# Package 'pvda'

July 23, 2025

```
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      ing data to the da() function, which executes the disproportionality analy-
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```

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add\_disproportionality

Add disproportionality estimates to data frame with expected counts

#### **Description**

Add disproportionality estimates to data frame with expected counts

### Usage

```
add_disproportionality(
  df = NULL,
  df_syms = NULL,
  da_estimators = c("ic", "prr", "ror"),
  rule_of_N = 3,
  conf_lvl = 0.95
)
```

add\_expected\_counts 3

#### **Arguments**

df Intended use is on the output tibble from add\_expected\_counts.

df\_syms A list built from df\_colnames through conversion to symbols.

da\_estimators Character vector specifying which disproportionality estimators to use, in case you don't need all implemented options. Defaults to c("ic", "prr", "ror").

rule\_of\_N Numeric value. Sets estimates for ROR and PRR to NA when observed counts are strictly less than the passed value of rule\_of\_N. Default value is 3, 5 is sometimes used as a more liberal alternative. Set to NULL if you don't want to apply any such rule.

conf\_lvl Confidence level of confidence or credibility intervals. Default is 0.95 (i.e. 95)

% confidence interval).

#### Value

The passed data frame with disproportionality point and interval estimates.

add expected counts Produces expected counts

# **Description**

Produces various counts used in disproportionality analysis.

### Usage

```
add_expected_counts(
    df = NULL,
    df_colnames = NULL,
    df_syms = NULL,
    expected_count_estimators = c("rrr", "prr", "ror")
)
```

### **Arguments**

df An object possible to convert to a data table, e.g. a tibble or data.frame, contain-

ing patient level reported drug-event-pairs. See header 'The df object' below for

further details.

df\_colnames A list of column names to use in df. That is, point da to the 'report id'-column

(report\_id), the 'drug name'-column (drug), the 'adverse event'-column (event) and optionally a grouping column group\_by to calculate disproportionality across.

See the vignette for further details.

df\_syms A list built from df\_colnames through conversion to symbols.

expected\_count\_estimators

A character vector containing the desired expected count estimators. Defaults to

c("rrr", "prr", "ror").

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#### Value

A tibble containing the various counts.

### The df object

The passed df should be (convertible to) a data table and at least contain three columns: report\_id, drug and event. The data table should contain one row per reported drug-event-combination, i.e. receiving a single additional report for drug X and event Y would add one row to the table. If the single report contained drug X for event Y and event Z, two rows would be added, with the same report\_id and drug on both rows. Column report\_id must be of type numeric or character. Columns drug and event must be of type character. If column group\_by is provided, it can be either numeric or character. You can use a df with column names of your choosing, as long as you connect role and name in the df\_colnames-parameter.

```
apply_rule_of_N
```

apply\_rule\_of\_N

#### **Description**

Internal function to set disproportionality cells for ROR and PRR to NA when observed count < 3

### Usage

```
apply_rule_of_N(
  da_df = NULL,
  da_estimators = c("ic", "prr", "ror"),
  rule_of_N = NULL
)
```

# **Arguments**

da\_df See the intermediate object da\_df in add\_disproportionality

da\_estimators Default is c("ic", "prr", "ror").

rule\_of\_N An length one integer between 0 and 10.

### **Details**

Sometimes, you want to protect yourself from spurious findings based on small observed counts combined with infinitesimal expected counts.

### Value

The input data frame (da\_df) with potentially some cells set to NA.

build\_colnames\_da 5

build_colnames_da	An internal function creating colnames for da confidence/credibility bounds
-------------------	---

# **Description**

Given the output from quantile\_prob, and a da\_name string, create column names such as PRR025, ROR025 and IC025

# Usage

```
build_colnames_da(
   quantile_prob = list(lower = 0.025, upper = 0.975),
   da_name = NULL
)
```

# **Arguments**

```
quantile_prob A list with two parameters, lower and upper. Default: list(lower = 0.025, upper = 0.975)
da_name A string, such as "ic", "prr" or "ror". Default: NULL
```

# Value

A list with two symbols, to be inserted in the dtplyr-chain

ci\_for\_ic Confidence intervals for Information Component (IC)

# Description

Mainly used in function ic. Produces quantiles of the posterior gamma distribution. Called twice in ic to create credibility intervals.

# Usage

```
ci_for_ic(obs, exp, conf_lvl_probs, shrinkage)
```

# Arguments

obs	A numeric vector with observed counts, i.e. number of reports for the selected drug-event-combination. Note that shrinkage (e.g. +0.5) is added inside the function and should not be included here.
exp	A numeric vector with expected counts, i.e. number of reports to be expected given a comparator or <i>background</i> . Note that shrinkage (e.g. +0.5) is added inside the function and should not be included here.

ci\_for\_prr

 ${\tt conf\_lvl\_probs} \quad The \ probabilities \ of \ the \ posterior, \ based \ on \ a \ passed \ confidence \ level \ ({\tt conf\_lvl})$ 

in ic. For instance, if sgn\_lvl = .95 in ic is used, quantiles will be extracted

at sgn\_lvl\_probs 0.025 and 0.975.

shrinkage A non-negative numeric value, to be added to observed and expected count.

Default is 0.5.

#### Value

The credibility interval specified by input parameters.

#### See Also

ic

ci\_for\_prr

Confidence intervals for Proportional Reporting Rate

### **Description**

Mainly for use in prr. Produces (symmetric, normality based) confidence bounds for the PRR, for a passed probability. Called twice in prr to create confidence intervals.

### Usage

```
ci_for_prr(
  obs = NULL,
  n_drug = NULL,
  n_event_prr = NULL,
  n_tot_prr = NULL,
  conf_lvl_probs = 0.95
)
```

### **Arguments**

obs Number of reports for the specific drug and event (i.e. the observed count).

n\_drug Number of reports with the drug of interest.

n\_event\_prr Number of reports with the event in the background.

n\_tot\_prr Number of reports in the background.

conf\_lvl\_probs The probabilities of the normal distribution, based on a passed confidence level

(conf\_lvl) in prr. If sgn\_lvl = .95 in prr, quantiles of the normal distribution

will be extracted at sgn\_lvl\_probs of 0.025 and 0.975.

### Value

The confidence interval specified by input parameters.

ci\_for\_ror 7

### See Also

prr

ci\_for\_ror

Confidence intervals for Reporting Odds Ratio

# Description

Mainly for use in ror. Produces (symmetric, normality based) confidence bounds for the ROR, for a passed probability. Called twice in ror to create confidence intervals.

# Usage

```
ci_for_ror(a, b, c, d, conf_lvl_probs)
```

# Arguments

a	Number of reports for the specific drug and event (i.e. the observed count).

b Number of reports with the drug, without the event

c Number of reports without the drug, with the event

d Number of reports without the drug, without the event

conf\_lvl\_probs The probabilities of the normal distribution, based on a passed confidence level (conf\_lvl) in ror. If sgn\_lvl = .95 in ror, quantiles of the normal distribution will be extracted at sgn\_lvl\_probs of 0.025 and 0.975.

# Value

The credibility interval specified by input parameters.

# See Also

ror

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```
conf_lvl_to_quantile_prob
```

Quantile probabilities from confidence level

# **Description**

Calculates equi-tailed quantile probabilities from a confidence level

### Usage

```
conf_lvl_to_quantile_prob(conf_lvl = 0.95)
```

# **Arguments**

conf\_lvl

Confidence level of confidence or credibility intervals. Default is 0.95 (i.e. 95 % confidence interval).

### Value

A list with two numerical vectors, "lower" and "upper".

# **Examples**

```
conf_lvl_to_quantile_prob(0.95)
```

count\_expected\_prr

Count expected for Proportional Reporting Rate

### **Description**

Internal function to provide expected counts related to the PRR

# Usage

```
count_expected_prr(count_dt)
```

# **Arguments**

count\_dt

A data table, output from count\_expected\_rrr

### Value

A data table with added columns for n\_event\_prr n\_tot\_prr and expected\_prr @export

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count\_expected\_ror

Count expected for Reporting Odds Ratio

### **Description**

Internal function to provide expected counts related to the ROR

# Usage

```
count_expected_ror(count_dt)
```

# **Arguments**

count\_dt

A data table, output from count\_expected\_rrr

# **Details**

**DETAILS** 

#### Value

A data table with added columns for n\_event\_prr, n\_tot\_prr and expected\_prr OUTPUT\_DESCRIPTION

#### See Also

mutate, select everything

count\_expected\_rrr

Count Expected for Relative Reporting Rate

# Description

Internal function to provide expected counts related to the RRR

# Usage

```
count_expected_rrr(df, df_colnames, df_syms)
```

# **Arguments**

df See documentation for add\_expected\_counts

df\_colnames See documentation for da

df\_syms A list built from df\_colnames through conversion to symbols.

### Value

A data frame with columns for obs, n\_drug, n\_event, n\_tot and (RRR) expected

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Disproportionality Analysis

Description

The function da executes disproportionality analyses, i.e. compares the proportion of reports with a specific adverse event for a drug, against an event proportion from a comparator based on the passed data frame. See the vignette for a brief introduction to disproportionality analysis. Furthermore, da supports three estimators: Information Component (IC), Proportional Reporting Rate (PRR) and the Reporting Odds Ratio (ROR).

### Usage

```
da(
    df = NULL,
    df_colnames = list(report_id = "report_id", drug = "drug", event = "event", group_by =
        NULL),
    da_estimators = c("ic", "prr", "ror"),
    sort_by = "ic",
    number_of_digits = 2,
    rule_of_N = 3,
    conf_lvl = 0.95,
    excel_path = NULL
)
```

#### **Arguments**

df

An object possible to convert to a data table, e.g. a tibble or data.frame, containing patient level reported drug-event-pairs. See header 'The df object' below for further details.

df\_colnames

A list of column names to use in df. That is, point da to the 'report id'-column (report\_id), the 'drug name'-column (drug), the 'adverse event'-column (event) and optionally a grouping column group\_by to calculate disproportionality across. See the vignette for further details.

da\_estimators

Character vector specifying which disproportionality estimators to use, in case you don't need all implemented options. Defaults to c("ic", "prr", "ror").

sort\_by

The output is sorted in descending order of the lower bound of the confidence/credibility interval for a passed da estimator. Any of the passed strings in "da\_estimators" is accepted, the default is "ic". If a grouping variable is passed, sorting is made by the sample average across each drug-event-combination (ignoring NAs).

number\_of\_digits

Round decimal columns to specified precision, default is two decimals.

rule\_of\_N

Numeric value. Sets estimates for ROR and PRR to NA when observed counts are strictly less than the passed value of rule\_of\_N. Default value is 3, 5 is sometimes used as a more liberal alternative. Set to NULL if you don't want to apply any such rule.

In

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conf\_lvl Confidence level of confidence or credibility intervals. Default is 0.95 (i.e. 95 % confidence interval).

excel\_path

Intended for users who prefer to work in excel with minimal work in R. To write the output of da to an excel file, provide a path to a folder. For instance, to write to your current working directory, pass getwd(). The excel file will by default be named da.xlsx. To control the excel file name, pass a path ending with the desired filename suffixed with .xlsx. If you do not want to export the output to an excel file, pass NULL (the default).

### Value

da returns a data frame (invisibly) containing counts and estimates related to supported disproportionality estimators. Each row corresponds to a drug-event pair.

# The df object

The passed df should be (convertible to) a data table and at least contain three columns: report\_id, drug and event. The data table should contain one row per reported drug-event-combination, i.e. receiving a single additional report for drug X and event Y would add one row to the table. If the single report contained drug X for event Y and event Z, two rows would be added, with the same report\_id and drug on both rows. Column report\_id must be of type numeric or character. Columns drug and event must be of type character. If column group\_by is provided, it can be either numeric or character. You can use a df with column names of your choosing, as long as you connect role and name in the df\_colnames-parameter.

# **Examples**

```
### Run a disproportionality analysis
da_1 <-
 tiny_dataset |>
 da()
### Run a disproportionality across subgroups
list_of_colnames <-</pre>
 list(
    report_id = "report_id",
    drug = "drug",
    event = "event",
    group_by = "group"
 )
 da_2 <-
 tiny_dataset |>
 da(df_colnames = list_of_colnames)
# If columns in your df have different names than the default ones,
# you can specify the column names in the df_colnames parameter list:
renamed_df <-
 tiny_dataset |>
```

drug\_event\_df

```
dplyr::rename(ReportID = report_id)
list_of_colnames$report_id <- "ReportID"
da_3 <-
    renamed_df |>
    da(df_colnames = list_of_colnames)
```

drug\_event\_df

A simulated ICSR database

# Description

drug\_event\_df is a simulated dataset, slightly larger than the "tiny\_dataset" which is also contained in this package.

### Usage

```
drug_event_df
```

#### **Format**

'drug\_event\_df' A data frame with 3,971 rows and 3 columns. In total 1000 unique report\_ids, i.e. the same report\_id can have several drugs and events.

Number of drugs per report\_id is sampled as 1 + Pois(3), with increasing probability as the drug letter closes in on Z. Every drug is assigned an event, with decreasing probability as the event index number increases towards 1000. See the DATASET.R file in the data-raw folder for details.

```
report_id A patient or report identifierdrug One of 26 fake drugs (Drug_A - Drug_Z)event Sampled events (Event_1 - Event_1000)
```

### **Source**

Simulated data.

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grouped\_da

Disproportionality Analysis by Subgroups

# Description

A package internal wrapper for executing da across subgroups

# Usage

```
grouped_da(
   df = NULL,
   df_colnames = NULL,
   df_syms = NULL,
   expected_count_estimators = NULL,
   da_estimators = NULL,
   sort_by = NULL,
   conf_lvl = NULL,
   rule_of_N = NULL,
   number_of_digits = NULL
)
```

# **Arguments**

df See the da function df\_colnames See the da function df\_syms A list built from df\_colnames through conversion to symbols. expected\_count\_estimators See the da function da\_estimators See the da function sort\_by See the da function conf\_lvl See the da function rule\_of\_N See the da function number\_of\_digits See the da function

### **Details**

See the da documentation

# Value

See the da function

ic ic

ic Information component

# Description

Calculates the information component ("IC") and credibility interval, used in disproportionality analysis.

# Usage

ic(obs = NULL, exp = NULL, shrinkage = 0.5, conf\_lvl = 0.95)

# **Arguments**

obs	A numeric vector with observed counts, i.e. number of reports for the selected drug-event-combination. Note that shrinkage (e.g. +0.5) is added inside the
	function and should not be included here.
exp	A numeric vector with expected counts, i.e. number of reports to be expected given a comparator or <i>background</i> . Note that shrinkage (e.g. +0.5) is added inside the function and should not be included here.
shrinkage	A non-negative numeric value, to be added to observed and expected count. Default is 0.5.
conf_lvl	Confidence level of confidence or credibility intervals. Default is 0.95 (i.e. 95 % confidence interval).

### **Details**

The IC is a log2-transformed observed-to-expected ratio, based on the relative reporting rate (RRR) for counts, but modified with an addition of "shrinkage" to protect against spurious associations.

$$\hat{IC} = log_2(\frac{\hat{O} + k}{\hat{E} + k})$$

where  $\hat{O}$  = observed number of reports, k is the shrinkage (typically +0.5), and expected  $\hat{E}$  is (for RRR, and using the entire database as comparator or *background*) estimated as

$$\hat{E} = \frac{\hat{N}_{drug} \times \hat{N}_{event}}{\hat{N}_{TOT}}$$

where  $\hat{N}_{drug}$ ,  $\hat{N}_{event}$  and  $\hat{N}_{TOT}$  are the number of reports with the drug, the event, and in the whole database respectively.

The credibility interval is created from the quantiles of the posterior gamma distribution with shape  $(\hat{S})$  and rate  $(\hat{R})$  parameters as

$$\hat{S} = \hat{O} + k$$

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$$\hat{R} = \hat{E} + k$$

using the stats::qgamma function. Parameter k is the shrinkage defined earlier. For completeness, a credibility interval of the gamma distributed X (i.e.  $X \sim \Gamma(\hat{S}, \hat{R})$  where  $\hat{S}$  and  $\hat{R}$  are shape and rate parameters) with associated quantile function  $Q_X(p)$  for a significance level  $\alpha$  is constructed as

$$[Q_X(\alpha/2), Q_X(1-\alpha/2)]$$

#### Value

A tibble with three columns (point estimate and credibility bounds).

#### **Further details**

From a bayesian point-of-view, the credibility interval of the IC is constructed from the poisson-gamma conjugacy. The shrinkage constitutes a prior of observed and expected of 0.5. A shrinkage of +0.5 with a gamma-quantile based 95 % credibility interval cannot have lower bound above 0 unless the observed count exceeds 3. One benefit of  $log_2$  is to provide a log-scale for convenient plotting of multiple IC values side-by-side.

#### References

Norén GN, Hopstadius J, Bate A (2011). "Shrinkage observed-to-expected ratios for robust and transparent large-scale pattern discovery." *Statistical Methods in Medical Research*, **22**(1), 57–69. doi:10.1177/0962280211403604, https://doi.org/10.1177/0962280211403604.

### **Examples**

```
ic(obs = 20, exp = 10) # Note that obs and exp can be vectors (of equal length, no recycling allowed) ic(obs = c(20, 30), exp = c(10, 10))
```

print.da

print function for da objects

# Description

print function for da objects

# Usage

```
## S3 method for class 'da'
print(x, n = 10, ...)
```

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# Arguments

```
    A S3 obj of class "da", output from pvda::da().
    Control the number of rows to print.
    For passing additional parameters to extended classes.
```

#### Value

Nothing, but prints the tibble da\_df in the da object.

### **Examples**

```
da_1 <-
tiny_dataset |>
da()
print(da_1)
```

prr

Proportional Reporting Rate

# **Description**

Calculates Proportional Reporting Rate ("PRR") with confidence intervals, used in disproportionality analysis.

# Usage

```
prr(
  obs = NULL,
  n_drug = NULL,
  n_event_prr = NULL,
  n_tot_prr = NULL,
  conf_lvl = 0.95
)
```

# Arguments

Number of reports for the specific drug and event (i.e. the observed count).

Number of reports with the drug of interest.

Number of reports with the event in the background.

Number of reports in the background.

Confidence level of confidence or credibility intervals. Default is 0.95 (i.e. 95 % confidence interval).

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#### **Details**

The PRR is the proportion of reports with an event in set of exposed cases, divided with the proportion of reports with the event in a background or comparator, which does not include the exposed.

The PRR is estimated from a observed-to-expected ratio, based on similar to the RRR and IC, but excludes the exposure of interest from the comparator.

$$P\hat{R}R = \frac{\hat{O}}{\hat{E}}$$

where  $\hat{O}$  is the observed number of reports, and expected  $\hat{E}$  is estimated as

$$\hat{E} = \frac{\hat{N}_{drug} \times (\hat{N}_{event} - \hat{O})}{\hat{N}_{TOT} - \hat{N}_{drug}}$$

where  $\hat{N}_{drug}$ ,  $\hat{N}_{event}$ ,  $\hat{O}$  and  $\hat{N}_{TOT}$  are the number of reports with the drug, the event, the drug and event, and in the whole database respectively.

A confidence interval is derived in Gravel (2009) using the delta method:

$$\hat{s} = \sqrt{1/\hat{O} - 1/(\hat{N}_{drug}) + 1/(\hat{N}_{event} - \hat{O}) - 1/(\hat{N}_{TOT} - \hat{N}_{drug})}$$

and

$$\begin{split} [\hat{C}I_{\alpha/2},\hat{C}I_{1-\alpha/2}] = \\ [\frac{\hat{O}}{\hat{E}} \times \exp(Q_{\alpha/2} \times \hat{s}), \frac{\hat{O}}{\hat{E}} \times \exp(Q_{1-\alpha/2} \times \hat{s})] \end{split}$$

where  $Q_{\alpha}$  denotes the quantile function of a standard Normal distribution at significance level  $\alpha$ .

Note: For historical reasons, another version of this standard deviation is sometimes used where the last fraction under the square root is added rather than subtracted, with negligible practical implications in large databases. This function uses the version declared above, i.e. with subtraction.

### Value

A tibble with three columns (point estimate and credibility bounds). Number of rows equals length of inputs obs, n\_drug, n\_event\_prr and n\_tot\_prr.

#### References

Montastruc J, Sommet A, Bagheri H, Lapeyre-Mestre M (2011). "Benefits and strengths of the disproportionality analysis for identification of adverse drug reactions in a pharmacovigilance database." *British Journal of Clinical Pharmacology*, **72**(6), 905–908. doi:10.1111/j.13652125.2011.04037.x, https://doi.org/10.1111/j.1365-2125.2011.04037.x.

Gravel C (2009). "Statistical Methods for Signal Detection in Pharmacovigilance." https://repository.library.carleton.ca/downloads/jd472x08w.

18 ror

### **Examples**

```
prr(
   obs = 5,
   n_drug = 10,
   n_event_prr = 20,
   n_tot_prr = 10000
)

# Note that input parameters can be vectors (of equal length, no recycling)
pvda::prr(
   obs = c(5, 10),
   n_drug = c(10, 20),
   n_event_prr = c(15, 30),
   n_tot_prr = c(10000, 10000)
)
```

ror

Reporting Odds Ratio

# **Description**

Calculates Reporting Odds Ratio ("ROR") and confidence intervals, used in disproportionality analysis.

### Usage

```
ror(a = NULL, b = NULL, c = NULL, d = NULL, conf_lvl = 0.95)
```

# **Arguments**

а	Number of reports for the specific drug and event (i.e. the observed count).
b	Number of reports with the drug, without the event
С	Number of reports without the drug, with the event
d	Number of reports without the drug, without the event
conf_lvl	Confidence level of confidence or credibility intervals. Default is 0.95 (i.e. 95 % confidence interval).

# **Details**

The ROR is an odds ratio calculated from reporting counts. The R for Reporting in ROR is meant to emphasize an interpretation of reporting, as the ROR is calculated from a reporting database. Note: the function is vectorized, i.e. a, b, c and d can be vectors, see the examples.

A reporting odds ratio is simply an odds ratio based on adverse event reports.

$$R\hat{O}R = \frac{a/b}{c/d}$$

where a = observed count (i.e. number of reports with exposure and outcome), b = number of reports with the drug and without the event, c = number of reports without the drug with the event and d = number of reports with neither of the drug and the event.

A confidence interval for the ROR can be derived through the delta method, with a standard deviation:

$$\hat{s} = \sqrt{1/a + 1/b + 1/c + 1/d}$$

with the resulting confidence interval for significance level  $\alpha$ 

$$[R\hat{O}R \times exp(\Phi_{\alpha/2} \times \hat{s}), R\hat{O}R \times exp(\Phi_{1-\alpha/2} \times \hat{s})]$$

### Value

A tibble with three columns (point estimate and credibility bounds). Number of rows equals length of inputs a, b, c, d.

#### References

Montastruc J, Sommet A, Bagheri H, Lapeyre-Mestre M (2011). "Benefits and strengths of the disproportionality analysis for identification of adverse drug reactions in a pharmacovigilance database." *British Journal of Clinical Pharmacology*, **72**(6), 905–908. doi:10.1111/j.13652125.2011.04037.x, https://doi.org/10.1111/j.1365-2125.2011.04037.x.

#### **Examples**

```
ror(
    a = 5,
    b = 10,
    c = 20,
    d = 10000
)

# Note that a, b, c and d can be vectors (of equal length, no recycling)
pvda::ror(
    a = c(5, 10),
    b = c(10, 20),
    c = c(15, 30),
    d = c(10000, 10000)
)
```

round\_and\_sort\_by\_lower\_da\_limit

Sort a disproportionality analysis by the lower da conf. or cred. limit

# Description

Sorts the output by the mean lower limit of a passed da estimator

### Usage

```
round_and_sort_by_lower_da_limit(
  df = NULL,
  df_colnames = NULL,
  df_syms = NULL,
  conf_lvl = NULL,
  sort_by = NULL,
  da_estimators = NULL,
  number_of_digits = 2
)
```

### **Arguments**

```
df See add_disproportionality
df_colnames See add_disproportionality
df_syms See add_disproportionality
conf_lvl See add_disproportionality
sort_by See add_disproportionality
da_estimators See add_disproportionality
number_of_digits
```

Numeric value. Set the number of digits to show in output by passing an integer. Default value is 2 digits. Set to NULL to avoid rounding.

### Value

The df object, sorted.

```
{\tt round\_columns\_with\_many\_decimals}
```

Rounds columns in da\_df with many decimals

# **Description**

Internal function containing a mutate + across

# Usage

```
round_columns_with_many_decimals(
  da_df = NULL,
  da_estimators = NULL,
  number_of_digits = NULL
)
```

summary.da 21

# **Arguments**

```
da_df See add_disproportionality
da_estimators See add_disproportionality
number_of_digits
See add_disproportionality
```

### Value

A df with rounded columns

summary.da

Summary function for disproportionality objects

### **Description**

Provides summary counts of SDRs and shows the top five DECs

# Usage

```
## S3 method for class 'da'
summary(object, print = TRUE, ...)
```

# Arguments

object A S3 obj of class "da", output from pvda::da().

print Do you want to print the output to the console. Defaults to TRUE.

... For passing additional parameters to extended classes.

### Value

Passes a tibble with the SDR counts invisibly.

tiny\_dataset

A 110 reports big, simulated ICSR database

# **Description**

The dataframe tiny\_dataset is used to demonstrate the functionality of the package in examples. The larger drug\_event\_df-dataset can also be used.

### Usage

```
tiny_dataset
```

22 write\_to\_excel

# **Format**

'tiny\_dataset' A data frame with 110 rows and 3 columns. In total 110 unique report\_ids. In particular, for Drug A and Event 1 the observed count will be 4 and exp\_rrr = 1.1

```
report_id A report identifier, 1-110.
```

drug Drugs named as Drug\_A - Drug\_Z.

event Events named as Event\_1 - Event\_97)

group In this example, sex of the patient, i.e. Male or Female.

### **Source**

Simulated data.

write\_to\_excel

Write to excel

# Description

Writes output from a disproportionality analysis to an excel file

# Usage

```
write_to_excel(df, write_path = NULL)
```

# Arguments

df The data frame to export. See '?da' for details.

write\_path A string giving the file path

# Value

Nothing.

# **Index**

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