EXPLORATIVE DATA STRUCTURE COMPARISON WITH APPLICATION TO ...

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Abstract

abstract...

Key words: keywords...

1. Introduction

Classical statistical methodology is aimed at analyzing data from designed experiments and historically statistical analyses have been done by researchers who knew the design and origin story of the data set well. The origin stories of data sets have changed over time and today a lot of data is accumulated without specific purpose. This is due to vast amounts of data being registered online and to a trend towards more open source research. The latter phenomenon in particular poses new challenges wrt. data quality assessment. When data are collected and made public without a specific end-point in mind, how do we ensure that differences in, say, choice measurement instruments, mode of administration, or sampling frame do not cause the data to be effectively divided into subsets that are simply not comparable?

Surveys often use mixed modes of administration, e.g. mail and telephone, and while this can improve response rates, the mode of administration can affect results Brambilla and McKinlay (1987); McHorney et al. (1994) and differences in response behavior can lead to biased results. Powers, Mishra and Young (2005) report effects of mode of administration on changes in mental health scores that are of a magnitude that is considered to be clinically meaningful.

The rapid growth of web surveys, due to low cost, timeliness, and other factors, generate large data sources that lack a sampling frame of the general population. However, it can be problematic to combine online panels (pre-recruited profiled pools of respondents) with intercept samples (a pool of respondents obtained through banners, ads, or promotions) Liu (2016).

Sophisticated methods for addressing this question are available when we are willing to assume a statistical model, but when these models are taken away, a remarkable void of methods is left behind. What is needed is a procedure that compares differences in overall data structures in two (or more) subsets of a dataset without assuming neither directional nor hierarchical relationships between the variables. We propose a new method for this task, namely Principal Component Analysis-based Data Structure Comparison (PCADSC). This method employs the principal component decomposition of the data matrix performed on two subsets of a dataset in order to create intuitive visualizations of data structure differences. Mention R package.

This manuscript is structured as follows: First, in Section 2, we present the data structure comparison problem in more detail and discuss what statistical methods are already available for solving similar challenges. Next, in Section 3, we move on to a description of the PCADSC procedure, including a brief introduction to principal component analysis (PCA) in general. In Section 4, we present a worked data example using the open source, online available PISA data (ref), which is an example of a dataset where multiple data collection methods Eller mske lande? have been employed.

2. Something about state of the art

2.1. More detailed description of the type of problem we wish to address

- Two subsets of a dataset, i.e. to datasets with the same variables, but different observations
- Wish to compare structures without specifying a model or even any variables of interest

- The most central example is the question of whether the two subsets can readily be combined in a (unknown) data analysis, or if the subset-inducing variable actually implies heterogeneity across the subset division
 - Examples: Large scale open source datasets such as the PISA data and ESS (European Social Survey) data and ...(?). In these datasets, the data producers are very far away from the majority of the data analysts. Therefore, problem-specific recommendations about potential instrument-induced challenges in the datasets are not available for the data analysts. How can data producers ensure that this will not be an issue, at least not related to known data gathering differences?
 - Other examples?
 - Perhaps a description of what happens if we are to combine the two subsets of the datasets without taking a e.g. an instrument-effect into account. When will it cause problems (maybe: causal graph style?)?
 - Mention somewhere: We want a solution that is largely independent of the sizes of the two subsets of data. Thereby, a lot of methods that compare each subset to the full dataset in some sense are excluded.
 - 2.2. Describe existing methods used to solve similar questions or parts of the question we are addressing
- The simplest case: variable-by-variable tests in distributional differences
 - Simple, but scales poorly
 - Only relates to marginal differences and not to the interplay between variables
- Karl's papers?
- Anne's papers: IRT-based methods for surveys
 - Moves beyond the marginal approach, but needs a model pre-specified
 - Thus, it is not a general data structure comparison method, but rather a fitted-model comparison method. It addresses the interplay between the model and the data, not the data alone. This is fundamentally a different (though related) question.

3. PCADSC - description of the method

Description from Anne's master's thesis. Rewrite.

As mentioned above, the purpose of PCADSC is comparing overall data structures in two or more subsets of a dataset. But before we can get further into describing this procedure, we must first define what exactly is meant by "overall structure". One such definition is the structure of the covariance matrix of the dataset. If we assume all variables in the dataset to be jointly normal with zero means, the covariance matrix is a sufficient statistic for describing the simultaneous distribution of all the variables. This gives it a very nice interpretation as a measure of the overall structure. If we do not accept the normality assumption, pairwise correlations and variable variances are still interesting quantities that say something about the interrelations between the variables, at least in terms of linear relationships. All in all, the empirical covariance matrix is a reasonable place to start looking for differences in "overall data structures".

Though the idea sounds appealing, it is quite difficult to assess similarity of matrices, and moreover, this becomes increasingly difficult for large numbers of variables and thus high dimensional covariance matrices. There is simply too much information to consider at once.

However, by use of linear algebra, we can construct a decomposition of the covariance matrix that makes it easier to gain an overview of the data. We propose a new method based on principal component analysis that seems to be able to identify differences in datasets based on intuitive, visual inspections. We refer to this method as principal component analysis-based data structure comparison (PCADSC) and we present the procedure below. But first, we give a minimal introduction to principal component analysis in general with reference to Koch (2014).

3.1. Principal component analysis

Consider n observations $x_1, \ldots, x_n \in \mathbb{R}^d$ of d variables, let $\bar{x} = \frac{1}{n} \sum_{i=1}^n x_n$ denote their average(s?) and let $S = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})(x_i - \bar{x})^\top \in \mathbb{R}^{d \times d}$ denote the empirical covariance Is $\bar{x} \in \mathbb{R}^d$ with $\bar{x} = (\bar{x}_1, \ldots, \bar{x}_d)^T$?. Suppose that we want to describe the observations by q numbers instead of the original d numbers. The associated rank-q-reconstruction error is defined as the minimal squared error that is achievable by linear subspaces $K_q \subset \mathbb{R}^d$ of dimension q < d, that is

$$\min_{K_q} \sum_{i=1}^n \min_{z \in K_q} ||x_i - \bar{x} - z||^2 = \min_{K_q} \sum_{i=1}^n ||x_i - \bar{x} - \operatorname{proj}_{K_q}(x_i - \bar{x})||^2.$$

Principal component analysis (PCA) ensures the existence of a subspace $\hat{K}_q \subset \mathbb{R}^d$ that attains this minimum, and it provides an explicit description of \hat{K}_q and the rank-q-reconstruction error. Thus, let $S = U\Lambda U^{\top}$ be the eigenvalue decomposition of S. Here $\Lambda \in \mathbb{R}^d$ is the diagonal matrix with the eigenvectors $\lambda_1 \geq \cdots \geq \lambda_d \geq 0$ in the diagonal, and $U \in \mathbb{R}^d$ is the orthogonal matrix with the associated eigenvectors $\eta_1, \ldots, \eta_d \in \mathbb{R}^d$ in the columns. The eigenvalues are uniquely defined, and the eigenvectors are uniquely defined up to a change of sign whenever the eigenvalues are different. If some of the eigenvalues are identical, e.g. $\lambda_i = \lambda_{i+1} = \cdots = \lambda_j$, then the associated eigenvectors $\eta_i, \eta_{i+1}, \ldots, \eta_j$ are uniquely defined up to a common rotation. In practice this only happens if n < d, in which case the last d - n eigenvalues will be zero Anne: What

about covariance matrices that are not of full rank (e.g. because of colinearity)? Won't we find eigenvalue multiplicities > 1 here as well?. It is a result from linear algebra that the rank-q-reconstruction error for q < d is achieved for

$$\hat{K}_q = \operatorname{span}\{\eta_1, \dots, \eta_q\}$$

and equals $\sum_{j=q+1}^{d} \lambda_{j}$. The eigenvectors $\eta_{j} \in \mathbb{R}^{d}$ are called *loadings*, and the eigenvalues $\lambda_{j} \geq 0$ may be understood as *variation components*. The projections $\eta_{j}^{\top}(x_{i} - \bar{x})$ of the observations onto the loadings are called *scores*. The *j*th loading can also be found iteratively as the unit vector $u \in \mathbb{R}^{d}$ orthogonal to \hat{K}_{j-1} , where the initial subspace is defined as $\hat{K}_{0} = \{0\}$, that maximizes the variation of the associated scores:

$$\eta_j = \operatorname{argmax}_{u \in \mathbb{R}^d : u \perp \hat{K}_{j-1}} \sum_{i=1}^n \|u^\top (x_i - \bar{x})\|^2, \qquad \lambda_j = \frac{1}{n-1} \sum_{i=1}^n \|\eta_j^\top (x_i - \bar{x})\|^2.$$

Anne: Should we maybe assume standardized xs here also? I feel like it is part of the PCA algorithm and subtracting the mean in the above seems off for that reason. It is worth emphasizing that the greedy approach of successively adding the next direction η_j explaining most of the remaining variation, also gives the sequence $\hat{K}_q = \hat{K}_{q-1} \oplus \text{span}\{\eta_q\}$ of subspaces minimizing the rank-q-reconstruction error. This strong interpretation of PCA, which is often overlooked in the literature, means that the sequence of loadings η_j and their associated variation components λ_j yield a simultaneous description of the structure of the data set for all approximating dimensions q. This implies that the loadings and variation components can be used to investigate the structure of the data set without the need to decide on an approximating dimension q.

3.2. PCA-based data structure comparisons

Above, we promised a method for intuitive, visual inspection of data structure similarities, but as of now, all intuition might have been lost in technicalities. The main point we want to emphasize from PCA is that whereas the scores describe the observations, the variation components and the accompanying loadings describe the usage of the variables. If two different datasets with the same variables, but different samples of observations, have similar loading patterns, then the variables appear to be measuring the same underlying quantities in both data situations. This can be the case while the two sets of scores could be arbitrarily different, which e.g. could happen if the two datasets were taken from two different populations of subjects. On the other hand, if the loading patterns are different in the two datasets, then this indicates that the variables are used differently in the two data situations, and hence it would be criticizable to use these variables for comparisons across the two datasets.

In this paper we propose three diagnostic plots for comparing the loading patterns in two datasets $x_{11}, \ldots, x_{1n_1} \in \mathbb{R}^d$ and $x_{21}, \ldots, x_{2n_2} \in \mathbb{R}^d$ in the same d variables. The construction of these plots, which we call CumEigenPlot, HairPlot and PancakePlot, proceeds in the following steps:

1. Standardized the observations x_{i1}, \ldots, x_{in_i} to have zero mean and unit standard deviation separately for each of the d variables within each datasets. Let $\tilde{x}_{ij} \in \mathbb{R}^d$ for i = 1, 2 and

 $j=1,\ldots,n_i$ be the standardized observations, and form the principal component analysis

$$S_i = \frac{1}{n_i - 1} \sum_{i=1}^{n_i} \tilde{x}_{ij} \tilde{x}_{ij}^{\top} = \sum_{k=1}^{d} \lambda_{ik} \eta_{ik} \eta_{ik}^{\top}$$
 for $i = 1, 2$.

2. Combine the standardized datasets into a joint dataset $\tilde{x}_1, \ldots, \tilde{x}_n \in \mathbb{R}^d$ with $n = n_1 + n_2$ observations, standardize this dataset, and form the principal component analysis

$$S = \frac{1}{n-2} \sum_{i=1}^{n} \tilde{x}_j \tilde{x}_j^{\top} = \sum_{k=1}^{d} \lambda_k \eta_k \eta_k^{\top}.$$

- 3. Make the diagnostic plot of interest. Details of the plots will be described below.
- 4. To investigate the sample variation randomly reallocate the combined and standardized dataset formed in step 2 to two datasets of n_1 and n_2 observations, respectively, and redo step 1 and 3. This is done several times, e.g. 10000 times, and compared to the plot for the original allocation.

Before describing the details of the CumEigenPlot, HairPlot and PancakePlot we make some general remarks. First, it should be noted that PCA is sensitive to scaling, as the procedure deconstructs the covariance matrix in components according to the most explained variance. This implies that if a variable has a very large sample variance (possibly because of its scale), this variable will be always be deemed highly influential, no matter the structure of the data. Therefore, the variables should always be scaled prior to performing PCA. Note that the covariance matrix for the standardized variables is the same as the correlation matrix for the original variables, so this simply corresponds to performing data structural comparisons of the correlation matrices rather than the covariance matrices. Do we still need more about this topic? Then we can add something like:

• Following the greedy interpretation of PCA, we know that the first eigenvector is chosen as

$$\eta_1 = \operatorname{argmax}_{u \in \mathbb{R}^d} \sum_{i=1}^n ||u^{\top}(x_i - \bar{x})||^2 = \operatorname{argmax}_{u \in \mathbb{R}^d} V(u^T X^T)$$

- Now, imagine that $V(x_1) >> V(x_j)$ for all $j \in 2, ..., d$. Note that $(1, 0, ..., 0)X^T = x_i$. Then, clearly a vector u that only picks out the first variable from the data matrix X, namely $u^T = (1, 0, ..., 0)$ will allow us to only look at the variance of x_1 , which is very large. This has nothing to do with the interplay between variables, as the phenomenon will disappear if we scale x_1
- A more formal explanation could come from this kind of intuition, if needed.

The standardization makes the variables comparable on the same scale, i.e. units of standard deviation, and it implies that the diagonal elements of S, S_1 , and S_2 all equals 1. This also holds

for the reallocated datasets, and in particular we have $\sum_{k=1}^{d} \lambda_k = \sum_{k=1}^{d} \lambda_{1k} = \sum_{k=1}^{d} \lambda_{2k} = d$. The random reallocations in step 4 provide resamples under the null hypothesis that the two sample populations have the same structure in the d variables, except that the two populations are allowed to have the own standard deviations for each of the variables. This exception is induced by the initial step 1, which performs a standardization within the two datasets before they are combined in step 2. These resamples can be used for a visualization of the sample variation, and if a test statistic can be defined in step 3, then a p-value for the null hypothesis may be computed. Such p-values are know as permutation tests for Goodness-of-Fit (reference?).

To do: Something about what to do if not all variables are numerical. Anne: Maybe leave this for the discussion?

3.2.1. Description of the CumEigenPlot

This plot compares the variation components. In order to investigate whether the same proportion of the total variation can be described by the same number of principal components in the two datasets we plot a piecewise linear curve connecting the points

$$(0,0), (\lambda_1, \lambda_{11} - \lambda_{12}), (\lambda_1 + \lambda_2, \lambda_{11} + \lambda_{12} - \lambda_{21} - \lambda_{22}), \ldots, \left(\sum_{j=1}^d \lambda_j, \sum_{j=1}^d \lambda_{1j} - \sum_{j=1}^d \lambda_{2j}\right).$$

This may be seen as a cumulated Bland-Altman plot for the variation components (reference to cumulated residuals and to Bland-Altman). Note that due to the standardization the last point will be equal to (d,0). Thus, this curve will begin and end at the x-axis. And the larger excursions it makes away from the x-axis the less alike the cumulated variation components for the two datasets are. Is this an informative x-axis? Should we maybe consider scaling it?

In step 3 we have implemented both the Kolmogorov-Smirnov and the $Cram\acute{e}r$ -von Mises test statistics given by

$$KS = \max_{k=1,\dots,d} \left| \sum_{j=1}^{k} \lambda_{1j} - \sum_{j=1}^{k} \lambda_{2j} \right|, \qquad CvM = \sum_{k=1}^{d-1} \frac{\lambda_k + \lambda_{k+1}}{2} \left(\sum_{j=1}^{k} \lambda_{1j} - \sum_{j=1}^{k} \lambda_{2j} \right)^2.$$

Maybe more here on what the null hypothesis of these tests is? From the random reallocations in step 4 we get p-values for the Kolmogorov-Smirnov and the Cramér-von Mises Goodness-of-Fit tests. Step 4 is also used to visualize the sampling variation in a display, where we plot the observed curve together with 20 of the resampled curves as well as a shaded region visualizing pointwise 95 % coverage intervals. If the observed curve is very different from the resampled curves or if it is substantially outside the shaded region, then this also indicates differences between the two datasets.

3.2.2. Description of the HairPlot

This plot simultaneously compares the variation components and the loadings. Let $\lambda_{\text{max}} = \max\{\lambda_{11}, \lambda_{21}\}$ be the largest variation component for the two datasets. The empirical

correlation matrix S_1 for the first dataset has the following orthogonal decomposition in the coordinate system of the second dataset

$$S_1 = \sum_{k=1}^d \lambda_{1k} \eta_{1k} \eta_{1k}^\top = \lambda_{\max} \sum_{k=1}^d \left(\sum_{j=1}^d \sqrt{\frac{\lambda_{1k}}{\lambda_{\max}}} (\eta_{1k} \eta_{2j}^\top) \eta_{2j} \right) \left(\sum_{j=1}^d \sqrt{\frac{\lambda_{1k}}{\lambda_{\max}}} (\eta_{1k} \eta_{2j}^\top) \eta_{2j} \right)^\top,$$

and we have a similar decomposition of S_2 in the coordinate system of the first dataset. We propose to visualize these two decompositions in a $d \times d$ grid display. In the jth row and kth column of this display we plot two arrows based at the lower left corner of the grid cell. The first arrow has length μ_{jk} and angle $\theta_{jk}/2$ anticlockwise from the diagonal, and the second arrow has length ν_{jk} and angle $\theta_{jk}/2$ clockwise from the diagonal. To facilitate the following description we will refer to the arrows drawn anticlockwise as the blue arrows, and the arrows drawn clockwise as the red arrows. The lengths μ_{jk} and ν_{jk} and the angle θ_{kj} are given by

$$\mu_{jk} = \sqrt{\frac{\lambda_{1k}}{\lambda_{\max}}} |\eta_{1k} \eta_{2j}^{\top}|, \qquad \qquad \nu_{jk} = \sqrt{\frac{\lambda_{2j}}{\lambda_{\max}}} |\eta_{2j} \eta_{1k}^{\top}|, \qquad \qquad \theta_{jk} = \arccos(|\eta_{1k} \eta_{2j}^{\top}|).$$

The absolute value of the projection $\eta_{1k}\eta_{2j}^{\top}$ is inserted due to the indeterminacy of the direction of loading vectors. This indeterminacy implies that the angle between loadings from the two datasets always can be chosen to be in the interval $[0, \pi/2]$, and hence the decomposition of S_1 and S_2 can be visualized in a joint plot by dividing the angles by two and using anticlockwise and clockwise shifts from the diagonal. Furthermore, the scaling of the lengths by λ_{max} is made so that the longest arrow has at most unit length.

In the *HairPlot* the blue arrows in the *k*th column of the grid display visualize the decomposition of the *k*th principal components for the first dataset in the coordinate system of the second dataset. Similarly, the red arrows in the *j*th row visualize the decomposition of the *j*th principal components for the second dataset in the coordinate system of the first dataset. If the structures of the two datasets are identical, then we will have coinciding blue and red arrows along the diagonal in the grid display, and nothing else as arrows in the off-diagonal cells would have zero length. Differences in the variation components are visualized as differences in the lengths of the blue and the red arrows, also in the diagonal. And loadings in other directions than the corresponding loading from the other dataset are visualized as angle separation of the blue and the red arrows in the diagonal cells as well as arrows of non vanishing length in the off-diagonal cells. The name *HairPlot* arise by seeing the arrows as *hairs*. Using this interpretation we may say that a more "hairy" plot indicates less agreement between the two loading patterns.

Along with the visual comparison we have also implemented numerical tests in step 3. Separately for the off-diagonal grid cells we may use the lengths μ_{jk} and ν_{jk} as test statistics, and for the on-diagonal grid cells we use the distance between the scaled loadings as test statistics, that is

$$\chi_k = \lambda_{\text{max}}^{-1} |\lambda_{1k} \eta_{1k} - \lambda_{2k} \eta_{2k}|, \text{ for } k = 1, \dots, d.$$

In total this gives $2d^2 - d$ tests, which should be corrected for the family wise type I error using a methods that doesn't assume independence among the tests. We also construct overall

Kolmogorov-Smirnov and Cramér-von Mises type test statistics by

$$KS_{1} = \max_{j,k=1,\dots,d} (1_{j \neq k} \mu_{jk} + 1_{j=k} \chi_{k}), \qquad CvM_{1} = \sum_{j=1}^{d} \sum_{k=1}^{d} (1_{j \neq k} \mu_{jk} + 1_{j=k} \chi_{k})^{2},$$

$$KS_{2} = \max_{j,k=1,\dots,d} (1_{j \neq k} \nu_{jk} + 1_{j=k} \chi_{k}), \qquad CvM_{2} = \sum_{j=1}^{d} \sum_{k=1}^{d} (1_{j \neq k} \nu_{jk} + 1_{j=k} \chi_{k})^{2}.$$

3.2.3. Description of the PancakePlot

This plot compares the loadings. The PancakePlot consists of two parallel vertical barplots, one for each of the datasets, with one bar for each of the d loading vector $\eta_{ij} \in \mathbb{R}^d$. In the bars the relative contributions of the d variables are visualized by separate colors for the different variables. Due to the indeterminacy of the sign all the signs are removed from the coefficients in the loadings, and the relative contribution is defined as the resulting coefficients divided by their total sum. The bars are ordered according to the variation components, and we also insert the cumulative variation components next to the barplots. The plots resulting from this procedure should be inspected focusing on two properties: Similarities in loading patterns, which will correspond to similar visual impressions, and similarities in variance contributions. Refer to example/show plot.

We not implemented a test statistic for the *PancakePlot*, but in order to visualize the sample variation the observed plot can be shown together with resampled plots generated in step 4.

4. Data example stuff

We will now turn to a concrete data example in order to illustrate the possibilities of each of the methods presented above more than just PCADSC?. We use data from the 2012 version of the European Social Survey (ESS) project, a very large dataset that is freely available online at www.europeansocialsurvey.org. As the name suggests, the data comes from a survey that was conducted primarily in Europe aiming to collect information about the social conditions of the citizens reference?. As with all international (or, simply, multi-center) studies, one might be concerned about whether or not the data from different countries can readily be combined. This is the question we will address using PCADSC in the current section.

All computations and figures presented in this section where created using our R package PCADSC, which is available online at www.github.com/AnnePetersen1/PCADSC maybe do a CRAN submission instead?.

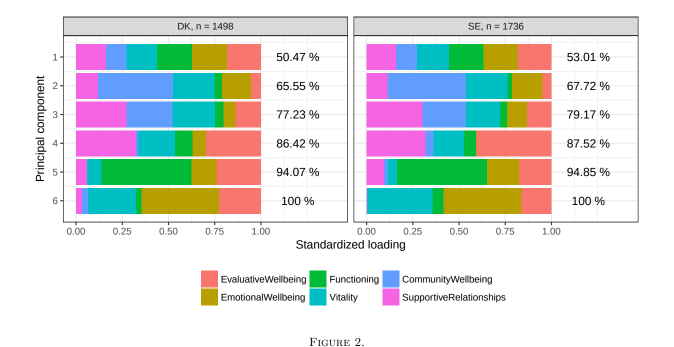
The ESS 2012 data contains a total of 626 variables collected from 54673 citizens in 29 countries. Here, we will only work with a subset of 35 questionnaire items that are all related to psychological well-being. These 35 items can be divided into 6 distinct scales (REF: ESS6 Topline Results Series 5), as illustrated in Figure 1. We represent each of these scales by a single variable, which is calculated as the average score within the items related to that variable (ref to someone

Table 1: Items from the wellbeing module grouped by the dimension of wellbeing they relate to

WELLBEING DIMENSION	ESS SURVEY ITEM
Evaluative wellbeing	How satisfied with life as a whole
	How happy are you
Emotional wellbeing	Felt sad, how often past week
	Felt depressed, how often past week
	Enjoyed life, how often past week
	Were happy, how often past week
	You felt anxious, how often past week
	You felt calm and peaceful, how often past week
Functioning	Free to decide how to live my life
	Little chance to show how capable I am
	Feel accomplishment from what I do
	Interested in what you are doing
	Absorbed in what you are doing
	Enthusiastic about what you are doing
	Feel what I do in life is valuable and worthwhile
	Have a sense of direction
	Always optimistic about my future
	There are lots of things I feel I am good at
	In general feel very positive about myself
Functioning	At times feel as if I am a failure
	When things go wrong in my life it takes a long time to get back to normal
	Deal with important problems
Vitality	Felt everything did an effort, how often past week
	Sleep was restless, how often past week
	Could not get going, how often past week
	Had lot of energy, how often past week
Community wellbeing	Most people can be trusted / can't be too careful
	People try to take advantage
	Most of the time people are helpful
	Feel people in local area help one another
	Feel close to the people in local area
Supportive relationships	How many with whom you can discuss intimate matters
	Feel appreciated by those you are close to
	Receive help and support
	Felt lonely, how often past week

Source: European Social Survey Round 6, 2012

FIGURE 1. Make a new table here



that says that is sensible?). For simplicity, we use only complete cases for this construction and thus exclude all participants that did not answer all the 35 questionnaire items used below.

something

In the following, we only compare two countries, namely Denmark and Bulgaria, which have $n_{BG} = 1798$ and $n_{DK} = 1498$ complete cases in the variables of interest, respectively. For these two countries, the ESS authors different term? particularly highlight differences in the relationship between the psychological well-being scales, at least on an nation-aggregated level (REF: ESS6 Topline Results Series 5). This might be the result of large, cultural and socio-economic differences between the two countries. Simply put, we do not expect happiness and psychological well-being to be the same phenomena in Bulgaria and Denmark more here about why, about the cultures?. Therefore, a successful method for data comparisons should be able to detect these differences by looking at the differences in the interplay between the 6 scales of psychological well-being.

4.2. PCADSC

Figure 2 illustrates the results of conducting PCADSC on the psychological well-being data from the ESS. While the first principal component, which is responsible for explaining 50-60 % of the variance in the data, is very similar for the two countries, we see quite large differences in the remaining components. In the second component, we see that the two countries disagree in the relative importance of the scales *Community wellbeing* and *Vitality*. In the third and fourth components, general disagreement is found. All in all, components 2-4, representing almost half of the variability in the data, are not very similar across the two countries. Moreover, the two

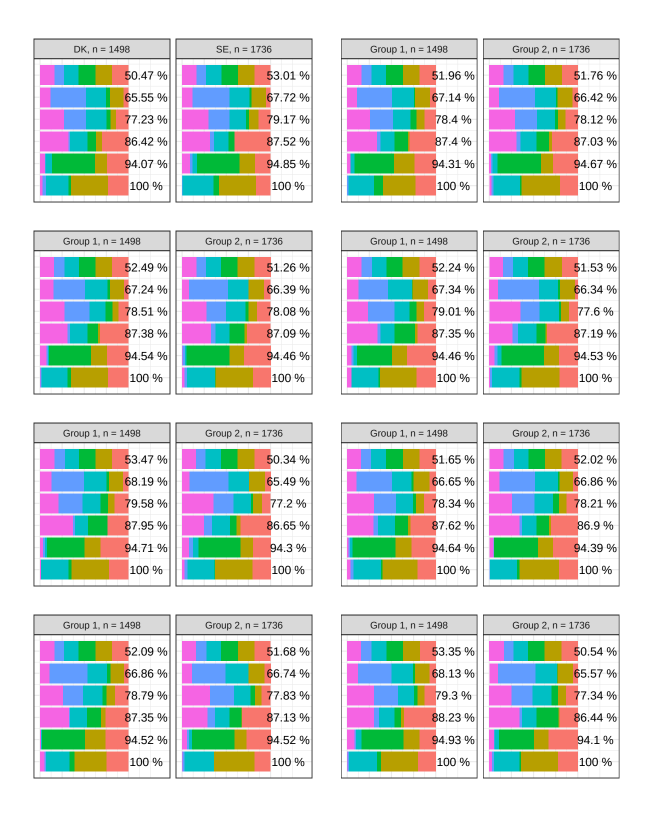


FIGURE 3. something.

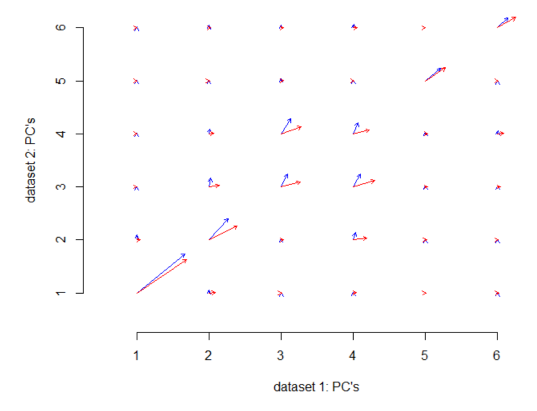


FIGURE 4.

The blue arrows show the principal components of the first dataset decomposed in the coordinate system of the principal components of the second dataset. And the red arrows the reverse.

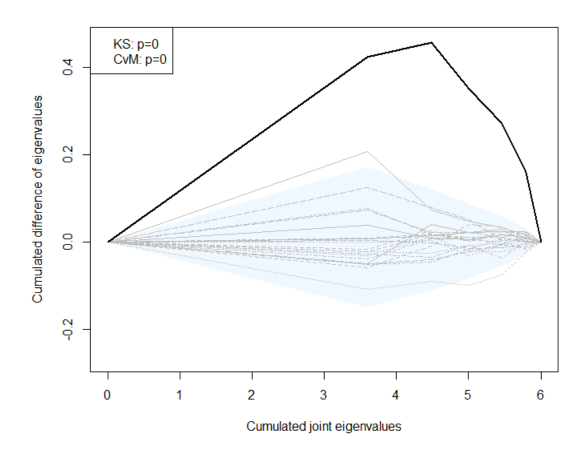


FIGURE 5. something.

subsets of the data also differ with respect to how much variance is explained by each component, and the difference is particularly big for the first component. This component has approximately 15 % more explanatory power in the Bulgarian subsample than it does in the Danish.

Figure 4 shows the hairplot. The blue arrows visualize the decomposition of the principal components for the first dataset in the coordinate system of the second dataset. We see that PC2 also loads on PC3, that PC3 also loads on PC4, and that PC4 also loads on PC2 and PC3. The red arrows visualize the decomposition of the principal components for the second dataset in the coordinate system of the first dataset. We see that PC2 also loads on PC4, that PC3 also loads on PC2 and PC4, and that PC4 also loads on PC3. For PC1, PC5 and PC6 the main difference is in the size of the variation component.

In summary, we find that the datasets are fundamentally different and that we should therefore be wary about combining them in a joint analysis, which was also the conclusion of the ESS authors, though based on country-level aggregated statistics.

But did we really illustrate a data structure difference due to country differences or did we just illustrate the variability of the results of the PCADSC method? In order to investigate this further, we look at a so-called Wally plot (ref: Claus Ekstrm). In this plot, we compare the results of PCADSC conducted with grouping by country with several random, but similar grouping variables. Specifically, we produce 7 PCADSC plots where the country variable was replaced by a randomly generated variable that divides the observations into two groups of the same sizes as the country samples. The results are illustrated in Figure 3. Here, we see that the differences in the second component from the original PCADSC results are not matched in any of the randomly grouped PCADSC runs. In fact, the 7 runs are remarkably similar, thereby illustrating that PCADSC seems to be very robust with respect to random groupings: The signal in the data is not blurred by the random subdivisions. When it comes to the differences in the third component for the two groups, we find much larger variability in the 7 random runs. more comments here... Wait until we are sure exactly what we think about the results and what other PCA-based methods, we will do before/after. Particularly, how do we deal with eigen value differences?

5. Discussion

- Generalizing the results to non-numeric variables?
- Generalizing the results to covariance matrices that are not of full rank?
- . 1

6. Concluding Remarks

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