Contamination-source based K-sample clustering

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Editor: Lorenzo Rosasco

Abstract

In this work, we investigate the K-sample clustering of populations subject to contamination phenomena. A contamination model is a two-component mixture model where one component is known (standard behaviour) and the second component, modeling a departure from the standard behaviour, is unknown. When K populations from such a model are observed we propose a semiparametric clustering methodology to detect which populations are impacted by the same type of contamination, with the aim of faciliting coordinated diagnosis and best practices sharing. We prove the consistency of our approach under the assumption of the existence of true clusters and demonstrate the performances of our methodology through an extensive Monte Carlo study. Finally, we apply our methodology, implemented in the \mathtt{admix}^1 R package, to a European countries COVID-19 excess of mortality dataset, aiming to cluster countries similarly impacted by the pandemic across different age groups.

Keywords: Admixture, Clustering, Hypothesis Testing, Semiparametric Mixture.

1. Introduction

Let us consider the two-component mixture model with the cumulative distribution function

$$L(x) = (1 - p)G(x) + pF(x), \tag{1}$$

for all $x \in \mathbb{R}$, where G is a known cumulative distribution function modeling a standard behavior, and the unknown parameters are the mixture proportion $p \in]0,1[$ along with the cumulative distribution function of the unknown component F, modeling a departure from the standard behaviour. This model, usually called contamination model, has been extensively studied over the last decades and is related to various applications, see Shen et al.

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^{1.} See https://CRAN.R-project.org/package=admix for more information about the package on CRAN.

(2018), for a comprehensive survey on this topic. This model is of particular interest when considering generic situations distorted by an unexpected event, such as: i) the mortality excess due to the COVID-19 crisis, see Milhaud et al. (2024); ii) the presence of diseased tissues in microarray analysis, see McLachlan et al. (2006) and Benjamini and Hochberg (1995) for the related multiple testing problem, iii) variables observation, such as metallicity and radial velocity of stars, in the background of the Milky Way, see Walker et al. (2009); iv) trees diameters modeling in the presence of extra varieties, see Podlaski and Roesch (2014).

In this paper, the data of interest is made of $K \geq 2$ independent and identically distributed samples $X^{(i)} = (X_1^{(i)}, \dots, X_{n_i}^{(i)})$, for $i = 1, \dots, K$, each having the respective cumulative distribution function

$$L_i(x) = (1 - p_i)G_i(x) + p_i F_i(x), (2)$$

for all $x \in \mathbb{R}$, where the p_i 's are the unknown mixture proportions and the F_i 's are the unknown cumulative distribution function component associated with the *i*th sample. In practice, the G_i 's correspond to a well-known population. For instance, in the real-life application of mortality excess due to COVID-19 discussed in Section 7.1, G_i represents the historical national mortality profile for a given country, while the unknown F_i 's correspond to a new subpopulation, such as the specific mortality profile associated with the pandemic. Generally speaking, the F_i 's represent emerging phenomena not previously modeled, making this model particularly interesting for generic crisis or population transformation modeling.

The aim of this paper is to provide a clustering methodology to detect subgroups among the K existing samples that may have similar unknown, sometimes called nodular, components. This is of obvious interest for coordinated diagnosis and best practices sharing, based on the type of contamination impacting each population. To address this original problem, we will adopt a testing approach in which samples will be grouped together if an ad hoc test at a significance level $0 < \alpha < 1$ (to be determined) cannot reject the equality of their unknown components. More formally, we assume the existence of N true clusters denoted by \mathcal{G}_s , $1 \le s \le N \le K$ and defined by

$$\mathcal{G}_s = \{ i_{s,1} \le j \le K : \ F_j = F_{i_{s,1}} \}, \tag{3}$$

where $F_{i_{s,1}}$ denotes the first representative of group \mathcal{G}_s , indexed increasingly through $\{1,\ldots,K\}$ in the family of nodular components $\mathcal{N}=\{F_i,\ i=1,\ldots,K\}$. For convenience we denote by n_s the cardinality of \mathcal{G}_s , allowing us to number the elements of \mathcal{G}_s as $\mathcal{G}_s=\{i_{s,1},i_{s,2},\ldots,i_{s,n_s}\}$. Clearly, we have the following partition

$$\{1,\ldots,K\} = \bigcup_{s=1}^N \mathcal{G}_s, \quad \mathcal{G}_s \cap \mathcal{G}_{s'} = \emptyset, \qquad (1 \le s \ne s' \le N),$$
 (4)

with the below group separation assumption.

(GS) There exists real-sets $A_{s,s'} \subseteq \mathbb{R}$ with $\mu(A_{s,s'}) \neq 0$ such that for all $x \in A_{s,s'}$:

$$F_{i_{s,1}}(x) \neq F_{i_{s',1}}(x), \qquad (1 \le s \ne s' \le N),$$

where μ denotes a reference measure on the support of the F_i 's (e.g. Lebesgue measure over \mathbb{R} , counting measure over \mathbb{N}). Given the above framework, our clustering strategy will

consist in identifying recursively over $s \in \{1, ..., N\}$ the first representative $F_{s,1}$ of group \mathcal{G}_s along with the whole group itself. In this work, similarly to Patra and Sen (2016) or Milhaud et al. (2024), we will consider situations where the G_i 's and F_i 's distributions are: either i) absolutely continuous with respect to the Lebesgue measure, supported over \mathbb{R} , \mathbb{R}^+ or intervals of \mathbb{R} ; or ii) finite discrete or \mathbb{N} -discrete distributions such as Poisson or Binomial. All our results will be still valid in such setups provided that the G_i 's are all distinct. If certain pairs (G_i, G_j) , $1 \le i \ne j \le K$, are possibly equal a distinct procedure will then be implemented (see Appendix D of the Supplement in Milhaud et al. (2024) for details). Given the above cluster modelling, we first adress the basic statistical problem

$$H_0: F_1 = \cdots = F_k$$
 against $H_1: F_i \neq F_j$ for some $1 \leq i \neq j \leq k$, (5)

without assigning any specific parametric family to the F_i 's. Our clustering methodology will be grounded on the above k-sample testing problem, with the value of k potentially evolving from 1 to K along an algorithmic scheme. When k=2, the above problem has been addressed in Milhaud et al. (2022) under restrictive shape constraints such as the zero-symmetry of the F_i 's. More recently the two-sample testing problem has been revisited by Milhaud et al. (2024), who propose the so-called IBM (Inversion-Best Matching) testing approach requiring very relaxed identifiability and regularity conditions making this methodology much more suitable for real-life applications. When data is directly observed, i.e. there is no uncertainty about the component label for each observation, there exists empirical distribution clustering method such as the EP-MEANS introduced in Henderson et al. (2015) or the method introduced in Paul et al. (2022), see also recent references therein, in the high dimensional case. The issue with adapting these methods to the semiparametric mixture model (1) is twofold. First, the most general and flexible estimation method proposed by Patra and Sen (2016), needed to recover at a pre-stage the hidden contaminant distributions to be clustered, can provide unstable and sometimes inaccurate results in challenging cases, see for example Figure 8 in Pommeret and Vandekerkhove (2019) related to the Carina dataset (Milky Way radial velocities). Second, these clustering methods are always based on distance choices or test statistics which can respectively be discussed or require at least identified \sqrt{n} -asymptotic distributions to derive proper testing procedure which is not the case when considering Patra and Sen (2016). These reasons motivate the creation of the concept of true clusters, as defined in (3), at the contaminated but unlabeled populations level. This framework provides a clear view of the problem we aim to address and allows us to develop useful asymptotic results for our k-sample testing-based clustering strategy despite the lack of identifiability specific to the semiparametric mixture model (1), as detailed in Section 3 of Milhaud et al. (2024).

Our contribution is twofold: i) on the one hand we aim to generalize the work of Milhaud et al. (2024) to the k-sample case, when k is greater than 2; ii) on the other hand our objective is to derive a handy clustering algorithm grounded on the previous k-sample testing procedure, as described in (3) and (4). For that purpose we develop a data-driven methodology, inspired from Schwarz (1978) or Kallenberg and Ledwina (1995), allowing to select the most different populations pairs among all the possible pairs. More precisely we introduce the following set of pair indices: $S(k) = \{(i,j) \in \mathbb{N}^2; 1 \leq i < j \leq k\}$. Clearly S(k) contains d(k) = k(k-1)/2 elements that can be lexicographically ordered as follows: we denote (i,j) < (i',j') if i < i', or if i = i' and j < j', and we denote by $r_k[(i,j)]$

the associated rank of (i, j) in the set S(k). This ordering will be used to sum the test statistics over all the pairs of populations, and can be considered as the natural ordering over the elements of an upper triangle $k \times k$ matrix. For instance we have across the first row $r_k[(1,2)] = 1$, $r_k[(1,3)] = 2$, etc., when across the second row we have $r_k[(2,3)] = k$, $r_k[(2,4)] = k+1$, and so on. For $(i,j) \in S(k)$ we denote by $T_{i,j}$ the two-sample statistic used in Milhaud et al. (2024) to compare populations i and j, for $1 \le i \ne j \le k$. For simplicity matters, we drop the dependence on n since the statistic $T_{i,j}$ stands for T_n defined in the paragraph following expression (12) in Milhaud et al. (2024).

We can then build-up a sequence of statistics by slicing the set of index as follows: for slices s numbered from 1 to k-1, we define couples of index delimiters $(b_s^-, b_s^+) = (1 + (s-1)k - \frac{s(s-1)}{2}), sk - \frac{s(s+1)}{2})$ with $b_{s+1}^- = b_s^+ + 1$. This enables to define the sequence of embedded statistics U_r , the n-dependence dropped again for simplicity matters, as follows

slice 1:
$$U_r = \sum_{i=1}^r T_{1,1+i}, \quad (b_1^- = 1 \le r \le k - 1 = b_1^+),$$

slice 2: $U_r = U_{k-1} + \sum_{i=1}^{r-(k-1)} T_{2,2+i}, \quad (b_2^- = k \le r \le 2k - 3 = b_2^+),$
 \vdots
slice $s: U_r = U_{b_{s-1}^+} + \sum_{i=1}^{r-(b_s^- - 1)} T_{s,s+i}, \quad (b_s^- \le r \le b_s^+),$
 \vdots
slice $k-1: U_r = U_{b_{k-2}^+} + T_{k-1,k}, \quad (r = d(k)).$

ously the first two pairs of populations (1,2) and (1,3), and more generally U_r with r in slice of index s compares simultaneously the populations from 1 to s-1 with populations of upper ranks pairwizely through $U_{b_{s-1}^+}$ and population s with upper ranks lying in $\{s+1,\ldots,s+r-(b_s^--1)\}$ through $\sum_{i=1}^{r-(b_s^--1)}T_{s,s+i}$. Clearly, since the test statistics $T_{i,j}$ are positive, each statistic U_r is a sum of such r positive quantities and we have with probability 1 that $U_1 \leq \cdots \leq U_{d(k)}$. We then propose a penalized rule inspired from Schwarz criteria (1978) to select the most sensitive rank r given by S(n) in expressions (9) or (10) of Section 3.1. Under the null, we prove that the asymptotic limit distribution of our procedure coincides with the one obtained in the two-sample case given by the less penalized statistic $T_{1,2}$. It is also shown that our test statistic goes to infinity with n under the alternative. Our procedure is then adapted to construct a data-driven clustering algorithm able to classify the populations with equal unknown components. In order to pre-select a natural cluster to be tested by the k-sample test, we investigate the "closest" populations based on their pairwise associated (distance-based) statistics. We propose in addition a self-tuning method for the penalisation term involved in our k-sample test statistic that yields to an automated and easy-to-implement clustering procedure. The only required parameter is the asymptotic test level used to accept or not a cluster. This method is illustrated through an

By construction U_1 compares the first two populations (1,2), U_2 compares simultane-

extensive Monte Carlo experiment including very diverse situations and applied to a real life dataset dealing with the post COVID-19 mortality rates across a panel of 29 European countries.

The paper is organized as follows: In Section 2 we review recent results about the two-sample case, making the paper self-contained. Section 3 is devoted to the penalized testing rule and contains the main results of the paper. In Section 4 we develop a tuning method that allows our approach to be data-driven. The clustering algorithm is described in Section 5. Section 6 is devoted to an extensive simulation study covering the empirical level and power behaviour of our k-sample test procedure along with the numerical performances of our test-based clustering method. This latter section also includes a comparison with a natural Patra and Sen (2016) based k-means clustering competitor method. Section 7 ends the paper with a study dealing with the excess of mortality due to COVID-19 over a panel of European countries during the early times of the pandemic. The proofs of our theoretical results are relegated in Appendix.

2. Mathematical background

The tools and results presented in this section are recent developments in the semiparametric mixture models theory for which basically no parametric distribution family is assigned to the mixture components. This new way has been considered first in the two-component d-variate case, with $d \geq 3$, by Hall and Zhou (2003), and extended with two separate approaches by Hunter et al. (2007) and Bordes et al. (2006b), to the two-shifted symmetric components case, with d=1. Shortly after the first semiparametric approach dealing with the contamination model (1) was proposed by Bordes et al. (2006a) under moments and symmetry conditions. We recommend the reading of Xiang et al. (2019) and Gassiat (2019), which are both two excellent surveys about semiparametric mixtures and hidden Markov Models, to have a global view about the advances made these recents years on semiparametric missing data models.

In this section along with Section 3 we consider k, $2 \leq k \leq K$, samples among the K original samples still denoted for simplicity and without loss of generality $X^{(i)} = (X_1^{(i)}, \ldots, X_{n_i}^{(i)})$, for $i = 1, \ldots, k$. In the spirit of Milhaud et al. (2024), we consider $n = \min_{1 \leq i \leq k} n_i$, and define $\kappa_i \geq 1$ such that $n_i = \kappa_i n$, for all $i = 1, \ldots, k$. For $i \neq j \in \{1, \ldots, k\}$, we denote $\theta_{ij} = (p_i, p_j) \in \Theta_{ij} = \Theta_i \times \Theta_j$ the pair of unknown proportions associated to the ith and jth populations, respectively.

(A0) Assume that Θ_i is a $[\delta_1, \delta_2]$ -type compact set satisfying $0 < \delta_1 < 1 < \delta_2$, for all i = 1, ..., k.

Similarly to Milhaud et al. (2024), we notice that the unknown component associated with sample i can be recovered under the correct parameter p_i by using, for all $x \in \mathbb{R}$, the following inversion formula

$$F_i(x, L_i, p_i) = \frac{L_i(x) - (1 - p_i)G_i(x)}{p_i}, \qquad (i = 1, \dots, k).$$
(7)

To compare populations i and j we define the sub-(i, j) testing problem

$$H_0(i,j): F_i = F_j$$
 against $H_1(i,j): F_i \neq F_j$,

and consider the following discrepancy measure and its empirical counterpart

$$d[i,j](\theta_{ij}) = \int_{\mathbb{R}} \left(F_i(x, L_i, p_i) - F_j(x, L_j, p_j) \right)^2 dH(x)$$

$$d_n[i,j](\theta_{ij}) = \int_{\mathbb{R}} \left(F_i(x, \widehat{L}_i, p_i) - F_j(x, \widehat{L}_j, p_j) \right)^2 dH(x),$$

for $\theta_{ij} = (p_i, p_j)$ fixed in $\Theta_{ij} = \Theta_i \times \Theta_j$, where H is a positive measure over \mathbb{R} that allows to weight the square of the difference between F_i and F_j along the real line, and \widehat{L}_i denotes the empirical cumulative distribution function associated to the sample $X^{(i)}$. In practice we choose for H a uniform distribution when the support of the L_i 's is bounded or a probability distribution having a density supported by \mathbb{R} in the unbounded case, see also Appendix F of the Supplement in Milhaud et al. (2024) for further discussion about the choice of H. In the discrete case we simply choose for H the counting measure over the observations support.

We introduce now two assumptions connected to the identifiability and definite positiveness of the d-Hessian matrix. These assumptions are based on a cross-model identifiability condition inspired from the identifiability Theorem 1 in Teicher (1963).

(A1) Under $H_0(i,j)$ ($F_i = F_j = F_{ij}$), there exists at least three points $(x_1[i,j], x_2[i,j], x_3[i,j]) \in \mathbb{R}^3$ such that

$$\det \begin{pmatrix} G_i(x_1[i,j]) & G_j(x_1[i,j]) & F_{ij}(x_1[i,j]) \\ G_i(x_2[i,j]) & G_j(x_2[i,j]) & F_{ij}(x_2[i,j]) \\ G_i(x_3[i,j]) & G_j(x_3[i,j]) & F_{ij}(x_3[i,j]) \end{pmatrix} \neq 0.$$

(A2) Under $H_1(i,j)$ ($F_i \neq F_j$), there exists at least four points $(x_1[i,j], x_2[i,j], x_3[i,j], x_4[i,j]) \in \mathbb{R}^4$ such that

$$\det \begin{pmatrix} G_i(x_1[i,j]) & G_j(x_1[i,j]) & F_i(x_1[i,j]) & F_j(x_1[i,j]) \\ G_i(x_2[i,j]) & G_j(x_2[i,j]) & F_i(x_2[i,j]) & F_j(x_2[i,j]) \\ G_i(x_3[i,j]) & G_j(x_3[i,j]) & F_i(x_3[i,j]) & F_j(x_3[i,j]) \\ G_i(x_4[i,j]) & G_j(x_4[i,j]) & F_i(x_4[i,j]) & F_j(x_4[i,j]) \end{pmatrix} \neq 0.$$

The above pairwise-model conditions are stated and discussed in Milhaud et al. (2024).

For all $i \neq j \in \{1, \dots, k\}$ we consider

$$(\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)}) = \arg\min_{\theta_{ij} \in \Theta_{ij}} d_n[i, j](\theta_{ij}),$$

which is the estimated pair of parameters (p_i, p_j) that makes the unknown components F_i and F_j look the more similar according to the d discrepancy measure, which is then basically evaluated by

$$d_n[i,j](\widehat{\theta}_{ij}^{(1)},\widehat{\theta}_{ij}^{(2)}) = \int_{\mathbb{R}} \left(F_i(x,\widehat{L}_i,\widehat{\theta}_{ij}^{(1)}) - F_j(x,\widehat{L}_j,\widehat{\theta}_{ij}^{(2)}) \right)^2 dH(x).$$

Remark 1. There exists covariance between $\widehat{\theta}_{ij}^{(1)}$ and $\widehat{\theta}_{ij}^{(2)}$ (see Milhaud et al. (2024), Theorem 2). If j change to j' while i remains fixed, then $\widehat{\theta}_{ij}^{(1)}$ and $\widehat{\theta}_{ij'}^{(1)}$ (similarly, $\widehat{\theta}_{ij}^{(2)}$ and $\widehat{\theta}_{ij'}^{(2)}$) are not the same estimators. Indeed, they do not share the same distribution, even though they are both unbiased. That is why we use the notation $\widehat{\theta}_{ij}^{(1)}$ and $\widehat{\theta}_{ij}^{(2)}$ instead of \widehat{p}_i and \widehat{p}_j .

Remark 2. Milhaud et al. (2024) prove under $H_0(i,j)$ that $(\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)}) \to \theta_{ij}^* = (p_i^*, p_j^*)$ almost surely, with $d(\theta_{ij}^*) = 0$, where p_i^* and p_j^* are respectively the true value of the proportions involved in the $X^{(i)}$ and $X^{(j)}$ models, see expression (2). In contrast under $H_1(i,j)$, $(\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)}) \to \theta_{ij}^c = (\theta_{ij}^{(1)}, \theta_{ij}^{(2)})$ almost surely, a local minima of $\theta \mapsto d(\theta)$ with $d(\theta_{ij}^c) > 0$ and generally $\theta_{ij}^c \neq \theta_{ij}^*$.

We recall here the main result of Milhaud et al. (2024), see Theorem 2, that we use to construct our k-sample test. For $(i, j) \in \mathcal{S}(k)$ we consider

$$T_{i,j} = nd_n[i,j](\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)}), \tag{8}$$

the estimator of the n-discrepancy measure between populations i and j.

Lemma 3. Assume that (A1-2) hold. Then,

- i) Under $H_0(i,j)$, the statistic $T_{i,j} = U_n^0(i,j)$ converges in distribution towards $U^0(i,j)$, as $n \to +\infty$, where the limiting random variable $U^0(i,j)$ is fully identified (closed form stochastic integral) and tabulated.
- ii) Under $H_1(i,j)$, the statistic $T_{i,j} = U_n^1(i,j) + V_n^1(i,j)$, where $U_n^1(i,j)$ converges in distribution towards $U^1(i,j)$, as $n \to +\infty$, where the limiting random variable $U^1(i,j)$ is fully identified and tabulated and $V_n^1(i,j) = \lambda[i,j] \times n + o_{a.s.}(n)$ is a drift term, where

$$\lambda[i,j] = \int_{\mathbb{R}} \left(F_i(x, L_i, \theta_{ij}^{(1)}) - F_j(x, L_j, \theta_{ij}^{(2)}) \right)^2 dH(x) > 0.$$

Remark 4. In order to get our n-asymptotic results, we need to slightly adapt the matrices involved in the identification of the covariance matrix $\Sigma_W[i,j] = M_{i,j}(\theta_{ij}^c, \cdot)\Sigma_{i,j}M_{i,j}(\theta_{ij}^c, \cdot)^T$, $1 \leq i < j \leq k$, of Theorem 2 in Milhaud et al. (2024). In the k-sample setup involving multiple n_i -sample sizes, we must define

$$\Sigma_{i,j}(x,y) = \begin{bmatrix} \Sigma_i(x,y) & 0_{3\times 3} \\ 0_{3\times 3} & \Sigma_j(x,y) \end{bmatrix}, \quad and \quad M_{i,j}(\theta_{ij}^c,\cdot) = L_{i,j}(\cdot,\theta_{ij}^c) J_{i,j}^{-1}(\theta_{ij}^c) \mathcal{C}_{i,j},$$

where, since $n^{1/2} = (\kappa_i n)^{1/2} \kappa_i^{-1/2} = n_i^{1/2} \kappa_i^{-1/2}$, i = 1, ..., K, we can denote $\zeta_i = \kappa_i^{-1/2}$ and get

$$C_{i,j} = \begin{bmatrix} -\zeta_i & 0 & 0 & 0 & -\zeta_j & 0 \\ 0 & -\zeta_i & 0 & -\zeta_j & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \end{bmatrix}, J_{i,j}(\theta) = \begin{bmatrix} \ddot{a}[i,j](\theta) & 0_{2\times 2} \\ 0_{2\times 2} & Id_{2\times 2} \end{bmatrix},$$

and

$$L_{i,j}(\cdot,\theta) = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ -\frac{L_i(\cdot) - G_i(\cdot)}{p_i^2} & \frac{L_j(\cdot) - G_j(\cdot)}{p_j^2} & \frac{\zeta_i}{p_i} & -\frac{\zeta_j}{p_j} \end{bmatrix}.$$

This way any sample size departures between samples can be automatically handled.

3. The k-sample test

3.1 Main results

Let us remind that we aim to test condition (5) based on the observation of any k samples, $X^{(i)} = (X_1^{(i)}, \dots, X_{n_i}^{(i)}), i = 1, \dots, k$, picked from the original K-sample.

To solve this problem we propose to generalize the two-sample case by considering series of embedded statistics defined by (8), each new of them including a new pair of populations to be compared. To choose automatically the appropriate number of pairs of populations we introduce the following penalization procedure, in the spirit of the Schwarz (1978) criteria procedure. The principle of the penalized rule consists in selecting the rank s for which the penalized statistic U_s is the greatest. We introduce more specifically a sensitive rank defined by

$$S(n) = \min \left\{ \underset{1 \le r \le d(k)}{\operatorname{arg max}} \left(U_r - r \sum_{(i,j) \in S(k)} \ell_n(i,j) \mathbb{I}_{\{r_k(i,j) = r\}} \right) \right\}, \tag{9}$$

where $\ell_n(i,j)$ is a penalty term, and $\mathbb{I}_{r_k(i,j)=r}$ is 1 if $r_k(i,j)=r$ and 0 otherwise, indicating that we consider only the pair (i,j) associated to the order r. In the sequel we consider a penalty term independent from the population, i.e. $\ell_n(i,j)=\ell_n$ for all $i,j=1,\ldots,k$. Finally, the data driven selection can simply be rewritten as

$$S(n) = \min \left\{ \underset{1 \le r \le d(k)}{\arg \max} (U_r - r\ell_n) \right\}, \tag{10}$$

each statistic U_r being penalized by ℓ_n and by the number r, as a scale factor, of pairs of populations in it, according to the standard parsimony principle introduced by Schwarz (1978). In this sense, the sensitive rank S(n) will select automatically the rank associated to the most significant group of $T_{i,j}$'s statistics incorporated cumulatively in $U_{S(n)}$, see slicing scheme (6). We assume now that

(B)
$$\ell_n = n^{\varepsilon}$$
, with $0 < \varepsilon < 1$.

Since under the null each test statistic is a $O_P(1)$ when $\ell_n \to +\infty$, as shown in Lemma 3, it is expected that only the first statistic will be kept. The following result shows that under the null as defined in problem (5), the penalty effectively allows to select the first element of S(k) asymptotically.

Theorem 5. Assume that (A1-2) and (B) hold. Under H_0 , S(n) converges in probability towards 1, as $n \to +\infty$.

Theorem 6. Assume that (A1-2) and (B) hold. Under H_0 , $U_{S(n)}$ converges in distribution towards $U^0(1,2)$ given in Lemma 3, as $n \to +\infty$.

Then our data driven test statistic is

$$\widetilde{U}_n = U_{S(n)}. (11)$$

From Theorem 6, the asymptotic distribution of \widetilde{U}_n under H_0 is exactly the null limit distribution studied in the two-sample case and given in Lemma 3 i). We can use a tabulation of the random variable $U^0(1,2)$ which corresponds to a parametrized closed form stochastic integral, see Theorem 2 in Milhaud et al. (2024), that can be easily and consistently sampled, see Section 5 in Milhaud et al. (2024). By considering an empirical sample based $(1-\alpha)$ -quantile, denoted $\widehat{q}_{1-\alpha}$, of the stochastic integral we decide to consider the following H_0 -rejection rule

$$\widetilde{U}_n \ge \widehat{q}_{1-\alpha} \quad \Rightarrow \quad H_0 \text{ is rejected.}$$
 (12)

3.2 Alternatives

We consider the following series of alternative hypothesis

$$H_1(1)$$
 : $F_1 \neq F_2$,
 $H_1(r)$: $F_i = F_j$ for $r_k(i,j) < r$ and $F_i \neq F_j$ for $r_k(i,j) = r$,

with $1 < r \le d(k)$. The hypothesis $H_1(r)$ means that the *i*th and *j*th populations such that $r_k(i,j) = r$ are the first (in the S(k) ordering sense) with different unknown components.

Theorem 7. Assume that **(A1-2)** and **(B)** hold. Under $H_1(r)$, S(n) converges in probability towards r, as $n \to +\infty$, and \widetilde{U}_n goes to $+\infty$ in probability, that is, $\mathbb{P}(\widetilde{U}_n < \xi) \to 0$ for all $\xi > 0$.

4. Real world and finite samples: test statistic tuning

Experiments show that using (12) with small samples often leads to unsatisfactory results. We thus present here additional tools to improve the quality of our testing procedure in cases where the asymptotic regime is clearly not achieved.

4.1 About the penalty term ℓ_n

Since all our results are asymptotic we can replace Assumption (B) by

$$\ell_n(C) = Cn^{\varepsilon}$$
, with $0 < \varepsilon < 1$,

where C > 0 is any positive constant that will be used as a tuning parameter to adjust the test level (type-I error). The choice of ε is important for small and moderate sample sizes. Indeed a value ε close to 1 will favour a smaller S(n) value and a smaller value of the test statistic, with a lower rejection rate, while a value close to 0 will clearly empower the test. In fact, in the latter case, the divergence of \widetilde{U}_n is less likely to be compensated by the penalty term. The limit case $\varepsilon = 0$ coincides with a constant penalty which is the Akaike procedure, see Akaike (1974). Following Inglot and Ledwina (2006), we propose a rule to select ε based on the data itself. To introduce this rule, consider first the two-sample case:

$$F_{1}(x, \widehat{L}_{1}, \widehat{\theta}_{12}^{(1)}) - F_{2}(x, \widehat{L}_{2}, \widehat{\theta}_{12}^{(2)}) = (F_{1}(x, \widehat{L}_{1}, \widehat{\theta}_{12}^{(1)}) - F_{1}(x, L_{1}, \theta_{12}^{(1)})) - (F_{2}(x, \widehat{L}_{2}, \widehat{\theta}_{12}^{(2)}) - F_{2}(x, L_{2}, \theta_{12}^{(2)})) + (F_{1}(x, L_{1}, \theta_{12}^{(1)}) - F_{2}(x, L_{2}, \theta_{12}^{(2)})) = A(x) - B(x) + C(x),$$

where, because we have only two sample to compare, $\theta_{12}^c = \theta^c = (\theta_{12}^{(1)}, \theta_{12}^{(2)})$ is the minimizer of the contrast $d(\cdot)$, see (10) and (11) in Milhaud et al. (2024), with the property $(\theta_{12}^{(1)}, \theta_{12}^{(2)})$ equal to the true value of the weight parameters $\theta^* = (p_1^*, p_2^*)$, under H_0 which makes C(x) = 0, for all $x \in \mathbb{R}$, under the null. For all $x \in \mathbb{R}$, a straightforward expansion of A(x) is

$$A(x) = \frac{1}{\theta_{12}^{(1)}} \left(\widehat{L}_1(x) - L_1(x) \right) + \frac{1}{\theta_{12}^{(1)} \widehat{\theta}_{12}^{(1)}} \left(\widehat{\theta}_{12}^{(1)} - \theta_{12}^{(1)} \right) \left(\widehat{L}_1(x) - G_1(x) \right),$$

where $(\theta_{12}^{(1)}, \widehat{\theta}_{12}^{(1)}) \in [\delta_1, \delta_2]^2$, see Assumption (A0) about the parametric space to which the proportion parameters belong, and \widehat{L}_1 , respectively G_1 , are cumulative distribution functions which difference in modulus is bounded by 1. We then obtain

$$\sup_{x \in \mathbb{R}} \left(n^{1/2} |A(x)| \right) \leq \frac{1}{\delta_1} \sup_{x \in \mathbb{R}} \left(n^{1/2} \left| \widehat{L}_1(x) - L_1(x) \right| \right) + \frac{1}{\delta_1^2} \left| n^{1/2} \left(\widehat{\theta}_{12}^{(1)} - \theta_{12}^{(1)} \right) \right|$$

$$= A_1 + A_2.$$

By the law of the iterated logarithm for empirical processes, see Shorack and Wellner (1986), we have $A_1 = O_P((\log \log(n))^{1/2})$ and by Theorem 1 of Milhaud et al. (2024), which establishes the central limit theorem of $\widehat{\theta}_{12}^{(1)}$ towards $\theta_{12}^{(1)}$, we have that $A_2 = o_P((\log \log(n))^{1/2})$. Similarly we obtain $\sup_{x \in \mathbb{R}} (n^{1/2}|B(x)|) = O_P((\log \log(n))^{1/2})$. It follows that under the null we have

$$\sup_{x \in \mathbb{R}} \left(n^{1/2} \left| F_1(x, \widehat{L}_1, \widehat{\theta}_{12}^{(1)}) - F_2(x, \widehat{L}_2, \widehat{\theta}_{12}^{(2)}) \right| \right) \leq \gamma \sup_{x \in \mathbb{R}} \left(n^{1/2} |A(x)| \right) + \sup_{x \in \mathbb{R}} \left(n^{1/2} |B(x)| \right) \\
= O_P \left((\log \log(n))^{1/2} \right).$$

Under $H_1(1)$ there exists at least a real x such that $C(x) \neq 0$. In that case we have for all $\gamma > 0$ and for all positive sequence b_n such that $b_n \to +\infty$

$$\mathbb{P}\left(\sup_{x\in\mathbb{R}}\left(b_n|C(x)|\right)\leq\gamma\right)\quad\to\quad0.$$

In particular, choosing $b_n = n^{1/2}(\log(n))^{-1}$, it follows that under $H_1(1)$ we have

$$\mathbb{P}\left(\sup_{x\in\mathbb{R}}\left(n^{1/2}\left|F_1(x,\widehat{L}_1,\widehat{\theta}_{12}^{(1)}) - F_2(x,\widehat{L}_2,\widehat{\theta}_{12}^{(2)})\right|\right) \leq \gamma\log(n)\right) \rightarrow 0,$$

as $n \to +\infty$, while this probability goes to 1 under H_0 .

To generalize this principle to the k-sample case we can consider

$$S_{i,j} = \sup_{x \in \mathbb{R}} \left(n^{1/2} \left| F_i(x, \widehat{L}_i, \widehat{\theta}_{ij}^{(1)}) - F_j(x, \widehat{L}_j, \widehat{\theta}_{ij}^{(2)}) \right| \right). \tag{13}$$

Therefore, the expected conclusion is that small values of $\max_{(i,j)} S_{i,j}$, over $(i,j) \in \mathcal{S}(k)$, indicate that the unknown distributions over the considered k-sample is close to the null hypothesis while large values indicate an $H_1(r)$ -type alternative. To take into account this information we set

$$I_n(\gamma) = \mathbb{I}\{\max_{(i,j)\in\mathcal{S}(k)} S_{i,j} \le \gamma \log(n)\},\tag{14}$$

for some positive constant $\gamma > 0$. Under the null, from the above computations we can see that $S_{i,j} = O_P(1)$ for all $(i,j) \in \mathcal{S}(k)$. Under $H_1(r)$ as seen previously $\mathbb{P}(S_{i,j} \leq \gamma \log(n)) \to 0$ for $r_k(i,j) = r$. We can deduce, as $n \to +\infty$, the convergence in probability

$$\begin{cases}
I_n(\gamma) \to 1 &, \text{ under } H_0, \\
I_n(\gamma) \to 0 &, \text{ under } H_1(r).
\end{cases}$$
(15)

We then define a new penalty term by

$$\ell_n(C,\gamma) = C\left(I_n(\gamma) \ n^{\varepsilon_0} + (1 - I_n(\gamma)) \ n^{\varepsilon_1}\right),\tag{16}$$

where $\varepsilon_0 \approx 1$ and ε_1 is small enough in order to keep acceptable test levels even in the case of a wrong $I_n(\gamma)$ selection, favoring respectively the null or the alternative. The corresponding new selection rule is

$$\widetilde{S}(n) = \min \left\{ \arg \max_{1 \le r \le d(k)} \{ U_r - r\ell_n(C, \gamma) \} \right\}.$$
(17)

In practice we have obtained very good performances with the following values $\varepsilon_0 = 0.99$ and $\varepsilon_1 = 0.75$. At this stage, it now remains to explain how to pick appropriate tuning parameters C and γ . To do this we will use the information given both by (15) and by Theorem 5.

4.2 Data-driven choice for the parameter γ

Assume that we want to test the equality of k populations. From (15)–(16), a small value of γ yields a smaller penalty and thus a more powerful test. But, in parallel, we also want under the null

$$I_n(\gamma) = 1, (18)$$

which is more likely achieved for large values of γ . Thus to optimize the power of the test we search for the smallest γ which guarantees (18) under H_0 . For this we create a dummy H_0 -setup as follows: by splitting a population in two we obtain two identical sub-populations. Since such sub-populations are identically distributed the γ associated to their test statistic should satisfy (18). To optimize the power we choose the smaller γ satisfying this equality. We repeat this procedure b times and we obtain Algorithm 1. In practice, we set b = 6.

Algorithm 1: Tuning of the parameter γ .

- 1 for i = 1, ..., k do
- Split randomly the *i*th sample $X^{(i)}$ into two subpopulations, namely $X^{(i_1)}$ and $X^{(i_2)}$, of equal size $n_i/2$.
- 3 Compute $q_i^* = \sup_{x \in \mathbb{R}} \left((n_i/2)^{1/2} \left| F_{i_1}(x, \widehat{L}_{i_1}, \widehat{\theta}_{i_1 i_2}^{(1)}) F_{i_2}(x, \widehat{L}_{i_2}, \widehat{\theta}_{i_1 i_2}^{(2)}) \right| \right)$, where index i_1 (resp. i_2) refers to subsample $X^{(i_1)}$ (resp. $X^{(i_2)}$). /* eq. (13) */
- Repeat b times steps 2 and 3 to get b subpopulations under the null, and b values of q_i^* for each sample $X^{(i)}$. Write \bar{q}_i^* the mean of the q_i^* over the b repetitions.
- 5 Now, we have obtained k mean values for q_i^* , i = 1, ..., k. Since all q_i^* are obtained under the null, based on (14)-(15) we estimate γ by: $\widehat{\gamma} = \max_{1 \le i \le k} (\overline{q}_i^* / \log(n_i/2))$.

4.3 Data-driven choice for the parameter C

While the tuning of γ is based on the property (15), the tuning of the parameter C will use the result given by Theorem 5. From (16), a smaller value of C coincides with a smaller penalty yielding a larger test statistic and finally a larger power. Moreover, from Theorem 5 under the null we would expect

$$\widetilde{S}(n) = 1. (19)$$

We then use this property by choosing the larger C such that (19) is satisfied. In this way we can split a population into k' sub-populations, creating an artificial null hypothesis for which we modify C in order to get Condition (19) satisfied. The simplest choice of k' is k' = 3, which gives d(k') = 3 and seems to tune correctly the test procedure given in Algorithm 2.

Algorithm 2: Tuning of the parameter C.

- 1 for i = 1, ..., k do
- Split randomly the *i*th sample $X^{(i)}$ into k' subpopulations, of equal size n_i/k' . We obtain k new k'-sample problems under the null.
- 3 | for j = 1, ..., d(k') do
- 4 Compute U_j^i , where *i* refers to the *i*th population and *j* plays the role of r in (6).
- 5 Choose C^i such that $U_1^i C^i(n_i/k')^{\varepsilon_0} > U_j^i jC^i(n_i/k')^{\varepsilon_0}$, for $j = 1, \ldots, d(k')$.

 /* Choose C^i such that S(n) = 1 in every case

 */
- 6 Equivalently, we have $C^i = \max_j \left(\frac{U_j^i U_1^i}{(j-1)(n_i/k')^{\varepsilon_0}} \right)$, for $j = 1, \dots, d(k')$.
- **7** Finally, choose $\widehat{C} = \min_{1 \le i \le k} C^i$.

In a nutschell, we first tune γ and C, which allows us to build-up the penalty term ℓ_n defined in (16). This way, we get the corresponding order S(n) through Equation (17). Finally, the test statistic given by Equation (11) is used in the test procedure (12).

Remark 8. The technique employed here, which involves splitting a given sample into multiple (at least two) subsamples, creates an artificial H₀ framework. This framework differs markedly from real-life scenarios where F_i would be equal to F_i when considering two distinct samples. Specifically, the known component G_i of the ith contamination model should generally differ from G_i . However, this technique results in $G_i = G_i$, since the observations originally come from the same sample. This discrepancy affects the estimation process and, consequently, the testing procedure, as discussed in Appendix D of the Supplement in Milhaud et al. (2024). Therefore, it is crucial to verify whether this artificial procedure significantly impacts the choice of parameters γ and C, compared to those selected by the tuning process in a real-life scenario under the null hypothesis. To address this, we repeat the following simulation scheme 100 times under H_0 : (i) simulate four samples (populations) following contamination/admixture models, (ii) use Algorithms 1 and 2 to obtain the distributions of γ and C under the artificial H_0 setting, and (iii) apply simplified versions of Algorithms 1 and 2 (omit step 2 and consider the samples themselves, since we are under the null) to get the distributions of γ and C without the technique. Finally, we compare the obtained distributions. As evidenced in many other frameworks, Figure 1 demonstrates that our tuning process, incorporating this technique, remains consistent. Although the distributions of the parameters γ and C are slightly different, they appear similar, with modes closely aligned.

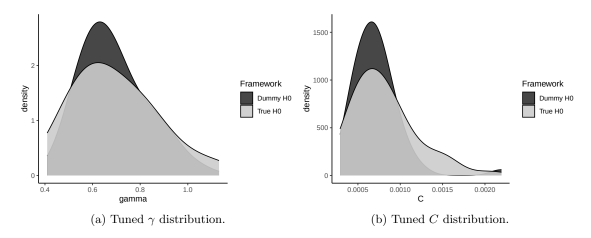


Figure 1: Distributions of selected tuning parameters obtained from Algorithms 1 and 2 over 100 repetitions, under the dummy H_0 framework and the true one.

5. Clustering strategy

In the sequel we propose to adapt the previous test procedure to obtain a data-driven method to cluster K populations into N subgroups characterized by a common unknown nodular component. The novelty here lies in the fact that we will be able to cluster unlabeled behaviours presenting similar distributions, contrary to classical existing clustering strategies that are based on directly/fully observed phenomenons. Moreover it is worth to notice that the number N of clusters is not assumed at the beginning but is automatically deduced at the end of the procedure.

Assume that we observe K independent samples $X^{(i)} = (X_1^{(i)}, \dots, X_{n_i}^{(i)})$, $i = 1, \dots, K$, made separately of independent and identically distributed observations. To build the first cluster we consider the two closest populations, i.e. leading to the smallest $T_{i,j}$ -statistic, $i \neq j \in \{1, \dots, K\}$. Two populations are thus proposed to be merged to create the first group \mathcal{G}_1 . We test their equality according to the testing procedure (12) to confirm the construction of such a cluster. We continue to add populations to the group until the test rejects equality. Once this first cluster \mathcal{G}_1 is fully identified: close the cluster, remove clustered populations from the initial collection of samples, and create a new cluster \mathcal{G}_2 . Then look for still unclustered neighboors from the last studied sample that led to reject H_0 . Again we select the biggest collection of samples, among the remaining pool, that is tested to share a common unknown component with the latter. This creates our second cluster. One can iterate this several times until every sample is associated with a cluster. Algorithm 3 describes our so-called KCMC (K-sample Contamination Model Clustering) algorithmic clustering strategy, with $S = \{1, \dots, K\}$ the set of population indices, c the cluster id and S_c the members of cluster c.

Algorithm 3: K-sample Contamination Model Clustering (KCMC).

```
1 Initialization: create the first cluster to be filled, i.e. c=1. By convention, S_0=\emptyset.
 2 Select (x,y) = \operatorname{argmin}\{nd_n[i,j](\widehat{\theta}_{ij}^{(1)},\widehat{\theta}_{ij}^{(2)}); i \neq j \in S \setminus \bigcup_{m=1}^c S_{m-1}\}.
 3 Test H_0 between x and y (two-sample test).
                                                                                          /* using (12)
 4 if H_0 is not rejected then
 S_1 = \{x, y\}
                                                          /* fill in the first cluster
                                                                                                                   */
 6 else
 7 | S_1 = \{x\}, S_{c+1} = \{y\} and then c = c+1 /* close, open new one
                                                                                                                   */
 8 while S \setminus \bigcup_{m=1}^{c} S_m \neq \emptyset do
         /* seek unclustered neighboors, select the closest one
                                                                                                                   */
        Select u = \operatorname{argmin}_{j} \{ nd_{n}[i, j](\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)}); i \in S_{c}, j \in S \setminus \bigcup_{m=1}^{c} S_{m} \}
 9
        Test H_0 the simultaneous equality of all the F_j, j \in S_c:
10
11
         if H_0 not rejected then
          | \text{ put } S_c = S_c \bigcup \{u\} 
12
13
          S_{c+1} = \{u\} \text{ and } c = c+1
14
```

The procedure stops when all populations are merged or when all populations have been considered by the algorithm. Note that since the first two populations selected are the closest, if the test rejects their equality then the algorithm stops and returns as many clusters as populations. The procedure is straightforward since the parameters γ and C are data-driven (up to the prior choice of b and k', see Algorithms 1 and 2 in Section 4.2). Furthermore, we deduce from Theorem 7 the following property.

Proposition 9. With a probability that tends to 1 as $n \to +\infty$, the number N^* of groups detected by Algorithm 3 satisfies $1 \le N \le N^* \le K$, where N denotes the true unknown number of groups (or clusters).

From Proposition 9 the number of clusters obtained from the KCMC algorithm is potentially greater than N. Moreover, from Theorem 7, asymptotically all distributions of each clusters are equal. Thus the only possible error is that a real group has been splitted into several other groups, which can happen because we have an asymptotic test level $\alpha = 5\%$, see testing rule (12). This parameter clearly reflects the threshold for accepting the creation of a group. One way to check the stability of the clusters is to change this threshold, for example by decreasing α to see if the groups merge then. We illustrate this point in our real world application, see Section 7. The tuning strategy of Sections 4.2 and 4.3 allows us to achieve very good performances across our simulation study. In particular the detected number N^* of clusters often does not exceed the actual number N.

6. Simulation study

All our numerical experiments were performed thanks to the R package admix², especially developed for estimation, test and clustering of populations generated from admixture models. To begin with, we test the influence of the number of populations under consideration, to check whether or not this impacts the quality of our k-sample testing procedure. For this purpose, we let k vary from 2 to 10. The populations are drawn from different distributions supported by various types of real-sets. We provide here the results for distributions supported over \mathbb{R} (Gaussian mixtures), but simulations were extended to other supports such as \mathbb{N} (Poisson mixtures) or \mathbb{R}^+ (Gamma mixtures) with very similar conclusions. The proportions of the unknown components are fixed all along the simulation scheme for easier comparisons. To evaluate the empirical level (and power) of the k-sample test, we use a Monte Carlo approach where each of the B experiments is performed in the same way. We also make the sample size vary to illustrate the asymptotic properties of our results. Unless otherwise stated, all our simulations were performed with fixed values $\varepsilon_0 = 0.99$ and $\varepsilon_1 = 0.75$ in (16) and (17), meaning that we use the tuning process described in Sections 4.1, 4.2 and 4.3. As expected, the tuning process reveals to be decisive to improve the power of the test, but has no real influence under the null. This is in line with common sense, since tuning parameters γ and C are estimated under the null. Once the quality of the k-sample test validated, we derive extra simulations to assess the performance of our clustering algorithm itself.

^{2.} See the package description at https://cran.r-project.org/web/packages/admix/index.html

6.1 Empirical level of our k-sample testing procedure

We draw k populations from two-component Gaussian mixtures, where the k simulated known components are distributed according to different Gaussian distributions. On the contrary, those k populations share the same unknown component distribution (namely a standard normal distribution). For each simulation being part of the Monte Carlo procedure, we implement the following steps: (i) generate the k populations, each one following an admixture model; (ii) perform the k-sample test; (iii) retrieve which penalty rule (similarly which ε , either ε_0 or ε_1) and which rank $\widetilde{S}(n)$ have been selected, along with the p-value of the test. We repeat this simulation scheme B=100 times in order to estimate the empirical level of our test procedure (12). Table 1 reports the parameters involved in each simulated population for three different sample sizes (about 400, 1000 and 3000 observations), along with the results related to the main indicators assessing the efficiency of our procedure. More comprehensively, Table 1 stores in its last four columns how often the right penalty rule (16) has been chosen (in percent), the 90%-percentile of the distribution of the selected order $\widetilde{S}(n)$ defined in (17), the mean of the 100 p-values obtained when testing, and finally the empirical level of the test.

Given that the selected sample-based quantile considered in (12) was fixed as the 95%-percentile of the tabulated distribution, it is expected that the empirical level of the test (last column) stays close to 5%. Looking at the results, our test procedure looks to perform globally well. Most of the time, the right penalty rule and the right testing rank have been selected. Indeed, in more than 90 out of 100 experiments, the selected order $\tilde{S}(n)$ has the correct value (equal to 1, since we are under H_0). Moreover, the number k of populations involved in the k-sample test does not seem to impact our testing procedure. Even when

Table 1: k-sample test. Reported $\widetilde{S}(n)$ corresponds to the 90%-percentile of the $\widetilde{S}(n)$ distribution over 100 experiments, and p-val is the average of the obtained p-values.

					S	amples								
i	1	2	3	4	5	6	7	8	9	10				
p_i	0.3	0.8	0.6	0.4	0.9	0.2	0.4	0.15	0.7	0.5	Pen.			Emp.
G_i	N(2, 0.7)	N(4, 1.1)	N(3, 0.8)	$\mathcal{N}(-1, 0.3)$	$\mathcal{N}(-3, 0.2)$	$\mathcal{N}(-5, 0.4)$	$\mathcal{N}(3.5, 0.1)$	$\mathcal{N}(-4, 0.7)$	$\mathcal{N}(-2.5, 1)$	$\mathcal{N}(1.5, 0.3)$	rule	$\widetilde{S}(n)$	p-	level
n_i	347	449	308	382	426	372	440	447	474	424	(%)	5(11)	value	(10^{-2})
	F_1	F_2	F_3	F_4	F_5	F_6	F_7	F_8	F_9	F_{10}				
k=2	N(0, 1)	N(0, 1)									100	1	0.53	5
k=4	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)							98	1	0.74	3
k=6	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)					96	1	0.76	4
k=8	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)			92	1	0.83	6
k=10	N(0, 1)	N(0, 1)	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	95	1	0.9	5
n_i	1011	1027	1077	1019	903	942	971	1065	1071	1068				
	F_1	F_2	F_3	F_4	F_5	F_6	F_7	F_8	F_9	F_{10}				
k=2	N(0, 1)	N(0, 1)									100	1	0.4	7
k=4	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)							100	1	0.77	2
k=6	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)					100	1	0.8	4
k=8	N(0, 1)	N(0, 1)	$\mathcal{N}(0,1)$	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)			87	1	0.8	8
k=10	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	N(0, 1)	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	86	1	0.83	6
n_i	3187	2847	3189	3175	3042	2989	3184	2868	2998	3193				
	F_1	F_2	F_3	F_4	F_5	F_6	F_7	F_8	F_9	F_{10}				
k=2	N(0, 1)	N(0, 1)									100	1	0.48	6
k=4	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)							100	1	0.71	3
k=6	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)					98	1	0.78	4
k=8	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0,1)$	N(0, 1)	N(0, 1)			93	1	0.81	6
k=10	N(0, 1)	N(0, 1)	$\mathcal{N}(0,1)$	N(0, 1)	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	N(0, 1)	$\mathcal{N}(0,1)$	$\mathcal{N}(0,1)$	94	1	0.92	3

some populations have overlapping components, the quality of the test remains satisfactory. Also, the same simulations were performed without using the tuning process, with almost no impact on test levels. In that latter case, we have set $\varepsilon = 0.87$ for the penalty given by Assumption (B) in (10), as this value lies exactly in the middle of $[\varepsilon_1, \varepsilon_0] = [0.75, 0.99]$. Setting ε to way lower values led to seriously deteriorate the test levels in finite sample size applications, which validates the need to keep a strong penalty under the null. Of course, this global picture may change depending on the chosen parameters to conduct the simulation study. For instance, much higher variances for the mixture components would clearly affect our results.

Remark 10. We observed higher empirical levels when the population to test is strongly under-represented. If the product $n_i p_i$ is low, say below 50 for our experiments, the estimation of the mixture proportion p_i deteriorates. This spreads out to the computation of supremum in (13), which mechanically increases and leads to the wrong choice in the penalization rule, i.e. taking ε_1 instead of ε_0 .

6.2 Empirical power

Now, we aim to study the power of our testing strategy, meaning how our k-sample test performs in detecting that (at least) two of the k populations have different unknown component distributions. For ease of comparison, we keep the same known and unknown component distributions as previously. The parameters involved in Gaussian mixtures are stored in Table 2, showing that some considered frameworks correspond to critical situations where mixture components can be highly overlapping, see for instance when k=2. As already mentioned, the tuning process is essential here to correctly detect the alternative. Indeed, the penalty term should not compensate the explosion of the test statistic, which means that taking $\varepsilon=0.75$ (i.e. $\varepsilon=\varepsilon_1$) instead of $\varepsilon=0.87$ leads to very distinct results. Let us focus here on the case k=10, and emphasize various possibilities depending on the number of different unknown component distributions involved across those k populations. Table 2 leads to several interesting conclusions.

First, the power of the test is much more sensitive to sample size than its level. This is not surprising, as detecting deviations from the null hypothesis requires substantial evidence that the two unknown component distributions differ. This can be challenging when dealing with mixtures with overlapping components and moderate sample sizes. However, once the product $n_i p_i$ reaches a sufficiently large value, say around 100, the power of the test approaches 1. With small sample sizes, selecting the appropriate penalty rule can be challenging in most cases (except when k = 2). In practice, ε_0 is often chosen over ε_1 . This is because dividing the population at the beginning of Algorithm 1 reduces the original sample size, likely increasing the variability of $S_{i,j}$. Consequently, the quantity $\max(S_{i,j})/\log(n)$ increases, leading to a higher value of γ , as detailed in Algorithm 1. This highlights the impact of sample size in (16), which generally raises the penalty term in (17), potentially compensating for the growth of the contrast function that typically signals a departure from the null hypothesis. Thus, the testing procedure tends to indicate we are more likely under the null, as seen in the 90%-percentile of $\tilde{S}(n)$, which sometimes equals 1 when k = 10.

Challenging our methodology with moderate sample sizes improves the overall quality of our testing procedure. Indeed, both the penalty rule and the correct order tend to be ap-

					Sa	mples								
i	1	2	3	4	5	6	7	8	9	10	-			
p_i	0.3	0.8	0.6	0.4	0.9	0.2	0.4	0.15	0.7	0.5	Pen.			Emp. power
G_i	N(2, 0.7)	N(4, 1.1)	N(3, 0.8)	$\mathcal{N}(-1, 0.3)$	$\mathcal{N}(-3, 0.2)$	$\mathcal{N}(-5, 0.4)$	$\mathcal{N}(3.5, 0.1)$	$\mathcal{N}(-4, 0.7)$	$\mathcal{N}(-2.5, 1)$	$\mathcal{N}(1.5, 0.3)$	rule	$\widetilde{S}(n)$	p-	Tune/NoTune
n_i	347	449	308	382	426	372	440	447	474	424	(%)	$\mathcal{O}(n)$	value	(10^{-2})
	F_1	F_2	F_3	F_4	F_5	F_6	F_7	F_8	F_9	F_{10}	-			
k=2	N(0, 1)	$\mathcal{N}(0.3, 1)$									n.a.	1	0.24	n.a. / 36
k=4	N(0, 1)	N(1, 1)	$\mathcal{N}(0.3, 1)$	N(0, 1)							15	6	0.46	20 / 11
k=7	N(0, 1)	N(0, 1)	N(1, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0.5, 1)$	N(0, 1)				30	21	0.7	26 / 3
k=10	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0.5, 0.5)$	N(0, 1)	N(0, 1)	N(0, 1)	30	1	0.8	7 / 1
k=10	N(0, 1)	$\mathcal{N}(-0.5, 1)$	$\mathcal{N}(0.3, 0.5)$	N(0, 1)	N(0, 1)	$\mathcal{N}(0.7, 0.7)$	N(0, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(-0.2, 1)$	25	1	0.72	9 / 1
k=10	$\mathcal{N}(0,1)$	$\mathcal{N}(-0.5, 1)$	$\mathcal{N}(0.3, 0.5)$	$\mathcal{N}(0,1)$	$\mathcal{N}(1,1)$	$\mathcal{N}(0.7, 0.7)$	N(0, 1)	N(-0.4, 1)	N(0.9, 3)	$\mathcal{N}(-0.2, 1)$	70	45	0.26	63 / 2
n_i	1011	1027	1077	1019	903	942	971	1065	1071	1068				
	F_1	F_2	F_3	F_4	F_5	F_6	F_7	F_8	F_9	F_{10}	•			
k=2	N(0, 1)	$\mathcal{N}(0.3, 1)$									n.a.	1	0.09	n.a. / 74
k=4	N(0, 1)	N(1, 1)	$\mathcal{N}(0.3, 1)$	N(0, 1)							41	6	0.22	50 / 18
k=7	N(0, 1)	N(0, 1)	N(1, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0.5, 1)$	N(0, 1)				95	21	0.04	95 / 1
k=10	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0.5, 0.5)$	N(0, 1)	N(0, 1)	N(0, 1)	95	45	0.23	72 / 1
k=10	N(0, 1)	$\mathcal{N}(-0.5, 1)$	$\mathcal{N}(0.3, 0.5)$	N(0, 1)	N(0, 1)	$\mathcal{N}(0.7, 0.7)$	N(0, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(-0.2, 1)$	85	45	0.13	84 / 2
k=10	$\mathcal{N}(0,1)$	$\mathcal{N}(-0.5, 1)$	$\mathcal{N}(0.3, 0.5)$	$\mathcal{N}(0,1)$	$\mathcal{N}(1,1)$	$\mathcal{N}(0.7, 0.7)$	$\mathcal{N}(0,1)$	N(-0.4, 1)	N(0.9, 3)	N(-0.2, 1)	100	45	0	99 / 1
n_i	3187	2847	3189	3175	3042	2989	3184	2868	2998	3193				
	F_1	F_2	F_3	F_4	F_5	F_6	F_7	F_8	F_9	F_{10}	-			
k=2	N(0, 1)	$\mathcal{N}(0.3, 1)$									n.a.	1	0.009	n.a. / 96
k=4	N(0, 1)	N(1, 1)	$\mathcal{N}(0.3, 1)$	N(0, 1)							99	6	0.001	99 / 12
k=7	N(0, 1)	N(0, 1)	N(1, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0.5, 1)$	N(0, 1)				100	21	0	98 / 1
k=10	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	N(0, 1)	$\mathcal{N}(0.5, 0.5)$	N(0, 1)	N(0, 1)	N(0, 1)	100	45	0	100 / 3

Table 2: k-sample test under H_1 , with emphasis on different settings when K = 10. Same interpretation as Table 1 for last columns (n.a. stands for not applicable).

propriately selected. As expected, the greater the number of distinct unknown components among the k populations, the more powerful the test. Note that when k=2, Algorithms 1 and 2 are unnecessary since they only influence the penalty rule, which helps detect the correct number of summands in the test statistics; in this case, there is only one summand by construction.

 $\mathcal{N}(0,1)$

 $\mathcal{N}(0,1)$

 $\mathcal{N}(-0.2, 1)$

Finally, Table 2 shows in its last column the improvement in test power achieved through the tuning process, confirming that the power of the k-sample test significantly depends on it. While results may vary with different simulation parameters, we aimed to cover a broad range of simulation setups to thoroughly test the robustness of our procedure.

6.3 Clustering

k=10 $\mathcal{N}(0,1)$ $\mathcal{N}(-0.5,1)$ $\mathcal{N}(0.3,0.5)$

N(0, 1)

N(0, 1)

 $\mathcal{N}(0.7, 0.7)$

Hereafter, we are willing to cluster the unknown components F_i 's over K populations under study, only based on the observation of samples drawn from L_i 's admixtures, with known G_i 's and unknown F_i 's. Let start with a close description of our clustering frameworks before discussing our results.

Clustering schemes description. We dedicated the previous section to the performance study of our k-sample testing procedure $(2 \le k \le K)$, since the quality and the robustness of our clustering algorithm strongly relies on it. Now, we would like to recover simulated clusters over K = 10 populations. Various frameworks are investigated, from the extreme cases of one single cluster up to ten clusters. In-between, we also study situations where we have both, sizewize speaking, unbalanced and balanced clusters. Figure 2 illustrates the simulated densities in the four considered settings, denoted further from (a) to (d). In the

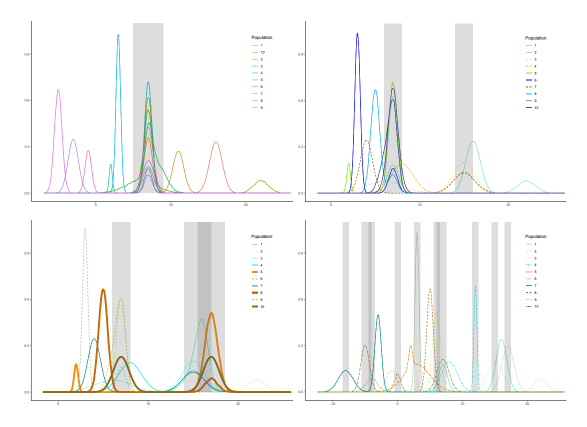


Figure 2: Simulated densities. From top left to bottom right: 1 cluster, 2 clusters ((1,2,5,6,8,9,10); (3,4,7)), 3 clusters ((1,3,4,7), (2,6,9)) and (5,8,10), 10 clusters.

first case (one single cluster) the common unknown component is distributed according to $F_i \sim \mathcal{N}(7, 0.5), i = 1, \dots, K$, whereas two clusters appears for the second case (grey areas). The densities of the populations associated with these two clusters are depicted through different line types (plain and dotted). In the third case, the densities associated to the three balanced clusters are displayed with different line types and widths. Clustering the unknown components of these populations just based on the extra knowledge of their known components, is not straightforward. Indeed, there is sometimes very strongly overlapping components among the various populations, see for instance the 3rd and 4th populations in the 1-cluster example. Moreover, some of the clusters can be close from one to another, see for instance the third case where two of the three clusters are not well separated because of close means and higher variances. All the parameters involved in those simulations are stored in Table 3.

Clustering performances. Once again, we use a Monte Carlo approach and perform the clustering task B times for each of the four cases aforementioned. As the clustering process is computationally intensive (it requires to perform many k-sample tests), we set B = 20. However, we also considered B = 50 for some of our examples, which led to minor modifications of the results without changing the global picture. Given the parameters

Table 3: Clustering setups: the 2-cluster is made of samples (1,2,5,6,8,9,10) and (3,4,7), when the 3-cluster is made of samples (1,3,4,7), (2,6,9) and (5,8,10).

Populations i	1	2	3	4	5	6	7	8	9	10
Weight p_i	0.3	0.8	0.6	0.4	0.9	0.2	0.4	0.15	0.7	0.5
Sample size n_i	312	271	293	322	289	282	279	280	294	324
1 cluster G_i	$\mathcal{N}(16, 0.7)$	$\mathcal{N}(22,1)$	$\mathcal{N}(6,2)$	N(8, 1.2)	$\mathcal{N}(2, 0.2)$	$\mathcal{N}(3, 0.3)$	$\mathcal{N}(-3, 0.4)$	$\mathcal{N}(-5, 0.5)$	$\mathcal{N}(-1, 0.1)$	$\mathcal{N}(11, 0.7)$
G_i (2&3 clusters)	$\mathcal{N}(16, 0.7)$	$\mathcal{N}(22,1) \ \mathcal{N}(7,0.5) \ \mathcal{N}(7,0.5)$	$\mathcal{N}(6,2)$	$\mathcal{N}(8, 1.2)$	$\mathcal{N}(2, 0.2)$	$\mathcal{N}(3, 0.3)$	$\mathcal{N}(4, 0.4)$	$\mathcal{N}(5, 0.5)$	$\mathcal{N}(6, 0.6)$	$\mathcal{N}(7, 0.7)$
F_i (2 clusters)	$\mathcal{N}(7, 0.5)$		$\mathcal{N}(15,1.1)$	$\mathcal{N}(15, 1.1)$	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(15, 1.1)$	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(7, 0.5)$
F_i (3 clusters)	$\mathcal{N}(15, 1.1)$		$\mathcal{N}(15,1.1)$	$\mathcal{N}(15, 1.1)$	$\mathcal{N}(17, 0.7)$	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(15, 1.1)$	$\mathcal{N}(17, 0.7)$	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(17, 0.7)$
G_i (10 clusters)	$\mathcal{N}(16, 0.7)$	$\mathcal{N}(22,1)$	$\mathcal{N}(6,2)$	$\mathcal{N}(8, 1.2)$	$\mathcal{N}(2, 0.2)$	$\mathcal{N}(3, 0.3)$	$\mathcal{N}(-3, 0.4)$	$\mathcal{N}(5, 0.5)$	$\mathcal{N}(-1, 0.1)$	$\mathcal{N}(7, 0.7)$
F_i (10 clusters)	$\mathcal{N}(7, 0.5)$	$\mathcal{N}(6,0.6)$	$\mathcal{N}(15,1.1)$	$\mathcal{N}(12, 0.05)$	$\mathcal{N}(3, 2)$	$\mathcal{N}(-4, 0.9)$	$\mathcal{N}(-8, 1.1)$	$\mathcal{N}(0, 0.5)$	$\mathcal{N}(17, 0.4)$	$\mathcal{N}(-5, 0.2)$

of the simulations (see Table 3) in the four studied frameworks (a) to (d), we expect our procedure to find respectively the following clusters: (1,2,3,4,5,6,7,8,9,10) in the 1-cluster case; (1,2,5,6,8,9,10) and (3,4,7) in the 2-cluster case; (1,3,4,7), (2,6,9) and (5,8,10) in the 3-cluster case; and finally (1),(2),(3),(4),(5),(6),(7),(8),(9),(10) in the 10-cluster framework.

In practice, there exists many ways for the clustering algorithm to reach wrong conclusions. Either it detects the right number of clusters but does not affect the right populations to the right clusters (which should not happen asymptotically), or it selects straight out a wrong number of clusters. In the latter case the algorithm tends to overestimate the correct number of clusters, leading to clusters with wrong number of elements and isolated populations. It is difficult to summarize all possible encountered wrong answers through one single indicator. In our case, we have chosen to measure the performance of the clustering algorithm through classification matrices (also called heatmaps). Indeed, it seems to us that it is an efficient and yet simple indicator.

Figures 3 and 4 display examples of such matrices for different sample sizes across our simulation setups. The reading of heatmaps is pretty straightforward. First, they are symmetric, with errors stored in the off-diagonal terms (of course a given population is always clustered with itself). For these non-diagonal terms, one counts how many times (among the B experiments) the clustering algorithm clustered each population with the other ones. Then, comparing this to the expected clusters, it is straightforward to deduce the percentage of correct classifications. To make the reading easier, we also report within each cell those percentages and use a colour gradient illustrating the quality of clustering (intense dark brown refers to best results).

To simplify, our heatmaps are organized by blocks, each block corresponding to an expected cluster. This means that a perfect clustering has 100% of right classifications for every blocks. This is the case for instance in Figure 3 in the fourth setting, see case (d). In the three other frameworks, the KCMC algorithm is sometimes mistaking, but results show that these errors remain reasonable most of time despite small sample sizes $(n_i \approx 300, i = 1, ..., K)$. The worst case seems to be the 8th population which is very often isolated over the 1, 2 or 3-cluster settings, see cases (a), (b) and (c). The reason for this, is very likely to be the quality of the estimation of the unknown proportion p_8 (the lowest weight, equal to 0.15), which deteriorates the estimation of the unknown component F_8 to be clustered (see further Figure 5, where the corresponding decontaminated estimated density \hat{f}_8 , obtained from (7), appears to be wrongly bimodal). Figure 4 illustrates the

phenomenon that was already observed when studying the performance of the k-sample test. Increasing the sample size has a strong beneficial influence on the clustering efficiency. Indeed, all the clusters are correctly identified under most of our simulations settings when the sample size reaches about 800 ($n_i \approx 800$, i = 1, ..., K). As already mentioned, what matters the most in reality is the value of the product $n_i p_i$, which should remain large enough to guarantee the asymptotic properties of our procedure (in this case, $n_i p_i \approx 120$ at the lowest).

The reader must keep in mind that using this clustering method can be tricky in applications having a low number of observations. Indeed, our Monte Carlo approach allows, through our B experiments, to recover the right clusters with a majority voting procedure, which would not be feasible in a one shot real life application. Finally, once the clusters are recovered, useful information can be deduced. For instance, knowing that the unknown weights are consistently estimated inside each cluster, one can retrieve the estimated weights of the unknown perturbation impacting the original populations. Table 4 provides such results based on our simulation parameters (not applicable with 10 clusters since weights are not consistently estimated when there are no similar unknown components) and reports the mean of estimated proportions. Indeed, a cluster containing populations i, j and k would lead to estimators $(\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)})$, $(\widehat{\theta}_{ik}^{(1)}, \widehat{\theta}_{ik}^{(2)})$ and $(\widehat{\theta}_{jk}^{(1)}, \widehat{\theta}_{jk}^{(2)})$ (see Remark 1). One can then compute the mean of $\widehat{\theta}_{ij}^{(1)}$ and $\widehat{\theta}_{ik}^{(1)}$ as an estimator of p_i , the mean of $\widehat{\theta}_{ij}^{(2)}$ and $\widehat{\theta}_{jk}^{(1)}$ for p_j , and the mean of $\widehat{\theta}_{ik}^{(2)}$ and $\widehat{\theta}_{jk}^{(2)}$ for p_k . Then, the corresponding decontaminated densities can be nicely illustrated, see Figure 5.

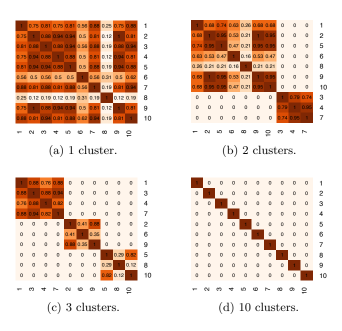


Figure 3: Heatmap describing the efficiency of our clustering algorithm ($n \approx 300$, see Table 3), with proportion of right predictions in each cell (value 0 means no error).

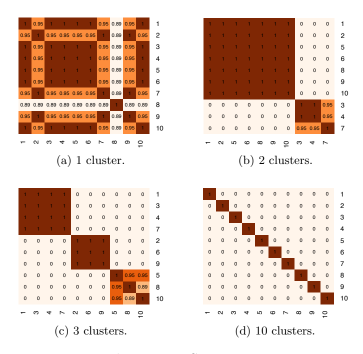


Figure 4: Heatmap with $n \approx 800$. Same interpretation as Figure 3.

Table 4: Mean of estimated unknown weights $(\widehat{\theta}_{ij}^{(1)}, \widehat{\theta}_{ij}^{(2)})$ over all pairs (i, j) of populations belonging to the same identified cluster (n = 300)

i	1	2	3	4	5	6	7	8	9	10	
Real weight p_i	0.3	0.8	0.6	0.4	0.9	0.2	0.4	0.15	0.7	0.5	
		Mean estimated weights									
Case of 1 cluster	0.271	0.798	0.669	0.424	0.894	0.159	0.378	0.174	0.676	0.390	
Case of 2 clusters	0.291	0.829	0.603	0.417	0.884	0.253	0.395	0.147	0.701	0.752	
Case of 3 clusters	0.367	0.821	0.581	0.441	0.903	0.216	0.448	0.151	0.740	0.494	

Comparison with a k-means competitor. The novelty of this work, which involves clustering unobserved distributions, precludes direct comparison of our results with existing techniques. Nonetheless, we implemented a natural competing algorithm that we call KMBD (k-means based densities). This method clusters estimated unknown densities by first estimating the unknown proportion as proposed by Patra and Sen (2016), and then applying the inversion formula (7) for densities, which is:

$$f_i(x) = \frac{\ell_i(x) - (1 - p_i)g_i(x)}{p_i}, \qquad (i = 1, \dots, K).$$

More specifically, we consider the density estimators \widehat{f}_i obtained by:

$$\widehat{f}_i(x) = \frac{\widehat{\ell}_i(x) - (1 - \widehat{p}_i)g_i(x)}{\widehat{p}_i}, \qquad (i = 1, \dots, K),$$

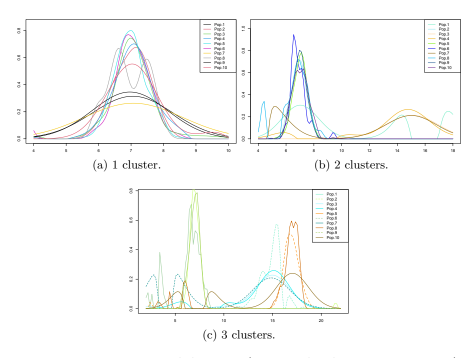


Figure 5: Decontaminated densities (recovered unknown components).

where $\hat{\ell}_i$ is a Gaussian kernel estimator based on the observations, using the Silverman rule (Silverman, 1986) for bandwidth selection, and \hat{p}_i is obtained as per Patra and Sen (2016). As an illustration, Figure 6 shows 10 densities for the case where K=10 with one single cluster, i.e., $f_1=\cdots=f_{10}$ (and $n_i\approx 300,\ i=1,...,K$).

Notably, significant variations among these curves are evident, attributable to the inherent instability of the estimators \hat{p}_i . To mitigate this instability, we employ a smoothing technique, summarizing each density with an M-vector representing mean values across a partition of size M of their support. Thus, we obtain K M-dimensional vectors, each representing a density. We then apply a k-means procedure to these vectors to cluster the densities. The GAP statistic (Tibshirani et al., 2001) is used to determine the optimal number of clusters. The choice of M is arbitrary and partitioning the supports is not straightforward. Due to the high variability illustrated in Figure 6, we experimented with several values of M using uniform partitioning to improve the KMBD algorithm. The results are shown in Figures 7 and 8 for all cases studied in Table 3. For small sample sizes $(n \approx 300)$, we managed to calibrate M to obtain clusters similar to our KCMC algorithm. However, as n increases, the KMBD approach's performance does not improve, and the results are markedly inferior to those obtained by the KCMC algorithm. This phenomenon reveals the competitor's weakness. Specifically, we lack the asymptotic guarantees for the estimators obtained by Patra and Sen (2016), unlike our approach, which leverages the convergence results given in Theorems 5 and 6, performing much better with moderate and large sample sizes. Furthermore, our method is automatic, whereas the proposed KMBD competitor requires a non-trivial partitioning of the density support.

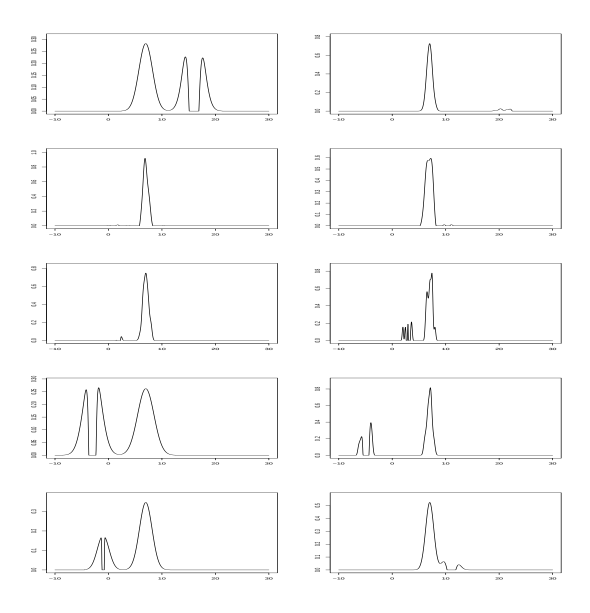


Figure 6: Estimation of densities f_1 to f_{10} (from top left to bottom right, 1 run simulation) in the 1-cluster case, based on Patra and Sen (2016)'s estimator for the p_i 's.

It is worth noting that we also explored k-means clustering based on cumulative distribution function (cdf) estimations. Each estimated cdf was represented by M values computed from a grid spanning their shared support and then subjected to k-means clustering. However, the resulting clustering performance was inferior to that of the KMBD method. Again, the considerable disparity among cdf estimations underscores the challenge of accurately estimating unknown unobserved distributions and choosing a suitable partitioning of the support for both density and cdf in clustering applications.

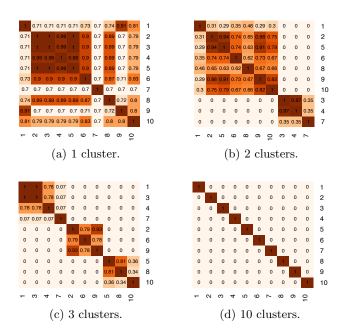


Figure 7: Performance of the KMBD concurrent clustering algorithm, with $n \approx 300$.

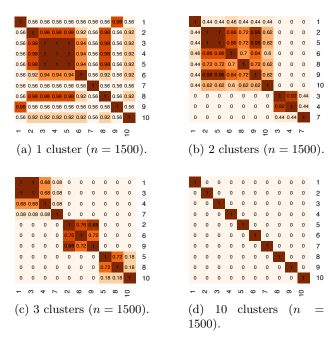


Figure 8: Performance of the KMBD concurrent clustering algorithm, with $n \approx 1500$.

7. Real world Application

7.1 Excess of mortality during pandemics

In Milhaud et al. (2024), a pairwise comparison of COVID-19 excess of mortality has been investigated. This is aimed at identifying countries experiencing similar impact of the COVID-19 pandemic on the mortality. More precisely, the age distribution of death is considered and is supposed to exhibit a common component over periods at the country level. This would correspond to the known component in model (1). The COVID-19 effect component is assumed to be unknown. For instance, the mortality change between 2019 and 2020 is considered as a baseline and serves to assess the excess of the mortality due to COVID-19. The current paper follows the same line as Milhaud et al. (2024) and proposes to cluster countries given the inherent impact of the COVID-19. The datasets of interest came from the Short-Term Mortality Fluctuations (STMF) data series compiled by the Human Mortality Database (HMD). It contains death records aggregated over age groups: 0-14, 15-64, 65-74, 75-85 and 85+. Here, we restrain our study to the four last age classes (given that experts agree to consider that the first one 0-14 was clearly not affected by the pandemic). First, we apply our clustering procedure to the same group of countries as considered in Milhaud et al. (2024) for the first wave. Formally, we study the similarities in terms of the changes for France, Belgium, Germany, Italy, Netherlands and Spain. The known distributions are multinomial ones with four categories here and we compare the unknown multinomial distributions caused by the COVID-19. We report the resulted clusters In Table 5.

Two clusters are the same as those already identified by the authors. Namely, France shows a proper COVID-19 impact on its mortality whereas Italy and the Netherlands share the same profile. However, our clustering methodology identifies a third cluster consisting of Germany, Spain and Belgium. In Milhaud et al. (2024), it is shown that Germany and Belgium, on one hand, and Belgium and Spain on the other have similar impacts but the test rejects the null hypothesis for Germany and Spain. This lack of transitivity of the pairwise testing procedure was already discussed in Milhaud et al. (2024) and the K-sample procedure offers an interesting yet robust generalization of the latter.

7.2 Europe-wide clustering of COVID-19 excess of mortality

In the following, we explore a larger clustering scheme and consider 29 European countries, i.e. Austria, Belgium, Bulgaria, Switzerland, Czech Republic, Germany, Denmark, Spain, Estonia, England and Wales, Finland, France, Greece, Croatia, Hungary, Iceland, Ireland, Italy, Lithuania, Luxembourg, Latvia, Netherlands, Norway, Poland, Portugal, Scotland, Slovakia, Slovenia and Sweden. We aim at exploring the impact of the pandemic over these countries in 2020 and identify the clusters. We adopt the same assumptions as described above and proceed to clustering the countries with regard to their shared COVID-19 excess

Table 5: Clustering of excess of mortality profile over 2020.

	France	Italy	Netherlands	Belgium	Germany	Spain
Cluster (id)	3	2	2	1	1	1

of mortality effect. The known and unknown distributions are multinomials with four categories (the four age classes). The sample sizes are given in the first row of Figure 9 and range from 1,141 for Iceland to 462,577 for Germany. We should recall that this comparison focuses on the distribution of the nodular effect of the COVID-19 rather than its dynamics. Since the countries under study suffered from the impact of the outbreak of the pandemic at different periods during the year of 2020, we are more concerned here with the impact (over classes of age) in the population rather than its magnitude (the mixture proportions p_i depending increasingly on the time of exposure to the pandemic). Hence, the following comparison should be root on the socio-demographic disparities that may exist among the populations as well as the healthcare capacities, public health measures and many other factors. The discussion of the implication of such an impact is, however, beyond the scope of this paper.

First, we set up the level α to 1% and explore the clusters that formed on the basis of the H_0 -rejection rule. At this level, we are left with 11 clusters, see the last row in Figure 9 and the map in Figure 10.

We can observe that some countries are single isolated clusters. This is the case for Spain, Iceland, Switzerland, Netherlands and Portugal. On the other hand, we have two large clusters that represent most countries from center and eastern European countries: Lithuania, Latvia, Poland, Hungry, Bulgaria, Slovakia and Estonia. This block is isolated from the geographically adjacent cluster constituted by the Czech Republic and Croatia. Some of the Northern European countries are gathered on two clusters. The largest is constituted of Finland, Austria, Germany, Northern Ireland, Scotland, Sweden and England & Wales. Surprisingly, a common factor, among other things, is the Protestant inheritance. Numerous studies, e.g. Kaklauskas et al. (2022) among others, validated the similarities between the English-speaking and the Protestant European clusters due to their closely related common histories, cultural interactions, similar development levels, and religions. Another cluster that appears quite clear is the one formed by Lithuania, Latvia, Poland, Hungry, Bulgaria, Slovakia and Estonia. These countries can be found in Figure 11, which illustrates the vaccination delays (or potential delays). This group clearly appears to have experienced a lag in their vaccination campaigns. More precisely, it appears that these countries have the lower percentages of people who received all doses prescribed by the initial

Pop	ulation size	10042	307605	2161	26928	39334	462577	7628	28146	41928	261626	1145	294612	62614	73827	32433
_	15%	SVN	ITA	LUX	FIN	AUT	GER	NIR	SCO	SWE	ENW	ISL	FRA	GRC	NLD	CHE
Level	5%	SVN	ITA	LUX	FIN	AUT	GER	NIR	sco	SWE	ENW	ISL	FRA	GRC	NLD	CHE
_	1%	SVN	ITA	LUX	FIN	AUT	GER	NIR	SCO	SWE	ENW	ISL	FRA	GRC	NLD	CHE
Pop	ulation size	18383	12878	209032	62346	55686	27103	7169	56903	25157	209431	55458	52715	27224	19712	
_	15%	LTU	LVA	POL	HUN	BGR	SVK	EST	CZE	HRV	ESP	PRT	BEL	DNK	NOR	
Level	5%	LTU	LVA	POL	HUN	BGR	SVK	EST	CZE	HRV	ESP	PRT	BEL	DNK	NOR	
	1%	LTU	LVA	POL	HUN	BGR	SVK	EST	CZE	HRV	ESP	PRT	BEL	DNK	NOR	

Figure 9: Clustering of the excess mortality profile (for the 29 countries) due to COVID-19 for different levels of α : 15% (top), 5% (middle) and 1% (bottom).



Figure 10: Clustering of the excess mortality profile due to COVID-19 during the year 2020 over 29 countries.

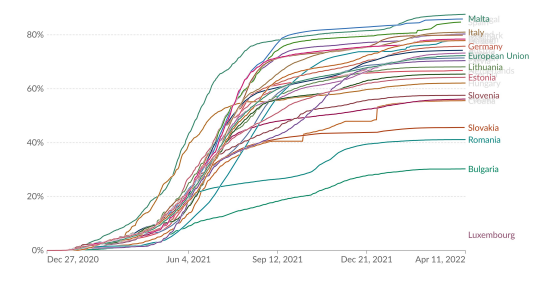


Figure 11: Share of people who received all doses prescribed by the initial COVID-19 vaccination protocol. Source: https://github.com/owid/covid-19-data/tree/master/public/data/vaccinations/locations.csv).

vaccination protocol at the end of 2021. These 7 countries are among the 8 countries with the lowest total vaccination coverage (all doses) reflecting a lack of incentive to vaccinate. Finally, in order to understand more closely the clusters we refer the reader to the plethora of studies that investigate the factors that influence mortality levels from COVID-19 such as well-functioning healthcare system, prevention measures (e.g. social distancing), and population age structure, among others.

Eventually, a way to check the stability of the clusters is to change the threshold of the test acceptance, for example by increasing α to see if the groups merge then. In Figure 9 we reported the clusters for different levels, respectively, 1%, 5% and 15%. As noted earlier, this parameter clearly reflects the threshold for accepting the composition of a group. Indeed, we see that these three levels lead respectively, to 11, 14 and 15 clusters. From Proposition 9 the number of groups determined by our algorithm is asymptotically greater than the true (unknown) number of groups. Since the sample sizes are large here we can conclude that 11 groups is a reasonable choice. If we want to obtain greater detail, a larger value for α will enable a more refined clustering, but may look artificial if too many groups are suddenly created.

Acknowledgments

The authors thank the two anonymous referees for their insightful and constructive comments. This work was conducted within the Research Chair DIALog under the aegis of the Risk Foundation, an initiative by CNP Assurances. D. Pommeret, Y. Salhi and P. Vandekerkhove would also like to acknowledge the support received from the Research Chair ACTIONS under the aegis of the Risk Foundation, an initiative by BNP Paribas Cardif and the French Institute of Actuaries.

Appendix

7.3 Proof of Theorem 5

Let us prove that $\mathbb{P}(S(n) \geq 2)$ vanishes as $n \to +\infty$. By definition of S(n) we have

$$\mathbb{P}\left(S(n) \geq 2\right) = \mathbb{P}\left(\text{it exists } 2 \leq r \leq d(k): \ U_r - r\ell_n \geq U_1 - \ell_n\right)$$

$$= \mathbb{P}\left(\text{it exists } 2 \leq r \leq d(k): \ U_r - U_1 \geq (r-1)\ell_n\right)$$

$$= \mathbb{P}\left(\text{it exists } 2 \leq r \leq d(k): \sum_{(i,j) \in S(k): \ 2 \leq r_k(i,j) \leq r} T_{i,j} \geq (r-1)\ell_n\right)$$

$$\leq \mathbb{P}\left(\text{it exists } (i,j) \text{ with } 2 \leq r_k(i,j) \leq d(k): \ T_{i,j} \geq \ell_n\right)$$

$$\leq \sum_{2 \leq r_k(i,j) \leq d(k)} \mathbb{P}\left(T_{i,j} \geq \ell_n\right),$$

where the penultimate inequality arises from the fact that if the sum of (r-1) positive terms is greater than $(r-1)\ell_n$ then at least one of the terms is greater than ℓ_n . From Lemma 3 we know that under H_0 , for all $\varepsilon > 0$, $n^{-\varepsilon}T_{i,j} = n^{-\varepsilon}nd_n[i,j](\widehat{\theta}_{ij}^{(1)},\widehat{\theta}_{ij}^{(2)})$ that goes to 0 in

probability, as $n \to +\infty$. Since d(k) = k(k-1)/2 is fixed we then obtain $\mathbb{P}(S(n) \ge 2) \to 0$ as $n \to +\infty$, which proves the wanted result.

7.4 Proof of Theorem 6

From Theorem 5 we have $\mathbb{P}(S(n)=1) \to 1$ as $n \to +\infty$, from which we can deduce that for all $\xi > 0$

$$\mathbb{P}(|U_{S(n)} - U_1| \ge \xi) = \mathbb{P}(|U_{S(n)} - U_1| \ge \xi \cap \{S(n) = 1\}) + \mathbb{P}(|U_{S(n)} - U_1| \ge \xi \cap \{S(n) > 1\})$$

$$= \mathbb{P}(|U_{S(n)} - U_1| \ge \xi \cap \{S(n) > 1\})$$

$$< \mathbb{P}(S(n) > 1) \to 0,$$

which implies that $U_{S(n)}$ has the same limiting distribution as $U_1 = T_{1,2}$, see Lemma 3.

7.5 Proof of Theorem 7

We want to prove that $\mathbb{P}(S(n) = r) = \mathbb{P}(S(n) \ge r) - \mathbb{P}(S(n) > r)$ tends to 1. Consider the general case $H_1(r)$ with r > 1, the particular case $H_1(1)$ being similar. We first show that $\mathbb{P}(S(n) \ge r)$ tends to 1 as $n \to +\infty$. Under $H_1(r)$, we have for all r' < r

$$\mathbb{P}\left(U_r - r\ell_n \ge U_{r'} - r'\ell_n\right) = \mathbb{P}\left(\left(U_r - U_{r'}\right) \ge (r - r')\ell_n\right) \\
= \mathbb{P}\left(\sum_{r' < r_k(i,j) \le r} T_{i,j} \ge (r - r')\ell_n\right) \\
\ge \mathbb{P}\left(T_{i,j}\mathbb{I}_{\{r_k(i,j) = r\}} \ge (r - r')\ell_n\right).$$

When $r_k(i,j) = r$, under $H_1(r)$ we have from Lemma 3 $T_{i,j} = U_n^1(i,j) + V_n^1(i,j)$ where $V_n^1(i,j) = \lambda[i,j] \times n + o_{a.s.}(n)$. From **(B)** we know that $\ell_n = n^{\varepsilon}$ with $\varepsilon < 1$, and we deduce that $\mathbb{P}\left(T_{i,j}\mathbb{I}_{\{r_k(i,j)=r\}} \geq (r-r')\ell_n\right) \to 1$, as n tends to infinity, which proves that $\mathbb{P}(S(n) \geq r)$ tends to 1.

We now prove that $\mathbb{P}(S(n) > r)$ tends to 0. For $r_k(i,j) = r$, since $U_r \geq T_{i,j}$, from the above we have $\mathbb{P}\Big(U_r \geq (r-r')\ell_n\Big) \to 1$, as n tends to infinity, which implies that for all $\xi > 0$, $\mathbb{P}(U_r - r\ell_n > \xi)$ tends to 1 as n tends to infinity. It implies that $\mathbb{P}(r \in \arg\max_{1 \leq s \leq d(k)} \{U_s - s\ell_n\}) \to 1$ which implies that $\mathbb{P}(S(n) > r)$ tends to 0.

7.6 Proof of Proposition 9

Let S be a given cluster and consider a wrong candidate population x, that is, x does not belong to S in reality. Then we want to test H_0 : " $\{x\} \cup S$ forms a new cluster". Let \widetilde{U}_n be the associated statistic to such a null hypothesis. From Theorem 7, since \widetilde{U}_n tends to infinity, for all $(1-\alpha)$ -quantile $\widehat{q}_{1-\alpha} > 0$ for a level α fixed, there exists n large enough such that $\mathbb{P}(\widetilde{U}_n < \widehat{q}_{1-\alpha}) < \alpha$ and then H_0 is rejected.

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