R for Clinical Study Reports and Submission

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Welcome

Preface

Folder structure

In clinical trial development, source code needs to be developed and maintained to generate and deliver Study Data Tabulation Model (SDTM), Analysis Dataset Model (ADaM) datasets and tables, listings, and figures (TLFs). A typical example is a Phase 3 clinical trial where hundreds of TLFs are required for submission. Considering the number of programs required for such an effort, a consistent and well-defined folder structure is crucial in managing a clinical trial analysis and reporting (A&R) project.

We recommend using the R package folder structure to organize all the A&R-related source code and documentation for a clinical trial A&R project. The R package folder structure is well defined and widely used in the R community through repositories (e.g., CRAN).

The consistent approach of using the R package folder structure simplifies the communication for all developers within and across organizations.

- For a new R developer, it is an essential step to develop R packages when you want to share your work with others. You will learn one folder structure widely used in the R community with outstanding tutorials and tools for free.
- For an experienced R developer, there is a minimal learning curve.
- For an organization, it simplifies the process, tool, template, and training development, because a unified folder structure is used to develop and maintain standard tool and analysis projects.

Preface

The workflow around an R package can also improve the traceability and reproducibility of an analysis project (Marwick, Boettiger, and Mullen 2018).

We will revisit folder structure when we discuss project management for a clinical trial project.

In addition, R package folder structure is also recommended to develop Shiny apps as discussed in Chapter 20 of the Mastering Shiny book and the Engineering Production-Grade Shiny Apps book.

In this book

This book is an intermediate-level book by assuming readers have R programming and clinical development knowledge. The assumption of each part is as below:

- Part 1, "Delivering TLFs in CSR" provides general information with examples to create tables, listings, and figures. In this part, we assume readers are individual contributors to a clinical project with some experience in R programming. We expect readers are familiar with data manipulation in R. Some good references include Hands-On Programming with R, R for Data Science and Data Manipulation with R.
- Part 2, "Clinical trial project" provides general information with examples to manage a clinical trial A&R project. In this part, we assume a reader is a project lead with experience in R package development. Some good references include R Packages and the tidyverse style guide.
- Part 3, "eCTD submission package" provides general information in preparing submission packages related to clinical study report (CSR) in electronic Common Technical Document (eCTD). In this part, we

assume a reader is a project lead of a clinical project with experience in R package development and submission.

Philosophy

We share the same philosophy described in Section 1.1 of the R Packages book and quote here.

- "Anything that can be automated, should be automated."
- "Do as little as possible by hand. Do as much as possible with functions."
- "The goal is to spend your time thinking about what you want to do rather than thinking about the minutiae of the package structure."

Authors and contributors

The document is maintained by a community. While reading the document, you can be a contributor as well. The quality of this document relies on you.

- Authors: contributed the majority of content to at least one chapter.

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- Contributors: contributed at least one commit to the source code.

We are grateful for all the improvements brought by these contributors (in chronological order): Yujie Zhao (@LittleBeannie), Aiming Yang, Steven Haesendonckx (@SHAESEN2), Howard Baek (@howardbaek), Xiaoxia Han (@echohan), Jie Wang (@ifendo).

Part I. Delivering TLFs in CSR

1.1. Background

In clinical trials, a critical step is to submit trial results to regulatory agencies. Electronic Common Technical Document (eCTD) has become a worldwide regulatory submission standard format. For example, the United States Food and Drug Administration (US FDA) requires new drug applications and biologics license applications must be submitted using the eCTD format. The Clinical Data Interchange Standards Consortium (CDISC) provides a pilot project following ICH E3 guidance.

Within eCTD, clinical study reports (CSRs) are located at module 5. ICH E3 guidance provides a compilation of the structure and content of clinical study reports.

A typical CSR contains full details on the methods and results of an individual clinical study. In support of the statistical analysis, a large number of tables, listings, and figures are incorporated into the main text and appendices. In the CDISC pilot project, an example CSR is also provided. If you are interested in more examples of clinical study reports, you can go to the European Medicines Agency (EMA) clinical data website.

Building CSRs is teamwork between clinicians, medical writers, statisticians, statistical programmers, and other relevant specialists such as experts on biomarkers. Here, we focus on the work and deliverables completed by statisticians and statistical programmers. In an organization, they commonly work together to define, develop, validate and deliver tables, listings, and figures (TLFs) required for a CSR to summarize the

efficacy and/or safety of the pharmaceutical product. Microsoft Word is widely used to prepare CSR in the pharmaceutical industry. Therefore, .rtf, .doc, .docx are commonly used formats in their deliverables.

In this chapter, our focus is to illustrate how to create tables, listings, and figures (TLFs) in RTF format that is commonly used in a CSR. The examples are in compliance with the FDA's Portable Document Format (PDF) Specifications.

Note

FDA's PDF specification is a general reference. Each organization can define more specific TLF format requirements that can be different from the examples in this book.

1.2. Structure and content

In the rest of this chapter, we are following the ICH E3 guidance on the structure and content of clinical study reports.

In a CSR, most of TLFs are located in

- Section 10: Study participants
- Section 11: Efficacy evaluation
- Section 12: Safety evaluation
- Section 14: Tables, listings, and figures referrals but not included in the text
- Section 16: Appendices

1.3. Datasets

We used publicly available CDISC pilot study data located in the CDISC GitHub repository.

For simplicity, we have downloaded all these datasets into the data-adam/ folder of this project and converted them from the .xpt format to the .sas7bdat format.

The dataset structure follows CDISC Analysis Data Model (ADaM).

1.4. Tools

In this part, we mainly use the R packages below to illustrate how to deliver TLFs in a CSR.

- tidyverse: prepare datasets ready for reporting.
- r2rtf: create RTF outputs

Note

There are other R packages to create TLFs in ASCII, RTF and Word format. For example, rtables, huxtable, pharmaRTF, gt, officer, flextable etc. Here we focus on r2rtf to illustrate the concept. Readers are encouraged to explore other R packages to find the proper tools to fit your purpose.

1.4.1. tidyverse

tidyverse is a collection of R packages to simplify the workflow to manipulate, visualize and analyze data in R. Those R packages share the tidy tools manifesto and are easy to use for interactive data analysis.

Posit provided outstanding cheatsheets and tutorials for tidyverse.

There are also books to introduce tidyverse. We assume the reader have experience in using tidyverse in this book.

- The tidyverse cookbook
- R for Data Science

1.4.2. r2rtf

r2rtf is an R package to create production-ready tables and figures in RTF format. This R package is designed to

- provide simple "verb" functions that correspond to each component of a table, to help you translate a data frame to a table in an RTF file:
- enable pipes (%);
- focus on the **table format** only. Data manipulation and analysis shall be handled by other R packages (e.g., tidyverse).

Before creating an RTF table, we need to

- figure out the table layout;
- split the layout into small tasks in the form of a computer program;
- execute the program.

We provide a brief introduction of r2rtf and show how to transfer data frames into table, listing, and figures (TLFs).

Other extended examples and features are covered on the r2rtf package website.

To explore the basic RTF generation verbs in r2rtf, we will use the dataset r2rtf_adae saved in the r2rtf package. This dataset contains adverse events (AEs) information from a clinical trial.

We will begin by loading the packages:

```
library(dplyr) # Manipulate data
library(tidyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
```

Below is the meaning of relevant variables. More information can be found on the help page of the dataset (?r2rtf_adae)

In this example, we consider three variables:

- USUBJID: Unique Subject Identifier
- TRTA: Actual Treatment
- AEDECOD: Dictionary-Derived Term

dplyr and tidyr packages within tidyverse are used for data manipulation to create a data frame that contains all the information we want to add in an RTF table.

```
tbl <- r2rtf_adae %>%
  count(TRTA, AEDECOD) %>%
  pivot_wider(names_from = TRTA, values_from = n, values_fill = 0)

tbl %>% head(4)
#> # A tibble: 4 x 4
```

#>		AEDECOD		Placebo	`Xanomeline	High	Dose`	`Xanomeline	Low	Dose`
#>		<chr></chr>		<int></int>			<int></int>			<int></int>
#>	1	ABDOMINAL	PAIN	1			2			3
#>	2	AGITATION		2			1			2
#>	3	ALOPECIA		1			0			0
#>	4	ANXIETY		2			0			4

Now we have a dataset tbl in preparing the final RTF table.

r2rtf aims to provide one function for each type of table layout. Commonly used verbs include:

- rtf_page(): RTF page information
- rtf_title(): RTF title information
- rtf_colheader(): RTF column header information
- rtf_body(): RTF table body information
- rtf_footnote(): RTF footnote information
- rtf_source(): RTF data source information

All these verbs are designed to enable the usage of pipes (%>%). A full list of all functions can be found in the r2rtf package function reference manual.

A minimal example below illustrates how to combine verbs using pipes to create an RTF table.

- rtf_body() is used to define table body layout.
- rtf_encode() transfers table layout information into RTF syntax.
- write_rtf() save RTF encoding into a file with file extension .rtf

```
head(tbl) %>%
  rtf_body() %>% # Step 1 Add table attributes
  rtf_encode() %>% # Step 2 Convert attributes to RTF encode
  write_rtf("tlf/intro-ae1.rtf") # Step 3 Write to a .rtf file
```

1.4. Tools

AEDECOD	Placebo	Xanomeline High Dose	Xanomeline Low Dose
ABDOMINAL PAIN	1	2	3
AGITATION	2	1	2
ALOPECIA	1	0	0
ANXIETY	2	0	4
APPLICATION SITE DERMATITIS	9	12	15
APPLICATION SITE ERYTHEMA	3	23	20

If we want to adjust the width of each column to provide more space to the first column, this can be achieved by updating the col_rel_width argument in the rtf_body() function.

In this example, the input of col_rel_width is a vector with the same length for the number of columns. This argument defines the relative width of each column within a pre-defined total column width.

In this example, the defined relative width is 3:2:2:2. Only the ratio of col_rel_width is used. Therefore it is equivalent to use col_rel_width = c(6, 4, 4, 4) or col_rel_width = c(1.5, 1, 1, 1).

```
head(tbl) %>%
  rtf_body(col_rel_width = c(3, 2, 2, 2)) %>%
  # define relative width
  rtf_encode() %>%
  write_rtf("tlf/intro-ae2.rtf")
```

In the previous example, we found the issue of a misaligned column header. We can fix the issue by using the rtf_colheader() function.

In rtf_colheader(), the colheader argument is used to provide the content of the column header. We use "|" to separate the columns.

In the example below, "Adverse Events | Placebo | Xanomeline High Dose | Xanomeline Low Dose" define a column header with 4 columns.

```
head(tbl) %>%
  rtf_colheader(
    colheader = "Adverse Events | Placebo | Xanomeline High Dose | Xanomel
    col_rel_width = c(3, 2, 2, 2)
) %>%
  rtf_body(col_rel_width = c(3, 2, 2, 2)) %>%
  rtf_encode() %>%
```

1.4. Tools

AEDECOD	Placebo		Xanomeline High Dose		Xanomeline Low Dose
ABDOMINAL PAIN		1		2	3
AGITATION		2		1	2
ALOPECIA		1		0	0
ANXIETY		2		0	4
APPLICATION SITE DERMATITIS		9		12	15
APPLICATION SITE ERYTHEMA		3		23	20

```
write_rtf("tlf/intro-ae3.rtf")
```

In rtf_*() functions such as rtf_body(), rtf_footnote(), the text_justification argument is used to align text. Default is "c" for center justification. To vary text justification by column, use character vector with length of vector equals to number of columns displayed (e.g., c("c", "l", "r")).

All possible inputs can be found in the table below.

```
r2rtf:::justification()
               name rtf_code_text rtf_code_row
     type
#> 1
       1
               left
                             \\ql
                                        \\trql
#> 2
                             \\qc
                                        \\trqc
             center
#> 3
                             \\qr
                                        \\trqr
       r
             right
                             \\qj
#> 4
       d
            decimal
        j justified
                             \\qj
```

Below is an example to make the first column left-aligned and centeraligned for the rest.

```
head(tbl) %>%
  rtf_body(text_justification = c("l", "c", "c", "c")) %>%
  rtf_encode() %>%
  write_rtf("tlf/intro-ae5.rtf")
```

In rtf_*() functions such as rtf_body(), rtf_footnote(), etc., border_left, border_right, border_top, and border_bottom control cell borders.

If we want to remove the top border of "Adverse Events" in the header, we can change the default value "single" to "" in the border_top argument, as shown below.

1.4. Tools

Adverse Events	Placebo	Xanomeline High Dose	Xanomeline Low Dose
ABDOMINAL PAIN	1	2	3
AGITATION	2	1	2
ALOPECIA	1	0	0
ANXIETY	2	0	4
APPLICATION SITE DERMATITIS	9	12	15
APPLICATION SITE ERYTHEMA	3	23	20

AEDECOD	Placebo	Xanomeline High Dose	Xanomeline Low Dose	
ABDOMINAL PAIN	1	2	3	
AGITATION	2	1	2	
ALOPECIA	1	0	0	
ANXIETY	2	0	4	
APPLICATION SITE	9	12	15	
DERMATITIS				
APPLICATION SITE	3	23	20	
ERYTHEMA				

r2rtf supports 26 different border types. The details can be found on the r2rtf package website.

In this example, we also demonstrate the possibility of adding multiple column headers.

```
head(tbl) %>%
  rtf_colheader(
    colheader = " | Treatment",
    col_rel_width = c(3, 6)
) %>%
  rtf_colheader(
    colheader = "Adverse Events | Placebo | Xanomeline High Dose | Xanomeline Low Dose",
    border_top = c("", "single", "single"),
    col_rel_width = c(3, 2, 2, 2)
) %>%
  rtf_body(col_rel_width = c(3, 2, 2, 2)) %>%
  rtf_encode() %>%
  write_rtf("tlf/intro-ae7.rtf")
```

In the r2rtf R package get started page, there are more examples to illustrate how to customize

- title, subtitle
- footnote, data source
- special character
- etc.

Those features will be introduced when we first use them in the rest of the chapters.

	Treatment			
Adverse Events	Placebo	Xanomeline High Dose	Xanomeline Low Dose	
ABDOMINAL PAIN	1	2	3	
AGITATION	2	1	2	
ALOPECIA	1	0	0	
ANXIETY	2	0	4	
APPLICATION SITE DERMATITIS	9	12	15	
APPLICATION SITE ERYTHEMA	3	23	20	

2. Disposition

Following ICH E3 guidance, a summary table needs to be provided to include all participants who entered the study in Section 10.1, Disposition of Participants.

The disposition of participants table reports the numbers of participants who were randomized, and who entered and completed each phase of the study. In addition, the reasons for all post-randomization discontinuations, grouped by treatment and by major reason (lost to follow-up, adverse event, poor compliance, etc.) are reported.

```
library(haven) # Read SAS data
library(dplyr) # Manipulate data
library(tidyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
```

In this chapter, we show how to create a typical disposition table.

The first step is to read in the relevant datasets into R. For a disposition table, all the required information is saved in a Subject-level Analysis Dataset (ADSL). This dataset is provided in sas7bdat format, which is a SAS data format currently used in many clinical trial analysis and reporting. The haven package is able to read the dataset, while maintaining its attributes (e.g., variable labels).

```
adsl <- read_sas("data-adam/adsl.sas7bdat")</pre>
```

2. Disposition

Disposition of Participants

	Placebo		Xanomeline Low Dose		Xanomeline High Dose	
	n	(%)	n	(%)	n	(%)
Participants in population	86		84		84	
Completed	58	67.4	25	29.8	27	32.1
Discontinued	28	32.6	59	70.2	57	67.9
Adverse Event	8	9.3	44	52.4	40	47.6
Death	2	2.3	1	1.2	0	0.0
I/E Not Met	1	1.2	0	0.0	2	2.4
Lack of Efficacy	3	3.5	0	0.0	1	1.2
Lost to Follow-up	1	1.2	1	1.2	0	0.0
Physician Decision	1	1.2	0	0.0	2	2.4
Protocol Violation	1	1.2	1	1.2	1	1.2
Sponsor Decision	2	2.3	2	2.4	3	3.6
Withdrew Consent	9	10.5	10	11.9	8	9.5

The following variables are used in the preparation of a simplified disposition of participants table:

- USUBJID: unique subject identifier
- TRT01P: planned treatment
- TRT01PN: planned treatment numeric encoding
- DISCONFL: discontinued from study flag
- DCREASCD: discontinued from study reason coded

```
adsl %>%
 select(USUBJID, TRT01P, TRT01PN, DISCONFL, DCREASCD) %>%
 head(4)
#> # A tibble: 4 x 5
    USUBJID TRT01P
                                     TRT01PN DISCONFL DCREASCD
     <chr>
               <chr>
                                       <dbl> <chr> <chr>
                                          0 ""
#> 1 01-701-1015 Placebo
                                                     Completed
                                           O "Y"
#> 2 01-701-1023 Placebo
                                                    Adverse Event
                                          81 ""
#> 3 01-701-1028 Xanomeline High Dose
                                                     Completed
#> 4 01-701-1033 Xanomeline Low Dose
                                          54 "Y"
                                                      Sponsor Decision
```

In the code below, we calculate the number of participants in the analysis population by treatment arms.

```
n_rand <- adsl %>%
  group_by(TRT01PN) %>%
  summarize(n = n()) %>%
  pivot_wider(
    names_from = TRT01PN,
    names_prefix = "n_",
    values_from = n
) %>%
  mutate(row = "Participants in population")
```

2. Disposition

```
n_rand
#> # A tibble: 1 x 4
     n_0 n_54 n_81 row
     <int> <int> <int> <chr>
                   84 Participants in population
#> 1
       86
             84
n_disc <- adsl %>%
  group_by(TRT01PN) %>%
  summarize(
   n = sum(DISCONFL == "Y"),
   pct = formatC(n / n() * 100,
     digits = 1, format = "f", width = 5
    )
  ) %>%
  pivot_wider(
   names_from = TRT01PN,
   values_from = c(n, pct)
  ) %>%
  mutate(row = "Discontinued")
n_disc
#> # A tibble: 1 x 7
     n_0 n_54 n_81 pct_0 pct_54 pct_81 row
#> <int> <int> <chr> <chr>
                                             <chr>
#> 1
       28
             59 57 " 32.6" " 70.2" " 67.9" Discontinued
```

In the code below, we calculate the number and percentage of participants who completed/discontinued the study for different reasons by treatment arms.

```
n_reason <- adsl %>%
  group_by(TRT01PN) %>%
```

```
mutate(n_total = n()) %>%
 group_by(TRT01PN, DCREASCD) %>%
 summarize(
   n = n(),
   pct = formatC(n / unique(n_total) * 100,
     digits = 1, format = "f", width = 5
   )
 ) %>%
 pivot_wider(
   id_cols = DCREASCD,
   names_from = TRT01PN,
   values_from = c(n, pct),
   values_fill = list(n = 0, pct = " 0.0")
 ) %>%
 rename(row = DCREASCD)
n_reason
\#> \# A tibble: 10 x 7
     row
                         n_0 n_54 n_81 pct_0 pct_54 pct_81
     <chr>
                       <int> <int> <chr> <chr>
                                                        <chr>
#>
                                     40 " 9.3" " 52.4" " 47.6"
#>
  1 Adverse Event
                          8
                                44
                                      27 " 67.4" " 29.8" " 32.1"
#> 2 Completed
                          58
                                25
#> 3 Death
                           2
                                 1
                                      0 " 2.3" " 1.2" "
                                                          0.0"
                                      2 " 1.2" " 0.0" "
#> 4 I/E Not Met
                           1
                                 0
                                      1 " 3.5" " 0.0" "
#> 5 Lack of Efficacy
                           3
                                 0
#> 6 Lost to Follow-up
                           1
                                 1
                                      0 " 1.2" " 1.2" "
#> 7 Physician Decision
                           1
                                0
                                      2 " 1.2" " 0.0" "
#> 8 Protocol Violation
                           1
                                 1
                                      1 " 1.2" " 1.2" "
#> 9 Sponsor Decision
                           2
                                 2
                                      3 " 2.3" "
                                                   2.4" "
                                                          3.6"
                                       8 " 10.5" " 11.9" " 9.5"
#> 10 Withdrew Consent
                                10
```

In the code below, we calculate the number and percentage of participants who complete the study by treatment arms. We split n_reason because

2. Disposition

we want to customize the row order of the table.

In the code below, we calculate the numbers and percentages of participants who discontinued the study for different reasons by treatment arms. For display purpose, pasteo(" ", row) is used to add leading spaces to produce indentation in the final report.

```
n reason <- n reason %>%
 filter(row != "Completed") %>%
 mutate(row = paste0("
                         ", row))
n_reason
#> # A tibble: 9 x 7
    row
                               n_0 n_54 n_81 pct_0 pct_54 pct_81
     <chr>
                             <int> <int> <int> <chr>
#>
#> 1 "
                                 8
                                           40 "
                                                 9.3" " 52.4" " 47.6"
         Adverse Event"
                                 2
                                            0 " 2.3" " 1.2" "
#> 2 "
         Death"
#> 3 "
                                            2 " 1.2" "
                                                         0.0" "
         I/E Not Met"
                                 1
#> 4 "
         Lack of Efficacy"
                                 3
                                      0
                                            1 "
                                                 3.5" "
                                                         0.0" "
#> 5 "
                                            0 "
                                                 1.2" "
         Lost to Follow-up"
                                 1
                                      1
                                                         1.2" "
                                                                 0.0"
                                            2 " 1.2" "
#> 6 "
         Physician Decision"
                                      0
                                                         0.0" "
                                                                 2.4"
                                 1
                                                         1.2" "
#> 7 "
         Protocol Violation"
                                 1
                                      1
                                            1 " 1.2" "
                                                                 1.2"
                                 2
                                            3 " 2.3" " 2.4" "
#> 8 "
         Sponsor Decision"
                                      2
                                                                 3.6"
                                            8 " 10.5" " 11.9" " 9.5"
#> 9 "
         Withdrew Consent"
                                 9
                                      10
```

Now we combine individual rows into one table for reporting purpose. tbl_disp is used as input for r2rtf to create final report.

```
tbl_disp <- bind_rows(n_rand, n_complete, n_disc, n_reason) %>%
  select(row, ends_with(c("_0", "_54", "_81")))
tbl_disp
#> # A tibble: 12 x 7
#>
     row
                                   n_0 pct_0
                                             #>
     <chr>
                                 <int> <chr>
                                              <int> <chr>
                                                            <int> <chr>
   1 "Participants in population"
                                    86 <NA>
                                                 84 <NA>
                                                               84 <NA>
#>
#>
   2 "Completed"
                                    58 " 67.4"
                                                 25 " 29.8"
                                                               27 " 32.1"
   3 "Discontinued"
                                    28 " 32.6"
                                                 59 " 70.2"
#>
                                                               57 " 67.9"
   4 "
          Adverse Event"
                                     8 " 9.3"
                                                 44 " 52.4"
                                                               40 " 47.6"
#>
   5 "
          Death"
                                     2 " 2.3"
                                                  1 " 1.2"
                                                               0 " 0.0"
#>
   6 "
                                                                2 "
          I/E Not Met"
                                     1 " 1.2"
                                                  0.0"
                                                                    2.4"
#>
                                     3 " 3.5"
   7 "
          Lack of Efficacy"
                                                  0.0"
                                                                1 " 1.2"
   8 "
          Lost to Follow-up"
                                     1 " 1.2"
                                                  1 " 1.2"
                                                                0 " 0.0"
                                     1 " 1.2"
                                                  0.0"
                                                                2 " 2.4"
   9 "
          Physician Decision"
#>
#> 10 "
          Protocol Violation"
                                     1 " 1.2"
                                                  1 " 1.2"
                                                                1 " 1.2"
                                                  2 " 2.4"
                                     2 " 2.3"
                                                                3 " 3.6"
#> 11 "
          Sponsor Decision"
#> 12 "
          Withdrew Consent"
                                     9 " 10.5"
                                                 10 " 11.9"
                                                               8 " 9.5"
```

In the below code, formatting of the final table is defined. Items that were not discussed in the previous sections, are highlighted below.

The rtf_title defines table title. We can provide a vector for the title argument. Each value is a separate line. The format can also be controlled by providing a vector input in text format.

```
tbl_disp %>%
    # Table title
    rtf_title("Disposition of Participants") %>%
    # First row of column header
```

2. Disposition

```
rtf_colheader(" | Placebo | Xanomeline Low Dose| Xanomeline High Dose",
  col_rel_width = c(3, rep(2, 3))
) %>%
# Second row of column header
rtf_colheader(" | n | (%) | n | (%) | n | (%)",
  col_rel_width = c(3, rep(c(0.7, 1.3), 3)),
  border_top = c("", rep("single", 6)),
  border_left = c("single", rep(c("single", ""), 3))
) %>%
# Table body
rtf_body(
  col_rel_width = c(3, rep(c(0.7, 1.3), 3)),
  text_justification = c("l", rep("c", 6)),
  border_left = c("single", rep(c("single", ""), 3))
) %>%
# Encoding RTF syntax
rtf_encode() %>%
# Save to a file
write_rtf("tlf/tbl_disp.rtf")
```

The procedure to generate a disposition table can be summarized as follows:

- Step 1: Read subject level data (i.e., ads1) into R.
- Step 2: Count participants in the analysis population and name the dataset n_rand.
- Step 3: Calculate the number and percentage of participants who discontinued the study by treatment arm, and name the dataset n_disc.
- Step 4: Calculate the numbers and percentages of participants who discontinued the study for different reasons by treatment arm, and name the dataset n_reason.
- Step 5: Calculate the number and percentage of participants who

Disposition of Participants

	P	Placebo		line Low Dose	Xanome	Xanomeline High Dose	
	n	(%)	n	(%)	n	(%)	
Participants in population	86		84		84		
Completed	58	67.4	25	29.8	27	32.1	
Discontinued	28	32.6	59	70.2	57	67.9	
Adverse Event	8	9.3	44	52.4	40	47.6	
Death	2	2.3	1	1.2	0	0.0	
I/E Not Met	1	1.2	0	0.0	2	2.4	
Lack of Efficacy	3	3.5	0	0.0	1	1.2	
Lost to Follow-up	1	1.2	1	1.2	0	0.0	
Physician Decision	1	1.2	0	0.0	2	2.4	
Protocol Violation	1	1.2	1	1.2	1	1.2	
Sponsor Decision	2	2.3	2	2.4	3	3.6	
Withdrew Consent	9	10.5	10	11.9	8	9.5	

2. Disposition

completed the study by treatment arm, and name the dataset ${\tt n_complete}.$

- Step 6: Bind n_rand, n_disc, n_reason, and n_complete by row.
- Step 7: Write the final table to RTF

3. Analysis population

Following ICH E3 guidance, we need to summarize the number of participants included in each efficacy analysis in Section 11.1, Data Sets Analysed.

```
library(haven) # Read SAS data
library(dplyr) # Manipulate data
library(tidyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
```

In this chapter, we illustrate how to create a summary table for the analysis population for a study.

The first step is to read relevant datasets into R. For the analysis population table, all the required information is saved in the ADSL dataset. We can use the haven package to read the dataset.

```
adsl <- read_sas("data-adam/adsl.sas7bdat")</pre>
```

We illustrate how to prepare a report data for a simplified analysis population table using variables below:

- USUBJID: unique subject identifier
- ITTFL: intent-to-treat population flag
- EFFFL: efficacy population flag
- SAFFL: safety population flag

3. Analysis population

Summary of Analysis Sets (All Participants Randomized)

	Placebo n (%)	Xanomeline line Low Dose n (%)	Xanomeline line High Dose n (%)
Participants in Population	86	84	84
Participants included in ITT population	86 (100.0)	84 (100.0)	84 (100.0)
Participants included in efficacy population	79 (91.9)	81 (96.4)	74 (88.1)
Participants included in safety population	86 (100.0)	84 (100.0)	84 (100.0)

```
ads1 %>%
  select(USUBJID, ITTFL, EFFFL, SAFFL) %>%
  head(4)
#> # A tibble: 4 x 4
     USUBJID
                 ITTFL EFFFL SAFFL
#>
     <chr>
                 <chr> <chr> <chr>
#> 1 01-701-1015 Y
                       Y
                              Υ
#> 2 01-701-1023 Y
                       Υ
                              Υ
#> 3 01-701-1028 Y
                       Υ
                              Υ
#> 4 01-701-1033 Y
                              Υ
```

3.1. Helper functions

Before we write the analysis code, let's discuss the possibility of reusing R code by writing helper functions.

As discussed in R for data science, "You should consider writing a function whenever you've copied and pasted a block of code more than twice".

In Chapter 2, there are a few repeating steps to:

- Format the percentages using the formatC() function.
- Calculate the numbers and percentages by treatment arm.

We create two ad-hoc functions and use them to create the tables in the rest of this book.

To format numbers and percentages, we create a function called fmt_num(). It is a very simple function wrapping formatC().

```
fmt_num <- function(x, digits, width = digits + 4) {
  formatC(
    x,</pre>
```

3. Analysis population

```
digits = digits,
  format = "f",
  width = width
)
}
```

The main reason to create the fmt_num() function is to enhance the readability of the analysis code.

For example, we can compare the two versions of code to format the percentage used in Chapter 2 and fmt_num().

```
formatC(n / n() * 100,
   digits = 1, format = "f", width = 5
)

fmt_num(n / n() * 100, digits = 1)
```

To calculate the numbers and percentages of participants by groups, we provide a simple (but not robust) wrapper function, count_by(), using the dplyr and tidyr package.

The function can be enhanced in multiple ways, but here we only focus on simplicity and readability. More details about writing R functions can be found in the STAT 545 course.

```
left_join(
    count(data, grp, var),
    count by(adsl, "PRTO1PN" "EFFFL") %>%
    select("ends, with(c("-54", "-81")))

#> #varte(n-0 pct_0 npct_0 var_label

#> chr> chr> num(150 * fchr) tot, digits chr),
#> 1 Nn = fmt_num(150 * fchr) tot, digits chr),
#> 2 Y npct = paste0(n, "" (", pct, ")") EFFFL

#> 2 Y npct = paste0(n, "" (", pct, ")") EFFFL

By using the count_by() function, we can simplify the analysis code as below. pivot_wider(
    id_cols = var,
        names_from = grp,
        values_fill = list(n = "0", pct = fmt_num(0, digits = 0))
    ) %>%
    mutate(var_label = var_label)
}
```

3.2. Analysis code

With the helper function count_by, we can easily prepare a report dataset as

```
# Derive a randomization flag
adsl <- adsl %>% mutate(RANDFL = "Y")
pop <- count_by(adsl, "TRT01PN", "RANDFL",</pre>
  var_label = "Participants in Population"
) %>%
  select(var_label, starts_with("n_"))
pop1 <- bind_rows(</pre>
  count_by(adsl, "TRT01PN", "ITTFL",
    var_label = "Participants included in ITT population"
  count_by(adsl, "TRT01PN", "EFFFL",
    var_label = "Participants included in efficacy population"
  count_by(adsl, "TRT01PN", "SAFFL",
    var_label = "Participants included in safety population"
  )
) %>%
  filter(var == "Y") %>%
  select(var_label, starts_with("npct_"))
```

Now we combine individual rows into one table for reporting purpose. tbl_pop is used as input for r2rtf to create the final report.

```
names(pop) <- gsub("n_", "npct_", names(pop))
tbl_pop <- bind_rows(pop, pop1)</pre>
```

We define the format of the output using code below.

```
rel_width \leftarrow c(2, rep(1, 3))
colheader <- " | Placebo | Xanomeline line Low Dose| Xanomeline line High Dose"
tbl_pop %>%
  # Table title
  rtf_title(
    "Summary of Analysis Sets",
    "(All Participants Randomized)"
  ) %>%
  # First row of column header
  rtf_colheader(colheader,
    col_rel_width = rel_width
  ) %>%
  # Second row of column header
  rtf_colheader(" | n (%) | n (%) | n (%)",
    border_top = "",
    col_rel_width = rel_width
  ) %>%
  # Table body
  rtf_body(
    col_rel_width = rel_width,
    text_justification = c("l", rep("c", 3))
```

3. Analysis population

```
) %>%
# Encoding RTF syntax
rtf_encode() %>%
# Save to a file
write_rtf("tlf/tbl_pop.rtf")
```

The procedure to generate an analysis population table can be summarized as follows:

- Step 1: Read data (i.e., adsl) into R.
- Step 2: Bind the counts/percentages of the ITT population, the efficacy population, and the safety population by row using the count_by() function.
- Step 3: Format the output from Step 2 using r2rtf.

3.2. Analysis code

Summary of Analysis Sets (All Participants Randomized)

	Placebo n (%)	Xanomeline line Low Dose n (%)	Xanomeline line High Dose n (%)
Participants in Population	86	84	84
Participants included in ITT population	86 (100.0)	84 (100.0)	84 (100.0)
Participants included in efficacy population	79 (91.9)	81 (96.4)	74 (88.1)
Participants included in safety population	86 (100.0)	84 (100.0)	84 (100.0)

4. Baseline characteristics

Following ICH E3 guidance, we need to summarize critical demographic and baseline characteristics of the participants in Section 11.2, Demographic and Other Baseline Characteristics.

In this chapter, we illustrate how to create a simplified baseline characteristics table for a study.

There are many R packages that can efficiently summarize baseline information. The table 1 R package is one of them.

```
library(table1)
library(r2rtf)
library(haven)
library(dplyr)
library(tidyr)
library(stringr)
library(tools)
```

As in previous chapters, we first read the adsl dataset that contains all the required information for the baseline characteristics table.

```
adsl <- read_sas("data-adam/adsl.sas7bdat")</pre>
```

For simplicity, we only analyze SEX, AGE and, RACE in this example using the table1 R package. More details of the table1 R package can be found in the package vignettes.

4. Baseline characteristics

Baseline Characteristics of Participants (All Participants Randomized)

		Xanomeline	Xanomeline	
	Placebo	High Dose	Low Dose	Overall
	(N=86)	(N=84)	(N=84)	(N=254)
SEX				
Female	53 (61.6%)	40 (47.6%)	50 (59.5%)	143 (56.3%)
Male	33 (38.4%)	44 (52.4%)	34 (40.5%)	111 (43.7%)
Age				
Mean (SD)	75.2 (8.59)	74.4 (7.89)	75.7 (8.29)	75.1 (8.25)
Median [Min, Max]	76.0 [52.0, 89.0]	76.0 [56.0, 88.0]	77.5 [51.0, 88.0]	77.0 [51.0, 89.0]
RACE				
Black or African American	8 (9.3%)	9 (10.7%)	6 (7.1%)	23 (9.1%)
White	78 (90.7%)	74 (88.1%)	78 (92.9%)	230 (90.6%)
American Indian or Alaska Native	0 (0%)	1 (1.2%)	0 (0%)	1 (0.4%)

The table 1 R package directly creates an HTML report.

```
ana <- adsl %>%
  mutate(
    SEX = factor(SEX, c("F", "M"), c("Female", "Male")),
    RACE = toTitleCase(tolower(RACE))
)

tbl <- table1(~ SEX + AGE + RACE | TRTO1P, data = ana)
tbl</pre>
```

	Placebo	Xanomeline High Dose	Xanomeline Low Dose	C
	(N=86)	(N=84)	(N=84)	(]
SEX				ļ
Female	53~(61.6%)	40~(47.6%)	50~(59.5%)	1
Male	$33\ (38.4\%)$	44~(52.4%)	$34\ (40.5\%)$	1
\mathbf{Age}				ļ
Mean (SD)	75.2 (8.59)	74.4 (7.89)	75.7 (8.29)	7
Median [Min, Max]	76.0 [52.0, 89.0]	76.0 [56.0, 88.0]	77.5 [51.0, 88.0]	7
RACE				
Black or African American	8 (9.3%)	9 (10.7%)	6 (7.1%)	2
White	78 (90.7%)	74 (88.1%)	78 (92.9%)	2
American Indian or Alaska Native	0 (0%)	1 (1.2%)	0 (0%)	1

The code below transfer the output into a dataframe that only contains ASCII characters recommended by regulatory agencies. tbl_base is used as input for r2rtf to create the final report.

```
tbl_base <- tbl %>%
  as.data.frame() %>%
  as_tibble() %>%
```

4. Baseline characteristics

```
mutate(across(
    everything(),
    ~ str_replace_all(.x, intToUtf8(160), " ")
  ))
names(tbl_base) <- str_replace_all(names(tbl_base), intToUtf8(160), " ")</pre>
tbl_base
#> # A tibble: 11 x 5
                        Placebo `Xanomeline High Dose` `Xanomeline Low Dose
#>
      <chr>
                        <chr>
                                 <chr>
                                                         <chr>
    1 ""
                        "(N=86~ "(N=84)"
                                                         "(N=84)"
                        11.11
                                 11-11
                                                         11.11
#>
    2 "SEX"
    3 " Female"
                        "53 (6~ "40 (47.6%)"
                                                         "50 (59.5%)"
                                                         "34 (40.5%)"
    4 " Male"
                        "33 (3~ "44 (52.4%)"
                                 11-11
    5 "Age"
                        11.11
                       "75.2 ~ "74.4 (7.89)"
                                                         "75.7 (8.29)"
    6 " Mean (SD)"
    7 " Median [Min,~ "76.0 ~ "76.0 [56.0, 88.0]"
                                                         "77.5 [51.0, 88.0]"
                        11.11
                                                         11 11
    8 "RACE"
    9 " Black or Afr~ "8 (9.~ "9 (10.7%)"
                                                         "6 (7.1%)"
                                                         "78 (92.9%)"
#> 10 " White"
                        "78 (9~ "74 (88.1%)"
                                                         "0 (0%)"
#> 11 " American Ind~ "0 (0%~ "1 (1.2%)"
```

We define the format of the output. We highlight items that are not discussed in previous discussion.

text_indent_first and text_indent_left are used to control the indent space of text. They are helpful when you need to control the white space of a long phrase, "AMERICAN INDIAN OR ALASKA NATIVE" in the table provides an example.

```
colheader1 <- paste(names(tbl_base), collapse = "|")
colheader2 <- paste(tbl_base[1, ], collapse = "|")</pre>
```

```
rel_width <- c(2.5, rep(1, 4))
tbl_base[-1, ] %>%
  rtf_title(
    "Baseline Characteristics of Participants",
    "(All Participants Randomized)"
  ) %>%
  rtf_colheader(colheader1,
    col_rel_width = rel_width
  ) %>%
  rtf_colheader(colheader2,
    border_top = "",
    col_rel_width = rel_width
  ) %>%
  rtf_body(
    col_rel_width = rel_width,
    text_justification = c("l", rep("c", 4)),
    text_indent_first = -240,
    text_indent_left = 180
  ) %>%
  rtf_encode() %>%
  write_rtf("tlf/tlf_base.rtf")
```

In conclusion, the procedure to generate demographic and baseline characteristics table is summarized as follows:

- Step 1: Read the data set.
- Step 2: Use table1::table1() to get the baseline characteristics table.
- Step 3: Transfer the output from Step 2 into a data frame that only contains ASCII characters.
- Step 4: Define the format of the RTF table by using the R package r2rtf.

4. Baseline characteristics

Baseline Characteristics of Participants (All Participants Randomized)

	Placebo (N=86)	Xanomeline High Dose (N=84)	Xanomeline Low Dose (N=84)	Overall (N=254)
SEX				
Female	53 (61.6%)	40 (47.6%)	50 (59.5%)	143 (56.3%)
Male	33 (38.4%)	44 (52.4%)	34 (40.5%)	111 (43.7%)
Age				
Mean (SD)	75.2 (8.59)	74.4 (7.89)	75.7 (8.29)	75.1 (8.25)
Median [Min, Max]	76.0 [52.0, 89.0]	76.0 [56.0, 88.0]	77.5 [51.0, 88.0]	77.0 [51.0, 89.0]
RACE				
Black or African American	8 (9.3%)	9 (10.7%)	6 (7.1%)	23 (9.1%)
White	78 (90.7%)	74 (88.1%)	78 (92.9%)	230 (90.6%)
American Indian or Alaska Native	0 (0%)	1 (1.2%)	0 (0%)	1 (0.4%)

Following ICH E3 guidance, primary and secondary efficacy endpoints need to be summarized in Section 11.4, Efficacy Results and Tabulations of Individual Participant.

```
library(haven) # Read SAS data
library(dplyr) # Manipulate data
library(tidyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
library(emmeans) # LS mean estimation
```

In this chapter, we illustrate how to generate an efficacy table for a study. For efficacy analysis, only the change from baseline glucose data at week 24 is analyzed.

5.1. Analysis dataset

To prepare the analysis, both adsl and adlbc datasets are required.

```
adsl <- read_sas("data-adam/adsl.sas7bdat")
adlb <- read_sas("data-adam/adlbc.sas7bdat")</pre>
```

First, both the population and the data in scope are selected. The analysis is done on the efficacy population, identified by EFFFL == "Y", and all records post baseline (AVISITN >= 1) and on or before Week 24 (AVISITN

ANCOVA of Change from Baseline Glucose (mmol/L) at Week 24 $$\operatorname{LOCF}$$ Efficacy Analysis Population

		Baseline	Week 24			Change from Baseline		
Treatment	N	Mean (SD)	N	Mean (SD)	N	Mean (SD)	LS Mean (95% CI) ^a	
Placebo	79	5.7 (2.23)	57	5.7 (1.83)	57	-0.1 (2.68)	0.07 (-0.27, 0.41)	
Xanomeline Low Dose	79	5.4 (0.95)	26	5.7 (1.26)	26	0.2 (0.82)	-0.11 (-0.45, 0.23)	
Xanomeline High Dose	74	5.4 (1.37)	30	6.0 (1.92)	30	0.5 (1.94)	0.40 (0.05, 0.75)	
Pairwise Comparison			Difference in LS Mean (95% CI) ^a				p-Value	
Xanomeline Low Dose - Placebo			-0.17 (-0.65, 0.30)				0.757	
Xanomeline High Dose - Placebo			0.33 (-0.16, 0.82)			0.381		
*Based on an ANCOVA mo	*Based on an ANCOVA model after adjusting baseline value LOCE approach is used to impute missing values							

"Based on an ANCOVA model after adjusting baseline value. LOCF approach is u: ANCOVA = Analysis of Covariance, LOCF = Last Observation Carried Forward CI = Confidence Interval, LS = Least Squares, SD = Standard Deviation

<= 24). Here the variable AVISITN is the numerical analysis visit. For
example, if the analysis visit is recorded as "Baseline" (i.e., AVISIT =
Baseline), AVISITN = 0; if the analysis visit is recorded as "Week 24"
(i.e., AVISIT = Week 24), AVISITN = 24; if the analysis visit is blank,
AVISITN is also blank. We will discuss these missing values in Section
6.4.</pre>

```
gluc <- adlb %>%
  left_join(adsl %>% select(USUBJID, EFFFL), by = "USUBJID") %>%
  # PARAMCD is parameter code and here we focus on Glucose (mg/dL)
  filter(EFFFL == "Y" & PARAMCD == "GLUC") %>%
  arrange(TRTPN) %>%
  mutate(TRTP = factor(TRTP, levels = unique(TRTP)))

ana <- gluc %>%
  filter(AVISITN > 0 & AVISITN <= 24) %>%
  arrange(AVISITN) %>%
  mutate(AVISIT = factor(AVISIT, levels = unique(AVISIT)))
```

Below is the first few records of the analysis dataset.

- AVAL: analysis value
- BASE: baseline value
- CHG: change from baseline

```
select(USUBJID, TRTPN, AVISIT, AVAL, BASE, CHG) %>%
 head(4)
#> # A tibble: 4 x 6
   USUBJID
               TRTPN AVISIT
                                       AVAL BASE
                                                     CHG
#>
    <chr>
          <dbl> <fct>
                                       <dbl> <dbl>
                                                    <dbl>
#> 1 01-701-1015 0 "
                              Week 2" 4.66 4.72 -0.0555
#> 2 01-701-1023
                 0 "
                               Week 2" 5.77 5.33 0.444
```

```
#> 3 01-701-1047 0 " Week 2" 5.55 5.55 0  
#> 4 01-701-1118 0 " Week 2" 4.88 4.05 0.833
```

5.2. Helper functions

To prepare the report, we create a few helper functions by using the fmt_num() function defined in Chapter 3.

• Format estimators

• Format confidence interval

5.3. Summary of observed data

First the observed data at Baseline and Week 24 are summarized using code below:

```
t11 <- gluc %>%
  filter(AVISITN %in% c(0, 24)) %>%
  group_by(TRTPN, TRTP, AVISITN) %>%
  summarise(
   n = n(),
   mean_sd = fmt_est(mean(AVAL), sd(AVAL))
) %>%
```

```
pivot_wider(
    id_cols = c(TRTP, TRTPN),
    names_from = AVISITN,
    values_from = c(n, mean_sd)
t11
#> # A tibble: 3 x 6
               TRTPN, TRTP [3]
#> # Groups:
     TRTP
                                  n_0 n_24 mean_sd_0
                          TRTPN
                                                             mean_sd_24
     <fct>
                          <dbl> <int> <int> <chr>
#>
                                                             <chr>
                                         57 " 5.7 ( 2.23)" "
#> 1 Placebo
                              0
                                   79
                                                                5.7 (1.83)
#> 2 Xanomeline Low Dose
                             54
                                   79
                                         26 " 5.4 ( 0.95)" "
                                                                5.7 (1.26)
#> 3 Xanomeline High Dose
                             81
                                   74
                                         30 " 5.4 ( 1.37)" "
                                                                6.0 (1.92)
```

Also the observed change from baseline glucose at Week 24 is summarized using code below:

```
t12 <- gluc %>%
  filter(AVISITN %in% 24) %>%
  group_by(TRTPN, AVISITN) %>%
  summarise(
    n_chg = n(),
    mean_chg = fmt_est(
        mean(CHG, na.rm = TRUE),
        sd(CHG, na.rm = TRUE)
    )
)

t12
#> # A tibble: 3 x 4
#> # Groups: TRTPN [3]
```

```
#>
     TRTPN AVISITN n_chg mean_chg
#>
     <dbl>
             <dbl> <int> <chr>
                       57 " -0.1 ( 2.68)"
#> 1
         0
                 24
#> 2
        54
                 24
                       26 " 0.2 ( 0.82)"
                       30 " 0.5 (1.94)"
#> 3
        81
                 24
```

5.4. Missing data imputation

In clinical trials, missing data is inevitable. In this study, there are missing values in glucose data.

```
count(ana, AVISIT)
#> # A tibble: 8 x 2
#>
     AVISIT
                              n
#>
     <fct>
                          <int>
#> 1 "
                 Week 2"
                             229
#> 2 "
                 Week 4"
                             211
#> 3 "
                 Week 6"
                            197
#> 4 "
                 Week 8"
                            187
#> 5 "
                Week 12"
                             167
#> 6 "
                Week 16"
                             147
#> 7 "
                Week 20"
                             126
#> 8 "
                Week 24"
                             113
```

For simplicity and illustration purpose, we use the last observation carried forward (LOCF) approach to handle missing data. LOCF approach is a single imputation approach that is **not recommended** in real application. Interested readers can find more discussion on missing data approaches in the book: The Prevention and Treatment of Missing Data in Clinical Trials.

```
ana_locf <- ana %>%
  group_by(USUBJID) %>%
  mutate(locf = AVISITN == max(AVISITN)) %>%
  filter(locf)
```

5.5. ANCOVA model

The imputed data is analyzed using the ANCOVA model with treatment and baseline glucose as covariates.

```
fit <- lm(CHG ~ BASE + TRTP, data = ana_locf)</pre>
summary(fit)
#>
#> Call:
#> lm(formula = CHG ~ BASE + TRTP, data = ana_locf)
#> Residuals:
               1Q Median
      Min
                               3Q
                                      Max
#> -6.9907 -0.7195 -0.2367 0.2422 7.0754
#>
#> Coefficients:
                           Estimate Std. Error t value Pr(>|t|)
#>
                                       0.39392
                                                7.637 6.23e-13 ***
#> (Intercept)
                            3.00836
#> BASE
                           -0.53483
                                       0.06267 -8.535 2.06e-15 ***
#> TRTPXanomeline Low Dose -0.17367
                                       0.24421 - 0.711
                                                          0.478
#> TRTPXanomeline High Dose 0.32983
                                       0.24846
                                                1.327
                                                          0.186
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#>
#> Residual standard error: 1.527 on 226 degrees of freedom
   (2 observations deleted due to missingness)
```

```
#> Multiple R-squared: 0.2567, Adjusted R-squared: 0.2468
#> F-statistic: 26.01 on 3 and 226 DF, p-value: 1.714e-14
```

The emmeans R package is used to obtain within and between group least square (LS) mean

```
fit_within <- emmeans(fit, "TRTP")</pre>
fit_within
#> TRTP
                                    SE df lower.CL upper.CL
#> Placebo
                                             -0.272
                          0.0676 0.172 226
#> Xanomeline Low Dose -0.1060 0.173 226
                                             -0.447
                                                       0.235
#> Xanomeline High Dose 0.3975 0.179 226
                                            0.045
                                                       0.750
#>
#> Confidence level used: 0.95
t13 <- fit_within %>%
  as_tibble() %>%
  mutate(ls = fmt_ci(emmean, lower.CL, upper.CL)) %>%
  select(TRTP, ls)
t13
#> # A tibble: 3 x 2
     TRTP
#>
                          ls
#>
     <fct>
                          <chr>
#> 1 Placebo
                          " 0.07 (-0.27, 0.41)"
#> 2 Xanomeline Low Dose "-0.11 (-0.45, 0.23)"
#> 3 Xanomeline High Dose " 0.40 ( 0.05, 0.75)"
fit_between <- pairs(fit_within, reverse = TRUE)</pre>
fit_between
#> contrast
                                                           SE df t.ratio p.value
                                               estimate
#> Xanomeline Low Dose - Placebo
                                                 -0.174 0.244 226 -0.711 0.7571
#> Xanomeline High Dose - Placebo
                                                  0.330 0.248 226
                                                                    1.327 0.3814
```

```
#>
   Xanomeline High Dose - Xanomeline Low Dose 0.504 0.249 226
                                                                    2.024
#>
#> P value adjustment: tukey method for comparing a family of 3 estimates
t2 <- fit_between %>%
  as_tibble() %>%
 mutate(
   ls = fmt_ci(
      estimate,
      estimate - 1.96 * SE,
      estimate + 1.96 * SE
    ),
   p = fmt_pval(p.value)
  ) %>%
  filter(stringr::str_detect(contrast, "- Placebo")) %>%
  select(contrast, ls, p)
t2
#> # A tibble: 2 x 3
     contrast
                                    ls
     <chr>
                                    <chr>
                                                          <chr>
#> 1 Xanomeline Low Dose - Placebo "-0.17 (-0.65, 0.30)" " 0.757"
#> 2 Xanomeline High Dose - Placebo " 0.33 (-0.16, 0.82)" " 0.381"
```

5.6. Reporting

t11, t12 and t13 are combined to get the first part of the report table

```
t1 <- cbind(
   t11 %>% ungroup() %>% select(TRTP, ends_with("0"), ends_with("24")),
```

```
t12 %>% ungroup() %>% select(ends_with("chg")),
 t13 %>% ungroup() %>% select(ls)
)
t1
#>
                     TRTP n_0
                                  mean_sd_0 n_24
                                                   mean_sd_24 n_chg
                                                                          mean_chg
#> 1
                  Placebo
                           79
                                5.7 (2.23)
                                                   5.7 (1.83)
                                                                  57 -0.1 (2.68)
                                              57
#> 2 Xanomeline Low Dose
                           79
                                5.4 (0.95)
                                              26
                                                   5.7 (1.26)
                                                                  26
                                                                       0.2 (0.82)
                                5.4 (1.37)
                                                   6.0 (1.92)
                                                                       0.5 (1.94)
#> 3 Xanomeline High Dose
                           74
                                              30
                                                                  30
#>
#> 1 0.07 (-0.27, 0.41)
#> 2 -0.11 (-0.45, 0.23)
#> 3 0.40 (0.05, 0.75)
```

Then r2rtf is used to prepare the table format for t1. We also highlight how to handle special characters in this example.

Special characters ^ and _ are used to define superscript and subscript of text. And {} is to define the part that will be impacted. For example, {^a} provides a superscript a for footnote notation. r2rtf also supports most LaTeX characters. Examples can be found on the r2rtf get started page. The text_convert argument in r2rtf_*() functions controls whether to convert special characters.

```
t1_rtf <- t1 %>%
  data.frame() %>%
  rtf_title(c(
    "ANCOVA of Change from Baseline Glucose (mmol/L) at Week 24",
    "LOCF",
    "Efficacy Analysis Population"
)) %>%
  rtf_colheader("| Baseline | Week 24 | Change from Baseline",
    col_rel_width = c(2.5, 2, 2, 4)
) %>%
```

```
rtf_colheader(
    paste(
      "Treatment |",
      paste0(rep("N | Mean (SD) | ", 3), collapse = ""),
      "LS Mean (95% CI){^a}"
    ),
    col_rel_width = c(2.5, rep(c(0.5, 1.5), 3), 2)
  ) %>%
  rtf_body(
    text_justification = c("l", rep("c", 7)),
    col_rel_width = c(2.5, rep(c(0.5, 1.5), 3), 2)
  ) %>%
 rtf_footnote(c(
    "{^a}Based on an ANCOVA model after adjusting baseline value. LOCF app
    "ANCOVA = Analysis of Covariance, LOCF = Last Observation Carried Forw
    "CI = Confidence Interval, LS = Least Squares, SD = Standard Deviation
  ))
t1_rtf %>%
  rtf_encode() %>%
  write_rtf("tlf/tlf_eff1.rtf")
```

We also use r2rtf to prepare the table format for t2

```
t2_rtf <- t2 %>%
  data.frame() %>%
  rtf_colheader("Pairwise Comparison | Difference in LS Mean (95% CI){^a}
  col_rel_width = c(4.5, 4, 2)
) %>%
  rtf_body(
  text_justification = c("l", "c", "c"),
  col_rel_width = c(4.5, 4, 2)
```

ANCOVA of Change from Baseline Glucose (mmol/L) at Week 24 LOCF Efficacy Analysis Population

	Baseline			Week 24		Change from Baseline			
Treatment	N	Mean (SD)	N	Mean (SD)	N	Mean (SD)	LS Mean (95% CI) ^a		
Placebo	79	5.7 (2.23)	57	5.7 (1.83)	57	-0.1 (2.68)	0.07 (-0.27, 0.41)		
Xanomeline Low Dose	79	5.4 (0.95)	26	5.7 (1.26)	26	0.2 (0.82)	-0.11 (-0.45, 0.23)		
Xanomeline High Dose	74	5.4 (1.37)	30	6.0 (1.92)	30	0.5 (1.94)	0.40 (0.05, 0.75)		

**Passed on an ANCOVA model after adjusting baseline value. LOCF approach is used to impute missing values. ANCOVA = Analysis of Covariance, LOCF = Last Observation Carried Forward CI = Confidence Interval, LS = Least Squares, SD = Standard Deviation

```
t2_rtf %>%
  rtf_encode() %>%
  write_rtf("tlf/tlf_eff2.rtf")
```

Finally, we combine the two parts to get the final table using r2rtf. This is achieved by providing a list of t1_rtf and t2_rtf as input for rtf_encode.

```
list(t1_rtf, t2_rtf) %>%
  rtf_encode() %>%
  write_rtf("tlf/tlf_eff.rtf")
```

In conclusion, the procedure to generate the above efficacy results table is summarized as follows.

- Step 1: Read the data (i.e., adsl and adlb) into R.
- Step 2: Define the analysis dataset. In this example, we define two analysis datasets. The first dataset is the efficacy population (gluc). The second dataset is the collection of all records post baseline and on or before week 24 (ana).
- Step 3: Impute the missing values. In this example, we name the ana dataset after imputation as ana_locf.
- Step 4: Calculate the mean and standard derivation of efficacy endpoint (i.e., gluc), and then format it into an RTF table.
- Step 5: Calculate the pairwise comparison by ANCOVA model, and then format it into an RTF table.
- Step 6: Combine the outputs from steps 4 and 5 by rows.

5.6. Reporting

Pairwise Comparison	Difference in LS Mean (95% CI) ^a	p-Value
Xanomeline Low Dose - Placebo	-0.17 (-0.65, 0.30)	0.757
Xanomeline High Dose - Placebo	0.33 (-0.16, 0.82)	0.381

ANCOVA of Change from Baseline Glucose (mmol/L) at Week 24 $$\operatorname{LOCF}$$ Efficacy Analysis Population

		Baseline	Week 24			Change from Baseline		
Treatment	N	Mean (SD)	N	Mean (SD)	N	Mean (SD)	LS Mean (95% CI) ^a	
Placebo	79	5.7 (2.23)	57	5.7 (1.83)	57	-0.1 (2.68)	0.07 (-0.27, 0.41)	
Xanomeline Low Dose	79	5.4 (0.95)	26	5.7 (1.26)	26	0.2 (0.82)	-0.11 (-0.45, 0.23)	
Xanomeline High Dose	74	5.4 (1.37)	30	6.0 (1.92)	30	0.5 (1.94)	0.40 (0.05, 0.75)	
Pairwise Comparison			Difference in LS Mean (95% CI) ^a				p-Value	
Xanomeline Low Dose - Placebo			-0.17 (-0.65, 0.30)				0.757	
Xanomeline High Dose - Placebo			0.33 (-0.16, 0.82)			0.381		
*Based on an ANCOVA mo	*Based on an ANCOVA model after adjusting baseline value LOCF approach is used to impute missing values							

"Based on an ANCOVA model after adjusting baseline value. LOCF approach is u: ANCOVA = Analysis of Covariance, LOCF = Last Observation Carried Forward CI = Confidence Interval, LS = Least Squares, SD = Standard Deviation

6. Efficacy figure

Following the ICH E3 guidance, primary and secondary efficacy endpoints need to be summarized in Section 11.4, Efficacy Results and Tabulations of Individual Participant.

```
library(haven) # Read SAS data
library(dplyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
library(survival) # Fit survival model
```

In this chapter, we illustrate how to create a simplified Kaplan-Meier plot in a study. For the survival analysis in efficacy, time to dermatologic event (TTDE) will be analyzed.

Note

R packages such as visR and survminer can create more informative Kaplan-Meier plots. Interested readers can find examples on their websites.

6.1. Analysis dataset

To prepare the analysis, the adtte dataset is required.

```
adtte <- read_sas("data-adam/adtte.sas7bdat")</pre>
```

6. Efficacy figure

First, to prepare the analysis ready data, filter all records for the efficacy endpoint of time to event of interest (TTDE) using PARAMCD (or PARAM, PRAMN), then select the survival analysis related variables:

- TRTP: treatment arm (using corresponding numeric code TRTAN to re-order the levels, "Placebo" will be the reference level)
- AVAL: time-to-event analysis value
- CNSR: event (censoring) status

```
adtte_ttde <- adtte %>%
  filter(PARAMCD == "TTDE") %>%
  select(TRTP, TRTAN, AVAL, CNSR) %>%
  mutate(
    TRTP = forcats::fct_reorder(TRTP, TRTAN), # Recorder levels
    AVAL_m = AVAL / 30.4367 # Convert Day to Month
)
```

6.2. Create Kaplan-Meier curve

The survival package is used to obtain the K-M estimate.

```
# Fit survival model, convert the time value from Days to Month
fit <- survfit(Surv(AVAL_m, 1 - CNSR) ~ TRTP, data = adtte_ttde)</pre>
```

We save the simplified K-M plot into a .png file using code below.

```
# Save as a PNG file
png(
  file = "tlf/fig_km.png",
  width = 3000,
  height = 2000,
  res = 300
```

```
plot(
   fit,
   xlab = "Time in Months",
   ylab = "Survival probability",
   mark.time = TRUE,
   lwd = 2,
   col = c(2, 3, 4),
   lty = c(1, 2, 3)
)

dev.off()
```

Now, we can use the r2rtf package to create a formatted RTF figure. More details can be found on the r2rtf website.

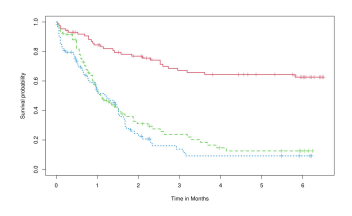
```
# Create RTF figure
rtf_read_figure("tlf/fig_km.png") %>% # Read the PNG file from the file path
  rtf_title(
    "Kaplan-Meier Plot for Time to First Dermatologic Event by Treatment Group",
    "All Participants"
) %>% # Add title or subtitle
  rtf_footnote("footnote") %>% # Add footnote
  rtf_source("[datasource: adam-adtte]") %>% # Add data source
  rtf_figure(fig_width = 6, fig_height = 4) %>% # Set proportional figure size to the ori
  rtf_encode(doc_type = "figure") %>% # Encode figure as rtf
  write_rtf(file = "tlf/tlf_km.rtf")
```

In conclusion, the steps to create a K-M plot are as follows.

- Step 1: Read the data adtte into R.
- Step 2: Define the analysis-ready dataset. In this example, we define the analysis dataset for the TTDE endpoint adtte_ttde.

6. Efficacy figure

Kaplan-Meier Plot for Time to First Dermatologic Event by Treatment Group All Participants



footnote

[datasource: adam-adtte]

6.2. Create Kaplan-Meier curve

- Step 3: Save figures into png files based on required analysis specification.
- Step 4: Create RTF output using the r2rtf package.

7. AE summary

Following ICH E3 guidance, we summarize number of participants that were included in each safety analysis in Section 12.2, Adverse Events (AEs).

```
library(haven) # Read SAS data
library(dplyr) # Manipulate data
library(tidyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
```

In this chapter, we illustrate how to summarize AEs information for a study.

The data used to summarize AE information is in adsl and adae datasets.

```
adsl <- read_sas("data-adam/adsl.sas7bdat")
adae <- read_sas("data-adam/adae.sas7bdat")</pre>
```

We first summarize participants in population by treatment arm.

```
pop <- adsl %>%
  filter(SAFFL == "Y") %>%
  rename(TRTAN = TRT01AN) %>%
  count(TRTAN, name = "tot")
pop
```

7. AE summary

Analysis of Adverse Event Summary (Safety Analysis Population)

	F	Placebo	Xanome	line Low Dose		neline High Dose
	n	(%)	n	(%)	n	(%)
Participants in population	86		84		84	
With one or more adverse events	69	(80.2)	77	(91.7)	79	(94.0)
With drug-related adverse events	44	(51.2)	73	(86.9)	70	(83.3)
With serious adverse events	0	(0.0)	1	(1.2)	2	(2.4)
With serious drug-related adverse events	0	(0.0)	1	(1.2)	1	(1.2)
Who died	2	(2.3)	1	(1.2)	0	(0.0)
Every subject is counted a single time for ea	ch applical	ble row and col	umn.			

```
#> # A tibble: 3 x 2
#> TRTAN tot
#> <dbl> <int>
#> 1 0 86
#> 2 54 84
#> 3 81 84
```

We transform the data to simplify the analysis of each required AE criteria of interest.

- With one or more adverse events
- With drug-related adverse events
- With serious adverse events
- With serious drug-related adverse events
- Who died

```
tidy_ae <- adae %>%
  mutate(
    all = SAFFL == "Y",
    drug = AEREL %in% c("POSSIBLE", "PROBABLE"),
    ser = AESER == "Y",
    drug_ser = drug & ser,
    die = AEOUT == "FATAL"
  ) %>%
  select(USUBJID, TRTAN, all, drug, ser, drug_ser, die) %>%
  pivot_longer(cols = c(all, drug, ser, drug_ser, die))
tidy_ae %>% head(4)
#> # A tibble: 4 x 4
    USUBJID
                 TRTAN name
                                value
                 <dbl> <chr>
#>
     <chr>
                                <lgl>
#> 1 01-701-1015
                     0 all
                                TRUE
#> 2 01-701-1015
                     0 drug
                                TRUE
```

7. AE summary

We summarize the number and percentage of participants who meet each AE criteria.

```
fmt_num <- function(x, digits, width = digits + 4) {</pre>
  formatC(
   х,
    digits = digits,
   format = "f",
   width = width
  )
}
ana <- tidy_ae %>%
 filter(value == TRUE) %>%
  group_by(TRTAN, name) %>%
  summarise(n = n_distinct(USUBJID)) %>%
  left_join(pop, by = "TRTAN") %>%
 mutate(
   pct = fmt_num(n / tot * 100, digits = 1),
   n = fmt_num(n, digits = 0),
   pct = paste0("(", pct, ")")
  )
ana %>% head(4)
#> # A tibble: 4 x 5
#> # Groups:
              TRTAN [2]
    TRTAN name n
                         tot pct
#>
    <dbl> <chr> <int> <chr>
       0 all " 69"
#> 1
                          86 (80.2)
```

```
#> 2  0 die " 2" 86 ( 2.3)
#> 3  0 drug " 44" 86 ( 51.2)
#> 4  54 all " 77" 84 ( 91.7)
```

We prepare reporting-ready dataset for each AE group.

```
t_ae <- ana %>%
 pivot_wider(
   id_cols = "name",
   names_from = TRTAN,
   values_from = c(n, pct),
   values_fill = list(
     n = "0"
     pct = "( 0.0)"
    )
  )
t_ae <- t_ae %>%
  mutate(name = factor(
    name,
    c("all", "drug", "ser", "drug_ser", "die"),
      "With one or more adverse events",
      "With drug-related adverse events",
      "With serious adverse events",
      "With serious drug-related adverse events",
      "Who died"
    )
  )) %>%
  arrange(name)
```

We prepare reporting-ready dataset for the analysis population.

7. AE summary

```
t_pop <- pop %>%
 mutate(
   name = "Participants in population",
   tot = fmt_num(tot, digits = 0)
  ) %>%
 pivot_wider(
   id_cols = name,
   names_from = TRTAN,
   names_prefix = "n_",
   values_from = tot
  )
t_pop
#> # A tibble: 1 x 4
                              #> name
    <chr>
                              <chr> <chr> <chr>
#>
#> 1 Participants in population " 86" " 84" " 84"
```

The final report data is saved in tbl_ae_summary.

```
tbl_ae_summary <- bind_rows(t_pop, t_ae) %>%
  select(name, ends_with("_0"), ends_with("_54"), ends_with("_81"))
tbl_ae_summary
#> # A tibble: 6 x 7
   name
                                             n_0 pct_0 n_54 pct_54 n_8
                                              <chr> <chr> <chr> <chr> <chr> <chr>
#> <chr>
#> 1 Participants in population
                                              " 8~ <NA> " 8~ <NA>
#> 2 With one or more adverse events
                                             " 6~ ( 80~ " 7~ ( 91.~ "
                                             " 4~ ( 51~ " 7~ ( 86.~ "
#> 3 With drug-related adverse events
                                             " ~ ( 0~ " ~ ( 1.~ "
#> 4 With serious adverse events
\#> 5 With serious drug-related adverse events " \ \ \sim ( 0 \sim " \ \ \sim ( 1. \sim "
```

```
#> 6 Who died " ~ ( 2~ " ~ ( 1.~ " ~ ( 0.~
```

We define the format of the output using code below:

```
tbl_ae_summary %>%
  rtf_title(
    "Analysis of Adverse Event Summary",
    "(Safety Analysis Population)"
  ) %>%
  rtf_colheader(" | Placebo | Xanomeline Low Dose| Xanomeline High Dose",
    col_rel_width = c(3.5, rep(2, 3))
  ) %>%
  rtf colheader(" | n | (%) | n | (%) | n | (%)",
    col_rel_width = c(3.5, rep(c(0.7, 1.3), 3)),
    border_top = c("", rep("single", 6)),
    border_left = c("single", rep(c("single", ""), 3))
  ) %>%
  rtf_body(
    col_rel_width = c(3.5, rep(c(0.7, 1.3), 3)),
    text_justification = c("l", rep("c", 6)),
    border_left = c("single", rep(c("single", ""), 3))
  ) %>%
  rtf_footnote("Every subject is counted a single time for each applicable row and column
  rtf_encode() %>%
  write_rtf("tlf/tlf_ae_summary.rtf")
```

The procedure to generate an AE summary table can be summarized as follows:

- Step 1: Read data (i.e., adae and adsl) into R.
- Step 2: Summarize participants in population by treatment arm, and name the dataset as t_pop.
- Step 3: Summarize participants in population by required AE criteria of interest, and name the dataset as t_ae.

7. AE summary

Analysis of Adverse Event Summary (Safety Analysis Population)

	F	Placebo	Xanome	line Low Dose		neline High Dose
	n	(%)	n	(%)	n	(%)
Participants in population	86		84		84	
With one or more adverse events	69	(80.2)	77	(91.7)	79	(94.0)
With drug-related adverse events	44	(51.2)	73	(86.9)	70	(83.3)
With serious adverse events	0	(0.0)	1	(1.2)	2	(2.4)
With serious drug-related adverse events	0	(0.0)	1	(1.2)	1	(1.2)
Who died	2	(2.3)	1	(1.2)	0	(0.0)
Every subject is counted a single time for each applicable row and column.						

• Step 4: Row-wise combine t_pop and t_ae and format it by using r2rtf.

8. Specific AE

Following ICH E3 guidance, we need to summarize number of participants for each specific AE in Section 12.2, Adverse Events (AEs).

```
library(haven) # Read SAS data
library(dplyr) # Manipulate data
library(tidyr) # Manipulate data
library(r2rtf) # Reporting in RTF format
```

In this chapter, we illustrate how to summarize simplified specific AE information for a study.

The data used to summarize AE information is in adsl and adae datasets.

```
adsl <- read_sas("data-adam/adsl.sas7bdat")
adae <- read_sas("data-adam/adae.sas7bdat")</pre>
```

For illustration purpose, we only provide counts in the simplified table. The percentage of participants for each AE can be calculated as shown in Chapter 7.

Here, we focus on the analysis script for two advanced features for a table layout.

• group content: AE can be summarized in multiple nested layers. (e.g., by system organ class (SOC, AESOC) and specific AE term (AEDECOD))

8. Specific AE

Analysis of Participants With Specific Adverse Events (Safety Analysis Population)

		Xanomeline Low	Xanomeline High
	Placebo	Dose	Dose
	n	n	n
Participants in population	86	84	84
Cardiac Disorders	13	13	18
Atrial Fibrillation	1	1	3
Atrial Flutter	0	1	1
Atrial Hypertrophy	1	0	0
Atrioventricular Block First Degree	1	1	0
Atrioventricular Block Second Degree	2	0	3
Bradycardia	1	0	0
Bundle Branch Block Left	1	0	0
Bundle Branch Block Right	1	1	0
Cardiac Disorder	0	0	1
Cardiac Failure Congestive	1	0	0
Myocardial Infarction	4	2	4
Palpitations	0	2	0
Sinus Arrhythmia	1	0	0
Sinus Bradycardia	2	7	8
Supraventricular Extrasystoles	1	1	1
Supraventricular Tachycardia	0	1	0
Tachycardia	1	0	0
Ventricular Extrasystoles	0	2	1
Ventricular Hypertrophy	1	0	0
Wolff-Parkinson-White Syndrome	0	1	0
Congenital, Familial and Genetic Disorders	0	1	2
Ventricular Septal Defect	0	1	2
Ear and Labyrinth Disorders	1	2	1
Cerumen Impaction	0	1	0
Ear Pain	1	0	0
Tinnitus	0	1	0
Vertigo	0	1	1
Eye Disorders	4	2	1
Conjunctival Haemorrhage	0	1	0
Conjunctivitis	2	0	0

• pagenization: there are many AE terms that can not be covered in one page. Column headers and SOC information need to be repeated on every page.

In the code below, we count the number of participants in each AE term by SOC and treatment arm, and we create a new variable order and set it as 0. The variable order can help with the data manipulation later.

```
fmt_num <- function(x, digits, width = digits + 4) {</pre>
  formatC(
    x,
    digits = digits,
    format = "f",
    width = width
}
ana <- adae %>%
  mutate(
    AESOC = tools::toTitleCase(tolower(AESOC)),
    AEDECOD = tools::toTitleCase(tolower(AEDECOD))
  )
t1 <- ana %>%
  group_by(TRTAN, AESOC) %>%
  summarise(n = fmt_num(n_distinct(USUBJID), digits = 0)) %>%
  mutate(AEDECOD = AESOC, order = 0)
t1 %>% head(4)
#> # A tibble: 4 x 5
#> # Groups: TRTAN [1]
     TRTAN AESOC
#>
                                               AEDECOD
                                                                            order
     <dbl> <chr>
                                        <chr> <chr>
#>
                                                                             <dbl>
```

8. Specific AE

In the code below, we count the number of subjects in each AE term by SOC, AE term, and treatment arm. Here we also create a new variable order and set it as 1.

```
t2 <- ana %>%
  group_by(TRTAN, AESOC, AEDECOD) %>%
  summarise(n = fmt_num(n_distinct(USUBJID), digits = 0)) %>%
  mutate(order = 1)
t2 %>% head(4)
#> # A tibble: 4 x 5
#> # Groups:
               TRTAN, AESOC [1]
     TRTAN AESOC
                             AEDECOD
                                                                   n
     <dbl> <chr>
                             <chr>
                                                                    <chr>
         O Cardiac Disorders Atrial Fibrillation
                                                                        1"
         O Cardiac Disorders Atrial Hypertrophy
                                                                        1"
#> 2
                                                                        1"
         O Cardiac Disorders Atrioventricular Block First Degree
#> 3
                                                                        2"
#> 4
         O Cardiac Disorders Atrioventricular Block Second Degree "
```

We prepare reporting data for AE information using code below:

```
t_ae <- bind_rows(t1, t2) %>%
  pivot_wider(
   id_cols = c(AESOC, order, AEDECOD),
   names_from = TRTAN,
   names_prefix = "n_",
   values_from = n,
   values_fill = fmt_num(0, digits = 0)
```

```
) %>%
 arrange(AESOC, order, AEDECOD) %>%
 select(AESOC, AEDECOD, starts_with("n"))
t_ae %>% head(4)
#> # A tibble: 4 x 5
    AESOC
                      AEDECOD
                                                n_54 n_81
                                          n_0
     <chr>
                      <chr>
#>
                                          <chr> <chr> <chr>
                                                    13" "
                                          " 13" "
#> 1 Cardiac Disorders Cardiac Disorders
#> 2 Cardiac Disorders Atrial Fibrillation "
                                            1" "
                                                     1" "
                                              0" "
                                                     1" "
                                                            1"
#> 3 Cardiac Disorders Atrial Flutter
#> 4 Cardiac Disorders Atrial Hypertrophy
                                              1" "
                                                     0" "
                                                            0"
```

We prepare reporting data for analysis population using code below:

```
count_by <- function(data, # Input data set</pre>
                     grp, # Group variable
                     var, # Analysis variable
                     var_label = var, # Analysis variable label
                     id = "USUBJID") { # Subject ID variable
  data <- data %>% rename(grp = !!grp, var = !!var, id = !!id)
  left_join(
    count(data, grp, var),
    count(data, grp, name = "tot"),
    by = "grp",
  ) %>%
    mutate(
      pct = fmt_num(100 * n / tot, digits = 1),
     n = fmt_num(n, digits = 0),
      npct = paste0(n, " (", pct, ")")
    ) %>%
```

```
pivot_wider(
     id_cols = var,
     names_from = grp,
     values_from = c(n, pct, npct),
     values_fill = list(n = "0", pct = fmt_num(0, digits = 0))
   ) %>%
   mutate(var_label = var_label)
}
t_pop <- adsl %>%
  filter(SAFFL == "Y") %>%
  count_by("TRT01AN", "SAFFL",
   var_label = "Participants in population"
  ) %>%
 mutate(
   AESOC = "pop",
   AEDECOD = var_label
  ) %>%
  select(AESOC, AEDECOD, starts_with("n_"))
t_pop
#> # A tibble: 1 x 5
#> AESOC AEDECOD
                                    <chr> <chr>
#>
                                    <chr> <chr> <chr>
#> 1 pop Participants in population " 86" " 84" " 84"
```

The final report data is saved in tbl_ae_spec. We also add a blank row between population and AE information in the reporting table.

```
tbl_ae_spec <- bind_rows(
  t_pop,
  data.frame(AESOC = "pop"),</pre>
```

```
t_ae
) %>%
 mutate(AEDECOD = ifelse(AEDECOD == AESOC,
    AEDECOD, pasteO(" ", AEDECOD)
 ))
tbl_ae_spec %>% head(4)
#> # A tibble: 4 x 5
    AESOC
                      AEDECOD
                                                     n_0
                                                            n_54
                                                                   n_81
#>
     <chr>
                      <chr>
                                                     <chr> <chr> <chr>
#> 1 pop
                       " Participants in population" " 86" "
                                                               84" "
#> 2 pop
                       <NA>
                                                      <NA>
                                                             <NA>
                                                                    <NA>
                                                               13" "
#> 3 Cardiac Disorders "Cardiac Disorders"
                                                        13" "
                                                                      18"
#> 4 Cardiac Disorders " Atrial Fibrillation"
                                                        1" "
                                                               1" "
```

We define the format of the output as below:

To obtain the nested layout, we use the page_by argument in the rtf_body function. By defining page_by="AESOC", r2rtf recognizes the variable as a group indicator.

After setting pageby_row = "first_row", the first row is displayed as group header. If a group of information is broken into multiple pages, the group header row is repeated on each page by default.

We can also customize the text format by providing a matrix that has the same dimension as the input dataset (i.e., tbl_ae_spec). In the code below, we illustrate how to display **bold** text for group headers to highlight the nested structure of the table layout.

```
n_row <- nrow(tbl_ae_spec)
n_col <- ncol(tbl_ae_spec)
id <- tbl_ae_spec$AESOC == tbl_ae_spec$AEDECOD
id <- ifelse(is.na(id), FALSE, id)</pre>
```

```
text_format <- ifelse(id, "b", "")</pre>
tbl_ae_spec %>%
  rtf_title(
    "Analysis of Participants With Specific Adverse Events",
    "(Safety Analysis Population)"
  ) %>%
  rtf_colheader(" | Placebo | Xanomeline Low Dose | Xanomeline High Dose",
    col_rel_width = c(3, rep(1, 3))
  ) %>%
  rtf_colheader(" | n | n | n ",
    border_top = "",
    border_bottom = "single",
    col_rel_width = c(3, rep(1, 3))
  ) %>%
  rtf_body(
    col_rel_width = c(1, 3, rep(1, 3)),
    text_justification = c("l", "l", rep("c", 3)),
    text_format = matrix(text_format, nrow = n_row, ncol = n_col),
    page_by = "AESOC",
    pageby_row = "first_row"
  ) %>%
  rtf_footnote("Every subject is counted a single time for each applicable
  rtf_encode() %>%
  write_rtf("tlf/tlf_spec_ae.rtf")
```

More discussion on page_by, group_by and subline_by features can be found on the r2rtf package website.

The procedure to generate a baseline characteristics table can be summarized as follows:

• Step 1: Read data (i.e., adae and adsl) into R.

Analysis of Participants With Specific Adverse Events (Safety Analysis Population)

		Xanomeline Low	Xanomeline High
	Placebo	Dose	Dose
	n	n	n
Participants in population	86	84	84
Cardiac Disorders	13	13	18
Atrial Fibrillation	1	1	3
Atrial Flutter	0	1	1
Atrial Hypertrophy	1	0	0
Atrioventricular Block First Degree	1	1	0
Atrioventricular Block Second Degree	2	0	3
Bradycardia	1	0	0
Bundle Branch Block Left	1	0	0
Bundle Branch Block Right	1	1	0
Cardiac Disorder	0	0	1
Cardiac Failure Congestive	1	0	0
Myocardial Infarction	4	2	4
Palpitations	0	2	0
Sinus Arrhythmia	1	0	0
Sinus Bradycardia	2	7	8
Supraventricular Extrasystoles	1	1	1
Supraventricular Tachycardia	0	1	0
Tachycardia	1	0	0
Ventricular Extrasystoles	0	2	1
Ventricular Hypertrophy	1	0	0
Wolff-Parkinson-White Syndrome	0	1	0
Congenital, Familial and Genetic Disorders	0	1	2
Ventricular Septal Defect	0	1	2
Ear and Labyrinth Disorders	1	2	1
Cerumen Impaction	0	1	0
Ear Pain	1	0	0
Tinnitus	0	1	0
Vertigo	0	1	1
Eye Disorders	4	2	1
Conjunctival Haemorrhage	0	1	0
Conjunctivitis	2	0	0

8. Specific AE

- Step 2: Count the number of participants by SOC and treatment arm (rows with bold text) and save into t1.
- Step 3: Count the number of participants in each AE term by SOC, AE term, and treatment arm (rows without bold text) and save into t2.
- Step 4: Bind t1 and t2 by row into t_ae.
- Step 5: Count the number of participants in each arm as t_pop.
- Step 6: Row-wise combine t_pop and t_ae into tbl_ae_spec.
- Step 7: Format the output by using r2rtf.

9. Assemble TLFs

```
library(r2rtf)
```

After TLFs are created and saved into individual files, we need to assemble them into one file in a pre-specified order.

There are two general approaches to achieving the goal.

- 1. Combine RTF source code in individual files into one large RTF file.
- 2. Leverage the Toggle Fields feature in Microsoft Word to embed RTF files using hyperlinks.

Let's illustrate the idea by using selected TLFs generated from previous chapters. Here, we assume files are saved in the tlf/ folder.

```
tlf_path <- c(
   "tlf/tbl_disp.rtf", # Disposition table
   "tlf/tlf_eff.rtf", # Efficacy table
   "tlf/tlf_km.rtf" # K-M plot
)</pre>
```

9. Assemble TLFs

Disposition of Participants

	P	Placebo		Xanomeline Low Dose		Xanomeline High Dose	
	n	(%)	n	(%)	n	(%)	
Participants in population	86		84		84		
Completed	58	67.4	25	29.8	27	32.1	
Discontinued	28	32.6	59	70.2	57	67.9	
Adverse Event	8	9.3	44	52.4	40	47.6	
Death	2	2.3	1	1.2	0	0.0	
I/E Not Met	1	1.2	0	0.0	2	2.4	
Lack of Efficacy	3	3.5	0	0.0	1	1.2	
Lost to Follow-up	1	1.2	1	1.2	0	0.0	
Physician Decision	1	1.2	0	0.0	2	2.4	
Protocol Violation	1	1.2	1	1.2	1	1.2	
Sponsor Decision	2	2.3	2	2.4	3	3.6	
Withdrew Consent	9	10.5	10	11.9	8	9.5	

9.1. Combine RTF Source Code

Note

The code below requires r2rtf version >= 1.0.0.

The r2rtf::assemble_rtf() function allows the user to combine RTF source code in individual files into one larger RTF file.



Caution

One limitation of combining RTF source code is that we are not able to specify the page orientation of each TLF in the combined document.

```
r2rtf::assemble_rtf(
  input = tlf_path,
  output = "tlf/rtf-combine.rtf"
)
```

9.2. Using Toggle Fields

Microsoft Word uses toggle fields to embed files into one Word document. The approach is a dynamic link between files by providing the absolute file path.



🕊 Tip

There is a slight learning curve on how toggle fields work in Microsoft Word. After you become familiar with the workflow, toggle fields can extend your capability to manage a large number of TLFs in RTF format.

The assemble_docx() function allows you to create a .docx file with toggle fields as below. One benefit is to control the page direction of each TLF as below.

```
r2rtf::assemble_docx(
   tlf_path,
   output = "tlf/rtf-combine-toggle.docx",
   landscape = c(FALSE, FALSE, TRUE)
)
```

After opening the generated .docx file, you will see a blank file because the file only contains fields with hyperlinks.

By using Alt + F9 to display the fields and you will see information similar to the screenshot below.

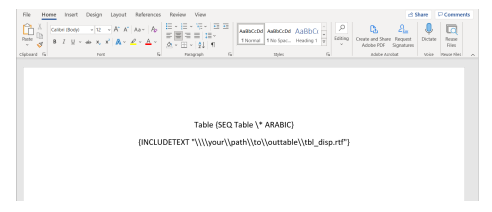


Figure 9.1.: Using Alt + F9 to display fields

? Tip

A typical error message is that system can not find the file if you only provide a relative path. Please double-check that the correct absolute file path is in the INCLUDETEXT field.

To test the toggle field, you can right-click an INCLUDETEXT filed and click Update Field.

If it works, you can see a table similar to the snapshot below by using Alt + F9. It is a shortcut to change between results and field display mode.

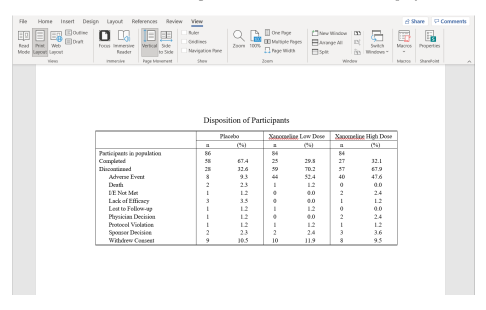


Figure 9.2.: Update fields

Now you can update all toggle fields to display all TLFs by selecting all fields (Ctrl + A), then press F9. We suggest testing one toggle field before updating all of them.

As the .docx file contain dynamic links, you can keep updating the TLFs

9. Assemble TLFs

if you need to refresh content in individual RTF files by selecting all fields (Ctrl + A), then press F9.



Tip

If you modify table content in the combined .docx file, you may get a weird table layout if you update all fields within a toggle field. To resolve the issue, please remove all * MERGEFORMAT in the filed mode using Alt + F9 before updating all toggle fields.

After the combined TLF is ready for delivery, you can also unlink toggle fields to save table contents, because the absolute path may only work for some. To unlink toggle fields, you can select all fields (Ctrl + A), then press Ctrl + Shift + F9.

Part II. Clinical trial project

10. Overview

In a late-stage clinical trial, the number of A&R deliverables can easily be in the hundreds. For an organization, it is common to have multiple ongoing clinical trials in a clinical program.

To deliver the A&R results of a clinical trial project, it is teamwork that typically requires collaborations from both statisticians and programmers. In this part, let's consider how to organize a clinical trial project as an A&R lead.

Chapter 11 will discuss how to organize source code, documents, and deliverables in an A&R clinical project. We recommend using the R package folder structure.

Chapter 12 will discuss a process or system development lifecycle to manage the A&R of a clinical project. We recommend following an agile management approach to define, develop, validate, and deliver work.

11. Project folder

A clearly defined clinical project folder structure can have many benefits to clinical development teams in an organization. Specifically, a well-defined project structure can achieve:

- Consistency: everyone works on the same folder structure.
- Reproducibility: analysis can be executed and reproduced by different team members months/years later.
- Automation: automatically check the integration of a project.
- Compliance: reduce compliance issues.

We will use the esubdemo R package to illustrate the project folder structure for the A&R project. You can clone the project using RStudio IDE with

```
git clone https://github.com/elong0527/esubdemo.git
```

For R users, you already benefit from a well-defined and consistent folder structure. That is the R package folder structure. Every R package developer is required to follow the same convention to organize their R functions before the R package can be disseminated through the Comprehensive R Archive Network (CRAN). As a user, you can easily install and use those R packages after downloading from CRAN. There are many good resources to guide developers on R package development, such as, the R Packages book by Hadley Wickham.

We recommend using the R package folder structure to organize analysis scripts for clinical trial development. Using the R package folder struc-

11. Project folder

ture to streamline data analysis work has also been proposed before (see Marwick, Boettiger, and Mullen (2018), Wu et al. (2021)).

11.1. Consistency

For consistency, a well-defined folder structure with potential templates ensures project teams organize the A&R work consistently across multiple projects. Consistent folder structure also reduces communication costs between study team members and enhances the transparency of projects.

In this book, we refer to an R package as a **project-specific R package** if the purpose of an R package is to organize analysis scripts for a clinical project.

We refer to an R package as a **standard R package** if the purpose of an R package is to share commonly used R functions to be hosted in a code repository such as CRAN.

Below is minimal sufficient folders and files for a project-specific R package based on the R package folder structure.

- *.Rproj: RStudio project file used to open RStudio project.
- DESCRIPTION: Metadata for a package including authors, license, dependencies, etc.
- R/: Project-specific R functions.
- vignettes/: Analysis scripts using R Markdown.
- man/: Manual of project-specific R functions.

A general discussion of the R package folder structure can be found in Chapter 4 of the R Packages book (Wickham (2015)).

We demonstrate the idea using the esubdemo project.

In the esubdemo project, we saved all TLF generation scripts in previous chapters into the vignettes/ folder.

Note

Under the vignettes/ folder, there are two folders: adam/ and tlf/. The adam/ folder contains ADaM datasets. The tlf/ folder contains output TLFs in RTF format. We put adam/ and tlf/ folders within the vignettes/ folder only for illustration purposes. In an actual A&R report, you may have a different location to save your input and output.

```
vignettes
data-adam
tlf
tlf-01-disposition.Rmd
tlf-02-population.Rmd
tlf-03-baseline.Rmd
tlf-04-efficacy.Rmd
tlf-05-ae-summary.Rmd
tlf-06-ae-spec.Rmd
```

While creating those analysis scripts, we also defined a few helper functions (e.g., fmt_num and count_by). Those functions are saved in the R/ folder.

```
R/
    count_by.R
    fmt.R
```

For a clinical trial project, it is also important to provide proper documentation for those help functions. We use roxygen2 package to document functions. For example, the header below defines each variable in fmt_est. More details can be found in Chapter 16 of the R Packages book.

11. Project folder

```
#' Format point estimator
#'

#' @param .mean mean of an estimator.
#' @param .sd sd of an estimator.
#' @param digits number of digits for `.mean` and `.sd`.
#'

#' @export

fmt_est <- function(.mean, .sd, digits = c(1, 2)) {
    .mean <- fmt_num(.mean, digits[1], width = digits[1] + 4)
    .sd <- fmt_num(.sd, digits[2], width = digits[2] + 3)
    pasteO(.mean, " (", .sd, ")")
}</pre>
```

The roxygen2 documentation will be converted into standard R documentation format, and saved as .Rd files in the man/ folder. This step is automatically handled by devtools::document().

```
man
count_by.Rd
fmt_ci.Rd
fmt_est.Rd
fmt_num.Rd
fmt_pval.Rd
```

The man/ folder is used to save documentation automatically generated by roxygen2. A typical workflow is to add roxygen2 documentation before each function in the R/ folder. Then devtools::document() is used to generate all the documentation files in the man/ folder. More details can be found in Chapter 16 of the R Packages book.

11.2. Reproducibility

Reproducibility of analysis is one of the most important aspects of regulatory deliverables. To ensure a successful reproduction, we need a controlled R environment, including the control of the R version and the R package versions. By using the R package folder structure and proper tools (e.g., renv, packrat), we illustrate how to achieve reproducibility for R and R package versions.



Tip

This is the same level of reproducibility in most SAS environments: https://support.sas.com/en/technical-support/servicespolicies.html#altos

11.2.1. R version

First, we introduce the control of the R version. In the esubdemo project, a reproducible environment is created when you open the esubdemo. Rproj from RStudio IDE. When we open the esubdemo project, RStudio IDE will execute the R code in .Rprofile automatically. So we can use .Rprofile to set up a reproducible environment. More details can be found in https: //rstats.wtf/r-startup.html. After we open the esubdemo project, the code in .Rprofile will automatically check the current R version is the same as we defined in .Rprofile.

```
# Set project R version
R_version <- "4.1.1"
```

If there is an R version mismatch, an error message is displayed as below.

Error: The current R version is not the same as the current project in R4.1.1

11. Project folder



.Rprofile is only for project-specific R packages. A standard R package should not use .Rprofile.

11.2.2. R package version

Next, we introduce the control of the R package version, which is controlled in two layers. Firstly, we define a snapshot date in .Rprofile. The snapshot date allows us to freeze the source code repository.

```
# set up snapshot date
snapshot <- "2021-08-06"

# set up repository based on the snapshot date
repos <- pasteO("https://packagemanager.posit.co/cran/", snapshot)

# define repo URL for project-specific package installation
options(repos = repos)</pre>
```

We can also define the package repository to be a specific snapshot date. For example, we used Posit Public Package Manager to define the snapshot date to be 2021-08-06. The snapshot date freezes the R package repository.

In other words, all R packages installed in this R project are based on the frozen R version at the snapshot date. Here it is 2021–08–06 by using the Posit Package Manager.

The information below will be displayed after a new R session is opened.

```
Current project R package repository:
    https://packagemanager.posit.co/cran/2021-08-06
```

Note

Posit Public Package Manager hosts daily CRAN snapshots for Mondays to Fridays of the week. Posit Package Manager, when deployed internally within an organization, provides a solution to host both publicly available and internally developed R packages.

Secondly, we use renv to lock R package versions and save them in the renv.lock file. renv provides a robust and stable approach to managing R package versions for project-specific R packages. An introduction of renv can be found on its website.

source("renv/activate.R")

The R code above in the .Rprofile initiates the renv running environment. As a user, you can use renv::init(), renv::snapshot(), and renv::restore() to initialize, save and restore R packages used for the current analysis project.

In the analysis project, the renv package will

- create a renv.lock file to save the state of package versions.
- create a renv/ folder to manage R packages for a project.

Caution

The renv.lock file and renv/ folder are only for project-specific R package. A standard R package should not use renv.

In summary, the R package version is controlled in two layers.

- Define a snapshot date in inst/startup.R.
- Using renv to lock R versions within a project.

11. Project folder

If the project is initiated properly, you should be able to see similar messages to inform how we control R package versions.

* Project '~/esubdemo' loaded. [renv 0.14.0]

Once R packages have been properly installed, the system will use the R packages located in the search path defined based on the order of .libPaths(). The startup message also provided the R package search path.

Below R package path are searching in order to find installed R packages in "/home/zhanyilo/github-repo/esubdemo/renv/library/R-4.1/x86_64-pc-linux-y"/rtmp/RtmpT3ljoY/renv-system-library"

? Tip

A cloud-based R environment (e.g., Posit Workbench) can enhance the reproducibility within an organization by using the same operating system, R version, and R package versions for an A&R project. More details can be found at https://environments.rstudio.com/.

Note

A container solution like Docker (Nüst et al. 2020) could further enhance the reproducibility across an organization at the operating system level but beyond the scope of this book.

In conclusion, to achieve reproducibility for a project-specific R package, a clinical project team can work under a controlled R environment in the same R version and R package versions defined by a repository snapshot date.

11.3. Automation

By using the R package folder structure, you will benefit from many outstanding tools to simplify and streamline your workflow.

We have learned a few functions in **devtools** to generate content automatically. Here is a list of tools that can enhance the workflow.

- devtools: make package development easier.
 - A good overview can be found in Chapter 2 of the R Packages book.
 - devtools::load_all(): load all functions in R/ folder and running environment.
 - devtools::document(): automatically create documentation using roxygen2.
 - devtools::check(): automatically perform compliance check as an R package.
 - devtools::build_site(): automatically run analysis scripts in batch and create a pkgdown website.
- usethis: automates repetitive tasks that arise during project setup and development.
- testthat: streamline testing code.
 - A discussion of using the testthat for an A&R project can be found in (Ginnaram et al. (2021)).
- pkgdown: generate static HTML documentation website for an R package
 - It also allows you to run all analysis code in batch.

You may further automatically execute routines by leveraging CI/CD workflow. For example, the esubdemo project will rerun all required checks and build a pkgdown website by using Github Actions.

11. Project folder

As the consistent folder is defined, it also becomes easier to create specific tools that fit the analysis and reporting purpose. Below are a few potential tools that can be helpful:

- Create project template using RStudio project templates;
- Add additional compliance checks for analysis and reporting;
- Save log files for running in batch.

11.4. Compliance

For a regulatory deliverable, it is important to maintain compliance. With a consistent folder structure, we can define specific criteria for compliance. Some compliance criteria can be implemented within the automatically checking steps.

For an R package, there are already criteria to ensure R package integrity. More details can be found in Chapter 20 of the R Packages book.

12. Project management

12.1. Setting up for success

A clinical data analysis project is not unlike typical data analysis projects or software projects. Therefore, the conventional wisdom and tricks for managing a successful project are also applicable here. At the same time, clinical projects also have unique traits, such as high standards for planning, development, validation, and delivery under strict time constraints.

Although many factors determine if a project can execute efficiently, we believe a few aspects are critical for long-term success, especially when managing clinical data analysis projects at scale.

12.1.1. Work as a team

As a general principle, all the team members involved in a project should take basic training on project management and understand how to work as a development team. Fitzpatrick and Collins-Sussman (2012) provides some valuable tips on this topic. As always, setting a clear goal and following a system development lifecycle (SDLC) is essential.

12. Project management

12.1.2. Design clean code architecture

Having a clean architecture design for your code improves the project's robustness and flexibility for future changes. For example, we should understand how to separate business logic from other layers; know what should be created as reusable components and what should be written as one-off analysis scripts; write low coupling, high cohesion code, and so on. Martin, Grenning, and Brown (2018) offers some helpful insights on this topic.

12.1.3. Set capability boundaries

Knowing what you can do is essential. Create a core capabilities list for your team.

Sometimes, it is also critical to understand **what not to do**. For example, the hidden cost of integrating with external systems or involving other programming languages can be prohibitively high. Remember, a simple, robust solution is almost always preferable to a complex solution that requires high maintenance and constant attention.

12.1.4. Contribute to the community

Every individual is limited in some way. The collective thinking from a community could benefit a project in the long term. When designing reusable components, make a plan to share with internal communities, or even better, with the open-source community.

12.2. The SDLC

For A&R deliverables in clinical project development, a clearly defined process or system development lifecycle (SDLC) is crucial to ensure regulatory compliance.

SDLC for the A&R deliverables can be defined in four stages.

- Planning: a planning stage to define the scope of a project.
- Development: a development stage to implement target deliverables.
- Validation: a validation stage to verify target deliverables.
- Operation: an operation stage to deliver work to stakeholders.

Importantly, we should not consider SDLC as a linear process. For example, if the study team identifies a new requirement in a development or validation stage, the team should return to the planning stage to discuss and align the scope. An agile project management approach is suitable and recommended for an A&R clinical development project. The goal is to embrace an iterative approach that continuously improves target deliverables based on frequent stakeholder feedback.

There are many good tools to implement agile project management strategy, for example:

- GitHub project board
- Jira

12.3. Planning

The planning stage is important in the SDLC lifecycle as the requirements for all A&R deliverables are gathered and documented.

12. Project management

In the planning stage, a project leader should identify all the deliverables, e.g., a list of tables, listings, and figures (TLFs). For each TLFs, the team should prepare the necessary specifications:

- mock-up tables
- validation level (e.g., independent review or double programming)
- etc.

The project leader should also align work assignments with team members. The purpose is to answer the question of "who is doing what?"

Warning in ensure_len_latex(background, nrows, off, include_thead, "white", The number of provided values in background does not equal to the number of rows.

Warning in ensure_len_latex(background, nrows, off, include_thead, "white", The number of provided values in background does not equal to the number of rows.

Warning in ensure_len_latex(background, nrows, off, include_thead, "white", The number of provided values in background does not equal to the number of rows.

The project lead should also set up a project folder, as discussed in Chapter 11. The project initiation can be simplified by creating an RStudio project template.

To enable reproducibility, the project leader should also review the startup file (i.e. .Rprofile discussed in Section 11.2) and define:

- R version
- Repository of R packages with a snapshot date
- Project package library path
- etc.

12.3. Planning

			rement/Specification
Program Name	Program Validation Category	Who	Status
count_by	3	Alice	С
fmt_ci	3	Alice	С
fmt_est	3	Alice	С
fmt_num	3	Alice	С
fmt_pval	3	Alice	С
tlf-01-disposition.Rmd	3	Bob	С
tlf-02-population.Rmd	3	Carol	С
tlf-03-baseline.Rmd	3	Dave	С
tlf-04-efficacy.Rmd	3	Alice	С
tlf-05-ae-summary.Rmd	3	Bob	С
tlf-06-ae-spec.Rmd	3	Carol	С



Caution

After project initiation, modifying .Rprofile will be a risk for reproducibility and should be handled carefully if necessary.

12.4. Development

After a project is initiated, the study team starts to develop TLFs based on pre-defined mock-up tables assigned to each team member.

The analysis code and relevant description can be saved in R Markdown files in the vignettes/ folder.

The use of R Markdown allows developers to assemble narrative text, code, and its comments in one place to simplify documentation. It would be helpful to create a template and define a name convention for all TLFs deliverables. For example, we can use the tlf_ prefix in the filename to indicate that the R Markdown file is for delivering TLFs. Multiple TLFs with similar designs can be included in one R Markdown file.

For example, in the esubdemo project, we have six R Markdown files to create TLFs.

If there are any project-specific R functions that need to be developed, the R functions can be placed in the R/ folder as discussed in Section 11.1.

12.5. Validation

Validation is a crucial stage to ensure the deliverables are accurate and consistent. After the development stage is completed, the project team needs to validate the deliverables, including R Markdown files for TLFs

deliverables and project-specific R functions. The level of validation is determined at the define stage.

In an R package development, the validation or testing is completed under the test/ folder. The testthat R package can be used to streamline the validation process. More details of the testthat package for R package validation can be found in Chapter 12 of the R package book.

It is recommended to have a name convention to indicate the type of validation. For example, we can use test-developer-test, test-independent-test, test-double-programming to classify the validation type.

It is recommended to follow the same organization for files in testthat folder as R/ folder and vignettes/ folder. Every single file in the R/ folder and vignettes/ folder should have a testing file saved in the tests/testthat/ folder to validate the content.

For example, in esubdemo project, we can have a list of testing files below.

tests/testthat

```
test-independent-test-tlf-01-disposition.R test-independent-test-tlf-02-population.R test-independent-test-tlf-03-baseline.R test-independent-test-tlf-04-efficacy.R test-independent-test-tlf-05-ae-summary.R test-independent-test-tlf-06-ae-spec.R test-independent-test-fmt.R
```

To validate the content of a table, we can save the last datasets ready for table generation as a .Rdata file. A validator can reproduce the TLF and compare it with the original result saved in the .Rdata file. A test is passed when the results match. Customers can directly review the formatting of the TLFs by comparing them with the mock-up.

12. Project management

To validate a figure, we can use the snapshot testing strategy.

After the validator completes the testing of project-specific functions and R Markdown files, the process to execute and report testing results is the same for a standard R package. The devtools::test() function automatically executes all testing cases and summarizes the testing results in a report.

After completing the validation, the validator updates the status in a validation tracker. The project lead reviews the tracking sheet to make sure all required activities in the SDLC are completed, and the tracking sheet has been filled correctly. The deliverables are ready for customer review after all the validation steps are completed. Any changes to the output requested by customers are documented.

12.6. Operation

After completion of development and required validation of all A&R deliverables, the project lead runs compliance checks for a project-specific R package similar to other R packages. devtools::check() is a convenient way to run compliance checks or R CMD check. R CMD check is an automated check of the contents in the R package for frequently encountered issues before submission to CRAN. Since the project-specific R package is not submitted to CRAN, some checks can be customized and skipped in devtools::check(). The project lead should work with the study team to ensure all reported errors, warnings, and notes by devtools::check() are fixed.

The project lead can also use the R package pkgdown to build a complete website for a project-specific R package. The pkgdown website is a convenient way to run all analyses in batch and integrate outputs in a website, which comprehensively covers project-specific R functions, TLF generation programs, outputs and validation tracking information, etc. For

example, in the esubdemo project, we created the pkgdown website at https://elong0527.github.io/esubdemo/.

Many of the tasks in SDLC can be completed automatically. An organization can leverage CI/CD workflow to automatically enable those tasks, such as running testing cases and creating a pkgdown website. For example, in the <code>esubdemo</code> project, we set up GitHub Actions for it. This can be done by using <code>usethis::use_github_action()</code>.

Part III. eCTD submission

13. Overview

The electronic Common Technical Document (eCTD) is a standard format for the electronic submission of applications, amendments, supplements, and reports from the applicant to the regulator. The eCTD offers a solution to submit documents stored in a standard directory structure, with file integrity validation mechanisms in place.

To submit TLFs created by R to regulatory agencies, we should follow the spirit of the existing eCTD submission guidelines to prepare the deliverables, and provide the essential details in the relevant documents for review.

The goal of the following two chapters is to provide guidance to follow Section 4.1.2.10 of the FDA Study Data Technical Conformance Guide:

Sponsors should provide the software programs used to create all ADaM datasets and generate tables and figures associated with primary and secondary efficacy analyses. Furthermore, sponsors should submit software programs used to generate additional information included in Section 14 CLINICAL STUDIES of the Prescribing Information (PI)26 if applicable. The specific software utilized should be specified in the ADRG. The main purpose of requesting the submission of these programs is to understand the process by which the variables for the respective analyses were created and to confirm the analysis algorithms. Sponsors should submit software programs in ASCII text format; however, executable file extensions should not be used.

13. Overview

Chapter 14 will focus on preparing proprietary R packages and analysis code into proper formats for submission.

Chapter 15 will discuss the recommendations to make the R code running environment reproducible for dry run tests and reviews.

14. Submission package

In this chapter, we will first give a high-level overview of what assets in the eCTD submission package we should focus on when submitting R code. Then, we will discuss how to prepare the proprietary R packages (if any), and make them be part of the submission package. In the end, we will provide reusable templates for updating Analysis Data Reviewer's Guide (ADRG) and Analysis Results Metadata (ARM) so that the reviewers receive proper instructions to reproduce the analysis results.

14.1. Prerequisites

This chapter uses pkglite (Zhao et al. 2023) to convert R source packages into text files and back.

```
install.packages("pkglite")
```

The demo project (R package) we will prepare for submission is called esubdemo, which is available on GitHub. You can download or clone it:

```
git clone https://github.com/elong0527/esubdemo.git
```

The demo submission package (not to be confused with the R package above) is ectddemo, which is also available on GitHub. You can download or clone it:

```
git clone https://github.com/elong0527/ectddemo.git
```

We assume the paths to the two folders are esubdemo/ and ectddemo/ below.

14.2. The whole game

In eCTD deliverable, the analysis datasets and source code are saved under the eCTD module 5 (clinical study reports) folder

```
ectddemo/m5/datasets/<study>/analysis/adam/
```

The files in two directories within the adam/ folder are critical for documenting analysis using R: datasets/ and programs/.

```
ectddemo/m5/datasets/ectddemo/analysis/adam/
  datasets
      adae.xpt
      ...
      adrg.pdf
      analysis-results-metadata.pdf
      define.xml
      define2-0-0.xsl
programs
      rOpkgs.txt
      tlf-01-disposition.txt
      tlf-02-population.txt
      tlf-03-baseline.txt
      tlf-04-efficacy.txt
      tlf-05-ae-summary.txt
      tlf-06-ae-spec.txt
```

The special considerations for each component are listed below.

14.2.1. datasets

Folder path: ectddemo/m5/datasets/ectddemo/analysis/adam/datasets/.

- ADaM data in .xpt format: created by SAS or R.
- define.xml: created by Pinnacle 21.
- ADRG (Analysis Data Reviewer's Guide)
 - "Macro Programs" section: provide R and R package versions with a snapshot date.
 - Appendix: provide step-by-step instructions for reviewers to reproduce the running environment and rerun analyses.
- ARM (Analysis Results Metadata): provide the links between TLFs and analysis programs in tables.

14.2.2. programs

Folder path: ectddemo/m5/datasets/ectddemo/analysis/adam/programs/.

- r0pkgs.txt: contains all internally developed proprietary R packages.
- Other .txt files: each contains R code for a specific analysis.

14.2.3. Notes

To verify if the submission package works, rerun all analyses following the instructions defined in ADRG.

A few things need to be paid attention to in order to pass compliance checks:

14. Submission package

- The file names under programs/ should be in lower case letters (with no underscores or other special characters).
- The .txt files should only contain ASCII characters. This can be verified by pkglite::verify_ascii()
- All .docx files should be converted to PDF files for formal submission.

Now you have a general idea about the relevant components of the submission package. We will prepare the proprietary R packages in the following sections.

14.3. Practical considerations for R package submissions

Before we start, there are a few aspects to figure out in order to accurately identify the R packages for submission.

14.3.1. Source location

There are a few common places to host R (source) packages:

- 1. CRAN
- 2. Public Git repository
- 3. Private Git repository (accessible externally)
- 4. Private Git repository (inaccessible externally)

For R packages hosted on CRAN or a public Git repository, you probably do not need to submit them as part of the submission package, as the reviewers can install them directly by following the instructions in ADRG.

For R packages hosted in private repositories, to avoid any complications in infrastructure, authentication, and communication, it is often recommended to submit them as part of the submission package.

14.3.2. Dependency locations

R package dependency is another major factor to consider before preparing your proprietary R package for submission.

For dependencies available from CRAN or public Git repositories, you can declare them directly using the regular Imports and Suggests syntax or the remotes dependency syntax in the DESCRIPTION file.

For dependencies hosted in private Git repositories, you should pack them with the primary R package(s) you want to submit, as pkglite supports packing multiple R packages into a single text file; then restore and install them in the order they are packed.

14.3.3. R version

Always use a consistent version of R for developing the TLFs and for submission. For example, you could enforce a rule to only use $R \times y \cdot z$ where z = 1, such as R 4.0.1 or R 4.1.1. This can be automatically checked using a startup script when the R project is opened.

14.3.4. Package repo version

Always use the same snapshot package repo for developing the TLFs and for submission. Again, this can be checked in the project startup script, as discussed in Section 11.2.

14.3.5. System environments

Introducing any extra external dependencies will likely increase the cost of qualification, validation, testing, and maintenance, especially under Windows. Therefore, it is recommended to keep the dependency chain simple, especially when involving compiled code (e.g., C, C++, Fortran).

14.4. Prepare R packages for submission

To prepare R packages for submission, one needs to pack the packages into text files, and then verify if the files only contain ASCII characters. With packed packages, one can unpack and install them from the text files, too.

14.4.1. Pack

Let's pack the esubdemo package into a text file. Assume the source package path is esubdemo/. You should be able to pack the package with a single pipe:

```
library("pkglite")

"esubdemo/" %>%
  collate(file_ectd(), file_auto("inst")) %>%
  pack(output = "r0pkgs.txt")
```

Let's open the generated text file:

```
file.edit("r0pkgs.txt")
```

What happened in the pipe? The function pkglite::collate() evaluates a specified scope of folders and files defined by a list of file specifications,

```
-- Packing into pkglite file -----
-- Reading package: esubdemo ------
Reading ".Rbuildignore"
Reading "DESCRIPTION"
Reading "NAMESPACE"
Reading "README.md"
Reading "R/count_by.R"
Reading "R/fmt.R"
Reading "R/utils-pipe.R"
Reading "R/zzz.R"
Reading "man/count_by.Rd"
Reading "man/fmt_ci.Rd"
Reading "man/fmt_est.Rd"
Reading "man/fmt_num.Rd"
Reading "man/fmt_pval.Rd"
Reading "man/pipe.Rd"
Reading "inst/pkgdown/assets/readme.txt"
Reading "inst/pkgdown/templates/readme.txt"
Reading "inst/startup.R"
Writing to: "r0pkgs.txt"
```

Figure 14.1.: Output of pkglite::pack()

and generates a **file collection** object. This file collection contains the metadata required to properly convert the files into text which is then used by pkglite::pack(). With this flow, you can define the scope of the R source package to be packed for submission in a flexible yet principled way.

To pack multiple R packages, simply feed multiple file collections as inputs:

```
pack(
   "/path/to/pkg1/" %>% collate(file_ectd()),
   "/path/to/pkg2/" %>% collate(file_ectd()),
   output = "r0pkgs.txt"
)
```

The R packages are always packed in the specified order and are always unpacked and installed in the same order. Therefore, make sure to pack

14. Submission package

```
r0pkgs.txt ×
                                                                                                                                                                                                                                    ___
        # Generated by pkglite: do not edit by hand
# Use pkglite::unpack() to restore the packages
        Package: esubdemo
File: .Rbuildignore
Format: text
Content:
*renv$
              10
       11
      12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
                   ^README\.Rmd$
^\.gitignore$
^\.DS_Store$
^\.git\$
^Meta$
^codecov\.yml$
^vignettes$
^tests/testthat/_snaps$
^library$
^data-raw$
^LICENSE\.md$
^output$
                    ^output$
^adam$
^\.github$
                    ^\.gitnus

^readme.md$

^esubdemo\.Rcheck$

^esubdemo.*\.tar\.gz$

^esubdemo.*\.tgz$
                Package: esubdemo
File: DESCRIPTION
Format: text
                Content:
Package: esubdemo
                    Type: Package
Title: A demo project for analysis and reporting of clinical trials
Version: 0.0.1
      41
42
  50:13
                                                                                                                                                                                                                             Text file $
```

Figure 14.2.: Preview the generated text file

the low-level dependencies first.

For more details on how to customize file specifications and operate on file collections, check out the vignette generate file specifications and curate file collections.

14.4.2. Verify

You should always verify if the text file only contains ASCII characters:

```
verify_ascii("r0pkgs.txt")
```

This should give TRUE if the file only contains ASCII characters, or FALSE with the affected lines otherwise.

14.4.3. Unpack

One can unpack and install the package from the text file, too. For example:

```
unpack("r0pkgs.txt", output = "/tmp/", install = TRUE)
```

If the test is successful, this command can be used in the ADRG instructions for restoring and installing the packed R package(s).

You can then proceed to move the file r0pkgs.txt to the folder ectddemo/m5/datasets/ectddemo/analysis/adam/programs/, or specify the output text file path above directly.

14.5. Prepare analysis programs for submission

Besides the R packages, we need to convert the R Markdown (.Rmd) files into .txt files and saved them in the programs/ folder. You can do this with knitr::purl():

```
input_path <- "esubdemo/vignettes/"
output_path <- "ectddemo/m5/datasets/ectddemo/analysis/adam/programs/"

convert_rmd <- function(filename, input_dir, output_dir) {
   knitr::purl(
     file.path(input_dir, paste0(filename, ".Rmd")),
     output = file.path(output_dir, paste0(filename, ".txt"))
   )
}

"tlf-01-disposition" %>% convert_rmd(input_path, output_path)
"tlf-02-population" %>% convert_rmd(input_path, output_path)
"tlf-03-baseline" %>% convert_rmd(input_path, output_path)
"tlf-04-efficacy" %>% convert_rmd(input_path, output_path)
"tlf-05-ae-summary" %>% convert_rmd(input_path, output_path)
"tlf-06-ae-spec" %>% convert_rmd(input_path, output_path)
"tlf-06-ae-spec" %>% convert_rmd(input_path, output_path)
```

Optionally, you can add a header to the individual .txt files to explain the context and help the reviewers rerun the code. For example:

```
# Note to Reviewer
# To rerun the code below, please refer to the ADRG appendix.
# After the required packages are installed,
# the path variable needs to be defined by using the example code below.
#
# path = list(adam = "/path/to/esub/analysis/adam/datasets") # Modify to use.
```

```
\# path\$outtable = path\$outgraph = "." \# Outputs saved to the current folder
```

To automate this process:

```
header <- readLines(textConnection("# Note to Reviewer
# To rerun the code below, please refer to the ADRG appendix.
# After the required packages are installed,
# the path variable needs to be defined by using the example code below.
# path = list(adam = \"/path/to/esub/analysis/adam/datasets\") # Modify to use actual loc
# path$outtable = path$outgraph = \".\" # Outputs saved to the current folder"))
append_header <- function(filename, output_dir, header) {</pre>
  file <- file.path(output_dir, paste0(filename, ".txt"))</pre>
  x <- readLines(file)
  y <- c(header, "", x)
  writeLines(y, con = file)
  invisible(file)
}
"tlf-01-disposition" %>% append_header(output_path, header)
"tlf-02-population" %>% append_header(output_path, header)
"tlf-03-baseline" %>% append_header(output_path, header)
"tlf-04-efficacy" %>% append_header(output_path, header)
"tlf-05-ae-summary" %>% append_header(output_path, header)
"tlf-06-ae-spec" %>% append_header(output_path, header)
```

14.6. Update ADRG

After we converted the R packages and R Markdown files into the appropriate formats and verified that they can be restored and executed correctly,

14. Submission package

we need update the ADRG to provide guidelines on how to use them.

Specifically, we need to update two sections in ADRG.

The first section is "Macro Programs", where R and R package versions with a snapshot date are provided. For example:

7.x Macro Programs

Submitted R programs have [specific patterns] in filenames. All internally developed R functions are saved in the r0pkgs.txt file. The recommended steps to unpack these R functions for analysis output programs are described in the Appendix.

The tables below contain the software version and instructions for executing the R analysis output programs:

Program Name	Output Table	Title
tlf-01-disposition.txt	Table x.y.z	Disposition of Patients
tlf-02-population.txt	Table x.y.z	Participants Accounting in Analysis Population (A
tlf-03-baseline.txt	Table x.y.z	Participant Baseline Characteristics (All Participa
tlf-04-efficacy.txt	Table x.y.z	ANCOVA of Change from Baseline Glucose (mmo
tlf-05-ae-summary.txt	Table x.y.z	Analysis of Adverse Event Summary (Safety Anal
tlf-06-ae-spec.txt	Table x.y.z	Analysis of Participants With Specific Adverse Ev

Open-Source R Analysis Package	Package Version	Analysis Package Description
pkglite	0.2.0	Prepare submission package
haven	2.4.3	Read SAS datasets
dplyr	1.0.7	Manipulate datasets
tidyr	1.1.3	Manipulate datasets
emmeans	1.6.2-1	Least-squares means estimation
r2rtf	0.3.0	Create RTF tables

Proprietary R Analysis Package	Package Version	Analysis Package Description
esubdemo	0.1.0	A demo package for analysis and reporting of clinical t

The second section (Appendix) should include step-by-step instructions to reproduce the running environment and rerun analyses. For example:

Appendix: Instructions to Execute Analysis Program in R

1. Install R

Download and install R 4.1.1 for Windows from https://cran.r-project.org/bin/windows/base/old/4.1.1/R-4.1.1-win.exe

2. Define working directory

Create a temporary working directory, for example, "C:\tempwork". Copy all submitted R programs into the temporary folder. All steps below should be executed in this working directory represented as "." in the example R code below.

3. Specify R package repository

The R packages are based on CRAN at 2021-08-06. To install the exact R package versions used in this project, run the code below to set the snapshot repository.

```
options(repos = "https://packagemanager.posit.co/cran/2021-08-06")
```

4. Install open-source R packages

In the same R session, install the required packages by running the code below.

```
install.packages(c("pkglite", "publicpkg1", "publicpkg2"))
```

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5. Install proprietary R packages

All internal R packages are packed in the file rOpkgs.txt. In the same R s restore the package structures and install them by running the code below. Adjust the output path as needed to use a writable local directory.

```
pkglite::unpack("rOpkgs.txt", output = ".", install = TRUE)
```

6. Update path to dataset and TLFs

INPUT path: to rerun the analysis programs, define the path variable

- Path for ADaM data: path\$adam

OUTPUT path: to save the analysis results, define the path variable

- Path for output TLFs: path\$output

All these paths need to be defined before executing the analysis program.

path = list(adam = "/path/to/esub/analysis/adam/datasets/") # Modify to us path\$outtable = path\$outgraph = "." # Outputs saved to the current folder

7. Execute analysis program

To reproduce the analysis results, rerun the following programs:

- tlf-01-disposition.txt
- tlf-02-population.txt
- tlf-03-baseline.txt
- tlf-04-efficacy.txt
- tlf-05-ae-summary.txt

```
- tlf-06-ae-spec.txt
```

An example ADRG following this template can be found in ectddemo.

14.7. Update ARM

The ARM (Analysis Results Metadata) should provide specific information related to R in two sections:

- Section 2: indicate the **Programming Language**;
- Section 3: document the details of the R programs listed in section 3.

For example, in ARM section 2, "Analysis Results Metadata Summary":

```
if (knitr::is_latex_output()) {
  df4 %>% kbl(format = "latex")
}
```

Table Reference	Table Title	Programming Language	Program Name (
[Ref. x.y.z: P001ZZZ9999: Table 1-1]	Disposition of Patients	R	tlf-01-disposition

```
if (knitr::is_html_output()) {
    df4 %>%
        kbl(format = "html") %>%
        kable_classic(full_width = FALSE, html_font = "'Times New Roman', Times, serif", font
        column_spec(1, extra_css = "border: 1px solid #000; text-align: center;") %>%
        column_spec(2, extra_css = "border: 1px solid #000; text-align: center;") %>%
        column_spec(3, extra_css = "border: 1px solid #000; text-align: center;") %>%
        column_spec(4, extra_css = "border: 1px solid #000; text-align: center;") %>%
```

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```
column_spec(5, extra_css = "border: 1px solid #000; text-align: center
row_spec(0, background = "#DFDFDF", bold = TRUE, extra_css = "border:
}
```

In ARM section 3, "Analysis Results Metadata Details":

x1	x2
Table Reference: [Ref. x.y.z: P001ZZZ9999: Table 1-1]	
Analysis Result	
Analysis Parameters (s)	
Analysis Reason	
Analysis Purpose	
Programming Statements	(R version 4.1.1), [P001ZZZ9999:

15. Running environment

In the previous chapter, we generated instructions to manually create the running environments for reproducing the A&R deliverables.

In this chapter, we focus on automating the creation of the R environments with R code to accelerate the dry run testing process, simplifying the ADRG instructions, and making it easy to recreate different environment settings with reproducible analysis results.

15.1. Prerequisites

cleanslate is an R package that offers a solution to create portable R environments.

i Note

As of Q4 2021, the cleanslate package used in this chapter is still under active development and validation. This chapter gives a preview of the planned APIs. They may change in the future.

Install cleanslate from CRAN (once available):

```
install.packages("cleanslate")
```

Or from GitHub (once available):

```
remotes::install_github("Merck/cleanslate")
```

15.2. Practical considerations

The cleanslate package supports:

- Creating a project folder with project-specific context (.Rproj, .Rprofile, .Renviron)
- Installing a specific version of R into the project folder
- Installing a specific version of Rtools into the project folder

An essential feature of cleanslate is that it does **not** require administrator privileges to run R and Rtools installers. This makes it easier to deploy under enterprise settings and avoids security and portability concerns.

As many of the A&R deliverables are currently created, validated, and delivered under Windows, the primary focus is Windows at the moment, while the support for other platforms might be added in future versions.

15.3. Create canonical environments

One can create a running environment with "canonical" settings with a single function call to use_cleanslate():

```
cleanslate::use_cleanslate(
   "C:/temp/",
   r_version = "4.1.1",
   from = "https://cran.r-project.org/",
   repo = "https://packagemanager.posit.co/cran/2021-08-06"
)
```

This will

- Create a project folder under C:/temp/ with a .Rproj file;
- Download R 4.1.1 installer from CRAN, and install it into C:/temp/R/;
- Not install Rtools (by default, rtools_version = NULL);
- Create a .Rprofile file under the project folder, set options(repos) to use the specified repo (a Posit Public Package Manager snapshot in this example), and give instruction to set the R binary path in RStudio IDE;
- Create a .Renviron file under the project folder and set the library path to be the library of the project-specific R installation.

As a principle, one should always double-click the .Rproj file to open the project. This will ensure some sanity checks in the .Rprofile, such as whether the R and library are located within the project folder.

15.4. Create tailored environments

To create a more customized running environment, one can use the specific functions to tailor each aspect, for example:

```
library("cleanslate")

"C:/temp/" %>%
   use_project() %>%
   use_rprofile() %>%
   use_renviron() %>%
   use_r = "4.1.1") %>%
   use_rtools(version = "rtools40")
```

The project context functions (use_project(), use_rprofile(), use_renviron) support custom templates using brew.

15. Running environment

The use_r_*() functions have variations that serve as shortcuts to use R versions defined by release lifecycles, for example, use_r_release(), use_r_oldrel(), and use_r_devel(). Note that to ensure better reproducibility, one should still use use_r_version() as the release, oldrel, and devel versions will shift as time goes by.

The helper functions version_*() and snapshot_*() can assist you in determining specific versions of R and Rtools that are currently available, besides generating and verifying the snapshot repo links.

15.5. Update ADRG

If you use cleanslate, remember to update the ADRG instructions for executing the analysis programs in R. Mostly, this can simplify the first three steps on creating a project, installing a specific version of R, and configuring the package repo location. For example:

```
Appendix: Instructions to Execute Analysis Program in R

1. Setup R environment

Open the existing R, install the required packages by running the code belinstall.packages("cleanslate")

Create a temporary working directory, for example, "C:\tempwork".

Copy all submitted R programs into the temporary folder.

In the same R session, run the code below to create a project with a portable R environment.

cleanslate::use_cleanslate(
    "C:/temp/",
```

```
r_{version} = "4.1.1",
  from = "https://cran.r-project.org/",
  repo = "https://packagemanager.posit.co/cran/2021-08-06"
2. Open the project
Go to the working directory created above, double click the .Rproj file
to open the project in RStudio IDE. Follow the instructions to select the
project-specific R version, then restart RStudio IDE. If successful,
the R version and package repo should be printed as defined above.
3. Install open-source R packages
In the new R session, install the required packages by running the code below.
install.packages(c("pkglite", "publicpkg1", "publicpkg2"))
4. Install proprietary R packages
All internal R packages are packed in the file rOpkgs.txt. In the same R session,
restore the package structures and install them by running the code below.
Adjust the output path as needed to use a writable local directory.
pkglite::unpack("r0pkgs.txt", output = ".", install = TRUE)
5. Update path to dataset and TLFs
INPUT path: to rerun the analysis programs, define the path variable
- Path for ADaM data: path$adam
```

15. Running environment

```
OUTPUT path: to save the analysis results, define the path variable

- Path for output TLFs: path$output

All these paths need to be defined before executing the analysis program.

path = list(adam = "/path/to/esub/analysis/adam/datasets/") # Modify to us path$outtable = path$outgraph = "." # Outputs saved to the current folder

6. Execute analysis program

To reproduce the analysis results, rerun the following programs:

- tlf-01-disposition.txt
- tlf-02-population.txt
- tlf-03-baseline.txt
- tlf-04-efficacy.txt
- tlf-05-ae-summary.txt
- tlf-05-ae-summary.txt
- tlf-06-ae-spec.txt
```

15.6. RStudio addin

To make it convenient to use cleanslate in experiments, one can also use its RStudio IDE addin. After cleanslate is installed, click Addins -> cleanslate -> Create portable R environment in RStudio IDE, or call cleanslate:::create_env_addin() to open it.

The addin provides a wizard-like interface to help create the environment with the most important options, yet with less flexibility compared to the functional API demonstrated above.

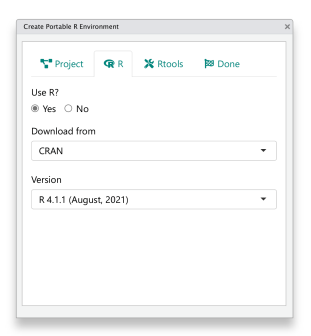


Figure 15.1.: cleanslate RStudio addin

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