You know it, I'm sure: You are developing a data transformation using lots of pipes. Then you need to remove one or more lines. You do this using comments.

So far so good. But if it's the last line you want to remove a simple comment will produce an error because of the trailing pipe symbol at the end of the second to last line.

Here's a solution:

Pipes

Magrittr

Let's say you are computing something like this:

15 ## # ... with 28 more rows

```
options(tidyverse.quiet = TRUE)
  library(tidyverse)
  options(dplyr.summarise.inform = FALSE)
 4
  starwars %>%
 5
   mutate(
    mean height = mean(height, na.rm = TRUE)
 7
 8
   group by(species, gender) %>%
   summarize(
10
    mean height species gender = mean(height, na.rm = TRUE),
11
    mean height = first(mean height)
12
   ) 응>응
13
   mutate(
14
    diff mean height = mean_height_species_gender - mean_height
15
16
   select(gender, species, diff mean height) %>%
17
    pivot wider(names_from = 'gender', values_from = 'diff_mean_height',
18 ' - values_fill = NA)
 1 ## # A tibble: 38 x 4
 2 ## # Groups: species [38]
 3 ##
     species masculine feminine `NA`
       4 ##
 5 ## 1 Aleena -95.4 NA NA
                                   NA
 6 ## 2 Besalisk
                  23.6
                          NA
                  23.6 NA NA NA 21.6 NA NA
 7 ## 3 Cerean
 8 ## 4 Chagrian
                  NA
                           -6.36 NA
 9 ## 5 Clawdite
10 ## 6 Droid
                 -34.4 -78.4
                                   NA
11 ## 7 Dug
                  -62.4
                                   NA
                          NA
12 ## 8 Ewok -86.4 NA
13 ## 9 Geonosian 8.64 NA
                                   NA
                                   NA
14 ## 10 Gungan 34.3 NA
                                  NA
```

Now you want to skip the last step pivoting the result. If you just comment out the last line, you'll get an error:

```
starwars %>%
   mutate(
2
   mean height = mean(height, na.rm = TRUE)
3
   group_by(species, gender) %>%
   summarize(
6
    mean height species gender = mean(height, na.rm = TRUE),
7
    mean height = first(mean height)
8
   ) 응>응
9
   mutate(
10
    diff mean height = mean_height_species_gender - mean_height
11
    12
   select(gender, species, diff mean height) %>%
13
   # pivot wider(names from = 'gender', values from = 'diff mean height',
  values fill = NA)
 ## Error: <text>:15:0: unexpected end of input
 ## 13: select(gender, species, diff mean height) %>%
          # pivot wider(names from = 'gender', values from =
 ## 14:
 'diff mean height', values fill = NA)
```

But if you use identity () as your last step everything is fine:

```
starwars %>%
   mutate(
    mean height = mean(height, na.rm = TRUE)
3
   ) 응>응
 4
   group by(species, gender) %>%
 5
   summarize(
 6
    mean height species gender = mean(height, na.rm = TRUE),
7
    mean height = first(mean height)
8
   ) %>%
9
   mutate(
10
    diff mean height = mean height species_gender - mean_height
11
    ) %>%
12
    select(gender, species, diff mean height) %>%
13
    # pivot wider(names from = 'gender', values from = 'diff mean height',
  values fill = NA) %>%
    identity()
1 ## # A tibble: 42 x 3
2 ## # Groups: species [38]
3 ## gender species diff_mean_height
     <chr> <chr> 1 masculine Aleena
 4 ##
                                    <dbl>
                                     -95.4
 5 ## 1 masculine Aleena
```

```
6 ## 2 masculine Besalisk
                                  23.6
 7 ## 3 masculine Cerean
                                  23.6
8 ## 4 masculine Chagrian
                                  21.6
9 ## 5 feminine Clawdite
                                   -6.36
10 ## 6 feminine Droid
                                 -78.4
                                 -34.4
11 ## 7 masculine Droid
12 ## 8 masculine Dug
                                  -62.4
13 ## 9 masculine Ewok
                                 -86.4
14 ## 10 masculine Geonosian
                                   8.64
15 ## # ... with 32 more rows
```

Base R (R >= 4.1.0)

This trick will also work with the new base R pipes:

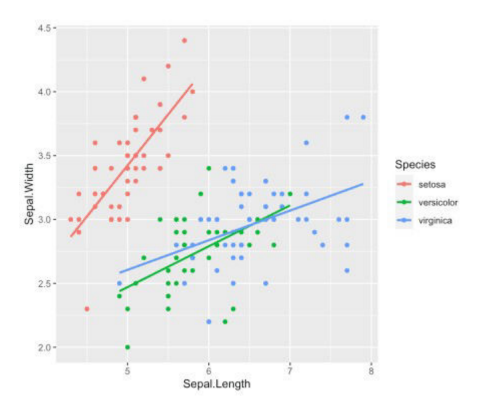
```
starwars |>
    mutate(
    mean height = mean(height, na.rm = TRUE)
 3
   group by (species, gender) |>
 summarize(
    mean height species gender = mean(height, na.rm = TRUE),
 7
    mean height = first(mean height)
 8
    ) |>
 9
   mutate(
10
    diff mean height = mean height species gender - mean height
11
12
   select(gender, species, diff mean height) |>
13
    # pivot wider(names from = 'gender', values from = 'diff mean height',
values_fill = NA) |>
    identity()
 1 ## # A tibble: 42 x 3
 2 ## # Groups: species [38]
 3 ## gender species diff mean height
       <chr> <chr>
 4 ##
                                    <dbl>
 5 ## 1 masculine Aleena
                                   -95.4
 6 ## 2 masculine Besalisk
                                    23.6
 7 ## 3 masculine Cerean
                                   23.6
 8 ## 4 masculine Chagrian
                                    21.6
 9 ## 5 feminine Clawdite
                                   -6.36
10 ## 6 feminine Droid
                                   -78.4
11 ## 7 masculine Droid
                                   -34.4
12 ## 8 masculine Dug
                                   -62.4
13 ## 9 masculine Ewok
                                   -86.4
14 ## 10 masculine Geonosian
                                   8.64
15 ## # ... with 32 more rows
```

ggplot2

The same problem may occur building ggplots:

```
iris %>%
ggplot(aes(Sepal.Length, Sepal.Width, color = Species, group = Species))

description
geom_point() +
geom_smooth(method = "lm", se = FALSE, formula = "y ~ x") +
description
formula = "y ~ x") +
NULL
```



As you can see, I've added ${\tt NULl}$ at the end of the ggplot2-pipe. So I comment out the line before without trouble.