dplyr tutorial

Warning: package 'knitr' was built under R version 3.1.3

What is dplyr?

dplyr is a powerful R-package to transform and summarize tabular data with rows and columns. For another explanation of dplyr see the dplyr package vignette: Introduction to dplyr

Why is it useful?

The package contains a set of functions (or "verbs") that perform common data manipulation operations such as filtering for rows, selecting specific columns, re-ordering rows, adding new columns and summarizing data.

In addition, dplyr contains a useful function to perform another common task which is the "split-apply-combine" concept. We will discuss that in a little bit.

How does it compare to using base functions R?

If you are familiar with R, you are probably familiar with base R functions such as split(), subset(), apply(), sapply(), lapply(), tapply() and aggregate(). Compared to base functions in R, the functions in dplyr are easier to work with, are more consistent in the syntax and are targeted for data analysis around data frames instead of just vectors.

How do I get dplyr?

To install dplyr

```
install.packages("dplyr")
```

To load dplyr

```
library(dplyr)
```

Warning: package 'dplyr' was built under R version 3.1.3

Data: mammals sleep

The msleep (mammals sleep) data set contains the sleeptimes and weights for a set of mammals and is available in the dagdata repository on github. This data set contains 83 rows and 11 variables.

Download the msleep data set in CSV format from here, and then load into R:

```
library(downloader)
```

Warning: package 'downloader' was built under R version 3.1.3

```
url <- "https://raw.githubusercontent.com/genomicsclass/dagdata/master/inst/extdata/msleep_ggplot2.csv"
filename <- "msleep_ggplot2.csv"
if (!file.exists(filename)) download(url,filename)
msleep <- read.csv("msleep_ggplot2.csv")
head(msleep)</pre>
```

##			nai	ne g	enus	vore		order	conservation
##	1		Cheeta	ah Acin	onyx	carni	(Carnivora	lc
##	2		Owl monk	ey A	otus	omni		Primates	<na></na>
##	3	N	Mountain beav	er Aplodo	ntia	herbi		Rodentia	nt
##	4	Greater shor	rt-tailed shr	ew Bla	rina	omni	Sor	icomorpha	lc
##	5		Co	υ	Bos	herbi	Art	iodactyla	domesticated
##	6	Tì	ree-toed slo	th Brad	ypus	herbi		Pilosa	<na></na>
##		sleep_total	sleep_rem sle	eep_cycle	awak	e brai	inwt	bodywt	
##	1	12.1	NA	NA	11.	9	NA	50.000	
##	2	17.0	1.8	NA	7.	0 0.01	1550	0.480	
##	3	14.4	2.4	NA	9.	6	NA	1.350	
##	4	14.9	2.3	0.1333333	9.	1 0.00	0029	0.019	
##	5	4.0	0.7	0.6666667	20.	0 0.42	2300	600.000	
##	6	14.4	2.2	0.7666667	9.	6	NA	3.850	

The columns (in order) correspond to the following:

column name	Description
name	common name
genus	taxonomic rank
vore	carnivore, omnivore or herbivore?
order	taxonomic rank
conservation	the conservation status of the mammal
$sleep_total$	total amount of sleep, in hours
$sleep_rem$	rem sleep, in hours
sleep_cycle	length of sleep cycle, in hours
awake	amount of time spent awake, in hours
brainwt	brain weight in kilograms
bodywt	body weight in kilograms

Important dplyr verbs to remember

dplyr verbs	Description
select()	select columns
filter()	filter rows
arrange()	re-order or arrange rows
<pre>mutate()</pre>	create new columns
<pre>summarise()</pre>	summarise values
<pre>group_by()</pre>	allows for group operations in the "split-apply-combine" concept

dplyr verbs in action

The two most basic functions are select() and filter() which selects columns and filters rows, respectively.

Selecting columns using select()

Select a set of columns: the name and the sleep_total columns.

```
sleepData <- select(msleep, name, sleep_total)
head(sleepData)</pre>
```

```
##
                            name sleep_total
## 1
                                         12.1
                         Cheetah
## 2
                      Owl monkey
                                         17.0
## 3
                                         14.4
                Mountain beaver
## 4 Greater short-tailed shrew
                                         14.9
                                          4.0
## 5
                             Cow
## 6
               Three-toed sloth
                                         14.4
```

To select all the columns *except* a specific column, use the "-" (subtraction) operator (also known as negative indexing)

```
head(select(msleep, -name))
```

```
##
          genus vore
                              order conservation sleep_total sleep_rem
## 1
       Acinonyx carni
                          Carnivora
                                              lc
                                                         12.1
                                                                     NA
## 2
                                                         17.0
                                                                    1.8
          Aotus omni
                          Primates
                                            <NA>
## 3 Aplodontia herbi
                          Rodentia
                                              nt
                                                         14.4
                                                                    2.4
        Blarina omni Soricomorpha
                                                         14.9
                                                                    2.3
## 4
                                              lc
## 5
            Bos herbi Artiodactyla domesticated
                                                          4.0
                                                                    0.7
## 6
       Bradypus herbi
                             Pilosa
                                            <NA>
                                                         14.4
                                                                    2.2
##
     sleep_cycle awake brainwt
                                 bodywt
## 1
              NA
                  11.9
                             NA
                                 50.000
## 2
                   7.0 0.01550
              NA
                                  0.480
## 3
              NA
                   9.6
                                  1.350
## 4
       0.1333333
                   9.1 0.00029
                                  0.019
## 5
       0.6666667
                  20.0 0.42300 600.000
       0.7666667
                                  3.850
## 6
                   9.6
                             NA
```

To select a range of columns by name, use the ":" (colon) operator

```
head(select(msleep, name:order))
```

```
##
                           name
                                     genus vore
                                                         order
## 1
                        Cheetah
                                  Acinonyx carni
                                                     Carnivora
## 2
                     Owl monkey
                                     Aotus omni
                                                      Primates
                Mountain beaver Aplodontia herbi
                                                      Rodentia
                                   Blarina omni Soricomorpha
## 4 Greater short-tailed shrew
                            Cow
## 5
                                       Bos herbi Artiodactyla
## 6
               Three-toed sloth
                                  Bradypus herbi
                                                        Pilosa
```

To select all columns that start with the character string "sl", use the function starts_with()

head(select(msleep, starts_with("sl")))

```
##
     sleep_total sleep_rem sleep_cycle
## 1
            12.1
                         NA
## 2
            17.0
                        1.8
                                      NA
## 3
            14.4
                        2.4
                                      NA
            14.9
                        2.3
                              0.1333333
## 4
## 5
             4.0
                        0.7
                              0.6666667
## 6
            14.4
                        2.2
                              0.7666667
```

Some additional options to select columns based on a specific criteria include

- 1. ends_with() = Select columns that end with a character string
- 2. contains() = Select columns that contain a character string
- 3. matches() = Select columns that match a regular expression
- 4. one_of() = Select columns names that are from a group of names

Selecting rows using filter()

Filter the rows for mammals that sleep a total of more than 16 hours.

filter(msleep, sleep_total >= 16)

##			name	genus	vore		order	conservation
##	1	(Owl monkey	Aotus	omni	P	rimates	<na></na>
##	2	Long-nosed	armadillo	Dasypus	carni	Ci	ngulata	lc
##	3	North America	an Opossum	Didelphis	omni	Didelphi	morphia	lc
##	4	Big	brown bat	Eptesicus	insecti	Chi	roptera	lc
##	5	Thick-taile	ed opposum	Lutreolina	carni	Didelphi	morphia	lc
##	6	Little	brown bat	Myotis	insecti	Chi	roptera	<na></na>
##	7	Giant	armadillo	Priodontes	insecti	Ci	ngulata	en
##	8	Arctic ground	d squirrel	Spermophilus	herbi	R	odentia	lc
##		sleep_total :	sleep_rem :	sleep_cycle a	wake bra	inwt body	wt	
##	1	17.0	1.8	NA	7.0 0.0	1550 0.4	80	
##	2	17.4	3.1	0.3833333	6.6 0.0	1080 3.5	00	
##	3	18.0	4.9	0.3333333	6.0 0.0	0630 1.7	00	
##	4	19.7	3.9	0.1166667	4.3 0.0	0.0 0.0	23	
##	5	19.4	6.6	NA	4.6	NA 0.3	70	
##	6	19.9	2.0	0.2000000	4.1 0.0	0025 0.0	10	
##	7	18.1	6.1	NA	5.9 0.0	8100 60.0	00	
##	8	16.6	NA	NA	7.4 0.0	0570 0.9	20	

Filter the rows for mammals that sleep a total of more than 16 hours and have a body weight of greater than 1 kilogram.

```
filter(msleep, sleep_total >= 16, bodywt >= 1)
```

```
##
                                  genus
                                           vore
                                                           order conservation
                        name
## 1
       Long-nosed armadillo
                                Dasypus
                                                                            lc
                                           carni
                                                       Cingulata
## 2 North American Opossum Didelphis
                                           omni Didelphimorphia
                                                                            lc
## 3
            Giant armadillo Priodontes insecti
                                                       Cingulata
                                                                            en
##
     sleep_total sleep_rem sleep_cycle awake brainwt bodywt
## 1
                        3.1
                              0.3833333
                                           6.6
                                                0.0108
                                                          3.5
            17.4
## 2
            18.0
                        4.9
                              0.3333333
                                           6.0
                                                0.0063
                                                          1.7
## 3
            18.1
                        6.1
                                           5.9 0.0810
                                                         60.0
                                     NA
```

Filter the rows for mammals in the Perissodactyla and Primates taxonomic order

```
filter(msleep, order %in% c("Perissodactyla", "Primates"))
```

##		name	genus	vore		ord	der con	servation
##	1	Owl monkey	Aotus	omni		Primat	tes	<na></na>
##	2	Grivet	Cercopithecus	omni		Primat	tes	lc
##	3	Horse	Equus	herbi	Peri	issodacty	yla dom	esticated
##	4	Donkey	Equus	herbi	Peri	issodacty	yla dom	esticated
##	5	Patas monkey	Erythrocebus	omni		Primat	tes	lc
##	6	Galago	Galago	omni		Primat	tes	<na></na>
##	7	Human	Homo	omni		Primat	tes	<na></na>
##	8	Mongoose lemur	Lemur	herbi		Primat	tes	vu
##	9	Macaque	Macaca	omni		Primat	tes	<na></na>
##	10	Slow loris	Nyctibeus	carni		Primat	tes	<na></na>
##	11	Chimpanzee	Pan	omni		Primat	tes	<na></na>
##	12	Baboon	Papio	omni		Primat	tes	<na></na>
##	13	Potto	Perodicticus	omni		Primat	tes	lc
##	14	Squirrel monkey	Saimiri	omni		Primat	tes	<na></na>
##	15	Brazilian tapir	Tapirus	herbi	Peri	issodacty	yla	vu
##		sleep_total slee	ep_rem sleep_c	ycle a	wake	${\tt brainwt}$	bodyw	t
##	1	17.0	1.8	NA	7.0	0.0155	0.48	0
##	2	10.0	0.7	NA	14.0	NA	4.75	0
##	3	2.9	0.6 1.000	0000	21.1	0.6550	521.00	0
##	4	3.1	0.4	NA	20.9	0.4190	187.00	0
##	5	10.9	1.1	NA	13.1	0.1150	10.00	0
##	6	9.8	1.1 0.550	0000	14.2	0.0050	0.20	0
##	7	8.0	1.9 1.500	0000	16.0	1.3200	62.00	0
##	8	9.5	0.9	NA	14.5	NA	1.67	0
##	9	10.1	1.2 0.750	0000	13.9	0.1790	6.80	0
##	10	11.0	NA	NA	13.0	0.0125	1.40	0
##	11	9.7	1.4 1.416	6667	14.3	0.4400	52.20	0
##	12	9.4	1.0 0.666	6667	14.6	0.1800	25.23	5
##	13	11.0	NA	NA	13.0	NA	1.10	0
##	14	9.6	1.4	NA	14.4	0.0200	0.74	3
##	15	4.4	1.0 0.900	0000	19.6	0.1690	207.50	1

You can use the boolean operators (e.g. >, <, >=, <=, !=, %in%) to create the logical tests.

Pipe operator: %>%

Before we go any futher, let's introduce the pipe operator: %>%. dplyr imports this operator from another package (magrittr). This operator allows you to pipe the output from one function to the input of another

function. Instead of nesting functions (reading from the inside to the outside), the idea of of piping is to read the functions from left to right.

Here's an example you have seen:

```
head(select(msleep, name, sleep_total))
```

##		name	sleep_total
##	1	Cheetah	12.1
##	2	Owl monkey	17.0
##	3	Mountain beaver	14.4
##	4	Greater short-tailed shrew	14.9
##	5	Cow	4.0
##	6	Three-toed sloth	14.4

Now in this case, we will pipe the msleep data frame to the function that will select two columns (name and sleep_total) and then pipe the new data frame to the function head() which will return the head of the new data frame.

```
msleep %>%
    select(name, sleep_total) %>%
    head
```

##		name	sleep_total
##	1	Cheetah	12.1
##	2	Owl monkey	17.0
##	3	Mountain beaver	14.4
##	4	Greater short-tailed shrew	14.9
##	5	Cow	4.0
##	6	Three-toed sloth	14.4

You will soon see how useful the pipe operator is when we start to combine many functions.

Back to dplyr verbs in action

Now that you know about the pipe operator (%>%), we will use it throughout the rest of this tutorial.

Arrange or re-order rows using arrange()

To arrange (or re-order) rows by a particular column such as the taxonomic order, list the name of the column you want to arrange the rows by

```
msleep %>% arrange(order) %>% head
```

```
##
                                      order conservation sleep_total sleep_rem
         name
                  genus
                        vore
## 1
                 Tenrec omni Afrosoricida
                                                                 15.6
       Tenrec
                                                    <NA>
                                                                            2.3
## 2
                    Bos herbi Artiodactyla domesticated
                                                                  4.0
                                                                            0.7
## 3 Roe deer Capreolus herbi Artiodactyla
                                                      lc
                                                                  3.0
                                                                             NA
                  Capri herbi Artiodactyla
                                                      lc
                                                                  5.3
                                                                            0.6
         Goat
                Giraffa herbi Artiodactyla
                                                                            0.4
## 5 Giraffe
                                                      cd
                                                                  1.9
```

```
## 6
                   Ovis herbi Artiodactyla domesticated
                                                                  3.8
                                                                            0.6
##
     sleep_cycle awake brainwt
                                 bodywt
## 1
              NA
                   8.4
                       0.0026
                                  0.900
                        0.4230 600.000
## 2
       0.6666667
                  20.0
## 3
              NA
                  21.0
                        0.0982
                                 14.800
## 4
                  18.7
                        0.1150
                                 33.500
              NA
## 5
                  22.1
                            NA 899.995
              NA
                  20.2 0.1750 55.500
## 6
              NA
```

Now, we will select three columns from msleep, arrange the rows by the taxonomic order and then arrange the rows by sleep_total. Finally show the head of the final data frame

```
msleep %>%
    select(name, order, sleep_total) %>%
    arrange(order, sleep_total) %>%
    head
```

```
##
                      order sleep_total
         name
## 1
       Tenrec Afrosoricida
                                    15.6
      Giraffe Artiodactyla
                                     1.9
## 2
## 3 Roe deer Artiodactyla
                                     3.0
## 4
        Sheep Artiodactyla
                                     3.8
## 5
          Cow Artiodactyla
                                     4.0
## 6
         Goat Artiodactyla
                                     5.3
```

Same as above, except here we filter the rows for mammals that sleep for 16 or more hours instead of showing the head of the final data frame

```
msleep %>%
    select(name, order, sleep_total) %>%
    arrange(order, sleep_total) %>%
    filter(sleep_total >= 16)
```

```
##
                                        order sleep_total
                        name
## 1
              Big brown bat
                                   Chiroptera
                                                     19.7
## 2
           Little brown bat
                                   Chiroptera
                                                     19.9
## 3
       Long-nosed armadillo
                                   Cingulata
                                                     17.4
## 4
            Giant armadillo
                                   Cingulata
                                                     18.1
## 5 North American Opossum Didelphimorphia
                                                     18.0
## 6
       Thick-tailed opposum Didelphimorphia
                                                     19.4
                                                     17.0
## 7
                 Owl monkey
                                    Primates
## 8 Arctic ground squirrel
                                    Rodentia
                                                     16.6
```

Something slightly more complicated: same as above, except arrange the rows in the sleep_total column in a descending order. For this, use the function desc()

```
msleep %>%
    select(name, order, sleep_total) %>%
    arrange(order, desc(sleep_total)) %>%
    filter(sleep_total >= 16)
```

```
##
                                        order sleep_total
                        name
## 1
           Little brown bat
                                  Chiroptera
                                                     19.9
              Big brown bat
                                                     19.7
## 2
                                  Chiroptera
## 3
            Giant armadillo
                                   Cingulata
                                                     18.1
## 4
       Long-nosed armadillo
                                   Cingulata
                                                     17.4
## 5
       Thick-tailed opposum Didelphimorphia
                                                     19.4
## 6 North American Opossum Didelphimorphia
                                                     18.0
## 7
                 Owl monkey
                                    Primates
                                                     17.0
## 8 Arctic ground squirrel
                                    Rodentia
                                                     16.6
```

Create new columns using mutate()

The mutate() function will add new columns to the data frame. Create a new column called rem_proportion which is the ratio of rem sleep to total amount of sleep.

```
msleep %>%
  mutate(rem_proportion = sleep_rem / sleep_total) %>%
  head
```

##				name	ge	enus	vore)	order	conservation
##	1		Ch	eetah	Acin	onyx	carni	_ (Carnivora	lc
##	2		Owl m	onkey	A	otus	omni	_	Primates	<na></na>
##	3		Mountain b	eaver	Aplodo	ntia	herbi	-	Rodentia	. nt
##	4	Greater sl	hort-tailed	shrew	Bla	rina	omni	Sor	icomorpha	lc
##	5			Cow		Bos	herbi	Art	iodactyla	domesticated
##	6		Three-toed	sloth	Brady	ypus	herbi	-	Pilosa	<na></na>
##		sleep_tota	al sleep_rem	slee	p_cycle	awak	ce bra	inwt	bodywt	rem_proportion
##	1	12	. 1 NA		NA	11.	9	NA	50.000	NA
##	2	17	.0 1.8	3	NA	7.	0.0	1550	0.480	0.1058824
##	3	14	.4 2.4	•	NA	9.	6	NA	1.350	0.1666667
##	4	14	.9 2.3	0.	1333333	9.	1 0.0	0029	0.019	0.1543624
##	5	4	.0 0.7	0.0	6666667	20.	0 0.4	12300	600.000	0.1750000
##	6	14	.4 2.2	0.	7666667	9.	6	NA	3.850	0.1527778

You can many new columns using mutate (separated by commas). Here we add a second column called bodywt_grams which is the bodywt column in grams.

```
##
                                      genus vore
                                                          order conservation
                            name
## 1
                                                      Carnivora
                                                                           lc
                         Cheetah
                                   Acinonyx carni
## 2
                     Owl monkey
                                      Aotus
                                             omni
                                                       Primates
                                                                         <NA>
## 3
                Mountain beaver Aplodontia herbi
                                                       Rodentia
                                                                           nt
## 4 Greater short-tailed shrew
                                    Blarina omni Soricomorpha
                                                                           lc
## 5
                                        Bos herbi Artiodactyla domesticated
                             Cow
## 6
               Three-toed sloth
                                                         Pilosa
                                                                         <NA>
                                   Bradypus herbi
##
     sleep_total sleep_rem sleep_cycle awake brainwt
                                                        bodywt rem_proportion
## 1
            12.1
                                         11.9
                                                    NA
                                                        50.000
                        NA
                                     NA
                                                                            NA
## 2
            17.0
                                          7.0 0.01550
                                                         0.480
                        1.8
                                     NA
                                                                     0.1058824
```

```
## 3
             14.4
                         2.4
                                       NA
                                            9.6
                                                      NA
                                                            1.350
                                                                        0.1666667
                                                           0.019
## 4
             14.9
                         2.3
                               0.1333333
                                            9.1 0.00029
                                                                        0.1543624
## 5
              4.0
                         0.7
                               0.6666667
                                           20.0 0.42300 600.000
                                                                        0.1750000
                         2.2
                               0.7666667
                                                                        0.1527778
## 6
             14.4
                                            9.6
                                                      NA
                                                            3.850
##
     bodywt grams
## 1
             50000
## 2
               480
## 3
              1350
## 4
                19
            600000
## 5
## 6
              3850
```

Create summaries of the data frame using summarise()

The summarise() function will create summary statistics for a given column in the data frame such as finding the mean. For example, to compute the average number of hours of sleep, apply the mean() function to the column sleep_total and call the summary value avg_sleep.

```
msleep %>%
    summarise(avg_sleep = mean(sleep_total))

## avg_sleep
## 1 10.43373
```

There are many other summary statistics you could consider such sd(), min(), max(), median(), sum(), n() (returns the length of vector), first() (returns first value in vector), last() (returns last value in vector) and n_distinct() (number of distinct values in vector).

```
## avg_sleep min_sleep max_sleep total
## 1 10.43373 1.9 19.9 83
```

Group operations using group_by()

The <code>group_by()</code> verb is an important function in dplyr. As we mentioned before it's related to concept of "split-apply-combine". We literally want to split the data frame by some variable (e.g. taxonomic order), apply a function to the individual data frames and then combine the output.

Let's do that: split the msleep data frame by the taxonomic order, then ask for the same summary statistics as above. We expect a set of summary statistics for each taxonomic order.

```
msleep %>%
   group_by(order) %>%
   summarise(avg_sleep = mean(sleep_total),
        min_sleep = min(sleep_total),
        max_sleep = max(sleep_total),
        total = n())
```

```
## Source: local data frame [19 x 5]
##
##
                order avg_sleep min_sleep max_sleep total
## 1
         Afrosoricida 15.600000
                                      15.6
                                                 15.6
## 2
         Artiodactyla 4.516667
                                       1.9
                                                 9.1
                                                          6
## 3
            Carnivora 10.116667
                                       3.5
                                                 15.8
                                                         12
## 4
              Cetacea 4.500000
                                       2.7
                                                 5.6
                                                          3
## 5
           Chiroptera 19.800000
                                                 19.9
                                                          2
                                      19.7
## 6
            Cingulata 17.750000
                                      17.4
                                                 18.1
                                                          2
## 7
      Didelphimorphia 18.700000
                                      18.0
                                                 19.4
                                                          2
## 8
        Diprotodontia 12.400000
                                      11.1
                                                 13.7
                                                          2
## 9
       Erinaceomorpha 10.200000
                                                 10.3
                                                          2
                                      10.1
## 10
           Hyracoidea 5.666667
                                       5.3
                                                  6.3
                                                          3
## 11
           Lagomorpha 8.400000
                                       8.4
                                                  8.4
                                                          1
## 12
          Monotremata 8.600000
                                       8.6
                                                  8.6
                                                          1
       Perissodactyla 3.466667
## 13
                                       2.9
                                                  4.4
                                                          3
## 14
               Pilosa 14.400000
                                      14.4
                                                 14.4
                                                          1
## 15
                                                 17.0
                                                         12
             Primates 10.500000
                                       8.0
## 16
                                                          2
          Proboscidea 3.600000
                                       3.3
                                                 3.9
## 17
             Rodentia 12.468182
                                                 16.6
                                                         22
                                       7.0
## 18
           Scandentia 8.900000
                                       8.9
                                                 8.9
                                                          1
## 19
         Soricomorpha 11.100000
                                       8.4
                                                 14.9
                                                          5
```