ggplot2 Presentation - Computational Biology

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Preliminaries:

Create a Data set:

```
# create sample IDs
ID \leftarrow c(1:200)
# create a factor called origin
Origin <- c(rep("local", 100),</pre>
             rep("California", 100))
# create a factor called flower type
FlowerType <- rep(c(rep("clover",25),</pre>
                     rep("goldenrod",25),
                     rep("treefoil",25),
                     rep("mixed",25)),2)
# create a variable called mass
Mass \leftarrow c(rnorm(n = 100,
                 mean=32,
                 sd = 8), rnorm(n = 100,
                                  mean=21.
                                  sd=4))
# create a variable called Nosema Load
NosemaLoad <-c(rnorm(n = 100,
                      mean=100000,
                      sd = 80000), rnorm(n = 100,
                                       mean=500000,
                                       sd=40000))
# create a variable called varroa load
VarroaLoad \leftarrow c(rnorm(n = 100,
                         mean=5,
                          sd = 2), rnorm(n = 100,
```

```
mean=9,
sd=3))

# create a variable called time
Time <- rep(c(rep("Time1", 50), rep("Time2", 50)),2)

# create data frame
DF <- data.frame(ID, Origin, FlowerType, Mass, NosemaLoad, VarroaLoad, Time)</pre>
```

Basic Graphics in ggplot:

Summary stats using ddply:

Bar Plot

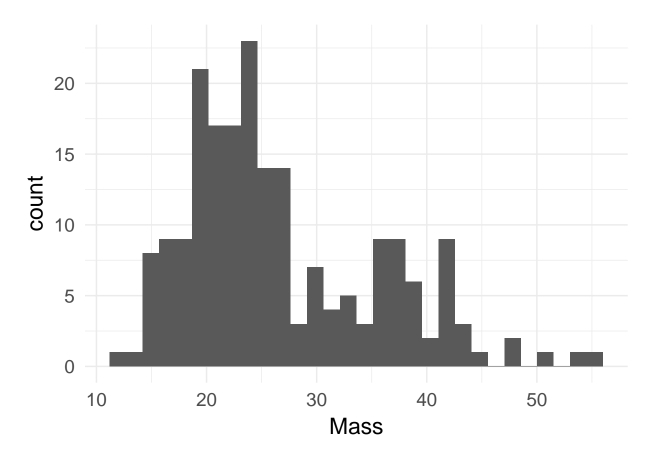


 $\# + coord_cartesian(ylim = c(0, 40))$

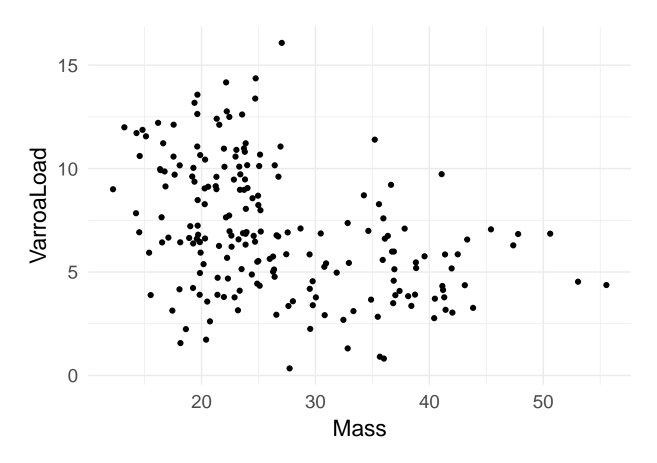
Histogram

```
plot2 <- ggplot(DF, aes(Mass))
plot2 + geom_histogram() + stat_bin(bins = 30) + theme_minimal(base_size = 17)</pre>
```

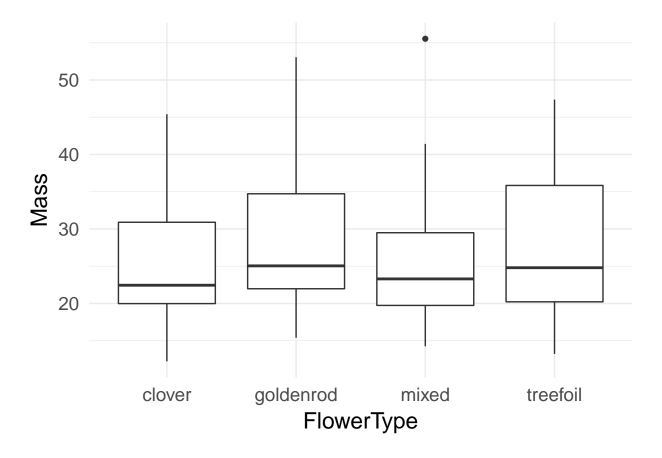
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



Scatterplot



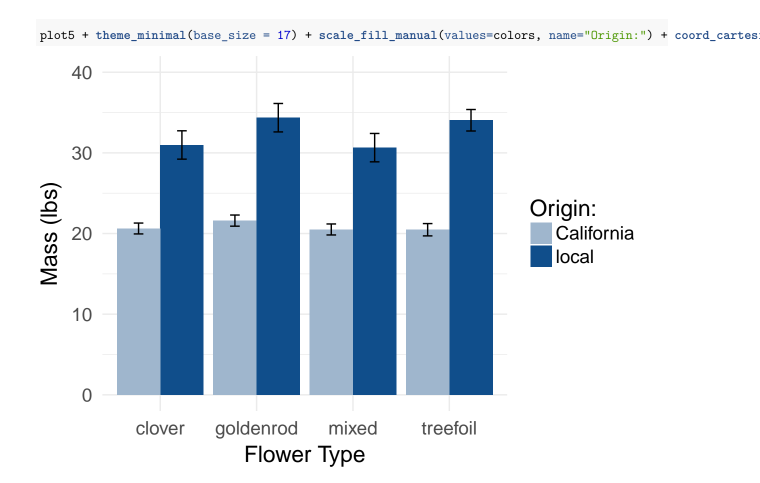
Boxplot



More Complicated Graphics in ggplot

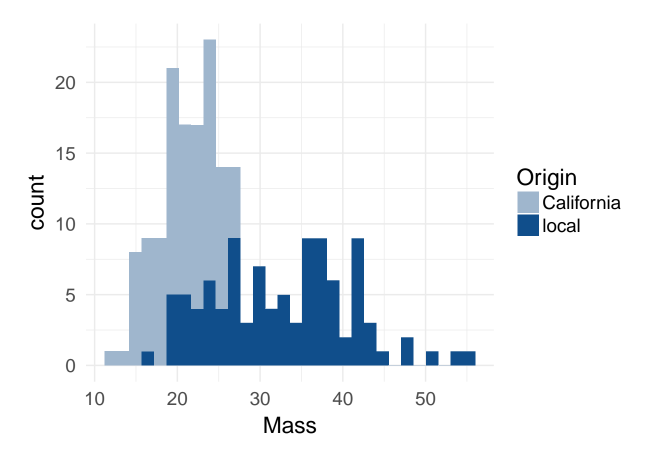
Summary stats using ddply:

Bar Plot



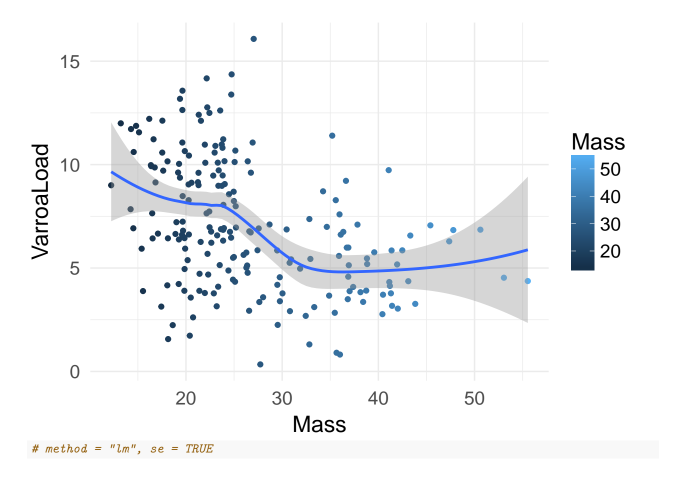
Histogram

```
plot6 <- ggplot(DF, aes(Mass, fill=Origin))
plot6 + geom_histogram() + stat_bin(bins = 30) + theme_minimal(base_size = 17) + scale_fill_manual(valu
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.</pre>
```

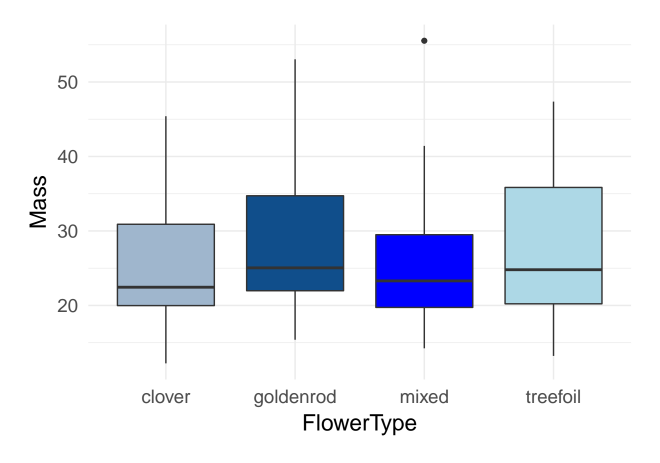


Scatterplot

`geom_smooth()` using method = 'loess'



Boxplot



Clean graphics for publication:

