# Software quality analysis of fourteen hydrological models

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### 1 Preliminary functions

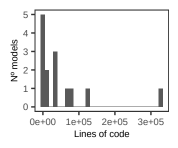
```
sensobol::load_packages(c("data.table", "tidyverse", "openxlsx", "scales",
                         "cowplot", "readxl", "ggrepel", "tidytext", "here"))
# Create custom theme -----
theme_AP <- function() {</pre>
 theme_bw() +
   theme(panel.grid.major = element_blank(),
         panel.grid.minor = element_blank(),
         legend.background = element_rect(fill = "transparent", color = NA),
         legend.key = element_rect(fill = "transparent", color = NA),
         strip.background = element_rect(fill = "white"),
         legend.text = element_text(size = 7.3),
         axis.title = element text(size = 10),
         legend.key.width = unit(0.4, "cm"),
         legend.key.height = unit(0.4, "cm"),
         legend.key.spacing.y = unit(0, "lines"),
         legend.box.spacing = unit(0, "pt"),
         legend.title = element_text(size = 7.3),
         axis.text.x = element_text(size = 7),
         axis.text.y = element_text(size = 7),
         axis.title.x = element_text(size = 7.3),
         axis.title.y = element_text(size = 7.3),
         plot.title = element_text(size = 8),
         strip.text.x = element_text(size = 7.4),
         strip.text.y = element_text(size = 7.4))
}
# Select color palette -----
color_languages <- c("fortran" = "steelblue", "python" = "lightgreen")</pre>
# Source all .R files in the "functions" folder -----
r_functions <- list.files(path = here("functions"),</pre>
                         pattern = "\\.R$", full.names = TRUE)
lapply(r_functions, source)
## [[1]]
## [[1]]$value
## function (plot, legend = NULL)
## {
##
      if (is.ggplot(plot)) {
```

```
##
            gt <- ggplotGrob(plot)</pre>
##
##
       else {
##
            if (is.grob(plot)) {
                gt <- plot
##
            }
##
##
            else {
##
                stop("Plot object is neither a ggplot nor a grob.")
##
       }
##
##
       pattern <- "guide-box"</pre>
##
       if (!is.null(legend)) {
            pattern <- paste0(pattern, "-", legend)</pre>
##
##
       indices <- grep(pattern, gt$layout$name)</pre>
##
       not_empty <- !vapply(gt$grobs[indices], inherits, what = "zeroGrob",</pre>
##
##
            FUN.VALUE = logical(1))
##
       indices <- indices[not_empty]</pre>
       if (length(indices) > 0) {
##
##
            return(gt$grobs[[indices[1]]])
##
       return(NULL)
##
## }
## [[1]]$visible
## [1] FALSE
```

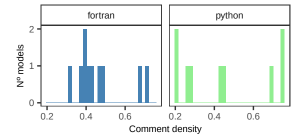
#### 2 Results

#### 2.1 Descriptive statistics

## `stat\_bin()` using `bins = 30`. Pick better value `binwidth`.

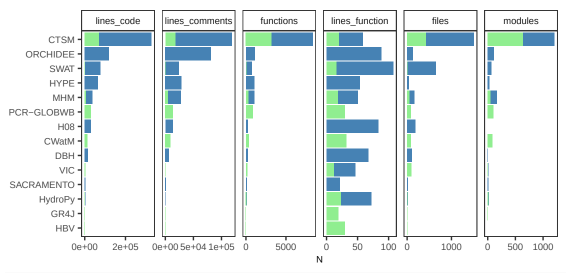


## `stat\_bin()` using `bins = 30`. Pick better value `binwidth`.



```
# Sort by model ------
model_ordered <- dt$descriptive_stats[, sum(lines), model] %>%
 .[order(V1)]
model ordered
##
         model
                 V1
##
        <char> <num>
## 1:
          \mathtt{HBV}
              180
## 2:
          GR4J
              423
## 3:
      HydroPy
              3739
## 4: SACRAMENTO
              5294
## 5:
          VIC
               5952
          DBH 24334
## 6:
## 7:
         CWatM 27745
## 8:
          H08 42917
## 9: PCR-GLOBWB 52686
          MHM 76286
## 10:
## 11:
          HYPE 89137
## 12:
          SWAT 99976
## 13:
     ORCHIDEE 211871
          CTSM 491592
## 14:
# Extract column names --
                            _____
col_names <- colnames(dt$descriptive_stats)</pre>
# Order facets -----
facet_order <- c("lines", "lines_code", "lines_comments", "functions",</pre>
             "lines_function", "files", "modules")
# Plot -----
plot_per_model <- melt(dt$descriptive_stats, measure.vars = col_names[-c(1, length(col_names))]</pre>
 .[, variable:= factor(variable, levels = facet_order)] %>%
 .[, model:= factor(model, levels = model_ordered[, model])] %>%
 .[!variable == "lines"] %>%
 ggplot(., aes(model, value, fill = language)) +
 geom_col() +
 coord_flip() +
 scale_y_continuous(breaks = breaks_pretty(n = 2)) +
```

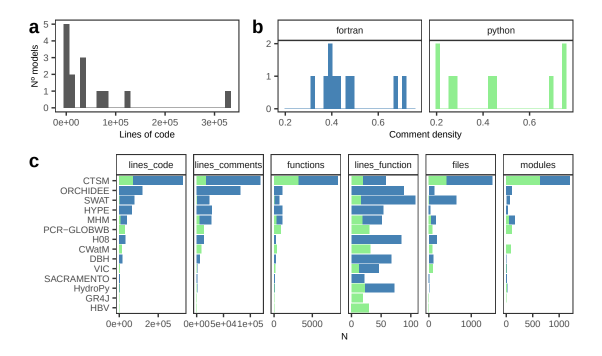
```
scale_fill_manual(values = color_languages) +
facet_wrap(~ variable, ncol = 7, scales = "free_x") +
labs(x = "", y = "N") +
theme_AP() +
theme(legend.position = "none")
```



```
## `stat_bin()` using `bins = 30`. Pick better value `binwidth`.
## `stat_bin()` using `bins = 30`. Pick better value `binwidth`.
p1 <- plot_grid(top, plot_per_model, ncol = 1, labels = c("", "c"), rel_heights = c(0.4, 0.6))</pre>
```

## Warning: Removed 3 rows containing missing values or values outside the scale range
## (`geom\_col()`).

p1

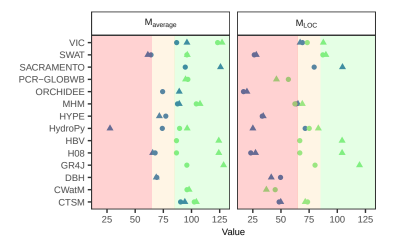


#### 2.2 Maintainability index

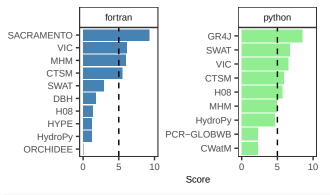
```
# Define vector of interpretation -----
vec_interpretation <- c("low", "moderate", "high")</pre>
# Calculate -----
dt$maintainability_index %>%
 melt(., measure.vars = c("M loc", "M average")) %>%
 .[, interpretativity:= ifelse(value > 85, vec_interpretation[3],
                            ifelse(value <=85 & value >= 65, vec_interpretation[2],
                                  vec_interpretation[1]))] %>%
 .[, .N, .(language, interpretativity, variable)] %>%
 dcast(., variable + language ~ interpretativity, value.var = "N") %>%
 .[, total:= rowSums(.SD, na.rm = TRUE), .SDcols = vec_interpretation] %>%
 .[, paste(vec_interpretation, "prop", sep = "_"):= lapply(.SD, function(x)
   x / total), .SDcols = vec_interpretation] %>%
 print()
## Key: <variable, language>
##
      variable language high
                             low moderate total low_prop moderate_prop
##
        <fctr>
               <char> <int> <int>
                                   <int> <num>
                                                 <num>
                                                              <num>
## 1:
        M loc fortran
                         1
                                           20
                                                  0.70
                                                              0.25
## 2:
        M_{loc}
              python
                               5
                                       9
                                           20
                                                  0.25
                                                              0.45
                         6
## 3: M_average fortran
                        9
                              3
                                      8
                                                  0.15
                                                              0.40
                                           20
## 4: M_average python
                        20
                              NA
                                     NA
                                           20
                                                   NA
                                                                NA
```

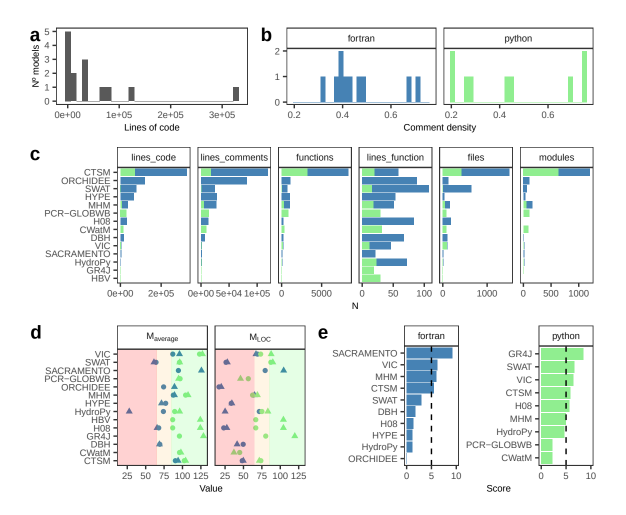
By combining the classic and extended versions of the maintainability index, our analysis reveals differences between Fortran and Python implementations. Using the weighted measure  $(M_{\rm LOC})$ , 70% of Fortran code falls into the "low" maintainability category, compared with only 15% when using the unweighted average  $(M_{\rm average})$ . This discrepancy indicates that a few complex, poorly maintainable routines dominate the overall profile of the Fortran codebase. In contrast, Python routines present a more favorable profile: 27% achieve high maintainability under  $M_{\rm LOC}$ , and all are classified as "highly maintainable" under  $M_{\rm average}$ .

```
plot_maintainability_index <- dt$maintainability_index %>%
 melt(., measure.vars = c("M_loc", "M_average")) %>%
 .[, variable:= factor(variable, levels = c("M_average", "M_loc"))] %>%
 ggplot(., aes(model, value, color = language, shape = type)) +
 geom_point() +
 annotate("rect", xmin = -Inf, xmax = Inf, ymin = -Inf, ymax = 65,
          fill = "red", alpha = 0.18) +
 annotate("rect", xmin = -Inf, xmax = Inf, ymin = 65, ymax = 85,
          fill = "orange", alpha = 0.1) +
 annotate("rect", xmin = -Inf, xmax = Inf, ymin = 85, ymax = Inf,
          fill = "green", alpha = 0.1) +
 facet_wrap(~variable, labeller = as_labeller(c(M_loc = "M[LOC]",
                                           M_average = "M[average]"),
                                         default = label parsed)) +
 labs(x = "", y = "Value") +
 scale_color_manual(values = color_languages, guide = "none") +
 theme_AP() +
 theme(legend.position = "none") +
 coord_flip()
plot_maintainability_index
```



#### 2.3 Score



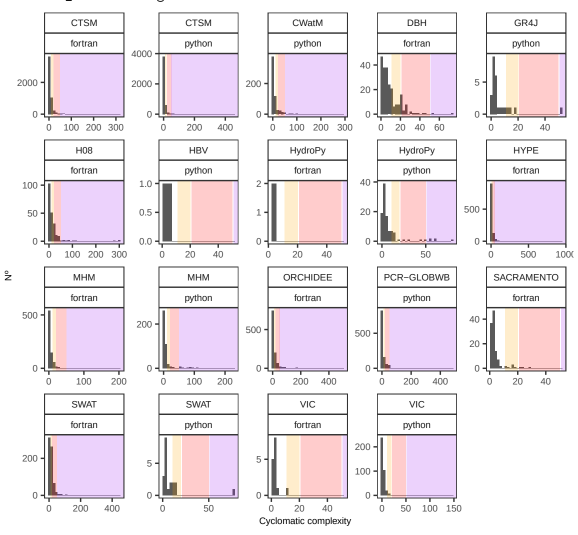


#### 2.4 Metrics at the function level

```
basename(func_metric_files)))
# Create function to combine files -----
make combined <- function(subset list, pattern) {</pre>
 rbindlist(subset_list[grep(pattern, names(subset_list))], idcol = "source_file")
}
# Combine files ------
metrics_combined <- list(file_fortran = make_combined(list_metrics$file_metrics, "fortran"),</pre>
                       file_python = make_combined(list_metrics$file_metrics, "python"),
                       func_fortran = make_combined(list_metrics$func_metrics, "fortran"),
                       func_python = make_combined(list_metrics$func_metrics, "python"))
# Functions to extract name of model and language from file ------
extract_model <- function(x)</pre>
  sub("^(file|func)_metrics_{d+_([A-Za-z0-9-]+)_(fortran|python).*", "\2", x)
extract_lang <- function(x)</pre>
  sub("^(file|func)_metrics_{d+_([A-Za-z0-9-]+)_(fortran|python).*", "\\3", x)
# Extract name of model and language -----
metrics_combined <- lapply(metrics_combined, function(dt) {</pre>
 dt[, source_file:= sub("\\.csv$", "", basename(source_file))]
 dt[, model:= extract_model(source_file)]
 dt[, language:= extract_lang(source_file)]
 dt
})
# Add column of complexity category -----
metrics_combined <- lapply(names(metrics_combined), function(nm) {</pre>
 dt <- as.data.table(metrics_combined[[nm]])</pre>
  if (grepl("^func_", nm) && "cyclomatic_complexity" %in% names(dt)) {
   dt[, complexity_category := cut(
     cyclomatic_complexity,
     breaks = c(-Inf, 10, 20, 50, Inf),
     labels = c("b1","b2","b3","b4")
   )]
 }
  dt
}) |> setNames(names(metrics_combined))
# Define labels -----
```

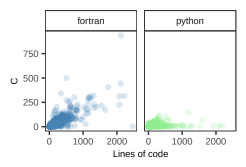
```
lab_expr <- c(</pre>
 b1 = expression(C %in% "(" * 0 * ", 10" * "]"),
 b2 = expression(C \%in\% "(" * 10 * ", 20" * "]"),
 b3 = expression(C %in% "(" * 20 * ", 50" * "]"),
 b4 = expression(C %in% "(" * 50 * ", " * infinity * ")")
# set output folder inside "datasets" -----
outdir <- file.path("datasets", "merged_results")</pre>
# write each slot to its own CSV ------
lapply(names(metrics_combined), function(nm) {
 out_file <- file.path(outdir, paste0(nm, ".csv"))</pre>
 fwrite(metrics_combined[[nm]], out_file)
})
## [[1]]
## NULL
##
## [[2]]
## NULL
##
## [[3]]
## NULL
##
## [[4]]
## NULL
# Cyclomatic complexity at the model level ------
metrics_combined[grep("^func_", names(metrics_combined))] %>%
 lapply(., function(x) x[, .(cyclomatic_complexity, model, language)]) %>%
 rbindlist() %>%
 ggplot(., aes(cyclomatic_complexity)) +
 geom_histogram() +
 annotate("rect",
        xmin = 11, xmax = 20,
         ymin = -Inf, ymax = Inf,
        fill = "orange", alpha = 0.2) +
 annotate("rect",
        xmin = 21, xmax = 50,
```

#### ## `stat\_bin()` using `bins = 30`. Pick better value `binwidth`.



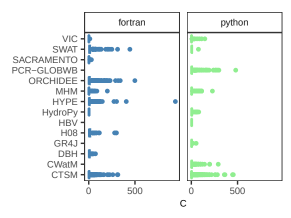
```
plot_scatterplot <- metrics_combined[grep("^func_", names(metrics_combined))] %>%
    lapply(., function(x)
        x[, .(model, language, `function`, cyclomatic_complexity, loc, bugs, type)]) %>%
    rbindlist() %>%
    ggplot(., aes(cyclomatic_complexity, loc, color = language)) +
```

```
geom_point(alpha = 0.2) +
scale_color_manual(values = color_languages) +
coord_flip() +
scale_y_continuous(breaks = pretty_breaks(n = 3)) +
facet_wrap(~language) +
labs(x = "C", y = "Lines of code") +
theme_AP() +
theme(legend.position = "none")
```



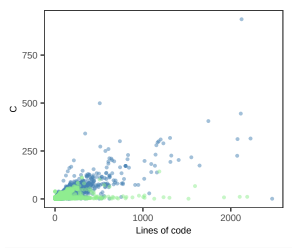
```
plot_c_model <- metrics_combined[grep("^func_", names(metrics_combined))] %>%
    lapply(., function(x)
        x[, .(model, language, `function`, cyclomatic_complexity, loc, bugs, type)]) %>%
    rbindlist() %>%
    ggplot(., aes(model, cyclomatic_complexity, fill = language, color = language)) +
    geom_boxplot(outlier.size = 1) +
    coord_flip() +
    scale_y_continuous(breaks = scales::breaks_pretty(n = 2)) +
    facet_wrap(~language) +
    labs(x = "", y = "C") +
    theme_AP() +
    scale_color_manual(values = color_languages) +
    theme(legend.position = "none")

plot_c_model
```



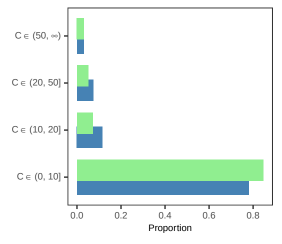
```
# Scatterplot cyclomatic vs lines of code ------

plot_c_vs_loc <- metrics_combined[grep("^func_", names(metrics_combined))] %>%
    lapply(., function(x) x[, .(loc, cyclomatic_complexity, language)]) %>%
    rbindlist() %>%
    ggplot(., aes(loc, cyclomatic_complexity, color = language)) +
    geom_point(alpha = 0.5, size = 0.7) +
    scale_x_continuous(breaks = breaks_pretty(n = 3)) +
    labs(x = "Lines of code", y = "C") +
    scale_color_manual(values = color_languages) +
    theme_AP() +
    theme(legend.position = "none")
```

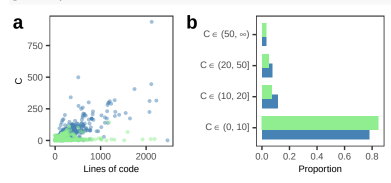


```
# Count & proportion -----
plot_bar_cyclomatic <- metrics_combined[grep("^func_", names(metrics_combined))] %>%
    lapply(., function(x) x[, .(complexity_category, language)]) %>%
    rbindlist() %>%
```

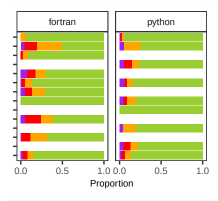
```
.[, .N, .(complexity_category, language)] %>%
.[, proportion := N / sum(N), language] %>%
ggplot(., aes(complexity_category, proportion, fill = language)) +
geom_bar(stat = "identity", position = position_dodge(0.6)) +
scale_fill_manual(values = color_languages) +
scale_y_continuous(breaks = scales::breaks_pretty(n = 4)) +
scale_x_discrete(labels = lab_expr) +
labs(x = "", y = "Proportion") +
coord_flip() +
theme_AP() +
theme(legend.position = "none")
```



plot cyclomatic



```
plot_bar_category <- metrics_combined[grep("^func_", names(metrics_combined))] %>%
  lapply(., function(x)
   x[, .(model, language, complexity_category)]) %>%
 rbindlist() %>%
  .[, .N, .(model, language, complexity category)] %>%
  .[, proportion := N / sum(N), .(language, model)] %>%
  ggplot(., aes(model, proportion, fill = complexity_category)) +
  geom_bar(stat = "identity") +
  scale_fill_manual(values = c("yellowgreen", "orange", "red", "purple"),
                    labels = lab_expr,
                    name = "") +
 facet_wrap(~language) +
  labs(x = "", y = "Proportion") +
  coord_flip() +
  scale_y_continuous(breaks = scales::breaks_pretty(n = 3)) +
  theme AP() +
  theme(legend.position = "none") +
  theme(axis.text.y = element_blank(),
        legend.text = element_text(size = 7))
plot_bar_category
```



```
di <- plot_grid(plot_scatterplot, plot_bar_cyclomatic, ncol = 1, labels = c("d", "e"))</pre>
```

```
legend <- get_legend_fun(plot_bar_category + theme(legend.position = "top"))

## Warning: `is.ggplot()` was deprecated in ggplot2 3.5.2.

## i Please use `is_ggplot()` instead.

## This warning is displayed once every 8 hours.

## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was

## generated.

dada <- plot_grid(plot_c_model, plot_bar_category, ncol = 2, rel_widths = c(0.61, 0.39))

dada2 <- plot_grid(legend, dada, ncol = 1, rel_heights = c(0.1, 0.9), labels = "f")

dada3 <- plot_grid(di, dada2, ncol = 2, rel_widths = c(0.4, 0.6))</pre>
```

dada4 <- plot\_grid(plot\_maintainability\_index, plot\_score, ncol = 2, labels = c("g", "h"))</pre> dada5 <- plot\_grid(p1, dada3, ncol = 1, rel\_heights = c(0.6, 0.4))</pre> plot\_grid(dada5, dada4, rel\_heights = c(0.73, 0.27), ncol = 1) 3 4 - 0 No models No 1 - 1 b python 0e+00 1e+05 2e+05 3e+05 0.2 0.4 0.6 0.2 0.4 0.6 Lines of code Comment density C lines\_code lines comments functions lines\_function modules files CTSM ORCHIDEE SWAT HYPE PCR-GLOBWB VIC SACRAMENTO HBV 0e+00 5e+04 1e+05 0 0e+00 2e+05 100 0 1000 500 1000 5000 0 50 0  $C \in (0, 10]$   $C \in (10, 20]$   $C \in (20, 50]$   $C \in (50, \infty)$ d fortran python 750 · 0 500 · 250 · fortran python python SWAT SACRAMENTO PCR-GLOBWB ORCHIDEE 1000 2000 1000 2000 0 Lines of code MHM HYPE HydroPy HBV H08 **e** C ∈ (50, ∞) - $C \in (20, 50]$  $C\in\left(10,\,20\right]$ CWatN C ∈ (0, 10] 0.2 0.4 0.6 500 0 500 0.0 0.5 1.00.0 0.5 1.0 Proportion Proportion С h g  $\mathsf{M}_{\mathsf{LOC}}$ Maverage fortran python SACRAMENTO -GR4J VIC SWAT SACRAMENTO PCR-GLOBWB ORCHIDEE MHM' HYPE HydroPy H08 GR4J DBH' CWatM CTSM VIC · SWAT MHM VIC CTSM · CTSM SWAT H08 DBH MHM H08 HydroPy HYPE -PCR-GLOBWB -HydroPy ORCHIDEE CWatM 75 100 125 50 75 100 125 Value Score