

## Managing Data

### Extended Materials

You can find the original, extended version of these materials from chapters [8](#), [13](#), and [14](#).

This week we will be diving deeper into operations needed for data analysis. We will look at how to manipulate column names, group data by its variables, and append datasets together.

### Data Preparation

We will be continuing with the same simulated Ebola outbreak dataset.

```
pacman::p_load(
  rio,          # importing data
  here,         # relative file pathways
  janitor,      # data cleaning and tables
  lubridate,    # working with dates
  matchmaker,   # dictionary-based cleaning
  epikit,       # age_categories() function
  tidyverse     # data management and visualization
)
linelist <- rio::import("data/case_linelists/linelist_cleaned.rds")

linelist <- import("linelist_cleaned.rds")
```

### Column names

In R, column *names* are the “header” or “top” value of a column. They are used to refer to columns in the code, and serve as a default label in figures.

Other statistical software such as SAS and STATA use “*labels*” that co-exist as longer printed versions of the shorter column names. While R does offer the possibility of adding column labels to the data, this is not emphasized in most practice. To make column names “printer-friendly” for figures, one typically adjusts their display within the plotting commands that create the outputs.

As R column names are used very often, so they must have “clean” syntax. We suggest the following:

- Short names

- No spaces (replace with underscores `_` )
- No unusual characters (`&`, `#`, `<`, `>`, `...`)
- Similar style nomenclature (e.g. all date columns named like **date\_\_onset**, **date\_\_report**, **date\_\_death...**)

Re-naming columns manually is often necessary, even after the standardization step above. Below, re-naming is performed using the **rename()** function from the **dplyr** package, as part of a pipe chain. **rename()** uses the style **NEW = OLD** - the new column name is given before the old column name.

Below, a re-naming command is added to the cleaning pipeline. Spaces have been added strategically to align code for easier reading.

Now you can see that the columns names have been changed:

[1] "case_id"	"generation"	"date_infection"
[4] "date_onset"	"date_hospitalisation"	"date_outcome"
[7] "outcome"	"gender"	"age"
[10] "age_unit"	"age_years"	"age_cat"
[13] "age_cat5"	"hospital"	"lon"
[16] "lat"	"infector"	"source"
[19] "wt_kg"	"ht_cm"	"ct_blood"
[22] "fever"	"chills"	"cough"
[25] "aches"	"vomit"	"temp"
[28] "time_admission"	"bmi"	"days_onset_hosp"

#### \* Rename by column position

You can also rename by column position, instead of column name, for example:

```
rename(newNameForFirstColumn = 1,
       newNameForSecondColumn = 2)
```

#### \* Rename via **select()** and **summarise()**

As a shortcut, you can also rename columns within the **dplyr** **select()** and **summarise()** functions. **select()** is used to keep only certain columns and **summarise()** is used when grouping data. These functions also uses the format **new\_name = old\_name**. Here is an example:

```
linelist_raw %>%
  select(# NEW name          # OLD name
         date_infection      = `infection date`,    # rename and KEEP ONLY these columns
         date_hospitalisation = `hosp date`)
```

## Other considerations with column names

### \* Empty Excel column names

R cannot have dataset columns that do not have column names (headers). So, if you import an Excel dataset with data but no column headers, R will fill-in the headers with names like “...1” or “...2”. The number represents the column number (e.g. if the 4th column in the dataset has no header, then R will name it “...4”).

You can clean these names manually by referencing their position number (see example above), or their assigned name (`linelist_raw$...1`).

### \* Merged Excel column names and cells

Merged cells in an Excel file are a common occurrence when receiving data. Merged cells can be nice for human reading of data, but are not “tidy data” and cause many problems for machine reading of data. R cannot accommodate merged cells.

One solution to deal with merged cells is to import the data with the function `readWorkbook()` from the package **openxlsx**. Set the argument `fillMergedCells = TRUE`. This gives the value in a merged cell to all cells within the merge range.

```
linelist_raw <- openxlsx::readWorkbook("linelist_raw.xlsx", fillMergedCells = TRUE)
```

## Revisiting select

Two weeks ago we learned to use `select()` to select the columns we wanted to keep.

```
# linelist dataset is piped through select() command, and names() prints just the column names
linelist %>%
  select(case_id, date_onset, date_hospitalisation, fever) %>%
  names() # display the column names
```

```
[1] "case_id"          "date_onset"       "date_hospitalisation"
[4] "fever"
```

Let's look at some more complicated scenarios when we need to think a bit deeper on how we're selecting or choosing columns in our data.

### “tidyselect” helper functions

These helper functions exist to make it easy to specify columns to keep, discard, or transform. They are from the package **tidyselect**, which is included in **tidyverse** and underlies how columns are selected in **dplyr** functions.

For example, if you want to re-order the columns, **everything()** is a useful function to signify “all other columns not yet mentioned”. The command below moves columns **date\_onset** and **date\_hospitalisation** to the beginning (left) of the dataset, but keeps all the other columns afterward. Note that **everything()** is written with empty parentheses:

```
# move date_onset and date_hospitalisation to beginning
linelist %>%
  select(date_onset, date_hospitalisation, everything()) %>%
  names()
```

[1] "date_onset"	"date_hospitalisation"	"case_id"
[4] "generation"	"date_infection"	"date_outcome"
[7] "outcome"	"gender"	"age"
[10] "age_unit"	"age_years"	"age_cat"
[13] "age_cat5"	"hospital"	"lon"
[16] "lat"	"infectior"	"source"
[19] "wt_kg"	"ht_cm"	"ct_blood"
[22] "fever"	"chills"	"cough"
[25] "aches"	"vomit"	"temp"
[28] "time_admission"	"bmi"	"days_onset_hosp"

Here are other “tidyselect” helper functions that also work *within* **dplyr** functions like **select()**, **across()**, and **summarise()**:

- **everything()** - all other columns not mentioned
- **last\_col()** - the last column
- **where()** - applies a function to all columns and selects those which are TRUE
- **contains()** - columns containing a character string
  - example: `select(contains("time"))`

- `starts_with()` - matches to a specified prefix
  - example: `select(starts_with("date_"))`
- `ends_with()` - matches to a specified suffix
  - example: `select(ends_with("_post"))`
- `matches()` - to apply a regular expression (regex)
  - example: `select(matches("[pt]al"))`
- `num_range()` - a numerical range like x01, x02, x03
- `any_of()` - matches IF column exists but returns no error if it is not found
  - example: `select(any_of(date_onset, date_death, cardiac_arrest))`

In addition, use normal operators such as `c()` to list several columns, `:` for consecutive columns, `!` for opposite, `&` for AND, and `|` for OR.

Use `where()` to specify logical criteria for columns. If providing a function inside `where()`, do not include the function's empty parentheses. The command below selects columns that are class Numeric.

```
# select columns that are class Numeric
linelist %>%
  select(where(is.numeric)) %>%
  names()
```

```
[1] "generation"      "age"              "age_years"        "lon"
[5] "lat"             "wt_kg"            "ht_cm"            "ct_blood"
[9] "temp"            "bmi"              "days_onset_hosp"
```

Use `contains()` to select only columns in which the column name contains a specified character string. `ends_with()` and `starts_with()` provide more nuance.

```
# select columns containing certain characters
linelist %>%
  select(contains("date")) %>%
  names()
```

```
[1] "date_infection"      "date_onset"          "date_hospitalisation"
[4] "date_outcome"
```

The function `matches()` works similarly to `contains()` but can be provided a regular expression, such as multiple strings separated by OR bars within the parentheses:

```
# searched for multiple character matches
linelist %>%
  select(matches("onset|hosp|fev")) %>% # note the OR symbol "|"
  names()
```

```
[1] "date_onset"          "date_hospitalisation" "hospital"
[4] "fever"               "days_onset_hosp"
```

## Deduplication

In a later week we will learn more about how to de-duplicate data. Only a very simple row de-duplication example is presented here.

The package **dplyr** offers the `distinct()` function. This function examines every row and reduce the data frame to only the unique rows. That is, it removes rows that are 100% duplicates.

When evaluating duplicate rows, it takes into account a range of columns - by default it considers all columns. As shown in the de-duplication page, you can adjust this column range so that the uniqueness of rows is only evaluated in regards to certain columns.

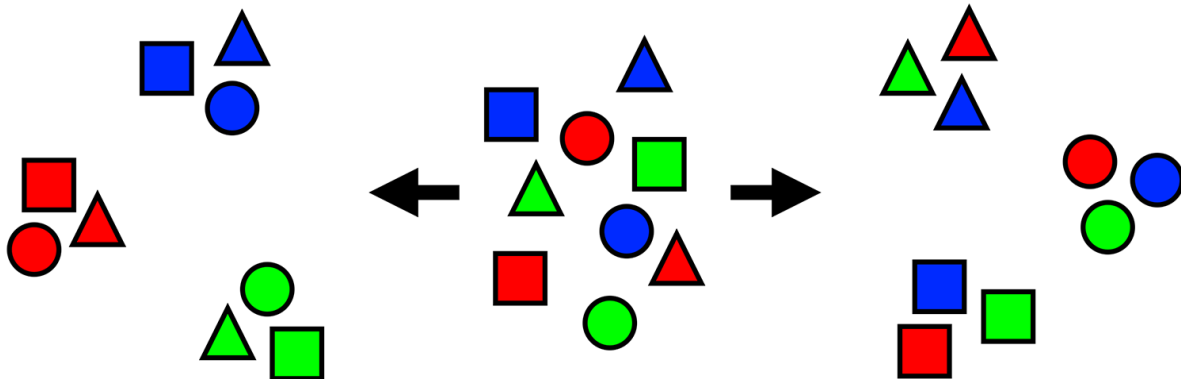
In this simple example, we just add the empty command `distinct()` to the pipe chain. This ensures there are no rows that are 100% duplicates of other rows (evaluated across all columns).

We begin with `nrow(linelist)` rows in `linelist`.

```
linelist <- linelist %>%
  distinct()
```

After de-duplication there are `nrow(linelist)` rows. Any removed rows would have been 100% duplicates of other rows.

## Grouping data



Grouping data is a core component of data management and analysis. Grouped data statistically summarised by group, and can be plotted by group. Functions from the **dplyr** package (part of the **tidyverse**) make grouping and subsequent operations quite easy.

We will be examining:

- Grouping data with the `group_by()` function
- Un-grouping data
- `summarise()` grouped data with statistics
- The difference between `count()` and `tally()`
- `arrange()` applied to grouped data
- `filter()` applied to grouped data
- `mutate()` applied to grouped data
- `select()` applied to grouped data
- The **base R** `aggregate()` command as an alternative

Let's load a clean version of the simulated Ebola epidemic dataset.

```
linelist <- import("linelist_cleaned.rds")
```

## Grouping

The function `group_by()` from **dplyr** groups the rows by the unique values in the column specified to it. If multiple columns are specified, rows are grouped by the unique combinations of values across the columns. Each unique value (or combination of values) constitutes a group. Subsequent changes to the dataset or calculations can then be performed within the context of each group.

For example, the command below takes the `linelist` and groups the rows by unique values in the column `outcome`, saving the output as a new data frame `ll_by_outcome`. The grouping column(s) are placed inside the parentheses of the function `group_by()`.

```
ll_by_outcome <- linelist %>%  
  group_by(outcome)
```

**Note that there is no perceptible change to the dataset** after running `group_by()`, *until* another **dplyr** verb such as `mutate()`, `summarise()`, or `arrange()` is applied on the “grouped” data frame.

You can however “see” the groupings by printing the data frame. When you print a grouped data frame, you will see it has been transformed into a **tibble class object** which, when printed, displays which groupings have been applied and how many groups there are - written just above the header row.

```
# print to see which groups are active  
ll_by_outcome  
  
# A tibble: 5,888 x 30  
# Groups:   outcome [3]  
  case_id generation date_infection date_onset date_hospitalisation  
  <chr>      <dbl> <date>          <date>      <date>  
1 5fe599      4 2014-05-08      2014-05-13 2014-05-15  
2 8689b7      4 NA              2014-05-13 2014-05-14  
3 11f8ea      2 NA              2014-05-16 2014-05-18  
4 b8812a      3 2014-05-04      2014-05-18 2014-05-20  
5 893f25      3 2014-05-18      2014-05-21 2014-05-22  
6 be99c8      3 2014-05-03      2014-05-22 2014-05-23  
7 07e3e8      4 2014-05-22      2014-05-27 2014-05-29  
8 369449      4 2014-05-28      2014-06-02 2014-06-03  
9 f393b4      4 NA              2014-06-05 2014-06-06  
10 1389ca     4 NA              2014-06-05 2014-06-07  
# i 5,878 more rows  
# i 25 more variables: date_outcome <date>, outcome <chr>, gender <chr>,
```



```
# age <dbl>, age_unit <chr>, age_years <dbl>, age_cat <fct>, age_cat5 <fct>,
# hospital <chr>, lon <dbl>, lat <dbl>, infector <chr>, source <chr>,
# wt_kg <dbl>, ht_cm <dbl>, ct_blood <dbl>, fever <chr>, chills <chr>,
# cough <chr>, aches <chr>, vomit <chr>, temp <dbl>, time_admission <chr>,
# bmi <dbl>, days_onset_hosp <dbl>
```

## Unique groups

The groups created reflect each unique combination of values across the grouping columns.

To see the groups *and the number of rows in each group*, pass the grouped data to `tally()`. To see just the unique groups without counts you can pass to `group_keys()`.

See below that there are **three** unique values in the grouping column `outcome`: “Death”, “Recover”, and NA. See that there were `nrow(linelist %>% filter(outcome == "Death"))` deaths, `nrow(linelist %>% filter(outcome == "Recover"))` recoveries, and `nrow(linelist %>% filter(is.na(outcome)))` with no outcome recorded.

```
linelist %>%
  group_by(outcome) %>%
  tally()
```

```
# A tibble: 3 x 2
  outcome      n
  <chr>   <int>
1 Death   2582
2 Recover 1983
3 <NA>    1323
```

You can group by more than one column. Below, the data frame is grouped by `outcome` and `gender`, and then tallied. Note how each unique combination of `outcome` and `gender` is registered as its own group - including missing values for either column.

```
linelist %>%
  group_by(outcome, gender) %>%
  tally()
```

```
# A tibble: 9 x 3
# Groups:   outcome [3]
  outcome gender      n
```

	<chr>	<chr>	<int>
1	Death	f	1227
2	Death	m	1228
3	Death	<NA>	127
4	Recover	f	953
5	Recover	m	950
6	Recover	<NA>	80
7	<NA>	f	627
8	<NA>	m	625
9	<NA>	<NA>	71

## New columns

You can also create a new grouping column *within* the `group_by()` statement. This is equivalent to calling `mutate()` before the `group_by()`. For a quick tabulation this style can be handy, but for more clarity in your code consider creating this column in its own `mutate()` step and then piping to `group_by()`.

```
# group dat based on a binary column created *within* the group_by() command
linelist %>%
  group_by(
    age_class = ifelse(age >= 18, "adult", "child")) %>%
  tally(sort = T)
```

```
# A tibble: 3 x 2
  age_class      n
  <chr>        <int>
1 child      3618
2 adult      2184
3 <NA>         86
```

## Add/drop grouping columns

By default, if you run `group_by()` on data that are already grouped, the old groups will be removed and the new one(s) will apply. If you want to add new groups to the existing ones, include the argument `.add = TRUE`.

```
# Grouped by outcome
by_outcome <- linelist %>%
  group_by(outcome)
```

```
# Add grouping by gender in addition
by_outcome_gender <- by_outcome %>%
  group_by(gender, .add = TRUE)
```

**\*\* Keep all groups\*\***

If you group on a column of class factor there may be levels of the factor that are not currently present in the data. If you group on this column, by default those non-present levels are dropped and not included as groups. To change this so that all levels appear as groups (even if not present in the data), set `.drop = FALSE` in your `group_by()` command.

## Un-group

Data that have been grouped will remain grouped until specifically ungrouped via `ungroup()`. If you forget to ungroup, it can lead to incorrect calculations! Below is an example of removing all groupings:

```
linelist %>%
  group_by(outcome, gender) %>%
  tally() %>%
  ungroup()
```

You can also remove grouping for only specific columns, by placing the column name inside `ungroup()`.

```
linelist %>%
  group_by(outcome, gender) %>%
  tally() %>%
  ungroup(gender) # remove the grouping by gender, leave grouping by outcome
```

### Tip

The verb `count()` automatically ungroups the data after counting.

## Summarise

The **dplyr** function `summarise()` (or `summarize()`) takes a data frame and converts it into a *new* summary data frame, with columns containing summary statistics that you define. On an ungrouped data frame, the summary statistics will be calculated from all rows. Applying `summarise()` to grouped data produces those summary statistics *for each group*.

The syntax of `summarise()` is such that you provide the name(s) of the **new** summary column(s), an equals sign, and then a statistical function to apply to the data, as shown below. For example, `min()`, `max()`, `median()`, or `sd()`. Within the statistical function, list the column to be operated on and any relevant argument (e.g. `na.rm = TRUE`). You can use `sum()` to count the number of rows that meet a logical criteria (with double equals `==`).

Below is an example of `summarise()` applied *without grouped data*. The statistics returned are produced from the entire dataset.

```
# summary statistics on ungrouped linelist
linelist %>%
  summarise(
    n_cases = n(),
    mean_age = mean(age_years, na.rm=T),
    max_age = max(age_years, na.rm=T),
    min_age = min(age_years, na.rm=T),
    n_males = sum(gender == "m", na.rm=T))
```

	n_cases	mean_age	max_age	min_age	n_males
1	5888	16.01831	84	0	2803

In contrast, below is the same `summarise()` statement applied to grouped data. The statistics are calculated for each outcome group. Note how grouping columns will carry over into the new data frame.

```
# summary statistics on grouped linelist
linelist %>%
  group_by(outcome) %>%
  summarise(
    n_cases = n(),
    mean_age = mean(age_years, na.rm=T),
    max_age = max(age_years, na.rm=T),
    min_age = min(age_years, na.rm=T),
    n_males = sum(gender == "m", na.rm=T))
```

```
# A tibble: 3 x 6
  outcome n_cases mean_age max_age min_age n_males
  <chr>    <int>    <dbl>   <dbl>   <dbl>   <int>
1 Death    2582     15.9     76      0    1228
2 Recover  1983     16.1     84      0     950
3 <NA>    1323     16.2     69      0     625
```

## Counts and tallies

`count()` and `tally()` provide similar functionality but are different. Read more about the distinction between `tally()` and `count()` [here](#)

### `tally()`

`tally()` is shorthand for `summarise(n = n())`, and *does not* group data. Thus, to achieve grouped tallies it must follow a `group_by()` command. You can add `sort = TRUE` to see the largest groups first.

```
linelist %>%  
  tally()
```

```
      n  
1 5888
```

```
linelist %>%  
  group_by(outcome) %>%  
  tally(sort = TRUE)
```

```
# A tibble: 3 x 2  
  outcome      n  
  <chr>   <int>  
1 Death    2582  
2 Recover  1983  
3 <NA>     1323
```

### `count()`

In contrast, `count()` does the following:

- 1) applies `group_by()` on the specified column(s)
- 2) applies `summarise()` and returns column `n` with the number of rows per group
- 3) applies `ungroup()`

```
linelist %>%
  count(outcome)
```

```
outcome      n
1   Death 2582
2 Recover 1983
3    <NA> 1323
```

Just like with `group_by()` you can create a new column within the `count()` command:

```
linelist %>%
  count(age_class = ifelse(age >= 18, "adult", "child"), sort = T)
```

```
age_class      n
1    child 3618
2    adult 2184
3    <NA>   86
```

`count()` can be called multiple times, with the functionality “rolling up”. For example, to summarise the number of hospitals present for each gender, run the following. Note, the name of the final column is changed from default “n” for clarity (with `name =` ).

```
linelist %>%
  # produce counts by unique outcome-gender groups
  count(gender, hospital) %>%
  # gather rows by gender (3) and count number of hospitals per gender (6)
  count(gender, name = "hospitals per gender" )
```

```
gender hospitals per gender
1      f                      6
2      m                      6
3    <NA>                    6
```

## Add counts

In contrast to `count()` and `summarise()`, you can use `add_count()` to *add* a new column `n` with the counts of rows per group *while retaining all the other data frame columns*.

This means that a group's count number, in the new column `n`, will be printed in each row of the group. For demonstration purposes, we add this column and then re-arrange the columns for easier viewing. See the section below on [filter on group size](#) for another example.

```
linelist %>%
  as_tibble() %>%                # convert to tibble for nicer printing
  add_count(hospital) %>%        # add column n with counts by hospital
  select(hospital, n, everything()) # re-arrange for demo purposes
```

# A tibble: 5,888 x 31

	hospital	n	case_id	generation	date_infection	date_onset
	<chr>	<int>	<chr>	<dbl>	<date>	<date>
1	Other	885	5fe599	4	2014-05-08	2014-05-13
2	Missing	1469	8689b7	4	NA	2014-05-13
3	St. Mark's Maternity Hosp~	422	11f8ea	2	NA	2014-05-16
4	Port Hospital	1762	b8812a	3	2014-05-04	2014-05-18
5	Military Hospital	896	893f25	3	2014-05-18	2014-05-21
6	Port Hospital	1762	be99c8	3	2014-05-03	2014-05-22
7	Missing	1469	07e3e8	4	2014-05-22	2014-05-27
8	Missing	1469	369449	4	2014-05-28	2014-06-02
9	Missing	1469	f393b4	4	NA	2014-06-05
10	Missing	1469	1389ca	4	NA	2014-06-05

# i 5,878 more rows

# i 25 more variables: date\_hospitalisation <date>, date\_outcome <date>,  
 # outcome <chr>, gender <chr>, age <dbl>, age\_unit <chr>, age\_years <dbl>,  
 # age\_cat <fct>, age\_cat5 <fct>, lon <dbl>, lat <dbl>, infector <chr>,  
 # source <chr>, wt\_kg <dbl>, ht\_cm <dbl>, ct\_blood <dbl>, fever <chr>,  
 # chills <chr>, cough <chr>, aches <chr>, vomit <chr>, temp <dbl>,  
 # time\_admission <chr>, bmi <dbl>, days\_onset\_hosp <dbl>

## Add totals

To easily add total *sum* rows or columns after using `tally()` or `count()` you can use the `tabyl` function from `janitor`. This package also offers functions like `adorn_totals()` and `adorn_percentages()` to add totals and convert to show percentages. Below is a brief example:

```
linelist %>%                                # case linelist
  tabyl(age_cat, gender) %>%                # cross-tabulate counts of two columns
  adorn_totals(where = "row") %>%          # add a total row
  adorn_percentages(denominator = "col") %>% # convert to proportions with column denominator
```

```

adorn_pct_formatting() %>%           # convert proportions to percents
adorn_ns(position = "front") %>%    # display as: "count (percent)"
adorn_title(                         # adjust titles
  row_name = "Age Category",
  col_name = "Gender")

```

Age Category	Gender					
	f		m		NA_	
0-4	640	(22.8%)	416	(14.8%)	39	(14.0%)
5-9	641	(22.8%)	412	(14.7%)	42	(15.1%)
10-14	518	(18.5%)	383	(13.7%)	40	(14.4%)
15-19	359	(12.8%)	364	(13.0%)	20	(7.2%)
20-29	468	(16.7%)	575	(20.5%)	30	(10.8%)
30-49	179	(6.4%)	557	(19.9%)	18	(6.5%)
50-69	2	(0.1%)	91	(3.2%)	2	(0.7%)
70+	0	(0.0%)	5	(0.2%)	1	(0.4%)
<NA>	0	(0.0%)	0	(0.0%)	86	(30.9%)
Total	2,807	(100.0%)	2,803	(100.0%)	278	(100.0%)

## Arranging grouped data

Using the **dplyr** verb `arrange()` to order the rows in a data frame behaves the same when the data are grouped, *unless* you set the argument `.by_group = TRUE`. In this case the rows are ordered first by the grouping columns and then by any other columns you specify to `arrange()`.

## Filter on grouped data

### \* `filter()`

When applied in conjunction with functions that evaluate the data frame (like `max()`, `min()`, `mean()`), these functions will now be applied to the groups. For example, if you want to filter and keep rows where patients are above the median age, this will now apply per group - filtering to keep rows above the *group's* median age.

### \* Slice rows per group

The **dplyr** function `slice()`, which [filters rows based on their position](#) in the data, can also be applied per group. Remember to account for sorting the data within each group to get the desired "slice".



For example, to retrieve only the latest 5 admissions from each hospital:

- 1) Group the linelist by column **hospital**
- 2) Arrange the records from latest to earliest **date\_hospitalisation** *within each hospital group*
- 3) Slice to retrieve the first 5 rows from each hospital

```
linelist %>%
  group_by(hospital) %>%
  arrange(hospital, date_hospitalisation) %>%
  slice_head(n = 5) %>%
  arrange(hospital) %>% # for display
  select(case_id, hospital, date_hospitalisation) # for display
```

```
# A tibble: 30 x 3
```

```
# Groups:   hospital [6]
```

	case_id	hospital	date_hospitalisation
	<chr>	<chr>	<date>
1	20b688	Central Hospital	2014-05-06
2	d58402	Central Hospital	2014-05-10
3	b8f2fd	Central Hospital	2014-05-13
4	acf422	Central Hospital	2014-05-28
5	275cc7	Central Hospital	2014-05-28
6	d1fafd	Military Hospital	2014-04-17
7	974bc1	Military Hospital	2014-05-13
8	6a9004	Military Hospital	2014-05-13
9	09e386	Military Hospital	2014-05-14
10	865581	Military Hospital	2014-05-15

```
# i 20 more rows
```

`slice_head()` - selects n rows from the top

`slice_tail()` - selects n rows from the end

`slice_sample()` - randomly selects n rows

`slice_min()` - selects n rows with highest values in `order_by` = column, use `with_ties = TRUE` to keep ties

`slice_max()` - selects n rows with lowest values in `order_by` = column, use `with_ties = TRUE` to keep ties

\* Filter on group size

The function `add_count()` adds a column `n` to the original data giving the number of rows in that row's group.

Shown below, `add_count()` is applied to the column `hospital`, so the values in the new column `n` reflect the number of rows in that row's hospital group. Note how values in column `n` are repeated. In the example below, the column name `n` could be changed using `name =` within `add_count()`. For demonstration purposes we re-arrange the columns with `select()`.

```
linelist %>%
  as_tibble() %>%
  add_count(hospital) %>%           # add "number of rows admitted to same hospital as this"
  select(hospital, n, everything())
```

```
# A tibble: 5,888 x 31
  hospital                n case_id generation date_infection date_onset
  <chr>                <int> <chr>         <dbl> <date>          <date>
1 Other                885 5fe599          4 2014-05-08      2014-05-13
2 Missing             1469 8689b7          4 NA              2014-05-13
3 St. Mark's Maternity Hosp~ 422 11f8ea          2 NA              2014-05-16
4 Port Hospital       1762 b8812a          3 2014-05-04      2014-05-18
5 Military Hospital    896 893f25          3 2014-05-18      2014-05-21
6 Port Hospital       1762 be99c8          3 2014-05-03      2014-05-22
7 Missing             1469 07e3e8          4 2014-05-22      2014-05-27
8 Missing             1469 369449          4 2014-05-28      2014-06-02
9 Missing             1469 f393b4          4 NA              2014-06-05
10 Missing            1469 1389ca          4 NA              2014-06-05
# i 5,878 more rows
# i 25 more variables: date_hospitalisation <date>, date_outcome <date>,
#   outcome <chr>, gender <chr>, age <dbl>, age_unit <chr>, age_years <dbl>,
#   age_cat <fct>, age_cat5 <fct>, lon <dbl>, lat <dbl>, infector <chr>,
#   source <chr>, wt_kg <dbl>, ht_cm <dbl>, ct_blood <dbl>, fever <chr>,
#   chills <chr>, cough <chr>, aches <chr>, vomit <chr>, temp <dbl>,
#   time_admission <chr>, bmi <dbl>, days_onset_hosp <dbl>
```

It then becomes easy to filter for case rows who were hospitalized at a “small” hospital, say, a hospital that admitted fewer than 500 patients:

```
linelist %>%
  add_count(hospital) %>%
  filter(n < 500)
```

## Mutate on grouped data

To retain all columns and rows (not summarise) and *add a new column containing group statistics*, use `mutate()` after `group_by()` instead of `summarise()`.

This is useful if you want group statistics in the original dataset *with all other columns present* - e.g. for calculations that compare one row to its group.

For example, this code below calculates the difference between a row's delay-to-admission and the median delay for their hospital. The steps are:

- 1) Group the data by hospital
- 2) Use the column `days_onset_hosp` (delay to hospitalisation) to create a new column containing the mean delay at the hospital of *that row*
- 3) Calculate the difference between the two columns

We `select()` only certain columns to display, for demonstration purposes.

```
linelist %>%
  # group data by hospital (no change to linelist yet)
  group_by(hospital) %>%

  # new columns
  mutate(
    # mean days to admission per hospital (rounded to 1 decimal)
    group_delay_admit = round(mean(days_onset_hosp, na.rm=T), 1),

    # difference between row's delay and mean delay at their hospital (rounded to 1 decimal)
    diff_to_group = round(days_onset_hosp - group_delay_admit, 1)) %>%

  # select certain rows only - for demonstration/viewing purposes
  select(case_id, hospital, days_onset_hosp, group_delay_admit, diff_to_group)
```

```
# A tibble: 5,888 x 5
```

```
# Groups:   hospital [6]
```

	case_id	hospital	days_onset_hosp	group_delay_admit	diff_to_group
	<chr>	<chr>	<dbl>	<dbl>	<dbl>
1	5fe599	Other	2	2	0
2	8689b7	Missing	1	2.1	-1.1
3	11f8ea	St. Mark's Maternity~	2	2.1	-0.1
4	b8812a	Port Hospital	2	2.1	-0.1

5	893f25	Military Hospital	1	2.1	-1.1
6	be99c8	Port Hospital	1	2.1	-1.1
7	07e3e8	Missing	2	2.1	-0.1
8	369449	Missing	1	2.1	-1.1
9	f393b4	Missing	1	2.1	-1.1
10	1389ca	Missing	2	2.1	-0.1

# i 5,878 more rows

## Select on grouped data

The verb `select()` works on grouped data, but the grouping columns are always included (even if not mentioned in `select()`). If you do not want these grouping columns, use `ungroup()` first.

## Appending Datasets

We often need to combine multiple sources of data. Later on we'll see more complex methods for combining data based on matching ID's or other values. However, to start we'll look at the case where we want to add additional observations to a dataset. You can also think of this as "appending" or "adding" rows.

## Bind rows

To bind rows of one data frame to the bottom of another data frame, use `bind_rows()` from **dplyr**. It is very inclusive, so any column present in either data frame will be included in the output. A few notes:

- Unlike the **base** R version `row.bind()`, **dplyr**'s `bind_rows()` does not require that the order of columns be the same in both data frames. As long as the column names are spelled identically, it will align them correctly.
- You can optionally specify the argument `.id =`. Provide a character column name. This will produce a new column that serves to identify which data frame each row originally came from.
- You can use `bind_rows()` on a **list** of similarly-structured data frames to combine them into one data frame.

One common example of row binding is to bind a "total" row onto a descriptive table made with **dplyr**'s `summarise()` function. Below we create a table of case counts and median CT values by hospital with a total row.

The function `summarise()` is used on data grouped by hospital to return a summary data frame by hospital. But the function `summarise()` does not automatically produce a “totals” row, so we create it by summarising the data *again*, but with the data not grouped by hospital. This produces a second data frame of just one row. We can then bind these data frames together to achieve the final table.

```
# Create core table
#####
hosp_summary <- linelist %>%
  group_by(hospital) %>%                                # Group data by hospital
  summarise(                                              # Create new summary columns of indicators
    cases = n(),                                         # Number of rows per hospital-outcome group
    ct_value_med = median(ct_blood, na.rm=T))           # median CT value per group
```

Here is the `hosp_summary` data frame:

Show 10 entries

Search:

hospital	cases	ct_value_med
Central Hospital	454	22
Military Hospital	896	21
Missing	1469	21
Other	885	22
Port Hospital	1762	22
St. Mark's Maternity Hospital (SMMH)	422	22

Showing 1 to 6 of 6 entries

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Create a data frame with the “total” statistics (*not grouped by hospital*). This will return just one row.

```
# create totals
#####
totals <- linelist %>%
  summarise(
    cases = n(), # Number of rows for whole dataset
    ct_value_med = median(ct_blood, na.rm=T)) # Median CT for whole dataset
```

And below is that **totals** data frame. Note how there are only two columns. These columns are also in **hosp\_summary**, but there is one column in **hosp\_summary** that is not in **totals** (**hospital**).

Show 10 entries

Search:

cases	ct_value_med
5888	22

Showing 1 to 1 of 1 entries

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```
cases ct_value_med
1  5888           22
```

Now we can bind the rows together with `bind_rows()`.

```
# Bind data frames together
combined <- bind_rows(hosp_summary, totals)
```

Now we can view the result. See how in the final row, an empty NA value fills in for the column `hospital` that was not in `hosp_summary`.

Show 10 entries

Search:

hospital	cases	ct_value_med
Central Hospital	454	22
Military Hospital	896	21
Missing	1469	21
Other	885	22
Port Hospital	1762	22
St. Mark's Maternity Hospital (SMMH)	422	22
	5888	22

Showing 1 to 7 of 7 entries

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```
# A tibble: 7 x 3
  hospital      cases ct_value_med
  <chr>      <int>      <dbl>
1 Central Hospital      454         22
2 Military Hospital      896         21
3 Missing      1469         21
4 Other          885         22
5 Port Hospital     1762         22
6 St. Mark's Maternity Hospital (SMMH)    422         22
7 <NA>          5888         22
```