



## ggESDA: An R Package for Exploratory Symbolic Data Analysis using ggplot2

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### Abstract

Exploratory data analysis (EDA) serves as a preliminary yet essential tool for summarizing the main characteristics of a data set before appropriate statistical modeling can be applied. Quite often, EDA employs the traditional graphical techniques such as the boxplot, histogram and scatterplot and are equipped with various dimension reduction methods and computer-aided interactive functionalities. EDA has been used to explore different data types. Examples were the cases of the survival data, the time series data, the functional data and the longitudinal data. Conventionally, these data set were tabulated by a table with  $p$  columns corresponding to  $p$  variables. Each subject is measured by a single numerical value for each variable. Nowadays the collected data keeps getting bigger and complex. The description of data was no longer stored by a form of a single value but the intervals, histograms and/or distributions. These are examples of the so-called symbolic data. This study develops an R package, namely **ggESDA**, as an extension of **ggplot2** package for exploring symbolic data through a wide variety of plots. SDA supplies various data descriptions and has great capacity for large and complex data. The **ggESDA** acts as an advanced graphical tool that supports the efficient, effective and practical exploration of symbolic data sets.

*Keywords:* data visualization, exploratory data analysis, interval-valued data, statistical graphics, symbolic data analysis.

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## 1. Introduction

For the conventional data analysis, the data set were usually presented by a classical data table where  $p$  random variables are represented by a single point in  $p$ -dimensional space. In contrast, symbolic data with measurements on  $p$  random variables are  $p$ -dimensional statistical units such as hypercubes or histograms in  $\mathbb{R}^p$ , or a Cartesian product of  $p$  distributions. The symbolic data analysis (SDA) was proposed by Billard and Diday (2003, 2007). The aim is

to provide an overview of the statistical methodologies for analyzing such data. Symbolic variables make possible to describe groups of individuals and concepts. In practice, the symbolic data is generated by aggregating the large or enormous datasets into intervals or histograms. The aggregation can produce a dataset of more manageable size in order to conduct appropriate analyses; or it can originate to some scientific questions of interest. These are very important characteristics of SDA so that it could be served as a potential solution for big data problem. The researches for interval-valued data such as the sign test for COVID-19 data ?, the prediction via regularized artificial neural network ?, a bivariate Bayesian method for regression models ?, etc. Figure ?? shows the number of the SDA related researches including clustering, classification, forecasting and so on is increasing among year 1998 to 2020 based on several database, which outstands the importance of it during the years. Among ScienceDirect, Engineering and Computer Science lead the subject areas obviously, shown in Figure ??.

Exploratory data analysis (EDA) was pioneered by Tukey (1977). The main goal of EDA is to obtain a general sense of the data by mostly employing a variety of graphical techniques such as scatterplots, boxplots and histograms. Quite often, the dimension reduction methods are also used to visualize multivariate data. There are some advantages of graphical approaches to data analysis over the traditional formal statistical modeling and inference, including (1) summaries for large, complicated data sets; (2) checking assumptions in statistical models; (3) interaction between the researcher and the data; (4) revealing structure, patterns, features, trends, anomalies, and relationships in data; (5) communicating the results of an analysis; (6) identifying the areas of interest. EDA is especially useful in early stages of data mining. Due to the popularity of data science and big data, EDA regains people's attention in the recent years for pattern discovery and statistical visualization.

The typical graphical techniques used in EDA such as the index plot, boxplot, scatter plot and histogram were developed for the classical data can not be applied directly to various symbolic data types such as interval-valued data, histogram-valued data, distribution-valued data and so on. On the other hand, when the dimensionality of symbolic data is high, one way to explore the insight structure of symbolic objects in the lower-dimensional space is to employ the DR methods. As a consequence, the development of an EDA tool with more visual methods and the advanced DR techniques for symbolic data is needed.

The R packages benefit users to analyze their data without difficulties. There have several R packages that related to the symbolic data analysis (see Table ??). The contexts include the formal statistical modeling and inference. However, little work has been conducted on the exploratory analysis and visualization. This motivates us to develop an R package, namely **ggESDA**, aiming at exploring symbolic data graphically. **ggESDA** is composed of the features to deal with the interval-valued data, statistics data frame, clustering results, and other components from **R6** ? class. The graphical techniques for interval-valued data can be divided into three categories: univariate, bivariate, and multivariate. **ggESDA** is designed to provides an overview of symbolic data sets and obtains a general understanding about the variables and their relationships.

This article is structured as follows. Section ?? introduces the **ggESDA** package design and features with some descriptive statistics for symbolic data. Section ?? describes the implementation of plots for exploring the real world interval-valued data. Principal component analysis (PCA) for interval-valued data with other functionalities are given in Section ?? and ??. We then conclude with our results in Section ??.

Package	Symbolic Object*	Description	Plots	Reference
<b>RSDA</b>	I	R to Symbolic Data Analysis	scatterplot, radar	Rojas <i>et al.</i>
<b>symbolicDA</b>	I	Analysis of Symbolic Data	radar	Dudek <i>et al.</i>
<b>HistDAWass</b>	H	Histogram-Valued Data Analysis	histogram, box, index	Irpino (2015)
<b>MAINT.Data</b>	I	Model and Analyse Interval Data	-	Silva & Brito
<b>iRegression</b>	I	Regression Methods for Interval-Valued Variables	-	Neto <i>et al.</i> (
<b>intReg</b>	I	Interval Regression	-	Toomet (201
<b>ISDA.R</b>	I	interval symbolic data analysis for R	histogram	Filho & Fag
<b>GPCSIV</b>	H	Generalized Principal Component of Symbolic Interval variables	-	Brahim (201
<b>GraphPCA</b>	H	Graphical Tools of Histogram PCA	-	Brahim & K
<b>ggESDA</b>	I	Exploratory Symbolic Data Analysis with 'ggplot2'	index, scatter, radar, etc.	Jiang (2022)

\*I for the interval-valued objects, H for the histogram-valued objects.

Table 1: The main features and the plots provided by the various R packages for symbolic data on CRAN.

## 2. The SDA-related packages and the ggESDA package design

### 2.1. The SDA-related Packages

There have several R packages that related to the symbolic data analysis (see Table ??). The contexts include the formal statistical modeling and inference. However, little work has been conducted on the exploratory analysis and visualization. This motivates us to develop an R package for symbolic data analysis. The most prominent packages on CRAN are commonly used for statistical or machine learning analyze. It can be briefly classified into two parts, one is focused on statistic analysis, and the other is general SDA packages including both analysis method and some graphical technology. Nevertheless, most of their graphical technology tends to use the basic graphics in R rather than **ggplot2**, or only visualizes univariate distribution which is difficult to present the relationship between variables.

Exploratory data analysis is cross-classified in two different ways where each method is either graphical or non-graphical. And then, each method is either univariate, bivariate or multivariate. **ggESDA** uses a high-level graphic system by **ggplot2** ? to solve the problem mentioned above and provides a variety of EDA methods in all kinds of the variate.

**ggESDA** is a R data visualization library based on **ggplot2**. It provides a high-level interface for drawing attractive and informative statistical graphics.

In Python, we can also find the SDA package such as **iardacil** ? which is available from the Github at <https://github.com/iardacil/SDA>. The zoomstart software in **iardacil** is provided with SODAS software project ?. It is a basic thinking for general radar plot, improved for distinct groups visualization by **ggESDA**, and implemented in R using **ggplot2**.

### 2.2. The ggESDA package designs

"Exploratory Data Analysis is majorly performed using the following methods: Univariate analysis:- provides summary statistics for each field in the raw data set (or) summary only on one variable. Ex:- CDF,PDF,Box plot, Violin plot.(don't worry, will see below what each of them is) Bivariate analysis:- is performed to find the relationship between each variable in the dataset and the target variable of interest (or) using 2 variables and finding the relationship between them.Ex:-Box plot, Violin plot. Multivariate analysis:- is performed to understand interactions between different fields in the dataset (or) finding interactions between variables

	Statistics
Univariate	internal mean, grand mean, range, sd, variance, quantile
Bivariate	covariance, correlation $(r, \rho, \tau)$

Table 2: The implemented summary statistics.

more than 2. Ex:- Pair plot and 3D scatter plot."

### *Descriptive statistics*

The main content of EDA relates to the basic numerical summaries of data (e.g., the central tendency measures, and variation or variability measures) and the basic graphical summaries of data. For example, the five-number summary of numerical data (minimum, 25% quartiles, median, 70% quartiles, and maximum) is used to construct a boxplot. Therefore, the basic descriptive statistics for symbolic data should be provided prior to the (some) graphing techniques. In addition, if we want to do the standardization or apply PCA to interval-valued data, the mean, variance and the covariance must be calculated first. There are many algorithms to calculate these descriptive statistics for interval-valued data. For examples, the methods we have introduced in Section ?? of part were the quantification methods, the quantile method, the distributional methods, and the interval arithmetic methods. As a consequence, choosing the appropriate methods to obtain the basic descriptive statistics for symbolic data is the first step to implement the exploratory symbolic data analysis.

The main content of EDA relates to the basic numerical summaries of data (e.g., the central tendency measures, and variation or variability measures) and the basic graphical summaries of data. For example, the five-number summary of numerical data (minimum, 25% quartiles, median, 70% quartiles, and maximum) is used to construct a boxplot. In the field of SDA, there are many algorithms to calculate descriptive statistics and frequency for interval-valued data, and we will illustrate the univariate and bivariate summaries respectively.

To build a statistic chart or analysis, descriptive statistics are necessary to be constructed, as well as the frequency occurring in each bin in a histogram chart. For a histogram chart, subdivisions of it into equidistant and non-equidistant will also be consider in this section.

For the quantile in interval-valued data, summarizing it may seem to be obvious to separate data into a minimum and maximum data table, then calculate quantiles of both data tables to build a new interval-valued quantile data table.

### *Graphics design*

The typical graphical techniques of EDA include the index plot, barplot, boxplot, and histogram for univariate data; the 2D scatterplot, joint histogram and double boxplot for bivariate data; and the satterplot matrix, and star plot for multivariate data. Of course, there are many other variants of graphical techniques have been proposed. Our aim in this study is to implement good statistical graphics to display symbolic data accurately and clearly.

A good picture is worth a 1,000 words. The typical graphical techniques of EDA include the index plot, barplot, boxplot, and histogram for univariate data; the 2D scatterplot, joint histogram and double boxplot for bivariate data; and the scatterplot matrix, and star plot for multivariate data. Of course, there are many other variants of graphical techniques have been proposed. Our aim in this study is to implement the good statistical graphics to display

symbolic data accurately and clearly. On the other hand, it's worth mentioning that there will be some advanced graphics implemented which are called the min-max plot, and center-range plot due to the characteristics of interval-valued data.

The concept behind **ggplot2** consists of three different fundamental parts:

**Plot = data + Aesthetics + Geometry.**

The principal components of every plot can be defined as follow:

1. data is a data frame.
2. Aesthetics is used to indicate  $x$  and  $y$  variables. It can also be used to control the color, the size or the shape of points, the height of bars, etc.
3. Geometry defines the type of graphics such as histogram, box plot, line plot, density plot, dot plot, and so on.

General **ggplot2** syntax

```
ggplot(data, aes(...)) + geom() + ... + stat() + ...
```

```
R> library(ggplot2)
```

```
R> ggplot(data = iris, aes(x = Sepal.Length, y = Sepal.Width, color = Species)) +  
  geom_point(size = 3)
```

The package **ggESDA** is developed to as an extension of **ggplot2** to the interval-valued data. It provides a general and customized transformation function **classic2sym** implemented for generating a symbolic data table from classical data frame by clustering algorithm using the **RSDA** package ? form and generalized **ggESDA** to other prominent SDA packages in R.

General **ggESDA** syntax

```
ggInterval_*(data = intervaldata, mapping = aes(...)) + geom() + ... + stat() + ...
```

\* The graphical types are shown in Table ??.

```
R> library(ggESDA)
```

```
R> ggInterval_index(data = facedata, aes(x = AD, fill = IND)) +  
  geom_point()
```

Table ?? lists the implemented summary statistics. The implemented graphics are given in Table ???. All are based on the content of the books: "Symbolic Data Analysis: Conceptual Statistics and Data Mining" and "Symbolic Data Analysis and the SODAS Software".

A possible source of interval data is the aggregation of a numerical dataset. That is, the data is clustered into groups by some criterion and the intervals are constructed by summarizing values into the minimum value and maximum value within each group. One of the advantage is that a large dataset can be summarized into a more manageable size, the interval-valued symbolic data still retaining as much knowledge inherent to the entire dataset as possible.

Variate	Graphics
Univariate	index plot, histogram, box plot, min-max plot, center-range plot
Bivariate	scatterplot, 2D-histogram
Multivariate	radar, scatterplot matrix, 2D-histogram matrix, image

Table 3: The implemented graphics.

The aggregation of an object with data.frame class into an object of the IntervalData class is by the function `classic2sym()`.

The **ggESDA** aims to convert the traditional data into the **ggESDA** object and visualizes the symbolic data using **ggplot2**, which is shown in Figure ???. Each row (observation) in the conventional data matrix  $X_{p \times p}$  in the figure contains a vector of numeric values,  $O_i = (x_1, x_2, \dots, x_n)$ , while each row of the interval-valued data matrix  $I_{k \times p}$  contains a vector of intervals (ranges),  $C_j = ([a_{j1}, b_{j1}], [a_{j2}, b_{j2}], \dots, [a_{jp}, b_{jp}])$ , called a CONCEPT (or UNIT). The CONCEPT describe the behavior of a group of observations. Thus, the aggregation method between them is an essential process in SDA. See Section ??? for details.

#### *Built-in datasets and object class*

In the following, we used the face recognition data to illustrate some graphing functions we have implemented so far.

The face recognition data is widely used in the SDA literature (??; ??; ?). The dataset gives six face measurements of nine men (Figure ???), each with three observations, resulting in a total 27 observations (Table ???). The measurements for each observation came from a sequence of over 1,000 images. They cover a range of values, hence interval-valued variables.

```
R> library(ggESDA)
R> data(facedata)
R> facedata
```

```

              AD              BC              AH
      <symblc_n>      <symblc_n>      <symblc_n>
1 [155.00 : 157.00] [58.00 : 61.01] [100.45 : 103.28]
2 [154.00 : 160.01] [57.00 : 64.00] [101.98 : 105.55]
3 [154.01 : 161.00] [57.00 : 63.00] [99.36 : 105.65]
...
```

In the following, to illustrate some graphing functions we have implemented so far, we will use the two well-known symbolic datasets. One is face recognition data (??; ??; ?). The dataset gives six face measurements of nine men (Figure ???), each with three observations, resulting in a total 27 observations. The measurements for each observation came from a sequence of over 1,000 images. They cover a range of values, hence interval-valued variables.

The other is the Environment questionnaire data (from the SODAS software package ?). Because of the demonstration of the performance dealing with modal multi-valued data, the same datasets were used in the radar plot. This dataset contains 14 objects and 17 variables and 4 of them are modal multi-valued, while the rest are interval-valued variables. Both of the part of example datasets are shown as follow:

Data	Reference	Usage
facedata	?; ?	iPCA
oils	?; ?	iPCA
mushroom	?; ?	iRegression
Cardiological (blood pressure)	?; ?	iRegression
AbaloneIdt	?	Dissimilarity measure
Environment	?	EDA

Table 4: Summaries of the built-in interval-valued data.

### 3. Descriptive statistics for interval-valued data

#### 3.1. Descriptive univariate statistics

##### *Quantification approaches*

In statistics, it may be more interesting to discuss mean and variance in a particular random variable  $Z$ ; see ?. The realization of  $Z$  for the observation  $W_u$  is the interval  $Z(W_u) = [a_u, b_u]$ , where  $u = 1, 2, \dots, m$  and  $m$  is the number of concepts.

##### *Distributional approaches*

First of all, assume that each object is equally likely to be observed with probability  $\frac{1}{m}$ , and the empirical density function of  $Z$  is defined as :

$$f(\xi) = \frac{1}{m} \sum_{u: \xi \in Z(W_u)} \left( \frac{1}{b_u - a_u} \right) \quad (1)$$

where  $\xi$  is the individual descriptions.

The Equation (??) is also equivalently to :

$$f(\xi) = \frac{1}{m} \sum_{u \in E} \frac{I_u(\xi)}{\|Z(u)\|}, \xi \in \mathbb{R} \quad (2)$$

where  $I_u(\cdot)$  is the indicator function that  $\xi$  is or is not in the interval  $Z(u)$ ,  $\|Z(u)\|$  is the length of that interval, and  $E = \{w_1, w_2, \dots, w_m\}$ .

Further, the symbolic sample mean from definition for  $Z$  is  $\bar{Z} = \int_{-\infty}^{\infty} \xi f(\xi) d\xi$ , which can be reduced as :

$$\bar{Z} = \frac{1}{m} \sum_{u \in E} \frac{a_u + b_u}{2} \quad (3)$$

Finally, after getting the sample mean, the symbolic sample variance can be defined as follow:

$$\begin{aligned} S^2 &= \int_{-\infty}^{\infty} (\xi - \bar{z})^2 f(\xi) d\xi \\ &= \int_{-\infty}^{\infty} \xi^2 f(\xi) d\xi - \bar{z} \end{aligned} \quad (4)$$

and substituting for  $f(\xi)$  from Equation (??), we have

$$\begin{aligned} \int_{-\infty}^{\infty} \xi^2 f(\xi) d\xi &= \frac{1}{m} \sum_{u \in E} \int_{-\infty}^{\infty} \xi^2 \frac{I_u(\xi)}{\|Z(u)\|} d\xi \\ &= \frac{1}{m} \sum_{u \in E} \int_{a_u}^{b_u} \frac{\xi^2}{(b_u - a_u)} d\xi \\ &= \frac{1}{3m} \sum_{u \in E} (b_u^3 - a_u^3) \end{aligned} \quad (5)$$

Hence,

$$S^2 = \frac{1}{3m} \sum_{u \in E} (a_u^2 + a_u b_u + b_u^2) - \frac{1}{4m^2} \left[ \sum_{u \in E} (a_u + b_u) \right]^2 \quad (6)$$

### Frequency distribution tables

"The frequency distribution table reflects how often an occurrence has taken place in the data. It gives a brief idea of the data and makes it easier to find patterns."

For the univariate histogram frequency, assume that we partition the interval  $I = [\min_{u \in E} a_u, \max_{u \in E} b_u]$  into  $r$  subintervals, and all of them in the histogram are equal distance. That is,  $I_g = [\zeta_{g-1}, \zeta_g), g = 1, 2, \dots, r$ , then  $\|I_j\| = \|I_k\|, j, k = 1, 2, \dots, r$ . As a consequence, the observed frequency of the interval-valued variate  $Z$  for the histogram subinterval  $I_g$  from the definitions is

$$f_g = \sum_{u \in E} \frac{\|Z(u) \cap I_g\|}{\|Z(u)\|} \quad (7)$$

Moreover, for the interval-valued variate  $Z$ , we can pool the  $a_u$  and  $b_u$  from the interval of all observations, and sort it as a new vector  $(x^{(1)}, x^{(2)}, \dots, x^{(2m)})$  to represent the cut of a histogram. The subinterval from the cut is then defined as  $I'_g = [x^{(j)}, x^{(j+1)})$ , where  $j = 1, 2, \dots, 2m - 1$ , and apply the Equation (??) to get frequency. In most cases,  $\|I'_g\|$  will not be equal to another, so we can get another histogram type, called non-equidistant-bin histogram.

### Standardization of interval-valued data

Before performing iSIR algorithm, we first standardize each interval-valued variable of the data to have zero mean and unit variance. This standardization prevents variables with larger measurement scale from dominating those with smaller scales. De Carvalho *et al.*, (2006) proposed three standardization approaches for interval-valued variables: dispersion of the interval centers, dispersion of the interval boundaries, and the global range.

In this study, the general rule for standardizing interval-valued variables  $\Xi_j, j = 1, \dots, p$ , consists in performing the same transformation separately to both the lower and upper bounds of all intervals  $[a_{ij}, b_{ij}]$ , which standardizes all point values between  $a_{ij}$  and  $b_{ij}$  in the same linear way

$$\xi_{ij}^s = \left[ \frac{a_{ij} - \bar{\xi}_j}{\varsigma_j}, \frac{b_{ij} - \bar{\xi}_j}{\varsigma_j} \right],$$



where  $\bar{\xi}_j$  is the sample mean of interval variable  $\Xi_j$  given in Equation (??) and  $\varsigma_j$  is the sample standard deviation of interval variable  $\Xi_j$ . The sample standard deviation  $\varsigma_j$  is estimated following different iSIR methods. With the exception of the distributional approaches, the general estimate of  $\varsigma_j$  is given by the standard deviation of the interval centers, i.e.,

$$\varsigma_j = \left[ \frac{1}{n} \sum_{i=1}^n \left( \frac{a_{ij} + b_{ij}}{2} - \bar{\xi}_j \right)^2 \right]^{1/2}.$$

For EJD, GQ, and SPT,  $\varsigma_j$  is obtained from Equation (??).

*An example*

```
R> mean(facedata)

      AD      BC      AH      DH      EH      GH
<dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
1    163.   60.0  113.   113.   59.8   57.7

R> sd(facedata$AD)

[1] 6.816762
```

### 3.2. Descriptive bivariate statistics

*The empirical joint density method (EJD)*

For bivariate interval-valued variables,  $Z_1$  and  $Z_2$ , the observations on the rectangle  $Z(u) = Z_1(u) \times Z_2(u) = ([a_{1u}, b_{1u}], [a_{2u}, b_{2u}])$  for each  $u \in E$ . Assume individual vectors  $x \in \text{vir}(d_u)$  are each uniformly distributed over the respective intervals  $Z_1(u)$  and  $Z_2(u)$ . Define the empirical joint density function for  $(Z_1, Z_2)$ :

$$f(\xi_1, \xi_2) = \frac{1}{m} \sum_{u \in E} \frac{I_u(\xi_1, \xi_2)}{\|Z(u)\|},$$

where  $I_u(\xi_1, \xi_2)$  is the indicator function that  $(\xi_1, \xi_2)$  is or is not in the rectangle  $Z(u)$  and where  $\|Z(u)\|$  is the area of this rectangle. The bivariate interval-valued variables covariance function is defined as

$$\begin{aligned} \text{cov}(Z_1, Z_2) &= S_{Z_1 Z_2} = \int_{-\infty}^{\infty} (\xi_1 - \bar{Z}_1)(\xi_2 - \bar{Z}_2) f(\xi_1, \xi_2) d\xi_1 d\xi_2 \\ &= \frac{1}{m} \sum_{u \in E} \frac{1}{(b_{1u} - a_{1u})(b_{2u} - a_{2u})} \\ &\quad \times \iint_{(\xi_1, \xi_2) \in Z(u)} \xi_1 \xi_2 d\xi_1 d\xi_2 - \bar{Z}_1 \bar{Z}_2 \\ &= \frac{1}{4m} \sum_{u \in E} (b_{1u} + a_{1u})(b_{2u} + a_{2u}) \\ &\quad - \frac{1}{4m^2} \left[ \sum_{u \in E} (b_{1u} + a_{1u}) \right] \left[ \sum_{u \in E} (b_{2u} + a_{2u}) \right]. \end{aligned}$$

Many of the principles developed for the univariate case can be expanded to a general  $p$ -variate case,  $p > 1$ . We shall focus attention on obtaining joint histograms for  $p = 2$ . The following will reveal the statistics and bivariate histogram.

#### Linear Correlation

For bivariate interval-valued variables,  $Z_1$  and  $Z_2$ , the observations  $u$ , where  $u \in E$ , on the rectangle  $Z(u) = Z_1(u) \times Z_2(u)$  is  $([a_{1u}, b_{1u}], [a_{2u}, b_{2u}])$ . Assume the individual description vectors  $\xi$  are each uniformly distributed over the respective intervals  $Z_1(u)$  and  $Z_2(u)$ . Define the empirical joint density function for  $(Z_1, Z_2)$ :

$$f(\xi_1, \xi_2) = \frac{1}{m} \sum_{u \in E} \frac{I_u(\xi_1, \xi_2)}{\|Z(u)\|} \quad (8)$$

where  $I_u(\xi_1, \xi_2)$  is the indicator function that  $(\xi_1, \xi_2)$  is or is not in the rectangle  $Z_u$  and where  $\|Z(u)\|$  is the area of this rectangle.

There are three methods to calculate bivariate interval-valued variables covariance function. For the first one, it can be derived by ? as following:

$$\text{cov}(Z_1, Z_2) = \frac{1}{4m} \sum_{u \in E} (b_{1u} + a_{1u})(b_{2u} + a_{2u}) - \frac{1}{4m^2} \left[ \sum_{u \in E} (b_{1u} + a_{1u}) \right] \left[ \sum_{u \in E} (b_{2u} + a_{2u}) \right] \quad (9)$$

#### The symbolic covariance method (GQ)

Second, an alternative expression of the symbolic sample variance for  $Z$  in Equation (??) can be expressed as

$$S^2 = \frac{1}{3m} \sum_{u \in E} \left[ (a_u - \bar{Z})^2 + (a_u - \bar{Z})(b_u - \bar{Z}) + (b_u - \bar{Z})^2 \right]$$

The above equation can be generalized by ? to formulate the form of the symbolic sample covariance for  $Z_j$  and  $Z_{j'}$  as

$$\text{cov}(Z_j, Z_{j'}) = \frac{1}{3m} \sum_{i=1}^m G_j G_{j'} [Q_j Q_{j'}]^{1/2}, \quad j, j' = 1, 2, \dots, p \quad (10)$$

where for  $J = j, j'$ ,

$$Q_J = (a_{iJ} - \bar{Z}_J)^2 + (a_{iJ} - \bar{Z}_J)(b_{iJ} - \bar{Z}_J) + (b_{iJ} - \bar{Z}_J)^2, \\ G_J = \begin{cases} -1, & \text{if } \xi_{iJ}^c \leq \bar{Z}_J \\ 1, & \text{if } \xi_{iJ}^c > \bar{Z}_J \end{cases}$$

and  $\xi_{iJ}^c$  is the midpoint of the interval  $[a_{iJ}, b_{iJ}]$ .

#### The total sum of products (SPT)

Last, it was further demonstrated by ? and ? that the sample variance in Equation (??) is a function of the total sum of squares (SST) and that the SST can be decomposed into the sum

of the internal (within) variation and the between variation. The total sum of products (SPT) is the sum of the within sum of products and the between sum of products. The Equation (??) was also extended to the bivariate case to obtain the sample covariance of  $Z_j$  and  $Z_{j'}$  based on the decomposition of the SPT as

$$\text{cov}(Z_j, Z_{j'}) = \frac{1}{6m} \sum_{i=1}^m \left[ 2(a_{ij} - \bar{Z}_j)(a_{ij'} - \bar{Z}_{j'}) + (a_{ij} - \bar{Z}_j)(b_{ij'} - \bar{Z}_{j'}) + (b_{ij} - \bar{Z}_j)(a_{ij'} - \bar{Z}_{j'}) + 2(b_{ij} - \bar{Z}_j)(b_{ij'} - \bar{Z}_{j'}) \right] \quad (11)$$

The definitions and calculations of the symbolic sample covariance in Equations (??)-(??) are consistent with the results in the classic data case if  $a_{ij} = b_{ij}$  for  $i = 1, 2, \dots, m$ ,  $j = 1, 2, \dots, p$ . If  $j = j'$ , the Equation (??) reduces to the sample variance of the interval-valued variable as given in Equation (??).

#### *Two-dimensional histogram*

Analogously with Equation (??), we can find the joint histogram for  $(Z_1, Z_2)$  by graphically plotting  $\{R_{g_1 g_2}, p_{g_1 g_2}\}$  over the rectangles  $R_{g_1 g_2} = \{[\zeta_{1,g_1-1}, \zeta_{1,g_1}) \times [\zeta_{2,g_2-1}, \zeta_{2,g_2})\}$ ,  $g_1 = 1, 2, \dots, r_1$ ,  $g_2 = 1, 2, \dots, r_2$ , where

$$f_{g_1 g_2} = \sum_{u \in E} \frac{\|Z(u) \cap R_{g_1 g_2}\|}{\|Z(u)\|} \quad (12)$$

i.e.,  $f_{g_1 g_2}$  is the number of observations that fall in the rectangle  $R_{g_1 g_2}$  and is not necessarily an integer value (except in the special case of classical data). The relative frequency that an arbitrary individual description vector lies in the rectangle  $R_{g_1 g_2}$  is therefore

$$p_{g_1 g_2} = \frac{f_{g_1 g_2}}{m} \quad (13)$$

#### *An example*

```
R> cov(facedata$AD, facedata$BC, method = "billard")
```

```
[1] 21.47015
```

## 4. Graphics for interval-valued symbolic data

### 4.1. Univariate plots

**Index plot** Another graphical technique, called index plot, is famous in conventional data as well. It is composed of minimum and maximum in interval-valued data and exhibits the concepts' behavior. Both of the function

```
ggInterval_index(data = NULL, mapping = aes(NULL), plotAll = FALSE)
```

can achieve this objective, but there is still a slight difference for visualizing.

Take the variable AD for example, and set option `mapping = aes(x = AD)`. The former function, shown in Figures ?? and ??, directly displays the interval which is similar to min-max plot. Among these plots, we classify the concepts by the same person by setting the aesthetics option `fill` and can discover the values of people LOT, KHA, INC, FRA in AD variable are less than others, which easily compares the difference.

Figure ?? shows two representations of index plots for all six variables. The index plot (also called the run sequence plot ( $x_i$  vs  $i$ )) takes a single argument which is a continuous variable and plots the values on the  $y$  axis, with the  $x$  coordinate determined by the position of the number in the vector.

Figure ?? displays the index plots for the face recognition dataset with different representations: (a) vertical representation; (b) vertical representation with color mapping; (c) vertical representation with ordering by the centers of each variable; (d) vertical representation with ordering by the ranges of each variable. According to the symmetric of facial characteristics for a normal person, we observed that two people, HUS and KHA, have different quantities of AH and DH measurements.

**Boxplot** Figure ?? shows the boxplot side-by-side for all six variables. For each quantile box, the locations of the bottom and top of the box are calculated from the minima and maxima of the intervals separately. The boxplot provides a quick view of a data distribution and particularly useful for comparing distributions between several groups of variables. As we can see that AD (the length of outer corner of eyes) is the longest length than other measurements. AH and DH (EH and GH) are similar as they are expected to be symmetric for the facial characteristics.

```
R> ggInterval_boxplot(facedata, plotAll = T)
```

**Bar chart** "The bar graph is very convenient while comparing categories of data or different groups of data. It helps to track changes over time. It is best for visualizing discrete data."

**Segmented or stacked bars**

**Quantile plot**

**Time series plot** Figure ??

**Pie charts** "Pie charts are mainly used to comprehend how a group is broken down into smaller pieces. The whole pie represents 100 percent, and the slices denote the relative size of that particular category."

**Histograms** "Histograms are similar to bar charts and display the same categorical variables against the category of data. Histograms display these categories as bins which indicate the number of data points in a range. It is best for visualizing continuous data."

A histogram of the interval data is constructed as follows.

1. Let  $I = [\min_{u \in E} a_u, \max_{u \in E} b_u]$  be the interval that spans all the observed values of  $Z$  in  $\mathcal{X}$ .
2. Partition  $I$  into  $r$  subintervals  $I_g = [\xi_{g-1}, \xi_g), g = 1, \dots, r-1$ , and  $I_r = [\xi_{r-1}, \xi_r]$ .
3. The observed frequency for the histogram subinterval  $I_g = [\xi_{g-1}, \xi_g), g = 1, \dots, r$ :

$$f_g = \sum_{u \in E} \frac{\|Z(u) \cap I_g\|}{\|Z(u)\|}$$

4. The relative frequency:  $p_g = f_g/m$ , that is,  $p_g$  is the probability an arbitrary individual description vector  $x$  lies in the interval  $I_g$ .
5. The histogram for  $Z$ :  $\{(I_g, f_g), g = 1, \dots, r\}$ .
6. If we want to plot the histogram with height  $f_g$  on the interval  $i_g$  so that the "area" is  $p_g$ , then  $p_g = (\xi_g - \xi_{g-1}) \times f_g$ .

Figure ?? shows the histograms with non-equidistant breaks for all six variables. The breaks are obtained by the boundaries of the intervals. That is, all the minima and maxima of the intervals for that variable are used to construct breaks. The phenomenon observed from these histograms is similar to those from above plots.

```
R> ggInterval_hist(facedata, plotAll = T, bins = 15) +
R>   theme_hc()+
R>   scale_y_continuous(breaks = c(0, 0.035, 0.07))+
R>   labs(title = "")
R> ggInterval_hist(this.data, plotAll = T,
R>   method = "unequal-bin") +
R>   theme_hc()+
R>   scale_y_continuous(breaks = c(0, 0.2, 0.4))+
R>   labs(title = "")
```

In the same way, demonstrating the distribution of data by histogram is also one of the most common ways. Construct the frequency of data by Section ?? for the first, which will be processed by the function

```
ggInterval_hist(data = NULL, mapping = aes(NULL), method = "equal-bin",
  bins = 10, plotAll = FALSE)
```

Option `method` is the key for setting whether the histogram bins are equal width, detail in Section ??, which can be set with "equal-bin" or "unequal-bin". In Figure ?? depicts the histograms with equidistant and non-equidistant breaks for all six variables. Both of them

present the location and scale of distribution of each variable, but the non-equidistant-bin histogram whose breaks are obtained by the boundaries of the intervals shows more details about the characteristic of data. As we did before, set `plotAll` and `method`, and it can be directly completed with one line command. In the figure, we may obtain some similar distribution information conclusion with Figure ??.

**Min-max plot** Figure ?? shows the min-max plot for face recognition data.

```
R> #One case for plotting all variables (Relative Position)
R> lapply(1:6, FUN = function(x){
R>   plot.var <- x
R>   ggInterval_minmax(facedata, aes(facedata[[plot.var]]))+
R>     scale_color_manual(values = c("darkblue", "darkred"))+
R>     guides(colour = F) +
R>     theme_bw() +
R>     coord_fixed(ratio = 1)}) %>%
R>   ggarrange(plotlist = ., nrow = 1, ncol = ncol(facedata), labels = "")

R> #The other case using 'plotAll = T' (Absolute Position)
R> ggInterval_minmax(facedata, plotAll = T, scaleXY = "global") +
R>   scale_color_manual(values = c("darkblue", "darkred")) +
R>     guides(colour = F) +
R>     theme_bw() +
R>     coord_fixed(ratio = 1)
```

For interval-valued data, one of the advanced graphics implemented is called the min-max plot, which shows the interval characteristics of the data. As Figure ?? shown, we visualize the Face data for example by the function

```
ggInterval_minmax(data = NULL, mapping = aes(NULL), scaleXY = "local",
  plotAll = FALSE)
```

For the top six of Figure ?? shows the difference between each location of concept in each variable through the 45-degree line, which is a relative position of the variable itself (by setting option `scaleXY = "local"`). And represents the range by the connected line between minimum and maximum. To design the widely-view exploratory graphic function like this instance, we apply the option `plotAll = TRUE` to extend one variable visualization to full variables in the same plot. For examples, the figure constructed by the code

However, when the unit is considered, the absolute position in the full sample space will display as the bottom six of Figure ?. It presents a different boundary in the axis from the previous figure, which makes researchers know about the actual location in each variable. In this case, we can clearly discover that the sample space changes along with the variables.

To complete the visualization, it actually needs the command `ggInterval_minmax(facedata, plotAll = T, scaleXY = "local")`, merely. Other aesthetics can be omitted. Because we now focus on the full variables exploration, the `mapping` option for a particular variable is not necessary as well. Thus, in the following, we will pay more attention to the option we changing.

**Center-range plot** Figure ?? shows the center-range plot for face recognition data.

```
R> ggInterval_centerRange(facedata, aes(size = 1.5),
R>                        plotAll = T)+
R>   theme_classic() +
R>   stat_ellipse(geom = "polygon", fill = "blue",
R>               alpha = 0.05, col = "grey")

R> face.scale <- scale_sym(facedata)$intervalData #scale

R> ggInterval_centerRange(face.scale, aes(size = 1.5),
R>                        plotAll = T)+
R>   theme_classic()+
R>   stat_ellipse(geom = "polygon", fill = "blue",
R>               alpha = 0.05, col = "grey")
```

In this field, it reveals the trend, distribution, how large the data spread, etc, for a particular variable. Moreover, another advanced graphics implemented is called center-range plot, which helps researchers to be able to grasp the relationship between center and range.

The side-by-side boxplot, which provides a quick view of data distribution, dispersion, and is particularly useful for comparing distributions between several groups of variables. Use the function

```
ggInterval_boxplot(data = NULL, mapping = aes(NULL), plotAll = FALSE)
```

By setting option `plotAll`, the top panel of Figure ?? will be easily implemented. In this Figure, the dispersion of six variables is obviously distinct. However, it is difficult to compare with their distribution because of the distinct sample space which leaves no room for a suitable presentation. We standardize the Face data, shown in the bottom panel of the figure, and take a deeper look at it.

Clearly, there is a vacancy in the middle of variable AD, AH, and next to the minimum of the variable DH, which indicates the frequency there may be quite low. Especially for AD and AH, this factor may result in a bimodal distribution.

In addition, the overlap of the variables BC, EH, and GH located in the middle of five quantile-box seem larger than others, which is possible to lead a bell-shaped for its distribution.

For the center-range plot, it is a special technique for the dispersion observing. The code

```
ggInterval_centerRange(data = NULL, mapping = aes(NULL), plotAll = FALSE)
```

Setting options are the same as the former. With the Figure ?? demonstration, we divide the exploration into two parts by standardization, shown in Figure ??.

With regard to the top panel of Figure ??, there is no significant difference in the range of all variables. But focusing on a particular variable, the dispersion is quite large. For the bottom panel of Figure ??, the relationship between center and range is clearly exposed. Among them, there is an obvious relationship in the variable BC, which presents a negative correlation, where others are not significant.

## 4.2. Bivariate plots

The bivariate functions reveal relationships between variables, trends of the regression line, and joint frequency across the sample space. In this field, we design the scatter plot, two-dimension histogram, and also implement the most well-known approach, the vertices method of principal component analysis (V-PCA) ?.

**The 2D scatterplot** Figure ?? shows two representations of 2D scatterplot for variables of AD and BC. The color within the rectangles of 2D scatterplot reflects the degree of overlapping. The darker blue presents the area where the most observations were occupied. From these scatterplot of AD and BC, we can see that the observations could be divided into two major groups. One group has both larger AD and BC measurements while the other group has both smaller AD and BC measurements.

```
R> ggInterval_scatter(facedata, aes(x = AD, y = BC, fill = "gray80")) +
R   theme_bw() +
R>   theme(legend.position = "None")
```

**The 2d histograms** The joint histogram of the bivariate interval-Valued variables is constructed as follows.

1. Define the joint histogram for  $Z_1$  and  $Z_2$  by plotting  $\{R_{g_1g_2}, p_{g_1g_2}\}$  over the rectangles  $R_{g_1g_2} = \{[\xi_{1,g_1-1}, \xi_{1g_1}) \times [\xi_{2,g_2-1}, \xi_{2g_2})\}$   $g_1 = 1, \dots, r_1, g_2 = 1, \dots, r_2$ , where

$$p_{g_1,g_2} = \frac{1}{m} \sum_{u \in E} \frac{\|Z(u) \cap R_{g_1g_2}\|}{\|Z(u)\|},$$

2.  $p_{g_1g_2}$  is the probability an arbitrary individual description vector lies in the rectangle  $R_{g_1g_2}$ .
3. Then, if the "volume" on the rectangle  $R_{g_1g_2}$  of the joint histogram represents the probability, its height would be

$$f_{g_1g_2} = [(\xi_{1g_1} - \xi_{1,g_1-1}) \times (\xi_{2g_2} - \xi_{2,g_2-1})]^{-1} p_{g_1g_2}.$$

Figure ?? shows the 2D histogram matrices for face recognition data.

```
R> ggInterval_2DhistMatrix(facedata, aes(col = "grey50"), xBins = 10,
R>                           yBins = 10, removeZero = T, addFreq = F) +
R>   theme_light() +
R>   theme(legend.position = "bottom") +
R>   scale_fill_gradient(low = "gray95",
R>                       high = "red",
R>                       limits = c(0, 1),
R>                       na.value = "red")
```

Another bivariate relationship plot is two-dimension histogram, which is to calculate joint histogram frequency (details in Section ??) and using matrix visualization (?, ?) to plot. The code



```
R> ggInterval_2Dhist(facedata, aes(x = AD, y = BC, col = "grey30"),
R>                      xBins = 25, yBins = 25) +
R>   theme(legend.position = "bottom") +
R>   scale_fill_gradient(low = "gray95", high = "red", limits = c(0, 0.15),
R>                      na.value = "red")
```

uses the joint frequency between the variables AD and BC and makes the XY-axis bins partial into 25 subintervals. As Figure ?? shown, we can observe that the concepts in these joint sample spaces concentrate on the top-right and bottom-left areas that seem like a slightly positive correlation.

The addition options `removeZero` and `addFreq` make zero value and frequency displaying remove, shown as Figure ?. Not only the relationship presentation, the graphic technique also depicts the details frequency. Among these, we might get a similar conclusion between variables relation, but for the calculation, this technique may consume a larger time-complexity than scatter plot.

### 4.3. Multivariate plots

**Scatterplot matrix** Figure ?? shows the the scatterplot matrix for face data. The rectangles with the same color present the same subject. The association between any pair of two variables can be investigated from the scatterplot matrix. Obviously, two variables, AH and DH (EH and GH), have strong positive correlation. In addition, the observations seems to be easy discriminant by varibales AD and AH. Additionally, these graphic techniques can visualize multiple variables at the same time by matrix type. For instance, the code of scatter matrix of Figure ??:

```
R> ggInterval_scaMatrix(data = facedata)
```

In Figure ??, we use grayscale to represent the concepts and the rectangle to display the intersection of two interval-valued observations. The relationship among this graph is clearly depicted. The variables AH and DH are an obvious positive correlation, whereas AD and AH seems to be an exponential relation. Of course, there is an easy way for scatter plot of two particular variables using the function `ggInterval_scatter`.

**Radar plot** Figure ?? shows the starplot for all observations of the face data. For each star, the radius of a circle is fixed. Therefore, the observed values for each subject across all variables are scaled into the range of  $[0, 1]$ . Therefore, we could compare the distributions of variables within each subject (star).

In the field of multivariate visualization, the most common graphic technique is radar plot, also known as star plot. In **ggESDA**, we provide the function

```
ggInterval_radar(data = NULL, layerNumber = 3, inOneFig = TRUE, Drift = 0.5,
  showLegend = TRUE, showXYLABs = FALSE, plotPartial = NULL, alpha = 0.5,
  base_circle = TRUE, base_lty = 2, addText = TRUE, type = "default",
  quantileNum = 4, addText_modal = TRUE, addText_modal.p = FALSE)
```

Except for the aesthetic options `layerNumber`, `showLegend`, `showXYLabs`, `alpha`, `base_circle`, `base_lty`, and `addText`, the visualizing key is to determine which concepts are presented through setting `plotPartial` with the rows of concepts. In addition, it will be decided to use area ("default"), rectangle ("rect"), or quantile ("quantile") to represent the interval by option `type`.

Above all, take the Environmental questionnaire dataset for example, and the variety of typical multivariate radar plots for a specific concept is shown in Figure ?? by the code

```
R> #Left fig : Remove the modal multi-valued variables for demo
R> ggInterval_radar(Environment[, 5:17], plotPartial = 2, showLegend = F,
R>                      addText = F)

R> #Right fig : Remain the modal multi-valued variables
R> ggInterval_radar(Environment, plotPartial = 2, showLegend = F,
R>                      base_circle = F, base_lty = 1) +
R>   scale_fill_manual(values = "gray50") +
R>   scale_color_manual(values = "gray50")
```

Use the color to represent the interval area, and it can clearly present the behavior of the particular concept in each variable. The figure in the left panel shows the relative position of the interval by the percentage circle, whereas the right one removes it and adds the frequency in the plot. Additionally, the kind of radar plot is generalized to visualize not only the interval-valued data but the modal multi-valued, which we design a cuboid to represent, and the height will correspond to the probability of the factor in the variable, as the right panel in the figure. The technique not just merges the distinct variable's type in one plot but even helps the researcher to compare with concepts easily under multiple variables. Figure ?? shows the three observations presented by the different plot.

Moreover, another advanced implemented graphic technique of radar plot is classification by different concepts using a color mapping. By the code:

```
R> area.fig <- ggInterval_radar(Environment, plotPartial = c(4, 6),
R>                      showLegend = F, base_circle = F,
R>                      base_lty = 1, addText = F,
R>                      addText_modal = F) +
R>   scale_fill_manual(values = c("darkred", "darkblue")) +
R>   scale_color_manual(values = c("darkred", "darkblue")) +
R>   labs(title = "") +
R>   theme_hc()

R> rect.fig <- ggInterval_radar(Environment, plotPartial = c(4, 6),
R>                      showLegend = F, base_circle = F,
R>                      base_lty = 1, addText = F, type = "rect",
R>                      addText_modal = F) +
R>   scale_fill_manual(values = c("darkred", "darkblue")) +
R>   scale_color_manual(values = c("darkred", "darkblue")) +
R>   labs(title = "") +
```

```
R> theme_hc()

R> gridExtra::marrangeGrob(list(area.fig, rect.fig),
R>                           nrow = 1, ncol = 2, top = "")
```

we can get the Figure ???. It's worth mentioning that whenever the users determine the option `plotPartial` with multiple observations and set `inOneFig = TRUE` which means if pool observations into the same figure, the mechanism in the function will automatically classify by color. In this case, the two visualized concepts are represented by blue and red color, and the modal multi-valued variables are stacked by the same factor. On the contrast, the graph displays the interval-valued variables in two ways. The left panel of the figure shows a typical area-presenting way, while the right one use rectangle to show the interval. These implement can be completed by setting the option `type` with "default" and "rect" respectively.

**Quantiles plot** Last but not least, we propose a graphic method based on quantiles. The benefit of this proposed method is that distributional details are visible for the graph. Under the option `type = "quantile"`, for example, we divide the quantile percentage of 25% to 75% into the equal subintervals represented as grayscale and shown in Figure ???. The variable WELFARE of the left panel in the figure depicts the rapid increase close to the 3rd quartiles, which implies it exists a left-skewed distribution. In this way, it becomes more straightforward to extend the comparison of variables to datasets.

**Data image** The dimension-free approach for visualizing symbolic data. The traditional statistical graphics use the geometric characteristics such as the scale, length, angles and size to display the information of objects/variables on the screen. As shown in Figure ??(a)(c), the index plots for face dataset were displayed as data images where the color strip represents the ranges of the measurements. Since the space required for this kind of displays is limited, the traditional graphics will become useless when graphing the ultra high-dimensional and large-scale dataset. In such case, we could consider the matrix visualization (MV) method (Chen, 2002; Chen et al., 2004) as a solution for this issue. MV is a dimension-free graphical technique that can simultaneously explore the associations of up to thousands of subjects, variables, and their interactions. MV heavily utilizes the colors to represent the numerical quantities and employs the effective ordering algorithms to discover patterns of data (Figure ??(b)(d)).

The latter function, shown in Figure ?? and ??, visualizes the interval values using a color mapping. As the legend says, the blue color corresponds to the low values, whereas the red color corresponds to the high. The only difference among these plots is the color bar extension (control by the option `full_strip = TRUE`), which will quickly discover concepts' behavior through the visual sense, as Figure ?? shown.

```
ggInterval_indexImage(data = NULL, mapping = aes(NULL), plotAll = FALSE,
  column_condition = TRUE, full_strip = FALSE)
```

For exploring data, we extend the index plot into full variables and visualize by color mapping as we did before using the code

```

R> #Column Condition
R> colCondition <- ggInterval_indexImage(facedata, plotAll = T,
                                         R> full_strip = T) +
R>   scale_colour_gradient2(low = "green", mid = "gray15", high = "red")

R> #Matrix Condition
R> #Prepare midpoint for scale_colour_gradient2
R> ggESDA.facedata <- RSDA2sym(facedata)
R> facedata.m <- mean(c(min(ggESDA.facedata$statisticsDF$min),
R>                       max(ggESDA.facedata$statisticsDF$max)))

R> matCondition <- ggInterval_indexImage(facedata, plotAll = T,
                                         full_strip = T,
R>                                         column_condition = F) +
R>   scale_colour_gradient2(low = "green", mid = "gray15",
R>                           high = "red", midpoint = facedata.m)

R> ggarrange(colCondition, matCondition, nrow = 1, ncol = 2)

```

Figure ?? shows two kinds of presentation ways for this method.

- *Column condition:* This technique is to standardize data by each variable, which will be able to compare the observations easily without the unit confounder. Figure ?? ,for example, shows the concept JPL has a large value in all variables in contrast with KHA and INC.
- *Matrix condition:* Instead of standardizing variables one by one, this method does it with the full data matrix. That will reveal the different performance between each variable, as Figure ?? shown. The values in variable AD are obviously larger than others, which means the interocular distance is the longest than other distances in the face dataset.

**2D histogram matrices** ?? The index plot and the image plot for the aggregated histograms of iris data.

**The scatterplots3d and dynamic graphics** Other than the static visualization, using computers and mobile devices to interact with charts and graphs can aid users to gain the relationships among variables/subjects of data in more details. PI would like to implement the immediately interaction such as brushing, highlighting, zooming, and/or identifying using Shiny (<http://shiny.rstudio.com>) by RStudio. Shiny is a web application framework for R which is easy to turn the exploratory analyses into interactive web applications.

```

R> scatterplot3d.i(idata.x[,1:3,], col=y.C)
R> scatterplot3d.i(idata.x[,1:3,], col=y.C, rgl=T)
R> play3d(spin3d(axis=c(1,0,0), rpm=10), duration=10)

```

## 5. Visualizing intervals in 2D Principal Component space

Among many research topics of SDA, the dimensionality reduction is one of the active research issues. So far the main thread has been focused on the extensions of the PCA in the literature. Most PCA extensions to the interval-valued data belong to the category of symbolic-numerical-symbolic approaches. That is, the interval-valued data in input are analyzed according to the classic approaches and the results are furnished in terms of intervals. The primary extended approaches of PCA for interval-valued data are the quantification methods. That is, the given interval-valued data table is transformed to a classical numerical data table so that the traditional PCA can be applied. The most well-known approaches were the vertices method of PCA (V-PCA) (Cazes *et al.*, 1997; Chouakria, 1998) and the center method of PCA (C-PCA) (Chouakria *et al.*, 2000). V-PCA decomposes the correlation matrix of vertices of hypercubes while C-PCA decomposes the correlation matrix of the centers of intervals. Several methods such as V-PCA, C-PCA, S-PCA, MR-PCA, and I-PCA have been compared and review (Lauro, Verde and Irpino, 2008; Douzal-Chouakria, Billard and Diday, 2011). Some of methods have been implemented in the SODAS software.

A main goal of dimension reduction is data visualization. Interval-valued data are projected into a two- or three-dimensional DR subspace to enable users to explore the data structure. Suppose there exists an interval object in 3D space, as shown in Figure ??(a). The linear projection of this interval object by the DR algorithms is equivalent to transforming the object from the input coordinates to the DR coordinates, as shown in Figure ??(b). Several methods have been proposed to visualize interval objects in lower-dimensional subspace, such as 2D-convex hull (Verde and De Angelis, 1997) and parallel edge connected shapes (PECS) (Irpino, Lauro and Verde, 2003). In the following, we introduce three additional types of graphical representation of the projected interval object on the DR subspace.

### 5.1. The maximum covering area rectangle (MCAR)

The maximum covering area rectangle (MCAR) (Cazes *et al.*, 1997; Chouakria, 1998) is widely used for graphical representation of interval objects on a DR subspace owing to its simplicity. It constructs observations  $\{z_{ik}\}_{i=1}^n$  as  $k$ -dimensional hyperrectangles (usually  $k = 2$ ) on a DR subspace, where the sides of hyperrectangles are parallel to the axes. As shown in Figure ??(c), the circumscribed rectangle with green dashed line in the first DR plane is an example of MCAR. The main drawback of MCAR is that the interval objects are over-sized in the DR subspace with respect to the real objects in  $\mathbb{R}^p$ . That is, the hyperrectangle in DR space may include data points that do not belong to the original observations. In this study, MCAR is adopted by CM, VM, and FV.

### 5.2. The polytopes representation

The polytopes representation for interval data was proposed by Le-Rademacher and Billard (2012). The vertices of interval-valued data are projected onto lower-dimensional subspace, and the edges of points are formed in the DR subspace by connecting the vertices, as in the original input space  $\mathbb{R}^p$ . That is, for the vertices of the  $i$ th interval-valued observation, the projections are

$$\mathbf{z}_{V_{ik}}^v = \hat{\beta}_k^T \mathbf{x}_{V_i}^v, \quad i = 1, \dots, n, \quad k = 1, \dots, K.$$

The edges with red dotted lines in the first DR plane in Figure ??(c) show an example of

the polytopes representation for an interval object. The polytopes representation improves MCAR and provides a true projection of the observed interval data. In this study, the polytopes representation is adopted by EJD, GQ and SPT.

Here we take the face data as an example to illustrate the visualization of interval-valued objects in PCA space. Suppose there is an interval-valued object in the 3D space as shown in Figure ??(a). The projection of the object on the PCA space is obtained by the linear combination of three axes of the original input space. This corresponds to the rotation of the original space in the DR space ( $dr1 \sim dr3$ ) as shown in Figure ??(b). The projection of the interval-valued object on the first DR plane of  $dr1$  and  $dr2$  can be constructed by several methods.

A more recent method, called the polytope representation, which is proposed by Le-Rademacher and Billard (2012) is the line connections among vertices in the DR space (the light blue-colored line in Figure ??(c)) as they were connected in the original input space (the red-colored line in Figure ??(c)). Applying vertices PCA to the face data, Figure ?? shows the projections of face recognition data in the first PCA plane with MACR (left) and polytope (right) representations. These two graphs shown very different visualization of the data in the lower-dimensional DR space. As for the quality assessment of the DR results, it needs further investigation.

Similar to PCA/SIR for interval-valued data, two key steps of PCA/SIR for histogram-valued data are required: (1) compute the sample means, sample (weighted) variance-covariance of symbolic variables to determine a PC space and (2) define a linear combination of symbolic variables. Chen, Wang and Qin (2014) defined a general probabilistic symbolic data to describe the intervals and histograms. This idea gives a clue to develop the visualization for histogram-valued data in the DR space.

The linear combination of  $p$  intervals Moore's (1966) linear combination of  $p$  intervals, the  $i$ th projected observation is:

$$z_{ik} = \hat{\beta}_k^T \xi_i = \sum_{j=1}^p \hat{\beta}_{kj} \xi_{ij} = [z_{ik}^a, z_{ik}^b],$$

where

$$z_{ik}^a = \sum_{j=1}^p \hat{\beta}_{kj} (\eta a_{ij} + (1 - \eta) b_{ij}) \quad \text{and} \quad z_{ik}^b = \sum_{j=1}^p \hat{\beta}_{kj} ((1 - \eta) a_{ij} + \eta b_{ij}),$$

with  $\eta = I(\hat{\beta}_{ij} > 0)$ . The linear combination algorithm is equivalent with the directly projection by vertices (Douzal-Chouakia, Billard and Diday, 2011).

With the technique, we combine the dimension reduction methods that **RSDA** provides, and implement the maximum covering area rectangle (MCAR) ? using the code:

```
R> pca.results <- RSDA::sym.pca(facedata, method = "top")
R> ggInterval_scatter(pca.results$Sym.Components, aes(Dim.1, Dim.2)) +
R>   scale_fill_manual(values = Concepts)
```

which is widely used for graphical representation of interval objects on a DR (dimension reduction) subspace, owing to its simplicity, shown as Figure ??.

Function	Description
<code>classic2sym</code>	Convert the classical data to symbolic data
<code>RSDA2sym</code>	Convert the <b>RSDA</b> object to <b>ggESDA</b> object
<code>scale_sym</code>	standardize the symbolic data
<code>cov</code>	Covariance of interval-valued variables

Table 5: Utility.

Nevertheless, the main drawback of MCAR is that the interval objects are oversized in the DR subspace with respect to the real objects in  $\mathbb{R}^p$ , hence, the polytopes representation ? for interval data was proposed. In **ggESDA**, we have also developed the graphic function for them using

```
ggInterval_PCA(data = NULL, mapping = aes(NULL), plot = TRUE, poly = FALSE,
               concepts_group = NULL)
```

The control option of PCA visualization method mentioned above is `poly`, which can be set with `TRUE` for the polytopes representation as shown in Figure ?? . By the dimension reduction and color mapping on each person, Figure ?? shows most of the concepts are clearly separated.

## 6. Other functions

As far as generalization and extension are concerned, the package provides a simple way for making connections with other useful packages, so that the result of common statistical or machine learning methods on other packages may be visualized as well using **ggESDA** if it is interval-valued. In addition, we may get the classical data most of the time, so summarization between classical to interval-valued data becomes important. The following will discuss these explicitly.

### 6.1. Generic functions

Support for common generic functions, such as `summary` or `print.write.table`

### 6.2. Customized plots and graphical parameters

using `+` `facet` according to a discrete group variable. log scales, color, fill with selected spectrum add text, lines, labeling, title, tick marks, font (give a example here)

Figure ??

```
R> ggInterval_index(d, aes(y = AD, fill = Concepts)) +
R>   geom_text(aes(x = 1:27,
R>                 y = .data$AD$min,
R>                 label = rownames(facedata),
R>                 vjust = 1.5), col = "red") +
R>   geom_line() +
R>   labs(title = "Customize plot", fill = "Concepts") +
```

```
R> scale_fill_brewer(palette = "Set1") +
R> facet_grid(. ~ d$g, scales = "free") +
R> theme_economist_white()
```

### 6.3. Generalization with prominent SDA packages

For the demonstration, we consider two famous R packages for SDA, **HistDAWass** ? and **MAINT.Data** ?. Both of these make lots of contributions to the statistics of SDA, so we tend to generalize these data into the **ggESDA** type, which may help us to visualize.

#### *General Principle*

Generally, it is not merely the well-known packages in R that can make a plot using it. As long as keeping some principle, it will be easily performed:

1. Understand the data structure clearly if it is an object from other packages.
2. Extract the min data and max data from it or make some necessary transformation.
3. Classify the data you extract belong.
4. Reorganized it and use `classic2sym` for the final transformation.
5. Visualize the front result using **ggESDA**.

Because of the interval-valued data, all SDA packages studying in the same field will store and deal with the min and max data. Hence, the transformation method in section ?? plays a important role. With the connection being built, it can be compatible with all other tools in R.

#### **HistDAWass**

We further use the principle to process BLOOD data in **HistDAWass**, first. With the method **HistDAWass** provided, it will be more convenient to get min and max data:

```
R> library(HistDAWass)
R> # Get min and max data
R> BLOOD <- HistDAWass::BLOOD
R> blood.min <- get.Math.stats(BLOOD, stat = "min")
R> blood.max <- get.Math.stats(BLOOD, stat = "max")
R> blood <- data.frame(blood.min, blood.max)

R> # Reorganized and Build ggESDA obj.
R> blood.sym <- classic2sym(blood, groupby = "customize",
R>                          minData = blood[, 2:4],
R>                          maxData = blood[, 6:8])

R> # Make names
```



```
R> blood.names <- get.MathH.main.info(BLOOD)$varnames
R> blood.i <- blood.sym$intervalData
R> colnames(blood.i) <- blood.names
R> head(as.data.frame(blood.i), 5)
```

	Cholesterol	Hemoglobin	Hematocrit
1	[80.00 : 240.00]	[12.00 : 15.00]	[35.00 : 47.00]
2	[80.00 : 240.00]	[10.50 : 14.00]	[31.00 : 44.00]
3	[95.00 : 245.00]	[10.50 : 14.00]	[31.00 : 43.50]
4	[105.00 : 260.00]	[10.50 : 14.00]	[31.00 : 42.50]
5	[115.00 : 260.00]	[10.80 : 13.60]	[31.00 : 42.50]

After getting the necessary data, classifying data belonging is vital for reorganization, which means that differentiating the min data and max data. For instance, `minData = blood[, 2:4]` represents the min data are the columns of 2,3,4 in this case.

### **MAINT.Data**

However, it is also a common way to store interval-valued data by median and range. In **MAINT.Data**, the data will exist in this form. Fortunately, a median-range form is not difficult to deal with. We can do the necessary conversion directly to get the data we expect:

```
R> library(MAINT.Data)
R> #get data interval-valued data in AbaloneIdt
R> AbaloneIdt <- MAINT.Data::AbaloneIdt
R> Aba.range <- AbaloneIdt@LogR
R> Aba.mid <- AbaloneIdt@MidP

R> #make a necessary transformation for build min max data
R> Aba <- data.frame(Aba.min = Aba.mid - exp(Aba.range) / 2,
R>                  Aba.max = Aba.mid + exp(Aba.range) / 2)

R> # Reorganized and Build ggESDA obj.
R> Aba.sym<- classic2sym(Aba, groupby = "customize",
R>                      minData = Aba[, 1:7],
R>                      maxData = Aba[, 8:14])

# Make names
R> colnames(Aba.sym$intervalData) <- AbaloneIdt@VarNames
R> Aba.i <- Aba.sym$intervalData %>%
R>   cbind(Aba.obs = AbaloneIdt@ObsNames) %>%
R>   column_to_rownames(var = "Aba.obs")
R> head(Aba.i[, 1:4], 5)
```

	Length	Diameter	Height	Whole_weight
F-10-12	[0.34 : 0.78]	[0.26 : 0.63]	[0.06 : 0.23]	[0.21 : 2.66]

```

F-13-15 [0.39 : 0.82] [0.30 : 0.65] [0.10 : 0.25] [0.27 : 2.51]
F-16-18 [0.40 : 0.74] [0.32 : 0.60] [0.10 : 0.24] [0.35 : 2.20]
F-19-21 [0.49 : 0.72] [0.36 : 0.58] [0.12 : 0.21] [0.68 : 2.12]
F-23-24 [0.45 : 0.80] [0.38 : 0.63] [0.14 : 0.22] [0.64 : 2.53]

```

In brief, following the general principle in section ?? may facilitate the integration, extend utilize of **ggESDA** and generalize to all SDA studies.

#### 6.4. Aggregate conventional data table to interval-valued data table

Modern statistical graphical methods for visualizing big data nowadays we encounter large data sets in many areas, including genomics, finance records, environmental research, and internet logs. In these cases, the size of data is getting larger and the structure is more complicated. Aggregation of these huge (volume) of data into symbolic objects such as histograms or distributions is sometimes necessary. As a consequence, the following will discuss the benefit of the symbolic data after aggregating.

In practice, the symbolic data is often generated by aggregating massive datasets into intervals in order to make the management easy and appropriate. An interval-valued symbolic random variable  $X$ , taking values in interval, can be denoted such as  $X = [a, b] \subset R^1$ , where  $a \leq b$ , and  $a, b \in R^1$ . Let the random variable  $X$ , for instance, be the weight, then  $X = [50, 100]$  represents the interval covering the weight of people. With the advent of big data analytic, interval-valued data is becoming more common and accessible than ever.

Another possibility of the data structure would be a classical data, we provide multiple method for dealing with this kind of transformation. The following would discuss the functionality of `classic2sym`.

We will apply the breast mass dataset, which is computed from a digitized image of a fine needle aspirate (FNA), to demonstrate how does a classical dataset transforms into a symbolic dataset. The breast mass dataset describe characteristics of the cell nuclei present in the image. It can be downloaded from the kaggle at <https://www.kaggle.com/uciml/breast-cancer-wisconsin-data?select=data.csv>. There are 569 observations and 32 variables in the dataset. We are going to store this dataset in `breastData` as data frame type in R, and the variables will be shown as follows:

```

colnames(breastData)

[1] "id"                "diagnosis"
[3] "radius_mean"       "texture_mean"
[5] "perimeter_mean"    "area_mean"
[7] "smoothness_mean"   "compactness_mean"
[9] "concavity_mean"    "concave points_mean"
...

```

Except for the first two variables, they are all composed of mean, standard error, and "worst" in their own field respectively.

#### *K-means*

K-means clustering is a method of vector quantization, originally from signal processing, that

aims to partition  $n$  observations into  $k$  clusters in which each observation belongs to the cluster with the nearest mean. In **ggESDA**, the algorithm will be based on the **stats** package, and the number of  $k$  is a parameter that user can define themselves:

```
R> breastData <- dplyr::select(breastData, -id)
R> breastData.sym <- classic2sym(breastData, groupby = "kmeans", k = 5)
R> breastData.sym.i <- breastData.sym$intervalData
R> as.data.frame(head(breastData.sym.i[, 1:4], 5))
```

	diagnosis	radius_mean	texture_mean	perimeter_mean
1	B:0.04 M:0.96	[13.81 : 19.59]	[11.89 : 39.28]	[91.56 : 132.40]
2	B:0.00 M:1.00	[15.50 : 24.25]	[10.38 : 32.47]	[102.90 : 166.20]
3	B:0.00 M:1.00	[20.73 : 28.11]	[17.25 : 31.12]	[135.70 : 188.50]
4	B:0.68 M:0.32	[11.84 : 16.30]	[10.89 : 30.72]	[77.93 : 109.80]
5	B:0.98 M:0.02	[6.98 : 13.05]	[9.71 : 33.81]	[43.79 : 85.09]

The `id` is unused in this case, so we remove it by **dplyr** ?. Then using **classic2sym** to aggregate **breastData**. It will return several result sets include clustering result and interval-valued data, etc. The interval-valued data can be extracted by `$intervalData`, and it will be presented by the package of **RSDA** type.

The `groupby` is a parameter that determine what kind of aggregation methods will be used. Whenever the K-means method is applied, the consequent `k` will become meaningful, whereas the other situation is not. It is also a default method when users have no input arguments in `groupby`.

For the conventional exploratory data analysis, it is always a severe challenge to deal with enormous datasets because conventional displays suffered from overstrikes of data points representing the value (scatterplot type displays) or overstrikes of line segments connecting values of neighboring variables. As a consequence, exploratory symbolic data analysis (SDA) becomes a preliminary yet essential tool for summarizing the main characteristics of a data set before appropriate statistical modeling can be applied. Besides escaping the problem mentioned above, SDA can effectively reduce observations in data, which will make the study focus on what we interesting instead of unnecessary information such as Figure ??.

In Figure ??, we can clearly visualize the scatter plot in the right hands, which is represented by symbolic data and aggregated by K-means ?, and effectively reduce the observations from 53940 to 5 groups.

### *Hierarchical clustering*

The second well-known clustering algorithm is called Hierarchical clustering ?, also called hierarchical cluster analysis or HCA. It can be performed with a distance matrix calculated by raw data and used to present the distance of each cluster. In basic R package, it is also realized by **stats**, which the **ggESDA** is based on for implementing HCA:

```
breastData.sym <- classic2sym(breastData, groupby = "hclust")
breastData.sym.i <- breastData.sym$intervalData
```

Remark that the `k` parameter is not meaningful in the case without K-means clustering. In `classic2sym`, the keywords of HCA is called `hclust`.

### *Variables aggregation*

Using a particular variable to merge different data is a common way for data analysis, too. **ggESDA** provides such as this concept in `classic2sym` to analyze different factors of category variables, and merge the same factor into the symbolic data type:

```
R> breastData.sym <- classic2sym(breastData, groupby = "diagnosis")
R> breastData.sym.i <- breastData.sym$intervalData
R> head(breastData.sym.i[, 1:4], 5)

      radius_mean      texture_mean      perimeter_mean
B [6.98 : 17.85]  [9.71 : 33.81]  [43.79 : 114.60]
M [10.95 : 28.11] [10.38 : 39.28] [71.90 : 188.50]
      area_mean
B [143.50 : 992.10]
M [361.60 : 2,501.00]
```

In `breastData`, the only category variable is `diagnosis`, which means the diagnosis of breast tissues (M = malignant, B = benign). We put it as an input argument in `groupby` for merging different diagnosis results, and the interval-valued data of result sets will display its factor levels in row names.

### *User-defined aggregation*

In general, users may not always use the aggregation methods we provide, thus, besides generating a particular variable for the group, **ggESDA** facilitates the process through the min data and max data that user-defined.

For the demonstration, we will build both min data and max data using `runif`. Generate a uniform random variable to make sure that all min data are smaller than max data:

```
R> minData <- runif(100, -100, -50)
R> maxData <- runif(100, 50, 100)
R> demoData <- data.frame(min = minData, max = maxData)
R> demoData.sym <- classic2sym(demoData, groupby = "customize",
R>                               minData = demoData$min,
R>                               maxData = demoData$max)

R> demoData.sym.i <- demoData.sym$intervalData
R> as.data.frame(head(demoData.sym.i, 5))
```

```
      V1
1 [-75.85 : 63.98]
2 [-93.71 : 85.33]
```

classic2sym					
groupby Args.	Transformation	Data Type	Moment	Cluster Result	Require Other Args.
<b>kmeans</b>	K-means	Numeric/Category	Smaller data	V	TRUE
<b>hclust</b>	Hierarchical	Numeric/Category	Smaller data	V	FALSE
variable name	Variables aggregation	Numeric/Category	Own the pre-clustering group	-	FALSE
<b>customize</b>	User-defined	Numeric	Connect with other packages	-	TRUE

Table 6: Summary for `classic2sym` function.

3 [-64.99 : 94.69]

4 [-57.34 : 66.03]

5 [-66.02 : 50.95]

Then choose the `customize` argument in `groupby`, input which data are `minData` or `maxData`, and the transformation will be simply completed.

In order to simplify the process and make the preprocessing friendly, we develop these methods and let the people who want to analyze symbolic data easier. Overall, the conversion and essential concepts can be summarized in table ??.

## 7. Conclusion and future development

We developed **ggESDA** to complementary the **ggESDA** package for symbolic data analysis, especially for interval-valued data. Being an extension of **ggESDA**, we inherit most of its plot-construction syntax and strict adherence to the rules governed by *The Grammar of Graphics* ?. So far, we have demonstrated various newly implementations of traditional graphics to symbolic data. As an answer to the challenges of big data and complex data that the traditional data encounter, **ggESDA** reduces and summarizes by classes into a structured data table with paired symbolic-valued variables, and visualizes symbolic data which are becoming increasingly important in this area.

We are committed to maintaining and developing the **ggESDA** package in the future. Future versions of the package will contain more multivariate exploratory graphical techniques and methodological improvement, including the revisions of 3D scatter plots, interactive tools, quantile representation, and so on. In order to overcome the constraints of the coordinate system of **ggESDA** in three dimensions, we will offer new geometries as well. The availability of implementation of these would be desirable for applications.

Modern statistical graphical methods for visualizing big data nowadays we encounter the large data sets in many areas, including genomics, finance records, environmental research and internet logs. In these cases, the size of data is getting larger and the structure is more complicated. Aggregation of these huge (*volume*) of data into the symbolic objects such as histograms or distributions is sometimes necessary. As a consequence, SDA is suitable in analyzing big datasets that consists of different sources (*variety*). The findings of SDA through the modern statistical graphical methods to big data will provide us with more comprehensive understanding of data and gain the *value* from it. Possibly it will speed up the scientific research.

As stated by Schweizer (1984), "Distributions are the numbers of the future. The present

is that future. We developed **graphics.SDA** to complementary the R base graphics package for symbolic data analysis, especially for interval-valued data. So far, we have seen several newly implementations of traditional graphics to symbolic data. In order to reveal structural information, the **seriation** (arrange all objects in a set in a linear order) of the objects/variables is quite important. Take the index plot of the face data as an example, one can rearrange the observations by its ranges of intervals of that variable. Hahsler, Hornik, Buchta (2008) developed an R Package, namely **seriation**, with various seriation methods and validation indices which is quite useful for ordering the numerical data table. The idea can be applied to the interval-valued and histogram-valued data after some suitable modifications.

SDA gives answers to big data and complex data challenges as big data can be reduced and summarized by classes and as complex data with multiple unstructured data tables and unpaired variables can be transformed into a structured data table with paired symbolic-valued variables. Graphics to symbolic data are becoming increasingly important in this era.

Most recently, *Big Data* has become a hot issue in the interdisciplinary researches and applications due to the popularization of the network communities and the cloud techniques. As described by Wikipedia: "Big data is an all-encompassing term for any collection of data sets so large and complex that it becomes difficult to process them using traditional data processing applications. . . . The challenges include analysis, capture, curation, search, sharing, storage, transfer, visualization, and privacy violations." It is common to use 3Vs (Laney, 2001) for describing the characteristics of big data: volume, variety and velocity. In 2012, Gartner (Laney, 2012) updated the definition: "Big data is high volume, high velocity, and/or high variety information assets that require new forms of processing to enable enhanced decision making, insight discovery and process optimization." The big data problems give rise to many statistical and technical issues (Jagadish *et al.*, 2014) to be studied and solved (Fan, Han and Liu, 2014; Chen, Mao and Liu, 2014).

Big data failed almost of the traditional graphical techniques of EDA. The statisticians are facing the challenges of analyzing the *big data* that are gathered rapidly from diverse resources with complex types. The newly efficient statistical and technical methods are needed to be developed for analyzing such data. PI believes that SDA has great capacity for big data. Due to the huge volumes of data, the big data failed almost of the traditional graphical techniques of EDA. The possible steps to conquer this issue is to summarize the dataset into the symbolic objects to a more manageable size for subsequent analysis. It raised the questions to be solved: how to display and explore these symbolic data effectively.

To do the quick and initial exploration for big data and to avoid the problems of memory and storage, computational cost and unstructured data, one possible solution is to summarize it into a moderate sized data of the form of interval-valued, histogram-valued or distributional-valued data. Then the proposed ESDA approach can be applied to these generated symbolic data without difficulties. Due to the display space required is limited, some graphical techniques described in Unwin, Theus, and Hofmann (2006) such as the smoothing and color smearing might be adopted to increase the visibility of plots.

Present the big data with various sources into symbolic objects. Big data is everywhere with many different ways, structured or non-structured. For examples, the vast amount of different types of data was gathered from archives of scanned documents, media data (images, vides, and audio), business apps, social media (Twitter, LinkedIn, Facebook etc) and sensor data.

Big data has become too complex and too dynamic to be able to process. How to convert them into the symbolic objects such as intervals, histograms and distributions is a big challenge. Currently, there is no standard rule for the conversion. Therefore, some pilot studies are needed to try.

EDA and data visualization are important for analyzing data. Data visualization help users to explore hidden structures in data. Analyzing data by numerical statistics without graphical exploration could be misleading about the data structure. Accordingly, development of the EDA and data visualization tool for exploring symbolic data is quite important. As noted in Borcard, Gillet, and Legendre (2011): "EDA is often neglected by people who are eager to jump to more sophisticated analyses. It should have an important place."

**why ggInterval** In order to implement the `symbolic_tbl` class data and the `symbolic_interval` class variables in **ggplot2**. It encounters some difficult issues to overcome. Specifically, **ggplot2** makes no consideration on what is required for a complex data to remain valid. We attempt to demonstrate this by using the `facedata` with the interval-valued variables input as following:

```
R> ggplot(facedata, aes(x = AD))
```

```
Error: `vec_math.symbolic_interval()` not implemented.
```

To escape the problem, two design thinking can be considered. One is using a new geometric to compatible the kind of variables. But the overloading in **ggplot2** internal and the extension with other layers is difficult to implement. The other improvement that has been the syntax of **ggESDA** is to generate the figure directly with the beginning command `ggInterval_` and the graphical name subsequently.

```
R> # (1) Difficult to implement the symbolic_interval class
R> ggplot(facedata) + geom_I(aes(x = AD))
R> # (2) Easy to implement the symbolic_interval class
R> ggInterval_foo(facedata, aes(x = AD))
```

Furthermore, unlike **ggplot** with a certain format for the data, the SDA packages in R have been full of the data form itself. There is no standard to stipulate this at all. So, the latter programming syntax may be better to generalize the data to the complex form and benefit to the future maintenance and extension.

## Computational details

The results in this paper were obtained using R 4.1.2 (R Core Team 2021). R itself and all packages used are available from the CRAN at <https://CRAN.R-project.org/>. The **ggESDA** is now available from the Comprehensive R Archive Network (CRAN) at <https://cran.r-project.org/web/packages/ggESDA/index.html>. All reference manual documented by exported functions and introduction vignettes can also be download here.

## Acknowledgments

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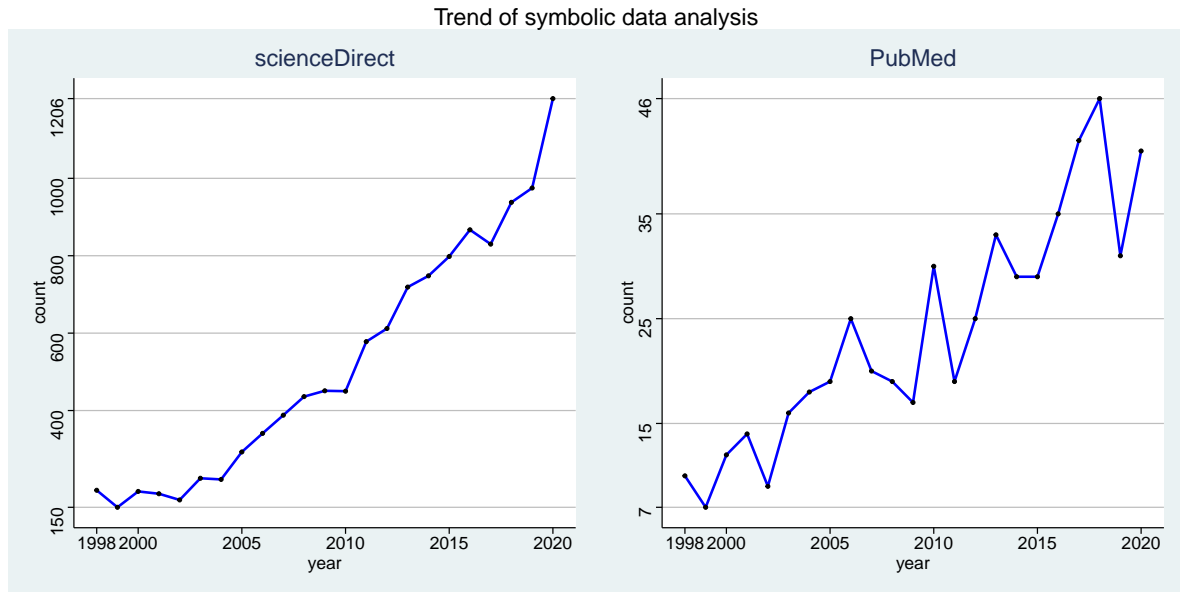


Figure 1: The number of "symbolic data analysis" or "interval-valued data" related articles in researches and applications according to PubMed and ScienceDirect online database over time from 1998 to 2020.

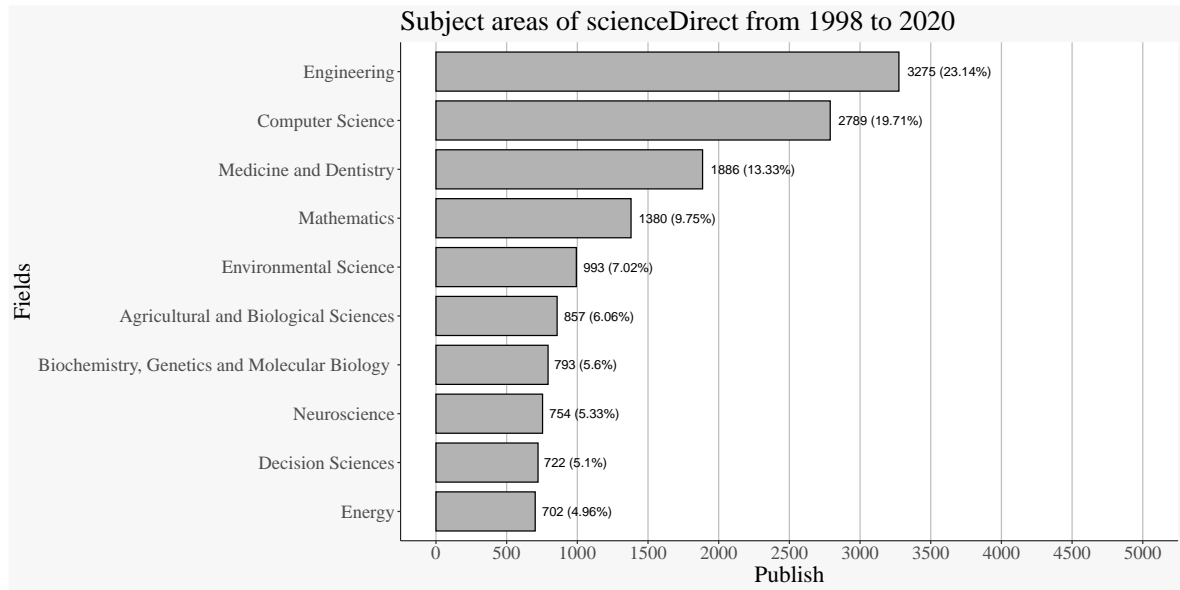


Figure 2: Top 10 researches and applications domains for SDA or interval-valued data (ScienceDirect) from 1998 to 2020

## A. Figures

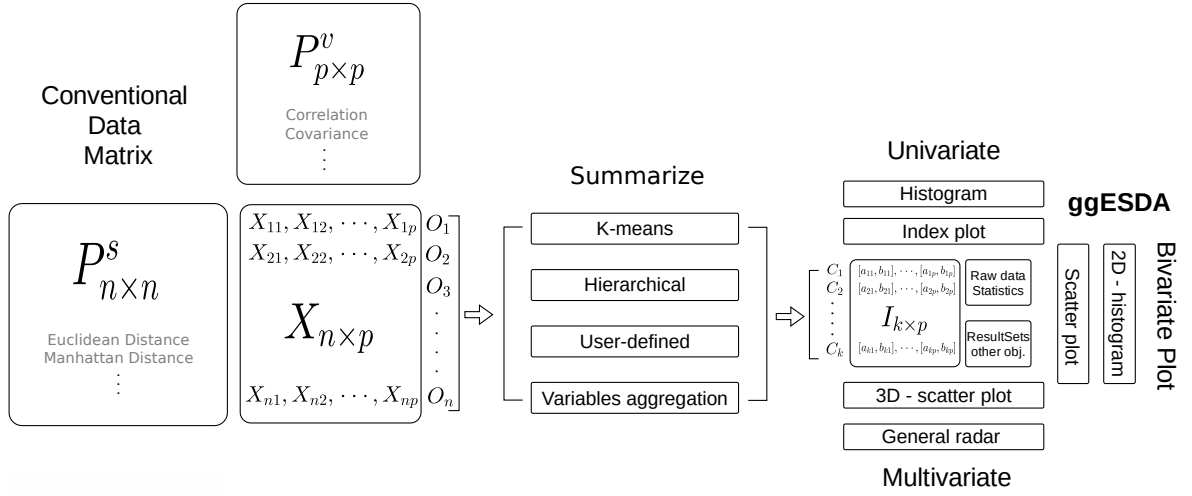


Figure 3: Package Structure and Diagram for the Transformation Flow

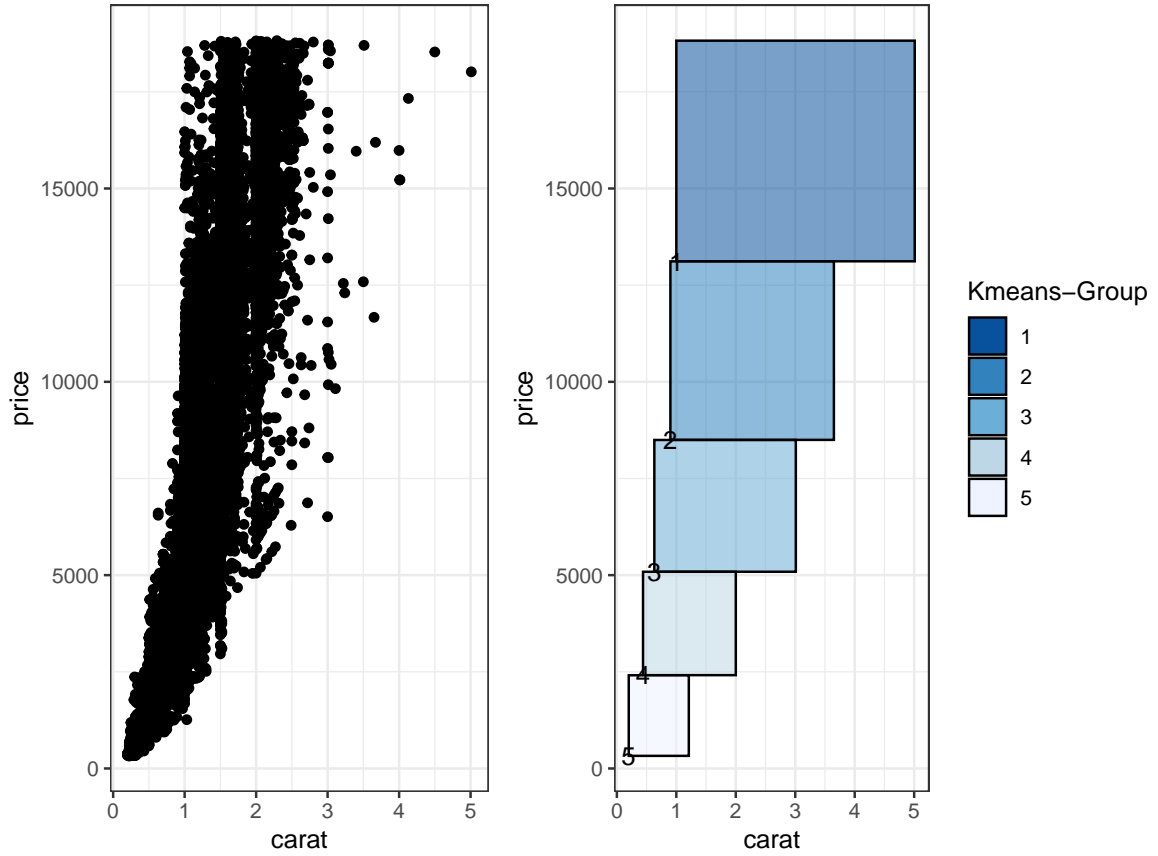


Figure 4: Compare classical data and symbolic data.

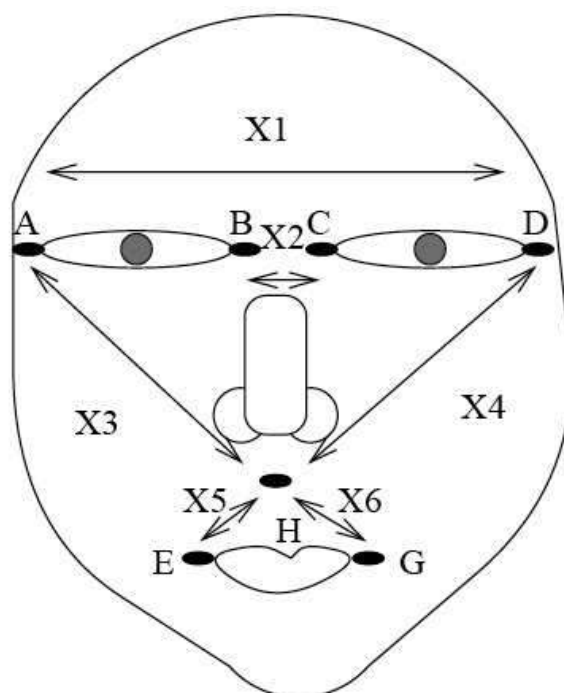


Figure 5: The six face measurements for the face recognition dataset (Douzal-Chouakria, Billard and Diday, 2011).

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 URL: <http://www.hmwu.idv.tw>

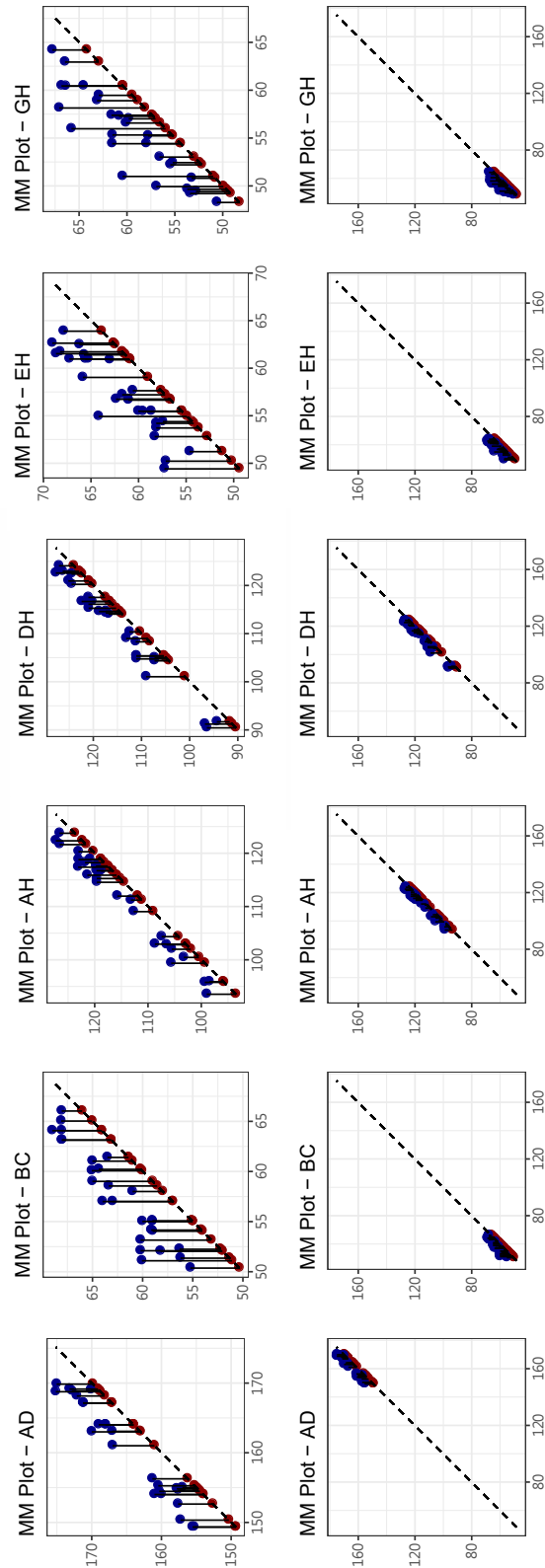


Figure 6: The min-max plot. The figure in the top six conditions the axis in their own variables, whereas the bottom six conditions it in all variables.

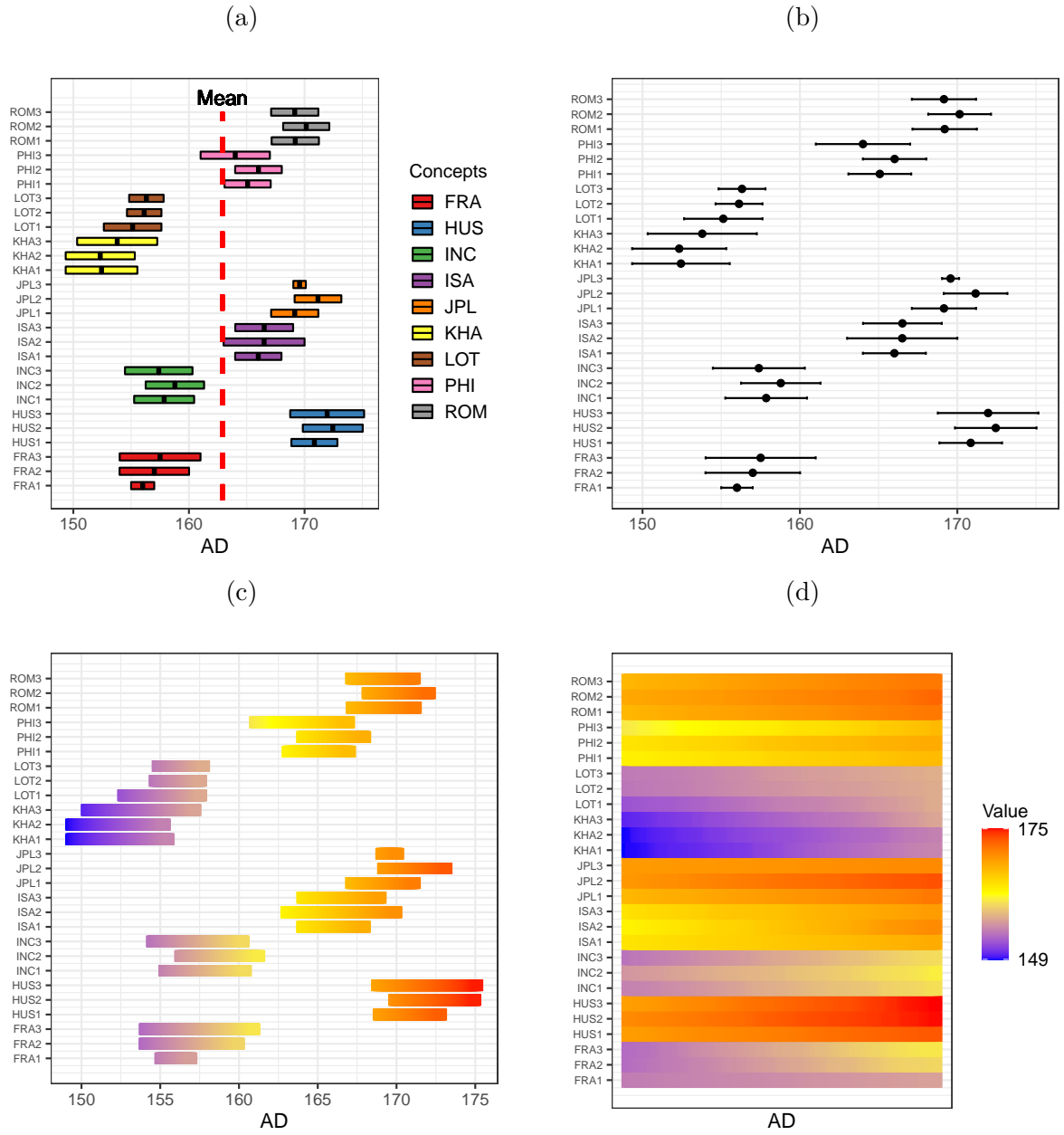


Figure 7: Four kinds of index plot for the variable AD. (a) Index plot, (b) Index plot with different group and common mean, (c) Index image, and (d) Index image with extended color.

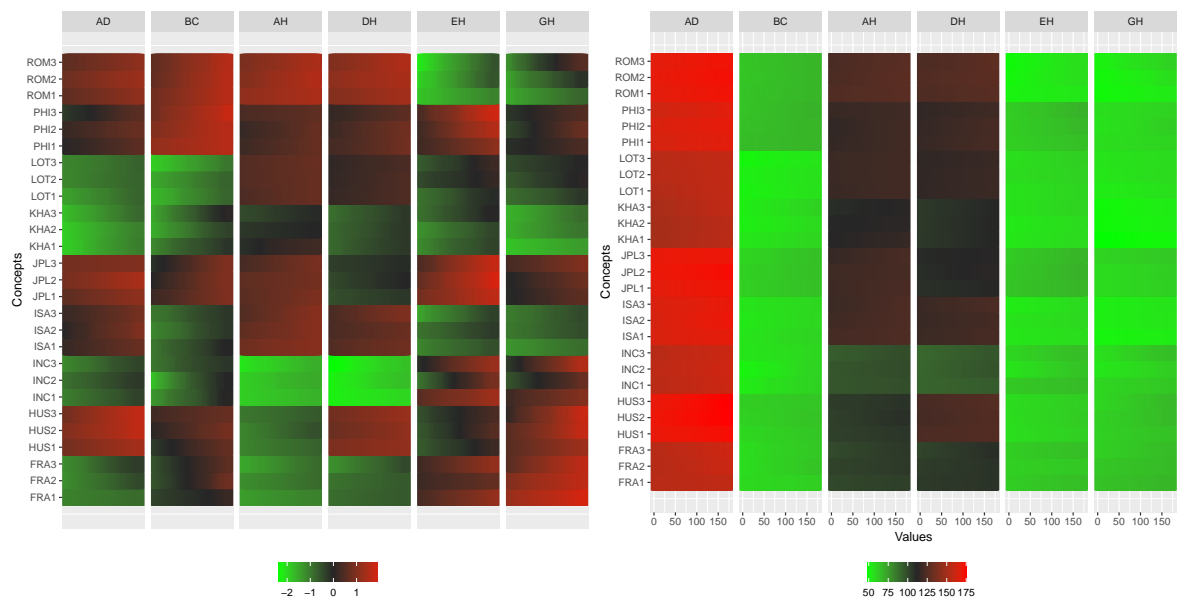


Figure 8: Index image using a heatmap type.

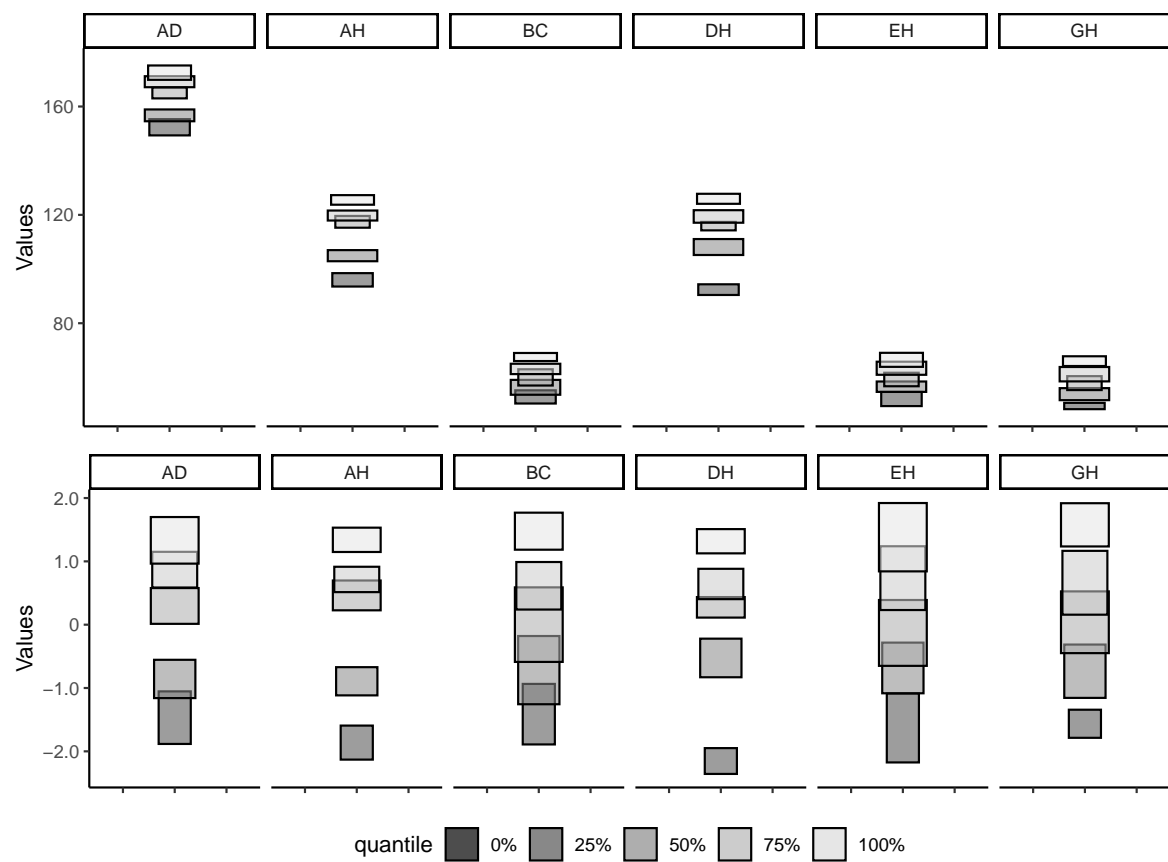


Figure 9: The side-by-side box plot. The figure in the top panel is the raw value of its variables, where the bottom panel is to standardize each variables.

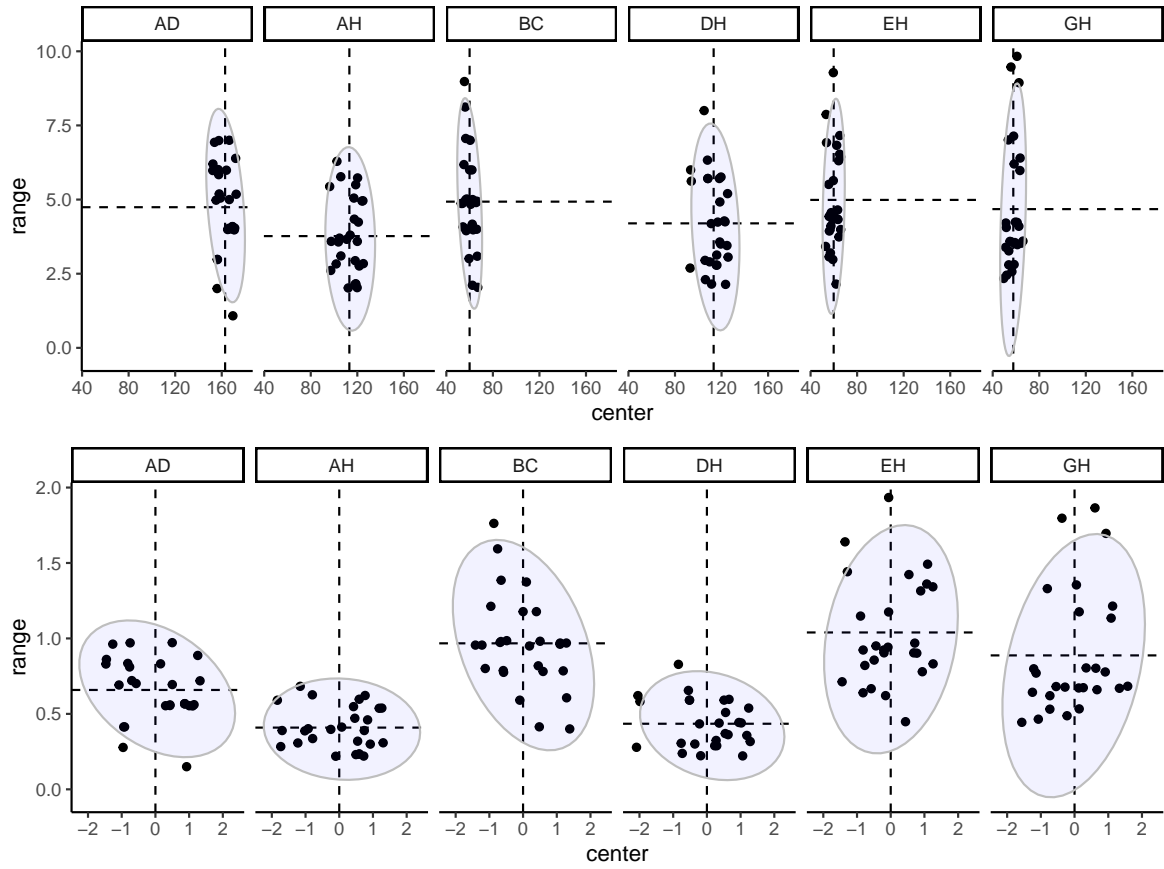


Figure 10: The center-range plot.

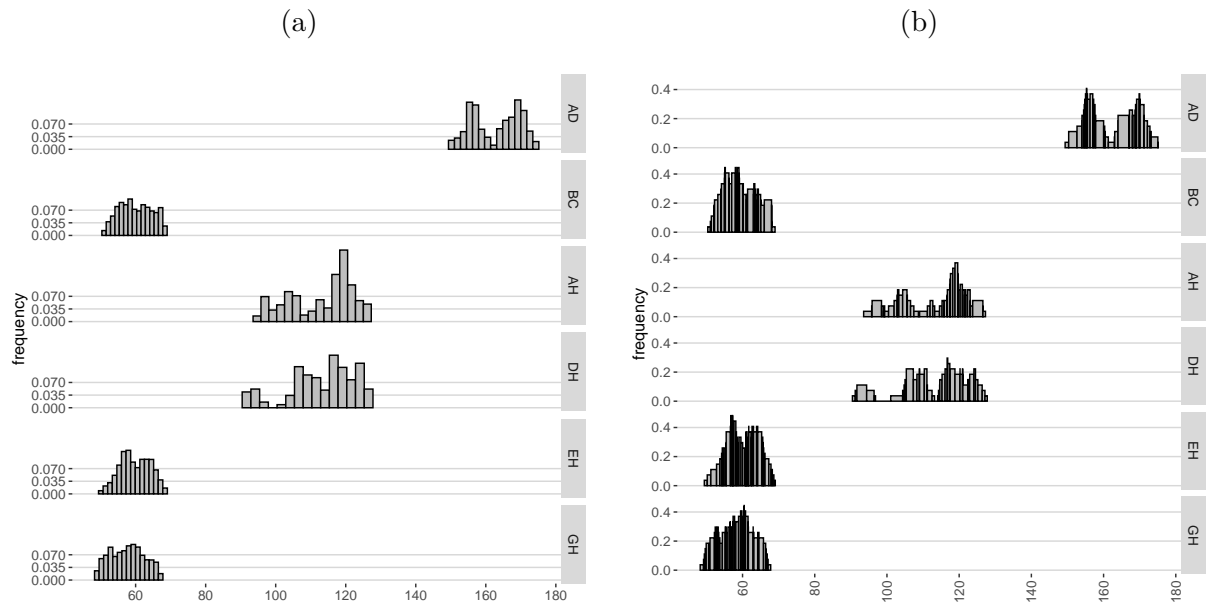


Figure 11: (a) Equidistant-bin histogram. (b) Non-equidistant-bin histogram.



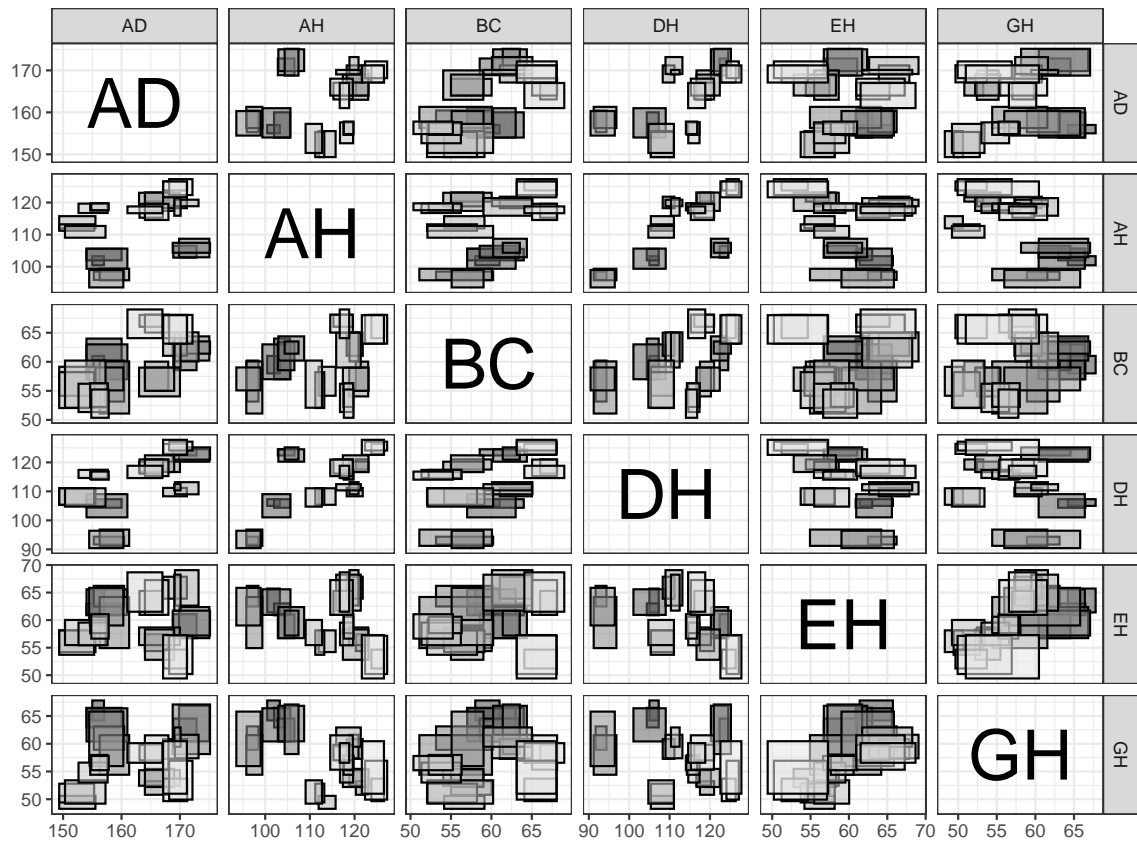


Figure 12: Scatter matrix

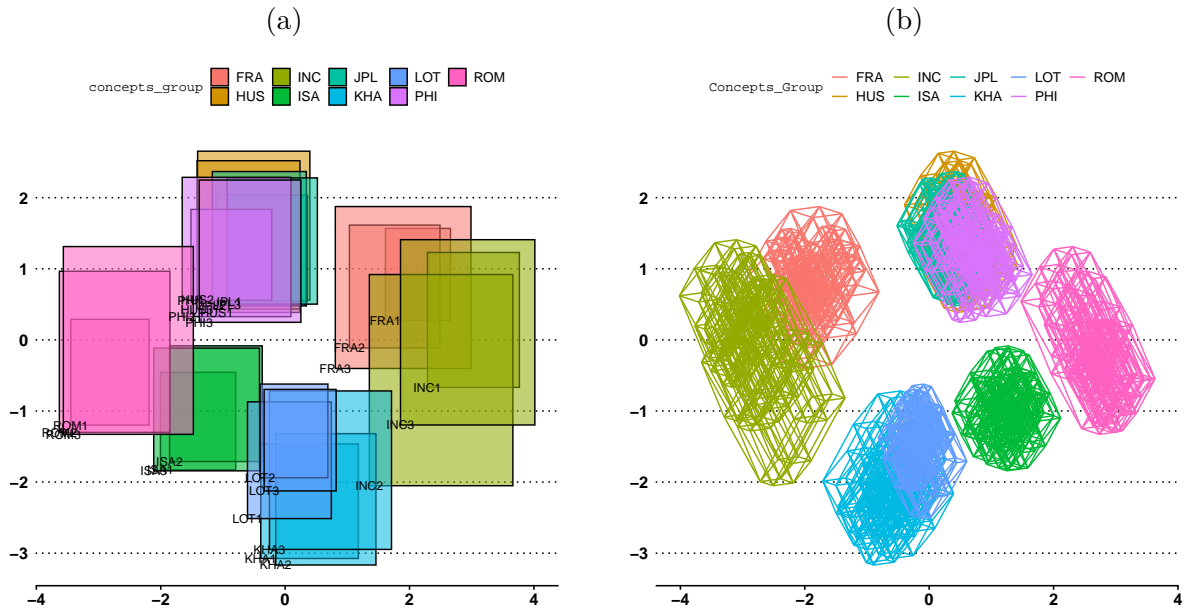


Figure 13: Two kinds of PCA for interval-valued data, where the XY-axis are represent the first principal component and the second principal component, respectively. (Vertex-PCA (MCAR)). (b) Vertex-PCA (Polytope)

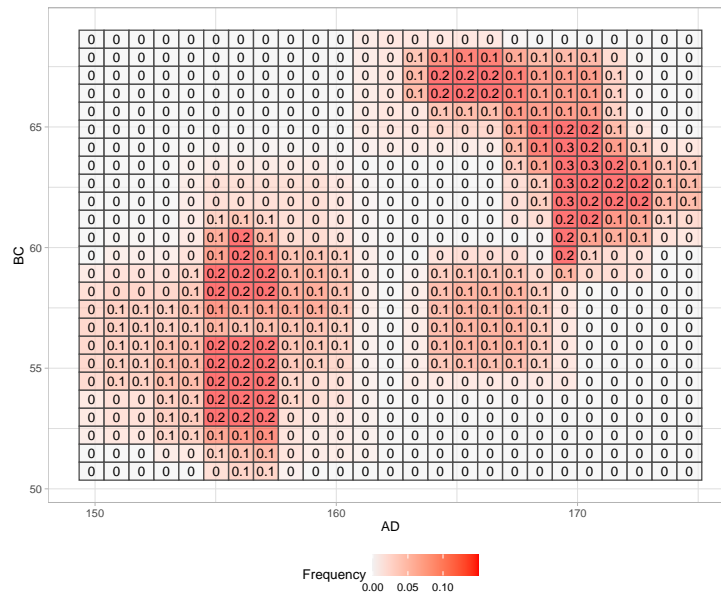


Figure 14: 2D histogram.

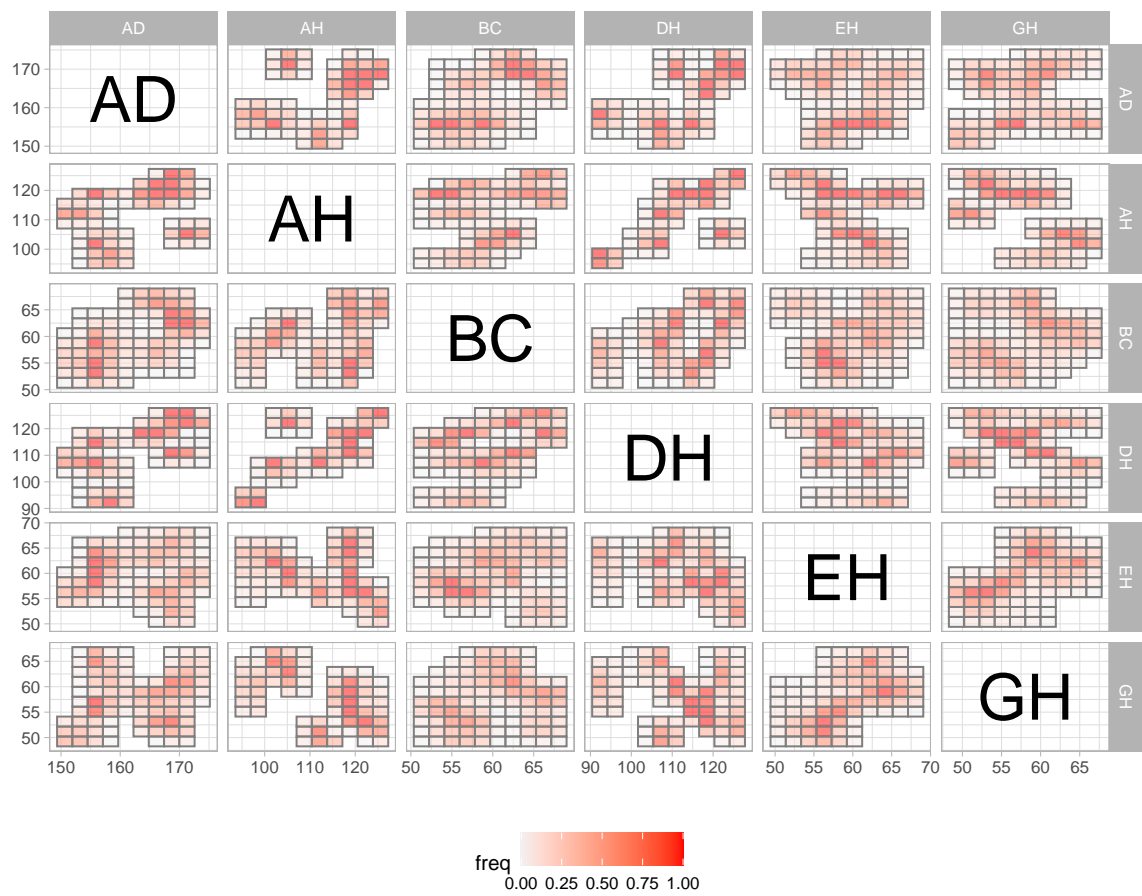


Figure 15: 2D histogram matrix.

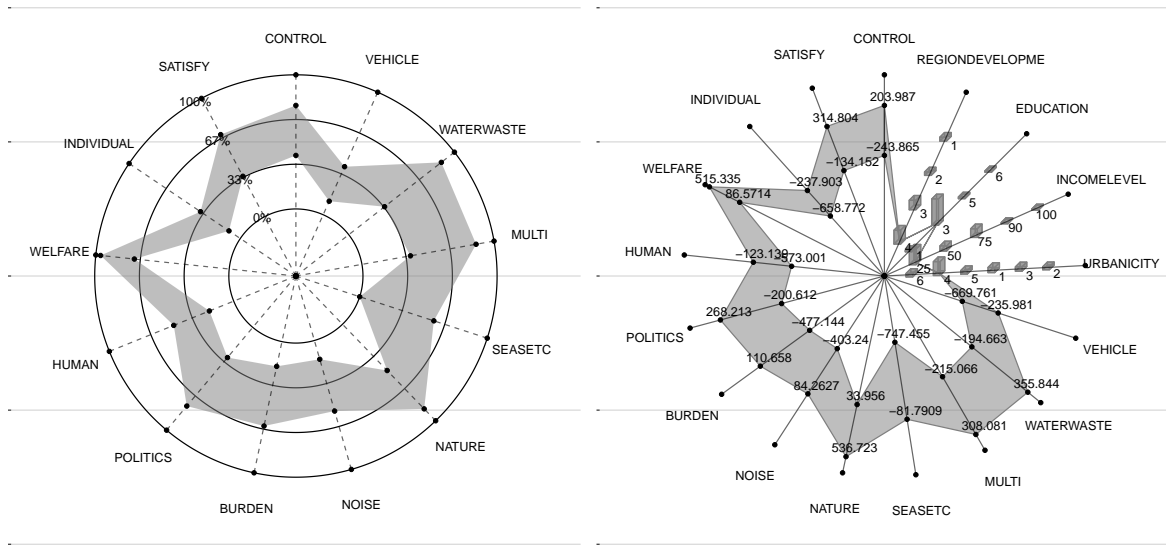


Figure 16: The variety of typical radar plot which is able to contain interval-valued and modal multi-valued variables.

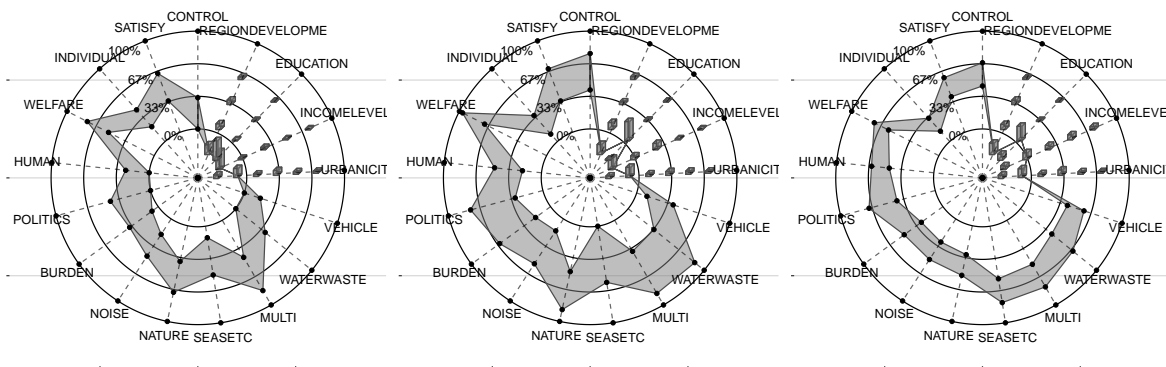


Figure 17: Compare three distinct observations by multiple variables.

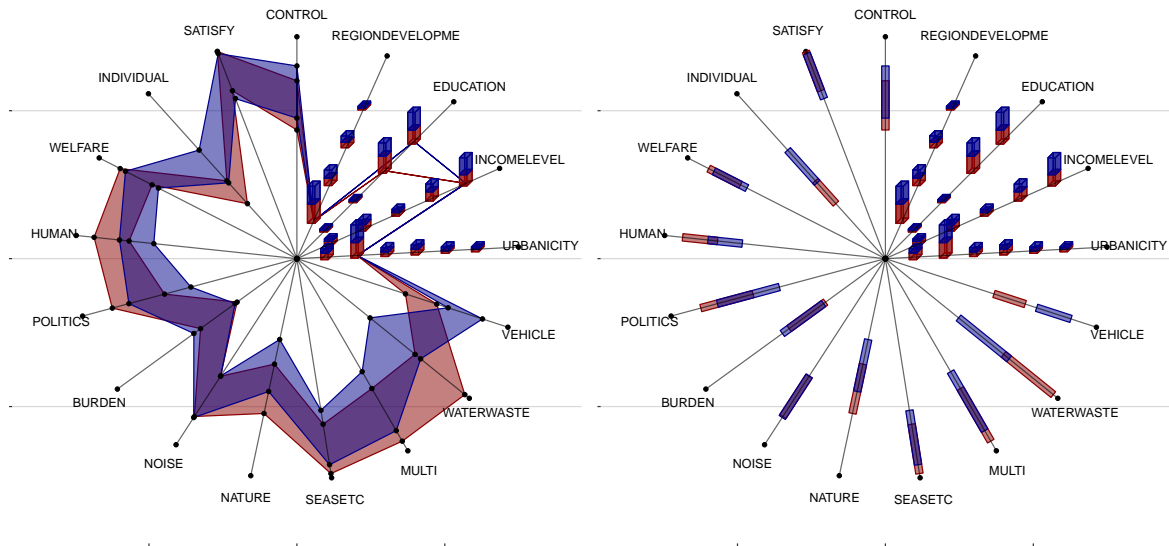


Figure 18: Compare the distinct observations in the same figure by color mapping.

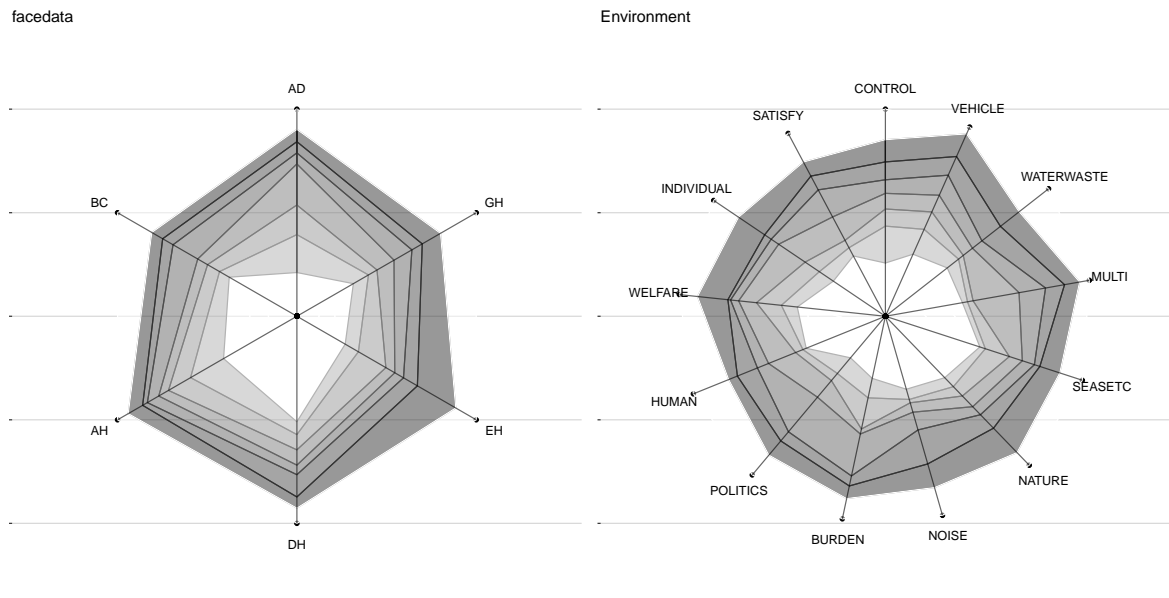


Figure 19: Quantile for two symbolic datasets.

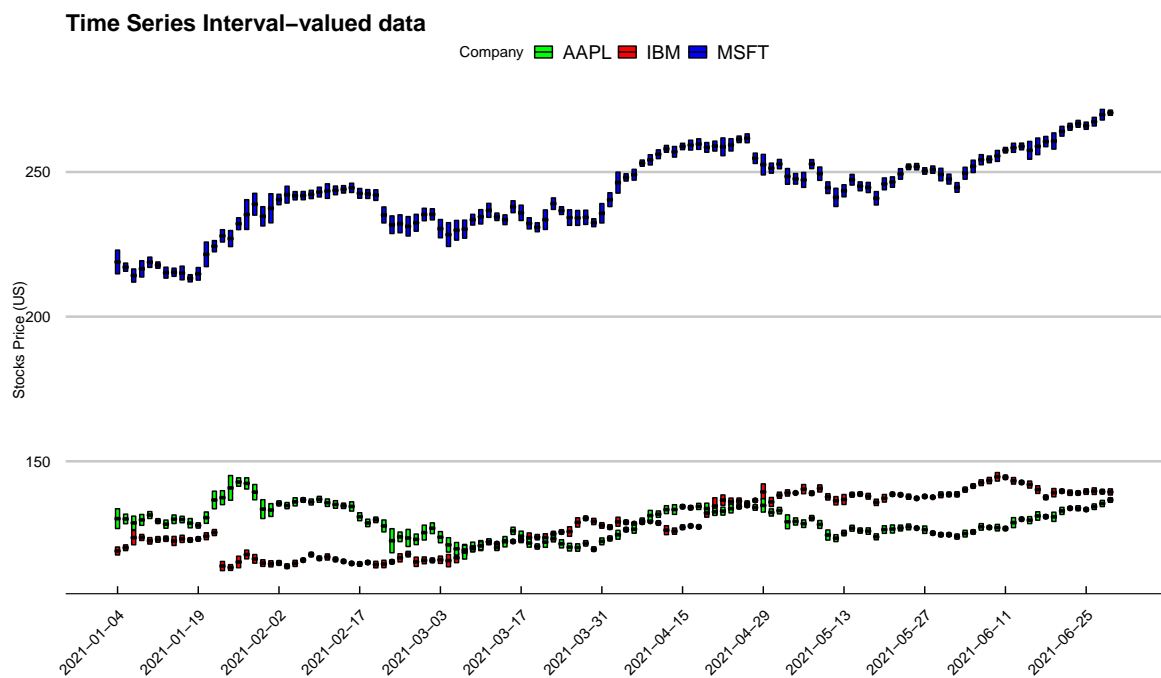


Figure 20:

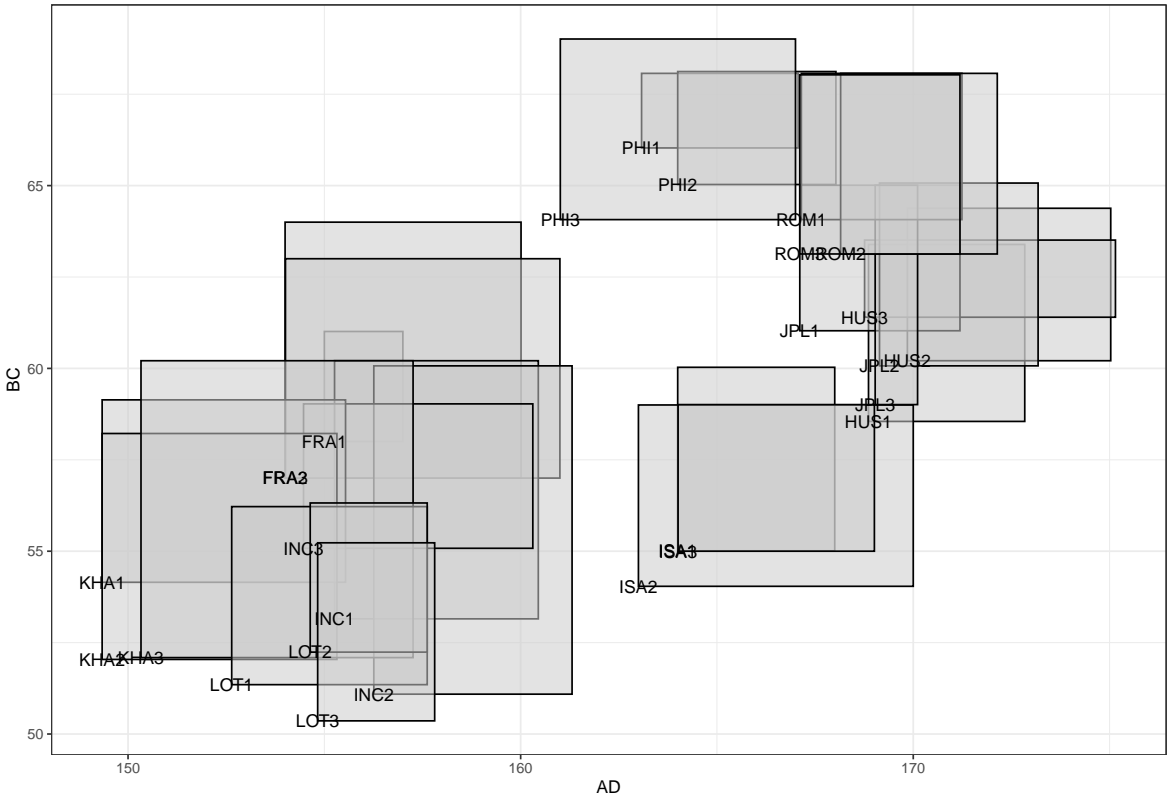


Figure 21:

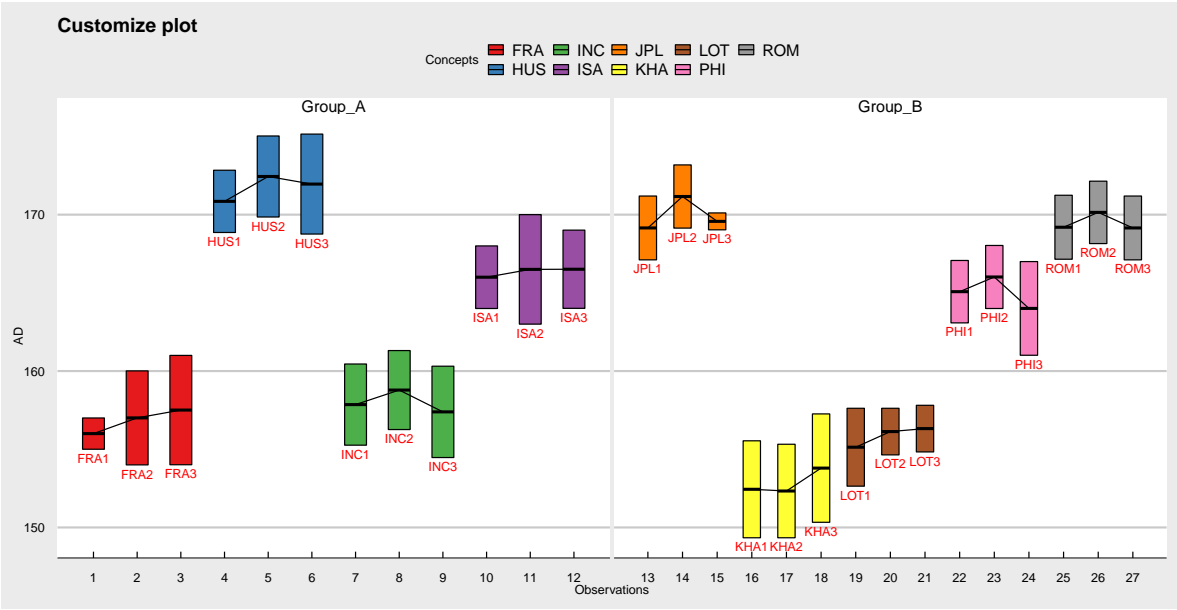


Figure 22:

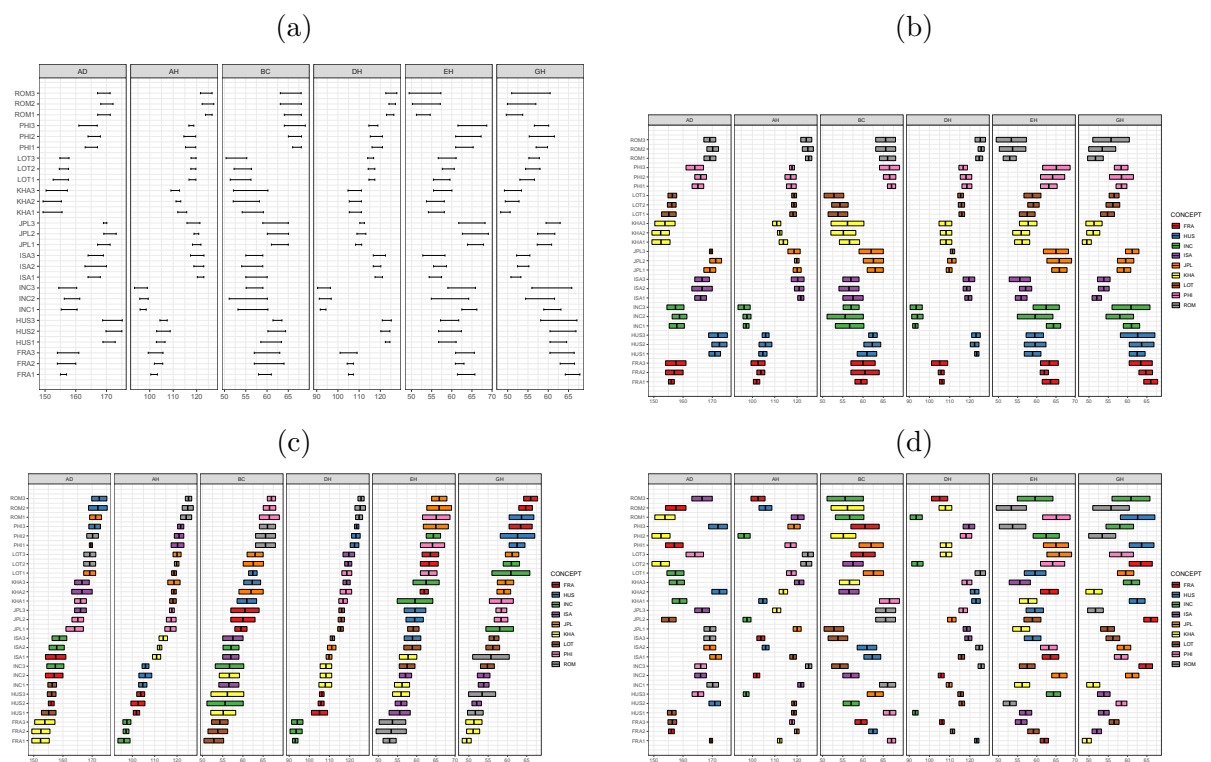


Figure 23: