Modifying ggplot objects - intermediate tutorial

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Getting started

If this is your first time using the ggplot2 library within R, please refer to the **Beginners Tutorial**.

```
# If you don't have ggplot2 installed already
install.packages("ggplot2")
```

Let us load up ggplot2 into our current environment.

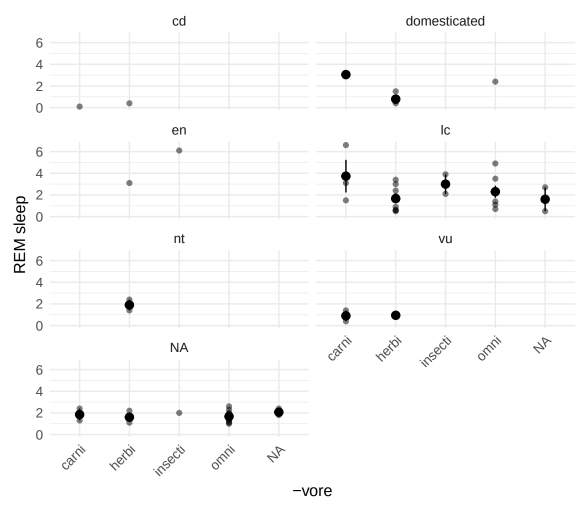
require(ggplot2)

ggplot - Review

Let us take a quick look at the plot script for the last ggplot figure we generated in the preceding tutorial.

```
# Get data we were working with earlier
data("msleep", package = "ggplot2")
# Plot the ggplot
ggplot(msleep, aes(x = vore, y = sleep_rem)) + geom_point(alpha = 0.5) +
    stat_summary() + theme_minimal() + theme(text = element_text(size = 12)) +
    ggtitle("I changed the title") + theme(plot.title = element_text(hjust = 0.5),
    axis.text.x = element_text(angle = 45, hjust = 1)) + labs(x = "-vore",
    y = "REM sleep") + facet_wrap(~conservation, ncol = 2)
```

I changed the title



There are 5 key points in this long line of code:

- The data object is msleep.
- We used theme_minimal() to apply a clean background (not grey!) to our plot.
- We modified the text size and plot title placement using theme().
- We use theme_minimal before we add any modifications to theme(), because otherwise

theme minimal() will reset any changes we added to our figure using theme().

• We separated different categories (in 'conservation') using facet_wrap(), and fixed the maximum number of 'columns' we wanted for the facet panels.

Here's a quick way to shorten this code a bit more. The *theme_minimal()* element has 2 options we can define in it - font size, and font family. Instead of defining these in *theme*, we can define them in **theme_minimal()**. Check to see if this gives the same figure as above.

```
ggplot(msleep, aes(x = vore, y = sleep_rem)) + geom_point(alpha = 0.5) +
    stat_summary() + theme_minimal(base_size = 12) + ggtitle("I changed the title") +
    theme(plot.title = element_text(hjust = 0.5), axis.text.x = element_text(angle = 45,
        hjust = 1)) + labs(x = "-vore", y = "REM sleep") + facet_wrap(~conservation,
    ncol = 2)
```

• hjust=0.5 means align text at the center, whereas hjust=1 means right, and hjust=0 means left.

Before we dive into today's tutorial, let me highlight an error message you may receive as you run the code above,

Warning message: Removed 22 rows containing missing values (geom_point).

This is because we have 'NA' values in our dataframe. We can overcome this by updating our msleep to get rid of rows that contain 'NA'. This is done using the function na_omit().

```
msleep = na.omit(msleep)
```

Fair warning, na.omit will remove rows that have an 'NA' value in *any* column, so if you have certain columns that are 'NA' more often than not, you might want to remove those first, or replace the value with 0.

Getting our data

Get Data

We will be working with a sample dataset available within the ggplot2 library, called *midwest*. This dataset contains statistics on populations in US Midwest.

```
data("midwest", package = "ggplot2")
```

We can take a peek at the data, using the head command.

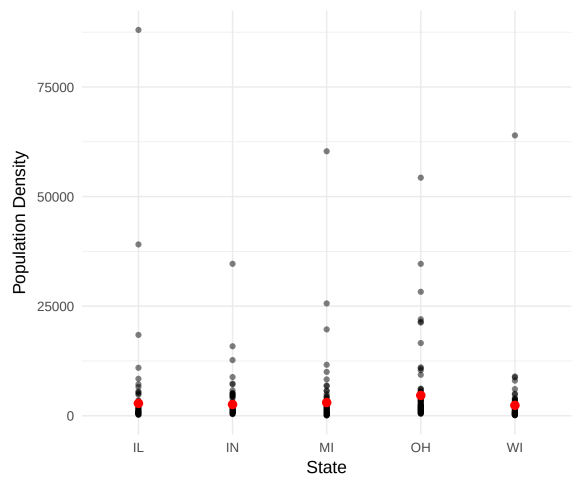
```
## Kable makes output table look pretty (requires package - knitr)
kable(head(midwest))
```

PID	county	state	area	poptotal	popdensity	popwhite	popblack	popamerindian	popasian	popotl
561	ADAMS	IL	0.052	66090	1270.9615	63917	1702	98	249	1
562	ALEXANDER	IL	0.014	10626	759.0000	7054	3496	19	48	
563	BOND	IL	0.022	14991	681.4091	14477	429	35	16	
564	BOONE	IL	0.017	30806	1812.1176	29344	127	46	150	11
565	BROWN	IL	0.018	5836	324.2222	5264	547	14	5	
566	BUREAU	IL	0.050	35688	713.7600	35157	50	65	195	2

Exploratory plots

I will generate some exploratory figures from our dataset first. In the next parts of the tutorial we will be modifying things like panels, font size, colour, panel colour, grid-lines - since most of our ggplot will retain the same structure (we will only be **stylizing** aspects of it), I will 'initialize' each new exploratory plot as a separate object, using <code>obj_name <- ggplot(something something)</code>. This comes in handy if we want to keep adding new 'geometric objects' on top of our original ggplot figure, without copying the code every single time.

```
# What does the state-wise population density look like? I will plot
# each data point (for all countys in each state), and use
# 'stat_summary' to also show the mean and standard deviation of the
# data
g1_population = "red") + theme_minimal(base_size = 13) +
    labs(x = "State", y = "Population Density")
plot(g1_population)
```



Notice how the y-axis has such a wide range but most of our data is in a tiny range - we can log transform the data to 'spread it out'- I will do this for visualizing the population of white people in the different states. - Instead of log transforming my column on the actual dataset, I can simply apply the function to the y variable within aes().

```
g1_popwhite <- ggplot(midwest, aes(x = state, y = log2(popwhite))) + geom_point(alpha = 0.5,
    ) + stat_summary(colour = "red", size = 4) + theme_minimal(base_size = 11) +
    labs(x = "State", y = "Log transformed - Population of Whites")</pre>
```

Maybe I also want to compare the population of whites against black people in the different countys of all states?

- We can fit a 'loess' curve to see specific trends for different states, with the stat_smooth() element.
- This can be modified with the method option in stat_smooth()

```
scatter_popbw <- ggplot(midwest, aes(x = log2(popblack), y = log2(popwhite),
    colour = state)) + geom_point(alpha = 0.5, ) + theme_bw(base_size = 11) +
    stat_smooth() + labs(x = "Log transformed - Population of Blacks",
    y = "Log transformed - Population of Whites")</pre>
```

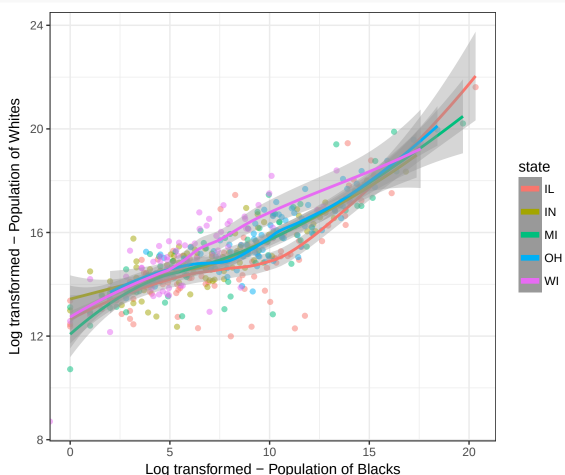
Try the following

- Remove the SD spread (set se = FALSE in stat_smooth()) - Plot the SD spread in the 90% confidence interval (set level = 0.90 in stat_smooth())

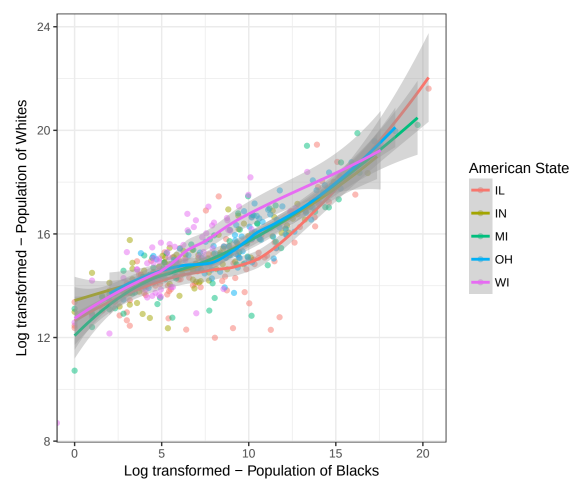
Modifying guides

Notice how, in the legend for colours in our last plot, the points appear faint - this is because by default, we plot them with alpha=0.5. However, we ideally want our points to appear with full opaqueness in our legend. For this, we modify the guides() element.

> Guides for different attributes can be set using this argument. Depending on the attribute we want to modify, we edit it using a nested attribute called guide_legend() (for fill, colour, line etc) or guide_colourbar (for bars of colour gradients).



What did we do? While there are several attributes we can modify using guide_legend, here we simply overwrote some of the default aesthetics for 'colour' in the colour legend for our plot. We can, ofcourse, go ahead and change other things for our colour legend.



Oops- notice how adding guides twice to a plot only keeps the last definition? Best to just define it once. We will also remove the annoying 'grey' colour around the line by setting fill=NA in the override.aes section.

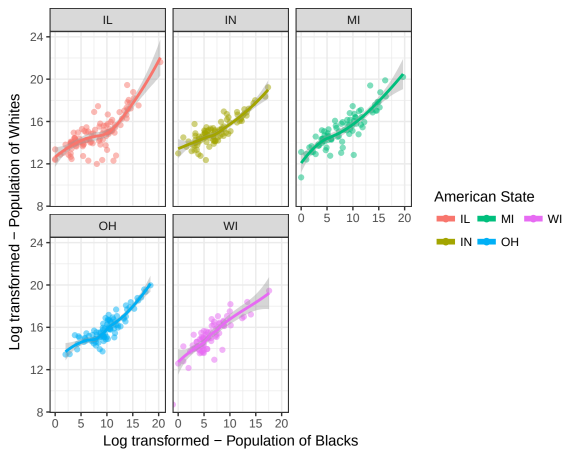
In the last plot, you may be having a hard time distinguishing the different states because of the high amount of overlap in them. We can separate these elements out into multiple ggplots using facet_wrap().

Facet Wrap and Facet Grid - plotting variables

The facet_wrap() function lets us split our data-points into separate groups based on a single column. We can also do this to compare two columns. If we are interested in visualizing all pair-wise comparisons between two columns (even when there may be no data points available for certain pairs of values), we use facet_grid(). The following two plots show the difference between the outputs of facet_wrap() and facet_grid().

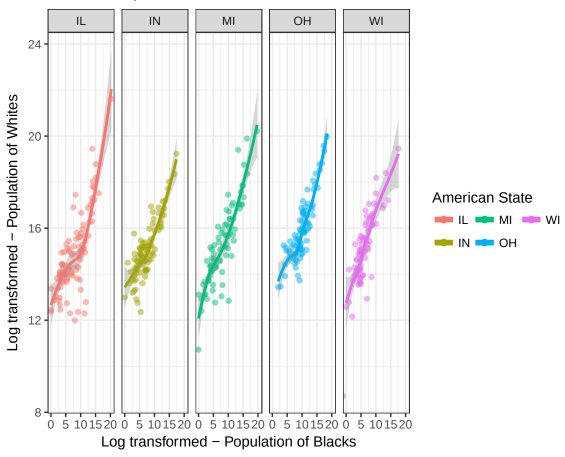
```
# Create Facet Wrap plot
g1_fw <- multiline_labels + ggtitle("Facet Wrap plot for states") + facet_wrap(~state)
# Print this
g1_fw</pre>
```

Facet Wrap plot for states



```
# Create Facet Grid plot
g1_fg <- multiline_labels + ggtitle("Facet Grid plot for states") + facet_grid(~state)
# Print this
g1_fg</pre>
```

Facet Grid plot for states

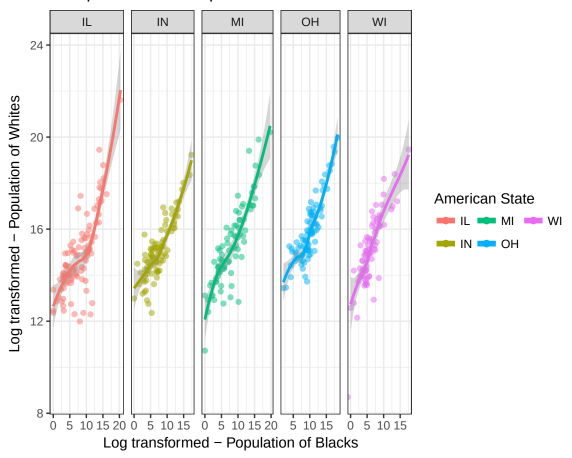


As you can see, facet_grid() is much more readable. We can improve readibility by removing entries along the x-axis that do not have any values in a given 'panel'. This is done using the scales option in facet_grid() - the possible values are "free_x", "free_y", "free". Depending on what you want your reader to focus on, one of these options may be chosen.

We can also make sure the panel width is complementary to the amount of space the data points *actually* take up. We can set this using the space option, which has the same set of possible values as scales.

```
# Create plot object
g1_fg_clean <- multiline_labels + ggtitle("Compact Facet Grid plot") +
    facet_grid(~state, scales = "free_x", space = "free_x")
# Print plot object
g1_fg_clean</pre>
```

Compact Facet Grid plot



removing the space option and see how the plot changes.

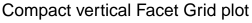
- What happens if you set scales = "free"? Do you see why including a free scale for 'y' might be misleading for readers?

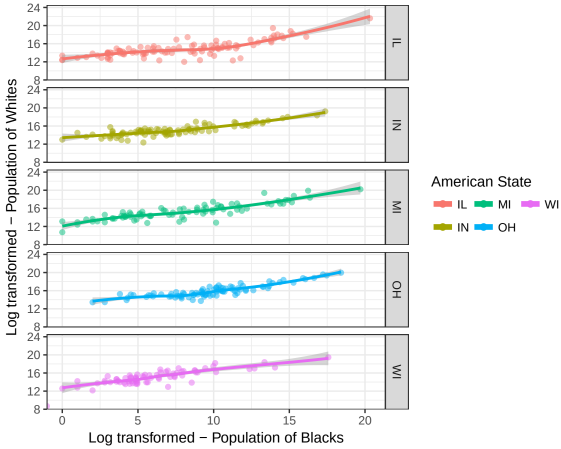
Try

- Do you also see how it is handy to create a new plot object?

Instead of having horizontally aligned columns in facet_grid, we can generate vertically aligned columns by defining the facetting relationship of the interesting variable against the rest of the data, state~., instead of everything versus variable, ~state.

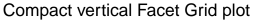
```
# Create plot object
g1_fg_clean_vertical <- multiline_labels + ggtitle("Compact vertical Facet Grid plot") +
    facet_grid(state ~ ., scales = "free_x", space = "free_x")
# Print plot object
g1_fg_clean_vertical</pre>
```

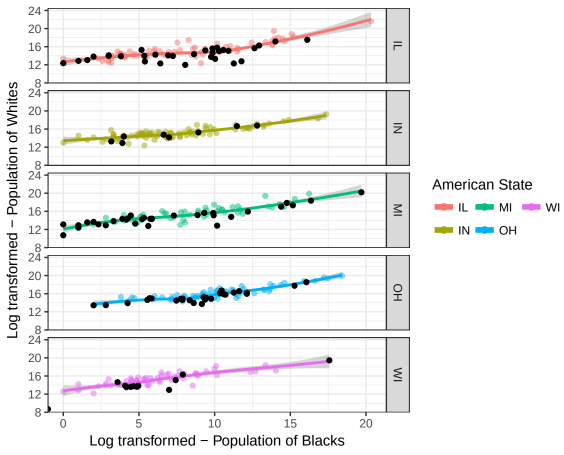




Highlighting a part of the plot

Sometimes we may want to draw attention to only specific entries in the plot. For example, in the plot above, we may want to focus specifically on data points (representative of State countys) where poverty is high (percbelowpoverty > 15). We can explicitly plot these points on top of our existing figure, as shown below:



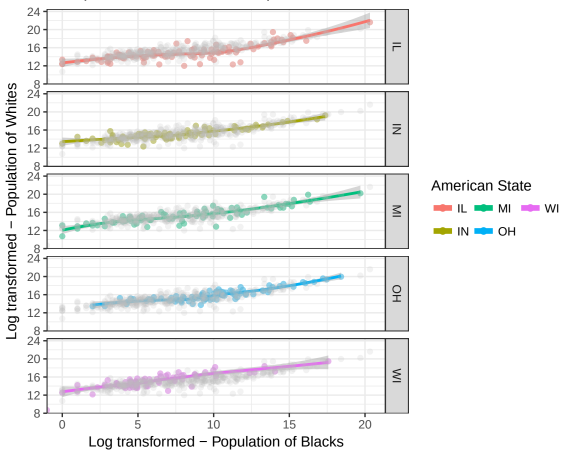


Does this seem to suggest that in areas where the relative proportion of blacks is lower than the whites, the percentage of people below poverty line is relatively higher? - find out by doing grad school!

What did we do here? We took our pre-existing plot where everything had been plotted. We then specified only a *part* of our original dataset to be passed to a new geometric function, a <code>geom_point()</code>, and we set the <code>colour="black"</code> for all the points that this <code>geom_point()</code> is plotting.

Optional: We can also go the other way, where we add **all** the points to **all** the panels as **background**. For this, however, we simply have to make sure that the new data we add into the new <code>geom_point()</code> does not contain the variable we facet on. Thus, we only keep the columns we need for our 'x' and 'y' axes.

Compact vertical Facet Grid plot



• What happens if you add "state" to the list of columns selected in the data attribute of the new geom_point() component?

We have managed to make a plot that is separated by different points of interest, and looks generally readable. However, our legend is taking up a lot of space, and our title doesn't really stand out. The next part, hence, is to stylize our plot.

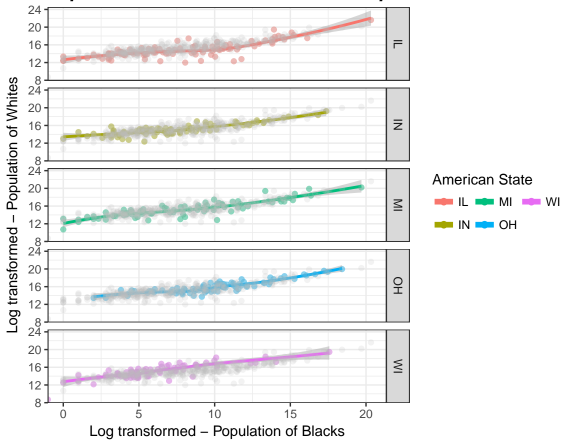
The theme() element

The theme() element in ggplot lets us stylize specific parts of the plot's layout - specifically the following 4:

- element_text(): axis and graph titles and text
- element_lines : components like axis and grid lines
- element_rect(): rectangular components like plots and panel background
- element blank: set other theme components to 'blank'

Firstly, I will edit the text styling for our plot title. This can be done by defining specific font-related attributes for plot.title in the theme component, as shown here:

Compact vertical Facet Grid plot

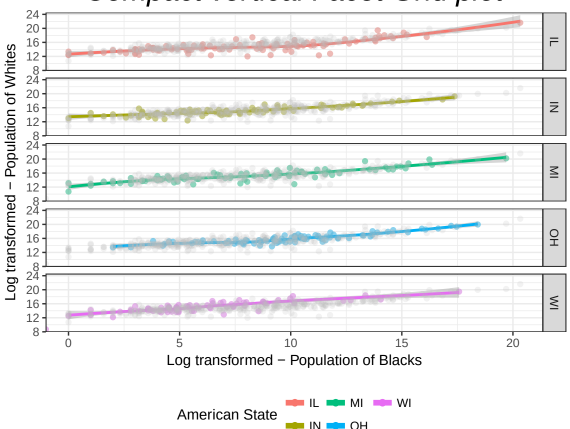


If you are typing the command in the RStudio console, pressing 'tab' while you are within the element_text() part reveals other attributes you can customize.

Follow this link to learn more about customizing the font on your ggplots. It requires an additional package called *extrafont*.

Next, we can change the position of the legend. Notice how this one doesn't require any element_??(). This attribute is defined as one of the following options, "none", "left", "right", "bottom", "top". Feel free to play around with this.

Compact vertical Facet Grid plot



Notice how the keys are in two rows. This goes back to our guides() section, where we defined our colour legend to be printed in 2 rows (nrow=2).

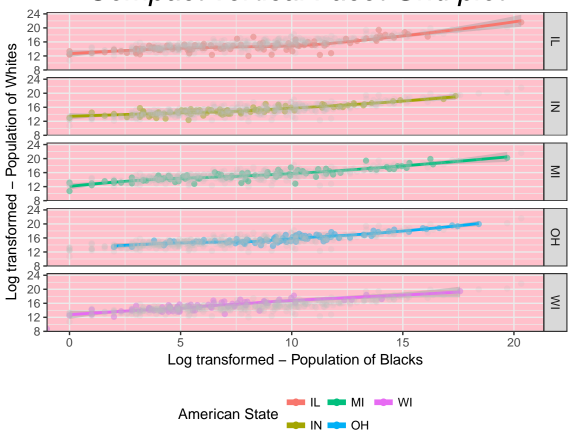
Lastly, say I want to make the background of my plot **pink**. I will modify the **panel.background** option within theme().

```
base_plt + theme(panel.background = "pink")
```

Does that work? Since panel.background is a 'rectangular attribute' that we are stylizing, we need to modify its characteristics using an element_rect option, like below:

```
base_plt + theme(panel.background = element_rect("pink"))
```

Compact vertical Facet Grid plot

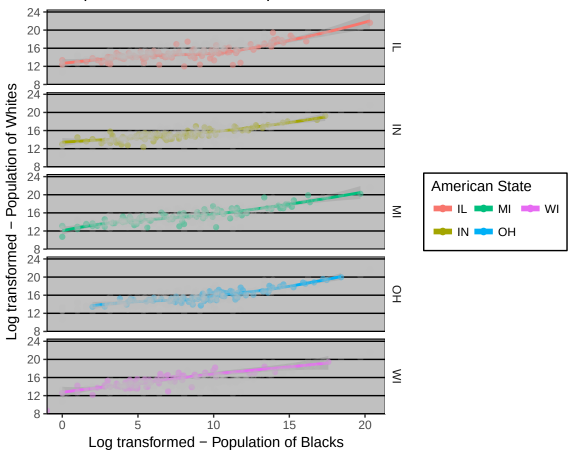


Make my plot look like it was made in Excel

Sometimes, all these fancy shmancy customizations just aren't enough. **ggthemes** is a handy package that lets you quickly customize colour palettes and overall plot appearance using simple commands. The following is a brief sampling of what it has to offer.

```
library(ggthemes)
# Going for an Excel look
base_plt + theme_excel()
```

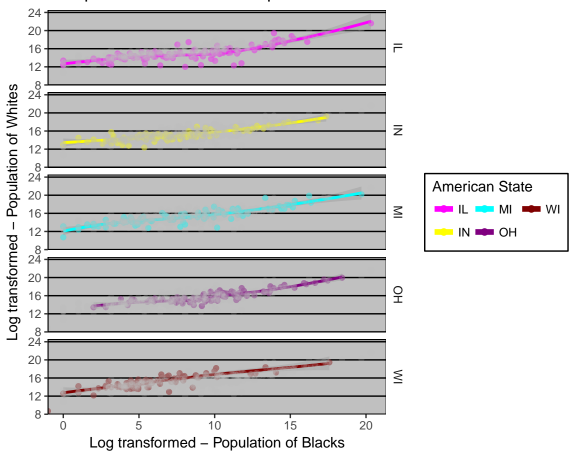
Compact vertical Facet Grid plot



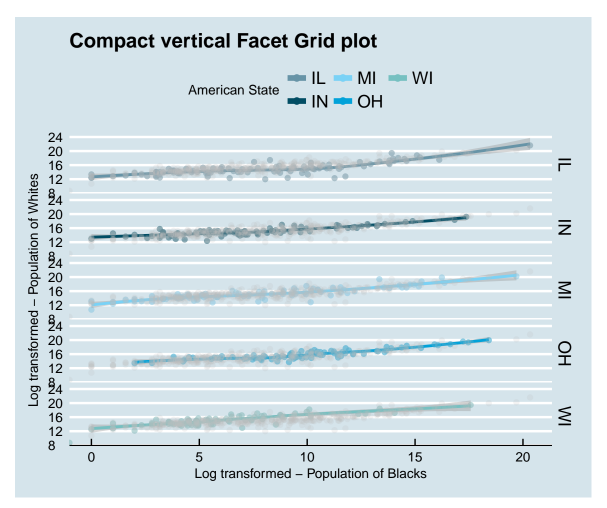
... as ugly as you would expect it to be.

```
# Going for an Excel look with Excel colours
base_plt + theme_excel() + scale_colour_excel()
```

Compact vertical Facet Grid plot



Going for an Economist look with different colours
base_plt + theme_economist() + scale_colour_economist()



Have fun!

Exercises

Can you trace back what the expanded plot definition in base_plt object is? That is, rewrite base_plt as the ggplot() definition it stands for.

Solution:

```
ggplot(midwest, aes(x = log2(popblack), y = log2(popwhite), colour = state)) +
    geom_point(alpha = 0.5, ) + theme_bw(base_size = 11) + stat_smooth() +
    labs(x = "Log transformed - Population of Blacks", y = "Log transformed - Population of Whites") +
    guides(colour = guide_legend(override.aes = list(alpha = 1, size = 2,
        fill = NA), title = "American State", nrow = 2)) + ggtitle("Compact vertical Facet Grid plot")    facet_grid(state ~ ., scales = "free_x", space = "free_x") + geom_point(data = midwest[,
        c("popwhite", "popblack")], colour = "grey", alpha = 0.2) + theme(plot.title = element_text(size = face = "italic", hjust = 0.5), legend.position = "bottom")
```

Take-aways

- 1. Saving ggplots as objects
- 2. Customize panels in ggplot
- 3. Highlight specific data points
- 4. Stylize parts of a plot
- 5. Automating ggplot makeovers with ggthemes package