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# Zigzag Expanded Navigation Plots in R: The R Package zenplots

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#### Abstract

We describe the features and implementation of the R package **zenplots** (zigzag expanded navigation plots) for displaying high-dimensional data according to the recently proposed zenplots.

By default, zenplots lay out alternating one- and two-dimensional plots in a zigzag-like pattern where adjacent axes share the same variate. Zenplots are especially useful when subsets of pairs can be identified as of particular interest by some measure, or as not meaningfully comparable, or when pairs of variates can be ordered in terms of potential interest to view, or the number of pairs is too large for more traditional layouts such as a scatterplot matrix. They also allow an essentially arbitrary layout of plots. A high-dimensional space can be explored in a zenplot (zenplot()) by navigating through lower dimensional subspaces along a zenpath (zenpath()) which orders the dimensions (i.e., variates) visited according to some measure of interestingness; see Hofert and Oldford (2018) for an application to S&P 500 constituent data.

The R package **zenplots** provides compact displays for high-dimensional data via the notion of zenplots, grouping of variates, and customizable displays of zigzag layouts. It accommodates different graphical systems including the base **graphics** package of R Core Team (2020b), the package **grid** of R Core Team (2020a) (and hence packages like **ggplot2** of Wickham *et al.* 2020), and the interactive graphical package **loon** of Waddell and Oldford (2020). **zenplots** handles groups of variates, partial and fully missing data, and more. One important feature is that **zenplot()** and its auxiliary functions in **zenplots** distinguish layout from plotting which allows one to freely choose and create one- and two-dimensional plot functions; predefined functions are exported for all graphical systems.

All R plots in this paper are reproducible with the vignette "selected\_features" (available in zenplots  $\geq 0.0-2$ ).

Keywords: R, data visualization, data analysis, graphics, grid, loon.

# 1. Introduction

Upper triangle scatterplot matrices for visualizing high-dimensional data through pairwise projections first appeared in Hartigan (1975). Tukey and Tukey (1981b) called them "generalized draftsman's views", the name "scatterplot matrix" (or, in short, "splom") is introduced in Tukey and Tukey (1983). The inherent limitations of scatterplot matrices for high-dimensional data were recognized soon thereafter:

"For data in several dimensions we can make triangular arrays of two-coordinatesat-a-time scatter plots. However, as the dimensionality of the data increases, two problems arise, one minor, the other more serious. First, the plots must be made smaller and smaller if they are to fit on a single page; second, and more importantly, they become less and less representative of the totality of all possible views. With increasing dimensionality the need for good methods of selection becomes ever more pressing."

(Tukey and Tukey 1981b, p. 210)

In Tukey and Tukey (1981b, Table 10.2, p. 195), "several dimensions" meant six to ten; "many" meant 11–20, "lots of" meant 21–40, and "high-dimensional" was reserved for  $d \ge 41$  dimensions. The scatterplot matrix works well for "data in several dimensions".

In our experience, beyond d = 30 to d = 50 dimensions, each cell in the scatterplot matrix becomes so small that, even after employing such tricks as sampling and alpha transparency, it can be challenging to extract useful information from this display; see, e.g., Hofert and Mächler (2014) or Hofert and Oldford (2018, Figure 3) (the latter only being able to show  $\binom{22}{2} = 231$  out of  $\binom{465}{2} = 107,880$  different pairs under consideration, so about 0.2%). In the extreme case, each cell might even be reduced to the size of a single pixel essentially rendering the scatterplot matrix as a pixel array. This as well as other limitations of scatterplot matrices have recently motivated the use of zenplots, see Hofert and Oldford (2018), for investigating high-dimensional data.

The R package **zenplots** (Hofert and Oldford 2020) presented here provides an implementation of zenplots (and accompanying tools) to help address both problems raised by Tukey and Tukey (1981b); see the quotation above. Package **zenplots** is available from the Comprehensive R Archive Network (CRAN) at https://CRAN.R-project.org/package=zenplots.

First, by providing a more compact layout, a zenplot will either accommodate many more (different) "two-coordinates-at-a-time", or "2d", plots in the same space, or, equivalently, give each individual 2d plot more relative space within any fixed size layout. Unlike scatterplot matrices, zenplots can also be easily broken over pages.

A zenplot accomplishes this by relaxing the common coordinate feature across columns (and rows) forced by the square (or triangular) layout of a scatterplot matrix (or generalized draftsman's display). With a zenplot, comparisons across a common coordinate are still available between any two 2d plots in the layout provided they have a single coordinate, or "1d", plot appearing between them. To effect this, a zenplot follows a rectilinear "zigzag" path over the display region alternating between 1d and 2d plots, all the while ensuring that neighbors along this (zen)path share a one-coordinate-at-a-time boundary.

When the objective is to visually search for plots that contain interesting patterns, the zenplot thus accommodates many more (or larger) plots and, by following the zenpath through the display, still preserves plot to plot comparisons. Moreover, the zenpath can be constrained to include only those pairs, or plot to plot comparisons, that are of interest.

The second problem raised by Tukey and Tukey (1981b) is that there are many more possible views of a high-dimensional data cloud than simply those chosen from all possible pairs of coordinates. For example, any two randomly chosen direction vectors on the (d-1)-dimensional unit sphere in  $\mathbb{R}^d$  define a projection plane which might reveal interesting structure in the data and that structure might be hidden from every one of the  $\binom{d}{2}$  2d plots defined by any two original coordinates. Moreover, there are exponentially many more of these planes than those defined only by pairs of coordinates (see Tukey and Tukey 1981a).

Zenplots address this problem in two important ways.

First, it is surprising how many "two-coordinates-at-a-time" plots can now actually be examined with modern computer displays. As a proof of concept, in a visual analysis of pairwise dependence for d = 465 dimensional data as defined by constituents of the S&P 500 stock prices, Hofert and Oldford (2018) examined all  $\binom{465}{2} = 107,880$  distinct 2d scatterplots via 164 zenplots (one per page) as a single PDF document in only 30 minutes.

Second, and more importantly, zenpaths may be constructed so that only the most interesting, or meaningful, plots are produced. Following Hurley and Oldford (2010, 2011b), imagine a graph G = (V, E) whose vertex set V is the set of all coordinates (variates) in the data and whose edge set E is the set of all meaningfully paired coordinates (variates) – typically this is a complete graph, but need not be. Weights could be attached to each edge which measure the "interestingness" of the respective two coordinates. For example, such measures were proposed as "cognostics" (for computer guided diagnostics) or "scagnostics" (scatterplot diagnostics) by Tukey and Tukey (1985) and formalized more recently by Wilkinson, Anand, and Grossman (2005). No matter how each node determines a coordinate (e.g., an original variate, some randomly or purposely chosen linear combination of original variates, or any other real-valued function of the variates), and however the weights on the edges between pairs of coordinates might be determined (e.g., statistical or scagnostic measures on that pair), a zenpath is any path on G, often it is one selected according to the weights along its edges.

In this way, following a selected zenpath provides a means to construct an interesting sequence of coordinates to be displayed within the corresponding zenplot. For example, a path of maximum (minimum) total weight would correspond to a solution to the "traveling salesman problem"; following the resulting path would display all coordinates exactly once while for each yet showing the most interesting 2d displays. The idea of a zenpath is to find those paths whose zenplot display reveals interesting structure via its coordinate sequence.

Finally, as in the original Hartigan (1975), there is no reason to restrict the 2d plots to be only scatterplots, or for the 1d plots to be, say, histograms. The pairs(...) function in R, e.g., has arguments (e.g., panel, lower.panel, diag.panel, ...) which allow the user to create essentially any display of the variates in that panel (or cell) of the scatterplot matrix. More recent authors such as Friendly (1999), Emerson *et al.* (2013), or Im, McGuffin, and Leung (2013) have undertaken generalizations of the scatterplot matrix to accommodate different pairwise and marginal displays depending on the variates in that panel of the scatterplot matrix.

Zenplots also accommodate any 1d or 2d plot. And the plot can be written in either of the R graphics packages graphics, see R Core Team (2020b), grid, see R Core Team (2020a) or

Murrell (2012) and hence **ggplot2**, see Wickham *et al.* (2020) or Wickham (2016), as well as in the more recent interactive visualization package **loon**, see Waddell and Oldford (2020).

The R package **zenplots** provides an implementation of zenplots and zenpaths with the functions **zenplot()** and **zenpath()**, respectively, as well as several other functions for exploratory data analysis and visualization. In this paper we focus on describing the functionality of the **zenplots** package, intentionally confining our illustrations to data only of "several dimensions", in the sense of Tukey and Tukey (1981b). A fuller appreciation of the package's functionality and value in visual data analysis for "high-dimensional" data is more readily had from Hofert and Oldford (2018) where the package is applied to a data example in d = 465 dimensions.

The paper is organized as follows. In Section 2 we present the structure, technical aspects and selected features of zenplot() as well as the related basic notion of a zenpath. Section 3 then focuses on zenpaths and describes how visual search can be conducted with the function zenpath(). Section 4 addresses the construction of customized zenplots and Section 5 presents more advanced features.

Some remarks concerning the history of high-dimensional data visualization are in order at this point. For several decades now, people have visualized high-dimensional space by connecting displays of low-dimensional projections. The connections are often made temporally so that one low-dimensional view dynamically and smoothly morphs into the next. The earliest versions of these would have been 3d point cloud rotations which date back to at least Ball and Hall (1970). The more general "tour" methods date to Asimov (1985) whereby arbitrary planes connected along geodesic paths are displayed over time. The earliest implementation of this was Buja, Hurley, and McDonald (1986). A series of re-implementations of these methods followed (namely XGobi of Swayne, Cook, and Buja 1998, GGobi of Swayne et al. 2008, and **rggobi** of Temple Lang, Swayne, Wickham, and Lawrence 2018), the most recent incarnations of which appear as **tourr** and **tourrGui**; see Huang, Cook, and Wickham (2012). Hurley and Oldford (2011b) constrained the planes to the orthogonal axes of the coordinate system defined in advance and proposed exploring high-dimensional spaces by sequences of planes temporally following navigation graphs. The first publicly available implementation of this strategy was **RnavGraph** of Waddell and Oldford (2011); it has most recently been implemented in the interactive and extensible data visualization package loon (see Waddell and Oldford 2020), which **zenplots** can also utilize if required. By default, zenplots accomplish the layout spatially (rather than temporally) and are therefore a novel contribution to highdimensional data visualization.

#### 2. Zenplots

We start by considering the olive data set of Forina, Armanino, Lanteri, and Tiscornia (1983); see also Azzalini and Torelli (2007). For convenience, this data set is available in the R package **zenplots**. It consists of n = 572 measurements of d = 10 variates (the geographical area and the region of origin of the olive oil, and measurements of eight fatty acid components in each of the oil specimens, namely palmitic, palmitoleic, stearic, oleic, linoleic, linolenic, arachidic and eicosenoic). As with scatterplot matrices, by default a zenplot will produce a two-dimensional point cloud (scatterplot without axes) for each variate pair and the variate's name for each one-dimensional plot. Unlike a scatterplot matrix, not all pairs of variates are displayed by default. Instead, variates are paired only if they appear next to one another in the column

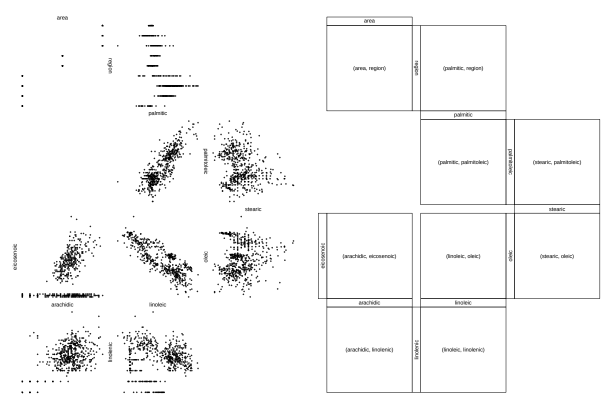


Figure 1: Zenplot of the olive data set (default, left) and again using "layout" (right).

order of the data; following column order allows variate pair sequences to be identified via column indices in the 'data.frame'. To keep the number of variate pairs low and the layout discussion simple, we consider only the nine pairs of variates as they appear in column order in the olive data; in Section 4.5, we discuss how sequences of "interesting" variate pairs might be determined and laid out for these data.

For (column ordered) variate pairs of the olive data, the zenplot is constructed as follows:

```
R> library("zenplots")
R> data("olive", package = "zenplots")
R> zenplot(olive)
```

and appears as the left-hand side of Figure 1.

The result is an alternating sequence of one-dimensional (1d) and two-dimensional (2d) plots laid out in a zigzag-like structure so that each consecutive pair of 2d plots has one of its variates (or coordinates) in common with that of the 1d plot appearing between them.

The right-hand side of Figure 1 is also a zenplot of these data but one constructed by choosing the value "layout" for arguments plot1d and plot2d:

```
R> zenplot(olive, plot1d = "layout", plot2d = "layout")
```

Now only a box containing the labels of the variates in the 1d and 2d plots is shown. This should help identify the variates that are available to construct each plot as well as the zigzag pattern of alternating plots in the display. Note also that in either display of Figure 1, the

labels of vertical 1d plots (e.g., "region", "palmitoleic" or "oleic") are oriented to reflect the left or right direction of the layout but that for horizontal 1d plots (e.g., "area", "palmitic" or "stearic") the natural reading direction is preserved for both up or down directions.

The entire set of arguments of zenplot() are seen by printing its structure:

```
R> str(zenplot)
```

```
function (x, turns = NULL, first1d = TRUE, last1d = TRUE,
 n2dcols = c("letter", "square", "A4", "golden", "legal"),
 n2dplots = NULL,
 plot1d = c("label", "points", "jitter", "density", "boxplot", "hist",
    "rug", "arrow", "rect", "lines", "layout"),
 plot2d = c("points", "density", "axes", "label", "arrow", "rect",
    "layout"),
 zargs = c(x = TRUE, turns = TRUE, orientations = TRUE, vars = TRUE,
   num = TRUE, lim = TRUE, labs = TRUE, width1d = TRUE,
   width2d = TRUE, ispace = match.arg(pkg) != "graphics"),
 lim = c("individual", "groupwise", "global"),
 labs = list(group = "G", var = "V", sep = ", ", group2d = FALSE),
 pkg = c("graphics", "grid", "loon"),
 method = c("tidy", "double.zigzag", "single.zigzag"),
 width1d = if (is.null(plot1d)) 0.5 else 1, width2d = 10,
 ospace = if (pkg == "loon") 0 else 0.02,
  ispace = if (pkg == "graphics") 0 else 0.037, draw = TRUE, ...)
```

This seemingly imposing number and variety of arguments group naturally as related to:

- Data: x.
- Plots: plot1d, plot2d, zargs, pkg, lim, labs, draw, ....
- Layout: turns, method, n2dplots, n2dcols, first1d, last1d, width1d, width2d.
- Spacing: ospace, ispace.

Each of these arguments will be discussed in some detail and illustrated in sections to come, but for now it is their grouping which merits attention because it helps answer the question: What is a zenplot?

A zenplot is first and foremost a *navigation plot*. By this we mean it is a sequence of lowdimensional plots which follow some trajectory through a higher dimensional data space for the purpose of revealing structure. Different trajectories and different 1d and 2d displays may reveal different features of the data. Choosing which trajectories and which displays amounts to navigating through the high-dimensional data space.

The trajectory is determined entirely by the data  $\mathbf{x}$ . This single argument,  $\mathbf{x}$ , provides the data and the order in which the data dimensions will be traversed in the low-dimensional trajectory. For example, in Figure 1 plots are laid out in the same numerical order in which the variates appear in the data frame. That is, for a *d*-dimensional data frame, with variates numbered  $1, \ldots, d$ , the variate pairs appear as  $(1, 2), (2, 3), \ldots, (d - 1, d)$ ; 2d plots would be constructed for each variate pair and 1d plots interspersed between them for the variate

they have in common. In this way, the determination of the trajectory is separated from its visualization. The code respects this separation in that the function <code>zenplot()</code> assumes the order given by the data x. Another function, <code>zenpath()</code>, is available to provide a variety of different orders which might be used to arrange that the data x produce a given trajectory.

The second grouping of arguments, the plots, determine the 1d and 2d plots that will be used to display the lower dimensional spaces along the trajectory. A number of plot types are built in as specific values of these arguments and, by using the information bundled as zargs, the user may also write their own display functions using any one of the R graphics packages given by pkg. Together with the trajectory, this specification of plots define the nature of the navigation through the high-dimensional space.

One could, as in Hurley and Oldford (2011b), Oldford and Waddell (2011) and Waddell and Oldford (2011, 2014, 2020) use dynamic graphics to move from one display to another thus linking the displays in time. Motion graphics very powerfully connect one low-dimensional display to the next, visually reinforcing the notion of navigation through high-dimensional space along low-dimensional trajectories. However, such temporal linking becomes increasingly burdensome on our short-term memory as the time over which the displays are transiently presented grows. And the presentation time can indeed grow quickly given that the number of variate pairs can grow quadratically with dimensionality d.

In contrast, zenplot() spatially links the displays along the trajectory by laying them out following a zigzag path pattern (hence the name "zigzag expanded navigation plot" or zenplot). Thus, it is in the third grouping of arguments, the layout arguments, which effectively define a zenplot and which distinguish it from other displays of high-dimensional data. The last grouping, spacing, is arguably also part of the layout but is not peculiar to the definition of a zenplot layout and so can be regarded as separate, more generic, layout arguments.

The next few sections deal with each of these groupings of arguments in some detail beginning with the layout arguments, being those which most distinguish a zenplot. Standard features are treated separately from those which are more customized so that the reader may skip the more advanced at first reading. Because of their simplicity, the spacing arguments will be mentioned and illustrated in the following section on the layout.

#### 2.1. Layout

Nearly all arguments of zenplot() are concerned with the physical layout of the plots. In this section, only those basic layouts (zigzag patterns) which conceptually distinguish a zenplot from other layouts are treated.

#### Zigzagging layout methods

Argument method provides a choice between the zigzag patterns "tidy", "double.zigzag", "single.zigzag" or "rectangular"; the default value is "tidy". To clearly contrast these options for method, more plots than those which appeared in Figure 1 are required. To this end, we simply double the number of columns in the olive data set so that there are now d = 20 variates to plot:

## R> olive2 <- cbind(olive, olive)</pre>

Using olive2 as the data x, the layout for the first three of the method values appears in Figure 2 produced as follows:

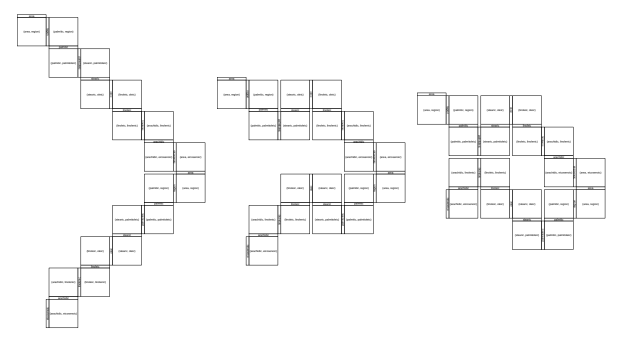


Figure 2: Zenplots of the layout of the doubled olive data set with method = "single.zigzag" (left), method = "double.zigzag" (middle), method = "tidy" (right) and n2dcols = 6.

```
R> zenplot(olive2, n2dcols = 6, plot1d = "layout", plot2d = "layout",
+ method = "single.zigzag")
R> zenplot(olive2, n2dcols = 6, plot1d = "layout", plot2d = "layout",
+ method = "double.zigzag")
R> zenplot(olive2, n2dcols = 6, plot1d = "layout", plot2d = "layout",
+ method = "tidy")
```

Both plot1d and plot2d have value "layout", and n2dcols = 6 so as to exaggerate the effect; the argument n2dcols determines the number of columns of 2d plots and is further explained in the next section. The leftmost layout is the "single.zigzag" method which follows the simplest pattern, zigzagging downwards left to right alternating 1d and 2d plots down and right as it goes. At most n2dcols columns can be used for 2d plots. Once this limit is reached, the right-hand side is reached, and the pattern then reverses to be from right to left, continuing downwards until ultimately it reaches the corresponding left side of the display area. Were there more plots to lay out, the pattern would continue repeating itself, reversing horizontal directions as each edge of the display is reached.

Note that when the display edge is reached on the right in Figure 2, the rightmost plot is a 2d plot and the change in direction is effected by moving down through a 1d plot followed by moving out left from the 2d plot below it. It is not possible to have a vertical 1d plot as the rightmost plot (unless it were the last plot of all). Minimally, the "single.zigzag" method must be able to tell that the edge has been reached and the layout must turn down and reverse directions horizontally.

The "single.zigzag" layout is easy to follow but wastes a considerable amount of display space. Had there been only ten 2d plots to display, the plots would move diagonally down

the display leaving the large areas on either side of the diagonal unused.

The "double.zigzag" method tries to make better use of this space by zigzagging up and down horizontally across the display, exhausting a double row of 2d plots before reaching the edge of the display, where it then moves down enough to reverse horizontal directions and continue as before.

The main challenge for the "double.zigzag" method is to not get "cornered" before it reverses horizontal direction. The middle display of Figure 2 shows a "double.zigzag" layout. Clearly there would be room at the top-right corner of this display for three more 2d plots to appear but using this space would trap the layout in that corner; there would be no room to turn and reverse horizontal direction. This is what we mean by getting "cornered". By the time the position of the rightmost 2d plot in the second row (from the top) is reached, the zenplot must determine that it is time to turn down and effect its horizontal reversal. This requires looking ahead a little farther than was done for reversal of the "single.zigzag" method.

While the "double.zigzag" layout is much more space efficient than is the "single.zigzag" one, there is still room for some improvement. For example, in the middle display of Figure 2, the blank space at the left of the row containing only two 2d plots seems wasted. This unused space was generated by the turning of the corner at the right of the display. Similarly, the very last 2d plot at the bottom seems to be poorly placed. This too is a consequence of changing horizontal directions, this time entirely due to anticipating the horizontal reversal. To effect the turn (on the left this time), the "double.zigzag" rule is to drop down so as to not get "cornered". However, the method will only get cornered if there are more than three 2d plots left to display. In the present case there is only one 2d plot left which could be easily accommodated by moving up instead of down.

The "tidy" method tries to make the most efficient use of the space. This requires looking ahead a little more before each turn. The rightmost display of Figure 2 shows a "tidy" layout. The first two rows are identical to that of the "double.zigzag"; so too are the rightmost two in row three, the rightmost three in row four, and the rightmost two in row five. Only the last three 2d plots are positioned in different places. For the "double.zigzag" method these are directed down and left anticipating the next possible reversal, whereas for the "tidy" method they are directed up and left to fill the empty space. Note that although there were only three 2d plots left, the "tidy" method was not in danger of being "cornered". As the position of the very last 1d plot shows, there is room for another 2d plot and for the corner to be turned.

The "tidy" method clearly has the most sophisticated look ahead and consequently compact display of the three methods. This becomes more important when there are a great many variates (dimensions) to display.

Finally, method = "rectangular" (see the vignette "selected\_features") produces a rectangular layout filled from left to right (then right to left, etc.) before moving downwards. This is an example of a method which leaves the zigzagging zenplot paradigm but can be useful for laying out 2d plots which are not necessarily connected through a variable; see zenplots for more details.

#### The number of columns containing 2d plots

As seen in the previous section, the layout of a zenplot (whichever the method) depends on

the width of the display area available. In zenplot() this width is essentially determined by the number of columns there are for 2d plots. This is specified by the value of the argument n2dcols. In Figure 2, we specified that this be n2dcols = 6 for illustrative purposes.

The default value of n2dcol is the string "letter" and was used, e.g., in Figure 1. The idea here is that the user imagines that the zenplot will be laid out to fit on a North American standard "letter" size display. Other possibilities corresponding to other standard formats are the strings "square", "A4", "golden" and "legal"; "golden" stands for the golden ratio, namely  $(1 + \sqrt{5})/2$  (interpreted as height/width).

Provided the total number of 2d plots, say  $n_{2dp}$ , is known, the number of columns and rows containing 2d plots can be determined to (approximately) respect the aspect ratio of any format. This approximation is based on the following reasoning. Let  $n_{2dr}$  and  $n_{2dc}$  denote the number of rows and columns of 2d plots. Since there is typically an empty 2d plot space at the end of a row in a zenplot, the total number of 2d plots displayed is about  $(n_{2dc}-1)n_{2dr}$ . A standard format suggests that the ratio of the number of rows to columns (i.e., height/width) be fixed at some specified scale  $s = n_{2dr}/n_{2dc}$ . This gives  $n_{2dr} = sn_{2dc}$  and hence, for known  $n_{2dp}$  and s,  $n_{2dc}$  is the solution to

$$n_{2dp} = (n_{2dc} - 1)sn_{2dc}$$
 or equivalently  $n_{2dc}^2 - n_{2dc} - n_{2dp}/s = 0.$ 

Solving the latter equation gives  $n_{2dc} = \frac{1+\sqrt{1+4n_{2dp}/s}}{2}$ . For n2dcols we essentially use the numerical value

$$n_{\rm 2dc} = \max\left\{3, \ \operatorname{round}\left(\frac{1+\sqrt{1+4n_{\rm 2dp}/s}}{2}\right)\right\},$$
 (1)

where  $n_{2dp}$  is the number of 2d plots and s is the scaling factor (height/width) given by the relevant standard. Furthermore, if necessary,  $n_{2dc}$  is increased by 1 to obtain an odd number of 2d columns in order to have a slightly more compact layout.

Alternatively, the user may supply any positive integer as the value of n2dcols.

#### The number of 2d plots displayed

In determining the layout, particularly the value of n2dcols, the total number of 2d plots to be displayed needs to be determined. This can be determined (by default) directly from the number of variates in the data given by the argument  $\mathbf{x}$ . Alternatively, the user may wish to specify a specific number of plots to be displayed only. This number must be less than or equal to d-1 where d is the number of variates appearing in  $\mathbf{x}$ .

This, and the remaining basic features of a zenplot layout are illustrated by the pair of zenplots displayed in Figure 3: The right-hand side plot shows the effect of the argument n2dplots of zenplot(). It allows to adjust the number of 2d plots displayed along the zenpath (with the obvious default to show all plots). Due to our choice n2dplots = 8, the right-hand side of Figure 3 shows only eight instead of all nine plots.

#### Omitting first or last 1d plots

Sometimes, the first or last of the 1d plots might be preferred to be omitted from the zenplot display. Examples are if there is no 2d plot after the last one (see Hofert and Oldford 2018,

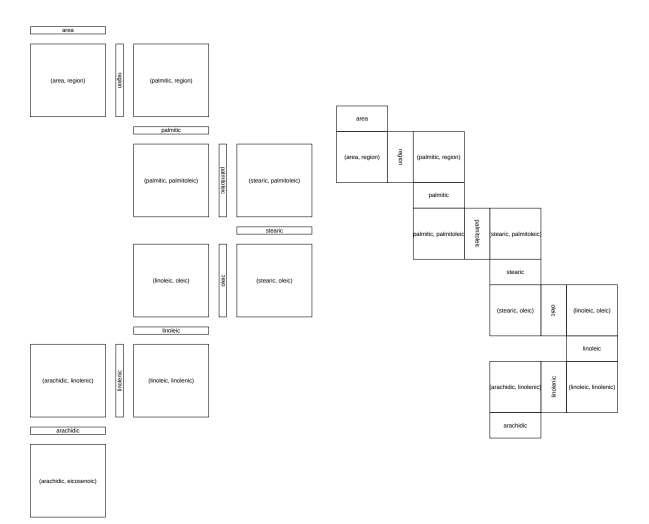


Figure 3: Zenplots of the layout of the olive data set with method = "double.zigzag", last1d = FALSE and ispace = 0.1 (left) and n2dcol = 4, n2dplots = 8, width1d = 2, and width2d = 4 (right).

Figure 9 for such an example) or to save space to provide a more compact plot. This is effected by the logical-valued arguments first1d and last1d. For example, the right-hand side of Figure 1 shows the last 1d plot, the left-hand side of Figure 3 does not.

#### Relative widths of 1d and 2d plots

The arguments width1d and width2d are non-negative integers whose ratio determines the relative space given to 1d and 2d plots. If a 2d plot occupies a square of side width2d, then every horizontal 1d plot will occupy a rectangle of width width2d and height width1d, and every vertical 1d plot will occupy a rectangle of width width1d and height width2d. See the right-hand side of Figure 3.

## Spacings between 1d or 2d plots and around whole zenplots

Users will also want to sometimes control the spacing around the individual 1d and 2d plots

as well as the spacing around the whole zenplot layout. To illustrate this, as well as some of the arguments of the previous sections, consider the two zenplots produced as follows and displayed as Figure 3:

```
R> zenplot(olive, plot1d = "layout", plot2d = "layout",
+ method = "double.zigzag", last1d = FALSE, ispace = 0.1)
R> zenplot(olive, plot1d = "layout", plot2d = "layout", n2dcol = 4,
+ n2dplots = 8, width1d = 2, width2d = 4)
```

The left-hand side of Figure 3 shows the same layout as the right-hand side of Figure 1, except for the changes described in the preceding sections and below.

The left-hand side of Figure 3 has a larger gap between all plots. This follows from ispace = 0.1, which sets the inner space (ispace to a number in [0,1]). For graphics (the default; more on that later), ispace defaults to 0, otherwise to some small positive fraction. This difference in behavior is because other graphics systems (like grid or loon) may not have a default extension of the plot region and so would lead to clipping of plot symbols near the margins of the plot region. Note that ispace can also be a vector of length four, in which case it provides the bottom, left, top and right inner space. To adjust the space around the whole zenplot, an analogous argument ospace is used to determine the zenplot's outer space (ospace).

Note that for **graphics** plots, **ispace** and **ospace** are internally set with the **par** arguments **plt** and **omd**, respectively; for **grid**-based zenplots they are set with respective viewports.

#### 2.2. Plots

A large number of plot functions for each of the arguments plot1d and plot2d can be specified via character strings. For 1d plots, these include:

"label": To show the label of the current plot variable.

"points": Scatterplots (of the data against their index).

"jitter": Jittered version of "points".

"density": Density plot based on density().

"boxplot": Box plots.

"hist": Histograms.

"rug": Rug plots.

"arrow": An arrow head indicating the direction of the zenpath.

"rect": A rectangle indicating the plot region.

"lines": Line spanning the plot region.

"layout": Showing the current plot variable and a box around it.

For 2d plots, the following are available:

"points": Scatterplots (of the two data columns under consideration).

"density": Density plot based on MASS's kde2d() (see Venables and Ripley 2002; Ripley 2020).

"axes": Coordinate axes.

"label": To show the labels of the two current plot variables.

"arrow": As for 1d plots (see above).

"rect": As for 1d plots (see above).

"layout": Showing the pair of current plot variables and a box around it.

It is also possible for the user to supply their own plot functions; this will be covered in more detail in Section 4.

#### Enforcing common plot limits

By construction, neighboring plots along a trajectory share a variate (on their common dimension/axis) and hence have common data limits on that variate. This allows the viewer to visually track the same location across adjacent plots that share that variate. By default, the limits of these shared dimensions are determined individually for each variate; this is effected by the lim argument's default value "individual".

Should the viewer wish that all dimensions have the same extent across all plots, then lim = "global" should be used. Alternatively, when the data argument x is a list (indicating groups of data and variates; see Section 2.3 and 4) setting lim = "groupwise" ensures that within each group all plots have identical data limit extents.

#### The graphical system

By default, the R package graphics is used for drawing zenplots; see pkg = "graphics". The underlying rectangular layout with rows and columns containing the 1d and 2d plots is constructed internally with layout() based on width1d and width2d; the same applies to pkg = "loon". For pkg = "grid", the layout is implemented with the more sophisticated grid.layout(). The 1d and 2d plots are placed (with placeGrob()) in a frame grob (with frameGrob()) which is returned invisibly. This grid object can further be modified, if required. The support of pkg = "grid" opens the door for other grid-based graphics systems; see, e.g., Figure 5 which shows a zenplot based on ggplot2 (the special layout will be discussed in Section 4.3). If pkg = "loon", then an interactive zenplot (e.g., with brushing, zooming, panning, and so on) is constructed via the R package loon. The resulting zenplot can then be linked to any other interactive plot in loon.

Note that plotting is done if draw = TRUE (the default). Either way, information about the underlying zenpath and layout is returned; more on this later.

#### Ellipsis arguments

All plots produced using the basic argument values described above for plot1d and plot2d are implemented using standard plot functions from the relevant package. The ellipsis arguments are passed on as extra arguments to these standard plot functions. So, e.g., if a zenplot is using pkg = "graphics" (the default), then plot arguments such as col, cex, pch and so on could reasonably be given to zenplot() whenever plot1d = "points" or plot2d = "points" were given as well; the extra arguments and their values would be passed on.

#### 2.3. Data

The data argument x of zenplot() is typically a 'matrix' or a 'data.frame'. In either case, each column/variate of x is taken to be a dimension and the trajectory moves through these dimensions in order as described earlier.

However, the value of x can also be a 'list' of objects of these types. Then each element of the list is interpreted as a group of variables which belong together. As such, each group should be visually separated from one another in the resulting zenplot. This case is covered in more detail in Section 4.

# 3. Zenpaths

Suppose the zenplot() argument x is a matrix named dataMat having d = 5 columns. By default, the zenplot will follow the trajectory through the data given by the path 1, 2, 3, 4, 5 as in

R> (path <- 1:5)

[1] 1 2 3 4 5

and the sequence of dimension pairs (1, 2), (2, 3), (3, 4), (4, 5) would determine the order and dimensions of the 2d plots.

This default sequence is not perhaps the best one to reveal interesting structure in the data. We might, e.g., wish to look at all pairs of dimensions at once in the zenplot. This could be the sequence

R> zenpath(5)

R> str(zenpath)

[1] 5 1 2 3 1 4 2 5 3 4 5

for example since every number appears beside every other number once. Or, if we had some way to measure the interestingness of any pair of dimensions, we might choose a path which visited only the most interesting pairs.

In any case, if **path** were a vector which contained the desired sequence then

R> zenplot(x = dataMat[, path])

would produce the zenplot that followed the navigation path of interest. We call such a navigation path a zenpath.

To construct zenpaths, R package **zenplots** provides a function of the same name, **zenpath()**, which takes a variety of arguments and returns a sequence of dimensions which can be used for the path.

```
function (x, pairs = NULL, method = c("front.loaded", "back.loaded",
    "balanced", "eulerian.cross", "greedy.weighted", "strictly.weighted"),
    decreasing = TRUE)
```

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Some of the methods to construct zenpaths simply depend upon the number of dimensions that are involved and ensure that all possible pairs of dimensions occur in the sequence. Since there are a great many possible sequences a few approaches are offered:

• "front.loaded": If x is an integer and method = "front.loaded" (the default), function zenpath() provides a sequence of numbers which, considering two consecutive numbers at a time, contains all pairs of the variables 1 to x sorted in such a way that the first variables appear the most frequently early in the sequence. Note that this sequence of numbers is typically not exactly of length  $\binom{d}{2}$  as the pairs have to be "connected" along a zenpath in the sense of consecutive numbers building pairs along the zenpath.

```
R> zenpath(5, method = "front.loaded")
```

[1] 5 1 2 3 1 4 2 5 3 4 5

• "back.loaded": Similar to method = "front.loaded" but with later variables appearing the most frequently later in the sequence.

R> zenpath(5, method = "back.loaded")

 $[1] \ 1 \ 2 \ 3 \ 1 \ 4 \ 2 \ 5 \ 3 \ 4 \ 5 \ 1$ 

• "balanced": Similar to method = "front.loaded" but with variables appearing in balanced blocks throughout the sequence (a so-called Hamiltonian decomposition, see Hurley and Oldford 2011b).

R> zenpath(5, method = "balanced")

[1] 1 2 3 5 4 1 3 4 2 5 1

• "eulerian.cross": For this method, two integers representing the sizes of two groups of variables must be passed to zenpath(). zenpath() returns a sequence of numbers, which, when interpreting two consecutive numbers at a time, sorts all pairs of variables such that each pair is formed with one variable from each group.

```
R> zenpath(c(3, 5), method = "eulerian.cross")
```

[1] 1 4 2 5 1 6 2 7 1 8 3 4 3 6 7 3 5 8 2

As an example, Figure 4 displays a zenplot of all pairs of olive acids only (no "region" or "area") based on an Eulerian zenpath showing the (default "front.loaded") ordering of all pairs of variables. Such a display would be of particular interest for very large data sets where simple scatterplots would have so many points overstriking one another as to obfuscate patterns in the density easily seen via contour plots. The code to generate the figure is as follows:

```
R> oliveAcids <- olive[, !names(olive) %in% c("area", "region")]
R> zpath <- zenpath(ncol(oliveAcids))
R> zenplot(oliveAcids[, zpath], plot1d = "hist", plot2d = "density")
```

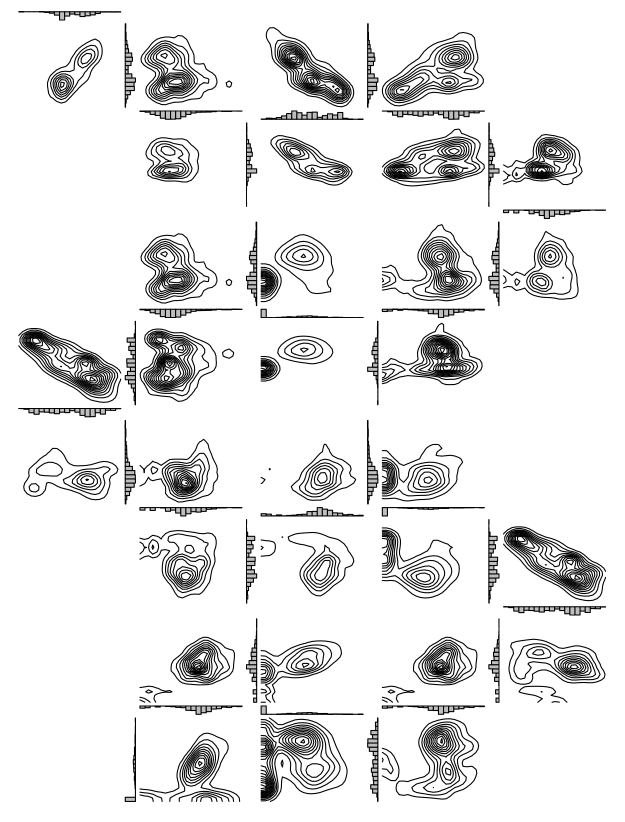


Figure 4: Zenplot of an Eulerian zenpath showing histograms (as 1d plots) and densities (as 2d plots) of all pairs of the acids of the olive data set.

The histograms point in the direction of the zigzag layout. Since plot1d = "hist" by default only shows histograms, the labeling of the variates known from the default plot1d = "label" is lost; a more sophisticated version of this plot using pkg = "grid" will be given in Section 4.1 to illustrate how more complex displays via simple plot1d and plot2d functions can be written.

Although it is possible to lay out hundreds of thousands of plots with a zenplot, see Hofert and Oldford (2018), one is often not interested in all of them. For example, Hofert and Oldford (2018, Figure 6) shows a zenplot containing only those 10 pairs of variables with largest and those 10 pairs with smallest pairwise tail dependence among all  $\binom{465}{2} = 107,880$  pairs of stocks in the S&P 500. Given a (numerical) measure of "interestingness" (such as correlation, tail dependence, etc.) assigned to all pairs of variables, one can sort all pairs according to this measure and plot the most (or also the least) interesting ones.

The zenpath() methods "greedy.weighted" and "strictly.weighted" provide the means for doing this, given that  $\mathbf{x}$  is a numeric vector or matrix or distance matrix of weights which encodes the interestingness of each pair. The argument pairs of zenpath() contains a two-column matrix, that, row-wise, contains the connected pairs of variables to be sorted according to the weights x; if pairs = NULL, a default is constructed in lexicographical order. For method = "greedy.weighted", zenpath() returns a sequence of numbers which sorts all pairs according to a greedy (heuristic) Euler path visiting each edge at least once (some edges may be visited twice since when the graph is not even, some number of edges will be duplicated to make it so and an Eulerian visiting each edge on the new graph precisely once will be constructed, see Hurley and Oldford 2011a); note that this method internally uses the function ftM2graphNEL() from the R package graph of Gentleman, Whalen, Huber, and Falcon (2020), available from Bioconductor. For method = "strictly.weighted", zenpath() returns a sequence of numbers strictly respecting the order given by the weights, so the first, second, third, etc. adjacent pair of numbers of the output of zenpath() corresponds to the pair with largest, second-largest, third-largest, etc. weight; should there be a disjunction in the path then plots will be grouped by contiguous sub-paths (see Section 4.5 for an example). If the argument decreasing of zenpath() is TRUE (the default), then the sorting by weight is done in decreasing order.

## 4. Build your own zenplots

The zenplot() arguments plot1d, plot2d and method provide sufficient functionality to satisfy a variety of common uses. Spacing arguments provide some simple aesthetic control over the layout and the use of zenpath() allows the analyst to select a trajectory through the high-dimensional data space. In this section, we describe some extended features of zenplot() which allow the user to begin to tailor a zenplot() to their particular analysis needs. This will require some familiarity with one of the graphics packages allowed by zenplot().

#### 4.1. Custom plot functions

Each named value of either argument plot1d or plot2d is implemented by calling a **zenplots** package function whose name has that argument value as prefix and the drawing package name as suffix. Between prefix and suffix the plot type 1d or 2d appears. For example, for arguments plot1d = "hist", plot1d = "points" and pkg = "grid", 1d plots are drawn by the function hist\_1d\_grid() and 2d plots by the function points\_2d\_grid().

That is, the plotting functions called within zenplot() are of the form \*\_1d\_graphics(), \*\_2d\_graphics() (for pkg = "graphics"), \*\_1d\_grid(), \*\_2d\_grid() (for pkg = "grid") and \*\_1d\_loon(), \*\_2d\_loon() for pkg = "loon"; here the \* stands for the respective string as given above. All these functions are exported from the zenplots package and so are also available to the user for direct use.

This means, e.g., that the effect of the call

```
R> zenplot(oliveAcids, plot1d = "hist", plot2d = "density",
+ pkg = "graphics")
```

would be identically produced by

```
R> zenplot(oliveAcids, plot1d = hist_1d_graphics,
+ plot2d = density_2d_graphics, pkg = "graphics")
```

Note that hist\_1d\_graphics and density\_2d\_graphics each end in "\_graphics" indicating that all of their drawing is done using only functions from the graphics package.

As the second case suggests, the user may also provide their own plot function using functionality exclusively from the named package (e.g., pkg = "graphics"). However, because it will be called within zenplot(), any user supplied plot function must also accept zargs as its first argument, the value of which will be constructed by zenplot(). The built-in plot functions named above may be helpful as templates and can also be called within any other function provided it passes on the zargs argument.

The zargs argument consists of a 'list' of further named arguments; which of these named arguments must appear in this list is determined by the zenplot() call and in particular by the value given to its argument zargs. For zenplot(), the zargs argument is merely a logical vector indicating which of the named set of arguments are to be passed to every 1d and 2d plot function as argument zargs.

For example, the default value of zargs (in the call to zenplot() is c(x = TRUE, turns = TRUE, orientations = TRUE, vars = TRUE, num = TRUE, lim = TRUE, labs = TRUE, width1d = TRUE, width2d = TRUE, ispace = match.arg(pkg) != "graphics"). If TRUE, the value of each argument (constructed within zenplot() itself), is passed to every 1d and 2d plot functions (in a named 'list') as the value of the plot function's argument zargs. The value and meaning of each argument contained in this zargs is as follows:

- x: The original data object x as provided by the user. Virtually any object can thus be passed to the underlying 1d and 2d plot functions as long as the latter two take care of appropriate plotting. As such, zenplot() completely distinguishes between layout and plotting, which provides great flexibility for data visualization. Although certainly a rare use case, setting x = FALSE in zargs avoids x being passed on. With appropriate plot1d and plot2d arguments, one can then even plot independently of the provided data object.
- turns: The character vector of turns (either computed or user-provided). This has the advantage of each 1d and 2d plot knowing all turns before plotting is done and thus the layout of the zenplot. Typically most helpful are the turns into and out of the current plot position. For example, one could have all plots colored blue (red) which turn down (up) out of the current position along the zenpath.

- orientations: The character vector consisting of the plot orientations ("s" for square (so 2d) plots, "h" for horizontal and "v" for vertical plots) as determined internally based on the turns. This information can be used, e.g., to determine the orientation of plot labels in a zenplot.
- vars: A two-column matrix containing, for each 1d and 2d plot (so each row), the (pairs of) variable(s) which is (are) to be plotted in the current plot; for 1d plots, the respective row simply shows the same variable twice.
- num: The current plot number among all 1d or 2d plots along the zenpath. This is especially helpful for indexing, say, a row in vars to find out which variable(s) was/is (were/are) to be plotted previously, currently and next.
- lim, labs, width1d, width2d, ispace: (Already discussed) arguments of zenplots() that are passed on to the underlying 1d and 2d plot functions.

Any user defined 1d or 2d plot function can then access any of these values from its zargs argument provided they were marked for use by the logical values of the argument zargs of the zenplots() call. This gives the user access to all of the functionality used by the built-in plotting functions.

Furthermore, plot1d or plot2d can also be NULL, in which case an "empty plot" is generated, so no visible plotting is done.

Besides **zargs**, the ellipsis arguments ("...") are also passed to the underlying 1d and 2d plot functions. If only one of the two functions shall obtain additional graphical parameters, say, one can proceed as already seen in the code for the right-hand side of Figure 6, namely by providing a corresponding 1d or 2d plot function.

#### 4.2. Custom layouts – as the plot turns

The defining layout of a zenplot is the "zigzag" pattern determined by the value of the zenplot() argument method. The various choices were described and illustrated in Section 2.1 and all zenplots so far considered have followed one of these zigzag layout methods. The attentive reader may have noticed, however, that one of the layout arguments mentioned in Section 2, namely the turns argument, did not appear in the basic layout discussion of Section 2.1. This is because the turns argument, if specified, will override the method argument and so can be used to produce a zenplot which need not follow a zigzag pattern; normally, the argument turns is not given but is determined within zenplot() based on the value of its method argument.

What is meant by turns? These are directional instructions "d", "l", "u" and "r", being short for the directions "down", "left", "up" and "right", which describe the next direction that the layout path follows upon completion of that plot. Each 1d or 2d plot has a directional instruction associated with it which describes the direction to be taken from that plot to the next. In this way, the entire layout path can be thought of as a sequence of directional instructions; this sequence is the value of turns.

For example, consider the most recent zenplot given in Figure 4. The zenplot begins with the histogram (a 1d plot) in the top-left corner. The next plot in the layout is immediately below the histogram so that the directional instruction from the histogram is "d", meaning move down. Moving down leads to the 2d density contour plot. From this plot the layout moves right to the next histogram so this first density contour has exiting directional instruction "r".

This leads to the next histogram from which again we move "right" to the next density plot from which we move down to the next histogram, and so on. The value of turns associated with this plot is the sequence "d", "r", "r", "d", "d", "r", "r", "u", "u", "r", "r", "d", "d", "r", "r", "d", "d", "l", "d", "d", "l", "l", "u", "u", "u", "d", "d", "1". Note that there is no turn associated with the last histogram at the lower-left of the Figure 4 because no plot follows this histogram in the layout.

Note the zigzag pattern apparent in both these turns and in the display layout. In the next section, we consider a layout which does not follow any of the zigzag patterns considered so far, though it is a zigzag spiral. In general, **turns** need not follow a zigzag of any sort.

#### 4.3. Custom layout and plots – a spiral of ggplots2 example

For most applications, the sequence of turns provided by zenplot() via the method argument will be all that is needed for a compact layout. The value of the turns argument is that it provides the user a means to construct new layouts which may be more appropriate to their analysis.

To illustrate this, and to also show how plots from **ggplot2** can be incorporated, we consider how plots might be laid out in a spiral. Though perhaps a somewhat fanciful layout, it will serve to show how much control the user has over layout should they desire it or their application warrant it. Even over-plotting of one plot on top of another could be possible (e.g., via the **grid** package), though would not be generally recommended.

Because the zenpath is laid out according to the given turns, the length of turns must match that of the zenpath. For the oliveAcids data we might therefore choose the following path and sequence of turns:

which already begins in a different way in that the layout moves "left" out from the first plot. The resulting layout starts at the center of the display area and moves in a zigzag pattern but now in a (counter-clockwise) spiral instead of any of the row-filling patterns seen earlier.

Note that there is no restriction on **turns** so that the layout is limited only to following a continuous path of alternating 1d and 2d plots. This flexibility allows analysts to tailor the layout to the needs of the problem.

As an example, we might prefer to use the plotting paradigm given by the R package **ggplot2**. In this case, we need to write custom 1d and 2d plotting functions. Here, we write a custom 2d plot function.

```
R> library("ggplot2")
R> stopifnot(packageVersion("ggplot2") >= "2.2.1")
R> ggplot2d <- function(zargs) {</pre>
```

```
+
     r <- extract 2d(zargs)</pre>
     num2d <- zargs$num/2</pre>
+
     df <- data.frame(x = unlist(r$x), y = unlist(r$y))</pre>
     p \leftarrow ggplot() + geom_point(data = df, aes(x = x, y = y), cex = 0.1) +
       theme(axis.line = element blank(), axis.ticks = element blank(),
         axis.text.x = element_blank(), axis.text.y = element_blank(),
         axis.title.x = element_blank(), axis.title.y = element_blank())
+
     if (num2d == 1) p <- p +
+
       theme(panel.background = element_rect(fill = "royalblue3"))
     if (num2d == (length(zargs$turns)-1)/2) p <- p +
+
       theme(panel.background = element_rect(fill = "maroon3"))
+
     ggplot_gtable(ggplot_build(p))
+
   7
+
```

Note the use of extract\_2d(zargs) to get the x and y coordinates. The principal additions to ggplot() are the removal of several default plot elements. Note also the use of num2d <- zargs\$num/2 to get the total number of 2d plots in the zenplot so that the first and last ones can be distinguished by background color (respectively, "royalblue3" and "maroon3"). Finally, because ggplot() is written using the grid package, a suitable grid object must be returned.

The spiral zenplot, beginning at the center ("royalblue3") and zigzagging counter-clockwise to end at the bottom-right ("maroon3") and using ggplot(), is now simply constructed as

```
R> zenplot(oliveAcids[,path], turns = turns, pkg = "grid",
+ plot2d = function(zargs) ggplot2d(zargs))
```

See Figure 5.

#### 4.4. Data groups

In Section 2.3 it was noted that the data argument x might also be a list. In such cases, zenplot() assumes that each element of the list is itself a data set comprised of its own set of variates (dimensions).

The Italian growing region (e.g., "Umbria", "Sicily") is known for each row (i.e., oil) of the olive data and these regions have been grouped into three broad geographic areas: "South", "Sardinia", and "Centre.North". It would be of interest then to study the relationships between the variates for olive oils within each area and also to see how these relationships might differ between areas. Zenplots provides a layout for observing relationships by group.

To this end, we take the **oliveAcids** data and split it into groups according to geographic area as follows:

```
R> oliveAcids.by.area <- split(oliveAcids, f = olive$area)
R> names(oliveAcids.by.area)[3] <- gsub("\\.", " ",
+ names(oliveAcids.by.area)[3])
R> names(oliveAcids.by.area)
```

[1] "South" "Sardinia" "Centre North"

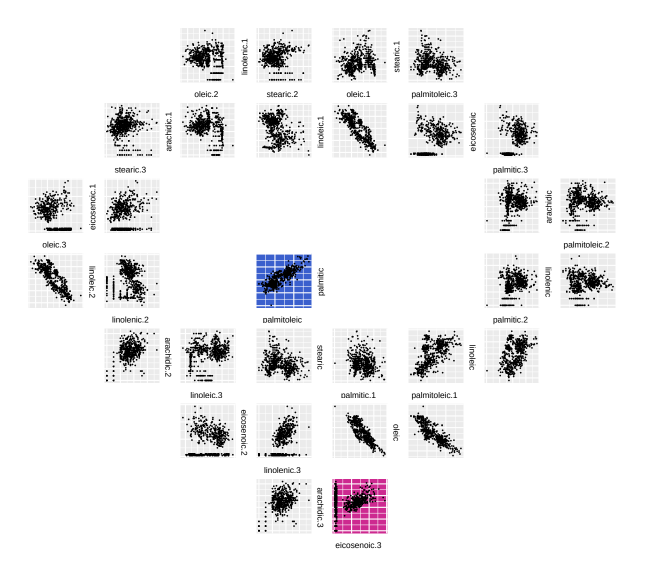


Figure 5: A zenplot displaying all pairs of acid components of the olive data at least once. In particular, this zenplot shows that one can use one's own turns (provided by the argument turns) and that plotting can also be done with **ggplot2**.

The object oliveAcids.by.area is a list having three components, the first, second and third containing 323, 98 and 151 measurements of the eight acid components, respectively. Zenplots of these data groups can now be produced as

```
R> zenplot(oliveAcids.by.area, labs = list(group = NULL))
R> zenplot(oliveAcids.by.area, lim = "groupwise", labs = list(sep = " - "),
+ plot1d = function(zargs) label_1d_graphics(zargs, cex = 0.8),
+ plot2d = function(zargs)
+ points_2d_graphics(zargs, group... = list(sep = "\n - \n")))
```

which appear as the left- and right-hand sides of Figure 6.

Following the zigzag in the left zenplot of Figure 6, we can see the groups separate first at the 2d plot containing only the geographic area names "South" and "Sardinia" and then farther

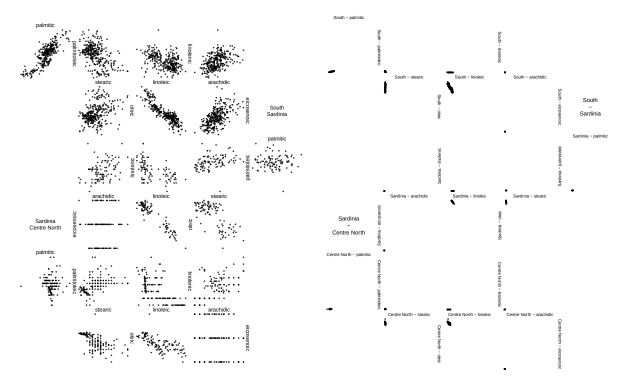


Figure 6: Zenplots showing the grouping feature (both left and right) and label feature (left: no group labels in 1d plots; right: groupwise scaling, label separators, downsizing labels).

on at the 2d plot containing only area names "Sardinia" and "Centre North". These two 2d plots mark the boundaries between the groups, which in order are "South", "Sardinia", and "Centre North". At a glance, we can see the greater number of observations in the "South" area, and the coarse granularity of the measurements apparent in much of the data from the "Centre North" (suggesting rounding or some other source of measurement error). Comparing relationships, one can also see for example that the acids palmitic and palmitoleic appear positively correlated in the "South" but possibly uncorrelated in "Sardinia" and even "Centre North". Other similarities and differences (e.g., measures on eicosenoic) between groups can also be readily observed.

In both zenplots of Figure 6, the displays follow the zigzag pattern for the variates that appear within each data group (each of which can have its own number of observations); the zigzag pattern continues on from one group to the next with a visually separating 2d plot showing the names of the groups to either side. This visual separator plot is constructed as a 2d plot function, group\_2d\_graphics() (and, for the other graphical systems, group\_2d\_grid() and group\_2d\_loon(), following the naming convention of Section 4.1), that simply displays the group labels in the order in which they appear indicating the transition from one group to the other (see "South" and "Sardinia" as well as "Sardinia" and "Centre North"; the first/top label indicates the preceding group along the zenpath, the second/bottom label indicates the next group). In the right-hand side zenplot, we use a more sophisticated option, namely the possibility to pass arguments to group\_2d\_graphics() with the argument group.... We use this to visually separate the group labels a bit better in this case. To this end, note that we need to provide a user-defined function (which just calls points\_2d\_graphics() with one argument different from the default).

In the right-hand side zenplot, every 1d plot has the group name appear before that variate name (separated by a hyphen). This was effected by specifying the argument labs = list(sep = " - "). In contrast, the left-hand side zenplot omitted the group labels from its 1d plots by specifying labs = list(group = NULL). Typically, labs is a list containing the components group (group label basename or labels for the groups; can be NULL in which case group labels are omitted), var (variable label basename or labels for the variables; can be NULL in which case variable labels are omitted), sep (separator between group and variable labels) and group2d (a logical indicating whether the group\_2d\_\*() functions should still plot group labels even if group = NULL).

Also note that because the resulting 1d plot labels would be too large when the group name was prepended, a user-defined function was provided as the argument of plot1d so that the size of this label could be reduced via the (ellipsis) argument cex = 0.8 passed on to the built-in plotting function label\_1d\_graphics() (see Section 4.1 for more complex examples). Each displayed 2d plot also has its own limits (lim = "individual", the default) as seen in the left-hand zenplot of Figure 6. In the right-hand zenplot, common limits are used within each group (lim = "groupwise"). The effect is that for these data the relationships between acid concentrations within any group have been obscured by the common scale across variates within groups (especially compared to when lim = "individual" as in the leftmost zenplot). Even though all proportions of olive acids must sum to 1 for each oil (or to 10,000 on the  $100 \times \%$  recorded values in the data), the oils are seen to contain far more **oleic** than any other fatty acid in the data. In all areas, oleic has so much higher concentration than the rest that on a common scale the others are compressed into the lower end of their axes. A third choice for the lim argument would be to have identical limits across all plots regardless of groups (lim = "global"). For the olive data, the effect would be nearly identical to that of lim = "groupwise".

#### 4.5. Custom zenpaths

The R package **zenplots** provides auxiliary functions for connecting and extracting pairs along a zenpath or for obtaining a list of matrices for plotting a zenplot with grouping feature. We will now briefly present these functions.

Say our measure of "interestingness" is "convexity" according to scatterplot diagnostics (scagnostics) and we are interested in the zenpath containing the six most convex pairs. We could proceed as follows. In a first step, we build a (d, d)-matrix, which, in the (i, j)th entry, contains the convexity measurement of the pair (i, j); this is the matrix M below.

```
R> library("scagnostics")
R> Y <- scagnostics(oliveAcids)
R > X <- Y["Convex", ]
R> d <- ncol(oliveAcids)</pre>
R> M <- matrix(, nrow = d, ncol = d)
R> M[upper.tri(M)] <- X
R> M[lower.tri(M)] <- t(M)[lower.tri(M)]</pre>
R > round(M, 5)
                                            [,5]
                                                     [,6]
         [,1]
                  [,2]
                           [,3]
                                   [,4]
                                                              [,7]
                                                                       [,8]
```

[2,]0.48952NA0.422760.504990.445910.358550.358460.31729[3,]0.463430.42276NA0.397000.363940.313160.295340.33709[4,]0.458870.504990.39700NA0.464540.366160.294510.34888[5,]0.439140.445910.363940.46454NA0.319770.314430.36750[6,]0.345830.358550.313160.366160.31977NA0.537260.34001[7,]0.312590.358460.295340.294510.314430.53726NA0.22231[8,]0.284130.317290.337090.348880.367500.340010.22231NA

Next, we use the entries in M to determine the zenpath of strictly ordered pairs (according to the ordering with respect to "convexity"); this can be done with zenpath(, method = "strictly.weighted") as described in Section 3.

```
R> zpath <- zenpath(M, method = "strictly.weighted")
R> head(M[do.call(rbind, zpath)])
```

[1] 0.5372599 0.5049945 0.4895179 0.4645377 0.4634277 0.4588675

Note that method = "strictly.weighted" returns a list of vectors of length two since there is no guarantee to obtain connected pairs when given weights have to be strictly respected. To obtain connected pairs along this list, we provide the function connect\_pairs(); see the last part of this section.

We can now conveniently use the function extract\_pairs() as follows to extract the pairs we are most interested in, namely the first six along the zenpath.

```
R> (ezpath <- extract_pairs(zpath, n = c(6, 0)))
[[1]]
[1] 7 6
[[2]]
[1] 4 2
[[3]]
[1] 2 1
[[4]]
[1] 5 4
[[5]]
[1] 3 1
[[6]]</pre>
```

[1] 4 1

The function graph\_pairs() (also depending on ftM2graphNEL() from the R package graph on Bioconductor) can be used to depict these six pairs in a graph, where an edge indicates that the adjacent nodes build a pair; see Figure 7.

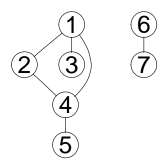


Figure 7: Graph representing those six pairs along the zenpath which exhibit largest convexity; an edge corresponds to a pair between the adjacent variables.

```
R> library("graph")
R> plot(graph_pairs(ezpath))
```

As we can see, some of the pairs are naturally connected: For example, the pairs with secondand third-largest convexity measure share the variable 2 and thus could be (zen)plotted in the same group of variables (4, 2, 1) (so plotting the pairs (4, 2) and (2, 1) with the variable 2 along the joint axis). How can we find connected pairs along a zenpath which still strictly respect the order given by "convexity"? This can be achieved with the function connect\_pairs(), which provides us with a "compactified" zenpath which then also leads to a more compact zenplot.

```
R> (cezpath <- connect_pairs(ezpath))
[[1]]
[1] 7 6
[[2]]
[1] 4 2 1
[[3]]
[1] 5 4
[[4]]
[1] 3 1 4</pre>
```

We can use the function groupData() to group our data into the groups determined by cezpath.

```
R> oliveAcids.grouped <- groupData(oliveAcids, indices = cezpath)
```

Lastly, we can plot the corresponding (grouped) zenplot, see Figure 8.

R> zenplot(oliveAcids.grouped)

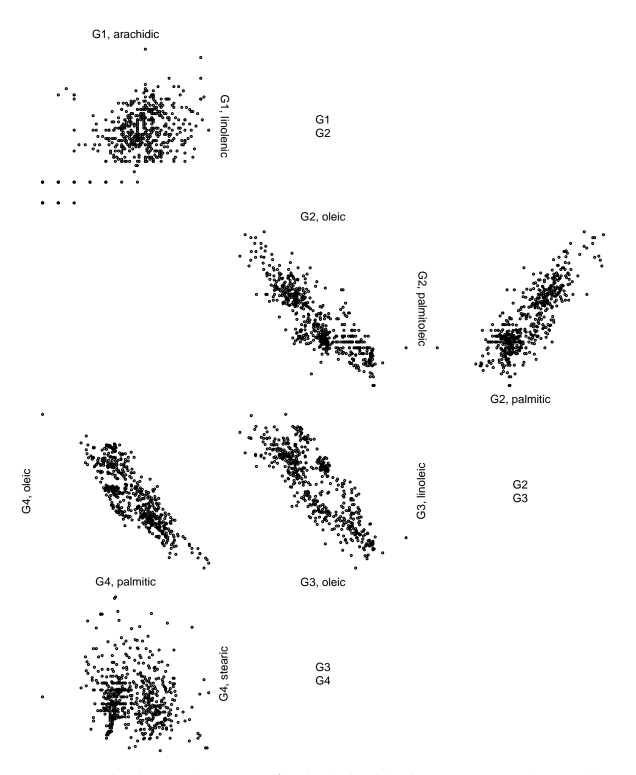


Figure 8: Zenplot showing the six pairs of acids which exhibit largest convexity; also note the U-turn at the end.

# 5. Advanced features

In the previous section, a variety of ways in which zenplots might be customized to suit the analysis were described and illustrated. There could be many more such illustrations given. In fact, an immense amount of control over the zenplot display is available to the user. This can be used to construct nearly arbitrary displays beyond any we have imagined here. To facilitate such creativity, a number of auxiliary software tools are provided in the **zenplots** package. This section provides detail on these tools so that the user can create their own novel layouts.

Section 5.1 describes the structure of a zenplot and highlights some of its important elements. In particular, the occupancy matrix is introduced which provides a potentially useful rectangular array description of the positions occupied by every plot in the display.

In Section 5.2 several important functions useful to create plots are described. Particularly important are those functions which extract the data so that it can be accessed by the plot functions. These include the function extract\_2d() seen earlier, its 1d counterpart extract\_1d(), as well as all of the components which each function returns. Two important working functions are burst() and unfold(). The first, as its name suggests, takes any of the data structures acceptable to zenplot() and "bursts" these data into an easy to use common form. The second, unfold(), is the major function underpinning zenplot(). This name is intended to suggest its functionality which is to take a high-dimensional hypercube (all coordinate pairs of the high-dimensional space) and "unfold" it so that it may be laid flat in the two-dimensional plane. The unfolding determines the layout of the 2d plots. The algorithmic details of this layout (and other functions on which it depends) are described in Appendix A. In that appendix, the entire top down structure of zenplot() and the functions on which it depends are given. The greatest detail, however, will be found in the source code itself.

#### 5.1. The structure of a zenplot

As mentioned before, zenplot(), besides plotting, invisibly returns a list containing information about the zenpath and layout. For Figure 1, e.g., we obtain:

```
R> res <- zenplot(olive, plot1d = "layout", plot2d = "layout", draw = FALSE)
R> str(res)
List of 2
 $ path :List of 3
              : chr [1:19] "d" "r" "r" "d" ...
  ..$ turns
  ..$ positions: num [1:19, 1:2] 1 2 2 2 3 4 4 4 5 6 ...
  ...- attr(*, "dimnames")=List of 2
  ....$ : NULL
  .....$ : chr [1:2] "x" "y"
  ..$ occupancy: chr [1:8, 1:6] "" "" "" ...
 $ layout:List of 6
  ...$ orientations : chr [1:19] "h" "s" "v" "s" ...
  ..$ dimensions : num [1:19] 1 2 1 2 1 2 1 2 1 2 ...
                  : num [1:19, 1:2] 1 1 2 3 3 3 4 5 5 5 ...
  ..$ vars
```

```
....- attr(*, "dimnames")=List of 2
.....$ : NULL
.....$ : chr [1:2] "x" "y"
..$ layoutWidth : num 33
..$ layoutHeight : num 44
..$ boundingBoxes: num [1:19, 1:4] 0 0 10 11 11 11 21 22 22 22 ...
....- attr(*, "dimnames")=List of 2
.....$ : NULL
.....$ : NULL
.....$ : chr [1:4] "left" "right" "bottom" "top"
```

Besides the components already discussed before (such as turns, orientations and vars), the return object of zenplot(), a list of length two, also contains the plot dimensions (as part of the layout component). Furthermore, it contains occupancy and positions:

R> res[["path"]][["occupancy"]]

	[,1]	[,2]	[,3]	[,4]	[,5]	[,6]
[1,]		"d"				
[2,]		"r"	"r"	"d"		
[3,]				"d"		
[4,]				"r"	"r"	"d"
[5,]						"d"
[6,]	"1"	"1"		"d"	"1"	"1"
[7,]		"u"		"d"		
[8,]		"u"	"1"	"1"		

R> head(res[["path"]][["positions"]])

x y [1,] 1 2 [2,] 2 2 [3,] 2 3 [4,] 2 4 [5,] 3 4 [6,] 4 4

The occupancy matrix (component occupancy) is a matrix which reflects the layout containing all 1d and 2d plots underlying a zenplot. Its elements are "" (for positions not occupied by a 1d or 2d plot), "1", "r", "d" or "u" for left, right, downward or upward turns out of the current position in the occupancy matrix, respectively. For example, the entry (2, 3) in the occupancy matrix is "r", so the turn out of the position (2, 3) is to the right. Note that zenplots also provides the auxiliary function convert\_occupancy() which allows to convert the occupancy matrix to contain different characters for the five possible states (with a default resembling arrow heads to easily spot the zenpath).

The *i*th row in the 2-column matrix **positions** provides the position of the *i*th plot in the occupancy matrix. For example, the fifth row is (3, 4), meaning that the fifth plot will be plotted in the position (3, 4) of the layout corresponding to the occupancy matrix.

Finally, let us remark that for specific graphical systems, more information might be returned. For example, for pkg = "grid", the whole plot as a grid object is also returned.

#### 5.2. Tools for writing 1d and 2d plot functions

We now present selected tools which help writing user-specific 1d and 2d plots. We also go into some more detail concerning the main underlying paradigm behind zenplots.

Bursting the data and extracting the current plot variables

As mentioned before, the data argument x of zenplot() is typically a 'vector', 'matrix', 'data.frame' or a 'list' of such. The auxiliary function burst() obtains the arguments x and labs (a list of length four as described in Section 4.4) and returns a list with the following components:

- **xcols**: A list containing the columns of  $\mathbf{x}$  (so  $\mathbf{x}$  "burst" into its columns). These columns are named and the names contain, by default, the combined group and variable labels.
- groups: The (group) number the respective column belongs to; all equal to 1 if x does not consist of different groups.
- vars: The (variable) number the respective column belongs to within its group.
- glabs: The group labels (NULL if x does not consist of different groups).

vlabs: The variable labels.

See ?burst for various examples showing how burst() works.

The functions extract\_1d() and extract\_2d() obtain the single argument zargs and extract (hence the name) useful information for most 1d and 2d plots, respectively. These are widely used in the exported \*\_1d\_\*() and \*\_2d\_\*() functions, respectively, and make use of burst(). In particular, extract\_1d() checks whether zargs contains x, orientations, vars, num, lim and labs (all discussed before) and returns a list with components:

x: The (named) data column to be plotted in the current 1d plot.

xcols, groups, vars, glabs, vlabs: These are the components as returned by burst().

horizontal: A logical indicating whether the current 1d plot is horizontal or not (so vertical).

xlim: The axis limits for the current 1d plot.

Similarly, extract\_2d() checks whether zargs contains x, vars, num, lim and labs and returns:

x: The (named) data column to be plotted as x variable in the current 2d plot.

y: The (named) data column to be plotted as y variable in the current 2d plot.

xcols, groups, vars, glabs, vlabs: These are the components as returned by burst().

xlim: The x axis limits for the current 2d plot.

ylim: The y axis limits for the current 2d plot.

same.group: A logical indicating whether the two data columns x and y belong to the same
group or not.

Simply passing zargs to the 1d and 2d plot functions \*\_1d\_\*() and \*\_2d\_\*() is very convenient. It also allows one to easily write new 1d and 2d plot functions without the need of having to deal with a lot of different arguments; we have already seen the short function function(zargs) label\_1d\_graphics(zargs, cex = 0.8) as plot1d in the code for producing the right-hand side of Figure 6. It is thus natural to have the extractor functions extract\_1d() and extract\_2d() available to extract the actual information to be used for plotting by most 1d and 2d plot functions. For example, here is the definition of points\_2d\_graphics():

```
R> points_2d_graphics
```

```
function (zargs, cex = 0.4, box = FALSE, add = FALSE, group... = NULL,
  plot... = NULL, ...)
Ł
  r <- extract_2d(zargs)</pre>
  xlim <- r$xlim</pre>
  ylim <- r$ylim
  x <- as.matrix(r$x)</pre>
  y <- as.matrix(r$y)</pre>
  same.group <- r$same.group</pre>
  if (same.group) {
    if (!add)
      plot_region(xlim = xlim, ylim = ylim, plot... = plot...)
    points(x = x, y = y, cex = cex, ...)
    if (box)
      box(...)
  }
  else {
    args <- c(list(zargs = zargs, add = add), group...)</pre>
    do.call(group_2d_graphics, args)
  }
}
<bytecode: 0x7f93230634c8>
<environment: namespace:zenplots>
```

As we can see, extract\_2d() is used to extract all the plotting information used for the current 2d plot. The component same.group is then used to determine whether a normal 2d scatterplot is produced via points() or whether a group-specific 2d plot is produced (via group\_2d\_graphics()) to indicate the change of groups.

Bursting x in every 1d and 2d plot involved in a zenplot is quite time-consuming and becomes prohibitive in larger zenplots. The extractor functions extract\_1d() and extract\_2d() nicely address this problem by setting up an environment containing the burst x the first time this needs to be done (which is by the first call of extract\_1d() if first1d = TRUE and by the first call of extract\_2d() if first1d = FALSE). Subsequent calls of extract\_1d() or extract\_2d() can simply perform a look-up.

Setting up the plot region and determining the plot indices

When writing 1d or 2d plot functions, one frequently needs to set up the plot region (of the 1d or 2d plots). To this end, **zenplots** provides the function plot\_region() in case the graphical system is **graphics**; see the exported functions \*\_\*\_graphics() such as the above points\_2d\_graphics() where plot\_region() is used.

```
R> plot_region
function (xlim, ylim, plot... = NULL)
{
    if (is.null(plot...)) {
        plot(NA, xlim = xlim, ylim = ylim, type = "n", ann = FALSE,
            axes = FALSE, log = "")
    }
    else {
        fun <- function(...) plot(NA, xlim = xlim, ylim = ylim,
            ...)
        do.call(fun, plot...)
    }
}
</pre>
```

Similarly for grid, where one often needs an auxiliary function for constructing the correct viewport, zenplots provides vport(); see the exported functions \*\_\*\_grid() for how to use it.

A short but useful helper function is plot\_indices(), which allows one to determine, based on zargs, the indices of the two variables to be plotted in the current 1d or 2d plot (as mentioned before, for 1d plots, the two are the same); see ?zenplot() for how to use plot\_indices() in constructing 1d and 2d plots.

```
R> plot_indices
```

function (zargs)
zargs\$vars[zargs\$num, ]
<bytecode: 0x7f93263d4cb0>
<environment: namespace:zenplots>

#### Unfolding the cube

The major helper function which zenplot() calls is unfold(). Although unfold() is probably rather rarely used directly by a user, it provides insight into how zenplots are constructed. Suppose, for example, we have only d = 3 variates, and hence dimensions. Then each 2d plot can be thought of as defining a face of a three-dimensional cube as shown in Figure 9. To arrive at all two-dimensional plots, we imagine beginning with the bottom most face of

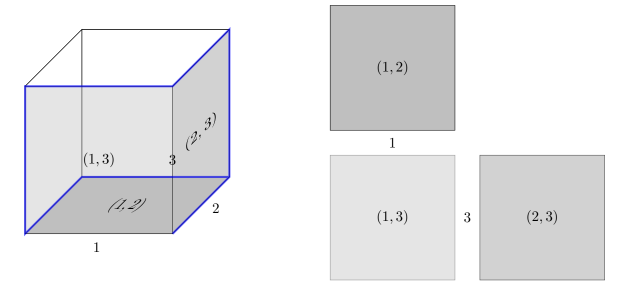


Figure 9: Three-dimensional (folded) hypercube with faces (1, 2), (1, 3) and (2, 3) (left) and unfolded faces after cutting along the blue line (right).

the cube, (1, 2)', and placing it in the display at the right of Figure 9. Now roll the cube forward about the edge marked '1', so that its (1, 3)' face is down; this face now appears in the right-hand display below (1, 2)' with the connecting edge '1' between them. With the cube face (1, 3)' now at the bottom, the cube is finally rolled left to right over the '3' edge to land on the '(2, 3)' face; the '3' edge and the '(2, 3)' face now appear in the display at right, reflecting that movement. It is in this sense that we imagine we are "unfolding the cube". In principle this could continue until all six faces appeared (though three would be redundant). Instead we have unfolded only **nfaces = 3** of the faces from the cube.

With some abuse of language, when the number of dimensions is d > 3 we still call the process unfolding (of the "cube") though really the rolling is now from one two-dimensional space to another about a shared axis (variate). It is this common axis that suggests continuing to use the "unfolding cube" metaphor. With d dimensions there are  $\binom{d}{2}$  distinct pairs of d plot variables and hence distinct two-dimensional spaces that might be visited, as ordered by the unfolding about shared axes.

The unfold() function takes, as first argument, nfaces, the number of 2d plots/spaces to be "unfolded" and produces the zenpath and zenplot layout required for the function zenplot(). Laying out these pairs with a zenplot means "unfolding" (at least a part of) the *d*-dimensional space. This is what unfold() does. It first constructs the zenpath (either based on given turns or based on the given method for constructing turns) with the (non-exported) helper function get\_path() and then, given the zenpath, determines the layout with the (non-exported) helper function get\_layout(). Both of these functions are described in more detail in the appendix.

Note that the (invisible) return value of zenplot() actually equals the return value of unfold(), which we can easily check:

```
R> n2dcols <- ncol(olive) - 1
R> stopifnot(identical(res, unfold(nfaces = n2dcols)))
```

# 6. Conclusion

We introduced the R package **zenplots** for visualization of high- (but also low-)dimensional data with the notion of zenplots and zenpaths. A zenpath provides an order of "interesting-ness" according to which the data can be sorted and then laid out and plotted with a zenplot. The package's main functions **zenpath()** and **zenplot()** achieve this task as presented in Sections 2 and 3, respectively.

Zenplots can be thought of as a variation on scatterplot matrices, where one sacrifices the (row, column) array look up and the row-wise or column-wise comparisons of a scatterplot matrix in exchange for having an arrangement of more and/or larger two-dimensional displays. With zenplots, comparisons along a common aligned scale are now restricted to a single pair of plots at a time. More importantly, however, unlike scatterplot matrices, zenplots directly address the problem of presenting possibly hundreds of plots, pre-selected for their importance or interesting structure, in a single layout. By accommodating up to a thousand or two plots in a single display, zenplots enable the analyst to consider the most interesting two-dimensional plots even from data having hundreds of dimensions (see Hofert and Oldford 2018 for an example). This means only those two-dimensional plots of greatest interest need to be considered by the analyst for further study, and these can be served up automatically.

With zenpaths, measures of interest, and groupings of variables, the R package **zenplots** adds a powerful new tool to the data analyst's arsenal when data dimensionality becomes high. As future work, we identify the need for developing and exporting more 1d and 2d plots specialized for particular analytic purposes such as dependence analysis for mixed categorical and continuous variates, or model selection and analysis in regression. More generally, an adaptation of zenplots to handle conditioning (analogous to but substantively different from coplots; Cleveland 1994) could provide new tools for multivariate data including multivariate time series.

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# A. Algorithms

In a "top-down" (top layer to bottom layer) manner, we now describe in terms of pseudoalgorithms how zenplots and zenpaths are constructed "under the hood". The presented algorithms only capture the main purpose, essential inputs (important for the algorithm overall, not necessarily required to be provided), optional inputs, the return value and the major steps; for more details (including more or more general features than captured here), consider the actual source code of **zenplots**. The reason these pieces of information are presented is to convey the intellectual challenge and contribution in constructing zenplots and zenpaths. The algorithms refer to functions which are used by **zenplots** (not all exported and thus visible to the user) but assume a more compact, mathematical notation.

The following algorithm describes the main structure of zenplot() as already discussed before. It mainly calls unfold() to determine the zenpath in the form of a list containing the vector of turns (denoted by t), a 2-column matrix of positions in the occupancy matrix (denoted by p) and the occupancy matrix itself (denoted by O). The function unfold() also determines the corresponding layout of the zenplot.

#### Algorithm A.1 (Main plot function; zenplot())

Purpose: Compute the zenplot and, invisibly, return the path and layout.

- Essential: An (n, d)-matrix or data frame (or list of such) X containing the n d-dimensional data vectors, the turns t (if not provided, they will be computed from the given number  $n_{2d} \geq 2$  of columns of 2d plots and the method ("tidy" (the default), "double zigzag", "single zigzag" or "rectangular") used for constructing the path, functions for creating the 1d and 2d plots, and an argument indicating the graphical system used for plotting.
- Optional: Logicals indicating whether the first or last 1d plot are to be plotted, the number of 2d plots along the zenpath, a vector of logicals indicating which pieces of information are passed to the 1d and 2d plot functions, how group and variable labels are determined, parameters for adjusting spaces in a zenplot and further parameters such as an ellipsis argument passed to the underlying 1d and 2d plot functions.
- Return: Nothing (R's invisible()) unless assigned to a variable in which case a list containing the path and layout is returned.

Major steps:

- (1) Call unfold() to compute the path and corresponding layout; see Algorithm A.2.
- (2) Determine the argument lists passed to the 1d and 2d plot functions, respectively.
- (3) Setting up the overall layout.
- (4) Loop over all plots and call the 1d and 2d plot functions with the respective arguments.

The following algorithm unfolds the hypercube along a (provided or determined) set of edges and returns the corresponding zenpath and a layout of 1d and 2d plots along it.

#### Algorithm A.2 (Unfolding the hypercube; unfold())

Purpose: Computes the path and corresponding layout by unfolding a cube (the sides of which are interpreted as the pairs of variables to be plotted in the 2d plots of a

zenplot).

Essential:	The number of faces of the hypercube (which equals the number $n_{2dp}$ of 2d plots
	in a zenplot, so typically the number of columns $d$ of $X$ minus one), the number
	$n_{\rm 2dc}$ of columns containing 2d plots and the method (see Algorithm A.1).

- Optional: The turns t, logicals indicating whether the first or last 1d plot are to be plotted and widths of 1d and 2d plots in the layout.
- Return: Returns the path (t, p, O) and corresponding layout (a list with orientations, plot dimensions, plot variables (2-column matrix), layout width and height and bounding boxes); see the output for Figure 1 we discussed earlier in the paper.

Major steps:

- (1) Construct the path via get\_path(); see Algorithm A.3.
- (2) Determine the layout (corresponding to the turns) via get\_layout(); see Algorithm A.6.

We now turn to determining the path. This is done via get\_path() (Algorithm A.3) which is an auxiliary function of unfold() (Algorithm A.2) and which itself has two more auxiliary functions discussed below; these are get\_zigzag\_turns() (Algorithm A.4) and next\_move\_tidy() (Algorithm A.5).

#### Algorithm A.3 (Determining the path; get\_path())

Purpose: Determine the path (turns t, positions p and occupancy matrix O).

- Essential: The turns t, the number  $n_{2dc}$  of columns containing 2d plots, the number  $n_{2dp}$  of 2d plots and the method (see Algorithm A.1).
- Optional: Logicals indicating whether the first or last 1d plot are to be plotted.

Returns: The path in the form of a list.

Major steps:

- (1) If the turns t are provided, say,  $n_{\rm t}$ -many, proceed as follows.
  - (1.1) Initialize the  $(n_t, 2)$ -matrix p of positions with zeros, set the current location  $p_{cur}$  to (0, 0) and the horizontal and vertical limits of the number of plots to (0, 0) each; the latter determine the extend of the occupancy matrix in horizontal and vertical direction.
  - (1.2) If  $n_t > 1$ , loop over  $i \in \{2, \ldots, n_t\}$ , set the current position  $p_{cur}$  to the position obtained by moving in the direction of turn i-1 (the turn into the current position) and set the *i*th row of p to  $p_{cur}$ . Then check whether the horizontal or vertical limits have to be extended and extend them if necessary.
  - (1.3) To obtain the final matrix of positions, "shift" all coordinates appropriately (by subtracting the minimum of each column of p from the respective column and adding 1 to each value).
  - (1.4) Initialize the occupancy matrix O with zeros (no position occupied); note that the maximum of the first and second column of p determine the dimensions of O. Then loop over all rows of p and set the respective entry in O to 1, 2, 3 or 4, depending on whether the turn out of the current position  $p_{cur}$  is "l", "r", "d" or "u", respectively.

- (2) If the turns t are not provided, proceed as follows. Let  $n_{\text{plots}}$  denote the number of all plots (1d or 2d).
  - (2.1) For method "rectangular", proceed as follows.
    - Set  $n_{\text{plots}} = 2n_{2\text{dp}} + 1 (1 \mathbb{1}_{\text{first1d}}) (1 \mathbb{1}_{\text{last1d}})$ , where  $\mathbb{1}_{\text{first1d}}$  ( $\mathbb{1}_{\text{last1d}}$ ) is the indicator of the event that the first (last) 1d plot is to be displayed.
    - Set the number  $n_{2dr}$  of rows with 2d plots to  $n_{2dr} = \lceil n_{2dp}/n_{2dc} \rceil$ .
    - Determine the turns to fill the rectangular layout (determined by  $n_{2dr}$  and  $n_{2dc}$ ), remove the first turn or append the last (depending on whether the first and last 1d are to be displayed) and extract the  $n_{plots}$ -many turns required for the zenplot.
    - Loop over the turns to determine the positions *p* and from there the occupancy matrix *O*.
    - Return the path (list of turns t, positions p and the occupancy matrix O).
  - (2.2) For all other methods, set  $n_{\text{plots}} = 2n_{2\text{dp}} + 1$  and proceed as follows.
    - If  $n_{\text{plots}} = 1$ , set turns t = ("d"), positions p = (1, 1), occupancy matrix O = (3).
    - If  $n_{\text{plots}} = 2$ , set turns t = (``d'', ``r''), positions  $p = \begin{pmatrix} 1 & 1 \\ 2 & 1 \end{pmatrix}$ , occupancy matrix

$$O = \binom{3}{2}.$$

• If  $n_{\text{plots}} = 3$ , set turns t = (``d'', ``r'', ``r''),

positions 
$$p = \begin{pmatrix} 1 & 1 \\ 2 & 1 \\ 2 & 2 \end{pmatrix}$$
, occupancy matrix  $O = \begin{pmatrix} 3 & 0 \\ 2 & 2 \end{pmatrix}$ .

- If  $n_{\text{plots}} \ge 4$ , proceed as follows.
  - (2.2.1) For method "double zigzag" or "single zigzag", the idea is to build all turns right away (the positions and the occupancy matrix can then be determined from the turns):
    - Compute the turns via get\_zigzag\_turns(); see Algorithm A.4.
    - Initialize the occupancy matrix O as an  $(1, 2n_{2d}-1)$ -matrix containing zeros and the positions p as an  $(n_{\text{plots}}, 2)$ -matrix containing zeros.
    - ► Encode the turns "l", "r", "d", "u" as the numeric values 1-4, respectively.
    - Set  $p_{cur} = (1, 1)$  and the first row of p to  $p_{cur}$ . Furthermore, set O at  $p_{cur}$  to the numeric value of the first turn.
    - ▶ Loop over  $i \in \{2, ..., n_{\text{plots}}\}$ , set the next position  $p_{\text{next}}$  to the value obtained by moving from  $p_{\text{cur}}$  in the direction of turn i 1 and set the *i*th row of *p* to  $p_{\text{cur}}$ . Then update  $p_{\text{cur}}$  to  $p_{\text{next}}$ . If  $p_{\text{cur}}$ 's first entry exceeds the maximal row number of *O* append another row of zeros to *O*. Then set *O* at  $p_{\text{next}}$  to the numeric value of the *i*th turn.
    - ▶ Trim off the last columns of zeros (if any) from O.
  - (2.2.2) For method "tidy":
    - Initialize the  $n_{\text{plots}}$ -vector turns t, and the  $(1, 2n_{2d} + 1)$  occupancy matrix O with zeros and the positions p as an  $(n_{\text{plots}}, 2)$ -matrix containing zeros. Furthermore, set the first turn to d, the first row of

p to (1,2) and the occupancy matrix O at (1,2) to 3 (numeric value corresponding to a turn "d").

- Loop over  $i \in \{2, \ldots, n_{\text{plots}}\}$ , determine the next position  $p_{\text{next}}$  and the turn  $t_{\text{next}}$  out of  $p_{\text{next}}$  via next\_move\_tidy(), see Algorithm A.5, set the *i*th turn to  $t_{\text{next}}$  and the *i*th row of *p* to  $p_{\text{next}}$ . If the first entry of  $p_{\text{next}}$  exceeds the maximal row number of *O* append another row to *O*. Then set *O* at  $p_{\text{next}}$  to the numeric value of the *i*th turn.
- If it exists, trim off the last column of zeros from O.
- ▶ If it exists, trim off the first column of zeros from *O* and subtract 1 from the second column of *p* (as all positions relative to the occupancy matrix changed).
- Depending on whether the first and last 1d plot are to be displayed, trim the path accordingly.
- Return the path (list of turns t, positions p and the occupancy matrix O).

# Algorithm A.4 (Compute the turns for zenplots with method "double zigzag" or "single zigzag"; get\_zigzag\_turns())

- Purpose: Compute the vector of turns for zenplots with method "double zigzag" or "single zigzag".
- Essential: The number of plots  $n_{\text{plots}}$ , the number  $n_{2d} \geq 2$  of columns of 2d plots, the method used for constructing the path ("double zigzag" or "single zigzag").

Returns: The turns t.

Major steps:

- Define the horizontal 2d pattern ("r",..., "r", "l"..., "l") (each of "r", "l" appears 2(n<sub>2d</sub>-1) times). So far this only contains horizontal movements, vertical movements will be added in Step (4) below.
- (2) Define the vertical 2d subpattern corresponding to the first part ("r",...,"r") of the horizontal 2d pattern in the following way:
  - For method "single zigzag", the vertical 2d subpattern is ("d",..., "d") (of length  $n_{2d} 1$ ).
  - For method "double zigzag", the vertical 2d subpattern can be constructed as follows: If  $n_{2d} = 2$ , it is ("d"). If  $n_{2d} > 2$ , take  $n_{2d} - 3$  elements from the alternating sequence "d", "u", "d", "u", … and append two "d"'s (so, ("d", "d") for  $n_{2d} = 3$ , ("d", "d", "d") for  $n_{2d} = 4$ , ("d", "u", "d", "d") for  $n_{2d} = 5$ , etc.).
- (3) Define the vertical 2d pattern corresponding to the horizontal 2d pattern from Step (1) by repeating the vertical 2d subpattern from Step (2) twice (i.e., vertical 2d subpattern, vertical 2d subpattern).
- (4) Set components of the horizontal 2d pattern from Step (1) with even indices to the values of the vertical 2d pattern from Step (3). With this merge, we obtain the 2d pattern which repeats itself until we run out of plots.
- (5) Repeat each element of this 2d pattern twice to obtain the overall pattern; this is to account for 1d plots, so the pattern now contains turns for both 1d plots and 2d plots.

(6) The vector of turns can now be obtained by taking "d" as first component (since the first turn is always "d") and then taking the first  $n_{\text{plots}}$  elements of an imaginary sequence consisting of the overall pattern repeated after each other; recall that  $n_{\text{plots}}$  denotes the total number of 1d and 2d plots.

Algorithm A.5 (Determine the next position and turn out of there; next\_move\_tidy())

- Purpose: Determine, for the method "tidy", the next position  $p_{\text{next}}$  to move to and the turn  $t_{\text{next}}$  out of this next position.
- Essential: The current plot number  $n_p$ , the total number of plots  $n_{plots}$  and the current path that has been determined up to and including  $n_p$ . Note that this function assumes that the zenplot starts with a 1d plot; get\_path() trims off the first 1d plot if necessary.
- Returns: A list containing the next position  $p_{\text{next}}$  (i.e., the position of plot numbered  $n_{\text{p}}+1$ ) and the turn  $t_{\text{next}}$  out of this next position.

Major steps:

- (1) Let  $n_{\text{left}} = n_{\text{plots}} n_{\text{p}}$  denote the number of remaining plots.
  - If  $n_p = 1$ , return  $p_{next} = (2, 2)$  and  $t_{next} = "r"$ .
  - If  $n_p = 2$ , return  $p_{next} = (2, 3)$  and  $t_{next} = "r"$ .
  - If  $n_p = 3$ , set  $t_{next} = u$  if  $n_{left} \le 2$  and set  $t_{next} =$  "d" otherwise. Return  $p_{next} = (2, 4)$  and  $t_{next}$ .
- (2) Determine the current position  $p_{cur}$  (corresponding to row  $n_p$  in the positions matrix p), the turn  $t_{in}$  into  $p_{cur}$  (element  $(n_p - 1)$  in the vector of turns t), the turn  $t_{out}$  out of  $p_{cur}$ (element  $n_p$  of t) and the next position  $p_{next}$  when moving from  $p_{cur}$  in the direction given by  $t_{out}$ .
- (3) If  $n_{\rm p}$  is even (i.e., the current plot is a 2d plot according to the above assumption), return  $p_{\rm next}$  and  $t_{\rm out}$ .
- (4) Now the current plot numbered  $n_p$  is a 1d plot. We proceed by determining the horizontal moving direction out of this plot. To this end, consider the turns at  $n_p 2$  and  $n_p 1$ . If "r" is among any of these two turns, the horizontal moving direction is "r", otherwise it is "l". Furthermore, determine the distance to the margin of the occupancy matrix O in the horizontal moving direction; if the latter is "r", the distance is the number of columns in O minus the second component of  $p_{cur}$  (distance to the right end of O); otherwise it is the second component of  $p_{cur}$  minus 1 (distance to the left end of O).
- (5) We now determine the turn  $t_{next}$  out of  $p_{next}$ . To this end, we distinguish two cases:
  - (5.1) If  $t_{out}$  is "d" or "u" (i.e., the current 1d plot is horizontal), proceed as follows.
    - (5.1.1) If  $n_{\text{left}} \leq 2$ , check the location of the 2d plot which comes after the next 2d plot in opposite horizontal moving direction.
      - If this location does not exist in O (which can only happen if  $t_{out} = "d"$  in which case O is missing a new row), proceed as follows. Put the last 1d plot in opposite horizontal moving direction if we are near the margin (i.e., set  $t_{next} = "l"$  if the horizontal moving direction is "r" and  $t_{next} = "r"$  otherwise; we follow this strategy here since we occupy an

additional column otherwise). Otherwise, put it in horizontal moving direction (i.e., set  $t_{\text{next}}$  to the horizontal moving direction; this can be done since we are away from the margin, i.e., "inside" O).

- If this location exists, check the occupancy matrix at this location to see whether it is occupied. If it is not occupied, set  $t_{\text{next}}$  to the opposite horizontal moving direction ("l" if the horizontal moving direction is "r" and vice versa). Otherwise, set  $t_{\text{next}}$  to the current horizontal moving direction.
- (5.1.2) If  $n_{\text{left}} \geq 3$  (so there are at least two more 2d plots left), change the horizontal moving direction if and only if the distance to the margin is less than or equal to two.
- (5.2) If  $t_{out}$  is "l" or "r" (i.e., the current 1d plot is vertical and the distance to the margin is at least two in this case), proceed as follows. Ideally, we would like to go up (in order to save space in the plotting region), but if there are too many plots left, we can only go up if we have the space to go down again. To explain this part of the algorithm, we utilize the concept of a U-turn (going up and back down again).
  - (5.2.1) Given  $p_{\text{cur}}$ , the horizontal moving direction and the occupancy matrix O so far, determine the number of plots which fit in the U-turn starting from  $p_{\text{cur}}$ , i.e., the length  $l_{\text{U-turn}}$  of the U-turn. This can be done as follows.
    - Let  $p_{\text{check}}$  denote the position obtained from  $p_{\text{cur}}$  by going up two rows and then one column in the horizontal moving direction. If  $p_{\text{check}}$  exists in the occupancy matrix O and is occupied, return  $l_{\text{U-turn}} = 1$  (as only one additional plot fits in along this U-turn). If it does not exist in O, return  $l_{\text{U-turn}} = 2$ .
    - Update  $p_{\text{check}}$  by going two columns further in the horizontal moving direction. If  $p_{\text{check}}$  does not exist in O or if it exists but is occupied, return  $l_{\text{U-turn}} = 4$ .
    - Update  $p_{\text{check}}$  by going two rows down. If  $p_{\text{check}}$  does not exist in O or if it exists but is occupied, return  $l_{\text{U-turn}} = 6$ .
    - Update  $p_{\text{check}}$  by going two columns further in the horizontal moving direction. If  $p_{\text{check}}$  does not exist in O or if it exists but is occupied, return  $l_{\text{U-turn}} = 8$ . Otherwise return  $l_{\text{U-turn}} = 10$  (which means that at least 10 plots fit in the U-turn and we can do a complete U-turn by going up and back down again).
  - (5.2.2) If the first component of the current position  $p_{cur}$  is less than or equal to two (i.e., we are in the second row of O and thus cannot go up) or if  $l_{\text{U-turn}} \leq 9$ (i.e., at most 9 plots fit in the U-turn starting from  $p_{cur}$ ) but  $n_{\text{left}} > l_{\text{U-turn}}$ (i.e., there are more plots left than fit in the U-turn), set  $t_{\text{next}} = \text{``d''}$ . Otherwise, set  $t_{\text{next}} = \text{``u''}$ .
- (6) Return  $p_{\text{next}}$  and  $t_{\text{next}}$ .

Finally, we describe how the layout can be constructed.

#### Algorithm A.6 (Determining the layout; get\_layout())

Purpose: Determine the layout (1d and 2d plot orientations, plot dimensions, plot variables, total width and height of the layout and bounding boxes).

- Essential: The turns t and number  $n_{2dp}$  of 2d plots.
- Optional: Logicals indicating whether the first or last 1d plot are to be plotted and widths of 1d and 2d plots in the layout.
- Returns: The layout (a list with orientations of 1d and 2d plots, their dimensions (either 1d or 2d), which variables are plotted in each 1d and 2d plot, layout widths and heights, and bounding boxes; see Section 5.1).

Major steps:

- (1) Set up an empty 4-column matrix containing the left, right, bottom and top coordinates (in units of the widths of 1d and 2d plots) of the bounding box of each plot. Furthermore, set up a 2-column matrix containing the variables of each 1d or 2d plot.
- (2) Loop over all 1d and 2d plots and determine, for each, its orientation, plot variables involved and coordinates of the bounding box. In particular, this loop requires as input the number of plots, the current and the last plot variable. Some more details about this loop:
  - Determining the current plot variables: For 1d plots, this is the current plot variable repeated twice. For 2d plots which turn left or right out of the current position in O, this is the last and current plot variable; for 2d plots turning down or up out of the current position in O, this is the current and the last plot variable (notice the difference in order).
  - Determining the current bounding box: This is a function of the last turn (i.e., the turn into the current plot position), the last bounding box and the width and height of the bounding box. The latter two are easy to determine based on the provided widths of the 1d and 2d plots. And the location of the bounding box for the current plot equals the bounding box of the previous plot shifted in the direction of the last turn (so the turn into the current plot position).

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