

A Tour of Trellis Graphics

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1. INTRODUCTION

Trellis displays are plots which contain one or more panels, arranged in a regular grid-like structure (a trellis). Each panel graphs a subset of the data. All panels in a Trellis display contain the same type of graph but these graphs are general enough to encompass a wide variety of 2-D and 3-D displays: histogram, scatter plot, dot plot, contour plot, wireframe, 3-D point cloud and more. The data subsets are chosen in a regular manner, conditioning on continuous or discrete variables in the data, thus providing a coordinated series of views of high-dimensional data.

This document leads you through Trellis graphics: it shows the functions in the Trellis library, it describes the common arguments that the functions share, and shows how Trellis displays are customized for various graphical devices. Other information is available about Trellis, including a user's manual and a journal article with data analysis examples. To find these and more, refer to the Trellis web page:

<http://netlib.att.com/projects/trellis/>

1.1 New Capabilities for S and S-PLUS Graphics

Graphics have always been a strong feature of S and S-PLUS (the commercial version of S, distributed by MathSoft). Its graphics provide device independence, high-level plotting functions that produce an entire display, low-level functions to augment existing displays or build new ones, and a collection of graphical parameters that provide a wide range of control over the details of plotting.

Graphical parameters in S provide the ability to produce several plots on a single page. However, producing a coordinated set of plots on a page, with control over aspect ratios and axes, has always taken more knowledge of the graphical functions than even a proficient user is likely to possess. In addition, graphics devices may vary in their capabilities, thus requiring adaptations in order to produce the best plot on each device.

The Trellis library is designed to remedy this situation. Besides providing a straightforward way to produce multiple panels on a single page, it also sets up a unifying framework for doing this. Trellis displays extend S graphics to handle multivariate data situations by using a powerful and general technique, *conditioning*. In addition, the Trellis software does an excellent job with single panel displays, making it a suitable vehicle for doing most high-level graphics in S.

While improving user control of graphics, the Trellis software also makes graphics functions behave just like any other S functions. The result of executing a Trellis expression is a Trellis object. Unless it is assigned a name or used in a further computation, the Trellis object is displayed.

The Trellis library is now distributed as a standard part of S-PLUS. (Hold onto your hats, jargon to follow!) S-PLUS prior to version 3.3 does not come with the Trellis library. In the PC environment, S-PLUS for Windows, Version 3.3 (and presumably anything later) comes with Trellis Version 2.0, as described in this document. Under the Unix operating system, S-PLUS Version 3.3 contains a slightly older version of Trellis. The next release, due in 1996, is scheduled to contain Trellis Version 2.0.

1.2 A Simple Scatterplot: Ethanol Data

Perhaps the easiest way to introduce Trellis displays is by examples. They illustrate the variety of Trellis displays that can be produced and also introduce the way that Trellis displays are controlled. This document gives an entry point to use of the software and explains common features. It does not have the space to explain, except in the most cursory way, the meaning or use of the graphical techniques. To find out much more about how to use Trellis displays to

understand data, read *Visualizing Data* by William S. Cleveland (1993).

Although Trellis graphics functions are capable of producing multiple panel displays, they are also excellent at doing basic single-panel graphs. For this example, we will use data from an experiment involving 88 trials of an engine running an ethanol mixture, contained in the data frame `ethanol`. There are three variables: emissions of oxides of nitrogen, `NOx`, equivalence ratio (a measure of the richness of the fuel/air mixture), `E`, and five values of compression ratio, `C`.

We can use the scatterplot function `xypLOT` on the `ethanol` data to produce a simple scatterplot:

```
xypLOT(NOx ~ E, data = ethanol) # Figure 1
```

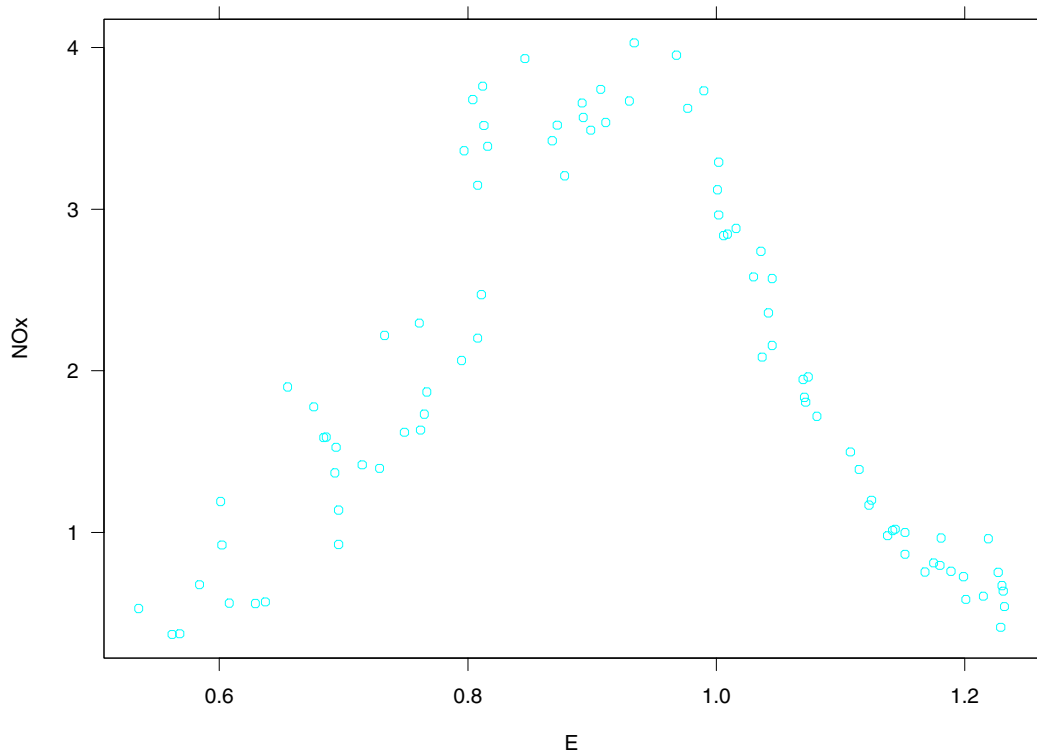


Figure 1. A simple scatterplot of the engine data, showing `NOx` emissions as a function of equivalence ratio.

The first argument to `xypLOT` and to most Trellis functions is a *formula* and the second tells where the data in the formula can be found. Both of these kinds of arguments were introduced in the book *Statistical Models in S* by Chambers & Hastie (1992). We have used this same paradigm for Trellis graphics.

1.3 Conditioning on Numeric Variables: Ethanol Data

A simple modification of the previous call to `xypLOT` produces a multi-panel display:

```
xypLOT(NOx ~ E | C, data = ethanol) # Figure 2
```

This produces Figure 2, which shows `NOx` emissions plotted against equivalence ratio, each panel showing data for one of the five values of compression ratio.

The Trellis display consists of 5 panels, each showing `NOx` on the vertical axis and `E` on the

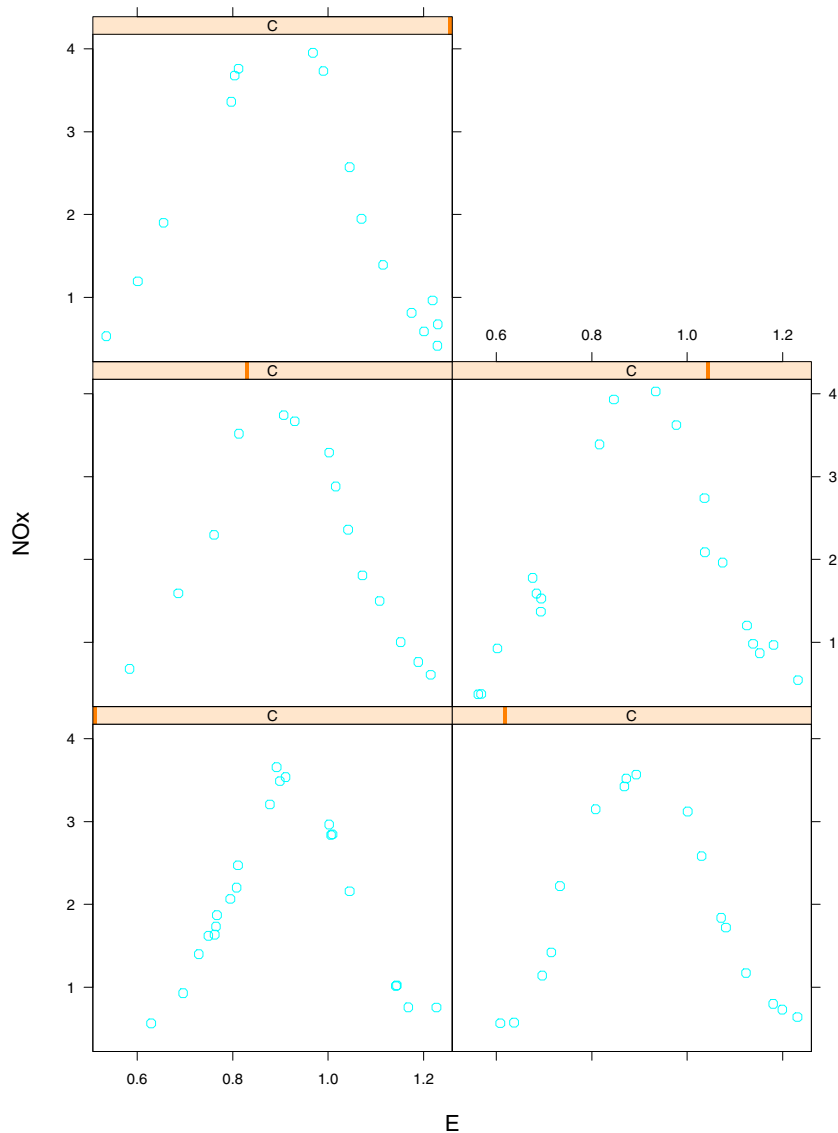


Figure 2. A Trellis display of engine emissions data.

horizontal axis. The value of C is shown by the *strip label* at the top of each panel; in this case, C takes on 5 discrete values as shown by the darkly-shaded region of the strip label atop each panel.

The formula,

$$\text{NOx} \sim \text{E} \mid \text{C}$$

is read aloud as “NO_x is plotted against E given C”. Note that the variable that goes on the vertical axis is mentioned first in the formula — conventionally the dependent variable is plotted on the vertical axis; the variable for the horizontal axis is given after the “~” operator, and *given* or *conditioning* variables are mentioned last.

The data = ethanol tells xypplot to look first in the data frame ethanol for the objects NO_x, E, and C in the formula. A data frame contains a set of related vectors and can be operated upon as if it were a matrix; however, data frames can hold data of various types, including

character and numeric vectors, factors, and shingles. Data mentioned in a formula can come from anywhere on the `S` search list. However, data frames are often a convenient way to keep related vectors together.

Suppose that we would like to control the layout of the five panels in Figure 2. We can do that with the `layout=` argument:

```
xyplot(NOx ~ E | C, data = ethanol, layout = c(3,2,1)) # Figure 3
```

This produces a layout with 3 columns and 2 rows on 1 page. Notice that `layout` specifies columns and then rows, unlike matrix (row, column) notation. We do this because we are dealing with graphs, and the convention with graphs is to have an origin in the lower left corner. We start there and proceed left to right, bottom to top, page to page. The number of panels to be produced in a Trellis display is determined by the number of levels in the given variables. However, if the layout argument allows less room than required for all of the panels, only the panels that will fit are produced. For example, using `layout=c(2,2,1)` with this example would produce just one page with 4 panels.

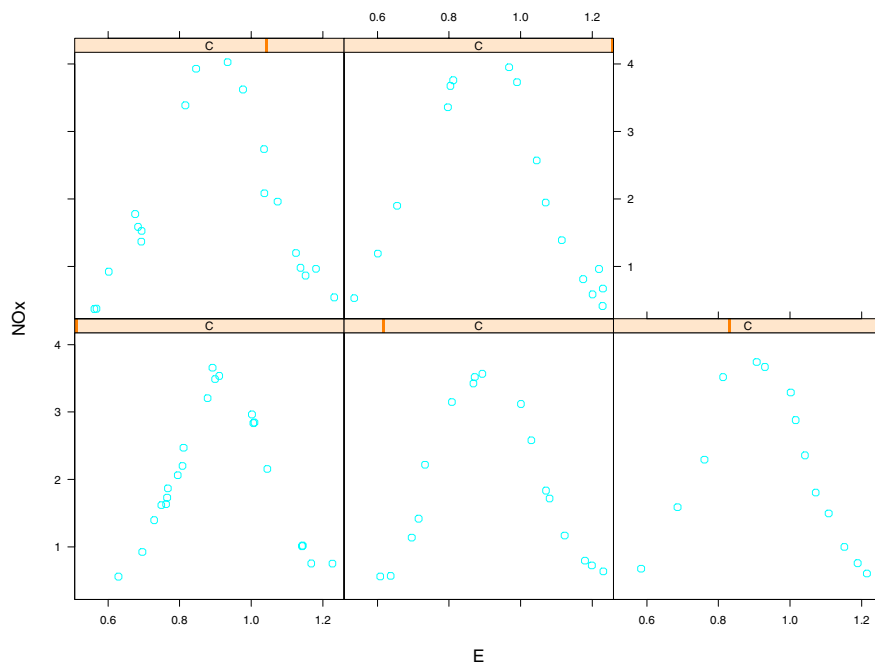


Figure 3. The engine data with `layout=c(3,2,1)`.

We have conditioned on compression ratio, a variable that has only 5 levels.

```
> sort(unique(ethanol$C))  
[1] 7.5 9.0 12.0 15.0 18.0
```

Suppose we want to see how `NOx` concentrations depend on `C` for various values of `E`. It won't work to condition on `E`, since there are 83 unique values of `E` for the 88 observations. However, we can do something similar by conditioning on *intervals* of `E`. The function `equal.count` constructs a data structure called a *shingle* from our data, with a specified number of intervals and a specified amount of overlap from one interval to another. For example,

```
EE <- equal.count(ethanol$E, number = 9, overlap = 1/4)
```

constructs a shingle with 9 intervals spanning the range of `E`, each containing approximately the

same number of observations, and each having about 25% of its points in common with the each of its two adjacent intervals.

```
> EE

Data:
 [1] 0.907 0.761 1.108 1.016 1.189 1.001 1.231 1.123 1.042 1.215 0.930 1.152
[13] 1.138 0.601 0.696 0.686 1.072 1.074 0.934 0.808 1.071 1.009 1.142 1.229
[25] 1.175 0.568 0.977 0.767 1.006 0.893 1.152 0.693 1.232 1.036 1.125 1.081
[37] 0.868 0.762 1.144 1.045 0.797 1.115 1.070 1.219 0.637 0.733 0.715 0.872
[49] 0.765 0.878 0.811 0.676 1.045 0.968 0.846 0.684 0.729 0.911 0.808 1.168
[61] 0.749 0.892 1.002 0.812 1.230 0.804 0.813 1.002 0.696 1.199 1.030 0.602
[73] 0.694 0.816 1.037 1.181 0.899 1.227 1.180 0.795 0.990 1.201 0.629 0.608
[85] 0.584 0.562 0.535 0.655

Intervals:
  min  max count
0.535 0.686   13
0.655 0.761   13
0.733 0.811   12
0.808 0.899   13
0.892 1.002   13
0.990 1.045   13
1.042 1.125   12
1.115 1.189   13
1.175 1.232   13

Overlap between adjacent intervals:
 [1] 4 3 3 3 4 3 3 4
```

We can see the intervals graphically by using the plot function on a shingle:

```
plot(EE, xlab="Range of E", ylab="Interval") # Figure 4
```

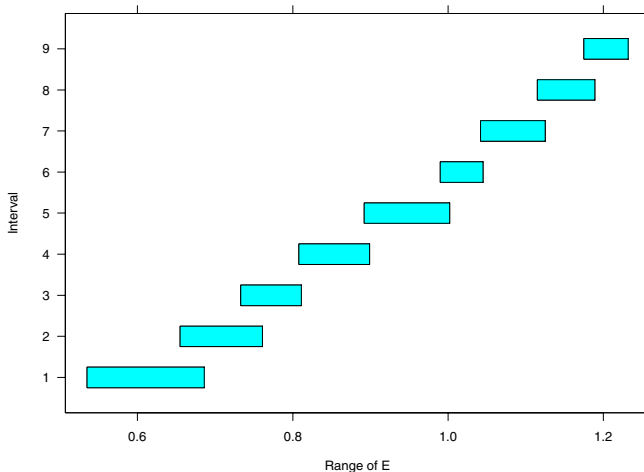


Figure 4. A plot of the shingle with 9 equal-count intervals constructed from the engine data.

Now that we have a set of intervals, we can use them to produce conditional scatterplots of NO_x vs. C for the various ranges of E:

```
xyplot(NOx ~ C | EE, data = ethanol, aspect = 2) # Figure 5
```

We made the panels tall by specifying aspect=2 in the call. (The reason for this will be made clear later.)

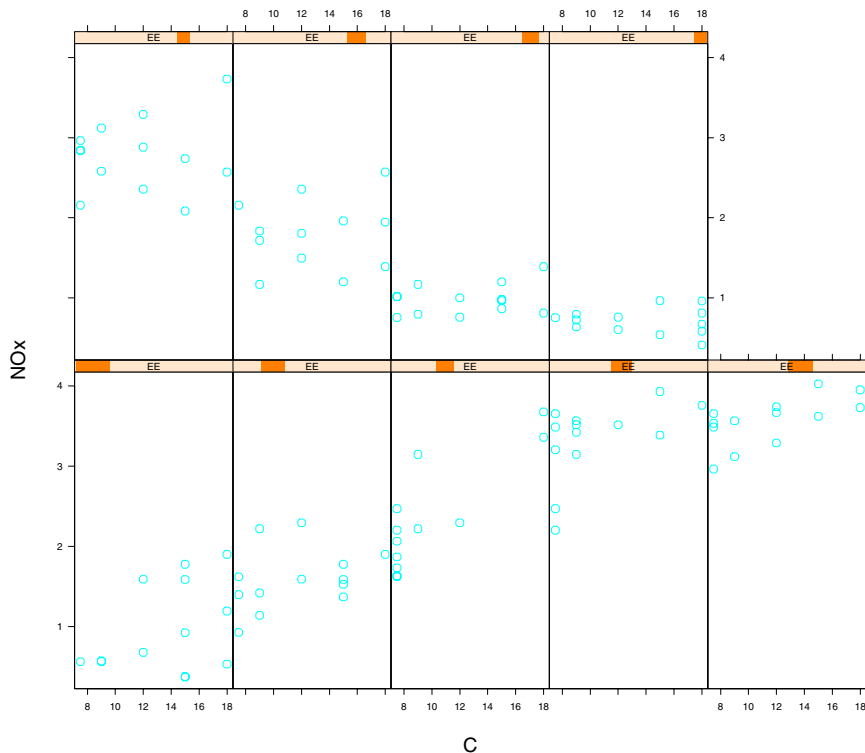


Figure 5. Scatterplots show NO_x emissions as a function of compression ratio for various values of equivalence ratio.

Suppose that for some reason we wanted to recreate Figure 2, but to show it only for compression ratios greater than 8. One way to do this would be

```
ii <- ethanol$C > 8
xyplot(NOx[ii] ~ E[ii] | C[ii], data = ethanol)
```

another possibility would be:

```
xyplot(NOx ~ E | C, data = ethanol[ethanol$C>8, ])
```

but an easier version can be accomplished using the `subset=` argument:

```
xyplot(NOx ~ E | C, data = ethanol, subset = C > 8) # Figure 6
```

In complicated expressions, `subset=` can save you from lots of subscripting and also allows you to refer to components of a data frame by name. In addition, using `subset=` causes an automatic operation on all factors used in the formula, to drop any levels that have no data associated with them. The next section describes how factors are handled in Trellis displays.

1.4 Conditioning on Factors: Barley Data

Let's leave the engine example and go on to something else, this time involving more than one conditioning variable and a dotplot rather than a scatterplot. The data frame `barley` describes the yield in bushels per acre of 10 varieties of barley, harvested at 6 sites in 2 different years. The expression

```
dotplot(variety ~ yield | year * site, # Figure 7
        data = barley,
        xlab = "Barley Yield (bushels/acre)")
```

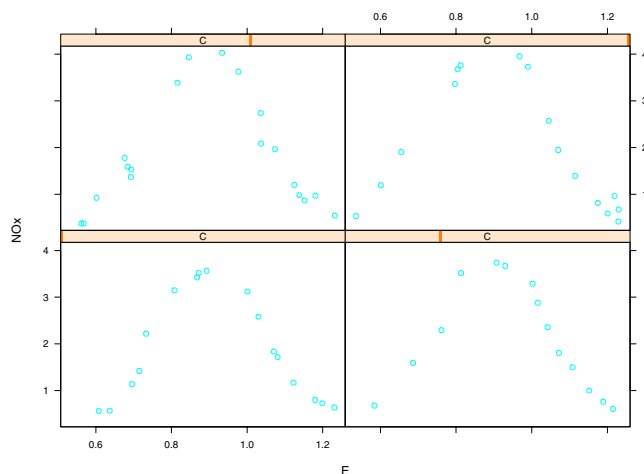



Figure 6. Engine emissions data for compression ratios larger than 8.

produces Figure 7. See Cleveland (1993) for a thorough analysis of the barley data.

1.5 Other Examples

We have reached the end of this introductory section and you should now have some familiarity with Trellis displays and how they are produced. However, there is no way in a document as short as this that we can do more than hint at the variety of displays you can produce with Trellis expressions. One way for you to get more experience is to explore the functions whose names start with `example` in the Trellis library. Execute

```
help(example.bwplot)
```

to get a full list. For example, you can execute

```
example.bwplot()
```

to produce a Trellis box-and-whisker display (boxplot) and execute

```
example.bwplot
```

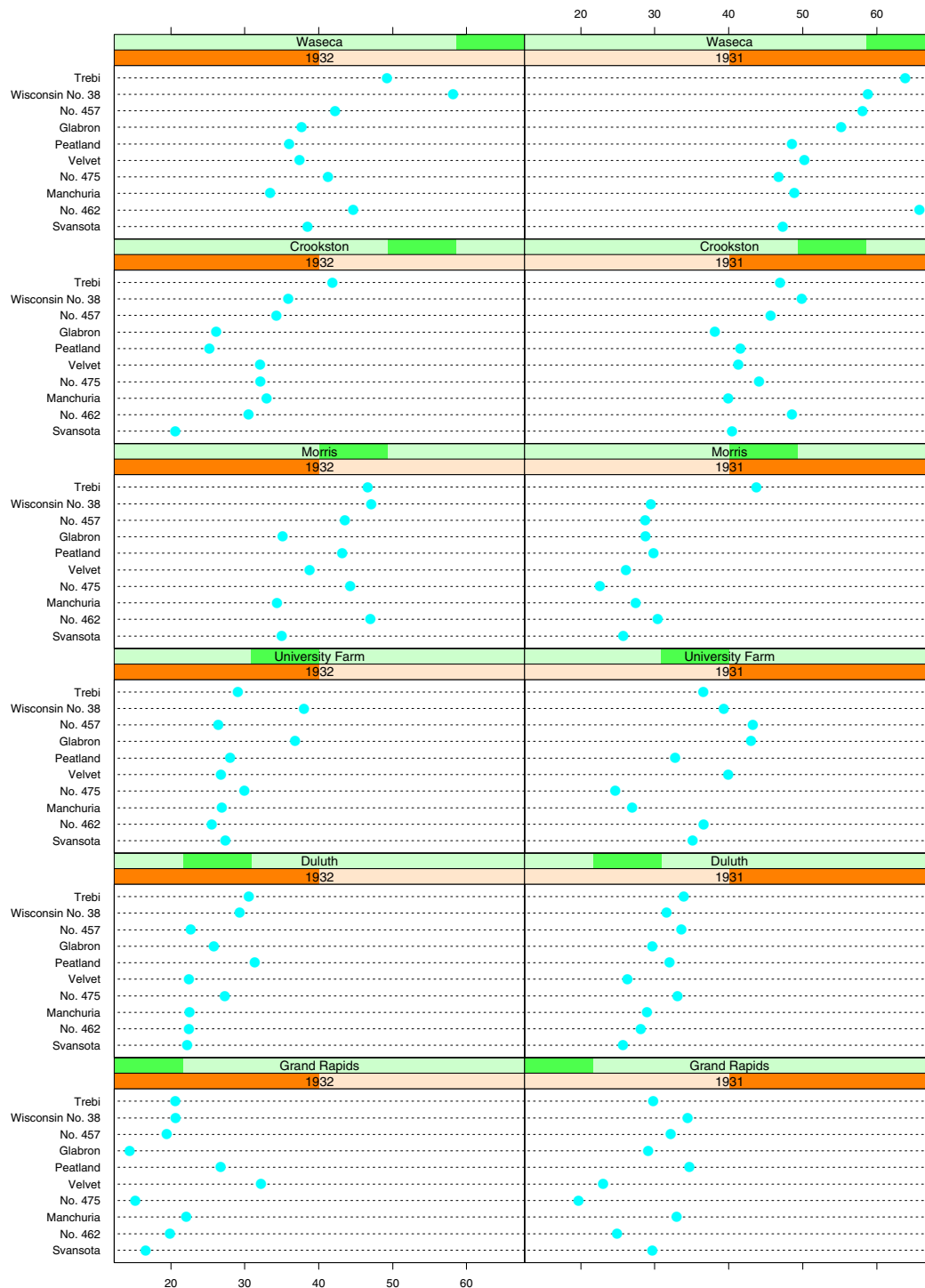
to print the function so you can understand how the display is created.

2. HOW TO USE TRELIS SOFTWARE

The previous section showed a few examples of Trellis displays and how they are specified and controlled in S. The purpose of this section is to give a broader look at the Trellis graphics software.

In general, Trellis displays consist of one or more panels, arranged in a regular grid-like structure of columns, rows, and pages. Simple displays are usually easy to create; multi-panel displays take little more effort. A wide range of graphs can be drawn inside each panel, although all panels in a particular Trellis display must be alike. Each panel displays a subset of the data, determined by the values of the *given* variables.

The Trellis software is structured so that there is one piece of software, the Trellis *print method*, that takes Trellis objects and produces all types of displays. It advances from panel to panel, sets up axes, computes appropriate aspect ratios, and generates overall labels. The print method calls a *panel function* once per panel to draw the graph inside the panel. This decoupling of the overall setup and panel drawing provides much of the power to the Trellis software. We



Barley Yield (bushels/acre)

Figure 7. A dotplot shows how the yield of 10 varieties of barley varied over 6 sites and 2 years. Notice that the 1931 yields were generally higher than 1932 except at the Morris site (a likely data transcription error).

supply panel functions for doing many kinds of Trellis displays, but you always have the option to write a custom panel function to draw just the right thing in each panel.

2.1 Display Functions

The Trellis graphics software comes with many *display functions* that produce various types of complete graphs. One way of classifying the display functions is according to the dimensionality of the incoming data. Here's a list:

Data Type	Function	Type of Plot
Univariate	barchart	bar plots
	bwplot	box and whisker plots
	densityplot	kernel density plot
	dotplot	dot plots
	histogram	histograms
	piechart	pie charts (yuck!)
	qqmath	quantile plots against mathematical distribution
	stripplot	1-dimensional scatter plot
Bivariate	qq	qq-plot for comparing two distributions
	timeplot	time series plot
	xyplot	scatter plot
Trivariate	contourplot	contour plot
	levelplot	level plots
Hypervariate	splom	scatterplot matrix
	parallel	parallel coordinate plots
3-D Displays	wireframe	function as a wire frame
	cloud	3-dimensional point cloud

2.2 Customization for Devices

Trellis displays are adapted to different graphical devices by the `trellis.device` function. When you are using Trellis graphics on your personal computer or workstation, you normally use `trellis.device` implicitly: when the first Trellis object is printed, `trellis.device` is called automatically to produce a window in which to display the graph. If you are running Windows, the `win.graph` function is invoked, and if you are on a workstation running the Unix operating system, the `motif` device function is executed.

However, there may be times when you will want to run `trellis.device` explicitly. For example, executing

```
trellis.device(postscript, file = "output.ps", color = TRUE)
```

will direct any further Trellis graphics to the `postscript` device function, leaving the file `output.ps` filled with PostScript® language commands. This file can then be sent to a printer or included in documents. The `color=TRUE` argument says that your output should be done in color. For a black and white printer, use `color=FALSE`.

Another reason why you may want to execute `trellis.device` explicitly is that you can use it in S-Plus to display multiple windows on one screen or you can set up an on-screen device in addition to a hardcopy device.

Not only does `trellis.device` set up the device, but it also initializes a list that controls graph characteristics such as plotting characters, line types, and colors. Gray scales and different

line styles and symbol types are used in the black and white version to distinguish areas, lines, and points. If color is available, the Trellis color scheme is used. It is designed to provide good discrimination amongst a fairly large group of colors. This is necessary because Trellis displays are often concerned with multivariate situations. The colors are also chosen to be relatively robust across various graphics devices, screens and hardcopy—they are simple mixtures of the cyan, magenta, and yellow printing primaries. The background and basic annotations are done monochromatically to contrast with the color assigned to data-based graphics. Black text is used because it is available on all devices and stands out well on white, gray, and color.

For hardcopy devices (e.g. those that support PostScript®), the background color will be the white of the paper; on a computer screen, the background color is light gray because white is too harsh and tends to hide light colors. The colors used are those that empirically offer good discrimination and are not difficult for the device. In particular, text is drawn in black and the colors cyan, magenta, green, orange, blue, yellow, and red are employed for various groups. The function `show.settings` gives a graphical display of the Trellis color (and other) parameter settings. See section 3.3 for an example plot and more details about device settings.

2.3 Panel Functions

Panel functions lie at the heart of Trellis displays. When any Trellis display function is executed, common code parses the formula, gets the data, and returns a Trellis object. If the object is “printed”, the `print` method, `print.trellis`, sets up the coordinate system, constructs appropriate axes, and then produces the panels, one by one. When the panel function is called, it is provided with two arguments, `x` and `y`, giving the horizontal and vertical coordinates of what should be plotted. The data passed to the panel function consists of subsets of the independent and dependent variables, the subsets determined by appropriate values of the given variables.

There is a default panel function for each of the high-level Trellis functions; the name is constructed by gluing `"panel."` onto the name of the Trellis function. Thus `panel.xyplot` corresponds to the `xyplot` function. Many of these functions are simple, basically a call to a plotting routine like `points` or `lines` with a few graphical parameters specified to mesh with the Trellis graphics customization rules (see section 3.3, “Device Settings”).

When a panel function is called, the Trellis code has already set up a coordinate system (based upon data values, and arguments `xlim=`, `ylim=`, and `scales=`). It has also extracted, from the entire collection of data, the data to be displayed upon the panel, the extraction based on the levels of the given variables. The panel function is passed arguments `x` and `y`, the coordinates for the points on the current panel. (If no points belong on the panel, the panel function is not called.)

The panel function is passed in as an argument to a high-level Trellis function, perhaps as an object (`panel.xyplot`), as a character string (`"panel.xyplot"`), or constructed on the fly in the argument list (`function(x,y) { ... }`).

Any arguments given to the top-level Trellis function but not recognized by it are passed, unchanged, down to the panel function. This can be a useful way of communicating extra information to the panel function, and is often used in conjunction with simple graphical parameters, like `col=` or `cex=`.

Because Trellis displays may take up many pages, the paradigm of producing a plot and then adding to it, will not work. (This was often the way things were done prior to Trellis graphics). However, the panel function can do something similar. This is relatively easy because panel functions have a nice synergy with the ability to define functions. For example, suppose you want panels with points and a smooth line. You can do this easily by combining two existing panel functions into one:

```
xyplot(NOx ~ E | C, data = ethanol, # Figure 8
       panel = function(x,y){
         panel.xyplot(x,y) # plot points
         panel.loess(x,y)  # add smooth line
       }
)
```

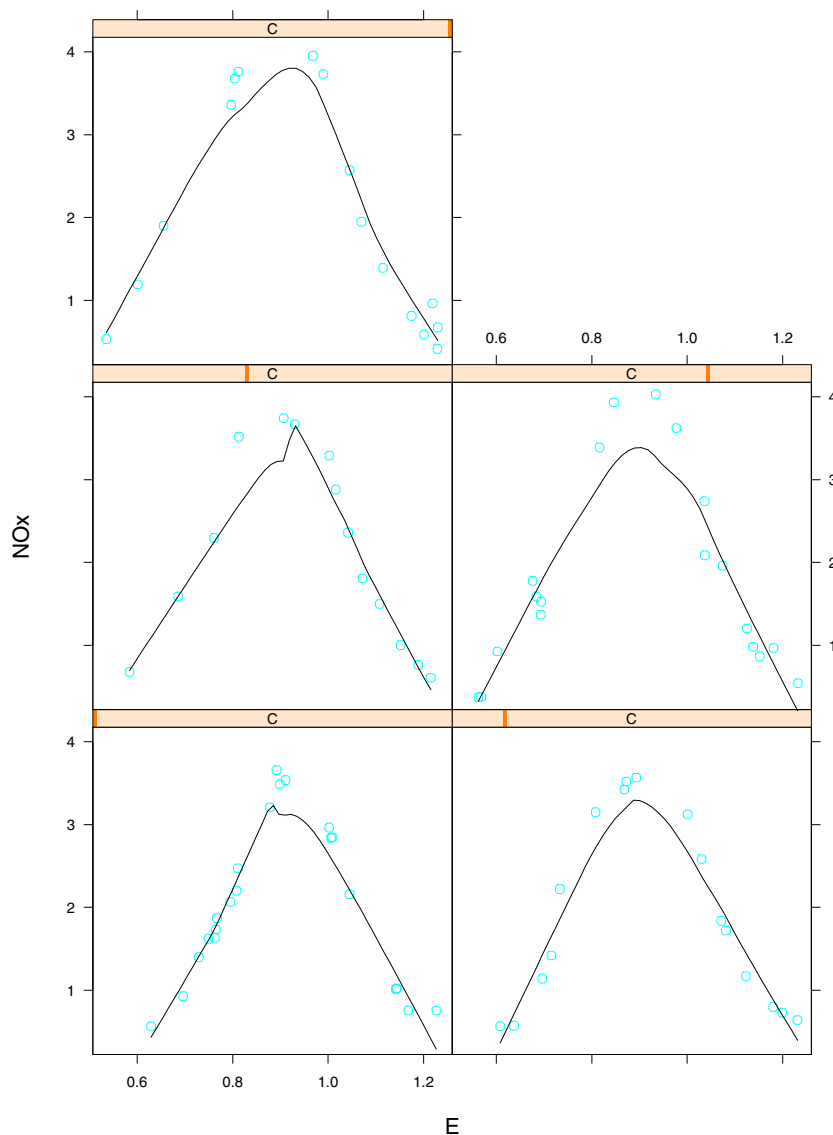


Figure 8. A Trellis display of engine emissions data. The smooth line was put on by the in-line panel function.

The previous example used two panel functions that come with the Trellis software: `panel.xyplot` and `panel.loess`. As described earlier, `panel.xyplot` is the default panel function for `xyplot`. The `panel.loess` function is one of a group of functions that are useful in constructing panel functions. This group includes: `panel.abline`, `panel.fill`, `panel.grid`, `panel.lmline`, `panel.loess`, and `panel.qqmathline`.

The `panel.abline` function plots one or more lines on the panel. The lines can be

horizontal, vertical, or the result of a linear fit expressed in the form $y = a + b * x$. The function uses Trellis line style conventions and will not generate warning messages if any of the lines fail to intersect the plot.

The function `panel.fill` is used to fill in the entire panel with a suitable color or gray level. Aside from making pretty colored panels, it also allows further plotting on the panel in the background color, `col=0`. For example, if you are on a device with a white background, you can fill each panel with gray and later use `panel.grid(col=0)` to draw a reference grid on top in white. The `panel.grid` function is designed to put a reference grid on each panel, to enhance the visual comparisons from one panel to another. It should generally be the first thing drawn on a panel (after `panel.fill`, though), so that the reference grid will not obscure any important information. Arguments control how many grid lines are used:

```
panel.grid(h=-1, v=4)
```

draws a horizontal grid aligned with the vertical axis ticks and places 4 vertical reference lines on the panel. Although conventional plots often have grids aligned with axes, there is no need for this. In many cases the grid is not there to help the viewer read-off numeric values — it provides reference lines against which patterns can be compared.

Functions `panel.lmline` and `panel.gqmathline` are designed to put fitted lines on the panel. The former draws a line fit with least-squares; the latter does a robust fit to a theoretical quantile plot.

As illustrated in Figure 7, the `panel.loess` function adds a smooth curve using the non-parametric `loess` procedure; various arguments to `loess` can be supplied.

The `panel.superpose` function is often used as a replacement for `panel.xyplot` when each panel is intended to contain multiple lines or points based upon the values of a grouping variable. The high-level function is given an argument, `groups=`, that specifies the grouping variable (as a factor or something that will be coerced to a factor). This `groups` argument will be passed down to `panel.superpose` to control the plotting symbols and colors used to display the specified groups. Examples using `panel.superpose` appear later in Figures 20 and 21.

One important thing to remember about the Trellis software — everything is written in the S/S-PLUS language. This means that you can read it and modify it. The panel functions are intended as the place for users to customize their plots, so they are generally short and easy to understand and modify. However, all of the Trellis functions are accessible and can often be understood with a little bit of work. In any case, the point is that you should not be deterred if the Trellis software doesn't do exactly what you want — you can change it yourself.

2.4 Formulas

The `formula` argument for Trellis functions often comes in the form:

```
y ~ x | g1 * g2 ...
```

Here, `x`, `y`, and the `g` variables are S objects or S function calls. The `y` variable describes the variable plotted on the vertical axis. In many cases, both the `x` and `y` variables are numeric, as in the examples of the `xyplot` function that produced Figure 1. However, for the univariate display functions such as `bwplot`, the `y` variable is treated as a factor (and is made into one if it isn't a factor already). For example, Figure 9 is produced from:

```
bwplot(voice.part ~ height, data = singer,  
       xlab="Height (inches)") # Figure 9
```

Notice that in producing Figure 9, we have not used any given variables, hence the Trellis display consists of only one panel.

When given variables are used in a formula, they are handled as factors or shingles. If a

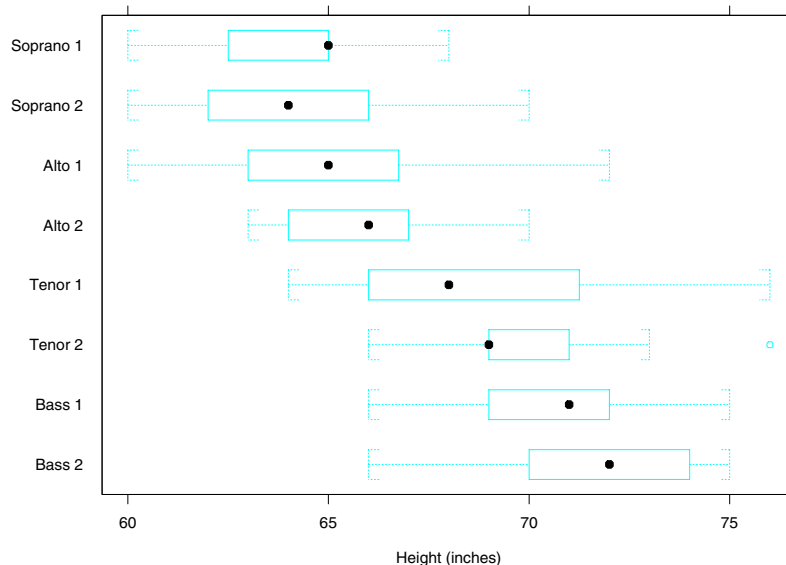


Figure 9. A box and whisker plot showing heights of members of a choral group, arranged according to the part they sing.

given variable is a character vector, it is automatically turned into a factor (with the sorted, unique values of the vector as the factor's levels); if it is numeric, it is turned into a shingle (with zero-width intervals at the unique values of the numeric variable; in this case, it may be better to create a shingle yourself, perhaps by using the `equal.count` function, so that you can control the intervals chosen.)

Certain Trellis functions take slightly different kinds of formulas. Univariate functions that produce a whole panel from one set of data omit the `y` variable and have a formula like

```
~ x | g1 * g2
```

A rationale for this is that, for example, a histogram plots the data values along the `x`-axis and internally computes the `y` values that determine the heights of the bars. So, Figure 10 is produced from:

```
histogram(~ height|voice.part, data = singer,  
          xlab="Height (inches)") # Figure 10
```

In Figure 10, each value of the given variable has produced a separate panel. This is a general principle: the `x` and `y` variables to the left of the vertical bar in a formula make up each panel and the given variables cause multiple panels to be produced.

The function `qq` takes a simple formula

```
y ~ x | g1 * g2 * ...
```

where the numeric response `x` is split into exactly two groups by the `y` variable. The resulting panels each show a quantile-quantile plot of the data from one group against the data from the other group.

The function `qqmath` produces quantile plots. Its formula is:

```
~ x | g1 * g2
```

It also takes an argument, `distribution`, that specifies a quantile function — a function of a vector of probabilities that produces a set of quantiles. The quantile function is often a standard

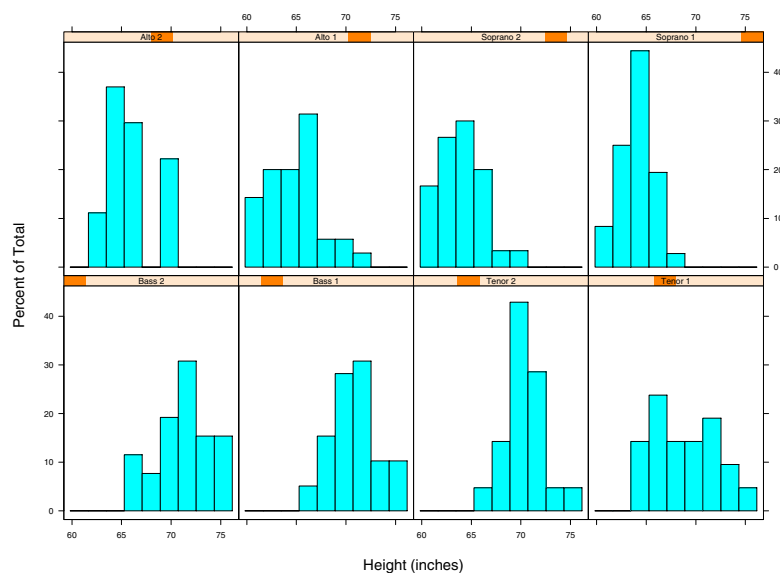


Figure 10. Histograms of singer heights by voice part.

reference distribution, `qnorm` (to compare the `y` data to the normal distribution) or `qunif` (to compare to a uniform distribution). As an example, we can see how the heights of singers within voice part compare to a normal distribution:

```
qqmath(~ height | voice.part, distribution = qnorm,
       data = singer) # Figure 11
```

The plot of Figure 11 could be improved, perhaps by adding a reference line and grid to make it easier to assess the quality of the fit and by adding a nicer x-label. The plots might also look better if they were made square (later, we can see if there is an even better aspect ratio). This is accomplished in Figure 12, produced by:

```
qqmath(~ height | voice.part, distribution = qnorm, # Figure 12
       data = singer,
       aspect = 1,
       prepanel = prepanel.qqmathline,
       panel = function(x, y, ...) {
         panel.grid()
         panel.qqmathline(y, distribution=qnorm, ...)
         panel.qqmath(x, y, ...)
       },
       xlab = "Unit Normal Quantile"
)
```

The `prepanel` function will be described later, in the “ADVANCED CONCEPTS” section.

Suppose we thought that the square roots of the singer heights, rather than the raw values, were more appropriate to compare to a normal distribution. We could have used the `sqrt` function inside the formula:

```
qqmath(~ sqrt(height) | voice.part, distribution = qnorm, data = singer,
      ...)
```

Functions and other S expressions can be used inside the formula. However, one disadvantage of using complicated expressions inside the formula is that the default labels are also more complicated and can therefore be more difficult to read. This is especially true for strip labels. An alternative, especially if you are using `equal.count` to generate shingles for given variables, is to

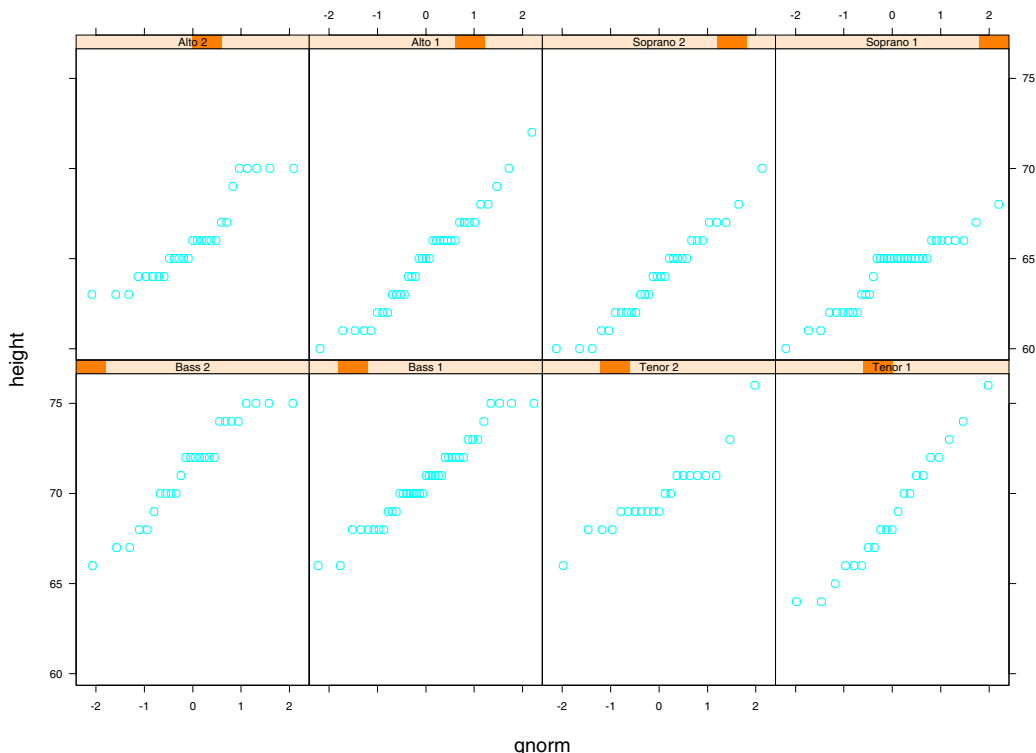


Figure 11. Singer heights within voice part compared to theoretical quantiles from a normal distribution.

create a new object with a nice-looking name (the variable’s name is used in labeling shingles) prior to calling the S function.

The `contourplot` and `wireframe` functions also take a different sort of formula. Since these plots are constructed from 3-D data, the formula is

$$z \sim x * y \mid g1 * g2 \dots$$

Here, x , y , and z are numeric vectors and x and y are evaluated on a regular grid. For example, the z values may represent a surface evaluated at 60 points, made up from 6 unique x values and 10 unique y values. If given variables are present, there should be a regular surface of data for each unique combination of values of the given variables, although the x and y values need not be the same for each surface. (For an example, see section 4.1, “3-D PLOTTING”.)

2.5 Trellis Objects

All of the high-level Trellis functions (those listed in Table 1) return an object of class `trellis` as their value. These objects are ordinarily plotted straight away, because the `print` method for this class, `print.trellis` actually plots the objects. However, the fact that Trellis functions return objects means that those objects can be stored and replotted later, perhaps when a different device is active. For example, in S

```
NOxplot <- xyplot(NOx ~ E | C, data = ethanol) # save result (no plotting)
trellis.device(iris4d) # graphics on the workstation
NOxplot # now plot it
trellis.device(postscript) # change devices
NOxplot # plot it again for new device
```

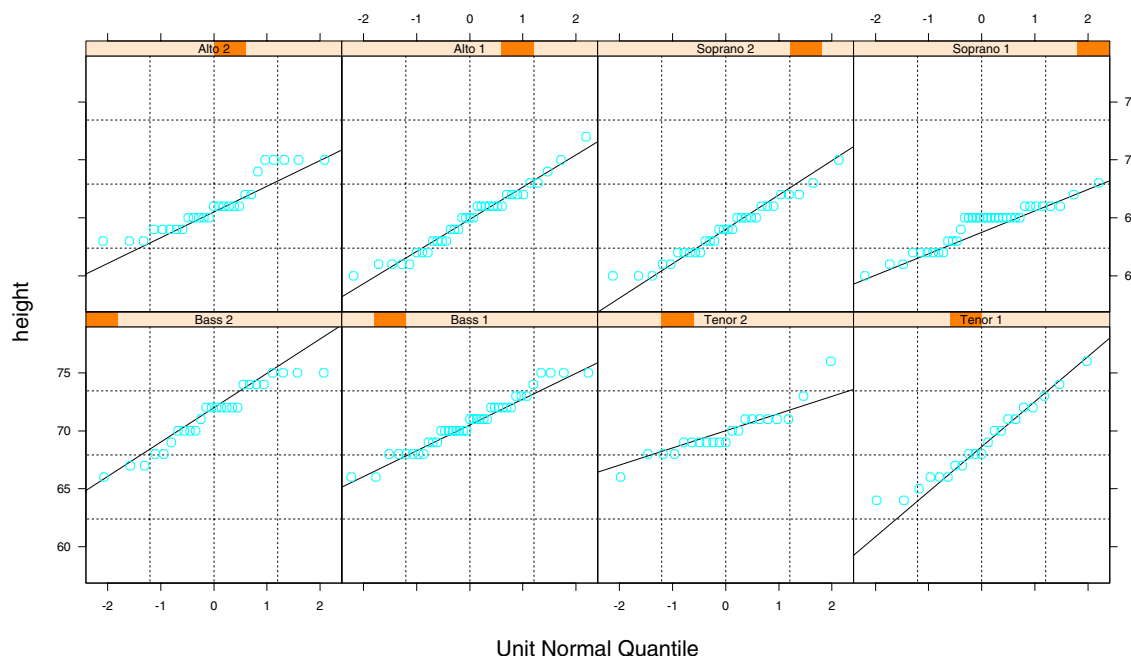


Figure 12. An improved version of the singer heights data with a comparison to the normal distribution.

S-Plus users have slightly different device functions and can have several functions active at once. Refer to the *Trellis Graphics User's Manual* MathSoft (1995), for more details.

It is sometimes useful to call `print.trellis` directly, because it has several arguments that can be used to position Trellis plots on a page. Suppose, for example, that you have two Trellis objects, `obj1` and `obj2` and would like to position them one above the other on a single page of output. If the two objects were identical in type, both scatterplots, for example, it might have been easier to use `xyplot` with a made-up given variable to do this. However, suppose `obj1` is a multi-panel Trellis display and `obj2` is a single display of a different type.

Let's be more concrete. Let

```
obj1 <- qqmath(~ sqrt(height) | voice.part, distribution = qnorm,
              data = singer, ...)
obj2 <- histogram(~ height, data = singer)
```

and now combine them using `print.trellis`:

```
print.trellis(obj1, split=c(1,2,1,2), more=TRUE) # Figure 13
print.trellis(obj2, split=c(1,1,1,2), more=FALSE)
```

The `split` argument to `print.trellis` is a vector of length 4, `c(x,y,nx,ny)` saying to use the `x,y` position of an `nx` by `ny` rectangular layout. The `more=` argument tells whether more plotting is to be done on the same page.

As some of our examples have shown, the description of some Trellis displays (through arguments to the high-level Trellis function) may take several lines. Suppose you are producing Trellis displays and are trying to adjust things to come up with just the right look. One way to do

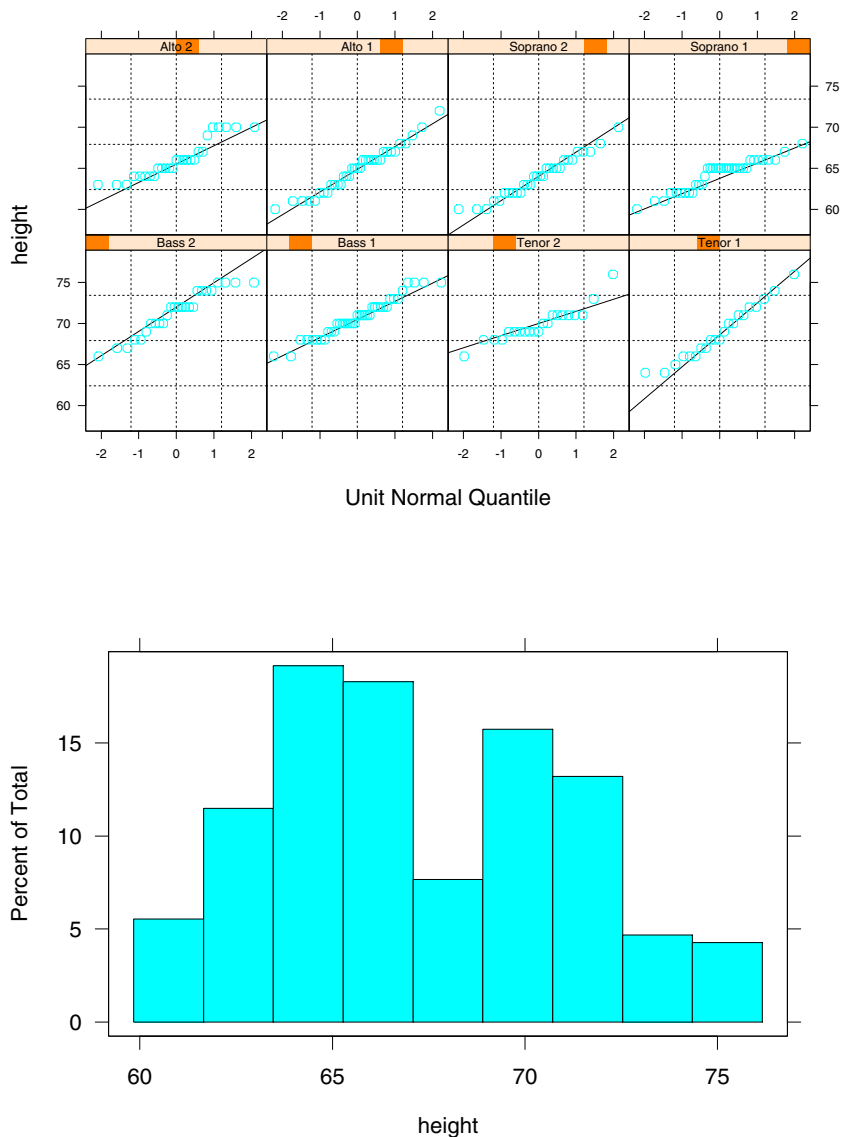


Figure 13. Two different Trellis displays on a single “page”, positioned by `print.trellis`.

this is to store the expression in a file, use your favorite editor to change the file, and then use `source` to execute the expression in the file. One thing to note about this plan: automatic printing of results is not done within a source file, so you should surround your expression with `print()`, for example, put the expression

```
print( xyplot(NOx ~ E | C, data = ethanol) )
```

in file `foo` and execute

```
source("foo")
```

An alternative is to create a Trellis object and then modify and redisplay it with the `update` function.

```
NOxplot <- xyplot(NOx ~ E | C, data = ethanol)  
NOxplot # display it
```

```
update(NOxplot, layout=c(3,2)) # display with new layout
NOxplot2 <- update(NOxplot, pch="X") # change plotting character
NOxplot2 # display it
```

You can also save the result of `update` and update that.

2.6 Layout

The formula given to a Trellis display determines the order in which subsets of the data are produced. The first *packet* of Trellis data corresponds to the first level of each of the given variables (or the first interval of a shingle). The second packet is at the second level of the first given variable and the first level of each of the other given variables. This goes on until the last level of the first given variable. Next, the second level of the second given variable is reached and all other variables go to their first level. All of the data packets are produced in this way. This determines the *packet order*. If there are N_1 , N_2 , and N_3 levels for 3 given variables, then there will be a total of $N_1 * N_2 * N_3$ packets.

How does this relate to the layout of panels on the page? The page is divided into columns and rows as specified by the `layout=` argument. The *panel order* is defined so that the first panel begins at the lower left corner and successive panels fill the bottom row. Next, panels fill the second row from left to right. The total number of panels in the panel order is determined by the layout specification. Remember that Trellis displays are filled as graphs, from the origin in the lower left, not top-down as in a table. That is also why the number of horizontal panels (columns) precedes the number of vertical panels (rows) in the layout specification.

When it comes time to display a Trellis graph, the packets and panels are associated with one another: the first packet (in the packet order) goes into the first panel (in panel order), and so on.

The important concept is that the packets are produced in packet order, and the panels, in panel order, are filled by those packets.

If you do not specify the `layout=` argument yourself, it has defaults that depend on how many given variables are in the formula. For one given variable, the default layout is chosen by a *layout optimization algorithm*. It chooses the number of rows and columns to maximize the page area devoted to the panels, taking into account the aspect ratio of the panels, the size of the device and the size of the strip labels. For two or more given variables, the default number of columns is determined by the number of levels of the first given variable, the rows by the second given variable, and the rest of the given variables vary across pages.

There is also a way to have the layout optimization algorithm assist you with 2 or more given variables. If you specify a layout argument of the form `layout=c(0, n)`, the optimization is carried out for n plots per page.

Occasionally, the layout of a Trellis display may be problematic. For example, suppose you have two factors, one with 14 levels and another with 10 and would like to have 3 columns and 5 rows on each page for 10 pages. If you use `layout=c(3, 5)` the result will not be satisfactory, because there will be 15 panels on a page, so the arrangement of factor levels will not be consistent from page to page. Things would be better if you could have 14 panels on the page, but 7 by 2 or 2 by 7 layouts cause the panels to be too tall or fat. You can solve the problem by using the 3 by 5 layout along with the `skip=` argument to skip the center panel or perhaps the upper right panel, when panels are traversed in panel order. The `skip=` argument is a logical vector that tells, for each panel on a page, whether it should be skipped. The `skip` vector is replicated as necessary and can even be longer than the number of panels on a page, so that complicated multi-page skip patterns can be expressed. For example, the combination of

```
layout=c(3, 5), skip=c(rep(F, 14), T)
```

would skip the upper right hand panel and

```
layout=c(3,5), skip=c(rep(F,7),T,rep(F,7))
```

would skip the middle panel.

Another similar situation might involve two given factors, A and B, with 5 and 2 levels respectively. As an exercise, consider how you could use `skip` to get 3 columns and 4 rows on a page filled out reasonably.

Sometimes, the interaction of the page size and the number of factor levels can make a nice layout difficult. Consider a plot of a fit to the ozone data where ozone is plotted as a function of radiation for 6 levels each of temperature and wind speed.

```
xyplot(env.fit ~ radiation | temperature * wind, # Figure 14
       data = twr.grid,
       panel = function(x,y) {
         panel.grid(h = 2)
         panel.xyplot(x, y, type = "l")
       },
       aspect = "xy",
       xlab = "Radiation (langleys)",
       ylab = "Cube Root Ozone (cube root ppb)")
```

The plot has an aspect ratio chosen by the slope of the lines in the panels and these banking computations (described in section 2.7, “Axes”) make it tall and thin.

Suppose we needed to show this data on a landscape-oriented piece of paper or on a display screen that was wider than high. How could we do this, preserve aspect ratios, and still make good use of the display area? Rather than a display with 6 rows and 6 columns, we display the panels in a layout with 12 columns and 3 rows, something that will fit the page quite well. The trick to doing this is to reorder the levels of the second given variable, `wind`.

Why does this work? Think of how the `layout` argument interacts with the levels of the given variable; in other words, think of the interaction between panel order and packet order. Trellis cycles through levels of the first given variable, `temperature` for the first level of the second given variable, `wind`. This uses 6 panels, starting from the lower left of the display and going horizontally. There are still 6 panels left in the bottom row, and those are taken up by 6 the temperature values and the second level of `wind`. We have reordered the levels of `wind` so that the resulting plot appears to be organized into a left half and a right half.

```
wlevels <- seq(4, 16, length = 6)[c(1,4,2,5,3,6)] # Figure 15
strip.shingle <- trellis.par.get("strip.background")
xyplot(env.fit ~ radiation | temperature * shingle(wind,wlevels),
       data = twr.grid,
       layout = c(12,3),
       between = list(x=c(0,0,0,0,0,1)),
       strip = function(...)strip.default(...,strip.names=FALSE),
       key = list(
         text = list(c("wind","temperature")),
         rectangle=Rows(strip.shingle,2:1),
         space="Top"
       ),
       panel = function(x,y) {
         panel.grid(h = 2)
         panel.xyplot(x, y, type = "l")
       },
       aspect = "xy",
       xlab = "Radiation (langleys)",
       ylab = "Cube Root Ozone (cube root ppb)")
```

You may find the panel order in Trellis displays somewhat peculiar, particularly the part about the rows of panels being filled from bottom to top. The reason for this is that the vertical

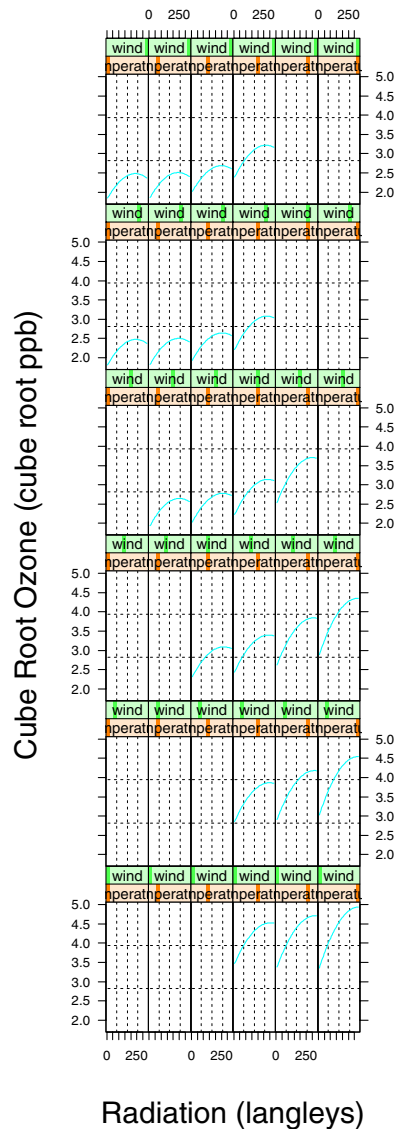


Figure 14. Loess fit to the ozone data with aspect ratio chosen by banking computations.

axis of a graph generally increases from bottom to top, and if given variables have their levels in increasing order, the standard Trellis panel order will make the given variables increase from bottom to top. That said, there are occasions when a top-to-bottom order might be convenient. This is called *table order*, left-to-right top-to-bottom, and is specified by using `as.table=TRUE` as an argument to a high-level Trellis function. For an example, see the function `example.calendar`.

Notice that the plot of Figure 17 contains a key that allows the reader to see how the shingle colors correspond to the given variables. (Also, for those of you who are looking carefully, the rectangle parameters given to `key` are done in reverse order, since `key` draws things top to bottom while strip labels are constructed bottom to top!) The in-line `strip=` function was used to suppress the strip names. Also, the `between=` argument allows us to insert space (in units of characters) between panels in the x- or y- direction; we used it to distinguish between the two major levels of wind speed.

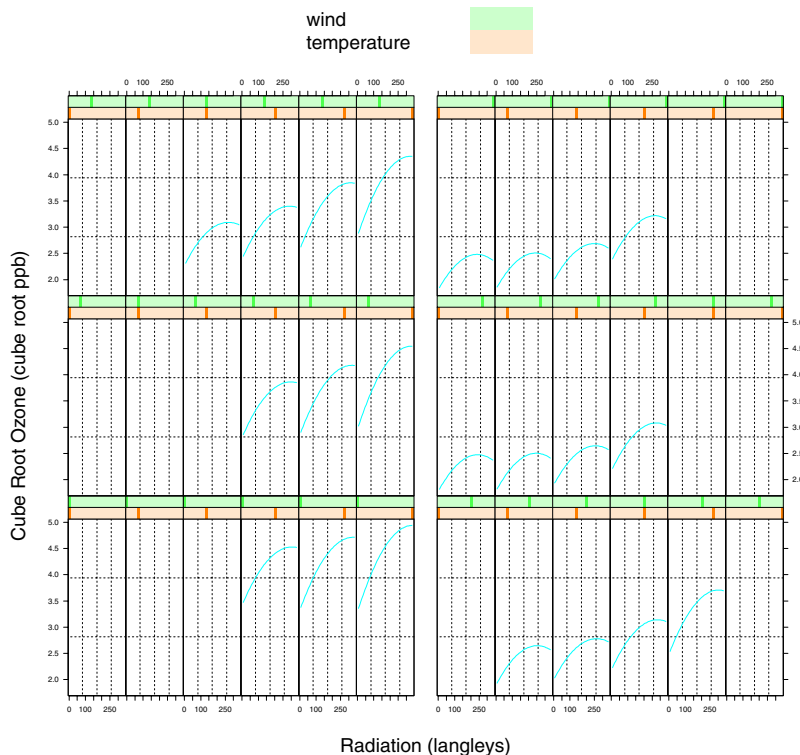


Figure 15. Loess fit to the ozone data. The key appears in color on a color device. The leftmost 6 columns show temperature for the lowest 3 levels of wind speed; the rightmost 6 columns are for the highest wind speeds.

Now that we have used the `panel=` and `strip=` functions, it should come as little surprise that there is another function accepted by high-level Trellis functions. A `page=` function is called at the end of each page of output and can be used to put on page numbers or other identifying information. For example

```
xyplot( ... ,
        page = function(x) if(x>1)
            mtext(paste("page", x), side = 1, line = par("oma")[1]-1,
                  outer = TRUE, adj = 1)
        )
```

would number each page but the first of a multi-page display, placing the page number at the bottom right of the page.

2.7 Axes

Alignment of axes is one characteristic that makes the layout of Trellis displays so powerful. Unlike arbitrary sized and positioned windows, such as those produced by many software packages, Trellis displays ensure that axes are aligned and can be readily compared to one another. In support of this capability, Trellis functions take several arguments that give control over the axes. Most basic are the `xlim=` and `ylim=` arguments, that allow specification of lower and upper limits for axes. For example, specifying, `xlim=c(0,100)` ensures that the x-axis will accommodate values between 0 and 100.

Although `xlim=` and `ylim=` are convenient, they are only a quick way of specifying one detail of the scales on a Trellis plot. Full axis control is available through the `scales=`

argument. It not only gives precise control over tick marks, labels, etc., but it also controls how the horizontal and vertical axes relate from panel to panel. In general, `scales=` is a list with components named `x=` and `y=`. Each of these components, in turn, is a list with components in `name=value` form, giving information about the `x` or `y` axis. If the `scales=` list contains other components, they are taken to apply to both the `x`- and `y`-axes. What kinds of things can you specify in the `scales=` list? Here's an example:

```
scales = list(
  x = list(col=2, tck=.5, at=c(.7, .9, .11),
    label=c("good", "better", "best"),
    alternating=FALSE),
  y = list("sliced", nticks=17, tick=FALSE, log=2),
  cex=.75
)
```

This would place 3 tick marks on the `x`-axis at coordinates `.7`, `.9` and `.11`, labeled with the words “good”, “better”, and “best”, would make the tick marks half their normal length, and would draw the axis line, tick marks and tick labels in color 2. In addition, the `x`-axis labels would not alternate from the bottom to the top of the page — they would all be on the bottom side. Each panel's `y`-axis would be logarithmic (base 2), would have approximately 17 tick labels (although no tick marks), and the `y`-axes would all have the same range (the “sliced” specification). On both axes, the character size would be `.75` of standard for that device.

Of course, most of the time you will not need to specify so much about axes, but the point is that the `scales=` argument gives you lots of control. In fact, it also lets you suppress the drawing of one or both axes:

```
scales=list(draw=FALSE)
```

suppresses both the `x`- and `y`-axes. This might be appropriate if, for example, the plot were a map or other recognizable entity that did not require a numeric scale.

One of the most common uses of the `scales=` argument is to specify axis relationships. By default, the scale argument takes the value `relation="same"`, which means that the horizontal axes will be identical on all panels and the vertical axes will be identical on all panels (the horizontal and vertical axes are not necessarily the same as one another, though). The value `relation="sliced"` means that the axes are set up to have the same units per inch — as if each axis was a slice from one consistent larger axis. This means that the maximum minus minimum value for each axis is identical. Finally, the value `relation="free"` can be used to allow each panel to have freedom in constructing an axis that just accommodates the data to be plotted there. Because it is used so often, the `relation=` component is the only one of the scale components that need not be named explicitly. For both “sliced” and “free” scaling, axes are drawn for each panel, taking up extra space on the display.

Suppose we had wanted to modify Figure 14 by allowing the `y`-axes to vary from plot to plot. If we include

```
scales=list(y="free"),
```

and remove `aspect="xy"` from the call, we generate Figure 16, which now makes most comparisons extremely difficult, since the vertical axes are now scaled differently.

When a Trellis display is finished, the coordinate system is left in a state that in general has no relation to the scales that were plotted on the page. The margins and outer margins are returned to their previous values and the multiple figure parameters are set for one plot per page. Because these parameters are reset, it is seldom reasonable to add to Trellis displays after they are produced. If you want to augment a Trellis display, do so by means of the `panel` or `page` functions.

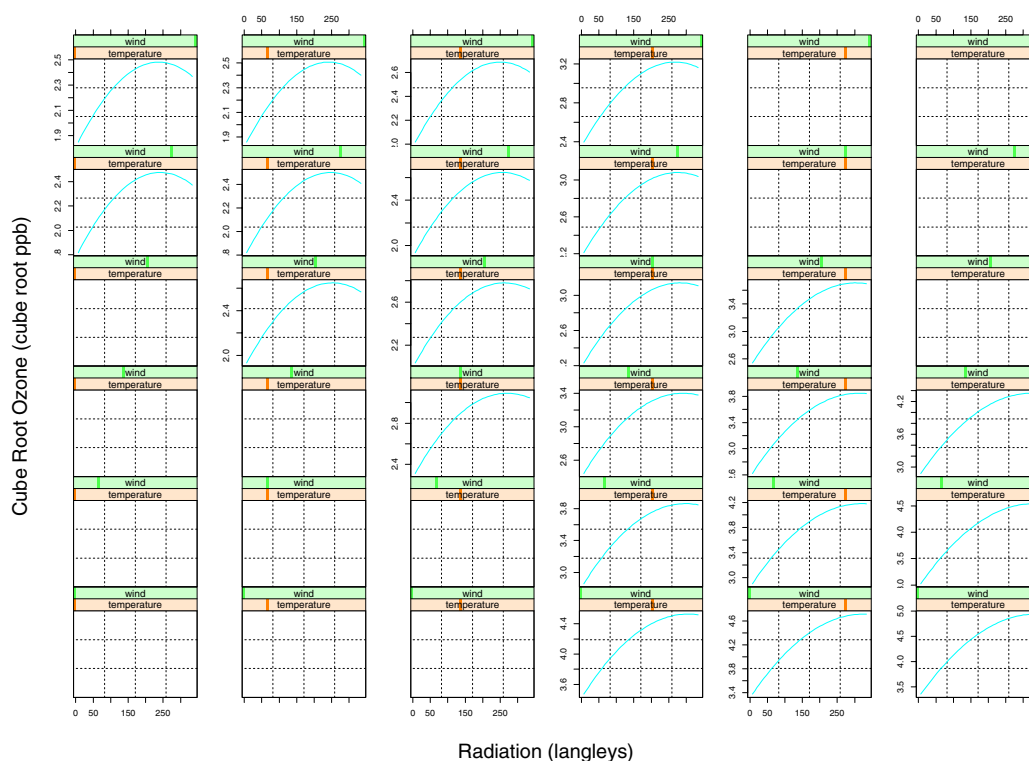


Figure 16. Loess fit to the ozone data, no aspect ratio control, vertical scales allowed to vary from panel to panel.

2.8 Aspect Ratio

One of the important capabilities of Trellis displays is the ability to control the panel aspect ratio in order to produce more understandable plots. The aspect ratio is the physical height of a panel divided by its physical width. (Physical measurements are in inches or centimeters, not in data units.) The aspect ratio can be specified numerically, or it can be computed by *banking* calculations. In particular, Cleveland has experimental evidence that angles near 45° are the easiest for the viewer to discriminate. Banking computes an appropriate aspect ratio to make the important characteristics of the display appear close to $\pm 45^\circ$. The `aspect=` argument controls the aspect ratio of Trellis displays. If a numeric value is given, that aspect ratio is used for all panels. For example, `aspect=1` says that each panel should be square. Another common use for numeric values is to ensure a physical relationship between the x- and y-coordinates:

```
aspect = diff(range(y))/diff(range(x))
```

would set up an aspect ratio that just matched the ratio of the y and x coordinates. This is what is required for making circles appear circular, for preserving physical shapes, etc. Finally, using `aspect="xy"` performs the banking to 45° computations on the x- and y-data for all panels. The sorted x-values and corresponding y-values are converted to fractions of the range, are differenced, scaled by their length and all thrown together into one computation that finds the aspect ratio that brings the segments closest to 45° .

Because control over the aspect ratio is best done with at least a hint from the user, the default is to use `aspect="fill"` which makes the aspect ratio appropriate so that the collection of panels fills the display area.

There is a lot of wasted space in Figure 14 because of the chosen aspect ratio. Suppose we had not tried to control the aspect ratio, but instead allowed the panels to fill the space (using the default `aspect="fill"`). Now we have a plot where the lines are too flat to compare easily (Figure 17).

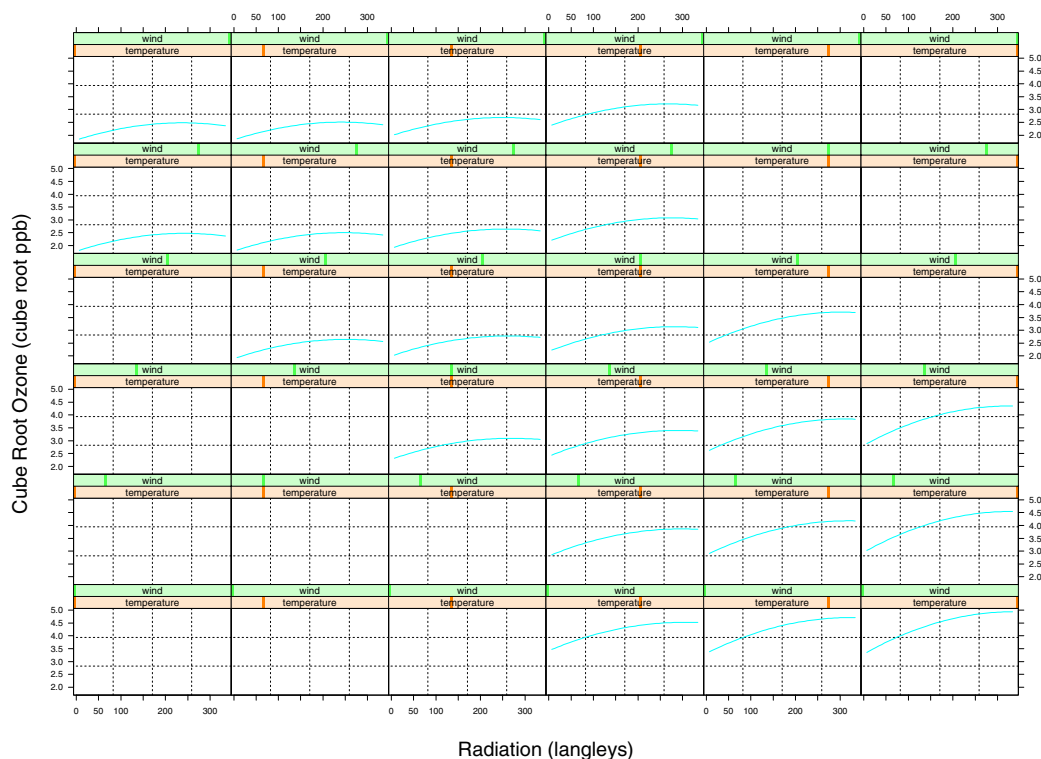


Figure 17. Loess fit to the ozone data, no aspect ratio control.

2.9 Data Structures

There are two data structures that are of particular importance to Trellis displays. These are *factors* and *shingles*. Factors were described in detail in Chambers & Hastie (1992). They are used to hold categorical values, for example, colors (red, green, blue), states (Alaska, Alabama, ...), etc. Factors are created by the `factor` function. The other data structure, a shingle, is a vector of numeric data and a set of intervals. Shingles get their name because, like shingles on a roof, the intervals can overlap one another. For each interval, some subset of the numeric values fall into that interval.

Two functions are available for dealing with shingles. The first is `shingle`, which creates a shingle by specifying the data and intervals. By default, the intervals are zero-width at the unique values of the data. This sort of shingle is almost like a factor, although the precise numeric value of each interval is preserved. Consider, for example, the engine data and the values of compression ratio. Treated as a factor, there would be 5 values. Treated as a shingle, there would also be 5 values, but the numerical spacing of the compression ratios would be preserved.

The second function for creating shingles is `equal.count`, where a numeric vector is divided up as equally as possible into a specified number of overlapping intervals; the amount of overlap between adjacent intervals can also be specified.

Why are we spending time discussing these data structures? One reason is that shingles are

new to Trellis displays. The other is that the ordered level names of factors are used to label Trellis plots, thus it is important to describe how these labels are created and manipulated. We chose this strategy to avoid having complicated labeling arguments for Trellis functions. Thus, to change the labels, you change the factor.

```
levels(month.observed) <- month.name # full names
...
levels(month.observed) <- month.abb # abbreviated names
```

This is a powerful notion and will come up in later examples.

Two other functions are useful for ordering the levels of a factor. The first is `ordered`:

```
ordered(grades) <- c("Poor", "Fair", "Good", "Excellent")
```

The second function useful for modifying factors is `reorder.factor`. It allows you to rearrange the levels of a factor based upon a computed value. For example, you could reorder a hypothetical factor `state` from its original alphabetical order to an order based upon the median income of the observations in that state:

```
state <- reorder.factor(state, income, median)
```

The function `make.groups` is often useful with Trellis displays. It constructs a data frame from several vectors and the data frame can be passed in to a Trellis function. Suppose that we have several vectors and want to see box and whisker plots or histograms of each. For example, we want to compare payoffs of the New Jersey Pick-It lottery from three time periods:

```
make.groups(lottery.payoff, lottery2.payoff, lottery3.payoff)
```

creates a data frame with two components: `data` and `which`. The `data` component is simply the combined numbers from all the `make.groups` arguments. The `which` component is a factor with 3 levels, giving the names of the original data vectors. So, we could use

```
bwplot(which ~ data, data = make.groups(lottery.payoff,
lottery2.payoff, lottery3.payoff))
```

to produce box and whisker plots or

```
histogram(~ data | which, data = make.groups(lottery.payoff,
lottery2.payoff, lottery3.payoff))
```

to produce histograms of the three sets of data.

Just as `make.groups` converts vectors to data frames for use with Trellis functions, there are also functions to facilitate graphics with *arrays* and *time series*. The functions `as.data.frame.array` and `as.data.frame.ts` convert arrays or time series into data frames (sorry about the long names, for those of you who don't like to type).

Consider the object `iris`, a 3-way array with a `dim` vector like this:

```
> dim(iris)
[1] 50 4 3
```

We can turn `iris` into a data frame in preparation for plotting by using:

```
iris.df <- as.data.frame.array(iris, col.dims = 2)
```

The resulting data frame has what used to be its second dimension turned into 4 columns, so that it looks like this:

```
> iris.df[1:5,]
  Sepal L. Sepal W. Petal L. Petal W. flower species
1     5.1     3.5     1.4     0.2     1 Setosa
2     4.9     3.0     1.4     0.2     2 Setosa
3     4.7     3.2     1.3     0.2     3 Setosa
4     4.6     3.1     1.5     0.2     4 Setosa
```

```
5      5.0      3.6      1.4      0.2      5      Setosa  
...
```

So, given a data frame like this, we can now use it with Trellis functions. Let's try a *parallel coordinate* plot. The function `parallel` takes a formula in the form

```
~ x | g1 * g2 ...
```

where `x` is a matrix. The expression

```
parallel(~iris.df[,1:4] | iris.df[,6], layout = c(3,1), main = "Iris Data")
```

produces Figure 18.

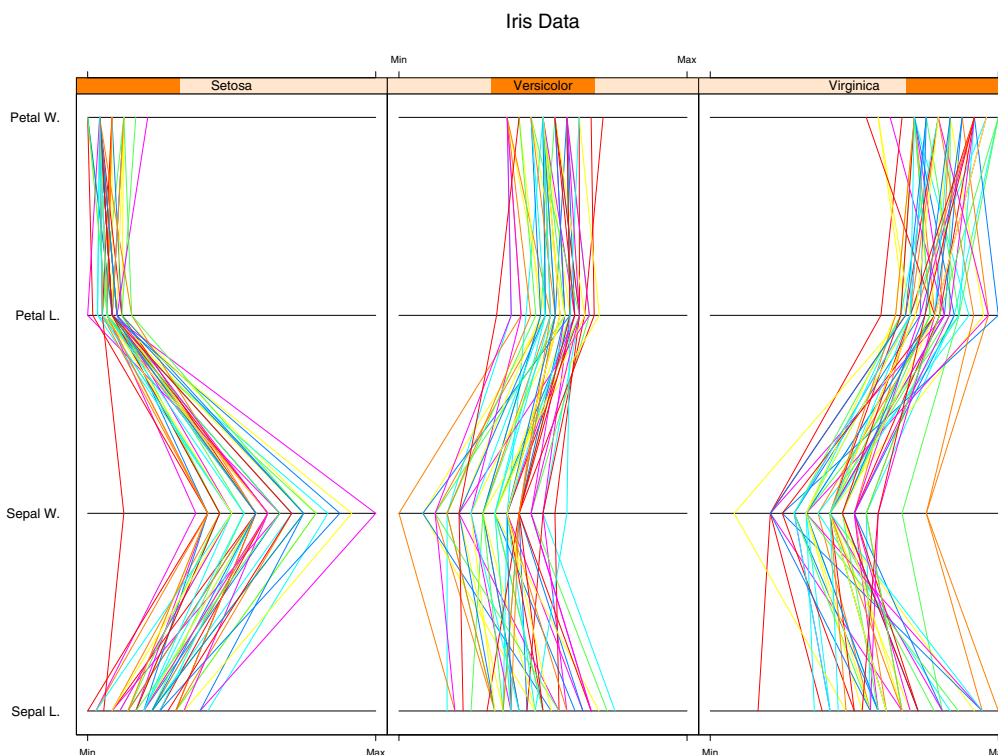


Figure 18. Parallel coordinate plot showing three groups of the iris data.

The function `as.data.frame.ts` takes one or more time series as arguments and produces a data frame with components named `series`, `which`, `time`, and `cycle`. The `series` component is just the data from all of the time series combined into one long vector. The `time` component gives the time associated with each of the points (measured in the same units as the original series, e.g. years), and `cycle` gives the periodic component of the time (e.g. 1=Jan, 2=Feb, ...). Finally, the `which` component is a factor that tells which of the time series the measurement came from. (In this case there was only one series, `hstart`, but in general `as.data.frame.ts` can take many arguments). For example:

```
> as.data.frame.ts(hstart)
  series which      time cycle
1  81.9 hstart 1966.000   Jan
2  79.0 hstart 1966.083   Feb
3 122.4 hstart 1966.167   Mar
4 143.0 hstart 1966.250   Apr
5 133.9 hstart 1966.333   May
```

...

Using `as.data.frame.ts`, we can produce a plot that shows housing starts from 1966 to 1974, broken down by their monthly levels.

```
timeplot(series ~ time|cycle, data = as.data.frame.ts(hstart), # Figure 19
type = "h", xlab="Year", ylab="Housing Starts by Month")
```

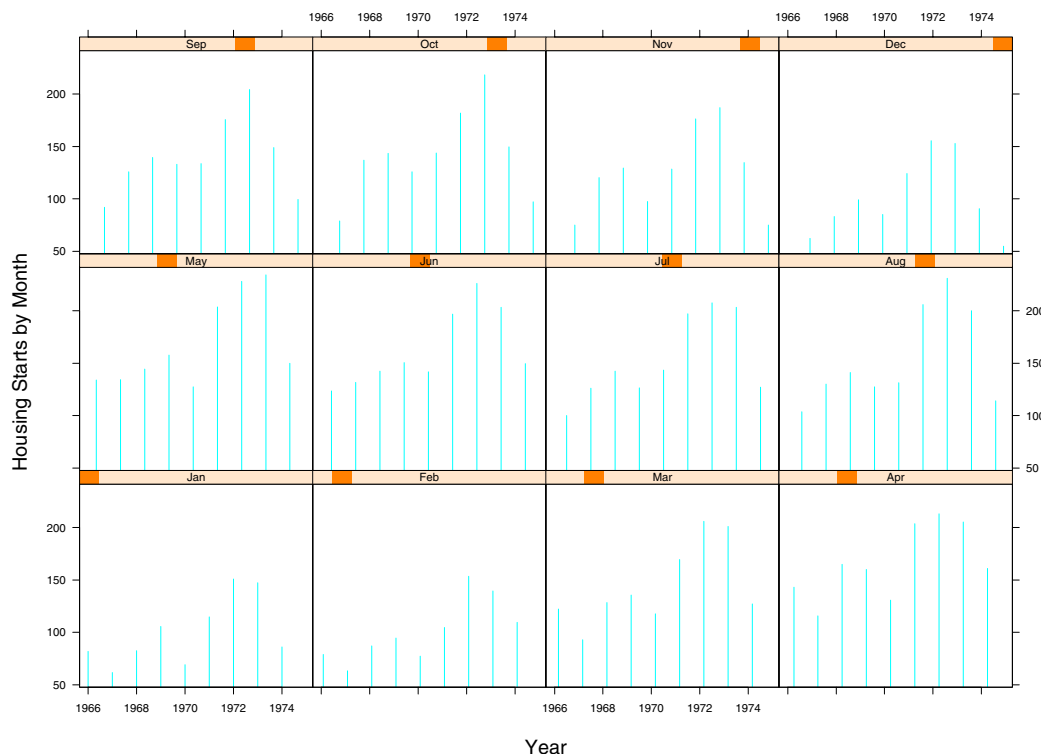


Figure 19. Each panel shows the housing starts for a particular month during the years from 1966 to 1974.

Notice that Figure 19 would change if `time` in the formula were replaced with `trunc(time)`; the vertical lines for each month would line up exactly at the year marks.

2.10 Labeling (Titles, Strip Labels, Keys)

Labeling is an important part of Trellis displays. The Trellis functions take arguments, `xlab=` and `ylab=` to control the x- and y-axis labels. By default, these labels are made up from the names of the variables (or the expressions) plotted there. Of course, you can often make the labels more meaningful by giving explicit values, for example:

```
xlab="Equivalence Ratio", ylab="NOx (micrograms/J) "
```

would do a nicer job of labeling Figure 1. Also, the arguments `main=` and `sub=` may be given to any high-level Trellis function to put a main title (at the top) and a subtitle (at the bottom) on each page. Any of these label arguments can be given as a list, so that the label comes along with associated graphical parameters, e.g.:

```
xlab=list("Equivalence Ratio", cex=1.5, col=2)
```

Strip labels are an important part of a Trellis display. By looking at the panel and its

associated strip labels, you can see the given variables and which levels are used for the particular panel. By default, strip labels for shingles give the name of the shingle variable and show by shading the fraction of the entire data range taken up by the current shingle interval. Strip labels for factors give the label corresponding to the factor level and are shaded to show the order of this level within the factor. By now you probably can guess that there are many ways to customize strip labels. At the most extreme level, you can pass an entire strip label drawing routine as the `strip=` argument to high-level Trellis plots (the default strip label routine is called `strip.default`). However, most of the time there is no need to do this much. The argument, `par.strip.text` allows you to pass in a list of graphical parameters that controls the rendering of the text strings in the strip labels. The most common use of this is to control the size of the text in the strip labels; the size of the strip label box changes to accommodate the characters. You can also use `par.strip.text` to control the color and font of the text:

```
par.strip.text=list(cex=.75)
```

By default, the text size varies with the layout; the more rows or columns on the display, the smaller the default strip label text size. If you want more control, you may want to construct your own strip label function or at least change the style given in the default strip function. It helps to know that `strip.default` takes two arguments you may want to change: `strip.names` and `style`. The `strip.names` argument to `strip.default` is a logical vector of length 2, for factors and shingles, respectively, that controls whether or not the variable names are included on strip labels. By default, names are present for shingles but not for factors. Similarly, there is a `style` argument to `strip.default`. So, for example,

```
strip=function(...)strip.default(..., strip.names=TRUE)
```

would put the variable name on both factors and shingles in the strip labels, and

```
strip=function(...)strip.default(..., style=4)
```

would use style 4 for strip labels (style 4 is described in the online documentation for `strip.default`). In some circumstances you may want to suppress the strip labels altogether. In this case, you can use `strip=FALSE`.

Trellis graphics provides a `key` function that allows great flexibility in labeling plots. The `key` argument to Trellis functions is a list that is passed down, almost unchanged, to the key function. Why, then, is it an argument to Trellis functions? Why not just call `key` directly? The reason is that the `key` argument is processed by Trellis functions so that they can leave sufficient space around the plots to hold both key and titles and they can also produce a legend on each page of a multi-page display.

One of the most common uses for a `key` is when you are using `panel.superpose` to distinguish between various groups of points on scatterplot panels. For example we can display a scatterplot matrix of the Anderson Iris data, using colors to encode the three different species.

```
new.iris <- iris.df[,1:4]
for(i in 1:4)
  new.iris[,i] <- jitter(new.iris[,i])
iris.variety <- iris.df[,6]
superpose.symbol <- trellis.par.get("superpose.symbol")
splom(~new.iris,
      panel = panel.superpose,
      groups = iris.variety,
      key = list(space="top", columns = 3,
                transparent = TRUE, text = list(levels(iris.variety)),
                points = Rows(superpose.symbol,1:3)))
```

The result is shown in Figure 20. The position of the key is controlled here by the `space=` component of the `key=` argument.

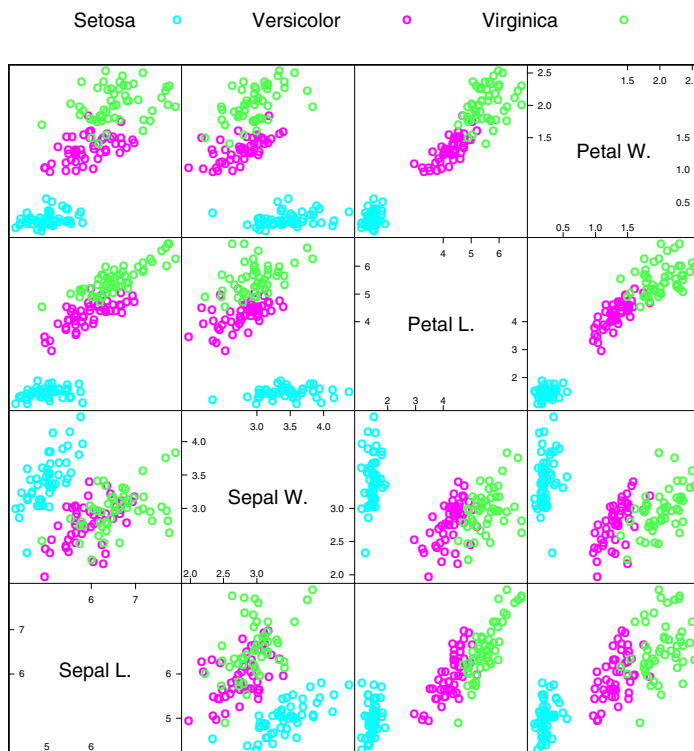


Figure 20. Scatterplot matrix of the jittered iris data with symbols coding the three species. Output from older versions of the Trellis library may not show the cute axes in the diagonal panels.

Precise control of key positioning could also have been done by using the `x=`, `y=`, and `center=` components to the `key=` argument. The coordinate system for `x` and `y` is a unit square surrounding the entire set of panels. If `x=` and `y=` are specified and `space=` is not specified, no additional space is left for the legend. This may be useful if the legend actually is superimposed on the panels (presumably in some unused space.)

Another thing to notice about the example is the use of the `superpose.symbol` customization list. The `panel.superpose` function uses `superpose.symbol` to encode the groups, so it is important to use the same parameters to build the key. Function `trellis.par.get` returns a list giving graphical parameters used to plot superposed symbols (see the online documentation); `Rows` chops it down to the first three values for each component of the list; and finally, those parameters are used by `key` to encode a column of points symbols.

The `panel.superpose` function can help us improve the Barley data plot of Figure 7. If we show the 1931 and 1932 values superimposed on a single panel, it will make the comparisons easier. We can use:

```
superpose.symbol <- trellis.par.get("superpose.symbol")
dot.line <- trellis.par.get("dot.line")

dotplot(variety ~ yield | site, # Figure 21
  data = barley,
  groups = year,
  panel = function(x, y, subscripts, ...){
    abline(h = y, lwd = dot.line$lwd, lty = dot.line$lty,
      col = dot.line$col)
    panel.superpose(x, y, subscripts, ...)
```

```

    },
    aspect = 0.5, layout = c(1,6),
    xlab = "Barley Yield (bushels/acre)",
    key = list(points = Rows(superpose.symbol, 1:2),
               text = list(levels(barley$year)),
               space = "Right", columns = 2))

```

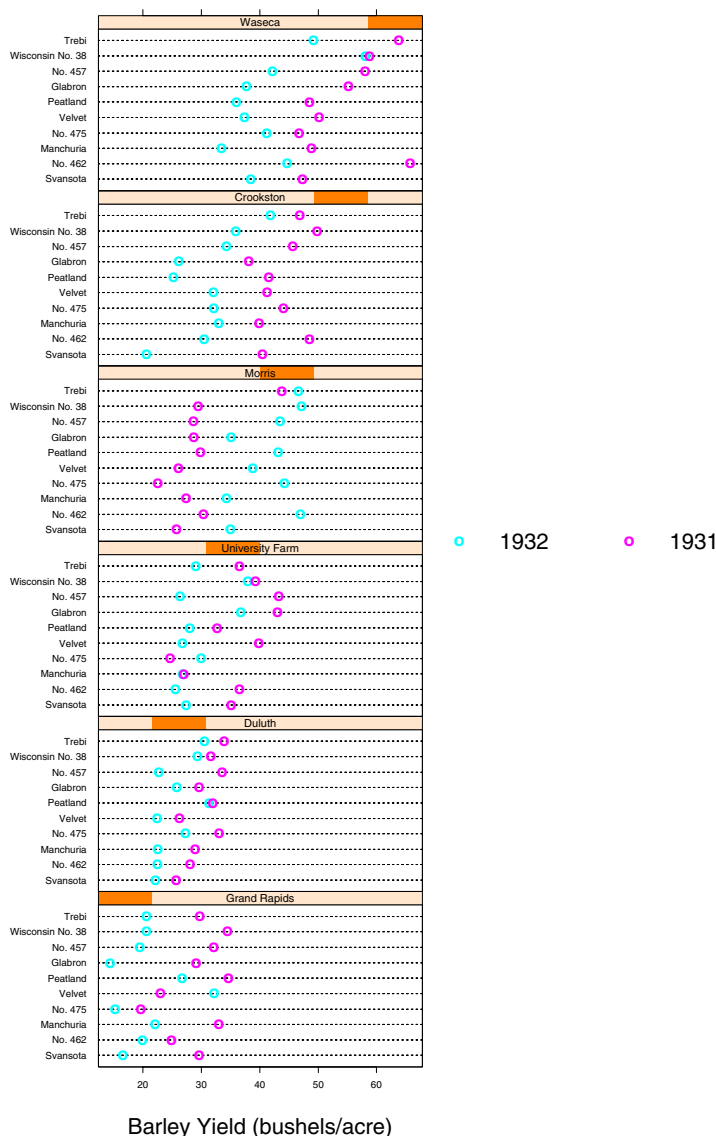


Figure 21. Barley Data: comparison of 1931 and 1932 on a single panel.

This figure makes the anomaly in Morris stand out boldly — it is the only panel where the 1932 symbols are furthest to the right. Notice how the panel function draws the horizontal dotted lines and then uses `panel.superpose` to draw the symbols.

It is important that you realize that the row, column, and strip labels are ordered according to how the data was initially read. For example, the data frame `barley` that we used earlier was created by reading a data file and making an intermediate data frame named `barley2`. The data file `barley.data`, looked like this:


```
yield      variety      year      site
27.00000   Manchuria    1931     University Farm
48.86667   Manchuria    1931     Waseca
27.43334   Manchuria    1931     Morris
39.93333   Manchuria    1931     Crookston
32.96667   Manchuria    1931     Grand Rapids
28.96667   Manchuria    1931     Duluth
43.06666   Glabron     1931     University Farm
. . .
```

The columns were separated by tab characters (important because sites like “University Farm” have embedded blanks), so we could, for example, use `read.table` to create the `barley2` data frame.

```
barley2 <- read.table("barley.data", sep="\t", header=TRUE)
```

When such a table is made, the default is to alphabetize the various factors. So, the expression:

```
dotplot(variety ~ yield | year * site, # Figure 22
        data = barley2,
        xlab="Barley Yield (bushels/acre)")
```

produces the difficult-to-understand Figure 22. The problem is that this ordering obscures the underlying pattern; the main-effects ordering that makes the Morris year anomaly stand out in Figure 7 is not present here. Notice, too, that `read.table` has produced a numeric vector for year and hence it is plotted as a shingle, rather than a factor.

In order to produce Figure 7, we changed the factors:

```
barley <- barley2
barley$variety <- reorder.factor(barley2$variety, barley2$yield, median)
barley$year <- ordered(barley$year, c(1932,1931)) # make it a factor
barley$site <- reorder.factor(barley2$site, barley2$yield, median)
```

For example, the levels of the `variety` factor were rearranged by `reorder.factor` by increasing order of barley yield.

3. ADVANCED CONCEPTS

3.1 Prepanel Functions

One of the strengths of a Trellis display is the independence between a) the common code that sets up the coordinate systems, labels, aspect ratios, and b) the panel function that takes care of the drawing for each individual panel. Generally, this decoupling allows you to customize the drawing in the panel without having to work with or understand the common code. Unfortunately, this is not always the case. The only thing that the common code can do in order to figure out how to set up coordinate systems is to look at the variables in the formula to determine what is going to be plotted on the horizontal and vertical axes.

Suppose, though, that your panel function draws more on the panel than just the x- and y-data. For example, suppose it puts on a fitted line or curve. How, then, can the Trellis software allocate enough space on the panel for the extra things plotted by the panel function, when the common code has no idea what extra plotting is going to take place? Another problem involves aspect ratio calculations. The Trellis code can bank to 45° based upon the x- and y-data, but suppose that the important thing for the banking computations is a fitted line, not the raw data. How can that be taken into account so that an appropriate aspect ratio can be computed? Remember, by the time the panel function is called, the coordinate system and aspect ratio are already locked in.

These problems are solved by the argument `prepanel`, which provides a *prepanel function*. The job of the `prepanel` function is to take the x- and y-data for a panel and return a list containing

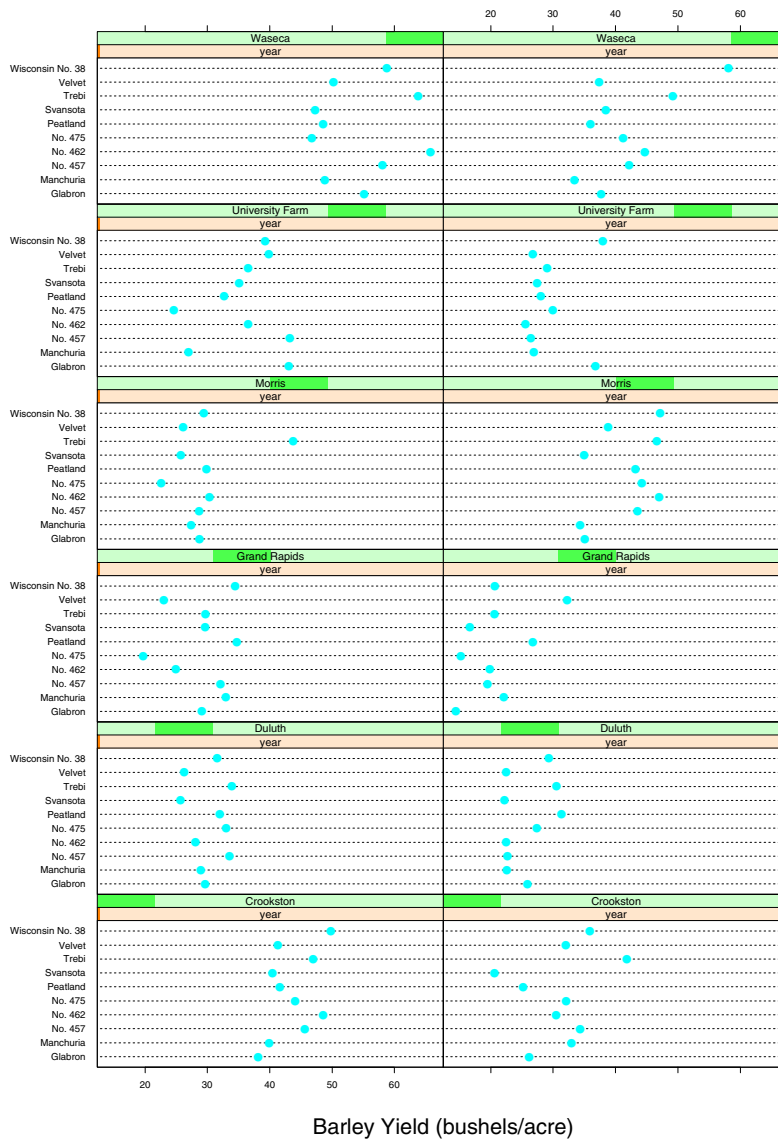


Figure 22.
The barley data, with factors in alphabetical order. Compare to Figure 7.

one or more of the following components: `xlim`, `ylim`, `dx`, and `dy`. The `xlim` and `ylim` values are vectors that give the minimum and maximum value on each axis. The `dx` and `dy` vectors describe (the run and rise of) line segments that should be banked to 45°.

Certain types of Trellis displays occur often enough that prepanel functions are already written for them. They are `prepanel.lmline`, `prepanel.qqmathline`, and `prepanel.loess`. If you want a least-squares line fit via the `lm` function to the x-y data in each panel, `prepanel.lmline` will carry out the computations so that the panels will have enough room to show the fitted line over the range of the x-data and so that the lines on the panels will be banked as closely as possible to 45°. The `prepanel.qqmathline` function does a similar thing, but the line is fit to a quantile plot and goes through the 25th and 75th percentiles. It is used in conjunction with `qqmath` and the `panel.qqmathline` functions.

Perhaps the most commonly used prepanel function is `prepanel.loess`. It does a loess smoothing on the data in each panel and banks the segments of the smooths to 45°. Let's give an example of this, applied to the engine data.

```
EE <- equal.count(ethanol$E, number = 9, overlap = 0.25) # Figure 23
xyplot(NOx ~ C | EE,
  data = ethanol,
  prepanel = function(x, y) prepanel.loess(x, y, span = 1),
  panel = function(x, y) {
    panel.grid(h = 2)
    panel.xyplot(x, y)
    panel.loess(x, y, span = 1)
  },
  aspect = 2.5,
  xlab = "Compression Ratio", ylab = "NOx (micrograms/J)")
```

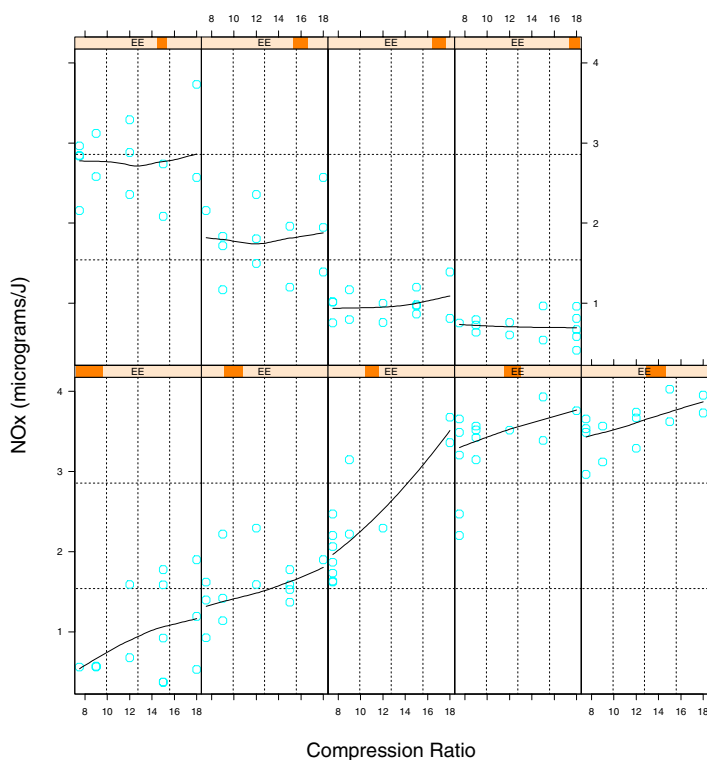


Figure 23. Engine data showing how NOx emissions depend on compression ratio for various intervals of equivalence ratio.

Studying Figure 23 and the `xyplot` expression that produced it gives a concrete example of many of the issues that we have been discussing. For example, notice how the `span` argument to `loess` is passed to both `panel.loess` and `prepanel.loess`.

As we mentioned earlier, the formulas given to Trellis functions are allowed to contain expressions. That means we could have made the call to `equal.count` as part of the formula. However, the problem with this is that there would then be no nice character string to use for the strip labels and we have found that names on strip labels are particularly important for shingles. Otherwise, the information about what variable is represented by the shingle needs to be placed in the caption or somewhere else in the figure.

As you can see, calls to Trellis functions may get a bit large, even though no particular part

is all that difficult. The way we structured the call, with a separate argument on each line and multi-line, indented panel functions, is designed to make it easier to read and understand. You may find this layout technique useful, too.

When we first produced Figure 23, we did not include the `aspect=2.5` argument. Because we used a `prepanel` function that returned `dx` and `dy` values, the banking computation was done automatically. Unfortunately, the resulting aspect ratio was far too big. To see how it looked, try the expression without using the `aspect=` argument. The problem is that many of the panels have basically horizontal fits, and only an extreme aspect ratio will make these panel fits non horizontal. This shows the occasional need for user adjustments to the automatically chosen values.

3.2 The `subscripts=` Argument

Suppose that we would like to produce the plot of Figure 3, but instead of plotting points, we would like to plot the observation number for each point. Since a panel function only gets the `x`- and `y`-values that belong on a panel, how are we to accomplish this? We solve this problem in a very general way by allowing the panel function to request one extra argument, named `subscripts`. The `subscripts` argument is a numeric vector that tells which observation in the original data is associated with the `x`- and `y`-values. Thus, we can neatly solve our problem by using

```
xyplot(NOx ~ E | C,  
       data = ethanol, # Figure 24  
       panel = function(x,y,subscripts)  
         text(x,y,subscripts)  
       )
```

This works because `xyplot` sees that the panel function expects an argument named `subscripts` and arranges to pass the appropriate information to the panel function.

We claimed that the `subscripts` argument was a very general solution to this and similar problems. Why is that so? Basically, once a panel function knows which of the original observations correspond to the `x`- and `y`-values, that information can be used to deal with any other data that is parallel to the original data. In particular, if `x` and `y` come from a data frame, any other variables in the data frame can be subscripted by the `subscripts` vector. To facilitate this even more, another argument, `groups`, can be given to all high-level Trellis functions. This argument will be passed down to the panel function, along with the `subscripts` argument. Inside the panel function, `groups[subscripts]` is a vector parallel to `x` and `y` that can be used for attribute colors, identifiers, etc. to the observations. The panel function `panel.superpose`, used earlier in several examples (Figures 18 and 20), uses exactly this mechanism to do its work.

When a high-level Trellis function can see that its panel function has a `subscripts` argument, it automatically arranges to call the panel function with arguments named `x`, `y`, and `subscripts`. The `subscripts` mechanism interacts properly with the `subset` argument: `subscripts` are done relative to the original data.

3.3 Device Settings

Before we get deeply into the concepts of device customization, a few words of reassurance. Don't worry — you normally do not need to write code like this yourself. Trellis graphics makes use of customization to produce excellent plots on all devices without requiring any changes on the part of the user to make it happen. If your device has color capabilities, Trellis functions will use them; they may also use various fonts, character sizes, line styles, area fills, etc. However, when you write your own panel functions, you may know for certain that you are producing the plot for a particular device and you know just what values the parameters should have. If so, just write the panel function with exactly the parameters you want.

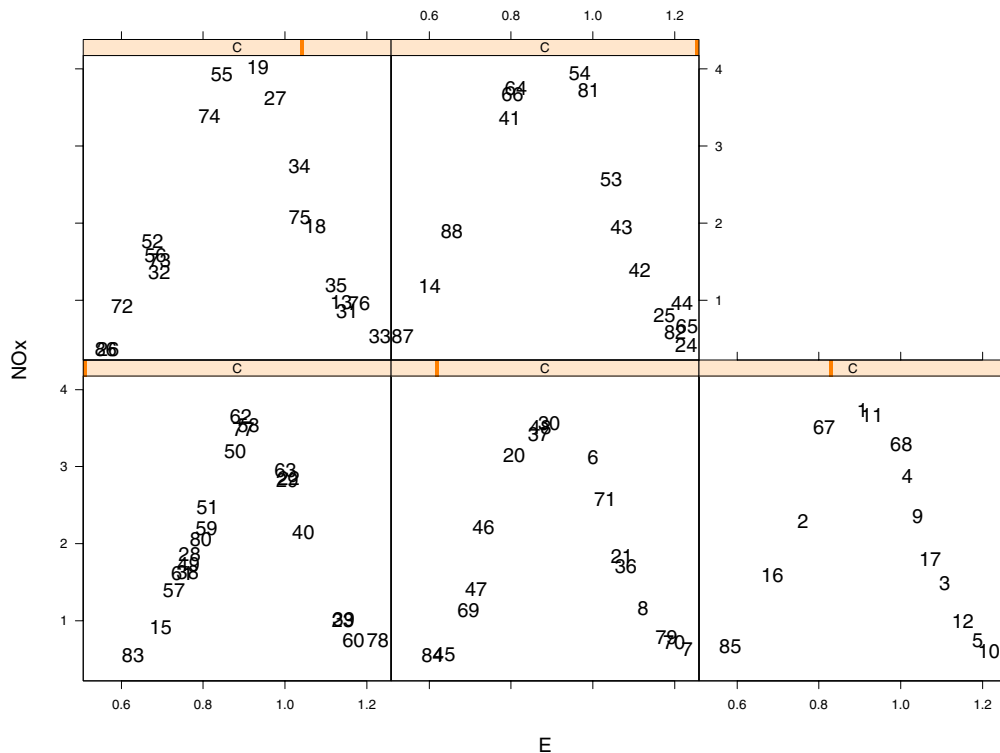


Figure 24. The ethanol data with each observation identified by observation number.

On the other hand, you may want to write your Trellis plot calls that adapt to device characteristics, too. In this case, an explanation of how Trellis functions deal with graphical parameters may help you to understand how device-specific customization works.

Basically, all Trellis functions are careful to specify graphical parameters in a symbolic way, by referring to objects that give the actual values of the parameters. For example, let's look at (a simplified version of) the most commonly used panel function, `panel.xyplot`:

```

panel.xyplot <- function(x, y, ...){
  plot.symbol <- trellis.par.get("plot.symbol")
  points(x, y,
         cex = plot.symbol$cex, pch = plot.symbol$pch,
         font = plot.symbol$font, col = plot.symbol$col, ...)
}

```

The object `plot.symbol` is a list with components `pch`, `cex`, `font`, and `col` to specify the plotting symbol, its size, font, and color. When the S device is set up by an explicit or implicit call to `trellis.device`, the `.Device` object is given an attribute that stores a set of Trellis parameter lists; `plot.symbol` is one such list. The function `trellis.par.get` gets the named list. As illustrated above, it is often convenient to save the list and then extract components from it in various graphics routines inside the panel function.

To see how customization is set up for on any particular device, start the device and then execute the function `show.settings`. Figure 25 shows an example.

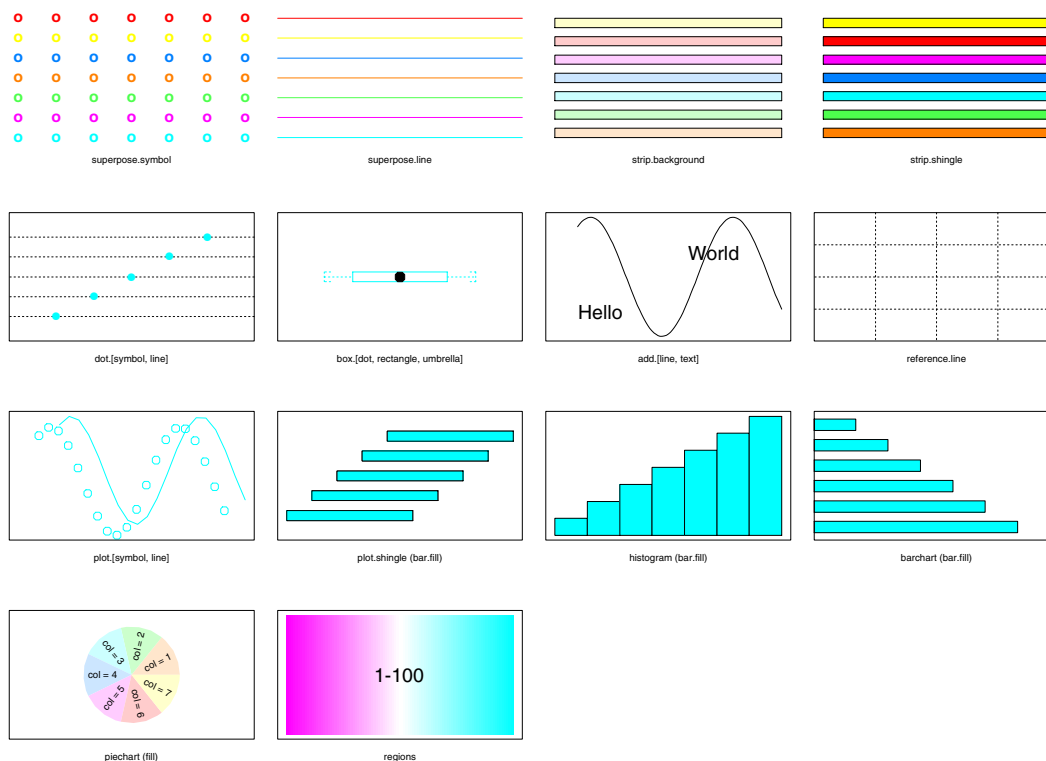


Figure 25. Sample output from the `show.settings` function. This shows the various customization settings done for a PostScript printer.

3.4 Finding the Data

In many of the examples so far, we have used the `data=` argument to present a data frame to be searched for objects named in a Trellis formula. There are a few subtleties of this that are addressed here. (These same issues affect the functions described in the book *Statistical Models in S.*) When the `data=` argument is given, the data frame or list specified by that argument is the first place searched for finding objects named in the formula or in the `subset=` or `groups=` arguments. This search of the `data=` argument *replaces* the normal search of the caller's frame. In any case, if an object is not found in the '`data=`' argument or in the caller's frame, the search will continue through frames 1, 0, and the databases explicitly on the search list.

Why does this matter? It matters when you write functions that call Trellis functions. For example:

```
myfun1 <- function(){
  histogram(~height | voice.part, data=singer)
}
```

would work and

```
myfun2 <- function(){
  histogram(~sqrt(height) | voice.part, data=singer)
}
```

would also work, but

```
myfun3 <- function(){
  sqrtht <- sqrt(singer$height)
  histogram(~sqrtht | voice.part, data=singer)
```

```
}
```

would fail with the error message

```
> myfun3()
Error in myfun3(): Object "sqrtht" not found
```

This is because the presence of the `data=` argument prevents `histogram` from looking into `myfun3`'s frame to find `sqrtht`.

Here are several ways to fix the problem. a) put the transformed object back into the data frame.

```
myfun3a <- function(){
  singer$sqrtht <- sqrt(singer$height)
  histogram(~sqrtht | voice.part, data=singer)
}
```

b) don't use the `data=` argument.

```
myfun3b <- function(){
  sqrtht <- sqrt(singer$height)
  histogram(~sqrtht | singer$voice.part)
}
```

c) explicitly put the data frame on the search list.

```
myfun3c <- function(){
  attach("singer")
  sqrtht <- sqrt(height)
  histogram(~sqrtht | voice.part)
}
```

4. HIGHER DIMENSIONS

This section describes plots for data of three or more dimensions.

4.1 3-D Plotting

Perspective Trellis displays are carried out by the function `cloud`, which produces a 3-dimensional point cloud, and the function `wireframe`, which draws a 3-D wireframe surface. Unfortunately, static cloud displays are typically difficult to understand because even though points in 3-space are displayed in perspective, there is too little structure for the viewer's eye to put together an integrated view when a static panel is drawn.

The going is a bit rough here, so we will give an extended example. The non-Trellis functions mentioned in this section are described in the book *Statistical Models in S*.

Suppose we are interested in learning about the environmental ozone data, where ozone concentrations are to be related to wind speed, temperature and solar radiation. One way to learn about the general pattern is to fit a smooth surface to the data, and we have previously determined that the cube-root of ozone concentration is an appropriate response variable. We can use the `loess` function compute the smooth fit:

```
attach(environmental)

ozo.m <- loess((ozone^(1/3)) ~ wind * temperature * radiation,
  parametric = c("radiation", "wind"), span = 1, degree = 2)
```

Notice how the formula given to `loess` looks just like the formulas given to Trellis functions. To produce a wireframe or surface plot, we need to evaluate the surface at a regular grid of values. We will evaluate the grid for 50 values of wind speed, ranging from the minimum to maximum observed speeds, 50 values of temperature, and 4 levels of radiation. We first form vectors

of the wind speed, temperature, and radiation marginal values, and then make a grid of all 50x50x4 combinations of those.

```
w.marginal <- seq(min(wind), max(wind), length = 50)
t.marginal <- seq(min(temperature), max(temperature), length = 50)
r.marginal <- seq(min(radiation), max(radiation), length = 4)
grid <- expand.grid(wind = w.marginal, temperature = t.marginal,
  radiation = r.marginal)
```

The `grid` object is just what the `predict` function needs to produce the fitted values:

```
fit <- predict(ozo.m, grid)
```

The `fit` object is a vector that matches the vectors in the `grid` data frame: `wind`, `temperature`, and `radiation`. Now, to produce the wireframe plots, we execute:

```
wireframe(fit ~ wind * temperature | radiation, # Figure 26
  data = grid,
  xlab = "Wind Speed (mph)",
  ylab = "Temperature (F)",
  zlab = "Cube Root Ozone \n(cube root ppb)")
```

The `wireframe` function also allows us to specify that *color draping* should be applied to the wireframe; this means that each patch of the wireframe can be drawn in a solid color controlled by the `z` value or by any other value given as the `drape=` argument (which must be the same length as `z`). For more information on color draping, see the online documentation for `wireframe`.

There are a few arguments that are unique to the 3-D functions. These specify parameters for the viewing transformation and provide for axis control. The default viewing perspective was shown in the wireframe example. Suppose, however, that you wish to look at the surface from other vantage points. You can do this with the argument `screen`, a list with named components, each describing a rotation. The name of the element tells which axis the rotation should be around; the value gives the degrees of rotation about the axis. For example, the default for `screen` is

```
screen=list(z = 40, x = -60)
```

which means that we rotate by 40° about the `z`-axis and then by -60° about the `x`-axis. To understand these rotations, we need to understand the 3-D axis orientations: the `x`-axis extends horizontally across the screen, the positive `y`-axis goes back into the screen, and the `z`-axis is vertical.

Another important viewing parameter is named `perspective`. This is a logical flag that tells whether the projection should be orthogonal (`perspective=FALSE`) or perspective. If a perspective projection is desired, the parameter `distance` controls the apparent distance from the eye to the object. A value of 0 implies an infinite viewing distance (an orthogonal view) while 1 has the eye point right up to the data; the default is 0.2.

There are two ways that the axes can be drawn in 3-D Trellis plots: either with 3-D axis lines with ticks and tick labels or by simple arrows that run parallel to the axes and point in the direction of increasing values. Arrows are the default; conventional axes by turning off the arrow axes via the `scale=list(arrows=FALSE)` argument. In either case, the axes or arrows are labeled by variable names, either taken from the formula or passed in through the arguments `xlab`, `ylab`, and `zlab`.

Another quick comment related to 3-D plotting: the notion of a simple panel function breaks down here. Unfortunately, the panel function does projections, axis drawing, etc., and thus it is huge and unwieldy. Don't expect to produce your own 3-D panel function without studying and modifying a copy of an existing one.

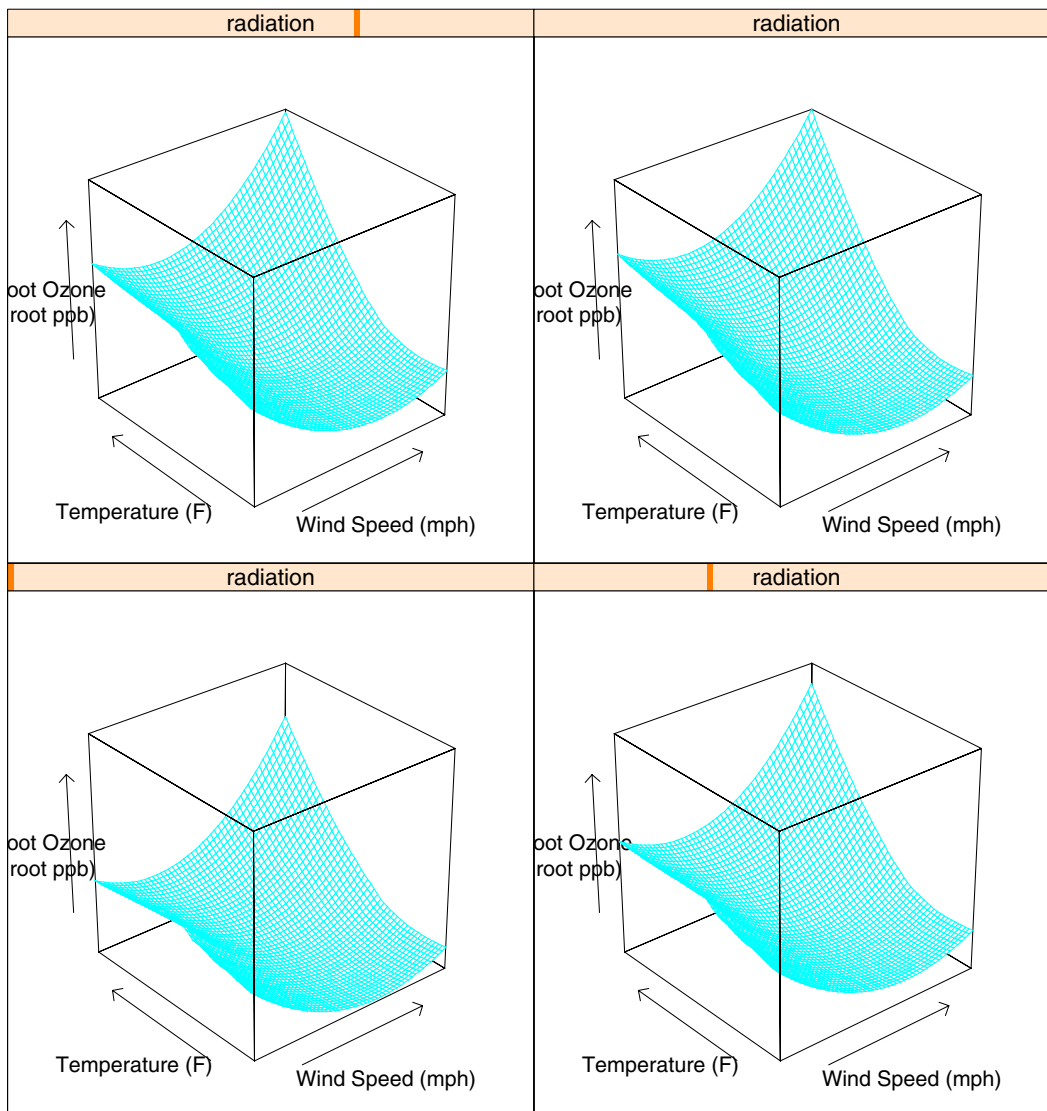


Figure 26. A Trellis wireframe display showing the loess fit of ozone data to wind speed and temperature, for four given levels of solar radiation. Notice the changes in height relative to the surrounding box, especially in the far corner and at intersections with axes.

4.2 Contour Plots

Wireframe and contour displays share many characteristics; the input data is often identical — only the way it is displayed changes. Once we have our data structures straight, a contour plot corresponding to the previous wireframe is simple to produce:

```
contourplot(fit ~ wind * temperature | radiation, # Figure 27
            data = grid,
            xlab = "Wind Speed (mph)",
            ylab = "Temperature (F)",
            main = "Cube Root Ozone (cube root ppb)")
```

If you compare the contour and wireframe figures you can probably see that the contour plot is preferable for quantitative information but the wireframe gives a good gestalt of the surface.

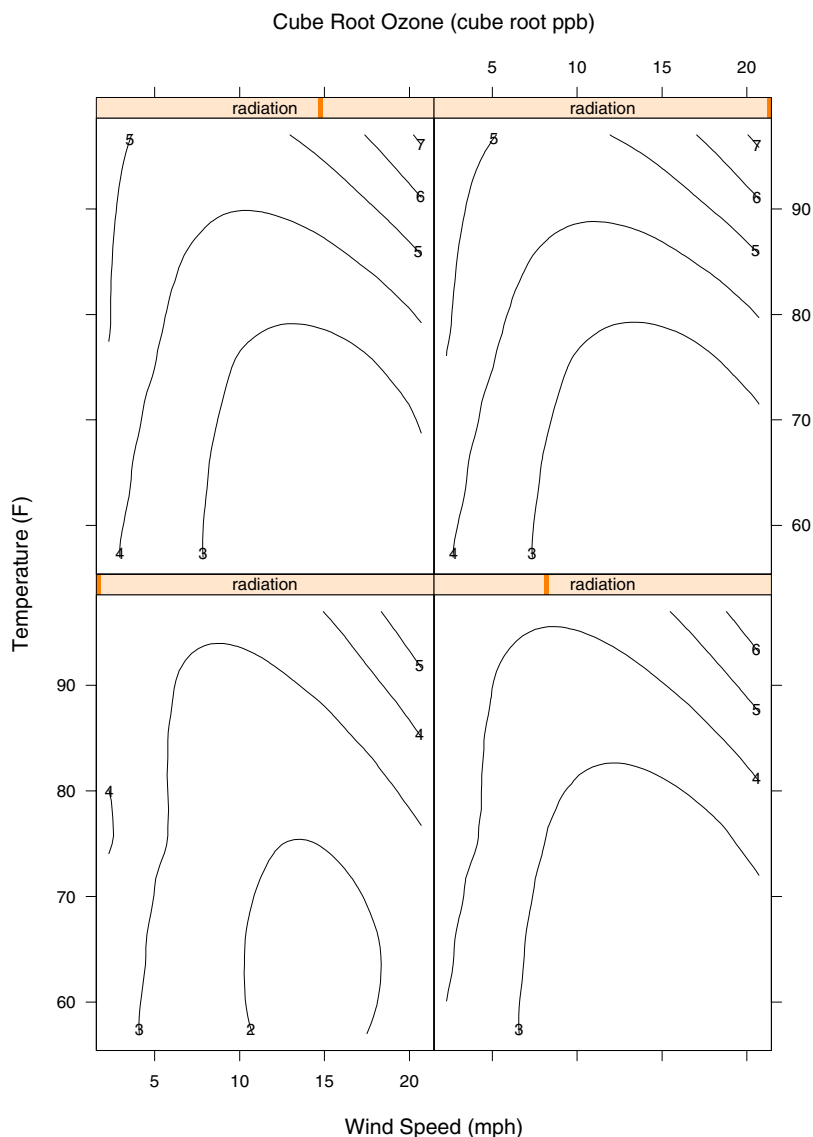


Figure 27. A Trellis contour display showing the loess fit of ozone data to wind speed and temperature, for four given levels of solar radiation. Compare to the wireframe display in Figure 24.

The function `levelplot` produces a color level plot, where color or gray levels are used to encode the value of a third variable. It is closely related to a contour plot — the boundaries between different regions of a levelplot are contours.

4.3 More Than Three Variables

Two functions are designed to work with more than three variables on each panel of a Trellis display: `splom` and `parallel`. You saw the result of `parallel` in Figure 18. The `splom` function produces a plot in which each column in an input matrix, `x`, is plotted against each other column. Figure 20 was an example of this. What might not be obvious is that the formula may also contain given variables,

```
splom( ~x | g1 * g2 * ... )
```

Here, `x` is a matrix, and the given vectors cause the packets to contain sub-matrices consisting of various rows of `x`. For each packet, a scatterplot matrix is plotted. The axes produced by `splom` are cleverly constructed to use minimal space; there is an argument, `pscales`, that can control these axes.

5. A GRAB BAG

This section contains descriptions of several Trellis functions that don't quite fit in with the general formula and data framework that has been introduced earlier. However, don't get the idea that they are not important, just because they are hard to place. Each of these functions does something non-trivial.

The function `rfs` produces a plot that shows the residual and fitted-values spreads (as described in Cleveland, 1993). It is used to assess the magnitude of the fit to the magnitude of the residuals. You initially fit a model, then give the model object to `rfs`. It produces a pair of plots with identical y-axis scaling (in units per inch). By comparing the sizes of the y-variation two plots, you can determine the importance of the fit.

The `tmd` function computes a Tukey mean and difference plot. It is unusual since it takes as an argument the output of one of the other plotting functions and it produces from that another similar object (of class `trellis`). However, for each plot it transforms the data so that the x-axis holds the mean of the original x- and y-variables and the y-axis gives the difference. The purpose of this is to allow more effective comparisons. Instead of comparing how well data fits the $y=x$ line, you can look at how well the `tmd`-modified data follows a horizontal line.

Finally, the function `oneway` is provided to do a one way analysis of variance. Suppose we believe that the heights of singers can be modeled by a typical value for each voice part with variation about that typical value. We can fit this model with `oneway` and assess the quality of the fit with `rfs`:

```
attach(singer)          # Figure 28
singer.model <- oneway(height ~ voice.part, spread = 1)
rfs(singer.model, aspect = 1, ylab = "Height (inches)")
```

The results are shown in Figure 28.

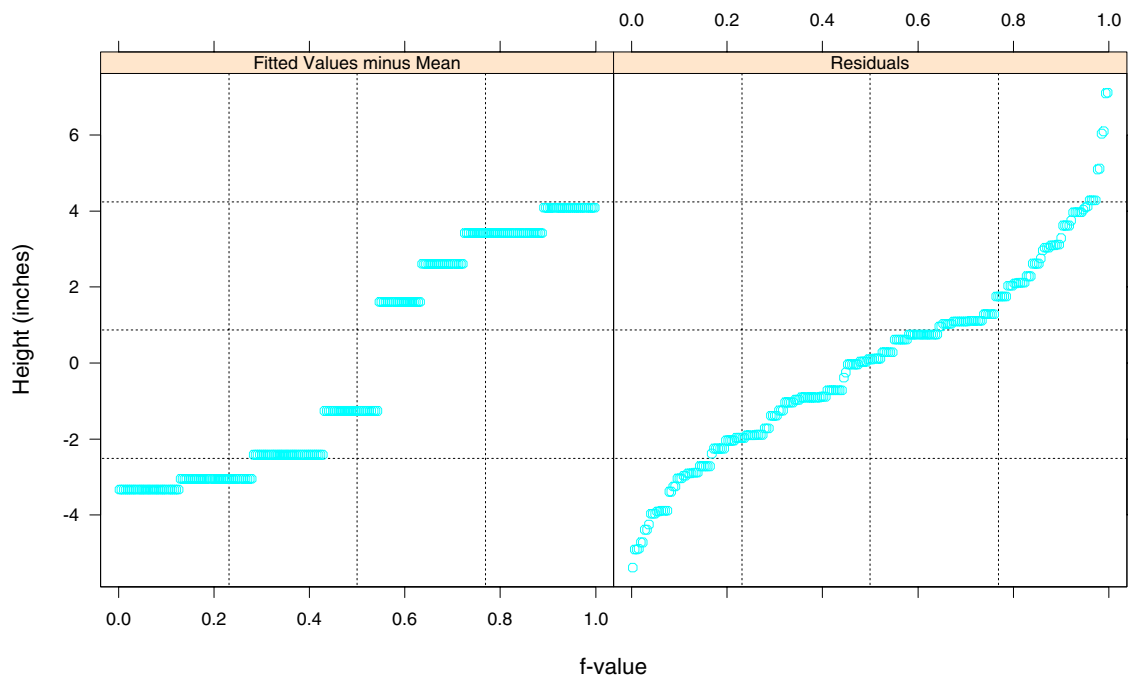


Figure 28. Residuals and fitted values from a one-way fit to the singer data.

REFERENCES

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