

# Package ‘vigicaen’

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**Title** 'VigiBase' Pharmacovigilance Database Toolbox

**Version** 1.0.0

**Description** Perform the analysis of the World Health Organization (WHO) Pharmacovigilance database 'VigiBase' (Extract Case Level version), [<https://who-umc.org/>](https://who-umc.org/) e.g., load data, perform data management, disproportionality analysis, and descriptive statistics. Intended for pharmacovigilance routine use or studies. This package is NOT supported nor reflect the opinion of the WHO, or the Uppsala Monitoring Centre. Disproportionality methods are described by Norén et al (2013) [<doi:10.1177/0962280211403604>](https://doi.org/10.1177/0962280211403604).

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<https://pharmacologie-caen.github.io/vigicaen/>

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---

add\_adr*Add adverse drug reaction column(s) to a dataset*

---

## Description

**[Stable]**Creates adr columns in vigibase datasets (demo, link, drug, and adr).

## Usage

```
add_adr(  
  .data,  
  a_code,  
  a_names = names(a_code),  
  adr_data,  
  data_type = deprecated()  
)
```

## Arguments

.data	The dataset to update (demo, link, drug, adr).
a_code	A named list of low level terms codes (llt_codes).
a_names	A character vector. Names for adr columns (must be the same length as adr_list), default to names(a_code)
adr_data	A data.frame containing the adr data (usually, it is adr)
data_type	<b>[Deprecated]</b> . Data_type is now detected internally.

## Details

Low-level term codes are the preferred level of requesting in Vigibase extract case level since they capture all possible codes for a given Preferred Term. Collect low-level terms with [get\\_llt\\_soc\(\)](#) and [get\\_llt\\_smq\(\)](#). You can add adr identification to a demo, a link, drug or even an adr dataset (in this latter case, you must provide adr twice, as .data and adr\_data). Column names of these dataset should not have been modified from the original vigibase dataset (as created with [tb\\_vigibase\(\)](#)).

## Value

A dataset with the new adr columns. Each element of a\_names will add a column with the same name in .data. The value can be 0 (the corresponding adr is absent) or 1 (the adr is present in the case if .data is demo or drug, or "this row correspond to this adr", if .data is adr or link).

## See Also

[add\\_drug\(\)](#), [get\\_llt\\_soc\(\)](#), [get\\_llt\\_smq\(\)](#)

## Examples

```
# create adr_colitis, adr_embolism and adr_pneumonitis columns in demo

# be careful, this example may overwrite your own demo dataset
demo <- demo_

a_pt_sel <- ex_$pt_sel

adr <- adr_

a_llt <-
  get_llt_soc(
    term_sel = a_pt_sel,
    term_level = "pt",
    meddra = meddra_
  )

demo <-
  demo |>
  add_adr(
    a_code = a_llt,
    adr_data = adr
  )

demo |>
  check_dm(names(a_pt_sel))
```

---

add\_dose

*Add drug dose column(s) to a dataset, in milligram per day*

---

## Description

**[Experimental]** `add_dose()` creates drug dose columns in vigibase datasets (`demo`, `link`, `adr`, `drug`, `ind`) for specified drugs in a dataset. It calculates daily dose values based on dose amount, frequency, and their corresponding units. The function is compatible with `demo`, `link`, `adr`, `drug` and `ind` datasets.

## Usage

```
add_dose(
  .data,
  d_code,
  d_dose_names = names(d_code),
  repbasis = "sci",
  method = c("DrecNo", "MedicinalProd_Id"),
  drug_data,
  verbose = TRUE
)
```

## Arguments

<code>.data</code>	The dataset used to identify individual reports (usually, it is demo)
<code>d_code</code>	A named list of drug codes (DrecNos or MPI). See Details.
<code>d_dose_names</code>	A character vector. Names for drug dose columns (must be the same length as <code>d_code</code> ), default to <code>names(d_code)</code> . Will be followed by a fixed suffix <code>"_dose_mg_per_day"</code> .
<code>repbasis</code>	Suspect, interacting and/or concomitant. Type initial of those you wish to select ("s" for suspect, "c" for concomitant and "i" for interacting ; default to all, e.g. "sci").
<code>method</code>	A character string. The type of drug code (DrecNo or MedicinalProd_Id). See details.
<code>drug_data</code>	A data.frame containing the drug data (usually, it is drug)
<code>verbose</code>	Logical, whether to display messages about added doses.

## Details

Actual supported dosage regimens are any combination of:

- Kilograms, grams, milligrams, micrograms, nanograms, or picograms
- Per minute, hour, day, week, month, or year.

Note that the result will be expressed in **milligrams per day**, whatever the aforementioned combination is. This may lead to very small or very large amounts in `drug_dose_mg_per_day` columns, depending on the actual dosage regimen. The function identifies drug doses in a dataset by cross-referencing with a `drug_data` table. If either the amount unit (grams, etc.) *or* the frequency (days, etc.) is missing in `drug_data`, the corresponding row will be omitted. Drugs may be filtered based on reputation bases (suspect, concomitant, or interacting). Either drug record numbers (e.g., from `get_drecno()`), or medicinalprod\_ids (e.g., from `get_atc_code()`) can be used to identify drugs. Default method is to DrecNos.

**It is very important to check the results**, as coding issues are common for dose data, and some results may seem unreliable.

## See Also

[add\\_drug\(\)](#), [get\\_drecno\(\)](#), [get\\_atc\\_code\(\)](#)

## Examples

```
# Example: Adding doses for paracetamol
d_code <- list(paracetamol = c(97818920, 97409107))
demo <-
  add_dose(
    .data = demo_,
    d_code = d_code,
    d_dose_names = "paracetamol",
    drug_data = drug_
  )
```

```

desc_cont(demo, "paracetamol_dose_mg_per_day")

# Use only drug dose where paracetamol had a "suspect" reputation base.
demo <-
  add_dose(
    .data = demo_,
    d_code = d_code,
    d_dose_names = "para_susp",
    repbasis = "s",
    drug_data = drug_
  )

desc_cont(demo, "para_susp_dose_mg_per_day")

```

---

add_drug	<i>Add drug column(s) to a dataset</i>
----------	--

---

## Description

**[Stable]** Creates drug columns. in vigibase datasets (demo, link, adr, drug, and ind).

## Usage

```

add_drug(
  .data,
  d_code,
  d_names = names(d_code),
  repbasis = "sci",
  method = c("DrecNo", "MedicinalProd_Id"),
  drug_data,
  data_type = deprecated()
)

```

## Arguments

.data	The dataset used to identify individual reports (usually, it is demo)
d_code	A named list of drug codes (DrecNos or MPI). See Details.
d_names	A character vector. Names for drug columns (must be the same length as d_code), default to names(d_code)
repbasis	Suspect, interacting and/or concomitant. Type initial of those you wish to select ("s" for suspect, "c" for concomitant and "i" for interacting ; default to all, e.g. "sci").
method	A character string. The type of drug code (DrecNo or MedicinalProd_Id). See details.
drug_data	A data.frame containing the drug data (usually, it is drug)
data_type	<b>[Deprecated]</b> . Data_type is now detected internally.

**Details**

d\_code is a named list containing drug codes. Either drug record numbers (e.g., from `get_drecno()`), or medicinalprod\_ids (e.g., from `get_atc_code()`). Default method is to DrecNos.

**Value**

A dataset with the new drug columns. Each element of d\_names will add a column with the same name in .data. The value can be 0 (the corresponding drug is absent) or 1 (the drug is present in the case if .data is demo or adr, or "this row correspond to this drug", if .data is drug or link).

**Argument** repbasis

Drugs can be reported according to one of three reputation bases:

- s for suspect
- c for concomitant
- i for interacting

in the occurrence of the adverse drug reaction. To study only one of these reputation basis, type only the corresponding letter in repbasis, e.g. "s" for suspects, or "si" for suspect **or** interacting.

You can add drug identification to a demo, link, adr, drug or ind dataset.(if working with drug, you must provide drug twice, as .data and drug\_data)

**See Also**

`add_adr()`, `get_drecno()`, `get_atc_code()`

**Examples**

```
# create a nivolumab column in demo_

d_sel_names <- list(nivolumab = "nivolumab")

d_drecno <- get_drecno(d_sel_names,
                      mp = mp_)

demo_ <-
  add_drug(
    .data = demo_,
    d_code = d_drecno,
    method = "DrecNo",
    repbasis = "sci",
    drug_data = drug_
  )

# remember to assign the result to your actual demo dataset

# do you want to work only with cases where nivolumab was a "suspected" drug?
# change argument repbasis to "s"

demo_ <-
```

```

add_drug(
  .data = demo_,
  d_code = d_drecno,
  d_names = "nivolumab_suspected",
  method = "DrecNo",
  repbasis = "s",
  drug_data = drug_
)

check_dm(demo_, cols = c("nivolumab", "nivolumab_suspected"))

```

---

add_ind	<i>Add indication column(s) to a dataset</i>
---------	--

---

## Description

**[Experimental]** Creates indication columns. in vigibase datasets (demo, link, adr, drug, or ind).

## Usage

```
add_ind(.data, i_list, i_names = names(i_list), drug_data, ind_data)
```

## Arguments

.data	The dataset used to identify individual reports (usually, it is demo)
i_list	A named list of indication terms. See Details.
i_names	A character vector. Names for indication columns (must be the same length as i_list), default to names(i_list)
drug_data	A data.frame containing the drug data (usually, it is drug)
ind_data	A data.frame containing the indication data (usually, it is ind)

## Details

Indication terms are issued from either MedDRA or International Classification of Diseases (ICD) - you need to use *both* dictionaries, should you wish to capture all terms related to a specific disease. Indication terms are not translated into codes in Vigibase ECL, unlike drug or adr terms. Therefore, there is no `get_*` step to collect such codes. The terms are passed directly to `i_list`, which should still be a *named* list containing indication terms.

## Value

A dataset with the new indication columns. Each element of `i_names` will add a column with the same name in `.data`. The value can be

- 0 The corresponding indication is absent.
- 1 The indication is present in the case if `.data` is demo or adr, or "this row correspond to this indication", if `.data` is drug, link or ind).
- NA There is no indication data for this case / drug.



**See Also**

[add\\_adr\(\)](#), [add\\_drug\(\)](#)

**Examples**

```
# Set up a list of indication terms

i_list <-
  list(
    melanoma = c("Malignant melanoma", "Metastatic malignant melanoma"),
    lung_cancer = c("Non-small cell lung cancer", "Lung adenocarcinoma")
  )

demo <-
  demo_ |>
  add_ind(i_list,
    drug_data = drug_,
    ind_data = ind_)

demo |> desc_facvar(names(i_list))
```

---

 cff

*Fast formatting of numbers*


---

**Description**

This is a formatting function for consistent number reporting.

**Usage**

```
cff(num, low_ci, up_ci, dig = 0, method = c("num_only", "num_ci", "ci"))
```

**Arguments**

num	A numeric. The number to format.
low_ci	A numeric. Lower end of a confidence interval
up_ci	A numeric. Upper end of a confidence interval
dig	A numeric. Number of digits
method	What sort of printing do you need? (see Details)

**Details**

Set method according to the printing you like: a unique number with `num_only` (default), the number and its confidence interval with `num_ci`, a `ci` only (for example a range of time to onset) The function properly returns NA when input is missing.

**Value**

A character vector with the formatted number(s)

**Examples**

```
num <- c(0.1, 0.02, 1.658)
```

```
cff(num)
```

```
cff(num, dig = 2)
```

```
cff(num = num[[1]],
     low_ci = num[[2]],
     up_ci = num[[3]],
     method = "num_ci",
     dig = 2)
```

---

check\_dm

*Check binary variables*

---

**Description**

**[Stable]** Quick check that your data management steps through [add\\_adr](#) or [add\\_drug](#) found cases.

**Usage**

```
check_dm(.data, cols)
```

**Arguments**

`.data` A data.frame to be checked

`cols` A character vector, name of columns to look at (usually will be d\_names, a\_names)

**Details**

It is a simple wrapper around `dplyr::summarise()`. Be careful not to supply factors with > 2 levels or continuous outcome (the function does NOT have a checker for this, so that it is faster). Also, the function WONT work with NAs. Use [desc\\_facvar\(\)](#). if you need more detailed description of your dataset.

**Value**

A transposed data.frame, with row.names equal to cols, and first column is the number of lines in .data where each col is equal to 1.

**See Also**

[desc\\_facvar\(\)](#), [add\\_adr\(\)](#), [add\\_drug\(\)](#)

**Examples**

```
# first create some new variables

demo <- demo_

demo <-
  demo |>
    add_adr(
      a_code = ex_$a_llt,
      adr_data = adr_
    )

# then check the number of reports with each feature

demo |>
  check_dm(names(ex_$a_llt))
```

---

compute_dispro	<i>Compute disproportionality</i>
----------------	-----------------------------------

---

**Description**

**[Stable]** Computes bivariate (reporting) Odds-Ratio and Information Component for a drug-adr pair.

**Usage**

```
compute_dispro(
  .data,
  y,
  x,
  alpha = 0.05,
  na_format = "-",
  dig = 2,
  export_raw_values = FALSE,
  min_n_obs = 0
)
```

**Arguments**

.data	The data.table to compute from.
y	A character vector, one or more variable to explain (usually an adr).
x	A character vector, one or more explaining variable (usually a drug).
alpha	Alpha risk.
na_format	Character string to fill NA values in ror and ci legends.
dig	Number of digits for rounding (this argument is passed to cff)

export\_raw\_values      A logical. Should the raw values be exported?

min\_n\_obs              A numeric, compute disproportionality only for pairs with at least min\_n\_obs cases.

## Details

Significance in pharmacovigilance analysis is only defined if the lower bound of the confidence/credibility interval is above 1 (i.e. `low_ci > 1`, or `ic_tail > 0`). Actually, the function computes an Odds-Ratio, which is not necessarily a **reporting** Odds-Ratio.

## Value

A data.table, with ROR, IC, and their confidence/credibility interval (at  $1 - \alpha$ ). Significance of both (as `signif_or` and `signif_ic`, if `export_raw_values` is TRUE).

A data.table with columns

- y and x, same as input
- n\_obs the number of observed cases
- n\_exp the number of expected cases
- or1 the formatted Odds-Ratio
- or\_ci the formatted confidence interval
- ic the Information Component
- ic\_tail the tail probability of the IC
- ci\_level the confidence interval level
- Additional columns, if `export_raw_values` is TRUE:
  - a, b, c, d the counts in the contingency table
  - std\_er the standard error of the log(OR)
  - or the Odds-Ratio
  - low\_ci the lower bound of the confidence interval
  - up\_ci the upper bound of the confidence interval
  - signif\_or the significance of the Odds-Ratio
  - signif\_ic the significance of the Information Component

## See Also

[compute\\_or\\_mod\(\)](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#)

## Examples

```
# Say you want to perform a disproportionality analysis between colitis and
# nivolumab among ICI cases

demo <-
  demo_ |>
```

```
add_drug(  
  d_code = ex_$d_drecno,  
  drug_data = drug_  
) |>  
add_adr(  
  a_code = ex_$a_llt,  
  adr_data = adr_  
)  
  
demo |>  
  compute_dispro(  
    y = "a_colitis",  
    x = "nivolumab"  
  )  
  
# You don't have to use the pipe syntax, if you're not familiar  
  
compute_dispro(  
  .data = demo,  
  y = "a_colitis",  
  x = "nivolumab"  
)  
  
# Say you want to compute more than one univariate ror at a time.  
  
many_drugs <-  
  names(ex_$d_drecno)  
  
demo |>  
  compute_dispro(  
    y = "a_colitis",  
    x = many_drugs  
  )  
  
# could do the same with adrs  
  
many_adrs <-  
  names(ex_$a_llt)  
  
demo |>  
  compute_dispro(  
    y = many_adrs,  
    x = many_drugs  
  )  
  
# Export raw values if you want to built plots, or other tables.  
  
demo |>  
  compute_dispro(  
    y = "a_colitis",  
    x = "nivolumab",
```

```

      export_raw_values = TRUE
    )

# Set a minimum number of observed cases to compute disproportionality

demo |>
  compute_dispro(
    y = "a_colitis",
    x = "nivolumab",
    min_n_obs = 5
  )

```

---

compute_interaction	<i>Compute interaction disproportionality</i>
---------------------	---

---

## Description

**[Experimental]** Returns the information component of interaction for a set of 3 variables, usually 2 drugs and an adr.

## Usage

```

compute_interaction(
  .data,
  y,
  x,
  z,
  alpha = 0.05,
  na_format = "-",
  dig = 2,
  export_raw_values = FALSE,
  min_n_obs = 0
)

```

## Arguments

.data	The data.table to compute from.
y	A character vector, one or more variable to explain.
x	A character vector, one or more explaining variable.
z	A character vector, one or more explaining variable.
alpha	Alpha risk.
na_format	Character string to fill NA values in ror and ci legends.
dig	Number of digits for rounding (this argument is passed to cff)
export_raw_values	A logical. Should the raw values be exported?
min_n_obs	A numeric, compute disproportionality only for pairs with at least min_n_obs cases.

**Details**

Significance is similar to usual disproportionality (see [compute\\_dispro\(\)](#)).

**Value**

A data.table, with Information Component (IC) of interaction, and its credibility interval (at 1 - alpha). Significance as signif\_ic, if export\_raw\_values is TRUE).

A data.table with columns

- y, x and z, same as input
- n\_obs the number of observed cases
- n\_exp the number of expected cases
- ic the Information Component
- ic\_tail the tail probability of the IC
- ci\_level the confidence interval level
- Additional columns, if export\_raw\_values is TRUE:
- a, b, c, d the counts in the contingency table
- signif\_ic the significance of the Information Component
- Additional columns, if export\_raw\_values is TRUE:
- n\_\* the counts of each setting
- signif\_ic the significance of the Information Component

**See Also**

[compute\\_dispro\(\)](#), [compute\\_or\\_mod\(\)](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#)

**Examples**

```
# Interaction on reporting of colitis with ipilimumab and nivolumab
demo <-
  demo_ |>
  add_drug(
    d_code = ex_$d_drecno,
    drug_data = drug_
  ) |>
  add_adr(
    a_code = ex_$a_llt,
    adr_data = adr_
  )

demo |>
  compute_interaction(
    y = "a_colitis",
    x = "nivolumab",
    z = "ipilimumab"
  )
```

---

compute_or_mod	<i>Compute (r)OR from a model summary</i>
----------------	---

---

## Description

**[Stable]** Compute and format Odds-Ratio from a model summary.

## Usage

```
compute_or_mod(.coef_table, estimate, std_er, p_val = NULL, alpha = 0.05)
```

## Arguments

.coef_table	A coefficient table, see details.
estimate	Quasiquoted name of estimate parameter.
std_er	Quasiquoted name of standard error parameter.
p_val	Quasiquoted name of p-value parameter. Optional.
alpha	alpha risk.

## Details

Helper to compute and format Odds-Ratio based on `summary(glm)$coefficients`, or any equivalent in other modelling packages. (see examples). Preferably, it is transformed into a `data.table` or `data.frame` before being evaluated in the function. Otherwise, `compute_or_mod()` will transform it. Significant OR-or column means `low_ci` is  $> 1$ . The `p_val` argument is only required if you wished to display a [nice\\_p\(\)](#).

Output is a `data.table`. Actually, the function computes an Odds-Ratio, which is not necessarily a *reporting* Odds-Ratio.

## Value

A `data.table` with OR, confidence intervals (at  $1 - \alpha$ ), significance (`low_ci`  $> 1$ ) and (optionally) p-value.

## See Also

[compute\\_dispro\(\)](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#)

## Examples

```
# Reporting Odds-Ratio of colitis with nivolumab among ICI cases.

demo <-
  demo_ |>
  add_drug(
    d_code = ex_$d_drecno,
    drug_data = drug_
```



```

) |>
add_adr(
  a_code = ex_$a_llt,
  adr_data = adr_
)

# Compute the model
mod <- glm(a_colitis ~ nivolumab, data = demo, family = "binomial")

# Extract coefficients
mod_summary <-
  mod |>
  summary()

coef_table <-
  mod_summary$coefficients

# Transform coefficients into ORs with their CI

coef_table |>
  compute_or_mod(
    estimate = Estimate,
    std_er = Std..Error,
    p_val = Pr...z..)

# Also works if you don't have a p_val column
coef_table |>
  compute_or_mod(
    estimate = Estimate,
    std_er = Std..Error)

```

---

create\_example\_tables *Example source tables for VigiBase and MedDRA*

---

## Description

**[Stable]** Write some example tables as source text/ascii/parquet files.

## Usage

```

create_ex_main_txt(path)

create_ex_sub_txt(path)

create_ex_who_txt(path)

create_ex_meddra_asc(path)

create_ex_main_pq(path)

create_ex_sub_pq(path)

```

**Arguments**

path                      Character string. A folder on your computer where the tables should be written.

**Details**

VigiBase tables and MedDRA tables are provided respectively as text files and ascii files. The `tb_*` family turns them into parquet files. These `create_example_*` functions are only used to produce example source files to illustrate the `tb_*` family, and parquet files for the same purpose.

**Value**

A set of text/ascii files, as received by the Uppsala Monitoring Centre or MedDRA

- For `create_ex_main_txt()`, DEMO.txt, DRUG.txt, LINK.txt, FOLLOWUP.txt, ADR.txt, OUT.txt, SRCE.txt, and IND.txt
- For `create_ex_sub_txt()`, AgeGroup\_Lx.txt, Dechallenge\_Lx.txt, Dechallenge2\_Lx.txt, Frequency\_Lx.txt, Gender\_Lx.txt, Notifier\_Lx.txt, Outcome\_Lx.txt, Rechallenge\_Lx.txt, Rechallenge2\_Lx.txt, Region\_Lx.txt, RepBasis\_Lx.txt, ReportType\_Lx.txt, RouteOfAdm\_Lx.txt, Seriousness\_Lx.txt, and SizeUnit\_Lx.txt
- For `create_ex_who_txt()`, ATC.txt, CCODE.txt, ING.txt, MP.txt, ORG.txt, PF.txt, PP.txt, PRT.txt, PRG.txt, SRCE.txt, STR.txt, SUN.txt, ThG.txt, and Unit-X.txt
- For `create_ex_meddra_asc()`, llt.asc, mdhier.asc, smq\_content.asc, smq\_list.asc
- For `create_ex_main_pq()`, demo.parquet, adr.parquet, drug.parquet, link.parquet, srce.parquet, ind.parquet, out.parquet, followup.parquet, suspdup.parquet
- For `create_ex_sub_pq()`, agegroup.parquet, dechallenge.parquet, dechallenge2.parquet, frequency.parquet, gender.parquet, notifier.parquet, outcome.parquet, rechallenge.parquet, rechallenge2.parquet, region.parquet, repbasis.parquet, reporttype.parquet, routeofadm.parquet, seriousness.parquet, and sizeunit.parquet

**Functions**

- `create_ex_sub_txt()`: sub txt tables
- `create_ex_who_txt()`: WHO txt tables
- `create_ex_meddra_asc()`: MedDRA txt tables
- `create_ex_main_pq()`: main parquet tables
- `create_ex_sub_pq()`: subsidiary parquet tables

**See Also**

[tb\\_vigibase\(\)](#), [tb\\_who\(\)](#), [tb\\_meddra\(\)](#)

**Examples**

```
path <- paste0(tempdir(), "/crex/")

dir.create(path)
```

```
# You may want to use different paths for each type of tables
create_ex_main_txt(path)

create_ex_sub_txt(path)

create_ex_who_txt(path)

create_ex_meddra_asc(path)

create_ex_main_pq(path)

create_ex_sub_pq(path)

# Remove temporary folders when you're done
unlink(path, recursive = TRUE)
```

demo\_

*Data of immune checkpoint inhibitors.*

## Description

Demo, drug, adr, link, ind, out, srce, and followup are the main table in Vigibase Extract Case Level data. In a regular workflow, you will work with those tables as R objects (e.g. demo, drug, adr, link, ind, out, srce, followup). These built-in example datasets use an underscore "\_" to avoid ambiguity with your own tables (e.g. demo\_, drug\_, adr\_, link\_, ind\_, out\_, srce\_, followup\_). This is a relational database, which means every table has a primary key variable (e.g., UMCReportId for demo\_. Keys will allow joints with other tables The full details on the original structure can be found in "VigiBase Extract Case Level - file description.pdf" in your VigiBase ECL folders. demo\_ will typically be your cornerstone table, since it contains one row per report. It is the preferred table to update for drugs and adrs identification before performing disproportionality analyses. These tables are subsets of the original ones, with some of the immune checkpoint inhibitor cases or immune-related adverse events. All data shown in these example data are **FAKE**, which means you shouldn't consider the counts and computations as accurate. Immune checkpoint inhibitors drugs include "Ipilimumab", "Atezolizumab", "Durvalumab", "Nivolumab", "Pembrolizumab", "Avelumab", "Cemiplimab", "REGN 2810", "Tremelimumab". More details on how to use vigibase tables can be found in the vignettes. `vignette("basic_workflow")`, `vignette("descriptive")`. To build your own tables, use `tb_vigibase()`. See `vignette("getting_started")`.

## Usage

```
data(demo_)
```

```
drug_
```

```
adr_
```

```
link_
```

followup\_

ind\_

out\_

srce\_

## Format

demo\_ is a data.table with 7 variables and 750 rows.

- UMCReportId Integer. The unique identifier of the case report.
- AgeGroup Character. The age group of the patient. Correspondence table is path\_sub/AgeGroup.parquet.
- Gender Character. Case gender. path\_sub/Gender.parquet
- DateDatabase Character (not date or numeric!). The date of the latest update of the report in the database.
- Type Character. The type of report. path\_sub/ReportType.parquet
- Region Character. The world region where the report comes from path\_sub/Region.parquet.
- FirstDateDatabase Character. The date the report was first submitted to the database.

drug\_ is a data.table with 10 variables and 3514 rows.

- UMCReportId Integer. See demo\_.
- Drug\_Id Integer. The unique identifier of each drug report.
- MedicinalProd\_Id Integer. The medicinalproduct identifier. See [get\\_atc\\_code\(\)](#).
- DrecNo Integer. Drug Record Number, pivotal to identify drugs with [get\\_drecno\(\)](#).
- Seq1, Seq2 Character. Seq 1 and 2 complement DrecNo, in WHODrug dictionary.
- Route Character. The route of administration of the drug.
- Basis Character. The reputation basis of the drug (suspect, concomitant, or interacting). path\_sub/RepBasis.parquet
- Amount Character. The amount of drug administered.
- AmountU Character. The unit of the amount of drug administered. path\_sub/SizeUnit.parquet
- Frequency Character. The frequency of drug administration.
- FrequencyU Character. The unit of the frequency of drug administration. path\_sub/Frequency.parquet

adr\_ is a data.table with 4 variables and 2133 rows.

- UMCReportId Integer. See demo\_.
- Adr\_Id Integer. The unique identifier of each adverse event report.
- MedDRA\_Id Integer. The MedDRA identifier of the adverse event. It is used in [get\\_llt\\_soc\(\)](#) and [get\\_llt\\_smq\(\)](#).
- Outcome Character. The outcome of the adverse event. path\_sub/Outcome.parquet

link\_ is a data.table with 3 variables and 3514 rows. The version built with `tb_vigibase()` is slightly different than the original one.

- Drug\_Id and Adr\_Id . Integers. Together, they are the key variable of link. See drug\_ and adr\_.
- Dechallenge1 and 2 Characters. Dechallenge action and outcome. path\_sub/Dechallenge.parquet, path\_sub/Dechallenge2.parquet
- Rechallenge1 and 2 Characters. Rechallenge action and outcome. path\_sub/Rechallenge.parquet, path\_sub/Rechallenge2.parquet
- TimeToOnsetMin and Max Numerics. The minimum and maximum time to onset of the adverse event.
- tto\_mean Numeric. The mean time to onset of the adverse event. It is the average of TimeToOnsetMin and Max.
- range Numeric. The incertitude around tto\_mean. See vignette("descriptive").
- UMCReportId Integer. See demo\_.

ind\_ is a data.table with 2 variables and 2426 rows.

- Drug\_Id Integer. See drug\_.
- Indication Character. The indication of the drug.

out\_ is a data.table with 3 variables and 747 rows.

- UMCReportId Integer. See demo\_.
- Seriousness Character. The seriousness criteria of the report. path\_sub/Seriousness.parquet
- Serious Character. Whether the case is serious or not ("N" No, "Y" Yes)

srce\_ is a data.table with 2 variables and 729 rows.

- UMCReportId Integer. See demo\_.
- Type Character. The Type of Reporter. path\_sub/Notifier.parquet

followup\_ is a data.table with 2 variables and 902 rows.

- UMCReportId Integer. See demo\_.
- ReplacedUMCReportId Integer. Previous version of the case, which is no longer available in demo\_.

An object of class data.table (inherits from data.frame) with 3514 rows and 12 columns.

An object of class data.table (inherits from data.frame) with 2133 rows and 4 columns.

An object of class data.table (inherits from data.frame) with 5136 rows and 11 columns.

An object of class data.table (inherits from data.frame) with 902 rows and 2 columns.

An object of class data.table (inherits from data.frame) with 2426 rows and 2 columns.

An object of class data.table (inherits from data.frame) with 747 rows and 3 columns.

An object of class data.table (inherits from data.frame) with 729 rows and 2 columns.

**Source**

None

**References**

There is none

**Examples**

```
data(demo_)
demo_ |> dplyr::count(AgeGroup)
```

---

desc_cont	<i>Summarise continuous variables</i>
-----------	---------------------------------------

---

**Description**

**[Stable]** Summarize continuous data and handle output format.

**Usage**

```
desc_cont(
  .data,
  vc,
  format = "median (q1-q3) [min-max]",
  digits = 1,
  export_raw_values = FALSE
)
```

**Arguments**

.data	A data.frame, where vc are column names of continuous variables
vc	A character vector, list of column names. Should only contain continuous variables
format	A character string. How would you like the output? See details.
digits	A numeric. How many digits? This argument calls internal formatting function
export_raw_values	A logical. Should the raw values be exported?

**Details**

Many other packages provide tools to summarize data. This one is just the package author's favorite. This makes it much easier to map to nice labeling thereafter. The format argument shows the output of the function. You can change square and round brackets, spaces, separators... Important format inputs are

- median the median value

- q1 the first quartile
- q3 the third quartile
- min the minimum value
- max the maximum value

The analogous for categorical variables is [desc\\_facvar\(\)](#).

## Value

A data.frame with columns

- var the variable name
- level NA, it is provided to have a consistent output with [desc\\_facvar\(\)](#)
- value the formatted value with possibly the median, interquartile range, and range (see details)
- n\_avail the number of cases with available data for this variable.

## See Also

[desc\\_facvar\(\)](#)

## Examples

```
df <-
  data.frame(
    smoke_status = c("smoker", "non-smoker",
                     "smoker", "smoker",
                     "smoker", "smoker",
                     "non-smoker"
                    ),
    age = c(60, 50, 56, 49, 75, 69, 85),
    bmi = c(18, 30, 25, 22, 23, 21, 22)
  )

# Use default formatting

desc_cont(.data = df, vc = c("age", "bmi"))

# Use custom formatting

desc_cont(.data = df,
          vc = c("age", "bmi"),
          format = "median (q1;q3)"
        )

# You might want to export raw values, to run plotting or
# other formatting functions

desc_cont(.data = df, vc = c("age", "bmi"),
          export_raw_values = TRUE)
```

---

desc_dch	<i>Dechallenge descriptive</i>
----------	--------------------------------

---

## Description

**[Stable]** Computes positive dechallenge counts over a set of adr and drug pairs.

## Usage

```
desc_dch(.data, drug_s = "drug1", adr_s = "adr1")
```

## Arguments

.data	A link data.table.
drug_s	A character vector, the drug column(s)
adr_s	A character vector, the adverse drug reaction column(s).

## Details

Counts are provided at the **case** level (not the drug-adr pair level). Positive dechallenge refers to cases where drug was withdrawn or dose-reduced and reaction abated (in part or in full). You will need a link data.table, see [link\\_](#), on which you have added drugs and adrs with [add\\_drug\(\)](#) and [add\\_adr\(\)](#).

## Value

A data.table with one row per drug-adr pair.

- drug\_s and adr\_s, same as input
- pos\_dch, number of positive dechallenge cases

## See Also

[link\\_](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#), [desc\\_tto\(\)](#), [desc\\_rch\(\)](#)

## Examples

```
link_ <-
link_ |>
add_drug(
  d_code = ex_$d_groups_drecno,
  drug_data = drug_
) |>
add_adr(
  a_code = ex_$a_llt,
  adr_data = adr_
)
```



```

desc_dch(link_,
  drug_s = "pd1",
  adr_s = "a_colitis")

# you can vectorize over multiple adrs and drugs

desc_dch(link_,
  drug_s = c("pd1", "pd11"),
  adr_s = c("a_colitis", "a_pneumonitis"))

```

---

desc_facvar	<i>Summarise categorical variables</i>
-------------	--

---

## Description

**[Stable]** Summarize categorical data and handle output format.

## Usage

```

desc_facvar(
  .data,
  vf,
  format = "n_/N_ (pc_%)",
  digits = 0,
  pad_width = 12,
  ncat_max = 20,
  export_raw_values = FALSE
)

```

## Arguments

.data	A data.frame, where vf are column names of categorical variables
vf	A character vector
format	A character string, formatting options.
digits	A numeric. Number of digits for the percentage (passed to interval formatting function).
pad_width	A numeric. Minimum character length of value output (passed to <code>stringr::str_pad()</code> ).
ncat_max	A numeric. How many levels should be allowed for all variables? See details.
export_raw_values	A logical. Should the raw values be exported?

## Details

Many other packages provide tools to summarize data. This one is just the package author's favorite. Important format inputs are

- `n_` number of patients with the categorical variable at said level
- `N_` the first quartile number of patients with an available value for this variable
- `pc_` percentage of  $n / N$

The format argument should contain at least the words "`n_`", "`N_`", and optionally "`pc_`". `ncat_max` ensures that you didn't provided a continuous variable to `desc_facvar()`. If you have many levels for one of your variables, set to `Inf` or high value. Equivalent for continuous data is `desc_cont()`.

## Value

A `data.frame` with columns

- `var` the variable name
- `level` the level of the variable
- `value` the formatted value with possible number of cases `n_`, number of available cases `N_`, and percentage `pc_`, depending on format argument.
- `n_avail` the number of cases with available data for this variable.

## See Also

[desc\\_cont\(\)](#)

## Examples

```
df1 <-
  data.frame(
    smoke_status = c("smoker", "non-smoker",
                     "smoker", "smoker",
                     "smoker", "smoker",
                     "non-smoker"
                    ),
    hypertension = c(1, 1, 0, 1, 1, 1, 1),
    age = c(60, 50, 56, 49, 75, 69, 85),
    bmi = c(18, 30, 25, 22, 23, 21, 22)
  )

# Use default formatting
desc_facvar(.data = df1, vf = c("hypertension", "smoke_status"))

# Use custom formatting
desc_facvar(.data = df1,
            vf = c("hypertension", "smoke_status"),
            format = "n_ out of N_, pc_%",
            digits = 1)

# You might want to export raw values, to run plotting or
```

```
# other formatting functions

desc_facvar(.data = df1,
            vf = c("hypertension", "smoke_status"),
            export_raw_values = TRUE)
```

---

desc_outcome	<i>Outcome descriptive</i>
--------------	----------------------------

---

## Description

**[Stable]** Compute outcome description over a set of adr and drugs.

## Usage

```
desc_outcome(.data, drug_s = "drug1", adr_s = "adr1")
```

## Arguments

.data	An adr data.table. See <a href="#">adr_</a>
drug_s	A character vector, the drug column(s)
adr_s	A character vector, the adverse drug reaction column(s).

## Details

You need an adr data.table. Be careful that you cannot directly filter adr data.table on drugs! You first have to add drug columns to adr, with [add\\_drug\(\)](#). The function reports the worst outcome into consideration for a given case, if many are reported. Outcomes, from best to worst are:

- Recovered/resolved
- Recovering/resolving
- Recovered/resolved with sequelae
- Not recovered/not resolved
- Fatal
- Died- unrelated to reaction
- Died- reaction may be contributory

See vignette("descriptive") for more details.

## Value

A data.table with one row per drug-adr pair.

- drug\_s and adr\_s, same as input
- n\_cas, number of cases for each category
- out\_label, the worst outcome for this drug-adr pair

See Also

[adr\\_](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#)

Examples

```
adr_ <-
  adr_ |>
  add_drug(
    d_code = ex_$d_groups_drecno,
    drug_data = drug_
  ) |>
  add_adr(
    a_code = ex_$a_llt,
    adr_data = adr_
  )

desc_outcome(
  adr_,
  drug_s = "pd1",
  adr_s = "a_colitis"
)

# you can vectorize over multiple adrs and drugs

desc_outcome(
  adr_,
  drug_s = c("pd1", "pd11"),
  adr_s = c("a_colitis", "a_pneumonitis")
)
```

---

desc_rch	<i>Rechallenge descriptive</i>
----------	--------------------------------

---

Description

**[Stable]** Computes counts of rechallenge cases, over a set of adr and drug pairs.

Usage

```
desc_rch(.data, drug_s = "drug1", adr_s = "adr1")
```

Arguments

- |        |  |
|--------|--|
| .data  | A link data.table. See <a href="#">link_</a> .   |
| drug_s | A character string. The name of the drug column. Drug columns can be created with <a href="#">add_drug</a> . |
| adr_s  | A character string. The name of the adr column. Adr columns can be created with <a href="#">add_adr</a> .    |

## Details

Counts are provided at the **case** level (not the drug-adr pair level). Description span from number of rechallenge cases to **informative** rechallenge cases (those cases where the outcome is known). You will need a link data.table, see [link\\_](#), on which you have added drugs and adrs with [add\\_drug\(\)](#) and [add\\_adr\(\)](#). Terminology

- Overall as opposed to rch for rechallenged ( $\text{rch} + \text{no\_rch} = \text{overall}$ ).
- Among rch, inf (informative) as opposed to non\_inf ( $\text{inf} + \text{non\_inf} = \text{rch}$ )
- Among inf, rec (recurring) as opposed to non\_rec ( $\text{rec} + \text{non\_rec} = \text{inf}$ )

## Value

A data.table with one row per drug-adr pair

- drug\_s and adr\_s, same as input.
- Counts of **overall**, **rch**, **inf**, and **rec** cases (see details).

## See Also

[link\\_](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#), [desc\\_dch\(\)](#), [desc\\_tto\(\)](#)

## Examples

```
link_ <-
  link_ |>
  add_drug(
    d_code = ex_$d_groups_drecno,
    drug_data = drug_
  ) |>
  add_adr(
    a_code = ex_$a_llt,
    adr_data = adr_
  )

desc_rch(.data = link_,
         drug_s = "pd1",
         adr_s = "a_colitis")

# You can vectorize over drugs and adrs

desc_rch(.data = link_,
         adr_s = c("a_colitis", "a_pneumonitis"),
         drug_s = c("pd1", "pd11")
       )
```

---

desc_tto	<i>Time to onset descriptive</i>
----------	----------------------------------

---

## Description

**[Stable]** desc\_tto() provides a drug-adr pair description of time to onset.

## Usage

```
desc_tto(.data, adr_s, drug_s, tto_time_range = 1, ...)
```

## Arguments

.data	A link data.table. See <a href="#">link_</a> .
adr_s	A character string. The name of the adr column. (see details)
drug_s	A character string. The name of the drug column. (see details)
tto_time_range	Incertitude range of Time to onset, in days. Defaults to 1 as recommended by umc
...	Additional parameters to be passed to <a href="#">desc_cont()</a> . E.g. format, digits...

## Details

Description of time (maximum available time) between drug initiation and event onset. This runs at the drug-adr pair level. Internally, it uses [extract\\_tto\(\)](#) and [desc\\_cont\(\)](#). You will need a link data.table, see [link\\_](#), on which you have added drugs and adrs with [add\\_drug\(\)](#) and [add\\_adr\(\)](#). you can supply extra arguments to [desc\\_cont\(\)](#) with ... Uppsala Monitoring Centre recommends to use only cases where the incertitude on time to onset is less than **1 day**. You can change this with tto\_time\_range.

## Value

A data.table with one row per drug-adr pair

- A descriptive of time to onsets for this combination (column tto\_max).
- It is Median (Quartile 1 - Quartile 3) and min-max by default, change with format arg (passed to [desc\\_cont\(\)](#)).

## See Also

[link\\_](#), [extract\\_tto\(\)](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#), [desc\\_dch\(\)](#), [desc\\_rch\(\)](#)

### Examples

```
link_ <-
  link_ |>
  add_drug(
    d_code = ex_$d_groups_drecno,
    drug_data = drug_
  ) |>
  add_adr(
    a_code = ex_$a_llt,
    adr_data = adr_
  )

desc_tto(.data = link_,
        adr_s = "a_colitis",
        drug_s = "pd1")

desc_tto(.data = link_,
        adr_s = c("a_colitis", "a_pneumonitis"),
        drug_s = c("pd1", "ctl4"))
```

---

dt\_parquet

Read parquet and convert to data.table

---

### Description

**[Stable]** Load data IN- our OUT- of memory. File extension can be omitted.

### Usage

```
dt_parquet(path_base, name = NULL, ext = ".parquet", in_memory = TRUE)
```

### Arguments

path_base	A character string, providing the path to read from.
name	Optional. A character string. The file name (if absent from path_base).
ext	Optional. A character string. The file extension.
in_memory	Logical, should data be loaded in memory?

### Details

Output is a data.table. For meddra and whodrug tables, it is still a good option to load data in-memory. This function is wrapping `arrow::read_parquet()`, `dplyr::collect()` and `data.table::as.data.table()` altogether. If you want to load **OUT** of memory, set arg `in_memory` to `FALSE`. **Be careful that doing so will change the function output format.** For this latter case, the output is not a data.table, so there is no practical benefit as compared to using `arrow::read_parquet()` directly, with `as_data_frame = FALSE`.

**Value**

A data.table if in\_memory is set to TRUE, a parquet Table if in\_memory is set to FALSE.

**See Also**

[tb\\_vigibase\(\)](#), [tb\\_who\(\)](#), [tb\\_meddra\(\)](#)

**Examples**

```
# Say you have a data.frame stored in a parquet format, such as this one
demo <-
  data.table::data.table(
    UMCReportId = c(1, 2, 3, 4),
    AgeGroup = c(1, 7, 7, 8)
  ) |>
  arrow::as_arrow_table()

tmp_folder <- paste0(tempdir(), "/dtparquetex")
dir.create(tmp_folder)
path_data <- paste0(tmp_folder, "/")

arrow::write_parquet(demo,
  sink = paste0(path_data, "demo.parquet")
)

# Now you have a new session without demo
rm(demo)

# You may import the file directly to data.table format with dt_parquet
demo <-
  dt_parquet(path_data, "demo")

# Clean up (required for CRAN checks)
unlink(tmp_folder, recursive = TRUE)
```

---

extract\_tto

*Time to onset extraction*

---

**Description**

**[Stable]** extract\_tto() collects all available time to onsets for a set of drug-adr pairs.

**Usage**

```
extract_tto(.data, adr_s, drug_s, tto_time_range = 1)
```



## Arguments

<code>.data</code>	A link data.table. See <a href="#">link_</a> .
<code>adr_s</code>	A character string. The name of the adr column. (see details)
<code>drug_s</code>	A character string. The name of the drug column. (see details)
<code>tto_time_range</code>	Incertitude range of Time to onset, in days. Defaults to 1 as recommended by umc

## Details

Extraction of (maximum available) time between drug initiation and event onset. This runs at the drug-adr pair level. You will need a link data.table, see [link\\_](#), on which you have added drugs and adrs with [add\\_drug\(\)](#) and [add\\_adr\(\)](#). Uppsala Monitoring Centre recommends to use only cases where the incertitude on time to onset is less than **1 day**. You can change this with `tto_time_range`. You might want to use [desc\\_tto\(\)](#) to obtain summary statistics of time to onset, but `extract_tto()` is useful to get the raw data and plot it, for instance with `ggplot2`.

## Value

A data.frame with

- All available time to onsets for this combination (column `tto_max`).
- `adr_s` and `drug_s`, same as input.
- `UMCReportId`, the unique identifier of the case.

## See Also

[link\\_](#), [desc\\_tto\(\)](#), [add\\_drug\(\)](#), [add\\_adr\(\)](#), [desc\\_dch\(\)](#), [desc\\_rch\(\)](#)

## Examples

```
link_ <-
  link_ |>
  add_drug(
    d_code = ex_$d_groups_drecno,
    drug_data = drug_
  ) |>
  add_adr(
    a_code = ex_$a_llt,
    adr_data = adr_
  )

extract_tto(.data = link_,
            adr_s = "a_colitis",
            drug_s = "pd1")
extract_tto(.data = link_,
            adr_s = c("a_colitis", "a_pneumonitis"),
            drug_s = c("pd1", "ctl4"))
```

---

ex\_*Data for the immune checkpoint inhibitors example*

---

## Description

These are a set of data to provide examples on the package.

- `smq_sel` is a named list of smq names
- `pt_sel` is a named list of pt names
- `a_llt` is a named list of meddra llt codes related to adrs from `smq_sel` and `pt_sel`
- `d_drecno` is a named list of drecnos for immune checkpoint inhibitors (some of them)
- `d_groups` is a named list of ici classes according to icis
- `d_groups_drecno` is a named list of drecnos for drug groups

## Usage

```
data(ex_)
```

## Format

An object of class `list`.

## Source

VigiBase Extract Case Level

## References

There is none

## Examples

```
data(ex_)
ex_$pt_sel
```

---

get_atc_code	<i>Get ATC codes (DrecNos or MPIs)</i>
--------------	--

---

### Description

**[Stable]** Collect Drug Record Numbers or MedicinalProd\_Ids associated to one or more ATC classes.

### Usage

```
get_atc_code(atc_sel, mp, thg_data, vigilyze = TRUE)
```

### Arguments

atc_sel	A named list of ATC codes. See Details.
mp	A modified MP data.table. See <a href="#">mp_</a>
thg_data	A data.table. Correspondence between ATC codes and MedicinalProd_Id (usually, it is thg)
vigilyze	A logical. Should ATC classes be retrieved using the vigilyze style? See details

### Details

get\_atc\_code() is an *ID collector* function. Provide atc\_sel in the same way as d\_sel in [add\\_drug\(\)](#), but remember to specify its method arg as MedicinalProd\_Id if vigilyze is set to FALSE. Vigilyze style means all conditioning of drugs will be retrieved after requesting an ATC class (i.e., drugs are identified with their DrecNos), even if a specific conditioning is not present in the ATC class. This is the default behavior in vigilyze.

### Value

A named list of integers. **DrecNos** if vigilyze is set to TRUE, or **MedicinalProd\_Ids** if vigilyze is set to FALSE.

### See Also

[mp\\_](#), [thg\\_](#), [add\\_drug\(\)](#), [get\\_drecno\(\)](#)

### Examples

```
# ## Find codes associated with one or more atc classes

# First, define which atc you want to use

atc_sel <-
  rlang::list2(penicillins_gcsf = c("L03AA", "J01CA"),
               ace_inhibitors = c("C09AA")
  )
```

```
# You can get DrecNos for you ATCs (if vigilyze is TRUE)

atc_drecno <-
  get_atc_code(atc_sel = atc_sel,
               mp = mp_,
               thg_data = thg_,
               vigilyze = TRUE)

# Or you can get MedicinalProd_Ids (if vigilyze is FALSE)

atc_mpi <-
  get_atc_code(atc_sel = atc_sel,
               mp = mp_,
               thg_data = thg_,
               vigilyze = FALSE)
```

get\_drecno

*Get DrecNo from drug names or MedicinalProd\_Id*

## Description

**[Stable]** Collect Drug Record Numbers associated to one or more drugs.

## Usage

```
get_drecno(
  d_sel,
  mp,
  allow_combination = TRUE,
  method = c("drug_name", "mpi_list"),
  verbose = TRUE,
  show_all = deprecated(),
  inspect = deprecated()
)
```

## Arguments

d_sel	A named list. Selection of drug names or medicinalprod_id. See details
mp	A modified MP data.table. See <a href="#">mp_</a>
allow_combination	A logical. Should fixed associations including the drug of interest be retrieved? See details.
method	Should DrecNo be found from drug names or from MedicinalProd_Id?
verbose	A logical. Allows you to see matching drug names in the console. Turn to FALSE once you've checked the matching.
show_all	<b>[Deprecated]</b> Use verbose instead.
inspect	<b>[Deprecated]</b> Use verbose instead.

## Details

get\_drecno() is an *ID collector* function. Collected IDs can be used to create drug columns in datasets like demo, link, etc. (see vignette("basic\_workflow"))

## Value

A named list of integers. DrecNos.

## Argument verbose

The verbose argument is here to let you check the result of get\_drecno(). This is an important step in your project setup: You must ensure that the drugs you are looking for are correctly matched.

## Argument d\_sel

d\_sel must be a named list of character vectors. To learn why, see vignette("basic\_workflow"). Names of d\_sel are automatically lowered and trimmed.

## Matching drugs

With "drug\_name" method, either exact match or perl regex match can be used. The latter is built upon lookarounds to ensure that a string does not match to composite drug names including the string, i.e. trastuzumab emtasine for trastuzumab, or close names like alitretinoin when looking for tretinoin.

Exact match is used for "mpi\_list" method.

## Choosing a method

"drug\_name" let you work with drug names. It's likely to be the appropriate method in most of the cases.

"mpi\_list" is used when you have a list of MedicinalProd\_Ids. A drug can have multiple MedicinalProd\_Ids, corresponding to different packagings. The MedicinalProd\_Id matching is typically used to identify DrecNo(s) contained in an ATC class (extracted from thg), since not all MPI of drugs are present in thg (explanations in [get\\_atc\\_code\(\)](#)).

## WHO names

WHO names are attributed to drugs by... the WHO. A drug only has one WHO name, but can have multiple international nonproprietary names (e.g. "tretinoin" and "all-trans retinoic acid").

You should use WHO names to ensure proper identification of drugs and DrecNos, especially if you work with combinations.

## Argument allow\_combination

Fixed associations of drugs refers to specialty containing more than one active ingredient (for example, acetylsalicylic acid and clopidogrel). In VigiLyze, the default is **NOT** to account for these fixed associations. For example, when you call "acetylsalicylic acid" in VigiLyze, you don't have the cases reported with the fixed-association "acetylsalicylic acid; clopidogrel" **unless the substances**

**were distinctly coded by the reporter.** Here, the default is to find a drug even if it is prescribed in a fixed association. Importantly, when retrieving fixed-association drugs, the non-of-interest drug alone drecno is not found, hence the cases related to this drug will not be added to those of the drug of interest.

### See Also

[add\\_drug\(\)](#), [get\\_atc\\_code\(\)](#)

### Examples

```
# ## Get drecnos for a list a drugs. Check spelling and use WHO name,
# in lowercase

d_sel_names <- list(
  nivolumab = "nivolumab",
  ipilimumab = "ipilimumab",
  nivo_ipi = c("nivolumab", "ipilimumab")
)

# Read mp with get_drecno(), to identify drugs without combinations

# Take the time to read the matching drugs. Did you forget a drug?

d_drecno <-
  get_drecno(d_sel_names,
             mp = mp_,
             allow_combination = FALSE,
             method = "drug_name")
d_drecno

# And DrecNos of drugs allowing for combinations

d_drecno <-
  get_drecno(d_sel = d_sel_names,
             mp = mp_,
             allow_combination = TRUE,
             method = "drug_name")
d_drecno
```

---

get\_llt\_smq

*Get low level term codes from SMQs*

---

### Description

**[Stable]** Collect llts from smq\_list and smq\_content data.tables, given an SMQ.

**Usage**

```
get_llt_smq(
  smq,
  smq_scope = c("narrow", "broad"),
  smq_list,
  smq_content,
  smq_list_content = deprecated()
)
```

**Arguments**

smq	A named list of character vector(s).
smq_scope	A character vector. One of "narrow" or "broad".
smq_list	A data.table. A list of SMQs.
smq_content	A data.table. A list of SMQs content.
smq_list_content	<b>[Deprecated]</b>

**Details**

get\_llt\_smq() is an *ID collector* function. SMQ stands for Standardized MedDRA query. get\_llt\_smq() only works with NON-algorithmic SMQs (this status is given in the smq\_list table). See [smq\\_list\\_](#) and [smq\\_content\\_](#). You can choose between the narrow and the broad scope of the SMQ. If you want to work with the SOC hierarchy, use [get\\_llt\\_soc\(\)](#).

**Value**

A named list of integers. Low-level term codes.

**See Also**

[get\\_llt\\_soc\(\)](#)

**Examples**

```
## Finding llt codes for Embolism (SMQ)

smq_sel <- rlang::list2(
  embolism = "Embolic and thrombotic events, venous (SMQ)"
)
get_llt_smq(smq_sel,
  smq_scope = "narrow",
  smq_list = smq_list_,
  smq_content = smq_content_
)

# You can query multiple SMQs in one item, and query high level SMQs
smq_sel2 <-
  rlang::list2(
```

```

    sepsis = c("Sepsis (SMQ)", "Toxic-septic shock conditions (SMQ)"),
    ischemic_heart_disease = c("Ischaemic heart disease (SMQ)")
  )

  get_llt_smq(smq_sel2,
    smq_scope = "narrow",
    smq_list = smq_list_,
    smq_content = smq_content_
  )

```

---

get\_llt\_soc

*Get low level term codes from soc classification*


---

## Description

**[Stable]** Collect llt codes from a meddra data.table, given another term of the MedDRA SOC Hierarchy.

## Usage

```

get_llt_soc(
  term_sel,
  term_level = c("soc", "hlgt", "hlt", "pt", "llt"),
  meddra,
  verbose = TRUE
)

```

## Arguments

term_sel	A named list of character vector(s). The terms to extract llts codes from. See details.
term_level	A character string. One of "soc", "hlgt", "hlt", "pt", or "llt"
meddra	A data.table. Built from meddra_builders functions
verbose	Logical. Allows you to see matching reactions in the console.

## Details

get\_llt\_soc() is an *ID collector* function. The function extracts low level term codes. get\_llt\_soc() is **case-sensitive**, and MedDRA terms always begin with a capital letter, in their native version. In term\_sel, all terms should come from the same hierarchical level, e.g. all preferred terms, all high level terms, etc.

## Value

A named list of integers. Low-level term codes.



**See Also**

`get_llt_smq()`

**Examples**

```
## Finding llt codes for colitis

pt_sel <- rlang::list2(
  colitis = c("Colitis",
              "Autoimmune colitis"),
  pneumonitis = c("Pneumonitis",
                  "Organising pneumonia")
)

hlt_sel <- rlang::list2(
  colitis = c("Gastrointestinal inflammatory disorders NEC"),
  pneumonitis = c("Pulmonary thrombotic and embolic conditions")
)

# Remember you can use more than one term to define each adverse reaction,
# but they should all be at the same hierarchical level in meddra.

# with preferred terms

get_llt_soc(
  term_sel = pt_sel,
  term_level = "pt",
  meddra = meddra_
)

# with high level terms

get_llt_soc(
  term_sel = hlt_sel,
  term_level = "hlt",
  meddra = meddra_
)
```

---

 ic\_tail

---

*Credibility interval limits for the information component*


---

**Description**

**[Stable]** Compute the Information Component credibility interval, typically the lower end of the 95% CI, also known as the IC025.

**Usage**

```
ic_tail(n_obs, n_exp, p = 0.025)
```

**Arguments**

n_obs	Number of observed cases
n_exp	Number of expected cases (see Details)
p	End of chosen credibility interval

**Details**

The ends of the credibility interval of the information component are estimated with the gamma distribution. `n_exp` is defined as `n_drug * n_event / n_total` for the basic IC (formula is different for interactions) Do not add `+.5` to `n_obs` and `n_exp` as it is automatically done in the function. By default, IC025 is computed. Change `p` for different ends. It may be easier to use [compute\\_dispro\(\)](#), which internally calls this function.

**Value**

A numeric vector. The lower end of the credibility interval

**See Also**

[compute\\_dispro\(\)](#)

**Examples**

```
ic_tail(n_obs = 12,
        n_exp = 5)
```

---

meddra\_

*Sample of MedDRA*

---

**Description**

Anonymized data from MedDRA, used to illustrate the package examples and vignettes. You can find term codes related to colitis, pneumonitis, hepatitis, a SMQ of embolisms. Compounds are `meddra_`, `smq_list_`, `smq_content_` and `smq_list_content_`. Create dedicated `.parquet` files using [tb\\_meddra\(\)](#). See examples in [get\\_llt\\_soc](#) and [get\\_llt\\_smq](#)

**Usage**

```
data(meddra_)

smq_list_content_

smq_list_

smq_content_
```

**Format**

meddra\_ is a data.table with 15 variables and 677 rows.

- The \*\_code columns. Integers. MedDRA code for the given term.
- The \*\_name columns. Characters. The name of the term.
- soc\_abbrev Character. The abbreviation of the SOC.
- null\_field Logical. Empty column.
- pt\_soc\_code Integer. The preferred term code of the SOC itself.
- primary\_soc\_fg Character. Whether the SOC is primary for this code. "Y" or "N", Yes or No.
- empty\_col Logical. Empty column.

smq\_list\_ is a data.table with 9 variables and 11 rows. It is the list of SMQ.

- smq\_code Integer. The code of the SMQ.
- smq\_name Character. The name of the SMQ.
- smq\_level Integer. The hierarchical level of the SMQ.
- smq\_description Character. The description of the SMQ.
- smq\_source Character. The source of the SMQ.
- smq\_note Character. Additional note on the SMQ.
- MedDRA\_version Numeric. The version of MedDRA.
- status Character. The status of the SMQ (active or not)
- smq\_algorithm Character. Whether the SMQ is algorithmic or not.
- empty\_col Logical. Empty column.

smq\_content\_ is a data.table with 9 variables and 3386 rows. It is the content of each SMQ.

- smq\_code Integer. The code of the SMQ.
- term\_code Integer. The low-level term code.
- term\_level Integer. The hierarchical level of the term.
- term\_scope Integer. The scope of the term (narrow 2 or broad 1)
- term\_category Character. In algorithmic SMQs, the category of the term.
- term\_weight Integer. The weight of the term (algorithmic SMQs).

- `term_status` Integer. The status of the term (active or not)
- `term_addition_version` Numeric. The version of the term addition.
- `term_last_modified_version` Numeric. The last MedDRA version the term was modified.
- `empty_col` Logical. Empty column.

`smq_list_content_` is a `data.table` with 19 variables and 3386 rows. It is a fusion of `smq_list` and `smq_content`, as created with `tb_meddra()`.

- `smq_code` Integer. The code of the SMQ.
- `smq_name` Character. The name of the SMQ.
- `smq_level` Integer. The hierarchical level of the SMQ.
- `smq_description` Character. The description of the SMQ.
- `smq_source` Character. The source of the SMQ.
- `smq_note` Character. Additional note on the SMQ.
- `MedDRA_version` Numeric. The version of MedDRA.
- `status` Character. The status of the SMQ (active or not)
- `smq_algorithm` Character. Whether the SMQ is algorithmic or not.
- `empty_col.x` Logical. Empty column.
- `term_code` Integer. The low-level term code.
- `term_level` Integer. The hierarchical level of the term.
- `term_scope` Integer. The scope of the term (narrow 2 or broad 1)
- `term_category` Character. In algorithmic SMQs, the category of the term.
- `term_weight` Integer. The weight of the term (algorithmic SMQs).
- `term_status` Integer. The status of the term (active or not)
- `term_addition_version` Numeric. The version of the term addition.
- `term_last_modified_version` Numeric. The last MedDRA version the term was modified.
- `empty_col.y` Logical. Empty column.

An object of class `data.table` (inherits from `data.frame`) with 3386 rows and 19 columns.

An object of class `data.table` (inherits from `data.frame`) with 11 rows and 9 columns.

An object of class `data.table` (inherits from `data.frame`) with 3386 rows and 9 columns.

## Source

None

## References

There is none

## Examples

```
data(meddra_)
```

mp\_

*Sample of WHODrug***Description**

A small part of WHODrug, used to illustrate the package examples and vignettes. You can find DrecNo related to immune checkpoint inhibitors, paracetamol, tramadol, tretinoin, anti-thrombin iii, and ATC classes L03AA Colony stimulating factors, C09AA ACE inhibitors, plain, J01CA Penicillins with extended spectrum. Compounds are thg\_ and mp\_. See examples in [get\\_drecno](#) and [get\\_atc\\_code](#)

**Usage**

```
data(mp_)
```

```
thg_
```

**Format**

mp\_ is a data.table with 8 variables and 14146 rows.

- MedicinalProd\_Id Integer. The medicinalproduct identifier.
- Sequence.number.1 and 2 Characters. Complement to DrecNo.
- DrecNo Character. Drug Record Number, pivotal to identify drugs with [get\\_drecno\(\)](#).
- drug\_name\_t Character. The name of the drug. Compared to the original drug\_name variable in mp table, this variable is trimmed for white spaces, and names are in lowercase.
- Create.date Character. The date the record was created.
- Date.changed Character. The date the record was last changed.
- Country Character. The country where the record was created.

thg\_ is a data.table with 5 variables and 4079 rows.

- Therapgroup\_Id Integer. The identifier of the therapeutic group.
- ATC.code Character. The ATC code of the drug.
- Create.date Character. The date the record was created.
- Official.ATC.code Character. Whether the ATC code is official (Yes/No).
- MedicinalProd\_Id Integer. The medicinalproduct identifier.

An object of class data.table (inherits from data.frame) with 4079 rows and 5 columns.

**Source**

None

## References

There is none

## Examples

```
data(mp_)
```

---

nice_p	<i>Nice printing of p-values</i>
--------	----------------------------------

---

## Description

**[Stable]** Formatting function for consistent p-value reporting.

You can choose to print the leading zero (e.g. 0.01) or not (e.g. .01) with `print_zero`.

## Usage

```
nice_p(p_val, print_zero = FALSE)
```

## Arguments

<code>p_val</code>	A numeric. The p-value to format.
<code>print_zero</code>	A logical. Should leading zero be printed? (see Details)

## Value

A character vector with the formatted p-value(s)

## Examples

```
pvals <-
  c(0.056548, 0.0002654, 0.816546, 0.0493321)
nice_p(pvals)

nice_p(pvals, print_zero = TRUE)
```

---

screen_adr	<i>Screening of adverse drug reactions</i>
------------	--

---

## Description

**[Stable]** Identify and rank the most frequently reported adverse drug reaction (ADR) terms in a dataset, based on a specified MedDRA term level. It allows users to filter terms by a frequency threshold or extract the top n most frequently occurring terms.

## Arguments

.data	An adr data.table. See <a href="#">adr_</a>
meddra	A meddra data.table. See <a href="#">meddra_</a>
term_level	A character string specifying the MedDRA hierarchy level. Must be one of "soc", "hlgt", "hlt", "pt", or "llt".
freq_threshold	A numeric value indicating the minimum frequency (as a proportion) of cases where a term must appear to be included in the results. For example, 0.05 means 5%. Defaults to NULL, meaning no threshold is applied unless top_n is different from NULL.
top_n	An integer specifying the number of most frequently occurring terms to return. Defaults to NULL. Overrides freq_threshold if both are provided.

## Details

- If freq\_threshold is set (e.g., 0.05), the function filters ADR terms appearing in at least 5% of unique reports in .data.
- If top\_n is specified, only the most frequent n terms are returned. If both freq\_threshold and top\_n are provided, only top\_n is applied (a warning is issued in such cases).
- Counts are computed at the *case* level, not the ADR level. This means frequencies reflect the proportion of unique reports (cases) where a term is mentioned, rather than the total mentions across all reports.

The function processes an ADR dataset (adr\_) and a MedDRA dataset (meddra\_) to generate results that are linked to a specific MedDRA hierarchy level (soc, hlgt, hlt, pt, or llt).

## Value

A data.frame with the following columns:

- **term:** The MedDRA term at the specified hierarchy level.
- **n:** The number of unique reports (cases) where the term appears.
- **percentage:** The percentage of total unique reports where the term appears.

The results are sorted in descending order of percentage.

## Examples

```
# Example 1: Filter terms appearing in at least 5% of reports
screen_adr(
  .data = adr_,
  meddra = meddra_,
  term_level = "pt",
  freq_threshold = 0.05
)

# Example 2: Get the top 5 most frequent terms
screen_adr(
  .data = adr_,
  meddra = meddra_,
  term_level = "hlt",
  top_n = 5
)
```

---

screen\_drug

*Screening of drugs*


---

## Description

**[Stable]** The `screen_drug()` function identifies and ranks the most frequently reported drugs (by active ingredient) in a dataset.

## Usage

```
screen_drug(.data, mp_data, freq_threshold = NULL, top_n = NULL)
```

## Arguments

<code>.data</code>	An drug data.table. See <a href="#">drug_</a>
<code>mp_data</code>	An MP data.table. See <a href="#">mp_</a>
<code>freq_threshold</code>	A numeric value indicating the minimum frequency (as a proportion) of cases where a drug must appear to be included in the results. Defaults to NULL.
<code>top_n</code>	An integer specifying the number of most frequently occurring drugs to return. Defaults to NULL.

## Details

- If `freq_threshold` is set (e.g., `0.05`), the function filters drugs appearing in at least 5% of unique reports in `.data`.
- If `top_n` is specified, only the most frequent `n` drugs are returned. If both `freq_threshold` and `top_n` are provided, only `top_n` is applied (a warning is raised in such cases).
- Counts are computed at the *case* level, not the drug mention level. This means frequencies reflect the proportion of unique reports (cases) where a drug is mentioned, rather than the total mentions across all reports.



**Value**

A data.frame with the following columns:

- Drug name: The drug name.
- DrecNo: The drug record number
- N: The number of unique reports (cases) where the drug appears.
- percentage: The percentage of total unique reports where the drug appears.

The results are sorted in descending order of percentage.

**Examples**

```
# Set up start
data.table::setDTthreads(1)

# Filter drugs appearing in at least 10% of reports
screen_drug(
  .data = drug_,
  mp_data = mp_,
  freq_threshold = 0.10
)

# Get the top 5 most reported drugs
screen_drug(
  .data = drug_,
  mp_data = mp_,
  top_n = 5
)

# nb: in the example datasets, not all drugs are recorded in mp_,
# leading to NAs in screen_drug output.

# Set up end
data.table::setDTthreads(0)
```

---

tb\_meddra

---

*Create MedDRA tables*


---

**Description**

**[Stable]** Transform MedDRA .ascii files to .parquet files

MedDRA is delivered as ascii files, that you should transform to a more efficient format. Parquet format from arrow has many advantages: It works with out-of-memory data, which makes it possible to process tables on a computer with not-so-much RAM. It is also lightweighted and standard across different languages. The function also creates variables in each table. You should note that NOT all MedDRA tables are processed with this function. Three tables are created: meddra\_hierarchy, that respects the System Organ Class hierarchic classification. smq\_list and smq\_content for Standardized MedDRA Queries. **Caution** There tends to be small variations in the MedDRA ascii files structure. Last verified version on which this function is working is **26.1**. Use `dt_parquet()` to load the tables afterward.

Usage

```
tb_meddra(path_meddra)
```

Arguments

path\_meddra      Character string, a directory containing MedDRA ascii tables. It is also the output directory.

Value

.parquet files into the path\_meddra directory. Three tables: meddra\_hierarchy, smq\_list, and smq\_content. Some columns are returned as integer (all \*\_code columns). All other columns are character.

See Also

[tb\\_vigibase\(\)](#), [tb\\_who\(\)](#), [tb\\_subset\(\)](#), [dt\\_parquet\(\)](#)

Examples

```
# Use the examples from tb_main if you want to see these functions in action.

path_meddra <- paste0(tempdir(), "/meddra_directory/")
dir.create(path_meddra)
create_ex_meddra_asc(path_meddra)

tb_meddra(path_meddra = path_meddra)

# Clear temporary files when you're done
unlink(path_meddra, recursive = TRUE)
```

---

tb_subset	<i>Extract of subset of Vigibase</i>
-----------	--------------------------------------

---

Description

**[Stable]** Create a subset of the VigiBase ECL datasets

Usage

```
tb_subset(
  wd_in,
  wd_out,
  subset_var = c("drecno", "medprod_id", "meddra_id", "age"),
  sv_selection,
  rm_suspdup = TRUE
)
```

**Arguments**

wd_in	Source directory pathway (character)
wd_out	Output directory pathway (character)
subset_var	One of "drecno", "medprod_id", "meddra_id", "age"
sv_selection	A named list or a vector containing the appropriate ids (according to the method, see details)
rm_suspdup	A logical. Should suspected duplicates be removed? TRUE by default

**Details**

You must select a subset variable with `subset_var` and provide an appropriate list according to this variable in `sv_selection`. Available `subset_var` :

- `drecno` will use Drug Record Number (DrecNo), from WHO Drug, and will subset from drug (see [get\\_drecno\(\)](#)).
- `medprod_id` will use MedicinalProd\_Id, also from drug. May be useful if requesting from ATC classes. (see [get\\_atc\\_code\(\)](#)).
- `meddra_id` will use MedDRA\_Id, subset from adr. (see [get\\_llt\\_soc\(\)](#) or See [get\\_llt\\_smq\(\)](#)).
- `age` will use AgeGroup from demo. See below.

Age groups ids are as follows:

- 1 0 - 27 days
- 2 28 days to 23 months
- 3 2 - 11 years
- 4 12 - 17 years
- 5 18 - 44 years
- 6 45 - 64 years
- 7 65 - 74 years
- 8 >= 75 years
- 9 Unknown

Example: To work with patients aged 18 to 74, provide `c(5, 6, 7)` as `sv_selection`.

Use [dt\\_parquet\(\)](#) to load the tables afterward.

**Value**

Parquet files in the output directory. All files from a vigibase ECL main folder are returned subsetting (including suspected duplicates).

**See Also**

[get\\_drecno\(\)](#), [get\\_atc\\_code\(\)](#), [get\\_llt\\_soc\(\)](#), [get\\_llt\\_smq\(\)](#), [dt\\_parquet\(\)](#)

## Examples

```
# --- technical steps ---- #

wd_in <- paste0(tempdir(), "/", "tbsubsetex")
dir.create(wd_in)
create_ex_main_pq(wd_in)

# Select a subset_var and corresponding data

# Subset on adr colitis codes

adr_llt <-
  list(
    colitis = "Colitis"
  ) |>
  get_llt_soc(term_level = "pt", meddra_, verbose = FALSE)

wd_out <- paste0(wd_in, "/", "colitis_subset", "/")

tb_subset(wd_in, wd_out,
          subset_var = "meddra_id",
          sv_selection = adr_llt)

# Subset on drug codes

d_drecno <-
  list(
    ipi = "ipilimumab" ) |>
  get_drecno(mp = mp_, verbose = FALSE)

wd_out <- paste0(wd_in, "/", "nivolumab_subset", "/")

tb_subset(wd_in, wd_out,
          subset_var = "drecno",
          sv_selection = d_drecno)

# Subset on age > 65 year-old

sv_selection <-
  c(7, 8)

wd_out <- paste0(wd_in, "/", "more_than_65_subset", "/")

tb_subset(wd_in, wd_out,
          subset_var = "age",
          sv_selection = sv_selection)

unlink(wd_in, recursive = TRUE)
```

---

tb_vigibase	<i>Create Vigibase ECL tables</i>
-------------	-----------------------------------

---

## Description

**[Stable]** Transform Vigibase .txt files to .parquet files.

## Usage

```
tb_vigibase(
  path_base,
  path_sub,
  force = FALSE,
  rm_suspdup = TRUE,
  overwrite_existing_tables = FALSE
)
```

## Arguments

path_base	Character string, a directory containing vigibase txt tables. It is also the output directory.
path_sub	Character string, a directory containing subsidiary tables.
force	Logical, to be passed to <code>cli::cli_progress_update()</code> . Used for internal purposes.
rm_suspdup	Logical, should suspected duplicates (from SUSPECTEDDUPLICATES.txt) be removed from main tables? Default is TRUE. Set to FALSE to keep all cases, including suspected duplicates.
overwrite_existing_tables	Logical, should existing parquet tables be overwritten? Default is FALSE.

## Details

Vigibase Extract Case Level is delivered as zipped text files, that you should transform to a more efficient format. Parquet format from arrow has many advantages: It works with out-of-memory data, which makes it possible to process Vigibase tables on a computer with not-so-much RAM. It is also lightweighted and standard across different languages. The function also creates variables in each table. The `suspectedduplicates` table will be added to the base directory. Use `dt_parquet()` to load the tables afterward.

The argument `overwrite_existing_tables` is especially useful if the function crashes or is interrupted: it allows you to resume the process without rebuilding tables that were already successfully created. If set to FALSE (the default), the function will skip the construction of any .parquet tables that already exist, so you do not have to start from scratch after a failure. Set to TRUE to force rebuilding all tables.

**Value**

- .parquet files of all main tables into the path\_base directory: demo, adr, drug, link, ind, out, srce, followup, and the suspdup (suspected duplicates) table. Check ?demo\_ for more information on the tables.
- The link table is augmented with tto\_mean and range, to analyze time to onset according to WHO's recommendations (see vignette("descriptive")).
- .parquet files of all other subsidiary tables into the path\_sub directory: AgeGroup, Dechallenge, Dechallenge2, Frequency, Gender, Notifier, Outcome, Rechallenge, Rechallenge2, Region, RepBasis, ReportType, RouteOfAdm, Seriousness, and SizeUnit.

.parquet files into the path\_base directory (**including suspected duplicates tables**). Some columns are returned as integer (UMCReportId, Drug\_Id, MedicinalProd\_Id, Adr\_Id, MedDRA\_Id), and some columns as numeric (TimeToOnsetMin, TimeToOnsetMax) All other columns are character.

**See Also**

[tb\\_who\(\)](#), [tb\\_meddra\(\)](#), [tb\\_subset\(\)](#), [dt\\_parquet\(\)](#)

**Examples**

```
# --- Set up example source files ---- #####

path_base <- paste0(tempdir(), "/", "main", "/")

path_sub  <- paste0(tempdir(), "/", "sub", "/")

dir.create(path_base)
dir.create(path_sub)

create_ex_main_txt(path_base)
create_ex_sub_txt(path_sub)

# ---- Running tb_vigibase

tb_vigibase(path_base = path_base,
            path_sub  = path_sub)

# Clear temporary files when you're done
unlink(path_base, recursive = TRUE)
unlink(path_sub, recursive = TRUE)
```

## Description

**[Stable]** Transform Vigibase WHO .txt files to .parquet files

WHODrug is delivered as zipped text files folder, that you should transform to a more efficient format. Parquet format from arrow has many advantages: It can work with out-of-memory data, which makes it possible to process tables on a computer with not-so-much RAM. It is also lightweighted and standard across different languages. The function also creates variables in each table. See [tb\\_vigibase\(\)](#) for some running examples, and try `?mp_` or `?thg_` for more details. Use [dt\\_parquet\(\)](#) to load the tables afterward.

## Usage

```
tb_who(path_who, force = FALSE)
```

## Arguments

path_who	Character string, a directory containing whodrug txt tables. It is also the output directory.
force	Logical, to be passed to <code>cli::cli_progress_update()</code> . Used for internal purposes.

## Value

.parquet files into the path\_who directory. Some columns are returned as integer (all Id columns, including MedicinalProd\_Id, with notable exception of DrecNo), and some columns as numeric (Quantity from ingredient table) All other columns are character.

## See Also

[tb\\_vigibase\(\)](#), [tb\\_meddra\(\)](#), [tb\\_subset\(\)](#), [dt\\_parquet\(\)](#)

## Examples

```
# Use the examples from tb_main if you want to see these functions in action.

path_who <- paste0(tempdir(), "/whodrug_directory/")
dir.create(path_who)
create_ex_who_txt(path_who)

tb_who(path_who = path_who)

# Clear temporary files when you're done
unlink(path_who, recursive = TRUE)
```

vigi\_routine

*Display routine pharmacovigilance summary***Description**

**[Maturing]** `vigi_routine()` draws an Information Component plot and a Time to Onset plot for a given drug-adr pair.

**Usage**

```
vigi_routine(
  demo_data,
  drug_data,
  adr_data,
  link_data,
  d_code,
  a_code,
  case_tto = NULL,
  vigibase_version,
  analysis_setting = "All reports",
  d_label = NULL,
  a_label = NULL,
  export_to = NULL,
  suspect_only = FALSE,
  d_code_2
)
```

**Arguments**

<code>demo_data</code>	A demo data.table.
<code>drug_data</code>	A drug data.table.
<code>adr_data</code>	An adr data.table.
<code>link_data</code>	A link data.table.
<code>d_code</code>	A named list. The drug code(s) to be used. There must be only one item in <code>d_code</code> .
<code>a_code</code>	A named list. The adr code(s) to be used. There must be only one item in <code>a_code</code> .
<code>case_tto</code>	A numeric. The time to onset of the studied case. See details.
<code>vigibase_version</code>	A character. The version of VigiBase used (e.g. "September 2024"). This is passed to the plot legend.
<code>analysis_setting</code>	A character. The setting of the analysis. See details.
<code>d_label</code>	A character. The name of the drug, as passed to the plot legend. Defaults to <code>names(d_code)</code> .



a_label	A character. The name of the adr, as passed to the plot legend. Defaults to names(a_code).
export_to	A character. The path to export the plot. If NULL, the plot is not exported. Should end by ".eps", ".ps", ".tex" (pictex), ".pdf", ".jpeg", ".tiff", ".png", ".bmp", ".svg" or ".wmf" (windows only).
suspect_only	Logical. If TRUE, only cases where the drug is suspected are used for IC analysis. If FALSE (default), all cases are used.
d_code_2	Optional named list. A second drug code for dual drug analysis. If provided, a single analysis is performed on cases exposed to both drugs.

## Details

See vignette("routine\_pharmacovigilance") for examples. The output can be exported. Time to onset data are bounded between 1 day and 10 years. Data outside this range are reassigned a 1 day and 10 years value, respectively. The function only works if there is one item in d\_code and a\_code. If you are working on a specific case, you can provide a case\_tto value. This value will be displayed on the Time to Onset plot. If your demo table was filtered on specific cases (e.g. older adults, a subset of all drugs), then you may want to indicate this setting on the plot legend, with arg analysis\_setting.

## Value

A ggplot2 graph, with two panels. The first panel, on top, is the Information Component (IC) plot. The arrow and "IC025 label" indicate the IC value for the selected drug-adr pair. The second panel, on the bottom, is the Time to Onset (TTO) density plot. It is derived only of cases where the drug was **suspected** to be responsible of the adr (irrespective of the suspect\_only argument). If you provide a case\_tto value, it is represented by the red line, and the label.

## Examples

```
# Say you want to perform a disproportionality analysis between colitis and
# nivolumab among ICI cases

# identify drug DrecNo, and adr LLT code

d_drecno <-
  ex$d_drecno["nivolumab"]

a_llt <-
  ex$a_llt["a_colitis"]

# But you could also use get_drecno() and get_llt_soc()

# load tables demo, drug, adr, and link (real tables with
# open_dataset() or dt_parquet("x", in_memory = FALSE))

demo <- demo_
adr <- adr_
drug <- drug_
link <- link_
```

```
# run routine

vigi_routine(
  demo_data = demo,
  drug_data = drug,
  adr_data = adr,
  link_data = link,
  d_code = d_drecno,
  a_code = a_llt,
  vigibase_version = "September 2024"
)

# if you're working on a case, you can provide his/her time to onset
# with arg `case_tto`

vigi_routine(
  case_tto = 150,
  demo_data = demo,
  drug_data = drug,
  adr_data = adr,
  link_data = link,
  d_code = d_drecno,
  a_code = a_llt,
  vigibase_version = "September 2024"
)

# Customize with d_name and a_name, export the plot with export_to

vigi_routine(
  case_tto = 150,
  demo_data = demo,
  drug_data = drug,
  adr_data = adr,
  link_data = link,
  d_code = d_drecno,
  a_code = a_llt,
  vigibase_version = "September 2024",
  d_label = "Nivolumab",
  a_label = "Colitis",
  export_to = paste0(tempdir(), "/", "vigicaen_graph.png")
)
```

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