riskCommunicator Manuscript Vignette

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Introduction to riskCommunicator

The riskCommunicator package facilitates the estimation of common epidemiological effect measures that are relevant to public health, but that are often not trivial to obtain from common regression models, like logistic regression. In particular, riskCommunicator estimates risk and rate differences, in addition to risk and rate ratios. The package estimates these effects using g-computation with the appropriate parametric model depending on the outcome (logistic regression for binary outcomes, Poisson regression for rate or count outcomes, negative binomial regression for overdispersed rate or count outcomes, and linear regression for continuous outcomes). Therefore, the package can handle binary, rate, count, and continuous outcomes and allows for dichotomous, categorical (>2 categories), or continuous exposure variables. Additional features include estimation of effects stratified by subgroup and adjustment of standard errors for clustering. Confidence intervals are constructed by bootstrap at the individual or cluster level, as appropriate.

This package operationalizes g-computation, which has not been widely adopted due to computational complexity, in an easy-to-use implementation tool to increase the reporting of more interpretable epidemiological results. To make the package accessible to a broad range of health researchers, our goal was to design a function that was as straightforward as the standard logistic regression functions in R (e.g. glm) and that would require little to no expertise in causal inference methods or advanced coding.
Getting started

Installation

The riskCommunicator R package is available from CRAN so can be installed using the following command:

```r
install.packages("riskCommunicator")
```

Load packages:

```r
library(riskCommunicator)
library(tidyverse)
#> -- Attaching packages --------------------------------------- tidyverse 1.3.1 --
#> v ggplot2 3.3.5   v purrr  0.3.4
#> v tibble  3.1.6   v dplyr  1.0.8
#> v tidyr  1.2.0   v stringr 1.4.0
#> v readr  2.1.2   v forcats 0.5.1
#> -- Conflicts ------------------------------------------ tidyverse_conflicts() --
#> x dplyr::filter() masks stats::filter()
#> x dplyr::lag() masks stats::lag()
library(printr)
#> Registered S3 method overwritten by 'printr':
#>   method from
#>   knit_print.data.frame rmarkdown
```

Description of main package function

The gComp function is the main function in the riskCommunicator package and allows you to estimate a variety of effects depending on your outcome and exposure of interest. The function is coded as follows:

```r
?gComp
#> Estimate difference and ratio effects with 95% confidence intervals.
#>
#> Usage:
#>
#>  gComp(
#>    data,
#>    outcome.type = c("binary", "count", "count_nb", "rate", "rate_nb", "continuous"),
#>    formula = NULL,
#>    Y = NULL,
#>    X = NULL,
#>    Z = NULL,
#>    subgroup = NULL,
#>    offset = NULL,
#>    rate.multiplier = 1,
#>    exposure.scalar = 1,
#>    R = 200,
#>    clusterID = NULL,
#>    parallel = "no",
#>    ncpus = getOption("boot.ncpus", 1L)
#>  )
#>
```
Arguments:

- data: (Required) A data.frame containing variables for 'Y', 'X', and 'Z' or with variables matching the model variables specified in a user-supplied formula. Data set should also contain variables for the optional 'subgroup' and 'offset', if they are specified.

- outcome.type: (Required) Character argument to describe the outcome type. Acceptable responses, and the corresponding error distribution and link function used in the 'glm', include:
  - binary (Default) A binomial distribution with link = 'logit' is used.
  - count A Poisson distribution with link = 'log' is used.
  - count_nb A negative binomial model with link = 'log' is used, where the theta parameter is estimated internally; ideal for over-dispersed count data.
  - rate A Poisson distribution with link = 'log' is used; ideal for events/person-time outcomes.
  - rate_nb A negative binomial model with link = 'log' is used, where the theta parameter is estimated internally; ideal for over-dispersed events/person-time outcomes.
  - continuous A gaussian distribution with link = 'identity' is used.

- formula: (Optional) Default NULL. An object of class "formula" (or one that can be coerced to that class) which provides the the complete model formula, similar to the formula for the glm function in R (e.g. 'Y ~ X + Z1 + Z2 + Z3'). Can be supplied as a character or formula object. If no formula is provided, Y and X must be provided.

- Y: (Optional) Default NULL. Character argument which specifies the outcome variable. Can optionally provide a formula instead of 'Y' and 'X' variables.

- X: (Optional) Default NULL. Character argument which specifies the exposure variable (or treatment group assignment), which can be binary, categorical, or continuous. This variable can be supplied as a factor variable (for binary or categorical exposures) or a continuous variable. For binary/categorical exposures, 'X' should be supplied as a factor with the lowest level set to the desired referent. Numeric variables are accepted, but will be centered (see Note). Character variables are not accepted and will throw an error. Can optionally provide a formula instead of 'Y' and 'X' variables.
Z: (Optional) Default NULL. List or single character vector which specifies the names of covariates or other variables to adjust for in the 'glm' function. All variables should either be factors, continuous, or coded 0/1 (i.e. not character variables). Does not allow interaction terms.

subgroup: (Optional) Default NULL. Character argument that indicates subgroups for stratified analysis. Effects will be reported for each category of the subgroup variable. Variable will be automatically converted to a factor if not already.

offset: (Optional, only applicable for rate/count outcomes) Default NULL. Character argument which specifies the variable name to be used as the person-time denominator for rate outcomes to be included as an offset in the Poisson regression model. Numeric variable should be on the linear scale; function will take natural log before including in the model.

rate.multiplier: (Optional, only applicable for rate/count outcomes). Default 1. Numeric variable signifying the person-time value to use in predictions; the offset variable will be set to this when predicting under the counterfactual conditions. This value should be set to the person-time denominator desired for the rate difference measure and must be inputted in the units of the original offset variable (e.g. if the offset variable is in days and the desired rate difference is the rate per 100 person-years, rate.multiplier should be inputted as 365.25*100).

exposure.scalar: (Optional, only applicable for continuous exposure) Default 1. Numeric value to scale effects with a continuous exposure. This option facilitates reporting effects for an interpretable contrast (i.e. magnitude of difference) within the continuous exposure. For example, if the continuous exposure is age in years, a multiplier of 10 would result in estimates per 10-year increase in age rather than per a 1-year increase in age.

R: (Optional) Default 200. The number of data resamples to be conducted to produce the bootstrap confidence interval of the estimate.

clusterID: (Optional) Default NULL. Character argument which specifies the variable name for the unique identifier for clusters. This option specifies that clustering should be accounted for in the calculation of confidence intervals. The 'clusterID' will be used as the level for resampling in the bootstrap procedure.

parallel: (Optional) Default "no." The type of parallel operation to be used. Available options (besides the default of no parallel processing) include multicore" (not available for Windows) or
Framingham Heart Study

We’ll demonstrate how to use the package with data from the Framingham Heart Study. The following information is from the official Framingham study documentation (https://biolincc.nhlbi.nih.gov/teaching/):

“The Framingham Heart Study is a long term prospective study of the etiology of cardiovascular disease among a population of free living subjects in the community of Framingham, Massachusetts. The Framingham Heart Study was a landmark study in epidemiology in that it was the first prospective study of cardiovascular disease and identified the concept of risk factors and their joint effects. The study began in 1948 and 5,209 subjects were initially enrolled in the study. Participants have been examined biennially since the inception of the study and all subjects are continuously followed through regular surveillance for cardiovascular outcomes. Clinic examination data has included cardiovascular disease risk factors and markers of disease such as blood pressure, blood chemistry, lung function, smoking history, health behaviors, ECG tracings, Echocardiography, and medication use. Through regular surveillance of area hospitals, participant contact, and death certificates, the Framingham Heart Study reviews and adjudicates events for the occurrence of Angina Pectoris, Myocardial Infarction, Heart Failure, and Cerebrovascular disease.

data(cvdd)

cvdd is a subset of the data collected as part of the Framingham study from 4,240 participants who conducted a baseline exam and were free of prevalent coronary heart disease when they entered the study. Participant clinic data was collected during three examination periods, approximately 6 years apart, from roughly 1956 to 1968. Each participant was followed for a total of 24 years for the outcome of the following events: Angina Pectoris, Myocardial Infarction, Atherothrombotic Infarction or Cerebral Hemorrhage (Stroke) or death.

NOTE: This is a “teaching” dataset. Specific methods were employed to ensure an anonymous dataset that protects patient confidentiality; therefore, this dataset is inappropriate for publication purposes.” The use of these data for the purposes of this package were approved on 11Mar2019 (request #7161) by NIH/NHLBI.

Binary outcome example

Research question: what is the effect of having diabetes at the beginning of the study on the 24-year risk of cardiovascular disease or death due to any cause?

Here, we will estimate the risk difference, risk ratio, odds ratio, and number needed to treat, adjusting for patient’s age, sex, body mass index (BMI), smoking status (current smoker or not), and prevalence of hypertension (if they are hypertensive or not at baseline). Logistic regression is used as the underlying parametric model for g-computation.

```r
## Specify the regression formula
cvdd.formula <- cvd_dth ~ DIABETES + AGE + SEX + BMI + CURSMOKE + PREVHYP
```

```r
## For reproducibility, we should always set the seed since the g-computation uses
```
## random resampling of the data to calculate confidence intervals and random sampling of the distribution when predicting outcomes.

```r
set.seed(1298)
```

## Call the gComp function

```r
binary.res <- gComp(data = cvdd,
                     formula = cvdd.formula,
                     outcome.type = "binary",
                     R = 1000)
```

binary.res

```r
#> Formula:
#> cvd_dth ~ DIABETES + AGE + SEX + BMI + CURSMOKE + PREVHYP
#>
#> Parameter estimates:
#> DIABETES1_v._DIABETES0 Estimate (95% CI)
#> Risk Difference 0.287 (0.196, 0.395)
#> Risk Ratio 1.700 (1.476, 1.966)
#> Odds Ratio 4.550 (2.771, 9.085)
#> Number needed to treat/harm 3.484
```

The result obtained from the gComp function is an object of class `gComp` which is a list containing the summary results, `results.df`, `n`, `R`, `boot.result`, `contrast`, `family`, `formula`, `predicted.outcome`, and `glm.result` (see `?gComp` or `help(gComp)` for a more detailed explanation of each item in the list).

```r
class(binary.res)
#> [1] "gComp" "list"
```

## The names of the different items in the list:

```r
names(binary.res)
#> [1] "summary" "results.df" "n"
#> [4] "R" "boot.result" "contrast"
#> [7] "family" "formula" "predicted.outcome"
#> [10] "glm.result"
```

## Sample size of the original data:

```r
binary.res$n
#> [1] 4240
```

## Contrast being compared in the analysis:

```r
binary.res$contrast
#> [1] "DIABETES1 v. DIABETES0"
```

### Rate outcome example

Research question: what is the effect of having diabetes at the beginning of the study on the rate of cardiovascular disease or death due to any cause?

Here, we will estimate the rate difference and rate ratio, adjusting for patient’s age, sex, body mass index (BMI), smoking status (current smoker or not), and prevalence of hypertension (if they are hypertensive or not at baseline). We have included `timeout` as the offset and a `rate.multiplier` of 365.25*100 so that the estimates are returned with units of 100 person-years. Poisson regression is used as the underlying
parametric model for g-computation. (Note: for overdispersed count/rate outcomes, the negative binomial distribution can be specified by setting `outcome.type` to “count_nb” or “rate_nb”.)

```r
## Modify the dataset to change the variable cvd_dth from a factor to a numeric variable since the outcome for Poisson regression must be numeric.
cvdd.t <- cvdd %>%
  dplyr::mutate(cvd_dth = as.numeric(as.character(cvd_dth)),
                timeout = as.numeric(timeout))

set.seed(6534)
rate.res <- gComp(data = cvdd.t,
                  Y = "cvd_dth",
                  X = "DIABETES",
                  Z = c("AGE", "SEX", "BMI", "CURSMOKE", "PREVHYP"),
                  outcome.type = "rate",
                  rate.multiplier = 365.25*100,
                  offset = "timeout",
                  R = 1000)
rate.res
#> Formula:
#> cvd_dth ~ DIABETES + AGE + SEX + BMI + CURSMOKE + PREVHYP + offset(log(timeout_adj))
#>
#> Parameter estimates:
#> DIABETES1_v._DIABETES0 Estimate (95% CI)
#> Incidence Rate Difference 2.189 (1.436, 3.063)
#> Incidence Rate Ratio 1.913 (1.603, 2.288)
```

Rate outcome with subgroups example

Research question: what is the effect of having diabetes at the beginning of the study on the rate of cardiovascular disease or death due to any cause, stratified by sex?

Here, we will estimate the same effects above, but in subgroups defined by sex.

```r
## Modify the dataset to change the variable cvd_dth from a factor to a numeric variable since the outcome for Poisson regression must be numeric.
set.seed(6534)
rate.res.subgroup <- gComp(data = cvdd.t,
                           Y = "cvd_dth",
                           X = "DIABETES",
                           Z = c("AGE", "SEX", "BMI", "CURSMOKE", "PREVHYP"),
                           subgroup = "SEX",
                           outcome.type = "rate",
                           rate.multiplier = 365.25*100,
                           offset = "timeout",
                           R = 1000)
rate.res.subgroup
#> Formula:
```
Checking model fit

To ensure that the parameter estimates from each bootstrap iteration are normally distributed, we can also look at the histogram and Q-Q plots of bootstrapped estimates by calling:

```r
plot(binary.res)
```

```
sessionInfo()
```

```
#> R version 4.1.2 (2021-11-01)
#> Platform: x86_64-apple-darwin17.0 (64-bit)
#> Running under: macOS Catalina 10.15.7
```
Matrix products: default
BLAS: /Library/Frameworks/R.framework/Versions/4.1/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/4.1/Resources/lib/libRlapack.dylib
locale:
attached base packages:
 [1] stats graphics grDevices utils datasets methods base
other attached packages:
[1] printr_0.2 forcats_0.5.1 stringr_1.4.0
[4] dplyr_1.0.8 purrr_0.3.4 readr_2.1.2
[7] tidyr_1.2.0 tibble_3.1.6 ggplot2_3.3.5
[10] tidyverse_1.3.1 riskCommunicator_1.0.0
loaded via a namespace (and not attached):
[1] tidyselect_1.1.1 xfun_0.29 haven_2.4.3 colorspace_2.0-2
[5] vctrs_0.3.8 generics_0.1.2 htmltools_0.5.2 yaml_2.2.2
[9] utf8_1.2.2 rlang_1.0.1 pillar_1.7.0 withr_2.4.3
[13] glue_1.6.1 DBI_1.1.2 dbplyr_2.1.1 modelr_0.1.8
[17] dbplyr_2.1.1 lifecycle_1.0.1 cellranger_1.1.0 munsell_0.5.0
[21] Rcpp_1.0.8 scales_1.1.1 backports_1.4.1
[25] broom_0.7.12 Rcpp_1.0.8 scales_1.1.1 backports_1.4.1
[33] jsonlite_1.7.3 farver_2.1.0 fs_1.5.2 gridExtra_2.3
[37] hms_1.1.1 digest_0.6.29 stringi_1.7.6 grid_4.1.2
[41] cli_3.1.1 tools_4.1.2 magrittr_2.0.2 crayon_1.4.2
[45] pkgconfig_2.0.3 MASS_7.3-55 ellipsis_0.3.2 xml2_1.3.3
[49] reprex_2.0.1 lubridate_1.8.0 assertthat_0.2.1 rmarkdown_2.11
[53] httr_1.4.2 rstudioapi_0.13 R6_2.5.1 boot_1.3-28
[57] compiler_4.1.2