

Data Visualization

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Basic Plots

We covered some basic plots previously, but we are going to expand the ability to customize these basic graphics first.

Read in Data

```
> death = read.csv("http://biostat.jhsph.edu/~ajaffe/files/  
+                  as.is=TRUE,header=TRUE, row.names=1)  
> print(death[1:2, 1:5])
```

	X1760	X1761	X1762	X1763	X1764
Afghanistan	NA	NA	NA	NA	NA
Albania	NA	NA	NA	NA	NA

We see that the column names were years, and R doesn't necessarily like to read in a column name that starts with a number and puts an X there.

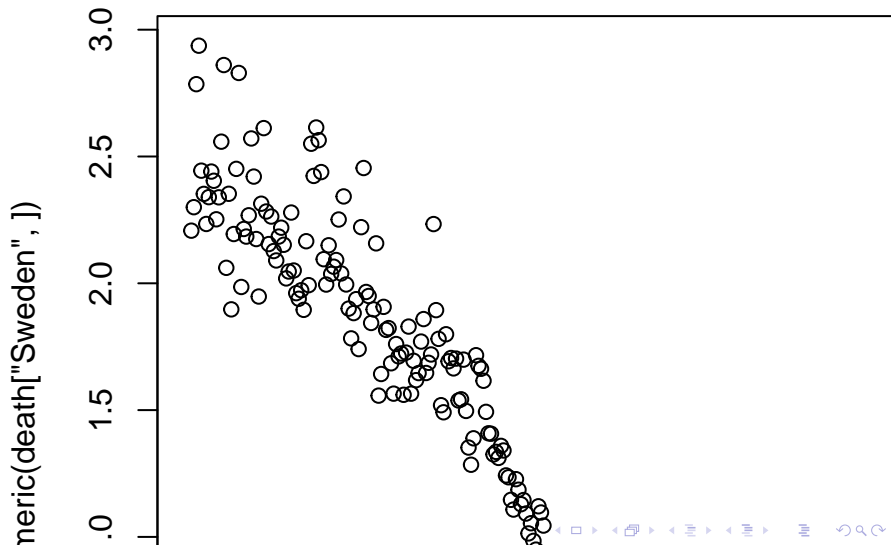
We'll just take off that X and get the years.

```
year = as.integer(gsub("X", "", names(death)))  
head(year)
```

```
[1] 1760 1761 1762 1763 1764 1765
```

Basic Plots

```
> plot(as.numeric(death["Sweden",])~year)
```



Basic Plots

The y-axis label isn't informative, and we can change the label of the y-axis using `ylab` (`xlab` for `x`), and `main` for the main title/label.

```
> plot(as.numeric(death["Sweden",])~year,  
+       ylab="# of deaths per family", main = "Sweden")
```

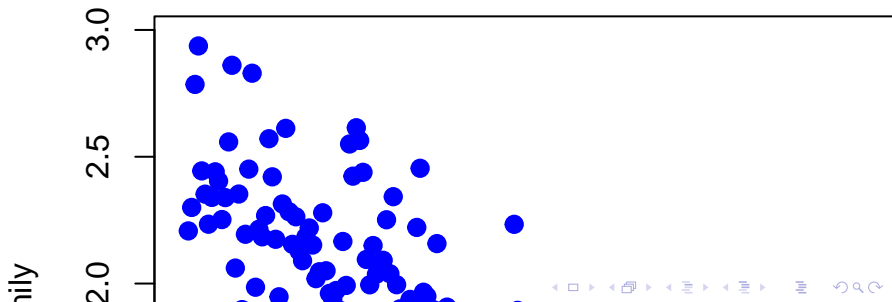


Basic Plots

Let's drop any of the projections and keep it to year 2012, and change the points to blue.

```
> plot(as.numeric(death["Sweden",])~year,  
+       ylab="# of deaths per family", main = "Sweden",  
+       xlim = c(1760,2012), pch = 19, cex=1.2,col="blue")
```

Sweden

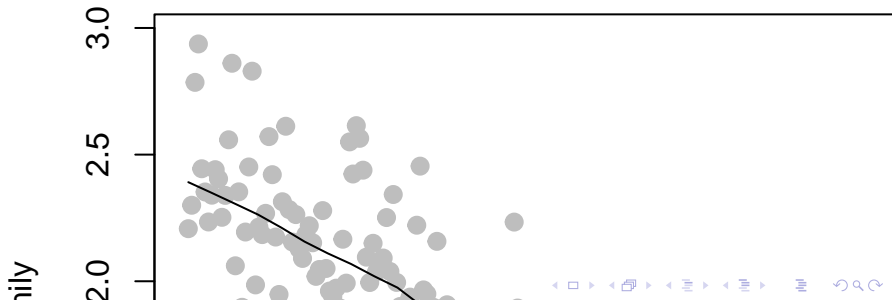


Basic Plots

Using `scatter.smooth` plots the points and runs a loess smoother through the data.

```
> scatter.smooth(as.numeric(death["Sweden",])~year,span=0.2,  
+               ylab="# of deaths per family", main = "Sweden",lwd=2,  
+               xlim = c(1760,2012), pch = 19, cex=0.9,col="grey")
```

Sweden



Basic Plots

`par(mfrow=c(1,2))` tells R that we want to set a parameter (par function) named `mfrow` (number of plots - 1 row, 2 columns) so we can have 2 plots side by side (Sweden and the UK)

```
> par(mfrow=c(1,2))
> scatter.smooth(as.numeric(death["Sweden",])~year,span=0.2,
+               ylab="# of deaths per family", main = "Sweden",lwd=
+               xlim = c(1760,2012), pch = 19, cex=0.9,col="grey")
> scatter.smooth(as.numeric(death["United Kingdom",])~year,span=0.2,
+               ylab="# of deaths per family", main = "United Kingdom",lwd=
+               xlim = c(1760,2012), pch = 19, cex=0.9,col="grey")
```

Sweden

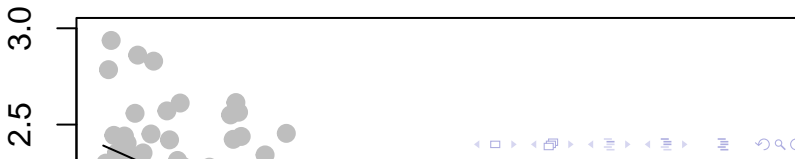


Basic Plots

We can set the y-axis to be the same.

```
> par(mfrow=c(1,2))
> yl = range(death[c("Sweden","United Kingdom"),])
> scatter.smooth(as.numeric(death["Sweden",])~year,span=0.2,
+               ylab="# of deaths per family", main = "Sweden",lwd=
+               xlim = c(1760,2012), pch = 19, cex=0.9,col="grey")
> scatter.smooth(as.numeric(death["United Kingdom",])~year,
+               ylab="", main = "United Kingdom",lwd=3,ylim=yl,
+               xlim = c(1760,2012), pch = 19, cex=0.9,col="grey")
```

Sweden

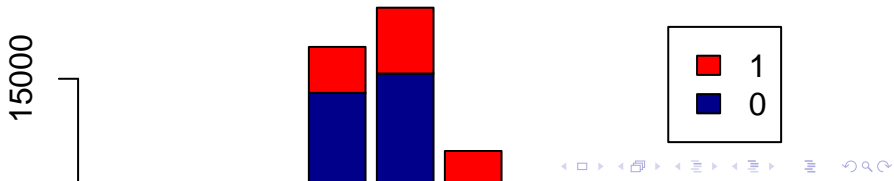


Bar Plots

- ▶ Stacked Bar Charts are sometimes wanted to show distributions of data

```
> ## Stacked Bar Charts
> cars = read.csv("http://biostat.jhsph.edu/~ajaffe/files/1
> counts <- table(cars$IsBadBuy, cars$VehicleAge)
> barplot(counts, main="Car Distribution by Age and Bad Buy
+   xlab="Vehicle Age", col=c("darkblue","red"),
+   legend = rownames(counts))
```

Car Distribution by Age and Bad Buy Status

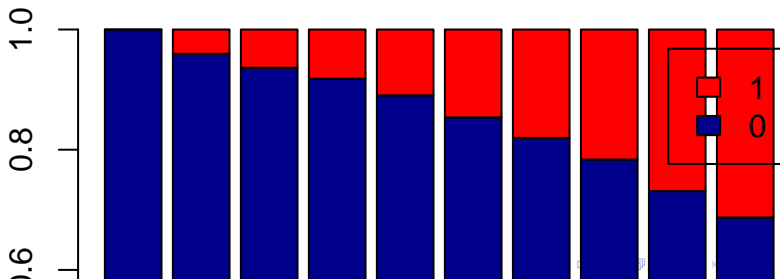


Bar Plots

`prop.table` allows you to convert a table to proportions (depends on margin - either row percent or column percent)

```
> ## Use percentages (column percentages)
> barplot(prop.table(counts, 2), main="Car Distribution by
+   xlab="Vehicle Age", col=c("darkblue","red"),
+   legend = rownames(counts))
```

Car Distribution by Age and Bad Buy Status

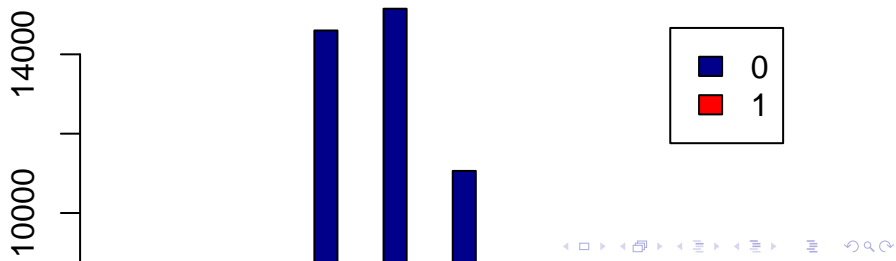


Bar Plots

Using the `beside` argument in `barplot`, you can get side-by-side barplots.

```
> # Stacked Bar Plot with Colors and Legend  
> barplot(counts, main="Car Distribution by Age and Bad Buy  
+   xlab="Vehicle Age", col=c("darkblue","red"),  
+   legend = rownames(counts), beside=TRUE)
```

Car Distribution by Age and Bad Buy Status



Graphics parameters

Set within most plots in the base 'graphics' package:

- * `pch` = point shape, http://voteview.com/symbols_pch.htm
- * `cex` = size/scale
- * `xlab`, `ylab` = labels for x and y axes
- * `main` = plot title
- * `lwd` = line density
- * `col` = color
- * `cex.axis`, `cex.lab`, `cex.main` = scaling/sizing for axes marks, axes labels, and title

Devices

By default, R displays plots in a separate panel. From there, you can export the plot to a variety of image file types, or copy it to the clipboard.

However, sometimes its very nice to save many plots made at one time to one pdf file, say, for flipping through. Or being more precise with the plot size in the saved file.

R has 5 additional graphics devices: `bmp()`, `jpeg()`, `png()`, `tiff()`, and `pdf()`

The syntax is very similar for all of them:

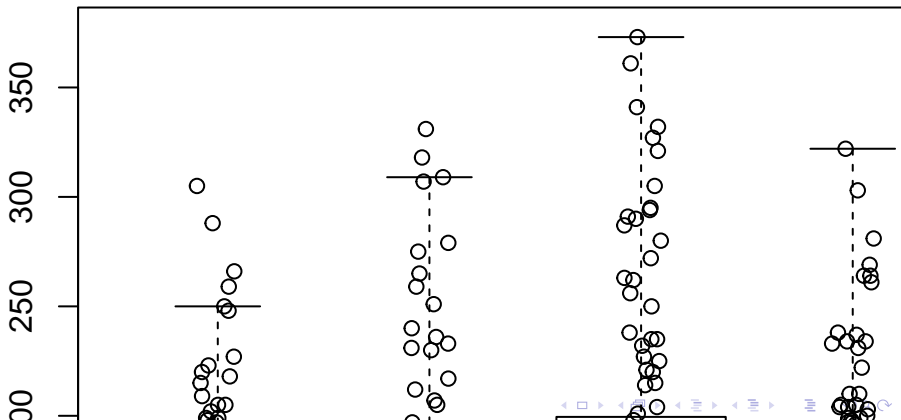
```
pdf("filename.pdf", width=8, height=8) # inches
plot() # plot 1
plot() # plot 2
# etc
dev.off()
```

Basically, you are creating a pdf file, and telling R to write any subsequent plots to that file. Once you are done, you turn the

Boxplots, revisited

These are one of my favorite plots. They are way more informative than the barchart + antenna...

```
> boxplot(weight ~ Diet, data=ChickWeight, outline=FALSE)  
> points(ChickWeight$weight ~ jitter(as.numeric(ChickWeight$Diet))
```



Formulas

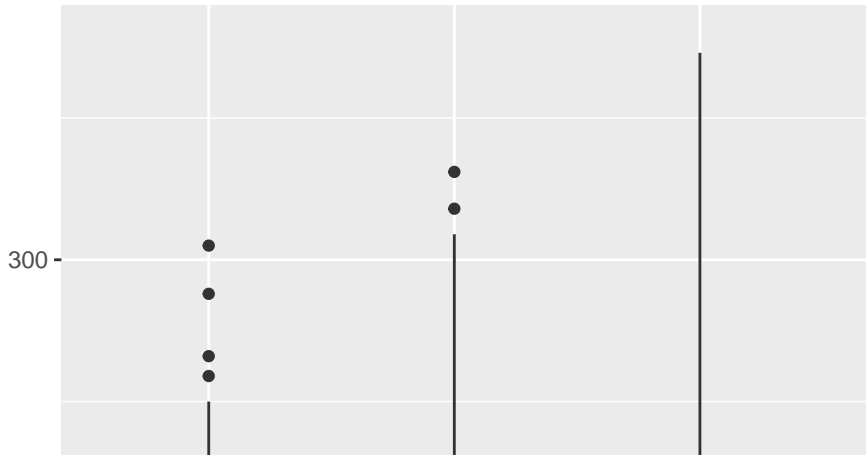
Formulas have the format of $y \sim x$ and functions taking formulas have a `data` argument where you pass the `data.frame`. You don't need to use `$` or referencing when using formulas:

```
boxplot(weight ~ Diet, data=ChickWeight, outline=FALSE)
```


ggplot2

ggplot2 is a package of plotting that is very popular and powerful.

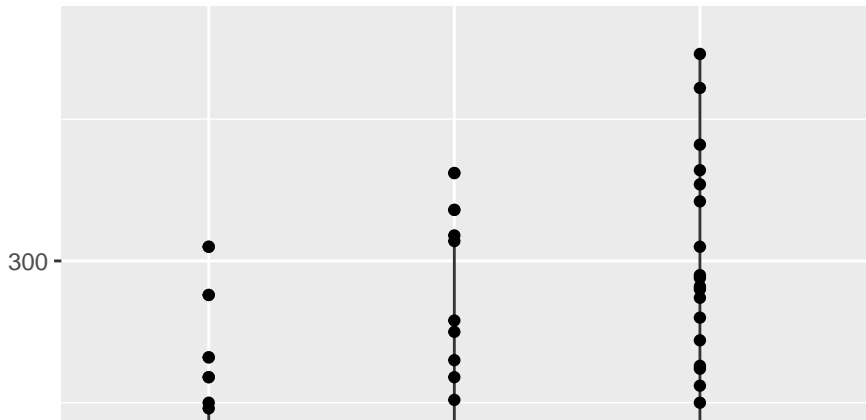
```
> library(ggplot2)
> qplot(factor(Diet), y= weight, data = ChickWeight, geom =
```



Boxplots revisited again

We can do the same plot, by just saying we want a boxplot and points (and jitter the points)

```
> qplot(factor(Diet), y= weight, data = ChickWeight, geom =  
+       position = c('identity', "jitter"))
```

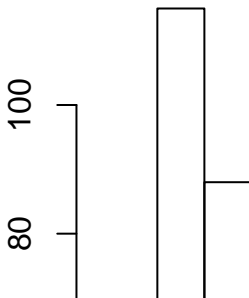


Histograms again

We can do histograms again using `hist`. Let's do histograms of weight at all time points for the chick's weights. We reiterate how useful these are to show your data.

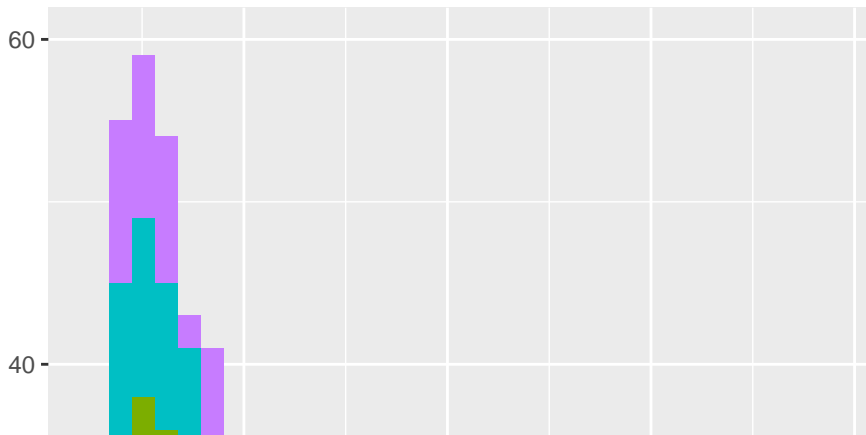
```
> hist(ChickWeight$weight, breaks=20)
```

Histogram of ChickWeight\$weight



Multiple Histograms

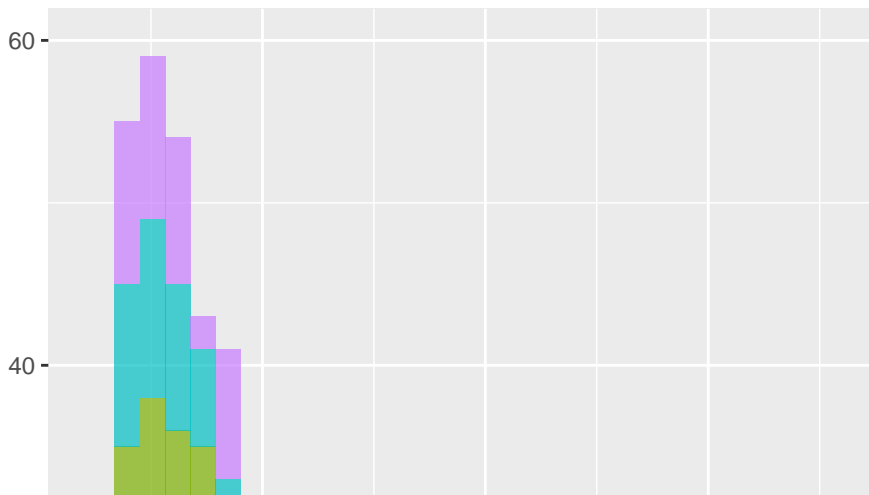
```
> qplot(x = weight,  
+       fill = factor(Diet),  
+       data = ChickWeight,  
+       geom = c("histogram"))
```



Multiple Histograms

Alpha refers to the opacity of the color, less is

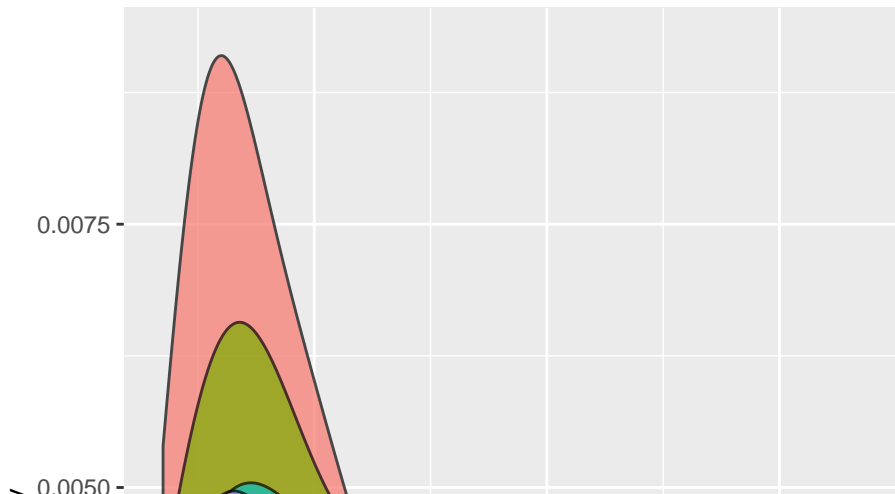
```
> qplot(x = weight, fill = Diet, data = ChickWeight, geom =
```



Multiple Densities

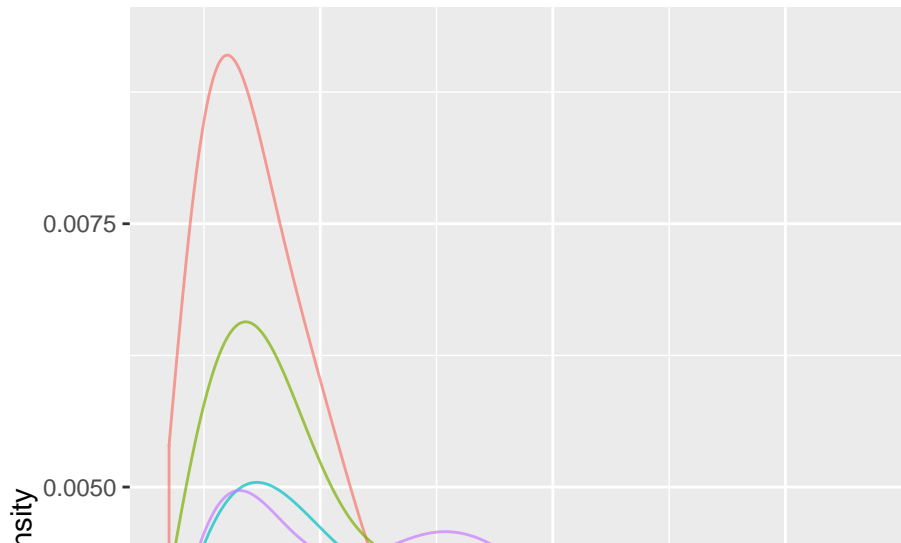
We could also do densities

```
> qplot(x= weight, fill = Diet, data = ChickWeight, geom =
```



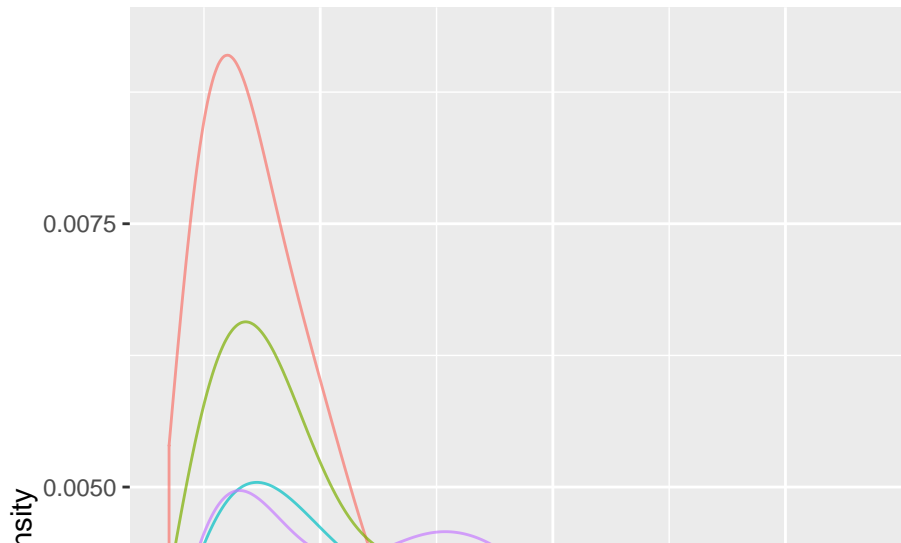
Multiple Densities

```
> qplot(x= weight, colour = Diet, data = ChickWeight, geom
```



Multiple Densities

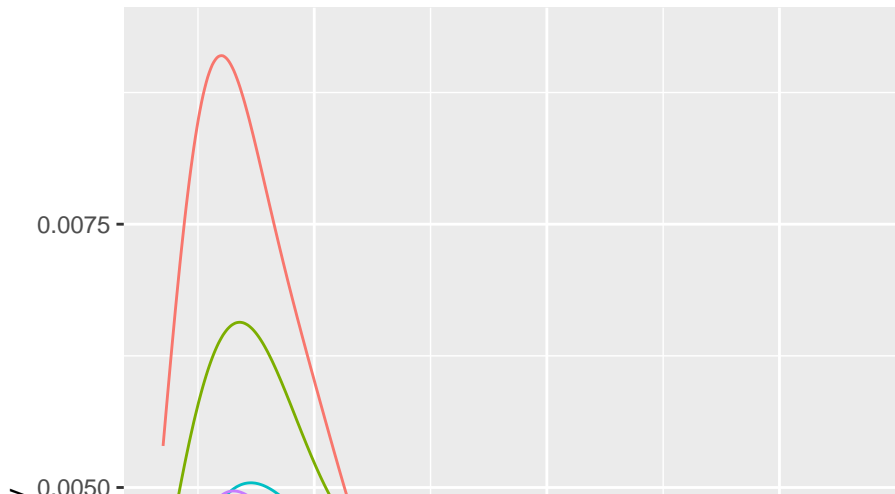
```
> ggplot(aes(x= weight, colour = Diet), data = ChickWeight)
```



Multiple Densities

You can take off the lines of the bottom like this

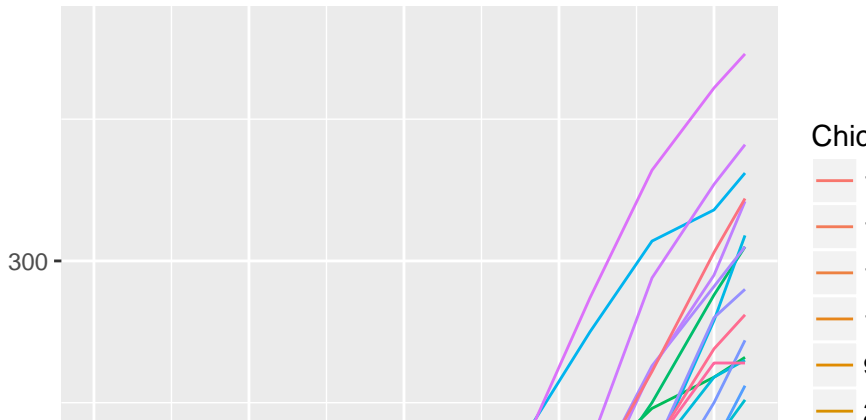
```
> ggplot(aes(x = weight, colour = Diet), data = ChickWeight
```



Spaghetti plot

We can make a spaghetti plot by telling ggplot we want a “line”, and each line is colored by Chick.

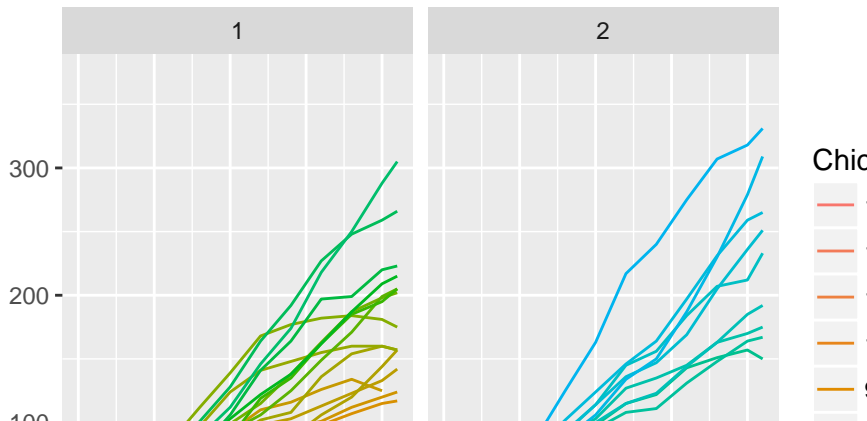
```
> qplot(x=Time, y=weight, colour = Chick,  
+       data = ChickWeight, geom = "line")
```



Spaghetti plot: Facets

In ggplot2, if you want separate plots for something, these are referred to as facets.

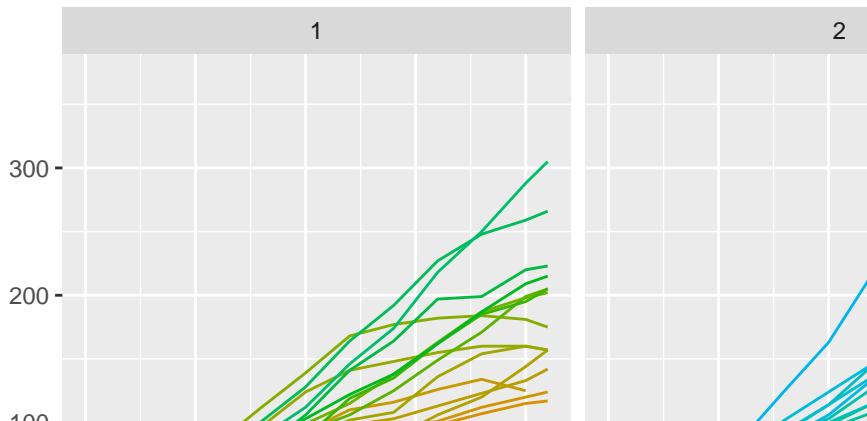
```
> qplot(x=Time, y=weight, colour = Chick, facets = ~ Diet,  
+       data = ChickWeight, geom = "line")
```



Spaghetti plot: Facets

We can turn off the legend (referred to a “guide” in ggplot2). (Note - there is different syntax with the +)

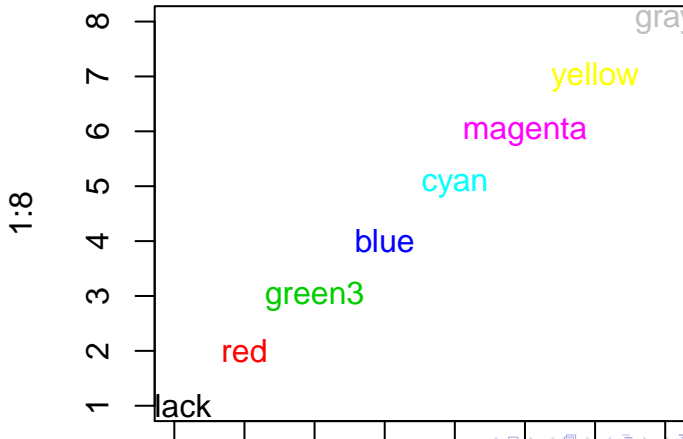
```
> qplot(x=Time, y=weight, colour = Chick, facets = ~ Diet,  
+       data = ChickWeight, geom = "line") + guides(colour = FALSE)
```



Colors

R relies on color 'palettes'.

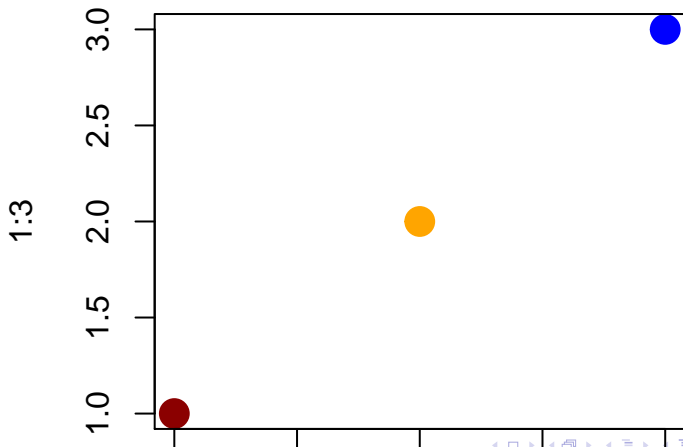
```
> palette("default")  
> plot(1:8, 1:8, type="n")  
> text(1:8, 1:8, lab = palette(), col = 1:8)
```



Colors

The default color palette is pretty bad, so you can try to make your own.

```
> palette(c("darkred", "orange", "blue"))  
> plot(1:3, 1:3, col=1:3, pch = 19, cex=2)
```



Colors

It's actually pretty hard to make a good color palette. Luckily, smart and artistic people have spent a lot more time thinking about this. The result is the 'RColorBrewer' package

`RColorBrewer::display.brewer.all()` will show you all of the palettes available. You can even print it out and keep it next to your monitor for reference.

The help file for `brewer.pal()` gives you an idea how to use the package.

You can also get a “sneak peek” of these palettes at: www.colorbrewer2.com . You would provide the number of levels or classes of your data, and then the type of data: sequential, diverging, or qualitative. The names of the RColorBrewer palettes are the string after 'pick a color scheme:'

Colors

```
> palette("default")
> plot(weight ~ Time, data= ChickWeight, pch = 19, col = D)
```



Adding legends

The `legend()` command adds a legend to your plot. There are tons of arguments to pass it.

`x, y=NULL`: this just means you can give (x,y) coordinates, or more commonly just give x, as a character string:

"top", "bottom", "topleft", "bottomleft", "topright", "bottomright".

`legend`: unique character vector, the levels of a factor

`pch, lwd`: if you want points in the legend, give a pch value. if you want lines, give a lwd value.

`col`: give the color for each legend level

```
> palette(brewer.pal(5,"Dark2"))
> plot(weight ~ jitter(Time,amount=0.2),data=ChickWeight,
+       pch = 19, col = Diet,xlab="Time")
> legend("topleft", paste("Diet",levels(ChickWeight$Diet)),
+       col = 1:length(levels(ChickWeight$Diet)),
+       lwd = 3, ncol = 2)
```

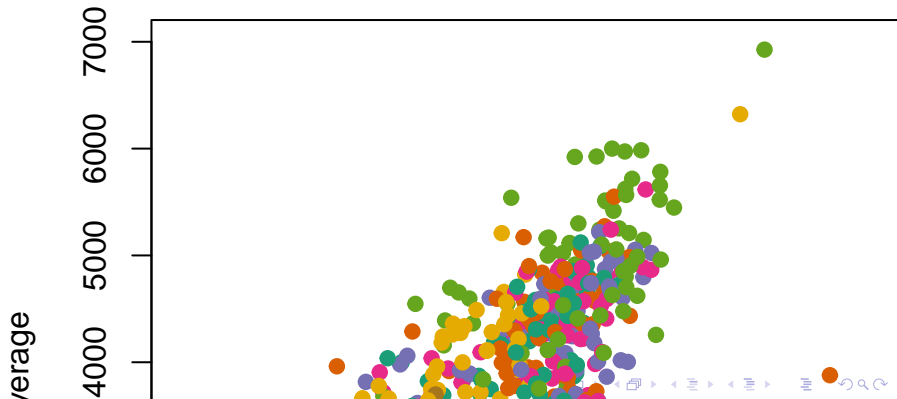
Coloring by variable

```
> destfile = tempfile(fileext = ".rda")
> download.file("http://www.aejaffe.com/winterR_2016/data/c")
> load(destfile)
> dat = circ
> dat2 = circ2
> palette(brewer.pal(7,"Dark2"))
> dd = factor(dat$day)
> plot(orangeAverage ~ greenAverage, data=dat,
+       pch=19, col = as.numeric(dd))
> legend("bottomright", levels(dd), col=1:length(dd), pch =
```



Coloring by variable

```
> dd = factor(dat$day, levels=c("Monday", "Tuesday", "Wednesday",  
+                               "Friday", "Saturday", "Sunday"))  
> plot(orangeAverage ~ greenAverage, data=dat,  
+       pch=19, col = as.numeric(dd))  
> legend("bottomright", levels(dd), col=1:length(dd), pch =
```



More powerful graphics

There are two very common packages for making very nice looking graphics.

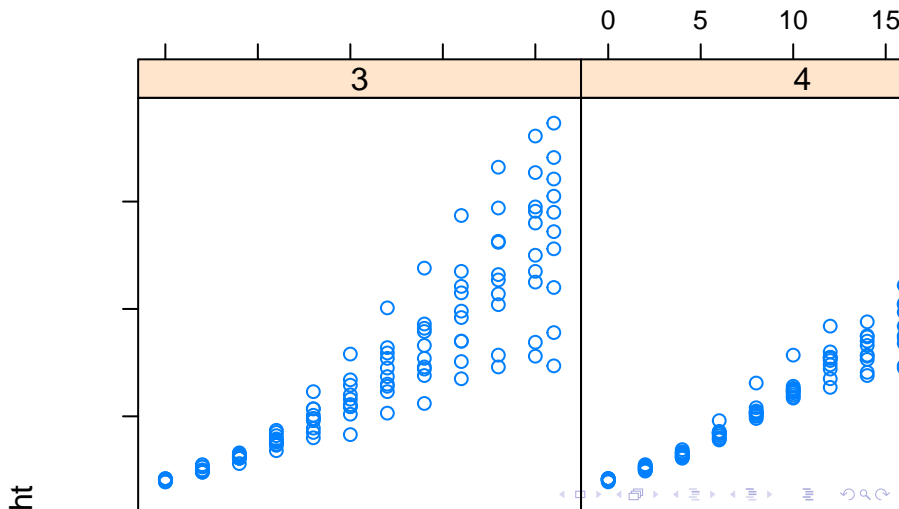
lattice: `http:`

`//lmdvr.r-forge.r-project.org/figures/figures.html`

ggplot2: `http://docs.ggplot2.org/current/index.html`

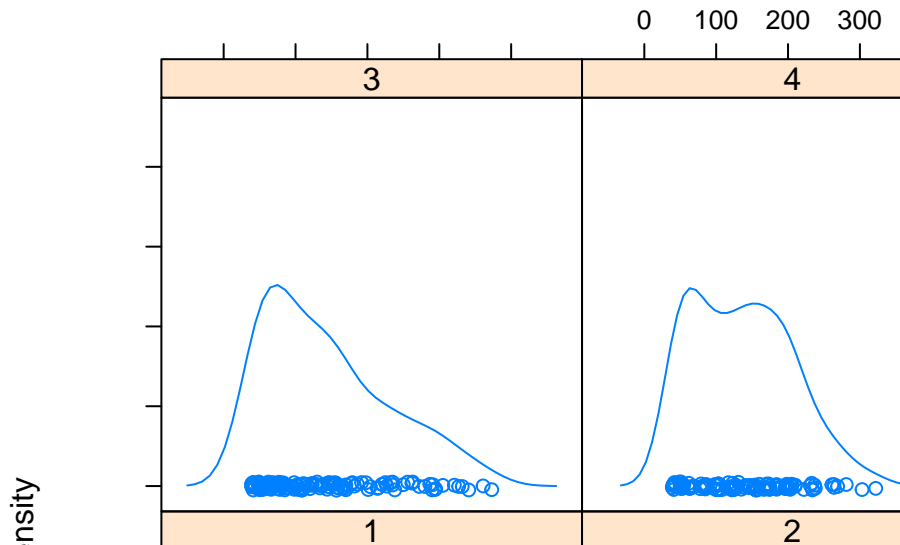
Lattice

```
> library(lattice)  
> xyplot(weight ~ Time | Diet, data = ChickWeight)
```



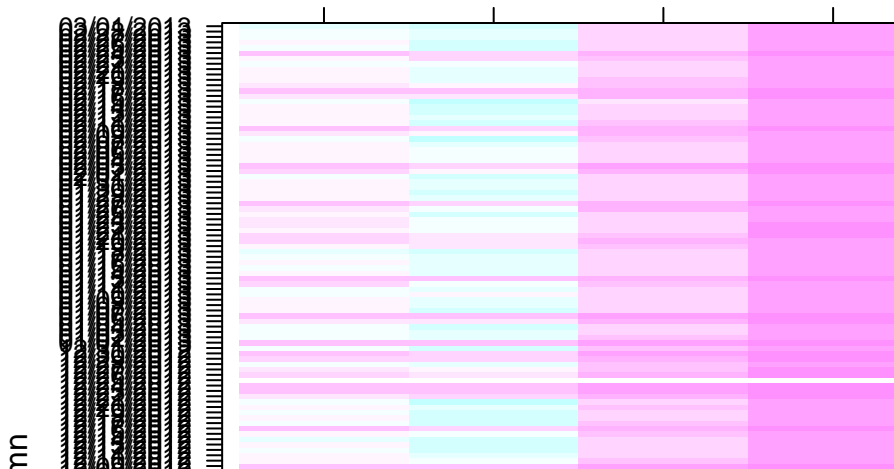
Lattice

```
> densityplot(~weight | Diet, data = ChickWeight)
```



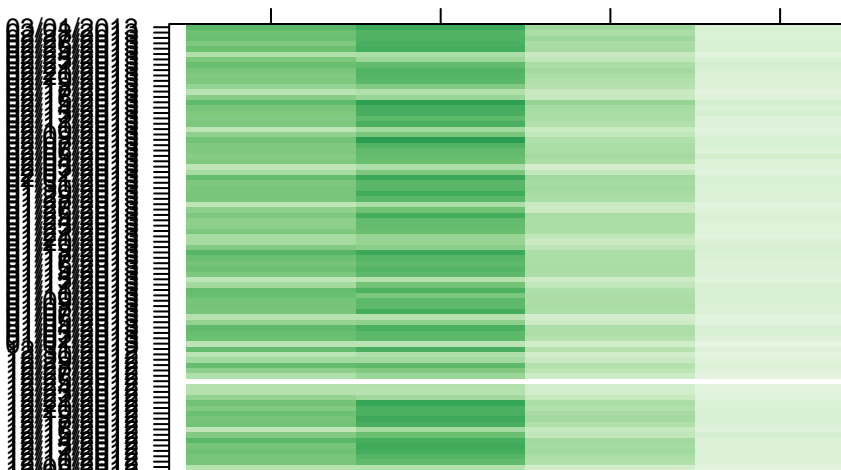
Lattice

```
> rownames(dat2) = dat2$date  
> mat = as.matrix(dat2[975:nrow(dat2),3:6])  
> levelplot(t(mat), aspect = "fill")
```



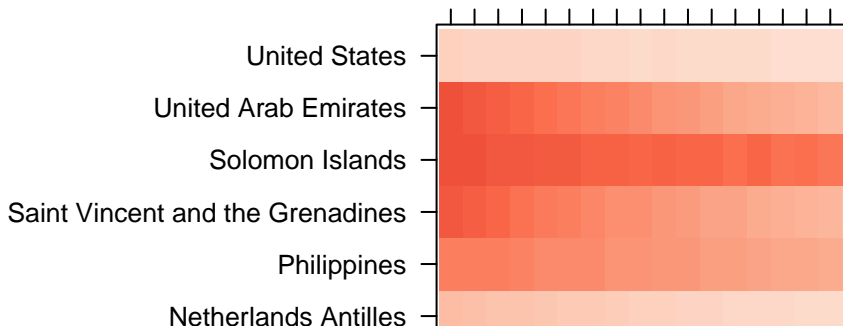
Lattice

```
> theSeq = seq(0,max(mat, na.rm = TRUE), by=50)  
> my.col <- colorRampPalette(brewer.pal(5,"Greens"))(length(theSeq))  
> levelplot(t(mat), aspect = "fill",at = theSeq,col.regions = my.col)
```



Lattice

```
> library(RColorBrewer)
> tmp=death[grepl("s$", rownames(death)), 200:251]
> yr = gsub("X", "", names(tmp))
> theSeq = seq(0,max(tmp,na.rm=TRUE), by=0.05)
> my.col <- colorRampPalette(brewer.pal(5,"Reds"))(length(theSeq))
> levelplot(t(tmp), aspect = "fill", at = theSeq, col.regions = my.col,
+           scales=list(x=list(label=yr, rot=90, cex=0.7)))
```



ggplot2

Useful links:

- ▶ <http://docs.ggplot2.org/0.9.3/index.html>
- ▶ <http://www.cookbook-r.com/Graphs/>