Univariate and multivariable regression

This page demonstrates the use of **base** R regression functions such as glm() and the **gt-summary** package to look at associations between variables (e.g. odds ratios, risk ratios and hazard ratios). It also uses functions like tidy() from the **broom** package to clean-up regression outputs.

Preparation

Load packages

```
pacman::p_load(
 rio,
              # File import
             # File locator
 here,
 tidyverse, # data management + ggplot2 graphics,
             # manipulate text strings
 stringr,
             # loop over objects in a tidy way
 purrr,
 gtsummary,  # summary statistics and tests
             # tidy up results from regressions
 broom,
             # likelihood-ratio tests
 lmtest,
 parameters, # alternative to tidy up results from regressions
              # alternative to visualise forest plots
 see
 )
```

Import data

We import the dataset of cases from a simulated Ebola epidemic. If you want to follow along, click to download the "clean" linelist (as .rds file).

```
# import the linelist
linelist <- import("linelist_cleaned.rds")</pre>
```

Clean data

* Store explanatory variables

We store the names of the explanatory columns as a character vector. This will be referenced later.

```
## define variables of interest
explanatory_vars <- c("gender", "fever", "chills", "cough", "aches", "vomit")</pre>
```

* Convert to 1's and 0's

Below we convert the explanatory columns from "yes"/"no", "m"/"f", and "dead"/"alive" to 1 / 0, to cooperate with the expectations of logistic regression models. To do this efficiently, used across() from dplyr to transform multiple columns at one time. The function we apply to each column is case_when() (also dplyr) which applies logic to convert specified values to 1's and 0's.

Note: the "." below represents the column that is being processed by across() at that moment.

* Drop rows with missing values

To drop rows with missing values, can use the **tidyr** function **drop_na()**. However, we only want to do this for rows that are missing values in the columns of interest.

The first thing we must to is make sure our explanatory_vars vector includes the column age (age would have produced an error in the previous case_when() operation, which was only for dichotomous variables). Then we pipe the linelist to drop_na() to remove any rows with missing values in the outcome column or any of the explanatory_vars columns.

Before running the code, the number of rows in the linelist is nrow(linelist).

```
## add in age_category to the explanatory vars
explanatory_vars <- c(explanatory_vars, "age_cat")

## drop rows with missing information for variables of interest
linelist <- linelist %>%
```

```
drop_na(any_of(c("outcome", explanatory_vars)))
```

The number of rows remaining in linelist is nrow(linelist).

Univariate

Your use case will determine which R package you use. We present two options for doing univariate analysis:

- Use functions available in **base** R to quickly print results to the console. Use the **broom** package to tidy up the outputs.
- Use the **gtsummary** package to model and get publication-ready outputs

base R

* Linear regression

The base R function lm() perform linear regression, assessing the relationship between numeric response and explanatory variables that are assumed to have a linear relationship.

Provide the equation as a formula, with the response and explanatory column names separated by a tilde ~. Also, specify the dataset to data =. Define the model results as an R object, to use later.

```
lm_results <- lm(ht_cm ~ age, data = linelist)</pre>
```

You can then run summary() on the model results to see the coefficients (Estimates), P-value, residuals, and other measures.

```
summary(lm_results)
```

Call:

```
lm(formula = ht_cm ~ age, data = linelist)
```

Residuals:

```
Min 1Q Median 3Q Max -128.579 -15.854 1.177 15.887 175.483
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 69.9051
                         0.5979
                                  116.9
                                          <2e-16 ***
                                  117.2
              3.4354
                         0.0293
                                          <2e-16 ***
age
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
Residual standard error: 23.75 on 4165 degrees of freedom
                                Adjusted R-squared: 0.7674
Multiple R-squared: 0.7675,
F-statistic: 1.375e+04 on 1 and 4165 DF, p-value: < 2.2e-16
```

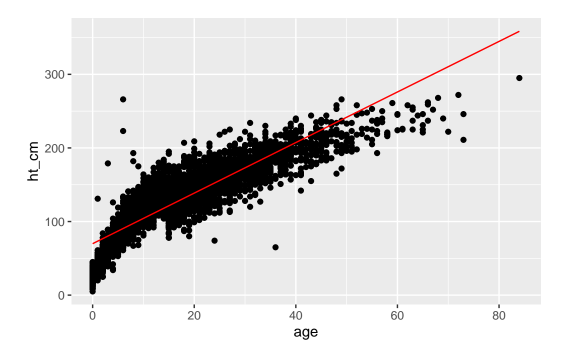
Alternatively you can use the tidy() function from the **broom** package to pull the results in to a table. What the results tell us is that for each year increase in age the height increases by 3.5 cm and this is statistically significant.

```
tidy(lm_results)
# A tibble: 2 x 5
              estimate std.error statistic p.value
  <chr>
                  <dbl>
                             <dbl>
                                       <dbl>
                                                <dbl>
1 (Intercept)
                  69.9
                           0.598
                                        117.
                                                    0
2 age
                   3.44
                           0.0293
                                        117.
                                                    0
```

You can then also use this regression to add it to a **ggplot**, to do this we first pull the points for the observed data and the fitted line in to one data frame using the **augment()** function from **broom**.

```
## pull the regression points and observed data in to one dataset
points <- augment(lm_results)

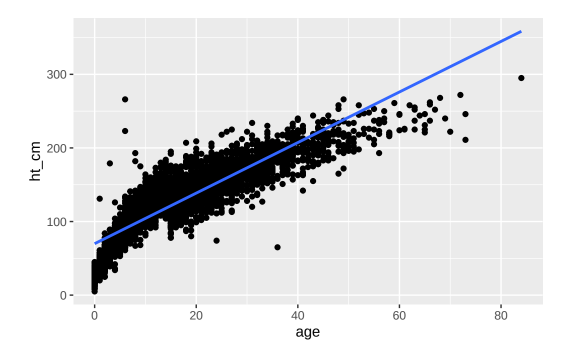
## plot the data using age as the x-axis
ggplot(points, aes(x = age)) +
    ## add points for height
geom_point(aes(y = ht_cm)) +
    ## add your regression line
geom_line(aes(y = .fitted), colour = "red")</pre>
```



It is also possible to add a simple linear regression straight straight in **ggplot** using the geom_smooth() function.

```
## add your data to a plot
ggplot(linelist, aes(x = age, y = ht_cm)) +
    ## show points
geom_point() +
    ## add a linear regression
geom_smooth(method = "lm", se = FALSE)
```

[`]geom_smooth()` using formula = 'y ~ x'



See the Resource section at the end of this chapter for more detailed tutorials.

* Logistic regression

The function glm() from the stats package (part of base R) is used to fit Generalized Linear Models (GLM).

glm() can be used for univariate and multivariable logistic regression (e.g. to get Odds Ratios). Here are the core parts:

```
# arguments for glm()
glm(formula, family, data, weights, subset, ...)
```

- formula = The model is provided to glm() as an equation, with the outcome on the left and explanatory variables on the right of a tilde ~.
- family = This determines the type of model to run. For logistic regression, use family = "binomial", for poisson use family = "poisson". Other examples are in the table below.
- data = Specify your data frame

If necessary, you can also specify the link function via the syntax family = familytype(link = "linkfunction")). You can read more in the documentation about other families and optional arguments such as weights = and subset = (?glm).

Family	Default link function
"binomial"	(link = "logit")
"gaussian"	(link = "identity")
"Gamma"	(link = "inverse")
"inverse.gaussian"	(link = "1/mu^2")
"poisson"	(link = "log")
"quasi"	<pre>(link = "identity", variance = "constant")</pre>
"quasibinomial"	(link = "logit")
"quasipoisson"	(link = "log")

When running glm() it is most common to save the results as a named R object. Then you can print the results to your console using summary() as shown below, or perform other operations on the results (e.g. exponentiate).

* Univariate glm()

In this example we are assessing the association between different age categories and the outcome of death (coded as 1 in the Preparation section). Below is a univariate model of outcome by age_cat. We save the model output as model and then print it with summary() to the console. Note the estimates provided are the *log odds* and that the baseline level is the first factor level of age_cat ("0-4").

```
model <- glm(outcome ~ age_cat, family = "binomial", data = linelist)
summary(model)</pre>
```

Call:

```
glm(formula = outcome ~ age_cat, family = "binomial", data = linelist)
```

Coefficients:

```
Estimate Std. Error z value Pr(>|z|)
(Intercept)
             0.233738
                        0.072805
                                   3.210 0.00133 **
age_cat5-9
            -0.062898
                        0.101733 -0.618
                                          0.53640
age_cat10-14 0.138204
                        0.107186
                                   1.289 0.19726
                        0.113343 -0.049
age_cat15-19 -0.005565
                                          0.96084
age_cat20-29 0.027511
                        0.102133
                                   0.269 0.78765
```

```
0.560 0.57517
age_cat30-49 0.063764
                         0.113771
age_cat50-69 -0.387889
                         0.259240 -1.496 0.13459
age_cat70+
             -0.639203
                         0.915770 -0.698 0.48518
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 5712.4 on 4166 degrees of freedom
Residual deviance: 5705.1 on 4159
                                    degrees of freedom
AIC: 5721.1
Number of Fisher Scoring iterations: 4
To alter the baseline level of a given variable, ensure the column is class Factor and move the
desired level to the first position with fct_relevel(). For example, below we take column
age_cat and set "20-29" as the baseline before piping the modified data frame into glm().
  linelist %>%
    mutate(age_cat = fct_relevel(age_cat, "20-29", after = 0)) %>%
    glm(formula = outcome ~ age_cat, family = "binomial") %>%
    summary()
Call:
glm(formula = outcome ~ age_cat, family = "binomial", data = .)
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
(Intercept)
             0.26125
                         0.07163
                                   3.647 0.000265 ***
age_cat0-4
             -0.02751
                         0.10213 -0.269 0.787652
age_cat5-9
             -0.09041
                         0.10090 -0.896 0.370220
                         0.10639
                                   1.040 0.298133
age_cat10-14 0.11069
age cat15-19 -0.03308
                         0.11259 -0.294 0.768934
age_cat30-49 0.03625
                         0.11302
                                  0.321 0.748390
age_cat50-69 -0.41540
                         0.25891
                                  -1.604 0.108625
age_cat70+
                         0.91568 -0.728 0.466546
            -0.66671
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
```

```
Null deviance: 5712.4 on 4166 degrees of freedom Residual deviance: 5705.1 on 4159 degrees of freedom
```

AIC: 5721.1

Number of Fisher Scoring iterations: 4

* Printing results

For most uses, several modifications must be made to the above outputs. The function tidy() from the package **broom** is convenient for making the model results presentable.

Here we demonstrate how to combine model outputs with a table of counts.

1) Get the *exponentiated* log odds ratio estimates and confidence intervals by passing the model to tidy() and setting exponentiate = TRUE and conf.int = TRUE.

```
model <- glm(outcome ~ age_cat, family = "binomial", data = linelist) %>%
    tidy(exponentiate = TRUE, conf.int = TRUE) %>%  # exponentiate and produce CIs
    mutate(across(where(is.numeric), round, digits = 2)) # round all numeric columns

Warning: There was 1 warning in `mutate()`.
i In argument: `across(where(is.numeric), round, digits = 2)`.
Caused by warning:
! The `...` argument of `across()` is deprecated as of dplyr 1.1.0.
Supply arguments directly to `.fns` through an anonymous function instead.

# Previously
    across(a:b, mean, na.rm = TRUE)

# Now
    across(a:b, \( \)(x) mean(x, na.rm = TRUE))
```

Below is the outputted tibble model:

Show 8 v entrie	S			Search:		
term			es	stimate 🖣		std.
(Intercept)	1.26	0.07	3.21	0	1.1	1.4
age_cat5-9	0.94	0.1	-0.62	0.54	0.77	1.1
age_cat10-14	1.15	0.11	1.29	0.2	0.93	1.4
age_cat15-19	0.99	0.11	-0.05	0.96	0.8	1.2
age_cat20-29	1.03	0.1	0.27	0.79	0.84	1.2
age_cat30-49	1.07	0.11	0.56	0.58	0.85	1.3
age_cat50-69	0.68	0.26	-1.5	0.13	0.41	1.1
age_cat70+	0.53	0.92	-0.7	0.49	0.07	3
4						•
Showing 1 to 8 of 8 e	ntries	Previo	ous 1	Next		

2) Combine these model results with a table of counts. Below, we create the a counts cross-table with the tabyl() function from **janitor**.

```
counts_table <- linelist %>%
  janitor::tabyl(age_cat, outcome)
```

Here is what this counts_table data frame looks like:

Show 8 v entries Search:					
age_cat					
0-4	338	427			
5-9	365	433			
10-14	273	396			
15-19	238	299			
20-29	345	448			
30-49	228	307			
50-69	35	30			
70+	3	2			
Showing 1 to 8 of 8 entries	Previous 1	Next			

Now we can bind the counts_table and the model results together horizontally with bind_cols() (dplyr). Remember that with bind_cols() the rows in the two data frames must be aligned perfectly. In this code, because we are binding within a pipe chain, we use . to represent the piped object counts_table as we bind it to model. To finish the process, we use select() to pick the desired columns and their order, and finally apply the base R round() function across all numeric columns to specify 2 decimal places.

Here is what the combined data frame looks like, printed nicely as an image with a function from **flextable**. The [Tables for presentation] explains how to customize such tables with **flextable**, or or you can use numerous other packages such as **knitr** or **GT**.

```
combined <- combined %>%
  flextable::qflextable()
```

* Looping multiple univariate models

Below we present a method using glm() and tidy() for a more simple approach, see the section on gtsummary.

To run the models on several exposure variables to produce univariate odds ratios (i.e. not controlling for each other), you can use the approach below. It uses str_c() from stringr to create univariate formulas (see [Characters and strings]), runs the glm() regression on each formula, passes each glm() output to tidy() and finally collapses all the model outputs together with bind_rows() from tidyr. This approach uses map() from the package purrr to iterate.

- 1) Create a vector of column names of the explanatory variables. We already have this as explanatory vars from the Preparation section of this page.
- 2) Use str_c() to create multiple string formulas, with outcome on the left, and a column name from explanatory_vars on the right. The period . substitutes for the column name in explanatory_vars.

```
explanatory_vars %>% str_c("outcome ~ ", .)
[1] "outcome ~ gender" "outcome ~ fever" "outcome ~ chills"
```

```
[4] "outcome ~ cough" "outcome ~ aches" "outcome ~ vomit"
[7] "outcome ~ age_cat"
```

- 3) Pass these string formulas to map() and set ~glm() as the function to apply to each input. Within glm(), set the regression formula as as.formula(.x) where .x will be replaced by the string formula defined in the step above. map() will loop over each of the string formulas, running regressions for each one.
- 4) The outputs of this first map() are passed to a second map() command, which applies tidy() to the regression outputs.
- 5) Finally the output of the second map() (a list of tidied data frames) is condensed with bind_rows(), resulting in one data frame with all the univariate results.

```
models <- explanatory_vars %>%
                                     # begin with variables of interest
  str_c("outcome ~ ", .) %>%
                                     # combine each variable into formula ("outcome ~ vari
  # iterate through each univariate formula
  map(
    .f = ~glm(
                                     # pass the formulas one-by-one to glm()
      formula = as.formula(.x),
                                   # within glm(), the string formula is .x
      family = "binomial",
                                     # specify type of glm (logistic)
      data = linelist)) %>%
                                     # dataset
  # tidy up each of the glm regression outputs from above
  map(
    .f = ~tidy(
      .x,
      exponentiate = TRUE,
                                   # exponentiate
                                  # return confidence intervals
      conf.int = TRUE)) %>%
  # collapse the list of regression outputs in to one data frame
  bind_rows() %>%
  # round all numeric columns
  mutate(across(where(is.numeric), round, digits = 2))
```

This time, the end object models is longer because it now represents the combined results of several univariate regressions. Click through to see all the rows of model.

Show 5 v e	ntries			Search:		
term			estim	ate 🖣		std.erro
(Intercept)	1.28	0.04	5.67	0	1.18	1.4
gender	1	0.06	-0.04	0.97	0.88	1.13
(Intercept)	1.28	0.07	3.44	0	1.11	1.48
fever	1	0.08	0.01	0.99	0.85	1.17
(Intercept)	1.28	0.03	6.98	0	1.19	1.37
4						+
Showing 1 to 5 c	of 20 entries		Previous	1	2 3	4 Next

As before, we can create a counts table from the linelist for each explanatory variable, bind it to models, and make a nice table. We begin with the variables, and iterate through them with map(). We iterate through a user-defined function which involves creating a counts table with dplyr functions. Then the results are combined and bound with the models model results.

```
## for each explanatory variable
univ_tab_base <- explanatory_vars %>%
 map(.f =
    ~{linelist %>%
                                  ## begin with linelist
        group_by(outcome) %>%
                                  ## group data set by outcome
        count(.data[[.x]]) %>%
                                  ## produce counts for variable of interest
        pivot_wider(
                                  ## spread to wide format (as in cross-tabulation)
          names_from = outcome,
          values_from = n) %>%
        drop_na(.data[[.x]]) %>%
                                         ## drop rows with missings
        rename("variable" = .x) %>%
                                         ## change variable of interest column to "variable"
        mutate(variable = as.character(variable))} ## convert to character, else non-diche
      ) %>%
  ## collapse the list of count outputs in to one data frame
  bind_rows() %>%
  ## merge with the outputs of the regression
  bind_cols(., models) %>%
  ## only keep columns interested in
  select(term, 2:3, estimate, conf.low, conf.high, p.value) %>%
  ## round decimal places
  mutate(across(where(is.numeric), round, digits = 2))
```

Below is what the data frame looks like.

Show 5	entries			Search:		
term				0 🌲	1 🕏	
(Intercept)	909	1168	1.28	1.18	1.4	0
gender	916	1174	1	0.88	1.13	0.97
(Intercept)	340	436	1.28	1.11	1.48	0
fever	1485	1906	1	0.85	1.17	0.99
(Intercept)	1472	1877	1.28	1.19	1.37	0
Showing 1 to 5	of 20 entr	ries	Prev	rious 1	2 3 4	4 Next

gtsummary package

Below we present the use of tbl_uvregression() from the gtsummary package. gtsummary functions do a good job of running statistics and producing professional-looking outputs. This function produces a table of univariate regression results.

We select only the necessary columns from the linelist (explanatory variables and the outcome variable) and pipe them into tbl_uvregression(). We are going to run univariate regression on each of the columns we defined as explanatory_vars in the data Preparation section (gender, fever, chills, cough, aches, vomit, and age cat).

Within the function itself, we provide the method = as glm (no quotes), the y = outcome column (outcome), specify to method.args = that we want to run logistic regression via family = binomial, and we tell it to exponentiate the results.

The output is HTML and contains the counts

${\bf Characteristic}$	${f N}$	\mathbf{OR}	95% CI	p-value
gender	4,167	1.00	0.88, 1.13	>0.9
fever	4,167	1.00	0.85, 1.17	> 0.9
chills	4,167	1.03	0.89, 1.21	0.7
cough	4,167	1.15	0.97, 1.37	0.11
aches	4,167	0.93	0.76, 1.14	0.5
vomit	4,167	1.09	0.96, 1.23	0.2
age_cat	4,167			
0-4				
5-9		0.94	0.77, 1.15	0.5
10-14		1.15	0.93, 1.42	0.2
15-19		0.99	0.80, 1.24	> 0.9
20-29		1.03	0.84, 1.26	0.8

Characteristic	N	OR	95% CI	p-value
30-49		1.07	0.85, 1.33	0.6
50-69		0.68	0.41, 1.13	0.13
70+		0.53	0.07, 3.20	0.5

There are many modifications you can make to this table output, such as adjusting the text labels, bolding rows by their p-value, etc. See tutorials here and elsewhere online.

Multivariable

For multivariable analysis, we again present two approaches:

- glm() and tidy()
- gtsummary package

The workflow is similar for each and only the last step of pulling together a final table is different.

Conduct multivariable

Here we use glm() but add more variables to the right side of the equation, separated by plus symbols (+).

To run the model with all of our explanatory variables we would run:

```
mv_reg <- glm(outcome ~ gender + fever + chills + cough + aches + vomit + age_cat, family
summary(mv_reg)</pre>
```

Call:

```
glm(formula = outcome ~ gender + fever + chills + cough + aches +
    vomit + age_cat, family = "binomial", data = linelist)
```

Coefficients:

```
Estimate Std. Error z value Pr(>|z|)
(Intercept)
              0.069054
                        0.131726
                                   0.524
                                             0.600
gender
              0.002448
                         0.065133
                                    0.038
                                             0.970
fever
              0.004309
                         0.080522
                                             0.957
                                    0.054
```

```
0.666
chills
              0.034112
                         0.078924
                                     0.432
cough
              0.138584
                         0.089909
                                     1.541
                                              0.123
                         0.104078 -0.679
                                              0.497
aches
             -0.070705
vomit
              0.086098
                         0.062618
                                     1.375
                                              0.169
age_cat5-9
             -0.063562
                         0.101851
                                  -0.624
                                              0.533
age_cat10-14 0.136372
                         0.107275
                                     1.271
                                              0.204
age_cat15-19 -0.011074
                         0.113640 -0.097
                                              0.922
age_cat20-29 0.026552
                         0.102780
                                     0.258
                                              0.796
age_cat30-49 0.059569
                         0.116402
                                     0.512
                                              0.609
age_cat50-69 -0.388964
                         0.262384
                                   -1.482
                                              0.138
             -0.647443
                         0.917375
                                   -0.706
                                              0.480
age_cat70+
```

(Dispersion parameter for binomial family taken to be 1)

```
Null deviance: 5712.4 on 4166 degrees of freedom Residual deviance: 5700.2 on 4153 degrees of freedom
```

AIC: 5728.2

Number of Fisher Scoring iterations: 4

If you want to include two variables and an interaction between them you can separate them with an asterisk * instead of a +. Separate them with a colon : if you are only specifying the interaction. For example:

```
glm(outcome ~ gender + age_cat * fever, family = "binomial", data = linelist)
```

Optionally, you can use this code to leverage the pre-defined vector of column names and recreate the above command using $str_c()$. This might be useful if your explanatory variable names are changing, or you don't want to type them all out again.

* Building the model

You can build your model step-by-step, saving various models that include certain explanatory variables. You can compare these models with likelihood-ratio tests using lrtest() from the package lmtest, as below:

NOTE: Using base anova (model1, model2, test = "Chisq) produces the same results

```
model1 <- glm(outcome ~ age_cat, family = "binomial", data = linelist)
model2 <- glm(outcome ~ age_cat + gender, family = "binomial", data = linelist)
lmtest::lrtest(model1, model2)</pre>
```

Likelihood ratio test

```
Model 1: outcome ~ age_cat
Model 2: outcome ~ age_cat + gender
#Df LogLik Df Chisq Pr(>Chisq)
1 8 -2852.6
2 9 -2852.6 1 2e-04 0.9883
```

Another option is to take the model object and apply the step() function from the stats package. Specify which variable selection direction you want use when building the model.

```
## choose a model using forward selection based on AIC
## you can also do "backward" or "both" by adjusting the direction
final_mv_reg <- mv_reg %>%
   step(direction = "forward", trace = FALSE)
```

You can also turn off scientific notation in your R session, for clarity:

```
options(scipen=999)
```

As described in the section on univariate analysis, pass the model output to tidy() to exponentiate the log odds and CIs. Finally we round all numeric columns to two decimal places. Scroll through to see all the rows.

```
mv_tab_base <- final_mv_reg %>%
broom::tidy(exponentiate = TRUE, conf.int = TRUE) %>% ## get a tidy dataframe of estimate mutate(across(where(is.numeric), round, digits = 2)) ## round
```

Here is what the resulting data frame looks like:

Show 10 v entri	es			Search:		
term			•	estimate 🖣		std.
(Intercept)	1.07	0.13	0.52	0.6	0.83	1.3
gender	1	0.07	0.04	0.97	0.88	1.1
fever	1	0.08	0.05	0.96	0.86	1.1
chills	1.03	0.08	0.43	0.67	0.89	1.2
cough	1.15	0.09	1.54	0.12	0.96	1.3
aches	0.93	0.1	-0.68	0.5	0.76	1.1
vomit	1.09	0.06	1.37	0.17	0.96	1.2
age_cat5-9	0.94	0.1	-0.62	0.53	0.77	1.1
age_cat10-14	1.15	0.11	1.27	0.2	0.93	1.4
age_cat15-19	0.99	0.11	-0.1	0.92	0.79	1.2
1						+
Showing 1 to 10 of	14 entries			Previous	1 2	Next

Combine univariate and multivariable

* Combine with gtsummary

The **gtsummary** package provides the **tbl_regression()** function, which will take the outputs from a regression (glm() in this case) and produce an nice summary table.

```
## show results table of final regression
mv_tab <- tbl_regression(final_mv_reg, exponentiate = TRUE)</pre>
```

Let's see the table:

```
mv_tab
```

Table printed with `knitr::kable()`, not {gt}. Learn why at https://www.danieldsjoberg.com/gtsummary/articles/rmarkdown.html
To suppress this message, include `message = FALSE` in code chunk header.

Characteristic	OR	95% CI	p-value
gender	1.00	0.88, 1.14	>0.9
fever	1.00	0.86, 1.18	> 0.9
chills	1.03	0.89, 1.21	0.7
cough	1.15	0.96, 1.37	0.12
aches	0.93	0.76, 1.14	0.5
vomit	1.09	0.96, 1.23	0.2
age_cat			
0-4			
5-9	0.94	0.77, 1.15	0.5
10-14	1.15	0.93, 1.41	0.2
15-19	0.99	0.79, 1.24	> 0.9
20-29	1.03	0.84, 1.26	0.8
30-49	1.06	0.85, 1.33	0.6
50-69	0.68	0.40, 1.13	0.14
70+	0.52	0.07, 3.19	0.5

You can also combine several different output tables produced by **gtsummary** with the **tbl_merge()** function. We now combine the multivariable results with the **gtsummary** univariate results that we created above:

Table printed with `knitr::kable()`, not {gt}. Learn why at https://www.danieldsjoberg.com/gtsummary/articles/rmarkdown.html
To suppress this message, include `message = FALSE` in code chunk header.

Characteristic	N	OR	95% CI	p-value	OR	95% CI	p-value
gender	4,167	1.00	0.88, 1.13	>0.9	1.00	0.88, 1.14	>0.9
fever	4,167	1.00	0.85, 1.17	> 0.9	1.00	0.86, 1.18	> 0.9
chills	4,167	1.03	0.89, 1.21	0.7	1.03	0.89, 1.21	0.7
cough	4,167	1.15	0.97, 1.37	0.11	1.15	0.96, 1.37	0.12
aches	4,167	0.93	0.76, 1.14	0.5	0.93	0.76, 1.14	0.5
vomit	4,167	1.09	0.96, 1.23	0.2	1.09	0.96, 1.23	0.2
age_cat	4,167						
0-4		_			_		
5-9		0.94	0.77, 1.15	0.5	0.94	0.77, 1.15	0.5
10-14		1.15	0.93, 1.42	0.2	1.15	0.93, 1.41	0.2
15-19		0.99	0.80, 1.24	> 0.9	0.99	0.79, 1.24	> 0.9
20-29		1.03	0.84, 1.26	0.8	1.03	0.84, 1.26	0.8
30-49		1.07	0.85, 1.33	0.6	1.06	0.85, 1.33	0.6
50-69		0.68	0.41, 1.13	0.13	0.68	0.40, 1.13	0.14
70+		0.53	0.07, 3.20	0.5	0.52	0.07, 3.19	0.5

* Combine with **dplyr**

An alternative way of combining the glm()/tidy() univariate and multivariable outputs is with the dplyr join functions.

- Join the univariate results from earlier (univ_tab_base, which contains counts) with the tidied multivariable results mv_tab_base
- Use select() to keep only the columns we want, specify their order, and re-name them
- Use round() with two decimal places on all the column that are class Double

```
## combine univariate and multivariable tables
left_join(univ_tab_base, mv_tab_base, by = "term") %>%
 ## choose columns and rename them
 select( # new name = old name
    "characteristic" = term,
    "recovered" = "0",
    "dead"
                   = "1",
    "univ or"
                   = estimate.x,
   "univ_ci_low" = conf.low.x,
    "univ_ci_high" = conf.high.x,
    "univ_pval"
                  = p.value.x,
    "mv_or"
                   = estimate.y,
   "mvv_ci_low" = conf.low.y,
"mv_ci_high" = conf.high.y,
    "mv_pval"
                   = p.value.y
  ) %>%
  mutate(across(where(is.double), round, 2))
```

A tibble: 20 x 11

	characteristic :	recovered	dead	univ_or	univ_ci_low	univ_ci_high	univ_pval
	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	(Intercept)	909	1168	1.28	1.18	1.4	0
2	gender	916	1174	1	0.88	1.13	0.97
3	(Intercept)	340	436	1.28	1.11	1.48	0
4	fever	1485	1906	1	0.85	1.17	0.99
5	(Intercept)	1472	1877	1.28	1.19	1.37	0
6	chills	353	465	1.03	0.89	1.21	0.68
7	(Intercept)	272	309	1.14	0.97	1.34	0.13
8	cough	1553	2033	1.15	0.97	1.37	0.11
9	(Intercept)	1636	2114	1.29	1.21	1.38	0
10	aches	189	228	0.93	0.76	1.14	0.51
11	(Intercept)	931	1144	1.23	1.13	1.34	0
12	vomit	894	1198	1.09	0.96	1.23	0.17
13	(Intercept)	338	427	1.26	1.1	1.46	0
14	age_cat5-9	365	433	0.94	0.77	1.15	0.54
15	age_cat10-14	273	396	1.15	0.93	1.42	0.2
16	age_cat15-19	238	299	0.99	0.8	1.24	0.96
17	age_cat20-29	345	448	1.03	0.84	1.26	0.79
18	age_cat30-49	228	307	1.07	0.85	1.33	0.58
19	age_cat50-69	35	30	0.68	0.41	1.13	0.13
20	age_cat70+	3	2	0.53	0.07	3.2	0.49
# 1	i 4 more variabl	es: mv_or	<dbl>,</dbl>	mvv_ci	_low <dbl>, n</dbl>	nv_ci_high <db< td=""><td>ol>,</td></db<>	ol>,

mv_pval <dbl>