# **ABG-VBG Analysis**

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## 1 Data Pre-processing

This code pulls in the master database (a STATA file) and does some initial cleaning - this will only need to be run once, and then the data can be accessed in the usual way.

```
# put this in your first R chunk
if (!requireNamespace("kableExtra", quietly = TRUE)) install.packages("kableExtra")
library(kableExtra)
library(gtsummary)

# globally tighten gtsummary/gt tables (smaller font + tighter padding)
gtsummary::theme_gtsummary_compact()
```

Setting theme "Compact"

```
# helper: turn any gtsummary table into a PDF-safe, auto-scaling LaTeX table
to_pdf_table <- function(tbl, font_size = 8, landscape = FALSE,</pre>
                         label_col_width = NULL) {
  kbl <- gtsummary::as_kable(</pre>
    tbl,
    format = "latex",
   booktabs = TRUE,
   longtable = TRUE # allows multipage tables; repeats header with kableExtra option below
  # optional: set a fixed width for the first (label) column to encourage wrapping
  if (!is.null(label_col_width)) {
   kbl <- kableExtra::column_spec(kbl, 1, width = label_col_width)</pre>
  }
  kbl <- kableExtra::kable_styling(</pre>
    kbl,
   latex_options = c("repeat_header", "hold_position", "scale_down"),
   font size
                 = font size
  )
  if (landscape) kbl <- kableExtra::landscape(kbl) # needs pdflscape (enabled above)
  kbl
# Consolidated package management -
required_pkgs <- c(</pre>
  "WeightIt", "broom", "cobalt", "codebookr", "dplyr", "flextable", "parallel",
  "gbm", "ggplot2", "gt", "gtsummary", "haven", "labelled", "scales",
  "modelsummary", "officer", "patchwork", "rms", "survey", "tibble", "lubridate", "sensitivitymw"
# Install any missing packages (with dependencies)
missing_pkgs <- setdiff(required_pkgs, rownames(installed.packages()))</pre>
```

```
if (length(missing_pkgs)) {
 install.packages(missing_pkgs, dependencies = TRUE)
# Load (or attach) all required packages
invisible(lapply(required_pkgs, require, character.only = TRUE))
Loading required package: WeightIt
Loading required package: broom
Loading required package: cobalt
 cobalt (Version 4.6.1, Build Date: 2025-08-20)
Loading required package: codebookr
Loading required package: dplyr
Attaching package: 'dplyr'
The following object is masked from 'package:kableExtra':
    group_rows
The following objects are masked from 'package:stats':
   filter, lag
```

```
The following objects are masked from 'package:base':
    intersect, setdiff, setequal, union
Loading required package: flextable
Attaching package: 'flextable'
The following object is masked from 'package:gtsummary':
    continuous_summary
The following objects are masked from 'package:kableExtra':
    as_image, footnote
Loading required package: parallel
Loading required package: gbm
Loaded gbm 2.2.2
This version of gbm is no longer under development. Consider transitioning to gbm3, https://github.com/gbm-
developers/gbm3
Loading required package: ggplot2
Loading required package: gt
Loading required package: haven
```

Loading required package: labelled Loading required package: scales Loading required package: modelsummary Loading required package: officer Loading required package: patchwork Loading required package: rms Loading required package: Hmisc Attaching package: 'Hmisc' The following object is masked from 'package:modelsummary': Mean The following objects are masked from 'package:gt': html, latex The following objects are masked from 'package:dplyr': src, summarize The following objects are masked from 'package:base': format.pval, units

```
Attaching package: 'rms'
The following object is masked from 'package:WeightIt':
    calibrate
Loading required package: survey
Loading required package: grid
Loading required package: Matrix
Loading required package: survival
Attaching package: 'survey'
The following object is masked from 'package:rms':
    calibrate
The following object is masked from 'package:Hmisc':
    deff
The following object is masked from 'package:WeightIt':
    calibrate
```

```
The following object is masked from 'package:graphics':
    dotchart
Loading required package: tibble
Loading required package: lubridate
Attaching package: 'lubridate'
The following objects are masked from 'package:base':
    date, intersect, setdiff, union
Loading required package: sensitivitymw
# ensure predictable, writable figure path + robust PNG device
knitr::opts_chunk$set(
 fig.path = "figs/",  # short local dir for figures
 dev
         = "png",
          = 144
  dpi
dir.create("figs", showWarnings = FALSE, recursive = TRUE)
\# on macOS and some setups this prevents device headaches
options(bitmapType = "cairo")
if (!requireNamespace("shapviz", quietly = TRUE) ||
    packageVersion("shapviz") < "0.2.0") {</pre>
 install.packages("shapviz") # or: remotes::install_github("ModelOriented/shapviz")
}
```

```
if (interactive() && !requireNamespace("fastshap", quietly = TRUE)) {
 options(repos = c(CRAN = "https://cran.rstudio.com/"))
 install.packages("fastshap")
if (interactive() && !requireNamespace("fastshap", quietly = TRUE)) {
  options(repos = c(CRAN = "https://cran.rstudio.com/"))
 install.packages("DALEX")
if (interactive() && !requireNamespace("fastshap", quietly = TRUE)) {
  options(repos = c(CRAN = "https://cran.rstudio.com/"))
 install.packages("shapviz")
# Make gt tables robust in PDF: full width, caption, small font
gt_pdf <- function(x, title = NULL, subtitle = NULL) {</pre>
  out <- x |>
    gt::tab_options(
                          = gt::pct(100),
= "left",
      table.width
      table.align
                        = gt::px(9),
      table.font.size
                           = gt::px(1),
      data_row.padding
      column labels.font.size = gt::px(9),
     heading.title.font.size = gt::px(10),
     heading.subtitle.font.size = gt::px(9)
    ) |>
    gt::opt align table header(align = "left")
                         out <- out |> gt::tab_caption(title)
  if (!is.null(title))
 if (!is.null(subtitle)) out <- out |> gt::tab source note(subtitle)
  out
```

Converts the data from a STATA format to rdata if the rdata file does not exist. If it does already exist, it just loads that.

```
# data_dir_name <- '/Users/blocke/Box Sync/Residency Personal Files/Scholarly Work/Locke Research Projects/abg-vbg-project/data
data_dir_name <- '/Users/reblocke/Research/abg-vbg-project/data'

rdata_file <- file.path(data_dir_name, "full_trinetx.rdata")
stata_file <- file.path(data_dir_name, "full_db.dta")

if (!dir.exists(data_dir_name)) {
    dir.create(data_dir_name)
    message("Directory 'data' created.")
} else {
    message("Directory 'data' already exists.")
}</pre>
```

Directory 'data' already exists.

```
if (file.exists(rdata_file)) {
    load(rdata_file)
    message("Loaded existing dataset from 'full_trinetx.rdata'.")
} else {
    message("RData file not found. Reading Stata dataset...")
    stata_data <- read_dta(stata_file)

    message("Extracting variable labels...")
    var_label(stata_data)

    message("Extracting value labels...")
    sapply(stata_data, function(x) if (is.labelled(x)) val_labels(x))

    save(stata_data, file = rdata_file)
    message("Dataset saved as 'full_trinetx.rdata'.")

load(rdata_file)
    message("Loaded newly saved dataset from 'full_trinetx.rdata'.")</pre>
```

```
Loaded existing dataset from 'full_trinetx.rdata'.
Creating subset_data
set.seed(123)
rows_to_keep <- round(nrow(stata_data) * 1) #0.1</pre>
subset_data <- stata_data[sample(nrow(stata_data), rows_to_keep), ]</pre>
subset_data <- subset_data %>%
 filter(encounter_type != 1)
table(subset_data$encounter_type)
     2
            3
171727 343559
dim(subset_data)
[1] 515286
              546
Generating Codebook for the Full Dataset
message("Generating codebook for the dataset...")
Generating codebook for the dataset...
```

Codebook saved as 'codebookr.docx' in the data directory.

New Variable - Death at 60 days

```
months_death_or_cens = case_when(
      !is.na(death_ym) ~ interval(enc_ym, death_ym) %/% months(1),
                       ~ interval(enc_ym, ref_ym) %/% months(1)
      TRUE
    ),
    ## 6. Remove impossible values
   months_death_or_cens = if_else(
      months_death_or_cens < 0 | months_death_or_cens > 16,
     NA_integer_, months_death_or_cens
    ),
    ## 7. Death within one or two months
    died_1mo = if_else(died == 1 & months_death_or_cens < 1, 1L, 0L),</pre>
   died_2mo = if_else(died == 1 & months_death_or_cens <= 1, 1L, 0L),</pre>
    ## 8. Month of death (missing if censored)
    death_time = if_else(died == 1, months_death_or_cens, NA_integer_),
    ## 9. Death within 60 days (new variable)
    death_60d = if_else(died == 1 & death_abs <= (encounter_date + days(60)), 1L, 0L)</pre>
  ) %>%
  select(-enc_ym, -death_ym)
subset_data <- subset_data %>%
  mutate(
    death_60d = if_else(died == 1 & death_abs <= (encounter_date + days(60)), 1L, 0L)</pre>
```

```
table(subset_data$death_60d, useNA = "ifany")
```

0 1 461485 53801

```
prop.table(table(subset_data$death_60d, useNA = "ifany"))
0.89559 0.10441
summary(subset_data$death_60d)
   Min. 1st Qu. Median
                          Mean 3rd Qu.
                                           Max.
 0.0000 0.0000 0.0000 0.1044 0.0000 1.0000
Table 1A and 1B:
# Robust derivation of analysis variables + helper for Table 1 production
# helper: label binary 0/1 → "No"/"Yes"
bin lab <- function(x) factor(x, levels = c(0, 1), labels = c("No", "Yes"))
subset_data <- subset_data %>%
  mutate(
    ## ensure 0/1 numerics (avoids factor-level coercion)
    across(c(has_abg, has_vbg, hypercap_on_abg, hypercap_on_vbg),
           ~ as.numeric(as.character(.))),
    ## derive ABG / VBG hypercapnia groups
    abg_group
    = case_when(
     has_abg == 0
                                          ~ "No ABG",
     has_abg == 1 & hypercap_on_abg == 0 ~ "ABG_NoHypercapnia",
      has_abg == 1 & hypercap_on_abg == 1 ~ "ABG_Hypercapnia",
                                          ~ "Missing"
      TRUE
```

```
),
vbg group = case when(
 has vbg == 0
                                      ~ "No VBG",
 has_vbg == 1 & hypercap_on_vbg == 0 ~ "VBG_NoHypercapnia",
 has_vbg == 1 & hypercap_on_vbg == 1 ~ "VBG_Hypercapnia",
                                      ~ "Missing"
  TRUE
),
## factorise groups with explicit NA/Missing level
abg_group = factor(
  abg_group,
 levels = c("No ABG", "ABG_NoHypercapnia", "ABG_Hypercapnia", "Missing")
),
vbg_group = factor(
 vbg_group,
 levels = c("No VBG", "VBG_NoHypercapnia", "VBG_Hypercapnia", "Missing")
),
## labelled covariates
sex_label = factor(sex, levels = c(0, 1), labels = c("Female", "Male")),
race_ethnicity_label
                       = factor(
 race ethnicity,
 levels = c(0, 1, 2, 3, 4, 5, 6),
  labels = c("White", "Black or African American", "Hispanic",
             "Asian", "American Indian", "Pacific Islander", "Unknown")
), location_label = factor(
  location,
 levels = c(0, 1, 2, 3),
  labels = c("South", "Northeast", "Midwest", "West")
), encounter_type_label = factor(
  encounter_type,
 levels = c(2, 3),
  labels = c("Emergency", "Inpatient")
),
```

```
= bin_lab(osa),
    osa_label
   asthma label = bin lab(asthma),
   copd_label
                 = bin_lab(copd),
   chf label
                = bin lab(chf),
   nmd_label
                 = bin_lab(nmd),
   phtn label = bin lab(phtn),
    ckd label
                = bin_lab(ckd),
   diabetes label = bin lab(dm)
# variables to summarise
vars <- c(</pre>
  "age_at_encounter", "curr_bmi", "sex_label", "race_ethnicity_label", "location_label",
  "osa_label", "asthma_label", "copd_label", "chf_label", "nmd_label",
  "phtn_label", "ckd_label", "diabetes_label", "encounter_type_label", "vbg_co2", "paco2"
# Table 1 constructor
make_table1 <- function(data, group_var, caption = "") {</pre>
 group_sym <- rlang::sym(group_var)</pre>
  data %>%
                                                 # drop explicit NA
   filter(!is.na(!!group_sym),
                                                 # drop "Missing" cohort
           !!group_sym != "Missing") %>%
    droplevels() %>%
                                                 # trim empty factor levels
    select(all_of(c(group_var, vars))) %>%
   gtsummary::tbl_summary(
      by = !!group_sym,
     type = list(sex_label ~ "categorical"),
      statistic = list(
        gtsummary::all_continuous() ~ "{mean} ± {sd}; {N miss}/{N obs} missing ({p_miss}%)",
       gtsummary::all_categorical() ~ "{n} ({p}%)"
      ),
      digits = list(gtsummary::all continuous() ~ 1),
```

```
missing = "no"  # no gtsummary missing column/row
) %>%
  gtsummary::modify_header(label = "**Variable**") %>%
  gtsummary::modify_caption(caption)
}

# build tables
table1A <- make_table1(subset_data, "abg_group", caption = "Table 1A: ABG cohorts")
table1B <- make_table1(subset_data, "vbg_group", caption = "Table 1B: VBG cohorts")</pre>
```

#### table1B

Generating Word Doc for Table 1A & 1B

```
ft_table1A <- as_flex_table(table1A)
ft_table1B <- as_flex_table(table1B)

doc <- read_docx() %>%
  body_add_par("Table 1A. Baseline Characteristics by ABG Group", style = "heading 1") %>%
  body_add_flextable(ft_table1A) %>%
  body_add_par("Table 1B. Baseline Characteristics by VBG Group", style = "heading 1") %>%
  body_add_par("Table 1B. Baseline Characteristics by VBG Group", style = "heading 1") %>%
  body_add_flextable(ft_table1B)

print(doc, target = "Table1_ABG_VBG.docx")
```

### NEW Table 1

```
# Status factors (column labels are taken from factor levels)
subset_data <- subset_data %>%
mutate(
```

Variable	No ABG $N = 328,044^{1}$	$ABG\_NoHypercapnia$ $N=129,429^1$	ABG_Hypercapnia
Age (years)	$58.1 \pm 18.1$ ; $0.0/328,044.0$ missing $(0.0\%)$	$60.8 \pm 17.1$ ; $0.0/129,429.0$ missing $(0.0\%)$	$\overline{62.1 \pm 16.4}$ ; 0.0/57,813.0
Current BMI kg/m2	$32.3 \pm 8.7$ ; $184,223.0/328,044.0$ missing (56.2%)	$28.6 \pm 6.9$ ; $75,826.0/129,429.0$ missing (58.6%)	$29.8 \pm 7.9; 33,496.0/57,813$
sex_label		, , ,	, , ,
Female	169,023~(52%)	57,767 (45%)	27,116 (479
Male	159,021 (48%)	71,662 (55%)	30,697 (539
race_ethnicity_label			
White	200,033 (61%)	81,357 (63%)	39,784 (699
Black or African American	62,418 (19%)	19,197 (15%)	8,082 (14%
Hispanic	$23,548 \ (7.2\%)$	$7,464 \ (5.8\%)$	2,757 (4.8%
Asian	4,880 (1.5%)	2,739(2.1%)	789 (1.4%)
American Indian	1,971 (0.6%)	1,768 (1.4%)	316 (0.5%
Pacific Islander	460 (0.1%)	162 (0.1%)	56 (<0.1%
Unknown	$34{,}73\overset{\checkmark}{4}(11\overset{\checkmark}{\%})$	$16,74\overset{\circ}{2}(13\overset{\circ}{\%})$	6,029 (10%
location_label		, , ,	,
South	138,843 (42%)	70,729 (55%)	32,694 (579
Northeast	93,209 (28%)	$23,\!262(18\%)$	12,975 (229
Midwest	22,924 (7.0%)	$10,703 \ (8.3\%)$	4,844 (8.4%
West	73,068 (22%)	24,735 (19%)	7,300 (13%
osa label	60,653 (18%)	17,709 (14%)	11,965 (219
asthma label	48,456 (15%)	13,049 (10%)	8,268 (14%
copd_label	60,214 (18%)	21,195 (16%)	18,846 (339
chf_label	59,770 (18%)	25,469 (20%)	16,219 (289
nmd_label	11,891 (3.6%)	5,861 (4.5%)	2,487 (4.3%
phtn label	$23,854 \ (7.3\%)$	10,513~(8.1%)	7,347 (13%
ckd label	54,528 (17%)	24,849 (19%)	11,769 (209
diabetes label	93,007 (28%)	37,426 (29%)	18,521 (329
encounter_type_label		, , ,	, ,
Emergency	142,713 (44%)	$19,196 \ (15\%)$	9,818 (17%
Inpatient	185,331 (56%)	$110,\!233(85\%)$	47,995 (839
VBG PCO2	$45.5 \pm 10.5$ ; $233,430.0/328,044.0$ missing (71.2%)	$42.0 \pm 11.2$ ; $91,782.0/129,429.0$ missing (70.9%)	$57.4 \pm 18.4; 40,411.0/57,813$
Arterial PCO2	$NA \pm NA$ ; 328,044.0/328,044.0 missing (100.0%)	$35.5 \pm 6.1$ ; $0.0/129,429.0$ missing $(0.0\%)$	$58.5 \pm 20.4$ ; $0.0/57,813.0$

 $<sup>^{7}</sup>$ Mean ± SD; N Missing/No. obs. missing (% Missing); n (%)

Variable	No VBG $N = 365,623^{1}$	$VBG\_NoHypercapnia\ N=105,646^1$	VBG_Hypercapnia
Age (years)	$59.4 \pm 17.8$ ; $0.0/365,623.0$ missing $(0.0\%)$	$58.1 \pm 17.8$ ; $0.0/105,646.0$ missing $(0.0\%)$	$\overline{61.0 \pm 16.7}$ ; 0.0/44,017.0
Current BMI kg/m2	$31.8 \pm 8.5$ ; $192,892.0/365,623.0$ missing (52.8%)	$28.7 \pm 7.2$ ; $69,615.0/105,646.0$ missing $(65.9\%)$	$29.3 \pm 7.9$ ; $31,038.0/44,017$
sex_label	, , ,	, , ,	
Female	184,619 (50%)	48,931 (46%)	20,356 (469
Male	181,004 (50%)	56,715 (54%)	23,661 (549)
race_ethnicity_label		. ,	
White	241,114 (66%)	55,100 (52%)	24,960 (579
Black or African American	61,814 (17%)	19,199 (18%)	8,684 (20%
Hispanic	$22,951 \ (6.3\%)$	$8,354 \ (7.9\%)$	2,464 (5.6%
Asian	$5{,}439 \; (1.5\%)$	$2,293\ (2.2\%)$	676 (1.5%
American Indian	$2{,}128\ (0.6\%)$	1,683 (1.6%)	244 (0.6%
Pacific Islander	543 (0.1%)	110 (0.1%)	25 (<0.1%
Unknown	$31{,}63\overset{\circ}{4}\ (8.7\overset{\circ}{\%})$	18,907 (18%)	6,964 (16%
location_label	, , ,	, , ,	,
South	196,774 (54%)	30,426 (29%)	15,066 (349
Northeast	65,537 (18%)	44,405 (42%)	19,504 (449)
Midwest	24,891 (6.8%)	$9,178 \ (8.7\%)$	4,402 (10%
West	78,421 (21%)	21,637 (20%)	5,045 (11%
osa_label	65,748 (18%)	15,634 (15%)	8,945 (20%
asthma label	49,810 (14%)	13,419 (13%)	6,544 (15%
copd_label	70,950 (19%)	16,459 (16%)	12,846 (299
chf_label	68,964 (19%)	20,573 (19%)	11,921 (279
nmd_label	14,796 (4.0%)	$3,754 \ (3.6\%)$	1,689 (3.8%
phtn_label	27,731 (7.6%)	8,534 (8.1%)	5,449 (12%
ckd_label	61,091 (17%)	21,290 (20%)	8,765 (20%
diabetes label	101,173 (28%)	33,665 (32%)	14,116 (329
encounter_type_label		. ,	
Emergency	124,405 (34%)	34,711 (33%)	12,611 (29%
Inpatient	241,218 (66%)	$70,935\ (67\%)$	31,406 (719
VBG PCO2	$NA \pm NA; 365,623.0/365,623.0 \text{ missing } (100.0\%)$	$40.1 \pm 6.6$ ; $0.0/105,646.0$ missing $(0.0\%)$	$60.2 \pm 12.6; 0.0/44,017.0$
Arterial PCO2	$42.4 \pm 15.5$ ; 233,430.0/365,623.0 missing (63.8%)	$38.6 \pm 15.4$ ; $68,334.0/105,646.0$ missing $(64.7\%)$	$52.7 \pm 19.6$ ; $26,280.0/44,017$

 $<sup>^{7}</sup>$ Mean ± SD; N Missing/No. obs. missing (% Missing); n (%)

```
abg_status = factor(has_abg, levels = c(0, 1),
                       labels = c("Did not get ABG", "Did get ABG")),
    vbg_status = factor(has_vbg, levels = c(0, 1),
                        labels = c("Did not get VBG", "Did get VBG"))
 )
# ABG table with "Everyone" column first
tbl1_abg <- subset_data %>%
  select(all_of(vars), abg_status) %>%
  gtsummary::tbl_summary(
   by = abg_status,
   type = list(sex_label ~ "categorical"),
   statistic = list(
      gtsummary::all_continuous() ~ "{mean} ± {sd}; {N_miss}/{N_obs} missing ({p_miss}%)",
     gtsummary::all_categorical() ~ "{n} ({p}%)"
   digits = list(gtsummary::all_continuous() ~ 1),
   missing = "no"
 ) %>%
  gtsummary::add overall(last = FALSE, col label = "Everyone") %>%
  gtsummary::modify_header(label = "**Variable**")
# VBG table (no "Everyone" here)
tbl1_vbg <- subset_data %>%
 select(all_of(vars), vbg_status) %>%
 gtsummary::tbl_summary(
    by = vbg_status,
   type = list(sex_label ~ "categorical"),
   statistic = list(
      gtsummary::all_continuous() ~ "{mean} ± {sd}; {N_miss}/{N_obs} missing ({p_miss}%)",
     gtsummary::all_categorical() ~ "{n} ({p}%)"
   ),
    digits = list(gtsummary::all_continuous() ~ 1),
   missing = "no"
```

```
gtsummary::modify_header(label = "**Variable**")

library(gtsummary)

tbl1 <- tbl_merge(
   tbls = list(tbl1_abg, tbl1_vbg)
) %>%
   modify_caption("**Table 1. Baseline summary: Everyone, ABG status, and VBG status**")

tbl1
```

#### NEW Table 2

```
# Hypercapnia factors within measured cohorts
subset_data <- subset_data %>%
 mutate(
   hyper_abg = factor(hypercap_on_abg, levels = c(1, 0),
                      labels = c("Got ABG & Hypercapnia", "Got ABG & No hypercapnia")),
   hyper_vbg = factor(hypercap_on_vbg, levels = c(1, 0),
                      labels = c("Got VBG & Hypercapnia", "Got VBG & No hypercapnia"))
# ABG cohort (has_abg == 1)
tbl2_abg <- subset_data %>%
 filter(has_abg == 1) %>%
 select(all_of(vars), hyper_abg) %>%
 gtsummary::tbl_summary(
    by = hyper_abg,
   type = list(sex_label ~ "categorical"),
   statistic = list(
      gtsummary::all_continuous() ~ "{mean} ± {sd}; {N_miss}/{N_obs} missing ({p_miss}%)",
     gtsummary::all_categorical() ~ "{n} ({p}%)"
    ),
```

		Table 1	
Variable	Everyone <sup>1</sup>	Did not get ABG $N = 328,044^{1}$	Did get ABG 1
Age (years)	$59.2 \pm 17.7$ ; $0.0/515,286.0$ missing $(0.0\%)$	$58.1 \pm 18.1$ ; $0.0/328,044.0$ missing $(0.0\%)$	$\phantom{00000000000000000000000000000000000$
Current BMI kg/m2	$31.1 \pm 8.4$ ; $293,545.0/515,286.0$ missing (57.0%)	$32.3 \pm 8.7$ ; $184,223.0/328,044.0$ missing (56.2%)	$29.0 \pm 7.2$ ; $109,322.0/187$ ,
sex_label			
Female	253,906 (49%)	169,023~(52%)	84,883 (4
Male	261,380 (51%)	159,021 (48%)	102,359 (
race_ethnicity_label			
White	321,174~(62%)	200,033 (61%)	121,141 (
Black or African American	89,697 (17%)	62,418 (19%)	27,279 (1
Hispanic	$33,769 \ (6.6\%)$	23,548 (7.2%)	10,221 (5
Asian	8,408 (1.6%)	4,880 (1.5%)	3,528 (1.
American Indian	$4,055\ (0.8\%)$	1,971 (0.6%)	2,084 (1.
Pacific Islander	678 (0.1%)	460 (0.1%)	218 (0.1
Unknown	$57,50\dot{5}$ $(11\%)$	$34,73\mathring{4}\ (11\%)$	22,771 (1
location_label	. ,	. , ,	
South	242,266 (47%)	$138,843 \ (42\%)$	103,423 (
Northeast	129,446 (25%)	93,209 (28%)	36,237 (1
Midwest	38,471 (7.5%)	$22,924 \ (7.0\%)$	15,547 (8
West	$105{,}103^{\circ}(20\%)$	73,068(22%)	32,035 (1
osa_label	90,327 (18%)	60,653 (18%)	29,674 (1
asthma_label	69,773 (14%)	48,456 (15%)	21,317 (1
copd_label	100,255 (19%)	60,214 (18%)	40,041 (2
chf_label	101,458 (20%)	59,770 (18%)	41,688 (2
nmd_label	$20,239 \ (3.9\%)$	11,891 (3.6%)	8,348 (4.
phtn_label	41,714 (8.1%)	$23,854 \ (7.3\%)$	17,860 (9
ckd label	91,146 (18%)	$54,528\ (17\%)$	36,618 (2
diabetes label	148,954 (29%)	93,007 (28%)	55,947 (3
encounter_type_label			, ,
Emergency	171,727 (33%)	142,713 (44%)	29,014 (1
Inpatient	343,559 (67%)	185,331 (56%)	158,228 (
VBG PCO2	$46.0 \pm 12.7$ ; $365,623.0/515,286.0$ missing (71.0%)	$45.5 \pm 10.5$ ; $233,430.0/328,044.0$ missing (71.2%)	$46.9 \pm 15.6$ ; $132,193.0/187$
Arterial PCO2	$42.6 \pm 16.3$ ; $328,044.0/515,286.0$ missing (63.7%)	$NA \pm NA$ ; 328,044.0/328,044.0 missing (100.0%)	$42.6 \pm 16.3$ ; $0.0/187,24$

 $<sup>^{1}</sup>$ Mean ± SD; N Missing/No. obs. missing (% Missing); n (%)

```
digits = list(gtsummary::all_continuous() ~ 1),
   missing = "no"
 ) %>%
 gtsummary::modify_header(
   label = "**Variable**",
   stat 1 = "**Got ABG & Hypercapnia**",
   stat_2 = "**Got ABG & No hypercapnia**"
# VBG cohort (has_vbg == 1)
tbl2_vbg <- subset_data %>%
 filter(has_vbg == 1) %>%
 select(all_of(vars), hyper_vbg) %>%
 gtsummary::tbl_summary(
   by = hyper_vbg,
   type = list(sex_label ~ "categorical"),
   statistic = list(
      gtsummary::all_continuous() ~ "{mean} ± {sd}; {N_miss}/{N_obs} missing ({p_miss}%)",
     gtsummary::all_categorical() ~ "{n} ({p}%)"
   ),
   digits = list(gtsummary::all_continuous() ~ 1),
   missing = "no"
 ) %>%
 gtsummary::modify_header(
   label = "**Variable**",
   stat_1 = "**Got VBG & Hypercapnia**",
   stat_2 = "**Got VBG & No hypercapnia**"
# Merge side-by-side (no spanners; 4 requested columns)
table2 <- gtsummary::tbl_merge(</pre>
 tbls = list(tbl2_abg, tbl2_vbg),
 tab_spanner = c(NULL, NULL)
) %>%
```

```
gtsummary::modify_caption("**Table 2. Baseline summary by hypercapnia within ABG and VBG cohorts**")
table2
```

Generating Word Docs for New Table 1 and 2

```
library(gtsummary)
library(officer)

# gtsummary objects (example: table1, table2)
ft1 <- as_flex_table(tbl1)
ft2 <- as_flex_table(table2)

doc <- read_docx() %>%
    body_add_par("Table 1", style = "heading 1") %>%
    body_add_flextable(ft1) %>%
    body_add_par("Table 2", style = "heading 1") %>%
    body_add_flextable(ft2)

print(doc, target = "Tables.docx")
```

## Unweighted, Hypercapnia (binary yes/no) Simple (1 predictor) Regressions:

Unweighted, ABG Group: hypercapnia treated as a binary (yes/no) predictor

```
logit_intubated_abg <- glm(imv_proc ~ hypercap_on_abg, data = subset_data, family = binomial)
summary(logit_intubated_abg)</pre>
```

```
Call:
glm(formula = imv_proc ~ hypercap_on_abg, family = binomial,
    data = subset_data)
```

	Tal		
Variable	Got ABG & Hypercapnia <sup>1</sup>	Got ABG & No hypercapnia <sup>1</sup>	Got VBG & Hyper
Age (years)	$62.1 \pm 16.4$ ; $0.0/57,813.0$ missing $(0.0\%)$	$60.8 \pm 17.1$ ; $0.0/129,429.0$ missing $(0.0\%)$	$61.0 \pm 16.7$ ; 0.0/44,017.0 m
Current BMI kg/m2	$29.8 \pm 7.9$ ; $33,496.0/57,813.0$ missing $(57.9\%)$	$28.6 \pm 6.9$ ; $75,826.0/129,429.0$ missing (58.6%)	$29.3 \pm 7.9$ ; $31,038.0/44,017.0$
sex_label		, ,	, , ,
Female	27,116 (47%)	57,767 (45%)	20,356 (46%)
Male	30,697 (53%)	71,662 (55%)	23,661 (54%)
race_ethnicity_label			
White	39,784 (69%)	81,357 (63%)	24,960 (57%)
Black or African American	8,082 (14%)	19,197 (15%)	8,684 (20%)
Hispanic	2,757 (4.8%)	7,464 (5.8%)	2,464 (5.6%)
Asian	789 (1.4%)	2,739(2.1%)	676 (1.5%)
American Indian	$316\ (0.5\%)$	1,768 (1.4%)	244 (0.6%)
Pacific Islander	56 (<0.1%)	162 (0.1%)	25~(<0.1%)
$\operatorname{Unknown}$	6,029 (10%)	$16,74\dot{2}\ (13\%)$	6,964 (16%)
location_label			
South	32,694 (57%)	70,729 (55%)	15,066 (34%)
Northeast	12,975 (22%)	23,262 (18%)	19,504 (44%)
Midwest	4,844 (8.4%)	$10,703 \ (8.3\%)$	4,402 (10%)
West	7,300 (13%)	24,735 (19%)	5,045 (11%)
osa_label	11,965 (21%)	17,709 (14%)	8,945 (20%)
asthma_label	8,268 (14%)	13,049 (10%)	6,544 (15%)
copd_label	18,846 (33%)	21,195 (16%)	12,846 (29%)
chf_label	16,219 (28%)	25,469 (20%)	11,921 (27%)
nmd_label	2,487 (4.3%)	5,861 (4.5%)	1,689 (3.8%)
phtn_label	7,347 (13%)	10,513~(8.1%)	5,449 (12%)
ckd_label	11,769 (20%)	24,849 (19%)	8,765 (20%)
diabetes_label	$18,521 \ (32\%)$	37,426 (29%)	14,116 (32%)
encounter_type_label			. ,
Emergency	9,818 (17%)	19,196~(15%)	12,611 (29%)
Inpatient	47,995 (83%)	$110,\!233(85\%)$	31,406 (71%)
VBG PCO2	$57.4 \pm 18.4$ ; $40,411.0/57,813.0$ missing (69.9%)	$42.0 \pm 11.2$ ; $91,782.0/129,429.0$ missing (70.9%)	$60.2 \pm 12.6$ ; $0.0/44,017.0$ m
Arterial PCO2	$58.5 \pm 20.4$ ; 0.0/57,813.0 missing (0.0%)	$35.5 \pm 6.1$ ; $0.0/129,429.0$ missing $(0.0\%)$	$52.7 \pm 19.6$ ; $26,280.0/44,017.0$

 $<sup>\</sup>overline{^{1}}$ Mean ± SD; N Missing/No. obs. missing (% Missing); n (%)

#### Coefficients:

```
Estimate Std. Error z value Pr(>|z|)

(Intercept) -2.275471 0.005086 -447.4 <2e-16 ***
hypercap_on_abg 1.259993 0.010700 117.8 <2e-16 ***
---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 362580 on 515285 degrees of freedom
Residual deviance: 350435 on 515284 degrees of freedom
AIC: 350439
```

Number of Fisher Scoring iterations: 5

```
tidy(logit_intubated_abg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept) hypercap_on_abg	$0.1027485 \\ 3.5253956$	0.0050863 $0.0106997$	-447.3684 117.7601	0	0.1017279 $3.4521736$	0.1037765 $3.6000445$

```
logit_niv_abg <- glm(niv_proc ~ hypercap_on_abg, data = subset_data, family = binomial)
summary(logit_niv_abg)</pre>
```

```
Call:
```

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)

(Intercept) -2.860418 0.006533 -437.84 <2e-16 \*\*\*
hypercap\_on\_abg 1.201665 0.013093 91.78 <2e-16 \*\*\*
--
Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 250639 on 515285 degrees of freedom
Residual deviance: 243461 on 515284 degrees of freedom

AIC: 243465

Number of Fisher Scoring iterations: 5

```
tidy(logit_niv_abg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept) hypercap_on_abg	0.0572448 $3.3256498$	0.0065330 $0.0130929$	-437.84069 91.77958	0	$0.0565151 \\ 3.2412752$	0.0579811 $3.4119744$

```
logit_death_abg <- glm(death_60d ~ hypercap_on_abg, data = subset_data, family = binomial)
summary(logit_death_abg)</pre>
```

```
Call:
```

```
glm(formula = death_60d ~ hypercap_on_abg, family = binomial,
    data = subset_data)
```

#### Coefficients:

```
Estimate Std. Error z value Pr(>|z|)

(Intercept) -2.258743 0.005052 -447.11 <2e-16 ***
hypercap_on_abg 0.756240 0.011903 63.53 <2e-16 ***
---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 344897 on 515285 degrees of freedom
Residual deviance: 341286 on 515284 degrees of freedom
AIC: 341290
```

Number of Fisher Scoring iterations: 5

```
tidy(logit_death_abg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.1044817	0.0050519	-447.10674	0	0.1034509	0.105520
hypercap_on_abg	2.1302521	0.0119030	63.53375	0	2.0810535	2.180455

```
logit_icd_abg <- glm(hypercap_resp_failure ~ hypercap_on_abg, data = subset_data, family = binomial)
summary(logit_icd_abg)</pre>
```

```
Call:
```

```
glm(formula = hypercap_resp_failure ~ hypercap_on_abg, family = binomial,
    data = subset_data)
```

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)

```
(Intercept) -3.355697 0.008192 -409.6 <2e-16 ***
hypercap_on_abg 2.175594 0.012779 170.2 <2e-16 ***
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 223278 on 515285 degrees of freedom
Residual deviance: 197920 on 515284 degrees of freedom
AIC: 197924
```

Number of Fisher Scoring iterations: 6

```
tidy(logit_icd_abg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.034885	0.0081920	-409.6314	0	0.034328	0.0354483
hypercap_on_abg	8.807416	0.0127795	170.2414	0	8.589528	9.0307810

Display the regression coefficients for the binary (hypercapnia yes/no) predictor logistic regressions

	Intubated	NIV	Death	ICD Hyper
hypercap_on_abg	3.53	3.33	2.13	8.81
	(3.45, 3.60)	(3.24, 3.41)	(2.08, 2.18)	(8.59, 9.03)

```
coef_omit = "(Intercept)",
gof_omit = ".*",  # drop all goodness-of-fit rows
fmt = 2,  # 2 decimal places everywhere
output = "gt"
) |>
gt_pdf(title = "Odds Ratios for ABG Hypercapnia (>45 mmHg)'s association with...")
```

Profiled confidence intervals may take longer time to compute.

Use `ci\_method="wald"` for faster computation of CIs.

Profiled confidence intervals may take longer time to compute.

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Use `ci\_method="wald"` for faster computation of CIs.

Profiled confidence intervals may take longer time to compute.

Use `ci\_method="wald"` for faster computation of CIs.

## Unweighted VBG Group

```
logit_intubated_vbg <- glm(imv_proc ~ hypercap_on_vbg, data = subset_data, family = binomial)
summary(logit_intubated_vbg)</pre>
```

#### Call:

```
glm(formula = imv_proc ~ hypercap_on_vbg, family = binomial,
    data = subset_data)
```

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)

```
(Intercept) -2.134026  0.004735 -450.69  <2e-16 ***
hypercap_on_vbg  0.648004  0.013168  49.21  <2e-16 ***
---
Signif. codes:  0 '***'  0.001 '**'  0.05 '.'  0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 362580  on 515285  degrees of freedom
Residual deviance: 360405  on 515284  degrees of freedom
AIC: 360409

Number of Fisher Scoring iterations: 4
```

```
tidy(logit_intubated_vbg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.1183598	0.0047351	-450.68580	0	0.1172651	0.1194621
hypercap_on_vbg	1.9117219	0.0131682	49.20969	0	1.8629159	1.9616038

```
logit_niv_vbg <- glm(niv_proc ~ hypercap_on_vbg, data = subset_data, family = binomial)
summary(logit_niv_vbg)</pre>
```

```
Call:
```

```
glm(formula = niv_proc ~ hypercap_on_vbg, family = binomial,
    data = subset_data)
```

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)
(Intercept) -2.735699 0.006091 -449.13 <2e-16 \*\*\*

```
hypercap_on_vbg 0.750555 0.015845 47.37 <2e-16 ***
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 250639 on 515285 degrees of freedom
Residual deviance: 248688 on 515284 degrees of freedom
AIC: 248692
```

Number of Fisher Scoring iterations: 5

```
tidy(logit_niv_vbg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept) hypercap_on_vbg	$0.0648486 \\ 2.1181752$	0.0060911 0.0158446	-449.12691 47.36965	0	$0.0640777 \\ 2.0532293$	$0.0656261 \\ 2.1848014$

```
logit_death_vbg <- glm(death_60d ~ hypercap_on_vbg, data = subset_data, family = binomial)
summary(logit_death_vbg)</pre>
```

```
Call:
```

```
glm(formula = death_60d ~ hypercap_on_vbg, family = binomial,
    data = subset_data)
```

#### Coefficients:

```
Estimate Std. Error z value Pr(>|z|)
(Intercept) -2.197765 0.004856 -452.56 <2e-16 ***
hypercap_on_vbg 0.479895 0.014131 33.96 <2e-16 ***
```

---

```
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 344897 on 515285 degrees of freedom Residual deviance: 343841 on 515284 degrees of freedom

AIC: 343845

Number of Fisher Scoring iterations: 4

```
tidy(logit_death_vbg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE) # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.1110511	0.0048563	-452.56297	0	0.1099977	0.1121119
hypercap_on_vbg	1.6159045	0.0141315	33.95924	0	1.5716549	1.6611746

```
logit_icd_vbg <- glm(hypercap_resp_failure ~ hypercap_on_vbg, data = subset_data, family = binomial)
summary(logit_icd_vbg)</pre>
```

#### Call:

```
glm(formula = hypercap_resp_failure ~ hypercap_on_vbg, family = binomial,
    data = subset_data)
```

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)
(Intercept) -3.136462 0.007293 -430.1 <2e-16 \*\*\*
hypercap\_on\_vbg 1.831609 0.013731 133.4 <2e-16 \*\*\*

```
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 223278 on 515285 degrees of freedom
Residual deviance: 208772 on 515284 degrees of freedom
AIC: 208776

Number of Fisher Scoring iterations: 6
```

```
tidy(logit_icd_vbg,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.0434362	0.0072930	-430.0654	0	0.0428184	0.0440602
hypercap_on_vbg	6.2439262	0.0137314	133.3879	0	6.0779584	6.4140808

Display model coefficients for binary hypercapnia on VBG logistic regression

```
modelsummary(
 list("Intubated" = logit intubated vbg,
      "NIV"
               = logit_niv_vbg,
      "Death" = logit_death_vbg,
      "ICD Hyper" = logit_icd_vbg),
 exponentiate = TRUE,
 conf_level = 0.95,
 estimate = "{estimate}",
 statistic = "({conf.low}, {conf.high})",
 coef_omit = "(Intercept)",
             = ".*",
 gof_omit
                                         # drop all goodness-of-fit rows
              = 2,
                                         # 2 decimal places everywhere
 fmt
```

	Intubated	NIV	Death	ICD Hyper
hypercap_on_vbg	1.91	2.12	1.62	6.24
	(1.86, 1.96)	(2.05, 2.18)	(1.57, 1.66)	(6.08, 6.41)

```
output = "gt"
) |>
  gt_pdf(title = "Odds Ratios for VBG Hypercapnia (>45 mmHg)'s association with...")
```

Profiled confidence intervals may take longer time to compute.

Use `ci\_method="wald"` for faster computation of CIs.

Profiled confidence intervals may take longer time to compute.

Use `ci\_method="wald"` for faster computation of CIs.

Profiled confidence intervals may take longer time to compute.

Use `ci\_method="wald"` for faster computation of CIs.

Profiled confidence intervals may take longer time to compute.  $% \left( 1\right) =\left( 1\right) \left( 1\right)$ 

Use `ci\_method="wald"` for faster computation of CIs.

Calculated ABG from VBG Using Farkas equation - binary predictor

```
subset_data <- subset_data %>%
  mutate(
    calc_abg = vbg_co2 - (0.22 * (93 - vbg_o2sat))
)
subset_data <- subset_data %>%
  mutate(
    hypercapnia_calc = ifelse(calc_abg > 45, 1, 0)
)
with(subset_data, table(hypercapnia_calc,niv_proc))
```

```
niv_proc
hypercapnia_calc 0 1
0 47526 2877
1 14146 2599
```

```
logit_intubated_calc <- glm(imv_proc ~ hypercapnia_calc, data = subset_data, family = binomial)
summary(logit_intubated_calc)</pre>
```

```
Call:
glm(formula = imv_proc ~ hypercapnia_calc, family = binomial,
   data = subset_data)
Coefficients:
                Estimate Std. Error z value Pr(>|z|)
(Intercept)
                -1.95271
                          0.01350 -144.62 <2e-16 ***
hypercapnia_calc 0.53248 0.02373 22.43 <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 54823 on 67147 degrees of freedom
Residual deviance: 54341 on 67146 degrees of freedom
  (448138 observations deleted due to missingness)
AIC: 54345
Number of Fisher Scoring iterations: 4
```

```
tidy(logit_intubated_calc,
    exponentiate = TRUE,  # turns log-odds \rightarrow OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.1418894	0.0135026	-144.61661	0	0.1381714	0.1456819
hypercapnia_calc	1.7031430	0.0237343	22.43489	0	1.6256101	1.7841152

```
logit_niv_calc <- glm(niv_proc ~ hypercapnia_calc, data = subset_data, family = binomial)
summary(logit_niv_calc)</pre>
```

## Call:

```
glm(formula = niv_proc ~ hypercapnia_calc, family = binomial,
    data = subset_data)
```

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)

(Intercept) -2.80453 0.01920 -146.07 <2e-16 \*\*\*

hypercapnia\_calc 1.11022 0.02871 38.67 <2e-16 \*\*\*

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 37944 on 67147 degrees of freedom Residual deviance: 36518 on 67146 degrees of freedom (448138 observations deleted due to missingness)

AIC: 36522

Number of Fisher Scoring iterations: 5

```
tidy(logit_niv_calc,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int  = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.0605353	0.0191993	-146.07446	0	0.0582873	0.0628436
hypercapnia_calc	3.0350373	0.0287066	38.67487	0	2.8689300	3.2106486

```
logit_death_calc <- glm(death_60d ~ hypercapnia_calc, data = subset_data, family = binomial)
summary(logit_death_calc)</pre>
```

```
Call:
glm(formula = death_60d ~ hypercapnia_calc, family = binomial,
   data = subset_data)
Coefficients:
                Estimate Std. Error z value Pr(>|z|)
(Intercept)
                -1.78892
                           0.01272 -140.681 < 2e-16 ***
hypercapnia_calc 0.15484 0.02447 6.327 2.49e-10 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 56349 on 67147 degrees of freedom
Residual deviance: 56310 on 67146 degrees of freedom
  (448138 observations deleted due to missingness)
AIC: 56314
Number of Fisher Scoring iterations: 4
```

```
tidy(logit_death_calc,
    exponentiate = TRUE,  # turns log-odds \rightarrow OR
    conf.int = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.1671414	0.0127161	-140.68135	0	0.1630151	0.1713468
hypercapnia_calc	1.1674692	0.0244711	6.32739	0	1.1126631	1.2246862

```
logit_icd_calc <- glm(hypercap_resp_failure ~ hypercapnia_calc, data = subset_data, family = binomial)
summary(logit_icd_calc)</pre>
```

```
Call:
```

```
glm(formula = hypercap_resp_failure ~ hypercapnia_calc, family = binomial,
    data = subset_data)
```

### Coefficients:

Estimate Std. Error z value Pr(>|z|)

(Intercept) -3.17657 0.02271 -139.85 <2e-16 \*\*\* hypercapnia\_calc 2.23074 0.02850 78.27 <2e-16 \*\*\*

---

Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 43605 on 67147 degrees of freedom Residual deviance: 36797 on 67146 degrees of freedom (448138 observations deleted due to missingness)

AIC: 36801

Number of Fisher Scoring iterations: 6

```
tidy(logit_icd_calc,
    exponentiate = TRUE,  # turns log-odds → OR
    conf.int  = TRUE)  # adds 95 % CI
```

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	0.0417287	0.0227148	-139.84562	0	0.0398994	0.0436152
hypercapnia_calc	9.3067708	0.0285021	78.26601	0	8.8025311	9.8430989

	Intubated	NIV	Death	ICD Hyper
hypercapnia_calc	1.70	3.04	1.17	9.31
	(1.63, 1.78)	(2.87, 3.21)	(1.11, 1.22)	(8.80, 9.84)

Display regression coefficients for binary Farkas adjustment (hypercapnia yes/no as predictor)

```
modelsummary(
 list("Intubated" = logit_intubated_calc,
      "NIV"
                 = logit_niv_calc,
              = logit death calc,
      "Death"
      "ICD Hyper" = logit_icd_calc),
 exponentiate = TRUE,
 conf level = 0.95,
 estimate = "{estimate}",
 statistic = "({conf.low}, {conf.high})",
 coef omit = "(Intercept)",
 gof_omit = ".*",
                                    # drop all goodness-of-fit rows
 fmt
             = 2,
                                       # 2 decimal places everywhere
             = "gt"
 output
) |>
 gt_pdf(title = "Odds Ratios for Calculated Hypercapnia (>45 mmHg)'s association with...")
```

Profiled confidence intervals may take longer time to compute.

Use `ci\_method="wald"` for faster computation of CIs.

Profiled confidence intervals may take longer time to compute.

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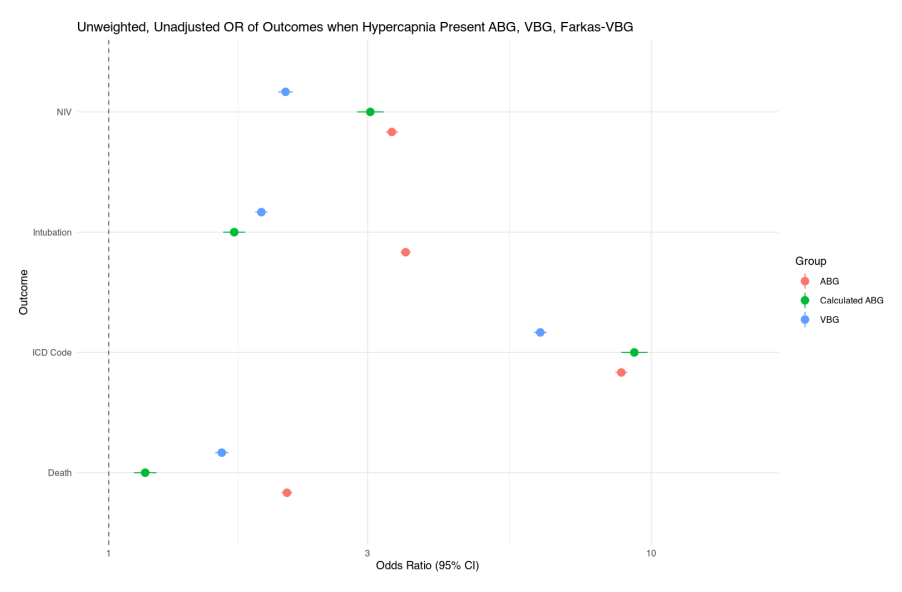
Use `ci\_method="wald"` for faster computation of CIs.

Odds Ratio Graph of all 3 simple, binary-predictor logistic regressions

```
tidy_with_labels <- function(model, group_label, outcome_label) {</pre>
 tidy(model, exponentiate = TRUE, conf.int = TRUE) %>%
    filter(term == "hypercap_on_abg" | term == "hypercap_on_vbg" | term == "hypercapnia_calc") %>%
   mutate(
      group = group_label,
      outcome = outcome label
}
# --- ABG Models ---
abg_intub <- tidy_with labels(glm(imv_proc ~ hypercap_on_abg, data = subset_data, family = binomial), "ABG", "Intubation")
abg_niv <- tidy_with labels(glm(niv_proc ~ hypercap_on abg, data = subset_data, family = binomial), "ABG", "NIV")
abg_death <- tidy_with_labels(glm(death_60d ~ hypercap_on_abg, data = subset_data, family = binomial), "ABG", "Death")
abg_icd <- tidy_with_labels(glm(hypercap_resp_failure ~ hypercap_on_abg, data = subset_data, family = binomial), "ABG", "ICI
# --- VBG Models ---
vbg_intub <- tidy_with_labels(glm(imv_proc ~ hypercap_on_vbg, data = subset_data, family = binomial), "VBG", "Intubation")</pre>
vbg_niv <- tidy_with_labels(glm(niv_proc ~ hypercap_on_vbg, data = subset_data, family = binomial), "VBG", "NIV")</pre>
vbg_death <- tidy_with_labels(glm(death_60d ~ hypercap_on_vbg, data = subset_data, family = binomial), "VBG", "Death")
vbg_icd <- tidy_with_labels(glm(hypercap_resp_failure ~ hypercap_on_vbg, data = subset_data, family = binomial), "VBG", "ICI
# --- Calculated ABG Models ---
calc_intub <- tidy_with_labels(glm(imv_proc ~ hypercapnia_calc, data = subset_data, family = binomial), "Calculated ABG", "Int
calc_niv <- tidy_with_labels(glm(niv_proc ~ hypercapnia_calc, data = subset_data, family = binomial), "Calculated ABG", "NIV</pre>
calc_death <- tidy_with_labels(glm(death_60d ~ hypercapnia_calc, data = subset_data, family = binomial), "Calculated ABG", "De
calc_icd <- tidy_with labels(glm(hypercap_resp_failure ~ hypercapnia_calc, data = subset_data, family = binomial), "Calculat
# --- Combine all model results ---
combined_or_df <- bind_rows(</pre>
  abg_intub, abg_niv, abg_death, abg_icd,
 vbg_intub, vbg_niv, vbg_death, vbg_icd,
 calc intub, calc niv, calc death, calc icd
```

```
ggplot(combined_or_df, aes(x = outcome, y = estimate, ymin = conf.low, ymax = conf.high, color = group)) +
    geom_pointrange(position = position_dodge(width = 0.5), size = 0.6) +
    geom_hline(yintercept = 1, linetype = "dashed", color = "gray40") +
    coord_flip() +
    labs(
        title = "Unweighted, Unadjusted OR of Outcomes when Hypercapnia Present ABG, VBG, Farkas-VBG ",
        x = "Outcome",
        y = "Odds Ratio (95% CI)",
        color = "Group"
    ) +
    scale_y_log10(limits = c(-0.5, 15)) + # optional log scale for better spacing
    theme_minimal(base_size = 10)
```

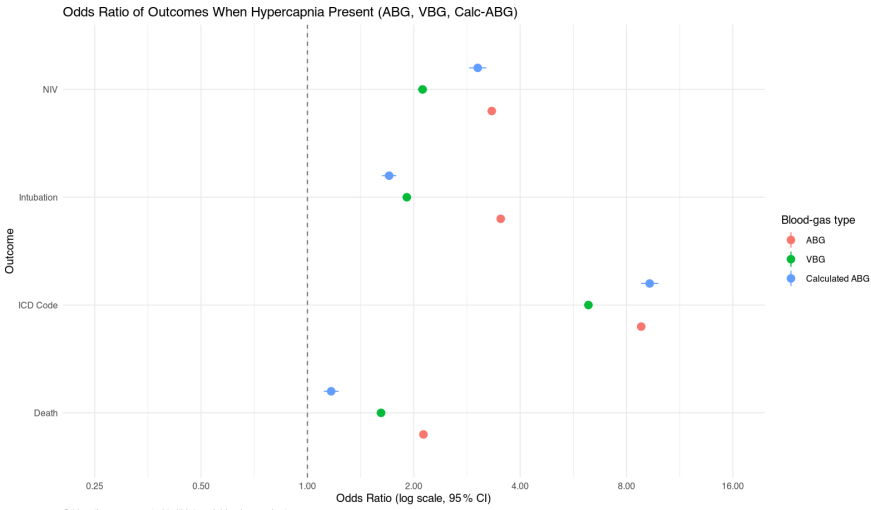
Warning in transform\$transform(limits): NaNs produced



combined\_or\_df\$group <- factor(combined\_or\_df\$group,
 levels = c("ABG", "VBG", "Calculated ABG"))</pre>

```
prerequisites
# order groups before plotting
combined_or_df$group <- factor(</pre>
 combined_or_df$group,
 levels = c("ABG", "VBG", "Calculated ABG")
# plot
ggplot(
  combined_or_df,
  aes(
   X
        = outcome,
        = estimate,
   ymin = conf.low,
   ymax = conf.high,
   color = group
 geom_pointrange(
   position = position_dodge(width = 0.6),
   size
         = 0.6
  ) +
  geom_hline(yintercept = 1, linetype = "dashed", colour = "grey40") +
  ## NOTE: scale_y_log10 applies to the axis that *becomes horizontal* after coord_flip()
  scale_y_log10(
   breaks = c(0.25, 0.5, 1, 2, 4, 8, 16),
   limits = c(0.25, 16),
   labels = number_format(accuracy = 0.01)
  ) +
 coord_flip() +
  labs(
   title = "Odds Ratio of Outcomes When Hypercapnia Present (ABG, VBG, Calc-ABG)",
          = "Outcome",
```

```
y = "Odds Ratio (log scale, 95 % CI)",
color = "Blood-gas type",
caption = paste(
   "Odds ratios are computed *within* each blood-gas cohort.",
   "Numerator = patients who received that blood-gas and **had** hypercapnia;",
   "denominator = patients who received the same blood-gas and **did not** have hypercapnia.",
   "Because the underlying cohorts differ (ABG, VBG, Calculated ABG),",
   "denominators are not identical across groups.",
   sep = "\n"
   )
) +
theme_minimal(base_size = 10) +
theme(plot.caption = element_text(hjust = 0))
```



Odds ratios are computed \*within\* each blood-gas cohort.

Numerator = patients who received that blood-gas and \*\*had\*\* hypercapnia;
denominator = patients who received the same blood-gas and \*\*did not\*\* have hypercapnia.

Because the underlying cohorts differ (ABG, VBG, Calculated ABG),
denominators are not identical across groups.

Now doing 3 groups instead of binary (above, normal and below)

```
subset_data <- subset_data %>%
  mutate(
    pco2_cat_abg = case_when(
      !is.na(paco2) & paco2 < 35 ~ "Below normal",
      !is.na(paco2) & paco2 > 45 ~ "Above normal",
      !is.na(paco2)
                                 ~ "Normal"
    ),
   pco2_cat_vbg = case_when(
      !is.na(vbg_co2) & vbg_co2 < 35 ~ "Below normal",
     !is.na(vbg_co2) & vbg_co2 > 50 ~ "Above normal",
     !is.na(vbg_co2)
                                    ~ "Normal"
    ),
    pco2_cat_calc = case_when(
      !is.na(calc_abg) & calc_abg < 35 ~ "Below normal",
     !is.na(calc_abg) & calc_abg > 45 ~ "Above normal",
     !is.na(calc_abg)
                                     ~ "Normal"
   )
 ) %>%
  mutate(
   across(starts_with("pco2_cat"),
           ~factor(.x, levels = c("Normal", "Below normal", "Above normal")))
  )
library(broom)
library(dplyr)
run_logit <- function(data, outcome, exposure, group_name) {</pre>
 f <- as.formula(paste(outcome, "~", exposure))</pre>
 glm(f, data = data, family = binomial) %>%
    tidy(exponentiate = TRUE, conf.int = TRUE) %>%
   filter(term != "(Intercept)") %>%
   mutate(
      outcome = outcome,
      group = group_name
```

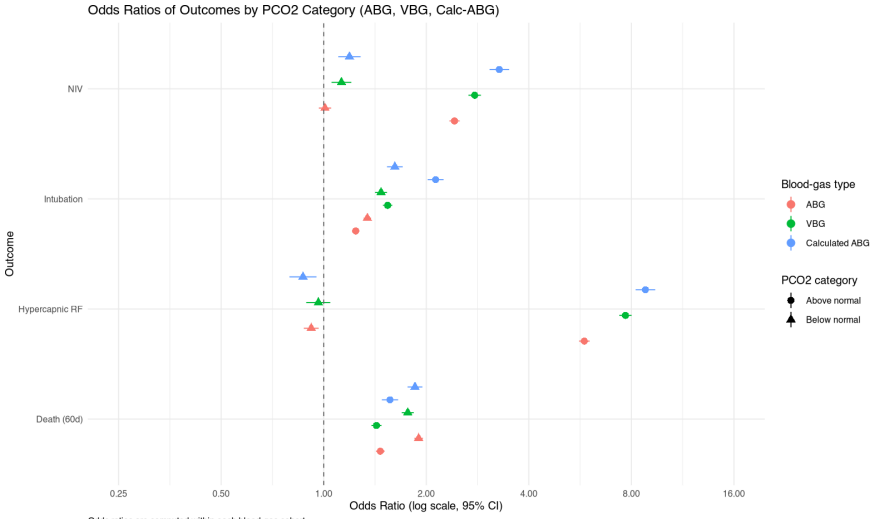
```
outcomes <- c("imv_proc", "niv_proc", "death_60d", "hypercap_resp_failure")</pre>
results <- bind rows(
 lapply(outcomes, function(o) run_logit(subset_data, o, "pco2_cat_abg", "ABG")),
 lapply(outcomes, function(o) run_logit(subset_data, o, "pco2_cat_vbg", "VBG")),
 lapply(outcomes, function(o) run_logit(subset_data, o, "pco2_cat_calc", "Calculated ABG"))
combined_or_df <- results %>%
 mutate(
    exposure = recode(term,
                      "pco2_cat_abgBelow normal" = "Below normal",
                      "pco2_cat_abgAbove normal" = "Above normal",
                      "pco2_cat_vbgBelow normal" = "Below normal",
                      "pco2 cat vbgAbove normal" = "Above normal",
                      "pco2_cat_calcBelow normal" = "Below normal",
                      "pco2 cat calcAbove normal" = "Above normal"),
    outcome = recode(outcome.
                     imv proc = "Intubation",
                    niv_proc = "NIV",
                     death 60d = "Death (60d)",
                    hypercap_resp_failure = "Hypercapnic RF")
 ) %>%
 select(outcome, group, exposure, estimate, conf.low, conf.high)
```

```
library(scales)

combined_or_df$group <- factor(
   combined_or_df$group,
   levels = c("ABG", "VBG", "Calculated ABG")
)</pre>
```

```
ggplot(
  combined_or_df,
  aes(
   X
          = outcome,
   y = estimate,
   ymin = conf.low,
   ymax = conf.high,
   color = group,
   shape = exposure
 geom_pointrange(
   position = position_dodge(width = 0.7),
            = 0.6
   size
 ) +
 geom_hline(yintercept = 1, linetype = "dashed", colour = "grey40") +
  scale_y_log10(
   breaks = c(0.25, 0.5, 1, 2, 4, 8, 16),
   limits = c(0.25, 16),
   labels = number_format(accuracy = 0.01)
 ) +
 coord_flip() +
 labs(
   title = "Odds Ratios of Outcomes by PCO2 Category (ABG, VBG, Calc-ABG)",
          = "Outcome",
          = "Odds Ratio (log scale, 95% CI)",
    color = "Blood-gas type",
    shape = "PCO2 category",
    caption = paste(
     "Odds ratios are computed within each blood-gas cohort.",
     "Reference = patients in the normal PCO2 range.",
     "Below normal: <35 mmHg. Above normal: >45 mmHg (ABG, Calc-ABG) or >50 mmHg (VBG).",
     "Because the underlying cohorts differ (ABG, VBG, Calculated ABG), denominators are not identical across groups.",
```

```
sep = "\n"
)
) +
theme_minimal(base_size = 10) +
theme(plot.caption = element_text(hjust = 0))
```



Odds ratios are computed within each blood-gas cohort.
Reference = patients in the normal PCO2 range.
Below normal: <35 mmHg. Above normal: >45 mmHg (ABG, Calc-ABG) or >50 mmHg (VBG).
Because the underlying cohorts differ (ABG, VBG, Calculated ABG), denominators are not identical across groups.

# Restricted Cubic Spline Regressions

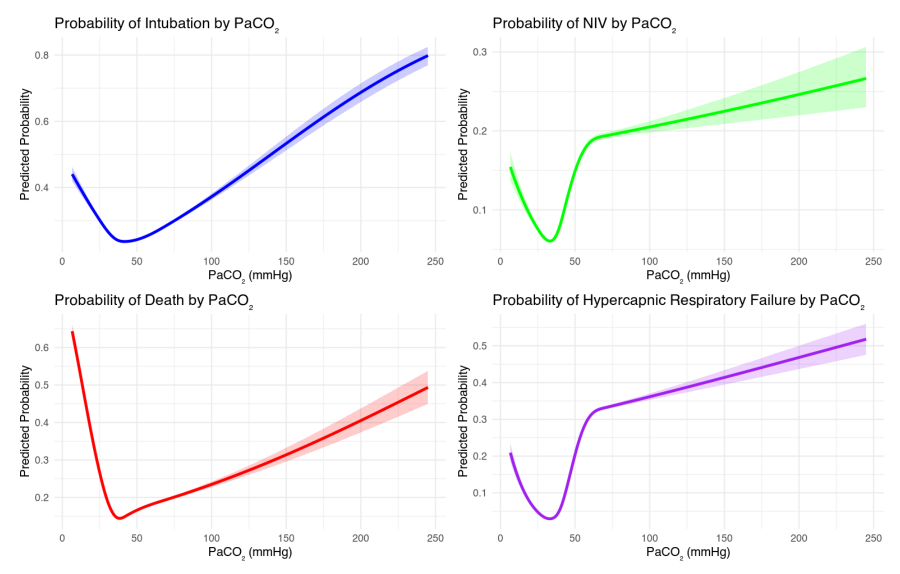
```
# ABG spline dataset
subset_data_abg <- subset_data %>%
   select(paco2, imv_proc, niv_proc, death_60d, hypercap_resp_failure) %>%
   filter(!is.na(paco2))

dd_abg <- datadist(subset_data_abg)
options(datadist = "dd_abg")</pre>
```

Unweighted, Restricted Cubic Spline Regression - ABG by PaCO2

Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0. i Please use `linewidth` instead.

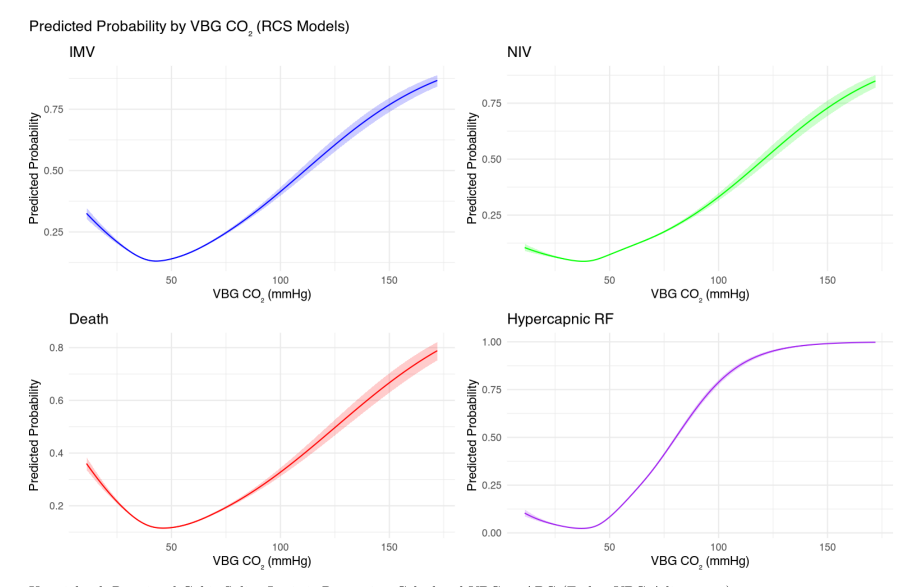
```
fit_death <- lrm(death_60d ~ rcs(paco2, 4), data = subset_data_abg)</pre>
pred death <- as.data.frame(Predict(fit death, paco2, fun = plogis))</pre>
plot_death <- ggplot(pred_death, aes(x = paco2, y = yhat)) +</pre>
  geom_line(color = "red", size = 1.2) +
  geom_ribbon(aes(ymin = lower, ymax = upper), fill = "red", alpha = 0.2) +
  labs(title = "Probability of Death by PaCO",
       x = "PaCO (mmHg)", y = "Predicted Probability") +
  theme_minimal()
fit hcrf <- lrm(hypercap_resp_failure ~ rcs(paco2, 4), data = subset_data_abg)
pred_hcrf <- as.data.frame(Predict(fit_hcrf, paco2, fun = plogis))</pre>
plot_hcrf <- ggplot(pred_hcrf, aes(x = paco2, y = yhat)) +</pre>
  geom_line(color = "purple", size = 1.2) +
  geom_ribbon(aes(ymin = lower, ymax = upper), fill = "purple", alpha = 0.2) +
  labs(title = "Probability of Hypercapnic Respiratory Failure by PaCO",
       x = "PaCO (mmHg)", y = "Predicted Probability") +
  theme_minimal()
(plot_imv | plot_niv) / (plot_death | plot_hcrf)
```



Unweighted, Restricted Cubic Spline - VBG

```
# --- VBG dataset ---
subset_data_vbg <- subset_data %>%
  dplyr::select(vbg_co2, imv_proc, niv_proc, death_60d, hypercap_resp_failure) %>%
  dplyr::filter(!is.na(vbg_co2) & complete.cases(.))
dd vbg <- datadist(subset data vbg)</pre>
                                      # create datadist for VBG
# activate when doing VBG models:
options(datadist = "dd vbg")
subset_data_vbg <- subset_data %>%
  select(vbg_co2, imv_proc, niv_proc, death_60d, hypercap_resp_failure) %>%
  filter(!is.na(vbg_co2) & complete.cases(.))
dd <- datadist(subset_data_vbg)</pre>
options(datadist = "dd")
fit_imv_vbg <- lrm(imv_proc ~ rcs(vbg_co2, 4), data = subset_data_vbg)</pre>
fit_niv_vbg <- lrm(niv_proc ~ rcs(vbg_co2, 4), data = subset_data_vbg)</pre>
fit_death_vbg <- lrm(death_60d ~ rcs(vbg_co2, 4), data = subset_data_vbg)</pre>
fit_hcrf_vbg <- lrm(hypercap_resp_failure ~ rcs(vbg_co2, 4), data = subset_data_vbg)</pre>
pred_imv_vbg <- as.data.frame(Predict(fit_imv_vbg, vbg_co2, fun = plogis))</pre>
pred niv vbg <- as.data.frame(Predict(fit niv vbg, vbg co2, fun = plogis))</pre>
pred_death_vbg <- as.data.frame(Predict(fit_death_vbg, vbg_co2, fun = plogis))</pre>
pred hcrf vbg <- as.data.frame(Predict(fit hcrf vbg, vbg co2, fun = plogis))</pre>
plot_imv_vbg <- ggplot(pred_imv_vbg, aes(x = vbg_co2, y = yhat)) +</pre>
  geom line(color = "blue") +
  geom ribbon(aes(ymin = lower, ymax = upper), fill = "blue", alpha = 0.2) +
  labs(title = "IMV", x = "VBG CO (mmHg)", y = "Predicted Probability") +
  theme minimal()
plot_niv_vbg <- ggplot(pred_niv_vbg, aes(x = vbg_co2, y = yhat)) +</pre>
  geom line(color = "green") +
```

```
geom_ribbon(aes(ymin = lower, ymax = upper), fill = "green", alpha = 0.2) +
 labs(title = "NIV", x = "VBG CO (mmHg)", y = "Predicted Probability") +
 theme_minimal()
plot_death_vbg <- ggplot(pred_death_vbg, aes(x = vbg_co2, y = yhat)) +</pre>
 geom line(color = "red") +
 geom_ribbon(aes(ymin = lower, ymax = upper), fill = "red", alpha = 0.2) +
 labs(title = "Death", x = "VBG CO (mmHg)", y = "Predicted Probability") +
 theme_minimal()
plot_hcrf_vbg <- ggplot(pred_hcrf_vbg, aes(x = vbg_co2, y = yhat)) +</pre>
 geom_line(color = "purple") +
 geom_ribbon(aes(ymin = lower, ymax = upper), fill = "purple", alpha = 0.2) +
 labs(title = "Hypercapnic RF", x = "VBG CO (mmHg)", y = "Predicted Probability") +
 theme_minimal()
((plot_imv_vbg | plot_niv_vbg) /
 (plot_death_vbg | plot_hcrf_vbg)) +
plot_annotation(title = "Predicted Probability by VBG CO (RCS Models)")
```

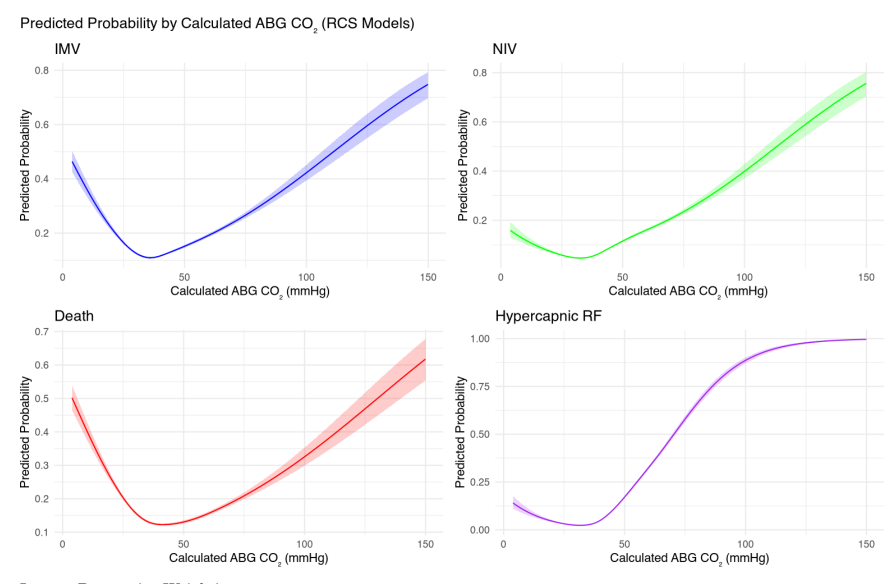


Unweighted, Restricted Cubic Spline Logistic Regressio - Calculated VBG to ABG (Farkas VBG Adjustment)

```
subset_data_calc <- subset_data %>%
  select(calc abg, imv proc, niv proc, death 60d, hypercap resp failure) %>%
  filter(!is.na(calc abg) & complete.cases(.))
dd <- datadist(subset data calc)</pre>
options(datadist = "dd")
fit_imv_abg <- lrm(imv_proc ~ rcs(calc_abg, 4), data = subset_data_calc)</pre>
fit_niv_abg <- lrm(niv_proc ~ rcs(calc_abg, 4), data = subset_data_calc)</pre>
fit death abg <- lrm(death_60d ~ rcs(calc_abg, 4), data = subset_data_calc)
fit hcrf abg <- lrm(hypercap resp failure ~ rcs(calc abg, 4), data = subset_data calc)
pred_imv_abg <- as.data.frame(Predict(fit_imv_abg, calc_abg, fun = plogis))</pre>
pred_niv_abg <- as.data.frame(Predict(fit_niv_abg, calc_abg, fun = plogis))</pre>
pred_death_abg <- as.data.frame(Predict(fit_death_abg, calc_abg, fun = plogis))</pre>
pred_hcrf_abg <- as.data.frame(Predict(fit_hcrf_abg, calc_abg, fun = plogis))</pre>
plot imv abg \leftarrow ggplot(pred imv abg, aes(x = calc abg, y = yhat)) +
  geom line(color = "blue") +
  geom ribbon(aes(ymin = lower, ymax = upper), fill = "blue", alpha = 0.2) +
  labs(title = "IMV", x = "Calculated ABG CO (mmHg)", y = "Predicted Probability") +
  theme minimal()
plot niv abg \leftarrow ggplot(pred niv abg, aes(x = calc abg, y = yhat)) +
  geom_line(color = "green") +
  geom ribbon(aes(ymin = lower, ymax = upper), fill = "green", alpha = 0.2) +
  labs(title = "NIV", x = "Calculated ABG CO (mmHg)", y = "Predicted Probability") +
  theme_minimal()
plot_death_abg <- ggplot(pred_death_abg, aes(x = calc_abg, y = yhat)) +</pre>
  geom_line(color = "red") +
  geom ribbon(aes(ymin = lower, ymax = upper), fill = "red", alpha = 0.2) +
  labs(title = "Death", x = "Calculated ABG CO (mmHg)", y = "Predicted Probability") +
  theme minimal()
```

```
plot_hcrf_abg <- ggplot(pred_hcrf_abg, aes(x = calc_abg, y = yhat)) +
    geom_line(color = "purple") +
    geom_ribbon(aes(ymin = lower, ymax = upper), fill = "purple", alpha = 0.2) +
    labs(title = "Hypercapnic RF", x = "Calculated ABG CO (mmHg)", y = "Predicted Probability") +
    theme_minimal()

((plot_imv_abg | plot_niv_abg) /
    (plot_death_abg | plot_hcrf_abg)) +
    plot_annotation(title = "Predicted Probability by Calculated ABG CO (RCS Models)")</pre>
```



# Inverse Propensity Weighting

IPW done using Gradient Boosting Methods (GBM) - a type of decision-tree based machine learning. "Random forests and GBM are designed to automatically include relevant interactions for variables included in the model. As such, using a GBM to

estimate the PS model, can reduce model misspecification, since the analyst is not required to identify relevant interactions or nonlinearities." from this citation: PMID: 39947224https://pmc.ncbi.nlm.nih.gov/articles/PMC11825193/

Current propensity score uses  $age_at_encounter + sex + race_ethnicity$  (remember - have to specify to use this as a factor variable)  $+ curr_bmi + copd + asthma + osa + chf + acute_nmd + phtn + location (as a factor variable)$ 

Note: for all these, I suggested new GBM adjustments that accomplish the following:

- 1. Smaller GBM & stopping rule  $\rightarrow$  faster fit, avoids over-fitting, lighter tails (which lead to extreme weights that are problematic).
- 2. bal.tab() documents balance; aim is to adjust spec until standard mean difference (SMD) < 0.1.
- 3. Weight stabilization (divide by mean) mitigates a few huge weights. I also winsorized, which is a way to avoid very extreme weights (ie you set <1st percentile to the 1st percentile value, and >99th percentile to 99th percentile.
- 4. Uses robust variance estimation (e.g. allows the variances to change by PaCO2) for IP-weighted GLM; works with splines via rcs(). This is a bit nuanced but I think good to change even though it adds complexity
- 5. Deterministic seed ensures result replication.

\*\*Removed lactate from weights, decreased n.trees, increased bagging

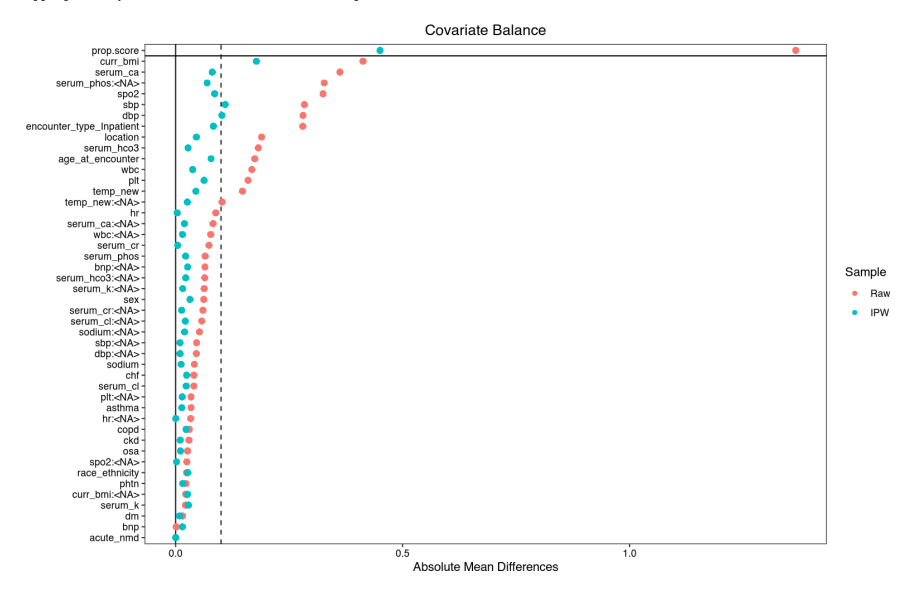
```
# 1. fit GBM propensity model, ABG
set.seed(42)
weight_model <- weightit(</pre>
 has_abg ~ age_at_encounter + sex + factor(race_ethnicity) + curr_bmi + copd + asthma + osa + chf + acute nmd + phtn + ckd +
              = subset_data,
 data
 method
              = "gbm",
              = "ATE",
  estimand
 missing
             = "ind",
 include.obj = TRUE,
                             # ← REQUIRED for importance/SHAP
 n.trees
              = 1500.
                             #decreased trees from 3000 to 1500
```

```
interaction.depth = 3,
 shrinkage = 0.01,
 bag.fraction= 0.8, #increased bagging 0.6 to 0.8 - less overfit extremes
 cv.folds = 5.
 stop.method = "es.mean",
 n.cores = parallel::detectCores()
w_abg <- weight_model # Canonical alias so later code can use `w_abg`</pre>
# 2. Winsorise / stabilise weights (two-sided)
w \leftarrow w / mean(w)
                                  # stabilise
cut <- quantile(w, c(0.01, 1), na.rm = TRUE)</pre>
w <- pmin(pmax(w, cut[1]), cut[2]) # two-tail Winsorisation
w \leftarrow w / mean(w)
                                   # re-stabilise so E[w]=1
# overwrite inside the object and attach to data
weight model$weights <- w</pre>
subset data$w abg <- w
# 3. balance diagnostics (only raw vs. IPW)
bal <- bal.tab(weight_model, un = TRUE, m.threshold = 0.1)</pre>
```

Warning: Missing values exist in the covariates. Displayed values omit these observations.

```
love.plot(
  bal,
  stats = "m",  # standardized mean differences only
  abs = TRUE,
  var.order = "unadjusted",
  sample.names = c("Raw", "IPW")
)
```

Warning: Standardized mean differences and raw mean differences are present in the same plot. Use the `stars` argument to distinguish between them and appropriately label the x-axis. See `?love.plot` for details.



```
# 4. survey design with the same weights
design <- svydesign(ids = ~1, weights = ~w_abg, data = subset_data)</pre>
    5. outcome models (examples)
fit niv <- svyglm(niv proc ~ has abg, design = design, family = quasibinomial())
fit_imv <- svyglm(imv_proc ~ has_abg, design = design, family = quasibinomial())</pre>
fit death <- svyglm(death 60d</pre>
                                    ~ has_abg, design = design, family = quasibinomial())
fit_icd <- svyglm(hypercap_resp_failure ~ has_abg, design = design, family = quasibinomial())</pre>
# quick effect estimates
lapply(list(IMV = fit_imv, NIV = fit_niv, Death = fit_death, ICD = fit_icd), function(m) {
  c(OR = exp(coef(m)[2]),
    LCL = exp(confint(m)[2,1]),
    UCL = exp(confint(m)[2,2]))
})
$IMV
OR.has_abg
                  LCL
                             UCL
  6.776346
             6.606385
                        6.950680
$NIV
OR.has abg
                  LCL
                             UCL
  1.881251
             1.834660
                       1.929026
$Death
OR.has_abg
                  LCL
                             UCL
             2.058994
  2.104033
                        2.150057
$ICD
OR.has_abg
                  LCL
                             UCL
  3.380855
             3.284563
                        3.479970
```

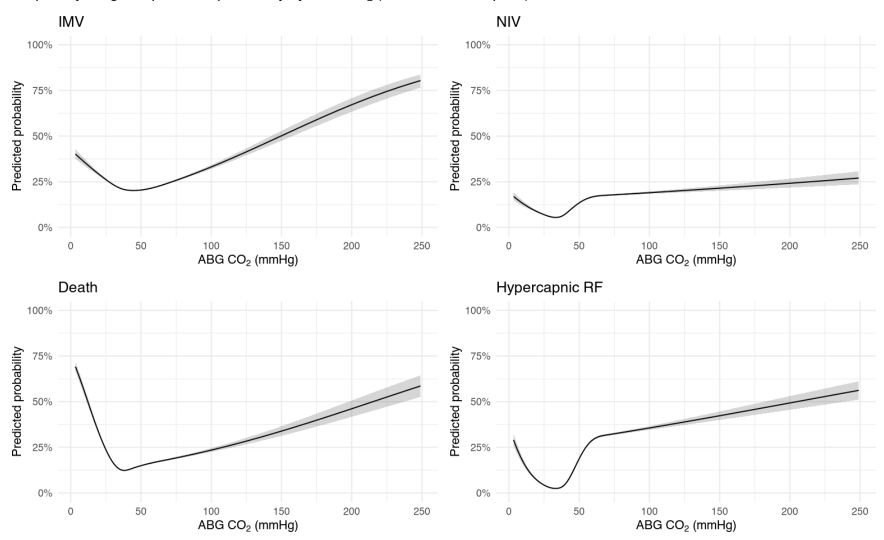
Inverse Propensity-Weighted Logistic Regressions with CO2 predictor represented as a restricted cubic spline.

```
# set.seed(42) # reproducible GBM fit
    1. inverse-probability weights for receiving an ABG
# # done in the last block, so not needed
# 2. analysis sample: rows with a measured PaCO
subset_data_abg <- subset_data %>%
 filter(!is.na(paco2)) %>%
                                               # implies has_abg == 1
  select(paco2, imv_proc, niv_proc, death_60d,
         hypercap_resp_failure, w_abg) %>%
 filter(complete.cases(.))
# 3. weighted logistic spline models with robust SEs
dd <- datadist(subset_data_abg); options(datadist = "dd")</pre>
fitfun <- function(formula)</pre>
 svyglm(
    formula.
   design = svydesign(ids = ~1, weights = ~w_abg, data = subset_data_abg),
   family = quasibinomial()
fit_imv_abg <- fitfun(imv_proc</pre>
                                    ~ rcs(paco2, 4))
fit_niv_abg <- fitfun(niv_proc ~ rcs(paco2, 4))</pre>
fit_death_abg <- fitfun(death_60d</pre>
                                                   ~ rcs(paco2, 4))
fit_hcrf_abg <- fitfun(hypercap_resp_failure ~ rcs(paco2, 4))</pre>
# 4. prediction helper
mkpred <- function(fit, data ref) {</pre>
  # 1. Grid of PaCO values
 newd <- data.frame(</pre>
```

```
paco2 = seq(min(data_ref$paco2, na.rm = TRUE),
                max(data_ref$paco2, na.rm = TRUE),
                length.out = 200)
  # 2. Design (model) matrix for the new data
  mm <- model.matrix(delete.response(terms(fit)),  # drop outcome</pre>
                      data = newd)
  # 3. Linear predictor and its standard error
  eta <- mm %*% coef(fit)
                                                    # 'x
  vcov <- vcov(fit)</pre>
                                                    # robust VCOV from svyglm
  se <- sqrt(rowSums((mm %*% vcov) * mm))
                                                    # √diag(X Σ X)
  # 4. Transform to probability scale
  transform(
    newd.
   yhat = plogis(eta),
   lower = plogis(eta - 1.96 * se),
    upper = plogis(eta + 1.96 * se)
pred_imv_abg <- mkpred(fit_imv_abg,</pre>
                                        subset_data_abg)
pred_niv_abg <- mkpred(fit_niv_abg,</pre>
                                        subset_data_abg)
pred_death_abg <- mkpred(fit_death_abg, subset_data_abg)</pre>
pred_hcrf_abg <- mkpred(fit_hcrf_abg, subset_data_abg)</pre>
# 5. plotting
xlab <- expression(paste("ABG CO"[2], " (mmHg)"))</pre>
plt <- function(dat, title)</pre>
  ggplot(dat, aes(paco2, yhat)) +
    geom line() +
```

```
geom_ribbon(aes(ymin = lower, ymax = upper), alpha = 0.2) +
   scale_y_continuous(limits = c(0, 1), labels = percent_format(accuracy = 1)) +
   labs(title = title, x = xlab, y = "Predicted probability") +
   theme_minimal()
(patchwork::wrap_plots(
   plt(pred_imv_abg,
                       "IMV"),
   plt(pred_niv_abg,
                       "NIV"),
   plt(pred_death_abg, "Death"),
                       "Hypercapnic RF"),
   plt(pred_hcrf_abg,
   ncol = 2
 plot_annotation(
   title = expression(
     paste("Propensity-weighted predicted probability by ABG CO"[2],
           " (restricted cubic spline)")
```

# Propensity-weighted predicted probability by ABG CO<sub>2</sub> (restricted cubic spline)

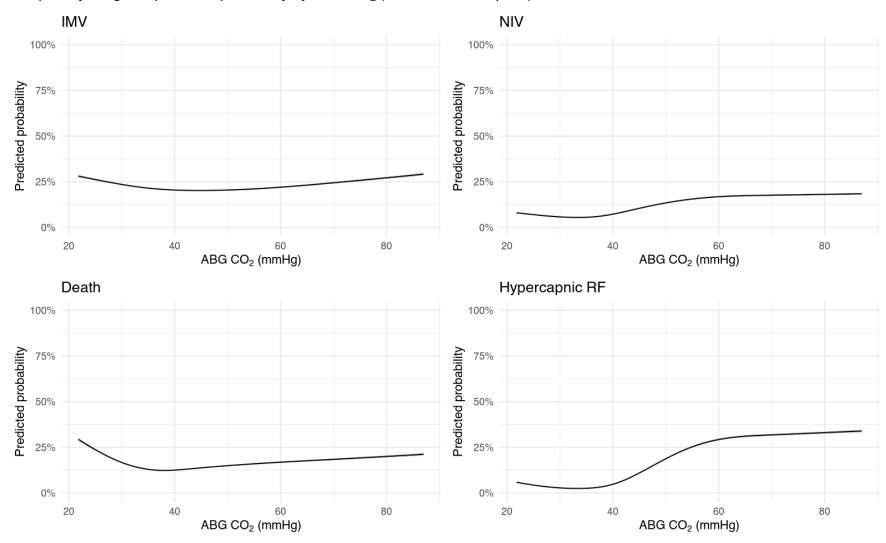


Restricting plots bewtween 0.02 and 0.98

```
subset_data_abg <- subset_data %>%
  filter(!is.na(paco2)) %>%
                                                # implies has abg == 1
  select(paco2, imv_proc, niv_proc, death_60d,
         hypercap_resp_failure, w_abg) %>%
  filter(complete.cases(.))
# 3. weighted logistic spline models with robust SEs
dd <- datadist(subset_data_abg); options(datadist = "dd")</pre>
fitfun <- function(formula)</pre>
  svyglm(
    formula,
    design = svydesign(ids = ~1, weights = ~w_abg, data = subset_data_abg),
    family = quasibinomial()
fit imv abg <- fitfun(imv proc</pre>
                                    \sim rcs(paco2, 4))
fit_niv_abg <- fitfun(niv_proc ~ rcs(paco2, 4))</pre>
fit_death_abg <- fitfun(death_60d</pre>
                                                  ~ rcs(paco2, 4))
fit_hcrf_abg <- fitfun(hypercap_resp_failure ~ rcs(paco2, 4))</pre>
# 4. prediction helper
mkpred <- function(fit, data ref) {</pre>
  # 1. Grid of PaCO values restricted to 2nd-98th percentile
  q <- quantile(data ref$paco2, probs = c(0.02, 0.98), na.rm = TRUE)
  newd <- data.frame(</pre>
    paco2 = seq(q[1], q[2], length.out = 200)
  # 2. Design (model) matrix for the new data
  mm <- model.matrix(delete.response(terms(fit)), data = newd)</pre>
  # 3. Linear predictor and its standard error
```

```
eta <- mm %*% coef(fit)
  vcov <- vcov(fit)</pre>
  se <- sqrt(rowSums((mm %*% vcov) * mm))
  # 4. Transform to probability scale
  transform(
    newd.
   yhat = plogis(eta),
   lower = plogis(eta - 1.96 * se),
    upper = plogis(eta + 1.96 * se)
pred_imv_abg <- mkpred(fit_imv_abg,</pre>
                                        subset_data_abg)
pred_niv_abg <- mkpred(fit_niv_abg,</pre>
                                         subset_data_abg)
pred_death_abg <- mkpred(fit_death_abg, subset_data_abg)</pre>
pred_hcrf_abg <- mkpred(fit_hcrf_abg, subset_data_abg)</pre>
    5. plotting
xlab <- expression(paste("ABG CO"[2], " (mmHg)"))</pre>
plt <- function(dat, title)</pre>
  ggplot(dat, aes(paco2, yhat)) +
    geom line() +
    geom_ribbon(aes(ymin = lower, ymax = upper), alpha = 0.2) +
    scale_y_continuous(limits = c(0, 1), labels = percent_format(accuracy = 1)) +
    labs(title = title, x = xlab, y = "Predicted probability") +
    theme_minimal()
(patchwork::wrap_plots(
    plt(pred_imv_abg,
                         "IMV"),
    plt(pred_niv_abg,
                         "NIV"),
    plt(pred_death_abg, "Death"),
    plt(pred_hcrf_abg,
                         "Hypercapnic RF"),
```

# Propensity-weighted predicted probability by ABG CO<sub>2</sub> (restricted cubic spline)



VBG - changed trees and bag fraction

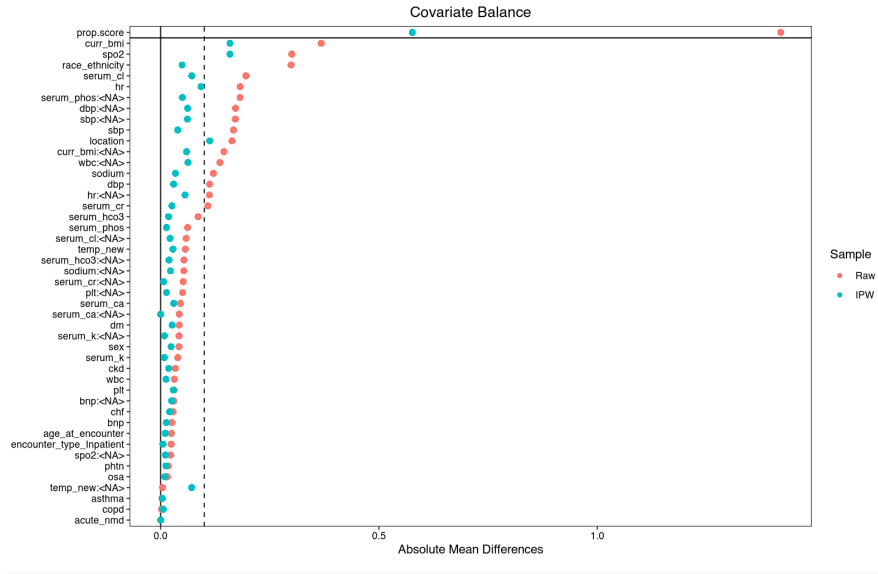
```
# Inverse-propensity weighting & outcome modelling for **VBG** cohort
  - mirrored 1-to-1 to the validated ABG workflow
set.seed(42)
# 1. IPW for VBG ------
w_vbg <- weightit(</pre>
 has_vbg ~ age_at_encounter + sex + factor(race_ethnicity) + curr_bmi +
   copd + asthma + osa + chf + acute_nmd + phtn + ckd + dm +
   factor(location) + factor(encounter_type) + temp_new + sbp + dbp + hr + spo2 + sodium + serum_cr + serum_hco3 + serum_cl +
   serum_k + wbc + plt + bnp + serum_phos + serum_ca,
            = subset_data,
 data
 method = "gbm",
 estimand = "ATE",
 missing = "ind",
 include.obj = TRUE,
                       # ← REQUIRED for importance/SHAP
 n.trees
          = 1500,
 interaction.depth = 3,
 shrinkage = 0.01,
 bag.fraction= 0.8,
 cv.folds = 5,
 stop.method = "es.mean",
 n.cores = parallel::detectCores()
# Stabilise & winsorise weights
w <- w_vbg$weights
w \leftarrow w / mean(w)
cut <- quantile(w, c(0.01, 1), na.rm = TRUE)
w <- pmin(pmax(w, cut[1]), cut[2])</pre>
w \leftarrow w / mean(w)
w_vbg$weights <- w</pre>
subset_data$w_vbg <- w</pre>
```

```
v_bal <- bal.tab(w_vbg, un = TRUE, m.threshold = 0.1)</pre>
```

Warning: Missing values exist in the covariates. Displayed values omit these observations.

```
love.plot(
  v_bal,
  stats = "m",  # standardized mean differences only
  abs = TRUE,
  var.order = "unadjusted",
  sample.names = c("Raw", "IPW")
)
```

Warning: Standardized mean differences and raw mean differences are present in the same plot. Use the `stars` argument to distinguish between them and appropriately label the x-axis. See `?love.plot` for details.



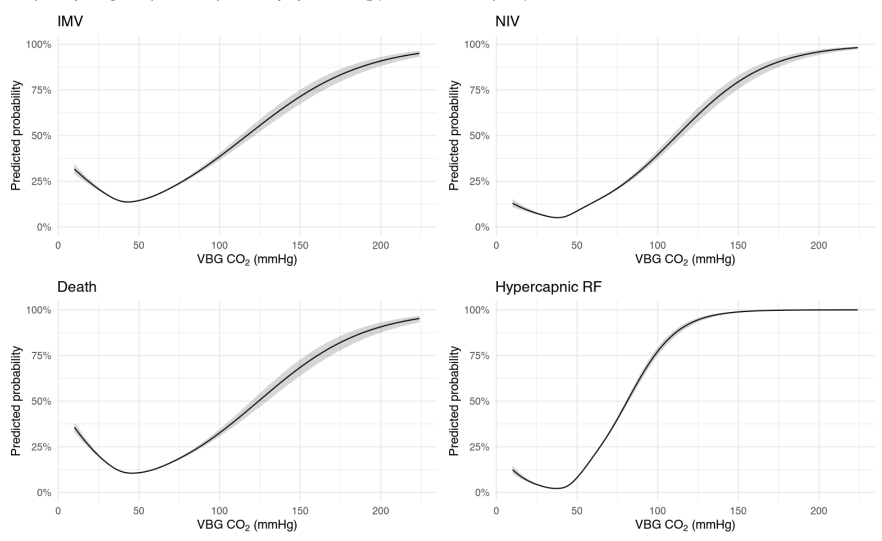
```
# 2. Analysis set (VBG only) ------
subset_data_vbg <- subset_data %>%
filter(!is.na(vbg_co2)) %>%
```

```
select(vbg_co2, imv_proc, niv_proc, death_60d,
        hypercap_resp_failure, w_vbg) %>%
 filter(complete.cases(.))
# 3. Weighted spline models -----
dd_vbg <- datadist(subset_data_vbg)</pre>
options(datadist = "dd_vbg")
fitfun <- function(formula)</pre>
  svyglm(
   formula,
   design = svydesign(ids = ~1, weights = ~w_vbg, data = subset_data_vbg),
   family = quasibinomial()
fit_death_vbg <- fitfun(death_60d ~ rcs(vbg_co2, 4))
fit_hcrf_vbg <- fitfun(hypercap_resp_failure ~ rcs(vbg_co2, 4))</pre>
# 4. Prediction helper -----
mkpred <- function(fit, data ref) {</pre>
 newd <- data.frame(</pre>
   vbg_co2 = seq(min(data_ref$vbg_co2, na.rm = TRUE),
                max(data_ref$vbg_co2, na.rm = TRUE),
                length.out = 200)
 mm <- model.matrix(delete.response(terms(fit)), newd)</pre>
  eta <- mm %*% coef(fit)
  vcov <- vcov(fit)</pre>
  se <- sqrt(rowSums((mm %*% vcov) * mm))
  transform(
   newd.
   yhat = plogis(eta),
```

```
lower = plogis(eta - 1.96 * se),
    upper = plogis(eta + 1.96 * se)
pred_imv_vbg <- mkpred(fit_imv_vbg,</pre>
                                        subset_data_vbg)
pred_niv_vbg <- mkpred(fit_niv_vbg,</pre>
                                        subset_data_vbg)
pred_death_vbg <- mkpred(fit_death_vbg, subset_data_vbg)</pre>
pred_hcrf_vbg <- mkpred(fit_hcrf_vbg, subset_data_vbg)</pre>
# 5. Plotting (gray scheme) -----
xlab <- expression(paste("VBG CO"[2], " (mmHg)"))</pre>
plt <- function(dat, title)</pre>
  ggplot(dat, aes(vbg_co2, yhat)) +
    geom_line() +
    geom ribbon(aes(ymin = lower, ymax = upper), alpha = 0.2) +
    scale_y_continuous(limits = c(0, 1), labels = percent_format(accuracy = 1)) +
    labs(title = title, x = xlab, y = "Predicted probability") +
    theme minimal()
(patchwork::wrap_plots(
    plt(pred_imv_vbg,
                         "IMV"),
   plt(pred_niv_vbg,
                         "NIV"),
    plt(pred_death_vbg, "Death"),
    plt(pred_hcrf_vbg, "Hypercapnic RF"),
    ncol = 2
  plot_annotation(
   title = expression(
      paste("Propensity-weighted predicted probability by VBG CO"[2],
            " (restricted cubic spline)")
```

)

# Propensity-weighted predicted probability by VBG CO<sub>2</sub> (restricted cubic spline)



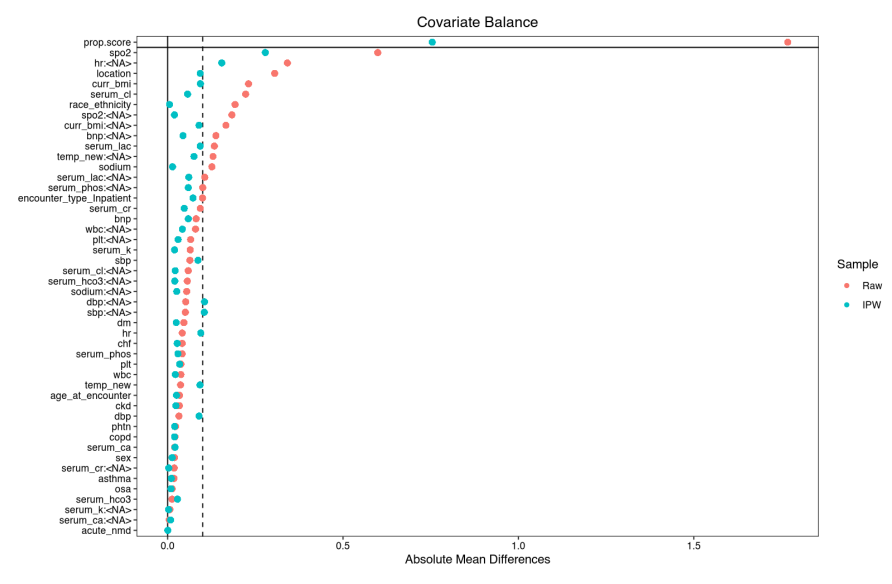
Calculated VBG to ABG / Farkas

```
Propensity-weighted spline models for **Calculated ABG CO **
  (weights still derive from propensity to receive a VBG)
# 1. define the new treatment variable -----
subset_data <- subset_data %>%
 mutate(
   has_vbg_co2_o2_sat = if_else(
     !is.na(vbg_co2) & vbg_co2 != 0 &
     !is.na(vbg_o2sat) & vbg_o2sat != 0,
     1, 0
   )
# quick sanity check
# table(subset_data$has_vbg_co2_o2_sat, useNA = "ifany")
# 2. fit the GBM propensity model -----
set.seed(42)
w_vbg_calc <- weightit(</pre>
 has_vbg_co2_o2_sat ~ age_at_encounter + sex + factor(race_ethnicity) + curr_bmi + copd + asthma + osa + chf + acute_nmd + ph
 data
            = subset_data,
           = "gbm",
 method
 estimand = "ATE",
 missing = "ind",
 include.obj = TRUE,
 n.trees
            = 3000,
 interaction.depth = 3,
 shrinkage = 0.01,
 bag.fraction= 0.6,
 cv.folds = 5,
 stop.method = "es.mean",
 n.cores
            = parallel::detectCores()
```

Warning: Missing values exist in the covariates. Displayed values omit these observations.

```
love.plot(
  v_calc_bal,
  stats = "m",  # standardized mean differences only
  abs = TRUE,
  var.order = "unadjusted",
  sample.names = c("Raw", "IPW")
)
```

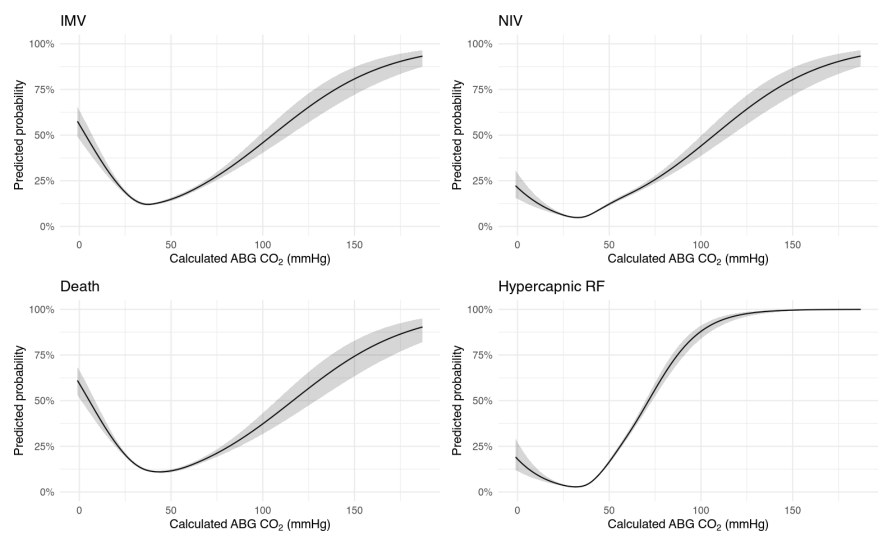
Warning: Standardized mean differences and raw mean differences are present in the same plot. Use the `stars` argument to distinguish between them and appropriately label the x-axis. See `?love.plot` for details.



```
select(calc_abg, imv_proc, niv_proc, death_60d,
         hypercap_resp_failure, w_vbg_calc) %>%
 filter(complete.cases(.))
# 3. Weighted logistic spline models with robust SEs -----
dd <- datadist(subset data calc); options(datadist = "dd")</pre>
fitfun <- function(formula)</pre>
  svyglm(
    formula,
   design = svydesign(ids = ~1, weights = ~w_vbg_calc, data = subset_data_calc),
   family = quasibinomial()
fit_imv_calc <- fitfun(imv_proc ~ rcs(calc_abg, 4))</pre>
fit_niv_calc <- fitfun(niv_proc ~ rcs(calc_abg, 4))</pre>
fit_death_calc <- fitfun(death_60d</pre>
                                                 ~ rcs(calc_abg, 4))
fit_hcrf_calc <- fitfun(hypercap_resp_failure ~ rcs(calc_abg, 4))</pre>
# 4. Prediction helper -----
mkpred <- function(fit, data_ref) {</pre>
  newd <- data.frame(</pre>
    calc_abg = seq(min(data_ref$calc_abg, na.rm = TRUE),
                   max(data_ref$calc_abg, na.rm = TRUE),
                   length.out = 200)
  )
 mm <- model.matrix(delete.response(terms(fit)), newd)</pre>
  eta <- mm %*% coef(fit)
  vcov <- vcov(fit)</pre>
  se <- sqrt(rowSums((mm %*% vcov) * mm))
  transform(
    newd.
   yhat = plogis(eta),
   lower = plogis(eta - 1.96 * se),
```

```
upper = plogis(eta + 1.96 * se)
  )
pred_imv_calc <- mkpred(fit_imv_calc,</pre>
                                           subset_data_calc)
pred_niv_calc <- mkpred(fit_niv_calc,</pre>
                                           subset_data_calc)
pred_death_calc <- mkpred(fit_death_calc, subset_data_calc)</pre>
pred_hcrf_calc <- mkpred(fit_hcrf_calc, subset_data_calc)</pre>
# 5. Plotting -----
xlab <- expression(paste("Calculated ABG CO"[2], " (mmHg)"))</pre>
plt <- function(dat, title)</pre>
  ggplot(dat, aes(calc_abg, yhat)) +
    geom_line() +
    geom_ribbon(aes(ymin = lower, ymax = upper), alpha = 0.2) +
    scale_y_continuous(limits = c(0, 1), labels = percent_format(accuracy = 1)) +
    labs(title = title, x = xlab, y = "Predicted probability") +
    theme_minimal()
(patchwork::wrap_plots(
    plt(pred_imv_calc,
                         "IMV"),
    plt(pred_niv_calc,
                         "NIV"),
    plt(pred_death_calc, "Death"),
    plt(pred_hcrf_calc, "Hypercapnic RF"),
    ncol = 2
  plot_annotation(
    title = expression(
      paste("Propensity-weighted predicted probability by Calculated ABG CO"[2],
            " (restricted cubic spline)")
```

## Propensity-weighted predicted probability by Calculated ABG CO<sub>2</sub> (restricted cubic spline)



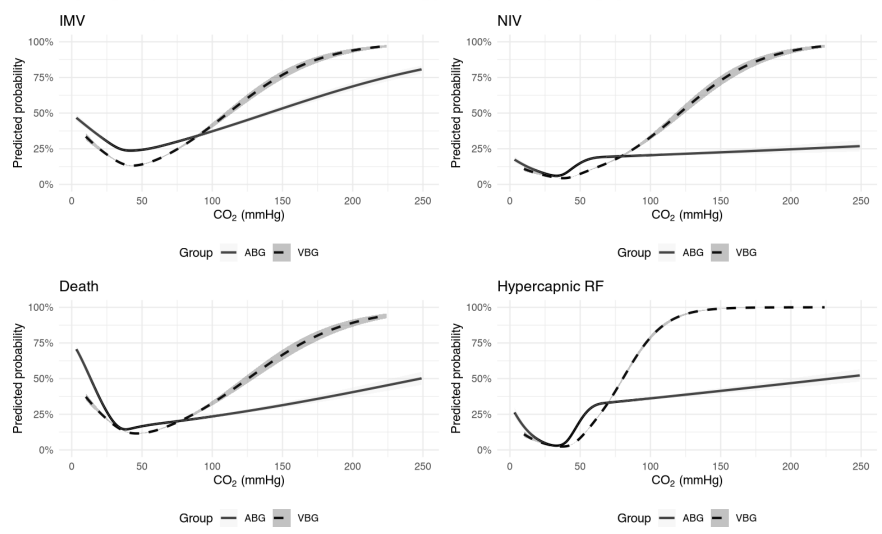
Superimposing ABG and VBG weighted restricted cubic splines

```
library(dplyr)
library(ggplot2)
library(patchwork)
library(scales)
library(rms)
# ABG spline fits (unweighted, rms::lrm)
fit_imv_abg <- lrm(imv_proc</pre>
                                           ~ rcs(paco2, 4), data = subset data abg)
fit_niv_abg <- lrm(niv_proc
                                           ~ rcs(paco2, 4), data = subset_data_abg)
fit death abg <- lrm(death 60d
                                           ~ rcs(paco2, 4), data = subset_data_abg)
fit_hcrf_abg <- lrm(hypercap_resp_failure ~ rcs(paco2, 4), data = subset_data_abg)</pre>
# VBG spline fits (mirror pattern)
                                           ~ rcs(vbg_co2, 4), data = subset_data_vbg)
fit_imv_vbg <- lrm(imv_proc
fit_niv_vbg <- lrm(niv_proc</pre>
                                           ~ rcs(vbg_co2, 4), data = subset_data_vbg)
fit_death_vbg <- lrm(death_60d
                                           ~ rcs(vbg_co2, 4), data = subset_data_vbg)
fit_hcrf_vbg <- lrm(hypercap_resp_failure ~ rcs(vbg_co2, 4), data = subset_data_vbg)</pre>
library(rms) # ensure lrm() and Predict() are available
# Helper to make predictions with standardized columns: co2, yhat, lower, upper, group
mkpred <- function(fit, data_ref, xvar, group_label, n = 200) {</pre>
  stopifnot(is.character(xvar), length(xvar) == 1, xvar %in% names(data_ref))
 xseq <- seq(min(data_ref[[xvar]], na.rm = TRUE),</pre>
              max(data_ref[[xvar]], na.rm = TRUE),
              length.out = n)
  if (inherits(fit, "lrm")) {
    # Predict() needs a datadist object visible by name set in options(datadist=)
    dd <- rms::datadist(data ref)</pre>
    old <- options(datadist = "dd")</pre>
    on.exit(options(old), add = TRUE)
    assign("dd", dd, envir = .GlobalEnv)
```

```
# IMPORTANT: name the model argument 'object', not 'fit'
    args <- c(list(object = fit, fun = plogis),</pre>
               stats::setNames(list(xseq), xvar))
    p <- do.call(rms::Predict, args)</pre>
    out <- as.data.frame(p)</pre>
    # standardize column names used by plotting code
    names(out)[names(out) == xvar] <- "co2"</pre>
    out$group <- group_label</pre>
    out[, c("co2", "yhat", "lower", "upper", "group")]
  } else {
    # glm/svyglm path
    newd <- stats::setNames(data.frame(xseq), xvar)</pre>
         <- stats::model.matrix(stats::delete.response(stats::terms(fit)), newd)</pre>
    beta <- stats::coef(fit)</pre>
    eta <- drop(X %*% beta)
    V <- stats::vcov(fit)</pre>
    se <- sqrt(rowSums((X %*% V) * X))
    data.frame(
      co2 = xseq,
      yhat = plogis(eta),
      lower = plogis(eta - 1.96 * se),
      upper = plogis(eta + 1.96 * se),
      group = group_label,
      check.names = FALSE
    Generate predictions
# VBG
pred_imv_vbg <- mkpred(fit_imv_vbg, subset_data_vbg, "vbg_co2", "VBG")</pre>
                                          subset_data_vbg, "vbg_co2", "VBG")
pred_niv_vbg <- mkpred(fit_niv_vbg,</pre>
pred_death_vbg <- mkpred(fit_death_vbg, subset_data_vbg, "vbg_co2", "VBG")</pre>
pred_hcrf_vbg <- mkpred(fit_hcrf_vbg, subset_data_vbg, "vbg_co2", "VBG")</pre>
```

```
# ABG
pred imv abg <- mkpred(fit imv abg,</pre>
                                         subset data abg, "paco2", "ABG")
pred_niv_abg <- mkpred(fit_niv_abg,</pre>
                                         subset_data_abg, "paco2", "ABG")
pred death abg <- mkpred(fit death abg, subset data abg, "paco2", "ABG")
pred_hcrf_abg <- mkpred(fit_hcrf_abg, subset_data_abg, "paco2", "ABG")</pre>
# Combine
pred imv <- bind rows(pred imv vbg,</pre>
                                       pred imv abg)
pred_niv <- bind_rows(pred_niv_vbg,</pre>
                                       pred_niv_abg)
pred_death <- bind_rows(pred_death_vbg, pred_death_abg)</pre>
pred_hcrf <- bind_rows(pred_hcrf_vbg, pred_hcrf_abg)</pre>
# Plotting function in grayscale with distinguishable ribbons
plt_gray <- function(dat, title) {</pre>
  ggplot(dat, aes(x = co2, y = yhat, linetype = group)) +
    geom_line(color = "black", linewidth = 1) +
    geom_ribbon(aes(ymin = lower, ymax = upper, fill = group),
                alpha = 0.3, color = NA) +
    scale_fill_manual(values = c("ABG" = "gray90", "VBG" = "gray20")) + # different gray shades
    scale linetype manual(values = c("ABG" = "solid", "VBG" = "dashed")) +
    scale_y_continuous(limits = c(0, 1),
                       labels = scales::percent format(accuracy = 1)) +
    labs(title = title,
         x = \exp(C0[2] \sim (mmHg)),
         y = "Predicted probability",
         fill = "Group",
         linetype = "Group") +
    theme_minimal() +
    theme(legend.position = "bottom")
   Patchwork layout with gray shades
(patchwork::wrap_plots(
```

# Propensity-weighted predicted probability by ABG vs VBG CO<sub>2</sub> (restricted cubic splines, gray scheme)



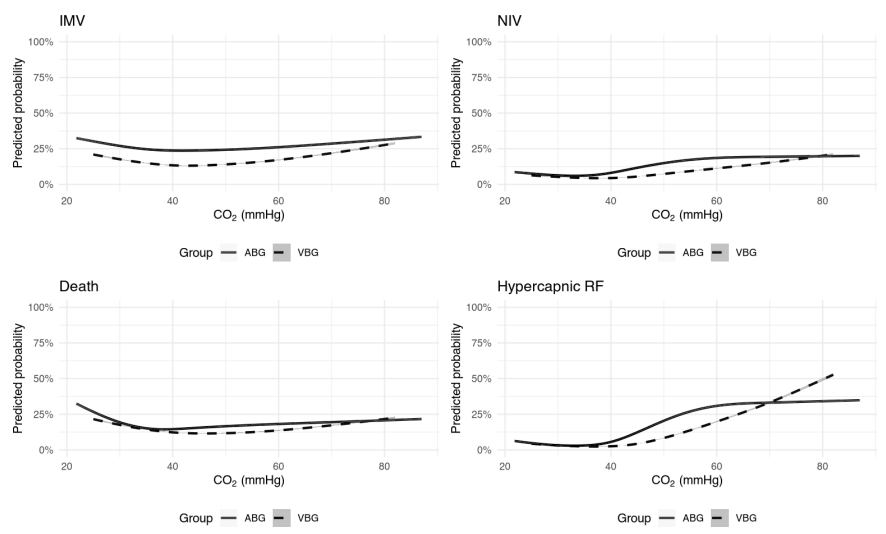
Restricting the plot to 0.02 to 0.99 (since this puts it at about  $100 \mathrm{mmHg}$  for CO2)

```
library(dplyr)
library(ggplot2)
library(patchwork)
library(scales)
library(rms)
# ABG spline fits (unweighted, rms::lrm)
fit_imv_abg <- lrm(imv_proc</pre>
                                           ~ rcs(paco2, 4), data = subset data abg)
fit_niv_abg <- lrm(niv_proc
                                           ~ rcs(paco2, 4), data = subset_data_abg)
fit_death_abg <- lrm(death_60d</pre>
                                           ~ rcs(paco2, 4), data = subset_data_abg)
fit_hcrf_abg <- lrm(hypercap_resp_failure ~ rcs(paco2, 4), data = subset_data abg)</pre>
# VBG spline fits (mirror pattern)
                                           ~ rcs(vbg_co2, 4), data = subset_data_vbg)
fit_imv_vbg <- lrm(imv_proc
fit_niv_vbg <- lrm(niv_proc</pre>
                                           ~ rcs(vbg_co2, 4), data = subset_data_vbg)
fit_death_vbg <- lrm(death_60d</pre>
                                           ~ rcs(vbg_co2, 4), data = subset_data_vbg)
fit hcrf vbg <- lrm(hypercap resp failure ~ rcs(vbg co2, 4), data = subset data vbg)
library(rms) # ensure lrm() and Predict() are available
# Helper to make predictions with standardized columns: co2, yhat, lower, upper, group
mkpred <- function(fit, data_ref, xvar, group_label, n = 200) {</pre>
  stopifnot(is.character(xvar), length(xvar) == 1, xvar %in% names(data_ref))
  # Restrict to 2nd and 98th percentiles
  rng <- quantile(data ref[[xvar]], probs = c(0.02, 0.98), na.rm = TRUE)</pre>
  xseq <- seq(rng[1], rng[2], length.out = n)</pre>
  if (inherits(fit, "lrm")) {
    dd <- rms::datadist(data ref)</pre>
    old <- options(datadist = "dd")</pre>
    on.exit(options(old), add = TRUE)
    assign("dd", dd, envir = .GlobalEnv)
```

```
args <- c(list(object = fit, fun = plogis),</pre>
               stats::setNames(list(xseq), xvar))
    p <- do.call(rms::Predict, args)</pre>
    out <- as.data.frame(p)</pre>
    names(out)[names(out) == xvar] <- "co2"</pre>
    out$group <- group label
    out[, c("co2", "yhat", "lower", "upper", "group")]
  } else {
    newd <- stats::setNames(data.frame(xseq), xvar)</pre>
    X <- stats::model.matrix(stats::delete.response(stats::terms(fit)), newd)</pre>
    beta <- stats::coef(fit)</pre>
    eta <- drop(X %*% beta)
    V <- stats::vcov(fit)</pre>
    se <- sqrt(rowSums((X %*% V) * X))</pre>
    data.frame(
      co2 = xseq,
      yhat = plogis(eta),
      lower = plogis(eta - 1.96 * se),
      upper = plogis(eta + 1.96 * se),
      group = group_label,
      check.names = FALSE
  }
    Generate predictions
# VBG
pred_imv_vbg <- mkpred(fit_imv_vbg,</pre>
                                        subset_data_vbg, "vbg_co2", "VBG")
pred_niv_vbg <- mkpred(fit_niv_vbg, subset_data_vbg, "vbg_co2", "VBG")</pre>
pred_death_vbg <- mkpred(fit_death_vbg, subset_data_vbg, "vbg_co2", "VBG")</pre>
pred_hcrf_vbg <- mkpred(fit_hcrf_vbg, subset_data_vbg, "vbg_co2", "VBG")</pre>
# ABG
pred imv abg <- mkpred(fit imv abg,</pre>
                                         subset data abg, "paco2", "ABG")
```

```
pred_niv_abg <- mkpred(fit_niv_abg,</pre>
                                         subset_data_abg, "paco2", "ABG")
pred death abg <- mkpred(fit death abg, subset data abg, "paco2", "ABG")</pre>
pred_hcrf_abg <- mkpred(fit_hcrf_abg, subset_data_abg, "paco2", "ABG")</pre>
# Combine
pred imv <- bind rows(pred imv vbg,</pre>
                                       pred imv abg)
pred_niv <- bind_rows(pred_niv_vbg,</pre>
                                       pred_niv_abg)
pred death <- bind rows(pred death vbg, pred death abg)</pre>
pred_hcrf <- bind_rows(pred_hcrf_vbg, pred_hcrf_abg)</pre>
# Plotting function in grayscale with distinguishable ribbons
plt_gray <- function(dat, title) {</pre>
  ggplot(dat, aes(x = co2, y = yhat, linetype = group)) +
    geom_line(color = "black", linewidth = 1) +
    geom_ribbon(aes(ymin = lower, ymax = upper, fill = group),
                alpha = 0.3, color = NA) +
    scale_fill_manual(values = c("ABG" = "gray90", "VBG" = "gray20")) + # different gray shades
    scale linetype manual(values = c("ABG" = "solid", "VBG" = "dashed")) +
    scale_y_continuous(limits = c(0, 1),
                       labels = scales::percent format(accuracy = 1)) +
    labs(title = title,
         x = expression(CO[2]~"(mmHg)"),
         y = "Predicted probability",
         fill = "Group",
         linetype = "Group") +
    theme_minimal() +
    theme(legend.position = "bottom")
   Patchwork layout with gray shades
(patchwork::wrap_plots(
   plt_gray(pred_imv,
                         "IMV").
                         "NIV").
    plt_gray(pred_niv,
    plt gray(pred death, "Death"),
```

## Propensity-weighted predicted probability by ABG vs VBG CO<sub>2</sub> (restricted cubic splines, gray scheme)



Feature importance: global contribution of a feature to the model's predictive performance on the training distribution. Quick global triage—which variables the model leaned on to fit propensity. Good for model debugging, feature pruning, and tracking drift across refits (qualitatively).

SHAP: a local, signed attribution for that subject: "by how much did feature j push this person's log-odds of receiving the test up or down vs baseline?, then global shap is mean absolute SHAP across subjects—i.e., the **typical magnitude** of a feature's contribution to predictions in your population. Good for auditability and **directional insight**—who is assigned higher/lower propensity by which features, spot proxies, and communicate fairness/operational drivers. Aggregate with mean |SHAP| for a **global ranking with direction available** when needed.

TODO: Can label y-axis in plots: contribution to the log odds of receiving an ABG or VBG for the SHAP values.

#### **ABG**

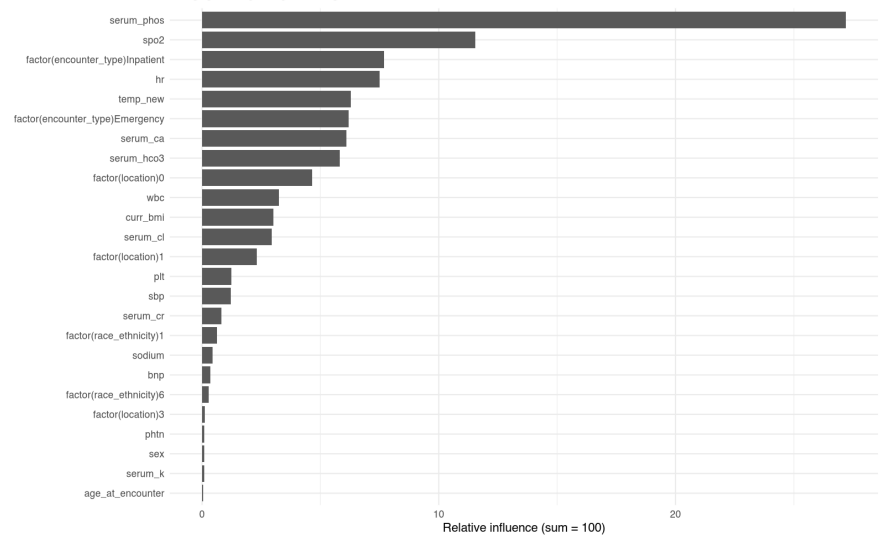
For the top SHAP-ranked predictors we computed partial- and accumulated-local-effects (ALE) to estimate the marginal change in predicted risk across clinically relevant ranges, robust to covariate correlation. We complemented this with SHAP dependence plots (colored by plausible interactions) and fitted transparent spline-logistic models to identify turn-points ('knees') where marginal log-odds slope changed.

```
# --- deps -----
library(WeightIt)
library(gbm)
library(dplyr)
library(ggplot2)
library(fastshap)
# --- 0) Canonicalize object name -----
# Your 9-26.qmd labeled the ABG propensity object `weight model`.
if (!exists("w abg", inherits = TRUE) && exists("weight model", inherits = TRUE)) {
  w abg <- weight model
stopifnot(exists("w abg", inherits = TRUE))
# --- 1) Ensure the WeightIt object stores the GBM + covariate matrix ------
ensure_gbm_obj <- function(W) {</pre>
 stopifnot(inherits(W, "weightit"))
 has_obj <- !is.null(W$obj) || !is.null(W$info$obj) || !is.null(W$info$model.obj)
 has_cov <- !is.null(W$covs)</pre>
 if (has_obj && has_cov) return(W)
```

```
cl <- as.list(W$call); cl[[1]] <- WeightIt::weightit</pre>
  if (!is.null(cl[["missing."]])) { cl$missing <- cl[["missing."]]; cl[["missing."]] <- NULL }</pre>
                                    cl$missing <- "ind"</pre>
  if (is.null(cl$missing))
  cl$include.obj <- TRUE</pre>
  if (is.null(cl$method))
                                    cl$method <- "gbm"</pre>
  if (is.language(cl$formula)) cl$formula <- eval(cl$formula, envir = .GlobalEnv)</pre>
  if (is.language(cl$data))
                                 cl$data <- eval(cl$data, envir = .GlobalEnv)</pre>
  do.call(WeightIt::weightit, cl[-1])
w_abg <- ensure_gbm_obj(w_abg)</pre>
# --- 2) Helpers: design alignment, importance, fast SHAP (logit scale) ------
prep_design <- function(W) {</pre>
  stopifnot(inherits(W, "weightit"))
  gbm_fit <- if (!is.null(W$obj)) W$obj else if (!is.null(W$info$obj)) W$info$obj else W$info$model.obj
  stopifnot(inherits(gbm_fit, "gbm"))
  stopifnot(!is.null(W$covs))
  X <- W$covs
  if (inherits(X, "tbl")) X <- as.data.frame(X)</pre>
  if (inherits(X, "Matrix")) X <- as.matrix(X)</pre>
  X <- as.data.frame(X, stringsAsFactors = FALSE)</pre>
  # conservative coercion: only numeric-like strings → numeric
  for (nm in names(X)) {
    if (is.factor(X[[nm]])) X[[nm]] <- as.character(X[[nm]])</pre>
    if (is.character(X[[nm]])) {
      ok <- grepl("^[-+]?[0-9.]+$", X[[nm]] %||% "")
      if (all(ok | is.na(X[[nm]]))) suppressWarnings(X[[nm]] <- as.numeric(X[[nm]]))</pre>
    }
  }
  vars <- gbm_fit$var.names</pre>
  miss <- setdiff(vars, colnames(X))</pre>
```

```
if (length(miss)) for (nm in miss) X[[nm]] <- 0</pre>
  X <- X[, vars, drop = FALSE]</pre>
  best_tree <- if (!is.null(W$info$best.tree)) W$info$best.tree else gbm_fit$n.trees
 list(X = X, gbm_fit = gbm_fit, best_tree = best_tree)
extract_gbm_importance <- function(W, top_n = 25) {</pre>
  mats <- prep_design(W)</pre>
  as.data.frame(summary(mats$gbm_fit, n.trees = mats$best_tree, plotit = FALSE)) |>
    arrange(desc(rel.inf)) |>
    slice_head(n = top_n)
plot_gbm importance <- function(imp_df, title = "GBM variable importance (relative influence)") {</pre>
  ggplot(imp_df, aes(x = rel.inf, y = reorder(var, rel.inf))) +
    geom_col(width = 0.85) +
    labs(x = "Relative influence (sum = 100)", y = NULL, title = title) +
    theme_minimal(base_size = 11)
}
# --- 3) Run: ABG selection model - importance + fast SHAP -----
imp_abg <- extract_gbm_importance(w_abg, top_n = 25)</pre>
p_imp_abg <- plot_gbm_importance(imp_abg, "ABG selection model - GBM relative influence")</pre>
p_imp_abg
```

## ABG selection model — GBM relative influence



# --- Build shapviz object robustly -----library(shapviz)

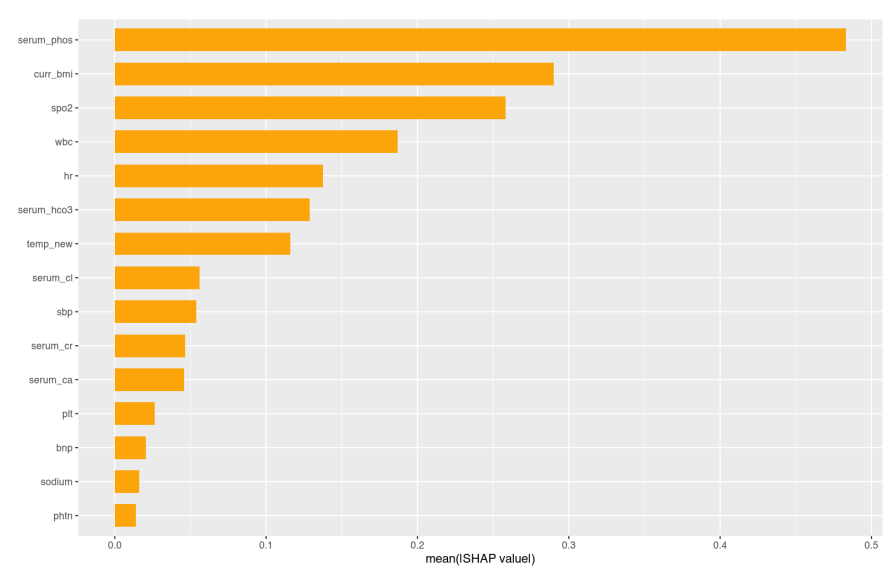
```
# fast SHAP on LOGIT scale; prunes zero-variance features; subsamples rows
compute_shap_fast <- function(W, top_k = 30, nsim = 64, frac_rows = 0.50,
                               \max \text{ rows} = 100000, \text{ seed} = 123)  {
 mats <- prep_design(W); X <- mats$X; gbm_fit <- mats$gbm_fit; best_tree <- mats$best_tree</pre>
  imp <- as.data.frame(summary(gbm_fit, n.trees = best_tree, plotit = FALSE))</pre>
 top_feats <- head(imp$var, min(top_k, nrow(imp)))</pre>
  top_feats <- intersect(top_feats, colnames(X))</pre>
  # drop constant features (avoid flat SHAP/plots)
 nzv <- sapply(X[, top feats, drop = FALSE], function(z) sd(z, na.rm = TRUE) > 0)
  top_feats <- top_feats[nzv]</pre>
 if (!length(top_feats)) stop("All candidate features are near-constant in this subset.")
 n <- nrow(X); target n <- min(n, max rows, ceiling(frac rows * n))</pre>
  set.seed(seed)
 Xsub <- if (target n < n) X[sample.int(n, target n), , drop = FALSE] else X</pre>
 # SHAP on logit scale for contrast/stability
 pfun <- function(object, newdata)</pre>
    predict(object, newdata = newdata, n.trees = best_tree, type = "link")
 fs formals <- names(formals(fastshap::explain))</pre>
 args <- list(object = gbm_fit, X = Xsub, pred_wrapper = pfun, nsim = nsim, adjust = TRUE)
 if ("feature_names" %in% fs formals) args$feature_names <- top_feats</pre>
  set.seed(seed)
  S <- do.call(fastshap::explain, args) # matrix or data.frame of SHAP
 list(shap = S, X = Xsub, top_feats = top_feats, imp = imp)
t0 <- Sys.time()
sh abg fast <- compute shap fast(w abg, top k = 100, nsim = 32, frac rows = 0.25, max rows = 100000)
```

```
t1 <- Sys.time(); message(sprintf("[compute_shap_fast] %.2f s", as.numeric(difftime(t1, t0, units="secs"))))
```

[compute\_shap\_fast] 6440.31 s

```
# 1) Take SHAP and design from your fast SHAP object
S <- as.matrix(sh_abg_fast$shap) # n x p SHAP matrix</pre>
X <- as.data.frame(sh_abg_fast$X) # matching rows, p columns</pre>
# 2) Make X numeric-only (handles factors / labelled types safely)
for (nm in names(X)) {
 if (inherits(X[[nm]], "haven_labelled")) {
    X[[nm]] <- labelled::to_factor(X[[nm]])</pre>
  }
  if (is.factor(X[[nm]])) X[[nm]] <- as.character(X[[nm]])</pre>
 if (is.character(X[[nm]])) suppressWarnings(X[[nm]]] <- as.numeric(X[[nm]]))</pre>
}
# 3) Align names/order between S and X (and give S names if missing)
if (is.null(colnames(S))) colnames(S) <- colnames(X)</pre>
S <- S[, intersect(colnames(S), colnames(X)), drop = FALSE]
X <- X[, colnames(S), drop = FALSE]</pre>
# 4) Construct shapviz object: PASS S POSITIONALLY (no name) to avoid dispatch bug
sv <- shapviz(S, X = as.matrix(X))</pre>
# --- Examples -----
# Bar plot of top 30 (global |SHAP|)
ord <- order(colMeans(abs(S), na.rm = TRUE), decreasing = TRUE)</pre>
topK <- colnames(S)[ord[1:min(30, ncol(S))]]
sv importance(sv, kind = "bar", v = topK)
```

Warning: `label` cannot be a <ggplot2::element\_blank> object.



```
library(shapviz)

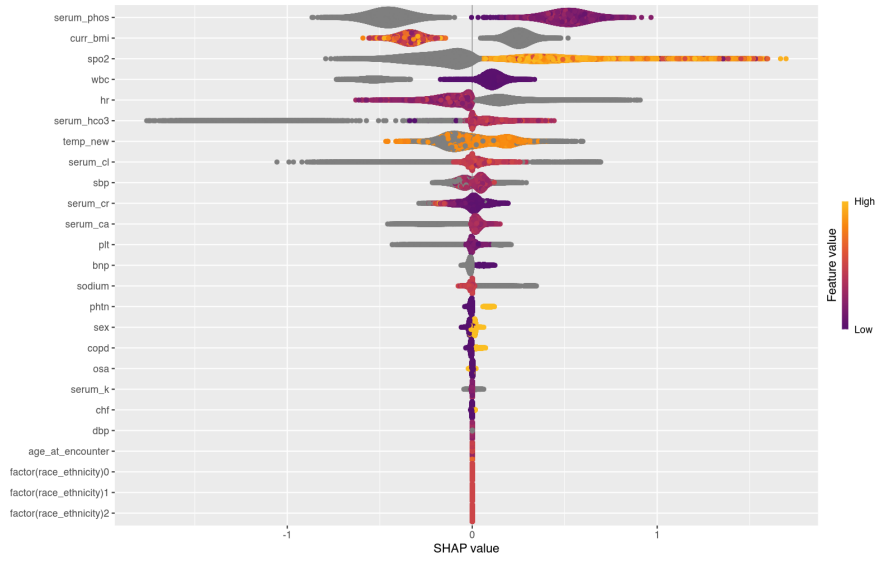
S <- as.matrix(sh_abg_fast$shap) # n x p SHAP values</pre>
```

```
X <- as.data.frame(sh_abg_fast$X) # same rows, p columns (features)

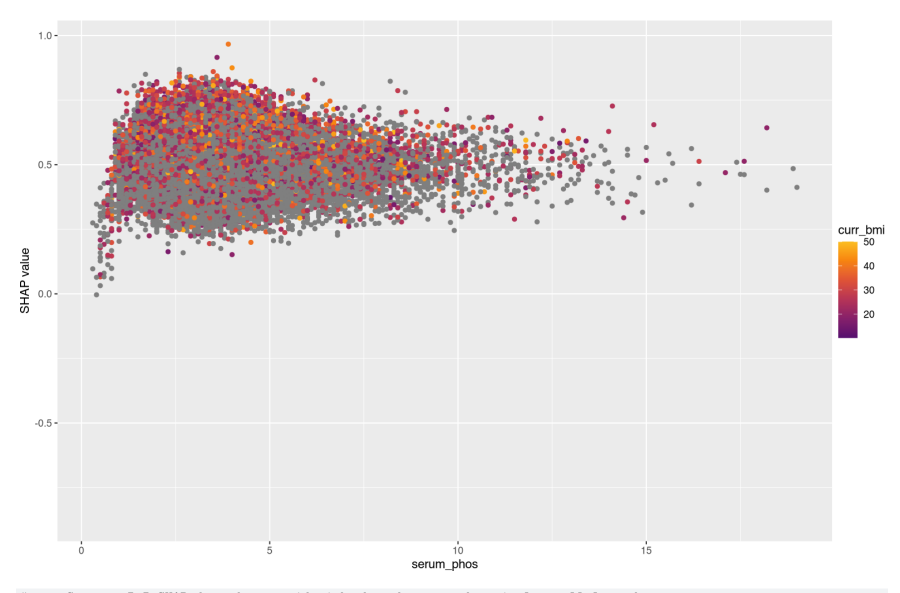
# Build shapviz object
sv <- shapviz(S, X = as.matrix(X))

# Beeswarm-style SHAP summary (like Python SHAP)
sv_importance(sv, kind = "beeswarm", max_display = 25) # overall</pre>
```

Warning: `label` cannot be a <ggplot2::element\_blank> object.



```
# Primary = top feature; color by next feature
imp_order <- colnames(S)[ord]
sv_dependence(sv, v = imp_order[1], color_var = imp_order[2], smooth = TRUE)</pre>
```



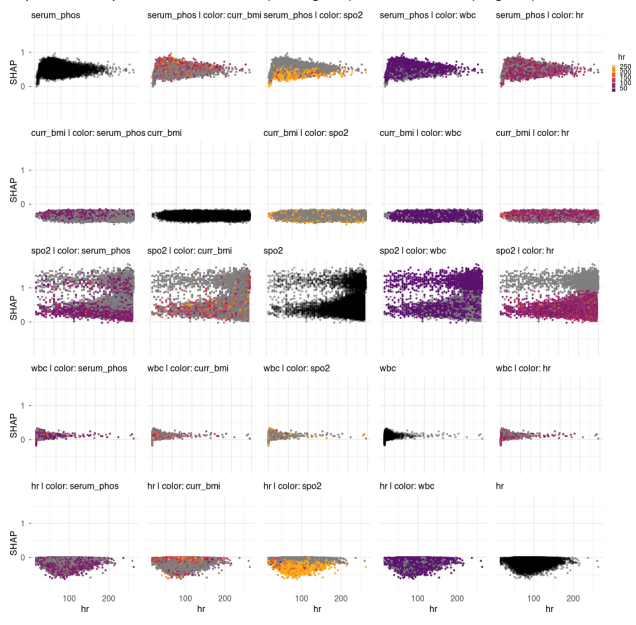
# --- Compact 5x5 SHAP dependence grid with shared axes and a single small legend ---library(shapviz)
library(ggplot2)

```
library(patchwork)
library(grid) # for unit()
# If needed, recover S and X sv from 'sv'
if (!exists("S")) S <- sv$S</pre>
if (!exists("X sv")) X sv <- as.data.frame(sv$X)</pre>
stopifnot(is.matrix(S), is.data.frame(X_sv))
# 1) Top-5 by global mean |SHAP|
ranked <- colnames(S)[order(colMeans(abs(S), na.rm = TRUE), decreasing = TRUE)]</pre>
top5 <- head(ranked, 5)
# 2) Shared y-range across all top-5 features
y rng <- range(unlist(lapply(top5, function(v) S[, v])), finite = TRUE)</pre>
# 3) Small theme helpers
theme axes compact <- function(show y = FALSE, show x = FALSE, base = 8) {
  theme_minimal(base_size = base) +
    theme(
      axis.title.y = if (show y) element text(size = base) else element blank(),
      axis.text.y
                     = if (show_y) element_text(size = base - 1) else element_blank(),
      axis.ticks.y = if (show_y) element_line(linewidth = 0.2) else element_blank(),
      axis.title.x = if (show_x) element_text(size = base) else element_blank(),
      axis.text.x
                     = if (show x) element text(size = base - 1) else element blank(),
                    = element_text(size = base, hjust = 0),
      plot.title
     legend.title = element_text(size = base - 1),
                     = element_text(size = base - 2),
      legend.text
      legend.key.height = unit(22, "pt"),
      legend.key.width = unit(3, "pt"),
                        = margin(0, 0, 0, 0, "pt"),
      legend.margin
     legend.box.margin = margin(0, 0, 0, 0, "pt")
}
```

```
# 4) One cell builder
cell_plot <- function(v_row, v_col, i, j, n) {</pre>
  show_y \leftarrow (j == 1) # y-axis only on first column
  show x \leftarrow (i == n) # x-axis only on bottom row
  if (identical(v row, v col)) {
    # diagonal: unshaded scatter (no legend)
    df <- data.frame(</pre>
     x = as.numeric(X_sv[[v_row]]),
      shap = as.numeric(S[, v_row])
    )
    df <- df[is.finite(df$x) & is.finite(df$shap), , drop = FALSE]</pre>
    ggplot(df, aes(x = x, y = shap)) +
      geom_point(alpha = 0.30, size = 0.45, na.rm = TRUE) +
      scale_y_continuous(limits = y_rng) +
      labs(title = v_row, x = v_row, y = "SHAP") +
      theme_axes_compact(show_y = show_y, show_x = show_x, base = 8) +
      theme(legend.position = "none")
  } else {
    # off-diagonal: color by partner feature
    p <- shapviz::sv_dependence(sv, v = v_row, color_var = v_col, size = 0.4) +
      scale_y_continuous(limits = y_rng) +
      labs(title = paste0(v_row, " | color: ", v_col),
           x = v_{row}, y = "SHAP")
    # keep a single, small legend on the top-right panel only
    keep\_legend \leftarrow (i == 1 \&\& j == length(top5))
    p +
      theme_axes_compact(show_y = show_y, show_x = show_x, base = 8) +
      guides(colour = guide_colorbar(
        barheight = unit(24, "pt"),
        barwidth
                     = unit(3, "pt"),
        title.position = "top",
```

```
title.hjust
                      = 0.5,
        label.position = "right"
      )) +
      theme(legend.position = if (keep_legend) "right" else "none")
  }
}
# 5) Build grid row-wise
n <- length(top5)</pre>
plots <- vector("list", n * n)</pre>
idx <- 1
for (i in seq_len(n)) {
 for (j in seq_len(n)) {
   vr <- top5[i]; vc <- top5[j]</pre>
    plots[[idx]] <- cell_plot(vr, vc, i, j, n)</pre>
    idx \leftarrow idx + 1
  }
}
# 6) Draw: 5 columns, shared layout; keep (not collect) legends so only the chosen one stays
patchwork::wrap_plots(plots, ncol = n, guides = "keep") +
  plot_annotation(title = "Top-5 SHAP dependence: interactions (off-diagonal) and main effects (diagonal)")
```

Top-5 SHAP dependence: interactions (off-diagonal) and main effects (diagonal)



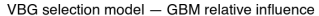
## VBG

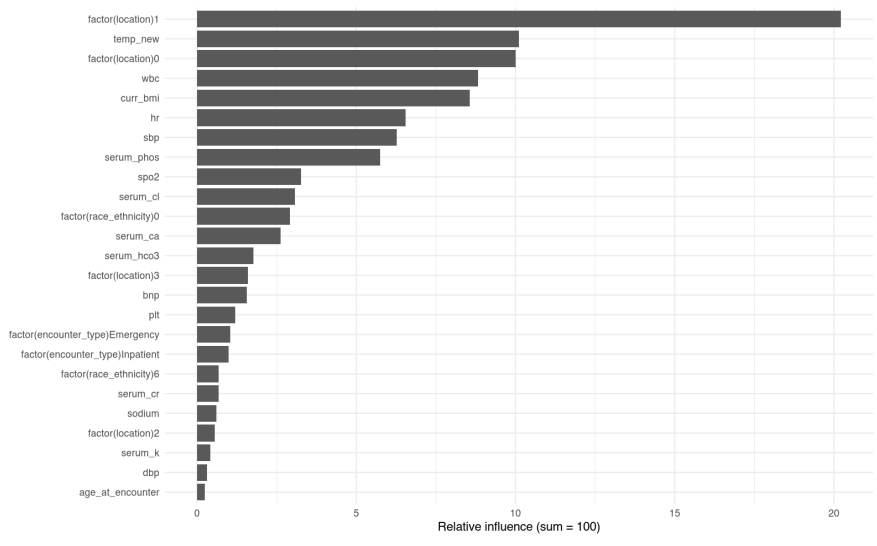
VBG Explainability

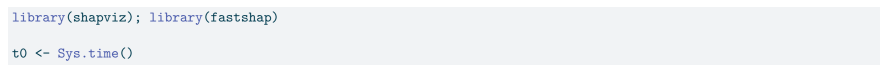
```
library(WeightIt); library(gbm); library(dplyr); library(ggplot2)

stopifnot(exists("w_vbg", inherits = TRUE))
w_vbg <- ensure_gbm_obj(w_vbg)

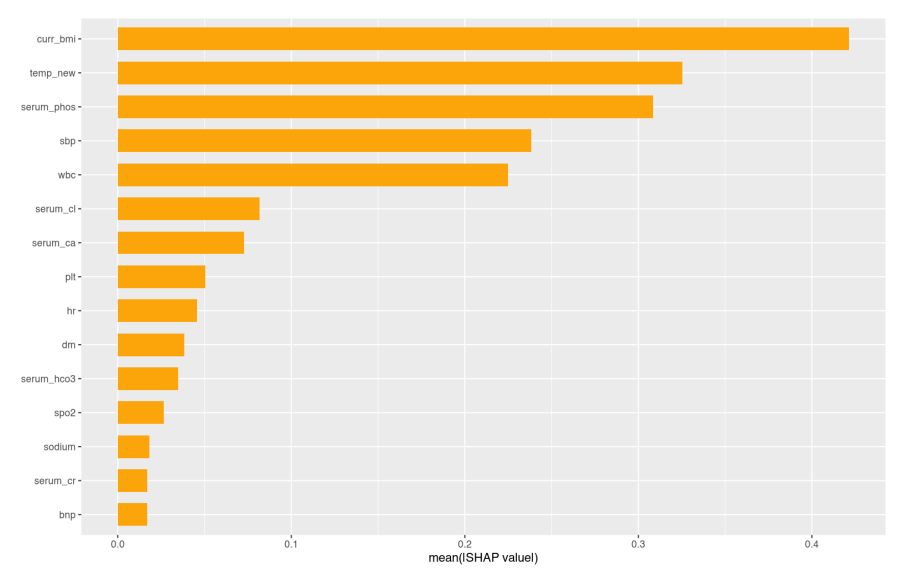
imp_vbg <- extract_gbm_importance(w_vbg, top_n = 25)
p_imp_vbg <- plot_gbm_importance(imp_vbg, "VBG selection model - GBM relative influence")
p_imp_vbg</pre>
```



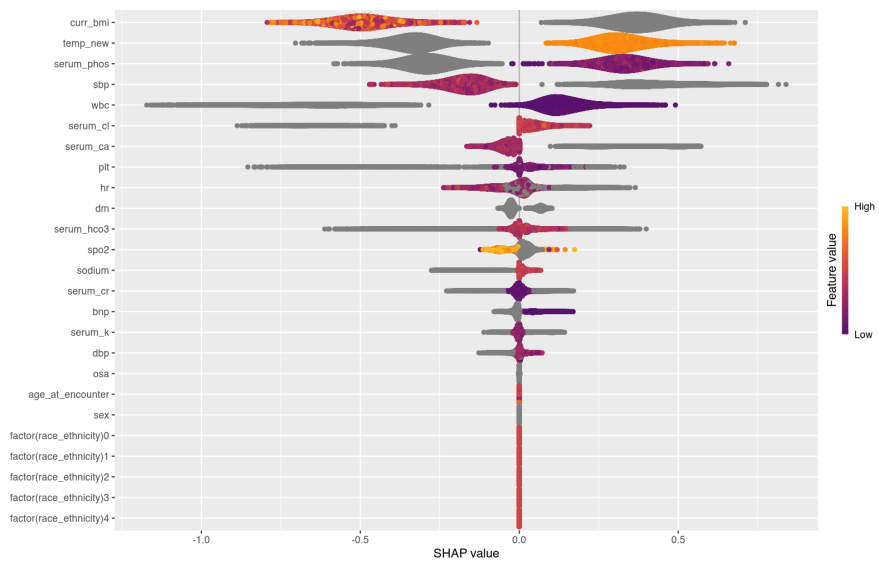




```
sh_vbg_fast <- compute shap_fast(w_vbg, top k = 100, nsim = 32, frac_rows = 0.25, max_rows = 100000)
t1 <- Sys.time(); message(sprintf("[compute_shap_fast VBG] %.2f s", as.numeric(difftime(t1, t0, units="secs"))))
S_vbg <- as.matrix(sh_vbg_fast$shap)</pre>
X_vbg <- as.data.frame(sh_vbg_fast$X)</pre>
for (nm in names(X_vbg)) {
  if (inherits(X_vbg[[nm]], "haven_labelled")) X_vbg[[nm]] <- labelled::to_factor(X_vbg[[nm]])</pre>
  if (is.factor(X_vbg[[nm]])) X_vbg[[nm]] <- as.character(X_vbg[[nm]])</pre>
  if (is.character(X_vbg[[nm]])) suppressWarnings(X_vbg[[nm]]) <- as.numeric(X_vbg[[nm]]))</pre>
}
if (is.null(colnames(S_vbg))) colnames(S_vbg) <- colnames(X_vbg)</pre>
S_vbg <- S_vbg[, intersect(colnames(S_vbg), colnames(X_vbg)), drop = FALSE]
X_vbg <- X_vbg[, colnames(S_vbg), drop = FALSE]</pre>
sv_vbg <- shapviz::shapviz(S_vbg, X = as.matrix(X_vbg))</pre>
ord vbg <- order(colMeans(abs(S vbg), na.rm = TRUE), decreasing = TRUE)
topK_vbg <- colnames(S_vbg)[ord_vbg[1:min(30, ncol(S_vbg))]]</pre>
sv_importance(sv_vbg, kind = "bar", v = topK_vbg)
```



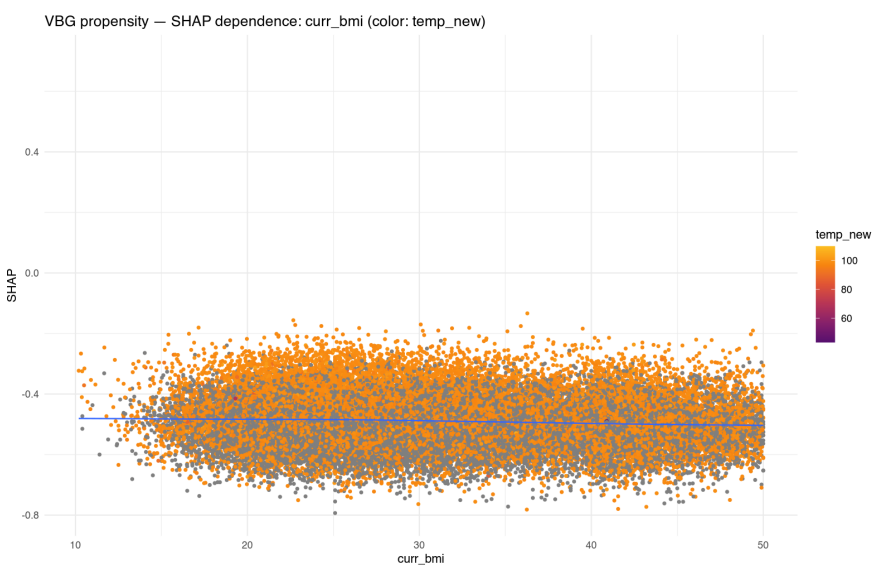
```
library(shapviz)
sv_importance(sv_vbg, kind = "beeswarm", max_display = 25)
```



```
library(ggplot2)
imp_order_vbg <- colnames(S_vbg)[order(colMeans(abs(S_vbg), na.rm = TRUE), decreasing = TRUE)]</pre>
```

```
pri_vbg <- imp_order_vbg[1]
aux_vbg <- imp_order_vbg[2]
if (identical(aux_vbg, pri_vbg) || !(aux_vbg %in% colnames(X_vbg))) aux_vbg <- imp_order_vbg[3]

shapviz::sv_dependence(sv_vbg, v = pri_vbg, color_var = aux_vbg, size = 1) +
    geom_smooth(se = FALSE, method = "loess", formula = y ~ x, linewidth = 0.6) +
    labs(title = sprintf("VBG propensity - SHAP dependence: %s (color: %s)", pri_vbg, aux_vbg),
        x = pri_vbg, y = "SHAP") +
    theme_minimal(base_size = 11)</pre>
```

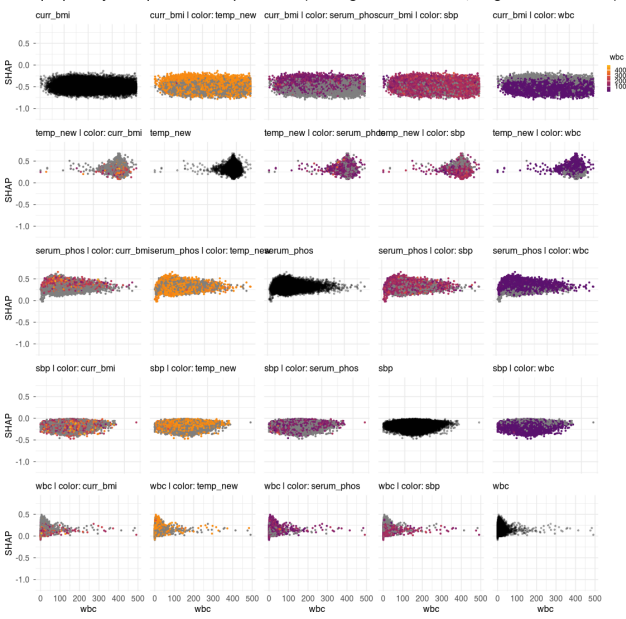


library(shapviz); library(ggplot2); library(patchwork); library(grid)
stopifnot(is.matrix(S\_vbg), is.data.frame(X\_vbg))

```
ranked_vbg <- colnames(S_vbg)[order(colMeans(abs(S_vbg), na.rm = TRUE), decreasing = TRUE)]</pre>
top5 vbg <- head(ranked vbg, 5)
y_rng_vbg <- range(unlist(lapply(top5_vbg, function(v) S_vbg[, v])), finite = TRUE)</pre>
theme_axes_compact <- function(show_y = FALSE, show_x = FALSE, base = 8) {</pre>
  theme minimal(base size = base) +
    theme(
      axis.title.y = if (show y) element text(size = base) else element blank(),
      axis.text.y
                     = if (show_y) element_text(size = base - 1) else element_blank(),
      axis.ticks.y
                    = if (show_y) element_line(linewidth = 0.2) else element_blank(),
                    = if (show_x) element_text(size = base) else element_blank(),
      axis.title.x
      axis.text.x
                     = if (show_x) element_text(size = base - 1) else element_blank(),
                    = element_text(size = base, hjust = 0),
      plot.title
     legend.title = element_text(size = base - 1),
                     = element_text(size = base - 2),
      legend.text
      legend.key.height = unit(22, "pt"),
      legend.key.width = unit(3, "pt"),
      legend.margin
                     = margin(0, 0, 0, 0, "pt"),
      legend.box.margin = margin(0, 0, 0, 0, "pt")
}
cell_plot_vbg <- function(v_row, v_col, i, j, n) {</pre>
  show y < - (j == 1); show x < - (i == n)
  if (identical(v_row, v_col)) {
    df <- data.frame(x = as.numeric(X_vbg[[v_row]]), shap = as.numeric(S_vbg[, v_row]))</pre>
    df <- df[is.finite(df$x) & is.finite(df$shap), , drop = FALSE]</pre>
    ggplot(df, aes(x = x, y = shap)) +
      geom_point(alpha = 0.30, size = 0.45, na.rm = TRUE) +
      scale_y_continuous(limits = y_rng_vbg) +
      labs(title = v_row, x = v_row, y = "SHAP") +
      theme_axes_compact(show_y, show_x, base = 8) +
      theme(legend.position = "none")
```

```
} else {
    keep_legend <- (i == 1 && j == length(top5_vbg))</pre>
    shapviz::sv_dependence(sv_vbg, v = v_row, color_var = v_col, size = 0.4) +
      scale_y_continuous(limits = y_rng_vbg) +
      labs(title = paste0(v_row, " | color: ", v_col), x = v_row, y = "SHAP") +
      theme axes compact(show y, show x, base = 8) +
      guides(colour = guide_colorbar(barheight = unit(24, "pt"), barwidth = unit(3, "pt"),
                                      title.position = "top", title.hjust = 0.5, label.position = "right")) +
      theme(legend.position = if (keep_legend) "right" else "none")
 }
}
n <- length(top5_vbg); plots <- vector("list", n * n); k <- 1</pre>
for (i in seq_len(n)) for (j in seq_len(n)) {
  plots[[k]] \leftarrow cell_plot_vbg(top5_vbg[i], top5_vbg[j], i, j, n); k \leftarrow k + 1
patchwork::wrap_plots(plots, ncol = n, guides = "keep") +
  plot_annotation(title = "VBG propensity - Top-5 SHAP dependence (off-diagonal interactions, diagonal main effects)")
```

#### VBG propensity — Top-5 SHAP dependence (off-diagonal interactions, diagonal main effects)

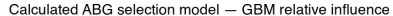


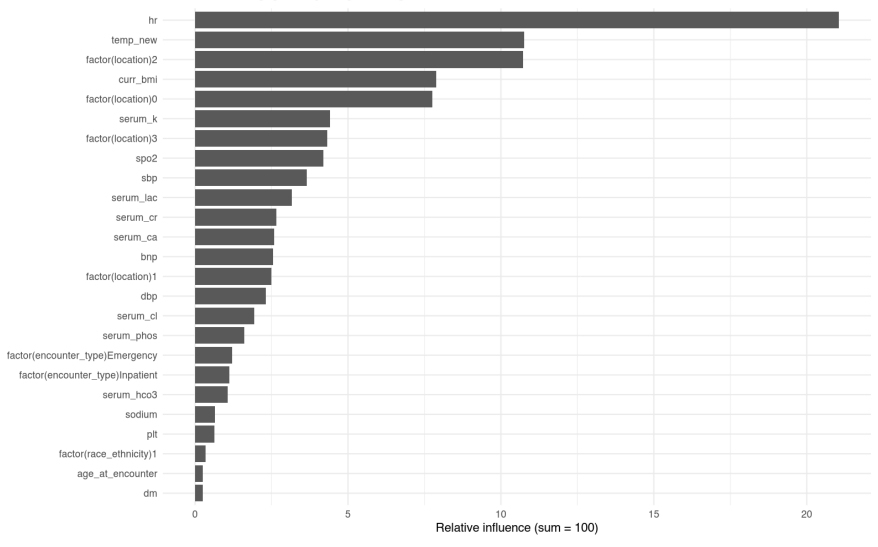
#### VBG-Calc

```
library(WeightIt); library(gbm); library(dplyr); library(ggplot2)

stopifnot(exists("w_vbg_calc", inherits = TRUE))
w_vbg_calc <- ensure_gbm_obj(w_vbg_calc)

imp_calc <- extract_gbm_importance(w_vbg_calc, top_n = 25)
p_imp_calc <- plot_gbm_importance(imp_calc, "Calculated ABG selection model - GBM relative influence")
p_imp_calc</pre>
```

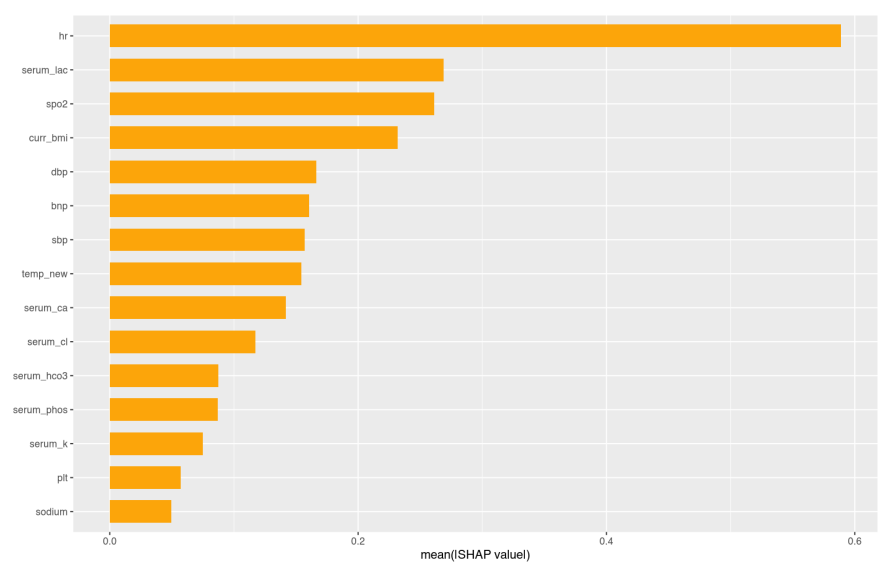




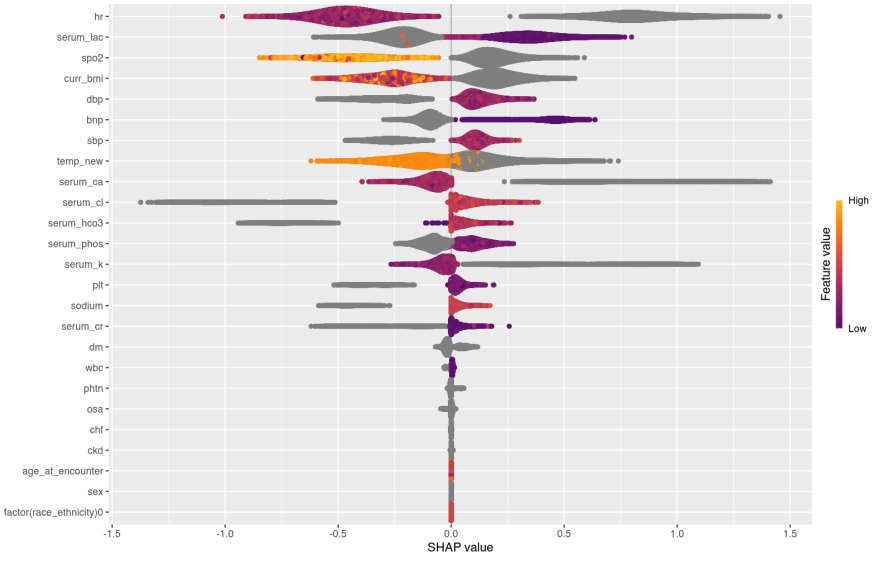
```
library(shapviz); library(fastshap)

t0 <- Sys.time()</pre>
```

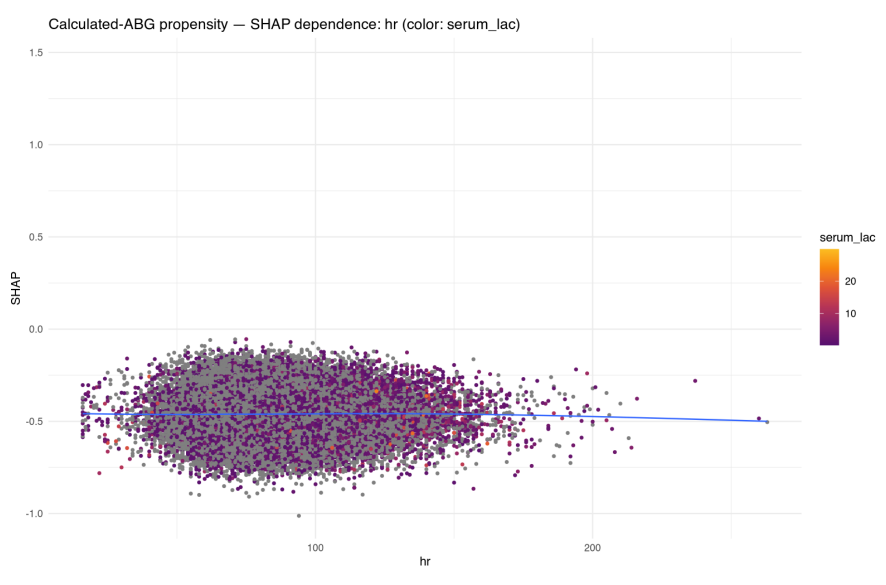
```
sh_calc_fast <- compute_shap_fast(w_vbg_calc, top_k = 100, nsim = 32, frac_rows = 0.25, max_rows = 100000)
t1 <- Sys.time(); message(sprintf("[compute_shap_fast Calc-ABG] %.2f s", as.numeric(difftime(t1, t0, units="secs"))))
S_calc <- as.matrix(sh_calc_fast$shap)</pre>
X_calc <- as.data.frame(sh_calc_fast$X)</pre>
for (nm in names(X_calc)) {
  if (inherits(X_calc[[nm]], "haven_labelled")) X_calc[[nm]] <- labelled::to_factor(X_calc[[nm]])</pre>
  if (is.factor(X_calc[[nm]])) X_calc[[nm]] <- as.character(X_calc[[nm]])</pre>
  if (is.character(X_calc[[nm]])) suppressWarnings(X_calc[[nm]] <- as.numeric(X_calc[[nm]]))</pre>
}
if (is.null(colnames(S_calc))) colnames(S_calc) <- colnames(X_calc)</pre>
S_calc <- S_calc[, intersect(colnames(S_calc), colnames(X_calc)), drop = FALSE]</pre>
X_calc <- X_calc[, colnames(S_calc), drop = FALSE]</pre>
sv_calc <- shapviz::shapviz(S_calc, X = as.matrix(X_calc))</pre>
ord calc <- order(colMeans(abs(S calc), na.rm = TRUE), decreasing = TRUE)
topK_calc <- colnames(S_calc)[ord_calc[1:min(30, ncol(S_calc))]]</pre>
sv_importance(sv_calc, kind = "bar", v = topK_calc)
```



```
library(shapviz)
sv_importance(sv_calc, kind = "beeswarm", max_display = 25)
```



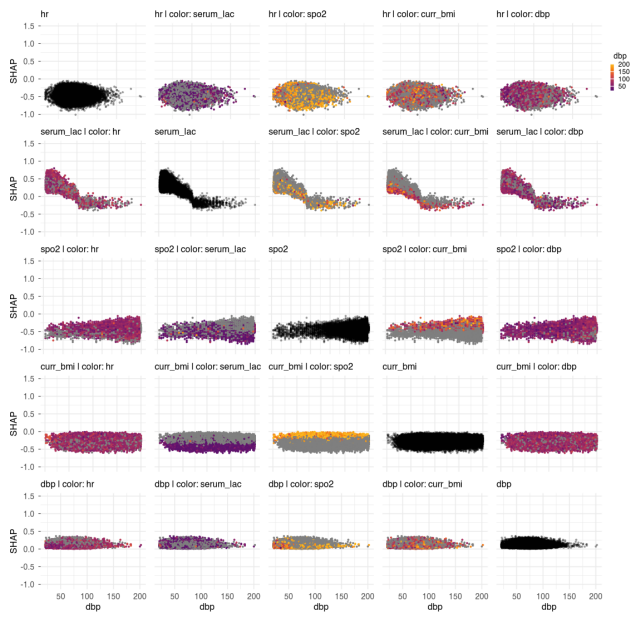
```
library(ggplot2)
imp_order_calc <- colnames(S_calc)[order(colMeans(abs(S_calc), na.rm = TRUE), decreasing = TRUE)]</pre>
```



```
library(shapviz); library(ggplot2); library(patchwork); library(grid)
stopifnot(is.matrix(S_calc), is.data.frame(X_calc))
```

```
ranked_calc <- colnames(S_calc)[order(colMeans(abs(S_calc), na.rm = TRUE), decreasing = TRUE)]
top5 calc <- head(ranked calc, 5)</pre>
y_rng_calc <- range(unlist(lapply(top5_calc, function(v) S_calc[, v])), finite = TRUE)</pre>
theme_axes_compact <- function(show_y = FALSE, show_x = FALSE, base = 8) {</pre>
  theme minimal(base size = base) +
    theme(
      axis.title.y = if (show y) element text(size = base) else element blank(),
      axis.text.y
                     = if (show_y) element_text(size = base - 1) else element_blank(),
      axis.ticks.y
                    = if (show_y) element_line(linewidth = 0.2) else element_blank(),
                    = if (show_x) element_text(size = base) else element_blank(),
      axis.title.x
      axis.text.x
                     = if (show_x) element_text(size = base - 1) else element_blank(),
                    = element_text(size = base, hjust = 0),
      plot.title
     legend.title = element_text(size = base - 1),
                     = element_text(size = base - 2),
      legend.text
      legend.key.height = unit(22, "pt"),
      legend.key.width = unit(3, "pt"),
      legend.margin
                     = margin(0, 0, 0, 0, "pt"),
      legend.box.margin = margin(0, 0, 0, 0, "pt")
}
cell_plot_calc <- function(v_row, v_col, i, j, n) {</pre>
  show y < - (j == 1); show x < - (i == n)
  if (identical(v_row, v_col)) {
    df <- data.frame(x = as.numeric(X_calc[[v_row]]), shap = as.numeric(S_calc[, v_row]))</pre>
    df <- df[is.finite(df$x) & is.finite(df$shap), , drop = FALSE]</pre>
    ggplot(df, aes(x = x, y = shap)) +
      geom_point(alpha = 0.30, size = 0.45, na.rm = TRUE) +
      scale_y_continuous(limits = y_rng_calc) +
      labs(title = v_row, x = v_row, y = "SHAP") +
      theme_axes_compact(show_y, show_x, base = 8) +
      theme(legend.position = "none")
```

### Calculated-ABG propensity — Top-5 SHAP dependence (off-diagonal interactions, diagonal mair

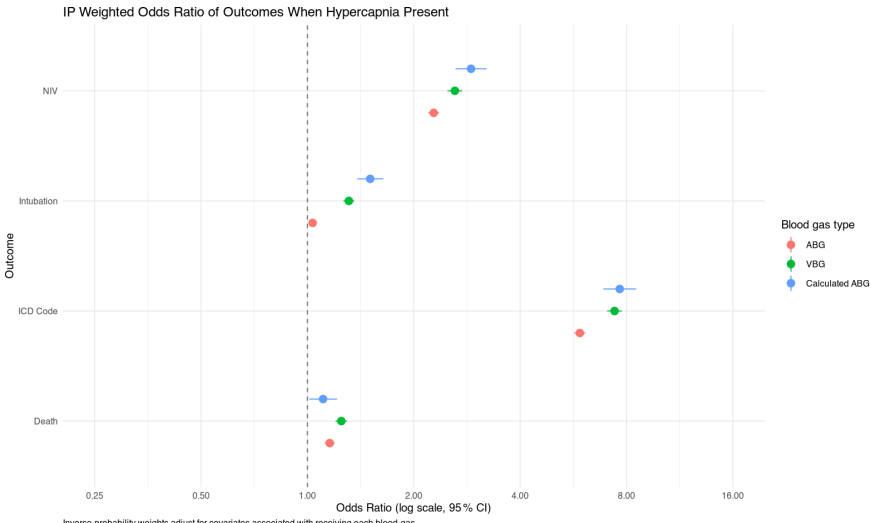


New weighted binary regression figures.

```
IP-weighted odds-ratio plot (ABG, VBG, Calculated-ABG)
   - exact analogue of the un-weighted figure
# weights already attached earlier:
# • w_abg - propensity for *ABG* (column in subset_data)
              - propensity for *VBG* (column in subset_data)
 w_vbg
# • w_vbg_calc - same weights, used for calculated ABG CO
# 1. helper to fit an IP-weighted GLM and return tidy OR -----
tidy_ipw <- function(data, outcome, exposure, weight_var,</pre>
                   group_label, outcome_label) {
 des <- svydesign(ids = ~1, weights = as.formula(paste0("~", weight_var)),</pre>
                 data = data)
 mod <- svyglm(</pre>
   as.formula(paste0(outcome, " ~ ", exposure)),
   design = des,
   family = quasibinomial()
 tidy(mod, exponentiate = TRUE, conf.int = TRUE) %>%
   filter(term == exposure) %>% # keep the exposure row
   mutate(group = group_label, outcome = outcome_label)
# 2. cohort-specific data frames ------
abg_df <- subset_data %>% filter(has_abg == 1)
vbg_df <- subset_data %>% filter(has_vbg == 1)
calc_df <- subset_data %>% filter(!is.na(calc_abg)) # implies VBG present
# 3. fit models & collect estimates -----
ipw_estimates <- bind_rows(</pre>
```

```
# ABG
 tidy ipw(abg df, "imv proc",
                                             "hypercap_on_abg", "w_abg",
                                                                              "ABG",
                                                                                               "Intubation"),
 tidy ipw(abg df, "niv proc",
                                             "hypercap_on_abg", "w_abg",
                                                                                               "NIV"),
                                                                              "ABG",
 tidy_ipw(abg_df, "death_60d",
                                                  "hypercap_on_abg", "w_abg",
                                                                               "ABG",
                                                                                                    "Death").
 tidy_ipw(abg_df, "hypercap_resp_failure", "hypercap_on_abg", "w_abg",
                                                                                               "ICD Code"),
                                                                              "ABG",
  # VBG
 tidy_ipw(vbg_df, "imv_proc",
                                                                              "VBG",
                                                                                               "Intubation"),
                                             "hypercap_on_vbg", "w_vbg",
 tidy_ipw(vbg_df, "niv_proc",
                                             "hypercap_on_vbg", "w_vbg",
                                                                                               "NIV"),
                                                                              "VBG",
                                                  "hypercap_on_vbg", "w_vbg",
                                                                                "VBG",
                                                                                                    "Death"),
 tidy_ipw(vbg_df, "death_60d",
 tidy ipw(vbg_df, "hypercap resp_failure", "hypercap on vbg", "w_vbg",
                                                                              "VBG",
                                                                                               "ICD Code"),
 # Calculated ABG
                                             "hypercapnia_calc", "w_vbg_calc", "Calculated ABG", "Intubation"),
 tidy_ipw(calc_df, "imv_proc",
 tidy_ipw(calc_df, "niv_proc",
                                             "hypercapnia_calc", "w_vbg_calc", "Calculated ABG", "NIV"),
                                                  "hypercapnia_calc", "w_vbg_calc", "Calculated ABG", "Death"),
 tidy_ipw(calc_df, "death_60d",
 tidy_ipw(calc_df, "hypercap_resp_failure", "hypercapnia_calc", "w_vbg_calc", "Calculated ABG", "ICD Code")
# 4. plotting -----
ipw_estimates$group <- factor(</pre>
 ipw_estimates$group,
 levels = c("ABG", "VBG", "Calculated ABG")
ggplot(
  ipw_estimates,
  aes(
          = outcome,
   X
         = estimate,
   ymin = conf.low,
   ymax = conf.high,
   color = group
```

```
) +
  geom_pointrange(position = position_dodge(width = 0.6), size = 0.6) +
  geom_hline(yintercept = 1, linetype = "dashed", colour = "grey40") +
  scale_y_log10(
   breaks = c(0.25, 0.5, 1, 2, 4, 8, 16),
   limits = c(0.25, 16),
   labels = number_format(accuracy = 0.01)
  ) +
  coord_flip() +
  labs(
    title = "IP Weighted Odds Ratio of Outcomes When Hypercapnia Present",
          = "Outcome",
    X
          = "Odds Ratio (log scale, 95 % CI)",
    color = "Blood gas type",
    caption = paste(
      "Inverse-probability weights adjust for covariates associated with receiving each blood-gas.",
      "Models are fitted within their respective cohorts:",
      "ABG (weights = w_abg), VBG (w_vbg), Calculated ABG (w_vbg_calc).",
      "Numerator = hypercapnic; denominator = normocapnic within cohort.",
      sep = "\n"
  ) +
  theme_minimal(base_size = 10) +
  theme(plot.caption = element_text(hjust = 0))
```



Inverse-probability weights adjust for covariates associated with receiving each blood-gas. Models are fitted within their respective cohorts:

ABG (weights=w\_abg), VBG (w\_vbg), Calculated ABG (w\_vbg\_calc).

Numerator = hypercapnic; denominator = normocapnic within cohort.

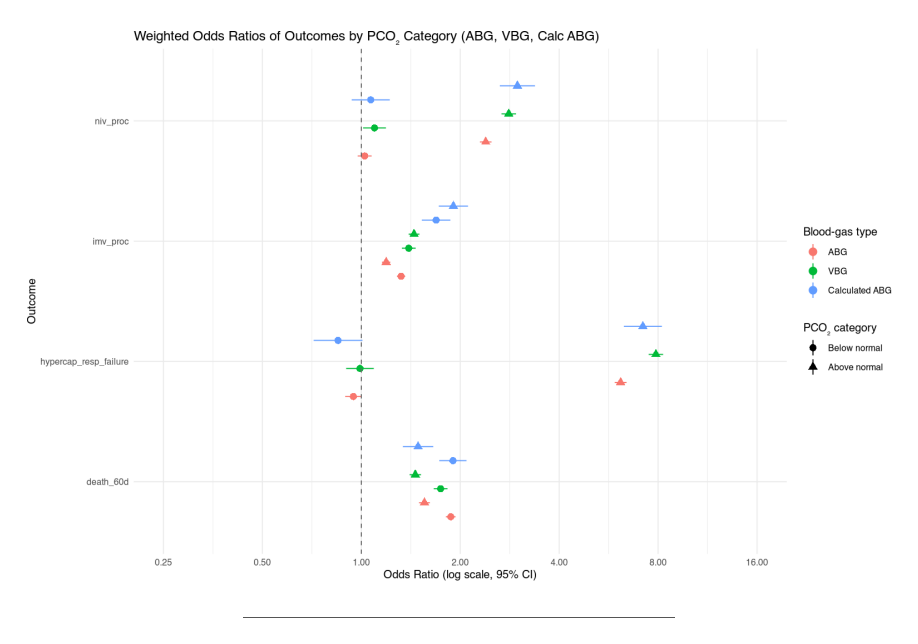
Three Groups with Weights

```
library(dplyr)
library(survey)
library(broom)
library(ggplot2)
library(scales)
# 1. Create PCO categories
subset_data <- subset_data %>%
  mutate(
    pco2_cat_abg = case_when(
      !is.na(paco2) & paco2 < 35 ~ "Below normal",
      !is.na(paco2) & paco2 >= 35 & paco2 <= 45 ~ "Normal",
      !is.na(paco2) & paco2 > 45 ~ "Above normal",
      TRUE ~ NA_character_
    ),
    pco2_cat_vbg = case_when(
      !is.na(vbg co2) & vbg co2 < 35 ~ "Below normal",
      !is.na(vbg_co2) & vbg_co2 >= 35 & vbg_co2 <= 50 ~ "Normal",
      !is.na(vbg_co2) & vbg_co2 > 50 ~ "Above normal",
      TRUE ~ NA character
    ),
    pco2_cat_calc = case_when(
      !is.na(calc_abg) & calc_abg < 35 ~ "Below normal",
      !is.na(calc_abg) & calc_abg >= 35 & calc_abg <= 45 ~ "Normal",
      !is.na(calc_abg) & calc_abg > 45 ~ "Above normal",
      TRUE ~ NA_character_
  )
# 2. Function: weighted logistic regression & OR extraction
run_weighted_or <- function(data, outcome, cat_var, weight_var, group_name) {</pre>
  dat <- data %>%
    filter(
      !is.na(.data[[cat_var]]),
```

```
!is.na(.data[[outcome]]),
      !is.na(.data[[weight_var]]),
      .data[[weight_var]] > 0
    ) %>%
    mutate(
      !!cat_var := factor(.data[[cat_var]],
                          levels = c("Normal", "Below normal", "Above normal"))
    ) %>%
    droplevels()
 design <- svydesign(</pre>
   ids = ~1,
   weights = as.formula(paste0("~", weight_var)),
   data = dat
 fit <- svyglm(as.formula(paste(outcome, "~", cat_var)),</pre>
                design = design, family = quasibinomial())
 tidy(fit, exponentiate = TRUE, conf.int = TRUE) %>%
    filter(term != "(Intercept)") %>%
   mutate(
      group
              = group_name,
      outcome = outcome,
      exposure = gsub(paste0(cat_var), "", term) %>%
                   gsub("`", "", .)
    3. Run across outcomes & cohorts
outcomes <- c("imv_proc", "niv_proc", "death 60d", "hypercap_resp_failure")
combined_or_df <- bind_rows(</pre>
 lapply(outcomes, function(out)
```

```
"ABG")),
    run_weighted_or(subset_data, out, "pco2_cat_abg", "w_abg",
 lapply(outcomes, function(out)
    run_weighted_or(subset_data, out, "pco2_cat_vbg", "w_vbg",
                                                                     "VBG")),
 lapply(outcomes, function(out)
    run_weighted_or(subset_data, out, "pco2_cat_calc", "w_vbg_calc", "Calculated ABG"))
# Ensure nice ordering
combined_or_df$group
                        <- factor(combined_or_df$group,</pre>
                                  levels = c("ABG", "VBG", "Calculated ABG"))
combined_or_df$exposure <- factor(combined_or_df$exposure,</pre>
                                  levels = c("Below normal", "Above normal"))
# 4. Plot weighted odds ratios
ggplot(
  combined_or_df,
 aes(
   x = outcome,
    y = estimate,
   ymin = conf.low,
   ymax = conf.high,
   color = group,
   shape = exposure
 geom pointrange(position = position_dodge(width = 0.7), size = 0.6) +
  geom hline(yintercept = 1, linetype = "dashed", colour = "grey40") +
 scale_y_log10(
   breaks = c(0.25, 0.5, 1, 2, 4, 8, 16),
   limits = c(0.25, 16),
   labels = number_format(accuracy = 0.01)
 ) +
 coord_flip() +
 labs(
```

```
title = "Weighted Odds Ratios of Outcomes by PCO Category (ABG, VBG, Calc ABG)",
    x = "Outcome",
    y = "Odds Ratio (log scale, 95% CI)",
    color = "Blood-gas type",
    shape = "PCO category"
) +
theme_minimal(base_size = 10) +
theme(plot.caption = element_text(hjust = 0))
```

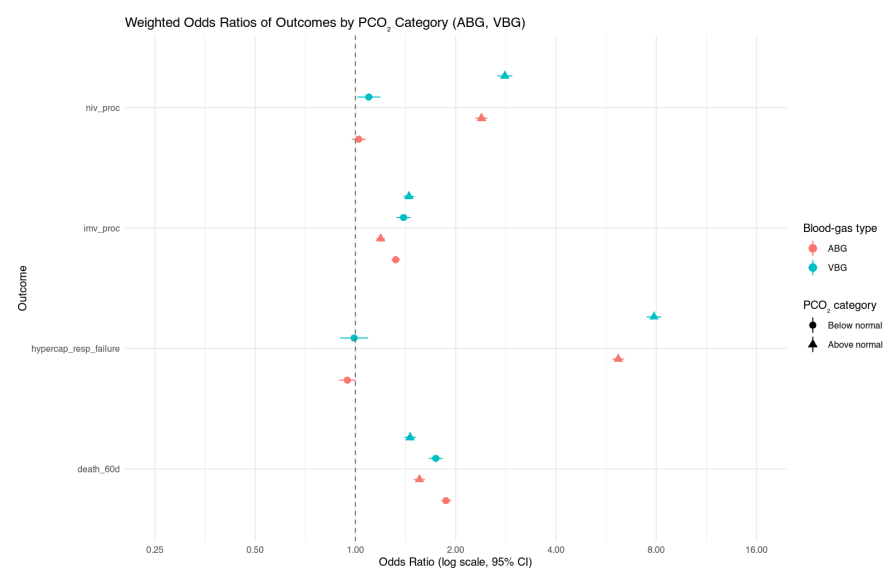


Three groups with weights: Just ABG and VBG  $\,$ 

```
library(dplyr)
library(survey)
library(broom)
library(ggplot2)
library(scales)
    2. Function: weighted logistic regression & OR extraction
run_weighted_or <- function(data, outcome, cat_var, weight_var, group_name) {</pre>
  dat <- data %>%
    filter(
      !is.na(.data[[cat_var]]),
      !is.na(.data[[outcome]]),
      !is.na(.data[[weight_var]]),
      .data[[weight_var]] > 0
    ) %>%
    mutate(
      !!cat_var := factor(.data[[cat_var]],
                          levels = c("Normal", "Below normal", "Above normal"))
    ) %>%
    droplevels()
  design <- svydesign(</pre>
    ids = ~1,
    weights = as.formula(paste0("~", weight_var)),
    data = dat
  fit <- svyglm(as.formula(paste(outcome, "~", cat_var)),</pre>
                design = design, family = quasibinomial())
  tidy(fit, exponentiate = TRUE, conf.int = TRUE) %>%
    filter(term != "(Intercept)") %>%
    mutate(
               = group_name,
      group
```

```
outcome = outcome,
      exposure = gsub(paste0(cat_var), "", term) %>%
                   gsub("`", "", .)
    3. Run across outcomes & cohorts
outcomes <- c("imv_proc", "niv_proc", "death_60d", "hypercap_resp_failure")</pre>
combined_or_df <- bind_rows(</pre>
 lapply(outcomes, function(out)
    run_weighted_or(subset_data, out, "pco2_cat_abg", "w_abg",
                                                                       "ABG")),
 lapply(outcomes, function(out)
   run_weighted_or(subset_data, out, "pco2_cat_vbg", "w_vbg",
                                                                       "VBG"))
# Ensure nice ordering
combined_or_df$group
                        <- factor(combined_or_df$group,</pre>
                                  levels = c("ABG", "VBG"))
combined_or_df$exposure <- factor(combined_or_df$exposure,</pre>
                                  levels = c("Below normal", "Above normal"))
   4. Plot weighted odds ratios
ggplot(
  combined_or_df,
  aes(
    X
           = outcome,
           = estimate,
   ymin = conf.low,
    ymax
          = conf.high,
    color = group,
    shape = exposure
```

```
geom_pointrange(position = position_dodge(width = 0.7), size = 0.6) +
geom_hline(yintercept = 1, linetype = "dashed", colour = "grey40") +
scale_y_log10(
 breaks = c(0.25, 0.5, 1, 2, 4, 8, 16),
 limits = c(0.25, 16),
 labels = number_format(accuracy = 0.01)
) +
coord_flip() +
labs(
 title = "Weighted Odds Ratios of Outcomes by PCO Category (ABG, VBG)",
        = "Outcome",
        = "Odds Ratio (log scale, 95% CI)",
 color = "Blood-gas type",
 shape = "PCO category"
theme_minimal(base_size = 10) +
theme(plot.caption = element_text(hjust = 0))
```



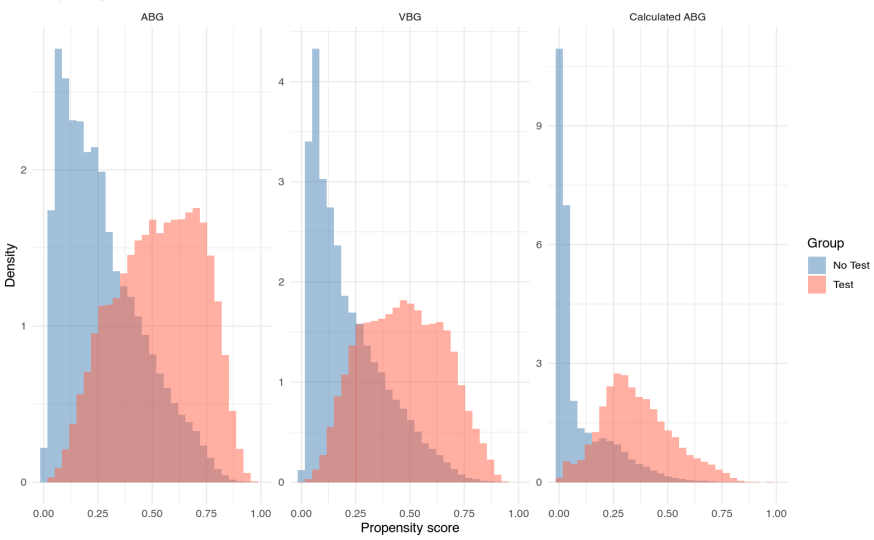
Plotting propensity scores

```
# --- Propensity score histograms (ABG / VBG / Calculated-ABG) ------
# ABG = arterial blood gas; VBG = venous blood gas
library(dplyr)
library(ggplot2)
library(scales)
# Resolve WeightIt objects regardless of naming used upstream
            <- if (exists("w_abg")) w_abg else if (exists("weight_model")) weight_model else NULL</pre>
w_abg_obj
w_vbg_obj <- if (exists("w_vbg")) w_vbg else NULL</pre>
w_vbg_calc_obj <- if (exists("w_vbg_calc")) w_vbg_calc else if (exists("w_vbg")) w_vbg_else NULL
if (is.null(w_abg_obj)) stop("ABG WeightIt object not found. Define `w_abg` or `weight model` before this block.")
if (!"has abg" %in% names(subset_data)) stop("`subset_data` must contain `has_abg` for ABG PS plotting.")
# Build list of per-cohort PS data frames conditionally (so missing cohorts don't error)
ps_dfs <- list(</pre>
 ABG = data.frame(
              = w_abg_obj$ps,
           = subset_data$has_abg,
   treat
    ScoreType = "ABG"
 )
if (!is.null(w_vbg_obj) && "has_vbg" %in% names(subset_data)) {
  ps_dfs$VBG <- data.frame(</pre>
              = w_vbg_obj$ps,
    ps
    treat
              = subset_data$has_vbg,
    ScoreType = "VBG"
} else if (is.null(w_vbg_obj)) {
 message("Note: VBG WeightIt object `w_vbg` not found; skipping VBG panel.")
}
```

```
# Calculated ABG uses the VBG selection model; prefer a dedicated `w_vbg_calc` if present
if (!is.null(w_vbg_calc_obj) && "has_vbg_co2_o2_sat" %in% names(subset_data)) {
  ps dfs$CalcABG <- data.frame(</pre>
             = w_vbg_calc_obj$ps,
    ps
    treat
            = subset_data$has_vbg_co2_o2_sat,
    ScoreType = "Calculated ABG"
} else if (is.null(w_vbg_calc_obj)) {
 message("Note: Calculated-ABG WeightIt object `w_vbg_calc` (or fallback `w_vbg`) not found; skipping Calc-ABG panel.")
# Bind, clean, and factorize for plotting
df_ps <- bind_rows(ps_dfs) %>%
 filter(!is.na(ps), !is.na(treat)) %>%
 mutate(
              = factor(treat, levels = c(0, 1), labels = c("No Test", "Test")),
   treat
   ScoreType = factor(ScoreType, levels = c("ABG", "VBG", "Calculated ABG"))
# Plot
ggplot(df_ps, aes(x = ps, fill = treat)) +
 geom_histogram(aes(y = ..density..), alpha = 0.5,
                position = "identity", bins = 30) +
 scale_fill_manual(values = c("No Test" = "steelblue", "Test" = "tomato")) +
 facet_wrap(~ScoreType, scales = "free_y") +
  coord_cartesian(xlim = c(0, 1)) +
 labs(
   title = "Propensity Score Distributions",
         = "Propensity score",
         = "Density",
   fill = "Group"
  ) +
  theme minimal(base size = 12)
```

Warning: The dot-dot notation (`..density..`) was deprecated in ggplot2 3.4.0. i Please use `after\_stat(density)` instead.

## Propensity Score Distributions



```
df_ps <- bind_rows(</pre>
  data.frame(
            = w_abg$ps,
    ps
   treat = subset_data$has_abg,
   ScoreType = "ABG"
  ),
  data.frame(
             = w_vbg$ps,
   ps
   treat = subset_data$has_vbg,
   ScoreType = "VBG"
 ),
  data.frame(
          = w_vbg_calc$ps,
    ps
   treat = subset_data$has_vbg_co2_o2_sat,
   ScoreType = "Calculated ABG"
 )
) %>%
 mutate(
   treat = factor(treat, levels = c(0,1), labels = c("No Test", "Test"))
 )
ggplot(df_ps, aes(x = ps, fill = treat)) +
  geom_histogram(aes(y = ..density..), alpha = 0.5,
                position = "identity", bins = 30) +
  scale fill manual(values = c("No Test" = "steelblue", "Test" = "tomato")) +
 facet_wrap(~ScoreType, scales = "free_y") +
  labs(
   title = "Propensity Score Distributions",
   x = "Propensity Score",
   y = "Density",
   fill = "Group"
  ) +
  theme_minimal(base_size = 12)
```

# Propensity Score Distributions

