Measure 4: Savings potentials, Normalization, Proportionalization and Creation of Plots and Tables

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Code for the calculation of the savings potential using the results of the LCA-studies. Normalization and Proportionalization is performed as well as the creation of plots and tables. The code needs to be adjusted to the specific measure in the parameter section. This is the example for measure 2.

To execute, change:

- Login data for Nextcloud
- Adjust download location
- Adjust upload location
- Adjust URLs for uploading

Load Packages:

```
# install.packages("httr")
# install.packages("readxl")
# install.packages("dplyr")
# install.packages("writexl")
# install.packages("ggplot2")
# install.packages("tidyr")
# install.packages("openxlsx")
# install.packages("flextable")
# install.packages("officer")
library(readxl)
library(httr)
library(dplyr)
library(writexl)
library(ggplot2)
library(tidyr)
library(openxlsx)
library(flextable)
library(officer)
```

Parameters

Parameters to be adjusted:

```
measure <- "04" # "02", "03" or "04"
sheet <- 5 # 3,4 or 5
# name for columns
name_with <- "with MS"</pre>
name without <- "without MS"</pre>
name SP <- "SP MS"
name SP prop <- "prop. SP MS"
name_with_prop <- "prop. with MS"</pre>
name_without_prop <- "prop. without MS"</pre>
name with norm <- "norm. with MS"
name_without_norm <- "norm. without MS"</pre>
name_SP_norm <- "norm. SP MS"</pre>
# vector with excluded categories due to missing normalization factor
excluded categories of <- c("Climate change: biogenic",
                              "Climate change: fossil",
                              "Climate change: land use and land use change",
                              "Ecotoxicity: freshwater, inorganics",
                              "Ecotoxicity: freshwater, organics",
                              "Human toxicity: carcinogenic, inorganics",
                              "Human toxicity: carcinogenic, organics",
                              "Human toxicity: non-carcinogenic, inorganics",
                              "Human toxicity: non-carcinogenic, organics")
# vector with excluded toxicity categories for Pareto rule
excluded_categories_pr <- c("Human toxicity: carcinogenic",</pre>
                              "Human toxicity: non-carcinogenic",
                              "Ecotoxicity: freshwater")
# storage location
location_download <- "C:/Users/Klene/Documents/Uni_Bremen/WS24_25/Masterarbei</pre>
t/R/Daten/"
location upload <- "C:/Users/Klene/Documents/Uni Bremen/WS24 25/Masterarbeit/</pre>
R/Results/"
# Login data for NextCloud
username <- "mkindler@uni-bremen.de" # username</pre>
password <- "DLCi3-Qc4iD-dcHzR-fJaXr-cyKHR" # password</pre>
# WebDAV-URL to file
nextcloud url <- "https://nc.uni-bremen.de/remote.php/dav/files/mkindler%40un</pre>
i-bremen.de/Masterarbeit/Masterarbeit_II/R/Daten/Ergebnisse_LCA_basic.xlsx"
nextcloud url 2 <- "https://nc.uni-bremen.de/remote.php/dav/files/mkindler%40</pre>
uni-bremen.de/Masterarbeit/Masterarbeit II/R/Results/"
```

Data preparation

Load data from NextCloud:

```
# download file
response <- GET(nextcloud_url, authenticate(username, password, type = "basic
"))
# Ckeck successful downloading
if (status_code(response) == 200) {
    # Save file</pre>
```

```
writeBin(content(response, "raw"), paste0(location_download, "Ergebnisse_LC
A basic.xlsx"))
 cat("The file was successfully downloaded and saved locally.\n")
} else {
  cat("Error downloading the file. Status code: ", status code(response), "\n
}
## The file was successfully downloaded and saved locally.
# Load results data from Excel file
results <- read_excel(paste0(location_download, "Ergebnisse_LCA_basic.xlsx"),
sheet = sheet)
# Check data type: DataFrame
if (is.data.frame(results)) {
  results <- results
} else {
  results <- as.data.frame(results)</pre>
# Rename columns
results <- results %>%
  rename(
    `impact category` = `Wirkungskategorien`,
    unit = Einheit
    )
# Load normalization factors from Excel file
NF <- read_excel(paste0(location_download, "Ergebnisse_LCA_basic.xlsx"), shee
t = 1
## New names:
## • `` -> `...2`
## • `` -> `...3`
# Check data type: DataFrame
if (is.data.frame(NF)) {
  NF <- NF
} else {
  NF <- as.data.frame(NF)</pre>
# Set column names right
NF <- NF[-1, ]
colnames(NF) <- as.character(NF[1, ])</pre>
NF <- NF[-1, ]
# Select relevant columns
NF <- NF %>%
  select(`impact category`,
         NF = `normalization factor\r\n(Matuštík 2024)`)
```

Calculations

Savings potential

Calculation of Savings Potentials:

```
results <- results %>%
  mutate(!!sym(name_SP) := !!sym(name_without) - !!sym(name_with))
```

Normalization

Calculation of normalized values (to the planetary boudaries (Matuštík 2024)):

Determine relevant impact categories

Calculation of proportion of total savings potential of each measure:

```
# Removing the toxicity categories
results_prop <- results_norm %>%
    filter(!`impact category` %in% excluded_categories_pr)

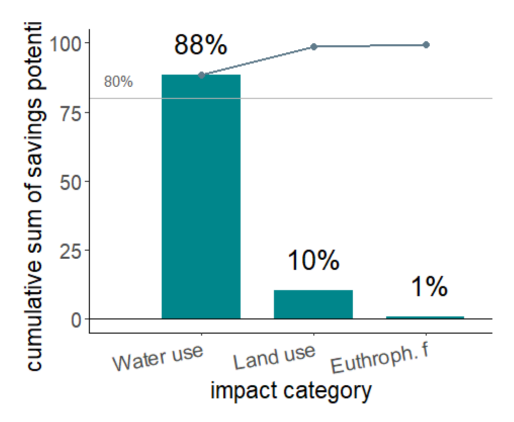
# Calculate the sum of each normalized column and divide the values by this s
um
results_prop <- results_prop %>%
    mutate(
    !!sym(name_without_prop) := !!sym(name_without_norm) / sum(abs(!!sym(name_without_norm)), na.rm = TRUE),
    !!sym(name_with_prop) := !!sym(name_with_norm) / sum(abs(!!sym(name_with_norm)), na.rm = TRUE),
    !!sym(name_SP_prop) := !!sym(name_SP_norm) / sum(abs(!!sym(name_SP_norm)))
na.rm = TRUE)
)
norm sum <- sum(abs(results_prop[[name_SP_norm]]))</pre>
```

Determine relevant impact categories:

Sort categories by proportion of total savings potential

```
results prop <- results prop %>%
  arrange(desc(abs(!!sym(name_SP_prop))))
# Calculation of the cumulative total and mark categories that contribute to
the sum of 0.8
results prop <- results prop %>%
  mutate(
    `cum. sum` = cumsum(abs(!!sym(name SP prop))),
    relevant = (`cum. sum` < 0.8) | lag(`cum. sum` < 0.8, default = FALSE)
# Mark at least the highest three impact categories
results prop$relevant[1:3] <- TRUE
Save as csv-file
write.csv(results prop, paste0(location upload, measure, " data.csv"), row.na
mes = FALSE)
Plot
Plot the relevant impact categories
# Ensure that the impact category retains the order in which it appears in th
e data
results_prop$`impact category` <- factor(results_prop$`impact category`, leve</pre>
ls = results prop$`impact category`)
# Prepare data
data plot <- results prop %>%
  filter(relevant == TRUE) %>%
  mutate(
    across(.cols = starts with("prop."), .fns = ~ . * 100),
    `cum. sum` = `cum. sum`*100) %>%
  select(`impact category`, name_SP_prop, `cum. sum`)
# Rename Long impact categories
data plot <- data plot %>%
  mutate(`impact category` = case_when(
    `impact category` == "Photochemical oxidant formation: human health" ~ "P
OF",
    `impact category` == "Eutrophication: freshwater" ~ "Euthroph. f",
    TRUE ~ `impact category`
  ))
# Ensure that the impact category retains the order in which it appears in th
e data
data_plot$`impact category` <- factor(data_plot$`impact category`, levels = d</pre>
ata plot$`impact category`)
```

```
# Plot relvant impact categories with their proportion of the total savings p
otential, kumulative sum
ggplot(data_plot, aes(x = `impact category`, y = !!sym(name_SP_prop))) +
  geom_bar(stat = "identity", width = 0.7, fill = "turquoise4", show.legend =
FALSE) +
  geom line(aes(y = `cum. sum`, group = 1), color = "lightskyblue4", size = 1
) +
  geom_point(aes(y = `cum. sum`), color = "lightskyblue4", size = 2) +
  geom_hline(yintercept = 80, color = "grey70", linetype = "solid") +
  geom hline(yintercept = 0, color = "black", linetype = "solid") +
  annotate("text", x = 0, y = 80, label = "80%", color = "grey35", size = 4,
hjust = -0.5, vjust = -1) +
  geom_text(aes(
    y = !!sym(name_SP_prop),
    label = ifelse(data_plot$`impact category` == "POF", paste0(format(round()))
!!sym(name_SP_prop), 1), nsmall = 0), "%"), paste0(format(round(!!sym(name_SP_
_prop), 0), nsmall = 0), "%"))
  ), vjust = -1, size = 7) +
  labs(
   x = "impact category",
   y = "cumulative sum of savings potentials"
  theme classic() +
  theme(
    axis.text.x = element_text(angle = 10, hjust = 1, size = 15),
    axis.text.y = element text(size = 15),
    axis.title.x = element_text(size = 17),
    axis.title.y = element text(size = 17)
 coord_cartesian(ylim = c(NA, 100))
```

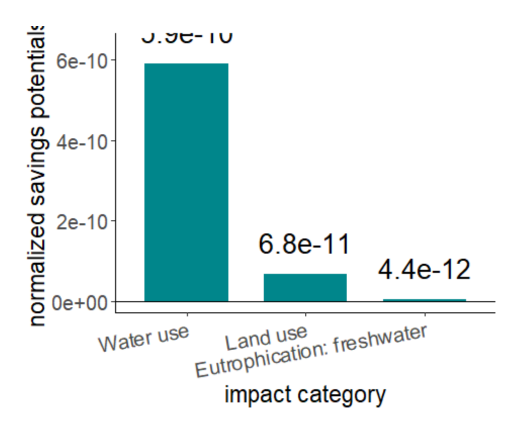


```
# Save plot
ggsave(paste0(location_upload, measure, "_relevant_SP_prop.png"), dpi = 600)
## Saving 5 x 4 in image
```

Plot relevant categories with their norm. savings potential:

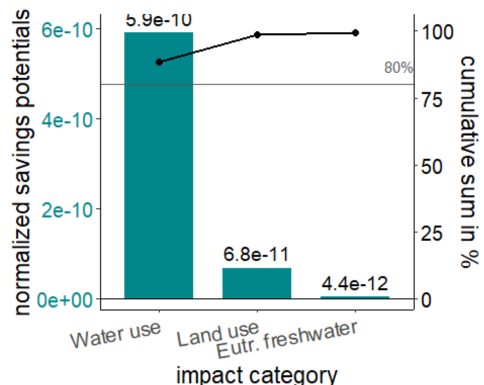
```
# Ensure that the impact category retains the order in which it appears in th
e data
results_norm$`impact category` <- factor(results_norm$`impact category`, leve</pre>
ls = results_norm$`impact category`)
results_relevant <- results_prop %>%
  select(`impact category`, relevant)
results_norm <- left_join(results_norm, results_relevant, by = c("impact cate
gory" = "impact category"))
# Prepare data
data_plot <- results_norm %>%
  filter(relevant == TRUE) %>%
  select(`impact category`, name_SP_norm) %>%
  arrange(desc(abs(!!sym(name_SP_norm))))
## Warning: Using an external vector in selections was deprecated in tidysele
ct 1.1.0.
## i Please use `all_of()` or `any_of()` instead.
     # Was:
```

```
##
     data %>% select(name_SP_norm)
##
##
     # Now:
##
     data %>% select(all of(name SP norm))
##
## See <https://tidyselect.r-lib.org/reference/faq-external-vector.html>.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
# Rename Long impact categories
data plot <- data plot %>%
  mutate(`impact category` = case_when()
    `impact category` == "Photochemical oxidant formation: human health" ~ "P
OF",
   TRUE ~ `impact category`
  ))
# Ensure that the impact category retains the order in which it appears in th
e data
data_plot$`impact category` <- factor(data_plot$`impact category`, levels = d</pre>
ata_plot$`impact category`)
ggplot(data_plot, aes(x = `impact category`, y = !!sym(name_SP_norm))) +
  geom_bar(stat = "identity", width = 0.7, fill = "turquoise4", show.legend =
FALSE) +
  geom hline(yintercept = 0, color = "black", linetype = "solid") +
  geom text(aes(
    y = !!sym(name_SP_norm),
    label = format(!!sym(name_SP_norm), scientific = TRUE, digits = 2)
    ), vjust = -1, size = 7) +
  labs(
    x = "impact category",
   y = "normalized savings potentials"
  ) +
  theme classic() +
  theme(
    axis.text.x = element_text(angle = 10, hjust = 1, size = 15),
    axis.text.y = element text(size = 15),
    axis.title.x = element_text(size = 17),
    axis.title.y = element text(size = 17)
  scale y continuous(expand = expansion(mult = c(0.05, 0.13)))
```



```
# Save plot
ggsave(paste0(location_upload, measure, "_relevant_SP_norm.png"), dpi = 600)
## Saving 5 x 4 in image
# Rename Long impact categories
data_plot <- data_plot %>%
  mutate(`impact category` = recode(`impact category`,
    "Photochemical oxidant formation: human health" = "POF",
    "Eutrophication: freshwater" = "Eutr. freshwater"
  ))
# create cumsum and arrange descending
data_plot <- data_plot %>%
  arrange(desc(abs(!!sym(name_SP_norm)))) %>%
    cumsum = cumsum(abs(!!sym(name_SP_norm) / norm_sum)) * 100
  )
# max. values for scaling
max_norm <- max(data_plot[[name_SP_norm]])</pre>
max_cumsum <- max(data_plot$cumsum)</pre>
# Conversion factor
factor <- max_norm / max_cumsum</pre>
ggplot(data_plot, aes(x = `impact category`)) +
  geom_bar(aes(y = !!sym(name_SP_norm)), stat = "identity",
           width = 0.7, fill = "turquoise4", show.legend = FALSE) +
```

```
geom_line(aes(y = cumsum * factor, group = 1), color = "black", size = 1) +
  geom_point(aes(y = cumsum * factor), color = "black", size = 2) +
  geom_hline(yintercept = 0, color = "black", linetype = "solid") +
  geom_hline(yintercept = 80 * factor, color = "grey35", linetype = "solid")
  annotate("text", x = 3.3, y = 80 * factor, label = "80%", color = "grey35",
size = 4, hjust = 0, vjust = -1) +
  geom_text(aes(
    y = !!sym(name SP norm),
    label = format(!!sym(name_SP_norm), scientific = TRUE, digits = 2)),
    vjust = -0.5, size = 5) +
  scale_y_continuous(
    name = "normalized savings potentials",
    sec.axis = sec_axis(~ . / factor, name = "cumulative sum in %"),
  expand = expansion(mult = c(0.04, 0.07))
  labs(x = "impact category") +
  theme_classic() +
  theme(
    axis.text.x = element_text(angle = 10, hjust = 1, size = 15),
    axis.text.y.left = element_text(color = "turquoise4", size = 15),
    axis.text.y.right = element_text(color = "black", size = 15),
    axis.title.x = element_text(size = 17),
    axis.title.y.left = element_text(size = 17),
    axis.title.y.right = element_text(size = 17)
```



```
# Save plot
ggsave(paste0(location_upload, measure, "_relevant_SP_norm_allin1.png"), dpi
= 600)
## Saving 5 x 4 in image
```

Tables

Prepare data:

```
# Select relevant columns
results prop <- results prop %>%
  select(`impact category`, unit, !!sym(name_without_prop), !!sym(name with p
rop),
         !!svm(name SP prop))
results norm <- results norm %>%
  select(`impact category`, unit, !!sym(name without norm), !!sym(name with n
orm),
         !!sym(name_SP_norm))
# Join of 'results' and 'results norm' according to the impact categories
results <- left join(results, results norm, by = c("impact category" = "impac
t category"))
results <- results %>%
  select(-unit.y) %>%
  rename(unit = unit.x)
# Join of 'results' and 'results prop' according to the impact categories
results <- left_join(results, results_prop, by = c("impact category" = "impac
t category"))
results <- results %>%
  select(-unit.y) %>%
  rename(unit = unit.x)
# Sort descending according to normalized savings potential
results <- results %>%
  arrange(desc(abs(!!sym(name_SP_norm))))
```

Create tables:

```
# Only results
table_results <- results %>%
    select("impact category", "unit", !!sym(name_without), !!sym(name_with), !!
sym(name_SP)) %>%
    arrange(desc(abs(!!sym(name_SP)))) %>%
    mutate(
        across(-all_of(c("impact category", "unit")), ~ formatC(.x, format = "e",
digits = 2)))
# Create flextable
ft <- flextable(table_results) %>%
    theme_vanilla() %>%
    autofit() %>%
    bold(part = "header") %>%
    border_outer() %>%
    border_inner_h() %>%
```

```
border_inner_v()
# Save as PNG
tf <- tempfile(fileext = ".png")</pre>
save_as_image(x = ft, path = paste0(location_upload, measure, "_results", ".p
ng"))
## [1] "C:/Users/Klene/Documents/Uni_Bremen/WS24_25/Masterarbeit/R/Results/04
results.png"
# Saving potentials
table results <- results %>%
  select(`impact category`, unit, !!sym(name_SP), !!sym(name_SP_norm), !!sym
(name SP prop)) %>%
  arrange(desc(abs(!!sym(name SP norm)))) %>%
  mutate(
    across(-all_of(c("impact category", "unit")), ~ formatC(.x, format = "e",
digits = 2)))
# Create flextable
ft <- flextable(table results) %>%
  theme vanilla() %>%
  autofit() %>%
  bold(part = "header") %>%
  border_outer() %>%
  border_inner_h() %>%
  border inner v()
# Save as PNG
tf <- tempfile(fileext = ".png")</pre>
save as image(x = ft, path = paste0(location upload, measure, " results SPs",
".png"))
## [1] "C:/Users/Klene/Documents/Uni Bremen/WS24 25/Masterarbeit/R/Results/04
results SPs.png"
# Normalized results
table results <- results %>%
  select("impact category", "unit", !!sym(name_without_norm), !!sym(name with
norm), !!sym(name SP norm)) %>%
  arrange(desc(abs(!!sym(name_SP_norm)))) %>%
  mutate(
    across(-all_of(c("impact category", "unit")), ~ formatC(.x, format = "e",
digits = 2)))
# Removing the categories without normalization factors
table results <- table results %>%
  filter(!`impact category` %in% excluded_categories_nf)
# Rename columns
table results <- table results %>%
  rename(!!sym(name_without) := !!sym(name_without_norm),
           !!sym(name_with) := !!sym(name_with_norm),
           !!sym(name_SP) := !!sym(name_SP_norm)
           )
# Create flextable
ft <- flextable(table results) %>%
```

```
theme_vanilla() %>%
  autofit() %>%
  bold(part = "header") %>%
  border outer() %>%
  border inner h() %>%
  border_inner_v()
# Save as PNG
tf <- tempfile(fileext = ".png")</pre>
save_as_image(x = ft, path = paste0(location_upload, measure, "_results_norm"
, ".png"))
## [1] "C:/Users/Klene/Documents/Uni Bremen/WS24 25/Masterarbeit/R/Results/04
results norm.png"
# Proportion data
table results <- results %>%
  select("impact category", "unit", !!sym(name_without_prop), !!sym(name_with
_prop), !!sym(name_SP_prop)) %>%
  arrange(desc(abs(!!sym(name SP prop)))) %>%
  mutate(
    across(-all_of(c("impact category", "unit")), ~ formatC(.x, format = "e",
digits = 2)))
# Removing the categories without normalization factor
table results <- table results %>%
  filter(!`impact category` %in% excluded categories nf)
# Removing the toxicity categories
table_results <- table_results %>%
  filter(!`impact category` %in% excluded categories pr)
# Rename columns
table results <-table results %>%
  rename(!!sym(name without) := !!sym(name without prop),
         !!sym(name with) := !!sym(name with prop),
         !!sym(name_SP) := !!sym(name_SP_prop),
# Create flextable
ft <- flextable(table_results) %>%
  theme vanilla() %>%
  autofit() %>%
  bold(part = "header") %>%
  border outer() %>%
  border inner h() %>%
  border inner v()
# Save as PNG
tf <- tempfile(fileext = ".png")</pre>
save_as_image(x = ft, path = paste0(location_upload, measure, "_results_prop"
, ".png"))
## [1] "C:/Users/Klene/Documents/Uni Bremen/WS24 25/Masterarbeit/R/Results/04
_results_prop.png"
```

Upload files

```
# # List of Excel files to be uploaded
# files_to_upload <- list.files(path = location_upload, pattern = "\\.xlsx$",
full.names = TRUE)
# # Loop over all files and upload
# for (file path in files to upload) {
  # Creating the file name
  file_name <- basename(file_path)</pre>
#
  # Upload file
#
  response <- PUT(
      url = paste0(nextcloud_url_2, file_name),
#
      authenticate(username, password),
#
#
      body = upload_file(file_path)
#
#
   # Check upload status
   if (status code(response) == 201) {
     print(paste(file_name, "was successfully uploaded."))
#
    } else if (status_code(response) == 204) {
#
      print(paste(file_name, "was successfully replaced (no content returned)
."))
#
   } else {
      print(paste("Error uploading", file_name, ". Status-Code:", status_code
(response)))
#
    }
# }
#
# # List of PNG files to be uploaded
# files_to_upload <- list.files(path = location_upload, pattern = "\\.png$",
full.names = TRUE)
# # Loop over all files and upload
# for (file_path in files_to_upload) {
  # Create the file name
# file name <- basename(file path)</pre>
# # Upload file
#
  response <- PUT(
#
      url = paste0(nextcloud_url_2, file_name),
      authenticate(username, password),
#
#
      body = upload file(file path)
#
  # Check upload status
   if (status code(response) == 201) {
      print(paste(file_name, "was successfully uploaded."))
#
    } else if (status_code(response) == 204) {
#
      print(paste(file name, "was successfully replaced (no content returned)
."))
#
    } else {
      print(paste("Error uploading", file_name, ". Status-Code:", status_code
(response)))
    }
# }
```