# Noname manuscript No. (will be inserted by the editor)

## My Example Computed Manuscript Created in Rmarkdown

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**Abstract** A mock computed manuscript created in RStudio using {Rmarkdown}. The {Bookdown} and {Rticles} packages were used to output the text in Springer Nature's desired manuscript format.

#### **Keywords**

#### 1 Introduction

"Literate programming" is a style of programming that uses computational notebooks to weave together code, explanatory text, data and results into a single document, enhancing scientific communication and computational reproducibility. 1–3 (These references were added into the document using RStudio's integration with the open-source Zotero reference manager<sup>4</sup> plus the Better BibTeX Zotero plugin.)

Several platforms for creating such documents exist.<sup>5</sup> Typically, these documents interleave code and text 'blocks' to build a computational narrative. But some, including R Markdown, Observable, and the Jupyter Book extension to the Jupyter ecosystem, also allow authors to include and execute code "inline" – that is, within the text itself.

This makes it possible to create fully executable manuscripts in which the document itself computes and inserts values and figures into the text rather than requiring authors to input them manually. This is in many ways the 'killer feature' of computed manuscripts: it circumvents the possibility that the author will enter an incorrect number, or forget to update a figure or value should

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new data arise. Among other uses, that allows authors to automatically time-stamp their documents, or insert the current version number of the software they use into their methods. For instance, this document was built at **02 Feb 2022 10:28:23 MST** and calls the following R packages: {tidyverse} ver. **1.3.0**, {ggbeeswarm} ver. **0.6.0** and {bookdown} ver. **0.24**.

In this manuscript, created in RStudio using the R Markdown language, we will demonstrate a more practical example. (A JavaScript version is available at https://observablehq.com/d/7a08166446057936.)

#### 2 Results

#### 2.1 Inline computation

Imagine we are analyzing data from a clinical trial. We have grouped subjects in three bins and measured the concentration of some metabolite. (These data are simulated.)

Rather than analyzing those data and then copying the results into our manuscript, we can use the programming language R to do that in the manuscript itself. Simply enclose the code inside backticks, with the letter r. For instance, we could calculate the circumference and area of a circle:

$$A = \pi r^2, C = 2\pi r$$

You could write "A = `r pi \* r^2` and C = `r 2 \* pi \* r`". Plugging in the radius r = 10, that evaluates to "A = 314.16 and C = 62.83".

Returning to our dataset, we can count the rows in our table to determine the number of samples, and insert that into the text. Thus, we have **99** (simulated) subjects in our study (see Table 1; see R/mock\_data.R in the GitHub repository for code to generate the mock dataset). Note that the tables, figures and sections in this document are numbered automatically thanks to the {bookdown} package.

The average metabolite concentration in our dataset is 185.36 (range: 78 to 298). We have 32 subjects in Group 1, 43 subjects in Group 2, and 24 in Group 3. (The numbers in **bold face type** throughout this document are computed values.)

#### 2.2 Incorporating new data

Now suppose we get another tranche of data (Table 2). There are **60** subjects in this new dataset. Their average concentration is **185.13** (range: **77 to 299**).

Combining the two datasets, we have a total of **159** subjects. The revised average metabolite concentration is **185.28** (range: **77 to 299**). We now have **55** subjects in Group 1, **60** subjects in Group 2, and **44** in Group 3. The concentration distribution for each group is shown graphically in Figure 1.

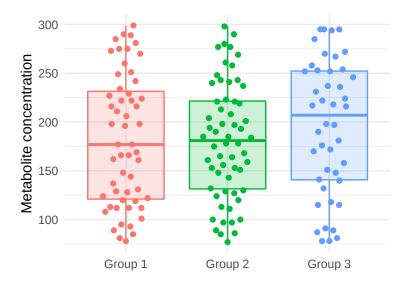


Fig. 1 Metabolite concentration of clinical trial subjects

#### 2.3 Incorporating a child document

Authors can break long manuscripts into more manageable pieces by using the R Markdown child option. The text in this section is pulled in from a second file, child\_doc.Rmd, and inserted into the final document. Citations that are created in the child automatically get inserted into the final document, like so: here's a reference for the R Markdown Cookbook.<sup>6</sup>

```
\# number of samples in our new dataset samples <- 50
```

Using the build\_mock\_dataset() function in the file R/mock\_data.R, we can create and add a third set of numbers to our growing dataset. By changing the samples value in the code cell above, you can make this dataset as big as you'd like, but for now, it contains 50 samples.

These final data are shown in Table 3. Folding them into our existing dataset gives us **209** subjects: **73** in Group 1, **70** in Group 2, and **66** in Group 3. The new concentration distribution is shown graphically in Figure 2.

#### 3 Code

The following code was used to load, merge, and plot the (simulated) clinical trial data in Figure 1:

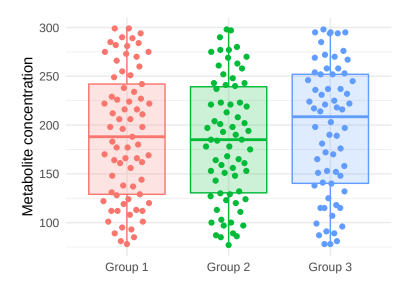


Fig. 2 Metabolite concentration of final clinical trial dataset

```
# load libraries
library(tidyverse)
library(ggbeeswarm)
library(bookdown)
# read in some initial data
df1 <- read_csv('data/example-data-1.csv')</pre>
# read new dataset
df2 <- read_csv('data/example-data-2.csv')</pre>
# merge datasets
final_data <- rbind(df1, df2)</pre>
# create a box-plot with overlaid points
create_plot <- function(mytable) {</pre>
  p <- mytable \%>\%
    ggplot(aes(x = class, y = conc, fill = class, color = class)) +
    geom_boxplot(outlier.shape = NA, alpha = 0.2) +
    ggbeeswarm::geom_quasirandom(width = 0.25) +
    xlab("") +
    ylab("Metabolite concentration") +
    theme_minimal() +
    theme(legend.position = "none")
```

 ${\bf Table~1}~{\rm Initial~subject~data}$ 

ID	Class	Conc	_	ID	Class	Conc	_	ID	Class	Conc
ID_1	Group 2	153		ID_34	Group 2	221		ID_67	Group 3	148
$ID_{-2}$	Group 1	224	_	$ID_{-}35$	Group 1	112	_	$ID_{-}68$	Group 1	281
$ID_3$	Group 2	127	_	$ID_{-}36$	Group 3	246	_	$ID_{-}69$	Group 3	295
$ID_{-4}$	Group 2	194	_	$ID_{-}37$	Group 2	190	_	$ID_{-}70$	Group 2	111
$ID_{-5}$	Group 1	251	_	$ID_{-}38$	Group 1	177	_	$ID_{-}71$	Group $2$	132
$ID_{-}6$	Group 1	81	_	$ID_39$	Group 1	148	_	${\rm ID}\_72$	Group $2$	261
$\mathrm{ID}_{\text{-}}7$	Group 2	100	_	$ID_{-}40$	Group 2	290	_	$ID_{-}73$	Group 1	122
ID8	Group 1	270	_	$ID_{-}41$	Group 2	151	_	$ID_{-}74$	Group 2	124
$ID_{-}9$	Group 2	100	_	$ID_{-}42$	Group 2	159	_	$ID_{-}75$	Group 1	234
$ID_{-}10$	Group 1	161	_	$ID_{-}43$	Group 2	113	_	$ID_{-}76$	Group 2	184
ID_11	Group 3	158	_	$ID_{-}44$	Group 1	249	_	$ID_{-}77$	Group 3	272
$ID_{-}12$	Group 3	118		$ID_{-}45$	Group 1	124	_	$ID_{-}78$	Group 1	242
$ID_{-}13$	Group 2	143	_	$ID_{-}46$	Group 3	87	_	$ID_{-}79$	Group 2	277
$ID_{-}14$	Group 2	258		$ID_{-}47$	Group 1	166	_	ID80	Group 3	236
$ID_{-}15$	Group 3	224	_	$ID_{-}48$	Group 1	196	_	ID_81	Group 1	101
ID_16	Group 3	254	_	$ID_{-}49$	Group 1	112	_	ID82	Group 3	218
$ID_{-}17$	Group 3	190		$ID_{-}50$	Group 1	289	_	ID83	Group 2	130
$ID_{-}18$	Group 2	148		$ID_{-}51$	Group 2	161	_	ID84	Group 1	128
$ID_{-}19$	Group 1	89		$ID_{-}52$	Group 3	270	_	ID85	Group 3	252
$ID_{-}20$	Group $2$	89	_	$ID_{-}53$	Group 2	237	_	ID86	Group 1	198
$ID_{-}21$	Group 3	253	_	$ID_{-}54$	Group 2	280	_	ID_87	Group 1	169
$ID_{-}22$	Group 3	231	_	$ID_{-}55$	Group 2	175	_	ID88	Group 2	185
$ID_{-}23$	Group 1	112		$ID_{-}56$	Group 2	223	_	ID89	Group 1	216
$ID_{-}24$	Group 2	277		$ID_{-}57$	Group 3	295	_	$ID_{-}90$	Group 2	185
$ID_{-}25$	Group $2$	197	_	$ID_{-}58$	Group 1	275	_	ID_91	Group 2	97
$ID_{-}26$	Group 2	208	_	$ID_{-}59$	Group 2	120	_	$ID_{-}92$	Group 2	165
$ID_{-}27$	Group 2	193	_	$ID_{-}60$	Group 1	78	_	$ID_{-}93$	Group 3	89
$ID_{-}28$	Group 3	141	_	$ID_{-}61$	Group 3	78	_	$ID_{-}94$	Group 2	221
$ID_{-}29$	Group 1	206	_	$ID_{-}62$	Group 3	140	_	$ID_{-}95$	Group 1	162
$ID_{-}30$	Group 2	168	_	${\rm ID}$ -63	Group $3$	294	_	$ID_{-}96$	Group 1	131
$ID_{-}31$	Group 2	298	_	$ID_{-}64$	Group 3	295	_	$ID_{-}97$	Group 1	93
$ID_{-}32$	Group 1	144	_	$ID_{-}65$	Group 3	285	_	$ID_{-}98$	Group 2	240
$ID_{-}33$	Group 2	241	_	$ID_{-}66$	Group 2	129	_	$ID_{-}99$	Group 2	86

```
p
}
# plot the data
create_plot(final_data)
```

### 4 Colophon

This manuscript was built at 02 Feb 2022 10:28:24 MST using the following computational environment and dependencies:

Table 2 New subject data

ID	Class	Conc	_	ID	Class	Conc	_	ID	Class	Conc
ID_100	Group 2	219	_	ID_120	Group 2	85	_	ID_140	Group 2	77
$ID_{-}101$	Group 2	243		$ID_{-}121$	Group 3	181	_	ID_141	Group 1	299
$ID_{-}102$	Group 2	213	_	$ID_{-}122$	Group 3	216	_	$ID_{-}142$	Group 3	222
ID <sub>-</sub> 103	Group 1	177	_	$ID_{-}123$	Group 1	222	_	$ID_{-}143$	Group 1	85
$ID_{-}104$	Group 3	197	_	$ID_{-}124$	Group 3	252	_	ID_144	Group 1	273
$ID_{-}105$	Group 2	198		$ID_{-}125$	Group 1	166	_	ID_145	Group 3	115
ID <sub>-</sub> 106	Group 1	120	_	$ID_{-}126$	Group 2	204	_	$ID_{-}146$	Group 1	290
ID_107	Group 3	170	_	$ID_{-}127$	Group 2	243	_	$ID_{-}147$	Group 2	269
ID_108	Group 3	78	_	$ID_{-}128$	Group 3	198	_	$ID_{-}148$	Group 2	97
$ID_{-}109$	Group 1	129	_	$ID_{-}129$	Group 1	119	_	ID_149	Group 1	229
ID_110	Group 1	137		ID_130	Group 1	198	_	ID_150	Group 3	176
$ID_{-}111$	Group 3	217		$ID_{-}131$	Group 3	151	_	$ID_{-}151$	Group 2	164
$ID_{-}112$	Group 1	227		$ID_{-}132$	Group 3	115	_	$ID_{-}152$	Group 3	172
ID_113	Group 3	81		ID_133	Group 3	237	_	$ID_{-}153$	Group 1	222
ID_114	Group 2	248		ID_134	Group 2	178	_	ID_154	Group 1	285
$ID_{-}115$	Group 1	211	_	$ID_{-}135$	Group 1	275	_	ID <sub>-</sub> 155	Group 2	153
$ID_{-}116$	Group 1	113		ID_136	Group 2	178	_	$ID_{-}156$	Group 3	132
ID_117	Group 1	216	_	$ID_{-}137$	Group 3	267	_	$ID_{-}157$	Group 2	156
ID_118	Group 3	91	_	$ID_{-}138$	Group 1	95	_	$ID_{-}158$	Group 1	260
ID <sub>-</sub> 119	Group 3	258		ID <sub>-</sub> 139	Group 1	108	_	ID <sub>-</sub> 159	Group 2	201

Table 3 Third subject data

ID	Class	Conc	_	ID	Class	Conc	_	ID	Class	Conc
ID_1	Group 3	245	_	ID_18	Group 3	237	_	ID_35	Group 3	125
$ID_{-2}$	Group 3	214	_	$ID_{-}19$	Group 3	115	_	$ID_{-}36$	Group 2	103
$ID_{-}3$	Group 2	270	_	$ID_{-}20$	Group 1	226	_	$ID_{-}37$	Group 2	87
$ID_{-4}$	Group 3	152	_	$ID_{-}21$	Group 1	284	_	$ID_{-}38$	Group 2	275
$ID_{-}5$	Group 1	164	_	$ID_{-}22$	Group 3	269	_	$ID_{-}39$	Group 2	223
$ID_{-}6$	Group 1	188	_	$ID_{-}23$	Group 3	225	_	$ID_{-}40$	Group 1	180
$ID_{-}7$	Group 3	165	_	$ID_{-}24$	Group 3	276	_	$ID_{-}41$	Group 2	297
ID8	Group 3	153	_	$ID_{-}25$	Group 3	258	_	$ID_{-}42$	Group 1	138
$ID_{-}9$	Group 1	212	_	$ID_{-}26$	Group 2	252	_	$ID_{-}43$	Group 3	189
$ID_{-}10$	Group 1	183	_	$ID_{-}27$	Group 1	282	_	$ID_{-}44$	Group $3$	138
ID_11	Group 1	156	_	$ID_{-}28$	Group 1	255	_	$ID_{-}45$	Group 3	221
$ID_{-}12$	Group 3	99	_	$ID_{-}29$	Group 1	294	_	$ID_{-}46$	Group 3	203
$ID_{-}13$	Group 1	299	_	$ID_{-}30$	Group 3	107	_	$ID_{-}47$	Group 1	266
$ID_{-}14$	Group 1	238	_	$ID_{-}31$	Group 2	123	_	$ID_{-}48$	Group 3	298
$ID_{-}15$	Group 3	294	_	$ID_{-}32$	Group 3	232	_	$ID_{-}49$	Group 1	206
ID_16	Group 1	170	_	ID_33	Group 2	202	_	ID_50	Group 1	101
ID_17	Group 3	96	_	$ID_{-}34$	Group 2	263	_		-	

```
## R version 4.0.4 (2021-02-15)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Big Sur 10.16
## Matrix products: default
          /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRlapack.dylib
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
                graphics grDevices utils
## [1] stats
                                               datasets methods
                                                                   base
##
## other attached packages:
    [1] bookdown_0.24
##
                         ggbeeswarm_0.6.0 forcats_0.5.1
                                                           stringr_1.4.0
##
   [5] dplyr_1.0.5
                         purrr_0.3.4
                                          readr_2.1.1
                                                           tidyr_1.1.3
## [9] tibble_3.1.6
                         ggplot2_3.3.3
                                          tidyverse_1.3.0
##
## loaded via a namespace (and not attached):
   [1] Rcpp_1.0.7
                         lubridate_1.8.0 assertthat_0.2.1 digest_0.6.29
##
##
    [5] utf8_1.2.2
                         R6_2.5.1
                                          cellranger_1.1.0 backports_1.2.1
   [9] reprex_2.0.0
                         evaluate_0.14
                                          highr_0.9
                                                           httr_1.4.2
## [13] pillar_1.6.4
                        rlang_0.4.12
                                          readxl_1.3.1
                                                           rstudioapi_0.13
                         rmarkdown_2.11
## [17] rticles_0.22
                                          labeling_0.4.2
                                                           bit_4.0.4
## [21] munsell_0.5.0
                         broom_0.7.6
                                          compiler_4.0.4
                                                           vipor_0.4.5
## [25] modelr_0.1.8
                        xfun_0.29
                                          pkgconfig_2.0.3
                                                           htmltools_0.5.2
## [29] tidyselect_1.1.1 fansi_1.0.0
                                          crayon_1.4.2
                                                           tzdb_0.2.0
                         withr_2.4.3
## [33] dbplyr_2.1.0
                                          grid_4.0.4
                                                           jsonlite_1.7.2
## [37] gtable_0.3.0
                         lifecycle_1.0.1 DBI_1.1.1
                                                           magrittr_2.0.1
## [41] scales_1.1.1
                         cli_3.1.0
                                          stringi_1.7.6
                                                           vroom_1.5.7
                         fs_1.5.2
## [45] farver_2.1.0
                                          xm12_1.3.3
                                                           ellipsis_0.3.2
## [49] generics_0.1.1 vctrs_0.3.8
                                          tools_4.0.4
                                                           bit64_4.0.5
## [53] glue_1.6.0
                         beeswarm_0.4.0
                                          hms_1.1.1
                                                           parallel_4.0.4
## [57] fastmap_1.1.0
                         yaml_2.2.1
                                          colorspace_2.0-0 rvest_1.0.2
## [61] knitr_1.37
                         haven_2.3.1
```

The current Git commit details are:

## [10dc11d] 2022-02-02: Add a child document.

#### References

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