institutions. The context of this paper is a project that aims at making analytic techniques such as clustering accessible to healthcare managers, within a tool that allows them to explore and understand the population under their care. We are working both with medical directors at large hospitals, with tens of thousands of admissions per year, but also with planners and policy-makers at regional healthcare agencies providing health services to a few million people. These professionals have little or no training in data science; their organizations often have a statistics department, but asking the department for analysis usually takes weeks, which hinders intuition and innovation, and leads in most cases to abandoning the analysis because of more pressing day-to-day matters. The goal of our project is to make the algorithms in the tool largely autonomous, requiring no ad-hoc tuning before every analysis required by the user, and efficient so that intuitions can be explored almost in real time, even for large populations. In the case of clustering, this setting clearly affects the algorithms that could be used. A literature scan for clustering applications shows that most of the examples that can be found in the literature use standard *distance based* clustering methods, like k-means (see for example Rixen et al., 1996; Kshetri, 2011;

Pérez et al., 2016), or linkage-based methods (see Cohen et al., 2010). These techniques are general purpose algorithms based on the notion of distance, a concept that may lose part of its meaning in high dimensional settings (see Aggarwal et al., 2001; Kriegel et al., 2009), especially when data are sparse and/or categorical as we anticipate to be the case. Additionally, the "right" notion of distance may be different for every application of the algorithm, i.e., may change if the manager is trying to cluster patients with diabetes, young or older population or any other possible profile. Furthermore, designing a distance function is too difficult or error-prone for our intended users, as it may involve non-trivial feature selection and engineering activity.

An alternative line of work uses nonnegative tensor factorization for phenotyping and clustering patients (see Ho et al., 2014b,a; Wang et al., 2015; Perros et al., 2017); the idea is to use several patient-associated sources of data (called modes), like diagnostics, medical procedures, age, to create a tensor that, for each patient, counts the co-occurrences of the observations along the various modes. Decomposing this tensor using nonnegative low-rank decomposition methods one finds k vectors of phenotypes summing up the behavior of the modes in the dataset, and assigns each patient to the cluster/phenotype that most characterizes his/her records. This approach does not assume any statistical generative model for the data, requires to work with several sources of information and needs a high degree of manual interaction to define the modes and interpret the results.

A different class of clustering techniques relies on the distributional properties of the observed variables, using mixture models (see Marin et al., 2005; Melnykov et al., 2010). Mixture models describe the observable data as outcomes of joint probability distributions and present many advantages over the cited techniques: they do not need an a-priori defined distance, their nature of probabilistic graphical models allows natural and objective interpretations and their flexibility allows to work with both single and multiple sources of data. In recent time, learning such models has received attention using methods of moments together with tensor decomposition techniques (see Anandkumar et al., 2014, and the references therein). While traditional techniques, like pure Expectation Maximization (EM) (Dempster et al., 1977), tend to scale too badly for the kind of interactive high-dimensional discovery that we envision, and often perform poorly, methods of moments provide guarantees of speed, stability and results quality that make them suitable for our purposes. This is the approach we take here, focusing on the case of mixture models where all the observable features are conditionally independent binary variables (mixture of independent Bernoulli). We are not aware of any previous application of such methods to patient clustering. In broad terms, these methods work by constructing from a sample some low-rank tensors representing the correlations among the data attributes, then decomposing them to recover the parameters of the mixture model that explains these correlations from a hidden variable. The key obstacle for the mixture of independent Bernoulli tends to be the first step, as there are several well-understood methods for the second. The problem of recovering the moments for mixtures with categorical features, was solved by Jain and Oh (2014) using an optimization algorithm, an approach that reduces the scalability of the method (the technique depends on a factor d^3 for both memory and time requirements, where d is the number of features). An alternative approach for these kind of data may be the so-called multi-view approach from Anandkumar et al. (2014); however this method is prone to produce unreliable results when the data are unbalanced, with some few features that have more weight than the others, requiring in this case a nontrivial preprocessing of the data. We refer to the appendix for a detailed discussion on the related works on spectral methods.

1.1 Our Contributions

The contributions of this paper are the following:

- We advocate for the use of methods moments for clustering datasets of patients. These methods can provide meaningful results, require little or no parameter tuning, do not need ad-hoc distance functions and scale in existing implementations to databases with at least tens of thousands of patients on commodity machines (few seconds on our example datasets, making interactive analysis a serious possibility). Also, unlike distance-based methods, they should not degrade when many irrelevant attributes are present.
- We present an heuristic based on the method of moments to recover the parameters of a mixture of independent Bernoulli. The proposed method, instead of using a tensor-decomposition algorithm to retrieve asymptotically convergent estimates of the unknown parameters, uses this decomposition procedure to generate approximate parameters that are then used to feed EM as starting point. In the practical applications, using this technique, EM converges fast, reaching in few steps surprisingly good optima; also the high scalability allows the usage of this method on high-dimensional datasets. While the main story of this paper would go through equally with several of the existing moments-based methods, the one presented here has a number of advantages (see appendix A.1 for a deeper comparison): it requires the desired number of clusters as sole input, it scales as d^2 (where d is the number of features) it only requires $d \ge k$ (where k is the number of clusters - the multi-view approach requires for example $d \ge 3k$), and is deterministic, not randomized — non-technical users distrust algorithms that provide different results on different runs.
- Finally, we apply our method to two real-world datasets obtained by collaboration with a healthcare agency. The datasets contain data from hospital admissions of two selected profiles: patients with Congestive Heart Failure, and "Tertiary" patients (patients with serious diseases that can only be treated by top specialists or with highly specialized equipment). The data contains essentially the list of ICD9 codes of the diagnostics at hospital admission time. Data is turned into a binary matrix where columns represent diagnostics, so highly sparse, as every patient has most often less than a dozen diagnostics out of several thousand possible ones. The proposed method finds well-characterized clusters in the data, which clinicians find meaningful, novel to some extent, and potentially useful for refining clinical guidelines.

Outline of the paper: Section 2 recalls how to use Naïve Bayes Models to perform clustering. Section 3 presents the proposed learning method and Section 5 contains our results on patient clustering.

2. Naïve Bayes Clustering

In this section we characterize the Naïve Bayes Model (NBM), together with an indication on how to use it to cluster data. Even if we are mainly interested on binary data, we keep the terms more general, as the presented results apply to all NBMs.

Definition 2.1 A Naïve Bayes model (NBM) is a set of d+1 variables (Y, x_1, \ldots, x_d) where Y is a hidden (unobservable) discrete variable with a finite number of possible outcomes: $Y \in \{1, \ldots, k\}$. We define $\omega_h := \mathbb{P}(Y = h)$ and $\omega := (\omega_1, \ldots, \omega_k)^\top$. The vector $X = (x_1, \ldots, x_d)$ is observable,

its outcomes depend on the value of the hidden variable Y and the random variables x_1, \ldots, x_d are conditionally independent given Y; we define $\mu_{i,j} = \mathbb{E}(x_i|Y = j)$, and $M = (\mu_{i,j})_{i,j} \in \mathbb{R}^{d \times k}$. Also we will denote with μ_i the set of columns of M: $M = [\mu_1|, ..., |\mu_k]$.

The generative model first generates an unobservable outcome for Y, say h, and then generates a set of observations x_1, \ldots, x_d , depending on h. If the parameters (M, ω) of a NBM are known, they can be used for clustering purposes, assigning an observation to the component with the highest probability of having generated it.

Example 2.1 (Mixture of independent Bernoulli) When the observables x_i are binary, their conditional expectations coincide with the probability of a positive outcome: $\mu_{i,j} = \mathbb{P}(x_i = 1 | Y = j)$. In this fashion, we can derive a closed form of the probability that a given observation X has been generated by a given hidden variable realization Y = j, $\mathbb{P}(Y = j | X)$, and perform clustering by assigning $Cluster(X) = argmax_{j=1,...,k}(\mathbb{P}(Y = j | X))$, where, in this case:

$$\mathbb{P}(Y = j | X) \propto \omega_j \prod_{i=1}^d \mu_{i,j}^{x_i} (1 - \mu_{i,j})^{1 - x_i}$$
(1)

Such Naïve Bayes Clustering procedure requires to know the conditional distribution of the observable data and the model parameters (M, ω) . However, in the real world, these parameters are never known, and need to be estimated from a sample of observations of the observable features. In all the paper, we will assume to deal with a dataset $\bar{X} \in \mathbb{R}^{N \times d}$ of size N of observations denoted as:

$$(\bar{X})_i = X^{(i)} = (x_1^{(i)}, \dots, x_d^{(i)}), \ i \in \{1, \dots, N\}.$$

In the next sections we will present a method of moments to infer the pair (M, ω) from a sample.

3. Learning NBM parameters

When learning the parameters of a Latent Variable Model (LVM) with a tensor method, the typical approach consists in two steps. First we need to manipulate the observable moments in order to express them in the form of symmetric-low rank tensors, to approximate the following operators:

$$M_1 := \sum_{i=1}^k \omega_i \mu_i, \quad M_2 := \sum_{i=1}^k \omega_i \mu_i \otimes \mu_i, \quad M_3 := \sum_{i=1}^k \omega_i \mu_i \otimes \mu_i \otimes \mu_i$$
(2)

where $M_1 \in \mathbb{R}^d$, $M_2 \in \mathbb{R}^{d \times d}$ and $M_3 \in \mathbb{R}^{d \times d \times d}$ and \otimes denotes the Kronecker product. Second, we have to use a tensor decomposition algorithm to get (M, ω) from the equations (2). The usage of the third order moments is necessary to guarantee the recoverability of (M, ω) with minimal assumptions on M (like M to be full rank). Methods dealing only with up to the second order moments exist, but present stronger requirements on the model parameters (Arora et al., 2013).

3.1 Overview of the proposed method

The classic approach to learn the parameters of a LVM consists in designing asymptotically (with the sample size) convergent estimates of M_1, M_2 and M_3 , and then to feed a tensor decomposition algorithm \mathcal{A} , to get asymptotically convergent estimates of (M, ω) . Unfortunately, the only

method we are aware of to estimate M_2 and M_3 for a NBM with binary outcomes is that of Jain and Oh (2014), that requires to deal with full tensors M_3 , compromising the scalability of the method. In this paper instead we will follow a different approach: we will find two estimators whose expectations \tilde{M}_2 and \tilde{M}_3 are close, but not equal, to M_2 and M_3 (Subsection 3.2), and obtain an approximate estimate $(\tilde{M}, \tilde{\omega})$ of (M, ω) using a tensor decomposition method (Subsection 3.3). The usage of the proposed approximated estimators \tilde{M}_2 and \tilde{M}_3 is crucial in order to allow the optimized implementation proposed in Appendix D, that allows to recover the parameters $(\tilde{M}, \tilde{\omega})$ without ever calculating explicitly \tilde{M}_3 , which is not possible with the method from Jain and Oh (2014). As the estimators \tilde{M}_2 and \tilde{M}_3 are biased, also the retrieved parameters $(\tilde{M}, \tilde{\omega})$ will suffer of a systematic bias; to get rid of it, we will refine the obtained parameters with EM, (Subsection 3.4), increasing the likelihood of the model. The intuition behind this approach is that EM is able to refine the parameters that we get, removing the systematic bias, if this bias is small. This intuition is confirmed in the experiments of Appendix E, where the proposed method outperforms many standard clustering methods when clustering binary data.

3.2 Recovering the moments

We now focus on how to estimate M_1 , M_2 and M_3 from a sample of observations distributed as a NBM. The estimation of M_1 is a trivial consequence of the conditional expectation: given a sample $(\bar{X})_i = X^{(i)} = (x_1^{(i)}, \ldots, x_d^{(i)})$, for $i \in \{1, \ldots, N\}$ we have, for each j, $(M_1)_j = \lim_{N \to \infty} \sum_{i=1}^N x_j^{(i)}/N$. We now focus on M_2 and M_3 . Define the following quantities:

$$\tilde{M}_2^{(N)} = \frac{\bar{X}^\top \bar{X}}{N} \in \mathbb{R}^{d \times d}, \quad \tilde{M}_3^{(N)} = \sum_{i=1}^N \frac{X^{(i)} \otimes X^{(i)} \otimes X^{(i)}}{N} \in \mathbb{R}^{d \times d \times d}$$
(3)

and their respective limits, $\tilde{M}_2 = \lim_{N \to \infty} \tilde{M}_2^{(N)}$ and $\tilde{M}_3 = \lim_{N \to \infty} \tilde{M}_3^{(N)}$. It is possible to demonstrate (see Appendix B for details) that \tilde{M}_2 and \tilde{M}_3 are close to M_2 and M_3 ; in particular, the estimates \tilde{M}_2 and \tilde{M}_3 always converge to the right value in the off-diagonal terms, while the diagonal entries instead converge to a value whose distance from the true value is bounded and can always be estimated from the sample. Even if biased, we will use \tilde{M}_2 and \tilde{M}_3 as input to a tensor decomposition algorithm, to estimate the parameters $(\tilde{M}, \tilde{\omega})$, which will consequently be biased. To reduce this bias, once learned the parameters of a NBM, we will update them with some iterations of EM, in order to increase the likelihood of the learned model.

3.3 Tensor Decomposition with SVTD

Given the true values of (2), we can use a decomposition algorithm to learn the NBM parameters. The literature provides a variety of algorithms to retrieve the desired pair (M, ω) from the estimated tensors, which differ one from another on how they exploit the spectral properties of the moment operators (Anandkumar et al., 2012a,b, 2014; Kuleshov et al., 2015). Even if the choice of the decomposition algorithm to be used is arbitrary, in this paper, we will base on SVTD, from Ruffini et al. (2016), for two reasons: first it is deterministic, not relying on any randomized matrix for its implementation; second, because it allows, with few simple modifications, to work without dealing explicitly with the tensor M_3 , that is often too large (cubic in the number of variables); these modifications, which reduce the complexity of the method, are presented in Appendix D where we embed the estimation procedure of the previous Subsection 3.2 with SVTD. A summary of the basis of

SVTD, as presented by Ruffini et al. (2016) can be found in Appendix C. When fed with the correct model parameters M_1 , M_2 , M_3 , SVTD produces the correct estimates (M, ω) . Conversely, when the inputs are biased, also the estimated $(\tilde{M}, \tilde{\omega})$ will suffer of a bias that depends on the bias of the inputs (Ruffini et al., 2016). To remove this bias, we will run few iterations of EM.

3.4 Expectation Maximization

In previous sections we have seen how to compute from data an estimate $(M, \tilde{\omega})$ of the parameters of a NBM. We can improve that estimate using Expectation Maximization (EM) (Dempster et al., 1977). EM, starting from an initial estimate $(\tilde{M}, \tilde{\omega})$, iteratively updates those parameters, increasing at each step the likelihood of the model, until convergence is reached. The issue of EM is that it may converge to poor local optima, if initialized poorly. Even if the results provided by the method described here suffer for the bias introduced in the estimates of moments, for sufficiently large datasets, those estimates are close to the correct estimates and so also the outputs will be close to the "correct" results (see Ruffini et al., 2016); the intuition is that the biased pair $(\tilde{M}, \tilde{\omega})$ of the previous paragraph, are good initializers for EM, and a few iterative steps will suffice to correct the initial bias, reaching a good optimum (See Marin et al., 2005, for details on EM).

4. Putting it all together

We are now ready to merge all the elements presented till now, and to show how to use them to perform Naïve Bayes Clustering. Starting from a dataset $\bar{X} \in \mathbb{R}^{N \times d}$, one can follow the following flow: *First*, estimate \tilde{M}_1 , \tilde{M}_2 , \tilde{M}_3 , as in Subsection 3.2 and retrieve the model parameters $(\tilde{M}, \tilde{\omega})$ with SVTD. We call *Approximate-SVTD*, or *ASVTD*, this method built on approximated tensor decomposition, and we refer to Algorithm 2 in Appendix D for an optimized implementation that, embedding the moments computation in the decomposition algorithm, allows to run the tensor decomposition without explicitly computing \tilde{M}_3 allowing the method to scale as d^2 instead of d^3 . *Then*, apply EM to improve the quality of the estimated parameters. *Last*, use Naïve Bayes Clustering to cluster the rows of \bar{X} . These steps will be applied in the next section on two medical datasets.

Remark 1 (Tuning the clustering results) The EM runs performed after the parameter retrieval, guarantee that the retrieved mixture model is (locally) optimal in terms of likelihood. Additional fine-tuning is possible re-running the algorithms in different configurations, for example modifying the number of clusters, adding/excluding certain features, or performing additional clustering within one of the clusters. The interpretation technique provided in the next section allows a user to visually inspect results and add expert considerations on the discovered patterns. Other type of manual fine-tuning (like manually assign a patient to a different cluster) may be possible, at the cost of adding human bias to the results of the model.

5. Patient Clustering

In this section we present the results of applying ASVTD to cluster patients form a real-world EHR¹.

^{1.} Due to privacy constraints, these datasets are not publicly available.

5.1 The datasets

We consider two datasets provided by Servei Català de la Salut, the major provider of public healthcare in Catalonia, Spain. The datasets are subsets of a bigger database containing all hospitalizations in Catalonia for the year 2016. The first dataset contains all the patients affected by Heart Failure, to be precise patients having a primary or secondary diagnostic 428 in the ICD-9 code. The second dataset contains "tertiary" patients, those with a serious disease that is supposed to be treated in one of the 5-6 large, reference hospitals in the area, because either the expertise or the required resources are only found there; criteria for tertiarism includes all oncology surgery, all heart surgery, all neurosurgery, major traumatism, acute ictus, myocardial infarct and sepsis, transplants, and a few rarer conditions. The datasets have the same format: each row represents a visit of a patient to an hospital, and the columns contain the codes of the up to 10 diagnostics that the doctor annotated in the patient history at the time of admission; their order is considered irrelevant. Diagnostics are coded in the international ICD-9 code (Geraci et al., 1997). The codes are hierarchical, of the form ddd.dd, with the .dd part being optional: the first 3 digits are a general diagnostic and extra digits if present make the diagnostic more precise. We kept only the first three digits of the ICD-9 codes, which are those with more consensus among doctors, obtaining a total vocabulary of 696 codes. In this way it is straightforward to map a dataset into a $N \times 696$ matrix with binary entries, where N is the number of records of each dataset, using an approach analogous to the bag-of-words: first associate each registered disease to a unique number between 1 and 696, and then populate a matrix X placing a 1 at position (i, j) if the record i presents diseases j, otherwise a zero.

5.2 Analysis of the results

In both the cases we run ASVTD, followed by EM, to then perform Naïve Bayes Clustering; the stopping criteria for EM was the moment in which convergence was reached; in particular, we stopped the algorithm when the norm of the variation on the estimated ω between an iteration and the next was less than 0.01. In both cases, to perform tensor decomposition, we removed the records with less than 3 diseases, obtaining two reduced datasets: for the Tertiarism dataset, \bar{X}_t with N = 16311 samples and for the Heart Failure dataset \bar{X}_c with N = 23154 samples. In order to analyze the results we plot two charts for each dataset: a heat-map and a disease-frequency chart. The heat-map, is in figure 1b. In that figure, we took the 40 most common diseases and plot the matrix \bar{X}_c only for the columns associated to those diseases. We ordered the rows of \bar{X}_c according to the cluster to which they belong, and draw them sequentially: each row of Figure 1b is a record, the black points indicate whether a disease is present. The background color of each row indicates the cluster (so: white background is the first cluster, clear gray the second, and so on). The purpose of this figure is to give a first visual inspection of the patterns present inside the clusters; heat-maps also give an idea of the dimension of the clusters with respect to the total dataset. Disease-frequency charts are instead used to give a meaning to those patterns, an example is at Figure 1a: the top-20 diseases in terms of frequency are displayed, each one in a different row. Then, for each one of the clusters, we show the relative frequencies of the considered diseases; for example, in Figure 1a, "Dyslipidemia" is present in the 50% of the patients belonging to the first cluster. To have an additional indication of the content of the clusters, we represent in Tables 1 and 2 the most relevant diagnostics, where the *relevance* (Sievert and Shirley, 2014) of a diagnostic i with respect to a

cluster j, given a weight parameter λ , is defined as

$$r(i,j) = \lambda \log(\mu_{i,j}) + (1-\lambda) \log(\mu_{i,j}/(\sum_{h=1}^{k} \mu_{i,h}\omega_h))$$

where $\mu_{i,j} = \mathbb{P}(x_i = 1 | Y = j)$. The relevance of a diagnostic with respect to a cluster j is an indicator that has a high value if the frequency of the considered diagnostic inside the cluster is much higher then its frequency on the full dataset; diagnostics with a high relevance are those that characterize a given cluster with respect to the others. An analysis of the content of a cluster can be performed merging the information contained in the disease-frequency charts with those provided by the tables of the highly relevant diseases: disease-frequency charts will show us the diseases that are frequent in a given cluster, regardless what happens in the other clusters; conversely, tables of the highly relevant diseases will highlight those diagnostics that characterize a cluster with respect to the others, providing in this way a complete view on the patterns that can be found in the data.

5.2.1 HEART FAILURE DATASET

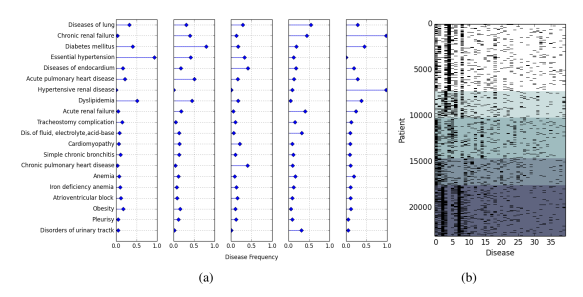


Figure 1: 1a Disease-frequency chart for the Heart Failure dataset. 1b The corresponding heatmap. The "Heart Failure" disease is not represented as it appears in all the records of the dataset.

The clusters obtained in the heart failure dataset are plotted in Figure 1a, with the associated table (1) of the highly relevant diseases. Clusters can be described with high coherence: Cluster 1 could be called the "purely metabolic": high prevalence of hypertension, and also dyslipidemia and diabetes mellitus. Cluster 2 contains the above complicated with kidney problems; Patients in Cluster 3 present valvular problems and pulmonary hypertension, well-known to be related. Cluster 4 is a mixture of kidney and pulmonary problems (lung diseases are pretty frequent here), with little metabolic implications. Cluster 5 contains the purely kidney sufferers, with diabetes mellitus (nephropaty being a common complication of diabetes) and dyslipidemia. Clinicians confirm that

Cluster	Most Relevant Diagnostics	Cluster Size
1	Essential hypertension; Dyslipidemia; Cardiac arrhythmias; Diabetes mellitus; Obesity;	7290
2	Diabetes mellitus; Atherosclerosis; Acute pulmonary heart disease; Retinal disorders; Hy-	2915
	pertensive heart and renal disease;	
3	Chronic pulmonary heart disease; Diseases of endocardial structures; Diseases of endo-	4480
	cardium; Hypertensive heart disease; Cardiac arrhythmias;	
4	Bacterial infection; Disorders of urethra - urinary tractk; Acute renal failure; Hypertensive	2936
	heart - renal disease; Disorders of fluid, electrolyte, acid-base balance; Diseases of lung;	
5	Hypertensive renal disease; Chronic and/or acute renal failure; Diabetes mellitus; Anemia;	5533

Table 1: The most relevant diseases for each cluster for the Heart failure dataset.

these clusters make sense once seen, and may be useful for guiding treatment. For example, medication that might be indicated for the purely cardiac clusters might be not advisable for the clusters with renal problems if it is suspected to be nephrotoxic.

5.2.2 TERTIARISM DATASET

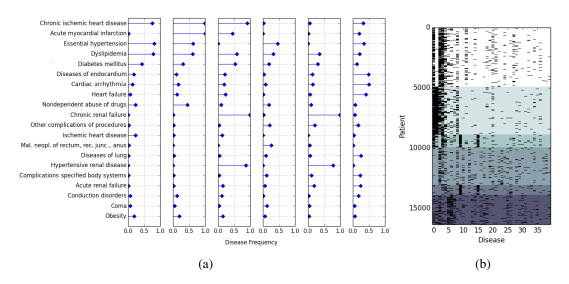


Figure 2: 2a Disease-frequency chart for the Tertiarism dataset. 2b The corresponding heatmap.

The clusters for the tertiarism dataset are plotted in Figure 2a, with the associated Table 2 of the high relevant diseases. Clinically, Cluster 1 might correspond to patients in need of coronary intervention (heart surgery or interventional cardiology), again with strong presence of metabolic anomalies. Cluster 2 is similar, but including myocardial infarction. Cluster 3 includes nephrology patients, who are also taken to tertiary hospitals when they need transplant or are in very advanced phase, and who tend to develop arteriopathy in the long term; interestingly, diabetes mellitus shows up strongly in this cluster, as it is the leading cause of entrance to dialysis programs. Cluster 4 has a surprising behavior: on a first hand, no clear signal comes out from the disease-frequency chart, besides the ever-present hypertension, dyslipidemia and a small signal of colo-rectum cancer. However, if we look at the table of high relevant diseases, we can see that the most relevant diagnostics for this cluster are neoplasms. Oncological problems are large constituents of tertiarism; they are

Cluster	Most Relevant Diagnostics	Cluster Size
1	Ischemic heart disease; Essential hypertension; Angina pectoris; Dyslipidemia; Chronic	4892
	ischemic heart disease;	
2	Acute myocardial infarction; Chronic ischemic heart disease; Nondependent abuse of	3982
	drugs; Essential hypertension; Dyslipidemia;	
3	Hypertensive renal disease; Chronic renal failure; Chronic ischemic heart disease; Dia-	1043
	betes mellitus; Acute myocardial infarction;	
4	Malignant neopl. of rectum, rectosigmoid junction, and anus; Secondary malignant neopl.	3133
	of respiratory and digestive systems; Malignant neopl. of trachea, bronchus, and lung;	
	Malignant neopl. of stomach; Diseases of white blood cells;	
5	Chronic renal failure; Hypertensive renal disease; Disorders involving the immune mech-	819
	anism; Anemia; Disorders of urethra and urinary tractk;	
6	Diseases of endocardial structures; Chronic pulmonary heart disease; Heart failure; Dis-	2442
	eases of endocardium; Diseases of lung;	

Table 2: The most representative diseases for each cluster for the Tertiarism dataset.

characterized by a large set of codes (almost 100 for neoplasms, by organ of origin mostly), so it is not surprising the fact that we do not find such a diagnostic in the disease-frequency plot, as there are many distinct diagnostics indicating oncological diseases; however, the fact that the most relevant diagnostics for this cluster are neoplasms allows us to consider this family of diagnostics as characterizing this cluster. Cluster 5 groups kidney patients without arteriopathy, so similar to the first and second clusters but without previous infarction. Cluster 6 includes patients with cardiac valvular disease; normally they need heart surgery or interventional cardiology, but their associated complications are quite different from those in Clusters 1 and 2.

Stability analysis. A clustering method is expected to be stable: if trained on two datasets of reasonable size generated by the same process, it should produce similar clusters. To study the stability of ASVTD we split each of the studied datasets into two non-disjoint subsets on which we run the clustering method; then we compare the clusterings obtained on the intersection between the two subsets with the Adjusted Rand Index (Hubert and Arabie, 1985). In both the cases, we got satisfying results, with an index higher than 0.9, indicating similarity between the two partitions and thus stability.

6. Conclusions

We have presented an efficient method for clustering high dimensional binary data, able to find interesting and potentially useful patterns even in the simplest format of EHR, namely, diagnostic codes only. A strong point noted by clinicians is that it provides a fresh look, without prejudice, based on an aseptic algorithm. A natural follow-up consists in including additional clinical information to the features, like demographics, lab results, vital signs or medications. Also, we believe that including higher-level diagnostic groups among the features may improve the clarity of some patterns; these are more clearly defined in the more recent coding scheme ICD10 than in ICD9.

Acknowledgments

Work by M. Ruffini and R. Gavaldà is partially funded by AGAUR project 2014 SGR-890 (MACDA) and by MINECO project TIN2014-57226-P (APCOM).

References

- Charu C Aggarwal, Alexander Hinneburg, and Daniel A Keim. On the surprising behavior of distance metrics in high dimensional space. In *International Conference on Database Theory*, pages 420–434. Springer, 2001.
- Anima Anandkumar, Yi-kai Liu, Daniel J Hsu, Dean P Foster, and Sham M Kakade. A spectral algorithm for Latent Dirichlet Allocation. In *NIPS*, pages 917–925, 2012a.
- Animashree Anandkumar, Daniel Hsu, and Sham M Kakade. A method of moments for mixture models and Hidden Markov models. In COLT, volume 1, page 4, 2012b.
- Animashree Anandkumar, Rong Ge, Daniel Hsu, Sham M Kakade, and Matus Telgarsky. Tensor decompositions for learning latent variable models. *Journal of Machine Learning Research*, 15 (1):2773–2832, 2014.
- Sanjeev Arora, Rong Ge, Yonatan Halpern, David Mimno, Ankur Moitra, David Sontag, Yichen Wu, and Michael Zhu. A practical algorithm for topic modeling with provable guarantees. In *International Conference on Machine Learning*, pages 280–288, 2013.
- Mitchell J Cohen, Adam D Grossman, Diane Morabito, M Margaret Knudson, Atul J Butte, and Geoffrey T Manley. Identification of complex metabolic states in critically injured patients using bioinformatic cluster analysis. *Critical Care*, 14(1):R10, 2010.
- Arthur P Dempster, Nan M Laird, and Donald B Rubin. Maximum likelihood from incomplete data via the em algorithm. *Journal of the Royal Statistical Society. Series B (methodological)*, pages 1–38, 1977.
- Jane M Geraci, Carol M Ashton, David H Kuykendall, Michael L Johnson, and Louis Wu. International classification of diseases, 9th revision, clinical modification codes in discharge abstracts are poor measures of complication occurrence in medical inpatients. *Medical care*, 35(6):589–602, 1997.
- Nathan Halko, Per-Gunnar Martinsson, and Joel A Tropp. Finding structure with randomness: Probabilistic algorithms for constructing approximate matrix decompositions. *SIAM review*, 53 (2):217–288, 2011.
- Joyce C Ho, Joydeep Ghosh, Steve R Steinhubl, Walter F Stewart, Joshua C Denny, Bradley A Malin, and Jimeng Sun. Limestone: High-throughput candidate phenotype generation via tensor factorization. *Journal of biomedical informatics*, 52:199–211, 2014a.
- Joyce C Ho, Joydeep Ghosh, and Jimeng Sun. Marble: high-throughput phenotyping from electronic health records via sparse nonnegative tensor factorization. In *Proceedings of the 20th ACM SIGKDD international conference on Knowledge discovery and data mining*, pages 115–124. ACM, 2014b.
- Lawrence Hubert and Phipps Arabie. Comparing partitions. *Journal of classification*, 2(1):193–218, 1985.

- Prateek Jain and Sewoong Oh. Learning mixtures of discrete product distributions using spectral decompositions. In *COLT*, pages 824–856, 2014.
- Hans-Peter Kriegel, Peer Kröger, and Arthur Zimek. Clustering high-dimensional data: A survey on subspace clustering, pattern-based clustering, and correlation clustering. *ACM Transactions on Knowledge Discovery from Data (TKDD)*, 3(1):1, 2009.
- Kanak Bikram Kshetri. *Modelling patient states in intensive care patients*. PhD thesis, Massachusetts Institute of Technology, 2011.
- Volodymyr Kuleshov, Arun Chaganty, and Percy Liang. Tensor factorization via matrix factorization. In *AISTATS*, pages 507–516, 2015.
- Jean-Michel Marin, Kerrie Mengersen, and Christian P Robert. Bayesian modelling and inference on mixtures of distributions. *Handbook of statistics*, 25:459–507, 2005.
- Volodymyr Melnykov, Ranjan Maitra, et al. Finite mixture models and model-based clustering. *Statistics Surveys*, 4:80–116, 2010.
- Fabian Pedregosa, Gaël Varoquaux, Alexandre Gramfort, Vincent Michel, Bertrand Thirion, Olivier Grisel, Mathieu Blondel, Peter Prettenhofer, Ron Weiss, Vincent Dubourg, et al. Scikit-learn: Machine learning in python. *Journal of Machine Learning Research*, 12(Oct):2825–2830, 2011.
- Laura M Pérez, Marco Inzitari, Terence J Quinn, Joan Montaner, Ricard Gavaldà, Esther Duarte, Laura Coll-Planas, Mercè Cerdà, Sebastià Santaeugenia, Conxita Closa, et al. Rehabilitation profiles of older adult stroke survivors admitted to intermediate care units: A multi-centre study. *PloS one*, 11(11):e0166304, 2016.
- Ioakeim Perros, Evangelos E Papalexakis, Fei Wang, Richard Vuduc, Elizabeth Searles, Michael Thompson, and Jimeng Sun. Spartan: Scalable parafac2 for large & sparse data. *arXiv preprint arXiv:1703.04219*, 2017.
- Dieter Rixen, John H Siegel, and Herman P Friedman. " sepsis/sirs," physiologic classification, severity stratification, relation to cytokine elaboration and outcome prediction in posttrauma critical illness. *Journal of Trauma and Acute Care Surgery*, 41(4):581–598, 1996.
- Matteo Ruffini, Marta Casanellas, and Ricard Gavaldà. A new spectral method for latent variable models. *arXiv preprint arXiv:1612.03409*, 2016.
- Carson Sievert and Kenneth E Shirley. Ldavis: A method for visualizing and interpreting topics. In *ACL workshop on interactive language learning, visualization, and interfaces*, 2014.
- Zhuoxin Sun, Ori Rosen, and Allan R Sampson. Multivariate bernoulli mixture models with application to postmortem tissue studies in schizophrenia. *Biometrics*, 63(3):901–909, 2007.
- Stefan Van Der Walt, S Chris Colbert, and Gael Varoquaux. The numpy array: a structure for efficient numerical computation. *Computing in Science & Engineering*, 13(2):22–30, 2011.
- Ulrike Von Luxburg. A tutorial on spectral clustering. *Statistics and computing*, 17(4):395–416, 2007.

Yichen Wang, Robert Chen, Joydeep Ghosh, Joshua C Denny, Abel Kho, You Chen, Bradley A Malin, and Jimeng Sun. Rubik: Knowledge guided tensor factorization and completion for health data analytics. In *Proceedings of the 21th ACM SIGKDD International Conference on Knowledge Discovery and Data Mining*, pages 1265–1274. ACM, 2015.

Appendix

Appendix A. Related works on spectral methods

A few examples of application of mixture models to the medical domain can be found in the literature (see Sun et al., 2007), but we are not aware of any of them using methods of moments and tensor decomposition. We are interested in performing clustering using a special case of mixture model, called Naïve Bayes Model (NBM), where all the observable features are conditionally independent binary variables. The key obstacle to learn such models with a spectral method consists in recovering from a sample a good approximation of the low rank tensors to be decomposed. In (Jain and Oh, 2014) the authors solve this problem for mixtures with categorical outcomes employing an optimization algorithm to reconstruct such tensors; then, they suggest to feed Tensor Power Method from Anandkumar et al. (2014) to recover the model parameters. Their approach works when the number of the features is significantly higher than the number of the clusters; the problem of that method is the fact that it requires the storage of the full three dimensional tensor in memory, reducing the scalability of the algorithm (depending on a factor d^3 for both memory and time requirements, where d is the number of features). An alternative approach for some mixture models, including NBMs was proposed in (Anandkumar et al., 2012b); this approach, which requires the number of features to be at least three times the number of clusters $(d \ge 3k)$, consists in treating a NBM as a three-views mixture model, by splitting the vector of the features into three subvectors, and independently learn the parameters of each one of the three views. The method proposed in that paper relies on the usage of randomized matrices, that highly affect the quality of the retrieved parameters; some of the same authors, presented in (Anandkumar et al., 2014) a different way to deal with the multi-view approach (and so with NBMs), together with the Tensor Power Method. This approach is robust, and seems to work fine in many different settings. We remark that it is possible to implement these techniques in an optimized way, reducing the complexity of the method to d^2 , allowing the management of large scale datasets.

A.1 Why a different technique?

The main drawback of the multi-view approach is the fact that it requires the three views to be fully informative about the latent variable: geometrically, this means that they are required to be full-rank; practically, it means that we can infer the status of a latent variable just by looking at one single view. With healthcare electronic records, this may not be the case. Consider a dataset of N patients, with d binary records. We can think at this dataset as a matrix \bar{X} with N rows and d columns, where at position (i, j) we have a 1 if the patient i has the disease j. In general, the frequency of the various diseases along the population of patients has a huge tail, with few most common diseases that are extremely spread along the population, and a very high portion of the remaining diseases that are pretty rare; however, even if a disease is rare for the whole population it might be characteristic of a small cluster of patients. In such a context, to implement a multi-view approach, we need to split the columns of the matrix X into three groups, obtaining three datasets; each one would represent a view on the latent status of the patient, which could be inferred by looking at an individual view. In our preliminary experiments, we tested the multi-view method on our databases of medical records, and we obtained results that were highly dependent on the way in which the split of the columns was made. Finding the "right" view, resulted to be a tricky task, highly depending on the analyzed dataset. A second issue with the multi-view approach, regards the ordering issues that emerge from

the asynchronous learning of the three views. Each view is a matrix whose column j represents the conditional expectation of the features of the view under cluster j. In general, when the views are returned by a tensor decomposition algorithm, the ordering of the columns is not consistent between the various views, and to obtain a proper reordering matching heuristics are required (see Section B.4 of Anandkumar et al., 2012b). With a broader view, for the specific application that we have in mind, one requires algorithms that: 1) need little hand-crafting (such as parameter tuning or data preprocessing like splitting the features into views, requiring long hours of collaboration between medical expert and data scientist), 2) are deterministic (doctors become highly suspicious of an algorithm that returns different results on different runs and same data), 3) have interpretable output and 4) are scalable, in the sense that can be run in seconds on large scale datasets. Besides the technical aspects on tensor decomposition method sketched above, our contribution is toward a clustering algorithm that is truly usable in this sense.

Appendix B. On the estimated tensors or moments

In the following theorem we show that the estimates proposed in Section 3.2 are close to the theoretical, unbiased ones.

Theorem B.1 Define the values of the conditional second and third order moments for an observable variable:

$$\mu_{i,j}^{(2)} = \mathbb{E}(x_i^2 | Y = j), \quad \mu_{i,j}^{(3)} = \mathbb{E}(x_i^3 | Y = j).$$

Then $(M_2)_{i,j} = (\tilde{M}_2)_{i,j}$ and $(M_3)_{i,j,l} = (\tilde{M}_3)_{i,j,l}$ for all $i \neq j \neq l$. Also, if $\mu_{i,j} \in [0,1] \ \forall i, j$, we have for all $i \neq j \neq l$:

$$|(M_2)_{i,i} - (\tilde{M}_2)_{i,i}| \le \sum_{j=1}^k \omega_j \mu_{i,j}^{(2)} - (M_1)_i^2,$$

$$|(M_3)_{i,i,l} - (\tilde{M}_3)_{i,i,l}| \le \sum_{j=1}^k \omega_j \mu_{i,j}^{(2)} \mu_{l,j} - \frac{(M_2)_{i,l}^2}{(M_1)_l}, \quad |(M_3)_{i,i,i} - (\tilde{M}_3)_{i,i,i}| \le \sum_{j=1}^k \omega_j \mu_{i,j}^{(3)} - (M_1)_i^3.$$

Notice that, because M_3 , \tilde{M}_3 , M_2 and \tilde{M}_2 are symmetric, the previous theorem states an error bound on all the entries of that operators.

Proof Recall the following notation

$$\mu_{i,j}^{(2)} = \mathbb{E}(x_i^2 | Y = j), \quad \mu_{i,j}^{(3)} = \mathbb{E}(x_i^3 | Y = j)$$

and observe that, because each $\mu_{i,j} \in [0,1]$,

$$\mu_{i,j}^{(2)} \ge \mu_{i,j}^2, \quad \mu_{i,j}^{(3)} \ge \mu_{i,j}^3.$$

We prove each claimed equation one by one.

1. $(M_2)_{i,j} = (\tilde{M}_2)_{i,j}$, and $(M_3)_{i,j,l} = (\tilde{M}_3)_{i,j,l}$ for all $i \neq j \neq l$. These equations are an immediate consequence of the conditional independence of the features.

2. $|(M_2)_{i,i} - (\tilde{M}_2)_{i,i}| \le \sum_{j=1}^k \omega_j \mu_{i,j}^{(2)} - (M_1)_i^2$ We consider

$$(\tilde{M}_2)_{i,i} = E(x_i^2) = \sum_{j=i}^k \omega_j \mu_{i,j}^{(2)} \ge \sum_{j=i}^k \omega_j \mu_{i,j}^2 =$$
$$= (M_2)_{i,i} \ge (\sum_{j=i}^k \omega_j \mu_{i,j})^2 = (M_1)_i^2.$$

From which we get the desired inequality.

3. $|(M_3)_{i,i,l} - (\tilde{M}_3)_{i,i,l}| \le \sum_{j=1}^k \omega_j \mu_{i,j}^{(2)} \mu_{l,j} - \frac{(M_2)_{i,l}^2}{(M_1)_l}$. We have $(\tilde{M}_3)_{i,i,l} = \sum_{j=1}^k \omega_j \mu_{i,j}^{(2)} \mu_{l,j} \ge \sum_{j=1}^k \omega_j \mu_{i,j}^2 \mu_{l,j} =$ $= (M_3)_{i,i,l} \ge \frac{(M_2)_{i,l}^2}{(M_1)_l},$

from which again we get the desired inequality.

4. $|(M_3)_{i,i,i} - (\tilde{M}_3)_{i,i,i}| \le \sum_{j=1}^k \omega_i \mu_{i,j}^{(3)} - (M_1)_i^3$. The proof of this point is identical to that of point 2.

Appendix C. Tensor Decomposition with SVTD: details

In this section we describe in depth the tensor decomposition method we used in the paper, SVTD. SVTD requires the matrix M to have rank k, with at least one row where all the entries are distinct. It is based on two key observations: 1) the matrix M_2 has rank $k \leq d$, k is the number of states that the latent variable can assume, and M_2 admits the following representation $M\Omega M^{\top}$, where $\Omega = diag(\omega)$; 2) for each $r \in \{1, ..., d\}$, the r - th slice of the tensor M_3 can be expressed as a rank-k matrix with the following structure:

$$M_{3,r} = M\Omega^{\frac{1}{2}} diag((\mu_{r,1},...,\mu_{r,k}))\Omega^{\frac{1}{2}}M^{\top}$$

The algorithm first performs a singular value decomposition (SVD) on M_2 , $M_2 = U_k S_k U_k^{\top}$, where $U_k \in \mathbb{R}^{d \times k}$ and $S_k \in \mathbb{R}^{k \times k}$ are the matrices of the singular vectors and values truncated at the k - th greatest singular value. Then defines $E_k = U_k S_k^{\frac{1}{2}} \in \mathbb{R}^{d \times k}$, and for each slice $M_{3,r}$ of M_3 , defines $H_r \in \mathbb{R}^{k \times k}$ as

$$H_r = E_k^{\dagger} M_{3,r} (E_k^{\top})^{\dagger},$$

where $E_k^{\dagger} = (S_k)^{-\frac{1}{2}} U_k^{\top}$ is the Moore-Penrose pseudo-inverse of E_k . Observing that there exists an orthonormal $k \times k$ matrix O, such that $M\Omega^{\frac{1}{2}} = E_k O$, one gets the following characterization of $M_{3,r}$:

$$M_{3,r} = M\Omega^{\frac{1}{2}} diag((\mu_{r,1},...,\mu_{r,k}))\Omega^{\frac{1}{2}}M^{\top}$$

$$= E_k Odiag((\mu_{r,1},...,\mu_{r,k}))O^{\top}E_k^{\top}$$

from which it follows that

$$H_r = Odiag((\mu_{r,1}, ..., \mu_{r,k}))O^+$$

Now, one gets the r-th row of M as the singular values of H_r . Repeating these steps for all the $r \in \{1, ..., d\}$ will provide the full matrix M. In order to avoid ordering issues with the columns of the retrieved M, one can use the same matrix O to diagonalize all the matrices H_r , as O does not depend on the considered row r. So, compute O as the singular vectors of H_r , for a certain feature r, and re-use it for all the other features. The main steps of SVTD are summed up in Algorithm 1.

Algorithm 1 SVTD

Require: M_1, M_2, M_3 , and the number of latent states k

- 1: Decompose M_2 as $M_2 = U_k S_k U_k^{\top}$ with a SVD.
- 2: Select a feature r and compute $M_{3,r}$
- 3: Compute O as the singular vectors of

$$H_r := (S_k)^{-\frac{1}{2}} U_k^{\top} M_{3,r} U_k (S_k)^{-\frac{1}{2}}$$

- 4: for $i = 1 \rightarrow d$ do
- 5: Compute $H_i := (S_k)^{-\frac{1}{2}} U_k^\top M_{3,i} U_k (S_k)^{-\frac{1}{2}}$
- 6: Obtain the *i*-th row of M as the diagonal entries of $O^{\top}H_iO$
- 7: **end for**
- 8: Obtain ω solving $M_1 = M\omega$
- 9: Return (M, ω)

With this algorithm, given the values of M_1 , M_2 , M_3 , one can retrieve the pair of unknown set of parameters (M, ω) . Note that step 2 of the algorithm requires to select a feature r to be used to compute the matrix O. This matrix will be then used to diagonalize all the matrices H_i in step 6. Even if theoretically one can decide to use any feature to compute O, it is shown by Ruffini et al. (2016) that the perturbations depend on the minimum distance between the elements of the r-th row of M, that are also the singular values of H_r . So, one repeats the step 3 of the algorithm for different features and selects the r that maximizes the minimum distance between the singular values of H_r . An explicit implementation of a method to select the feature r from which to compute the matrix O is presented in Appendix D.

Appendix D. Algorithmic details: implementing ASVTD

The usage of the estimates proposed in Section 3.2 allows to modify the structure of SVTD in order to reduce the complexity and deal with perturbed matrices. In this section, we propose a modification of the method proposed by Ruffini et al. (2016) to reach these goals. The algorithm we are going to present will produce the same results of SVTD when fed with the approximated tensors of Section 3.2, but in a faster way. First, we want to exploit the nice form of the operators introduced at equations (3) to improve the scalability of the method. Pure SVTD in fact works with all the explicit slices of the $d \times d \times d$ tensor M_3 ; we want to avoid this dependence, working with matrices with a lower dimension. Second, we need to indicate a way to select the feature r from which to compute matrix O of singular vectors used to perform the simultaneous diagonalization; in fact, in the unperturbed version of the theorem we can just randomly select a feature among the d available. Here, with perturbed matrices, we want to do this choice in order to minimize the perturbations on the output. These modifications are presented in Algorithm 2, named Approximate-SVTD (ASVTD).

Algorithm 2 Approximate - SVTD

Require: The sample $\bar{X} \in \mathbb{R}^{N \times d}$, the number of latent states k 1: Compute $\tilde{M}_2^{(N)} = \frac{\bar{X}^\top \bar{X}}{N} \in \mathbb{R}^{d \times d}$ 2: Decompose $\tilde{M}_2^{(N)}$ as $\tilde{M}_2^{(N)} = U_k S_k U_k^\top$ with a SVD. 3: Project \bar{X} on $\mathbb{R}^{N \times k}$: $\bar{X}_k = \bar{X} U_k(S_k)^{-\frac{1}{2}}$ 4: Initialize $\alpha_{min} = -\infty$ and $O = \mathbb{I}_k$ as the $k \times k$ identity matrix. 5: for $i = 1 \rightarrow d$ do Compute $H_i := \frac{\bar{X}_k^\top diag((x_i^{(1)}, \dots, x_i^{(N)}))\bar{X}_k}{N}$ 6: Compute the singular values of H_i , $(s_1, ..., s_k)$ and the left singular vectors O_i . 7: if $\min_{i\neq j}(|s_i - s_j|) > \alpha_{min}$ then 8: Set: $\alpha_{min} = \min_{i \neq j} (|s_i - s_j|)$, and $O = O_i$ 9: end if 10: 11: end for 12: for $i = 1 \rightarrow d$ do Obtain the *i*-th row of \tilde{M} as the diagonal entries of $O^{\top}H_iO$ 13: end for 14: 15: Obtain $\tilde{\omega}$ solving $(\frac{\sum_{i} x_{j}^{(i)}}{N})_{j=1..d} = (\tilde{M}\tilde{\omega})_{j=1..d}$ 16: Return $(\tilde{M}, \tilde{\omega})$

Comparing SVTD (Algorithm 1) and its approximate version (Algorithm 2), we notice that the main differences are in the loop from row 5 to 11 of Algorithm 2. In fact, standard SVTD computes the matrices H_i , at its line 5, using directly the i - th slice of M_3 :

$$H_i = (S_k)^{-\frac{1}{2}} U_k^{\top} M_{3,i} U_k (S_k)^{-\frac{1}{2}}$$

this operation is pretty expensive, as it requires $O(d^2k)$ operations, and is repeated d times, for a total cost of $O(d^3k)$. Instead, in ASVTD, we never compute explicitly $\tilde{M}_3^{(N)}$, but we compute directly H_i , exploiting the fact that, in this case

$$H_i = \frac{((S_k)^{-\frac{1}{2}} U_k^\top \bar{X}^\top) diag((x_i^{(1)}, ..., x_i^{(N)}))(\bar{X} U_k(S_k)^{-\frac{1}{2}})}{N}.$$

So, in row 3, we compute the matrix $\bar{X}_k = \bar{X}U_k(S_k)^{-\frac{1}{2}}$ and we use it during all the algorithm; the calculation of each H_i now costs only $O(k^2N)$, which in many cases is a substantial improvement (especially when the number of the features is high). The second main difference between SVTD and ASVTD is the fact that we use the loop from row 5 to 11 of Algorithm 2 to compute the value of the orthornomal matrix O to be used to diagonalize the various H_i , selecting the feature that maximizes the difference between the two nearest singular values. It is shown in (Ruffini et al., 2016, Thm. 5.1) that in this way, we are going to maximize the stability of the resulting matrices $(\tilde{M}, \tilde{\omega})$. We now analyze the complexity requirements of ASVTD. It is easy to see that the steps

from 1 to 3 have a time complexity of $O(d^2N)$ for step 1, $O(d^2k)$ for step 2, using randomized techniques (Halko et al., 2011) and O(Ndk) for step 3. In the loop from step 5 to 11 we have the calculation of H_i , costing $O(k^2N)$, and a $k \times k$ SVD, costing $O(k^3)$, giving a total time complexity of $O(d^2N + dk^2N + dk^3)$ in the realistic cases where d, k < N. Also the memory requirements are mild, as the only storage is for the matrices H_i , costing $O(dk^2)$, $\tilde{M}_2^{(N)}$, costing $O(d^2)$ and \bar{X} , for O(dN). The total memory is thus $O(dk^2 + d^2 + dN)$.

Appendix E. Experimental comparison of ASVTD with other methods

In this paper we have described how to recover the parameters (M, ω) of a NBM, presenting a simple heuristic. The advantage of our method is mainly practical, as it does not require any expert knowledge added in the preprocessing of the data. In this paragraph we want to compare experimentally the clustering accuracy of the proposed method with that of state-of-the-art methods. We proceed as follows: First we generate a synthetic dataset, distributed as a mixture of independent Bernoulli, registering for each sample, the cluster to which it belongs (i.e. the latent variable generating the observation); the synthetic parameters (M, ω) have been generated as exponentially distributed random numbers, normalized to be between 0 and 1. Then we cluster the samples in an unsupervised way, and we compare the obtained clusters with the theoretical ground-truth, registered when generating the data. We then repeat this procedure for various model and sample sizes. In order to analyze the clustering accuracy, we use the Adjusted Rand Index from Hubert and Arabie (1985) between the theoretical ground-truth partition and the one obtained with the clustering algorithm. Adjusted Rand Index is an indicator that compares whether two partitioning of a dataset are or not similar; an Adjusted Rand Index of 1 indicates that two clusterings are identical, while a random labeling should have a value close to 0. Note that many well-known measures of cluster fitness such as the silhouette coefficient are not applicable as they rely on notions of distance among instances. We run the clustering with many different algorithms:

- ASVTD, as described in this paper.
- Naïve Bayes clustering with the three-view approach from Anandkumar et al. (2012b) (labeled "HMM12" in the figures).
- Naïve Bayes clustering with the three-view approach from Anandkumar et al. (2014) (labeled "TPM" in the figures).
- K-Means.
- Spectral Clustering from Von Luxburg (2007), using a linear kernel ("SC-lin" in the figures).
- PCA clustering, where k-means is used to cluster the the sample where the dimensionality has been reduced using a PCA.

We have not run pure EM with random initialization, as its final results and running time strongly depend on the random initialization. For K-Means and the Spectral clustering methods we used the implementation present in Scikit-learn Python library (Pedregosa et al., 2011). In order to avoid ordering issues for the multi-view clustering methods, we just used the true parameters of the models to recover the correct ordering; the views have been built by dividing the set of the features into three subsets: the first view are the features from 1 to d/3, the second those from d/3 + 1 to d2/3 and

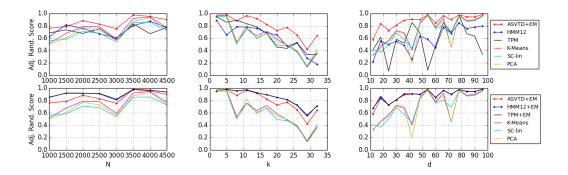


Figure 3: Adjusted Rand Index of the compared methods, when EM is not used after the multi-view methods (top three figures) and when is used (bottom three figures).

the last are those from d2/3 + 1 to d; this was an easy task, as the entries of M are well balanced. After ASVTD, we always run EM until convergence is reached, stopping the algorithm when the norm of the variation on the estimated ω between an iteration and the next was less than 0.01; we always use EM, because ASVTD is an approximate method, without theoretical guaranteers, and EM is an integral part of its usage. In a first experiment we do not run any EM iteration after the multi-view clustering methods, while in the second one, we do run EM after those methods, with the same stopping criteria used for ASVTD².

The results are displayed in figure 3. The top three figures represent the experiment where EM is not run after the multi-view clusterings, while the three figures below represent the experiment where EM is used. In the leftmost figures we fixed the number of features d = 99 and the number of latent states k = 12, varying the sample size from N = 100 to N = 10000. In the central figures, the sample size N = 10000 and the number of features d = 99 are fixed, while the number of latent states varies from k = 2 to k = 33. In the rightmost figure, the sample size N = 10000and the number of latent states k = 4 are fixed, while the number of features varies from d = 12to d = 99. It is interesting to notice that all the reference methods (K-means, Spectral Clustering and PCA) are outperformed from tensor methods; also, seems that the intuition of using a mixed tensor + EM method provides good results, as ASVTD, despite the lack of theoretical guarantees, performs better than the pure (i.e. without EM) multi-view approaches, both with Tensor Power method and with the method from (Anandkumar et al., 2012b). If we plug some EM iterations after the pure tensor methods (three figures below), we can see interesting results. EM provides a huge improvement to the quality of the results of TPM and of Anandkumar et al. (2012b) method. In particular, we can see that now ASVTD gets almost the same results as these other methods, without requiring any domain-dependent preprocessing of the data; in this case, this preprocessing activity was trivial, but, as explained in paragraph A.1, with certain unbalanced datasets, it may be very challenging to find a proper splitting of the features. In addition, if we look at the experiments of Section 5, we can see that they are in the case where N is large, d is large as well and k is

^{2.} All the algorithms have been implemented in Python 2.7, using *numpy* (Van Der Walt et al., 2011) library for linear algebra operations. All the experiments have run on a MacBook Pro, with an Intel Core i5 processor.

Method	Time (seconds)
ASVTD + EM	1.3
TPM + EM	3.68
HMM12 + EM	0.77
KMeans	0.76
SC - lin	7.48
PCA	0.38

Table 3: The average clustering time spent for each method.

small; these are exactly the configurations where ASVTD performs better, providing results at least as good as the competing methods; these motivations justify our choice of using ASVTD for the experimental results of Section 5.

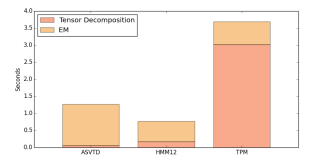


Figure 4: Average running times of the tensor-based methods. The red portion of the bars represents the average time spent in doing tensor decomposition; the orange portion, the time for EM.

We now briefly discuss the running times. For each run of the experiment described in the previous paragraph we registered the time employed by each of the clustering methods. We then average all these records, in order to get, for each method, the average time needed to perform the clustering. Results are displayed in Table 3. It is immediate to see that Spectral Clustering method is much slower than the competing techniques. This is a consequence of dealing with an $N \times N$ affinity matrix, that requires a number of operations quadratic in the sample size. PCA and Kmeans are pretty fast, but we have seen that they perform poorly on binary data. Looking at tensor methods, TPM seems to be the slowest method, consequence of a worst scalability on the number of latent components, while ASVTD and HMM12 have similar performances. Focusing on the tensor based methods, it is interesting to understand the amount of time spent in performing tensor decomposition and the amount spent in refining parameters with EM, to reach convergence. This analysis is displayed in Figure 4; the red portion of each bar represents the time for decomposition, the orange portion, the time for EM; the total bars are the average time spent by each algorithm to perform the full reconstruction (decomposition and EM). As expected, ASVTD is significantly faster than other decomposition methods, due to the better scalability. However, as it does not provide a guaranteed estimation of the model parameters, EM takes a bit more time to converge. If we look at the other algorithms we can see a significantly longer time in decomposing tensors, and a shorter time in doing EM.