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# BiplotGUI

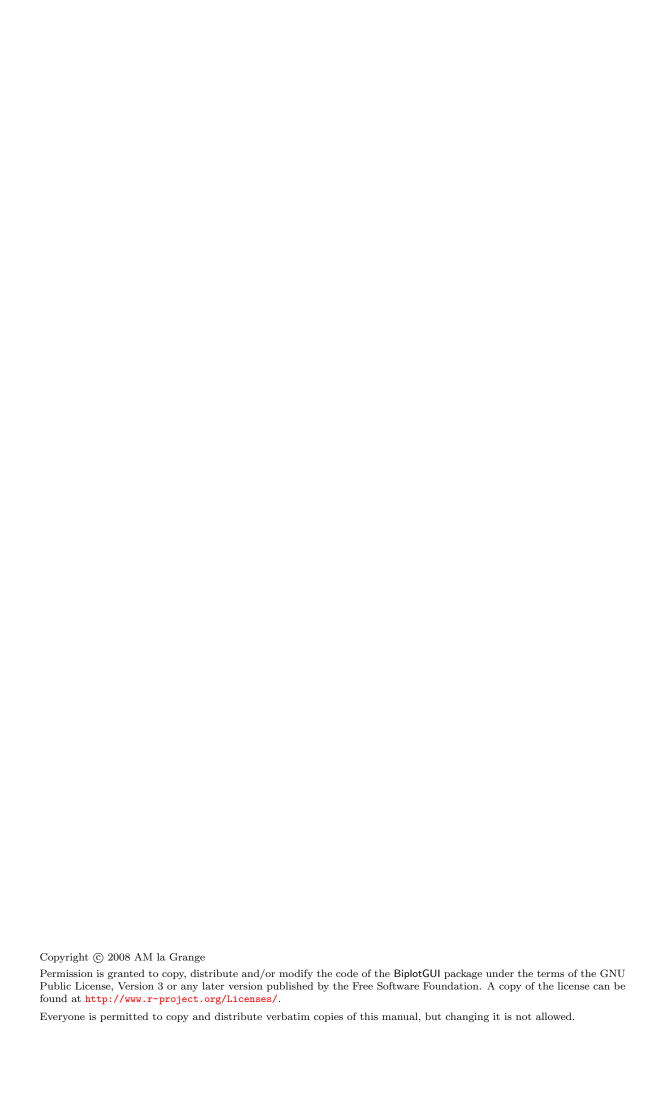
Interactive Biplots in R

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Version 0.0-2

 $4~{\rm August}~2008$ 



### **Preface**

This manual documents the features of version 0.0-2 of the BiplotGUI package, the first release on CRAN. BiplotGUI makes it easy for users to construct and interact with biplots in R.

Chapter 1 gives an overview of biplots and introduces the new package. The setup of the package and some known issues are also discussed. Chapter 2 introduces the main features of the package through the exploration of three data sets, while all the features of the graphical user interface (GUI) are documented in full in Chapter 3. An appendix summarises the history of the package.

While the statistical output provided by the BiplotGUI has been checked for accuracy, the GUI itself occassionally gives console warnings or even crashes. It's quick to simply restart the GUI. If the behaviour is replicable, please contact me at the address below with the details. Otherwise, visit <a href="http://biplotgui.r-forge.r-project.org/">http://biplotgui.r-forge.r-project.org/</a> for more information, for help, to report bugs, or to contribute to the package.

The BiplotGUI package was written as part of my Masters degree at Stellenbosch University in South Africa. I would like to thank my two supervisors, Prof Niël le Roux and Dr Sugnet Gardner-Lubbe, for their continued support. I would also like to thank Prof Patrick Groenen, whose suggestions inadvertently gave rise to the project.

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# List of Keyboard Shortcuts

A	Shows points determined by PCO.
В	Shows points determined by metric MDS (identity transformation).
$\mathbf{C}$	Shows points determined by non-metric MDS (monotone regression).
D	Shows points determined by semi-metric MDS (monotone spline transformation).
0	No biplot axes.
1	Constructs a PCA biplot.
2	Constructs a covariance/correlation biplot.
3	Constructs a CVA biplot.
4	Shows regression biplot axes.
5	Shows Procrustes biplot axes.
6	Shows circular non-linear biplot axes.
Ctrl+N	Interpolates a new sample.
Ctrl+L	Clears all additional descriptors from the biplot.
F11	Shows the currently displayed biplot region in an external window.
F12	Shows the currently displayed biplot in 3D in an external window.
Ctrl++	Shows the next set of legend entries.
Ctrl+-	Shows the previous set of legend entries.
Ctrl+G	Allows many graphical parameters to be set for the different groups of points.
Ctrl+A	Allows many graphical parameters to be set for the different biplot axes.
Ctrl+R	Reverts all the graphical parameters to their default values.
Ctrl+S	Saves the currently displayed biplot region in the currently selected file format.
Ctrl+C	Copies the currently displayed biplot region to the clipboard.
Ctrl+P	Prints the currently displayed biplot region.
F1	Enables pop-up help messages for the components of the main GUI window.

## List of Abbreviations

AOD analysis of distance

cv canonical variate

CVA canonical variate analysis
GDP gross domestic product
GUI graphical user interface
IM iterative majorisation
MDS multidimensional scaling

PC principal component

PCA principal component analysis
PCO principal coordinates analysis

PPP purchasing price parity
RAE relative absolute error

## Chapter 1

## Introduction

### 1.1 Overview

In this section we give a brief overview of biplots, existing biplot software, and the statistical programming language and environment R. In Section 1.2 we set out the main aims of the BiplotGUI package, while its most important features are showcased in Chapter 2 through the exploration of three data sets. The present version of the package is 0.0-2.

The style of the article is intentionally non-mathematical. However, detailed references are provided for those who wish to gain a fuller understanding of the underlying theory. This approach allows the focus to lie firmly with the package and its features.

### 1.1.1 Biplots

Introduced by Gabriel (1971), the biplot is described by Gower and Hand (1996) in their authoritative monograph as the multivariate analogue of the ordinary scatter plot. As such, biplots are representations of multivariate data in which information on both samples and variables is given simultaneously in two or three dimensions: samples (observations) are represented as points, while variables are represented as labelled and calibrated axes. The axes are either linear and oblique, or non-linear. This new approach to biplots differs from the more traditional approach in which samples and variables are represented as points and/or uncalibrated vectors.

Some dimension-reduction technique is typically used to represent the samples as points, often principal component analysis (PCA) (Pearson, 1901; Hotelling, 1933) or canonical variate analysis (CVA) (Hotelling, 1935, 1936). More generally, scaling techniques such as principal coordinates analysis (PCO) (Torgerson, 1952; Gower, 1966) or metric or non-metric multidimensional scaling (MDS) (Kruskal, 1964a,b; Sammon, 1969) are used. Jolliffe (2002) dedicates a monograph to PCA, while Krzanowski (2000) covers general multivariate topics in tune with the spirit of biplots. Cox and Cox (2001) and Borg and Groenen (2005) are standard references for scaling techniques.

The placement of the axes depends partly on the mechanism used in the placement of the points. The PCA biplot provides linear axes for points placed by PCA (Gower and Hand, 1996, Chapter 2); similarly the CVA biplot provides linear axes for points placed by CVA (Gabriel, 1972; Gower and Hand, 1996, Chapter 5). The regression biplot (Gower and Hand, 1996, Chapter 3) gives approximate linear axes for any ordination of points. So too does the Procrustes biplot (Gower and Hand, 1996, Chapter 3). The regression and Procrustes biplots correspond to the PCA biplot for points determined by PCO based on Pythagorean

<sup>&</sup>lt;sup>1</sup>This chapter and the next are based on work submitted for publication by La Grange, Le Roux and Gardner-Lubbe.

distance. The covariance biplot (Greenacre, 1984; Underhill, 1990) adjusts the points and axes of the PCA biplot so that the cosines of the angles between the axes approximate the correlations between the corresponding variables. The correlation biplot is similar, except that the variables are standardised to have unit variances.

The placement of the axes may also depend on how they are to be used. *Predictive* axes are positioned and calibrated so that the orthogonal projection of a point onto an axis 'predicts' as best as is graphically possible the value of the corresponding sample on the corresponding variable. *Interpolative* axes, on the other hand, are positioned and calibrated so that a new sample may be added to an existing configuration of points at the most appropriate position graphically possible. Interpolation can be either by the centroid or vector sum of the positions on the axes corresponding to the respective variable values of the new sample.

For additive inter-sample dissimilarities (Gower and Hand, 1996, p. 105), biplots with non-linear axes (or trajectories) may be constructed for points determined by PCO. The PCO solution itself requires the inter-sample dissimilarities to be Euclidean-embeddable (Gower, 1982); dissimilarity measures for which this is the case are discussed by Gower and Legendre (1986). Non-linear predictive axes may make use of circular projection (Gower and Hand, 1996, Chapter 6), while non-linear interpolative axes (Gower and Harding, 1988; Gower and Hand, 1996, Chapter 6) are used in the same way as the linear variety. Non-linear biplots are often most useful to gauge what is otherwise approximated by linear biplots.

While very many examples of biplots of the traditional approach may be found in the literature, there are fewer examples of biplots of the new approach. An important reason has been the lack of software, as is discussed in Sections 1.1.2 and 1.1.3. The value of biplots of the new approach, however, has often been demonstrated. In an easy to read introduction, for example, Le Roux and Gardner (2005) cite and showcase many examples of the uses of linear biplots, from such diverse fields as archaeology, agriculture, antiques, education, financial management, mineralogy and process control. Other recent fields of application include cephalometry (Naidoo, Harris, Swanevelder and Lombard, 2006), chemistry (Alves, Cunha, Amaral, Pereira and Oliveira, 2005) and mineralogy (Jemwa and Aldrich, 2006). Examples of non-linear biplots may be found in Gower and Harding (1988), Gower and Hand (1996) and Gower and Ngouenet (2005).

Given the ubiquity of multivariate data and the usefulness of biplots in describing such data, there is still much scope for the further popularisation of the technique.

#### 1.1.2 Other biplot software

Many statistical packages can be used to produce at least the simplest of biplots of the traditional approach. These include the major statistical packages Minitab (Minitab Inc, 2007), SPSS (SPSS Inc, 2007), STATA (StataCorp LP, 2007), STATISTICA (StatSoft Inc, 2007) and various products from SAS (2008). However, functionality is often limited, and results hard to obtain. Greater functionality is provided by the three dedicated biplot programs XLS-Biplot (Udina, 2005a,b), GGEBiplot (Yan and Kang, 2006) and BiPlot (Lipkovich and Smith, 2002a,b). XLS-Biplot is based on XLisp-Stat (Tierney, 1990) and has many useful features including a related web-server which can be used to construct biplots online. GGEbiplot is aimed mainly at agronomists, crop scientists and geneticists. It supplements the book by Yan and Kang (2003). BiPlot is an add-on for Excel, and although therefore potentially widely useful, it unfortunately has some minor but serious shortcomings (see Udina, 2005b).

The Genstat package (VSN International Ltd, 2007) can be used to calculate the coordinates of the elements of a biplot. These can then be drawn using a procedure from an add-on library. Other packages, offering some traditional biplot functionality, include Manet (Hofmann, 2000), for Macintosh only, and ViSta (Young, 2001). Some packages are aimed at ecologists—brodgar (Highland Statistics Ltd, 2007) with R, Canoco (Plant Research International, 2002) with

CanoDraw (Smilauer, 2003), MVSP (Kovach Computing Services, 2008) and PC-ORD (MjM Software Design, 2007)—while the Excel add-on BrandMap (WRC Research Systems Inc, 2007) is aimed at marketers. All the software mentioned are for purchase, except XLS-Biplot, BiPlot, Manet and ViSta which are available free of charge. So too is R.

#### **1.1.3** R

R (R Development Core Team, 2008) is a free statistical programming language and environment capable of producing high-quality graphics. Initiated by Ihaka and Gentleman (1996), it has become 'the *de facto* standard for statistical computing' (Greenacre, 2007, p. 213). It is an open-source implementation of the S programming language, available for download for all the major platforms from the R Project homepage at <a href="http://www.r-project.org">http://www.r-project.org</a>. The R core is updated regularly with minor version revisions released roughly every six months. The current version (as of June 2008) is R 2.7.1. Updates are relatively painless. R is easily extensible: a large number of user-written packages is available for download from repositories such as CRAN and BioConductor. These repositories can be accessed via the R Project homepage. As R has increased in popularity, so too has the number of books devoted to it. Recent general-topic books on R include Braun and Murdoch (2007), Chambers (2007) and Spector (2008). The book by Murrel (2005) deals specifically with graphics in R. Many more resources are freely available from the R Project homepage.

As far as biplots are concerned, the biplot method in R can be used to produce two variations of Gabriel's (1971) classical biplot. The classical biplot is most similar to the covariance/correlation biplot described earlier. Packages with support for traditional biplots include ade4 (Dray and Dufour, 2007, 2008), ade4TkGUI (Thioulouse and Dray, 2007a,b) and vegan (Oksanen, Kindt, Legendre, O'Hara, Simpson, Stevens and Wagner, 2008). In addition, the calibrate package (Graffelman, 2007) can be used to calibrate both scatter plot and biplot axes as described by Graffelman and van Eeuwijk (2005). In general, however, these calibrations do not correspond to those of Gower and Hand (1996).

As opposed to the many solutions for biplots of the traditional approach listed in this section and in the previous one, software for biplots of the new approach has not been readily available. To produce biplots of the new approach, users have had to do their own programming in a suitable environment (for example, Gardner, 2001). To many potential users, such a task represents a major obstacle.

### 1.2 A new package

The primary aim with the BiplotGUI package is to make it easy to construct biplots of the kind advocated by Gower and Hand (1996) – biplots in which samples are represented as points and variables are represented as calibrated axes. The package goes beyond this, however, allowing users to interact with the data through the biplots which are produced. Naturally, the graphical output should be of a high quality and easily customisable. Its characteristics make R the ideal environment for the development of such a package.

The main features of the package can be summarised as follows:

- The package has a graphical user interface (GUI), making it easy to use even to the those new to R.
- Different types of biplots are supported: PCA, covariance/correlation, CVA, regression, Procrustes and circular non-linear.
- The points in regression and Procrustes biplots are determined by a separate scaling technique: PCO and various forms of MDS are available. In circular non-linear biplots, the points are fixed by PCO.

- Various distance metrics and data transformations are available.
- New samples can be interpolated onto existing biplots, and sample group means can be incorporated.
- Additional descriptors can be shown: convex hulls, alpha-bags, point densitites and classification regions.
- Various diagnostic graphs are easily accessible: graphs of convergence; point, group and axis predictivities; and Shepard diagrams.
- The package is highly interactive with support for dynamic variable value prediction, point and axis drag and drop, zooming, and the exploration of biplots in 3D.
- Biplots can be finely tailored. Amongst the elements that can be customised are titles, labels, legends, and more than 80 well-organised graphical parameters.
- Data can be imported from R objects or Excel files. Graphs can be saved to eight file formats. An array of context-specific numerical quantities are available for export back to R for further manipulation.
- The package is free and open-source. Therefore the source code is available to all users, making it easy for others to adapt it, and to include new developments.

Many of these features are showcased in Chapter 2, and all of them are documented in Chapter 3

### 1.3 Setup

R can be downloaded from CRAN via the R Project homepage at <a href="http://www.r-project.org">http://www.r-project.org</a> for any of the three major platforms: Linux, MacOS X and Windows. BiplotGUI version 0.0-2 requires at least R 2.7.

At present, the package is intented to be run under Windows. To run, BiplotGUI requires a number of other packages to be installed.

The essential packages are

- MASS (Venables and Ripley, 2002)
- tcltk (R Development Core Team, 2008)
- tcltk2 (Grosjean, 2008)
- tkrplot (Tierney, 2008)

The first two of these should already be in place after a basic installation of R. The recommended packages are

- KernSmooth (Wand, 2008), for the estimation of point densities
- rgl (Adler and Murdoch, 2008), for the creation of dynamic 3D biplots
- vcd (Meyer, Zeileis and Hornik, 2008), for the colours to be used in the representation
  of point densities
- xlsReadWrite (Suter, Treetron and Switzerland, 2006), for the import of data from Excel

BiplotGUI runs marginally better when the R console is set to SDI mode rather than the default MDI mode.

### 1.4 Known issues

- The graphs in diagnostic tabs occassionally do not show when the tabs are opened. Click to another tab and back.
- Interpolative circular non-linear biplots do not always initialise correctly in 3D. If this happens, first display a 3D predictive circular non-linear biplot.
- Extensive use of the GUI leads to memory leaks. If the system becomes noticably slower, close the R console after saving, and re-open.

Avoid performing actions—besides changing the diagnostic tabs—while the progress bar shows the GUI to be busy.

## Chapter 2

## **Exploring**

### 2.1 Introduction

In the next two sections, the most important features of the BiplotGUI package are illustrated. This is done through the exploration of three data sets. Further features are highlighted in Section 2.5. A systematic account of all features is given in the 'Details' chapter of the package manual. This manual can be accessed via the 'Help' menu from within the package.

#### 2.2 Countries

In this section we introduce a country-comparative data set. It is used to show how the GUI may be initialised, how its features are laid out, and how it may be used to explore multivariate data using, amongst other things, PCA and regression biplots.

Table 2.1 gives measurements of eight variables for the countries with the 15 largest economies (by purchasing price parity (PPP) gross domestic product (GDP)) in 2007. These data have been derived largely from the 2007 CIA World Factbook (Agency, 2007), and are for illustrative purposes only. The variables are: PPP GDP in trillions of US dollars (GDP); HIV/Aids prevalence as a percentage of the population (HIV.Aids); life expectancy in years (Life exp.); military spending as a percentage of GDP (Mil.); oil consumption in barrels per annum per capita (Oil cons.); population in millions (Pop.); number of fixed line telephones per capita (Tel.); and percentage unemployed (Unempl.). The aim is to represent these data in two or three dimensions so that a single, multivariate visual impression may be obtained, with the calibrated biplot axes incorporating information on the original variables.

### 2.2.1 Getting started

After R and the required packages have been downloaded and installed, the BiplotGUI package itself needs to be installed (more detail is given in ??). This process needs to be performed only once. To then load the BiplotGUI into R, the following command is entered at the R prompt, followed as usual by the enter key:

```
library (BiplotGUI)
```

If the user is acquainted with R, data may be entered at the keybord or be imported into R and saved as a matrix or a data frame. The countries data have already been included in the package as a data frame, and may be viewed from within R by typing the commands

```
data (Countries)
```

at the R prompt. To initialise the GUI with the countries data, the command Biplots (Data = Countries)

Country	GDP	HIV.Aids	Life exp.	Mil.	Oil cons.	Pop.	Tel.	Unempl.
Brazil	8710	0.7	72.2	2.6	4.0	190	204.2	9.6
Canada	35370	0.3	80.3	1.1	25.1	33	622.3	6.4
China	7724	0.1	72.9	4.3	1.8	1322	278.4	4.2
France	29852	0.4	80.6	2.6	11.3	64	543.5	8.7
Germany	31941	0.1	79.0	1.5	11.7	82	657.8	7.1
India	3685	0.9	68.6	2.5	0.8	1130	44.0	7.8
Indonesia	4041	0.1	70.2	3.0	1.8	235	63.2	12.5
Italy	30199	0.5	79.9	1.8	11.8	58	430.8	7.0
Japan	33100	0.1	82.0	0.8	16.0	127	432.8	4.1
Mexico	10570	0.3	75.6	0.5	6.6	109	182.7	3.2
Russia	12350	1.1	65.9	2.7	6.5	141	283.6	6.6
S Korea	24386	0.1	77.2	2.7	16.0	49	547.8	3.3
Spain	27418	0.7	79.8	1.2	14.2	40	454.5	8.1
UK	31723	0.2	78.7	2.4	11.0	61	552.9	2.9
USA	43369	0.6	78.0	4.1	25.1	301	571.2	4.8

Table 2.1. The countries data set. Eight variables measured on the countries with the 15 largest economies (PPP GDP) in 2007; countries listed in alphabetical order.

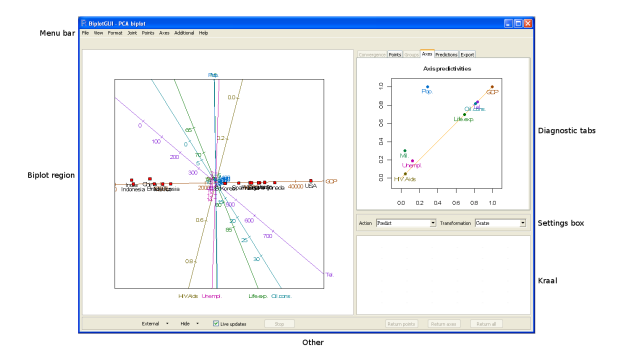
is entered. Alternatively, a data set may be saved as an Excel 1997-2003 file, with its samples as rows and its variables as columns. If the countries data were saved as an Excel file named 'Countries.xls', the call would be

Biplots (excel = "Countries.xls")

If the Excel file is not in the working directory, the full path must be given in the usual way. Either of the previous two instructions launches the GUI and no further R commands are necessary.

#### 2.2.2 The layout

- 2.1 shows the layout of the GUI after it has launched. Six regions are indicated:
  - The menu bar, in addition to the settings box, contains the most important options. The three most important drop-down menus are 'Joint', 'Points' and 'Axes'. The biplots listed under 'Joint' have both their points and axes determined according to a single, joint mechanism. The other biplots have their points determined from the 'Points' menu and their axes determined from the 'Axes' menu.
  - The biplot region is where the biplot and optional title and legend are displayed. This space is responsive to mouse clicks and motion.
  - The settings box may be used to set the action of the biplot axes, either predictive, centroid interpolative or vector sum interpolative. Various data transformations may be effected.
  - The diagnostics tabs show output related to the currently displayed biplot. The convergence tab shows a graph of convergence; the points, groups and axes tabs show context-specific graphs of goodness-of-fit for points, groups and axes, respectively; the predictions tab shows dynamically predicted variable values; while the export tab allows various objects to be exported to R.
  - The kraal is where points and axes may be kept, temporarily removing them from consideration.



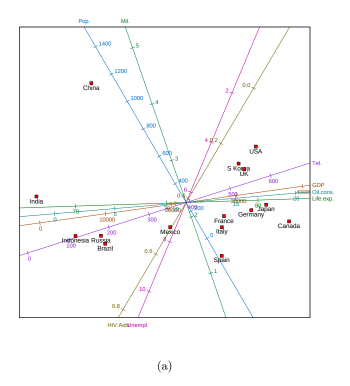
Screenshot 2.1: The BiplotGUI window as it initially appears. A predictive PCA biplot of the countries data is shown towards the left. The axis predictivities are shown top right.

• Other. The options in this section can be used to show the currently displayed biplot in an external window (in two or three dimensions), to control the biplot region or to control the kraal. While the GUI is busy, a progress bar is shown towards the left of this area.

The 'Show pop-up help' option in the 'Help' menu activates pop-up help messages which appear when the mouse cursor is hovered over the components of the main GUI window.

#### 2.2.3 A first look

The PCA biplot with predictive axes is shown by default. For the countries data, this is the biplot shown towards the left of the screenshot in 2.1. As should be the case for all biplots, a unit aspect ratio is used to ensure that distances within the biplot are properly represented. In this biplot, the points representing the countries lie ordered along a virtually straight line. In fact, the imagined line corresponds very closely to the biplot axis for GDP, and importantly, the line is almost horizontal. The reason for this becomes clear by looking at the GDP column of the countries data set. The values of GDP are orders larger than those of the other variables. Therefore, that linear combination of the variables that has the largest possible variation (the first principal component) is heavily weighted towards GDP. In effect, GDP drowns out the other variables. To avoid this, we choose the 'Centre, scale' transformation from the available transformations in the settings box. This transformation independently transforms each variable to unit variance and automatically updates the biplot to the one shown in the top left panel of Figure 2.1. Irrespective of the chosen transformation, however, the axes are always calibrated in terms of the original variable values. The first principal component in this new figure still ranks the countries from least to most wealthy, in some more complicated sense. The developed countries of the West, together with Japan and newly-industrialised South



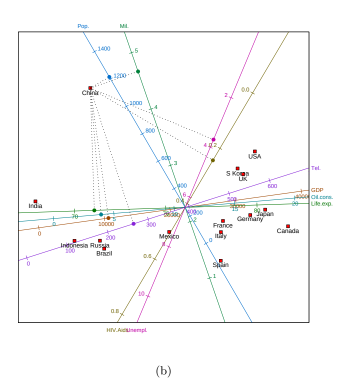
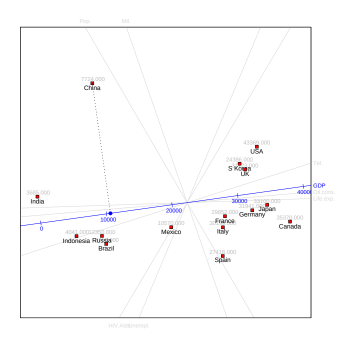


Figure 2.1. In (a), a predictive PCA biplot of the centred, scaled countries data. In (b), a predictive PCA biplot of the centred, scaled countries data with China projected onto all the biplot axes. In (c), a predictive PCA biplot of the centred, scaled countries data with GDP highlighted and China projected. In (d), PCA point predictivities of the centred, scaled countries data. In (e),PCA axis predictivities of the centred, scaled countries data.



(c)



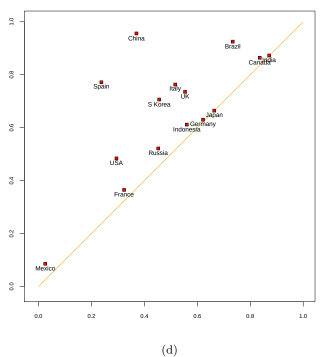


Figure 2.1. (continued)

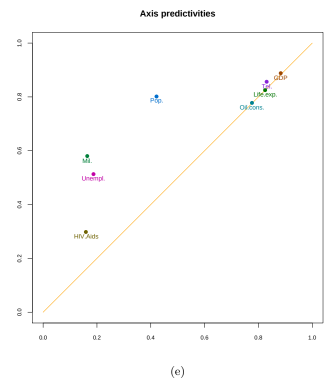


Figure 2.1. (continued)

Korea, cluster in the south-east quadrant. Brazil, Russia and Indonesia lie more towards the west, with Mexico straddling the divide. India, and especially China, lie further away.

While the relative positions of the points are interesting, biplots come into their own when the points are related to their original variable values through the axes. By right clicking inside the predictive linear biplot and selecting 'Predict cursor positions' from the pop-up menu, an array of orthogonally projecting lines emanates from, and follows, the cursor as it moves over the biplot. If 'Predict points closest to cursor positions' is selected instead, the lines project from the point closest to the cursor as it moves, rather than from the cursor itself. So for example, the image in the top right panel of Figure 2.1 was created by hovering the cursor closer to the point for China than to any other point. These orthogonally projecting lines intersect the axes at the positions at which the optimal approximations to the original variables values are to be read off. It can be seen from the image that China scores relatively low on all the variables except population and military spending. As the cursor moves, these predictions are also given numerically, in real time, in the predictions tab. Dynamic prediction is disabled by right clicking inside the biplot and selecting 'Don't predict' from the pop-up menu. (Some numerical predictions are given below.)

With many variables, a biplot may become crowded. A particular axis can be highlighted by right clicking it and then selecting 'Highlight' from the pop-up menu. Doing so greys the other axes, and displays the true variable values of the highlighted axis above the corresponding points. The displays in the diagnostic tabs are shaded accordingly and orthogonal projections are drawn to the highlighted axis only. An example is shown in the centre panel of Figure 2.1, where GDP is highlighted and China is predicted.

The question of course, is how good the biplot approximation is. This depends on both the points and the axes. As for the points, the 'quality' of the PCA approximation is found by clicking to the export tab. For the countries data, 69.3% of the variation in the samples is accounted for by the first two principal components. Point and axis predictivities (Gower, Le

Roux and Gardner-Lubbe, 2008) may also be calculated. Diagrams of these are available in the points and axes tabs, respectively. Those for the countries data are shown in the bottom panels of Figure 2.1. They were saved by right clicking them in the GUI and then making use of the save as options in the pop-up menu. Points or axes further to the right of these diagrams are better represented in the first (or horizontal) biplot dimension. The closer to the top of the diagram, the better the point or axis is represented overall. India, Canada and Brazil are therefore relatively well represented in the first dimension. Japan, Germany and Indonesia are represented reasonably in the first dimension, but poorly in the second. France, the United States and Russia are poorly represented overall, and Mexico extremely poorly. China is the best represented country overall. The axes may be similarly interpreted. Predictivities are also available numerically from the export tab.

Another measure of the goodness of the approximation is the relative absolute error, which may be calculated for any sample on any variable. The relative absolute error is defined to be the absolute difference between the predicted and actual values, expressed as a percentage of the range  $(\max - \min)$  of the particular variable. For GDP, for example, the following output is obtained for the countries data from the export tab:

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	Prediction	Actual	RelAbsErr%
Brazil	9330.3	8710	1.6
Canada	37282.7	35370	4.8
China	10606.7	7724	7.3
France	27669.1	29852	5.5
Germany	31869.4	31941	0.2
India	40.2	3685	9.2
${\tt Indonesia}$	5054.5	4041	2.6
Italy	27130.3	30199	7.7
Japan	34209.8	33100	2.8
Mexico	19392.0	10570	22.2
Russia	8865.5	12350	8.8
S Korea	30946.1	24386	16.5
Spain	26507.7	27418	2.3
UK	31644.7	31723	0.2
USA	33889.0	43369	23.9

Although the United States, Mexico and South Korea predict poorly on the GDP axis, the overall configuration is optimal. By taking means over the samples, mean relative absolute errors may be obtained for the different variables. From the export tab these are:

GDP	HIV.Aids Lif	e.exp.	Mil. Oi	l.cons.	Pop.	Tel.	Unempl.
7 7	20 1	98	14 4	11 2	11 3	88	15 1

These error rates reinforce what is conveyed by the axis predictivities: that HIV/Aids prevalence, unemployment and military spending are relatively poorly represented, the other variables better. Mean relative absolute errors are useful as a measure of the loss of information in biplots since they can be calculated for any type of biplot. Predictivities are defined only when certain orthogonal decompositions exist (Gower *et al.*, 2008), as they do in the case of PCA, CVA and analysis of distance (AOD) (Krzanowski, 2004; Gardner, Le Roux, Rypstra and Swart, 2005) biplots.

For a biplot to be usable in printed form, it must necessarily be two-dimensional. However, assisted by a computer, a user may easily interact with a biplot in three dimensions. Three-dimensional, non-MDS biplots may be obtained in the BiplotGUI package by clicking the

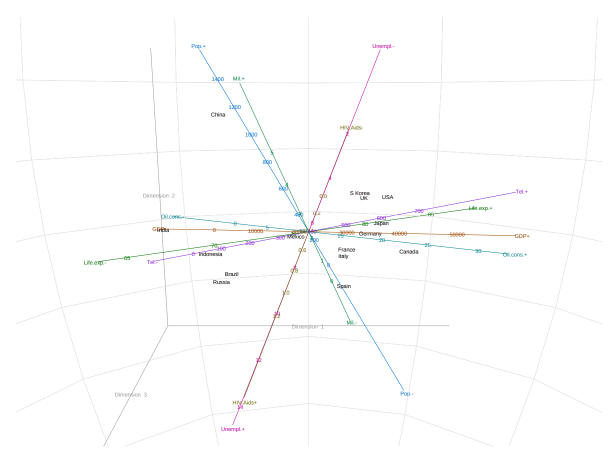
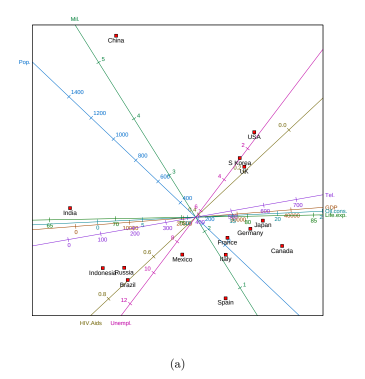


Figure 2.2. The predictive PCA biplot of the centred, scaled countries data, in three dimensions. This figure corresponds to the two-dimensional biplot at the top left of Figure 2.1.

External' menu button at the bottom left of the GUI and then selecting the 'In 3D' option. Alternatively, the user may simply press the F12 shortcut key shown alongside the option. Doing so renders the three-dimensional version of the currently displayed two-dimensional biplot in an external window. This feature makes use of the rgl package (Adler and Murdoch, 2008) and allows the biplot to be rotated and enlarged dynamically. Figure 2.2 shows the three-dimensional predictive PCA biplot of the countries data that corresponds to the two-dimensional version at the top left of Figure 2.1. A further 12.9% of the total variation in the samples is accounted for in the additional dimension. A 360 degree 'fly-by' of the biplot is given by default as the window opens.

A PCA approximation results from the projection of samples onto the plane of best fit. In a covariance biplot ('Joint  $\rightarrow$  Covariance/Correlation'), these 'scores' are adjusted so that the cosines of the angles between the biplot axes approximate the correlations between the corresponding variables. The correlation biplot is the same as the covariance biplot, but with the variables first transformed to have unit variances (via the settings box, in the usual manner). The correlation biplot of the countries data is shown in the top left panel of  $\ref{eq:condition}$ . Three groups of variables are seen to be highly positively correlated (the angles between them are small):

- life expectancy, oil consumption, GDP, the number of telephone lines;
- HIV/Aids prevalence, unemployment;
- population, military spending.



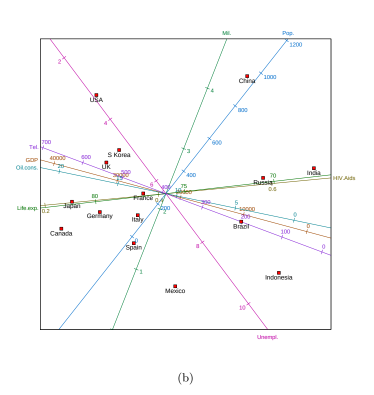
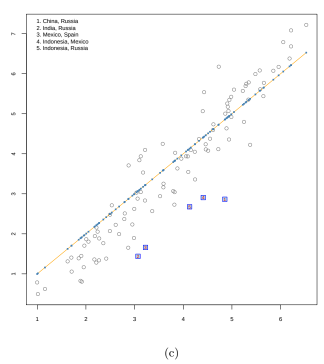


Figure 2.3. In (a), a predictive correlation biplot of the centred, scaled countries data. In (b), a predictive regression biplot for the metric MDS representation of the centred, scaled countries data. The MDS representation is in terms of its principal axes. In (c), the Shepard diagram corresponding to the regression biplot of (a). In (d), a predictive regression biplot for the metric MDS representation of the centred, scaled country data, but with China removed. In (e), the metric MDS of the centred, scaled data with China removed.





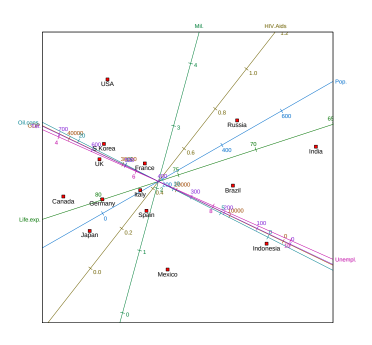
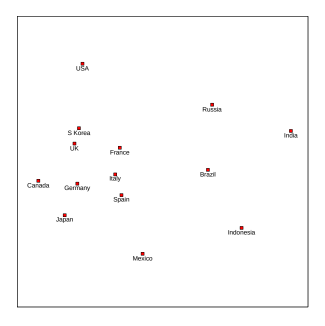


Figure 2.3. (continued)

(d)



(e)

Figure 2.3. (continued)

Notice that the labels of the axes are attached to those ends of the axes that have the higher calibrations. This is the default option for all linear biplots. Alternatively, the axis labels may be given in a legend underneath the biplot, or no axis labels may be given whatsoever. These and similar options may be set via the 'View' menu. Notice also that the option 'Joint  $\rightarrow$  CVA' is disabled, as canonical variate analysis requires the specification of group membership. This has not been provided in the case of the countries data. We return to CVA biplots in Section 2.3.

As opposed to dimension reduction by projection, in MDS the points are chosen so that the squared differences between the inter-sample disparities and the inter-point distances are explicitly minimised. The 'Points  $\rightarrow$  MDS' menu gives various options. These include taking the inter-sample disparities to be the inter-sample dissimilarities themselves (the identity transformation); retaining merely the order of the inter-sample dissimilarities by optimally transforming them into disparities (monotone regression, Kruskal, 1964b); or monotonically smoothing the inter-sample dissimilarities into disparities (the monotone spline transformation, Ramsey, 1982, 1988). Therefore metric, non-metric and semi-metric MDS representations are available. The inter-sample dissimilarities are calculated according to the chosen distance metric. Four such metrics are currently available from the 'Points  $\rightarrow$  Distance metric' menu: Pythagoras distance, Square-root-of-Manhattan distance, Clark distance and Mahalanobis distance (see the package manual for the formulae). Inter-point distances are always Pythagorean. An iterative majorisation (IM) algorithm (De Leeuw, 1977; De Leeuw and Heiser, 1980) is used to find the MDS solutions; for the implementations, see Borg and Groenen (2005, Chapters 8, 9). The IM algorithm converges uniformly, and usually leads to a local minimum, although in theory a saddle-point cannot be ruled out.

The top right panel of ?? shows a metric MDS of the countries data, expressed in terms of its principal axes, with approximate regression biplot axes superimposed (from 'Points  $\rightarrow$  MDS  $\rightarrow$  In terms of principal axes', thereafter 'Points  $\rightarrow$  MDS  $\rightarrow$  Identity transformation').

The default distance metric, Pythagoras distance, is retained. In this representation, the relative distances between the points are directly related to the corresponding dissimilarities between the countries. The United Kingdom and South Korea, therefore, are more similar to one another than they are to the other countries with respect to the eight variables. As the algorithm converges, updates of the configuration are shown in the biplot region, together with updates of the graphs in the diagnostic tabs. The 'Live updates' option, however, may be disabled to increase the speed at which the algorithm runs (the checkbox in question is amongst the buttons at the bottom of the GUI). A graph of the stress values over iterations is given in the convergence tab; in this instance, from the export tab, convergence is reached after 96 iterations, with a final stress value of 45.8. A Shepard diagram (Borg and Groenen, 2005, Section 3.3; Shepard, 1962) can be found in the points tab and is shown at the centre of ??. Each circle in the Shepard diagram represents a pair of samples. The horizontal axis indicates the inter-sample dissimilarity; the vertical axis indicates the corresponding inter-point distance. The blue dots on the yellow line (which generalises to a step function or a curve) indicate the disparities. Thus the closer the circles are to the line (or step function or curve), the better the overall fit. The five worst-fitting point pairs are identified in the top left corner of the diagram. The dissimilarity between China and Russia, therefore, is most poorly approximated by the points. The 'Points  $\rightarrow$  MDS  $\rightarrow$  Random initial configuration' option forces the algorithm to start from a random configuration at each run. Otherwise, the last PCO or MDS solution is taken to be the new initial configuration, as is the case for ??.

To conclude with the countries data, suppose that we feel that China is in many ways atypical, and that we would like to see what the effect would be of removing it from consideration. To do so we need simply 'drag' the point representing China from the biplot into the kraal. We may also right click the point representing China and select 'Send to kraal' from the pop-up menu. The biplot region is then automatically updated as if China were never part of the data set. The updated biplot is given in the bottom left panel of ??. Russia's position relative to the other countries seems to have been most greatly affected. There has also been a re-alignment amongst the axes, most notably the axes for HIV/Aids, population and unemployment. Axes may also be removed to the kraal. Points and axes which have been removed to the kraal may be dragged back onto the biplot, or the kraal may be emptied of its points only, its axes only, or of both its points and axes simultaneously by making use of the buttons below it, or by right clicking inside it and selecting the desired option from the pop-up menu. At any stage, the points and/or axes of any representation may be hidden by clicking on the options in the hide menu button at the bottom of the window. The figure at the bottom right of ?? is the same as the one in the bottom left panel, but with the biplot axes hidden as described.

### 2.3 Antique furniture

It is often of great interest to collectors, auctioneers and cultural historians to be able to correctly identify the type of wood used to make antique furniture. In the period between 1652 and 1900, wood from both the indigenous  $Ocotea\ bullata$  ('Stinkwood'), and the imported  $Ocotea\ porosa$  ('Imbuia'), were used to make Old-Cape furniture in South Africa. Being from the same genus and family (Lauraceae), it is often difficult to distinguish between the two types of wood based solely on a traditional analysis of colour, smell, and other observable characteristics. Burden, Gardner, Le Roux and Swart (2001) and Le Roux and Gardner (2005) make use of CVA biplots of anatomical measurements to distinguish between the species. A third species,  $Ocotea\ kenyensis$ , is also included in the analyses. The microscopically measured variables are: tangential vessel diameter in  $\mu$ m (VesD); vessel element length in  $\mu$ m (VesL); fibre length in  $\mu$ m (FibL); ray height in  $\mu$ m (RayH); ray width in  $\mu$ m (RayW);

and the number of vessels per mm<sup>2</sup> (NumVes). The 37 observations are the mean values over fifty repeat-measurements made on 20 samples of *Ocotea bullata*, 10 of *Ocotea porosa*, and 7 of *Ocotea kenyensis*. The data are included in the BiplotGUI package as the data frame AntiqueFurniture, of which the first column contains the group specifications. The data may be viewed from within R by entering the following instructions at the R prompt of the R console:

```
data (Antique Furniture)
Antique Furniture
```

To initialise the GUI with the antique furniture data, the following command may be entered at the R prompt:

```
Biplots (Data = AntiqueFurniture [ \, , \, \, -1], \ groups = AntiqueFurniture [ \, , \, \, 1])
```

In other words, the data consist of all the columns of AntiqueFurniture except the first, while the group specifications are precisely the contents of the first column. Alternatively, if the data were saved as an Excel 1997–2003 file 'AntiqueFurniture.xls', with the group specifications given in, say, the first column, the call

```
Biplots \, (\, excel \, = \, "\, AntiqueFurniture \, . \, xls \, " \, , \  \, ExcelGroupsCol \, = \, 1)
```

could have been made instead.

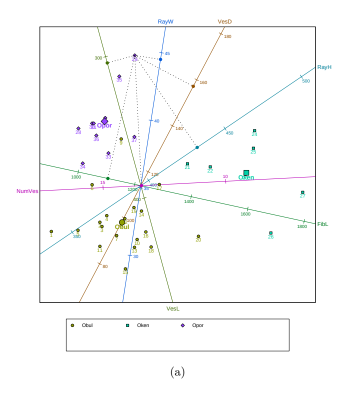
As was mentioned earlier, upon initialisation of the GUI, the predictive PCA biplot is shown by default. To show the CVA biplot instead, the user simply needs to click the option 'Joint  $\rightarrow$  CVA'. This option is now available since, in the call to the Biplots function, groups were defined. The predictive CVA biplot of the antique furniture data is shown in the top left panel of ??. The positions of the points are determined by the first two canonical variates – those linear combinations of the original variables that maximally separate the group means, subject to certain restrictions (Krzanowski, 2000, Section 11.1). The group means themselves are shown as larger but corresponding symbols (activated by clicking 'Additional  $\rightarrow$  Interpolate  $\rightarrow$  Sample group means', retaining the default options). Since there is more than one group, an optional legend is included below the biplot by default. The mechanism for prediction is the same as before and is illustrated in the case of sample 29.

The top right panel of ?? shows the same biplot, now overlain onto a two-dimensional density estimate of the points. The density estimate is obtained by clicking 'Additional → Point densities' and accepting the default options (amongst other things, for the point densities to be estimated for all points, as opposed to certain groups of points only). The point densities are calculated using the default arguments to the bkde2D function of the KernSmooth package (Wand, 2008). Similar biplots can be found in Blasius, Eilers and Gower (2008).

Sometimes it is helpful to zoom into or out of portions of a biplot. This is done by right clicking on a focal point inside the biplot, and selecting the 'Zoom in' or 'Zoom out' option from the pop-up menu which then appears. The bottom panel of ?? shows the CVA biplot of antique furniture, enlarged around the mean of the species *Ocotea bullata*. The original view can be restored by choosing the 'Reset zoom' option from the pop-up menu.

A screenshot of the GUI is shown in 2.2. To the left, a CVA biplot of the antique furniture data appears. From the settings box, it can be seen that the axes are not predictive; in fact they are vector sum interpolative. Also, the data have not been transformed, except for the obligatory centring of the columns to have zero means. In any case, CVA biplots are unaffected by the scaling of the variables to unit variance.

Sample 18 has been dragged from the biplot into the kraal. It has therefore not been taken into account in the construction of the biplot. However, using its variable values, it has subsequently been interpolated onto the biplot towards the bottom of the image (using the 'Additional  $\rightarrow$  Interpolate  $\rightarrow$  A New Sample' option). This is the most appropriate position for the sample in the existing biplot. It is reassuring that the positions assigned to sample 18 in Figures ?? and 2.2 correspond so closely. This need not have been the case. Also



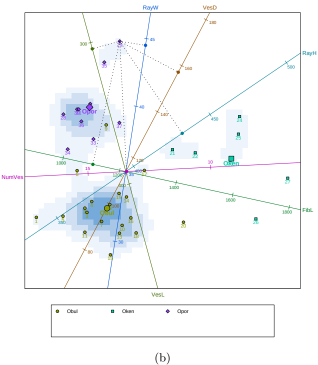


Figure 2.4. In (a), a predictive CVA biplot of the antique furniture data with sample 29 projected onto all the biplot axes. The group means are also shown. In (b), as in the top left panel, but with the biplot overlain onto the two-dimensional density estimate of the points. In (c), the same biplot as the other two, but zoomed in around the mean of the species *Ocotea bulluta*.

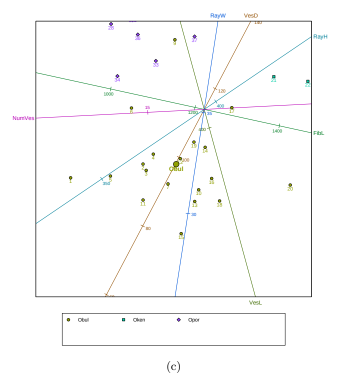
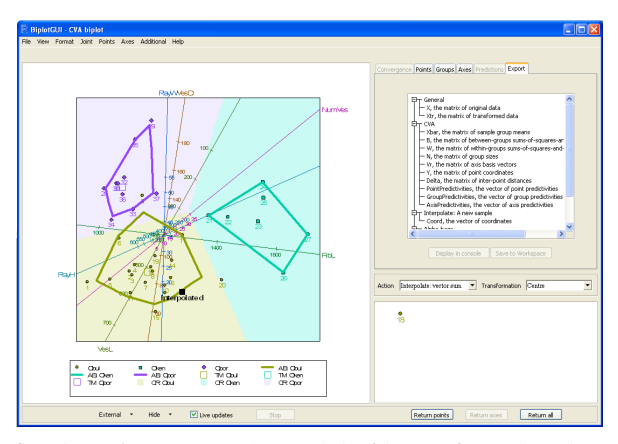


Figure 2.4. (continued)

notice that, notwithstanding the removal of sample 18, the calibrations and *directions* of the predictive and interpolative biplot axes differ. This is in general the case for CVA biplots.

The biplot in 2.2 also sports colour-coded classification regions. These are the regions in the display space plane closest to the respective group means in a specified number of canonical dimensions, here the maximum number possible for the particular data set, six. The classification regions are included by selecting 'Classification regions' from the 'Additional' menu. They may be used for the classification of interpolated samples. For more on the links between biplots and discrimination, see Gardner and Le Roux (2005). Furthermore, by clicking 'Additional → Alpha-bags', alpha-bags (Gardner, 2001; Aldrich, Gardner and Le Roux, 2004) and Tukey medians have been superimposed for the species Ocotea bullata and Ocotea porosa (there are too few samples for an alpha-bag for Ocotea kenyensis to be constructed; with an appropriate warning, a convex-hull is displayed instead). Alpha-bags are closely related to the bagplots of Rousseeuw, Ruts and Tukey (1999) and enclose regions that contain approximately the inner  $100\alpha\%$  of samples, here 90% of the samples for the two species separately. The alpha-bags and convex hull do not overlap. This emphasises the high degree of separation between the species. For CVA biplots, group predictivities (Gower et al., 2008) may also be calculated, in addition to the point and axis predictivities discussed earlier. A diagram of these is available in the groups tab. Finally, 2.2 also shows the export tab. As has been explained, various objects are available for export from this tab, and may be either displayed in the R console, or saved to the current R workspace. The list of available objects depends on what is shown in the biplot.

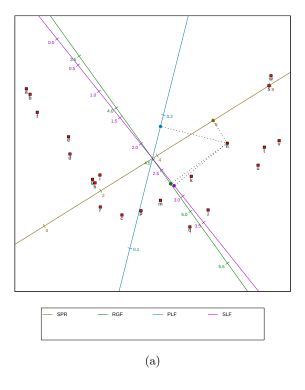


Screenshot 2.2: A vector sum interpolative CVA biplot of the antique furniture data is shown towards the left, with sample 18 removed to the kraal. Sample 18 has then been interpolated to give its implied position. Classification regions are shown, as well as 90% alpha-bags for the species Ocotea bullata and Ocotea porosa. A convex hull surrounds the points of the species Ocotea kenyensis. The export tab is shown top right.

### 2.4 Fighter aircraft

Measurements of four variables on 22 types of fighter aircraft were extracted by Cook and Weisberg (1982) from a report by Stanley and Miller (1979). Following Gower and Hand (1996), we consider only the first 21 of these aircraft in the biplots below. The four variables are: specific power, proportional to power per unit weight (SPR); flight range factor (RGF); payload as a fraction of gross weight (PLF); and sustained load factor (SLF). These data can be found in the FighterAircraft data frame included in the BiplotGUI package. The GUI is initialised in the same way it was for the countries data in Section 2.2.1.

The left panel of  $\ref{eq:property}$  shows a regression biplot of the fighter aircraft data with the points determined by PCO and the inter-sample dissimilarities calculated according to the Square-root-of-Manhattan distance metric. The figure is obtained by clicking 'Points  $\rightarrow$  Distance metric  $\rightarrow$  Square-root-of-Manhattan' and then 'Axes  $\rightarrow$  Regression'. Making use of orthogonal projection, the variable values for aircraft 'n' are predicted to be 7.342, 4.95, 0.202 and 3.25, respectively. These can be compared to the actual values, 5.855, 4.53, 0.172 and 2.50. The right panel of  $\ref{eq:property}$  shows the corresponding circular non-linear biplot (obtained by clicking 'Axes  $\rightarrow$  Circular non-linear'). Here prediction is performed by completing the circle which has, as diagonal, the line stretching from the origin of the biplot to the point to be predicted. The predicted values are read off at the points at which the circle intersects the axes (Gower and Hand, 1996, Section 6.3.2). If a particular axis is intersected at more than one position,



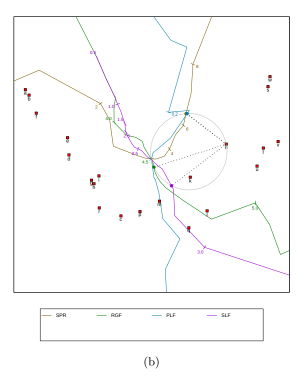


Figure 2.5. In (a), a predictive regression biplot of the fighter aircraft data, with points determined by PCO based on the Square-root-of-Manhattan distance metric. The orthogonal prediction of the variables values of aircraft 'n' is shown. In (b), a predictive circular non-linear biplot of the fighter aircraft data, with points determined by PCO based on the Square-root-of-Manhattan distance metric. The circular prediction of the variable values of aircraft 'n' is shown.

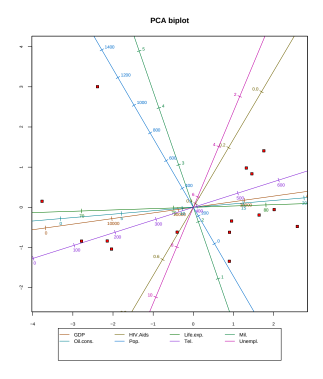


Figure 2.6. A modified version of the biplot given in the top left panel of Figure 2.1: a predictive PCA biplot of the countries data with a title, hidden point labels, axis labels in a legend, and the axes of the display space calibrated.

the position closest to the point being predicted is used. If an axis isn't intersected at all, no prediction for the corresponding variable can be made. For aircraft 'n', the valid points of intersection are shown in the figure as small, filled circles on the circumference of the larger circle. From the predictions or export tabs, the circular non-linear predictions for aircraft 'n' are 6.090, 4.54, 0.174 and 2.90, respectively (these values depend on how finely the non-linear axes are constructed; by default 20 positions are taken into account from each calibrated marker to the next). Except for the fourth variable, the non-linear predictions are very close to the actual values. The non-linear biplot axes represent an appreciable improvement over the approximate linear axes in this case. This need not have been so.

### 2.5 Further features

In this section, we briefly touch upon some further features of the BiplotGUI package which haven't yet received much attention – the customisation of biplots and other graphs, and how they may be exported.

There are two main ways in which graphs may be customised in the BiplotGUI package. Basic customisation can be performed using the options of the 'View' menu, while the 'Format' menu options can be used to alter a large number of graphical parameters.

Figure 2.6 shows the same predictive PCA biplot of the countries data as was shown in the top left panel of Figure 2.1. However, the biplot in Figure 2.6 has been modified by changing the default selections in the 'View' menu. The 'Title' option places a title above the biplot; by default the title reflects the type of biplot, but it may be changed via 'Format  $\rightarrow$  Title'. Furthermore, the point labels have been disabled by deselecting 'Show point labels'. Instead of showing the axis labels around the edges of the biplot as in Figure 2.1, the labels in Figure 2.6

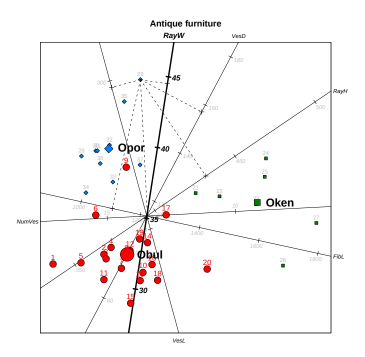
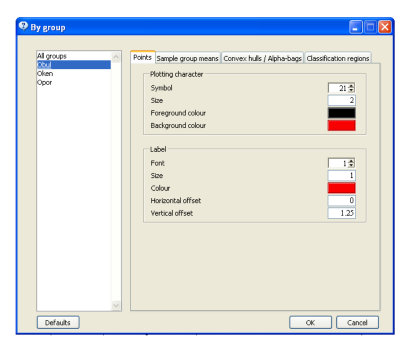


Figure 2.7. A modified version of the biplot given in the left panel of ??: a predictive CVA biplot of the antique furniture data with sample 29 projected onto all the biplot axes; the group means are shown.

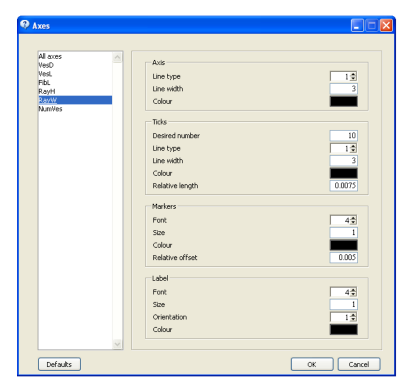
are shown in a legend ('Show axis labels in legend'). The 'Calibrate display space axes' option calibrates the two dimensions of the biplot, but this is generally undesirable in biplots of the new approach (Gower and Hand, 1996, Section 2.6).

The 'Format' menu allows virtually all the graphical parameters used internally by the package to be altered. The biplot in Figure 2.7 serves as an example. This biplot is the same as the one in the left panel of ??, but with some of the default graphical parameters changed. The 'ByGroup' option allows the graphical parameters which relate to points, sample group means, convex hulls / alpha-bags and classification regions to be set for all groups simultaneously, or for a single group at a time. 2.3 shows the dialogue box as it appears when customising the points of the species Ocotea bullata. The parameter values appear as they have been set for Figure 2.7. The 'Axis' option similarly allows the graphical parameters for axes to be set, simultaneously for all axes or individually. 2.4 shows the dialogue box of the graphical parameters for the axis 'RayW', again with the parameters as they have been set for Figure 2.7. The graphical parameters used in variable value prediction and in the highlighting of axes can also be modified by clicking 'Format → Interactive', while diagnostic tab customisation may be performed by clicking the 'Diagnostic tabs' option. The 'Reset all' option reverts all the graphical parameters back to their default values. In all, more than 80 different graphical parameters may be set, often-times differently for different groups or axes. All these parameters are documented in detail in the package manual.

Naturally users would want to export the graphs produced using the BiplotGUI package. Biplots and diagnostic tab graphs can be saved in various file formats. These are: PDF, Postscript, Metafile, Bmp, Png, Jpeg (50%, 75% and 100% quality) and PicTeX. Any graph can be saved by right clicking it and navigating the 'Save as' menu. The biplot region may also be saved via the 'File  $\rightarrow$  Save as' menu. While the images shown onscreen are by necessity Metafile images, the images that appear in this article were saved in the PDF file format.



Screenshot 2.3: The Format  $\rightarrow$  By group dialogue box as it appears for the biplot in Figure 2.7.



Screenshot 2.4: The  $Format \rightarrow Axes$  dialogue box as it appears for the biplot in Figure 2.7.

Together with Postscript, such images are of the highest quality. 'Copy' and 'Print' options are also available.

### 2.6 Conclusion

In this paper, the BiplotGUI package for R was introduced. Its features were illustrated using three data sets. Ideas for future releases were briefly explored.

The package makes it possible to easily construct many types of biplots and to interact with them in various ways. The package is free and its source code shared. Amongst linear biplots, the PCA, covariance/correlation, CVA, regression and Procrustes biplots are supported. Circular non-linear biplots can be created. In addition, PCO and MDS representations can be displayed on their own, without added biplot axes. Additional descriptors can be superimposed, and three-dimensional biplots can be explored using the rgl package. Various goodness-of-fit measures are easily accessible.

## Chapter 3

### Detail

In this chapter, the features of the BiplotGUI package are documented, starting with some general comments in Section 3.1. The structure of the GUI can be seen from Section 3.2. This section also serves as a table of contents to Section 3.3, where the features are presented. Known issues are listed in ??.

#### 3.1 General

While more than one instance of the GUI may be opened simultaneously from within the same R console (by issuing additional calls to Biplots), this often results in a console warning as the GUIs are closed. It is safe to use the console for other commands while running an instance of the GUI. GUIs can be run simultaneously from different instances of R.

For more information on Biplots, the single function of the package, see the R help file for it by entering ?Biplots at the R prompt of the R console. This chapter focuses on the features of the GUI itself.

#### 3.2 List of Features

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                                        Show point values
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                                        Show group labels in legend
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#### 3.3 **Features**

In this section, the features of the BiplotGUI are systematically documented. Section 3.2 serves as a table of contents. Features are indented as in Section 3.2. The symbols used, from the outer-most to inner-most levels of indentation, are  $\triangleright$ ,  $\diamond$ ,  $\triangleright$ ,  $\diamond$  and  $\diamond$ , respectively. Information on the options in dialogue boxes is given in framed sections. After an introductory sentence, each feature is discussed under standard headings. These headings are listed below, together with a short description of each. Headings appear only when applicable.

DETAIL More detailed information on the feature For graphical parameters, the default value DEFAULT VALUE

A keyboard shortcut to the feature SHORTCUT

Mutually exclusive features ALTERNATIVE TO

For dialogue boxes, a reference to a screenshot **SREENSHOT** 

References to external material REFERENCES

### ▶ Menu bar

The primary interface of the GUI.

DETAIL · The menu bar is located at the top of the GUI. It consists of a number of drop-down menus, each with a number of entries. Entries may have sub-entries. The entries of menus are arranged around common themes of action. The Joint menu is for biplots with a joint mechanism for determining the positions of both points and axes. For other biplots, the Points menu determines the positions of the points while the Axes menu determines the positions of the axes. Many entries contained in the menu bar can also be accessed from the pop-up menus which appear when various parts of graphs are right clicked.

SHORTCUTS · Often-used entries have dedicated keyboard shortcuts displayed to their right. These keyboard shortcuts are listed on page vi. Menu bar entries may also be accessed via the keyboard by pressing the Alt key together with the underlined letter in the name of the corresponding drop-down menu, followed by the sequence of underlined letters in the path to the entry.

### • File

Contains export and other options.

DETAIL · The File menu lets the user save, copy or print the currently displayed biplot region. In addition, general options may be set, or the user may exit the GUI.

#### Save as

Lets the user save the currently displayed biplot region in a number of widely-used file formats.

DETAIL · Depending on how the GUI has been resized onscreen, the proportions within a saved biplot region may differ from what is shown onscreen. To ensure consistency, saved biplot regions always have both their width and height set to 20.32 centimeters (8 inches). The PDF and Postscript file formats offer the highest quality, better even than that shown onscreen.

SHORTCUT · Ctrl+S. Saves the currently displayed biplot region in the currently selected file format.

### • **PDF...**

Saves the currently displayed biplot region as a PDF file.

DETAIL · Together with Postscript, the PDF file format offers the highest quality.

ALTERNATIVE TO · Postscript, Metafile, Bmp, Png, Jpeg 50% quality, Jpeg 75% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the PDF file.

# • Postscript...

Saves the currently displayed biplot region as a Postscript file (.ps).

DETAIL · Together with PDF, the Postscript file format offers the highest quality.

ALTERNATIVE TO · PDF, Metafile, Bmp, Png, Jpeg 50% quality, Jpeg 75% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the Postscript file.

### o Metafile...

Saves the currently displayed biplot region as a Windows Metafile (.wmf). ALTERNATIVE TO · PDF, Postscript, Bmp, Png, Jpeg 50% quality, Jpeg 75% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the Windows Metafile.

### ∘ **Bmp...**

Saves the currently displayed biplot region as a bitmap (.bmp).

ALTERNATIVE TO · PDF, Postscript, Metafile, Png, Jpeg 50% quality, Jpeg 75% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the bitmap file.

## $\circ$ Png...

Saves the currently displayed biplot region as a png file.

ALTERNATIVE TO · PDF, Postscript, Metafile, Bmp, Jpeg 50% quality, Jpeg 75% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the png file.

### Jpeg

Lets the user save the currently displayed biplot region as a Jpeg file (.jpg, .jpeg).

## ♦ **50%** quality...

Saves the currently displayed biplot region as a Jpeg file at 50% quality. ALTERNATIVE TO · PDF, Postscript, Metafile, Bmp, Png, Jpeg 75% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the 50% quality Jpeg file.

# ♦ 75% quality...

Saves the currently displayed biplot region as a Jpeg file at 75% quality. ALTERNATIVE TO · PDF, Postscript, Metafile, Bmp, Png, Jpeg 50% quality, Jpeg 100% quality, PicTeX

A standard dialogue box appears which lets the user specify the name and location of the 75% quality Jpeg file.

# ♦ 100% quality...

Saves the currently displayed biplot region as a Jpeg file at 100% quality. ALTERNATIVE TO  $\cdot$  PDF, Postscript, Metafile, Bmp, Png, Jpeg 50% quality, Jpeg 75% quality, PicTeX

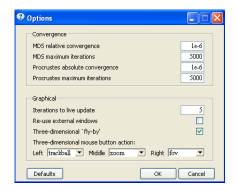
A standard dialogue box appears which lets the user specify the name and location of the 100% quality Jpeg file.

## o PicTeX...

Saves the currently displayed biplot region in the PicTeX format (.tex).

ALTERNATIVE TO · PDF, Postscript, Metafile, Bmp, Png, Jpeg 50% quality, Jpeg 75% quality, Jpeg 100% quality

REFERENCES  $\cdot$  See the R help file of the pictex function for the limitations of the conversion to this format.



Screenshot 3.1: The  $File \rightarrow Options$  dialogue box.

A standard dialogue box appears which lets the user specify the name and location of the PicTeX file.

# $\triangleright$ Copy

Copies the currently displayed biplot region to the clipboard as a Windows Metafile. DETAIL · Depending on how the GUI has been resized onscreen, the proportions within a copied biplot region may differ from what is shown onscreen. To ensure consistency, copied biplot regions always have both their width and height set to 20.32 centimeters (8 inches). Biplots saved as PDF or Postscript files are of a higher quality than those copied to the clipboard.

 ${\tt SHORTCUT} \cdot Ctrl + C$ 

## ⊳ Print…

Prints the currently displayed biplot region.

DETAIL · Depending on how the GUI has been resized onscreen, the proportions within a printed biplot region may differ from what is shown onscreen. To ensure consistency, printed biplot regions always have both their width and height set to 20.32 centimeters (8 inches).

SHORTCUT · Ctrl+P

A standard dialogue box appears which lets the user set various options before printing.

## ⊳ Options...

Lets the user set various general options.

SCREENSHOT  $\cdot$  3.1

### Convergence

Lets the user set various convergence criteria.

# MDS relative convergence

If still running, the  ${\tt IM}$  algorithm for  ${\tt MDS}$  stops when the proportional decrease in stress becomes smaller than this value.

default value  $\cdot~10^{-6}$ 

REFERENCES · Borg and Groenen (2005, pp. 191–192, 204–205)

#### MDS maximum iterations

If still running, the IM algorithm for MDS stops when the number of iterations reaches this value.

Default value  $\cdot$  5000

REFERENCES · Borg and Groenen (2005, pp. 191–192, 204–205)

### Procrustes absolute convergence

If still running, the Procrustes algorithm for interpolative biplots stops when the absolute decrease in stress first becomes smaller than this value.

DEFAULT VALUE  $\cdot~10^{-6}$ 

REFERENCES · Gower and Hand (1996, Section A.10.2), Gower and Dijksterhuis (2004, p. 57)

### Procrustes maximum iterations

If still running, the Procrustes algorithm for interpolative biplots stops when the number of iterations reaches this value.

DEFAULT VALUE · 5000

REFERENCES · Gower and Hand (1996, Section A.10.2), Gower and Dijksterhuis (2004, p. 57)

# Graphical

Lets the user set general graphical options.

# Iterations to live update

The biplot region and relevant graphs in the diagnostic tabs are updated after this number of MDS iterations, and multiples of it.

DETAIL · The smaller the value of this option, the smoother the transitions in the graphs; the larger the value of this option, the smaller the time to convergence. Irrespective of the value of this option, the biplot region and the relevant graphs in the diagnostic tabs are only updated during MDS convergence if Live updates is checked below the biplot region.

default value  $\cdot$  5

# Reuse external windows

Toggles between using the same or new windows for external graphics.

DETAIL · If checked, the same window is used to display any two-dimensional external graphs that are produced. Such graphs can be created by clicking External  $\rightarrow$  As is below the biplot region, or by right clicking the graphs of the diagnostic tabs and selecting External from the pop-up menu. If clear, new windows are used to show any additional two-dimensional graphs, retaining previous external graphs, if any. Similar comments hold for three-dimensional external biplots created by clicking External  $\rightarrow$  In 3D below the biplot region. The windows of the two- and three-dimensional graphs operate independently.

DEFAULT VALUE  $\cdot$  clear

# Three-dimensional 'fly-by'

Toggles between including an initial 'fly-by' in three-dimensional biplots, or not including one.

 ${\tt DEFAULT\ VALUE\cdot checked}$ 

# Three-dimensional mouse button action: left

The action of the left mouse button in three-dimensional biplots.

DEFAULT VALUE  $\cdot$  trackball

REFERENCES · See the R help file of the par3d function of the rgl package (Adler and Murdoch, 2008) for details on the choices.

### Three-dimensional mouse button action: middle

The action of the middle mouse button in three-dimensional biplots.

DEFAULT VALUE · zoom

REFERENCES · See the R help file of the par3d function of the rgl package (Adler and Murdoch, 2008) for details on the choices.

# Three-dimensional mouse button action: right

The action of the right mouse button in three-dimensional biplots.

DEFAULT VALUE · fov

REFERENCES · See the R help file of the par3d function of the rgl package (Adler and Murdoch, 2008) for details on the choices.

### **Defaults**

Reverts the options to their default values.

#### OK

Saves the options and returns to the GUI.

#### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

#### ⊳ Exit

Lets the user exit the GUI.

DETAIL  $\cdot$  Upon exiting, all actions performed and changes made in the GUI are lost. The user is returned to the R prompt of the R console.

A standard dialogue box appears which asks the user to confirm whether or not to exit the GUI.

### • View

Allows the users to customise the biplot region.

### **⊳** Show title

Shows a title above the biplot.

DETAIL  $\cdot$  The biplot retains the same size and relative position, irrespective of whether or not a title is shown.

# 

Sets the biplot borders around the points, ignoring the axes.

DETAIL · If selected, the biplot borders are set just large enough to show all the points and their labels. Non-linear axes may not be visible in their entirety.

ALTERNATIVE TO · Clip around points and axes

# ▷ Clip around points and axes

Sets the biplot borders around the points and axes.

DETAIL  $\cdot$  If selected, the biplot borders are set large enough to show all the points, their labels, and the axes in their entirety. Available only for non-linear axes and when the axes are not hidden.

ALTERNATIVE TO · Clip around points

# Show point labels ■ Show point labels ■

Shows the point labels in the biplot.

DETAIL  $\cdot$  If selected, the biplot borders are enlarged so that all the point labels fit. Not available when the points are hidden.

# > Show point values

Shows the variable values of a highlighted axis alongside the corresponding points in the biplot.

DETAIL · Only available when an axis has been highlighted.

# > Show group labels in legend

Shows the group labels in the legend.

DETAIL · Not available when there is only one group of samples, or when the points are hidden.

# ▷ Don't show axis labels

Suppresses the axis labels.

DETAIL · Not available if there are no axes, or when the axes are hidden.

ALTERNATIVE TO · Show clinging axis labels, Show axis labels in legend

# 

Shows the axis labels at those edges of the axes with the higher calibrations.

DETAIL · Available only for linear axes when the axes are not hidden.

ALTERNATIVE TO · Don't show axis labels, Show axis labels in legend

# 

Shows the axis labels in the legend.

DETAIL · Not available if there are no axes, or when the axes are hidden.

ALTERNATIVE TO · Don't show axis labels, Show clinging axis labels

# > Show Additional labels in legend

Shows the labels of descriptors from the Additional menu in the legend.

### > Show next legend entries

Shows the succeeding set of 16 legend entries, if any, in the legend.

DETAIL · The legend can contain up to a maximum of 16 entries at once. This option updates the legend to show the succeeding set of at most 16 entries. The option does not show the first set of entries after the last set. Not available when there are no succeeding entries, or when no legend is shown.

SHORTCUT · Ctrl++

# > Show previous legend entries

Shows the preceding set of 16 legend entries, if any, in the legend.

DETAIL  $\cdot$  The legend can contain up to a maximum of 16 entries at once. This option updates the legend to show the preceding set of 16 entries. The option does not show the last set of entries before the first set. Not available when there are no preceding entries, or when no legend is shown.

SHORTCUT · Ctrl+-

# ▷ Calibrate display space axes

Calibrates the two biplot dimensions.

DETAIL · This is counter to the spirit of biplots of the new approach.

REFERENCES · Gower and Hand (1996, Section 2.6)

# • Format

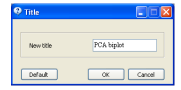
Lets the user customise the internally used graphical parameters.

### ⊳ Title...

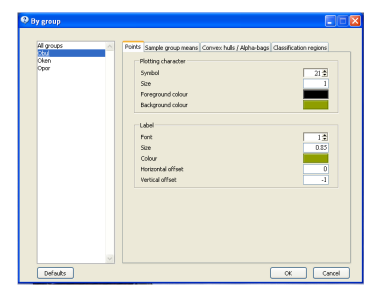
Lets the user change the title.

DETAIL · The title is shown only if View  $\rightarrow$  Show title is selected.

SCREENSHOT  $\cdot$  3.2



Screenshot 3.2: The Format  $\rightarrow$  Title dialogue box.



Screenshot 3.3: The Points tab of the Format  $\rightarrow$  By group dialogue box.

# New title

The new title.

DEFAULT VALUE  $\cdot$  Depends on the currently displayed biplot; can be seen from the title bar of the GUI.

#### Default

Reverts the option to its default value.

#### OK

Saves the title, returns to the GUI, and redraws the biplot region.

# Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

# ⊳ By group...

Allows graphical parameters for points, sample group means, convex hulls / alphabags, and classification regions to be set for a particular group, or for all groups simultaneously.

DETAIL  $\cdot$  When  $All\ groups$  is selected from the list to the left of the dialogue box, changes made to the parameter values to the right of the dialogue box affect all groups. Changes to particular groups can be made by clicking the group names in the list to the left of the dialogue box. Parameter values which differ amongst groups are left blank when  $All\ groups$  is selected.

SHORTCUT · Ctrl+G

SCREENSHOT  $\cdot$  3.3

Only the Points tab is discussed here. The other tabs are discussed together with the options of the Additional menu.

# Plotting character

Lets the user change the graphical parameters of the points that represent the samples, by group.

# Symbol

Sets the symbol used to represent the points, by group.

DETAIL · Corresponds to R's pch argument. Possible values: NA,  $0, \ldots, 25$ .

DEFAULT VALUE  $\cdot$  If there is only one group, 22. Otherwise the values are recycled by group with the values 21 to 25.

#### Size

Sets the size of the points, by group.

DETAIL · Corresponds to R's cex argument. Larger values result in larger points. Possible values: positive real numbers.

Default value  $\cdot$  1

# Foreground colour

Sets the exterior colour of the points, by group.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "black"

# Background colour

Sets the interior colour of the points, by group.

DETAIL · Corresponds to R's bg argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE  $\cdot$  If there is only one group, "red". Otherwise equally spaced colours from the hcl spectrum.

#### Label

Lets the user change the graphical parameters of the point labels, by group.

#### Font

Sets the font of the point labels, by group.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

default value  $\cdot$  1

# Size

Sets the size of the point labels, by group.

 $\texttt{DETAIL}\cdot \texttt{Corresponds}$  to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

DEFAULT VALUE · 0.85

#### Colour

Sets the colour of the point labels, by group.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

<code>DEFAULT VALUE</code>  $\cdot$  If there is only one group, "black". Otherwise equally spaced colours from the <code>hcl</code> spectrum.

## Horizontal offset

Sets the horizontal offset of the point labels from the corresponding points, by group.

 $DETAIL \cdot$  The point labels are shifted horizontally from the corresponding

points with this multiple of the width of the letter 'x' (in the selected font size). Positive values shift the labels to the right; negative values shift the labels to the left.

DEFAULT VALUE · 0

# Vertical offset

Sets the vertical offset of the point labels from the corresponding points, by group.

DETAIL · The point labels are shifted vertically from the corresponding points with this multiple of the height of the letter 'x' (in the selected font size). Positive values shift the labels towards the top; negative values shift the labels towards the bottom.

DEFAULT VALUE  $\cdot -1$ 

# **Defaults**

Reverts the options of all the tabs of the dialogue box to their default values.

#### $\mathbf{OK}$

Saves the options of all the tabs of the dialogue box, returns to the GUI, redraws the currently displayed biplot region and other graphs if necessary.

#### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

#### Axes...

Allows graphical parameters for axes to be set for a particular axis, or for all axes simultaneously.

DETAIL  $\cdot$  When All axes is selected from the list to the left of the dialogue box, changes made to the parameter values to the right of the dialogue box affect all axes. Changes to particular axes can be made by clicking the variable names from the list to the left of the dialogue box. Parameter values which differ amongst axes are left blank when All axes is selected.

 $_{\rm SHORTCUT}\cdot Ctrl{+}A$ 

SCREENSHOT  $\cdot$  3.4

## Axis

Lets the user change the graphical parameters associated with the axis that represents the variable.

### Line type

Sets the type of line used to draw the axis.

DETAIL · Corresponds to R's 1ty argument. Possible values: 0, ..., 6.

default value  $\cdot$  1

# Line width

Sets the width of the axis.

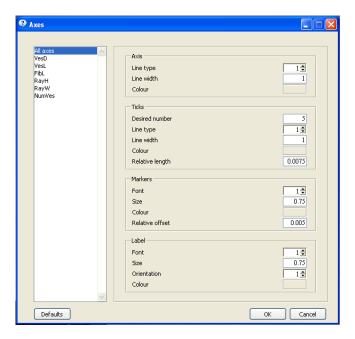
DETAIL · Corresponds to R's lwd argument. Larger values result in wider axes. Possible values: positive real numbers.

DEFAULT VALUE · 1

### Colour

Sets the colour of the axis.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the



Screenshot 3.4: The  $Format \rightarrow Axes$  dialogue box.

colour, producing a standard colour selection dialogue box.

DEFAULT VALUE  $\cdot$  Equally spaced colours from the hcl spectrum.

### **Ticks**

Lets the user change graphical parameters associated with the ticks of the axis.

DETAIL  $\cdot$  The ticks, together with the markers, calibrate the axis. The ticks are the short lines orthogonal to the axis, alongside which the markers (the numbers) are given. For non-linear axes, the slopes of the ticks are approximated numerically.

### $Desired\ number$

Sets the desired number of ticks on the axis.

DETAIL · Corresponds to the argument n of R's pretty function. This number is not strictly adhered to; it is only approximate.

default value  $\cdot$  5

REFERENCES · See the R help file of the pretty function.

### Line type

Sets the type of line used to draw the ticks on the axis.

DETAIL · Corresponds to R's 1ty argument. Possible values:  $0, \ldots, 6$ .

default value  $\cdot$  1

## Line width

Sets the width of the ticks of the axis.

DETAIL · Corresponds to R's lwd argument. Larger values result in wider ticks. Possible values: positive real numbers.

DEFAULT VALUE · 1

#### Colour

Sets the colour of the ticks of the axis.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the

colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · The same as the default colour of the axis.

# Relative length

Sets the relative length of the ticks of the axis.

DETAIL  $\cdot$  The length of the ticks is twice this fraction of the width of the biplot. DEFAULT VALUE  $\cdot$  0.0075

#### Markers

Lets the user change graphical parameters associated with the markers of the axis.

DETAIL · The markers, together with the ticks, calibrate the axis. The markers are the numbers, in terms of the original variable values, given alongside the ticks (the short lines orthogonal to the axis). The markers are positioned on that side of the axis that would place them below the axis if the axis were to be rotated to be horizontal (with the marker values increasing from left to right).

#### **Font**

Sets the font of the markers of the axis.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

DEFAULT VALUE · 1

#### Size

Sets the size of the markers of the axis.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

DEFAULT VALUE · 0.75

### Colour

Sets the colour of the markers of the axis.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE  $\cdot$  The same as the default colour of the axis.

#### Relative offset

Sets the relative offset of the markers from the corresponding ticks of the axis.

 $DETAIL \cdot The markers are shifted this fraction of the width of the biplot from the edge of the corresponding ticks.$ 

Default value  $\cdot$  0.005.

### Label

Lets the user change the graphical parameters of the axis label.

#### Font

Sets the font of the axis label.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

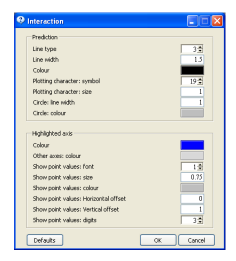
Default value  $\cdot$  1

### Size

Sets the size of the axis label.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

DEFAULT VALUE · 0.75



Screenshot 3.5: The  $Format \rightarrow Interaction$  dialogue box.

### Orientation

Sets the orientation of clinging axis labels.

DETAIL · Corresponds to R's las argument. Possible values: 0: always parallel to the biplot dimension; 1: always horizontal; 2: always perpendicular to the biplot dimension; 3: always vertical.

DEFAULT VALUE · 1

#### Colour

Sets the colour of the axis label.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · The same as the default colour of the axis.

### **Defaults**

Reverts the options to their default values.

#### OK

Saves the options, returns to the GUI, redraws the currently displayed biplot region and other graphs if necessary.

# Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

# ▷ Interaction...

Allows graphical parameters for variable value prediction and highlighted axes to be set.

SCREENSHOT  $\cdot$  3.5

### Prediction

Lets the user change the graphical parameters associated with variable value prediction.

## Line type

Sets the line type used for projection.

Detail · Corresponds to R's 1ty argument. Possible values:  $0, \ldots, 6$ . Default value · 3

#### Line width

Sets the width of the projection lines.

DETAIL · Corresponds to R's lwd argument. Larger values result in wider ticks. Possible values: positive real numbers.

DEFAULT VALUE · 1.5

#### Colour

Sets the colour of the projection lines.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "black"

# $Plotting\ character:\ symbol$

Sets the symbol used to represent the projected points on the axes.

DETAIL · Corresponds to R's pch argument. Possible values: NA,  $0, \ldots, 25$ .

DEFAULT VALUE · 19

## Plotting character: size

Sets the size of the projected points on the axes.

DETAIL · Corresponds to R's cex argument. Larger values result in larger points. Possible values: positive real numbers.

DEFAULT VALUE · 1

#### Circle: line width

Sets the width of the circle used in circular projection in the circular non-linear biplot.

DETAIL · Corresponds to R's lwd argument. Larger values result in wider ticks. Possible values: positive real numbers.

default value  $\cdot$  1

# Circle: colour

Sets the colour of the circle used in circular projection in the circular non-linear biplot.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "gray75"

### Highlighted axis

Lets the user change the graphical parameters associated with highlighted axes.

DETAIL · The graphical parameters prefixed with 'Show point values' are for the customisation of the variable values which are shown when an axis is highlighted and Show point values is selected.

### Colour

Sets the colour of the highlighted axis.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "blue"

# Other axes: colour

Sets the colour of the non-highlighted axes.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "gray85"

# Show point values: font

Sets the font of the point values.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

DEFAULT VALUE · 1

### Show point values: size

Sets the size of the point values.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

Default value  $\cdot~0.75$ 

# Show point values: colour

Sets the colour of the point values.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "gray75"

# Show point values: horizontal offset

Sets the horizontal offset of the point values from the corresponding points.

DETAIL · The point values are shifted horizontally from the corresponding points with this multiple of the width of the letter 'x' (in the selected font size). Positive values shift the values to the right; negative values shift the values to the left.

default value  $\cdot$  0

# Show point values: vertical offset

Sets the vertical offset of the point values from the corresponding points.

DETAIL · The point values are shifted vertically from the corresponding points with this multiple of the height of the letter 'x' (in the selected font size). Positive values shift the values towards the top; negative values shift the values towards the bottom.

Default value  $\cdot$  1

# Show point values: digits

Sets the number of decimal places to show in the point values.

DETAIL · Possible values:  $0, \ldots, 8$ .

DEFAULT VALUE · 3

#### **Defaults**

Reverts the options to their default values.

#### OK

Saves the options, returns to the GUI, and redraws the currently displayed biplot region.

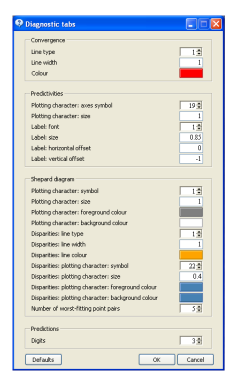
### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

### ▷ Diagnostic tabs...

Allows graphical parameters of the graphs of the diagnostic tabs to be set.

SCREENSHOT  $\cdot$  3.6



Screenshot 3.6: The  $Format \rightarrow Diagnostics$  dialogue box.

# Convergence

Lets the user change the graphical parameters of the graph of the convergence tab.

# Line type

Sets the line type.

DETAIL · Corresponds to R's 1ty argument. Possible values: 0, ..., 6.

DEFAULT VALUE · 1

#### Line width

Sets the line width.

 ${\tt DETAIL}$   $\cdot$  Corresponds to R's  ${\tt lwd}$  argument. Possible values: positive real numbers.

DEFAULT VALUE · 1

#### Colour

Sets the line colour.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "red"

### **Predictivities**

Lets the user change the graphical parameters of the points, groups and axes predictivities graphs of the points, group and axes tabs, respectively.

# Plotting character: axes symbol

Sets the symbol used to represent axes in the axes tab.

DETAIL · Corresponds to R's pch argument. Possible values: NA,  $0, \ldots, 25$ .

DEFAULT VALUE · 19

# Plotting character: size

Sets the size of the plotting characters.

DETAIL · Corresponds to R's cex argument. Larger values result in larger plotting characters. Possible values: positive real numbers.

DEFAULT VALUE · 1

### Label: font

Sets the font of the labels.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

DEFAULT VALUE · 1

#### Label: size

Sets the size of the labels.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

DEFAULT VALUE · 0.85

# Label: horizontal offset

Sets the horizontal offset of the labels from the corresponding plotting characters.

DETAIL · The labels are shifted horizontally from the corresponding plotting characters with this multiple of the width of the letter 'x' (in the selected font size). Positive values shift the labels to the right; negative values shift the labels to the left.

DEFAULT VALUE · 0

# Label: vertical offset

Sets the vertical offset of the labels from the corresponding plotting characters. Detail  $\cdot$  The labels are shifted vertically from the corresponding plotting characters with this multiple of the height of the letter 'x' (in the selected font size). Positive values shift the labels towards the top; negative values shift the labels towards the bottom.

DEFAULT VALUE  $\cdot -1$ 

# Shepard diagram

Lets the user change the graphical parameters of the Shepard diagram of the points tab.

# Plotting character: symbol

Sets the symbol used to represent the inter-point distances.

DETAIL · Corresponds to R's pch argument. Possible values: NA, 0, ..., 25.

default value  $\cdot$  1

# Plotting character: size

Sets the size of the plotting character used for inter-point distances.

DETAIL · Corresponds to R's cex argument. Larger values result in larger plotting characters. Possible values: positive real numbers.

DEFAULT VALUE · 1

# Plotting character: foreground colour

Sets the exterior colour of the plotting character used for inter-point distances. DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "gray50"

# Plotting character: background colour

Sets the interior colour of the plotting character used for inter-point distances.

DETAIL · Corresponds to R's bg argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "white"

# Disparities: line type

Sets the line type used for inter-sample disparities.

DETAIL · Corresponds to R's 1ty argument. Possible values: 0, ..., 6.

Default value  $\cdot$  1

# Disparities: line width

Sets the width of the line used for inter-sample disparities.

DETAIL · Corresponds to R's lwd argument. Larger values result in a wider line. Possible values: positive real numbers.

Default value  $\cdot$  1

# Disparities: line colour

Sets the colour of the line used for inter-sample disparities.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "orange"

# Disparities: plotting character: symbol

Sets the symbol used to represent the inter-sample disparities.

DETAIL · Corresponds to R's pch argument. Possible values: NA,  $0, \ldots, 25$ .

DEFAULT VALUE  $\cdot$  22

### Disparities: plotting character: size

Sets the size of the plotting character used for inter-samples disparities.

DETAIL · Corresponds to R's cex argument. Larger values result in larger plotting characters. Possible values: positive real numbers.

DEFAULT VALUE · 0.4

# Disparities: plotting character: foreground colour

Sets the exterior colour of the plotting character used for inter-samples disparities.

 $DETAIL \cdot Corresponds$  to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE  $\cdot$  "steelblue"

# Disparities: plotting character: background colour

Sets the interior colour of the plotting character used for inter-samples disparities.

DETAIL · Corresponds to R's bg argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "steelblue"

### Number of worst-fitting point pairs

Sets the number of worst-fitting point pairs to show at the top left of the Shepard diagram.

DETAIL · The degree of misfit between a pair of points is quantified as the difference between the inter-point distance and the corresponding inter-sample disparity. Therefore as the vertical distance from the plotting character that

represents the inter-point distance to the line representing the inter-sample disparities.

Default value  $\cdot$  5

#### **Predictions**

Lets the user change the settings associated with the predictions tab.

# Digits

Sets the number of decimal places to show in the variable value predictions of the predictions tab.

DETAIL · Possible values:  $0, \ldots, 8$ .

DEFAULT VALUE · 3

#### **Defaults**

Reverts the options to their default values.

### $\mathbf{OK}$

Saves the options, returns to the GUI, and redraws the the graphs of the diagnostic tabs.

#### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

# ⊳ Reset all...

Reverts all the parameters that can be changed in the Format menu to their default values.

SHORTCUT · Ctrl+R

A standard dialogue box appears asking the user to confirm whether or not to reset the graphical parameters of the Format menu.

DETAIL · Upon confirmation, the graphical parameters are reset and the bipot region and the graphs of the diagnostic tabs are redrawn.

### • Joint

For the construction of joint-mechanism biplots.

DETAIL · The biplots of this menu have both their points and axes determined according to a single, joint mechanism. Other biplots have their points determined from the Points menu and their axes determined from the Axes menu.

# ⊳ PCA

Constructs a PCA biplot.

DETAIL · The points correspond to the scores of the first two PCs.

SHORTCUT · 1

ALTERNATIVE TO · Covariance/Correlation, CVA, None, Regression, Procrustes, Circular non-linear

REFERENCES · Gower and Hand (1996, Chapter 2)

# 

Contructs a covariance/correlation biplot.

 $\mathtt{DETAIL}$  · If the data are only centred, a covariance biplot is produced. If the data are centred and scaled, a correlation biplot is produced.

Shortcut  $\cdot$  2

ALTERNATIVE TO · PCA, CVA, None, Regression, Procrustes, Circular non-linear REFERENCES · Gabriel (1971), Greenacre (1984), Gardner (2001, Section 2.3.2),

Gower and Hand (1996, Section 11.5.1)

#### $\triangleright$ CVA

Constructs a CVA biplot.

DETAIL  $\cdot$  The points correspond to the first two CVs. The group means can be included by clicking Additional  $\rightarrow$  Interpolate  $\rightarrow$  Sample group means. CVA biplots are invariant to the scaling of data to have unit variances. Only available when there is more than one group.

Shortcut  $\cdot$  3

ALTERNATIVE TO · PCA, Covariance/Correlation, None, Regression, Procrustes, Circular non-linear

REFERENCES · Gower and Hand (1996, Chapter 5 except Section 5.5)

### • Points

Fixes the positions of the points of those biplots which have their axes determined from the Axes menu.

DETAIL · As opposed to the biplots of the Joint menu, the Points and Axes menus are used together. The options of the Points menu fix the points, while the options of the Axes menu fix the axes.

### ▷ Distance metric

Sets the distance metric to be used in PCO and MDS.

DETAIL  $\cdot$  All the distance metrics that are available are Euclidean-embeddable. Therefore points exist in Euclidean space which have the same distances between them in Pythagorean distance as the samples have between them in the Euclidean-embeddable distance metric. If the axes previously selected in the Axes menu are no longer available because of the change in distance metric, the most appropriate replacement is automatically chosen.

REFERENCES · Gower and Legendre (1986), Gower and Hand (1996, Section A.5.1)

# o Pythagoras

The distance metric is set to Pythagoras distance.

DETAIL · 'Pythagoras distance' and 'Euclidean distance' are synomomous. However, to avoid confusion with the class of 'Euclidean-embeddable' distances, the terminology 'Pythagoras distance' is used instead. Under this distance metric, the distance between a vector  $\mathbf{x} : p \times 1$  and a vector  $\mathbf{y} : p \times 1$  is given by  $((\mathbf{x} - \mathbf{y})'(\mathbf{x} - \mathbf{y}))^{1/2}$ .

ALTERNATIVE TO · Square-root-of-Manhattan, Clark, Mahalanobis REFERENCES · Loomis (1968), Cox and Cox (2001, p. 11), Borg and Groenen (2005, p. 122)

# $\circ$ Square-root-of-Manhattan

The distance metric is set to Square-root-of-Manhattan distance.

DETAIL · Distances under this metric are the square root of those under the Manhattan or City Block distance metric. Therefore the distance between a vector  $\mathbf{x} : p \times 1$  and a vector  $\mathbf{y} : p \times 1$  is given by  $(\sum_{i=1}^{p} |x_i - y_i|)^{1/2}$ .

ALTERNATIVE TO · Pythagoras, Clark, Mahalanobis

REFERENCES · Gower and Harding (1988), Gower and Hand (1996, p. 103), Cox and Cox (2001, p. 11), Borg and Groenen (2005, p. 122)

#### o Clark

The distance metric is set to Clark distance.

DETAIL · Under this distance metric, the distance between a vector  $\mathbf{x}: p \times 1$ 

and a vector  $\mathbf{y}: p \times 1$  is given by  $\sum_{i=1}^{p} \left(\frac{x_i - y_i}{x_i + y_i}\right)^2$ .

ALTERNATIVE TO · Pythagoras, Square-root-of-Manhattan, Mahalanobis REFERENCES · Clark (1952), Gower and Ngouenet (2005)

#### Mahalanobis

The distance metric is set to Mahalanobis distance.

DETAIL · Under this distance metric, the distance between a vector  $\mathbf{x} : p \times 1$  and a vector  $\mathbf{y} : p \times 1$  is given by  $((\mathbf{x} - \mathbf{y})'\mathbf{S}^{-1}(\mathbf{x} - \mathbf{y}))^{1/2}$  where  $\mathbf{S}$  is the covariance matrix of all the vectors of observations. Mahalanobis distance is also known as 'statistical distance'.

ALTERNATIVE TO  $\cdot$  Pythagoras, Square-root-of-Manhattan, Clark References  $\cdot$  Mahalanobis (1936), Cox and Cox (2001, p. 11)

#### ▷ PCO

The points are fixed by PCO, with inter-sample dissimilarities calculated according to the chosen distance metric.

DETAIL · The coordinates of the points are taken to be the first two principal coordinates. PCO is also known as 'classical scaling'.

SHORTCUT · A

ALTERNATIVE TO · MDS Identity transformation, MDS Monotone regression, MDS Monotone spline transformation.

REFERENCES · Gower (1966), Gower and Hand (1996, Section A.5.2), Cox and Cox (2001, Section 2.2), Borg and Groenen (2005, Chapter 12)

#### $\triangleright$ MDS

The points are fixed by MDS, with inter-sample dissimilarities calculated according to the chosen distance metric.

REFERENCES · Cox and Cox (2001), Borg and Groenen (2005)

# • Random initial configuration

The next MDS algorithm will start from a random initial configuration.

DETAIL · If selected, the initial MDS configuration is taken to have points uniformly distributed over [-1,1] in both dimensions. If not selected, the initial configuration is taken to be the previous configuration of points. Useful to avoid local minima.

# • Identity transformation

Performs an MDS with the identity transformation.

DETAIL  $\cdot$  The inter-sample disparities are taken to be the inter-sample dissimilarities, calculated according to the chosen distance metric. Using an IM algorithm, the points are chosen so that stress, the sum of squared differences between the inter-point distances and the inter-sample disparities, is minimised.

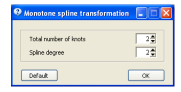
SHORTCUT · B

ALTERNATIVE TO · PCO, Monotone regression, Monotone spline transformation REFERENCES · Borg and Groenen (2005, Section 8.6)

### Monotone regression

Performs a non-metric MDS.

DETAIL · The inter-sample disparities are calculated by the up-and-down-blocks algorithm from the inter-sample dissimilarities. In essense, merely the order of the inter-sample dissimilarities is preserved. Using an IM algorithm, the points are chosen so that stress, the sum of squared differences between the inter-point



Screenshot 3.7: The Points  $\rightarrow MDS \rightarrow Monotone$  spline transformation dialogue box.

distances and the inter-sample disparities, is minimised. The Primary approach to ties or the Secondary approach to ties can be used.

SHORTCUT · C

ALTERNATIVE TO · PCO, Identity transformation, Monotone spline transformation,

REFERENCES · Borg and Groenen (2005, Sections 9.1, 9.2)

# • Monotone spline transformation...

Performs an MDS with a monotone spline transformation

DETAIL · The inter-sample dissimilarities are smoothed using monotone splines to obtain the inter-sample disparities. Using an IM algorithm, the points are chosen so that stress, the sum of squared differences between the inter-point distances and the inter-sample disparities, is minimised.

SHORTCUT · D

SCREENSHOT  $\cdot$  3.7

ALTERNATIVE TO · PCO, Identity transformation, Monotone regression

REFERENCES · Borg and Groenen (2005, Sections 9.1, 9.6)

Lets the user select the spline transformation.

# Total number of knots

The total number of spline knots, including the two exterior knots.

DETAIL  $\cdot$  Between each successive pair of knots, a polynomial of degree  $Spline\ degree$  is fit.

DEFAULT VALUE · 2

REFERENCES · Borg and Groenen (2005, Section 9.6)

#### Spline degree

The degree of the polynomial of the spline transformation.

DETAIL  $\cdot$  Between each successive pair of knots, a polynomial of degree  $Spline\ degree$  is fit.

default value  $\cdot$  2

REFERENCES · Borg and Groenen (2005, Section 9.6)

#### Default

Reverts the options to their default values.

### OK

Saves the options, returns to the GUI, and starts the MDS iteration process.

#### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the GUI.

#### • Primary approach to ties

If selected, equality in dissimilarities need not translate to equality in disparities

in monotone regression.

DETAIL · Only available when Monotone regression is selected.

ALTERNATIVE TO · Secondary approach to ties

REFERENCES · Borg and Groenen (2005, Section 9.4)

### Secondary approach to ties

If selected, equality in dissimilarities implies equality in disparities in monotone regression.

DETAIL · Only available when Monotone regression is selected.

ALTERNATIVE TO · Primary approach to ties

REFERENCES · Borg and Groenen (2005, Section 9.4)

# o In terms of principal axes

If selected, newly converged MDS configurations are rotated to be in terms of their principal axes.

DETAIL  $\cdot$  Since scaling techniques are invariant to rotation (amongst other things), expressing MDS solutions in terms of their principal axes can make different configurations more comparable.

REFERENCES · Borg and Groenen (2005, Section 24.1)

#### • Axes

Determines the axes to be shown along with the points determined from the Points menu.

DETAIL · As opposed to the biplots of the Joint menu, the Points and Axes menus are used together. The options of the Points menu fix the points, while the options of the Axes menu fix the axes. The available axes depend on the choice of distance metric and the scaling technique from the Points menu.

#### ⊳ None

No biplot axes are displayed, only points.

SHORTCUT · 0

ALTERNATIVE TO · PCA, Covariance/Correlation, CVA, Regression, Procrustes, Circular non-linear

### ▷ Regression

Regression biplot axes are shown.

DETAIL · Not available when the distance metric is Mahalanobis and the scaling technique is PCO.

SHORTCUT · 4

ALTERNATIVE TO  $\cdot$  PCA, Covariance/Correlation, CVA, None, Procrustes, Circular non-linear

REFERENCES  $\cdot$  Gower and Hand (1996, Sections 3.3.2, 3.4.3), Cox and Cox (2001, Section 3.7)

# > Procrustes

Procrustes biplot axes are shown.

DETAIL · Not available when the distance metric is Mahalanobis and the scaling technique is PCO.

SHORTCUT  $\cdot$  5

ALTERNATIVE TO · PCA, Covariance/Correlation, CVA, None, Regression, Circular non-linear

REFERENCES · Gower and Hand (1996, Sections 3.3.1, 3.4.2, A.10.2), Gower and Dijksterhuis (2004, p. 57)



Screenshot 3.8: The Additional  $\rightarrow$  Interpolate  $\rightarrow$  A new sample dialogue box.

### 

Circular non-linear biplot axes are shown.

DETAIL · Only available when the points are determined by PCO, except when the distance metric is Mahalanobis.

SHORTCUT · 6

ALTERNATIVE TO · PCA, Covariance/Correlation, CVA, None, Regression, Procrustes

REFERENCES · Gower and Harding (1988), Gower and Hand (1996, Chapter 6), Gower and Ngouenet (2005)

#### ▷ Default

Selects the default biplot type from the Axes menu.

DETAIL · The default biplot type depends on the choice of distance metric and scaling technique from the Points menu.

#### • Additional

Additional descriptors can be included in biplots.

DETAIL · When the entries of the Additional menu appear in the legend, the group names are preceded by short prefixes. The prefixes are as follows: SGM for Sample Group Mean; CH for Convex Hull; AB for Alpha-Bag; TM for Tukey Median; and CR for Classification Region.

### > Interpolate

Allows for the interpolation of additional points onto biplots.

### • A new sample...

Allows a new sample to be interpolated onto the currently displayed biplot.

DETAIL · Not available if there are no axes (as opposed to the axes being hidden).

SHORTCUT · Ctrl+N

SCREENSHOT  $\cdot$  3.8

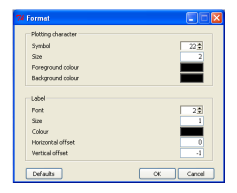
REFERENCES · Gower and Hand (1996, Sections 2.3, 6.3.1)

# Original variable values

The variable values of the sample which is to be interpolated.

DETAIL · The variable values are given in terms of the units of the original variables, and in that order. The variable values are separated by commas, without spaces. The variable values of variables that have been dragged to the kraal are not entered. If axes are dragged back onto the biplot from the kraal, the option is disabled.

DEFAULT VALUE · The mean values of the non-kraal variables.



Screenshot 3.9: The Additional  $\rightarrow$  Interpolate  $\rightarrow$  A new sample  $\rightarrow$  Format dialogue box.

### Label

The label to be used to indicate the interpolated point in the biplot or the legend.

 ${\tt DEFAULT\ VALUE\cdot Interpolated}$ 

# Label in biplot

If checked, shows the interpolated point's label in the biplot itself. Otherwise the label is shown in the legend.

DEFAULT VALUE · checked

### **Defaults**

Reverts the options to their default values.

### **Format**

See the next framed section.

# $\mathbf{OK}$

Saves the options, returns to the GUI, and redraws the currently displayed biplot region with the interpolated point.

### Off

Disables the option, and returns to the GUI, not showing the interpolated point.

### **Format**

Allows various relevant graphical parameters to be customised.

SCREENSHOT  $\cdot$  3.9

# Plotting character

Lets the user change the graphical parameters of the of the interpolated point.

# Symbol

Sets the symbol used to represent the interpolated point.

DETAIL · Corresponds to R's pch argument. Possible values: NA,  $0, \ldots, 25$ .

# default value $\cdot$ 22

Size

Sets the size of the interpolated point.

DETAIL · Corresponds to R's cex argument. A larger value results in a larger point. Possible values: positive real numbers.

default value  $\cdot$  2

# Foreground colour

Sets the exterior colour of the interpolated point.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "black"

# Background colour

Sets the interior colour of the interpolated point.

DETAIL · Corresponds to R's bg argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "black"

#### Label

Lets the user change the graphical parameters of the interpolated point label.

#### **Font**

Sets the font of the interpolated point label.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

DEFAULT VALUE  $\cdot$  2

### Size

Sets the size of the interpolated point label.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

DEFAULT VALUE · 1

#### Colour

Sets the colour of the interpolated point label.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "black"

# Horizontal offset

Sets the horizontal offset of the interpolated point label from the interpolated point.

DETAIL · The interpolated point label is shifted horizontally from the interpolated point with this multiple of the width of the letter 'x' (in the selected font size). Positive values shift the label to the right; negative values shift the label to the left.

DEFAULT VALUE · 0

## Vertical offset

Sets the vertical offset of the interpolated point label from the interpolated point.

DETAIL · The interpolated point label is shifted vertically from the interpolated point with this multiple of the height of the letter 'x' (in the selected font size). Positive values shift the label towards the top; negative values shift the label towards the bottom.

DEFAULT VALUE  $\cdot$  -1

#### **Defaults**

Reverts the options to their default values.

#### OK

Saves the options and returns to the A new sample dialogue box.



Screenshot 3.10: The Additional  $\rightarrow$  Interpolate  $\rightarrow$  Sample group means dialogue box.

#### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the A new sample dialogue box.

# o Sample group means...

Interpolates one or more of the sample group means as points.

DETAIL · This option is especially important for CVA biplots, where the optimality criterion is in terms of separation of the group means. If the sample group mean labels are shown in the legend, the group names are preceded by the prefix SGM for 'Sample Group Mean'. Not available if there are no axes (as opposed to the axes being hidden).

SCREENSHOT  $\cdot$  3.10

REFERENCES · Gower and Hand (1996, Sections 2.3, 6.3.1)

# Sample group mean(s) for

Sets the group(s) of points for which sample group means must be displayed.

DETAIL  $\cdot$  Possible options: All samples (the single overall mean is shown), all groups (the mean of each group is shown), or a specific group.

DEFAULT VALUE · All groups

# Label(s) in biplot

If checked, shows the sample group mean labels in the biplot itself. Otherwise the labels are shown in the legend.

DEFAULT VALUE  $\cdot$  checked

### Defaults

Reverts the options to their default values.

# Format

See the next framed region.

# $\mathbf{OK}$

Saves the options, returns to the GUI, and redraws the currently displayed biplot region with the sample group mean(s).

#### Off

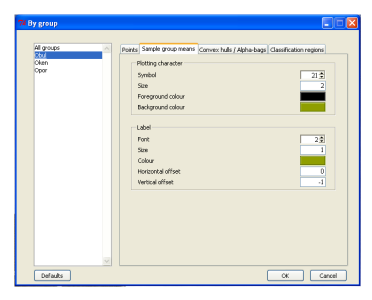
Disables the option, and returns to the GUI, not showing sample group mean(s).

#### **Format**

Allows various graphical parameters to be set by group.

DETAIL · The same dialogue box can be obtained from the Sample group means tab of the Format  $\rightarrow$  By group dialogue box.

SCREENSHOT  $\cdot$  3.11



Screenshot 3.11: The Additional  $\rightarrow$  Interpolate  $\rightarrow$  Sample group means  $\rightarrow$  Format dialogue box. This is also the Sample group means tab of the Format  $\rightarrow$  By group dialogue box

# Plotting character

Lets the user change the graphical parameters of the point that represent the sample group mean, by group.

#### Symbol

Sets the symbol used to represent the sample group mean, by group.

DETAIL · Corresponds to R's pch argument. Possible values: NA,  $0, \ldots, 25$ . DEFAULT VALUE · If there is only one group, 22. Otherwise the values are recycled by group with the values 21 to 25.

#### Size

Sets the size of the point that represents the sample group mean, by group.  $\texttt{DETAIL} \cdot \texttt{Corresponds}$  to R's cex argument. A larger value results in a larger point. Possible values: positive real numbers.

default value  $\cdot$  2

## Foreground colour

Sets the exterior colour of the point that represents the sample group mean, by group.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · "black"

### Background colour

Sets the interior colour of the sample group means, by group.

DETAIL · Corresponds to R's bg argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE  $\cdot$  If there is only one group, "red". Otherwise equally spaced colours from the hcl spectrum.

# Label

Lets the user change the graphical parameters of the sample group mean labels, by group.

#### **Font**

Sets the font of the sample group mean label, by group.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold; 3: italic; 4: bold italic.

DEFAULT VALUE  $\cdot$  2

#### Size

Sets the size of the sample group mean label, by group.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

Default value  $\cdot$  1

#### Colour

Sets the colour of the sample group mean label, by group.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE  $\cdot$  If there is only one group, "black". Otherwise equally spaced colours from the hcl spectrum.

# Horizontal offset

Sets the horizontal offset of the sample group mean label from the corresponding sample group mean, by group.

DETAIL  $\cdot$  The sample group mean label is shifted horizontally from the sample group mean with this multiple of the width of the letter 'x' (in the selected font size). Positive values shift the labels to the right; negative values shift the labels to the left.

DEFAULT VALUE · 0

# Vertical offset

Sets the vertical offset of the sample group mean label from the corresponding sample group mean, by group.

DETAIL · The sample group mean label is shifted vertically from the sample group mean with this multiple of the height of the letter 'x' (in the selected font size). Positive values shift the label towards the top; negative values shift the label towards the bottom.

DEFAULT VALUE  $\cdot -1$ 

# **Defaults**

Reverts the options of all the tabs of the dialogue box to their default values.

#### $\mathbf{OK}$

Saves the options of all the tabs of the dialogue box, and returns to the Sample group means dialogue box.

## Cancel

Cancels the changes made since the dialogue box was opened, and returns to the Sample group means dialogue box.

#### Convex hulls...

Lets the user superimpose convex hulls around one or more of the groups of points. DETAIL · Convex hulls may be drawn around all the points, around each group, or around a specific group. If the convex hull labels are shown in the legend, the group names are preceded by the prefix CH for 'Convex Hull'.

SCREENSHOT  $\cdot$  3.12



Screenshot 3.12: The Additional  $\rightarrow$  Convex hulls dialogue box.



Screenshot 3.13: The Additional  $\rightarrow$  Alpha-bags dialogue box.

ALTERNATIVE TO · Alpha-bags

REFERENCES · See the R help file of the chull function.

# Convex hull(s) for

The group(s) of points for which convex hulls must be drawn.

DETAIL  $\cdot$  Possible options: All points (a single convex hull is drawn around all points), All groups (a convex hull is drawn around each group), or a specific group.

DEFAULT VALUE · All groups

# Defaults

Reverts the options to their default values.

### **Format**

The same dialogue box can be obtained from the Convex hulls / Alpha-bags tab of the  $Format \rightarrow By$  group dialogue box. Discussed under Alpha-bags.

#### $\mathbf{OK}$

Saves the option, returns to the GUI, and redraws the currently displayed biplot region with the convex hull(s).

# Off

Disables the option, and returns to the GUI, not showing convex hulls.

### 

Lets the user superimpose alpha-bags around one or more of the groups of points. DETAIL · Alpha-bags may be drawn around all the points, around each group, or around a specific group. Alpha-bags are closely related to bagplots which can be considered to be two-dimensional boxplots. Alpa-bags enclose approximately the inner  $100\alpha\%$  of the points of a group. If the alpha-bag labels are shown in thelegend, the group names are preceded by the prefix AB for 'Alpha-Bag'. If the Tukey median labels are shown in the legend, the group names are preceded by the prefix TM for 'Tukey Median'.

SCREENSHOT  $\cdot$  3.13

ALTERNATIVE TO · Convex hulls

REFERENCES · Rousseeuw et al. (1999), Gardner (2001), Wurz, Le Roux, Gardner and Deacon (2003)

# Alpha-bag(s) for

The group(s) of points for which alpha-bags must be drawn.

DETAIL · Possible options: All points (a single alpha-bag is drawn around all points), All groups (an alpha-bag is drawn around each group), or a specific group.

DEFAULT VALUE · All groups

### Alpha

The value of alpha for the alpha-bags to be shown.

DETAIL · Approximately the inner  $100\alpha\%$  of the points of the group are enclosed in an alpha-bag.

default value  $\cdot 0.9$ 

# $Tukey \ median(s)$

If checked, the tukey median(s) of the groups are also shown.

DEFAULT VALUE  $\cdot$  checked

REFERENCES · Rousseeuw et al. (1999), Gardner (2001)

# Tukey median label(s) in biplot

If checked and Tukey median(s) are shown, the Tukey median labels are shown in the biplot, rather than in the legend.

DETAIL · Approximately the inner  $100\alpha\%$  of the points of the group are enclosed in an alpha-bag.

DEFAULT VALUE  $\cdot$  clear

# Defaults

Reverts the options to their default values.

### **Format**

See the next framed section.

#### $\mathbf{OK}$

Saves the options, returns to the GUI and redraws the biplot region with the alpha-bags as chosen.

# Off

Disables the option, and returns to the GUI, without alpha-bags.

# **Format**

Allows various graphical parameters to be set by group.

DETAIL · The same dialogue box can be obtained from the Convex hulls / Alphabags tab of the Format  $\rightarrow$  By group dialogue box. The dialogue box is the same for convex hulls.

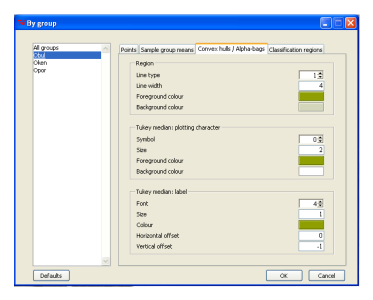
SCREENSHOT  $\cdot$  3.14

### Region

Lets the user change graphical parameters associated with convex hulls or alpha-bags.

# Line type

Sets the type of line used to draw a convex hull or alpha-bag around a group



Screenshot 3.14: The Additional  $\rightarrow$  Convex hulls / Alpha-bags  $\rightarrow$  Format dialogue box. This is also the Convex hulls / Alpha-bags tab of the Format  $\rightarrow$  By group dialogue box.

of points, by group.

DETAIL · Corresponds to R's 1ty argument. Possible values:  $0, \ldots, 6$ .

DEFAULT VALUE · 1

### Line width

Sets the width of the convex hull or alpha-bag, by group.

DETAIL · Corresponds to R's lwd argument. Larger values result in wider border for convex hulls and alpha-bags. Possible values: positive real numbers.

DEFAULT VALUE · 4

# Foreground colour

Sets the exterior colour of the convex hull or alpha-bag, by group.

DETAIL · Corresponds to R's col argument.

DEFAULT VALUE  $\cdot$  If there is only one group, #919191. Otherwise equally spaced colours from the hcl spectrum.

### Background colour

Sets the interior colour of the convex hull or alpha-bag, by group.

DETAIL · Corresponds to R's bg argument.

DEFAULT VALUE  $\cdot$  If there is only one group, #D4D4D4. Otherwise equally spaced colours from the hcl spectrum.

# Tukey median: Plotting character

Lets the user change the graphical parameters of the Tukey medians, by group.

## Symbol

Sets the symbol used to represent the Tukey medians, by group.

DETAIL · Corresponds to R's pch argument. Possible values: NA, 0, ..., 25.

DEFAULT VALUE · 0

#### Size

Sets the size of the Tukey medians, by group.

DETAIL · Corresponds to R's cex argument. A larger value results in a larger

point. Possible values: positive real numbers.

DEFAULT VALUE  $\cdot$  2

# Foreground colour

Sets the exterior colour of the Tukey medians, by group.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

<code>DEFAULT VALUE</code>  $\cdot$  If there is only one group, "red". Otherwise equally spaced colours from the <code>hcl</code> spectrum.

# Background colour

Sets the interior colour of the Tukey medians, by group.

DETAIL · Corresponds to R's bg argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · NA

# Tukey median: label

Lets the user change the graphical parameters of the Tukey median labels, by group.

### **Font**

Sets the font of the Tukey median label, by group.

DETAIL · Corresponds to R's font argument. Possible values: 1: regular; 2: bold: 3: italic: 4: bold italic.

DEFAULT VALUE · 4

### Size

Sets the size of the Tukey median label, by group.

DETAIL · Corresponds to R's cex argument. Larger values result in larger text. Possible values: positive real numbers.

default value  $\cdot$  1

# Colour

Sets the colour of the Tukey median label, by group.

DETAIL · Corresponds to R's col argument. Can be changed by clicking the colour, producing a standard colour selection dialogue box.

DEFAULT VALUE · If there is only one group, "black". Otherwise equally spaced colours from the hcl spectrum.

# Horizontal offset

Sets the horizontal offset of the Tukey median label from the corresponding Tukey median, by group.

DETAIL · The Tukey median label is shifted horizontally from the Tukey median with this multiple of the width of the letter 'x' (in the selected font size). Positive values shift the labels to the right; negative values shift the labels to the left.

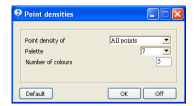
DEFAULT VALUE · 0

# Vertical offset

Sets the vertical offset of the Tukey median label from the corresponding Tukey median.

DETAIL · The Tukey median label is shifted vertically from the Tukey median with this multiple of the height of the letter 'x' (in the selected font size). Positive values shift the label towards the top; negative values shift the label towards the bottom.

DEFAULT VALUE  $\cdot$  -1



Screenshot 3.15: The Additional  $\rightarrow$  Point densities dialogue box.

### **Defaults**

Reverts the options of all the tabs of the dialogue box to their default values.

#### OK

Saves the options of all the tabs of the dialogue box, and returns to the Alpha-bags dialogue box.

#### Cancel

Cancels the changes made since the dialogue box was opened, and returns to the Alpha-bags dialogue box.

### ▷ Point densities...

Lets the user have point density estimates as a background to the biplot.

SCREENSHOT  $\cdot$  3.15

ALTERNATIVE TO · Classification regions

REFERENCES · Wand (2008), Blasius et al. (2008)

# Point density of

Sets the group of points on which the point densities are to be based.

DETAIL · Possible options: All points, or a specific group.

DEFAULT VALUE · All points

### Palette

The colour palette in which the point densities are represented.

DETAIL · Possible options: Terrain, Heat, 1, ..., 8

DEFAULT VALUE · 7

### Number of colours

The number of distinct colours to include in the palette.

 $\texttt{DETAIL}\cdot \texttt{The}$  higher the number, the smoother the image. The lower the image, the easier to distinguish between different levels.

DEFAULT VALUE · 5

# **Defaults**

Reverts the options to their default values.

#### OK

Saves the options, returns to the GUI and redraws the biplot region with the point density estimates as chosen.

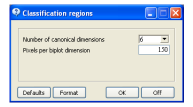
#### Off

Disables the option, and returns to the GUI, without point density estimates.

# ▷ Classification regions...

Draws classification regions in the background to a CVA biplot.

DETAIL · Pixels are coloured in according to the closest group mean in a specified



Screenshot 3.16: The Additional  $\rightarrow$  Classification regions dialogue box.

number of canonical dimensions. If the classification region labels are shown in the legend, the group names are preceded by the prefix CR for 'Classification Region'. The option is only available for CVA biplots.

SCREENSHOT  $\cdot$  3.16

ALTERNATIVE TO · Point densities

REFERENCES · Gower and Hand (1996, Section 7.3.2), Gardner and Le Roux (2005)

# Number of canonical dimensions

The number of canonical dimensions in which classification are to be made. DETAIL  $\cdot$  From 1, ..., min(the number of variables, one less than the number of groups).

DEFAULT VALUE · The maximum value.

# Pixels per biplot dimension

This number of pixels squared are classified into one of the groups and colour-coded.

DETAIL  $\cdot$  The larger the number, the finer the image, but the longer the time taken to produce the image and the more memory required. The smaller the number, the quicker the image is produced, but the grainier the image becomes.

DEFAULT VALUE · 150

#### Default

Reverts the option to its default value.

# **Format**

See the next framed region.

#### OK

Saves the options, returns to the GUI and redraws the biplot region with the classification regions shown.

#### Off

Disables the option, and returns to the GUI, without classification regions.

## **Format**

Allows various graphical parameters to be set by group.

DETAIL · The same dialogue box can be obtained from the Classification regions tab of the Format  $\rightarrow$  By group dialogue box.

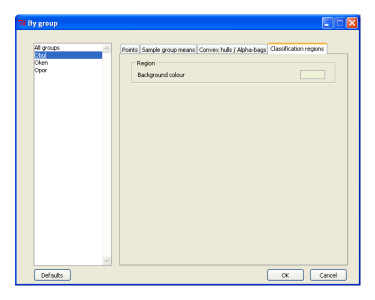
 ${\tt SCREENSHOT}\,\cdot\,3.17$ 

# Region

# $Background\ colour$

The background colour of the group's classification region.

DETAIL · Equally spaced colours from the hcl spectrum.



Screenshot 3.17: The Additional  $\rightarrow$  Classification regions  $\rightarrow$  Format dialogue box. This is also the Classification regions tab in the Format  $\rightarrow$  By group dialogue box.

#### ▷ Clear all

Disables all additional descriptors.

SHORTCUT · Ctrl+L

A standard dialogue box appears. If the user confirms, all the additional descriptors are disabled. Otherwise, the user is returned to the GUI with no changes made.

#### • Help

Some help options.

## 

Opens this manual if possible in Windows.

SHORTCUT · F1

## 

Opens the package home page at <a href="http://biplotgui.r-forge.r-project.org/">http://biplotgui.r-forge.r-project.org/</a> if an Internet connection is available.

## ⊳ Show pop-up help

If selected, pop-up help is shown as the mouse cursor is hovered over the major elements of the main view of the GUI.

#### ⊳ About…

Shows information about the author, version and license.

## ▶ Biplot region

Contains the biplot, optional title, and optional legend.

DETAIL · The biplot region is located towards the left of the GUI. The biplot itself is positioned at the centre of the biplot region. An optional title is shown above the biplot when  $View \rightarrow Show$  title is selected. An optional legend may be shown below the biplot.

The biplot region is responsive to mouse clicks and motion. Different pop-up menus appear when right clicking inside the biplot, on a point, on an axis, or outside the biplot.

Depending on how the GUI has been resized onscreen, the proportions within the biplot region onscreen may differ from the proportions in biplot regions that have been saved,

copied, printed or shown in external windows. To ensure consistency, saved and copied biplot regions always have both their width and height set to 20.32 centimeters (8 inches). Two-dimensional external-window biplot regions always have both their width and height set to 17.78 centimeters (7 inches). This is the default size of R graphs.

#### • Title

Provides a title to the biplot.

DETAIL  $\cdot$  The title is only shown if View  $\rightarrow$  Show title is selected. The default title depends on the currently displayed biplot. The biplot retains the same size and relative position, irrespective of whether or not a title is shown.

## • Biplot

A graph that displays samples as points, and variables as calibrated axes.

DETAIL · This is the graph of main interest. Right clicking on a point, on an axis, or on white space produces pop-up menus. Points and axes may be dragged from the biplot into the kraal, and be dragged back onto the biplot from the kraal.

REFERENCES · Gabriel (1971), Gower and Hand (1996)

## • Legend

A legend to the elements which appear in the biplot.

DETAIL · If necessary, a legend is provided below the biplot. The legend can identify point groups, axes, an interpolated point, sample group means, convex hulls, alpha-bags, Tukey medians and classification regions, in that order. All these elements can separately be included or excluded from the legend. The legend can contain a maximum of 16 entries at any one time. If there are more than 16 entries, only the first 16 are shown initially. The succeeding set of entries can be shown by clicking View  $\rightarrow$  Show next legend entries or right clicking inside the biplot region but outside of the biplot, and choosing the Show next legend entries option. The keyboard shortcut is Ctrl++. Similar options are available for the preceding entries. All legend entries are restricted to be 14 characters long.

When the entries of the Additional menu appear in the legend, the group names are preceded by short prefixes. The prefixes are as follows: SGM for Sample Group Mean; CH for Convex Hull; AB for Alpha-Bag; TM for Tukey Median; CR for Classification Region.

## • Inner pop-up menu

Provides access to biplot options.

DETAIL  $\cdot$  This pop-up menu appears when the user right clicks inside the biplot (as opposed to the biplot region), but not on a point or on an axis.

## 

Zooms in around the right clicked position.

DETAIL · The biplot is enlarged and the focus is shifted so that the position which was right clicked to produce the pop-up menu appears at the centre of the biplot.

#### 

Zooms out from the right clicked position.

DETAIL · The biplot is reduced in size and the focus is shifted so that the position which was right clicked to produce the pop-up menu appears at the centre of the biplot.

## 

Resets the biplot to its original zoom factor.

## ▷ Don't predict

No dynamic variable value prediction takes place.

DETAIL  $\cdot$  Not shown if there are no axes (as opposed to the axes being hidden).

ALTERNATIVE TO · Predict cursor positions, Predict points closest to cursor positions

## ▷ Predict cursor positions

The variable values of the point under the cursor position are dynamically predicted. DETAIL · For linear biplots, the orthogonal projections onto the biplot axes are shown. For non-linear biplots, circular projections are shown. The predicted variable values themselves are shown in the predictions tab. Not available when the axes are hidden, or if the biplot is interpolative rather than predictive. Not shown when there are no axes.

ALTERNATIVE TO · Don't predict, Predict points closest to cursor positions REFERENCES · (Gower and Hand, 1996, Sections 2.3, 6.3.2)

## > Predict points closest to cursor positions

The variable values of the point closest to the cursor position are dynamically predicted.

DETAIL · For linear biplots, orthogonal projections onto the biplot axes are shown. For non-linear biplots, circular projections are shown. The predicted variable values themselves are shown in the Predictions tab. Not available when the points are hidden, the axes are hidden, or if the biplot is interpolative rather than predictive. Not shown when there are no axes.

ALTERNATIVE TO · Don't predict, Predict cursor positions REFERENCES · (Gower and Hand, 1996, Sections 2.3, 6.3.2)

#### > Remove axis highlight

If an axis has been highlighted, this option removes the highlight.

DETAIL · Only available if an axis has been highlighted. Axes are highlighted by right clicking them and selecting Highlight from the pop-up menu. Not shown when there are no axes (as opposed to the axes being hidden).

## 

The same as Format  $\rightarrow$  By group in the menu bar.

## ⊳ Format axes...

The same as  $Format \rightarrow Axes$  in the menu bar.

DETAIL · Not shown when there are no axes (as opposed to the axes being hidden).

## > Save as

The same as  $File \rightarrow Save$  as in the menu bar

#### $\triangleright$ Copy

The same as  $File \rightarrow Copy$  in the menu bar

#### ⊳ Print...

The same as  $File \rightarrow Print$  in the menu bar

## • Point pop-up menu

Provides access to options associated with points.

DETAIL · This pop-up menu appears when the user right clicks a point inside the biplot. A point or axis is under the mouse if the mouse cursor changes from an arrow to a hand.

#### > Send to kraal

Sends the point which was right clicked to the kraal.

DETAIL · Sending a point to the kraal temporarily removes it from consideration. The biplot region and diagnostic tabs will automatically be updated. Points and axes may also be dragged from the biplot to the kraal.

#### ⊳ Format...

Opens the *Points* tab of the Format  $\rightarrow$  By group dialogue box at the group of the point that was right clicked.

#### • Axis pop-up menu

Provides access to options associated with axes.

DETAIL · This pop-up menu appears when the user right clicks an axis inside the biplot. A point or axis is under the mouse if the mouse cursor changes from an arrow to a hand.

## 

Highlights the axis which was right clicked.

DETAIL · This option highlights the axis which was right clicked by greying the other axes in the biplot region and diagnostic tabs. By default, the variable values of the highlighted axis are also then shown in the biplot. An axis highlight may be removed by right clicking inside the biplot and selecting Remove axis highlight from the pop-up menu. The highlight is automatically removed if the highlighted axis is moved to the kraal.

#### > Send to kraal

Sends the axis which was right clicked to the kraal.

DETAIL  $\cdot$  Sending an axis to the kraal temporarily removes it from consideration. The biplot region and diagnostic tabs are automatically updated. At least three axis must be left in the biplot.

#### $\triangleright$ Format...

Opens the Format  $\rightarrow$  Axes dialogue box at the axis that was right clicked.

#### • Outer pop-up menu

Provides access to biplot options.

#### **⊳** Show title

The same as  $View \rightarrow Show$  title in the menu bar

#### ⊳ Format title...

The same as  $Format \rightarrow Title$  in the menu bar

#### Show group labels in legend

The same as  $View \rightarrow Show$  group labels in legend in the menu bar.

## ▷ Don't show axis labels

The same as  $View \rightarrow Don't$  show axis labels in the menu bar.

DETAIL · Not shown when there are no axes (as opposed to the axes being hidden).

## 

The same as  $View \rightarrow Show$  clinging axis labels in the menu bar.

DETAIL · Not shown when there are no axes (as opposed to the axes being hidden).

## 

The same as  $View \rightarrow Show$  axis labels in legend in the menu bar.

DETAIL  $\cdot$  Not shown when there are no axes (as opposed to the axes being hidden).

## > Show Additional labels in legend

The same as  $View \rightarrow Show Additional labels$  in legend in the menu bar

#### > Show next legend entries

The same as  $View \rightarrow Show$  next legend entries in the menu bar.

## > Show previous legend entries

The same as  $View \rightarrow Show$  previous legend entries in the menu bar.

## ⊳ Save as

The same as  $File \rightarrow Save$  as in the menu bar.

## 

The same as  $File \rightarrow Copy$  in the menu bar.

> Print...

The same as  $File \rightarrow Print$  in the menu bar.

## ► Settings box

Allows the action of the biplot axes to specified, and data transformations to be effected.

DETAIL · The settings box is located towards the centre right of the GUI.

#### Action

Lets the user choose the action of the biplot axes.

DETAIL · Disabled when there are no axes, or the axes are hidden.

#### > Predict

A predictive biplot is shown.

DETAIL · Predictive biplot axes are positioned and calibrated to optimally predict variable values from the positions of points in a biplot.

ALTERNATIVE TO · Interpolate: centroid, Interpolate: vector sum

REFERENCES · (Gower and Hand, 1996, Sections 2.3, 6.3.2)

## > Interpolate: centroid

A centroid interpolative biplot is shown.

DETAIL · Centroid interpolative biplot axes are positioned and calibrated to allow for interpolation of new samples onto an existing biplot by finding a centroid.

ALTERNATIVE TO · Predict, Interpolate: vector sum

REFERENCES · Gower and Hand (1996, Section 5.3)

## ▷ Interpolate: vector sum

A vector sum interpolative biplot is shown.

 $DETAIL \cdot Vector$  sum interpolative biplot axes are positioned and calibrated to allow for interpolation of new samples onto an existing biplot by finding a vector sum.

ALTERNATIVE TO · Predict, Interpolate: centroid

REFERENCES · (Gower and Hand, 1996, Sections 2.3, 6.3.1)

## • Transformation

Lets the user choose from various data transformations.

DETAIL · These transformations are performed on the data passed to the BiplotGUI through the Biplots function (the Data argument). Data are always centred to have zero column means. Log-transformations are only available if all the non-kraal variable values of the non-kraal points are strictly positive. Irrespective of the transformation, the biplot axes are always calibrated in terms of the original variable values. In order to have biplot calibrated in terms of transformed variable values, the transformation must be performed outside of the BiplotGUI package in R, and passed to the BiplotGUI package.

#### ▷ Centre

Transforms the data to have zero column means.

 $DETAIL \cdot The mean of each column is subtracted from the elements of that column.$  The data are always centred to have zero column means.

ALTERNATIVE TO · Centre, scale; Unitise, centre; Log, centre; Log, centre, scale; Log, unitise, centre

#### > Centre, scale

Transforms the data to have zero column means, and then scales the data so that each column has unit variance.

DETAIL · The mean of each column is first substracted from the elements of that column. Each column is then divided by its standard deviation.

ALTERNATIVE TO · Centre; Unitise, centre; Log, centre; Log, centre, scale; Log, unitise, centre

#### □ Unitise, centre

Scales and shifts each column to have minimum value 0 and maximum value 1; thereafter centres to have zero column means.

DETAIL · The minimum of each column is subtracted from the elements of that column, the answers divided by the range of the (original) column.

ALTERNATIVE TO · Centre; Centre, scale; Log, centre; Log, centre, scale; Log, unitise, centre

#### 

Takes natural logarithms, then transforms the data to have zero column means.

DETAIL · Natural logarithms are taken of all elements. Thereafter the mean of each column is subtracted from the elements of that column. Only available if all the non-kraal variable values of the non-kraal points are strictly positive.

ALTERNATIVE TO · Centre; Centre, scale; Unitise, centre; Log, centre, scale; Log, unitise, centre

## 

Takes natural logarithms, transforms to have zero column means, then scales so that each column has unit variance.

DETAIL  $\cdot$  Natural logarithms are taken of all elements. Thereafter the mean of each column is first substracted from the elements of that column. Each column is then divided by its standard deviation. Only available if all the non-kraal variable values of the non-kraal points are strictly positive.

ALTERNATIVE TO · Centre; Centre, scale; Unitise, centre; Log, centre; Log, unitise, centre

## 

Takes natural logarithms, scales and shifts each column to have minimum value 0 and maximum value 1; thereafter centres to have zero column means.

DETAIL · Natural logarithms are taken of all elements. Thereafter the minimum of each column is subtracted from the elements of that column, the answers divided by the range of the (post-log) column. Only available if all the non-kraal variable values of the non-kraal points are strictly positive.

ALTERNATIVE TO · Centre; Centre, scale; Unitise, centre; Log, centre; Log, centre, scale;

## ▶ Diagnostic tabs

Consists of tabs for diagnostic graphs, dynamic variable value predictions, and quantities for export back to R.

DETAIL  $\cdot$  The diagnostic tabs are located towards the top right of the GUI.

Note: A known issue affects the diagnostic tabs of the BiplotGUI package. After a new biplot has been drawn, the graphs of the diagnostic tabs do not always display when the tabs are changed to them. The graphs can be seen by clicking to other tabs and back.

## • Convergence

If the points are determined by MDS, a graph of stress values over iterations is shown. DETAIL · If Live updates is checked, the graph is updated live. The tab is only available when the points are determined by MDS.

## ⊳ Pop-up menu

Makes various options available with repspect to the graph of the convergence tab. DETAIL · The convergence pop-up menu is obtained by right clicking the graph in the convergence tab.

#### o Show title

The same as  $View \rightarrow Show$  title in the menu bar. However, it applies to the graph in the tab rather than the biplot region.

#### o Format...

The same as  $Format \rightarrow Diagnostic\ tabs\ in\ the\ menu\ bar$ .

#### o Save as

The same as  $File \rightarrow Save$  as in the menu bar. However, it it applies to the graph in the tab rather than the biplot region.

DETAIL · Diagnostic graphs are saved with the same proportions that would be used if the currently displayed biplot region were to be saved. This ensures that the graphs can be used alongside one another. Specifically, if the biplot region has a legend, in the saved version of diagnostic graphs, an equivalent open space will be left. To remove the space, temporarily remove the legend from the biplot and save the diagnostic graph once more.

#### o Copy

The same as  $File \rightarrow Copy$  in the menu bar. However, it applies to the graph in the tab rather than the biplot region.

DETAIL · Diagnostic graphs are copied with the same proportions that would be used if the currently displayed biplot region were to be copied. This ensures that the graphs can be used alongside one another. Specifically, if the biplot region has a legend, in the copied version of diagnostic graphs, an equivalent open space will be left. To remove the space, temporarily remove the legend from the biplot and copy the diagnostic graph once more.

#### o Print...

The same as  $File \rightarrow Print$  in the menu bar. However, it applies to the graph in the tab rather than the biplot region.

DETAIL · Diagnostic graphs are printed with the same proportions that would be used if the currently displayed biplot region were to be printed. This ensures that the graphs can be professionally used alongside one another. Specifically, if the biplot region has a legend, in the printed version of diagnostic graphs, an equivalent open space will be left. To remove the space, temporarily remove the legend from the biplot and print the diagnostic graph once more.

#### o External

The same as  $External \rightarrow As$  is below the biplot region. However, it applies to the graph in the tab rather than the biplot region.

DETAIL · External diagnostic graphs are have the same proportions that would be used if the currently displayed biplot were to be placed in an external window. This ensures that the graphs can be professionally used alongside one another. Specifically, if the biplot has a legend, in the diagnostic graph will have an equivalent open space. To remove the space, temporarily remove the legend and choose the External option for the diagnostic graph once more.

#### • Points

Shows graphs of goodness of fit of the points.

DETAIL · For PCA and CVA biplots, graphs of point predictivities are shown. The first axis shows the point predictivities in the first biplot dimension. Therefore, the closer a point is to the right of the graph, the better it is represented in the first dimension of the currently displayed biplot. The distance above the diagonal line represents the contribution of the second biplot dimension. Therefore, the second axis of the point predictivities graph shows the point predictivities of the first two biplot dimensions jointly. The closer a point is to the top of the graph, the better represented it is in the two dimensions of the currently displayed biplot.

When the points in a biplot are based either on PCO or MDS, a Shepard diagram is shown in this tab. In a Shepard diagram, inter-sample dissimilarities are shown on the x-axis. On the y-axis, inter-point disparities are shown on the line, while inter-point distances are shown as points. The closer the inter-point distances are to the inter-point disparities, the better the fit. By default, the five worst-fitting point pairs are identified. A Shepard diagram is only shown if there are fewer than 250 non-kraal samples.

For covariance/correlation biplots, the tab is disabled.

REFERENCES · Borg and Groenen (2005, Section 3.3), Gower et al. (2008)

#### ▶ Pop-up menu

The same as the convergence pop-up menu. However, it acts on the graphs of the points tab.

#### • Groups

Shows graphs of goodness of fit of the groups.

DETAIL · For CVA biplots, graphs of group predictivities are shown. The first axis shows the group predictivities in the first biplot dimension. Therefore, the closer a point is to the right of the graph, the better represented the corresponding group is in the first dimension of the currently displayed biplot. The distance above the diagonal line represents the contribution of the second biplot dimension. Therefore, the second axis of the group predictivities graph shows the group predictivities of the first two biplot dimensions jointly. The closer a point is to the top of the graph, the better represented the corresponding group is in the two dimensions of the currently displayed biplot.

In other cases, the tab is disabled.

REFERENCES · Gower et al. (2008)

#### ▶ Pop-up menu

The same as the convergence pop-up menu. However, it acts on the graph of the groups tab.

## • Axes

Shows graphs of goodness of fit of the axes.

DETAIL · For PCA and CVA biplots, graphs of axis predictivities are shown. The first axis shows the axis predictivities in the first biplot dimension. Therefore, the closer a point is to the right of the graph, the better represented the corresponding axis is in the first dimension of the currently displayed biplot. The distance above the diagonal line represents the contribution of the second biplot dimension. Therefore, the second axis of the axis predictivities graph shows the axis predictivities of the first two biplot dimensions jointly. The closer a point is to the top of the graph, the better represented the corresponding axis is in the two dimensions of the currently displayed biplot.

In other cases, the tab is disabled.

REFERENCES · Gower et al. (2008)

#### ⊳ Pop-up menu

The same as the convergence pop-up menu. However, it acts on the graph of the axes tab.

#### • Predictions

The tab in which live variable value predictions are shown.

DETAIL · Dynamic variable value prediction is enabled by right clicking inside the biplot and choosing either Predict cursor positions or Predict points closest to cursor positions from the pop-up menu. If Predict cursor positions is chosen, the second column of the table in the predictions tab contains the variable value predictions corresponding to the position of the mouse cursor as it moves over the biplot. If Predict points closest to cursor positions is chosen, the variable values of the point closest to the mouse cursor as it moves over the biplot, are predicted. In this case, the third column contains the actual variable values of the points. The fourth column contains the RAE, which is calculated as (prediction - actual)/(maximum actual for variable - minimum actual for variable) ×100. Biplots are not exact, but are optimal.

Not available when there are no axes, the axes are hidden, or the biplot is interpolative rather than predictive.

REFERENCES · Gower and Hand (1996, Sections 2.3, 6.3.2)

#### • Export

Allows various quantities to be exported back to R for further manipulation or inspection. DETAIL · Depending on the representation in the biplot region, various quantities are available for export back to R. A quantity can be displayed in the R console by clicking it and clicking the Display in console button. Alternatively, a quantity can be saved to the current R workspace by clicking it and clicking the Save to workspace button.

REFERENCES · Gower and Hand (1996), Borg and Groenen (2005)

#### > Display in console

Displays the quantity highlighted in the predictions tab in the R console.

DETAIL · Only available if a quantity has been highlighted in the export tab.

## 

Saves the quantity highlighted in the predictions tab in the current R workspace under the number given in the list of quantities.

DETAIL · Only available if a quantity has been highlighted in the export tab.

#### ► Kraal

Points and axes may be dragged to the kraal, temporarily removing them from consideration.

DETAIL · The kraal is located towards the bottom right of the GUI. A point or axis can be removed to the kraal by first hovering the mouse cursor over it so that the cursor changes from an arrow to a hand, and then dragging the point or axis beyond the biplot borders. Alternatively, the point or axis can be right clicked, and Send to kraal selected from the pop-up menu. The biplot region and diagnostic tabs are then automatically updated as if the corresponding sample or variable were never part of the data set; the point or axis itself appears in the kraal. Many points and axes can simultaneously be kept in the kraal, but at least three axes must remain in the biplot.

The kraal itself is responsive to mouse clicks and motion. Different pop-up menus appear when right clicking inside the kraal, on a point, or on an axis. Points and axes may be dragged around in the kraal over a grid of possible positions, or back onto the biplot. Alternatively, the buttons below the kraal may be used to send back the points only, the axes only, or both the points and axes.

## • Kraal pop-up menu

Shows options associated with the kraal.

DETAIL  $\cdot$  The kraal pop-up menu is obtained by right clicking the kraal, excluding points and axes in the kraal.

#### > Return points

Returns all the points in the kraal to the biplot and updates the biplot and the diagnostic tabs.

DETAIL  $\cdot$  Not available when there are no points in the kraal.

#### 

Returns all the axes in the kraal to the biplot and updates the biplot and the diagnostic tabs.

DETAIL · Not available when there are no axes in the kraal.

#### > Return all

Returns all the points and axes in the kraal to the biplot and updates the biplot and diagnostic tabs.

DETAIL · Not available when there are no points or axes in the kraal.

## ⊳ Format by group...

The same as Format  $\rightarrow$  By group in the menu bar.

#### ⊳ Format axes...

The same as  $Format \rightarrow Axes$  in the menu bar.

#### • Kraal point pop-up menu

Shows options associated with a point in the kraal.

DETAIL · The kraal point pop-up menu is obtained by right clicking a point in the kraal. When the mouse cursor is above a point or axis, the cursor changes from an arrow to a hand.

#### > Return to biplot

Returns the right clicked point to the biplot and updates the biplot and diagnostic tabs.

#### ⊳ Format...

Opens the points tab of the Format  $\rightarrow$  By group dialogue box at the group of the point that was right clicked.

## • Kraal axis pop-up menu

Shows options associated with an axis in the kraal.

DETAIL · The kraal axis pop-up menu is obtained by righ clicking an axis in the kraal. When the mouse cursor is above a point or axis, the cursor changes from an arrow to a hand.

## > Return to biplot

Returns the right clicked axis to the biplot and updates the biplot and diagnostic tabs.

#### ⊳ Format...

Opens the Format  $\rightarrow$  Axes dialogue box at the axis that was right clicked.

## ▶ Other

Contains miscellaneous features.

DETAIL · These features are located in the strip at the bottom of the GUI.

#### • Progress bar

Shows the progress while performing an action.

DETAIL · A progress bar appears at the bottom left of the screen while an action is being

performed. While the GUI is busy, no new actions should be initiated. Changing tabs is not considered an action.

#### • External

Options to display the currently displayed biplot region in an external window. DETAIL  $\cdot$  Whether external windows are re-used or not can be set in File  $\rightarrow$  Options.

#### $\triangleright$ As is

Shows the currently displayed biplot region in an external R graph.

DETAIL  $\cdot$  Depending on how the GUI has been resized onscreen, the proportions within an external biplot region may differ from what is shown onscreen. To ensure consistency, external biplot regions always have both their width and height set to 17.78 centimeters (7 inches). This is the default size of R graphs.

SHORTCUT · F11

#### ⊳ In 3D

Creates a three-dimensional version of the currently displayed biplot which can be dynamically navigated.

DETAIL · This option is only available for non-MDS representations so that the first two dimensions of the three-dimensional biplot corresponds to the two-dimensional biplot shown in the GUI. Various options can be set in File  $\rightarrow$  Options. By default the left mouse button acts as a trackball for navigation, the middle mouse button allows zooming in and out, and the right mouse button sets the field of view. Not all options from the View menu or graphical parameters from the Format menu transfer to the three-dimensional biplots.

SHORTCUT · F12

REFERENCES · See the R help file of the par3d function of the rgl package (Adler and Murdoch, 2008) for details on the choices.

#### • Hide

Hides either the points or the axes of a biplot.

## > Points

Hides the points of the biplot.

## > Axes

Hides the axes of the biplot.

DETAIL  $\cdot$  Not available when there are no axes.

#### • Live updates

Toggles between showing live updates of MDS convergence (both in the biplot and diagnostic tabs), and merely showing the graphs for the final, converged configuration. DETAIL  $\cdot$  It can be informative to see the MDS configuration converges. However, if checked, the biplot region and Shepard diagram is redrawn at each update, taking more time. To speed up convergence, the checkbutton can be cleared. The number of iterations to the next update can be changed in File  $\rightarrow$  Options.

## • Stop

Stops an MDS convergence.

DETAIL  $\cdot$  Available while an MDS configuration is being found. Clicking it stops the iterations and proceeds with the current configuration, fitting biplot axes and all the rest. Users should be aware that such representations do not meet the optimality criteria and may therefore be misleading.

#### • Return points

Returns all the points in the kraal to the biplot and updates the biplot and the diagnostic tabs.

DETAIL · Not available when there are no points in the kraal.

## • Return axes

Returns all the axes in the kraal to the biplot and updates the biplot and the diagnostic tabs.

DETAIL  $\cdot$  Not available when there are no axes in the kraal.

## • Return all

Returns all the points and axes in the kraal to the biplot and updates the biplot and diagnostic tabs.

DETAIL  $\cdot$  Not available when there are no points or axes in the kraal.

# Appendix A

# History

## A.1 Version 0.0-1

29 July 2008. The first release. Contains support for

- Biplots: PCA, covariance/correlation, CVA, regression, Procrustes, circular non-linear
- Transformations: centre; centre, scale; unitise, centre; log, centre; log, centre, scale; log, unitise, centre
- Scaling methods: PCO, metric MDS (identity transformation), non-metric MDS (monotone regression), semi-metric MDS (monotone spline transformation)
- Distance metrics: Pythagoras, Square-root-of-Manhattan, Clark, Mahalanobis
- Additional descriptors: interpolation of new samples and sample group means, convex hulls, alpha-bags, point densities, classification regions
- Miscellaneous: interactive prediction, zooming, point and axis drag-and-drop, 3D biplots, extensive biplot customisation

## A.2 Version 0.0-2

- 4 August 2008. The first CRAN release.
  - $\bullet$  The Help  $\to$  Home page option is added.
  - Slight changes are made to the package help files.
  - The CountryData data set is renamed the Countries data set.
  - Two chapters are added to the manual.

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