# Script 1

Data missingness

Peter Kamerman

 $22 \ January \ 2020$ 

#### Contents

Objective	]
Analysis notes  Definitions of missingness	
Import data	2
Quick look	2
Number of participants with/without complete pain data	3
Demographic variables Process data Ancestry Sex Education Employment status	
Clinical variables  CD4 T-cell count	11 12
Proportion missing pain data by study site	12 14
Session information	16

## Objective

To determine whether the degree of missingness or data inconsistencies were associated with any demographic variables (age, sex, ancestry, education, employment status), study variables (study site and group allocation), or clinical variables (CD4, viral load).

## Analysis notes

#### Definitions of missingness

Data were regarded as **missing** when *pain in the last week* data were not present for one or more of weeks 0, 12, 24, 36, 48. Data also were classified as **missing** when there were inconsistencies in the data across the variables collected within a week.

#### Definition of data inconsistencies

Pain was defined as pain in the last week being 'Yes', and pain at its worst being > 0. These two measurements were then the "gatekeeper" measurements, such that the two measurements both had to be positive ('Yes' and '> 0', respectively) in order for there to be any entries for site of pain and site of worst pain. Were the data were inconsistent (e.g., when there was no pain in the last week and pain at its worst = 0, but there were entries for site of pain and site of worst pain), then the site of pain and site of worst pain entries were marked as **inconsistent**.

Data also were considered **inconsistent** when pain in the last week = 'Yes', but site of worst pain = 'None'.

Lastly, data were considered **inconsistent** when *site of worst pain* was not listed as one of the pain locations for a given measurement week.

For analysis purposes, missing data in the *site of pain* columns were changed to 'No' (pain not present at the site). This approach was conservative, but we believed that the approach would have the least effect on the outcome, while still retaining as many participants as possible.

#### Import data

```
df <- read_rds('data-cleaned/data-ADVANCE.rds')</pre>
```

### Quick look

```
head(df)
## # A tibble: 6 x 32
     ranid interval name site name pain in the las... where does it h...
##
     <chr> <ord>
                          <chr>
                                     <chr>
                                                       <chr>
## 1 01-0... 0 weeks
                          Wits RHI... No
                                                       <NA>
## 2 01-0... 12 weeks
                          Wits RHI... No
                                                       <NA>
## 3 01-0... 24 weeks
                          Wits RHI... No
                                                       <NA>
## 4 01-0... 36 weeks
                          Wits RHI... No
                                                       <NA>
## 5 01-0... 48 weeks
                          Wits RHI... No
                                                       <NA>
## 6 01-0... 0 weeks
                          Wits RHI... No
                                                       <NA>
## # ... with 27 more variables: pain_worst <dbl>, pain_now <dbl>,
       head_pain <chr>, cervical_pain <chr>, shoulder_pain <chr>,
## #
## #
       arm_pain <chr>, hand_pain <chr>, chest_pain <chr>,
## #
       abdominal_pain <chr>, low_back_pain <chr>, buttock_pain <chr>,
## #
       hip_groin_pain <chr>, leg_pain <chr>, genital_pain <chr>,
## #
       foot_pain <chr>, site_worst <chr>, age <dbl>, sex <chr>,
## #
       ancestry <chr>, education <chr>, employment_status <chr>, group <chr>,
## #
       cd4_cells.ul <dbl>, viral_load_cp.ml <dbl>, general_health <dbl>,
## #
       interval_numeric <dbl>, any_missing <chr>
glimpse(df)
## Observations: 5,265
```

```
## Variables: 32
## $ ranid
                              <chr> "01-0001", "01-0001", "01-0001", "01-000...
## $ interval_name
                              <ord> 0 weeks, 12 weeks, 24 weeks, 36 weeks, 4...
## $ site_name
                              <chr> "Wits RHI Yeoville Research Centre", "Wi...
                              <chr> "No", "No", "No", "No", "No", "No", "Yes...
## $ pain_in_the_last_week
## $ where_does_it_hurt_most <chr> NA, NA, NA, NA, NA, NA, "Hip/groin left"...
## $ pain_worst
                              <dbl> 0, 0, 0, 0, 0, 0, 3, 3, 5, 0, 0, 0, 0...
## $ pain_now
                              <dbl> NA, O, NA, O, NA, NA, O, 2, 4, NA, NA, O...
## $ head_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
```

```
<chr> "No", "No", "No", "No", "No", "No", "No"...
## $ cervical_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ shoulder_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ arm pain
## $ hand_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ chest_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ abdominal_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ low_back_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ buttock_pain
                              <chr> "No", "No", "No", "No", "No", "No", "Yes...
## $ hip_groin_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ leg_pain
                              <chr> "No", "No", "No", "No", "No", "No",
## $ genital_pain
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
## $ foot_pain
                              <chr> "None", "None", "None", "None", "None", ...
## $ site_worst
                              <dbl> 30, 30, 30, 30, 30, 34, 34, 34, 34, 34, ...
## $ age
## $ sex
                              <chr> "Male", "Male", "Male", "Male", "Male", ...
## $ ancestry
                              <chr> "Black", "Black", "Black", "Black", "Bla...
## $ education
                              <chr> "Secondary", "Secondary", "Secondary", "...
                             <chr> "Employed", "Employed", "Employed", "Emp...
## $ employment_status
                              <chr> "DTG + TAF + FTC", "DTG + TAF + FTC", "D...
## $ group
## $ cd4 cells.ul
                              <dbl> 642, NA, 525, NA, 668, 241, NA, 364, NA,...
## $ viral_load_cp.ml
                              <dbl> 641, 50, 50, 50, 50, 3851, 50, 50, 50, 5...
## $ general_health
                              <dbl> 4, 4, 5, 5, 4, 3, 5, 3, 3, 3, 4, 5, 5, 5...
## $ interval_numeric
                              <dbl> 0, 12, 24, 36, 48, 0, 12, 24, 36, 48, 0,...
## $ any_missing
                              <chr> "No", "No", "No", "No", "No", "No", "No"...
```

### Number of participants with/without complete pain data

Table 1: Number of participants with/without complete pain data

any_missing	count	n	proportion
No	787	1053	0.747
Yes	266	1053	0.253

## Demographic variables

#### Process data

```
# Extract demographic data
df_demo <- df %>%
```

#### Ancestry

```
# Check counts
df_combined %>%
group_by(ancestry) %>%
summarise(count = n()) %>%
knitr::kable(caption = 'Count within each category of self-identified ancestry')
```

Table 2: Count within each category of self-identified ancestry

ancestry	count
Black	1051
Coloured	2

Only 2 out of 1053 participants did not identify and Black African, and therefore no analysis done on ancestry.

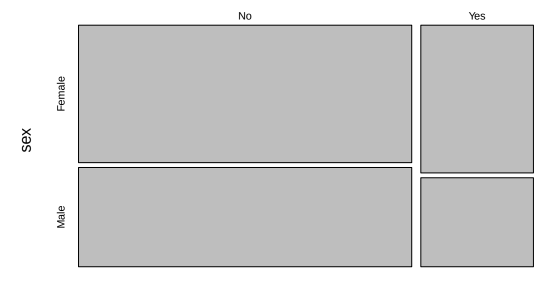
#### Sex

Table 3: Missing pain data by sex

sex	any_missing	count	total	proportion
Female	No	457	623	0.734
Female	Yes	166	623	0.266
Male	No	330	430	0.767
Male	Yes	100	430	0.233

```
# Tabulate, plot, and test
tab_sex <- xtabs(~any_missing + sex, data = df_combined)
mosaicplot(tab_sex, main = 'Counts by sex')</pre>
```

# Counts by sex

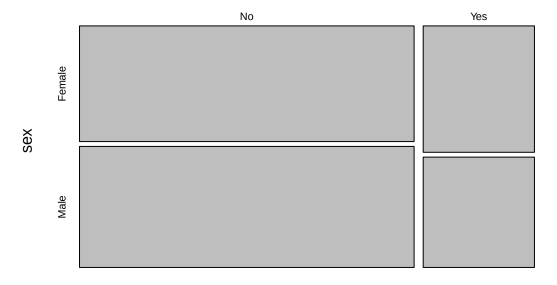


### any\_missing

```
prop_sex <- prop.table(tab_sex, 2)

mosaicplot(prop_sex, main = 'Proportions by sex')</pre>
```

## **Proportions by sex**



any\_missing

```
fisher.test(tab_sex)
```

```
##
## Fisher's Exact Test for Count Data
##
## data: tab_sex
## p-value = 0.2207
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 0.6196133 1.1201888
## sample estimates:
```

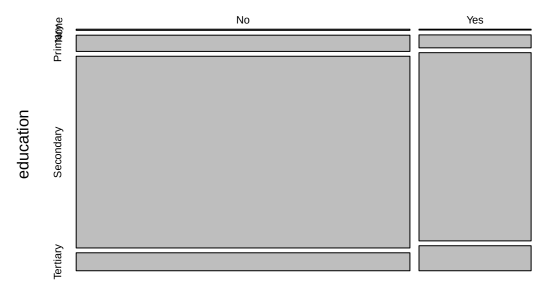
```
## odds ratio
## 0.8343897
```

#### Education

Table 4: Missing pain data by level of education

education	any_missing	count	total	proportion
No schooling	No	4	5	0.800
No schooling	Yes	1	5	0.200
Primary	No	56	71	0.789
Primary	Yes	15	71	0.211
Secondary	No	661	879	0.752
Secondary	Yes	218	879	0.248
Tertiary	No	62	91	0.681
Tertiary	Yes	29	91	0.319
(Missing)	No	4	7	0.571
(Missing)	Yes	3	7	0.429

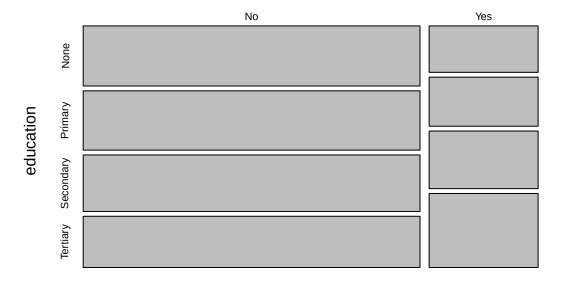
# Counts by level of education



any\_missing

```
prop_edu <- prop.table(tab_edu, 2)
mosaicplot(prop_edu, main = 'Proportions by level of education')</pre>
```

## Proportions by level of education



any\_missing

```
fisher.test(tab_edu)
```

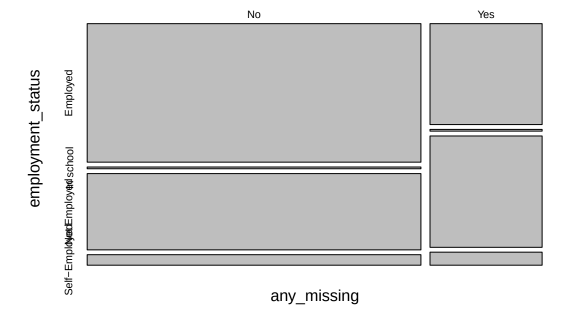
```
##
## Fisher's Exact Test for Count Data
##
## data: tab_edu
## p-value = 0.4006
## alternative hypothesis: two.sided
```

#### Employment status

Table 5: Missing pain data by employment status

employment_status	any_missing	count	total	proportion
Employed	No	474	590	0.803
Employed	Yes	116	590	0.197
Not Employed	No	261	389	0.671
Not Employed	Yes	128	389	0.329
Schooling	No	6	8	0.750
Schooling	Yes	2	8	0.250
Self-Employed	No	36	51	0.706
Self-Employed	Yes	15	51	0.294
(Missing)	No	10	15	0.667
(Missing)	Yes	5	15	0.333

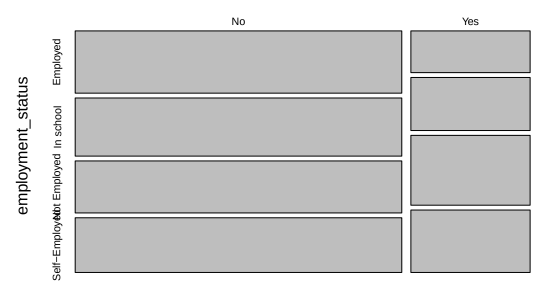
## Counts by employment status



```
prop_employ <- prop.table(tab_employ, 2)

mosaicplot(prop_employ, main = 'Proportions by employment status')</pre>
```

### **Proportions by employment status**



#### any\_missing

```
##
## Fisher's Exact Test for Count Data
##
## data: tab_employ
```

## alternative hypothesis: two.sided

Those who were unemployed had the greatest proportion of missing values.

#### Clinical variables

fisher.test(tab\_employ)

## p-value = 3.88e-05

#### CD4 T-cell count

low\_CD4 defined as the lowest CD4 T-cell count measured during the course of the first 48 weeks of the study.

```
# Process the CD4 data and join with missingness data

df_CD4 <- df %>%
    select(ranid, interval_name, cd4_cells.ul) %>%
    # Determine highest VL per participant
    group_by(ranid) %>%
    summarise(low_CD4 = min(cd4_cells.ul, na.rm = TRUE))

df_CD4 <- df_combined %>%
    select(ranid, any_missing) %>%
    left_join(df_CD4)
```

## Joining, by = "ranid"

```
# Tabulate and print
df_CD4 %>%
    group_by(any_missing) %>%
    select(any_missing, low_CD4) %>%
    skim() %>%
    skimr::kable(caption = 'Data missingness by lowest CD4')
```

Skim summary statistics

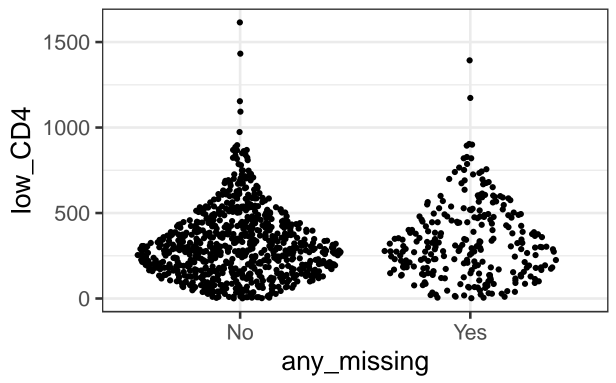
n obs: 1053 n variables: 2

Variable type: numeric

any_missing	variable	missing	complete	n	mean	$\operatorname{sd}$	p0	p25	p50	p75	p100
No	$low\_CD4$	0	787	787	315.84	201.84	1	171	286	424	1615
Yes	$low\_CD4$	0	266	266	335.17	221.53	2	170.75	300	476.5	1393

```
# Plot, and test
ggplot(data = df_CD4) +
   aes(x = any_missing,
        y = low_CD4) +
   geom_sina() +
   labs(subtitle = 'Density plot: data missingness vs lowest CD4')
```

# Density plot: data missingness vs lowest CD



```
wilcox.test(low_CD4 ~ any_missing, data = df_CD4)
```

Wilcoxon rank sum test with continuity correction

data: low CD4 by any missing W = 100308, p-value = 0.309 alternative hypothesis: true location shift is not equal to 0

#### Viral load

high\_VL defined as the highest viral load measured during the course of the first 48 weeks of the study.

```
# Process the VL data and join with missingness data
df_VL <- df %>%
    select(ranid, interval_name, viral_load_cp.ml) %>%
    # Determine highest VL per participant
    group_by(ranid) %>%
    summarise(high_VL = max(viral_load_cp.ml, na.rm = TRUE))
df_VL <- df_combined %>%
    select(ranid, any_missing) %>%
    left_join(df_VL)
## Joining, by = "ranid"
# Tabulate and print
df_VL %>%
    group_by(any_missing) %>%
    select(any_missing, high_VL) %>%
    skim() %>%
    skimr::kable(caption = 'Data missingness by greatest viral load')
```

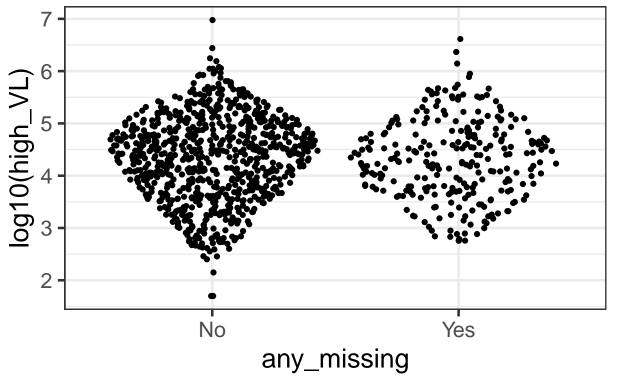
Skim summary statistics

n obs: 1053 n variables: 2

Variable type: numeric

any_missing	variable	missing	complete	n	mean	sd	p0	p25	p50	p75	p100
No	high_VL	0	787	787	99267.18	386828.79	50	5791	26333	85912.5	9475772
Yes	$high\_VL$	0	266	266	96926.88	318787.27	570	5935.5	20859.5	66562	4117370

# Density plot: data missingness vs highest viral I



```
wilcox.test(high_VL ~ any_missing, data = df_VL)
```

Wilcoxon rank sum test with continuity correction

data: high\_VL by any\_missing W = 107520, p-value = 0.5064 alternative hypothesis: true location shift is not equal to 0

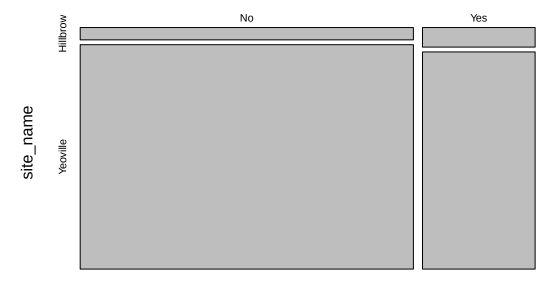
## Study variables

#### Proportion missing pain data by study site

Table 8: Proportion missing pain data by study site

site_name	any_missing	count	total	proportion
Wits RHI Shandukani Hillbrow Johannesburg	No	41	63	0.651
Wits RHI Shandukani Hillbrow Johannesburg	Yes	22	63	0.349
Wits RHI Yeoville Research Centre	No	746	990	0.754
Wits RHI Yeoville Research Centre	Yes	244	990	0.246

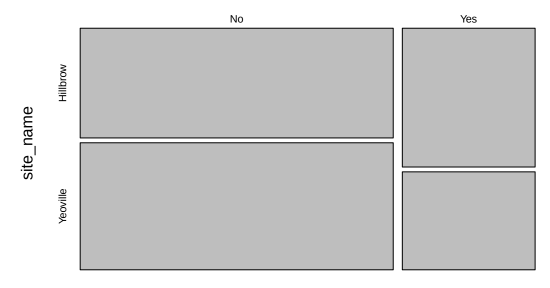
## Counts by study site



#### any\_missing

```
prop_site <- prop.table(tab_site, 2)
mosaicplot(prop_site, main = 'Proportion of counts by study site')</pre>
```

## Proportion of counts by study site



any\_missing

```
chisq.test(tab_site)

##

## Pearson's Chi-squared test with Yates' continuity correction
##

## data: tab_site
## X-squared = 2.7898, df = 1, p-value = 0.09487
```

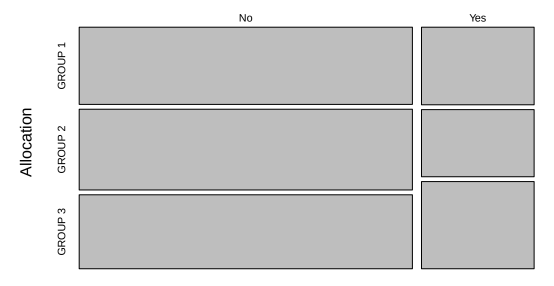
#### Proportion missing pain data by group allocation

```
GROUP 1: DTG + TAF + FTC
GROUP 2: DTG + TDF + FTC
GROUP 3: EFV + TDF + FTC
```

Table 9: Proportion missing pain data by group allocation

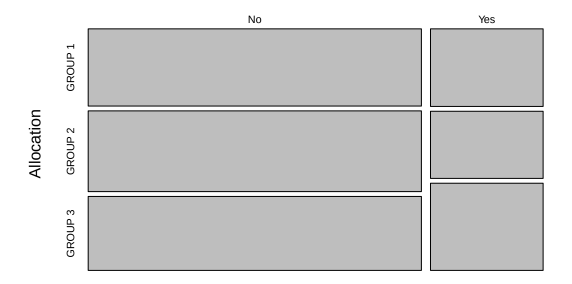
group	any_missing	count	total	proportion
$\overline{\mathrm{DTG} + \mathrm{TAF} + \mathrm{FTC}}$	No	262	351	0.746
DTG + TAF + FTC	Yes	89	351	0.254
DTG + TDF + FTC	No	274	351	0.781
DTG + TDF + FTC	Yes	77	351	0.219
EFV + TDF + FTC	No	251	351	0.715
EFV + TDF + FTC	Yes	100	351	0.285

## Counts by group allocation



#### any\_missing

## **Proportions by group allocation**



any\_missing

```
##
## Fisher's Exact Test for Count Data
##
## data: tab_group
## p-value = 0.1361
## alternative hypothesis: two.sided
```

#### Session information

#### sessionInfo()

```
## R version 3.6.1 (2019-07-05)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS Mojave 10.14.6
##
## Matrix products: default
## BLAS:
         /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en US.UTF-8/en US.UTF-8/en US.UTF-8/C/en US.UTF-8/en US.UTF-8
## attached base packages:
                graphics grDevices utils
## [1] stats
                                               datasets methods
                                                                   base
##
## other attached packages:
## [1] ggforce_0.3.1
                        skimr_1.0.7
                                        forcats_0.4.0
                                                        stringr_1.4.0
## [5] dplyr_0.8.3
                        purrr_0.3.3
                                        readr_1.3.1
                                                        tidyr_1.0.0
## [9] tibble_2.1.3
                        ggplot2_3.2.1
                                        tidyverse_1.2.1
##
## loaded via a namespace (and not attached):
## [1] tidyselect_0.2.5 xfun_0.10
                                          haven_2.1.1
                                                           lattice_0.20-38
   [5] colorspace_1.4-1 vctrs_0.2.0
##
                                          generics_0.0.2
                                                           htmltools 0.4.0
## [9] yaml_2.2.0
                        utf8_1.1.4
                                          rlang_0.4.2
                                                           pillar_1.4.2
## [13] glue_1.3.1
                        withr_2.1.2
                                          tweenr_1.0.1
                                                           modelr_0.1.5
## [17] readxl_1.3.1
                        lifecycle_0.1.0 munsell_0.5.0
                                                           gtable_0.3.0
## [21] cellranger_1.1.0 rvest_0.3.4
                                          evaluate_0.14
                                                           labeling_0.3
## [25] knitr 1.25
                        fansi 0.4.0
                                          highr 0.8
                                                           broom 0.5.2
                         scales_1.0.0
                                          backports_1.1.5 jsonlite_1.6
## [29] Rcpp_1.0.3
## [33] farver_1.1.0
                        \mathtt{hms}_{\mathtt{0.5.1}}
                                          digest_0.6.23
                                                           stringi_1.4.3
## [37] polyclip_1.10-0 grid_3.6.1
                                          cli_2.0.0
                                                           tools_3.6.1
## [41] magrittr_1.5
                         lazyeval_0.2.2
                                          crayon_1.3.4
                                                           pkgconfig_2.0.3
## [45] zeallot_0.1.0
                         MASS_7.3-51.4
                                          xm12_1.2.2
                                                           lubridate_1.7.4
                                          httr_1.4.1
## [49] assertthat_0.2.1 rmarkdown_1.16
                                                           rstudioapi_0.10
## [53] R6_2.4.1
                        nlme_3.1-141
                                          compiler_3.6.1
```