Package ‘interfacer’

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Title Define and Enforce Contracts for Dataframes as Function Parameters

Version 0.2.3

Description A dataframe validation framework for package builders who use dataframes as function parameters. It performs checks on column names, coerces data-types, and checks grouping to make sure user inputs conform to a specification provided by the package author. It provides a mechanism for package authors to automatically document supported dataframe inputs and selectively dispatch to functions depending on the format of a dataframe much like S3 does for classes. It also contains some developer tools to make working with and documenting dataframe specifications easier. It helps package developers to improve their documentation and simplifies parameter validation where dataframes are used as function parameters.

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as.list.iface Cast an iface to a plain list.

Description
Cast an iface to a plain list.

Usage
## S3 method for class 'iface'
as.list(x, ..., flatten = FALSE)

Arguments
x
...  
flatten

Value
a list representation of the iface input.

Examples
my_iface = iface(
  col1 = integer + group_unique ~ "an integer column"
)

as.list(my_iface, flatten=TRUE)

format.iface Format an iface specification for printing

Description
Format an iface specification for printing

Usage
## S3 method for class 'iface'
format(x, ...)


iclip

Arguments

| x | an iface specification |
| ... | not used. |

Value

a formatted string representation of an iface

Examples

```r
my_iface = iface(
  col1 = integer + group_unique ~ "an integer column"
)

print(my_iface)
knitr::knit_print(my_iface)
```

---

Create an iface specification from an example dataframe

Description

When developing with interfacer it is useful to be able to base a function input off a prototype that you are for example using as testing. This function generates an interfacer::iface specification for the supplied data frame and copies it to the clipboard so that it can be pasted into the package code you are working on.

Usage

```r
iclip(df, df_name = deparse(substitute(df)))
```

Arguments

| df | a prototype dataframe |
| df_name | an optional name for the parameter (defaults to i_<df name>) |

Details

If the dataframe contains one or more list columns with nested dataframes the nested dataframes are also defined using a second iface specification.

Value

nothing, populates clipboard

Examples

```r
if (interactive()) iclip(iris)
```
iconvert

Convert a dataframe to a format compatible with an interface specification

Description

This function is called by invalidate and is not generally intended to be used directly by the end user. It may be helpful in debugging during package development to interactive test a iface spec. iconvert is an interactive version of invalidate.

Usage

iconvert(
  df,
  iface,
  .imap = interfacer::imapper(),
  .dname = "<unknown>",
  .fname = "<unknown>",
  .has_dots = TRUE,
  .prune = FALSE,
  .env = rlang::current_env()
)

Arguments

df the dataframe to convert
iface the interface spec as an iface
.imap an optional imapper mapping
dname the name of the parameter value (optional).
.fname the name of the function (optional).
.has_dots internal library use only. Changes the nature of the error message.
.prune do you want to remove non matching columns?
.env internal use only

Value

the input dataframe coerced to be conformant to the iface specification, or an informative error is thrown.

Examples

i_diamonds = iface(
  color = enum(D,E,F,G,H,I,J,extra) ~ "the colour",
  price = integer ~ "the price"
)
idispatch

Dispatch to a named function based on the characteristics of a dataframe

Description

This provides a dataframe analogy to S3 dispatch.

Usage

idispatch(x, ..., .default = NULL)

Arguments

x: a dataframe

...: a set of function name=interfacer::iface pairs

.default: a function to apply in the situation where none of the rules can be matched. The default results in an error being thrown.

Details

If multiple possible dataframe formats are possible for a function, each with different processing requirements, then the choice of function can be made based on matching the input dataframe to a set of iface specifications. The first matching iface specification determines which function is used for dispatch.

Value

the result of dispatching the dataframe to the first function that matches the rules in .... Matching is permissive in that the test is passed if a dataframe can be coerced to the iface specified format.

Examples

i1 = iface( col1 = integer ~ "An integer column" )
i2 = iface( col2 = integer ~ "A different integer column" )

# this is an example function that would typically be inside a package, and
# is exported from the package.
extract_mean = function(df, ...) {
  idispatch(df,
    extract_mean.i1 = i1,
    extract_mean.i2 = i2
  )
}

iconvert(ggplot2::diamonds, i_diamonds,.prune = TRUE)
# this is expected to be an internal package function
# the naming convention here is based on S3 but it is not required
extract_mean.i1 = function(df = i1, ...) {
  message("using i1")
  # input validation is not required in functions that are being called using
  # `idispatch` as the validation occurs during dispatch.
  mean(df$col1)
}

extract_mean.i2 = function(df = i2, uplift = 1, ...) {
  message("using i2")
  mean(df$col2)+uplift
}

# this input matches `i1` and the `extract_mean` call is dispatched
# via `extract_mean.i1`
test = tibble::tibble( col2 = 1:10 )
exttract_mean(test, uplift = 50)

# this input matches `i2` and the `extract_mean` call is dispatched
# via `extract_mean.i2`
test2 = tibble::tibble( col1 = 1:10 )
exttract_mean(test2, uplift = 50)

# This input does not match any of the allowable input specifications and
# generates an error.
test3 = tibble::tibble( wrong_col = 1:10 )
try(extract_mean(test3, uplift = 50))

---

**idocument**  
*Document an interface contract for inserting into roxygen2*

**Description**

This function is expected to be called within the documentation of a function as inline code in the parameter documentation of the function. It details the expected columns that the input dataframe should possess. This has mostly been superseded by the @iparam <name> <description> roxygen2 tag which does this automatically, however in some circumstances (particularly multiple dispatch) you may want to assemble dataframe documentation manually.

**Usage**

idocument(fn, param = NULL)

**Arguments**

- fn  
  the function that you are documenting
- param  
  the parameter you are documenting (optional. if missing defaults to the first argument of the function)
Value

a markdown snippet

Examples

```r
# @param df `r idocument(x, df)`
x = function(df = iface( col1 = integer ~ "an integer column" )) {} 
cat(idocument(x, df))
```

### iface

**Construct an interface specification**

**Description**

An iface specification defines the expected structure of a dataframe, in terms of the column names, column types, grouping structure and uniqueness constraints that the dataframe must conform to. A dataframe can be tested for conformance to an iface specification using iconvert.

**Usage**

```r
iface(..., .groups = NULL, .default = NULL)
```

**Arguments**

`...` The specification of the interface (see details), or an unnamed iface object to extend, or both.

`.groups` either FALSE for no grouping allowed or a formula of the form `~ var1 + var2 + ...` which defines what columns must be grouped in the dataframe (and in which order). If NULL (the default) then any grouping is permitted. If the formula contains a dot e.g. `~ . + var1 + var2` then the grouping must include `var1` and `var2` but other groups are also allowed.

`.default` a default value to supply if there is nothing given in a function parameter using the iface as a formal. This is either NULL in which case there is no default, TRUE in which case the default is a zero row dataframe conforming to the specification, or a provided dataframe, which is checked to conform, and used as the default.

**Details**

An iface specification is designed to be used to define the type of a parameter in a function. This is done by using the iface specification as the default value of the parameter in the function definition. The definition can then be validated at runtime by a call to ivalidate(...) inside the function. When developing a function output an iface specification may also be used in ireturn(...) to enforce that the output of a function is correct.

iface definitions can be printed and included in roxygen2 documentation and help us to document input dataframe parameters and dataframe return values in a standardised way by using the @iparam roxygen2 tag.
iface specifications are defined in the form of a named list of formulae with the structure `column_name = type ~ "documentation"`. 

- type can be one of anything, character, complete, date, default, double, enum, factor, finite, group_unique, in_range, integer, logical, not_missing, numeric, of_type, positive_double, positive_integer, proportion (e.g. `enum(level1,level2,...)`, `in_range(min,max)`) or alternatively anything that resolves to a function e.g. `as.ordered`.

- If type is a function name, then the function must take a single vector parameter and return a single vector of the same size. The function must also return a zero length vector of an appropriate type if passed `NULL`.

- type can also be a concatenation of rules separated by `+`, e.g. `integer + group_unique` for an integer that is unique within a group.

**Value**

- the definition of an interface as a `iface` object

**Examples**

```r
# test_df = tibble::tibble(
#   grp = c(rep("a",10),rep("b",10)),
#   col1 = c(1:10,1:10)
# ) %>% dplyr::group_by(grp)

my_iface = iface(
  col1 = integer + group_unique ~ "an integer column",
  .default = test_df
)

print(my_iface)
# the function x defines a formal `df` with default value of `my_iface`
# this default value is used to validate the structure of the user supplied
# value when the function is called.
x = function(df = my_iface, ...) {
  df = ivalidate(df,...)
  return(df)
}

# this works
x(tibble::tibble(col1 = c(1,2,3)))

# this fails as x is of the wrong type
try(x(tibble::tibble(col1 = c("a","b","c"))))

# this fails as x has duplicates
try(x(tibble::tibble(col1 = c(1,2,3,3))))

# this gives the default value
x()
```
my_iface2 = iface(
    first_col = numeric ~ "column order example",
    my_iface,
    last_col = character ~ "another col", .groups = ~ first_col + col1
)  
print(my_iface2)

my_iface_3 = iface(
    col1 = integer + group_unique ~ "an integer column",
    .default = test_df_2
)  
x = function(d = my_iface_3) {invalidate(d)}

# Doesn't work as test_df_2 hasn't been defined
try(x())

test_df_2 = tibble::tibble(
    grp = c(rep("a",10),rep("b",10)),
    col1 = c(1:10,1:10)
)  
%>% dplyr::group_by(grp)

# now it works as has been defined
x()

# it still works as default has been cached.
rm(test_df_2)

x()

---

if_col_present

Execute a function or return a value if a column in present in a dataframe

Description

The simple use case. For more complex behaviour see switch_pipeline().

Usage

if_col_present(df, col, if_present, if_missing = ~.x)

Arguments

- **df**: a dataframe
- **col**: a column name
- **if_present**: a purrr style function to execute on the dataframe if the column is present (or a plain value)
- **if_missing**: a purrr style function to execute on the dataframe if the column is missing (or a plain value)
**Value**

either the value of `if_present/if_absent` or the result of calling `if_present/if_absent` as functions on `df`.

**Examples**

```r
iris %>% if_col_present(Species, ~ .x %>% dplyr::rename(new = Species)) %>%
  colnames()
```

# in contrast to `purrr` absolute values are not interpreted as function names

```r
iris %>% if_col_present(Species2, "Yes", "No")
```

---

**Description**

This function is designed to be used by a package author within an enclosing function. The enclosing function is assumed to take as input a dataframe and have an `iface` specified for that dataframe.

**Usage**

```r
igroup_process(df = NULL, fn, ...)
```

**Arguments**

- `df`  
  a dataframe from an enclosing function in which the grouping may or may not have been correctly supplied.

- `fn`  
  a function to call with the correctly grouped dataframe as specified by the `iface` in the enclosing function.

- `...`  
  passed onto `iconvert` this could be used to supply `.prune` parameters. triple dot parameters in the enclosing function will be separately handled and automatically passed to `fn` so in general should not be passed to `igroup_process` as an intermediary although it probably won’t hurt. This behaviour is similar to `NextMethod` in S3 method dispatch.

**Details**

This function detects when the grouping of the input has additional groups over and above those in the specification and intercepts them, regrouping the dataframe and applying `fn` group-wise using an equivalent of a `dplyr::group_modify`. The parameters provided to the enclosing function will be passed to `fn` and they should have compatible method signatures.

**Value**

the result of calling `fn(df, ...)` on each unexpected group
Examples

# This specification requires that the dataframe is grouped only by the color
# column
i_diamond_price = interfacer::iface(
    color = enum(`D`, `E`, `F`, `G`, `H`, `I`, `J`, .ordered=TRUE) ~ "the color column",
    price = integer ~ "the price column",
    .groups = ~ color
)

# An example function which would be exported in a package
ex_mean = function(df = i_diamond_price, extra_param = ".") {
    # When called with a dataframe with extra groups `igroup_process` will
    # regroup the dataframe according to the structure
    # defined for `i_diamond_price` and apply the inner function to each group
    # after first calling `ivalidate` on each group.
    igroup_process(df,
        # the real work of this function is provided as an anonymous inner
        # function (but can be any other function e.g. package private function)
        # or a purrr style lambda.
        function(df, extra_param) {
            message(extra_param, appendLF = FALSE)
            return(df %>% dplyr::summarise(mean_price = mean(price)))
        }
    )
}

# The correctly grouped dataframe. The `ex_mean` function calculates the mean
# price for each `color` group.
ggplot2::diamonds %>%
    dplyr::group_by(color) %>%
ex_mean(extra_param = "without additional groups...") %>%
dplyr::glimpse()

# If an additionally grouped dataframe is provided by the user. The `ex_mean`
# function calculates the mean price for each `cut`, `clarity`, and `color`
# combination.

ggplot2::diamonds %>%
dplyr::group_by(cut, color, clarity) %>%
ex_mean() %>%
dplyr::glimpse()

# The output of this is actually grouped by cut then clarity as
# color is consumed by the igroup_dispatch summarise.
Specify mappings that can make dataframes compatible with an iface specification

Description

When a function uses `ivalidate` internally to check a dataframe conforms to the input it can attempt to rescue an incorrectly formatted dataframe. This is a pretty advanced idea and is not generally recommended.

Usage

```
imapper(...)```

Arguments

```
...               a set of `dplyr::mutate()` specifications that when applied to a dataframe will rename or otherwise fix missing columns
```

Details

This function is expected to be used only in the context of a `.imap = imapper(...)` parameter to an `ivalidate` call to make sure that certain columns are present or are a set value. Anything provided here will overwrite existing dataframe columns and its use is likely to make function behaviour obtuse. It may be deprecated in the future. The `...` input expressions should almost certainly check for the values already existing before overwriting them.

If you are considering using this for replacing missing values check using the `default(...)` iface type definition instead.

Value

a set of mappings

Examples

```
x = function(df = iface(col1 = integer ~ "an integer column" ), ...) {
  df = ivalidate(df,...)
}
input=tibble::tibble(col2 = c(1,2,3))
# This fails because col1 is missing
try(x(input))
# This fixes it for this input
x(input, .imap=imapper(col1 = col2))
```
iproto

Generate a zero length dataframe conforming to an iface specification

Description

This function is used internally for default values for a dataframe parameter. It generates a zero length dataframe that conforms to an iface specification, in terms of column names, data types and groupings. Such a dataframe is not guaranteed to be fully conformant to the iface specification if, for example, completeness constraints are applied.

Usage

iproto(iface)

Arguments

iface the specification

Value

a dataframe conforming to iface

Examples

i = interfacer::iface(
  col1 = integer ~ "A number",
  col2 = character ~ "A string"
)
iproto(i)

ireturn

Check a return parameter from a function

Description

This is intended to be used within a function to check the validity of a dataframe being returned from a function against an iface which is provided.

Usage

ireturn(df, iface, .prune = FALSE)
is.iface

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>df</td>
<td>a dataframe - if missing then the first parameter of the calling function is assumed to be a dataframe.</td>
</tr>
<tr>
<td>iface</td>
<td>the interface specification that df should conform to.</td>
</tr>
<tr>
<td>.prune</td>
<td>get rid of excess columns that are not in the spec.</td>
</tr>
</tbody>
</table>

Value

a dataframe based on df with validity checks passed, data-types coerced, and correct grouping applied to conform to iface

Examples

```r
input = iface(col_in = integer ~ "an integer column" )
output = iface(col_out = integer ~ "an integer column" )

x = function(df = input, ...) {
  df = invalidate(...)
  tmp = df %>% dplyr::rename(col_out = col_in)
  return(tmp, output)
}
x(tibble::tibble(col_in = c(1,2,3)))
output
```

is.iface

Check if an object is an interface specification

Description

Check if an object is an interface specification

Usage

```r
is.iface(x, ...)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>x</td>
<td>the object to check</td>
</tr>
<tr>
<td>...</td>
<td>ignored</td>
</tr>
</tbody>
</table>

Value

a boolean.
is_col_present  
*Check for existence of a set of columns in a dataframe*

**Description**

Check for existence of a set of columns in a dataframe

**Usage**

    is_col_present(df, ...)

**Arguments**

- **df**  
a dataframe to test
- **...**  
the column names (unquoted)

**Value**

TRUE if the columns are all there, false otherwise

**Examples**

    is_col_present(iris, Species, Petal.Width)

---

itest  
*Test dataframe conformance to an interface specification.*

**Description**

ivalidate throws errors deliberately however sometimes dealing with invalid input may be desirable. itest is generally designed to be used within a function which specifies the expected input using iface, and allows the function to test if its given input is conformant to the interface.

**Usage**

    itest(df = NULL, iface = NULL, .imap = imapper())

**Arguments**

- **df**  
a dataframe to test. If missing the first parameter of the calling function is assumed to be the dataframe to test.
- **iface**  
an interface specification produced by iface(). If missing this will be inferred from the current function signature.
- **.imap**  
an optional mapping specification produced by imapper()
ivalidate

Value

TRUE if the dataframe is conformant, FALSE otherwise

Examples

```r
if (rlang::is_installed("ggplot2")) {
  i_diamonds = iface(
    color = enum(D,E,F,G,H,I,J,extra) ~ "the colour",
    price = integer ~ "the price"
  )

  # Ad hoc testing
  itest(ggplot2::diamonds, i_diamonds)

  # Use within function:
  x = function(df = i_diamonds) {
    if(itest()) message("PASS!")
  }

  x(ggplot2::diamonds)
}
```

ivalidate

Perform interface checks on dataframe inputs using enclosing function formal parameter definitions

Description

ivalidate(...) is intended to be used within a function to check the validity of a data frame parameter (usually the first parameter) against an ispec which is given as a default value of a formal parameter.

Usage

```r
ivalidate(df = NULL, ..., .imap = imapper(), .prune = FALSE, .default = NULL)
```

Arguments

df a dataframe - if missing then the first parameter of the calling function is assumed to be a dataframe.

... not used but ivalidate should be included in call to inherit .imap from the caller function.

.imap a set of mappings as an imapper object.

.prune get rid of excess columns that are not in the spec.

.default a default dataframe conforming to the specification. This overrides any defaults defined in the interface specification
Value

a dataframe based on \( df \) with validity checks passed and \( .imap \) mappings applied if present

Examples

```r
x = function(df = iface(col1 = integer ~ "an integer column"), ...) {
  df = invalidate(...)
  return(df)
}
input = tibble::tibble(col1 = c(1,2,3))
x(input)

# This fails because col1 is not coercable to integer
input2 = tibble::tibble(col1 = c(1.5,2,3))
try(x(input2))
```

---

### knit_print.iface

Format an iface specification for printing

**Description**

Format an iface specification for printing

**Usage**

```r
knit_print.iface(x, ...)
```

**Arguments**

- **x**: an iface specification
- **...**: not used.

**Value**

A formatted string representation of an iface

**Examples**

```r
my_iface = iface(
  col1 = integer + group_unique ~ "an integer column"
)

print(my_iface)
knitr::knit_print(my_iface)
```
Format an iface specification for printing

Usage

## S3 method for class 'iface'
print(x, ...)

Arguments

x an iface specification

... not used.

Value

a formatted string representation of an iface

Examples

my_iface = iface(
  col1 = integer + group_unique ~ "an integer column"
)

print(my_iface)
knitr::knit_print(my_iface)

Parser for @iparam tags

Description

The @iparam <name> <description> tag can be used in roxygen2 documentation of a function to describe a dataframe parameter. The function must be using interfacer::iface to define the input dataframe parameter format. The @iparam tag will then generate documentation about the type of dataframe the function is expecting.

Usage

## S3 method for class 'roxy_tag_iparam'
roxy_tag_parse(x)
Arguments

x A tag

Value

a roxygen2 object with the val field set to the parsed value

Examples

# This provides support to `roxygen2` and only gets executed in the context
# of `devtools::document()`. There is no interactive use of this function.

Description

The @iparam tag can be used in roxygen2 documentation of a function to describe a dataframe parameter. The function must be using interfacer::iface to define the input dataframe parameter format. The @iparam tag will then generate documentation about the type of dataframe the function is expecting.

Usage

## S3 method for class 'roxy_tag_iparam'
roxy_tag_rd(x, base_path, env)

Arguments

x The tag
base_path Path to package root directory.
env Environment in which to evaluate code (if needed)

Value

an roxygen2::rd_section (see roxygen2 documentation)

Examples

# An example function definition:
fn_definition <- "
# This is a title
# This is the description.
# @iparam df the input"
switch_pipeline

# @export
f <- function(df = interfacer::iface(
    id = integer ~ "an integer `ID`",
    test = logical ~ "the test result"
)) {
  invalidate(df)
}

# For this example we manually parse the function specification in `fn_definition`
# creating a .Rd block - normally this is done by `roxygen2` which then
# writes this to an .Rd file. This function is not intended to be used
# outside of a call to `devtools::document`.

tmp = roxygen2::parse_text(fn_definition)
print(tmp)

---

**switch_pipeline**  
*Branch a dplyr pipeline based on a set of conditions*

**Description**

Branch a dplyr pipeline based on a set of conditions

**Usage**

`switch_pipeline(.x, ...)`

**Arguments**

- `.x`  
a dataframe
- `...`  
a list of formulae of the type `predicate ~ purrr function` using `.x` as the single parameter

**Value**

the result of applying `purrr` function to `.x` in the case where `predicate` evaluates to true. Both `predicate` and `function` can refer to the pipeline dataframe using `.x`

**Examples**

```r
iris %>% switch_pipeline(  
is_col_present(.x, Species) ~ .x %>% dplyr::rename(new = Species)  
) %>% dplyr::glimpse()
```
**type.anything**  
*Coerce to an unspecified type*

**Description**
Coerce to an unspecified type

**Usage**
type.anything(x)

**Arguments**
x any vector

**Value**
the input (unless x is NULL in which case a character())

**type.character**  
*Coerce to a character.*

**Description**
Coerce to a character.

**Usage**
type.character()

**Value**
the input as a character.
type.complete

**Description**

This test checks either for factors that all factor levels are present in the input, or for numerics if the sequence from minimum to maximum by the smallest difference are not all (approximately) present. Empty values are ignored.

**Usage**

```
type.complete(x)
```

**Arguments**

- `x` any vector, factor or numeric

**Value**

the input or error if not complete

---

**type.date**

**Description**

Coerce to a Date.

**Usage**

```
type.date(x, ...)
```

**Arguments**

- `x` an object to be converted.
- `...` further arguments to be passed from or to other methods.

**Value**

the input as a date vector, error if this would involve data loss.
**type.default**

*Set a default value for a column*

**Description**

Any NA values will be replaced by this value. N.b. default values must be provided before any other rules if the validation is not to fail.

**Usage**

type.default(value)

**Arguments**

value a length one item of the correct type.

**Value**

a validation function that switches NAs for default values

**type.double**

*Coerce to a double.*

**Description**

Coerce to a double.

**Usage**

type.double()

**Value**

the input as a double, error if this would involve data loss.
type.enum

*Define a conformance rule to match a factor with specific levels.*

**Description**
Define a conformance rule to match a factor with specific levels.

**Usage**

```
type.enum(..., .drop = FALSE, .ordered = FALSE)
```

**Arguments**

- `...`: the levels (no quotes, backticks if required)
- `.drop`: should levels present in the data and not specified cause an error (FALSE the default) or be silently dropped to NA values (TRUE).
- `.ordered`: must the factor be ordered

**Value**

A function that can check and convert input into the factor with specified levels. This will re-level factors with matching levels but in a different order.

**Examples**

```
f = type.enum(one,two,three)
f(c("three","two","one"))
f(factor(rep(1:3,5), labels = c("one","two","three")))
```

type.factor

*Coerce to a factor.*

**Description**

Coerce to a factor.

**Usage**

```
type.factor(x)
```

**Arguments**

- `x`: any vector

**Value**

The input as a factor, error if this would involve data loss.
**type.finite**  
*Check for non-finite values*

**Description**

Any non finite values will cause failure of validation.

**Usage**

```r
type.finite(x)
```

**Arguments**

- `x`  
  any vector that can be coerced to numeric

**Value**

the input coerced to a numeric value, or an error if any non-finite values detected

---

**type.group_unique**  
*Coerce to a unique value.*

**Description**

Coerce to a unique value.

**Usage**

```r
type.group_unique(x)
```

**Arguments**

- `x`  
  any vector

**Value**

the input or error if any of x is not unique.
**type.integer**

*Coerce to integer*

**Description**

Coerce to integer

**Usage**

\[ \text{type.integer}(x) \]

**Arguments**

- \( x \) any vector

**Value**

the input as an integer, error if this would involve data loss.

---

**type.in_range**

*Define a conformance rule to confirm that a numeric is in a set range*

**Description**

This is anticipated to be part of a iface rule e.g.

**Usage**

\[ \text{type.in_range}(\text{min}, \text{max}, \text{include.min} = \text{TRUE}, \text{include.max} = \text{TRUE}) \]

**Arguments**

- \( \text{min} \) the lower limit
- \( \text{max} \) the upper limit
- \( \text{include.min} \) is lower limit open (default \text{TRUE})
- \( \text{include.max} \) is upper limit open (default \text{TRUE})

**Details**

\[ \text{iface}(\text{test}\_\text{col} = \text{integer} + \text{in}\_\text{range}(-10,10) \sim \text{"An integer from -10 to 10"}) \]

**Value**

a function which checks the values and returns them if OK or throws an error if not
Examples

```
type.in_range(0, 10, TRUE, TRUE)(0:10)
try(type.in_range(0, 10, TRUE, FALSE)(0:10))
try(type.in_range(0, 10, FALSE, FALSE)(0:10))
type.in_range(0, 10, FALSE, TRUE)(1:10)
type.in_range(0, 10, TRUE, FALSE)(0:9)
type.in_range(0, Inf, FALSE, FALSE)(1:9)
try(type.in_range(0, 10)(1:99))
```

type.logical

*Coerce to a logical*

**Description**

Coerce to a logical

**Usage**

```
type.logical(x)
```

**Arguments**

- `x` any vector

**Value**

the input as a logical, error if this would involve data loss.

type.not_missing

*Check for missing values*

**Description**

Any NA values will cause failure of validation.

**Usage**

```
type.not_missing(x)
```

**Arguments**

- `x` any vector, factor or numeric

**Value**

the input if no missing values detected, otherwise an error
type.numeric

Coerce to a numeric.

description

Coerce to a numeric.

Usage

type.numeric()

Value

the input as a numeric, error if this would involve data loss.

type.of_type

Check for a given class

description

Any values of the wrong class will cause failure of validation. This is particularly useful for custom vectors of for list types (e.g. list(of_type(lm)))

Usage

type.of_type(type, .not_null = FALSE)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>type</td>
<td>the class of the type we are checking as a symbol</td>
</tr>
<tr>
<td>.not_null</td>
<td>are NULL values allowed (for list column entries only)</td>
</tr>
</tbody>
</table>

Value

a function that can check the input is of the correct type.
type.positive_double  Coerce to a positive double.

Description
Coerce to a positive double.

Usage
type.positive_double(x)

Arguments
x  object to be coerced or tested.

Value
the input as a positive double, error if this would involve data loss.

type.positive_integer  Coerce to a positive integer.

Description
Coerce to a positive integer.

Usage
type.positive_integer(x)

Arguments
x  any vector

Value
the input as a positive integer, error if this would involve data loss.
**type.proportion**  
Coerce to a number between 0 and 1

**Description**
Coerce to a number between 0 and 1

**Usage**
type.proportion(x)

**Arguments**

- **x**  
  object to be coerced or tested.

**Value**
the input as a number from 0 to 1, error if this would involve data loss.

**use_dataframe**  
Use a dataframe in a package including structure based documentation

**Description**
Using the interfacer framework you can document data during development. This provides the basic documentation framework for a dataset based on a dataframe in the correct format into the right place.

**Usage**
use_dataframe(
  df,
  name = deparse(substitute(df)),
  output = "R/data.R",
  pkg = "."  
)

**Arguments**

- **df**  
  the data frame to use
- **name**  
  the name of the variable you wish to use (defaults to whatever the function is called with)
- **output**  
  where to write data documentation code (defaults to R/data.R)
- **pkg**  
  the package (defaults to current)
Details

If this is your only use case for `interfacer` then you will not need to import `interfacer` in your package, as none of the generated code will depend on it.

Value

nothing, used for side effects.

Examples

```r
# example code
if (interactive()) {
  # This is not run as it is designed for interactive use only and will
  # write to the userspace after checking that is what the user wants.
  use_dataframe(iris)
}
```

Description

Generating and documenting an iface for a given dataframe would be time consuming and annoying if you could not do it automatically. In this case as you interactively develop a package using a test dataframe, the structure of which can be explicitly documented and made into a specific contract within the package. This supports development using test dataframes as a prototype for function ensuring future user input conforms to the same expectations as the test data.

Usage

```r
use_iface(
  df,
  name = deparse(substitute(df)),
  output = "R/interfaces.R",
  use_as_default = FALSE,
  pkg = "."
)
```

Arguments

df the data frame to use
name the name of the variable you wish to use (defaults to whatever the dataframe was called)
output where within the current package to write data documentation code (defaults to `R/interfaces.R`)
use iface

use_as_default if this is set to true the current dataframe is saved as package data and the
interfacer::iface specification is created referring to the package copy of
the current dataframe as the default value.

pkg the package (defaults to current)

Value

nothing, used for side effects.

Examples

# example code
if (interactive()) {
  # This is not run as it is designed for interactive use only and will
  # write to the userspace after checking that is what the user wants.
  use_iface(iris)
}

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