

Package ‘DFplyr’

October 2, 2025

Title A `DataFrame` (`S4Vectors`) backend for `dplyr`

Version 1.3.2

Description Provides `dplyr` verbs (`mutate`, `select`, `filter`, etc...) supporting `S4Vectors::DataFrame` objects. Importantly, this is achieved without conversion to an intermediate `tibble`. Adds grouping infrastructure to `DataFrame` which is respected by the transformation verbs.

biocViews DataRepresentation, Infrastructure, Software

License GPL-3

Encoding UTF-8

RoxygenNote 7.3.2

URL <https://github.com/jonocarroll/DFplyr>

BugReports <https://github.com/jonocarroll/DFplyr/issues>

Depends dplyr

Imports BiocGenerics, methods, rlang, S4Vectors, tidyselect

Suggests BiocStyle, GenomeInfoDb, GenomicRanges, IRanges, knitr, rmarkdown, sessioninfo, testthat (>= 3.0.0), tibble

VignetteBuilder knitr

Config/testthat/edition 3

Roxygen list(markdown = TRUE)

git_url <https://git.bioconductor.org/packages/DFplyr>

git_branch devel

git_last_commit 2aa58c3

git_last_commit_date 2025-09-26

Repository Bioconductor 3.22

Date/Publication 2025-10-02

Author Jonathan Carroll [aut, cre] (ORCID:
<<https://orcid.org/0000-0002-1404-5264>>),
Pierre-Paul Axisa [ctb]

Maintainer Jonathan Carroll <rpkg@jcarroll.com.au>

Contents

DFplyr-package	2
arrange.DataFrame	3
bindROWS,DataFrame-method	5
count.DataFrame	5
desc	7
distinct.DataFrame	8
filter.DataFrame	9
format.DataFrame	11
group_by.DataFrame	13
group_by_drop_default.DataFrame	16
group_data	17
group_data.DataFrame	17
group_vars.DataFrame	18
inner_join.DataFrame	19
mutate.DataFrame	20
pull.DataFrame	22
rename,DataFrame-method	23
rename2	24
select.DataFrame	24
slice.DataFrame	28
summarise.DataFrame	31
summarize.DataFrame	33
tally.DataFrame	35
tbl_vars.DataFrame	36
ungroup.DataFrame	37
[,DataFrame-method	39
Index	41

DFplyr-package	<i>Treat a S4Vectors::DataFrame as a dplyr data source</i>
----------------	--

Description

Add **dplyr** compatibility to S4Vectors::DataFrame for use with a selection of **dplyr** verbs.

Arguments

x A S4Vectors::DataFrame object

Author(s)

Maintainer: Jonathan Carroll <rpkg@jcarroll.com.au> ([ORCID](#))

Other contributors:

- Pierre-Paul Axisa [contributor]

See Also

Useful links:

- <https://github.com/jonocarroll/DFplyr>
- Report bugs at <https://github.com/jonocarroll/DFplyr/issues>

Examples

```
library(S4Vectors)
library(dplyr)

d <- as(mtcars, "DataFrame")

mutate(d, newvar = cyl + hp)

mutate_at(d, vars(starts_with("c")), ~ .^2)

group_by(d, cyl, am) %>%
  tally(gear)

count(d, gear, am, cyl)

select(d, am, cyl)

select(d, am, cyl) %>%
  rename2(foo = am)

arrange(d, desc(hp))

rbind(DataFrame(mtcars[1, ], row.names = "MyCar"), d) %>%
  distinct()

filter(d, am == 0)

slice(d, 3:6)
```

arrange.DataFrame	<i>Order rows using column values</i>
-------------------	---------------------------------------

Description

`arrange()` orders the rows of a data frame by the values of selected columns.

Unlike other dplyr verbs, `arrange()` largely ignores grouping; you need to explicitly mention grouping variables (or use `.by_group = TRUE`) in order to group by them, and functions of variables are evaluated once per data frame, not once per group.

Usage

```
## S3 method for class 'DataFrame'
arrange(.data, ...)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>...</code>	<data-masking> Variables, or functions of variables. Use <code>desc()</code> to sort a variable in descending order.

Details

Missing values:

Unlike base sorting with `sort()`, NA are:

- always sorted to the end for local data, even when wrapped with `desc()`.
- treated differently for remote data, depending on the backend.

Value

An object of the same type as `.data`. The output has the following properties:

- All rows appear in the output, but (usually) in a different place.
- Columns are not modified.
- Groups are not modified.
- Data frame attributes are preserved.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

See Also

Other single table verbs: `filter()`, `mutate()`, `reframe()`, `rename()`, `select()`, `slice()`, `summarise()`

Examples

```
arrange(mtcars, cyl, disp)
arrange(mtcars, desc(dis))

# grouped arrange ignores groups
by_cyl <- mtcars %>% group_by(cyl)
by_cyl %>% arrange(desc(wt))
# Unless you specifically ask:
by_cyl %>% arrange(desc(wt), .by_group = TRUE)

# use embracing when wrapping in a function;
# see ?rlang::args_data_masking for more details
tidy_eval_arrange <- function(.data, var) {
  .data %>%
    arrange({{ var }})
}
tidy_eval_arrange(mtcars, mpg)

# Use `across()` or `pick()` to select columns with tidy-select
```

```
iris %>% arrange(pick(starts_with("Sepal")))
iris %>% arrange(across(starts_with("Sepal"), desc))
```

bindROWS,DataFrame-method

rbind DataFrames

Description

rbind DataFrames

Usage

```
## S4 method for signature 'DataFrame'
bindROWS(x, objects = list())
```

Arguments

x	a DataFrame
objects	a list of DataFrames

Value

a new DataFrame combining the inputs by rows

count.DataFrame

Count the observations in each group

Description

count() lets you quickly count the unique values of one or more variables: `df %>% count(a, b)` is roughly equivalent to `df %>% group_by(a, b) %>% summarise(n = n())`. `count()` is paired with `tally()`, a lower-level helper that is equivalent to `df %>% summarise(n = n())`. Supply `wt` to perform weighted counts, switching the summary from `n = n()` to `n = sum(wt)`.

`add_count()` and `add_tally()` are equivalents to `count()` and `tally()` but use `mutate()` instead of `summarise()` so that they add a new column with group-wise counts.

Usage

```
## S3 method for class 'DataFrame'
count(
  x,
  ...,
  wt = NULL,
  sort = FALSE,
  name = "n",
  .drop = group_by_drop_default(x)
)
```

Arguments

<code>x</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>).
<code>...</code>	<code><data-masking></code> Variables to group by.
<code>wt</code>	<code><data-masking></code> Frequency weights. Can be <code>NULL</code> or a variable: • If <code>NULL</code> (the default), counts the number of rows in each group. • If a variable, computes <code>sum(wt)</code> for each group.
<code>sort</code>	If <code>TRUE</code> , will show the largest groups at the top.
<code>name</code>	The name of the new column in the output. If omitted, it will default to <code>n</code> . If there's already a column called <code>n</code> , it will use <code>nn</code> . If there's a column called <code>n</code> and <code>nn</code> , it'll use <code>nnn</code> , and so on, adding <code>ns</code> until it gets a new name.
<code>.drop</code>	Handling of factor levels that don't appear in the data, passed on to <code>group_by()</code> . For <code>count()</code> : if <code>FALSE</code> will include counts for empty groups (i.e. for levels of factors that don't exist in the data). [Deprecated] For <code>add_count()</code> : deprecated since it can't actually affect the output.

Value

An object of the same type as `.data`. `count()` and `add_count()` group transiently, so the output has the same groups as the input.

Examples

```
# count() is a convenient way to get a sense of the distribution of
# values in a dataset
starwars %>% count(species)
starwars %>% count(species, sort = TRUE)
starwars %>% count(sex, gender, sort = TRUE)
starwars %>% count(birth_decade = round(birth_year, -1))

# use the `wt` argument to perform a weighted count. This is useful
# when the data has already been aggregated once
df <- tribble(
  ~name,    ~gender,    ~runs,
  "Max",    "male",      10,
  "Sandra", "female",    1,
  "Susan",  "female",    4
)
# counts rows:
df %>% count(gender)
# counts runs:
df %>% count(gender, wt = runs)

# When factors are involved, `.drop = FALSE` can be used to retain factor
# levels that don't appear in the data
df2 <- tibble(
  id = 1:5,
  type = factor(c("a", "c", "a", NA, "a"), levels = c("a", "b", "c"))
)
df2 %>% count(type)
```

```
df2 %>% count(type, .drop = FALSE)

# Or, using `group_by()`:
df2 %>% group_by(type, .drop = FALSE) %>% count()

# tally() is a lower-level function that assumes you've done the grouping
starwars %>% tally()
starwars %>% group_by(species) %>% tally()

# both count() and tally() have add_ variants that work like
# mutate() instead of summarise
df %>% add_count(gender, wt = runs)
df %>% add_tally(wt = runs)
```

desc

Descending order

Description

Transform a vector into a format that will be sorted in descending order. This is useful within [arrange\(\)](#).

Usage

```
desc(x)
```

Arguments

x vector to transform

Value

the input vector in a format that will be sorted in descending order.

Examples

```
desc(1:10)
desc(factor(letters))

first_day <- seq(as.Date("1910/1/1"), as.Date("1920/1/1"), "years")
desc(first_day)

starwars %>% arrange(desc(mass))
```

distinct.DataFrame	<i>Keep distinct/unique rows</i>
--------------------	----------------------------------

Description

Keep only unique/distinct rows from a data frame. This is similar to `unique.data.frame()` but considerably faster.

Usage

```
## S3 method for class 'DataFrame'
distinct(.data, ..., .keep_all = FALSE)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from dbplyr or dtplyr). See <i>Methods</i> , below, for more details.
<code>...</code>	<data-masking> Optional variables to use when determining uniqueness. If there are multiple rows for a given combination of inputs, only the first row will be preserved. If omitted, will use all variables in the data frame.
<code>.keep_all</code>	If TRUE, keep all variables in <code>.data</code> . If a combination of <code>...</code> is not distinct, this keeps the first row of values.

Value

An object of the same type as `.data`. The output has the following properties:

- Rows are a subset of the input but appear in the same order.
- Columns are not modified if `...` is empty or `.keep_all` is TRUE. Otherwise, `distinct()` first calls `mutate()` to create new columns.
- Groups are not modified.
- Data frame attributes are preserved.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

Examples

```
df <- tibble(
  x = sample(10, 100, rep = TRUE),
  y = sample(10, 100, rep = TRUE)
)
nrow(df)
nrow(distinct(df))
nrow(distinct(df, x, y))

distinct(df, x)
```

```

distinct(df, y)

# You can choose to keep all other variables as well
distinct(df, x, .keep_all = TRUE)
distinct(df, y, .keep_all = TRUE)

# You can also use distinct on computed variables
distinct(df, diff = abs(x - y))

# Use `pick()` to select columns with tidy-select
distinct(starwars, pick(contains("color")))

# Grouping -----

df <- tibble(
  g = c(1, 1, 2, 2, 2),
  x = c(1, 1, 2, 1, 2),
  y = c(3, 2, 1, 3, 1)
)
df <- df %>% group_by(g)

# With grouped data frames, distinctness is computed within each group
df %>% distinct(x)

# When `...` are omitted, `distinct()` still computes distinctness using
# all variables in the data frame
df %>% distinct()

```

filter.DataFrame

Keep rows that match a condition

Description

The `filter()` function is used to subset a data frame, retaining all rows that satisfy your conditions. To be retained, the row must produce a value of `TRUE` for all conditions. Note that when a condition evaluates to `NA` the row will be dropped, unlike base subsetting with `[]`.

Usage

```

## S3 method for class 'DataFrame'
filter(.data, ..., .preserve = FALSE)

```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>...</code>	<data-masking> Expressions that return a logical value, and are defined in terms of the variables in <code>.data</code> . If multiple expressions are included, they are combined with the <code>&</code> operator. Only rows for which all conditions evaluate to <code>TRUE</code> are kept.
<code>.preserve</code>	Relevant when the <code>.data</code> input is grouped. If <code>.preserve = FALSE</code> (the default), the grouping structure is recalculated based on the resulting data, otherwise the grouping is kept as is.

Details

The `filter()` function is used to subset the rows of `.data`, applying the expressions in `...` to the column values to determine which rows should be retained. It can be applied to both grouped and ungrouped data (see `group_by()` and `ungroup()`). However, `dplyr` is not yet smart enough to optimise the filtering operation on grouped datasets that do not need grouped calculations. For this reason, filtering is often considerably faster on ungrouped data.

Value

An object of the same type as `.data`. The output has the following properties:

- Rows are a subset of the input, but appear in the same order.
- Columns are not modified.
- The number of groups may be reduced (if `.preserve` is not `TRUE`).
- Data frame attributes are preserved.

Useful filter functions

There are many functions and operators that are useful when constructing the expressions used to filter the data:

- `==`, `>`, `>=` etc
- `&`, `|`, `!`, `xor()`
- `is.na()`
- `between()`, `near()`

Grouped tibbles

Because filtering expressions are computed within groups, they may yield different results on grouped tibbles. This will be the case as soon as an aggregating, lagging, or ranking function is involved. Compare this ungrouped filtering:

```
starwars %>% filter(mass > mean(mass, na.rm = TRUE))
```

With the grouped equivalent:

```
starwars %>% group_by(gender) %>% filter(mass > mean(mass, na.rm = TRUE))
```

In the ungrouped version, `filter()` compares the value of `mass` in each row to the global average (taken over the whole data set), keeping only the rows with `mass` greater than this global average. In contrast, the grouped version calculates the average `mass` separately for each gender group, and keeps rows with `mass` greater than the relevant within-gender average.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

See Also

Other single table verbs: [arrange\(\)](#), [mutate\(\)](#), [reframe\(\)](#), [rename\(\)](#), [select\(\)](#), [slice\(\)](#), [summarise\(\)](#)

Examples

```
# Filtering by one criterion
filter(starwars, species == "Human")
filter(starwars, mass > 1000)

# Filtering by multiple criteria within a single logical expression
filter(starwars, hair_color == "none" & eye_color == "black")
filter(starwars, hair_color == "none" | eye_color == "black")

# When multiple expressions are used, they are combined using &
filter(starwars, hair_color == "none", eye_color == "black")

# The filtering operation may yield different results on grouped
# tibbles because the expressions are computed within groups.
#
# The following filters rows where `mass` is greater than the
# global average:
starwars %>% filter(mass > mean(mass, na.rm = TRUE))

# Whereas this keeps rows with `mass` greater than the gender
# average:
starwars %>% group_by(gender) %>% filter(mass > mean(mass, na.rm = TRUE))

# To refer to column names that are stored as strings, use the `.data` pronoun:
vars <- c("mass", "height")
cond <- c(80, 150)
starwars %>%
  filter(
    .data[[vars[[1]]]] > cond[[1]],
    .data[[vars[[2]]]] > cond[[2]]
  )
# Learn more in ?rlang::args_data_masking
```

format.DataFrame

*Encode in a Common Format***Description**

Format an R object for pretty printing.

Usage

```
## S3 method for class 'DataFrame'
format(x, ...)
```

Arguments

x any R object (conceptually); typically numeric.
... further arguments passed to or from other methods.

Details

format is a generic function. Apart from the methods described here there are methods for dates (see `format.Date`), date-times (see `format.POSIXct`) and for other classes such as `format.octmode` and `format.dist`.

`format.data.frame` formats the data frame column by column, applying the appropriate method of `format` for each column. Methods for columns are often similar to `as.character` but offer more control. Matrix and data-frame columns will be converted to separate columns in the result, and character columns (normally all) will be given class "`AsIs`".

`format.factor` converts the factor to a character vector and then calls the default method (and so `justify` applies).

`format.AsIs` deals with columns of complicated objects that have been extracted from a data frame. Character objects and (atomic) matrices are passed to the default method (and so `width` does not apply). Otherwise it calls `toString` to convert the object to character (if a vector or list, element by element) and then right-justifies the result.

Justification for character vectors (and objects converted to character vectors by their methods) is done on display width (see `nchar`), taking double-width characters and the rendering of special characters (as escape sequences, including escaping backslash but not double quote: see `print.default`) into account. Thus the width is as displayed by `print(quote = FALSE)` and not as displayed by `cat`. Character strings are padded with blanks to the display width of the widest. (If `na.encode = FALSE` missing character strings are not included in the width computations and are not encoded.)

Numeric vectors are encoded with the minimum number of decimal places needed to display all the elements to at least the digits significant digits. However, if all the elements then have trailing zeroes, the number of decimal places is reduced until at least one element has a non-zero final digit; see also the argument documentation for `big.*`, `small.*` etc, above. See the note in `print.default` about `digits >= 16`.

Raw vectors are converted to their 2-digit hexadecimal representation by `as.character`.

`format.default(x)` now provides a "minimal" string when `isS4(x)` is true.

While the internal code respects the option `getOption("OutDec")` for the 'decimal mark' in general, `decimal.mark` takes precedence over that option. Similarly, `scientific` takes precedence over `getOption("scipen")`.

Value

An object of similar structure to `x` containing character representations of the elements of the first argument `x` in a common format, and in the current locale's encoding.

For character, numeric, complex or factor `x`, `dims` and `dimnames` are preserved on matrices/arrays and names on vectors: no other attributes are copied.

If `x` is a list, the result is a character vector obtained by applying `format.default(x, ...)` to each element of the list (after `unlisting` elements which are themselves lists), and then collapsing the result for each element with `paste(collapse = ", ")`. The defaults in this case are `trim = TRUE`, `justify = "none"` since one does not usually want alignment in the collapsed strings.

References

Becker, R. A., Chambers, J. M. and Wilks, A. R. (1988) *The New S Language*. Wadsworth & Brooks/Cole.

See Also

[format.info](#) indicates how an atomic vector would be formatted.

[formatC](#), [paste](#), [as.character](#), [sprintf](#), [print](#), [prettyNum](#), [toString](#), [encodeString](#).

Examples

```
format(1:10)
format(1:10, trim = TRUE)

zz <- data.frame("(row names)"= c("aaaaa", "b"), check.names = FALSE)
format(zz)
format(zz, justify = "left")

## use of nsmall
format(13.7)
format(13.7, nsmall = 3)
format(c(6.0, 13.1), digits = 2)
format(c(6.0, 13.1), digits = 2, nsmall = 1)

## use of scientific
format(2^31-1)
format(2^31-1, scientific = TRUE)
## scientific = numeric scipen (= {sci}entific notation {pen}alty) :
x <- c(1e5, 1000, 10, 0.1, .001, .123)
t(sapply(setNames(,-4:1),
          \(sci) sapply(x, format, scientific=sci)))

## a list
z <- list(a = letters[1:3], b = (-pi+0i)^((-2:2)/2), c = c(1,10,100,1000),
          d = c("a", "longer", "character", "string"),
          q = quote( a + b ), e = expression(1+x))
## can you find the "2" small differences?
(f1 <- format(z, digits = 2))
(f2 <- format(z, digits = 2, justify = "left", trim = FALSE))
f1 == f2 ## 2 FALSE, 4 TRUE

## A "minimal" format() for S4 objects without their own format() method:
cc <- methods::getClassDef("standardGeneric")
format(cc) ## "<S4 class .....>"
```

group_by.DataFrame	<i>Group by one or more variables</i>
--------------------	---------------------------------------

Description

Most data operations are done on groups defined by variables. `group_by()` takes an existing `tbl` and converts it into a grouped `tbl` where operations are performed "by group". `ungroup()` removes grouping.

Usage

```
## S3 method for class 'DataFrame'
group_by(.data, ..., add = FALSE, .drop = group_by_drop_default(.data))
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>...</code>	In <code>group_by()</code> , variables or computations to group by. Computations are always done on the ungrouped data frame. To perform computations on the grouped data, you need to use a separate <code>mutate()</code> step before the <code>group_by()</code> . Computations are not allowed in <code>nest_by()</code> . In <code>ungroup()</code> , variables to remove from the grouping.
<code>add</code>	When <code>FALSE</code> , the default, <code>group_by()</code> will override existing groups. To add to the existing groups, use <code>.add = TRUE</code> . This argument was previously called <code>add</code> , but that prevented creating a new grouping variable called <code>add</code> , and conflicts with our naming conventions.
<code>.drop</code>	Drop groups formed by factor levels that don't appear in the data? The default is <code>TRUE</code> except when <code>.data</code> has been previously grouped with <code>.drop = FALSE</code> . See <code>group_by_drop_default()</code> for details.

Value

A grouped data frame with class `grouped_df`, unless the combination of `...` and `add` yields a empty set of grouping columns, in which case a tibble will be returned.

Methods

These function are **generics**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

Methods available in currently loaded packages:

- `group_by()`: no methods found.
- `ungroup()`: no methods found.

Ordering

Currently, `group_by()` internally orders the groups in ascending order. This results in ordered output from functions that aggregate groups, such as `summarise()`.

When used as grouping columns, character vectors are ordered in the C locale for performance and reproducibility across R sessions. If the resulting ordering of your grouped operation matters and is dependent on the locale, you should follow up the grouped operation with an explicit call to `arrange()` and set the `.locale` argument. For example:

```
data %>%
  group_by(chr) %>%
  summarise(avg = mean(x)) %>%
  arrange(chr, .locale = "en")
```

This is often useful as a preliminary step before generating content intended for humans, such as an HTML table.

Legacy behavior:

Prior to `dplyr` 1.1.0, character vector grouping columns were ordered in the system locale. If you need to temporarily revert to this behavior, you can set the global option `dplyr.legacy_locale`

to TRUE, but this should be used sparingly and you should expect this option to be removed in a future version of dplyr. It is better to update existing code to explicitly call `arrange(.locale = )` instead. Note that setting `dplyr.legacy_locale` will also force calls to `arrange()` to use the system locale.

See Also

Other grouping functions: `group_map()`, `group_nest()`, `group_split()`, `group_trim()`

Examples

```
by_cyl <- mtcars %>% group_by(cyl)

# grouping doesn't change how the data looks (apart from listing
# how it's grouped):
by_cyl

# It changes how it acts with the other dplyr verbs:
by_cyl %>% summarise(
  disp = mean(disp),
  hp = mean(hp)
)
by_cyl %>% filter(disp == max(disp))

# Each call to summarise() removes a layer of grouping
by_vs_am <- mtcars %>% group_by(vs, am)
by_vs <- by_vs_am %>% summarise(n = n())
by_vs
by_vs %>% summarise(n = sum(n))

# To removing grouping, use ungroup
by_vs %>%
  ungroup() %>%
  summarise(n = sum(n))

# By default, group_by() overrides existing grouping
by_cyl %>%
  group_by(vs, am) %>%
  group_vars()

# Use add = TRUE to instead append
by_cyl %>%
  group_by(vs, am, .add = TRUE) %>%
  group_vars()

# You can group by expressions: this is a short-hand
# for a mutate() followed by a group_by()
mtcars %>%
  group_by(vsam = vs + am)

# The implicit mutate() step is always performed on the
# ungrouped data. Here we get 3 groups:
mtcars %>%
  group_by(vs) %>%
  group_by(hp_cut = cut(hp, 3))

# If you want it to be performed by groups,
```

```
# you have to use an explicit mutate() call.
# Here we get 3 groups per value of vs
mtcars %>%
  group_by(vs) %>%
  mutate(hp_cut = cut(hp, 3)) %>%
  group_by(hp_cut)

# when factors are involved and .drop = FALSE, groups can be empty
tbl <- tibble(
  x = 1:10,
  y = factor(rep(c("a", "c"), each = 5), levels = c("a", "b", "c"))
)
tbl %>%
  group_by(y, .drop = FALSE) %>%
  group_rows()
```

```
group_by_drop_default.DataFrame
```

Default value for .drop argument of group_by

Description

Default value for .drop argument of group_by

Usage

```
## S3 method for class 'DataFrame'
group_by_drop_default(.tbl)
```

Arguments

.tbl A data frame

Value

TRUE unless .tbl is a grouped data frame that was previously obtained by group_by(.drop = FALSE)

Examples

```
group_by_drop_default(iris)

iris %>%
  group_by(Species) %>%
  group_by_drop_default()

iris %>%
  group_by(Species, .drop = FALSE) %>%
  group_by_drop_default()
```

group_data

*Set and Get Group Data on a DataFrame***Description**

The location of group data is an internal implementation detail, so these get and set methods enable interfacing with that data.

Usage

```
set_group_data(x, g, .drop = group_by_drop_default(x))
```

```
get_group_data(x)
```

Arguments

x	A S4Vectors::DataFrame on which to set group data.
g	Group data (a <code>data.frame</code>).
.drop	Drop groups formed by factor levels that don't appear in the data?

Value

For `set_group_data`, the input x with group data set as metadata. For `get_group_data`, the group data that is set on x.

group_data.DataFrame *Grouping metadata***Description**

This collection of functions accesses data about grouped data frames in various ways:

- `group_data()` returns a data frame that defines the grouping structure. The columns give the values of the grouping variables. The last column, always called `.rows`, is a list of integer vectors that gives the location of the rows in each group.
- `group_keys()` returns a data frame describing the groups.
- `group_rows()` returns a list of integer vectors giving the rows that each group contains.
- `group_indices()` returns an integer vector the same length as `.data` that gives the group that each row belongs to.
- `group_vars()` gives names of grouping variables as character vector.
- `groups()` gives the names of the grouping variables as a list of symbols.
- `group_size()` gives the size of each group.
- `n_groups()` gives the total number of groups.

See [context](#) for equivalent functions that return values for the *current* group.

Usage

```
## S3 method for class 'DataFrame'
group_data(.data)
```

Arguments

`.data` a `S4Vectors::DataFrame()`

Value

a data.frame of group data

group_vars.DataFrame *Grouping metadata*

Description

This collection of functions accesses data about grouped data frames in various ways:

- `group_data()` returns a data frame that defines the grouping structure. The columns give the values of the grouping variables. The last column, always called `.rows`, is a list of integer vectors that gives the location of the rows in each group.
- `group_keys()` returns a data frame describing the groups.
- `group_rows()` returns a list of integer vectors giving the rows that each group contains.
- `group_indices()` returns an integer vector the same length as `.data` that gives the group that each row belongs to.
- `group_vars()` gives names of grouping variables as character vector.
- `groups()` gives the names of the grouping variables as a list of symbols.
- `group_size()` gives the size of each group.
- `n_groups()` gives the total number of groups.

See [context](#) for equivalent functions that return values for the *current* group.

Usage

```
## S3 method for class 'DataFrame'
group_vars(x)
```

Arguments

`x` a `S4Vectors::DataFrame()`, likely grouped

Value

the grouping variables as a character vector

inner_join.DataFrame *Mutating joins*

Description

Mutating joins

Usage

```
## S3 method for class 'DataFrame'
inner_join(
  x,
  y,
  by = NULL,
  copy = FALSE,
  suffix = c(".x", ".y"),
  ...,
  keep = NULL
)
```

Arguments

x	a DataFrame
y	a DataFrame or data.frame
by	columns to use for joining the objects. If NULL, the function will look for common columns.

Value

a DataFrame

Examples

```
library(dplyr)
library(S4Vectors)
da <- starwars[, c("name", "mass", "species")][1:10, ]
db <- starwars[, c("name", "homeworld")]

Da <- as(da, "DataFrame")
Db <- as(db, "DataFrame")

Res_inner <- inner_join(Da, Db[1:3, ])
```

mutate.DataFrame	Create, modify, and delete columns
------------------	------------------------------------

Description

mutate() creates new columns that are functions of existing variables. It can also modify (if the name is the same as an existing column) and delete columns (by setting their value to NULL).

Usage

```
## S3 method for class 'DataFrame'
mutate(.data, ...)
```

Arguments

.data	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from dbplyr or dtplyr). See <i>Methods</i> , below, for more details.
...	<data-masking> Name-value pairs. The name gives the name of the column in the output. The value can be: • A vector of length 1, which will be recycled to the correct length. • A vector the same length as the current group (or the whole data frame if ungrouped). • NULL, to remove the column. • A data frame or tibble, to create multiple columns in the output.

Value

An object of the same type as .data. The output has the following properties:

- Columns from .data will be preserved according to the .keep argument.
- Existing columns that are modified by ... will always be returned in their original location.
- New columns created through ... will be placed according to the .before and .after arguments.
- The number of rows is not affected.
- Columns given the value NULL will be removed.
- Groups will be recomputed if a grouping variable is mutated.
- Data frame attributes are preserved.

Useful mutate functions

- +, -, log(), etc., for their usual mathematical meanings
- lead(), lag()
- dense_rank(), min_rank(), percent_rank(), row_number(), cume_dist(), ntile()
- cumsum(), cummean(), cummin(), cummax(), cumany(), cumall()
- na_if(), coalesce()
- if_else(), recode(), case_when()

Grouped tibbles

Because mutating expressions are computed within groups, they may yield different results on grouped tibbles. This will be the case as soon as an aggregating, lagging, or ranking function is involved. Compare this ungrouped mutate:

```
starwars %>%
  select(name, mass, species) %>%
  mutate(mass_norm = mass / mean(mass, na.rm = TRUE))
```

With the grouped equivalent:

```
starwars %>%
  select(name, mass, species) %>%
  group_by(species) %>%
  mutate(mass_norm = mass / mean(mass, na.rm = TRUE))
```

The former normalises mass by the global average whereas the latter normalises by the averages within species levels.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

Methods available in currently loaded packages: no methods found.

See Also

Other single table verbs: [arrange\(\)](#), [filter\(\)](#), [reframe\(\)](#), [rename\(\)](#), [select\(\)](#), [slice\(\)](#), [summarise\(\)](#)

Examples

```
# Newly created variables are available immediately
starwars %>%
  select(name, mass) %>%
  mutate(
    mass2 = mass * 2,
    mass2_squared = mass2 * mass2
  )

# As well as adding new variables, you can use mutate() to
# remove variables and modify existing variables.
starwars %>%
  select(name, height, mass, homeworld) %>%
  mutate(
    mass = NULL,
    height = height * 0.0328084 # convert to feet
  )

# Use across() with mutate() to apply a transformation
# to multiple columns in a tibble.
starwars %>%
  select(name, homeworld, species) %>%
  mutate(across(!name, as.factor))
```

```
# see more in ?across

# Window functions are useful for grouped mutates:
starwars %>%
  select(name, mass, homeworld) %>%
  group_by(homeworld) %>%
  mutate(rank = min_rank(desc(mass)))
# see `vignette("window-functions")` for more details

# By default, new columns are placed on the far right.
df <- tibble(x = 1, y = 2)
df %>% mutate(z = x + y)
df %>% mutate(z = x + y, .before = 1)
df %>% mutate(z = x + y, .after = x)

# By default, mutate() keeps all columns from the input data.
df <- tibble(x = 1, y = 2, a = "a", b = "b")
df %>% mutate(z = x + y, .keep = "all") # the default
df %>% mutate(z = x + y, .keep = "used")
df %>% mutate(z = x + y, .keep = "unused")
df %>% mutate(z = x + y, .keep = "none")

# Grouping -----
# The mutate operation may yield different results on grouped
# tibbles because the expressions are computed within groups.
# The following normalises `mass` by the global average:
starwars %>%
  select(name, mass, species) %>%
  mutate(mass_norm = mass / mean(mass, na.rm = TRUE))

# Whereas this normalises `mass` by the averages within species
# levels:
starwars %>%
  select(name, mass, species) %>%
  group_by(species) %>%
  mutate(mass_norm = mass / mean(mass, na.rm = TRUE))

# Indirection -----
# Refer to column names stored as strings with the `.data` pronoun:
vars <- c("mass", "height")
mutate(starwars, prod = .data[[vars[[1]]]] * .data[[vars[[2]]]])
# Learn more in ?rlang::args_data_masking
```

pull.DataFrame

Extract a single column

Description

pull() is similar to \$. It's mostly useful because it looks a little nicer in pipes, it also works with remote data frames, and it can optionally name the output.

Usage

```
## S3 method for class 'DataFrame'
pull(.data, var = -1, name = NULL, ...)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from dbplyr or dtplyr). See <i>Methods</i> , below, for more details.
<code>var</code>	A variable specified as: • a literal variable name • a positive integer, giving the position counting from the left • a negative integer, giving the position counting from the right. The default returns the last column (on the assumption that's the column you've created most recently). This argument is taken by expression and supports quasiquotation (you can unquote column names and column locations).
<code>name</code>	An optional parameter that specifies the column to be used as names for a named vector. Specified in a similar manner as <code>var</code> .
<code>...</code>	For use by methods.

Value

A vector the same size as `.data`.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

Examples

```
d <- S4Vectors::DataFrame(mtcars)
pull(d, cyl)
```

rename, DataFrame-method

Rename Columns of a DataFrame

Description

Rename Columns of a DataFrame

Usage

```
## S4 method for signature 'DataFrame'
rename(x, ...)
```

Arguments

<code>x</code>	a DataFrame
<code>...</code>	NSE syntax; <code>new_name = old_name</code> to rename selected variables

Value

a DataFrame with columns renamed

rename2	<i>Rename Columns of a DataFrame (deprecated)</i>
---------	---

Description

Rename Columns of a DataFrame (deprecated)

Usage

```
rename2(.data, ...)
```

Arguments

.data	a DataFrame
...	columns to be renamed with syntax new = old

Value

Deprecated - use rename

select.DataFrame	<i>Keep or drop columns using their names and types</i>
------------------	---

Description

Select (and optionally rename) variables in a data frame, using a concise mini-language that makes it easy to refer to variables based on their name (e.g. `a:f` selects all columns from `a` on the left to `f` on the right) or type (e.g. `where(is.numeric)` selects all numeric columns).

Overview of selection features:

Tidyverse selections implement a dialect of R where operators make it easy to select variables:

- `:` for selecting a range of consecutive variables.
- `!` for taking the complement of a set of variables.
- `&` and `|` for selecting the intersection or the union of two sets of variables.
- `c()` for combining selections.

In addition, you can use **selection helpers**. Some helpers select specific columns:

- `everything()`: Matches all variables.
- `last_col()`: Select last variable, possibly with an offset.
- `group_cols()`: Select all grouping columns.

Other helpers select variables by matching patterns in their names:

- `starts_with()`: Starts with a prefix.
- `ends_with()`: Ends with a suffix.
- `contains()`: Contains a literal string.

- `matches()`: Matches a regular expression.
- `num_range()`: Matches a numerical range like x01, x02, x03.

Or from variables stored in a character vector:

- `all_of()`: Matches variable names in a character vector. All names must be present, otherwise an out-of-bounds error is thrown.
- `any_of()`: Same as `all_of()`, except that no error is thrown for names that don't exist.

Or using a predicate function:

- `where()`: Applies a function to all variables and selects those for which the function returns TRUE.

Usage

```
## S3 method for class 'DataFrame'
select(.data, ...)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from dbplyr or dtplyr). See <i>Methods</i> , below, for more details.
<code>...</code>	<tidy-select> One or more unquoted expressions separated by commas. Variable names can be used as if they were positions in the data frame, so expressions like <code>x:y</code> can be used to select a range of variables.

Value

An object of the same type as `.data`. The output has the following properties:

- Rows are not affected.
- Output columns are a subset of input columns, potentially with a different order. Columns will be renamed if `new_name = old_name` form is used.
- Data frame attributes are preserved.
- Groups are maintained; you can't select off grouping variables.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

Examples

Here we show the usage for the basic selection operators. See the specific help pages to learn about helpers like `starts_with()`.

The selection language can be used in functions like `dplyr::select()` or `tidyr::pivot_longer()`. Let's first attach the tidyverse:

```
library(tidyverse)

# For better printing
iris <- as_tibble(iris)
```

Select variables by name:

```
starwars %>% select(height)
#> # A tibble: 87 x 1
#>   height
#>   <int>
#> 1    172
#> 2    167
#> 3     96
#> 4    202
#> # i 83 more rows
```

```
iris %>% pivot_longer(Sepal.Length)
#> # A tibble: 150 x 6
#>   Sepal.Width Petal.Length Petal.Width Species name      value
#>   <dbl>         <dbl>      <dbl> <fct>   <chr>    <dbl>
#> 1         3.5         1.4        0.2 setosa Sepal.Length  5.1
#> 2          3         1.4        0.2 setosa Sepal.Length  4.9
#> 3         3.2         1.3        0.2 setosa Sepal.Length  4.7
#> 4         3.1         1.5        0.2 setosa Sepal.Length  4.6
#> # i 146 more rows
```

Select multiple variables by separating them with commas. Note how the order of columns is determined by the order of inputs:

```
starwars %>% select(homeworld, height, mass)
#> # A tibble: 87 x 3
#>   homeworld height  mass
#>   <chr>      <int> <dbl>
#> 1 Tatooine    172    77
#> 2 Tatooine    167    75
#> 3 Naboo       96     32
#> 4 Tatooine    202   136
#> # i 83 more rows
```

Functions like `tidyr::pivot_longer()` don't take variables with dots. In this case use `c()` to select multiple variables:

```
iris %>% pivot_longer(c(Sepal.Length, Petal.Length))
#> # A tibble: 300 x 5
#>   Sepal.Width Petal.Width Species name      value
#>   <dbl>         <dbl> <fct>   <chr>    <dbl>
#> 1         3.5         0.2 setosa Sepal.Length  5.1
#> 2         3.5         0.2 setosa Petal.Length  1.4
#> 3          3         0.2 setosa Sepal.Length  4.9
#> 4          3         0.2 setosa Petal.Length  1.4
#> # i 296 more rows
```

Operators::

The `:` operator selects a range of consecutive variables:

```
starwars %>% select(name:mass)
```

```
#> # A tibble: 87 x 3
#>   name      height mass
#>   <chr>      <int> <dbl>
#> 1 Luke Skywalker  172    77
#> 2 C-3PO          167    75
#> 3 R2-D2           96    32
#> 4 Darth Vader    202   136
#> # i 83 more rows
```

The ! operator negates a selection:

```
starwars %>% select(!(name:mass))
#> # A tibble: 87 x 11
#>   hair_color skin_color eye_color birth_year sex gender homeworld species
#>   <chr>      <chr>      <chr>      <dbl> <chr> <chr>   <chr>   <chr>
#> 1 blond     fair        blue        19  male masculine Tatooine Human
#> 2 <NA>      gold        yellow      112 none masculine Tatooine Droid
#> 3 <NA>      white, blue red        33  none masculine Naboo   Droid
#> 4 none      white        yellow      41.9 male masculine Tatooine Human
#> # i 83 more rows
#> # i 3 more variables: films <list>, vehicles <list>, starships <list>
```

```
iris %>% select(!c(Sepal.Length, Petal.Length))
#> # A tibble: 150 x 3
#>   Sepal.Width Petal.Width Species
#>   <dbl>      <dbl> <fct>
#> 1      3.5      0.2 setosa
#> 2      3        0.2 setosa
#> 3      3.2      0.2 setosa
#> 4      3.1      0.2 setosa
#> # i 146 more rows
```

```
iris %>% select(!ends_with("Width"))
#> # A tibble: 150 x 3
#>   Sepal.Length Petal.Length Species
#>   <dbl>      <dbl> <fct>
#> 1      5.1      1.4 setosa
#> 2      4.9      1.4 setosa
#> 3      4.7      1.3 setosa
#> 4      4.6      1.5 setosa
#> # i 146 more rows
```

& and | take the intersection or the union of two selections:

```
iris %>% select(starts_with("Petal") & ends_with("Width"))
#> # A tibble: 150 x 1
#>   Petal.Width
#>   <dbl>
#> 1      0.2
#> 2      0.2
#> 3      0.2
#> 4      0.2
#> # i 146 more rows
```

```
iris %>% select(starts_with("Petal") | ends_with("Width"))
#> # A tibble: 150 x 3
#>   Petal.Length Petal.Width Sepal.Width
#>       <dbl>       <dbl>    <dbl>
#> 1         1.4         0.2        3.5
#> 2         1.4         0.2         3
#> 3         1.3         0.2        3.2
#> 4         1.5         0.2        3.1
#> # i 146 more rows
```

To take the difference between two selections, combine the & and ! operators:

```
iris %>% select(starts_with("Petal") & !ends_with("Width"))
#> # A tibble: 150 x 1
#>   Petal.Length
#>       <dbl>
#> 1         1.4
#> 2         1.4
#> 3         1.3
#> 4         1.5
#> # i 146 more rows
```

See Also

Other single table verbs: [arrange\(\)](#), [filter\(\)](#), [mutate\(\)](#), [reframe\(\)](#), [rename\(\)](#), [slice\(\)](#), [summarise\(\)](#)

slice.DataFrame

Subset rows using their positions

Description

slice() lets you index rows by their (integer) locations. It allows you to select, remove, and duplicate rows. It is accompanied by a number of helpers for common use cases:

- slice_head() and slice_tail() select the first or last rows.
- slice_sample() randomly selects rows.
- slice_min() and slice_max() select rows with the smallest or largest values of a variable.

If .data is a [grouped_df](#), the operation will be performed on each group, so that (e.g.) slice_head(df, n = 5) will select the first five rows in each group.

Usage

```
## S3 method for class 'DataFrame'
slice(.data, ..., .preserve = FALSE)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>...</code>	For <code>slice()</code> : <data-masking> Integer row values. Provide either positive values to keep, or negative values to drop. The values provided must be either all positive or all negative. Indices beyond the number of rows in the input are silently ignored. For <code>slice_*</code> (), these arguments are passed on to methods.
<code>.preserve</code>	Relevant when the <code>.data</code> input is grouped. If <code>.preserve = FALSE</code> (the default), the grouping structure is recalculated based on the resulting data, otherwise the grouping is kept as is.

Details

Slice does not work with relational databases because they have no intrinsic notion of row order. If you want to perform the equivalent operation, use [filter\(\)](#) and [row_number\(\)](#).

Value

An object of the same type as `.data`. The output has the following properties:

- Each row may appear 0, 1, or many times in the output.
- Columns are not modified.
- Groups are not modified.
- Data frame attributes are preserved.

Methods

These function are **generics**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

Methods available in currently loaded packages:

- `slice()`: no methods found.
- `slice_head()`: no methods found.
- `slice_tail()`: no methods found.
- `slice_min()`: no methods found.
- `slice_max()`: no methods found.
- `slice_sample()`: no methods found.

See Also

Other single table verbs: [arrange\(\)](#), [filter\(\)](#), [mutate\(\)](#), [reframe\(\)](#), [rename\(\)](#), [select\(\)](#), [summarise\(\)](#)

Examples

```

# Similar to head(mtcars, 1):
mtcars %>% slice(1L)
# Similar to tail(mtcars, 1):
mtcars %>% slice(n())
mtcars %>% slice(5:n())
# Rows can be dropped with negative indices:
slice(mtcars, -(1:4))

# First and last rows based on existing order
mtcars %>% slice_head(n = 5)
mtcars %>% slice_tail(n = 5)

# Rows with minimum and maximum values of a variable
mtcars %>% slice_min(mpg, n = 5)
mtcars %>% slice_max(mpg, n = 5)

# slice_min() and slice_max() may return more rows than requested
# in the presence of ties.
mtcars %>% slice_min(cyl, n = 1)
# Use with_ties = FALSE to return exactly n matches
mtcars %>% slice_min(cyl, n = 1, with_ties = FALSE)
# Or use additional variables to break the tie:
mtcars %>% slice_min(tibble(cyl, mpg), n = 1)

# slice_sample() allows you to random select with or without replacement
mtcars %>% slice_sample(n = 5)
mtcars %>% slice_sample(n = 5, replace = TRUE)

# you can optionally weight by a variable - this code weights by the
# physical weight of the cars, so heavy cars are more likely to get
# selected
mtcars %>% slice_sample(weight_by = wt, n = 5)

# Group wise operation -----
df <- tibble(
  group = rep(c("a", "b", "c"), c(1, 2, 4)),
  x = runif(7)
)

# All slice helpers operate per group, silently truncating to the group
# size, so the following code works without error
df %>% group_by(group) %>% slice_head(n = 2)

# When specifying the proportion of rows to include non-integer sizes
# are rounded down, so group a gets 0 rows
df %>% group_by(group) %>% slice_head(prop = 0.5)

# Filter equivalents -----
# slice() expressions can often be written to use `filter()` and
# `row_number()`, which can also be translated to SQL. For many databases,
# you'll need to supply an explicit variable to use to compute the row number.
filter(mtcars, row_number() == 1L)
filter(mtcars, row_number() == n())
filter(mtcars, between(row_number(), 5, n()))

```

summarise.DataFrame	<i>Summarise each group down to one row</i>
---------------------	---

Description

summarise() creates a new data frame. It returns one row for each combination of grouping variables; if there are no grouping variables, the output will have a single row summarising all observations in the input. It will contain one column for each grouping variable and one column for each of the summary statistics that you have specified.

summarise() and summarize() are synonyms.

Usage

```
## S3 method for class 'DataFrame'
summarise(.data, ...)
```

Arguments

.data	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from dbplyr or dtplyr). See <i>Methods</i> , below, for more details.
...	<data-masking> Name-value pairs of summary functions. The name will be the name of the variable in the result. The value can be: • A vector of length 1, e.g. min(x), n(), or sum(is.na(y)). • A data frame, to add multiple columns from a single expression. [Deprecated] Returning values with size 0 or >1 was deprecated as of 1.1.0. Please use reframe() for this instead.

Value

An object *usually* of the same type as .data.

- The rows come from the underlying [group_keys\(\)](#).
- The columns are a combination of the grouping keys and the summary expressions that you provide.
- The grouping structure is controlled by the .groups= argument, the output may be another [grouped_df](#), a [tibble](#) or a [rowwise](#) data frame.
- Data frame attributes are **not** preserved, because summarise() fundamentally creates a new data frame.

Useful functions

- Center: [mean\(\)](#), [median\(\)](#)
- Spread: [sd\(\)](#), [IQR\(\)](#), [mad\(\)](#)
- Range: [min\(\)](#), [max\(\)](#),
- Position: [first\(\)](#), [last\(\)](#), [nth\(\)](#),
- Count: [n\(\)](#), [n_distinct\(\)](#)
- Logical: [any\(\)](#), [all\(\)](#)

Backend variations

The data frame backend supports creating a variable and using it in the same summary. This means that previously created summary variables can be further transformed or combined within the summary, as in `mutate()`. However, it also means that summary variables with the same names as previous variables overwrite them, making those variables unavailable to later summary variables.

This behaviour may not be supported in other backends. To avoid unexpected results, consider using new names for your summary variables, especially when creating multiple summaries.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

See Also

Other single table verbs: `arrange()`, `filter()`, `mutate()`, `reframe()`, `rename()`, `select()`, `slice()`

Examples

```
# A summary applied to ungrouped tbl returns a single row
mtcars %>%
  summarise(mean = mean(displacement), n = n())

# Usually, you'll want to group first
mtcars %>%
  group_by(cyl) %>%
  summarise(mean = mean(displacement), n = n())

# Each summary call removes one grouping level (since that group
# is now just a single row)
mtcars %>%
  group_by(cyl, vs) %>%
  summarise(cyl_n = n()) %>%
  group_vars()

# BEWARE: reusing variables may lead to unexpected results
mtcars %>%
  group_by(cyl) %>%
  summarise(displacement = mean(displacement), sd = sd(displacement))

# Refer to column names stored as strings with the `.data` pronoun:
var <- "mass"
summarise(starwars, avg = mean(.data[[var]], na.rm = TRUE))
# Learn more in ?rlang::args_data_masking

# In dplyr 1.1.0, returning multiple rows per group was deprecated in favor
# of `reframe()`, which never messages and always returns an ungrouped
# result:
mtcars %>%
  group_by(cyl) %>%
  summarise(qs = quantile(displacement, c(0.25, 0.75)), prob = c(0.25, 0.75))
# ->
```

```
mtcars %>%
  group_by(cyl) %>%
  reframe(qs = quantile(displacement, c(0.25, 0.75)), prob = c(0.25, 0.75))
```

summarize.DataFrame	<i>Summarise each group down to one row</i>
---------------------	---

Description

`summarise()` creates a new data frame. It returns one row for each combination of grouping variables; if there are no grouping variables, the output will have a single row summarising all observations in the input. It will contain one column for each grouping variable and one column for each of the summary statistics that you have specified.

`summarise()` and `summarize()` are synonyms.

Usage

```
## S3 method for class 'DataFrame'
summarize(.data, ...)
```

Arguments

<code>.data</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>). See <i>Methods</i> , below, for more details.
<code>...</code>	<data-masking> Name-value pairs of summary functions. The name will be the name of the variable in the result. The value can be: • A vector of length 1, e.g. <code>min(x)</code>, <code>n()</code>, or <code>sum(is.na(y))</code>. • A data frame, to add multiple columns from a single expression. [Deprecated] Returning values with size 0 or >1 was deprecated as of 1.1.0. Please use reframe() for this instead.

Value

An object *usually* of the same type as `.data`.

- The rows come from the underlying [group_keys\(\)](#).
- The columns are a combination of the grouping keys and the summary expressions that you provide.
- The grouping structure is controlled by the `.groups=` argument, the output may be another [grouped_df](#), a [tibble](#) or a [rowwise](#) data frame.
- Data frame attributes are **not** preserved, because `summarise()` fundamentally creates a new data frame.

Useful functions

- Center: `mean()`, `median()`
- Spread: `sd()`, `IQR()`, `mad()`
- Range: `min()`, `max()`,
- Position: `first()`, `last()`, `nth()`,
- Count: `n()`, `n_distinct()`
- Logical: `any()`, `all()`

Backend variations

The data frame backend supports creating a variable and using it in the same summary. This means that previously created summary variables can be further transformed or combined within the summary, as in `mutate()`. However, it also means that summary variables with the same names as previous variables overwrite them, making those variables unavailable to later summary variables.

This behaviour may not be supported in other backends. To avoid unexpected results, consider using new names for your summary variables, especially when creating multiple summaries.

Methods

This function is a **generic**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

The following methods are currently available in loaded packages: no methods found.

See Also

Other single table verbs: `arrange()`, `filter()`, `mutate()`, `reframe()`, `rename()`, `select()`, `slice()`

Examples

```
# A summary applied to ungrouped tbl returns a single row
mtcars %>%
  summarise(mean = mean(displacement), n = n())

# Usually, you'll want to group first
mtcars %>%
  group_by(cyl) %>%
  summarise(mean = mean(displacement), n = n())

# Each summary call removes one grouping level (since that group
# is now just a single row)
mtcars %>%
  group_by(cyl, vs) %>%
  summarise(cyl_n = n()) %>%
  group_vars()

# BEWARE: reusing variables may lead to unexpected results
mtcars %>%
  group_by(cyl) %>%
  summarise(displacement = mean(displacement), sd = sd(displacement))
```

```
# Refer to column names stored as strings with the `.data` pronoun:
var <- "mass"
summarise(starwars, avg = mean(.data[[var]], na.rm = TRUE))
# Learn more in ?rlang::args_data_masking

# In dplyr 1.1.0, returning multiple rows per group was deprecated in favor
# of `reframe()`, which never messages and always returns an ungrouped
# result:
mtcars %>%
  group_by(cyl) %>%
  summarise(qs = quantile(displacement, c(0.25, 0.75)), prob = c(0.25, 0.75))
# ->
mtcars %>%
  group_by(cyl) %>%
  reframe(qs = quantile(displacement, c(0.25, 0.75)), prob = c(0.25, 0.75))
```

tally.DataFrame

Count the observations in each group

Description

`count()` lets you quickly count the unique values of one or more variables: `df %>% count(a, b)` is roughly equivalent to `df %>% group_by(a, b) %>% summarise(n = n())`. `count()` is paired with `tally()`, a lower-level helper that is equivalent to `df %>% summarise(n = n())`. Supply `wt` to perform weighted counts, switching the summary from `n = n()` to `n = sum(wt)`.

`add_count()` and `add_tally()` are equivalents to `count()` and `tally()` but use `mutate()` instead of `summarise()` so that they add a new column with group-wise counts.

Usage

```
## S3 method for class 'DataFrame'
tally(x, wt = NULL, sort = FALSE, name = NULL)
```

Arguments

<code>x</code>	A data frame, data frame extension (e.g. a tibble), or a lazy data frame (e.g. from <code>dbplyr</code> or <code>dtplyr</code>).
<code>wt</code>	<code><data-masking></code> Frequency weights. Can be <code>NULL</code> or a variable: • If <code>NULL</code> (the default), counts the number of rows in each group. • If a variable, computes <code>sum(wt)</code> for each group.
<code>sort</code>	If <code>TRUE</code> , will show the largest groups at the top.
<code>name</code>	The name of the new column in the output. If omitted, it will default to <code>n</code> . If there's already a column called <code>n</code> , it will use <code>nn</code> . If there's a column called <code>n</code> and <code>nn</code> , it'll use <code>nnn</code> , and so on, adding <code>ns</code> until it gets a new name.

Value

An object of the same type as `.data`. `count()` and `add_count()` group transiently, so the output has the same groups as the input.

Examples

```
# count() is a convenient way to get a sense of the distribution of
# values in a dataset
starwars %>% count(species)
starwars %>% count(species, sort = TRUE)
starwars %>% count(sex, gender, sort = TRUE)
starwars %>% count(birth_decade = round(birth_year, -1))

# use the `wt` argument to perform a weighted count. This is useful
# when the data has already been aggregated once
df <- tribble(
  ~name,    ~gender,    ~runs,
  "Max",    "male",      10,
  "Sandra", "female",    1,
  "Susan",  "female",    4
)
# counts rows:
df %>% count(gender)
# counts runs:
df %>% count(gender, wt = runs)

# When factors are involved, `.drop = FALSE` can be used to retain factor
# levels that don't appear in the data
df2 <- tibble(
  id = 1:5,
  type = factor(c("a", "c", "a", NA, "a"), levels = c("a", "b", "c"))
)
df2 %>% count(type)
df2 %>% count(type, .drop = FALSE)

# Or, using `group_by()``
df2 %>% group_by(type, .drop = FALSE) %>% count()

# tally() is a lower-level function that assumes you've done the grouping
starwars %>% tally()
starwars %>% group_by(species) %>% tally()

# both count() and tally() have add_ variants that work like
# mutate() instead of summarise
df %>% add_count(gender, wt = runs)
df %>% add_tally(wt = runs)
```

tbl_vars.DataFrame	<i>List variables provided by a tbl.</i>
--------------------	--

Description

tbl_vars() returns all variables while tbl_nongroup_vars() returns only non-grouping variables. The groups attribute of the object returned by tbl_vars() is a character vector of the grouping columns.

Usage

```
## S3 method for class 'DataFrame'
tbl_vars(x)
```

Arguments

x A tbl object

Value

all variables, with a groups attribute when grouped.

See Also

[group_vars\(\)](#) for a function that returns grouping variables.

ungroup.DataFrame	<i>Group by one or more variables</i>
-------------------	---------------------------------------

Description

Most data operations are done on groups defined by variables. `group_by()` takes an existing `tbl` and converts it into a grouped `tbl` where operations are performed "by group". `ungroup()` removes grouping.

Usage

```
## S3 method for class 'DataFrame'
ungroup(x, ...)
```

Arguments

x A `tbl()`

... In `group_by()`, variables or computations to group by. Computations are always done on the ungrouped data frame. To perform computations on the grouped data, you need to use a separate `mutate()` step before the `group_by()`. Computations are not allowed in `nest_by()`. In `ungroup()`, variables to remove from the grouping.

Value

A grouped data frame with class `grouped_df`, unless the combination of ... and `add` yields a empty set of grouping columns, in which case a tibble will be returned.

Methods

These function are **generics**, which means that packages can provide implementations (methods) for other classes. See the documentation of individual methods for extra arguments and differences in behaviour.

Methods available in currently loaded packages:

- `group_by()`: no methods found.
- `ungroup()`: no methods found.

Ordering

Currently, `group_by()` internally orders the groups in ascending order. This results in ordered output from functions that aggregate groups, such as `summarise()`.

When used as grouping columns, character vectors are ordered in the C locale for performance and reproducibility across R sessions. If the resulting ordering of your grouped operation matters and is dependent on the locale, you should follow up the grouped operation with an explicit call to `arrange()` and set the `.locale` argument. For example:

```
data %>%
  group_by(chr) %>%
  summarise(avg = mean(x)) %>%
  arrange(chr, .locale = "en")
```

This is often useful as a preliminary step before generating content intended for humans, such as an HTML table.

Legacy behavior:

Prior to dplyr 1.1.0, character vector grouping columns were ordered in the system locale. If you need to temporarily revert to this behavior, you can set the global option `dplyr.legacy_locale` to `TRUE`, but this should be used sparingly and you should expect this option to be removed in a future version of dplyr. It is better to update existing code to explicitly call `arrange(.locale = )` instead. Note that setting `dplyr.legacy_locale` will also force calls to `arrange()` to use the system locale.

See Also

Other grouping functions: `group_map()`, `group_nest()`, `group_split()`, `group_trim()`

Examples

```
by_cyl <- mtcars %>% group_by(cyl)

# grouping doesn't change how the data looks (apart from listing
# how it's grouped):
by_cyl

# It changes how it acts with the other dplyr verbs:
by_cyl %>% summarise(
  disp = mean(disp),
  hp = mean(hp)
)
by_cyl %>% filter(disp == max(disp))

# Each call to summarise() removes a layer of grouping
by_vs_am <- mtcars %>% group_by(vs, am)
by_vs <- by_vs_am %>% summarise(n = n())
by_vs
by_vs %>% summarise(n = sum(n))

# To removing grouping, use ungroup
by_vs %>%
  ungroup() %>%
  summarise(n = sum(n))
```

```

# By default, group_by() overrides existing grouping
by_cyl %>%
  group_by(vs, am) %>%
  group_vars()

# Use add = TRUE to instead append
by_cyl %>%
  group_by(vs, am, .add = TRUE) %>%
  group_vars()

# You can group by expressions: this is a short-hand
# for a mutate() followed by a group_by()
mtcars %>%
  group_by(vsam = vs + am)

# The implicit mutate() step is always performed on the
# ungrouped data. Here we get 3 groups:
mtcars %>%
  group_by(vs) %>%
  group_by(hp_cut = cut(hp, 3))

# If you want it to be performed by groups,
# you have to use an explicit mutate() call.
# Here we get 3 groups per value of vs
mtcars %>%
  group_by(vs) %>%
  mutate(hp_cut = cut(hp, 3)) %>%
  group_by(hp_cut)

# when factors are involved and .drop = FALSE, groups can be empty
tbl <- tibble(
  x = 1:10,
  y = factor(rep(c("a", "c"), each = 5), levels = c("a", "b", "c"))
)
tbl %>%
  group_by(y, .drop = FALSE) %>%
  group_rows()

```

[,DataFrame-method *Subset a DataFrame*

Description

Subset a DataFrame

Usage

```
## S4 method for signature 'DataFrame'
x[i, j, ..., drop = FALSE]
```

Arguments

x	a DataFrame to be subset
i	rows to subset

<code>j</code>	columns to subset
<code>...</code>	other params, passed to regular S4Vectors subsetting
<code>drop</code>	drop dimensions?

Value

a `DataFrame` subset by rows and/or columns

Index

- * **internal**
 - group_data, [17](#)
 - inner_join.DataFrame, [19](#)
- +, [20](#)
- ==, [10](#)
- >, [10](#)
- >=, [10](#)
- [,DataFrame-method, [39](#)
- &, [10](#)
- all(), [31](#), [34](#)
- all_of(), [25](#)
- any(), [31](#), [34](#)
- any_of(), [25](#)
- arrange, [11](#), [21](#), [28](#), [29](#), [32](#), [34](#)
- arrange(), [7](#), [14](#), [15](#), [38](#)
- arrange.DataFrame, [3](#)
- as.character, [12](#), [13](#)
- AsIs, [12](#)
- between(), [10](#)
- bindROWS,DataFrame-method, [5](#)
- case_when(), [20](#)
- cat, [12](#)
- coalesce(), [20](#)
- contains(), [24](#)
- context, [17](#), [18](#)
- count.DataFrame, [5](#)
- cumall(), [20](#)
- cumany(), [20](#)
- cume_dist(), [20](#)
- cummax(), [20](#)
- cummean(), [20](#)
- cummin(), [20](#)
- cumsum(), [20](#)
- dense_rank(), [20](#)
- desc, [7](#)
- desc(), [4](#)
- DFplyr (DFplyr-package), [2](#)
- DFplyr-package, [2](#)
- distinct.DataFrame, [8](#)
- encodeString, [13](#)
- ends_with(), [24](#)
- everything(), [24](#)
- filter, [4](#), [21](#), [28](#), [29](#), [32](#), [34](#)
- filter(), [29](#)
- filter.DataFrame, [9](#)
- first(), [31](#), [34](#)
- format.DataFrame, [11](#)
- format.Date, [12](#)
- format.info, [13](#)
- format.POSIXct, [12](#)
- formatC, [13](#)
- get_group_data (group_data), [17](#)
- getOption, [12](#)
- group_by(), [6](#), [10](#)
- group_by.DataFrame, [13](#)
- group_by_drop_default(), [14](#)
- group_by_drop_default.DataFrame, [16](#)
- group_cols(), [24](#)
- group_data, [17](#)
- group_data.DataFrame, [17](#)
- group_keys(), [31](#), [33](#)
- group_map, [15](#), [38](#)
- group_nest, [15](#), [38](#)
- group_split, [15](#), [38](#)
- group_trim, [15](#), [38](#)
- group_vars(), [37](#)
- group_vars.DataFrame, [18](#)
- grouped_df, [14](#), [28](#), [31](#), [33](#), [37](#)
- if_else(), [20](#)
- inner_join.DataFrame, [19](#)
- IQR(), [31](#), [34](#)
- is.na(), [10](#)
- isS4, [12](#)
- lag(), [20](#)
- last(), [31](#), [34](#)
- last_col(), [24](#)
- lead(), [20](#)
- log(), [20](#)
- mad(), [31](#), [34](#)
- matches(), [25](#)

max(), [31, 34](#)
mean(), [31, 34](#)
median(), [31, 34](#)
min(), [31, 34](#)
min_rank(), [20](#)
mutate, [4, 11, 28, 29, 32, 34](#)
mutate(), [32, 34](#)
mutate.DataFrame, [20](#)

n(), [31, 34](#)
n_distinct(), [31, 34](#)
na_if(), [20](#)
nchar, [12](#)
near(), [10](#)
nth(), [31, 34](#)
ntile(), [20](#)
num_range(), [25](#)

paste, [13](#)
percent_rank(), [20](#)
prettyNum, [13](#)
print, [13](#)
print.default, [12](#)
pull.DataFrame, [22](#)

quasiquotation, [23](#)

recode(), [20](#)
reframe, [4, 11, 21, 28, 29, 32, 34](#)
reframe(), [31, 33](#)
rename, [4, 11, 21, 28, 29, 32, 34](#)
rename.DataFrame-method, [23](#)
rename2, [24](#)
row_number(), [20, 29](#)
rowwise, [31, 33](#)

S4Vectors::DataFrame, [17](#)
S4Vectors::DataFrame(), [18](#)
sd(), [31, 34](#)
select, [4, 11, 21, 29, 32, 34](#)
select.DataFrame, [24](#)
set_group_data(group_data), [17](#)
slice, [4, 11, 21, 28, 32, 34](#)
slice.DataFrame, [28](#)
sprintf, [13](#)
starts_with(), [24, 25](#)
summarise, [4, 11, 21, 28, 29](#)
summarise(), [14, 38](#)
summarise.DataFrame, [31](#)
summarize.DataFrame, [33](#)

tally.DataFrame, [35](#)
tbl(), [37](#)
tbl_vars.DataFrame, [36](#)

tibble, [31, 33](#)
toString, [12, 13](#)

ungroup(), [10](#)
ungroup.DataFrame, [37](#)
unique.data.frame(), [8](#)
unlist, [12](#)

where(), [25](#)

xor(), [10](#)