Programming in R Workshop

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Load packages

Load purrr, tidyverse and dplyr packages.

```
library(purrr)
library(tidyverse)
library(conflicted)
library(dplyr)
```

Load the Eukaryotes dataset - only have to run this once to get the data

```
eukaryotes <- read_tsv(
   file = "ftp://ftp.ncbi.nlm.nih.gov/genomes/GENOME_REPORTS/eukaryotes.txt",
   na = c("", "na", "-")
)

# Reformat dataset headers
names_new <- names(eukaryotes) |>
   str_replace_all("[#%()]", "") |>
   str_replace_all("[/]", "_") |>
   str_to_lower()
```

```
eukaryotes <- eukaryotes |>
   set_names(names_new)

# Save tibble
write_tsv(eukaryotes, "eukaryotes.tsv")
```

Load the saved dataset

```
eukaryotes <- read_tsv(
   "https://raw.githubusercontent.com/swuyts/purrr_tutorial/master/data/eukaryotes.tsv"
)

Rows: 11508 Columns: 19
-- Column specification -------
Delimiter: "\t"
chr (10): organism_name, bioproject_accession, group, subgroup, assembly_ac...
dbl (7): taxid, bioproject_id, size_mb, gc, scaffolds, genes, proteins
date (2): release_date, modify_date

i Use `spec()` to retrieve the full column specification for this data.
i Specify the column types or set `show_col_types = FALSE` to quiet this message.</pre>
```

How many different organisms are there in our dataset?

```
eukaryotes |>
  pull(organism_name) |>
  n_distinct()
```

[1] 6111

Subset dataframe by selecting variables for the analysis:

```
eukaryotes_subset <- eukaryotes |>
select(organism_name, group, subgroup)
```

Using n_distinct to each variable of eukaryotes_subset

```
map(eukaryotes_subset, n_distinct)
```

```
$organism_name
[1] 6111
$group
[1] 5
$subgroup
[1] 19
eukaryotes_subset |>
 map(n_distinct)
$organism_name
[1] 6111
$group
[1] 5
$subgroup
[1] 19
Nested Tibbles
Split the eukaryotes dataset according to groups defined in the group variable:
eukaryotes |> pull(group) |> unique()
[1] "Other"
               "Protists" "Plants"
                                       "Fungi"
                                                  "Animals"
eukaryotes_nested <- eukaryotes |>
  group_by(group) |>
 nest()
eukaryotes_nested
# A tibble: 5 x 2
```

Groups: group [5]

data

t>

group

<chr>

Split the "eukaryotes_nested" into 5 smaller dataframes.

eukaryotes_nested |> pull(data)

[[1]]

```
# A tibble: 51 x 18
  organism_name
                       taxid bioproject_accession bioproject_id subgroup size_mb
   <chr>
                       <dbl> <chr>
                                                           <dbl> <chr>
                                                                            <dbl>
1 Pyropia yezoensis
                        2788 PRJNA589917
                                                          589917 Other
                                                                            108.
2 Thalassiosira pse~ 296543 PRJNA191
                                                             191 Other
                                                                             32.4
3 Guillardia theta ~ 905079 PRJNA53577
                                                           53577 Other
                                                                             87.1
4 Cyanidioschyzon m~ 280699 PRJNA10792
                                                           10792 Other
                                                                             16.5
5 Galdieria sulphur~ 130081 PRJNA13023
                                                           13023 Other
                                                                             13.7
6 Phaeodactylum tri~ 556484 PRJNA13152
                                                           13152 Other
                                                                             27.5
7 Bigelowiella nata~ 753081 PRJNA47111
                                                          47111 Other
                                                                             91.4
8 Ectocarpus silicu~
                        2880 PRJEA42625
                                                          42625 Other
                                                                            196.
                                                           36595 Other
9 Thalassiosira oce~ 159749 PRJNA36595
                                                                             92.2
10 Fragilariopsis cy~ 635003 PRJNA32761
                                                          32761 Other
                                                                             80.5
# i 41 more rows
# i 12 more variables: gc <dbl>, assembly_accession <chr>, replicons <chr>,
    wgs <chr>, scaffolds <dbl>, genes <dbl>, proteins <dbl>,
    release_date <date>, modify_date <date>, status <chr>, center <chr>,
    biosample_accession <chr>
```

[[2]]

A tibble: 888 x 18

	organism_name	taxid	${\tt bioproject_accession}$	bioproject_id	subgroup	size_mb
	<chr></chr>	<dbl></dbl>	<chr></chr>	<dbl></dbl>	<chr></chr>	<dbl></dbl>
1	Emiliania huxleyi~	280463	PRJNA77753	77753	Other P~	168.
2	Leishmania major ~	347515	PRJNA10724	10724	Kinetop~	32.9
3	Trypanosoma bruce~	679716	PRJEA40697	40697	Kinetop~	22.1
4	Trypanosoma cruzi	5693	PRJNA11755	11755	Kinetop~	89.9
5	Entamoeba histoly~	294381	PRJNA142	142	Other P~ $$	20.8
6	Giardia intestina~	5741	PRJNA561185	561185	Other P~ $$	11.5
7	Eimeria tenella	5802	PRJEB4863	224694	Apicomp~	51.9
8	Cryptosporidium p~	353152	PRJNA144	144	Apicomp~	9.10

```
9 Toxoplasma gondii~ 508771 PRJNA28893
                                                          28893 Apicomp~
                                                                           65.7
10 Plasmodium berghei
                       5821 PRJEB11993
                                                         305111 Apicomp~
                                                                           18.8
# i 878 more rows
# i 12 more variables: gc <dbl>, assembly_accession <chr>, replicons <chr>,
   wgs <chr>, scaffolds <dbl>, genes <dbl>, proteins <dbl>,
   release_date <date>, modify_date <date>, status <chr>, center <chr>,
   biosample accession <chr>
[[3]]
# A tibble: 1,304 x 18
  organism_name
                      taxid bioproject_accession bioproject_id subgroup size_mb
  <chr>
                       <dbl> <chr>
                                                          <dbl> <chr>
                                                                           <dbl>
1 Arabidopsis thali~ 3702 PRJNA10719
                                                          10719 Land Pl~
                                                                            120.
2 Glycine max
                       3847 PRJNA19861
                                                          19861 Land Pl~
                                                                            979.
3 Medicago truncatu~ 3880 PRJNA10791
                                                          10791 Land Pl~
                                                                            413.
4 Solanum lycopersi~ 4081 PRJNA119
                                                            119 Land Pl~
                                                                            828.
5 Hordeum vulgare s~ 112509 PRJEB34217
                                                        576847 Land Pl~
                                                                           4341.
6 Oryza sativa Japo~ 39947 PRJNA12269
                                                         12269 Land Pl~
                                                                           374.
7 Triticum aestivum 4565 PRJNA392179
                                                        392179 Land Pl~ 15419.
8 Zea mays
                      4577 PRJNA10769
                                                         10769 Land Pl~
                                                                           2135.
9 Coffea arabica
                                                        506972 Land Pl~
                      13443 PRJNA506972
                                                                         1094.
10 Lotus japonicus
                                                          28941 Land Pl~
                      34305 PRJDA28941
                                                                            394.
# i 1,294 more rows
# i 12 more variables: gc <dbl>, assembly_accession <chr>, replicons <chr>,
   wgs <chr>, scaffolds <dbl>, genes <dbl>, proteins <dbl>,
   release date <date>, modify date <date>, status <chr>, center <chr>,
   biosample_accession <chr>
[[4]]
# A tibble: 6,064 x 18
                      taxid bioproject_accession bioproject_id subgroup size_mb
  organism_name
                       <dbl> <chr>
                                                          <dbl> <chr>
                                                                           <dbl>
1 Pneumocystis cari~ 1.41e6 PRJNA223511
                                                         223511 Ascomyc~
                                                                            7.66
2 Schizosaccharomyc~ 4.90e3 PRJNA13836
                                                         13836 Ascomyc~
                                                                           12.6
3 Saccharomyces cer~ 5.59e5 PRJNA43747
                                                          43747 Ascomyc~
                                                                           12.2
4 Aspergillus nidul~ 2.27e5 PRJNA130
                                                            130 Ascomyc~
                                                                           30.3
5 Aspergillus fumig~ 3.31e5 PRJNA131
                                                            131 Ascomyc~
                                                                           29.4
6 Neurospora crassa~ 3.67e5 PRJNA13841
                                                         13841 Ascomyc~
                                                                          41.1
7 Phanerochaete chr~ 5.31e3 PRJNA343563
                                                        343563 Basidio~
                                                                         39.2
8 Candida albicans ~ 2.38e5 PRJNA10701
                                                         10701 Ascomyc~
                                                                          14.3
9 Encephalitozoon c~ 2.85e5 PRJNA13833
                                                         13833 Other F~
                                                                           2.50
10 Aspergillus terre~ 3.42e5 PRJNA15631
                                                         15631 Ascomyc~
                                                                           29.4
```

i 6,054 more rows

```
# i 12 more variables: gc <dbl>, assembly_accession <chr>, replicons <chr>,
   wgs <chr>, scaffolds <dbl>, genes <dbl>, proteins <dbl>,
   release_date <date>, modify_date <date>, status <chr>, center <chr>,
   biosample_accession <chr>
# A tibble: 3,201 x 18
  organism_name
                      taxid bioproject_accession bioproject_id subgroup size_mb
   <chr>
                      <dbl> <chr>
                                                         <dbl> <chr>
 1 Caenorhabditis br~ 6238 PRJNA10731
                                                         10731 Roundwo~
                                                                          108.
 2 Caenorhabditis el~ 6239 PRJNA13758
                                                                          100.
                                                         13758 Roundwo~
                                                        10729 Roundwo~
3 Brugia malayi
                     6279 PRJNA10729
                                                                         93.7
                      7159 PRJNA392114
4 Aedes aegypti
                                                        392114 Insects
                                                                         1279.
5 Aedes albopictus
                     7160 PRJNA552090
                                                        552090 Insects
                                                                         2538.
6 Anopheles gambiae~ 180454 PRJNA1438
                                                          1438 Insects
                                                                       265.
7 Drosophila melano~ 7227 PRJNA13669
                                                         13669 Insects
                                                                         144.
8 Apis mellifera
                       7460 PRJNA477511
                                                        477511 Insects
                                                                          225.
9 Ciona intestinalis 7719 PRJDA65419
                                                         65419 Other A~ 115.
10 Danio rerio
                       7955 PRJNA11776
                                                         11776 Fishes
                                                                         1679.
# i 3,191 more rows
# i 12 more variables: gc <dbl>, assembly_accession <chr>, replicons <chr>,
   wgs <chr>, scaffolds <dbl>, genes <dbl>, proteins <dbl>,
   release_date <date>, modify_date <date>, status <chr>, center <chr>,
   biosample accession <chr>>
```

Combine nested tibbles and map

Count number of rows for each sub data frames

```
map(eukaryotes_nested$data, nrow)
```

[[1]] [1] 51 [[2]] [1] 888 [[3]] [1] 1304

[[4]]

```
[1] 6064
[[5]]
[1] 3201
```

Create a new column using mutate()

```
eukaryotes_nested |>
mutate(n_row = map_int(data, nrow))
```

```
# A tibble: 5 x 3
# Groups:
           group [5]
 group
          data
                                n_row
 <chr>
          t>
                                <int>
1 Other
          <tibble [51 x 18]>
                                   51
2 Protists <tibble [888 x 18]>
                                  888
3 Plants <tibble [1,304 x 18]> 1304
          <tibble [6,064 x 18]>
4 Fungi
                                 6064
5 Animals <tibble [3,201 x 18]>
                                 3201
```

How many different organisms are there per group?

There are two different ways:

```
# Define a custom function
n_distinct_organisms <- function(data) {
    data |>
        pull(organism_name) |>
        n_distinct()
}

# Define a custom function as a formula
# .x is the notation for the object that is given as an input to this function.
n_distinct_organisms2 <- ~ .x |>
    pull(organism_name) |>
        n_distinct()
```

Apply the function to our nested data:

```
eukaryotes_nested |>
  mutate(
    n_organisms = map_dbl(data, n_distinct_organisms),
    n_organisms2 = map_dbl(data, n_distinct_organisms2)
)
```

```
# A tibble: 5 x 4
# Groups:
           group [5]
                                 n_organisms n_organisms2
 group
           data
  <chr>
                                                     <dbl>
           t>
                                       <dbl>
1 Other
           <tibble [51 x 18]>
                                          35
                                                       35
2 Protists <tibble [888 x 18]>
                                         490
                                                      490
3 Plants
          <tibble [1,304 x 18]>
                                                      673
                                         673
           <tibble [6,064 x 18]>
4 Fungi
                                        2926
                                                     2926
5 Animals <tibble [3,201 x 18]>
                                        1987
                                                     1987
```

We can define the functions on the fly:

```
eukaryotes_nested |>
  mutate(
    n_organisms = map_dbl(data, ~ .x |> pull(organism_name) |> n_distinct()),
    n_centers = map_dbl(data, ~ .x |> pull(center) |> n_distinct()),
    n_subgroups = map_dbl(data, ~ .x |> pull(subgroup) |> n_distinct())
)
```

```
# A tibble: 5 x 5
# Groups:
            group [5]
 group
           data
                                 n_organisms n_centers n_subgroups
  <chr>
           t>
                                       <dbl>
                                                  <dbl>
                                                              <dbl>
1 Other
           <tibble [51 x 18]>
                                          35
                                                     34
                                                                  1
2 Protists <tibble [888 x 18]>
                                                                  3
                                         490
                                                    265
          <tibble [1,304 x 18]>
                                                                  3
3 Plants
                                                    492
                                         673
           <tibble [6,064 x 18]>
                                                                  3
4 Fungi
                                        2926
                                                    950
5 Animals <tibble [3,201 x 18]>
                                        1987
                                                    769
                                                                  9
```

pmap and walk2 functions

We will explain by the following example for pmap:

```
df <- data.frame(</pre>
  x = c(
    "ATTTTACTGGGAGGGAA",
    "TATTTTTTAAAGGGCCC",
    "GCGCGCCCCAAATTATAGGC",
    "TGCCACATTTTATCCGCGCA"
  ),
  pattern = c("A", "T", "G", "C"),
  replacement = c("a", "t", "g", "c"),
  stringsAsFactors = FALSE
pmap(df, gsub)
[[1]]
[1] "aTTTTaCTGGGaGGGaa"
[[2]]
[1] "tAttttttAAAGGCCC"
[[3]]
[1] "gCgCgCCCCAAATTATAggC"
[[4]]
[1] "TGccAcATTTTATccGcGcA"
Example for walk2:
df1 <- data.frame(</pre>
  index = c(1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10),
  value = c(
    40,
    20,
    10,
    5,
    50,
    80,
    30,
    20,
    80,
```

90,

```
33,
    21,
    56,
    66,
    43,
    89,
    66,
    80,
    30,
    10
  ),
  category = c(
    "A",
    "B",
    "B",
    "B",
    "B",
    "B",
    "B",
    "B",
    "B",
    "B",
    "B"
)
df1 %>%
  split(.$category) %>%
  .[order(names(.))] %>%
  walk2(
    paste('Plot', names(.)),
    ~ plot(value ~ index, data = .x, type = "l", main = .y)
```

