An introduction to ggplot

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1 Getting started with qplot

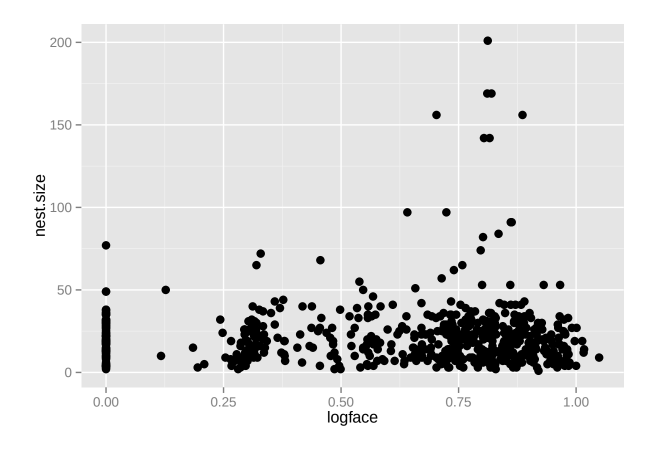
ggplot2 is a powerful plotting system for R. ggplot2 documentation is available at docs.ggplot2.org. qplot is the basic plotting function in the ggplot2 package. qplot is similar to the base R function plot; however, you cannot pass any type of R object to qplot.

Basic usage: qplot(x, y, data...), where data is the data frame to use and ... means other aesthetics passed for each layer.

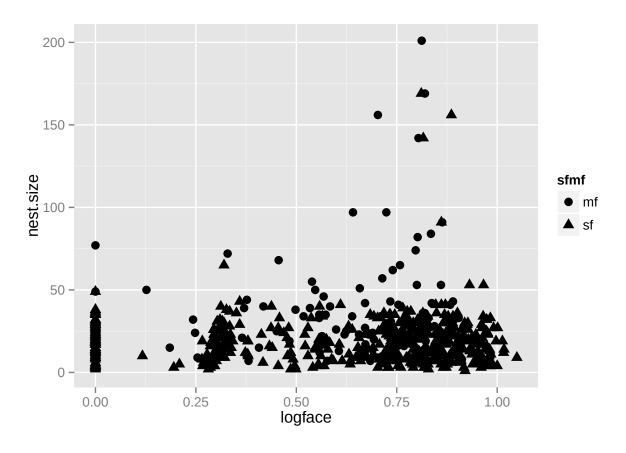
1.1 Using qplot to create scatterplots, histograms and boxplots

1.1.1 Scatterplots

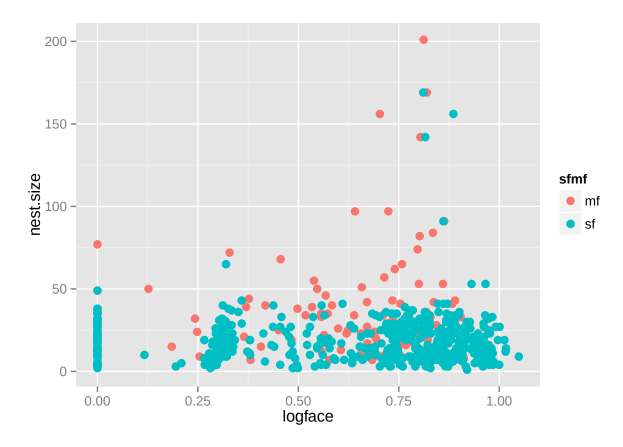
```
# Retrieve a published dataset
# Copy the file's raw GitHub URL
polistesdat.url <- "https://raw.githubusercontent.com/flopezo/atd/master/Tibbetts_et_al_2015_data.csv"
polistesdat.url <- getURL(polistesdat.url)</pre>
polistes.data <- read.csv(textConnection(polistesdat.url))</pre>
# Show the variables included in the dataset
summary(polistes.data)
##
                    day.collected
                                         weight
                                                        \operatorname{\mathfrak{sfmf}}
         year
##
    Min.
           :2011
                   Min.
                           :-6.00
                                    Min.
                                            :0.05100
                                                        mf:106
                   1st Qu.: 1.00
                                    1st Qu.:0.08400
                                                        sf:505
##
   1st Qu.:2011
   Median:2011
                   Median :25.00
                                    Median :0.09600
##
    Mean
           :2011
                   Mean
                           :20.96
                                    Mean
                                            :0.09692
##
    3rd Qu.:2012
                   3rd Qu.:31.00
                                    3rd Qu.:0.10900
##
   Max.
           :2012
                   Max.
                           :53.00
                                    Max.
                                            :0.15300
##
      nest.size
                         logface
##
   Min.
          : 1.00
                     Min.
                             :0.0000
##
   1st Qu.: 10.00
                      1st Qu.:0.3315
  Median : 19.00
                      Median :0.7270
           : 22.28
##
  Mean
                      Mean
                             :0.6014
    3rd Qu.: 27.00
                      3rd Qu.:0.8470
##
           :201.00
##
   Max.
                     Max.
                             :1.0490
# Create a scatter diagram
qplot(logface, nest.size, data=polistes.data) + geom_point(size=3)
```



qplot(logface, nest.size, data=polistes.data, shape=sfmf) + geom_point(size=3)



Add colors based on a categorical variable
qplot(logface, nest.size, data=polistes.data, colour=sfmf) + geom_point(size=3)

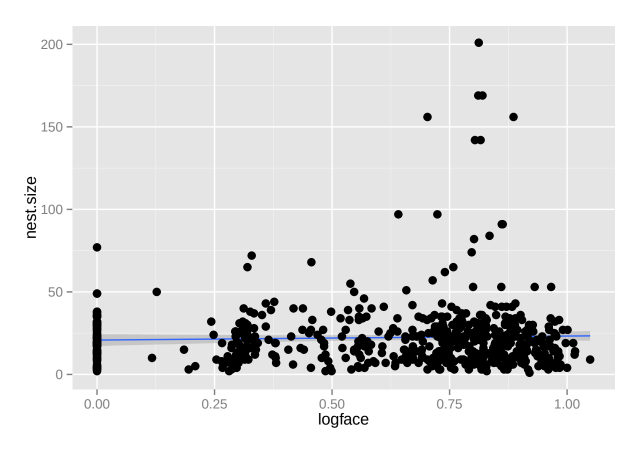


In ggplot, geoms, short for geometric objects, describe the type of plot you will produce. For example, geom_point is used for scatterplots. geom_point is the default in qplot if x and y are specified. If only x is specified, qplot defaults to geom_histogram. In the scatter diagrams above, the size argument in geom_point controls the size of data points.

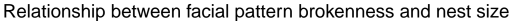
1.1.2 Fit a linear model

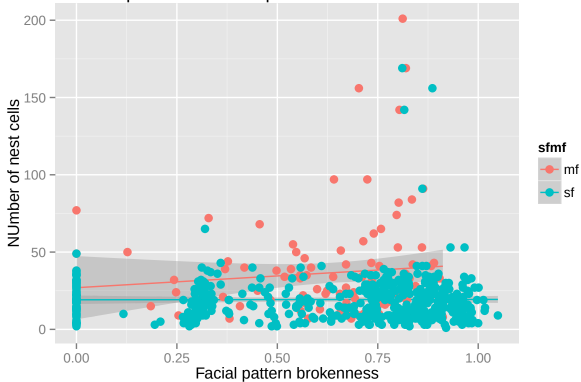
Add a smoothed line and fit linear models. 1m is used to fit linear models and carry out regressions. A typical 1m model has the form response ~ terms where response is the (numeric) response vector and terms specifies a linear predictor for response.

```
# Fit a linear model (by default includes 95% confidence region)
qplot(logface, nest.size, data=polistes.data, geom=c("point", "smooth"),
   method="lm") + geom_point(size=3)
```



```
# Create separate regressions for each factor and add labels
qplot(logface, nest.size, data=polistes.data, geom=c("point", "smooth"),
   method="lm", colour=sfmf,
   main="Relationship between facial pattern brokenness and nest size",
   xlab="Facial pattern brokenness",
   ylab="NUmber of nest cells") + geom_point(size=3)
```





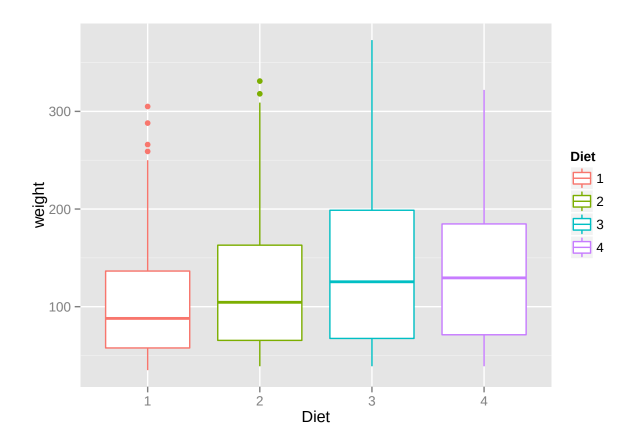
1.1.3 Boxplots and jittered points

```
# Load the ChickWeight dataset from the base R packages
# Results from an experiment on the effect of diet on early growth of chicks
cw <- ChickWeight
summary(cw)
```

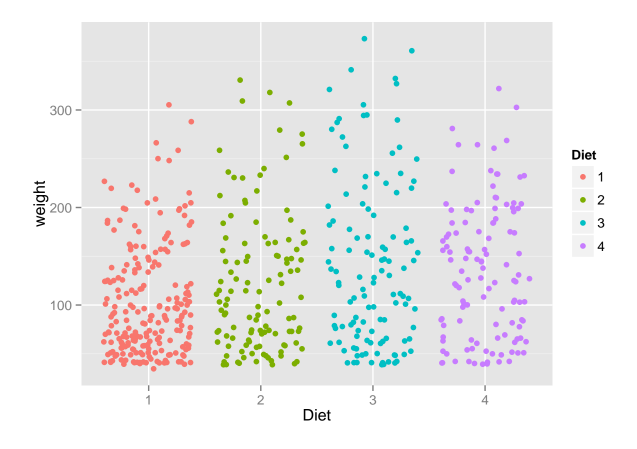
```
##
        weight
                         Time
                                        Chick
                                                   Diet
                                                   1:220
   Min. : 35.0
                    Min.
                           : 0.00
                                            : 12
   1st Qu.: 63.0
                    1st Qu.: 4.00
                                            : 12
                                                   2:120
##
                                    9
##
   Median :103.0
                    Median :10.00
                                    20
                                            : 12
                                                   3:120
##
   Mean
          :121.8
                    Mean
                           :10.72
                                    10
                                            : 12
                                                   4:118
   3rd Qu.:163.8
                    3rd Qu.:16.00
                                    17
                                            : 12
           :373.0
                           :21.00
                                            : 12
##
   Max.
                    Max.
                                    19
##
                                     (Other):506
```

```
tapply(cw$weight, cw$Diet, FUN=mean)
```

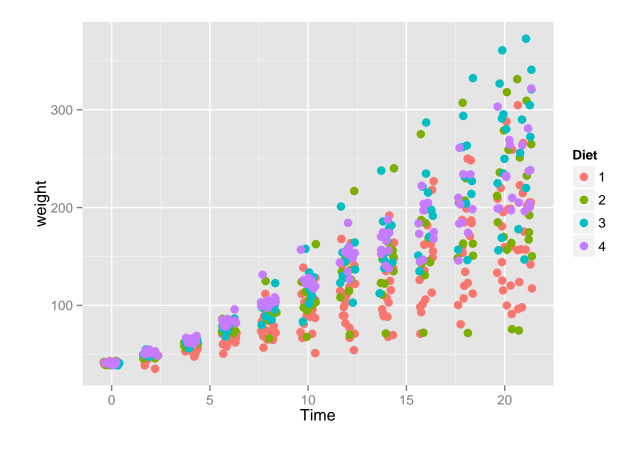
```
## 1 2 3 4
## 102.6455 122.6167 142.9500 135.2627
```



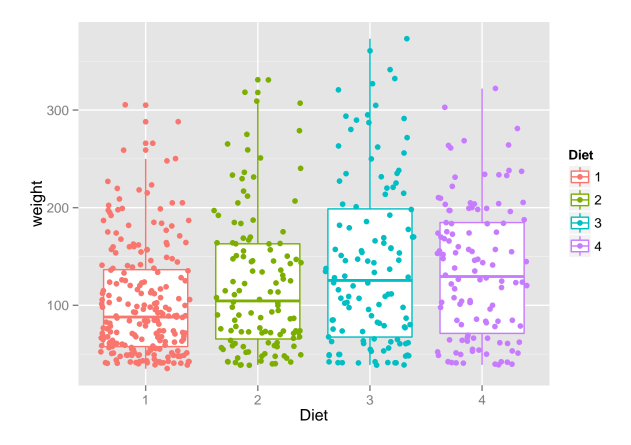
qplot(Diet, weight, data=cw, geom="jitter", colour=Diet)



Use I() to manually set the aesthetics, e.g., colour = I("red") or size = I(3) qplot(Time, weight, data=cw, geom="jitter", colour=Diet, size=I(3))



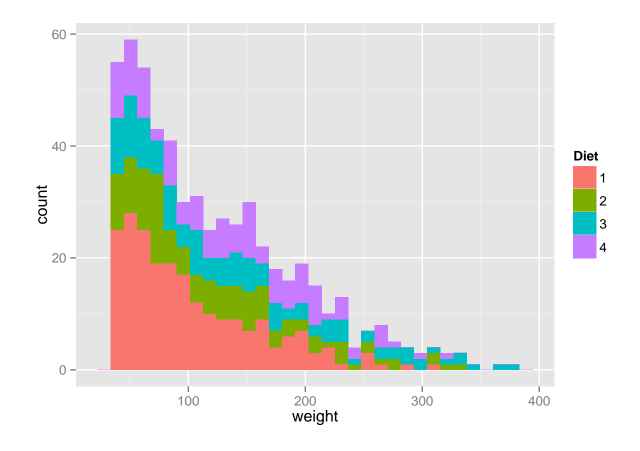
qplot(Diet, weight, data=cw, geom=c("boxplot","jitter"), colour=Diet)



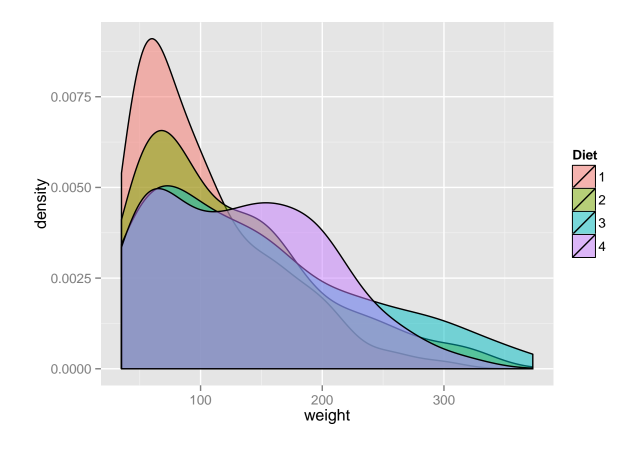
1.1.4 Histograms and density plots

```
qplot(weight, data=cw, geom="histogram", fill=Diet)
```

stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.



qplot(weight, data=cw, geom="density", fill=Diet, alpha=I(0.5))

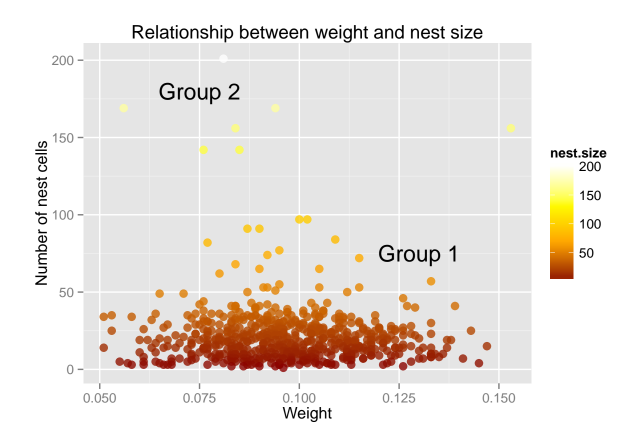


2 ggplot

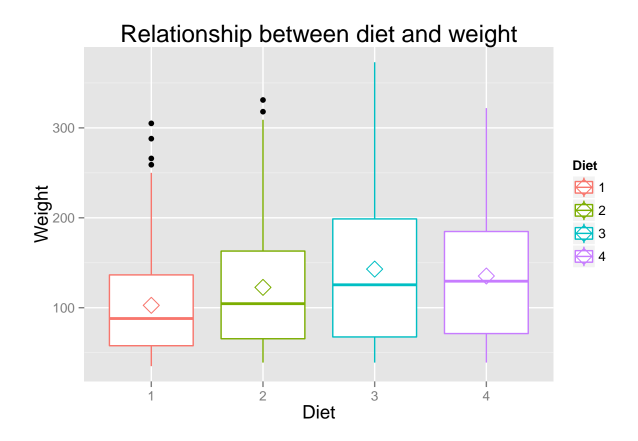
qplot does not show the power of ggplot. ggplot2 functions can be chained with "+" signs. All the options that can be chained are available at docs.ggplot2.org. Let us remake the previous graphs using a few of the wide variety of options available in ggplot.

2.1 Scatterplots

```
ggplot(polistes.data, aes(x=weight, y=nest.size, colour=nest.size)) +
    geom_point(size=3, alpha=.8) +
    # Add main and axis titles
labs(title="Relationship between weight and nest size",
        x="Weight", y="Number of nest cells") +
    # Add text to graph
    annotate("text", x=0.130, y=75, label="Group 1", size=6) +
    annotate("text", x=0.075, y=180, label="Group 2", size=6) +
    # Use a manually defined palette
    scale_colour_gradientn(colours=c("darkred", "orange", "yellow", "white"))
```

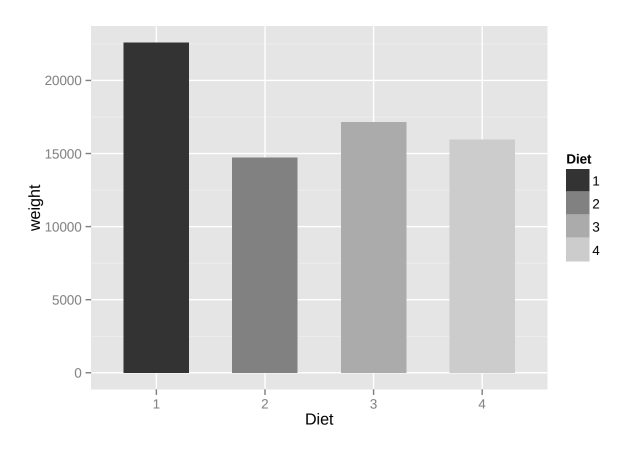


2.2 Boxplots

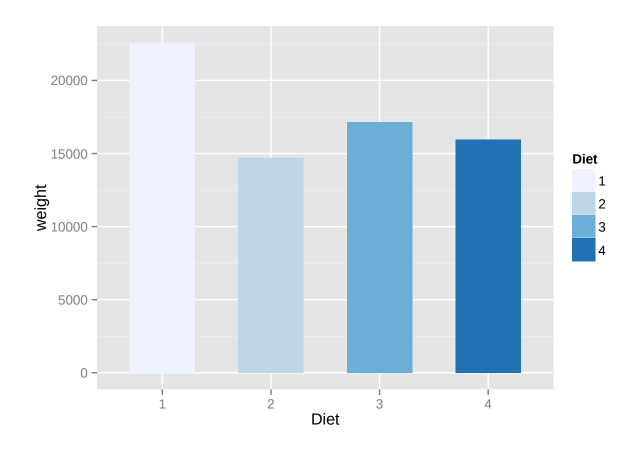


2.3 Bargraphs

```
# Use stat="identity" if you want the heights of the bars to represent
# values in the data
ggplot(cw, aes(x=Diet, y=weight, fill=Diet)) +
geom_bar(stat="identity", width=0.6) +
scale_fill_grey()
```

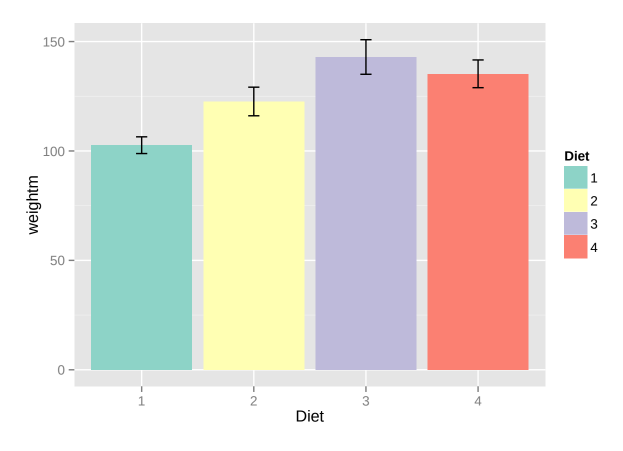


```
ggplot(cw, aes(x=Diet, y=weight, fill=Diet)) +
geom_bar(stat="identity", width=0.6) +
scale_fill_brewer(palette="Blues")
```

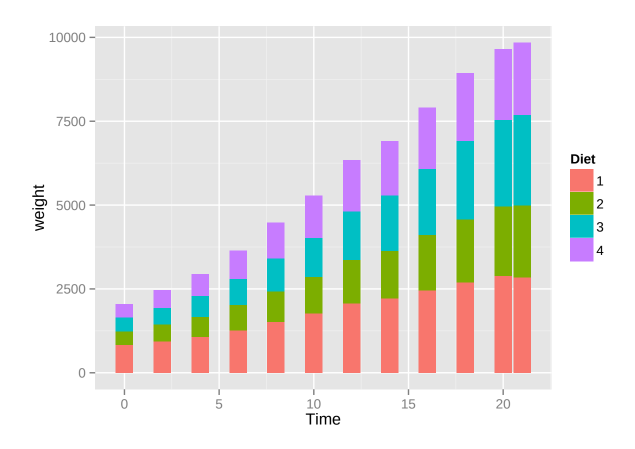


```
# Add error bars
\# Split data frame, apply function, and return results in a data frame
library(plyr)
cwse <- ddply(cw, "Diet", summarise,</pre>
  weightm=mean(weight, na.rm=TRUE),
  sd=sd(weight, na.rm=TRUE),
  n=sum(!is.na(weight)),
  se=sd/sqrt(n))
cwse
     Diet weightm
##
                         sd
                             n
## 1
        1 102.6455 56.65655 220 3.819784
## 2
        2 122.6167 71.60749 120 6.536840
        3 142.9500 86.54176 120 7.900146
## 3
        4 135.2627 68.82871 118 6.336197
## 4
cwse$Diet <- as.factor(cwse$Diet)</pre>
ggplot(cwse, aes(x=Diet, y=weightm, fill=Diet)) +
  geom_bar(stat="identity") +
  geom_errorbar(aes(ymin=weightm-se, ymax=weightm+se), width=.1) +
```

scale_fill_brewer(palette="Set3")

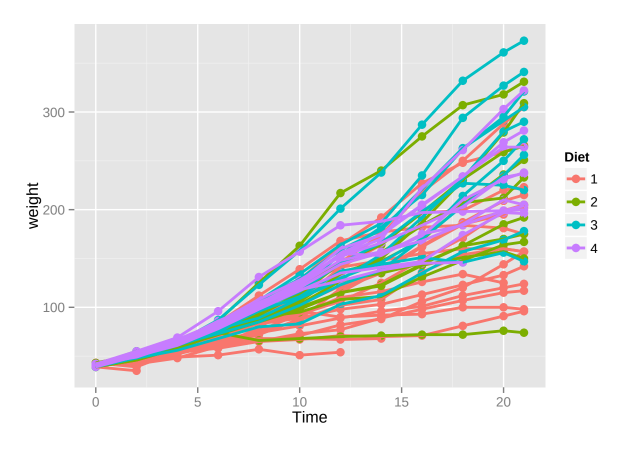


```
# Stacked bar graph
ggplot(cw, aes(x=Time, y=weight, fill=Diet)) +
geom_bar(stat="identity")
```



2.4 Line graphs and stacked area graphs

```
# If your line graph looks wrong, specify the grouping variable with `group`.
# Problems occur with line graphs because ggplot() is unable to determine
# how to group the variables
# A sawtooth pattern results from improper grouping
ggplot(cw, aes(x=Time, y=weight, group=Chick, colour=Diet)) +
    geom_point(size=3) + geom_line(size=1)
```



```
cwd12 <- subset(cw, Diet==1:2)
summary(cwd12)</pre>
```

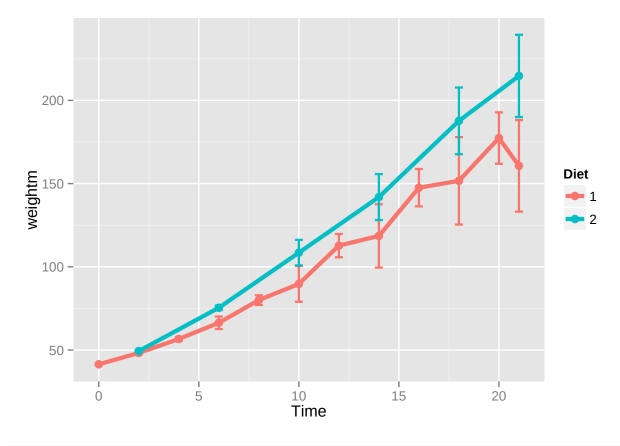
```
weight
                                       Chick
                                                 Diet
##
                        Time
   Min. : 39.0
                          : 0.00
##
                   Min.
                                   13
                                         : 6
                                                 1:110
   1st Qu.: 61.0
                   1st Qu.: 6.00
                                          : 6
                                                 2: 60
   Median : 95.5
                   Median :10.00
                                   20
                                          : 6
##
##
   Mean :111.3
                   Mean :10.86
                                   10
                                          : 6
   3rd Qu.:148.8
                   3rd Qu.:17.50
                                          : 6
##
                                   8
          :331.0
                          :21.00
                                   17
##
   Max.
                   Max.
                                         : 6
                                   (Other):134
##
```

```
cwse12 <- ddply(cwd12, c("Diet", "Time"), summarise,
  weightm=mean(weight, na.rm=TRUE),
  sd=sd(weight, na.rm=TRUE),
  n=sum(!is.na(weight)),
  se=sd/sqrt(n))
cwse12</pre>
```

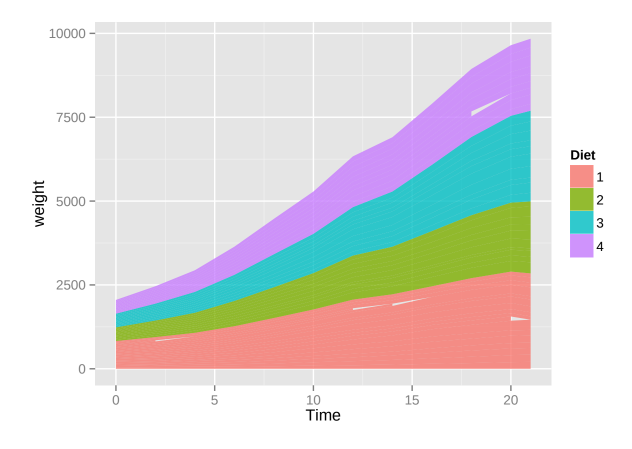
```
sd n
     Diet Time
                 weightm
                                             se
## 1
        1
             0 41.41667 1.164500 12
                                      0.3361622
## 2
             2 48.25000 2.549510 8
                                      0.9013878
## 3
             4 56.63636 3.722169 11
                                      1.1222763
## 4
             6 66.37500 10.595653 8 3.7461290
## 5
             8 80.00000 9.643651 11 2.9076701
        1
```

```
10 89.75000 30.471298 8 10.7732307
## 6
            12 112.72727 23.242594 11 7.0079058
## 7
## 8
            14 118.57143 50.345000 7 19.0286215
## 9
            16 147.54545 37.264899 11 11.2357899
## 10
            18 151.66667 64.301374
                                   6 26.2509259
## 11
            20 177.36364 51.252849 11 15.4533155
## 12
            21 160.66667 67.408209 6 27.5192862
        2
            2 49.40000 2.875181 10 0.9092121
## 13
## 14
        2
            6 75.40000 4.168666 10 1.3182480
## 15
        2 10 108.50000 24.295633 10 7.6829537
## 16
        2 14 141.90000 43.697063 10 13.8182247
        2
           18 187.70000 63.331667 10 20.0272315
## 17
## 18
            21 214.70000 78.138126 10 24.7094449
```

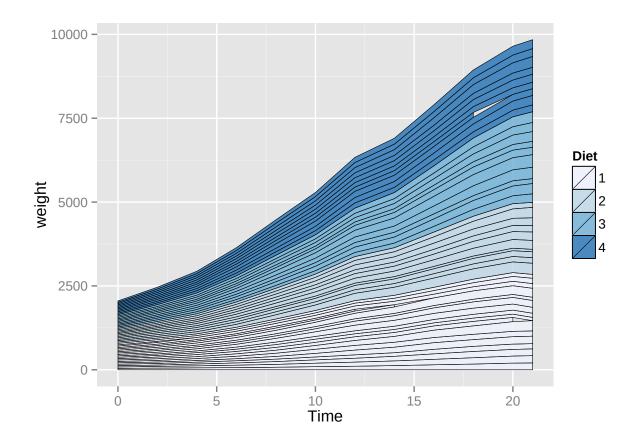
```
ggplot(cwse12, aes(x=Time, y=weightm, colour=Diet)) +
   geom_errorbar(aes(ymin=weightm-se, ymax=weightm+se), width=.4, size=0.8) +
   geom_line(size=1.5) +
   geom_point(size=3)
```



```
# Stacked area graph
# Area graphs represent cumulated totals over time
#tapply(cw$weight, cw$Diet, FUN=sum)
#cwd4 <- subset(cw, Diet==4)
#tapply(cwd4$weight, cwd4$Time, FUN=sum)
ggplot(cw, aes(x=Time, y=weight, group=Chick, fill=Diet)) +
    geom_area(alpha=0.8)</pre>
```



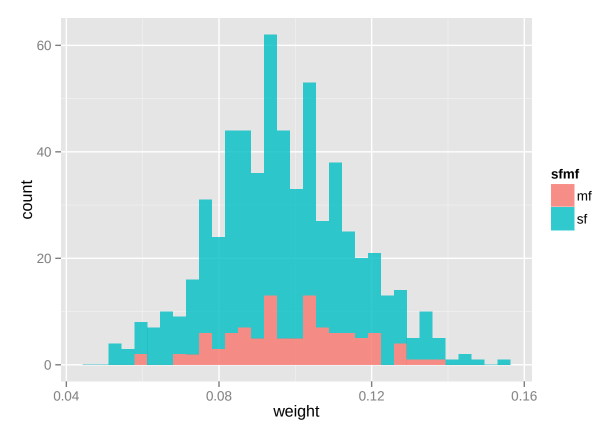
```
ggplot(cw, aes(x=Time, y=weight, group=Chick, fill=Diet)) +
  geom_area(colour="black", size=.2, alpha=.8) +
  scale_fill_brewer(palette="Blues")
```



2.5 Histograms

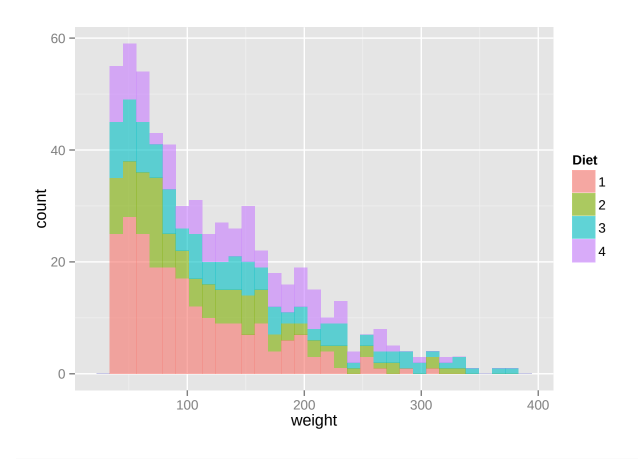
```
ggplot(polistes.data, aes(x=weight, fill=sfmf)) + geom_histogram(alpha=.8)
```

stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.



ggplot(cw, aes(x=weight, fill=Diet)) + geom_histogram(alpha=.6)

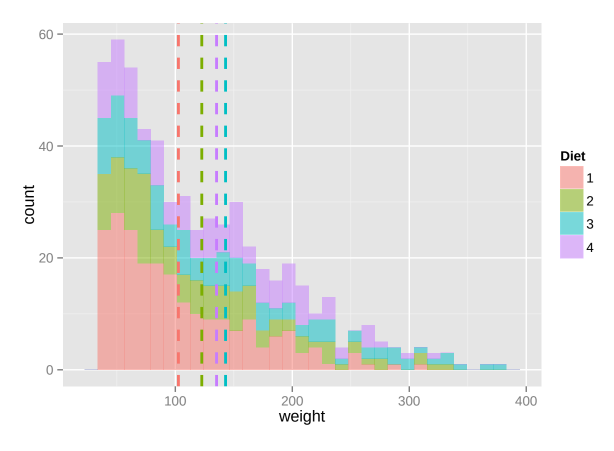
stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.



```
library(plyr)
# For each subset of a data frame, ddply applies a function and then combines
# results into a data frame
mwt <- ddply(cw, "Diet", summarise, weight.mean=mean(weight))</pre>
mwt
##
     Diet weight.mean
             102.6455
## 1
        1
## 2
        2
             122.6167
## 3
        3
             142.9500
             135.2627
## 4
        4
# Overlaid histograms with means
ggplot(cw, aes(x=weight, fill=Diet)) + geom_histogram(alpha=.5) +
    geom_vline(data=mwt, aes(xintercept=weight.mean, colour=Diet),
      linetype="dashed", size=1)
```

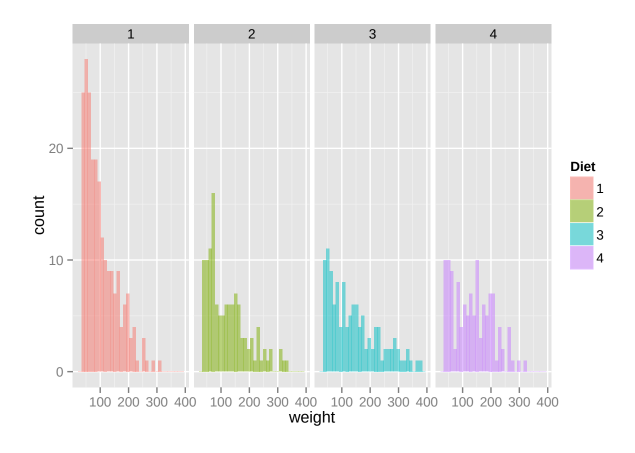
stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.

Find the mean of each group



```
# Use facets to display subsets of the dataset in different panels
ggplot(cw, aes(x=weight, fill=Diet)) + geom_histogram(alpha=.5) +
facet_grid(. ~ Diet)
```

```
## stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this. ## stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this. ## stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this. ## stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.
```

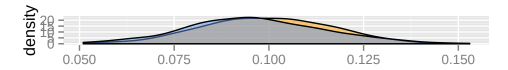


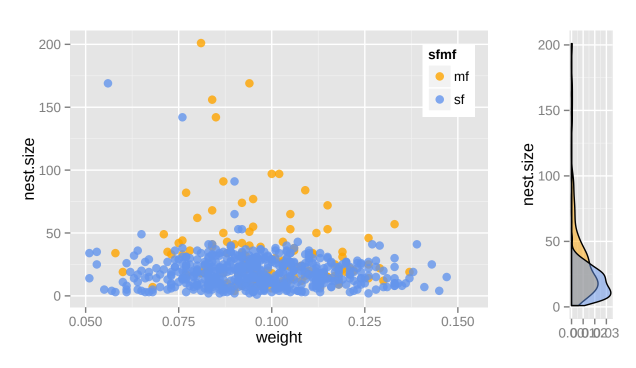
2.6 Scatterplots with marginal density plots

```
empty.plot <- ggplot() + geom_point(aes(1,1), colour="white") +</pre>
 theme(plot.background=element blank(),
    panel.grid.major=element_blank(),
    panel.grid.minor=element blank(),
    panel.border=element_blank(),
    panel.background=element_blank(),
    axis.title.x=element blank(),
    axis.title.y=element_blank(),
    axis.text.x=element_blank(),
    axis.text.y=element_blank(),
    axis.ticks=element_blank())
xyscatter <- ggplot(polistes.data, aes(x=weight, y=nest.size,</pre>
  colour=sfmf)) + geom_point(size=3, alpha=.8) +
  scale_color_manual(values=c("orange", "cornflowerblue")) +
  theme(legend.position=c(1,1), legend.justification=c(1,1))
xdensity.top <- ggplot(polistes.data, aes(weight, fill=sfmf)) +</pre>
  geom_density(alpha=0.5) +
  scale_fill_manual(values=c("orange", "cornflowerblue")) +
  theme(legend.position="none", axis.title.x=element_blank())
```

```
ydensity.right <- ggplot(polistes.data, aes(nest.size, fill=sfmf)) +
  geom_density(alpha=0.5) +
  coord_flip() +
  scale_fill_manual(values=c("orange", "cornflowerblue")) +
  theme(legend.position="none", axis.title.x=element_blank())

# Arrange the plots together
grid.arrange(xdensity.top, empty.plot, xyscatter, ydensity.right,
  ncol=2, nrow=2, widths=c(4, 1), heights=c(1, 4))</pre>
```



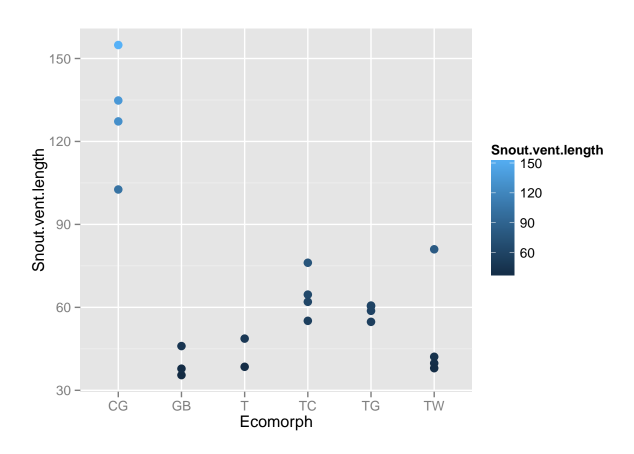


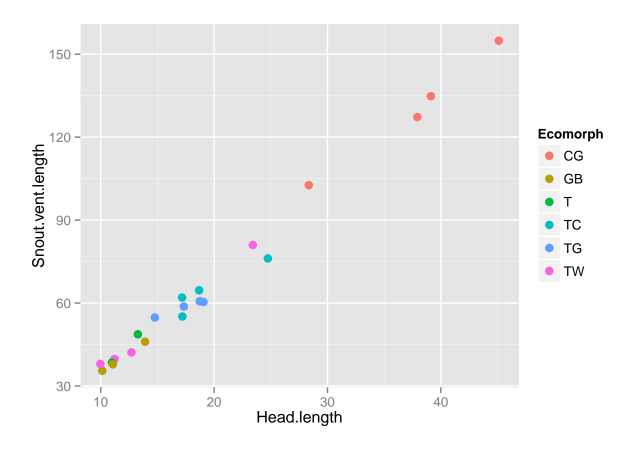
3 More ggplot and an introduction to rOpenSci packages

rOpenSci is an inititative to create R packages for accessing data repositories. The full list of packages is available here

```
# Here we will use the 'rdryad' package to retrieve a dataset from Dryad
# Retrieve the data using the Dryad identifier '10255/dryad.34389'
# which is at the end of the URL where the dataset is found
# http://datadryad.org/handle/10255/dryad.34389
# the original publication is Kolbe et al. 2011, Evolution 65(12): 3608-3624
anolis.data <- download_url("10255/dryad.34389")
anolis.data <- dryad_getfile(anolis.data)
# create plot with discrete and continous variables</pre>
```

```
anolis.scatter <- ggplot(anolis.data, aes(x=Ecomorph, y=Snout.vent.length,
   colour=Snout.vent.length)) + geom_point(size=3)
anolis.scatter</pre>
```





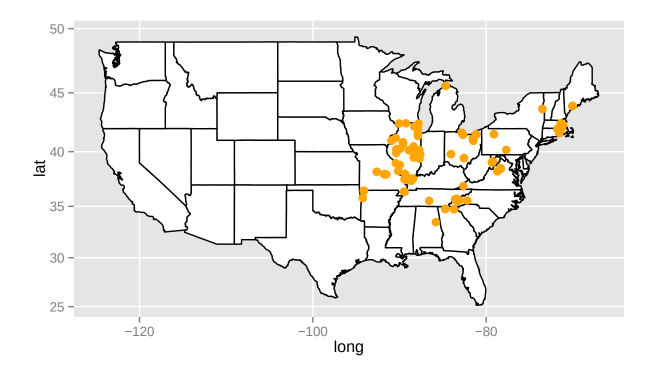
4 ggplot and rgbif

4.1 Plot GBIF occurrences of a species

```
# search for occurrences on GBIF
?occ_search
apicea.occ <- occ_search(scientificName="Aphaenogaster picea", limit=1000,
    return='data', hasCoordinate=TRUE)
# this dataset includes more than 100 columns
# show first 10 lines for columns 1 to 4
head(apicea.occ, n=10L)[,1:4]</pre>
```

##			name	key	decimalLatitude	decimalLongitude
##	1	Aphaenogaster	picea	899067422	41.98968	-71.78827
##	2	Aphaenogaster	picea	899067424	41.98968	-71.78827
##	3	Aphaenogaster	picea	899067434	41.98968	-71.78827
##	4	Aphaenogaster	picea	910640573	41.63587	-71.55863
##	5	Aphaenogaster	picea	910640541	41.63587	-71.55863
##	6	${\tt Aphaenogaster}$	picea	910640582	41.63587	-71.55863
##	7	${\tt Aphaenogaster}$	picea	910640546	41.63587	-71.55863
##	8	Aphaenogaster	picea	910640559	41.63587	-71.55863
##	9	Aphaenogaster	picea	891147998	36.89260	-82.62960
##	10	${\tt Aphaenogaster}$	picea	1142484688	34.76128	-84.70864

```
# select four columns from the complete data set
apicea.lat.lon <- apicea.occ[,c("name", "decimalLatitude", "decimalLongitude",
 "countryCode")]
# show the unique country codes in the data set
unique(apicea.lat.lon$countryCode)
## [1] "US" "MX"
# select only US occurrences
apicea.lat.lon <- subset(apicea.lat.lon, countryCode=="US")</pre>
summary(apicea.lat.lon)
##
                      decimalLatitude decimalLongitude countryCode
       name
## Length:300
                      Min.
                             :33.49 Min.
                                             :-94.25
                                                      Length:300
                                                     Class : character
## Class :character
                      1st Qu.:38.23 1st Qu.:-88.51
## Mode :character Median :40.11 Median :-83.89 Mode :character
##
                      Mean :39.79 Mean :-83.52
##
                      3rd Qu.:41.50 3rd Qu.:-79.32
                      Max. :45.57 Max. :-70.04
##
# get map data for world
#world map <- map data("world")</pre>
# show unique regions
#sort(unique(world_map$region))
# get US map data
states.map <- map_data("state")</pre>
head(states.map)
                   lat group order region subregion
##
         long
                        1 1 alabama
## 1 -87.46201 30.38968
                                                <NA>
## 2 -87.48493 30.37249
                                2 alabama
                                                <NA>
## 3 -87.52503 30.37249
                           1
                                3 alabama
                                                <NA>
## 4 -87.53076 30.33239
                                4 alabama
                                                <NA>
                           1
                                                <NA>
## 5 -87.57087 30.32665
                           1
                                 5 alabama
## 6 -87.58806 30.32665
                                 6 alabama
                                                <NA>
#states.map <- subset(states.map, long > -100 & lat < 50)
states.map <- ggplot(states.map, aes(x=long, y=lat, group=group)) +</pre>
 geom_polygon(fill="white", colour="black") + coord_map("mercator")
#states.map
states.map +
geom_point(aes(x=decimalLongitude, y=decimalLatitude, group=name),
 colour="orange", size=3, data=apicea.lat.lon)
```



```
#gbifmap(apicea.lat.lon)
```

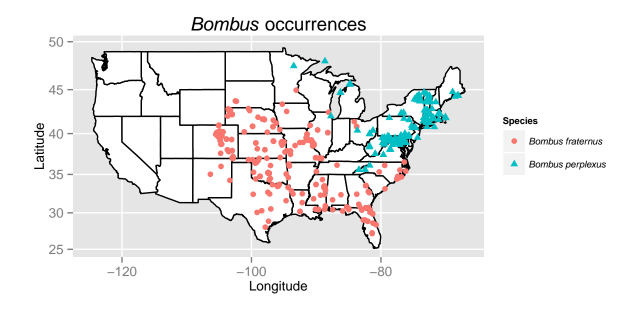
4.2 Plot GBIF occurrences of two or more species

```
## Length Class Mode
## 1340432 107 data.frame list
## 1340406 122 data.frame list
```

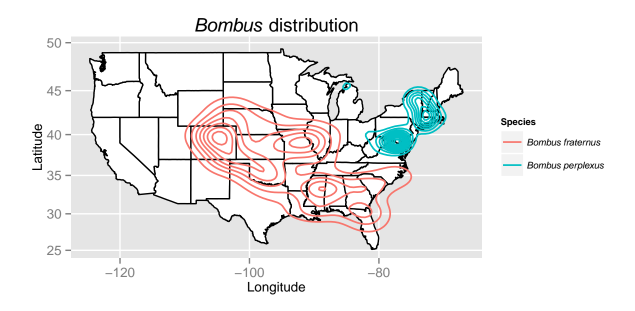
head(bombus.occ\$`1340432`, n=10L)[,1:4]

```
## 1 Bombus fraternus 1098920735 33.72666 -95.68521
## 2 Bombus fraternus 1143521876 33.53094 -95.91432
## 3 Bombus fraternus 899954560 32.87881 -93.81990
```

```
## 4 Bombus fraternus 923923296
                                         33.51451
                                                          -95.91873
                                                          -95.99989
## 5 Bombus fraternus 923923624
                                         33.48436
## 6 Bombus fraternus 923923680
                                         33.48885
                                                          -95.97723
## 7 Bombus fraternus 1065596357
                                         33.69256
                                                          -96.76983
## 8 Bombus fraternus 1143569911
                                         33.48602
                                                          -95.95263
## 9 Bombus fraternus 1024185501
                                         33.55605
                                                          -95.92782
## 10 Bombus fraternus 1024191102
                                                          -97.46752
                                         32.71433
bfraternus.lat.lon <- bombus.occ$`1340432`[,c("name", "decimalLatitude",
  "decimalLongitude", "countryCode")]
head(bombus.occ$`1340406`, n=10L)[,1:4]
##
                              key decimalLatitude decimalLongitude
                  name
## 1 Bombus perplexus 1143520430
                                         46.64574
                                                          -60.95197
## 2 Bombus perplexus 1143520142
                                         44.23823
                                                          -68.54755
## 3 Bombus perplexus 1143521611
                                         43.70909
                                                          -73.03349
## 4 Bombus perplexus 1143532609
                                         44.35068
                                                          -72.51181
## 5 Bombus perplexus 1122964996
                                                          -73.34100
                                         44.69461
## 6 Bombus perplexus 1135213313
                                         44.73630
                                                          -73.33179
## 7 Bombus perplexus 1143530758
                                         44.69111
                                                          -73.34442
## 8 Bombus perplexus 1135350545
                                         45.65765
                                                          -65.01472
## 9 Bombus perplexus 1143522783
                                         46.71553
                                                          -60.39149
## 10 Bombus perplexus 1136227380
                                         39.34410
                                                          -77.75340
bperplexus.lat.lon <- bombus.occ$`1340406`[,c("name", "decimalLatitude",
 "decimalLongitude", "countryCode")]
bperplexus.lat.lon <- subset(bperplexus.lat.lon, decimalLatitude < 50)</pre>
bombus.lat.lon <- rbind(bfraternus.lat.lon, bperplexus.lat.lon)
bombus.lat.lon <- subset(bombus.lat.lon, countryCode=="US")</pre>
states.map <- map_data("state")</pre>
#states.map <- subset(states.map, long > -120 & lat < 50)
states.map <- ggplot(states.map, aes(x=long, y=lat, group=group)) +</pre>
  geom_polygon(fill="white", colour="black") + coord_map("mercator")
#states.map
# create a title for applot that italicizes only the genus name
bombus.occ.title <- expression(paste(italic("Bombus"), " occurrences"))</pre>
bombus.map.occ <- geom_point(aes(x=decimalLongitude, y=decimalLatitude,</pre>
  group=name, colour=name, shape=name), size=2, data=bombus.lat.lon)
bombus.maplabels <- labs(title=bombus.occ.title, x="Longitude", y="Latitude",
  colour="Species", shape="Species")
bombus.maplegend <- theme(plot.title=element_text(face="italic",</pre>
  size=rel(1.2))) +
  theme(legend.title=element_text(size=rel(0.6))) +
  theme(legend.text=element text(face="italic", size=rel(0.6))) +
  theme(axis.title.x=element text(size=rel(0.8))) +
  theme(axis.title.y=element text(size=rel(0.8)))
states.map + bombus.map.occ + bombus.maplabels + bombus.maplegend
```

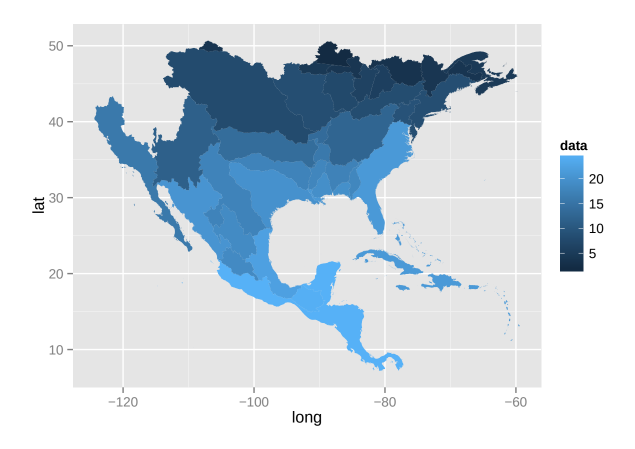


```
bombus.map.den <- stat_density2d(aes(x=decimalLongitude, y=decimalLatitude,
   group=name, colour=name), size=0.6, data=bombus.lat.lon, geom="density2d")
bombus.den.title <- expression(paste(italic("Bombus"), " distribution"))
bombus.maplabels <- labs(title=bombus.den.title, x="Longitude", y="Latitude",
   colour="Species", shape="Species")
states.map + bombus.map.den + bombus.maplabels + bombus.maplegend</pre>
```



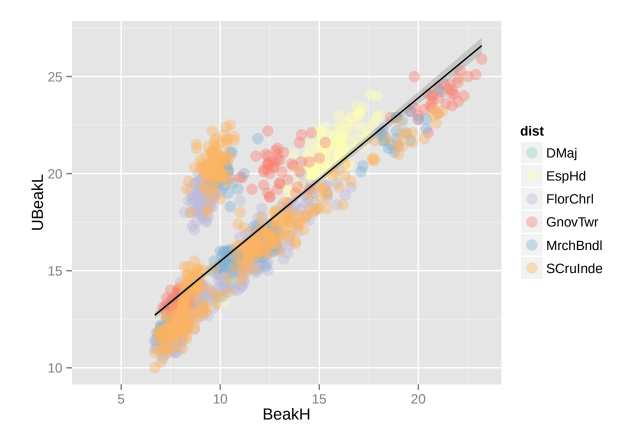
4.3 Map environmental data

```
# Create a directory for kml files
dir.create("~/Desktop/kmltemp")
options(kmlpath="~/Desktop/kmltemp")
#http://data.worldbank.org/developers/climate-data-api
#http://unstats.un.org/unsd/methods/m49/m49alpha.htm
#http://data.worldbank.org/sites/default/files/climate_data_api_basins.pdf
# Create a vector of World Bank basin IDs
nam <- 328:365
# Download map for vector of basins
nam.basin <- create_map_df(nam)</pre>
# Retrieve historical precipitation data
temp.dat <- get_historical_temp(nam, "decade")</pre>
# Create a subset using only data from one year
temp.dat <- subset(temp.dat, temp.dat$year==2000)</pre>
# Create maps of climate data
nam.map <- climate_map(nam.basin, temp.dat, return_map=TRUE)</pre>
nam.map
```



5 Other examples using morphological data for Darwin's finches

```
data(finch.ind)
tr <- traits.finch; sp <- sp.finch; dist <- ind.plot.finch</pre>
tr$sp <- sp; tr$dist <- dist</pre>
head(tr)
       WingL BeakH UBeakL N.UBkL
##
                                              sp dist
## 165
          69 10.8
                    15.2 10.6 Geospiza_fortis DMaj
                           9.8 Geospiza_fortis DMaj
## 166
          65 10.3
                     14.2
                           9.9 Geospiza_fortis DMaj
          65 10.0
                     13.6
## 167
## 168
          68
              9.5
                     13.5
                           9.9 Geospiza_fortis DMaj
## 169
                     15.0
                            10.7 Geospiza_fortis DMaj
          66 11.0
          68 10.4
## 170
                     15.0
                            10.2 Geospiza_fortis DMaj
ggplot(tr, aes(x=BeakH, y=UBeakL, colour=dist)) +
  geom_point(size=4, alpha=0.4) +
  stat_smooth(method=lm, colour="black", level=0.95) +
  scale_colour_brewer(palette="Set3")
```



```
mbh <- ddply(tr, "dist", summarise, bh.mean=mean(BeakH, na.rm=TRUE))
mbh</pre>
```

```
## dist bh.mean
## 1 DMaj 10.286047
## 2 EspHd 11.711345
## 3 FlorChrl 9.715032
## 4 GnovTwr 10.447945
## 5 MrchBndl 11.663877
## 6 SCruInde 9.890248
```

```
ggplot(tr, aes(x=BeakH, fill=dist)) + geom_density(alpha=0.4) +
scale_fill_brewer(palette="Set1") +
geom_vline(data=mbh, aes(xintercept=bh.mean, colour=dist),
linetype="dashed", size=1)
```

