# Package 'ProAE'

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Title PRO-CTCAE Scoring, Analysis, and Graphical Tools Version 1.0.3 Description A collection of tools to facilitate standardized analysis and graphical procedures when using the National Cancer Institute's Patient-Reported Outcomes version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE) and other PRO measurements. License GPL-3 **Encoding** UTF-8 LazyData true Imports ggplot2, ggnewscale, ggtext, dplyr, Hmisc, magrittr, ggpattern, DescTools, gridExtra, kableExtra Suggests knitr, rmarkdown, testthat VignetteBuilder knitr **Depends** R (>= 4.0.0) RoxygenNote 7.2.3 NeedsCompilation no Author Blake Langlais [aut, cre], Brie Noble [ctb], Mia Truman [ctb], Molly Voss [ctb], Amylou Dueck [aut] Maintainer Blake Langlais <langlais.blake@mayo.edu> **Repository** CRAN

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ProAE

ProAE.

#### Description

This package is a collection of tools for analyzing and plotting PRO-CTCAE data

PROCTCAE\_table PRO-CTCAE variable / label crosswalk

#### Description

A crosswalk / look-up table of expected variable names for associated PRO-CTCAE symptom items.

# Format

A data frame with 124 rows and 2 variables

# Details

- name. Expected variable name item number/letter corresponds to the NCI-PRO-CTCAE (English) Item Library Version 1.0
- short\_label. Item label including the item symptom group and frequency, severity, interference, or presence component

#### Author(s)

Blake Langlais

#### References

https://healthcaredelivery.cancer.gov/pro-ctcae/instrument-pro.html

Create longitudinal mean score line plots for PRO-CTCAE data with modified Area Under the Curve (AUC) estimates showing descriptive symptomatic adverse event burden worsening and improvement from baseline.

#### Description

toxAUC

Data format should be in 'long' format, where each PRO-CTCAE item is a variable/column. AUC calculations will only include subjects with non-missing baseline values (within each PRO-CTCAE item).

#### Usage

```
toxAUC(
  dsn,
  id_var,
  cycle_var,
  baseline_val,
  arm_var = NA,
  auc = "above",
  cycle_limit = NA,
 y_limit = 4,
  tab_ymin = NA,
  tab_ymax = NA,
  round_dec = 2,
  permute_tests = FALSE,
  permute_n = 2000,
  bootstrap_ci = FALSE,
  bootstrap_ci_alpha = 0.05,
 bootstrap_n = 2000,
  arm_colors = NA,
  x_label = NA,
  add_item_title = FALSE,
  cycle_label = FALSE,
  cycle_vals = NA,
  cycle_labs = NA
)
```

#### Arguments

| dsn          | A data.frame object with PRO-CTCAE data   |
|--------------|---|
| id_var       | A character string. Name of ID variable differentiating each unique patient.  |
| cycle_var    | A character string. Name of variable differentiating one longitudinal/repeated. PRO-CTCAE survey from another, within an individual ID. |
| baseline_val | A number indicating the expected baseline cycle/time point.   |

| arm_var                   | A character string. Name of arm variable differentiating treatment groups. Must<br>be character or factor class. Overall frequencies will be reported if no arm/grouping<br>variable is provided. Defaults to NA.   |
|---------------------------|---|
| auc                       | A character string. Specifies the partitioning of area shown. options include:<br>"above" = accumulated area above the baseline horizontal can be interpreted as<br>worsening severity from baseline. "below" = accumulated area below the base-<br>line horizontal can be interpreted the as the amount of decreased severity from<br>baseline. "both" = accumulated area above and below the baseline horizontal<br>are shown. Defaults to "above".   |
| cycle_limit               | A number. Limit the number of cycles to be use to calculate the AUC metrics up<br>to and including a given cycle number. All available cycle time points are used<br>if no cycle number is provided. Defaults to NA.  |
| y_limit                   | A number. Y axis limit for plots. Defaults to 4.  |
| tab_ymin                  | A number. Y axis coordinate for adjusting the vertical placement of the AUC table within the figure. Defaults to NA.  |
| tab_ymax                  | A number. Y axis coordinate for adjusting the vertical placement of the AUC table within the figure. Defaults to NA.  |
| round_dec                 | A number. Number of decimal places to be shown within the AUC table. Defaults to 2.   |
| permute_tests             | Logical. Calls to calculate p values comparing the difference in AUC between<br>two arms using a permutation test. Typical two- sided null hypothesis for a per-<br>mutation test is applied. That is, assigning subjects to groups is interchangeable<br>when calculating AUC. Computation time may be extensive depending on data<br>size, number of PRO-CTCAE items, and number of permutations called. Con-<br>sider staring out an open window or crafting a haiku during this time. Defaults<br>to FALSE. |
| permute_n                 | A number. The number of permutations to be used for permutation tests. De-<br>faults to 2000.   |
| bootstrap_ci              | Logical. Calls to construct alpha-level confidence intervals for the difference in AUC between arms. Similar considerations for computation time as permute_tests are recommended here. Defaults to FALSE.  |
| <pre>bootstrap_ci_a</pre> |   |
|                           | A number. Specifies the alpha level for bootstrap confidence intervals. Must be between 0 and 1. Defaults to 0.05.  |
| bootstrap_n               | A number. The number of bootstrap iterations to be used for bootstrap confidence intervals. Defaults to 2000.   |
| arm_colors                | A column vector of valid colors. Allows the user to define the colors of arms<br>shown in the returned figure. Column vector must have length greater than or<br>equal to the number of arms. Default colors provided.  |
| x_label                   | A character string. Label for the x axis of the plot. Defaults to NA resulting in x axis labeled with cycle_var variable name.  |
| add_item_title            | Logical. Adds the item short label to the title of each figure. Defaults to FALSE.  |
| cycle_label               | Logical. Assign custom labels to cycles/time point. If TRUE, the cycle_vals and cycle_labs must also be specified.  |

# toxFigures

| cycle_vals | Numeric column vector. Vector of values seen within the cycle_var variable.<br>Must be same length of cycle_labs. Defaults to NA.       |
|------------|---|
| cycle_labs | Character column vector. Vector of labels to be mapped to the associated cycle_vals. Must be same length of cycle_vals. Defaults to NA. |

# Examples

```
## Not run:
AUC=toxAUC(dsn = ProAE::tox_acute,
id_var = "id",
cycle_var = "Cycle",
baseline_val = 1)
AUC[[1]]
```

## End(Not run)

| toxFigures | Create PRO-CTCAE severity frequency distribution figures for indi- |
|------------|--|
|            | vidual survey items and composite scores                           |

# Description

Data format should be in 'long' format, where each PRO-CTCAE item is a variable/column.

#### Usage

```
toxFigures(
 dsn,
  id_var,
  cycle_var,
 baseline_val,
 arm_var = NA,
 plot_limit = NA,
  colors = 1,
 bar_label = 0,
  cycle_label = FALSE,
  cycle_vals = NA,
  cycle_labs = NA,
  summary_only = FALSE,
  summary_highlight = FALSE,
  cycles_only = TRUE,
  x_lab_angle = 0,
 x_lab_vjust = 1,
  x_lab_hjust = 0,
 x_label = "Randomized Treatment Assignment",
 y_label = "Percent of Total Frequency",
  footnote_break = FALSE,
```

```
suppress_legend = FALSE,
add_item_title = FALSE
)
```

# Arguments

| dsn                         | A data.frame object with PRO-CTCAE data  |
|-----------------------------|--|
| id_var                      | A character string.Name of ID variable differentiating each PRO-CTCAE survey/participant entered as a quoted string.   |
| cycle_var                   | A character string. Name of variable differentiating one longitudinal/repeated PRO-CTCAE survey from another, within an individual ID.   |
| baseline_val                | A number indicating the expected baseline cycle/time point.  |
| arm_var                     | A character string. Name of arm variable differentiating treatment groups. Must be character or factor class. Overall AUC will be reported if no arm/grouping variable is provided. Defaults to NA.  |
| plot_limit                  | A number. Limit the number of cycles to be plotted up to and including a given cycle number. All available cycle time points are plotted if no cycle number is provided. Defaults to NA.   |
| colors                      | A number. Specify the coloring scheme of symptom scores within frequency bars. Options include: $1 =$ Blue and red color shading, $2 =$ qualitative color shades (color blind friendly), $3 =$ black and white. Defaults to 1.   |
| bar_label                   | A number. Label frequency bars with sample size (n) or percent shown on the y-axis. Label options include: $1 =$ sample size (n) within each cycle (symptom score 0 or higher), $2 =$ sample size (n) within each cycle with present symptoms (symptom score > 0), $3 =$ sample size (n) within each cycle with severe symptoms (symptom score >= 3), $4 =$ percent of subjects within each cycle with present symptoms (symptom score > 0), $5 =$ percent of subjects within each cycle with not specified. Defaults to NA. |
| cycle_label                 | Logical. Assign custom labels to cycles/time point. If TRUE, the cycle_vals and cycle_labs must also be specified.   |
| cycle_vals                  | Numeric column vector. Vector of values seen within the cycle_var variable.<br>Must be same length of cycle_labs. Defaults to NA.  |
| cycle_labs                  | Character column vector. Vector of labels to be mapped to the associated cycle_vals.<br>Must be same length of cycle_vals. Defaults to NA.   |
| summary_only                | Logical. Only display the summary measures in figures / Suppress the individual time points from plotting. Defaults to FALSE.  |
| <pre>summary_highligh</pre> | -  |
|                             | Logical. Add black box around summary measure bar chart. Defaults to FALSE.  |
| cycles_only                 | Logical. Only display the longitudinal time points in figures / Suppress the summary measures from plotting. Defaults to TRUE.   |
| x_lab_angle                 | A integer between 0 and 360. Allows the user to rotate the x axis labels in order to fit long arm names (0 or 45 recommended). Defaults to 0.  |
| x_lab_vjust                 | A number. A ggplot2 object option. Allows the user to vertically adjusts the x axis labels in order to fit arm names. Defaults to 1.   |

#### toxScores

| x_lab_hjust   | A number. A ggplot2 object option. Allows the user to horizontally adjusts the x axis labels in order to fit arm names. Defaults to $0$ .                                       |  |  |  |
|---|---|--|--|--|
| x_label   | A character string. Label for the x axis of the plot. Defaults to "Randomized Treatment Assignment" if arm_var is specified, defaults to "Overall" if not arm_var is specified. |  |  |  |
| y_label   | A character string. Label for the y axis of the plot. Defaults to "Percent of Total Frequency".   |  |  |  |
| footnote_break Logical. Add a line break to the footnote Defaults to FALSE. |   |  |  |  |
|   | Logical. Suppress the legend from appearing in figure. Defaults to FALSE.   |  |  |  |
| add_item_title  | Logical. Add the symptom item name as a title to the figure. Defaults to FALSE.   |  |  |  |

#### Value

A list object. The returned object is a  $(k \times 2)$  or  $(k \times 3)$  nested list. Where k is the number of PRO-CTCAE item groups (e.g. pain, fatigue, nausea); list[[1 ... i ... k]]. For each list item there are 2 or 3 elements. The 1st element of each list item is the name of the PRO-CTCAE item group returned as a string. The 2nd element is the PRO-CTCAE figure as a ggplot object. These objects can be modified as such.

#### Examples

```
## Not run:
fig_acute = toxFigures(dsn = ProAE::tox_acute,
    cycle_var = "Cycle",
    baseline_val = 1,
    arm_var = "arm",
    id_var = "id",
    x_lab_angle = -45,
    x_lab_vjust = .3,
    x_lab_hjust = .2,
    colors = 2)
fig_acute[[1]]
```

toxScores

## End(Not run)

*Re-code PRO-CTCAE text responses, apply a zero-imputation procedures, and construct PRO-CTCAE composite scores.* 

#### Description

This function takes in a data frame set with PRO-CTCAE survey text fields/responses and returns a data frame with appropriate numerical re-coding. This function will accept 1 or up to all 124 PRO-CTCAE survey fields. All PRO-CTCAE variable names MUST conform to a pre-specified naming structure. PRO-CTCAE variable names are made up of FOUR components: 1)'PROCTCAE', 2) number [1,2,3, ..., i, ..., 80], 3) 'A', 'B', or 'C' component of the i-th PRO-CTCAE field, 4) and 'SCL' (if severity, interference, or frequency) or 'IND' (if yes/no variable). Each component must be delimited by an underscore (\_)

#### Usage

```
toxScores(
   dsn,
   reformat = FALSE,
   impute = FALSE,
   composites = FALSE,
   short_labels = FALSE
)
```

## Arguments

| dsn          | A data.frame object with PRO-CTCAE data  |
|--------------|--|
| reformat     | Reformat PRO-CTCAE text responses to numeric scores. Defaults to FALSE.                      |
| impute       | Apply zero-imputation where appropriate. Defaults to FALSE.                                  |
| composites   | Construct composite score using available PRO-CTCAE variables within dsn. Defaults to FALSE. |
| short_labels | Add PRO-CTCAE short labels to available PRO-CTCAE variables within re-<br>turned object      |

#### Details

- [EX1] Question 1 of PRO-CTCAE should be: PROCTCAE\_1A\_SCL
- [EX2] Question 48 of PRO-CTCAE should be: PROCTCAE\_48A\_SCL, PROCTCAE\_48B\_SCL, PROCTCAE\_48C\_SCL
- [EX3]Question 73 of PRO-CTCAE should be: PROCTCAE\_73A\_IND

This function also constructs PRO-CTCAE composite scores. Composite score variables for respective PRO-CTCAE item groups are created and named as PROCTCAE\_##\_COMP.

- 1. https://healthcaredelivery.cancer.gov/pro-ctcae/pro-ctcae\_english.pdf
- 2. Ethan Basch, et al. Development of a Composite Scoring Algorithm for the National Cancer Institute's Patient-Reported Outcomes version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE). ISOQOL 2019
- Basch E, et al. Composite Grading Algorithm for the National Cancer Institute's Patient-Reported Outcomes version of the Common Terminology Criteria for Adverse Events (PRO-CTCAE). Clinical Trials 2020.

Data format should be in 'long' format, where each PRO-CTCAE item is a variable/column.

#### Value

A data.frame object.

# Examples

```
tox_acute_comp = toxScores(dsn = ProAE::tox_acute, composites = TRUE)
```

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toxSummary

# Description

Data format should be in 'long' format, where each PRO-CTCAE item is a variable/column.

#### Usage

```
toxSummary(
   dsn,
   id_var,
   cycle_var,
   summary_measure,
   baseline_val = NA,
   arm_var = NA
)
```

# Arguments

| dsn             | A data.frame object with PRO-CTCAE data.   |
|-----------------|--|
| id_var          | A character string. Name of ID variable differentiating each PRO-CTCAE survey/participant entered as a quoted string.  |
| cycle_var       | A character string. Name of variable differentiating one longitudinal/repeated. PRO-CTCAE survey from another, within an individual ID.  |
| summary_measure |  |
|                 | A character string. Type of summary statistic to be used. Please consult current literature for appropriate interpretations of the summary measure selected and suitable analysis procedures for comparing groups. Options include: "max" = Use subjects' maximum score. "max_post_bl" = Use subjects' maximum score post-baseline visit. "bl_adjusted" = Use subjects' baseline adjusted score over the study period. The baseline adjusted score is derived by the following: If the maximum score post-baseline is more severe than the baseline score, then the use maximum score post-baseline is used as the adjusted score. Otherwise, if the maximum score post-baseline is the same or less serve than the baseline score, then zero (0) is used as the adjusted score. "toxicity_index" = Construct patient-level toxicity index. "AUC_worsening" = Calculate group-level AUC describing |
| baseline_val    | A number indicating the expected baseline cycle/time point.  |
| arm_var         | A character string. Name of arm variable differentiating treatment arms or other grouping factor. Required for group-level summary measures.   |

# Value

A data.frame with only the id and PRO-CTCAE variables being summarized. Each subject will now only have 1 observation (PRO-CTCAE variables are now the summary measure value).

# Examples

```
toxSummary(dsn=ProAE::tox_acute,
id_var="id",
cycle_var="Cycle",
baseline_val=1,
summary_measure = "max")
```

| toxT | ables |
|------|-------|
| LOXI | antes |

*Create toxicity tables for individual and composite PRO-CTCAE survey items.* 

# Description

Data format should be in 'long' format, where each PRO-CTCAE item is a variable/column.

#### Usage

```
toxTables(
   dsn,
   id_var,
   cycle_var,
   baseline_val,
   type = "bl_adjusted",
   test = "c",
   riskdiff = FALSE,
   risk_ci = "wald",
   risk_ci_alpha = 0.05,
   arm_var = NA,
   cycle_limit = NA
)
```

# Arguments

| dsn          | A data.frame object with PRO-CTCAE data.   |
|--------------|--|
| id_var       | A character string. Name of ID variable differentiating each PRO-CTCAE survey/participant entered as a quoted string.  |
| cycle_var    | A character string. Name of variable differentiating one longitudinal/repeated. PRO-CTCAE survey from another, within an individual ID.  |
| baseline_val | A number indicating the expected baseline cycle/time point.  |
| type         | A character string. Type of summary measure to be be used. Options in-<br>clude: "max_post_bl" = Use subjects' maximum score post-baseline visit.<br>"bl_adjusted" = Use subjects' baseline adjusted score over the study period.<br>"max" = Use subjects' maximum score over the study period. The baseline ad-<br>justed score is derived by the following: If the maximum score post-baseline is<br>more severe than the baseline score, then the use maximum score post-baseline<br>is used as the adjusted score. Otherwise, if the maximum score post-baseline |

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|               | is the same or less serve than the baseline score, then zero (0) is used as the adjusted score. Defaults to "bl_adjusted".   |
|---------------|--|
| test          | A character string. Specify the statistical test to apply where comparing rates among arms. Options include: " $c$ " = chi square, "f" = fisher's exact. Defaults to "c".                                    |
| riskdiff      | Logical. Calculates risk differences between two arms. Valid if there are only two arms in the data.frame specified. This option will countermand options called with the test parameter. Defaults to FALSE. |
| risk_ci       | A character string. Specify the confidence interval type to be constructed for risk differences. Options include: "wald", and "agresti-caffo". Defaults to "wald".   |
| risk_ci_alpha | A number between 0 and 1. Specify the alpha level of the risk difference confidence intervals. Defaults to 0.05.   |
| arm_var       | A character string. Name of arm variable differentiating treatment groups. Must be character or factor class. Overall frequencies will be reported if no arm/grouping variable is provided. Defaults to NA.  |
| cycle_limit   | A number. Limit the data to be analyzed up to and including a given cycle number or time point. Defaults to NA.  |

# Value

A list object with data.frame elements for individual items and composite scores.

#### Examples

toxTables(dsn=ProAE::tox\_acute, id\_var="id", cycle\_var="Cycle", baseline\_val=1)

| tox_acute |
|-----------|
|-----------|

PRO-CTCAE data reflecting acute drug toxicity

#### Description

Simulated example data where the drug group experiences acute toxicity followed by symptom abatement over the course of treatment.

# Format

A data frame with 1400 rows and 5 variables

#### Details

- id. Subject identifier
- Cycle. Time variable denoting visits/cycles (1-10)
- arm. Treatment groups include drug and placebo
- PROCTCAE\_9A\_SCL. PRO-CTCAE frequency item for nausea
- PROCTCAE\_9B\_SCL. PRO-CTCAE severity item for nausea
- PROCTCAE\_9\_COMP. PRO-CTCAE composite score for nausea

#### Author(s)

Blake Langlais

tox\_chronic

#### PRO-CTCAE data reflecting chronic drug toxicity

# Description

Simulated example data where the drug group experiences chronic toxicity over the course of treatment.

#### Format

A data frame with 1400 rows and 5 variables

#### Details

- id. Subject identifier
- Cycle. Time variable denoting visits/cycles (1-10)
- arm. Treatment groups include drug and placebo
- PROCTCAE\_9A\_SCL. PRO-CTCAE frequency item for nausea
- PROCTCAE\_9B\_SCL. PRO-CTCAE severity item for nausea
- PROCTCAE\_9\_COMP. PRO-CTCAE composite score for nausea

#### Author(s)

Blake Langlais

tox\_cumulative PRO-CTCAE data reflecting cumulative drug toxicity

# Description

Simulated example data where drug toxicity is cumulative over the course of treatment.

#### Format

A data frame with 1400 rows and 5 variables

# tox\_cyclic

# Details

- id. Subject identifier
- Cycle. Time variable denoting visits/cycles (1-10)
- arm. Treatment groups include drug and placebo
- PROCTCAE\_9A\_SCL. PRO-CTCAE frequency item for nausea
- PROCTCAE\_9B\_SCL. PRO-CTCAE severity item for nausea
- PROCTCAE\_9\_COMP. PRO-CTCAE composite score for nausea

#### Author(s)

Blake Langlais

tox\_cyclic

#### PRO-CTCAE data reflecting cyclical drug toxicity

# Description

Simulated example data where the drug group experiences cyclically toxicity post-treatment administration

#### Format

A data frame with 1400 rows and 5 variables

# Details

- id. Subject identifier
- Cycle. Time variable denoting visits/cycles (1-10)
- arm. Treatment groups include drug and placebo
- PROCTCAE\_9A\_SCL. PRO-CTCAE frequency item for nausea
- PROCTCAE\_9B\_SCL. PRO-CTCAE severity item for nausea
- PROCTCAE\_9\_COMP. PRO-CTCAE composite score for nausea

#### Author(s)

Blake Langlais

tox\_late

# Description

Simulated example data where the drug group experiences late incipient toxicity towards the end of the treatment period.

#### Format

A data frame with 1400 rows and 5 variables

#### Details

- id. Subject identifier
- Cycle. Time variable denoting visits/cycles (1-10)
- arm. Treatment groups include drug and placebo
- PROCTCAE\_9A\_SCL. PRO-CTCAE frequency item for nausea
- PROCTCAE\_9B\_SCL. PRO-CTCAE severity item for nausea
- PROCTCAE\_9\_COMP. PRO-CTCAE composite score for nausea

# Author(s)

Blake Langlais

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