Graphics with ggplot2

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Introduction



Today we will be covering some of the basics of how to use the R package **ggplot2**. The goal is to introduce you to some of the terminology used in **ggplot2** and then go through first some introductory and then more advanced examples to give you some exposure to the **ggplot2** style of coding.

We will spend some time preparing data for graphical display. In **ggplot2** there is a strong relationship between the dataset you are working with and the graphics you make. While you will see some code for data manipulation using packages **tidyr** and **dplyr**, we will not discuss it in detail as the focus today is on creating graphics.

Online resources

There are many resources for **ggplot2** help online. Once you get started making graphics, you will be able to find answers to many of questions you have through online searches. Here are a few websites that I have found to be most helpful.

- For nice coverage of some of the basics, check out the Cookbook for R website: http://www.cookbook-r.com/Graphs.
- All the functions, complete with examples, are on the **ggplot2** website: http://ggplot2.tidyverse.org/reference/.
- Folks on Stack Overflow answer questions about **ggplot2** code pretty much daily. You can look at (and search in) all the questions tagged with the [ggplot2] tag: http://stackoverflow.com/questions/tagged/ggplot2.
- The newer RStudio Community website, https://community.rstudio.com/, is another place to look for and ask for help that can be less intimidating than Stack Overflow.
- The active development of **ggplot2** is done on GitHub. To report bugs or check for updates, see the GitHub repository: https://github.com/tidyverse/ggplot2.

• In recent years, **ggplot2** has become easier to *extend* to make additional kinds of plots. A webpage that tracks packages that extend **ggplot2** to make other kinds of plots is here: https://exts.ggplot2.tidyverse.org/gallery/

Loading the package

The first thing we need to do is to load package **ggplot2**. If you are not working on a computer with a current version of **ggplot2**, which is version 3.3.0, you will need to install it first.

Check if your package version is up-to-date with packageVersion().

```
packageVersion("ggplot2")
```

```
[1] '3.3.0'
```

To install the package you can type install.packages("ggplot2") in your R Console or go the RStudio Packages pane, click Install, type the name of all the packages you want to install, and press enter.

Once the package you want to use is installed, you can load it into R via the library() function. Let's do that for ggplot2 now.

```
library(ggplot2)
```

Introduction to ggplot2 and exploratory graphics

The goal of the first part of the workshop is to show you some of the basic **ggplot2** syntax while making graphics. We will be making these relatively simple graphics using the built-in R dataset mtcars. Information about this dataset is available in the R help files. The variables we will be using from this dataset are:

```
mpg (miles per US gallon),
wt (car lbs/1000),
cyl (number of cylinders),
am (type of transmission), and
disp (engine displacement).
```

We will treat mpg as if it were our response variable of interest.

Let's take a quick look at the first six lines and structure of this dataset. You should recognize that cyl and am are both categorical variables. However, R reads them as numeric variables in the dataset because their categories are expressed as numbers.

head(mtcars)

```
mpg cyl disp hp drat
                                         wt qsec vs am gear carb
                21.0 6 160 110 3.90 2.620 16.46 0 1
Mazda RX4
                21.0 6 160 110 3.90 2.875 17.02 0 1
Mazda RX4 Wag
                                                               4
Datsun 710
                22.8
                      4 108 93 3.85 2.320 18.61 1 1
                                                          4
                                                              1
Hornet 4 Drive
                21.4 6 258 110 3.08 3.215 19.44 1 0
                                                          3
                                                              1
Hornet Sportabout 18.7
                       8 360 175 3.15 3.440 17.02 0 0
                                                          3
                                                               2
                       6 225 105 2.76 3.460 20.22 1 0
                                                          3
Valiant
                 18.1
                                                               1
```

str(mtcars)

```
'data.frame': 32 obs. of 11 variables:

$ mpg : num 21 21 22.8 21.4 18.7 18.1 14.3 24.4 22.8 19.2 ...

$ cyl : num 6 6 4 6 8 6 8 4 4 6 ...

$ disp: num 160 160 108 258 360 ...

$ hp : num 110 110 93 110 175 105 245 62 95 123 ...
```

```
$ drat: num 3.9 3.9 3.85 3.08 3.15 2.76 3.21 3.69 3.92 3.92 ...
$ wt : num 2.62 2.88 2.32 3.21 3.44 ...
$ qsec: num 16.5 17 18.6 19.4 17 ...
$ vs : num 0 0 1 1 0 1 0 1 1 1 ...
$ am : num 1 1 1 0 0 0 0 0 0 0 ...
$ gear: num 4 4 4 3 3 3 3 4 4 4 ...
$ carb: num 4 4 1 1 2 1 4 2 2 4 ...
```

Defining the dataset in ggplot()

Let's start by diving right into the ggplot() function. When building a plot, ggplot() is going to be your first line of code.

Defining the dataset is an important part of the ggplot() philosophy. It allows us to refer to columns in the dataset directly by name. A common mistake for new ggplot2 users is to use dollar sign notation to refer to column names (e.g., mtcars\$mpg); try to avoid this.

The dataset is the first argument in ggplot(). If we only define the dataset, we get a completely blank graph. By default the plot background is grey.

```
ggplot(data = mtcars)
```

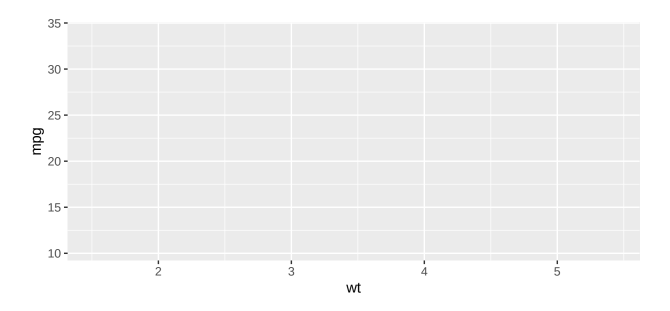
Defining the variables for the axes

A standard way to use ggplot() is to define the dataset and the axis variables within the ggplot() function. The axis variables are defined within the aes() function inside ggplot(). The aes() stands for aesthetics, which we will be talking more about in a few minutes.

Here is how we use ggplot() to define the dataset and axes. We will now get a blank graph with axes, since we haven't chosen a plot type yet.

Now you can see the default plot background, grey with white grid lines

```
ggplot(data = mtcars, aes(x = wt, y = mpg) )
```



Adding a geom layer to make a plot

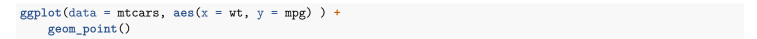
In **ggplot2** if we are talking about different *geoms* (i.e., *geometric objects*) we are talking about different types of plots. Different geom functions is how we ask for a certain kind of plot. We add these geom functions as *layers* to our initial **ggplot()** call.

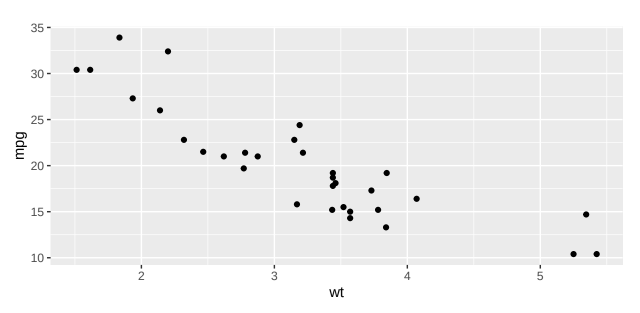
Scatterplot

We add layers by using whatever geom function we want along with the + sign. In this case we are going to start by making a scatterplot of mpg vs wt, so we will add the geom_point() layer to our ggplot() call.

It is standard coding practice to put new layers on new lines. This makes the code more readable, which is important when a collaborator (such as your future self) needs to understand what you did. Putting spaces in your code, much like you would while typing a sentence, also makes your code more readable.

Here is a scatterplot we get using geom_point(). The data are now plotted as points on the default background.

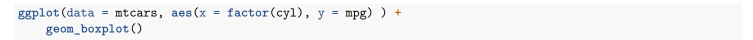


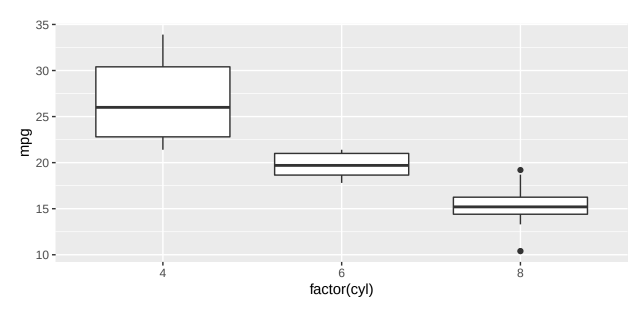


Boxplot

Now let's make boxplots of mpg for each level of the categorical variable cyl. The x axis variable is usually going to be categorical when making boxplots, so we have to change cyl to categorical using factor(cyl) (remember that cyl is numeric in mtcars).

We again define the dataset and axis variables within ggplot(), but this time we add the boxplot geom, geom_boxplot(), instead of geom_point().





Mapping aesthetics to variables

Package **ggplot2** does not support 3D graphics. However, you can display more than two dimensions in a plot by assigning variables in the dataset to colors, shapes, line types, etc. These are all examples of *aesthetic attributes*. An aesthetic attribute is a visual property that affects the way observations are displayed in the plot. The x and y positions are aesthetic attributes.

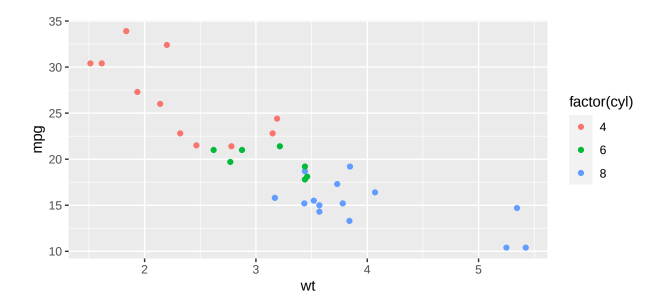
Assigning a variable from the dataset to an aesthetic is always done within the aes() function. A term commonly used for the process of assigning variables to aesthetics is aesthetic mapping. We'll use this term throughout the rest of the workshop.

Let's make our scatterplot again, but this time we will map a different color to points for cars in different cylinder categories. We'll do this by mapping the color aesthetic to the cyl variable. Again we will need to use factor(cyl) to indicate that cyl should be considered categorical instead of continuous.

One of the convenient things about **ggplot2** is that we when map aesthetics to variables we get legends automatically. When making quick exploratory graphics I always use the default colors that **ggplot2** chooses. Later today we'll see that we can change these defaults by using some of the **scale_*()** functions, which is an important part of making a final version of a graph.

You can see I've stopped writing out the data argument, since the dataset is always the first argument in ggplot().

```
ggplot(mtcars, aes(x = wt, y = mpg, color = factor(cyl) ) ) +
   geom_point()
```



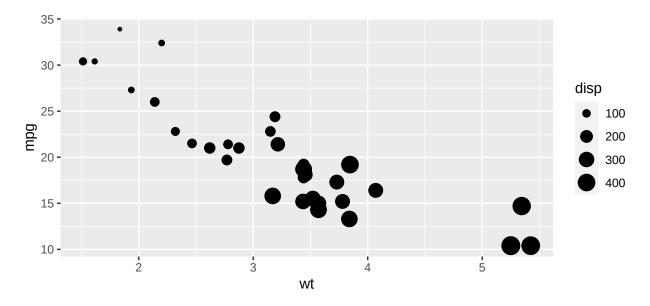
Continuous variables and aesthetic mapping

In the plots we are making today we will be only mapping categorical variables to aesthetics. You can map some aesthetics to continuous variables, though. For example, heat maps and bubble plots are created by mapping the aesthetics color and size, respectively, to continuous variables.

Note that some aesthetics can never be used with continuous variables. For example, shape and linetype can only be used with categorical variables as those aesthetics don't have a continuous nature. You will get an error message if you attempt to use continuous variables with either of those.

To see how aesthetic mapping with a continuous variable works, we'll make a bubble plot by mapping the size aesthetic to the continuous variable disp in our scatterplot. As always, the aesthetic mapping takes place inside aes().



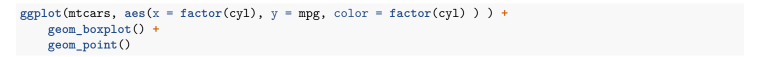


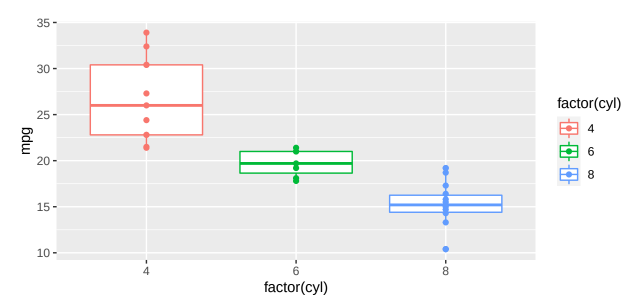
Adding more layers

Not only can we add additional aesthetics to a plot, we can also add additional geoms to a plot by adding more layers. This gives us a lot of freedom to make really informative graphics.

Let's add the data points on top of the boxplots we made by adding a geom_point() layer. In addition, let's map color to the number of cylinders. We put the aesthetic mapping in ggplot(), which causes both the boxplots and points to be colored by cyl and put into the legend. This is called mapping an aesthetic globally, as it affects all layers in the plot.

Notice that when we use color for plots such as boxplots, the color changes the outline color and not the color of the inside of the box. We need the fill aesthetic for changing the color of the inside of the boxes.





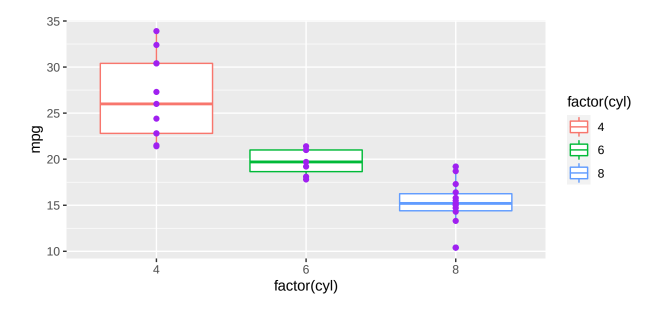
Setting aesthetics to constants within layers

We can add arguments to individual geom layers. In the next graphic, we will map the colors with cyl again for the boxplots, but we will set the color of the points to purple. We can do this by adding the color argument to geom_point(). When we add an aesthetic to a geom that we've already defined within ggplot(), the global ggplot() aesthetic gets overridden for that geom only.

It is important to recognize that the argument within <code>geom_point()</code> sets the aesthetic to a constant value (purple) instead of mapping to a variable. This means that we use <code>color</code> outside of the <code>aes()</code> function for the first time. When we set aesthetics to constants the legends are not affected.

Knowing when to define aesthetics like color inside or outside the aes() function can be confusing when you first start working with ggplot2. Generally speaking, if you are mapping an aesthetic to a variable from your dataset you do it inside aes() but if you are setting an aesthetic to a constant value you do it outside aes().

```
ggplot(mtcars, aes(x = factor(cyl), y = mpg, color = factor(cyl))) +
    geom_boxplot() +
    geom_point(color = "purple")
```



Mapping aesthetics separately for different geoms

So far we've been using aesthetic mapping only within ggplot(). We can map aesthetics to variables within specific geoms, as well.

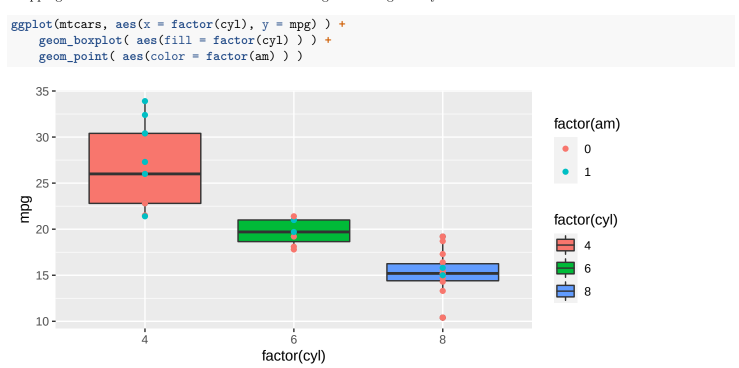
As I mentioned earlier, when we map aesthetics in ggplot() they are global, so those aesthetics mappings are used in all the layers throughout the graphic. We just saw that we can override global aesthetics within a geom if we need to.

There are times when making complicated graphics that we might not want to map aesthetics globally because we'd have to override them in too many subsequent layers. In this case, we'd do the aesthetic mapping within specific geom layers instead.

In the next graph we will map the fill color for the boxplots to cyl and the color of the points to am (transmission type). We don't want to color the boxplot outline by am, though, so we map the aesthetics separately within the geom layers rather than globally in ggplot(). Notice we do the mapping within the aes() function still, but we did it within each geom layer instead of in ggplot().

In this example, x and y are mapped globally while fill is mapped to cyl only for the boxplot layer and color is mapped to am only for the points layer.

Mapping aesthetics to different variables means we'll get more legends by default.



Edit the dataset to change the graphic

This is a good place for us to talk a bit about the relationship between the dataset and the graphic. In the philosophy behind **ggplot2**, the dataset and the graphic go hand in hand. Making changes to the dataset can be the most straightforward way to make changes to the resulting graphic. The answer to the common help forum question "How do I change the appearance of xxx in **ggplot2**?" is very often "Make changes to your dataset."

In the example dataset we've been using, the two categorical variables we've used are considered numeric in the mtcars dataset so we've had to define them as factors every time we use them. In addition, the levels of the am variable are not useful for discerning which color represents an automatic vs manual transmission and so that legend is particularly uninformative.

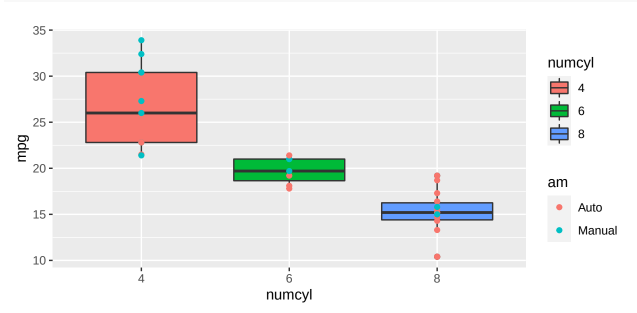
Let's take a moment to go back to the mtcars dataset to define cyl and am as factors and make informative names for the levels of am (remember 0 in am stands for an automatic transmission). We'll call the new cylinder variable numcyl.

```
mtcars$numcyl = factor(mtcars$cyl)
mtcars$am = factor(mtcars$am, labels = c("Auto", "Manual") )
str(mtcars)
```

```
'data.frame':
               32 obs. of 12 variables:
        : num
               21 21 22.8 21.4 18.7 18.1 14.3 24.4 22.8 19.2 ...
$ mpg
               6 6 4 6 8 6 8 4 4 6 ...
$ cyl
        : num
        : num 160 160 108 258 360 ...
$ disp
$ hp
        : num
               110 110 93 110 175 105 245 62 95 123 ...
               3.9 3.9 3.85 3.08 3.15 2.76 3.21 3.69 3.92 3.92 ...
$ drat
        : num 2.62 2.88 2.32 3.21 3.44 ...
$ wt
  qsec : num 16.5 17 18.6 19.4 17 ...
  VS
        : num 0 0 1 1 0 1 0 1 1 1 ...
        : Factor w/ 2 levels "Auto", "Manual": 2 2 2 1 1 1 1 1 1 1 ...
$
       : num 4 4 4 3 3 3 3 4 4 4 ...
  gear
$ carb : num 4 4 1 1 2 1 4 2 2 4 ...
$ numcyl: Factor w/ 3 levels "4","6","8": 2 2 1 2 3 2 3 1 1 2 ...
```

After those changes, let's recreate the last graphic we were working on using the edited variables.

```
ggplot(mtcars, aes(x = numcyl, y = mpg) ) +
    geom_boxplot( aes(fill = numcyl) ) +
    geom_point( aes(color = am) )
```



Choosing colors for color and fill

We won't be working with colors when we switch from exploratory graphics to creating polished graphics, so I thought I'd give an example here of changing the color scheme away from the default colors. You can change the *values* for any aesthetic by using the appropriate *scale* function. For example, to change the fill and color colors away from the defaults, we can use scale_fill_manual() and scale_color_manual(), respectively.

We change the default colors to the desired colors using the values argument. New colors are assigned in the order of the factor variable levels that we mapped the aesthetic to. The levels of the numcyl variable are ordered 4, 6, 8 and the levels of the am variable are Auto, Manual. See the limits argument for changing the order of the factor levels within the scale functions (we will not cover this today).

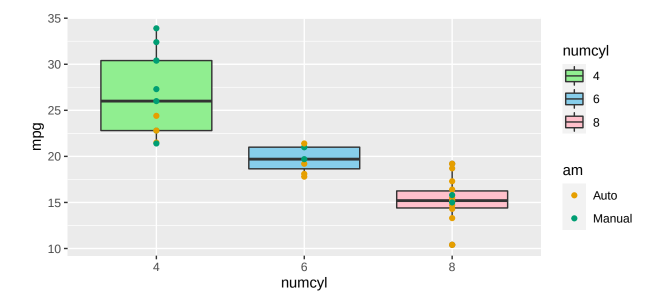
There is some nice information about available colors and color palettes at the Cookbook to R site, http://www.cookbook-r.com/Graphs/Colors_(ggplot2)/ and package colorspace overview, http://colorspace.r-forge.r-project.org/articles/hcl_palettes.html. You can set some nice default color schemes by using scale_color_viridis_d() instead of manually choosing colors.

Let's change the fill colors to light green, sky blue, and pink using color names, and the point colors to a green and an orange using hexadecimal string names I pulled out of the Cookbook to R link I gave above. If you don't want to rely on the factor order for setting colors to each factor level you can give named color vectors, shown within scale_color_manual() below.

You'll see the colors change both within the graph and in the legend.

Important: Make sure you use the correct scale_*() function for the aesthetic you are using in your graphic. If you map a variable to fill, scale_color_manual() won't change the fill colors.

```
ggplot(mtcars, aes(x = numcyl, y = mpg) ) +
    geom_boxplot(aes(fill = numcyl) ) +
    geom_point(aes(color = am) ) +
    scale_fill_manual(values = c("light green", "sky blue", "pink") ) +
    scale color manual(values = c(Manual = "#009E73", Auto = "#E69F00") )
```



Dot plot example

There are many, many other geoms available in **ggplot2**, far too many to cover today. Let's talk about a few of the ones we commonly use in data exploration.

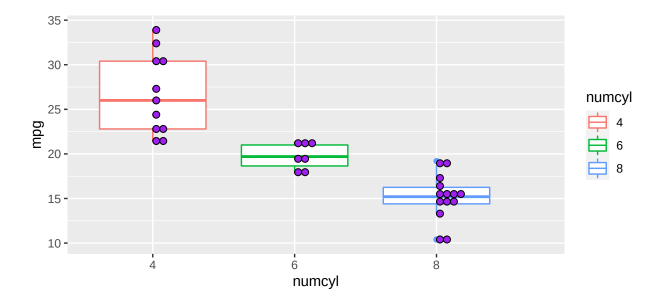
One useful graphic not everyone has seen before is the dot plot. Dot plots are essentially a type of histogram made using dots. This is an alternative way to show the raw data values along with a boxplot (which shows summary statistics).

Let's put a dot plot on top of our boxplot using geom_dotplot(). Here we need to bin along the y axis, so this will look like a histogram turned on its side. You always get a message about the binwidth for dot plots and histograms. This is information about the plot, and does not indicate an error.

We'll set the dot fill color to purple, which is done outside of aes() since this involves setting an aesthetic to a constant value. In dot plots, color changes the outline color of the dots and fill changes the inside color.

```
ggplot(mtcars, aes(x = numcyl, y = mpg) ) +
    geom_boxplot(aes(color = numcyl) ) +
    geom_dotplot(fill = "purple", binaxis = "y")
```

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

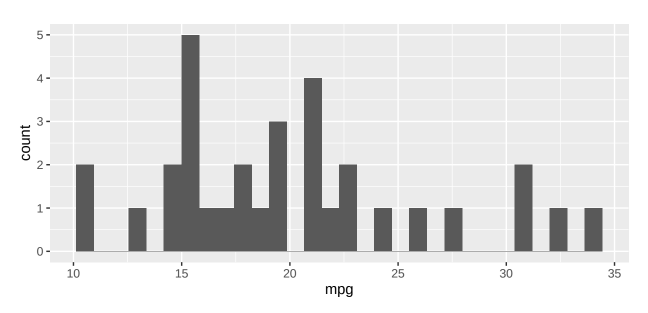


Histograms and density plots

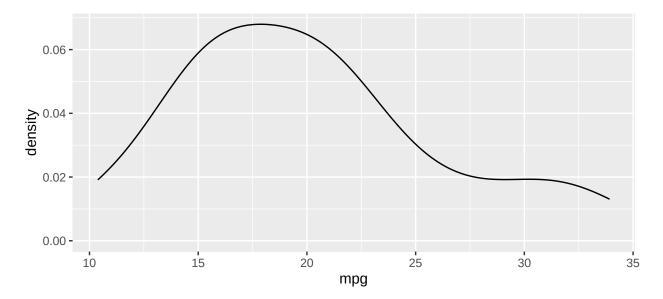
Histograms and density plots are both commonly used when making exploratory graphics of single variables. You can think of a density plot of a sort of smoothed histogram (although this is not the formal definition).

We only need to define the x axis for these plots, as the y axis is either a count or a density and **ggplot2** calculates those for us. Let's make a histogram and then a density plot of mpg.

```
ggplot(mtcars, aes(x = mpg) ) +
   geom_histogram()
```

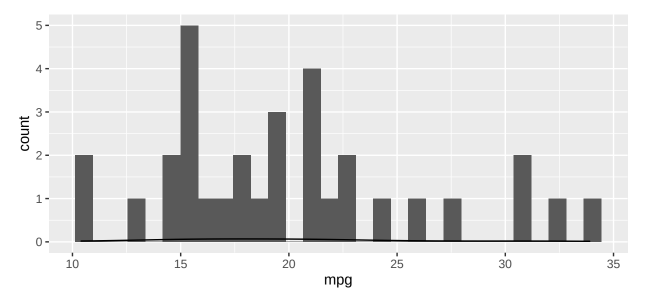


```
ggplot(mtcars, aes(x = mpg) ) +
  geom_density()
```



A next step might be to make a histogram with a density overlay. We need to be careful with this, though. Look at what happens when I try to add geom_density() on top of geom_histogram(); we can barely see the density line.

```
ggplot(mtcars, aes(x = mpg) ) +
    geom_histogram() +
    geom_density()
```

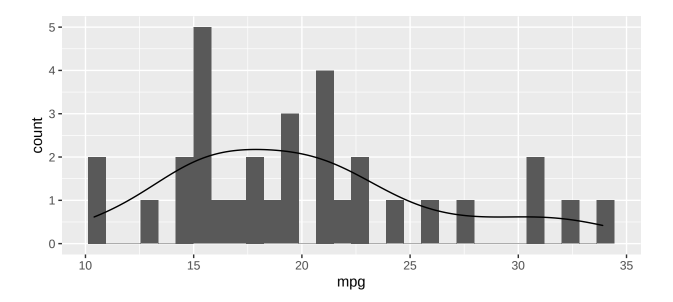


By default, histograms and density plots are on different scales. If we want both on the same plot we'd need to get both layers on the same scale, showing either counts or densities. The histogram and density geoms compute count and density variables, among others, that we can use for this. The variables computed by each geom are listed in the "Computed variables" section of the documentation. Follow this link for the computed variables from geom_histogram().

We can map our y variable to one of these special variables via after_stat(). As this involves aesthetic mapping, we have to do this inside aes().

Let's put the density curve on the scale of counts and try the plot again.

```
ggplot(mtcars, aes(x = mpg) ) +
   geom_histogram() +
   geom_density(aes(y = after_stat(count) ) )
```

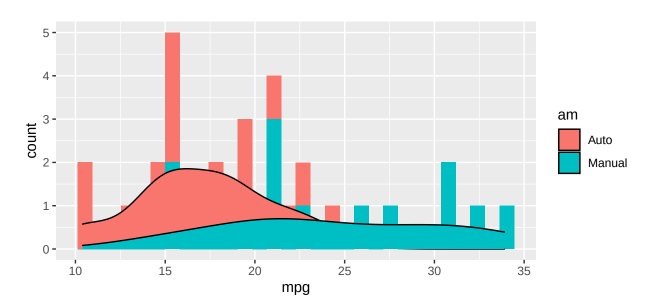


Layer order

The order we add the geom layers matters. Look what happens when we map fill to am for all the layers and change the order in which we add the histogram and density layers. The layer added last is always on top.

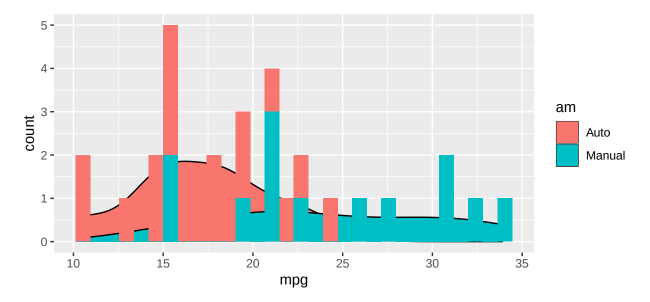
Here the density layer comes last so it is on top of the histogram.

```
ggplot(mtcars, aes(x = mpg, fill = am) ) +
    geom_histogram() +
    geom_density(aes(y = after_stat(count) ) )
```



But if we use the histogram layer last the histogram is plotted on top of the density plot.

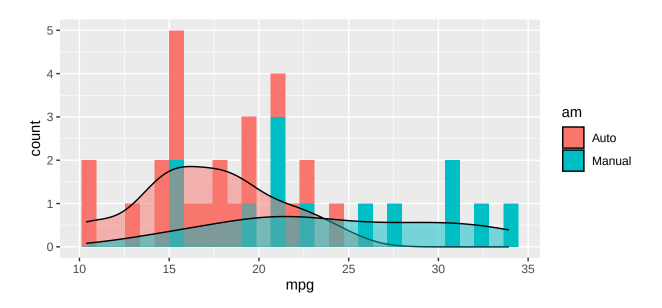
```
# Change the order of the geom layers
ggplot(mtcars, aes(x = mpg, fill = am)) +
    geom_density(aes(y = after_stat(count))) +
    geom_histogram()
```



These filled graphics are a little difficult to read. The graphic become slightly more useful if we allow the color of the fill to be more transparent in the density layer. We control transparency using the alpha aesthetic. We can set alpha to be between 0 and 1, with 0 being completely transparent and 1 completely opaque.

We will set alpha to a constant inside geom_density(). Because we are not aesthetic mapping, we will do this outside of aes(). You can map the alpha aesthetic to variables, although we won't do this today.

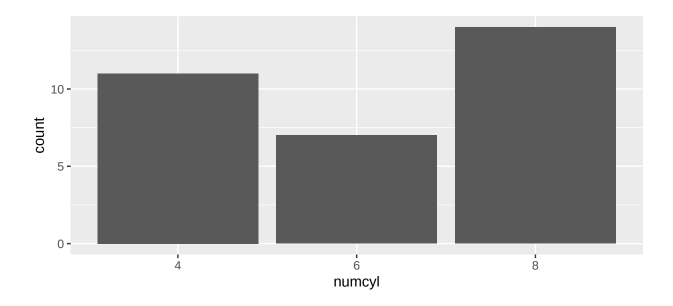
```
ggplot(mtcars, aes(x = mpg, fill = am)) +
    geom_histogram() +
    geom_density(aes(y = after_stat(count)), alpha = .5)
```



Bar graphs

Bar graphs are closely related to histograms, but involve graphing counts of a categorical variable instead of binning a continuous variable. You can map values other than counts to the bars using the y aesthetic, but geom_bar() defaults to counts. See geom_col() for a bar chart mapping y to a variable instead of graphing the count per group.

```
ggplot(mtcars, aes(x = numcyl) ) +
  geom_bar()
```



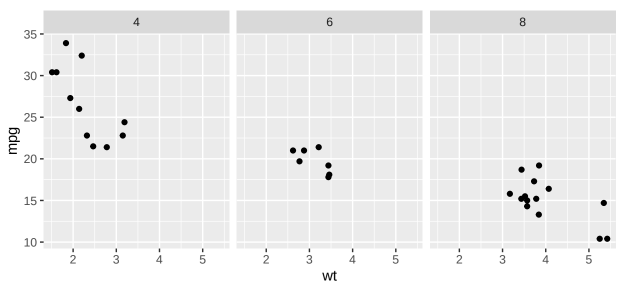
Facets

Sometimes we want to create plots for different groups in different panes within the same graphic. In **ggplot2** lingo this is called *faceting*. There are two faceting functions, **facet_wrap()** and **facet_grid()**. They actually can be quite different, so if you're doing faceting you should review the help page examples for each.

Today I'll demonstrate a basic plot using facet_wrap(). While there's a ton you can do with this function, I won't get into any of the bells and whistles here.

Inside facet_wrap() we list the variable that contains the groups that we want to plot as separate panels after the tilde, ~. Here is the mpg vs wt scatterplot, with a separate plot for each level of numcyl.

```
ggplot(mtcars, aes(x = wt, y = mpg) ) +
    geom_point() +
    facet_wrap(~numcyl)
```

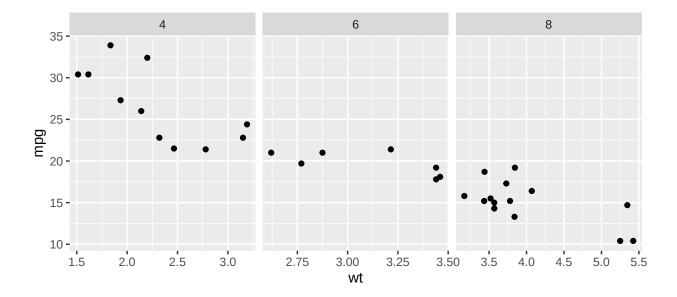


If you wanted to

facet by multiple variables you add them with a + sign. For example, facet_wrap(~am + numcyl).

The three panes have the same x and y axis by default. That behavior isn't always desirable, and one or both axes can be allowed to vary using the scales argument with "free", "free_x", or "free_y". Let's allow the x axis to vary among facet panels.

```
ggplot(mtcars, aes(x = wt, y = mpg) ) +
    geom_point() +
    facet_wrap(~numcyl, scales = "free_x")
```



Adding layers using summary statistics

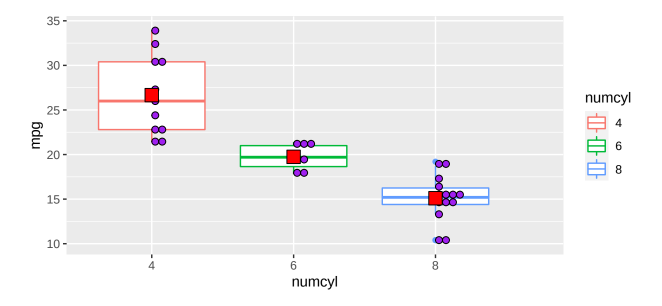
The function stat_summary() can be used to add layers based on simple summary statistics. To see how this works, let's add mean mpg by group to our boxplot/dot plot graphic as a large red square.

Sine we are working with a categorical x axis, stat_summary() automatically summarizes the y variable when we define the function we want to summarize with in fun. If x were continuous this would work differently. In addition to defining the summary f unction we need to define the geom we want to use to plot the summary statistics. In this case we'll use the point geom to add the means as points. Notice that all aesthetics we use in stat_summary() are set to constants and not mapped to variables so they are defined outside of aes().

Summary information can also be added to a graphic by creating a summary dataset and then using that summary dataset when adding additional layers to a graphic. I find this is often easier to do than to try to figure out how to do complicated summaries with stat_summary(). We will see this approach later in the workshop.

I generally can't remember which numbers correspond to which shapes in R. There is a nice cheat sheet on Cookbook for R: http://www.cookbook-r.com/Graphs/Shapes_and_line_types/. Note that shapes 21 through 25 are *fillable*, meaning you can color the inside with fill and the outline with color.

```
ggplot(mtcars, aes(x = numcyl, y = mpg) ) +
    geom_boxplot( aes(color = numcyl) ) +
    geom_dotplot(fill = "purple", binaxis = "y") +
    stat_summary(fun = mean, geom = "point", size = 5, shape = 22, fill = "red")
```



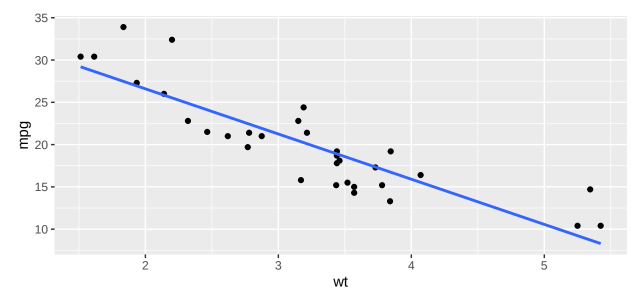
Adding regression lines

Along the same lines as summary statistics, it is straightforward to add regression lines to graphics using <code>geom_smooth()</code> or <code>stat_smooth()</code>. Let's add linear regression lines to a scatter plot using <code>geom_smooth()</code>. We have to set the <code>method</code> to "lm" to get regression lines. The default method of <code>geom_smooth()</code> is a "loess" line, which can also be useful when exploring your data.

By default geom_smooth() calculates a confidence envelope around the line, which isn't usually reasonable when working on exploratory graphics. We'll remove it using se = FALSE.

```
ggplot(mtcars, aes(x = wt, y = mpg) ) +
    geom_point() +
    geom_smooth(method = "lm", se = FALSE)
```

`geom_smooth()` using formula 'y ~ x'

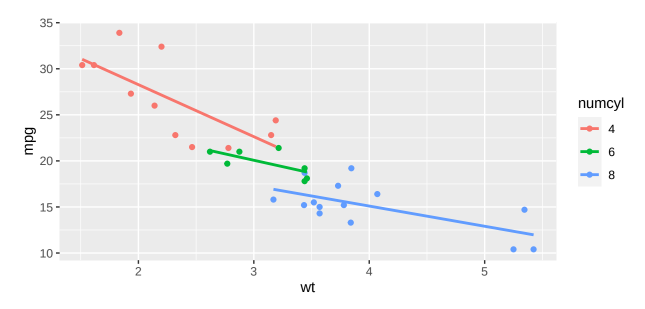


We can fit regression lines separately for each group by mapping an aesthetic such as color to a categorical variable. We'll map color to numcyl to get a separate regression line for each cylinder category.

Let's remove the confidence envelope in the next plot by using the argument se = FALSE. We could always add it back after we checked our assumptions and knew it was appropriate.

```
ggplot(mtcars, aes(x = wt, y = mpg, color = numcyl) ) +
   geom_point() +
   geom_smooth(method = "lm", se = FALSE)
```

`geom_smooth()` using formula 'y ~ x'



Polishing ggplot2 graphics

Now that we've covered some of the basics of **ggplot2**, we are going to spend time making two "final" graphics. These are graphics that you might include in a thesis or manuscript or in a presentation. Such graphics need look nicer than the basic exploratory graphics you make just for yourself, and you will have to spend time tweaking the appearance of a graph until is just right.

The second half of this workshop focuses on some of the ways we can control the overall appearance of a **ggplot2** graphic. I decided to present you two examples that end up being fairly complex. While I could have chosen to make simpler graphs, seeing some graphics with complex elements may give you an idea if **ggplot2** will be a useful tool for you when creating this sort of graphic.

I'm not advocating either of these as what any of your own final graphics should look like, as a lot of that depends on personal preference and, potentially, manuscript requirements. These examples are really just to show you some more **ggplot2** options.

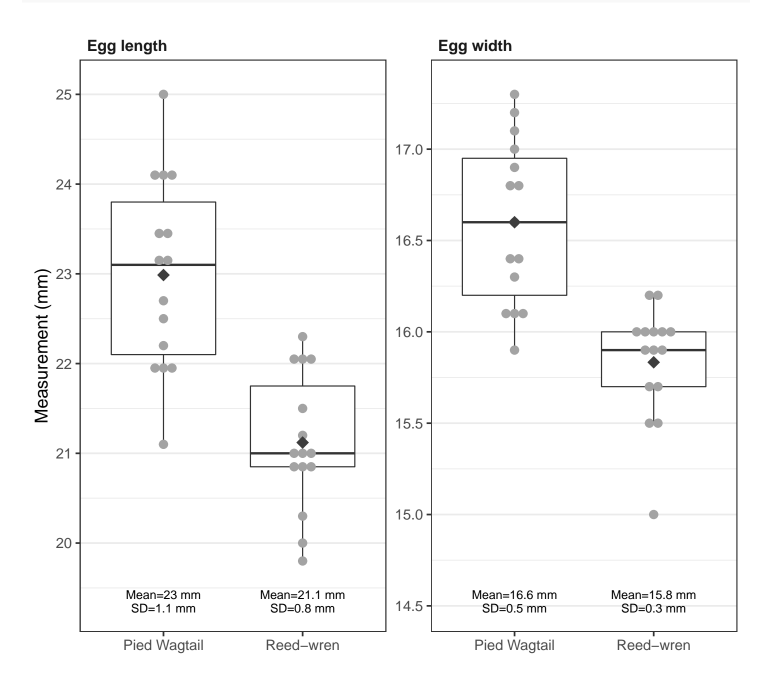
Polished graphic #1: A plot of the raw data

The first polished graphic we will create will be based on some data used to compare mean egg length and mean egg width of two bird species using two two-sample t-tests.

While a display of the results from simple tests like these would be unnecessary, we can give our audience a nice picture of what the data look like in a well thought-out graphic.

Here is the dataset.

```
16.9, 15.9, 16.3, 17.2, 16.1, 17, 17.3, 15, 16,
                               16.2, 15.7, 16.2, 15.5, 16, 15.9, 15.5,
                               15.9, 16, 15.7, 15.9, 16, 16),
                     species = structure(c(1L, 1L, 1L, 1L, 1L, 1L, 1L, 1L, 1L,
                                           1L, 1L, 1L, 1L, 1L, 2L, 2L, 2L,
                                           .Label = c("Pied Wagtail", "Reed-wren"), class = "factor") ),
                class = "data.frame", row.names = c(NA, -30L) )
head(eggs)
   id length width
                       species
       23.1 16.4 Pied Wagtail
1 198
2 199
       23.5 16.8 Pied Wagtail
      24.1 17.1 Pied Wagtail
3 200
4 201 23.4 16.4 Pied Wagtail
5 202
      23.2 16.8 Pied Wagtail
6 203
      22.5 16.6 Pied Wagtail
str(eggs)
'data.frame':
               30 obs. of 4 variables:
        : int 198 199 200 201 202 203 204 205 206 207 ...
 $ length: num 23.1 23.5 24.1 23.4 23.2 22.5 21.9 21.9 25 24.1 ...
 $ width : num 16.4 16.8 17.1 16.4 16.8 16.6 16.1 16.1 16.9 15.9 ...
 $ species: Factor w/ 2 levels "Pied Wagtail",..: 1 1 1 1 1 1 1 1 1 1 ...
You can see the final graphic we are working towards below.
library(tidyr)
library(dplyr)
eggs2 = pivot_longer(eggs, cols = c(length, width),
                    names_to = "type",
                    values_to = "measurement",
                    names_ptypes = list(type = factor() ) )
levels(eggs2$type) = c("Egg length", "Egg width")
sumdat = eggs2 %>%
        group_by(type, species) %>%
        summarise(Mean = round(mean(measurement), 1),
                 SD = round(sd(measurement), 1) )
loc = eggs2 %>%
      group by(type) %>%
      summarise(yloc = min(measurement) - .5)
sumdat = sumdat %>%
    inner_join(loc, by = "type") %>%
    mutate(label = paste0("Mean=", Mean, " mm", "\n", "SD=", SD," mm") )
ggplot(eggs2, aes(x = species, y = measurement) ) +
    geom_boxplot() +
    facet_wrap(~type, scales = "free_y") +
    geom_dotplot(binaxis = "y", stackdir = "center",
                color = "grey64", fill = "grey64", dotsize = .5) +
    stat_summary(fun = mean, geom = "point", shape = 18,
                size = 5, color = "grey24") +
    theme_bw(base_size = 16) +
    theme(panel.grid.major.x = element_blank(),
          strip.background = element_blank(),
          strip.text = element_text(hjust = 0,
                                   face = "bold",
                                   size = 14) ) +
```



Reshaping a dataset prior to graphing

You can see that I used faceting to put measurement type (length or width) in separate panes. However, the current dataset doesn't have a variable for us to facet with because length and width are in separate columns. We will need to reshape this wide dataset into long format. I will use gather() from package tidyr for this today.

Using gather(), we will create a new categorical variable representing measurement type in one column and all the quantitative measurement values in another. We define the names of the new columns we'll are making using the key and value arguments, and then list which columns we want to take and put into a single column.

Reshaping is often needed in order to take full advantage of **ggplot2**. Being comfortable with data manipulation in R is one key to success for creating **ggplot2** graphics.

We'll name the reshaped dataset eggs2. It is in long format, so it has twice as many rows as the original dataset eggs.

```
library(tidyr)
eggs2 = gather(eggs, key = type, value = measurement,
            length, width, factor_key = TRUE)
head(eggs2)
   id
           species
                     type measurement
1 198 Pied Wagtail length
                                  23.1
2 199 Pied Wagtail length
                                  23.5
3 200 Pied Wagtail length
                                  24.1
4 201 Pied Wagtail length
                                  23.4
5 202 Pied Wagtail length
                                  23.2
6 203 Pied Wagtail length
                                  22.5
```

str(eggs2)

```
'data.frame': 60 obs. of 4 variables:

$ id : int 198 199 200 201 202 203 204 205 206 207 ...

$ species : Factor w/ 2 levels "Pied Wagtail",..: 1 1 1 1 1 1 1 1 1 1 1 ...

$ type : Factor w/ 2 levels "length", "width": 1 1 1 1 1 1 1 1 1 1 ...

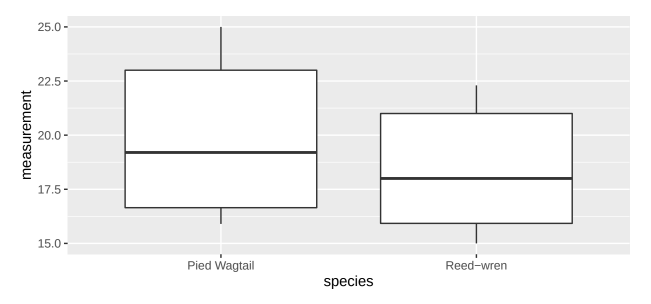
$ measurement: num 23.1 23.5 24.1 23.4 23.2 22.5 21.9 21.9 25 24.1 ...
```

I'm going to change the style of how I create the polished graphics for teaching purposes. You will likely see this style on help forums or in help documents, although I personally rarely use this style in my own graphics code.

This is how it will go: First I will create and name a basic graphic by defining the dataset and the x and y axes in ggplot() and adding a geom layer. This first graphical object will be named g1. Then we will start to add layers to g1, renaming the graphic as g1 each time we add a layer. This will allow us to see how the graphic changes layer by layer.

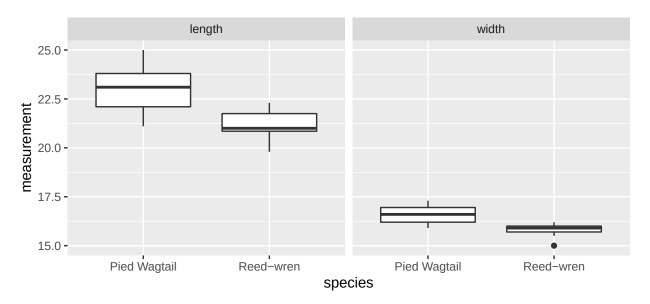
We will start with a simple boxplot, putting species on the x axis and measurement on the y axis. The extra pair of parentheses prints the graphic to the plotting pane so we can see the plot change as we go.

```
( g1 = ggplot(eggs2, aes(x = species, y = measurement) ) +
    geom boxplot() )
```



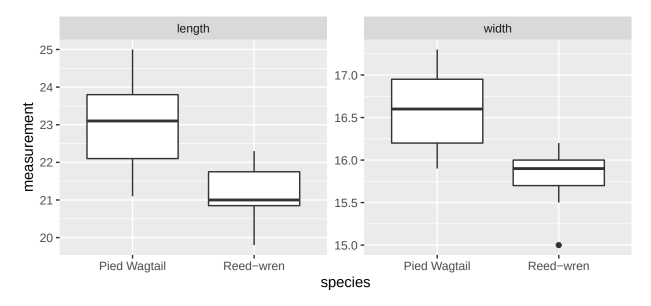
Let's add faceting by type to the graphic g1.

```
g1 + facet_wrap(~type)
```



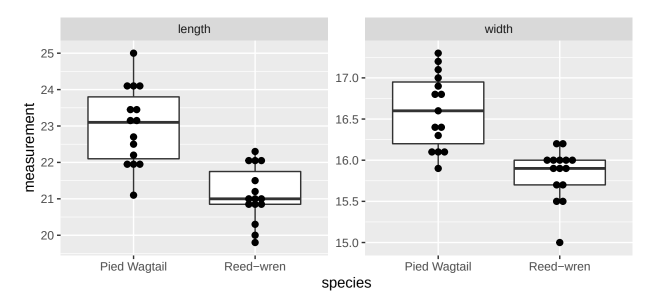
Not surprisingly, length and width cover fairly different ranges. It makes sense to allow the y axis to have different limits for the two different panes (i.e., we want to free the y axis scale). The x axis is the same for each pane, though, so we don't need to allow that to vary.



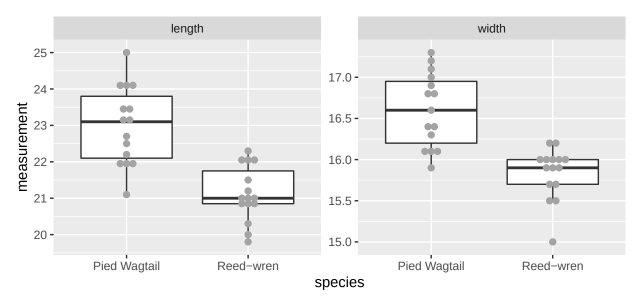


Adding the raw data on top of the boxplot as a dot plot will give a more complete picture of what the distribution of these data looks like. I use stackdir = "center" to make a centered dot plot, which is a little bit like a violin plot and a histogram.

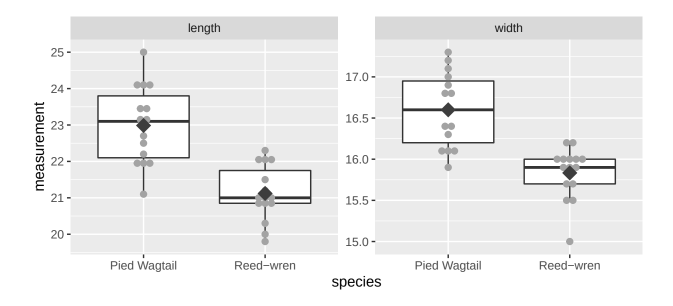
```
g1 + geom_dotplot(binaxis = "y", stackdir = "center")
```



I thought the black dots were too dark, and decided to change the color to a lighter gray. Color choice is often a lot of trial and error for me, which I'm not going to show you here. I settled on a gray color called grey64. As we'll set both fill and color aesthetics to constants, these go outside of aes.



Boxplots show medians but not means. It can be nice to add the means, especially when the statistical test used to analyze these data was about differences in means. Let's add the mean value for each species and type as diamonds, and fill them with a darker gray color, grey24. I didn't like how small these points were at first, so we'll increase the size of the diamonds, as well.



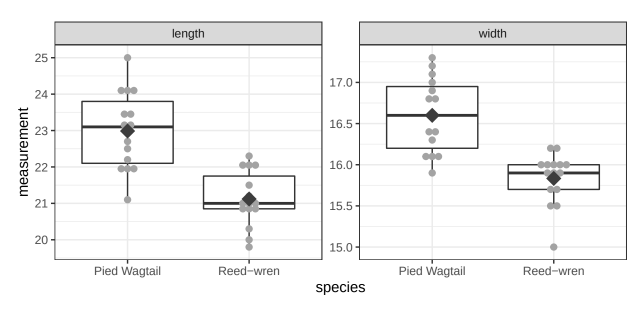
Changing plot appearance with theme()

So far this is pretty much a review of things we did in the first part of the workshop. Let's change our focus to the overall appearance of the graphic, particularly the appearance of the panels and axes.

The default appearance of **ggplot2** graphics is called **theme_gray()**. The default gray background of **ggplot2** can be nice for seeing differences in colors, but is not as useful if printing in black and white. If I'm going to include something in a printed document, I often use the built-in theme theme_bw() to change the overall appearance of the graphic. This theme is called "theme black-and-white". You also might be interested in the built-in themes theme_minimal() or theme_classic(), which are a standard plot format in some fields. There are many themes out there - check out package **ggthemes** for more options.

While I could change all elements of the panel manually, I find theme bw() gives a nice starting point for further changes.

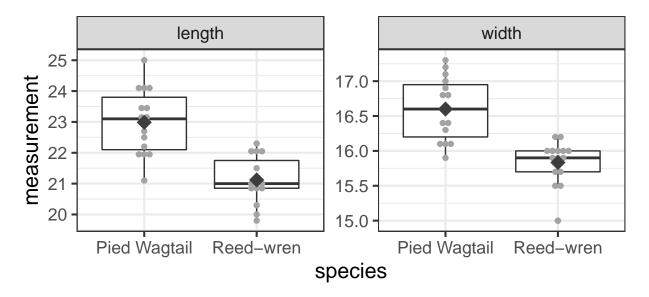




Although the result isn't actually black and white, as the grid lines and axis tick labels are still grey, you can see this changes the panel background to white.

The default text size in **ggplot2** themes is often too small for graphs that will be in a presentation or publication. I can make all these bigger by changing the base_size in the overall theme_*() function. I have used 18 for presentations before; here I'll use 16 (the default is 11)

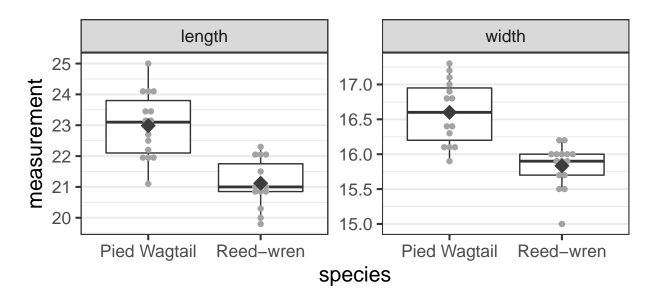




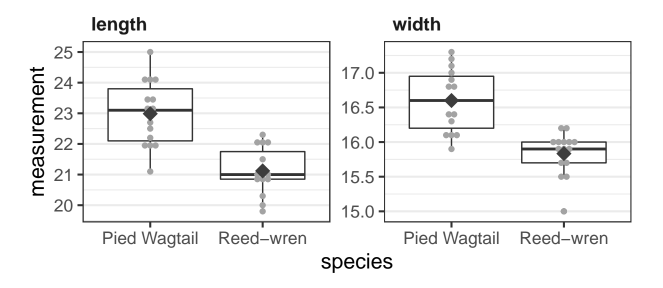
Having grid lines along the y axis is useful for reading the graph, but grid lines along a categorical axis like the x axis seems unnecessary. Control of panel elements, including the removal of grid lines, can be done in theme(). See the help page for theme() via ?theme for the many, many elements of the graphic that you can change.

Because we only want to remove the grid lines on the x axis we will use the panel.grid.major.x argument set to element_blank() within theme().

g1 + theme(panel.grid.major.x = element_blank())



While we're working in theme(), let's make changes to the strips of the facets. We can change the color of the background (or remove it all together by setting it to element_blank()) with strip.background and adjust the text placement and size with strip.text. Note we cannot actually change the labels in strip.text, only adjust how the text element is displayed.



Changing facet labels

At this point I realized I'd made an error. Changing the text in the facet strips is something that is not particularly easy, although some work can be done via the labeller argument in facet_wrap(). I should have changed the levels of the variable type in the dataset to begin with to make them look nicer. Let's do that now, by assigning new levels to the type variable.

```
# Changing the factor levels is easy
levels(eggs2$type)

[1] "length" "width"

# We can change the names by assigning new levels
levels(eggs2$type) = c("Egg length", "Egg width")
levels(eggs2$type)
```

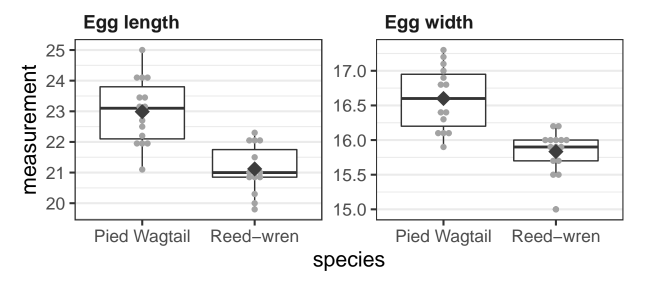
[1] "Egg length" "Egg width"

Creating the same plot using a different or edited dataset

Does that mean we have to rerun all the code we've done so far? We could - in a situation where we weren't going through things one step at a time, running the code again would be a piece of cake.

We do have another option, though, with the %+% function. This function allows us to recreate a graphic we've already made (and named) using a different dataset. This can be convenient if you want identical graphic appearance from data from separate datasets. Here we'll use the newly edited eggs2 dataset with g1 to recreate the graphic we've made so far.

```
( g1 = g1 %+% eggs2 )
```



The last appearance change we really need is the axis labels. Axis and other labels can be changed in labs(). I decided to suppress the x axis label because it didn't seem necessary. By using NULL instead of "" I removed all of the space between the tick labels and the bottom of the plot. The labels on the y axis needed cleaning up, with capitalization and the units of measurement added.



Adding summary statistics as text

I could have stopped here and had a perfectly nice graphic. However, I like adding additional information to graphics so I decided to walk you through the process I went through to add summary statistics as text to this graphic.

To do this, we first need to calculate the summary statistics and figure out where to place them along the x and y axes. Then we'll add the information the graphic as *labels* with <code>geom_text()</code>. The other way to do this, which I use more often, is to add tables to graphical objects using the <code>gridExtra</code> package. This package is also useful for putting two graphical objects in one pane.

We are going to add the mean and standard deviation of each type of egg measurement for each species as text in the graphic. First let's calculate the statistics we want for each group. I often use group_by() and summarise() from the dplyr package for this kind of work. We'll name the new summary data frame sumdat. Notice that we need to round the results here to a reasonable number of significant digits.

```
# Groups:
            type [2]
                           Mean
                                    SD
 type
             species
  <fct>
             <fct>
                          <dbl> <dbl>
1 Egg length Pied Wagtail 23
                                   1.1
2 Egg length Reed-wren
                           21.1
                                   0.8
3 Egg width Pied Wagtail 16.6
                                  0.5
4 Egg width Reed-wren
                           15.8
                                  0.3
```

To make the labels for plotting, I added the name of each statistic as well as a line break between them using pasteo()

```
[1] "Mean=23 mm\nSD=1.1 mm" "Mean=21.1 mm\nSD=0.8 mm" "Mean=16.6 mm\nSD=0.5 mm" [4] "Mean=15.8 mm\nSD=0.3 mm"
```

You can see in the final plot that I decided to add the summary statistics beneath each boxplot. This means we can still use species as the variable that defines where the text will be placed on the x axis, and type will define which facet it will be in. However, we'll need to calculate a new variable to define where the text will be positioned on the y axis.

After some trial and error (which you don't get to see), I decided the minimum y value from each facet minus 0.5 would be a good place to put the text in the plot. We can calculate this number separately for each type using group_by()/summarise() again. We'll name the y position variable yloc. We can then add yloc to sumdat using inner_join() from dplyr to join or merge the two datasets.

```
( loc = eggs2 \%>%
      group_by(type) %>%
      summarise(yloc = min(measurement) - .5) )
# A tibble: 2 x 2
  type
              yloc
  <fct>
             <dbl>
1 Egg length 19.3
2 Egg width
              14.5
( sumdat = inner_join(sumdat, loc, by = "type") )
# A tibble: 4 x 6
            type [2]
# Groups:
             species
                           Mean
                                    SD label
                                                                   yloc
 type
             <fct>
                           <dbl> <dbl> <chr>
                                                                  <dbl>
  <fct>
1 Egg length Pied Wagtail
                           23
                                   1.1 "Mean=23 mm\nSD=1.1 mm"
                                                                   19.3
2 Egg length Reed-wren
                           21.1
                                   0.8 "Mean=21.1 mm\nSD=0.8 mm"
                                                                   19.3
3 Egg width Pied Wagtail
                           16.6
                                   0.5 "Mean=16.6 mm\nSD=0.5 mm"
                                                                   14.5
                                   0.3 "Mean=15.8 mm\nSD=0.3 mm"
                            15.8
4 Egg width Reed-wren
                                                                   14.5
```

If you don't know data manipulation in R very well, this may seem like a huge amount of work in order to get what you want. I am showing you this, though, because I wanted to drive the point home that working with a dataset outside of **ggplot2** is often required in order to take full advantage of the strengths of this package. If you aren't comfortable with this sort of work in R then **ggplot2** likely won't be a useful tool for you for making complicated final graphics. Another option is to add such information outside R using a different software package.

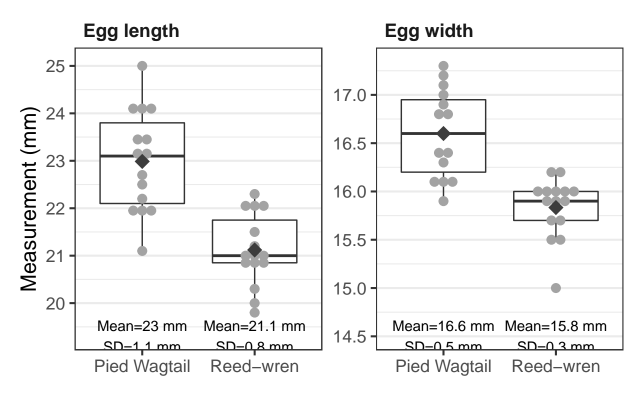
Defining a new dataset for plotting for a specific layer

We can now add text to a graphic using <code>geom_text()</code>. For the first time we'll be defining a new dataset for plotting within a specific geom layer. Defining a new dataset in a geom will override the global dataset for the graphic that we defined in <code>ggplot()</code>. The rest of the graphic will still plot information from the dataset <code>eggs2</code>, but <code>geom_text()</code> will plot what is in the <code>sumdat</code> dataset. When defining a dataset within a layer it is safest to type out the <code>data</code> argument, as the dataset is not the first argument in the <code>geom</code> layers like it is in <code>ggplot()</code>.

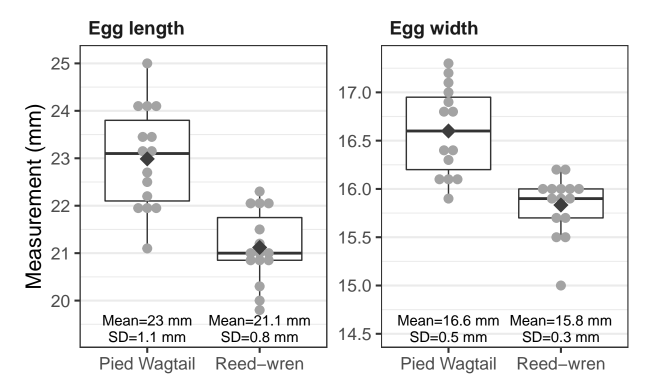
Notice we have to set the y position for geom_text() because we're using a different variable for y than we had for the rest of the graphic. We map the y axis position to yloc within aes.

The geom_text() geom requires us to define a label variable within aes(). The help files for individual geoms will tell you if there are required aesthetics, which you should check for you are using a geom for the first time. We will map the variable label to the label aesthetic.

```
g1 + geom_text(data = sumdat, aes(label = label, y = yloc) )
```

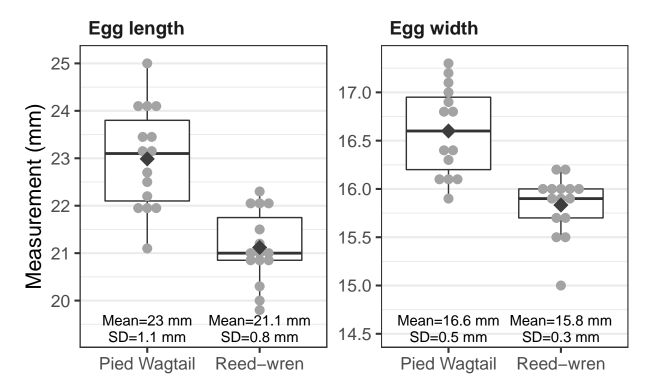


As often happens, we need to do some adjustments to the text we've added to make the graphic look nicer. Here we'll adjust the line height, change the vertical justification, and change the size of the text within <code>geom_text()</code>.



This is what the code looks like when it is put all together instead of adding layers line by line.

```
( g1 = ggplot(eggs2, aes(x = species, y = measurement) ) +
   geom_boxplot() +
   facet_wrap(~type, scales = "free_y") +
   geom_dotplot(binaxis = "y", stackdir = "center",
                color = "grey64", fill = "grey64") +
   stat_summary(fun = mean, geom = "point", shape = 18,
                size = 5, color = "grey24") +
   theme_bw(base_size = 16) +
   theme(panel.grid.major.x = element_blank(),
         strip.background = element_blank(),
         strip.text = element_text(hjust = 0,
                                    face = "bold",
                                    size = 14) ) +
   labs(x = NULL,
        y = "Measurement (mm)") +
   geom_text(data = sumdat, aes(label = label, y = yloc),
             lineheight = .9, vjust = .3, size = 4) )
```



In making this graph, I've relied on the graphic displayed in the plotting window to help decide on spacing and point sizes and such. Be careful using this method, because we haven't actually set the plotting window size to anything. In fact, you can't set the RStudio plotting window to a specific size at this time. Depending on the size and shape we save the graphic as, we may decide we need to make a few more adjustments. One way to check how things will look at different sizes is to preview the plot at difference sizes using the Export drop down menu in the RStudio Plots pane. You can also just save the graphic to your desired size and take a look at it.

Saving a plot with ggsave()

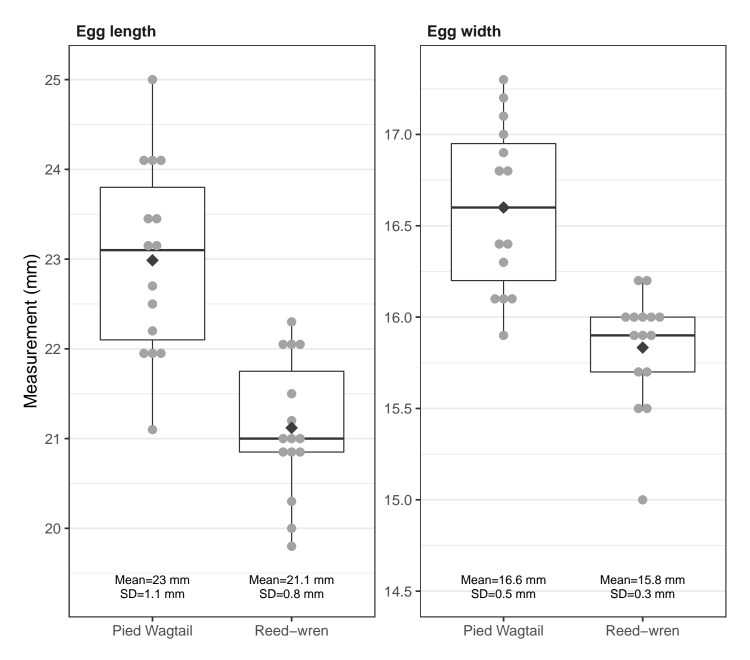
We'll save the plots using ggsave() from ggplot2. You can save graphics in all kinds of formats, but here we'll save the graphic as a PDF and as a PNG file. The sizes I chose are arbitrary, and are just to demonstrate some of the ggsave() options.

By default ggsave() saves the last plot made to whatever size your plot window is. You can set the graphic size via width and height.

It can be safer to define the graphic you want saved by name using the plot argument. If you want the plot to look like it does in the plotting window, simply leave height and width at their default values.

```
ggsave("final plot 1.pdf", width = 7, height = 7) # setting width and height (default inches)
ggsave("final plot 1.png", plot = g1) # using default size based on plotting window
```

At this point I decided that the dot size was too big. I edited the geom_dotplot() argument dotsize and then re-saved the final plot at a final size, which was 9 inches wide and 8 inches high. The graphic at this size is shown below.



Notice we can change dpi (dots per inch) for the PNG format

```
ggsave("final plot 1.pdf", plot = g1, width = 9, height = 8)
ggsave("final plot 1.png", dpi = 600, width = 9, height = 8)
```

Polished graphic #2: A "results" plot

The last graphic we are going to make is one that displays the results from an analysis of a dataset that had two factors. The results we will be graphing are the tests of differences in mean growth of each group against the control group. I pulled the estimated differences in means and the associated confidence intervals from the statistical model and saved the results.

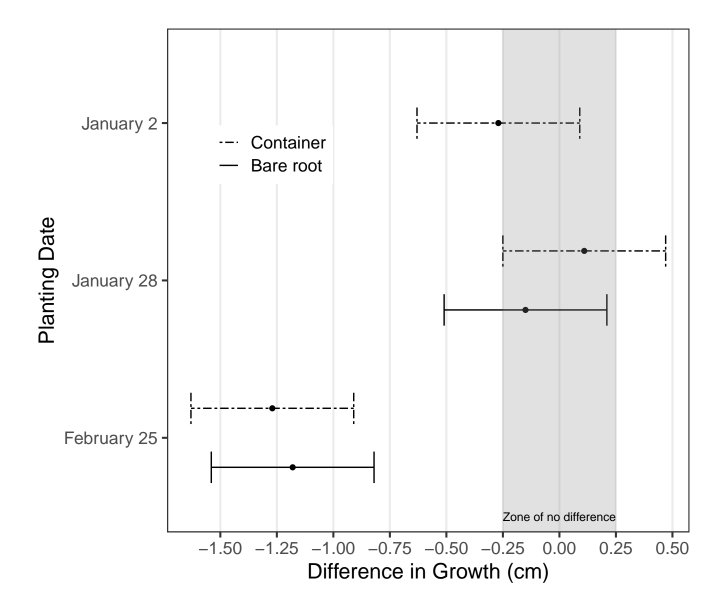
```
res = structure(list(Diffmeans = c(-0.27, 0.11, -0.15, -1.27, -1.18)
),
                    Lower.CI = c(-0.63, -0.25, -0.51, -1.63, -1.54),
                    Upper.CI = c(0.09, 0.47, 0.21, -0.91, -0.82),
                    plantdate = structure(c(2L, 3L, 3L, 1L, 1L),
                      .Label = c("Feb25", "Jan2", "Jan28"), class = "factor"),
                    stocktype = structure(c(2L, 2L, 1L, 2L, 1L),
                      .Label = c("bare","cont"), class = "factor") ),
                    class = "data.frame", row.names = c(NA, -5L) )
str(res)
'data.frame':
                5 obs. of 5 variables:
 $ Diffmeans: num -0.27 0.11 -0.15 -1.27 -1.18
 $ Lower.CI : num -0.63 -0.25 -0.51 -1.63 -1.54
 $ Upper.CI : num 0.09 0.47 0.21 -0.91 -0.82
 $ plantdate: Factor w/ 3 levels "Feb25","Jan2",..: 2 3 3 1 1
 $ stocktype: Factor w/ 2 levels "bare", "cont": 2 2 1 2 1
```

```
Diffmeans Lower.CI Upper.CI plantdate stocktype
1
      -0.27
                -0.63
                          0.09
                                     Jan2
                                                cont
2
                -0.25
                          0.47
                                    Jan28
       0.11
                                                cont
3
      -0.15
                -0.51
                          0.21
                                    Jan28
                                                bare
4
      -1.27
                -1.63
                          -0.91
                                    Feb25
                                                cont
5
      -1.18
                -1.54
                         -0.82
                                    Feb25
                                                bare
```

res

I added the two factor variables to the results dataset before I saved it so I would be able to tell which groups were being compared to the control on each row. The two factors are plantdate, which has three levels that indicate the date that trees were planted, and stocktype, which has two levels and indicates the kind of stock (either bare root or in a container). The control group was bare root trees planted on January 2. There were six groups including the control group, and so there are five comparisons in this table of results.

The graphic we are working towards is shown below.



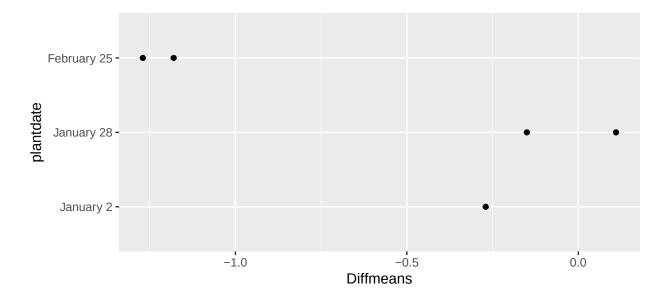
Setting the factor order to control axis order

We now know that it is often easiest to clean up the dataset before we are ready to begin plotting it, so let's make some changes to the variable plantdate. The levels of this variable aren't in date order because R by default orders levels of factors alphanumerically. We can always change the display order of a factor in **ggplot2** by changing the order of the factor levels in the dataset. We will relabel the levels of the factor to make them look nicer while we're at it to save us some work later.

This example turns out to be more complex than I was originally planning on because of the odd number of comparisons. Two of the plant dates have comparisons between both stock types and the control but January 2 only has one stock type comparison. However, I decided to stick with the complex graphic so you can see the kinds of problems you have might run into as your graphic complexity increases.

We'll start by plotting the estimated differences in means as points with plantdate on the y axis since we want horizontal error bars

```
ggplot(res, aes(x = Diffmeans, y = plantdate) ) +
  geom_point()
```



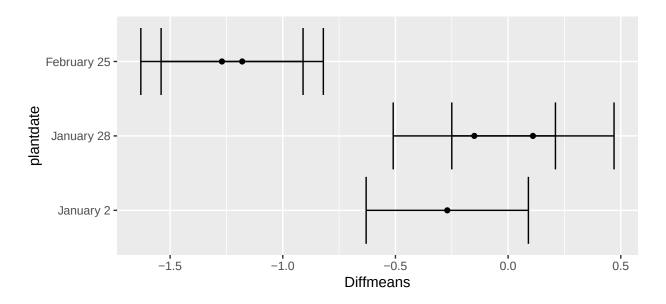
Adding error bars to a plot

We can add the 95% confidence intervals around the estimated means as error bars with <code>geom_errorbar()</code>. Since we want horizontal error bars, we map the aesthetics <code>xmin</code> and <code>xmax</code> to variables that represent the <code>ends</code> of the error bars within <code>aes()</code>. I added the upper and lower confidence limits to the dataset for this purpose.

If we were making vertical error bars we'd use ymin and ymax. These (xmin/xmax or ymin/ymax) are required aesthetics in geom_errorbar().

Being able to make horizontal error bars directly like this was introduced in **ggplot2** version 3.3.0. If your version of the package is older then this code won't work.

```
ggplot(res, aes(x = Diffmeans, y = plantdate) ) +
    geom_point() +
    geom_errorbar( aes(xmin = Lower.CI, xmax = Upper.CI) )
```



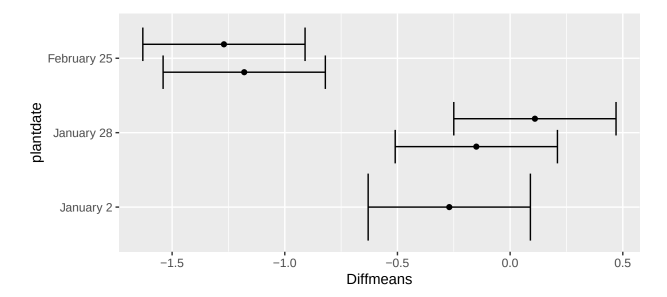
Dodging to avoid overlap

When we have two different stock types at a single planting date, the error bars overlap and the graphic is difficult to understand. To avoid this, we will do what is called *dodging*. In order to *dodge* the overlapping points, we need to define which variable identifies that we have two values for two groups at a single planting date. In this case that variable is

stocktype, so we map the group aesthetic in aes() to stocktype. A common issue when trying to dodge overlapping geoms apart is failing to make a variable to dodge on.

We want to dodge whenever we get an overlap along the x axis, so in addition to adding group we need to use the position argument. This is done *outside* aes() in each layer. We choose the amount we want to dodge by in position_dodge() with the width argument.

Notice we have to dodge both the points and the error bars, and that we do it by the same width amount so they stay lined up. Common dodging amounts are 0.9 and 0.75.

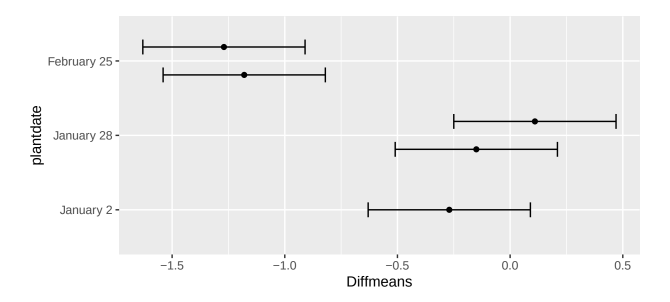


Setting the width of error bars for unbalanced factors

Now we have gotten rid of the overlap problem. But this is where things start getting complicated. The width of the error bar is twice as big for the single stock type on January 2 as it is when we have two stock types on the other dates. We can control this using the width argument in geom_errorbar().

Usually we set width to a constant and all the widths would be changed the same way. Here we'll need to set the width of the first error bar to *half* the value of the others. This means we'll have to set width to a vector defining the width of all five error bars (so the vector has a length of 5). Order matters; the first value is the one for January 2.

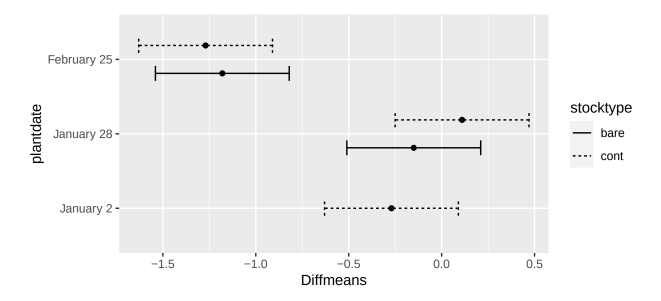
I picked widths using trial and error. Don't be surprised if you don't get widths you like on your first try.



Adding a legend with linetype.

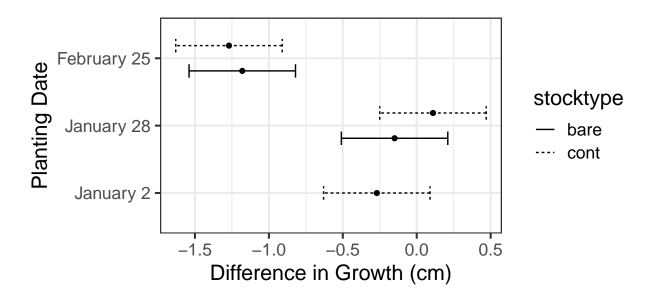
That looks much nicer. But we can't tell which line is for which stock type. We'll fix that by mapping linetype to stocktype within geom_errorbar(). This is aesthetic mapping, so goes inside aes().

This is the basic plot we'll build on, so we'll assign it a name.



Changing appearance with overall theme

Now that we have the basic graphic built, let's start polishing it up by setting the theme to theme_bw(), change the base text size, and clean up the axis labels. This was all done in the previous example.

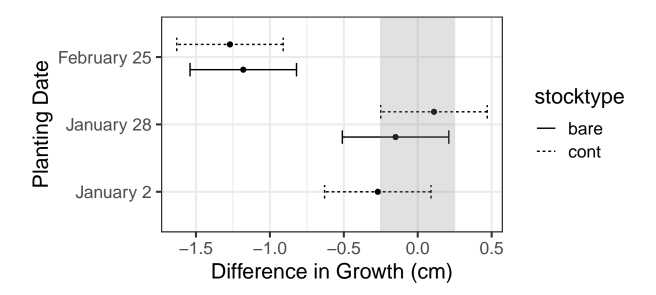


Adding a shaded rectangle

It can be informative to give some indication of what a practical difference would be on the graphic. In this case, a difference in mean growth of 0.25 cm compared to the control would be practically meaningful to growers. In the past I've indicated the practically important limits using horizontal lines with <code>geom_hline()</code>. Today we'll add a shaded rectangle to indicate this "Zone of no difference" using the <code>rect</code> geom.

If you want to add many rectangles based on a dataset you'll likely use <code>geom_rect()</code>. However, if you want to add a single rectangle with transparency like we'll do today you'll often use <code>annotate()</code>. In <code>annotate()</code> we need to define which geom we are using; in this case, "rect".

When making rectangles we need to define where the *edges* of the rectangle should be drawn. Since these aesthetics are set to constants this is done outside aes(). The x axis values are set to the practically important difference (-0.25, 0.25). Because we want the rectangle to go all the way across the y axis we use -Inf/Inf for ymin and ymax. We'll also set alpha to make the rectangle see-through and set the fill color to some gray shade with fill.

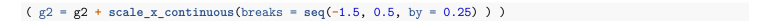


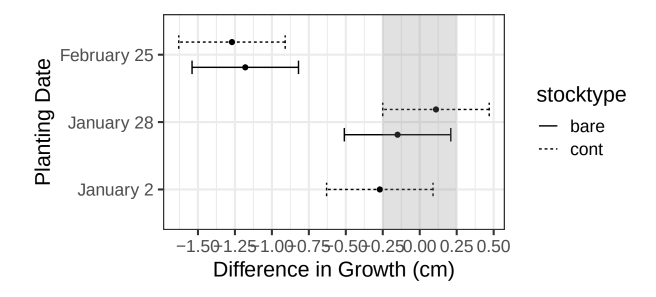
Changing the x axis breaks

If 0.25 cm is the practically important difference, we may want to show finer tick marks or *breaks* on the x axis. Notice that right now the breaks are set for every 0.5 cm change. We can control axes with the appropriate scale functions. In this case, the x axis is continuous and we'll use $scale_x_continuous()$.

We'll make axis breaks every 0.25 cm instead of 0.5 cm using a sequence of numbers from the -1.5 to 0.5 in 0.25 increments (see ?seq for more info).

The result looks a little busy when printed here in this document, but we're not quite to the final plot yet and this will look better in the end.



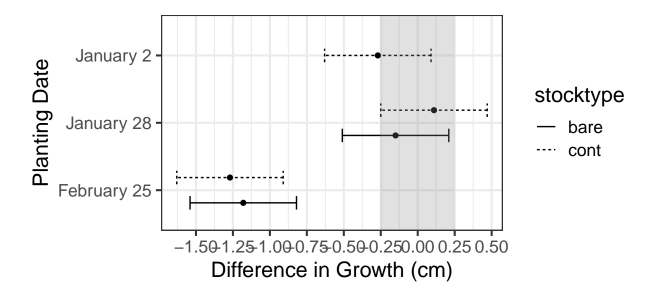


Reversing the y axis

Since y is categorical, the y axis is a *discrete* axis. By default the factor levels on a discrete axis are listed bottom to top. Let's change these so they go top to bottom; i.e., January 2 should be listed first.

This can be done in scale_y_discrete() using the limits argument. We reverse limits with rev.

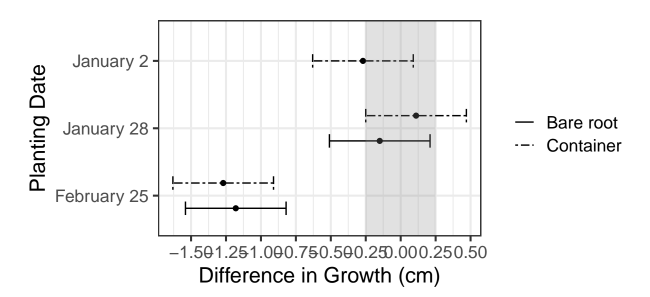




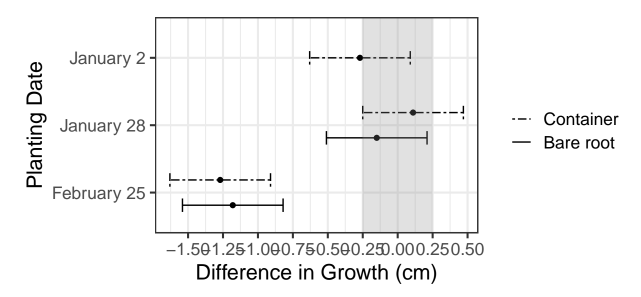
Changing the appearance of the legend

There are a few final appearance changes we need to make on this graphic. First, I didn't like the dotted linetype that we got by default. We can change the type of lines we get in the plot and legend with the function scale_linetype_manual(). Most aesthetics have a scale function that you can use to change the default aesthetic settings (we saw color and fill scales earlier in the workshop).

We'll set the values of the line types in scale_linetype_manual(); you can see the different types of lines available here. We'll also change the title and the labels of the groups that are displayed in the legend with this function, as well, via name and labels, respectively. We remove the names all together.



It seems more aesthetically pleasing to have the order of the lines in the legend match the order that the lines show up in the plot. In this case the dashed line/Container group comes first in the plot. We can control this inside scale_linetype_manual() as well, using the guide argument with guide_legend(). We set reverse to TRUE to reverse the order of the legend. The guide_legend() function allows us to control many aspects of the legend.



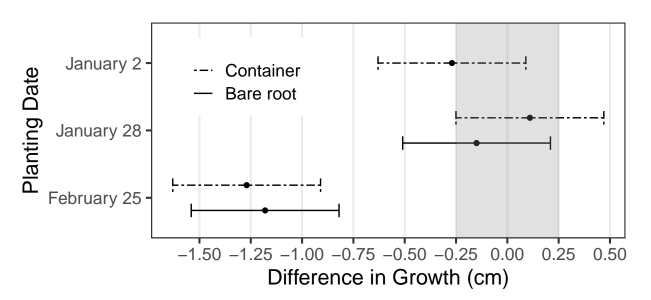
Moving the legend inside the plot panel

We have plenty of blank space within the graphic to fit the legend. Let's move the legend inside the plot panel. This can be done in theme().

We use legend.position to move the legend inside the plot panel. The legend position here is based on coordinates between 0 and 1 on each axis, and took trial and error to get a nice placement. You can also use options such as, e.g., "bottom" in legend.position to leave the legend outside the plot panel but move it to a different side of the plot. See the help page for theme() for more options.

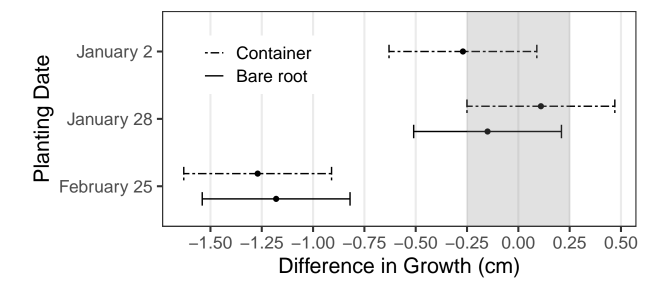
I'll keep the legend vertical but if we wanted to make it horizontal we would do so using legend.direction.

I decided to remove the y axis grid lines all together along with the minor x axis gridlines, which is also done in theme().



Once I moved the legend inside the plot I thought the space around the legend could use a couple more tweaks. This led me down a rabbit hole figuring out how to change the margins (with legend.margin) and fix the spacing above the legend. I ended up using legend.spacing.y after reading this GitHub issue.

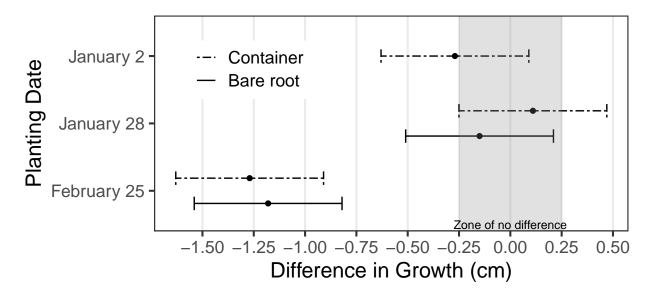
As you can see, there are a lot of things you can change in theme().



Adding a label with annotate()

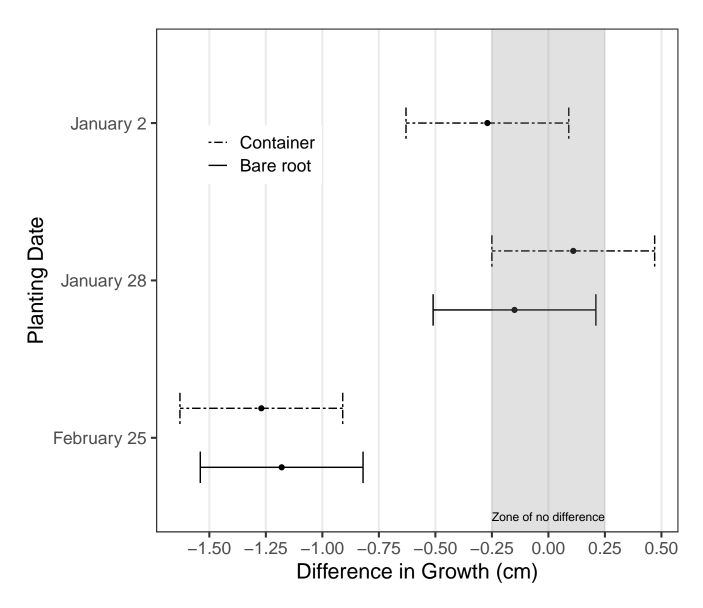
The last thing we want to do is add the label to our "Zone of no difference" rectangle. Unlike adding multiple labels all positioned separately by group like in the last graphic, we want to add a single label. This is a clue that we should add the text using annotate() instead of geom_text(), much like we used annotate() earlier for making the rectangles instead of geom_rectangle(). This time we'll use the "text" geom in annotate().

In addition to placing the label we'll also shrink the size of the text to make it fit inside the rectangle (once the plot is at its final size). Because we are setting all aesthetics to constants, they are outside aes(). The x and y aesthetics are based on the limits of each axis.



And there we have it, the final graphic. Here's what the code looks like all together. The graphic is displayed at the size of the final graphic I showed you before we began working on the plot, which is 7 x 6 inches.

```
( g2 = ggplot(res, aes(x = Diffmeans, y = plantdate,
                       group = stocktype) ) +
     geom_point(position = position_dodge(width = 0.75) ) +
     geom_errorbar( aes(xmin = Lower.CI, xmax = Upper.CI,
                        linetype = stocktype),
                     position = position_dodge(width = 0.75),
                     width = c(.2, .4, .4, .4, .4)) +
     theme_bw(base_size = 16) +
     labs(x = "Difference in Growth (cm)",
          y = "Planting Date") +
     annotate(geom = "rect",
              xmin = -0.25, xmax = 0.25,
              ymin = -Inf, ymax = Inf,
              fill = "grey54", alpha = 0.25) +
     scale_x_continuous(breaks = seq(-1.5, 0.5, by = 0.25)) +
     scale_y_discrete(limits = rev) +
     scale_linetype_manual(values = c("solid", "twodash"),
                            name = NULL,
                            labels = c("Bare root", "Container"),
                            guide = guide_legend(reverse = TRUE) ) +
     theme(legend.position = c(0.2, 0.75),
           legend.spacing.y = unit(0, "pt"),
           legend.margin = margin(t = 4, r = 5, b = 5, 1 = 5),
           panel.grid.major.y = element_blank(),
           panel.grid.minor.y = element_blank(),
           panel.grid.minor.x = element_blank() ) +
     annotate(geom = "text", x = 0, y = 0.5,
               label = "Zone of no difference", size = 3) )
```



You would save this graphic in the same way as before with ggsave().

```
ggsave("final plot 2.pdf", plot = g2, width = 7, height = 6)
```

That is all the material we will cover today. There is much more that you can do in **ggplot2** that we didn't have time to review, but this should give you a start on making your own graphics with **ggplot2** in R.