Data Manipulation with R

bmRn CSM: An introduction to the dplyr package

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2020-12-13

Objectives



- 1) Common data manipulation tasks
- 2) dplyr's verbs
- 3) the pipe %>%

Materials



Follow along with the exercises:

https://mjfrigaard.github.io/data-manip-intro-exercises/Index.html

A web version of these slides is located:

https://mjfrigaard.github.io/data-manip-intro/Index.html

The RStudio.Cloud project:

https://rstudio.cloud/project/1918833

What are common data manipulation tasks?



- 1. Viewing the dataset
- 2. Choosing columns/rows
- 3. Ordering rows

- 4. Changing existing columns
- 5. Creating or calculating new columns



dplyr = a grammar for data manipulation

dplyr = "dee + ply + ARRRR"



Pliers are tools for grasping or manipulating common objects

The dplyr package has a variety of verbs for performing common data manipulations



The starwars dataset



These data come from the Star Wars API:



Read more about the data here:

Load the starwars dataset



The starwars data comes from the dplyr package, so we can access it using the code below:

```
install.packages("dplyr")
library(dplyr)
dplyr::starwars
```

We'll use a smaller version of this dataset (original_starwars) to show dplyr's common data manipluation verbs

Import original_starwars

Import the data using the url and readr

```
library(readr)
original_starwars <- read_csv("https://bit.ly/mini-strwrs")</pre>
```

This loads the dataset into our *Environment* pane





dplyr verbs



The primary verbs for data manipulation in dplyr:

```
glimpse()
select()
filter()
arrange()
mutate()
```



Viewing the data = glimpse()

We need to view the data we're manipulating to see if the changes are correct

View = glimpse()

Take a look at the entire dataset using dplyr::glimpse()

```
glimpse(original_starwars)

## Rows: 6
## Columns: 6
```

glimpse() transposes the data and prints as much of it to the screen as
possible



View the data in the Console

Enter the name of the dataset to print it to the Console

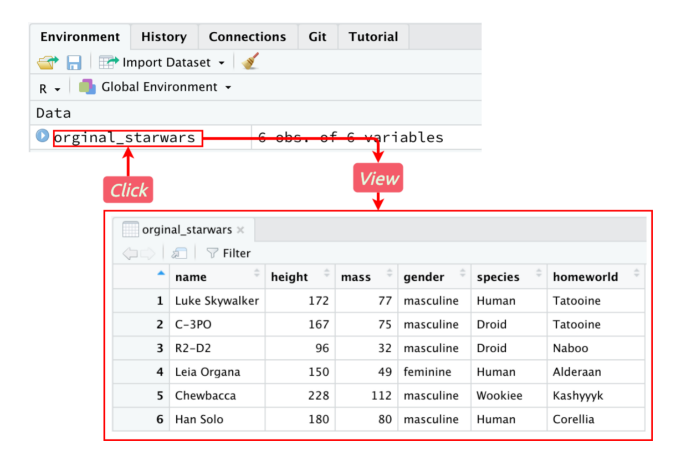
original_starwars

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
Luke Skywalker	172	77 blond	Human	Tatooine
C-3PO	167	75 NA	Droid	Tatooine
R2-D2	96	32 NA	Droid	Naboo
Leia Organa	150	49 brown	Human	Alderaan
Chewbacca	228	112 brown	Wookiee	Kashyyyk
Han Solo	180	80 brown	Human	Corellia
6 rows				



View the data in the Data Viewer

View the original_starwars dataset using RStudio's data editor







Choosing columns = select()

Choose columns with select()



select() allows us to pick specific columns out of a dataset

select(original_starwars, name, homeworld, species)

name	homeworld	species
<chr></chr>	<chr></chr>	<chr></chr>
Luke Skywalker	Tatooine	Human
C-3PO	Tatooine	Droid
R2-D2	Naboo	Droid
Leia Organa	Alderaan	Human
Chewbacca	Kashyyyk	Wookiee
Han Solo	Corellia	Human
6 rows		

Choose columns with select()



We can use negation (-) to remove columns

select(original_starwars, -c(mass, height, hair_color))

name	species	homeworld
<chr></chr>	<chr></chr>	<chr></chr>
Luke Skywalker	Human	Tatooine
C-3PO	Droid	Tatooine
R2-D2	Droid	Naboo
Leia Organa	Human	Alderaan
Chewbacca	Wookiee	Kashyyyk
Han Solo	Human	Corellia
6 rows		

select() helpers



select() comes with 'helpers' to make chosing columns easier (and reduces
typing!)

Helper	Outputs
starts_with()	choose columns starting with
ends_with()	choose columns ending with
contains	choose columns with names containing
<pre>matches()</pre>	choose columns matching regex
one_of()	choose columns from a set of names
num_range()	choose columns from a numerical index

Choose columns with select()



Select columns using matches()

select(original_starwars, name, matches("_"))

name	hair_color
<chr></chr>	<chr></chr>
Luke Skywalker	blond
C-3PO	NA
R2-D2	NA
Leia Organa	brown
Chewbacca	brown
Han Solo	brown
6 rows	



See the select() exercises for more examples!



filter() lets us pull out rows based on logical conditions

filter(original_starwars, species == "Human")

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
Luke Skywalker	172	77 blond	Human	Tatooine
Leia Organa	150	49 brown	Human	Alderaan
Han Solo	180	80 brown	Human	Corellia
3 rows				



filter() logical conditions include:

Logical Test	Outputs
<	Less than
>	Greater than
==	Equal to
<=	Less than or equal to
>=	Greater than or equal to
!=	Not equal to
%in%	Group membership
is.na()	is NA (missing)
!is.na()	is not NA (non-missing)



Combine logical condtions with & or,

this gets the same results...

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
Luke Skywalker	172	77 blond	Human	Tatooine
Leia Organa	150	49 brown	Human	Alderaan
Han Solo	180	80 brown	Human	Corellia
3 rows				





Combine logical condtions with & or,

...as this

```
filter(original_starwars,
     species == "Human" , !is.na(hair_color))
```

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
Luke Skywalker	172	77 blond	Human	Tatooine
Leia Organa	150	49 brown	Human	Alderaan
Han Solo	180	80 brown	Human	Corellia
2				

3 rows



Remember that *any* logical condition works for **filter()**ing, so we can borrow functions from other packages to help us

The stringr::str_detect() function returns a logical condition, so we can use it *inside* filter()

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
C-3PO	167	75 NA	Droid	Tatooine
R2-D2	96	32 NA	Droid	Naboo
2 rows				



See the filter() exercises for more examples!



Sorting rows with arrange()

Sort rows with arrange()



arrange() sorts the contents of a dataset (ascending or descending)

arrange(original_starwars, height)

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
R2-D2	96	32 NA	Droid	Naboo
Leia Organa	150	49 brown	Human	Alderaan
C-3PO	167	75 NA	Droid	Tatooine
Luke Skywalker	172	77 blond	Human	Tatooine
Han Solo	180	80 brown	Human	Corellia
Chewbacca	228	112 brown	Wookiee	Kashyyyk
6 rows				

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Sort rows with arrange()



arrange()'s default is to sort ascending--include desc() to sort descending

arrange(original_starwars, desc(height))

name	height	mass hair_color	species	homeworld
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>
Chewbacca	228	112 brown	Wookiee	Kashyyyk
Han Solo	180	80 brown	Human	Corellia
Luke Skywalker	172	77 blond	Human	Tatooine
C-3PO	167	75 NA	Droid	Tatooine
Leia Organa	150	49 brown	Human	Alderaan
R2-D2	96	32 NA	Droid	Naboo
6 rows				



See the arrange () exercises for more examples!

Create columns with mutate()

mutate() allows us to create new columns



name	height	m hair_color	species	homewo	bmi
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>	<dbl></dbl>
Luke Skywalker	172	77 blond	Human	Tatooine	26.02758
C-3PO	167	75 NA	Droid	Tatooine	26.89232
R2-D2	96	32 NA	Droid	Naboo	34.72222
Leia Organa	150	49 brown	Human	Alderaan	21.77778
Chewbacca	228	112 brown	Wookiee	Kashyyyk	21.54509
Han Solo	180	80 brown	Human	Corellia	24.69136
6 rows					

Create columns with mutate()

mutate() allows us to change existing columns, too



name	height	mass hair_color	species	homeworld	bmi
<chr></chr>	<dbl></dbl>	<dbl> <chr></chr></dbl>	<chr></chr>	<chr></chr>	<dbl></dbl>
Luke Skywalker	172	77 blond	Human	Tatooine	26
C-3PO	167	75 NA	Droid	Tatooine	27
R2-D2	96	32 NA	Droid	Naboo	35
Leia Organa	150	49 brown	Human	Alderaan	22
Chewbacca	228	112 brown	Wookiee	Kashyyyk	22
Han Solo	180	80 brown	Human	Corellia	25
6 rows					

6 rows



See the mutate() exercises for more examples!



Write clearer code with the pipe

%>%

The pipe (%>%)



The pipe comes from the magrittr package:

https://magrittr.tidyverse.org/

The pipe makes our code easier to read (and write)

Create pipes easily with keyboard shortcuts

Mac Windows

Cmd + Shift + M Crtl + Shift + M

How the pipe (%>%) works



Without the pipe, we have to constantly assign the output to new object:

```
first_output <- first_function(input)
second_output <- second_function(first_output)</pre>
```

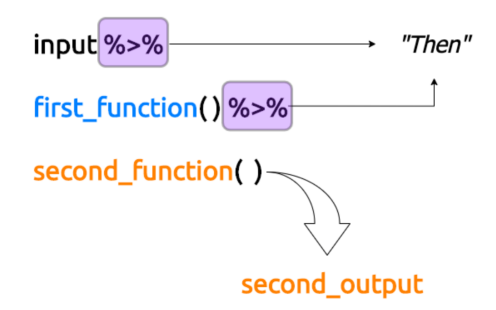
Or use nested functions:

```
second_output <- second_function(first_function(input), first_output)</pre>
```

How the pipe (%>%) works



The pipe allows us to pass the output from functions left-to-right



%>% can be read as "then"

Creating pipelines of functions



Review the code below and think about what each object contains:

- 1. Filter original_starwars to only brown-haired characters over 100 cm tall
- 2. Create a bmi column using: mass / ((height / 100) ^ 2)
- 3. Select name, bmi, and homeworld
- 4. Arrange the data by bmi, descending

Creating pipelines of functions

Re-write these functions into a pipeline, ending with a single output (new_original_starwars)

- 1. Filter original_starwars to only brown-haired characters over 100 cm tall
- 2. Create a bmi column using: mass / ((height / 100) ^ 2)
- 3. Select name, bmi, and homeworld
- 4. Arrange the data by bmi, descending

```
original_starwars %>%
  filter(hair_color == "____" & height > ___) %>%
  mutate(___ = mass / ((height / 100) ^ 2)) %>%
  select(___, bmi, ____) %>%
  arrange(___(bmi)) -> new_original_starwars
```



Creating pipelines of functions

The answer is below:

```
original_starwars %>%
  filter(hair_color == "brown" & height > 100) %>%
  mutate(bmi = mass / ((height / 100) ^ 2)) %>%
  select(name, bmi, homeworld) %>%
  arrange(desc(bmi)) -> new_original_starwars
new_original_starwars
```

name	bmi homeworld
<chr></chr>	<dbl> <chr></chr></dbl>
Han Solo	24.69136 Corellia
Leia Organa	21.77778 Alderaan
Chewbacca	21.54509 Kashyyyk
3 rows	





See the pipe exercises for more examples!

Resources for Data Manipluation

- 1. R for Data Science
- 2. Data Wrangling with R
- 3. Stack Overflow questions tagged with dplyr
- 4. RStudio Community posts tagged dplyr

