



2. Graphics

Ken Rice

Thomas Lumley

Universities of Washington and Auckland

Seattle, July 2012

Important pre-takeoff announcement:

We are assuming you know;

- ... that graphics are useful! (and may be worth ≤ 1000 words)
- How to make some simple plots e.g. making a scatterplot with `plot()`, adding to existing plots using `points()`, `lines()`, `text()`, and `legend()`
- That these functions can take many three letter arguments; `lwd`, `lty`, `pch` and many others, which can be looked up via `?par`
- That, ultimately, we want PDFs, JPEGs and other output formats – not just a window in an R session

Plotting large & high-dimensional data

'Simple' plots involve two-dimensional data, which we measure on the x and y axes.

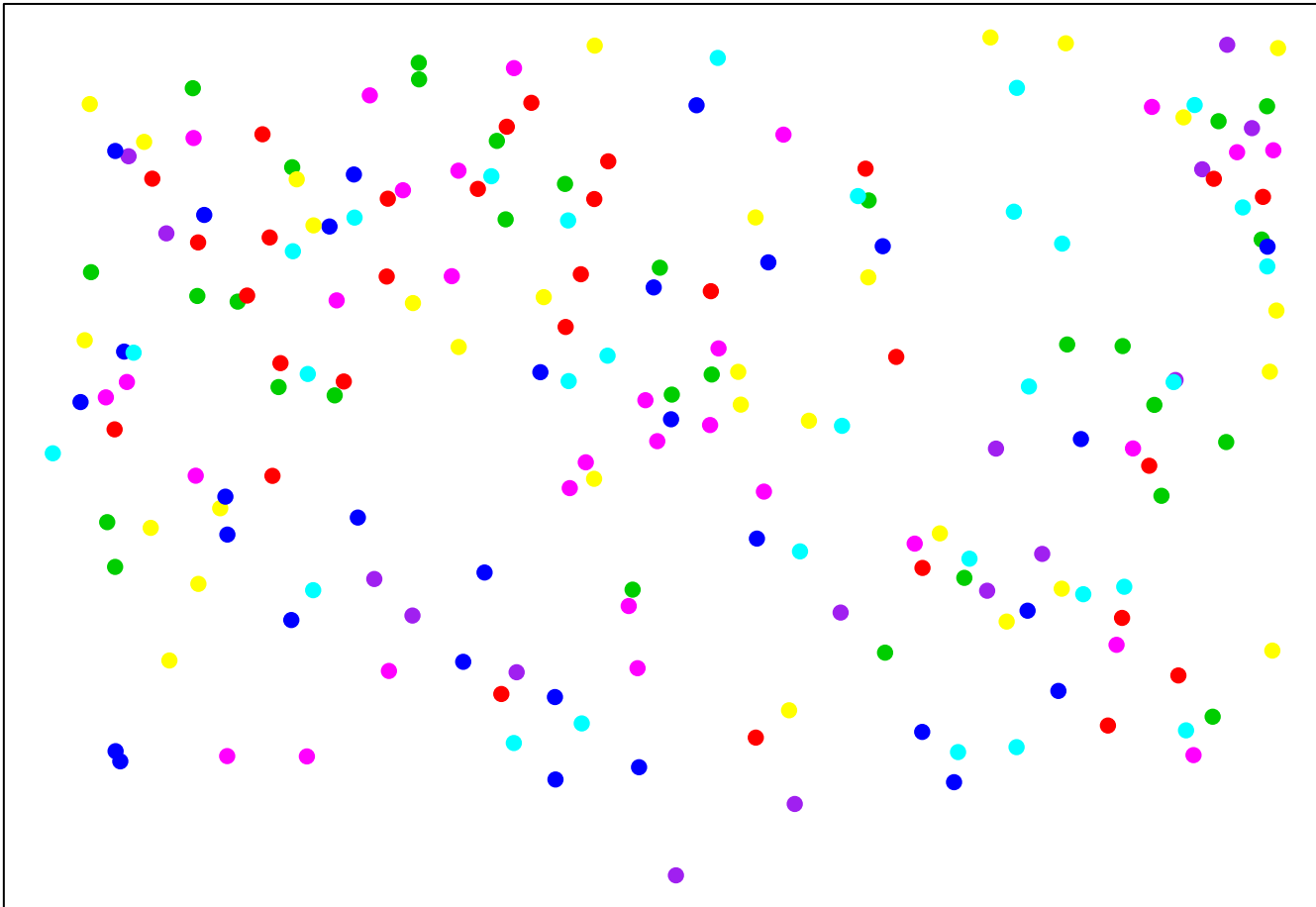
For higher-dimensions, some traditional approaches are;

- Different colors for e.g. men, women (`col`)
- Different-shaped symbols (`pch`), or different sizes (`cex`)

For ≤ 100 's of data points, **modest** use of these is fine. But your eye is not good at concentrating e.g. just on the purple points, in a fully Technicolor plot;

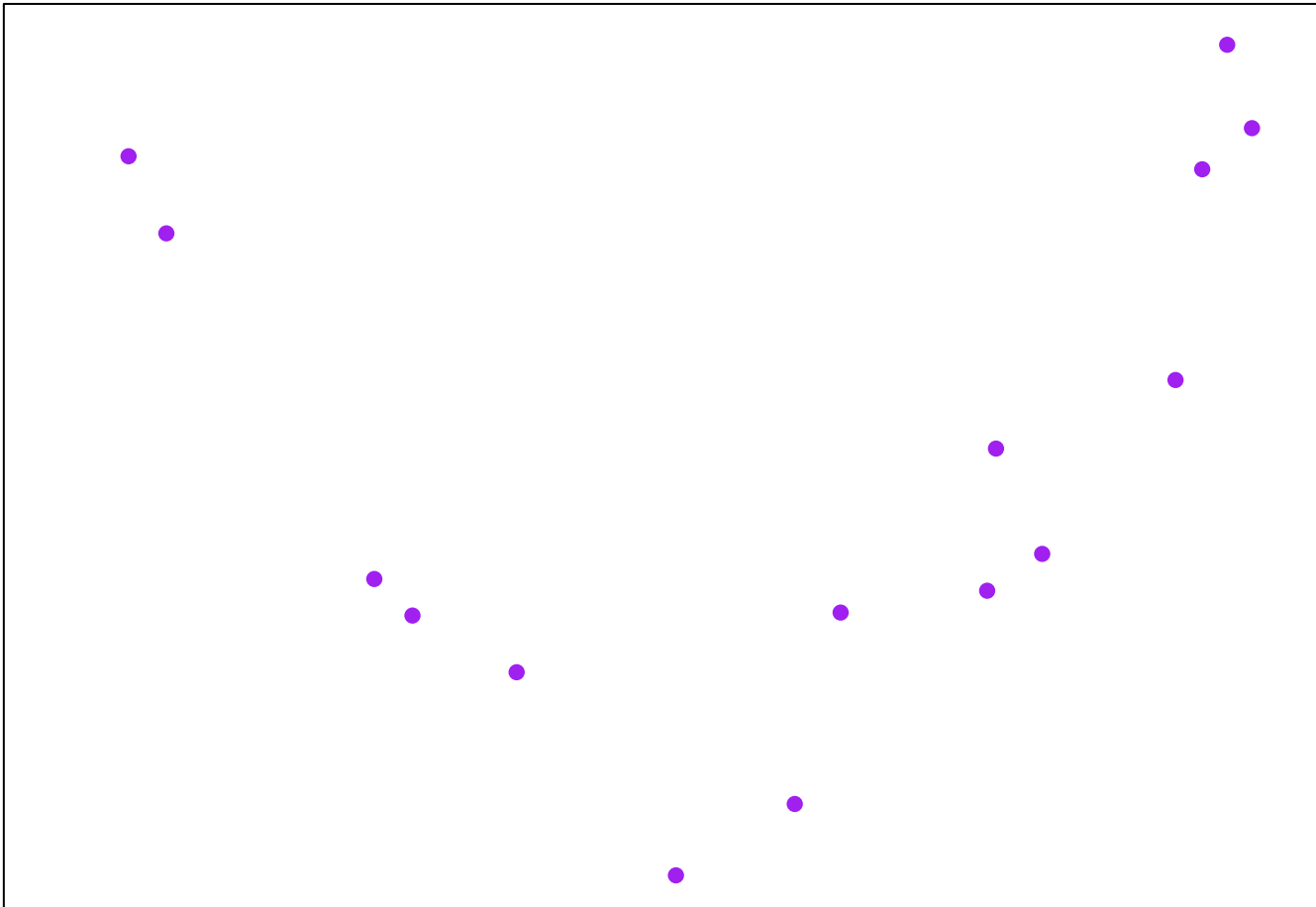
Plotting large & high-dimensional data

Some of these points are not like the others...



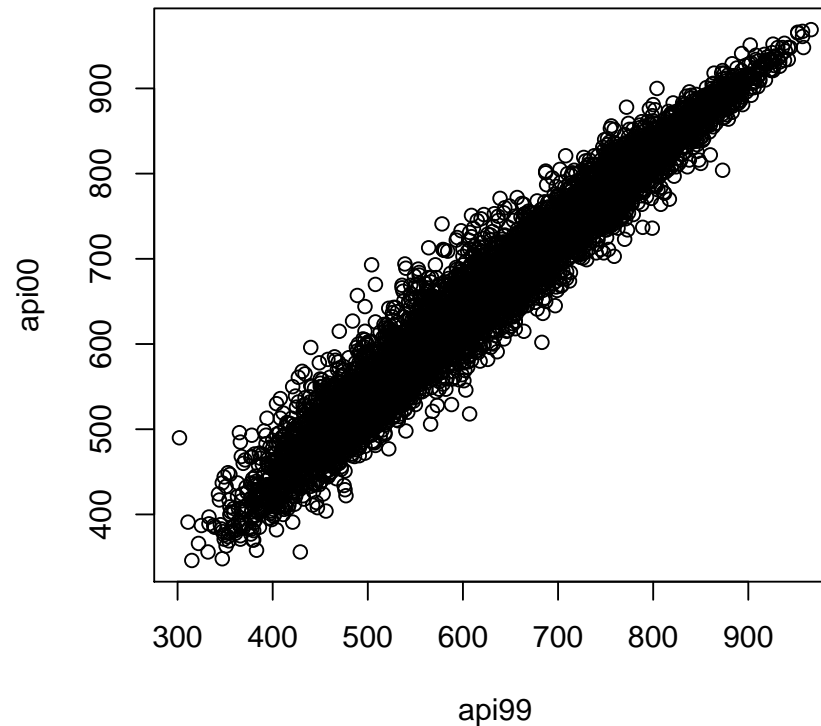
Plotting large & high-dimensional data

Some of these points are not like the others...



Plotting large & high-dimensional data

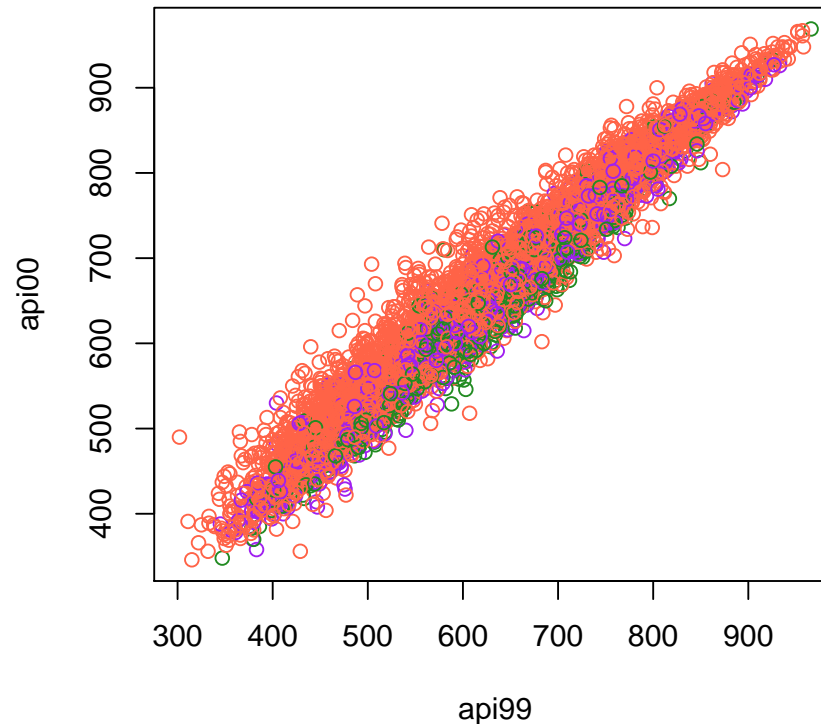
For large(ish) data, 'overlap' is a fundamental problem...



(California Academic Performance Index on 6194 schools)

Plotting large & high-dimensional data

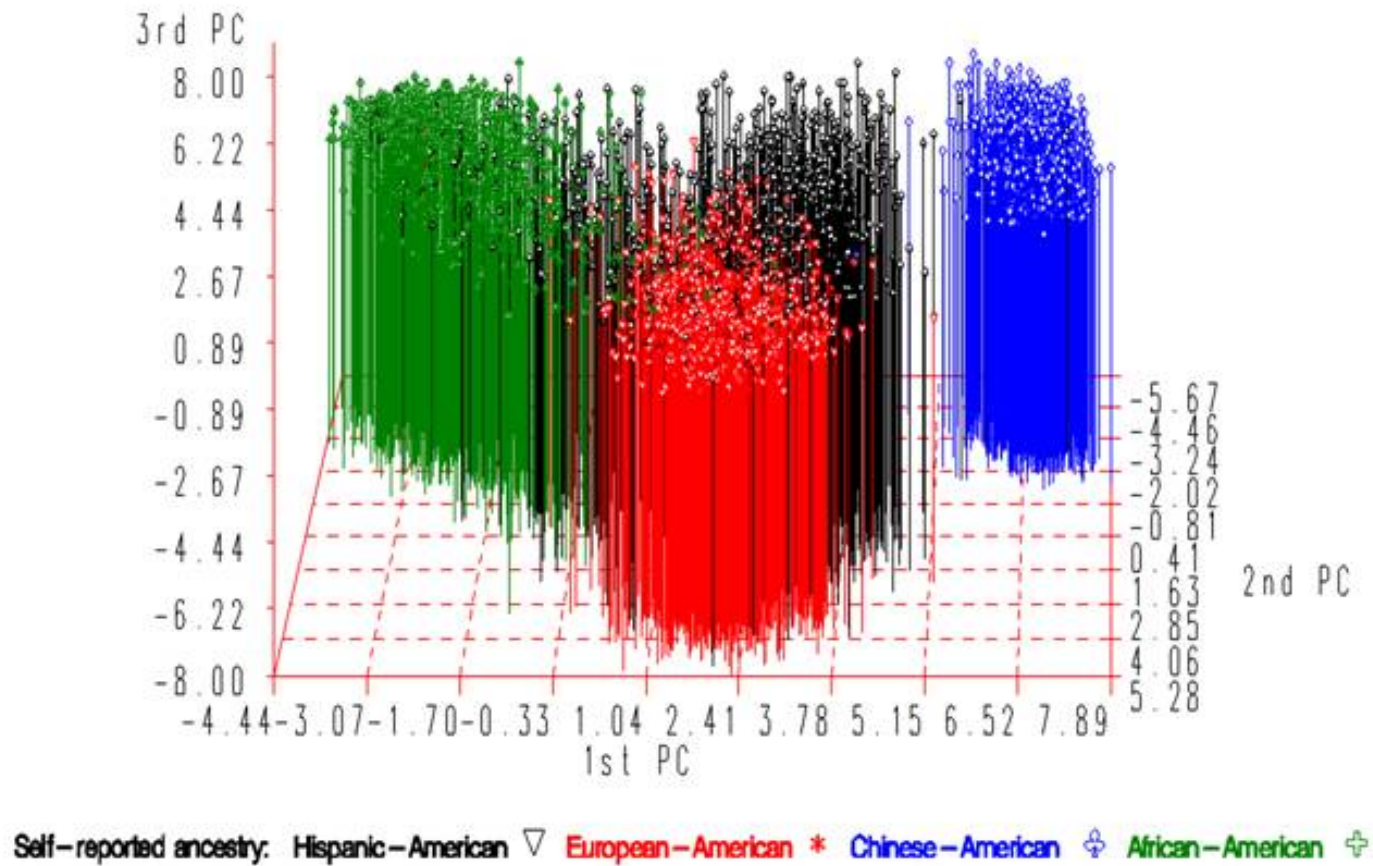
... which remains, when we color-code.



Colors denote Elementary, Middle & High Schools

Plotting large & high-dimensional data

With three dimensions + color-codes, this can happen;



(R does have `persp()`, for occasional use)

Conditioning plots

A typical goal for measuring Z is to see whether the $Y - X$ relationship changes at different values of Z . For example, we might want to see if a Blood Pressure/genotype association varies by Body Mass Index (weight/height²)

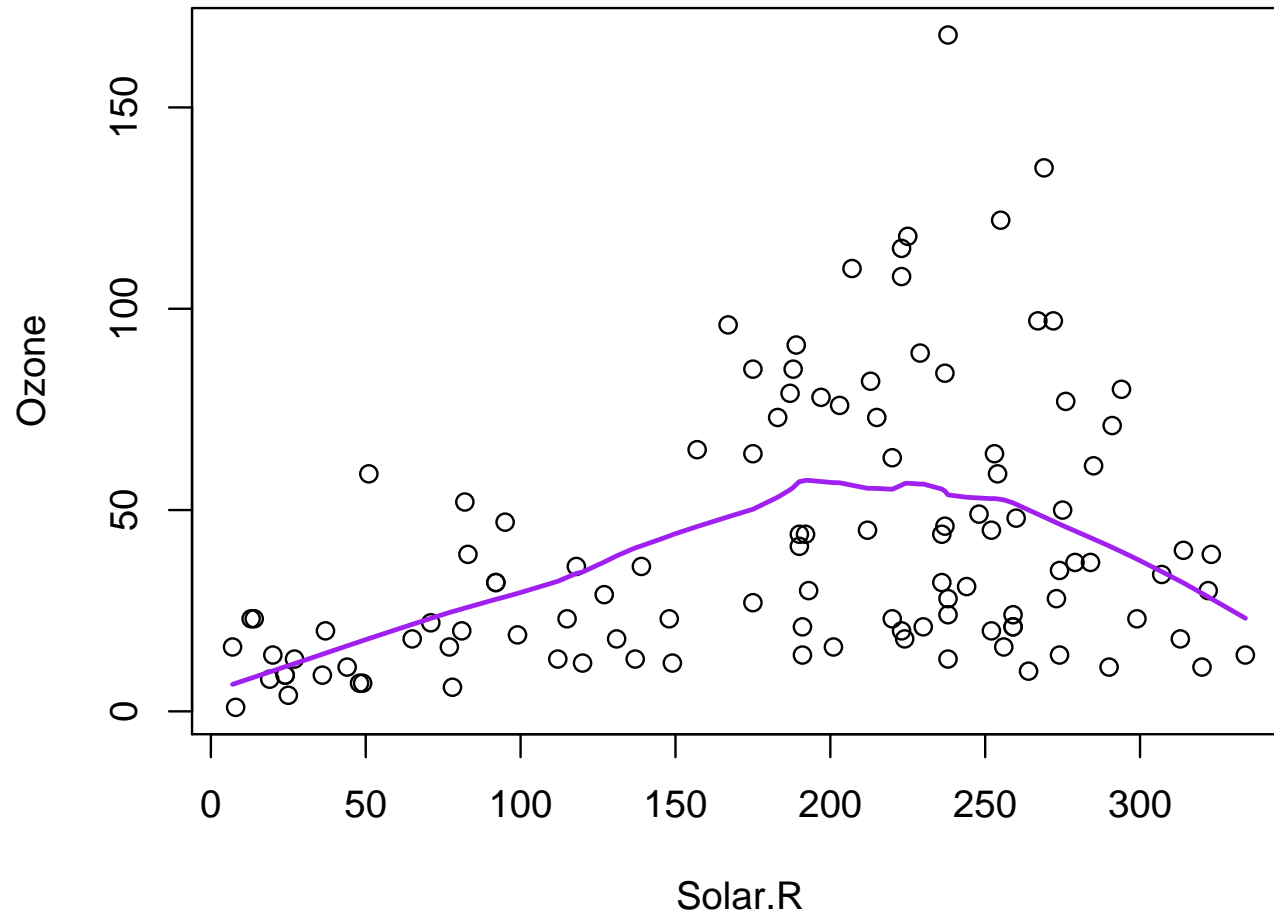
In this case, it's useful to show plots of Y against X conditioned on the value of Z , i.e. Y versus X for all data with Z in a small range. This is known as a **conditioning plot**, and can be produced with `coplot()`.

Conditioning plots

Ozone is a **secondary pollutant**, it is produced from organic compounds and atmospheric oxygen in reactions catalyzed by nitrogen oxides and powered by sunlight.

However, looking at ozone concentrations in NY in summer (Y) we see a non-monotone relationship with sunlight (X)

Conditioning plots

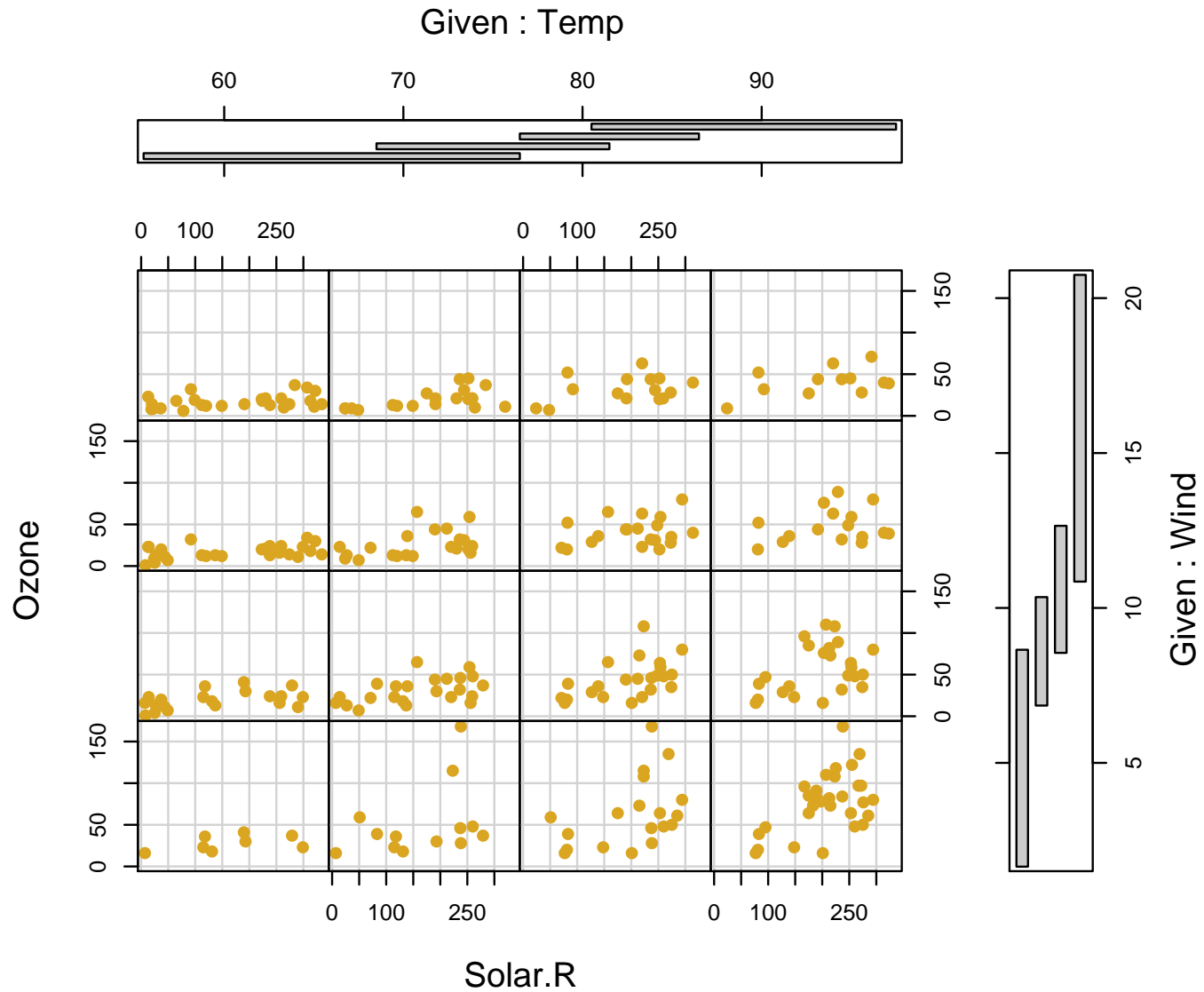


Conditioning plots

Here we draw a scatterplot of `Ozone` vs `Solar.R` for various subranges of `Temp` and `Wind`. For more examples like this, see the commands in the `lattice` package.

```
data(airquality)
coplot(Ozone ~ Solar.R | Temp * Wind, number = c(4, 4),
      data = airquality,
      pch = 21, col = "goldenrod", bg = "goldenrod")
```

Conditioning plots

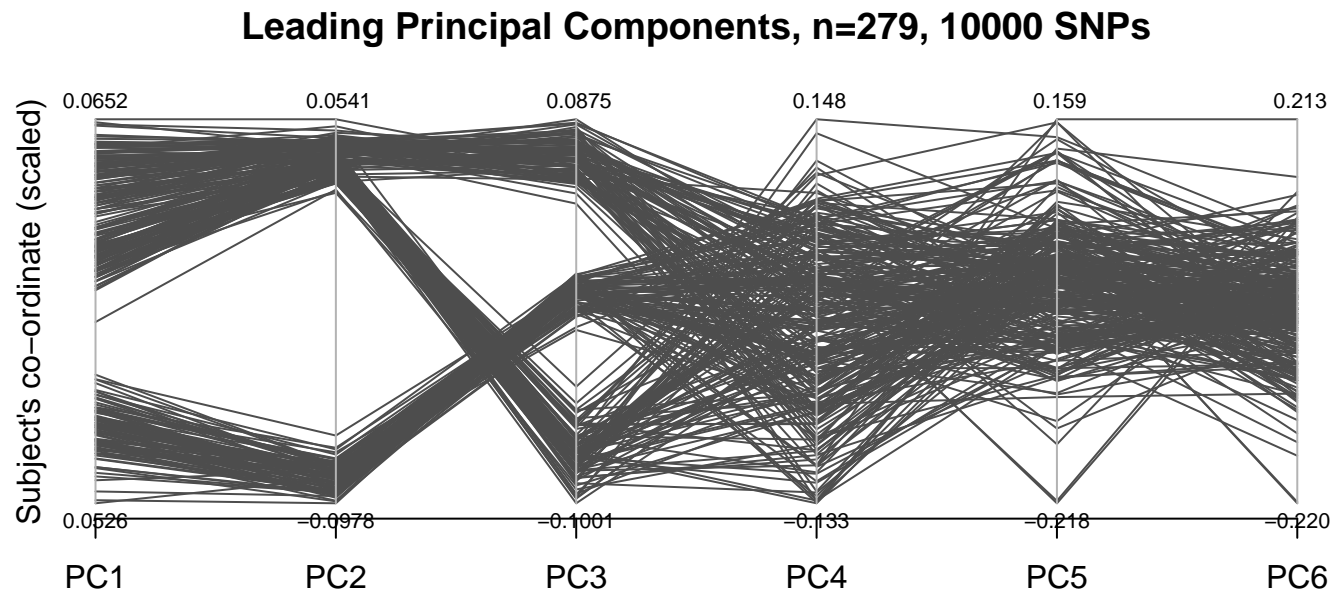


Conditioning plots

- A 4-D relationship is illustrated; the Ozone/sunlight relationship changes in strength depending on both the Temperature and Wind
- The vertical bar | is statistician-speak for 'conditioning on' (nb this is different to use of |'s meaning as Boolean 'OR')
- The horizontal/vertical 'shingles' tell you which data appear in which plot. The overlap can be set to zero, if preferred
- `coplot()`'s default layout is a bit odd; try setting `rows`, `columns` to different values
- For more plotting commands that support conditioning, see `library(help="lattice")`

Parallel Coordinate Plots

For even higher-dimensional data, scatterplots can not provide adequate summaries. For data where the dimensions can be ordered, the **parallel co-ordinates plot** is useful;

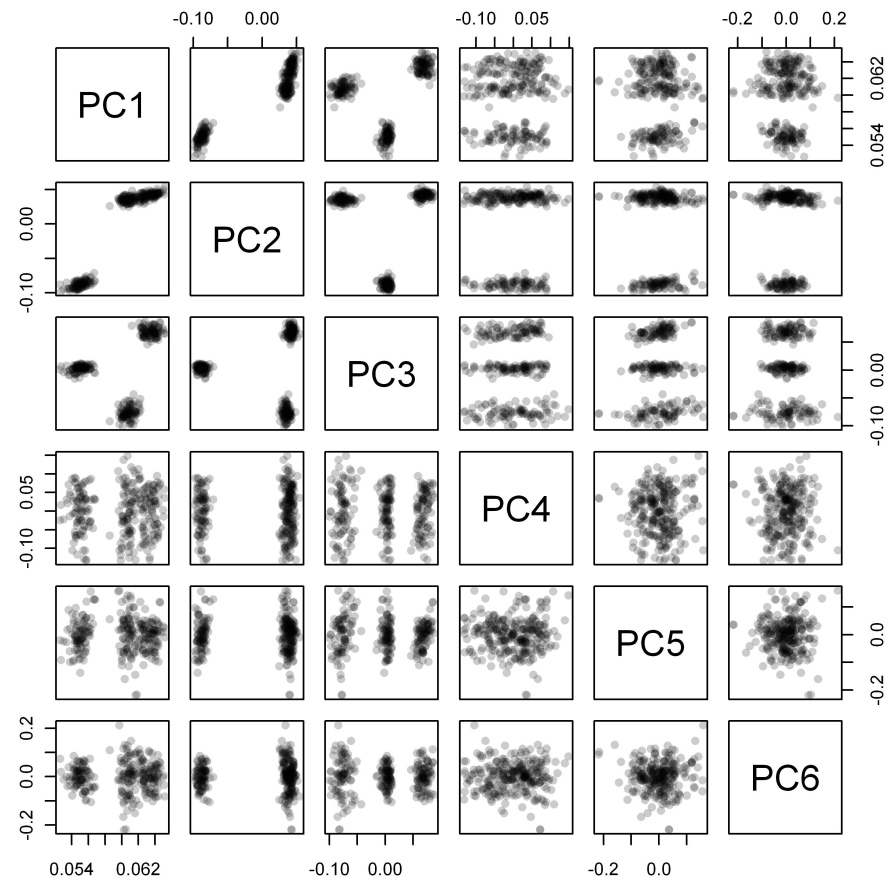


Parallel Coordinate Plots

- Each multi-dimensional data point (i.e. each person) is represented by a line – not a point
- `parcoord()` in the MASS package is one simple implementation – writing your own version is not a big job
- Coloring the lines also helps (example later)
- Scaling of axes, and their vertical positions are arbitrary
- Doing ‘Principal Components Analysis’ is just choosing axes for your data so that their variance is maximized on axis 1, then axis 2, ...

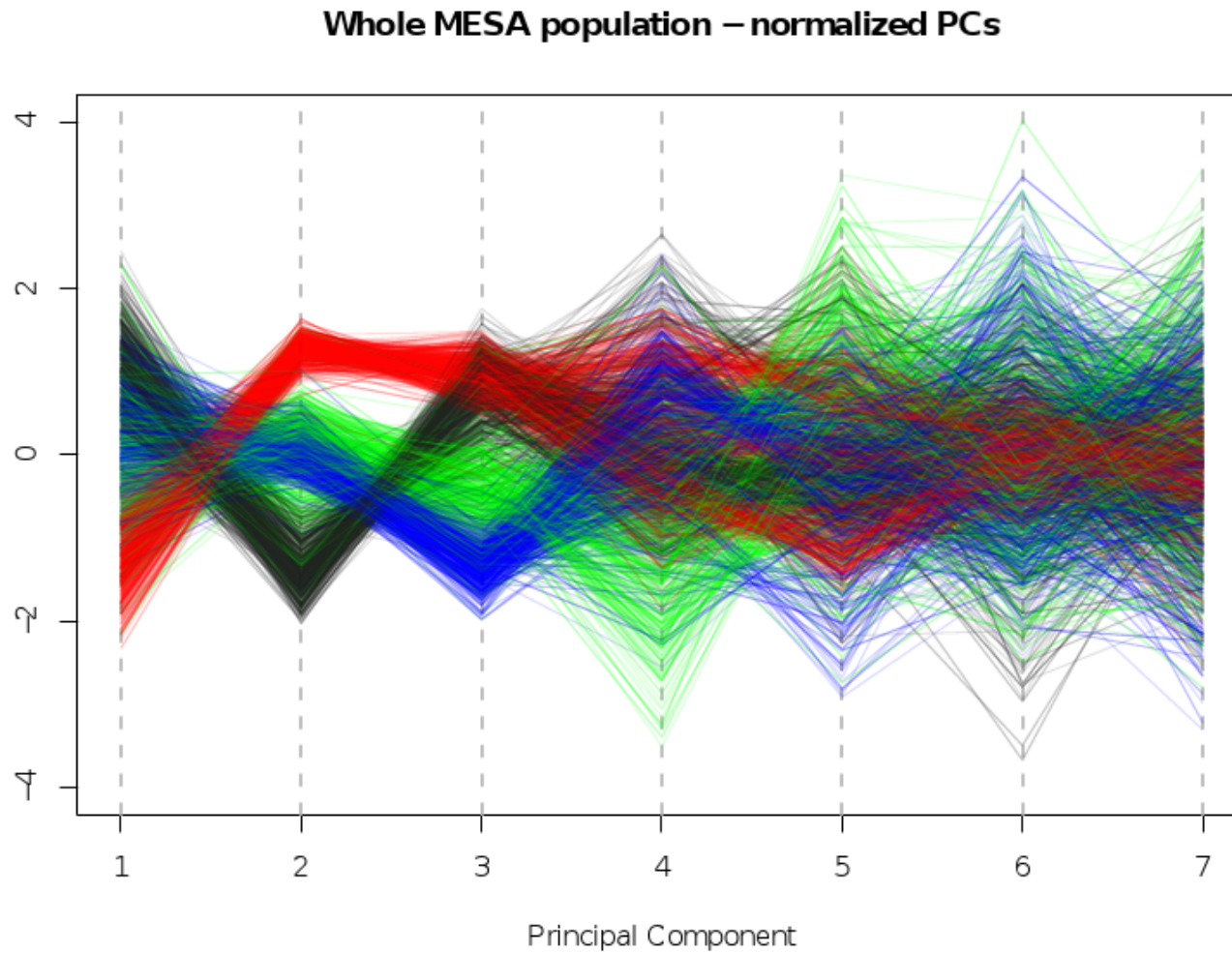
Parallel Coordinate Plots

A pairs() plot of the same thing; (nasty!)



Parallel Coordinate Plots

The pin cushion data++ : colors indicate self-report ancestry



Transparency

The colors in the last examples were **transparent**. As well as specifying e.g. `col=2` or `col="red"`, you can also specify

```
col="#FF000033"
```

– coded as RRGGBB in hexadecimal, with transparency 33 (also hexadecimal). This is a ‘pale’ red – $33/FF \approx 20\%$.

Get from color names to RGB with `col2rgb()`, and from base 10 to base 16 using `format(as.hexmode(11), width=2)`

Transparency

An example; (also shows other graphics commands)

```
curve(0.8*dnorm(x), 0, 6, col="blue", ylab="density", xlab="z")
curve(0.2*dnorm(x,3,2), 0, 6, col="red", add=T)

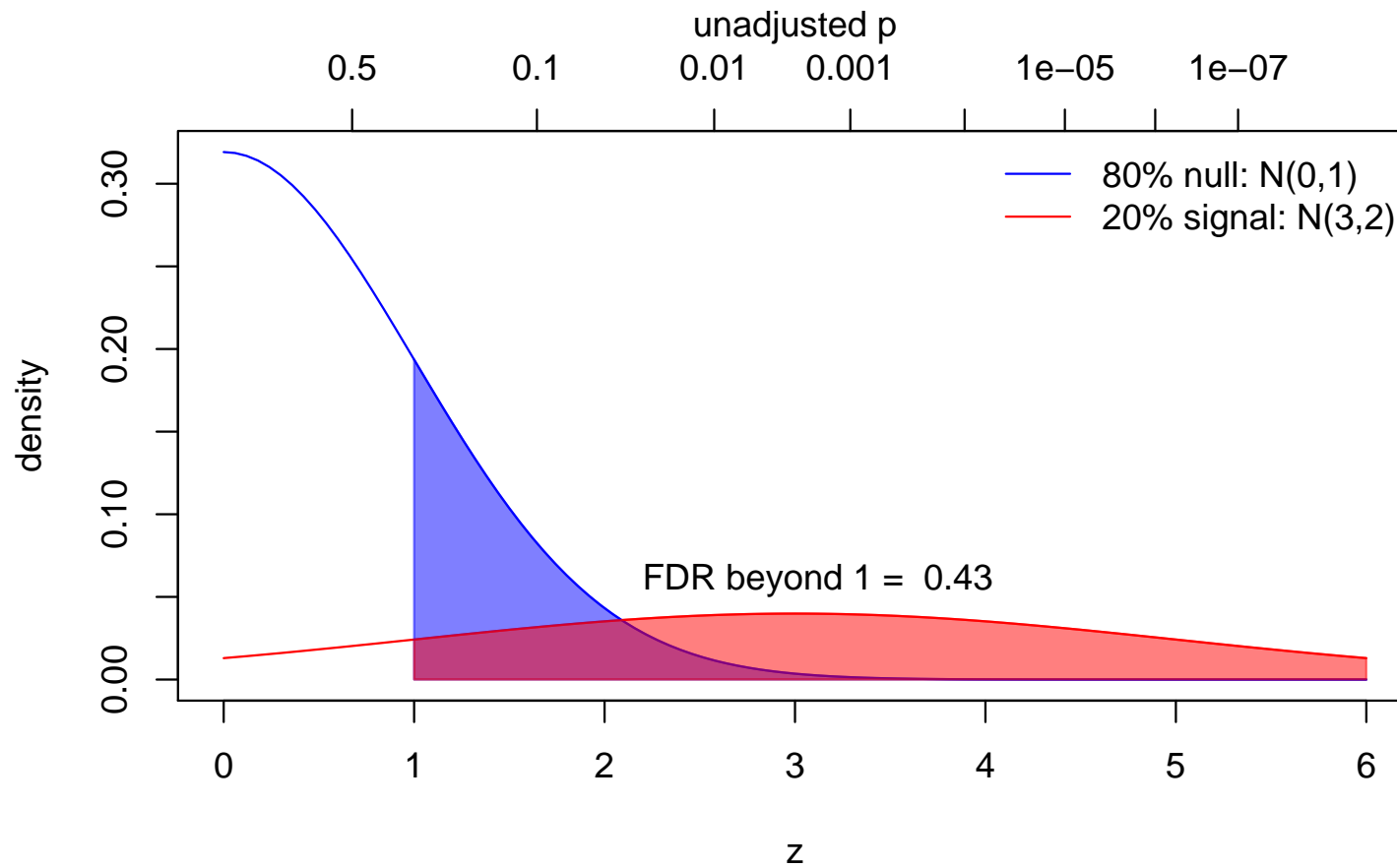
xvals <- seq(1, 6, l=101)
polygon(
  c(xvals,6,1), c(0.8*dnorm(xvals), 0,0),
  density=NA, col="#0000FF80" ) # transparent blue
polygon(
  c(xvals,6,1), c(0.2*dnorm(xvals,3,2), 0,0),
  density=NA, col="#FF000080" ) # transparent red

legend("topright", bty="n", lty=1, col=c("blue","red"),
  c("80% null: N(0,1)", "20% signal: N(3,2)"))
axis(3, at=qnorm(c(0.25, 0.5*10^(-1:-7))), lower=F), c(0.5, 10^(-1:-7)) )
mtext(side=3, line=2, "unadjusted p")

text(2.2, 0.07, adj=c(0,1), paste("FDR beyond 1 = ",
  round(0.8*pnorm(1,lower=F)/(0.8*pnorm(1,lower=F) + 0.2*pnorm(1,3,2,lower=F)),3)))
```

Transparency

Here's the output;



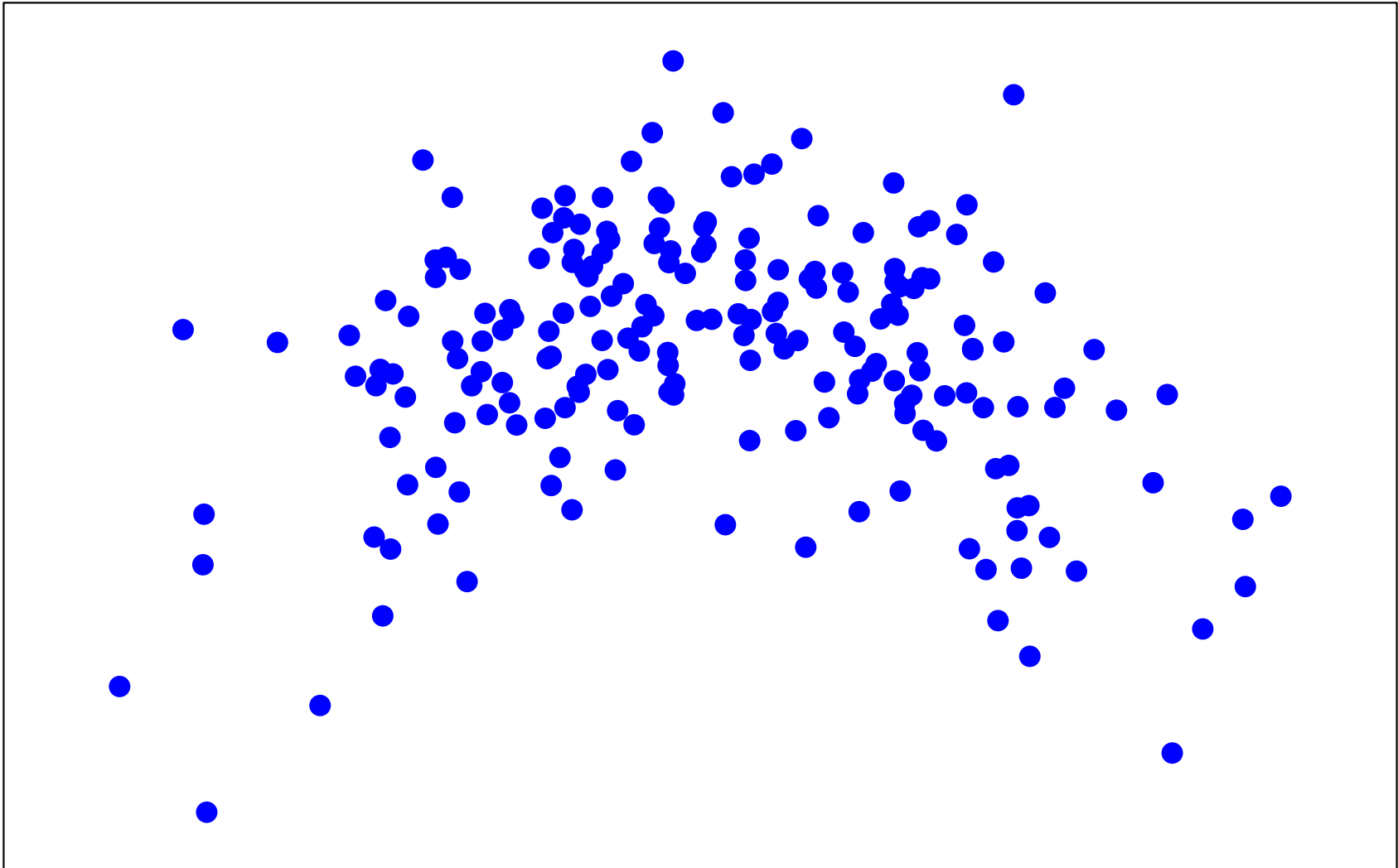
Hexagonal binning

Using transparent plotting symbols is a quick-and-dirty way to adapt scatterplots for use with large datasets.

A better method is 'hexagonal binning'; this is a 2D analog of a histogram – where you would count the number of data in one area, and then draw a bar with height proportional to that count.

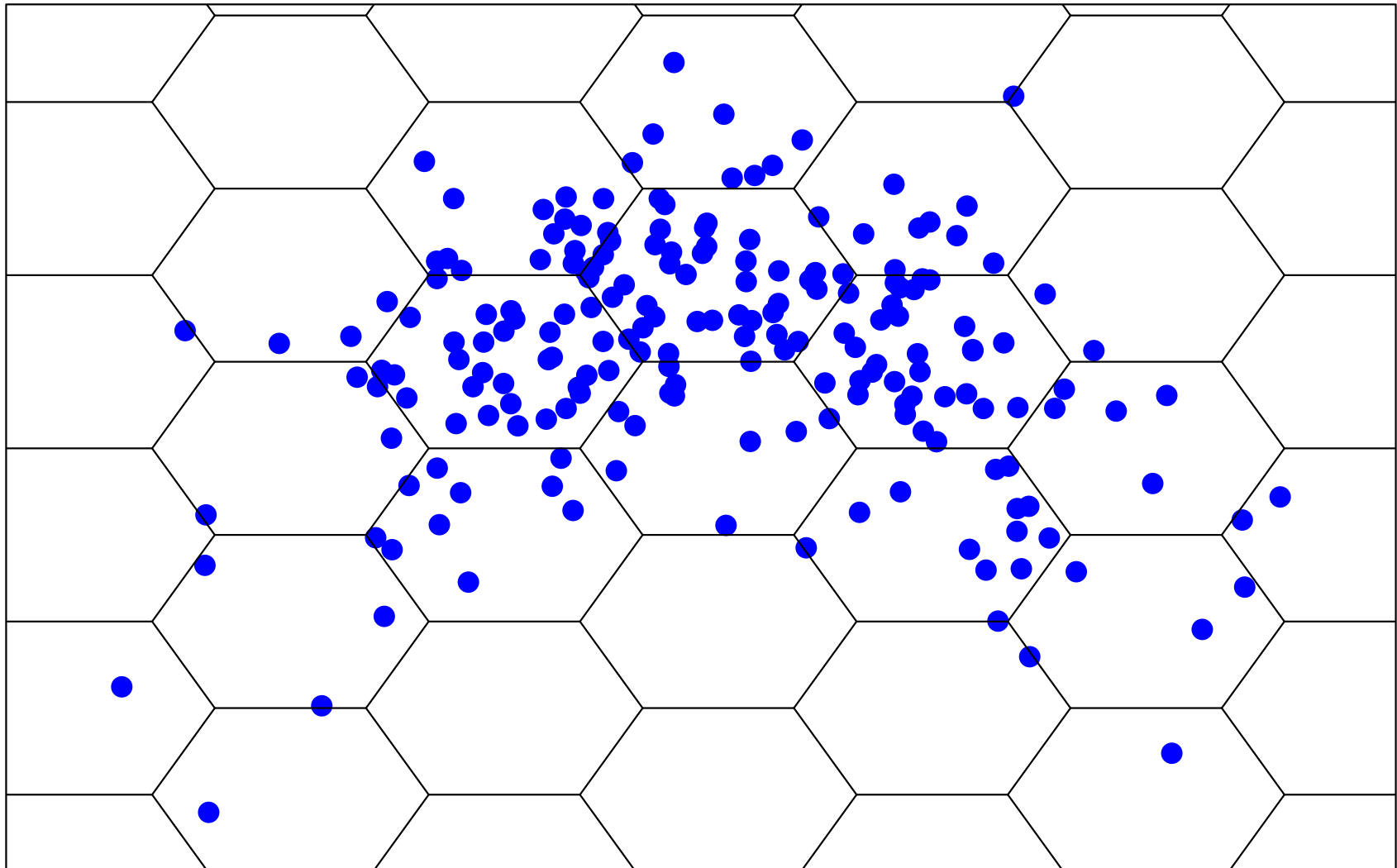
Hexagonal binning

Binning in two dimensions;



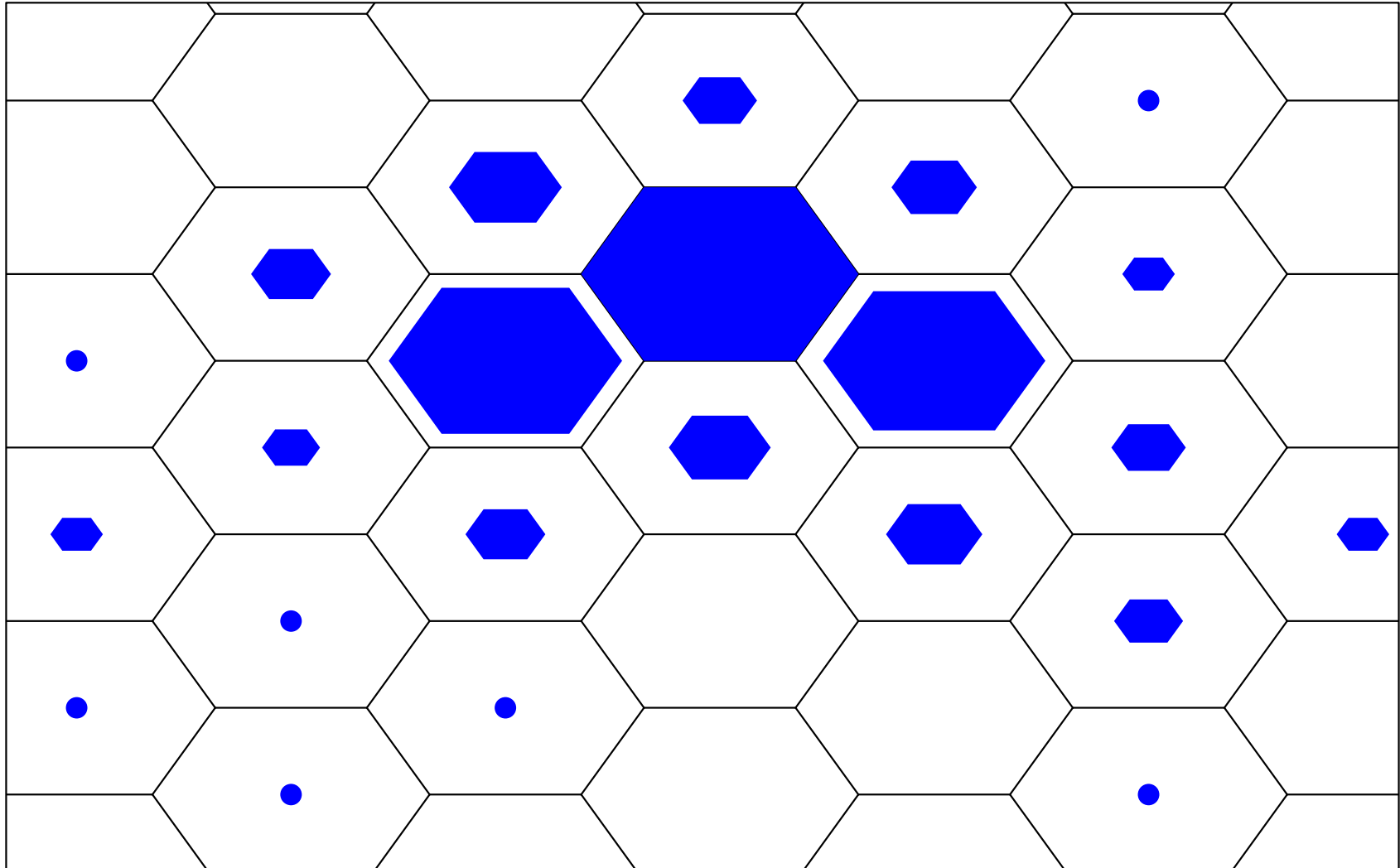
Hexagonal binning

Binning in two dimensions;



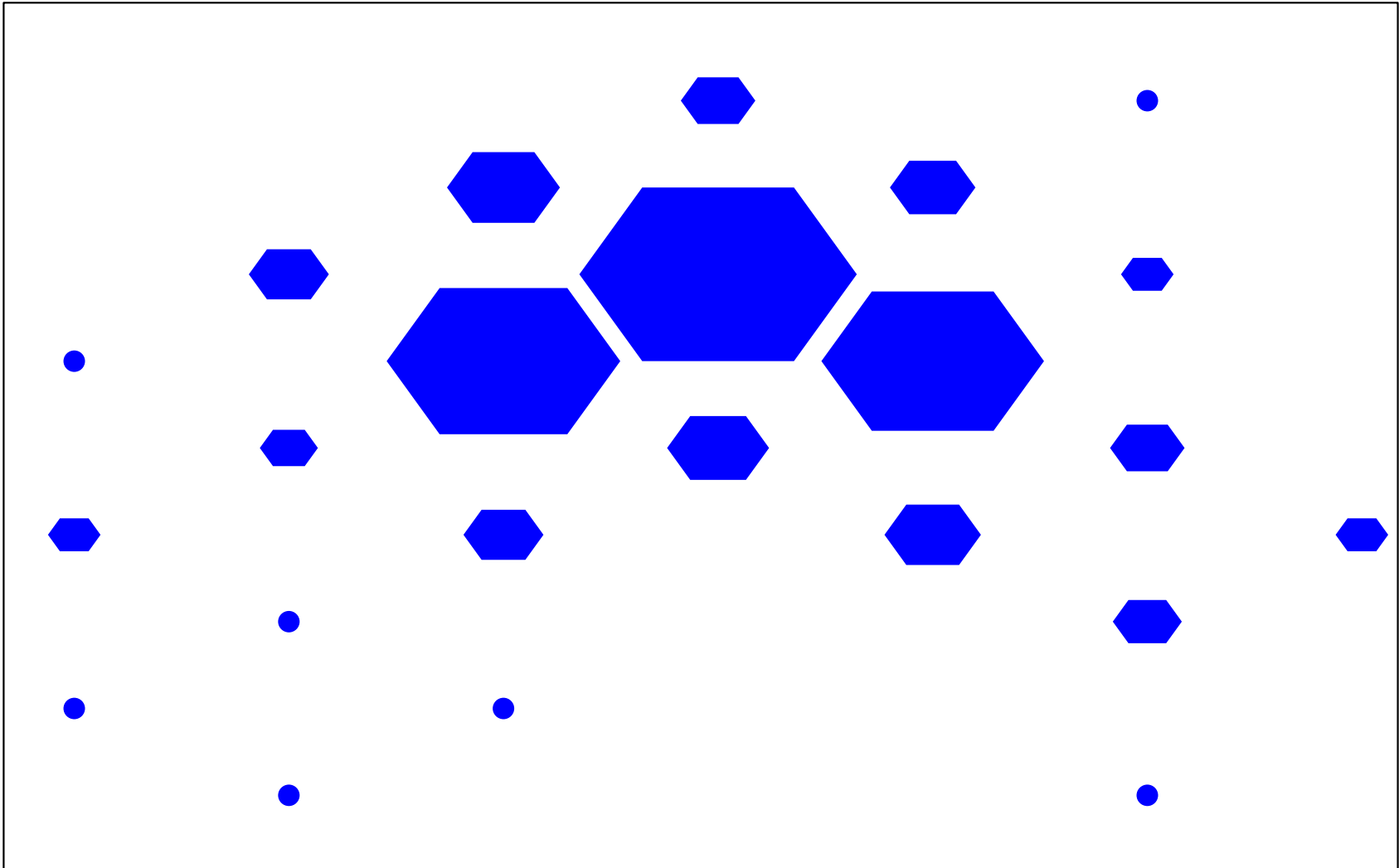
Hexagonal binning

Binning in two dimensions;



Hexagonal binning

Binning in two dimensions;

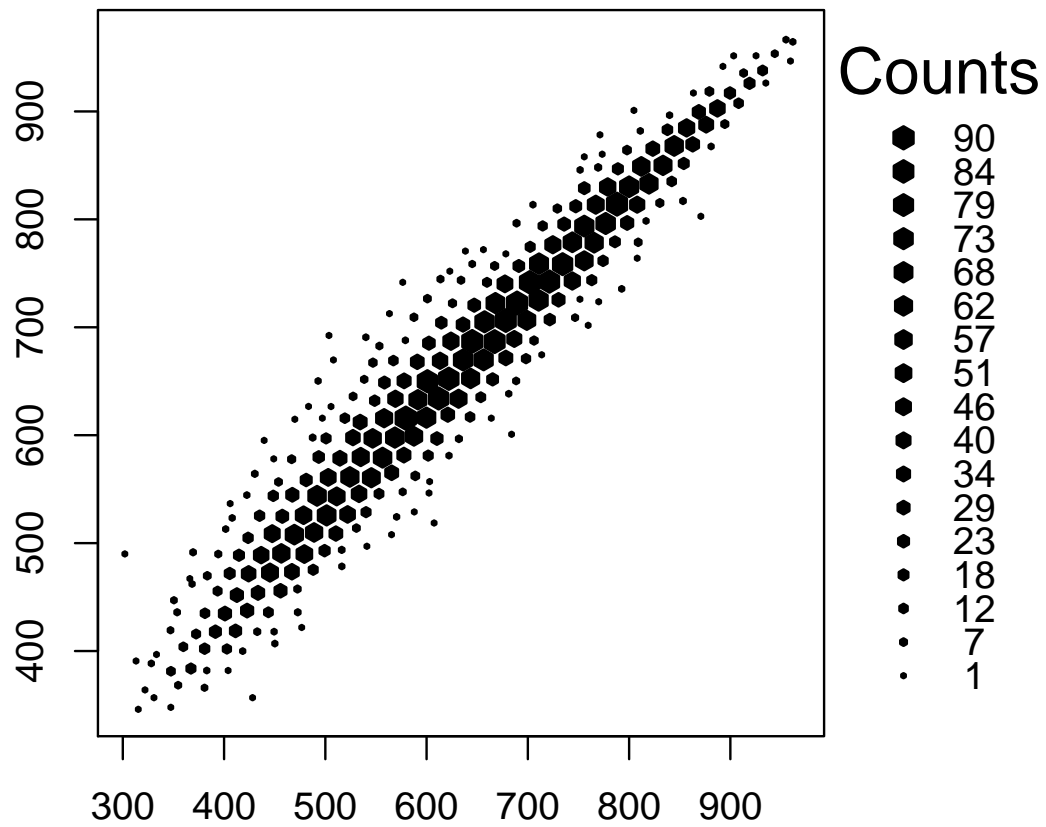


Hexagonal binning

The `hexbin()` package does all the bin construction, and counting. It has a `plot` method for its `hexbin` objects;

```
install.packages(c("hexbin", "survey"))  
library("hexbin")  
library("survey")# for apipop data frame  
  
with(apipop, plot(hexbin(api99, api00), style="centroids"))
```

Hexagonal binning



Hexagonal binning

Hexbin is used when you don't *really* care about the exact location of every single point

- Singleton points are plotted 'as usual'; you do (perhaps) care about them
- `hexbin` centers the 'ink' at the cell data's 'center of gravity'
- `style="centroids"` gives the center-of-gravity version; the default style is `colorscale` – usually grayscale. See `?gplot.hexagons` for more options

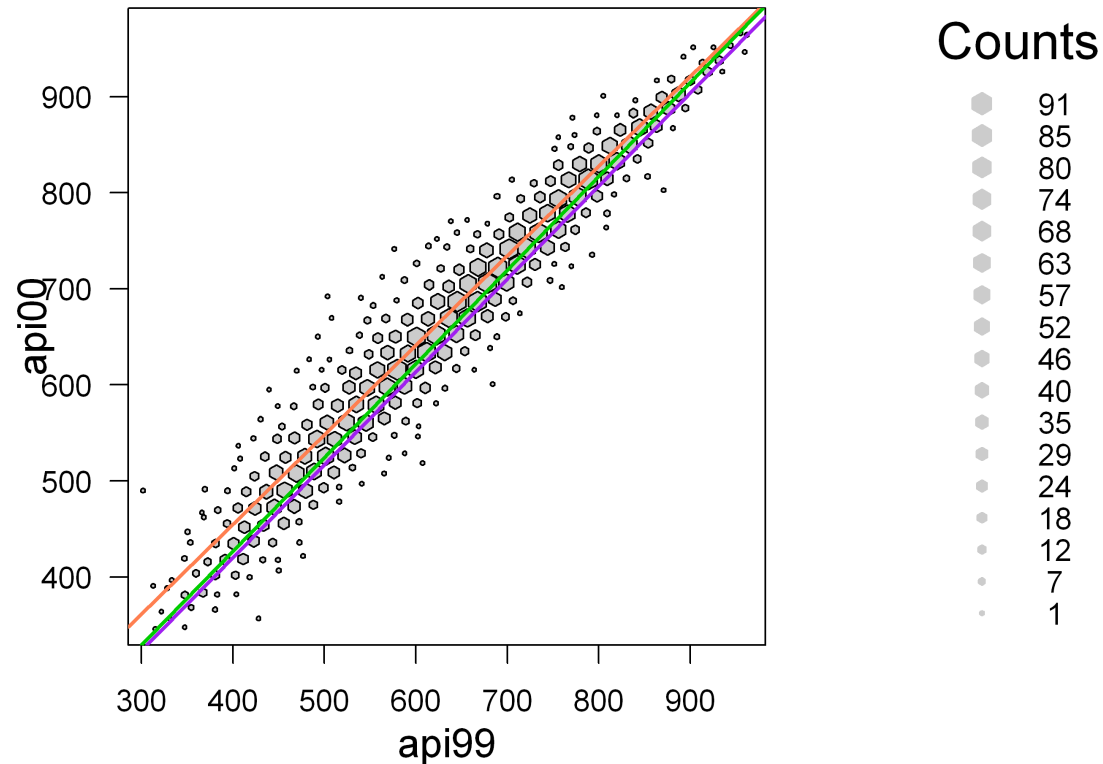
Hexagonal binning

For keen people: the `hexbin` package doesn't use the standard R graphics plotting devices; instead, it operates through the `Grid` system (in the `grid` package) which defines rectangular regions on a graphics device; these `viewport` regions can have a number of coordinate systems. To add lines to a hexbin plot, the options are;

- Use `hexVP.abline()` to add these directly
- Move everything into 'standard' graphics – not `Grid` graphics (see `?Grid`). The `Grid` system lets you alter graphics *after* plotting them
- Write your own plot method for hexbin objects, with standard R graphics commands
- Make do with `hexBinning()` in the `fMultivar` package

Hexagonal binning

An example; color-coded lines of best fit, by school type;



```
lm.e <- coef(lm(api00~api99, data=apipop, subset=stype=="E"))  
lm.m <- coef(lm(api00~api99, data=apipop, subset=stype=="M"))  
lm.h <- coef(lm(api00~api99, data=apipop, subset=stype=="H"))  
  
hexVP.abline(vp1$plot.vp, lm.e[1], lm.e[2], col="coral")
```

File formats

Ultimately, we want to output the graph in an appropriate file format. (Cut-and-paste is possible, but not recommended)

R knows more about font sizes and spacing than most users – so first design the graph at the size it will end up, eg:

```
## on Windows  
windows(height=4,width=6)  
## on Unix  
x11(height=4,width=6)
```

... and, when that's done, write a version to a file

File formats

For example, for a 6×4 PDF file;

```
pdf("myprettypic.pdf", height=4, width=6) # inches
... plotting commands here ...
dev.off() # close the file
```

Some other formats: (see ?Devices for a full list)

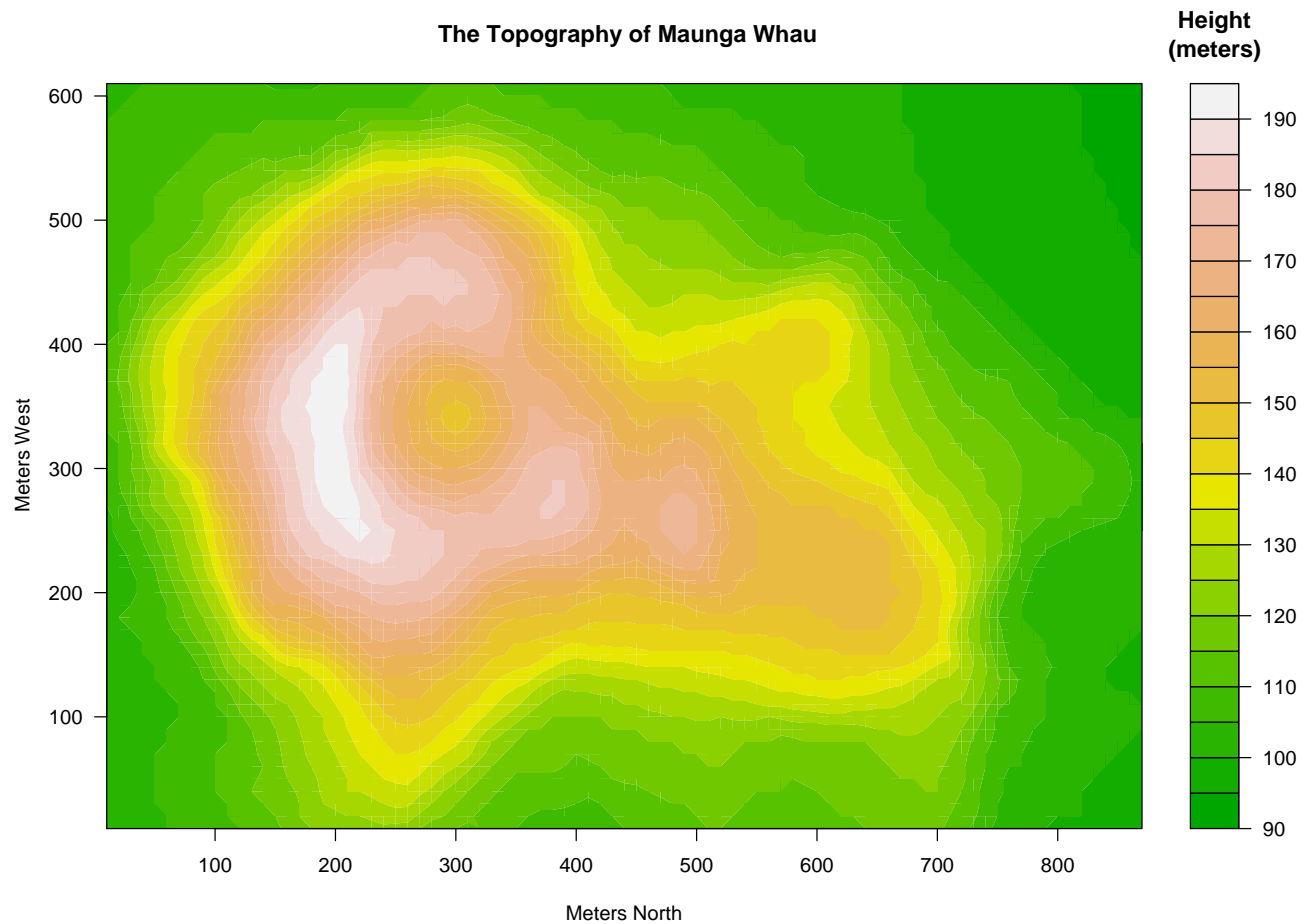
- `jpeg("mypic.jpg", w=6*288, h=4*288, res=288)` – lossy
- `png("mypic.png", w=6*288, h=4*288, res=288)` – lossless

– point size of text can also be manipulated, which can be useful when making posters

PowerPoint, or Word, or \LaTeX can all rescale graphs. But when the graph gets smaller, so do the axis labels...

File formats

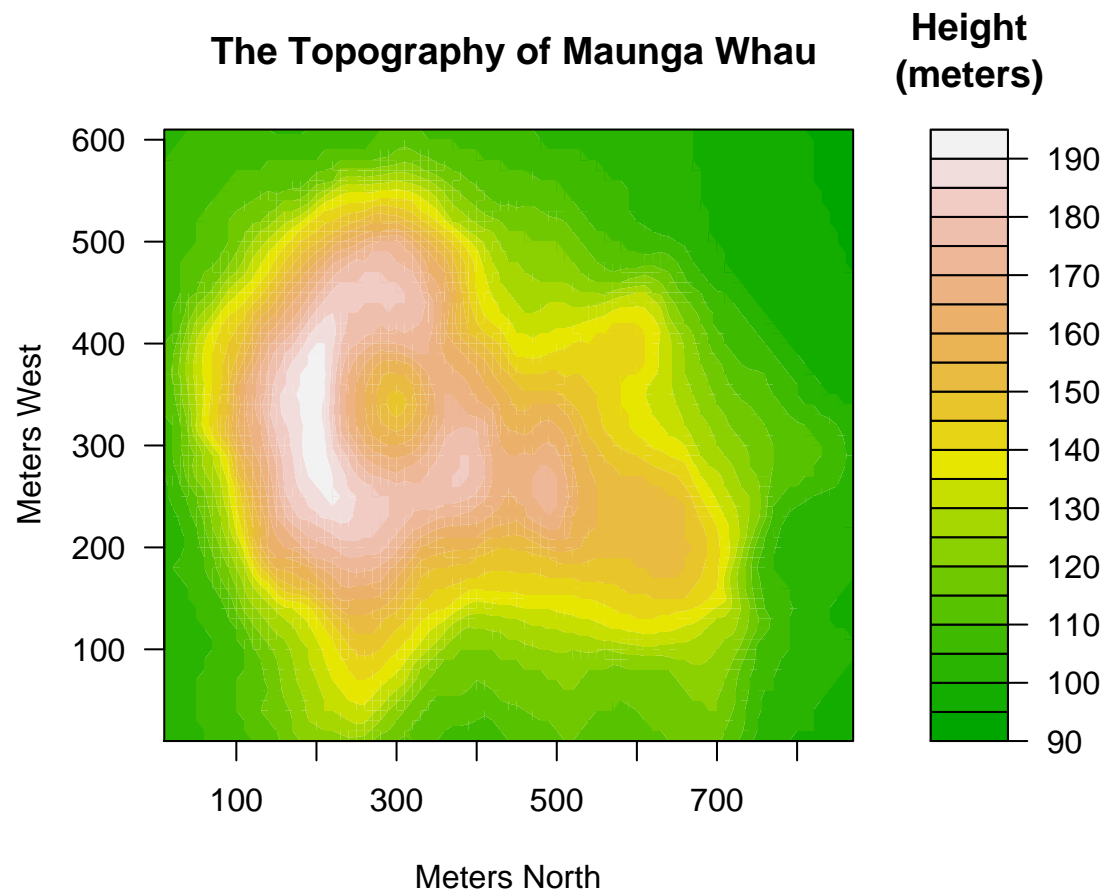
Created at full-page size (11×8.5 inches)



filled.contour(.) from R version 2.5.1 (2007-06-27)

File formats

Created at 6×5 inches



filled.contour(.) from R version 2.5.1 (2007-06-27)

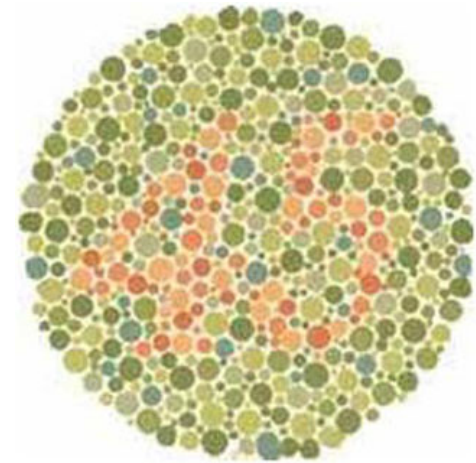
Color schemes

Color choice is best left to experts, or people with taste.

<http://www.colorbrewer.org> has color schemes designed for the National Cancer Atlas, also in package `RColorBrewer`

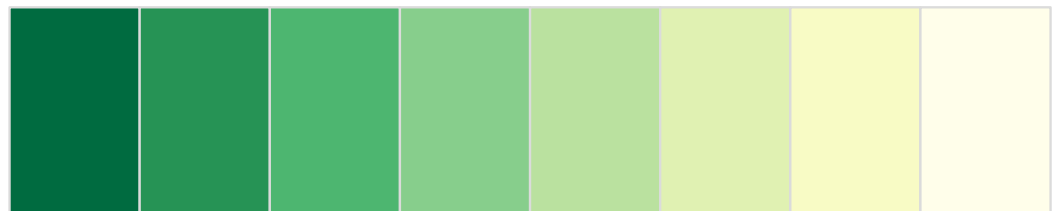
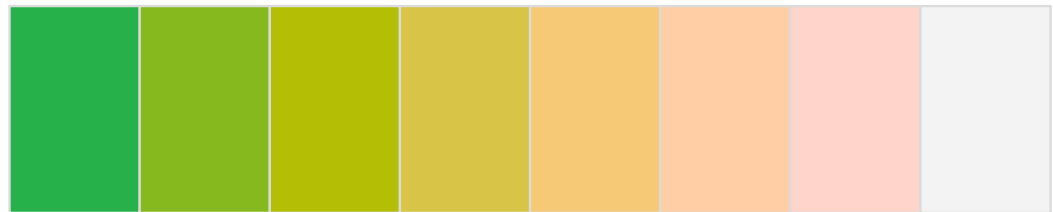
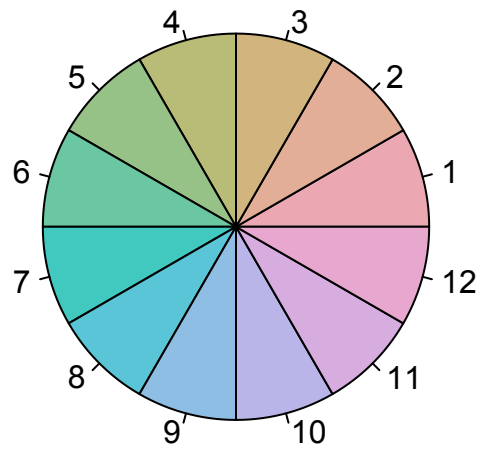
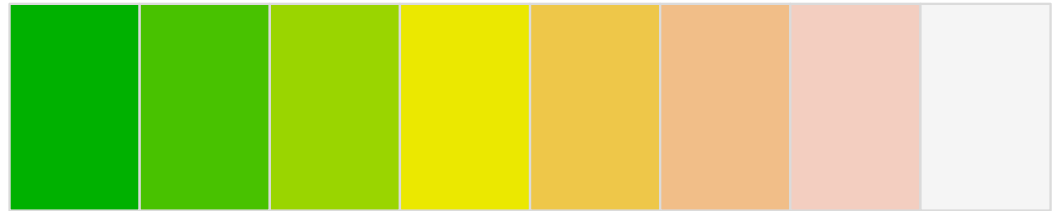
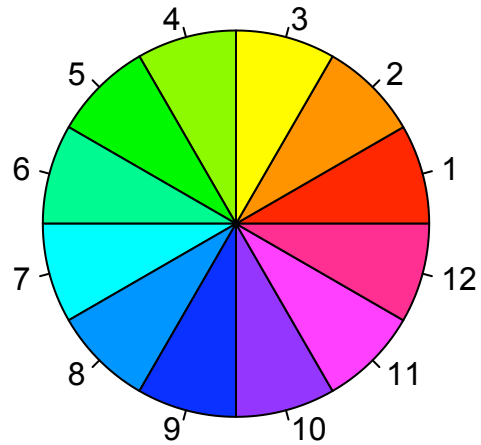
`colorspace` package has color schemes based on straight lines in a perceptually-based color space (rather than RGB).

`dichromat` package attempts to show the impact of red:green color blindness on your R color schemes.



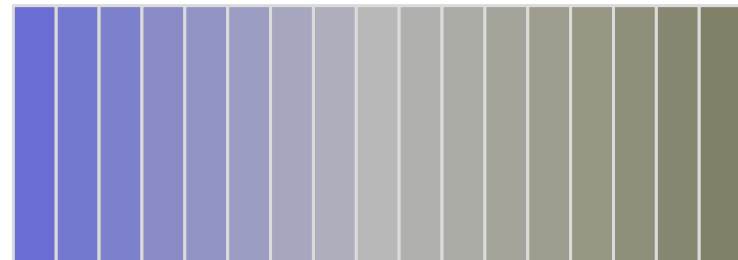
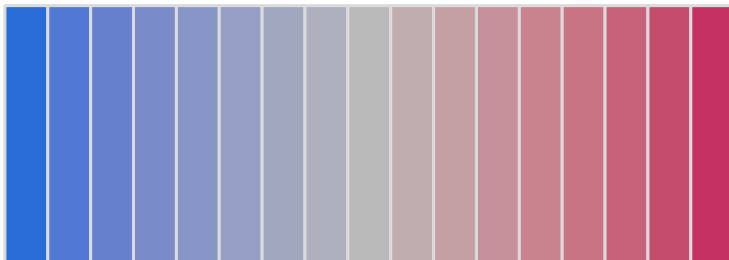
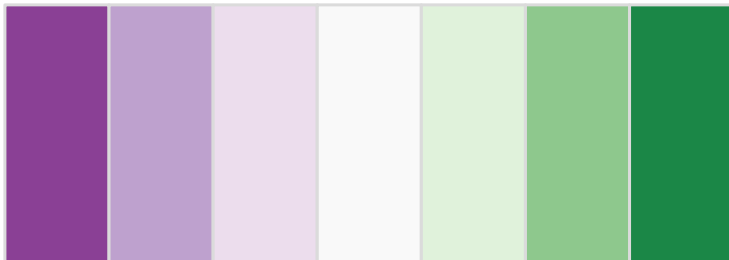
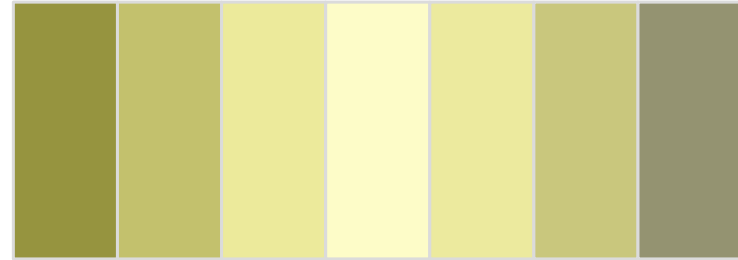
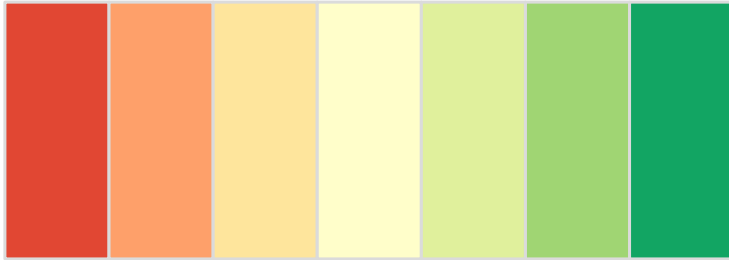
[Code for examples is in file `colorpalettes.R` on course website]

Color choice



(nb B&W printed copies of this slide may not be helpful!)

Color blindness



(nb B&W printed copies of this slide may not be helpful!)