Mental health and subjective well-being of trans and non-binary population in Colombia

Supplementary Material: Code and analyses

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Description

This document contains all code, and step by step explanations for all analyses, figures and tables (including supplementary figures and tables) for:

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Data are available on the Open Science Framework (OSF): https://doi.org/10.17605/OSF.IO/HJ7VB. The analyses were designed by Juan David Leongómez and María Fernanda Reyes and performed by Juan David Leongómez. This document and all its underlying code were created in R Markdown by Juan David Leongómez using R and LATEX.

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1 Preliminaries

1.1 Load packages

This file was created using knitr (Xie, 2014), mostly using tidyverse (Wickham et al., 2019) syntax. As such, data wrangling was mainly done using packages such as dplyr (Wickham et al., 2023), and most figures were created or modified using ggplot2 (Wickham, 2016). Tables were created using knitr::kable and kableExtra (Zhu, 2021).

Multi-model inference and model averaging was achieved using MuMIn (Bartoń, 2025), and model assumptions were performed using performance (Lüdecke et al., 2021).

All packages used in this file can be directly installed from the Comprehensive R Archive Network (CRAN). For a complete list of packages used to create this file, and their versions, see section 6, at the end of the document.

```
library(ltm)
library(psych) # For statistical functions (e.g., Cronbach's alpha)
library(MuMIn) # For model selection and averaging
library(performance) # For model performance metrics
library(readr) # For reading data files
library(scales) # For percent formatting
library(knitr)
library(kableExtra)
library(car)
library(tidyverse) # For data manipulation and piping
library(ggpubr)
library(gtsummary)
library(Hmisc)
library(magick)
library(ggplotify)
library(pdftools)
```

1.2 Custom functions

1.2.1 pval.lev and pval.stars

These functions take p-values and formats them, either in LaTeX and highlighting significant p-values in bold and representing all in an appropriate level, or as stars.

```
# Function to format p-values for LaTeX output, highlighting significant values in bold
pval.lev <- function(pvals) {
  ifelse(pvals < 0.0001, "\\textbf{< 0.0001}", # Highlight very small p-values
   ifelse(pvals < 0.001, "\\textbf{< 0.001}", # Bold p-values < 0.001
      ifelse(pvals < 0.05, paste0("\\textbf{", round(pvals, 4), "}"), # Bold p-values < 0.05
      round(pvals, 2) # Round non-significant values to two decimal places
   )
  )
}

# Function to add significance stars based on p-value thresholds
pval.stars <- function(pvals) {
  ifelse(pvals < 0.0001, "****", # Four stars for p < 0.0001
   ifelse(pvals < 0.001, "***", # Three stars for p < 0.001
   ifelse(pvals < 0.01, "**", # Two stars for p < 0.01
   ifelse(pvals < 0.05, "*", NA) # One star for p < 0.05, NA otherwise
  )
  )
}</pre>
```

1.2.2 corr.stars

This function creates a correlation matrix, and displays significance (function corr.stars modified from http://myowelt.blogspot.com/2008/04/beautiful-correlation-tables-in-r.html).

```
# Function to create a correlation matrix with significance levels in LaTeX format
corr.stars <- function(x) {
  require(Hmisc) # Load Hmisc package for correlation and p-value calculations
  x <- as.matrix(x) # Ensure input is a matrix
  R <- rcorr(x, type = "spearman")$r # Compute correlation coefficients
  p <- rcorr(x, type = "spearman")$P # Extract p-values for significance testing
  # Define symbols for significance levels, using LaTeX formatting for bold and stars</pre>
```

```
mystars <- ifelse(p < .001, paste0("\\textbf{", round(R, 2), "***}"), # p < 0.001
    ifelse(p < .05, paste0("\\textbf{", round(R, 2), "**}"), # p < 0.01
    ifelse(p < .05, paste0("\\textbf{", round(R, 2), "**}"), # p < 0.05
    ifelse(p < .10, paste0(round(R, 2), "$^{\\dagger}$"), # p < 0.10 (trend level)
        format(round(R, 2), nsmall = 2) # Format non-significant values with two decimals
    )
    )
    )
    # Construct a new matrix with correlation values and significance symbols
    Rnew <- matrix(mystars, ncol = ncol(x))
    # Ensure diagonal values remain the original correlation values (without significance symbols)
    diag(Rnew) <- paste(diag(R), " ", sep = "")
    # Assign row and column names for the formatted matrix
    rownames(Rnew) <- colnames(x)
    colnames(Rnew) <- paste(colnames(x), "", sep = "")
    # Remove the upper triangle of the matrix (including the diagonal) for a clean presentation
    Rnew <- as.matrix(Rnew)
    Rnew[upper.tri(Rnew, diag = TRUE)] <- ""
    # Convert to a data frame for better handling and remove the last empty column
    Rnew <- as.data.frame(Rnew)
    Rnew <- cbind(Rnew[1:length(Rnew) - 1])
    return(Rnew) # Return formatted correlation table
}</pre>
```

1.2.3 dredge.plot

This function generates a model selection plot from a MuMIn::dredge() model selection table. It visualizes the top 100 candidate models, showing which terms are included in each model and their relative importance based on Akaike weights. The function also dynamically creates a caption summarizing the number of fitted models, suitable for use as a LaTeX or R Markdown fig.cap. Additionally, it saves the plot as a PDF file in a PDF_plots/subdirectory (relative to the project root), with the filename based on the input object's name

```
dredge.plot <- function(mod.sel.table) {</pre>
  # Ensure the input is a dredge object
  if (!inherits(mod.sel.table, "model.selection")) {
    stop("Input must be a MuMIn::dredge() model selection table.")
  model_call <- attr(mod.sel.table, "model.calls")[[1]]</pre>
  response_var <- all.vars(model_call$formula)[1]</pre>
  response_label <- gsub("_", " ", response_var)</pre>
  dir.create("PDF_plots", showWarnings = FALSE)
  # Define output file path
  file_path <- file.path("PDF_plots", pasteO(response_var, ".pdf"))</pre>
  pdf(file_path, width = 8, height = 6)
  plot_object <- plot(mod.sel.table[1:100, ], col = "#A92395FF",</pre>
                       main = NULL)
  dev.off()
  plot(mod.sel.table[1:100, ], col = "#A92395FF",
       main = NULL, par.vlab = NULL)
```

```
caption <- paste0(
   "Model selection plot. Each row represents one of the top 100 models ",
   "(ranked by $AICc$) out of ", formatC(nrow(mod.sel.table), big.mark = ","),
   " fitted models. Purple cells indicate included terms, with row height corresponding",
   " to each model's relative support based on its Akaike weight ($\\\omega$)"
)
# Return both the plot object and the caption
return(list(plot = plot_object, caption = caption))
}</pre>
```

1.2.4 avg.mod.summary

This function creates a formatted summary table for model-averaged coefficients obtained from MuMIn::model.avg(). It includes coefficient estimates and 95% confidence intervals, significance levels (formatted for LaTeX output), a dynamically generated caption including the dependent variable name, and a footnote specifying reference levels for categorical predictors.

```
avg.mod.summary <- function(avg_model, data) {</pre>
  # Function to extract relevant reference levels
 extract_relevant_reference_levels <- function(avg_model, data) {</pre>
   factor_vars <- names(data)[sapply(data, is.factor)]</pre>
    ref_levels <- sapply(factor_vars, function(var) levels(data[[var]])[1])</pre>
   model_terms <- rownames(summary(avg_model)$coefmat.full)</pre>
    relevant_factors <-
      factor_vars[sapply(factor_vars,
                         function(var) any(grepl(paste0("^", var), model_terms)))]
    relevant_refs <- ref_levels[names(ref_levels) %in% relevant_factors]
    if (length(relevant_refs) > 0) {
      paste0(
        "Reference levels: ",
        pasteO(names(relevant_refs),
               " = ", relevant_refs,
               collapse = "; "
   } else {
      "No factor variables in the average model"
 cis <- confint(avg_model, full = TRUE) |>
   as.data.frame() |>
   mutate(across(everything(), round, 3)) |>
    unite(col = "CI", `2.5 %`: `97.5 %`, sep = " - ")
  # Full average summary table
  summ_table <- summary(avg_model)$coefmat.full |>
    as.data.frame() |>
   bind_cols(cis) |>
    rownames_to_column(var = "term") |>
```

```
mutate(term = term |>
           str_replace_all("Gender", "Gender: ") |>
           str_replace_all("Housing", "Housing: ") |>
           str_replace_all("Ethnicity", "Ethnicity: ") |>
           str replace all("Education", "Education: ") |>
           str_replace_all("Discrimination", "Discrimination: ") |>
           str_replace_all("Self_efficacy", "Self-efficacy") |>
           str_replace_all("_", " ")) |>
  select(term, Estimate, CI, `z value`, `Pr(>|z|)`) |>
  mutate(`Pr(>|z|)` = pval.lev(`Pr(>|z|)`)) |>
  arrange(term)
kable_tab <- kable(</pre>
  summ_table,
  digits = 3,
 booktabs = TRUE,
  linesep = "",
  align = c("l", rep("c", 4)),
  caption = paste0(
   as.character(avg_model$formula)[2] |>
      str_replace_all("_", " "),
  col.names = c("Term", "$B$", "$95 \\% CIs$", "$z$", "$p$"),
  escape = FALSE
  kable_styling(latex_options = "HOLD_position") |>
  footnote(
    general = paste0(
      "The full model average ensures that all variables are included in every model, ",
      "$B$ represents unstandardized model coefficients. ",
      extract_relevant_reference_levels(avg_model, data), # Now using the internal function
    threeparttable = TRUE,
    footnote_as_chunk = TRUE,
    escape = FALSE
return(list(table = kable_tab,
            coefficients = summ_table,
            reference_levels = extract_relevant_reference_levels(avg_model, data)))
```

1.2.5 avg.mod.plot

This function visualizes model-averaged estimates from MuMIn::model.avg(), displaying point estimates with 95% confidence intervals, predictor importance (mapped to point size and colour), and a dynamically generated annotation for the number of averaged model.

```
avg.mod.plot <- function(avg_mod) {</pre>
 x <- summary(avg_mod)$coefmat.full |>
   as_tibble(rownames = "key") |> # Convert row names to a "key" column
   bind cols(
     confint(avg_mod, full = TRUE) |> as_tibble(), # Add confidence intervals
     summary(avg_mod)$coef.nmod |>
       as_tibble() |>
        pivot_longer(cols = everything(), names_to = "model", values_to = "value")
   mutate(
     avmod = deparse(substitute(avg_mod)) |>
        factor(), # Store model name as a factor
     value = value / max(value, na.rm = TRUE), # Normalize 'value' column
     sig = pval.stars(`Pr(>|z|)`) |>
        str_replace("\\.", "†"), # Convert p-values into significance stars
     key = key |>
       str_replace_all("Gender", "Gender: ") |>
        str_replace_all("Housing", "Housing: ") |>
        str_replace_all("Ethnicity", "Ethnicity: ") |>
        str_replace_all("Education", "Education: ") |>
        str_replace_all("Discrimination", "Discrimination: ") |>
        str_replace_all("Self_efficacy", "Self-efficacy") |>
        str_replace_all("_", " ")
 x <- x |>
   mutate(key = factor(key, levels = as.character(unique(x$key))))
  nMods <- dim(avg_mod$msTable)[1]
 ggplot(x, aes(x = key, y = Estimate)) +
   geom_hline(yintercept = 0, color = "grey") +
   geom_point(aes(size = value, color = value), alpha = 0.7) +
   geom_errorbar(aes(ymin = ^2.5 \%, ymax = ^97.5 \%),
                 colour = "black", width = 0.1
   geom_point(size = 1) +
   theme_bw() +
   labs(x = NULL, y = "Estimate") +
    theme(axis.text.x = element_text(angle = 60, hjust = 1)) +
   scale_size_continuous(
     range = c(2, 8),
     breaks = seq(0, 1, by = 0.2)
    guides(
```

```
size = guide_legend(title = "Importance"),
  color = guide_legend(title = "Importance")
scale x discrete(
  labels = levels(x$key),
  expand = c(0, 0.5)
scale_colour_viridis_c(
  option = "plasma", direction = -1,
  breaks = seq(0, 1, by = 0.2)
geom_text(aes(label = sig), y = x$^97.5 %, vjust = -0.4) +
annotate(
  "text",
 x = Inf, y = Inf,
 label = paste("Models averaged = ", nMods),
 hjust = 1.1, vjust = 1.8,
  size = 4,
  fontface = "bold.italic"
```

1.3 Load data

Load raw CSV data

```
data_RAW <- read_csv("data/data.csv")</pre>
```

1.3.1 Define PANAS Subscales (Positive & Negative Affect)

The Positive and Negative Affect Schedule (PANAS) is a widely used self-report questionnaire that measures two independent dimensions of affect:

- Positive Affect (PA): Reflects the extent to which a person feels enthusiastic, active, and alert.
- Negative Affect (NA): Represents distress and unpleasurable engagement, including emotions such as fear, anger, and nervousness.

The following lists define the PANAS subscales based on their respective items.

```
# List of PANAS Positive Affect (PANAS_P) items
PANAS_P <- c(
    "PANASB_1", "PANASB_3", "PANASB_5", "PANASB_9",
    "PANASB_10", "PANASB_12", "PANASB_14", "PANASB_16",
    "PANASB_17", "PANASB_19"
)

# List of PANAS Negative Affect (PANAS_N) items
PANAS_N <- c(
    "PANASB_2", "PANASB_4", "PANASB_6", "PANASB_7",
    "PANASB_8", "PANASB_11", "PANASB_13", "PANASB_15",
    "PANASB_18", "PANASB_20"
)</pre>
```

1.4 Internal consistency

1.4.1 Calculate Cronbach's Alpha for Different Scales

To measure the internal consistency of these tests, we used standardized Cronbach's alpha (α or Tau-equivalent reliability: ρ_T) coefficients, using the function cronbach.alpha from the package ltm (Rizopoulos, 2006).

```
# Compute Cronbach's alpha for the Self-efficacy (EAG) scale
alpha_EAG <- data_RAW |>
 mutate(across(where(is.numeric), ~ na if(., 99))) |> # Replace 99 with NA (missing values)
  select(starts_with("EAG_")) |> # Select all columns starting with "EAG_"
  drop na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE) # Compute Cronbach's alpha
alpha_SWLS <- data_RAW |>
  mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
  select(starts_with("SWLS_")) |> # Select all columns starting with "SWLS_"
 drop_na() |>
 cronbach.alpha(CI = TRUE, standardized = TRUE)
# Compute Cronbach's alpha for the Resilience (EBR) scale
alpha EBR <- data RAW |>
 mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
  select(starts_with("EBR_")) |> # Select all columns starting with "EBR_"
 drop_na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE)
alpha_EBD <- data_RAW |>
 mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
  select(starts_with("EBD_")) |> # Select all columns starting with "EBD_"
 mutate(across(everything(), ~ ifelse(is.na(.x), NA, .x - 1))) |> # Adjust values
 drop_na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE)
alpha_MOS2 <- data_RAW |>
 mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
  select(starts_with("MOS2_")) |> # Select all columns starting with "MOS2_"
  drop_na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE)
alpha_PANAS_P <- data_RAW |>
 mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
  select(all_of(PANAS_P)) |> # Select PANAS_P variables
 drop_na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE)
# Compute Cronbach's alpha for PANAS Negative Affect (PANAS_N)
alpha_PANAS_N <- data_RAW |>
 mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
 select(all_of(PANAS_N)) |> # Select PANAS_N variables
 drop_na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE)
```

```
alpha_PCPS3 <- data_RAW |>
  mutate(across(where(is.numeric), ~ na_if(., 99))) |> # Replace 99 with NA
  select(starts_with("PCPS3_")) |> # Select all columns starting with "PCPS3_"
  drop_na() |>
  cronbach.alpha(CI = TRUE, standardized = TRUE)
```

1.4.2 Table S1. Internal consistency of measured scales

The internal consistency of the measured scales was generally strong, with Cronbach's α values ranging from 0.767 to 0.977. In particular, the Social Support (MOS2) and Self-efficacy (EAG) scales exhibited excellent internal consistency, while the Depression (EBD) and Community Cohesion (PCPS3) scales had acceptable reliability, suggesting a slightly lower but still adequate level of internal consistency.

```
tibble(
 Scale = c(
   "Self-efficacy$^1$",
   "Life Satisfaction$^2$",
    "Resilience$^3$",
    "Depression$^4$"
   "Social Support$^5$",
   "PANAS Positive$^6$",
    "PANAS Negative$^6$",
    "Community Cohesion$^7$"
   alpha_EAG$p, alpha_SWLS$p, alpha_EBR$p, alpha_EBD$p, alpha_MOS2$p,
   alpha_PANAS_P$p, alpha_PANAS_N$p, alpha_PCPS3$p
 n = c(
    alpha_EAG$n, alpha_SWLS$n, alpha_EBR$n, alpha_EBD$n, alpha_MOS2$n,
   alpha_PANAS_P$n, alpha_PANAS_N$n, alpha_PCPS3$n
 alpha = c(
    alpha_EAG$alpha, alpha_SWLS$alpha, alpha_EBR$alpha, alpha_EBD$alpha,
    alpha_MOS2$alpha, alpha_PANAS_P$alpha, alpha_PANAS_N$alpha, alpha_PCPS3$alpha
  ),
  ci2.5 = c(
    alpha_EAG$ci[1], alpha_SWLS$ci[1], alpha_EBR$ci[1], alpha_EBD$ci[1],
   alpha_MOS2$ci[1], alpha_PANAS_P$ci[1], alpha_PANAS_N$ci[1], alpha_PCPS3$ci[1]
 ci97.5 = c(
    alpha_EAG$ci[2], alpha_SWLS$ci[2], alpha_EBR$ci[2], alpha_EBD$ci[2],
    alpha_MOS2$ci[2], alpha_PANAS_P$ci[2], alpha_PANAS_N$ci[2], alpha_PCPS3$ci[2]
 mutate(across(starts_with("ci"), round, 3)) |>
 unite(col = "CI", ci2.5:ci97.5, sep = " - ") |>
 kable(
   digits = 3,
   booktabs = TRUE,
   linesep = "",
   align = c("l", rep("c", 4)),
    caption = "Internal consistency of measured scales",
   col.names = c("Variable", "Items", "$n$", "$\\alpha$", "$95\\% CI$"),
    escape = FALSE
```

Table S1. Internal consistency of measured scales

Variable	Items	n	α	95%CI
Self-efficacy ¹	10	223	0.905	0.874 - 0.929
Life Satisfaction ²	5	253	0.869	0.838 - 0.896
$Resilience^3$	4	278	0.861	0.822 - 0.892
$Depression^4$	7	223	0.767	0.712 - 0.812
Social Support ⁵	19	195	0.977	0.971 - 0.982
PANAS Positive ⁶	10	285	0.884	0.857 - 0.905
PANAS Negative ⁶	10	282	0.827	0.795 - 0.866
Community Cohesion ⁷	3	281	0.769	0.704 - 0.822

Note: 95% confidence intervals were calculated with 1,000 bootstrap samples. Standardized Cronbach's alpha (α) coefficients were computed. ¹Baessler and Schwarzer, 1996; ²Diener et al., 1985; ³Sinclair and Wallston, 2004; ⁴Andresen et al., 1994; ⁵Sherbourne and Stewart, 1991; ⁶Watson et al., 1988; ⁷Stanton et al., 2017.

2 Data Preprocessing

2.1 Renaming, recoding, and filtering

This section contains the code to load the raw dataset (data_RAW) and create a final dataset (data) for analysis. Steps include:

- Renaming variables to meaningful labels
- Handling missing values (99 \rightarrow NA)
- Recoding categorical variables into more readable labels
- Estimating participant scores for item-based scales (Self-efficacy, Life Satisfaction, Resilience, Depression, Social Support, Positive Affect, Negative Affect, Community Cohesion)
- Filtering out ineligible cases (e.g., participants under 18)

To estimate participant scores for item-based scales (i.e., Self-efficacy, Life Satisfaction, Resilience, Depression, Social Support, Positive Affect, Negative Affect, and Community Cohesion), and to minimise missing data in multimodel inference procedures, we computed scores in accordance with recommended practices (see Graham, 2012), using Cronbach's alpha levels to guide acceptable thresholds for missing data. For scales with Cronbach's alpha values ≥ 0.90 (e.g., Self-efficacy and Social Support), scores were calculated when at least 50% of the items had been completed. For scales with alpha values $\geq 0.80 < 0.90$ (e.g., Life Satisfaction, Resilience, and PANAS), up to 20% of items could be missing. For scales with alpha values ≥ 0.70 and < 0.80 (e.g., Depression and Community Cohesion), scores were calculated only when participants had one or no missing item. Scores of participants who did not meet these criteria for a given scale were treated as missing data (i.e., their scale score was set to NA).

This procedure implies that the number of items used to compute scale scores may vary across participants. To account for this, all scale scores were calculated as the mean of available items, even for scales that are typically computed as a sum: Self-efficacy, Life Satisfaction, Resilience, Depression, and Social Support. For these scales, however, an alternative, more traditional sum score was also computed for participants with complete item responses, and this version was used to report more readable descriptive statistics in Table S3.

```
data <- data_RAW |>
  rename(
    Age = SD1,
    City = SD2,
    Gender = SD3,
    Sexualientation = SD4,
    Sex = SD5,
    Ethnicity = SD6,
    Farmer = SD7,
    Marital_Status = SD8,
    SES = SD9, # Socioeconomic Status
    Education = SD10,
    Children = SD11,
    Housing = SD12,
    Cohabitant = SD13,
    Monthly_Income = SD14,
    Income_Source = SD15,
    Employment = SD16,
    Job = SD17,
    Hearing_Difficulties = SD18_1,
    Speaking_Difficulties = SD18_2,
    Seeing_Difficulties = SD18_3,
    Moving_Difficulties = SD18_4,
    Grabing_Difficulties = SD18_5,
    Understanding_Difficulties = SD18_6,
    Interacting_Difficulties = SD18_7,
    LP_Alcohol = SD19_1_A,
    LP_Cigarette = SD19_2_A,
    LP_Cannabis = SD19_3_A,
    LP_Cocaine = SD19_4_A,
    LP_Basuco = SD19_5_A,
    LP_Inhalant = SD19_6_A,
    LP_Ecstasy = SD19_7_A,
    LP_Psilocybin = SD19_8_A,
    LP_LSD = SD19_9_A,
    LP_Tranquilizer = SD19_10_A,
    LP_Popper = SD19_11_A,
    LP_Anfetamines = SD19_12_A,
    LP_Heroine = SD19_13_A,
    LM_Alcohol = SD19_1_B,
    LM_Cigarette = SD19_2_B,
    LM_Cannabis = SD19_3_B,
    LM_Cocaine = SD19_4_B,
    LM_Basuco = SD19_5_B,
    LM_Inhalant = SD19_6_B,
    LM_Ecstasy = SD19_7_B,
    LM_Psilocybin = SD19_8_B,
    LM_LSD = SD19_9_B,
    LM_TRAN = SD19_10_B,
    LM_Popper = SD19_11_B,
    LM_Anfetamines = SD19_12_B,
    LM_Heroine = SD19_13_B,
```

```
LW_Alcohol = SD19_1_C,
  LW_Cigarette = SD19_2_C,
 LW_Cannabis = SD19_3_C,
 LW Cocaine = SD19 4 C,
 LW_Basuco = SD19_5_C,
 LW_Inhalant = SD19_6_C,
 LW_Ecstasy = SD19_7_C,
 LW_Psilocybin = SD19_8_C,
  LW_LSD = SD19_9_C,
 LW_Tranquilizer = SD19_10_C,
 LW_Popper = SD19_11_C,
 LW_Anfetamines = SD19_12_C,
 LW_Heroine = SD19_13_C,
 Health = SD20_1,
 Illness = SD21,
 Disease_Other = SD22_13_TEXT,
 PCPS1_4_Other = PCPS1_4_texto,
  eed1_7_Other = EED1_7_TEXT
# Replace character "99" with NA for missing values
mutate(across(where(is.character), ~ na_if(., "99"))) |>
mutate(across(where(is.numeric), ~ na_if(., 99))) |>
mutate(
  Gender = recode(
    Gender,
    "1" = "Male",
    "2" = "Female",
    "3" = "Androgynous",
    "4" = "Trans woman",
    "9" = "Non-binary",
   "10" = "Don't know",
mutate(Gender = if_else(Gender %in% c(
if_else(Gender %in% c("Man", "Trans masculine", "Trans man"),
  "Trans man",
  "Non-binary"
mutate(
 Housing = recode(
    Housing,
```

```
"2" = "Renting (entire home)",
    "5" = "Without permanent housing"
mutate_at(
    "LP_Basuco",
    "LP_Ecstasy",
    "LP_Psilocybin",
    "LP_Popper",
    "LP_Heroine",
    "LM_Alcohol",
    "LM_Cigarette",
    "LM_Cannabis",
    "LM_Cocaine",
    "LM_Basuco",
    "LM_Inhalant",
    "LM_Ecstasy",
    "LM_Psilocybin",
    "LM_LSD",
    "LM_TRAN",
    "LM_Popper",
    "LM_Anfetamines",
    "LM_Heroine",
    "LW_Alcohol",
    "LW_Cigarette",
    "LW_Cannabis",
    "LW_Cocaine",
    "LW_Basuco",
    "LW_Ecstasy",
    "LW_Psilocybin",
    "LW_LSD",
    "LW_Tranquilizer",
    "LW_Popper",
    "LW_Anfetamines",
    "LW_Heroine"
  ~ recode(.x, "1" = 1, "2" = 0)
select(
    Codigo,
```

```
ends_with("_TEXT"),
    Sexualientation,
    ends_with("_texto")
 Ethnicity = recode(
    Ethnicity,
    "1" = "Indigenous",
    "2" = "Rrom",
    "6" = "Afro-Colombian",
mutate(Farmer = recode(
  Farmer,
  "5" = NA_character_
mutate(
 Marital_Status = recode(
    Marital_Status,
    "3" = "Widow/er",
mutate(
 SES = recode_factor(
   SES,
    "1" = "Low",
    "4" = "Middle-high",
mutate(
 Education = recode_factor(
    Education,
```

```
"7" = "Technical degree",
    "8" = "University (unfinished)",
    "10" = "Postgraduate studies"
mutate(across(
   SD22_1:SD22_13,
   EED1_1,
   EED1_2,
   EED1_3,
   EED1_4,
   EED1_5,
   EED1_6,
   EED1_7,
   EED2_1:EED2_5
  ~ as.numeric(
   recode(
      as.character(.x),
     "2" = "0",
      .default = NA_character_,
      .missing = NA_character_
mutate(across(
  ends_with("_Difficulties"),
  ~ case_when(.x == 99 ~ NA_real_, is.na(.x) ~ NA_real_, .x == 4 ~ 1, TRUE ~ 0)
mutate(Difficulty_Dichotomous = if_else(
  rowSums(across(ends_with("_Difficulties"), ~ is.na(.))) > 0,
 NA_real_, # Assign NA if any difficulty is missing
  if_else(rowSums(across(ends_with("_Difficulties"), ~ . == 1)) == 7,
mutate(across(PCPS1 1:PCPS1 5, ~ case when(
 TRUE ~ NA_real_ # Missing or other values
))) |>
mutate(across(PCPS2_1:PCPS2_5, ~ case_when(
```

```
. %in% 2:5 ~ 1, # Some engagement
 TRUE ~ NA_real_ # Missing or other values
mutate(across(starts_with("EBD_"), ~ ifelse(is.na(.x), NA, .x - 1))) |>
# Compute aggregate variables summarizing different aspects
 Polyconsumption_Month = rowSums(across(LM_Alcohol:LM_Heroine, ~.), na.rm = TRUE),
  Disease_Burden = rowSums(across(SD22_1:SD22_13, ~.), na.rm = TRUE),
  Group_Membership = rowSums(across(PCPS1_1:PCPS1_5, ~.), na.rm = TRUE),
  Community_Engagement = rowSums(across(PCPS2_1:PCPS2_5, ~.), na.rm = TRUE),
 Discrimination = rowSums(across(EED1_1:EED1_7, ~.), na.rm = TRUE),
 Discrimination = ifelse(Discrimination >= 1, "Yes", "No"),
  Self_efficacy = if_else(rowSums(!is.na(across(starts_with("EAG_")))) >= 5,
    rowMeans(across(starts_with("EAG_")), na.rm = TRUE),
   NA_real_
 Life_Satisfaction = if_else(rowSums(!is.na(across(starts_with("SWLS_")))) >= 3,
    rowMeans(across(starts_with("SWLS_")), na.rm = TRUE),
   NA_real_
  Resilience = if_else(rowSums(!is.na(across(starts_with("EBR_")))) >= 3,
    rowMeans(across(starts_with("EBR_")), na.rm = TRUE),
   NA_real_
  Depression = if_else(rowSums(!is.na(across(starts_with("EBD_")))) >= 6,
    rowMeans(across(starts_with("EBD_")), na.rm = TRUE),
   NA real
  Social_Support = if_else(rowSums(!is.na(across(starts_with("MOS2_")))) >= 10,
    rowMeans(across(starts_with("MOS2_")), na.rm = TRUE),
    NA_real_
 PANAS_Positive_Affect = if_else(rowSums(!is.na(across(all_of(PANAS_P)))) >= 8,
    rowMeans(across(all_of(PANAS_P)), na.rm = TRUE),
   NA_real_
  PANAS_Negative_Affect = if_else(rowSums(!is.na(across(all_of(PANAS_N)))) >= 9,
    rowMeans(across(all_of(PANAS_N)), na.rm = TRUE),
   NA real
  Community_Cohesion = if_else(rowSums(!is.na(across(starts_with("PCPS3_")))) >= 2,
    rowMeans(across(starts_with("PCPS3_")), na.rm = TRUE),
   NA real
  ),
  Self_efficacy_sum = if_else(rowSums(!is.na(across(starts_with("EAG_")))) >= 1,
    rowSums(across(starts_with("EAG_")), na.rm = TRUE),
   NA_real_
  Life_Satisfaction_sum = if_else(rowSums(!is.na(across(starts_with("SWLS_")))) >= 1,
```

```
rowSums(across(starts_with("SWLS_")), na.rm = TRUE),
   NA_real_
 Resilience_sum = if_else(rowSums(!is.na(across(starts_with("EBR_")))) >= 1,
    rowSums(across(starts_with("EBR_")), na.rm = TRUE),
   NA_real_
  ),
  Depression_sum = if_else(rowSums(!is.na(across(starts_with("EBD_")))) >= 1,
    rowSums(across(starts_with("EBD_")), na.rm = TRUE),
   NA_real_
 Social_Support_sum = if_else(rowSums(!is.na(across(starts_with("MOS2_")))) >= 1,
    rowSums(across(starts_with("MOS2_")), na.rm = TRUE),
    NA_real_
# Select relevant variables
select(
  Age, Gender, Ethnicity, Marital_Status, SES, Education, Housing,
 Health, Polyconsumption_Month:Social_Support_sum
# Convert categorical variables Housing to Job into factors
mutate(Housing = as.factor(Housing)) |>
mutate_if(is.character, as.factor) |>
filter(Age >= 18)
```

2.2 Missing Data

Create a summary of missing data for each variable in the final dataset

```
Missing_data <- data |>
    select(-ends_with("_sum")) |>
    # Summarize across all columns, counting the number of NA values in each column
    summarise(across(everything(), ~ sum(is.na(.)))) |>
    # Convert the summary from wide format (one row, many columns) to long format
    pivot_longer(
        everything(), # Select all columns
        names_to = "Variable", # Store column names in a new variable "Variable"
        values_to = "NA_count" # Store the count of NAs in a new variable "NA_count"
) |>
    # Compute the proportion of missing values for each variable
    mutate(Proportion = NA_count / dim(data)[1]) # Divide NA count by total number of rows
```

2.2.1 Fig. S1. Proportion of missing data

To apply multi-model inference techniques such as dredge and model averaging, models must be fitted with complete data. Therefore, assessing the proportion of missing data per variable was crucial. While excessive missingness could lead to unreliable models, imputing missing values might reduce data credibility. Since no variable had an unacceptably high proportion of missing data, we opted not to impute missing values.

```
Missing_data |>
  mutate_at("Variable", str_replace_all, "Self_efficacy", "Self-efficacy") |>
  mutate_at("Variable", str_replace_all, "_", " ") |>
  ggplot(aes(
    x = fct_reorder(Variable, Proportion, .desc = TRUE), # Reorder variables
    y = Proportion,
```

```
fill = Proportion # Use fill color to indicate proportion of missing data
)) +
geom_col() + # Create bar plot
# Add percentage labels on top of bars
geom_text(aes(label = percent(Proportion, accuracy = 1)),
    vjust = -0.5, size = 2
) +
# Apply color gradient: Green (low missing data), Yellow (moderate), Red (high missing data)
scale_fill_viridis_c(
    option = "plasma", # Define color range
    direction = -1, # Reverse the color scale
    labels = percent_format(accuracy = 1), # Convert legend values to percentage format
    alpha = 0.7
) +
# Convert Y-axis (proportion of missing data) to a percentage scale
scale_y_continuous(labels = scales::percent_format(accuracy = 1)) +
# Add axis labels
labs(
    y = "Percentage of Missing Data", # Label for Y-axis
    x = "Variable" # Label for X-axis
) +
# Use a minimal theme for a cleaner visual appearance
theme_minimal() +
# Rotate X-axis labels for better readability
theme(axis.text.x = element_text(angle = 60, hjust = 1))
```

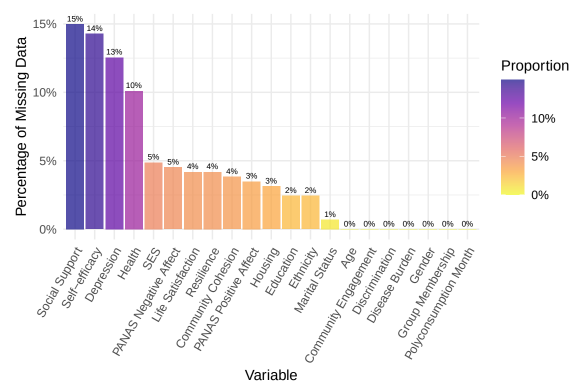


Figure S1. Proportion of missing data per variable. Variables are ordered from highest to lowest proportion of missing values. The color gradient indicates the proportion of missingness, with darker shades representing higher percentages.

3 Descriptives

3.1 Table S2. Socio-demographic characteristics by gender

```
data |>
 select(Age:Housing) |>
 rename_with(~ gsub("_", " ", .x)) |>
 tbl_summary(by = Gender,
             statistic = all_continuous() ~ "{mean}±{sd}",
             digits = all_continuous() ~ 2,
             missing_text = "Missing data") |>
 add_n() |> # add column with total number of non-missing observations
 bold_labels() |>
 remove_footnote_header(columns = all_stat_cols()) |>
 as kable extra(
   format = "latex",
   linesep = "",
   booktabs = TRUE,
   caption = "Sociodemographic characteristics of the study participants
 kable_styling(latex_options = c("scale_down", "HOLD_position")) |>
 footnote(
   general = "Age is reported as mean ± SD; all other variables are shown as frequency
   threeparttable = TRUE,
   footnote_as_chunk = TRUE,
   escape = FALSE
```

Table S2. Sociodemographic characteristics of the study participants by gender identity

Characteristic	N	Non-binary N = 70	Trans man N = 28	Trans woman N = 189
Age	287	30.80 ± 10.57	28.96 ± 8.36	36.61 ± 11.75
Ethnicity	280			
Afro-Colombian		18 (26%)	10(37%)	57 (31%)
Indigenous		5 (7.4%)	1(3.7%)	17(9.2%)
None		43 (63%)	15 (56%)	108 (58%)
Rrom		2(2.9%)	1(3.7%)	3 (1.6%)
Missing data		$\overline{2}$	1	4
Marital Status	285			
Civil union		7(10%)	6 (21%)	27 (14%)
Divorced		1 (1.4%)	0 (0%)	2 (1.1%)
Married		8 (11%)	4 (14%)	11 (5.9%)
Single		48 (69%)	13 (46%)	138 (74%)
Stable relationship		3(4.3%)	4 (14%)	6 (3.2%)
Widow/er		3(4.3%)	1(3.6%)	3(1.6%)
Missing data		0	0	$\stackrel{\cdot}{2}$
SES	273			
Low		54 (81%)	20~(74%)	146 (82%)
Middle-low		12 (18%)	4 (15%)	29 (16%)
Middle-high		1 (1.5%)	2(7.4%)	2 (1.1%)
High		0 (0%)	1(3.7%)	2(1.1%)
Missing data		3	1	10
Education	280			
No studies, illiterate		1(1.5%)	1(3.7%)	3(1.6%)
No studies, literate		0 (0%)	0 (0%)	3~(1.6%)
Primary school (unfinished)		3(4.4%)	0 (0%)	18 (9.7%)
Primary school		1 (1.5%)	0 (0%)	15~(8.1%)
Secondary school (unfinished)		12 (18%)	4 (15%)	54 (29%)
Secondary school		29 (43%)	9(33%)	57 (31%)
Technical degree		13~(19%)	6~(22%)	23 (12%)
University (unfinished)		7(10%)	5(19%)	9~(4.9%)
University		1 (1.5%)	2(7.4%)	3~(1.6%)
Postgraduate studies		1 (1.5%)	0 (0%)	0 (0%)
Missing data		2	1	4
Housing	278			
Home-owner		5(7.4%)	2(7.4%)	10~(5.5%)
Living with family		4 (5.9%)	3 (11%)	11 (6.0%)
Renting (entire home)		30 (44%)	17~(63%)	86 (47%)
Shared rental (room)		27 (40%)	4 (15%)	66 (36%)
Without permanent housing		2(2.9%)	1(3.7%)	10 (5.5%)
Missing data		2	1	6

Note: Age is reported as mean \pm SD; all other variables are shown as frequency (%).

3.2 Table S3. Measured variables by gender

```
data |>
   select(
    Gender,
    Disease_Burden,
   Group_Membership,
   Community_Engagement,
   Discrimination,
```

```
Polyconsumption_Month,
 Self_efficacy_sum,
 Life_Satisfaction_sum,
  Resilience_sum,
  Depression_sum,
  Social_Support_sum,
 PANAS_Positive_Affect,
  PANAS_Negative_Affect,
  Community_Cohesion
rename_with(~ gsub("Self_efficacy", "Self-efficacy", .x)) |>
rename_with(~ gsub("_sum", " (sum)", .x)) |>
rename_with(~ gsub("_", " ", .x)) |>
tbl_summary(by = Gender,
            statistic = all_continuous() ~ "{mean} ± {sd}",
            missing_text = "Missing data") |>
add n() |> # add column with total number of non-missing observations
bold_labels() |>
remove_footnote_header(columns = all_stat_cols()) |>
as_kable_extra(
 format = "latex",
 linesep = "",
  booktabs = TRUE,
  caption = "Other sociodemographic, health, and psychosocial characteristics
            of the study participants by gender identity"
kable_styling(latex_options = c("scale_down", "HOLD_position")) |>
footnote(
  general = "Continuous variables are reported as mean ± SD, and categorical variables
            as frequency (\\\\)). Missing data indicate the number of participants with
            insufficient responses to compute the corresponding score. For interpretability,
            Self-efficacy, Life Satisfaction, Resilience, Depression, and Social Support are
            shown as sum scores, calculated only for participants with complete item
  threeparttable = TRUE,
  footnote_as_chunk = TRUE,
  escape = FALSE
```

Table S3. Other sociodemographic, health, and psychosocial characteristics of the study participants by gender identity

		37 11				
Characteristic	\mathbf{N}	Non-binary	Trans man	Trans woman		
		N = 70	N = 28	N = 189		
Disease Burden	287					
0		35~(50%)	11 (39%)	91 (48%)		
1		20~(29%)	11 (39%)	58 (31%)		
2		9 (13%)	3 (11%)	19 (10%)		
3		2(2.9%)	3 (11%)	8 (4.2%)		
4		2(2.9%)	0 (0%)	8 (4.2%)		
5		1 (1.4%)	0 (0%)	3~(1.6%)		
6		1 (1.4%)	0 (0%)	2(1.1%)		
Group Membership	287					
0		29 (41%)	10 (36%)	65 (34%)		
1		32 (46%)	14 (50%)	84 (44%)		
2		4~(5.7%)	2(7.1%)	23~(12%)		
3		1 (1.4%)	2(7.1%)	9 (4.8%)		
4		3 (4.3%)	0 (0%)	6 (3.2%)		
5		1 (1.4%)	0 (0%)	2(1.1%)		
Community Engagement	287					
0		21 (30%)	4 (14%)	27 (14%)		
1		9 (13%)	2(7.1%)	43~(23%)		
2		6~(8.6%)	6 (21%)	44~(23%)		
3		$11 \ (16\%)$	5 (18%)	39~(21%)		
4		15 (21%)	8 (29%)	$28 \ (15\%)$		
5		8 (11%)	3 (11%)	8~(4.2%)		
Discrimination	287	60~(86%)	21~(75%)	169 (89%)		
Polyconsumption Month	287	2.16 ± 2.11	1.82 ± 2.02	2.37 ± 3.13		
Self-efficacy (sum)	251	30 ± 8	26 ± 9	33 ± 7		
Missing data		8	7	21		
Life Satisfaction (sum)	284	21 ± 8	21 ± 9	23 ± 10		
Missing data		1	0	2		
Resilience (sum)	280	14.6 ± 5.0	12.5 ± 4.1	15.4 ± 4.6		
Missing data		3	1	3		
Depression (sum)	286	9.9 ± 4.4	9.7 ± 4.6	9.5 ± 4.2		
Missing data		0	0	1		
Social Support (sum)	250	60 ± 24	57 ± 22	60 ± 24		
Missing data		8	7	22		
PANAS Positive Affect	277	3.53 ± 1.09	3.31 ± 0.85	3.94 ± 0.91		
Missing data		2	1	7		
PANAS Negative Affect	274	2.31 ± 0.94	2.43 ± 0.89	2.50 ± 1.01		
Missing data		2	2	9		
Community Cohesion	276	4.15 ± 1.74	3.98 ± 1.44	4.68 ± 1.45		
Missing data		5	0	6		

Note: Continuous variables are reported as mean \pm SD, and categorical variables as frequency (%). Missing data indicate the number of participants with insufficient responses to compute the corresponding score. For interpretability, Self-efficacy, Life Satisfaction, Resilience, Depression, and Social Support are shown as sum scores, calculated only for participants with complete item responses.

3.3 Figure S2. Distribution of participants' categorical and ordinal variables by gender

```
custom_palette <- viridis::viridis(3, option = "plasma", begin = 0, end = 0.75)
p desc1 <- data |>
  select(Gender,
        Ethnicity,
        Marital_Status,
        SES,
        Education,
        Housing,
        Disease_Burden,
        Group_Membership,
        Community_Engagement,
        Discrimination) |>
 mutate_all(as.factor) |>
 pivot_longer(
   cols = -Gender,
   names_to = "Variable", values_to = "Response"
 mutate(
   Response = as.factor(Response),
   Variable = str_wrap(Variable, width = 20)
  group_by(Gender, Variable, Response) |>
  summarise(n = n(), .groups = "drop") |>
  group_by(Gender, Variable) |>
 mutate(Percentage = n / sum(n) * 100,
         Variable = str_replace_all(Variable, "_", " ")) |>
  ggplot(aes(x = Response, y = Percentage, fill = Gender)) +
  geom_col(position = position_dodge(width = 0.9)) +
  facet_wrap(~Variable, scales = "free_x", ncol = 5) +
 scale_y_continuous(labels = scales::percent_format(scale = 1)) + # Show axis in %
  scale_fill_manual(values = custom_palette) +
 labs(x = NULL, y = "Percentage") +
  theme_minimal() +
  theme(legend.position = "bottom",
        axis.text.x = element_text(angle = 52, hjust = 1))
p_desc1
```

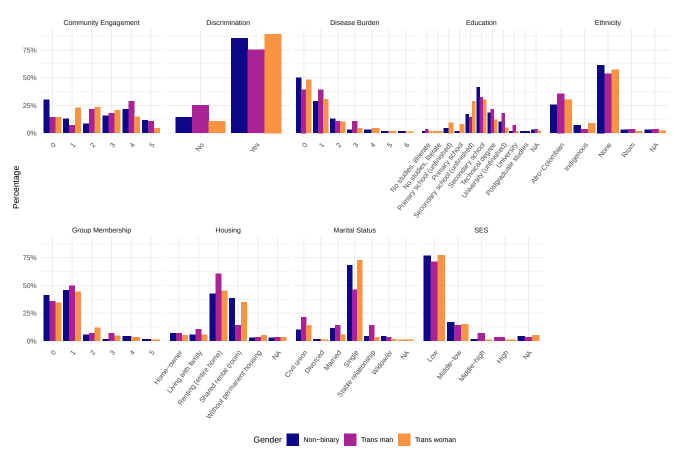


Figure S2. Distribution of categorical and ordinal variables by gender. Values represent the percentage of responses in each category. To facilitate comparison across genders, percentages were calculated within each gender group. Additional descriptive statistics are provided in Table S2.

3.4 Figure S3. Distribution of participants' measured numeric variables by gender

```
desc_long_data <- data |>
 select(
   Gender,
    Age,
   Polyconsumption_Month,
   Self_efficacy_sum,
   Life_Satisfaction_sum,
   Resilience_sum,
   Depression_sum,
   Social_Support_sum,
   PANAS_Positive_Affect,
   PANAS_Negative_Affect,
    Community_Cohesion
 pivot_longer(
    cols = -Gender,
   names_to = "Variable", values_to = "Value"
 mutate(
    Variable = str_replace_all(Variable, "Self_efficacy", "Self-efficacy"),
```

```
Variable = str_replace_all(Variable, "_sum", " (sum)"),
    Variable = str_replace_all(Variable, "_", " "),
    Variable = str_wrap(Variable, width = 30)
desc_means <- desc_long_data |>
  group_by(Gender, Variable) |>
  summarise(Mean = mean(Value, na.rm = TRUE), .groups = "drop")
p_desc2 <- ggplot(desc_long_data, aes(Value, fill = Gender, colour = Gender)) +</pre>
  geom_density(alpha = 0.6) +
  geom_vline(data = desc_means,
             aes(xintercept = Mean,
                 colour = Gender),
             linetype = "dashed", size = 0.8) +
  facet_wrap(~Variable, scales = "free", ncol = 5) +
  labs(y = "Density", x = NULL) +
  scale_fill_manual(values = custom_palette) +
  scale_colour_manual(values = custom_palette) +
  theme minimal() +
  theme(legend.position = "bottom")
 _desc2
```

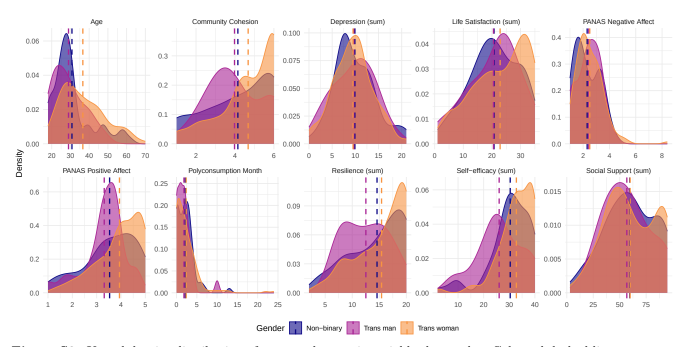


Figure S3. Kernel density distribution of measured numeric variables by gender. Coloured dashed lines represent mean values. For interpretability, Self-efficacy, Life Satisfaction, Resilience, Depression, and Social Support are shown as sum scores, calculated only for participants with complete item responses. Detailed descriptives are found in Table S3.

3.5 Table S4. Correlations

```
# Compute correlations for all participants combined dat.corr.ALL <- data |>
```

```
select(Age, Polyconsumption_Month:Community_Cohesion, -Discrimination) |> # Numeric variables
  corr.stars() |>
  tail(-1) |>
 rownames to column(var = " ")
dat.corr.NB <- data |>
  filter(Gender == "Non-binary") |> # Select only non binary
 select(Age, Polyconsumption_Month:Community Cohesion, -Discrimination) |> # Numeric variables
 corr.stars() |> # Compute correlation matrix with significance stars
 tail(-1) |>
 rownames_to_column(var = " ") # Move row names to a column
dat.corr.TM <- data |>
 filter(Gender == "Trans man") |> # Select only trans men
 select(Age, Polyconsumption_Month:Community_Cohesion, -Discrimination) |> # Numeric variables
  corr.stars() |>
 tail(-1) |>
 rownames_to_column(var = " ")
dat.corr.TW <- data |>
  filter(Gender == "Trans woman") |> # Select only trans women
 select(Age, Polyconsumption_Month:Community_Cohesion, -Discrimination) |> # Numeric variables
  corr.stars() |>
 tail(-1) |>
 rownames_to_column(var = " ")
bind_rows(dat.corr.ALL, dat.corr.NB, dat.corr.TM, dat.corr.TW) |>
 rename_with(~ gsub("Self_efficacy", "Self-efficacy", .x)) |>
  rename_with(~ gsub("_", " ", .x)) |>
 mutate_at(" ", str_replace_all, "Self_efficacy", "Self-efficacy") |>
 mutate_at(" ", str_replace_all, "_", " ") |>
 kable(
   digits = 2, booktabs = TRUE,
   align = c("l", rep("c", 12)),
   linesep = "",
   caption = "Correlations between measured variables", escape = FALSE
 pack_rows("All participants",
   start_row = 1, end_row = 12, bold = FALSE,
   background = "lightgray"
 pack_rows("Non binary",
    start_row = 13, end_row = 24, bold = FALSE,
   background = "lightgray"
 pack_rows("Trans men",
   start_row = 25, end_row = 36, bold = FALSE,
   background = "lightgray"
 pack_rows("Trans women",
    start_row = 37, end_row = 48, bold = FALSE,
```

```
background = "lightgray"
) |>
# Apply LaTeX styling
kable_styling(latex_options = c("HOLD_position", "scale_down")) |>
column_spec(2:10, width = "2.2cm") |> # Adjust column widths
# Add footnote explaining correlation significance levels
footnote(
    general = paste0(
        "Values represent Spearman correlation coefficients ($\\\\rho$). ",
        "For significance, $^{\\\\dagger}p$ < 0.1, *$p$ < 0.05, ",
        "**$p$ < 0.01, ***$p$ < 0.001. ",
        "Significant correlations are in bold."
),
        threeparttable = TRUE, footnote_as_chunk = TRUE, escape = FALSE
) |>
landscape() # Rotate table for better readability in LaTeX
```

Table S4. Correlations between measured variables

	Age	Polyconsumption Month	Disease Burden	Group Membership	Community Engagement	Self-efficacy	Life Satisfaction	Resilience	Depression	Social Support	PANAS Positive Affect	PANAS Negative Affect
All participants												
Polyconsumption Month	-0.32***											
Disease Burden	0.16**	-0.03										
Group Membership	0.17**	-0.04	0.11^{\dagger}									
Community Engagement	-0.01	0.01	0.07	0.42***								
Self-efficacy	0.23***	-0.18**	-0.12^{\dagger}	0.13*	0.06							
Life Satisfaction	0.11^{\dagger}	-0.21***	-0.23***	0.22***	0.07	0.4***						
Resilience	0.17**	-0.14*	-0.07	0.29***	0.14*	0.49***	0.49***					
Depression	0.05	0.10	0.27***	0.13*	0.10	-0.10	-0.21***	-0.05				
Social Support	0.14*	-0.06	-0.08	0.24***	0.12^{\dagger}	0.33***	0.33***	0.26***	-0.06			
PANAS Positive Affect	0.18**	-0.10	-0.10	0.26***	0.02	0.45***	0.46***	0.42***	-0.02	0.33***		
PANAS Negative Affect	-0.11^{\dagger}	0.19**	0.11^{\dagger}	-0.06	-0.12*	-0.09	-0.18**	-0.19**	0.38***	0.04	-0.02	
Community Cohesion	0.19**	-0.09	0.06	0.26***	0.25***	0.28***	0.25***	0.29***	0.23***	0.26***	0.3***	-0.03
Non binary												
Polyconsumption Month	-0.28*											
Disease Burden	0.22^{\dagger}	0.01										
Group Membership	0.12	-0.02	0.06									
Community Engagement	-0.05	0.04	0.21^{\dagger}	0.39***								
Self-efficacy	0.35**	-0.3*	-0.05	0.03	0.21							
Life Satisfaction	0.33**	-0.34**	-0.10	0.05	0.00	0.26*						
Resilience	0.19	-0.21^{\dagger}	-0.04	0.17	0.19	0.56***	0.48***					
Depression	0.31*	0.15	0.47***	0.32**	0.20	0.11	0.01	0.13				
Social Support	0.14	0.13	0.08	0.12	0.05	0.15	0.32*	0.05	0.15			
PANAS Positive Affect	0.27*	-0.06	-0.06	0.21^{\dagger}	0.10	0.49***	0.27*	0.27*	0.20	0.22^{\dagger}		
PANAS Negative Affect	0.01	0.21^{\dagger}	0.10	-0.12	-0.02	-0.17	-0.13	-0.26*	0.35**	0.02	0.18	
Community Cohesion	0.27*	0.02	0.17	0.31*	0.39**	0.3*	0.14	0.06	0.41**	0.23^{\dagger}	0.20	0.09
Trans men												
Polyconsumption Month	-0.24											
Disease Burden	-0.04	-0.19										
Group Membership	-0.07	-0.07	0.47*									
Community Engagement	-0.15	0.18	0.22	0.55**								
Self-efficacy	0.07	-0.24	0.02	0.52*	0.05							
Life Satisfaction	0.19	0.21	-0.11	0.06	-0.19	0.21						
Resilience	-0.08	0.15	0.10	0.34^{\dagger}	-0.16	0.57*	0.12					
Depression	-0.16	0.05	0.35	0.49*	0.62**	0.14	-0.21	-0.22				
Social Support	0.13	-0.37^{\dagger}	-0.05	0.4^{\dagger}	0.05	0.35	0.15	0.27	-0.14			
PANAS Positive Affect	0.27	-0.11	-0.10	0.31	-0.28	0.34	0.30	0.61***	-0.29	0.44^{\dagger}		
PANAS Negative Affect	-0.27	0.16	0.27	0.17	0.13	-0.01	-0.29	-0.34^{\dagger}	0.58**	-0.15	-0.32	
Community Cohesion	-0.15	-0.01	0.14	0.4*	0.09	0.5*	-0.06	0.34^{\dagger}	0.22	0.27	0.25	0.16
Trans women												
Polyconsumption Month	-0.37***											
Disease Burden	0.21**	-0.03										
Group Membership	0.18*	-0.03	0.09									
Community Engagement	0.07	-0.02	-0.02	0.42***								
Self-efficacy	0.08	-0.14^{\dagger}	-0.13^{\dagger}	0.13^{\dagger}	0.05							
Life Satisfaction	-0.03	-0.22**	-0.29***	0.28***	0.16*	0.46***						
Resilience	0.12	-0.16*	-0.09	0.31***	0.18*	0.42***	0.51***					
Depression	0.03	0.09	0.19*	0.02	-0.01	-0.15^{\dagger}	-0.3***	-0.07				
Social Support	0.14^{\dagger}	-0.10	-0.14^{\dagger}	0.27***	0.14^{\dagger}	0.43***	0.35***	0.32***	-0.11			
PANAS Positive Affect	0.06	-0.13^{\dagger}	-0.12	0.24***	0.05	0.41***	0.5***	0.41***	-0.05	0.37***		
PANAS Negative Affect	-0.18*	0.19*	0.11	-0.09	-0.2**	-0.06	-0.21**	-0.13^{\dagger}	0.37***	0.07	-0.10	
Community Cohesion	0.16*	-0.13^{\dagger}	0.01	0.2**	0.23**	0.21**	0.27***	0.33***	0.18*	0.27***	0.3***	-0.10

 $Note: \qquad \text{Values represent Spearman correlation coefficients } (\rho). \text{ For significance, } ^\dagger p < 0.1, *p < 0.05, **p < 0.01, ***p < 0.01. ***p < 0.001. Significant correlations are in bold.}$

4 Multi-model Inference

This section outlines the multi-model inference approach used to examine predictors of various outcome variables. The strategy follows a systematic model selection process, accounting for model uncertainty.

For each outcome variable, we repeated the following steps:

1. Data Preparation

- Relevant predictors are selected from the cleaned dataset.
- Missing values are removed to ensure complete case analysis ¹

2. Global Model Specification

- A linear model (1m) is specified, including a broad set of demographic, psychological, and social predictors.
- This full model represents all hypothesized influences on the outcome.

3. Model Selection via dredge()

- The MuMIn::dredge() function generates all possible nested models from the global model.
- Models are ranked by Akaike's Information Criterion corrected for small sample sizes (AICc).
- This helps identify the most parsimonious models while considering model uncertainty.

4. Model Averaging (model.avg)

- Models with $\triangle AICc < 2$ (within two AICc units of the top model) are averaged.
- This produces shrinkage estimates, ensuring robust coefficient estimates.

5. Results Interpretation

- Tables summarize the coefficient estimates, importance values, and significance levels.
- Figures visualize term estimates, confidence intervals, and predictor importance.

This process is repeated for each outcome variable in the following sections.

4.1 Life Satisfaction model

This section details the modelling process for Life Satisfaction, including data preparation, model selection using Akaike's Information Criterion corrected for small sample sizes (AICc), and model averaging based on the best-ranked models.

The global model includes a broad set of predictors representing demographic, psychological, and social factors, allowing us to explore their relative contributions to Life Satisfaction. The model selection procedure identifies the most parsimonious models while accounting for model uncertainty.

```
dat_LS <- data |>
    select(
        Life_Satisfaction, Age, Gender, Ethnicity, Marital_Status, Education, Housing,
        Self_efficacy, Community_Cohesion, Depression, Social_Support, Polyconsumption_Month,
        Disease_Burden, Discrimination, Group_Membership, Community_Engagement
) |>
        drop_na()

global_LS <- lm(
        Life_Satisfaction ~ Age + Gender + Ethnicity + Marital_Status + Education +
        Housing + Self_efficacy + Community_Cohesion + Depression +
        Social_Support + Polyconsumption_Month + Disease_Burden + Discrimination +
        Group_Membership + Community_Engagement,</pre>
```

¹To maximize the number of non-missing cases, we created separate datasets for each outcome variable. Since models generated by dredge() must not contain missing values (NA) to remain comparable, this approach ensures that each model includes the largest possible sample size.

```
data = dat_LS,
  na.action = "na.fail"
)
```

4.1.1 Dredge

To identify the best-fitting models, we perform automated model selection using MuMIn::dredge(), which fits all possible models nested within the global model and ranks them based on AICc. This helps determine the most important predictors of Life Satisfaction while considering model uncertainty.

```
dr_LS <- dredge(
  global_LS,
  # trace = 2 # uncomment to see progress bar
)</pre>
```

4.1.1.1 Fig. S4. Dredge results of the Life Satisfaction model The plot below visualizes the model selection results, displaying the top 100 models ranked by AICc. Each row represents an individual model, with blue cells indicating included predictors. The height of each row reflects the model's Akaike weight (ω), representing its relative support in the model set.



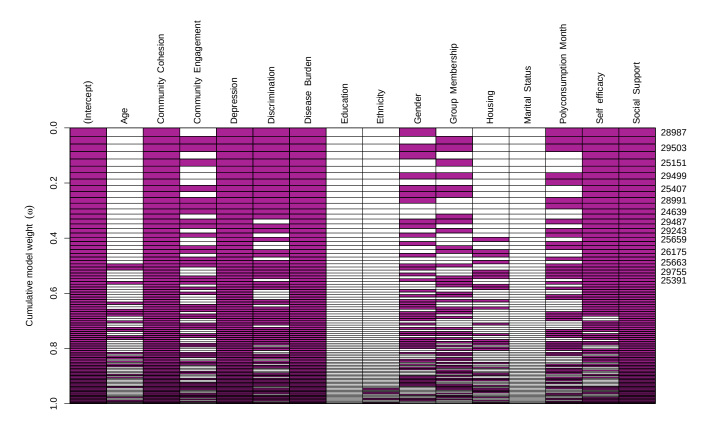


Figure S4. Model selection plot. Each row represents one of the top 100 models (ranked by AICc) out of 32,768 fitted models. Purple cells indicate included terms, with row height corresponding to each model's relative support based on its Akaike weight (omega)

4.1.2 Average model

To account for model uncertainty, we compute an averaged model that combines estimates from all models with $\Delta AICc < 2$ (i.e., models within two AICc units of the best-supported model). This approach produces more robust and conservative parameter estimates, reducing overconfidence in any single model.

```
avg_LS <- model.avg(dr_LS, subset = delta < 2, fit = TRUE)</pre>
```

4.1.2.1 Table S5. Term estimates for the Life Satisfaction model The table below presents coefficient estimates from the full average model, including 95% confidence intervals, standard errors, and significance levels.

tab_avg_LS <- avg.mod.summary(avg_LS, data = dat_LS)
tab_avg_LS\$table</pre>

Table S5.	Coefficient	estimates	for	the Li	fe	Satisfaction	model.

Term	В	95%CIs	z	p
(Intercept)	2.935	1.394 - 4.476	3.733	< 0.001
Age	0.000	-0.004 - 0.003	0.090	0.93
Community Cohesion	0.237	0.086 - 0.389	3.073	0.0021
Community Engagement	-0.061	-0.212 - 0.091	0.784	0.43
Depression	-0.572	-0.943 — -0.202	3.028	0.0025
Discrimination: Yes	-0.541	-1.276 - 0.195	1.441	0.15
Disease Burden	-0.234	-0.3970.071	2.815	0.0049
Gender: Trans man	0.281	-0.504 - 1.067	0.702	0.48
Gender: Trans woman	0.247	-0.327 - 0.82	0.843	0.4
Group Membership	0.075	-0.136 - 0.285	0.696	0.49
Housing: Living with family	-0.144	-1.033 - 0.746	0.316	0.75
Housing: Renting (entire home)	-0.043	-0.419 - 0.332	0.227	0.82
Housing: Shared rental (room)	-0.066	-0.538 - 0.405	0.275	0.78
Housing: Without permanent housing	-0.107	-0.827 - 0.613	0.291	0.77
Polyconsumption Month	-0.031	-0.105 - 0.044	0.800	0.42
Self-efficacy	0.413	0.043 - 0.784	2.189	0.0286
Social Support	0.313	0.121 - 0.505	3.190	0.0014

Note: This table includes estimates based on a full model average. The full model average ensures that all variables are included in every model, with coefficients set to zero when absent. As a consequence, it acts as a shrinkage estimator, making estimates more conservative. B represents unstandardized model coefficients. Reference levels: Gender = Non-binary; Housing = Home-owner; Discrimination = No. Significant effects are in bold.

4.1.2.2 Fig. S5. Term estimates for the Life Satisfaction model This figure represents parameter estimates from the model-averaged coefficients.

```
fig_avg_LS <- avg.mod.plot(avg_LS)
fig_avg_LS</pre>
```

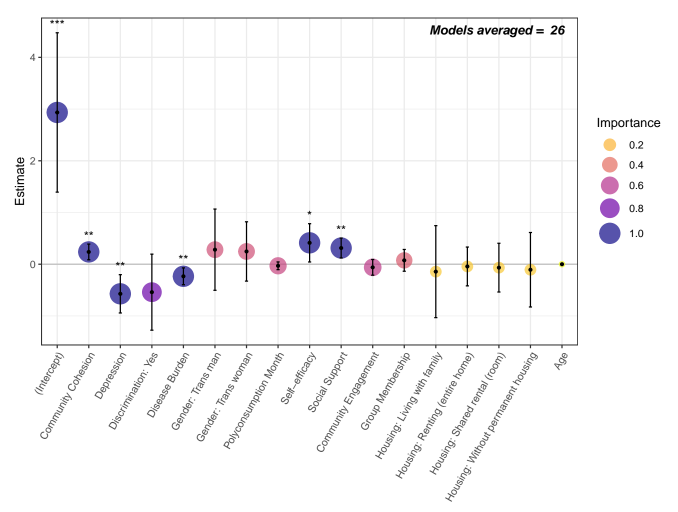


Figure S5. Term estimates with 95% confidence intervals for the final model average of the best (AICc Δ < 2) models predicting Life Satisfaction. Importance is the proportion of averaged models in which a term appears (i.e. an Importance of 0.5 means that the term is included in half the averaged models, and an Importance of 1.0 means that all the averaged models included that term). Larger and darker points indicate higher importance. Confidence intervals show uncertainty in effect size estimates. For term significance: † p < 0.10; * p < 0.05; *** p < 0.01; **** p < 0.001; **** p < 0.001.

4.2 PANAS Positive Affect model

This section details the modelling process for PANAS Positive Affect, including data preparation, model selection using Akaike's Information Criterion corrected for small sample sizes (AICc), and model averaging based on the best-ranked models.

The global model includes a broad set of predictors representing demographic, psychological, and social factors, allowing us to explore their relative contributions to PANAS Positive Affect. The model selection procedure identifies the most parsimonious models while accounting for model uncertainty.

```
dat_PANAS_P <- data |>
  select(
    PANAS_Positive_Affect, Age, Gender, Ethnicity, Marital_Status, Education, Housing,
    Self_efficacy, Community_Cohesion, Depression, Social_Support, Polyconsumption_Month,
    Disease_Burden, Discrimination, Group_Membership, Community_Engagement
) |>
    drop_na()
```

```
global_PANAS_P <- lm(
   PANAS_Positive_Affect ~ Age + Gender + Ethnicity + Marital_Status + Education +
        Housing + Self_efficacy + Community_Cohesion + Depression +
        Social_Support + Polyconsumption_Month + Disease_Burden + Discrimination +
        Group_Membership + Community_Engagement,
        data = dat_PANAS_P,
        na.action = "na.fail"
)</pre>
```

4.2.1 Dredge

To identify the best-fitting models, we perform automated model selection using MuMIn::dredge(), which fits all possible models nested within the global model and ranks them based on AICc. This helps determine the most important predictors of PANAS Positive Affect while considering model uncertainty.

```
dr_PANAS_P <- dredge(
  global_PANAS_P,
  # trace = 2 # uncomment to see progress bar
)</pre>
```

4.2.1.1 Fig. S6. Dredge results of the PANAS Positive Affect model The plot below visualizes the model selection results, displaying the top 100 models ranked by AICc. Each row represents an individual model, with blue cells indicating included predictors. The height of each row reflects the model's Akaike weight (ω) , representing its relative support in the model set.

PANAS_P_dredge_plot <- dredge.plot(dr_PANAS_P) # Run the function and store the result

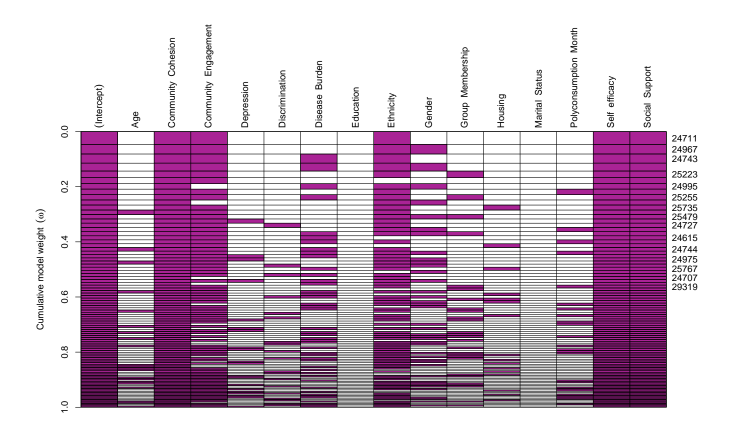


Figure S6. Model selection plot. Each row represents one of the top 100 models (ranked by AICc) out of 32,768 fitted models. Purple cells indicate included terms, with row height corresponding to each model's relative support based on its Akaike weight (omega)

4.2.2 Average model

To account for model uncertainty, we compute an averaged model that combines estimates from all models with $\Delta AICc < 2$ (i.e., models within two AICc units of the best-supported model). This approach produces more robust and conservative parameter estimates, reducing overconfidence in any single model.

```
avg_PANAS_P <- model.avg(dr_PANAS_P, subset = delta < 2, fit = TRUE)</pre>
```

4.2.2.1 Table S6. Term estimates for the PANAS Positive Affect model The table below presents coefficient estimates from the full average model, including 95% confidence intervals, standard errors, and significance levels.

```
tab_avg_PANAS_P <- avg.mod.summary(avg_PANAS_P, data = dat_PANAS_P)
tab_avg_PANAS_P$table</pre>
```

Term B95%CIszp(Intercept) 0.649-0.118 - 1.4161.658 0.1Community Cohesion 0.1270.044 - 0.212.9980.0027Community Engagement -0.073-0.167 - 0.021.535 0.12Disease Burden -0.021-0.099 - 0.0560.5380.59Ethnicity: Indigenous 0.311 -0.134 - 0.7561.369 0.17Ethnicity: None 0.248-0.051 - 0.5471.6250.1Ethnicity: Rrom 0.8170.014 - 1.621.995 0.046Gender: Trans man 0.028-0.252 - 0.3090.1980.84Gender: Trans woman 0.098-0.204 - 0.3990.6340.53Group Membership 0.009-0.053 - 0.070.2810.78Housing: Living with family 0.004 -0.156 - 0.1640.051 0.96 Housing: Renting (entire home) 0.027-0.208 - 0.2620.2240.82Housing: Shared rental (room) 0.011 -0.136 - 0.1580.1440.89 $\textbf{-}0.225 - \hspace{-0.04cm} -0.272$ Housing: Without permanent housing 0.0240.1880.85Polyconsumption Month -0.001-0.013 - 0.0110.1530.88Self-efficacy 0.5840.379 - 0.789< 0.00015.5890.185Social Support 0.077 - 0.2943.356< 0.001

Table S6. Coefficient estimates for the PANAS Positive Affect model.

Note: This table includes estimates based on a full model average. The full model average ensures that all variables are included in every model, with coefficients set to zero when absent. As a consequence, it acts as a shrinkage estimator, making estimates more conservative. B represents unstandardized model coefficients. Reference levels: Gender = Non-binary; Ethnicity = Afro-Colombian; Housing = Home-owner. Significant effects are in bold.

4.2.2.2 Fig. S7. Term estimates for the PANAS Positive Affect model This figure represents parameter estimates from the model-averaged coefficients.

fig_avg_PANAS_P <- avg.mod.plot(avg_PANAS_P)
fig_avg_PANAS_P</pre>

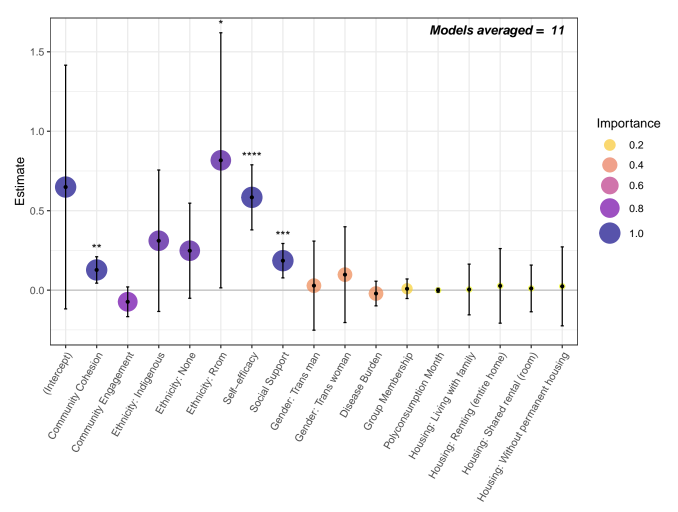


Figure S7. Term estimates with 95% confidence intervals for the final model average of the best (AICc Δ < 2) models predicting PANAS Positive Affect. Importance is the proportion of averaged models in which a term appears (i.e. an Importance of 0.5 means that the term is included in half the averaged models, and an Importance of 1.0 means that all the averaged models included that term). Larger and darker points indicate higher importance. Confidence intervals show uncertainty in effect size estimates. For term significance: † p < 0.10; * p < 0.05; *** p < 0.01; **** p < 0.001; **** p < 0.001.

4.3 PANAS Negative Affect model

This section details the modelling process for PANAS Negative Affect, including data preparation, model selection using Akaike's Information Criterion corrected for small sample sizes (AICc), and model averaging based on the best-ranked models.

The global model includes a broad set of predictors representing demographic, psychological, and social factors, allowing us to explore their relative contributions to PANAS Negative Affect. The model selection procedure identifies the most parsimonious models while accounting for model uncertainty.

```
dat_PANAS_N <- data |>
    select(
        PANAS_Negative_Affect, Age, Gender, Ethnicity, Marital_Status, Education, Housing,
        Self_efficacy, Community_Cohesion, Depression, Social_Support, Polyconsumption_Month,
        Disease_Burden, Discrimination, Group_Membership, Community_Engagement
    ) |>
        drop_na()
```

```
global_PANAS_N <- lm(
   PANAS_Negative_Affect ~ Age + Gender + Ethnicity + Marital_Status + Education +
        Housing + Self_efficacy + Community_Cohesion + Depression +
        Social_Support + Polyconsumption_Month + Disease_Burden + Discrimination +
        Group_Membership + Community_Engagement,
        data = dat_PANAS_N,
        na.action = "na.fail"
)</pre>
```

4.3.1 Dredge

To identify the best-fitting models, we perform automated model selection using MuMIn::dredge(), which fits all possible models nested within the global model and ranks them based on AICc. This helps determine the most important predictors of PANAS Negative Affect while considering model uncertainty.

```
dr_PANAS_N <- dredge(
  global_PANAS_N,
  # trace = 2 # uncomment to see progress bar
)</pre>
```

4.3.1.1 Fig. S8. Dredge results of the PANAS Negative Affect model The plot below visualizes the model selection results, displaying the top 100 models ranked by AICc. Each row represents an individual model, with blue cells indicating included predictors. The height of each row reflects the model's Akaike weight (ω) , representing its relative support in the model set.

PANAS_N_dredge_plot <- dredge.plot(dr_PANAS_N) # Run the function and store the result

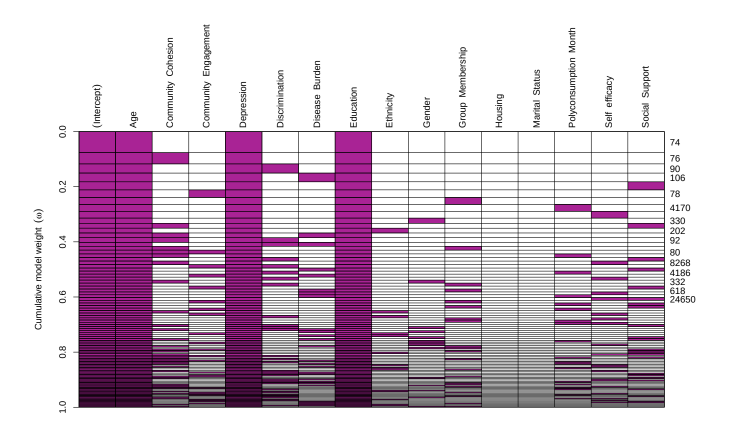


Figure S8. Model selection plot. Each row represents one of the top 100 models (ranked by AICc) out of 32,768 fitted models. Purple cells indicate included terms, with row height corresponding to each model's relative support based on its Akaike weight (omega)

4.3.2 Average model

To account for model uncertainty, we compute an averaged model that combines estimates from all models with $\Delta AICc < 2$ (i.e., models within two AICc units of the best-supported model). This approach produces more robust and conservative parameter estimates, reducing overconfidence in any single model.

```
avg_PANAS_N <- model.avg(dr_PANAS_N, subset = delta < 2, fit = TRUE)</pre>
```

4.3.2.1 Table S7. Term estimates for the PANAS Negative Affect model The table below presents coefficient estimates from the full average model, including 95% confidence intervals, standard errors, and significance levels.

```
tab_avg_PANAS_N <- avg.mod.summary(avg_PANAS_N, data = dat_PANAS_N)
tab_avg_PANAS_N$table</pre>
```

Term B95%CIszp(Intercept) 2.768 3.932 4.663< 0.00011.605 -< 0.001 -0.019-0.03 — -0.008 3.426Age -0.063 - 0.045Community Cohesion -0.0090.3210.75Depression 0.6920.481 - 0.9046.415< 0.0001 Discrimination: Yes 0.024 -0.163 - 0.2110.2490.8 Disease Burden -0.053 - 0.042-0.0050.2250.82Education: No studies, literate 0.247-1.306 - 1.80.3120.76Education: Primary school -1.679 - 0.4960.29 -0.5911.066 Education: Primary school (unfinished) -0.177-1.239 - 0.8850.3260.74Education: Secondary school -0.638-1.649 - 0.3721.238 0.22Education: Secondary school (unfinished) -0.679-1.696 - 0.3371.310 0.19Education: Technical degree -2.378 - -0.275-1.3272.4720.0134Education: University -1.619-2.87 — -0.368 2.537 0.0112Education: University (unfinished) -2.169 - 0.007-1.0811.948 0.05Social Support 0.004-0.041 - 0.050.1900.85

Table S7. Coefficient estimates for the PANAS Negative Affect model.

Note: This table includes estimates based on a full model average. The full model average ensures that all variables are included in every model, with coefficients set to zero when absent. As a consequence, it acts as a shrinkage estimator, making estimates more conservative. B represents unstandardized model coefficients. Reference levels: Education = No studies, illiterate; Discrimination = No. Significant effects are in bold.

4.3.2.2 Fig. S9. Term estimates for the PANAS Negative Affect model This figure represents parameter estimates from the model-averaged coefficients.

fig_avg_PANAS_N <- avg.mod.plot(avg_PANAS_N)
fig_avg_PANAS_N</pre>

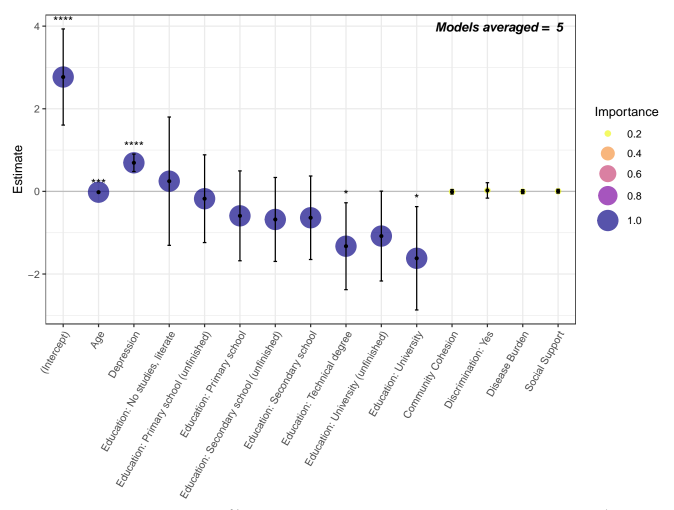


Figure S9. Term estimates with 95% confidence intervals for the final model average of the best (AICc Δ < 2) models predicting PANAS Negative Affect. Importance is the proportion of averaged models in which a term appears (i.e. an Importance of 0.5 means that the term is included in half the averaged models, and an Importance of 1.0 means that all the averaged models included that term). Larger and darker points indicate higher importance. Confidence intervals show uncertainty in effect size estimates. For term significance: † p < 0.10; * p < 0.05; *** p < 0.01; **** p < 0.001; **** p < 0.001.

5 Final figures and tables

5.1 Fig. 1. Distribution of participants' measured variables by gender

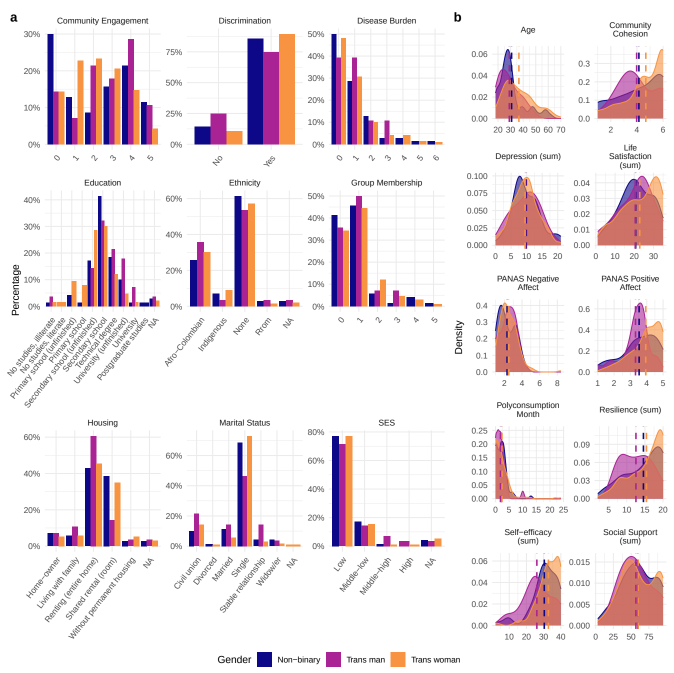


Figure 1. Distribution of variables by gender. (a) Categorical and ordinal variables. Values represent the percentage of responses in each category. To facilitate comparison across genders, percentages were calculated within each gender group. (b) Kernel density distribution of measured numeric variables. Coloured dashed lines represent mean values. For interpretability, Self-efficacy, Life Satisfaction, Resilience, Depression, and Social Support are shown as sum scores, calculated only for participants with complete item responses. Additional descriptive statistics are provided in Tables S2 and S3.

5.2 Fig. 2. Distribution of participants' measured variables by gender

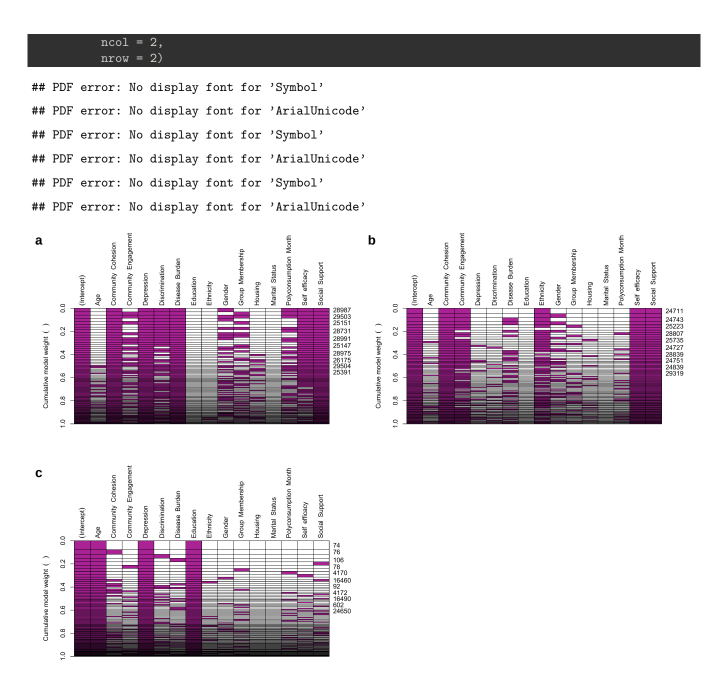


Figure 2. Model selection plot. (a) Life Satisfaction; (b) PANAS Positive Affect; (c) PANAS Negative Affect. In all cases, each row represents one of the top 100 candidate models (ranked by AICc) out of 32,768 fitted models. Purple cells indicate included terms, with row height corresponding to each model's relative support based on its Akaike weight (ω). Numbers on the right identify the top models.

```
bla <- ggplot() +
  background_image(image_read_pdf("PDF_plots/Life_Satisfaction.pdf", density = 600)) +
  theme(plot.margin = margin(t=0.2, l=1, r=0, b=0, unit = "cm"))

## PDF error: No display font for 'Symbol'
## PDF error: No display font for 'ArialUnicode'</pre>
```

5.3 Tab. 1. Term estimates for the three final averaged model

```
tab_avg_LS$coefficients |>
 full_join(tab_avg_PANAS_P$coefficients, by = "term") |>
 full_join(tab_avg_PANAS_N$coefficients, by = "term") |>
 arrange(term) |>
 kable(digits = 3,
       booktabs = TRUE,
       linesep = "",
       align = c("l", rep("c", 12)),
       caption = "Coefficient estimates for the three final average models",
       col.names = c("Term",
                      rep(c("\$B\$", "\$95 \% CIs\$", "\$z\$", "\$p\$"), times = 3)),
       escape = FALSE
 kable_styling(latex_options = c("scale_down", "HOLD_position")) |>
 add_header_above(c(" ",
                     "Life Satisfaction" = 4,
                     "PANAS Positive Affect" = 4,
 column_spec(c(5, 9), border_right = TRUE) |>
 footnote(
   general = paste0(
      "This table includes estimates based on full model averages. ",
     averaged model are included, with coefficients set to zero when absent.
     "estimator, making estimates more conservative. ",
      "$B$ represents unstandardized model coefficients for each of the three models:
     Life Satisfaction (", tab_avg_LS$reference_levels, "),
     PANAS Positive Afffect(", tab_avg_PANAS_P$reference_levels, "),
     and PANAS Negative Afffect(", tab_avg_PANAS_N$reference_levels, ").",
   threeparttable = TRUE,
   footnote_as_chunk = TRUE,
   escape = FALSE
```

Life Satisfaction PANAS Positive Affect PANAS Negative Affect В 95%CIsВ 95%CIsВ 95%CIsp(Intercept) 2.935 1.394 -4.476< 0.001 -0.118 — 1.416 0.1 2.768 1.605 -3.9324.663 < 0.0001 < 0.001 0.000 -0.004 - 0.003 0.090 0.93-0.019 -0.03 -0.008 3 426 0.044 - 0.210.0027 Community Cohesion 0.2370.0860.3893.073 0.0021 0.1272.998 0.75-0.009-0.063-0.0450.321Community Engagement -0.061 -0.2120.091 0.784 0.43 -0.073-0.167 — 0.02 1.535 0.12 -0.572 -0.943 -0.202 3.028 0.0025 0.6920.4810.904 6.415< 0.0001 Depression Discrimination: Yes -0.541-1.276-0.1951.441 0.15 0.024-0.1630.211 0.249 0.0049 -0.021 -0.099 -- 0.056 0.538 0.59 0.225 0.82 Disease Burden -0.234-0.397-0.0712.815 -0.005-0.0530.042 Education: No studies, literate 0.247-1.3060.76 0.312 1.8 Education: Primary school -0.591 0.29 -1.6790.4961.066 Education: Primary school (unfinished) -0.177-1.2390.885 0.3260.74Education: Secondary school (unfinished) -0.638-1.6490.3721.238 0.22-0.679 -1.6960.337 1.310 0.19Education: Technical degree -1.327-2.378-0.2750.0134 Education: University -1.619 -2.87 --0.368 2.537 0.0112 Education: University (unfinished) -1.081-2.169 --0.0071.948 0.05 0.17 Ethnicity: Indigenous 0.311 -0.1340.7561.369 Ethnicity: None 0.248 -0.051 0.547 1.625 Ethnicity: Rrom 0.817 0.014 1.62 1.995 0.046Gender: Trans man Gender: Trans woman 0.281-0.504-1.0670.7020.48 0.028 -0.2520.309 0.1980.840.247 -0.327 0.82 0.843 0.098 -0.2040.399 0.634 0.53 0.4Group Membership 0.075 -0.1360.2850.696 0.490.009 -0.053 0.07 0.281 0.78Housing: Living with family -1.033 0.7460.96-0.1440.316 0.004-0.1560.051Housing: Renting (entire home) -0.043-0.419 0.332 0.2270.82 0.027-0.208 0.262 0.224 0.82 0.89 Housing: Shared rental (room) -0.066-0.5380.405 0.2750.78 0.011-0.1360.1580.144Housing: Without permanent housing -0.827 0.6130.2910.77 0.024 -0.2250.2720.188 0.85-0.1070.001 0.88 Polyconsumption Month -0.031-0.1050.044-0.0130.011 < 0.0001 Self-efficacy 0.413 0.0430.784 2.189 0.0286 0.584 0.3790.7895.589 0.004 -0.041 - 0.05Social Support 3.1900.077 0.3130.1210.5050.00140.1850.2943.356 < 0.001

Table 1. Coefficient estimates for the three final average models

Note: This table includes estimates based on full model averages. The full model average ensures that all predictors present in any averaged model are included, with coefficients set to zero when absent. As a consequence, it acts as a shrinkage estimator, making estimates more conservative. B represents unstandardized model coefficients for each of the three models: Life Satisfaction (Reference levels: Gender = Non-binary; Housing = Home-owner; Discrimination = No), PANAS Positive Afffect(Reference levels: Gender = Non-binary; Ethnicity = Afro-Colombian; Housing = Home-owner), and PANAS Negative Afffect(Reference levels: Education = No studies, illiterate; Discrimination = No). Significant effects are in bold.

5.4 Fig. 3. Term estimates for the three final averaged model

This figure represents parameter estimates from the model-averaged coefficients.

```
ggarrange(
  fig_avg_LS +
    labs(subtitle = "Life Satistaction") +
    theme(legend.position = "none"),
  fig_avg_PANAS_P +
    labs(subtitle = "PANAS Possitive Affect") +
    theme(legend.position = "none"),
  fig_avg_PANAS_N +
    labs(subtitle = "PANAS Negative Affect") +
    theme(legend.position = "none"),
    as_ggplot(get_legend(fig_avg_LS)),
    ncol = 2,
    nrow = 2,
    labels = c("a", "b", "c", NULL)
)
```

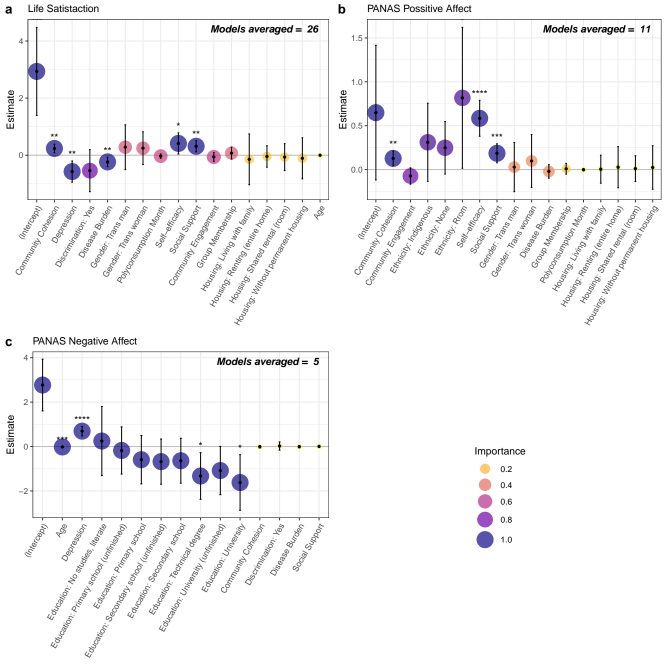


Figure 3. Term estimates with 95% confidence intervals for the final model averages of the best (AICc $\Delta < 2$) models predicting (a) Life Satisfaction, (b) PANAS Positive Affect, and (c) PANAS Negative Affect. Importance is the proportion of averaged models in which a term appears (i.e. an Importance of 0.5 means that the term is included in half the averaged models, and an Importance of 1.0 means that all the averaged models included that term). Larger and darker points indicate higher importance. Confidence intervals show uncertainty in effect size estimates. For term significance: † p < 0.10; * p < 0.05; *** p < 0.01; **** p < 0.001; **** p < 0.0001.

6 Session info (for reproducibility)

```
# Display session information for reproducibility
# - Uses `pander()` for better formatting
```

```
# - `locale = FALSE` to exclude locale-specific info
library(pander)
pander(sessionInfo(), locale = FALSE)
```

R version 4.5.0 (2025-04-11 ucrt)

Platform: x86_64-w64-mingw32/x64

attached base packages: stats, graphics, grDevices, utils, datasets, methods and base

other attached packages: pander(v.0.6.6), pdftools(v.3.5.0), ggplotify(v.0.1.2), magick(v.2.8.6), Hmisc(v.5.2-3), gtsummary(v.2.2.0), ggpubr(v.0.6.0), lubridate(v.1.9.4), forcats(v.1.0.0), stringr(v.1.5.1), dplyr(v.1.1.4), purrr(v.1.0.4), tidyr(v.1.3.1), tibble(v.3.2.1), ggplot2(v.3.5.2), tidyverse(v.2.0.0), car(v.3.1-3), carData(v.3.0-5), kableExtra(v.1.4.0), scales(v.1.3.0), readr(v.2.1.5), performance(v.0.13.0), MuMIn(v.1.48.11), psych(v.2.5.3), ltm(v.1.2-0), polycor(v.0.8-1), msm(v.1.8.2), MASS(v.7.3-65) and knitr(v.1.50)

loaded via a namespace (and not attached): mnormt(v.2.1.1), gridExtra(v.2.3), rlang(v.1.1.6), magrittr(v.2.0.3), compiler(v.4.5.0), systemfonts(v.1.2.2), vctrs(v.0.6.5), pkgconfig(v.2.0.3), crayon(v.1.5.3), fastmap(v.1.2.0), backports(v.1.5.0), labeling(v.0.4.3), rmarkdown(v.2.29), tzdb(v.0.5.0), bit(v.4.6.0), xfun(v.0.52), broom(v.1.0.8), parallel(v.4.5.0), cluster(v.2.1.8.1), R6(v.2.6.1), stringi(v.1.8.7), rpart(v.4.1.24), Rcpp(v.1.0.14), bookdown(v.0.43), base64enc(v.0.1-3), Matrix(v.1.7-3), splines(v.4.5.0), nnet(v.7.3-20), timechange(v.0.3.0), tidyselect(v.1.2.1), viridis(v.0.6.5), rstudioapi(v.0.17.1), abind(v.1.4-8), yaml(v.2.3.10), admisc(v.0.38), qpdf(v.1.3.5), lattice(v.0.22-6), withr(v.3.0.2), askpass(v.1.2.1), evaluate(v.1.0.3), foreign(v.0.8-90), gridGraphics(v.0.5-1), survival(v.3.8-3), xml2(v.1.3.8), pillar(v.1.10.2), checkmate(v.2.3.2), stats4(v.4.5.0), insight(v.1.2.0), generics(v.0.1.3), vroom(v.1.6.5), hms(v.1.1.3), munsell(v.0.5.1), glue(v.1.8.0), tools(v.4.5.0), data.table(v.1.17.0), ggsignif(v.0.6.4), fs(v.1.6.6), mvtnorm(v.1.3-3), cowplot(v.1.1.3), grid(v.4.5.0), cards(v.0.6.0), colorspace(v.2.1-1), nlme(v.3.1-168), htmlTable(v.2.4.3), Formula(v.1.2-5), cli(v.3.6.5), expm(v.1.0-0), viridisLite(v.0.4.2), svglite(v.2.1.3), gtable(v.0.3.6), rstatix(v.0.7.2), yulab.utils(v.0.2.0), digest(v.0.6.37), htmlwidgets(v.1.6.4), farver(v.2.1.2), htmltools(v.0.5.8.1), lifecycle(v.1.0.4) and bit64(v.4.6.0-1)

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