

# Package ‘tidyFlowCore’

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**Type** Package

**Title** tidyFlowCore: Bringing flowCore to the tidyverse

**Version** 0.99.2

**Description** tidyFlowCore bridges the gap between flow cytometry analysis using the flowCore Bioconductor package and the tidy data principles advocated by the tidyverse. It provides a suite of dplyr-, ggplot2-, and tidyr-like verbs specifically designed for working with flowFrame and flowSet objects as if they were tibbles; however, your data remain flowCore data structures under this layer of abstraction. tidyFlowCore enables intuitive and streamlined analysis workflows that can leverage both the Bioconductor and tidyverse ecosystems for cytometry data.

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**Encoding** UTF-8

**URL** <https://github.com/keyes-timothy/tidyFlowCore>,  
<https://keyes-timothy.github.io/tidyFlowCore/>

**BugReports** <https://github.com/keyes-timothy/tidyFlowCore/issues>

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---

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

---

## Description

Order rows using column values

## Usage

```
## S3 method for class 'flowFrame'
arrange(.data, ..., .by_group = FALSE)
```

## Arguments

<code>.data</code>	A <a href="#">flowFrame</a>
<code>...</code>	Variables, or functions of variables, to arrange by.
<code>.by_group</code>	Unused.

## 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. \* The [flowFrame](#)'s `identifier` will be preserved.

## Examples

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::arrange(feature_1)
```

---

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

---

### Description

Order rows using column values

### Usage

```
## S3 method for class 'flowSet'
arrange(.data, ..., .by_group = FALSE)
```

### Arguments

.data	A <a href="#">flowSet</a>
...	Variables, or functions of variables, to arrange by.
.by_group	Unused.

### 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. \* The [flowSet](#)'s [pData](#) will be preserved.

### Examples

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::arrange(feature_1)
```

---

as_flowFrame	<i>Coerce an object into a <a href="#">flowFrame</a></i>
--------------	--

---

### Description

Coerce an object into a [flowFrame](#)  
 Coerce a data.frame, tbl\_df, or tof\_tbl into a [flowFrame](#)

### Usage

```
as_flowFrame(x, ...)

## S3 method for class 'tof_tbl'
as_flowFrame(x, ...)
```

**Arguments**

x                    A data.frame, tbl\_df, or tof\_tbl.  
 ...                  Unused.

**Value**

A [flowFrame](#)

A [flowFrame](#). Note that all non-numeric columns in 'x' will be removed.

**Examples**

```
NULL
```

```
NULL
```

---

```
as_flowSet
```

```
Coerce an object into a flowSet
```

---

**Description**

Coerce an object into a [flowSet](#)

Coerce a tof\_tbl into a [flowSet](#)

**Usage**

```
as_flowSet(x, ...)
```

```
## S3 method for class 'tof_tbl'  

as_flowSet(x, group_cols, ...)
```

**Arguments**

x                    A tof\_tbl.  
 ...                  Unused.  
 group\_cols          Unquoted names of the columns in 'x' that should be used to group cells into separate [flowFrames](#). Supports tidyselect helpers. Defaults to NULL (all cells are written into a single [flowFrame](#)). Note that the metadata column name "name" is a special value in the [flowSet](#) class, so if any of 'group\_cols' refers to a column named "name," an error will be thrown.

**Value**

A [flowSet](#)

A [flowSet](#) in which cells are grouped into constituent [flowFrames](#) based on the values in 'group\_cols'. If no 'group\_cols' are specified, a [flowFrame](#) will be returned instead. Note that all non-numeric columns in will be removed.

**Examples**

```
NULL
```

```
NULL
```

---

as\_tof\_tbl

*Coerce flowFrames or flowSets into tibbles.*


---

### Description

Coerce flowFrames or flowSets into tibbles.

### Usage

```
as_tof_tbl(
  flow_data,
  .name_method = c("tidyFlowCore", "featureNames", "colnames"),
  sep = "|",
  ...
)
```

### Arguments

flow_data	A flowFrame or flowSet
.name_method	A string indicating how tidyFlowCore should extract column names from ‘flow_data’. Available options are "tidyFlowCore" (the default), which uses tidyFlowCore’s internal heuristic to name columns; "featureNames", which uses <a href="#">featureNames</a> to name the columns; and "colnames", which uses <a href="#">colnames</a> to name the columns. Note that, in most cases, "featureNames" and "colnames" will give identical results.
sep	A string indicating which symbol should be used to separate antigen names and channel names in the columns of the output tof_tbl when .name_method = 'tidyFlowCore'.
...	Optional method-specific arguments.

### Value

A cytometry-specialized tibble called a ‘tof\_tbl’.

### Examples

```
input_file <- system.file("extdata", "0877408774.B08", package="flowCore")
input_flowframe <- flowCore::read.FCS(input_file)
tof_tibble <- as_tof_tbl(input_flowframe)
```

---

as_tof_tbl.flowSet	Convert an object into a tibble-flowCore abstraction (a 'tof_tbl')
--------------------	--

---

## Description

Convert an object into a tibble-flowCore abstraction (a 'tof\_tbl')

## Usage

```
## S3 method for class 'flowSet'
as_tof_tbl(
  flow_data,
  .name_method = c("tidyFlowCore", "featureNames", "colnames"),
  sep = "|",
  ...,
  include_metadata = FALSE,
  include_tidyFlowCore_identifier = FALSE
)
```

## Arguments

flow_data	A FlowSet
.name_method	A string indicating how tidyFlowCore should extract column names for the output tof_tbl from 'flow_data'. Available options are "tidyFlowCore" (the default), which uses tidyFlowCore's internal heuristic to name columns; "featureNames", which uses <a href="#">featureNames</a> to name the columns; and "colnames", which uses <a href="#">colnames</a> to name the columns.
sep	A string to use to separate the antigen name and its associated channel name in the column names of the output tibble. Defaults to " ".
...	Currently unused.
include_metadata	A boolean value indicating if the metadata for each .fcs file read by flowCore (stored in <a href="#">pData</a> ) should be merged into the final result. Defaults to FALSE.
include_tidyFlowCore_identifier	A boolean value indicating if tidyFlowCore's internal identifier for each flowFrame in the flowSet should be included in the output tof_tbl result. Defaults to FALSE.

## Value

A cytometry-specialized tibble called a 'tof\_tbl'.

---

count.flowFrame	<i>Count the observations in each group.</i>
-----------------	--

---

## Description

Count the observations in each group.

## Usage

```
## S3 method for class 'flowFrame'  
count(x, ..., wt = NULL, sort = FALSE, name = NULL)
```

## Arguments

x	A <a href="#">flowFrame</a>
...	Variables to group by, named according to <a href="#">featureNames</a>
wt	If NULL (the default), counts the number of rows in each group. If a variable, computes sum(wt) for each group.
sort	If TRUE, will show the largest groups at the top.
name	If omitted, it will default to n. If there's already a column called n, it will use nn. If there's a column called n and nn, it'll use nnn, and so on, adding ns until it gets a new name.

## Value

A data.frame containing the groupwise counts.

## Examples

```
my_flowframe <-  
  simulate_cytometry_data()$flowframe |>  
  dplyr::mutate(  
    random_group =  
      sample(  
        c("a", "b"),  
        size = nrow(simulate_cytometry_data()$flowframe),  
        replace = TRUE  
      )  
  )  
  
my_flowframe |>  
  dplyr::count(random_group)
```



---

count.flowSet	<i>Count the observations in each group.</i>
---------------	--

---

## Description

Count the observations in each group.

## Usage

```
## S3 method for class 'flowSet'  
count(x, ..., wt = NULL, sort = FALSE, name = NULL)
```

## Arguments

x	A <a href="#">flowSet</a>
...	Variables to group by, named according to <a href="#">featureNames</a> or the columns of the flowSet's <a href="#">pData</a>
wt	If NULL (the default), counts the number of rows in each group. If a variable, computes sum(wt) for each group.
sort	If TRUE, will show the largest groups at the top.
name	If omitted, it will default to n. If there's already a column called n, it will use nn. If there's a column called n and nn, it'll use nnn, and so on, adding ns until it gets a new name.

## Value

A data.frame containing the groupwise counts. If no columns are specified in '...', the grouping is performed by experiment in the flowSet. Otherwise, the columns specified by '...' will be used for grouping.

## Examples

```
my_flowset <- simulate_cytometry_data()$flowset
```

```
my_flowset |>  
  dplyr::count()
```

```
my_flowset |>  
  dplyr::count(cell_type)
```

---

filter.flowFrame	<i>Keep rows that match a condition.</i>
------------------	--

---

**Description**

Keep rows that match a condition.

**Usage**

```
## S3 method for class 'flowFrame'
filter(.data, ..., .by = NULL, .preserve = FALSE)
```

**Arguments**

.data	A <a href="#">flowFrame</a>
...	Expressions that return a logical value, and are defined in terms of the variables in the <a href="#">featureNames</a> of .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.
.by	Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> .
.preserve	Unused.

**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 [flowFrame](#)'s [identifier](#) will be preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::filter(feature_1 > 50)
```

---

filter.flowSet	<i>Keep rows that match a condition.</i>
----------------	--

---

**Description**

Keep rows that match a condition.

**Usage**

```
## S3 method for class 'flowSet'
filter(.data, ..., .by = NULL, .preserve = FALSE)
```

**Arguments**

<code>.data</code>	A <a href="#">flowSet</a>
<code>...</code>	Expressions that return a logical value, and are defined in terms of the variables in the <a href="#">featureNames</a> of the <a href="#">flowFrames</a> in <code>.data</code> . If multiple expressions are included, they are combined with the <code>&amp;</code> operator. Only rows for which all conditions evaluate to TRUE are kept.
<code>.by</code>	Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> .
<code>.preserve</code>	Unused.

**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 [flowSet](#)'s [pData](#) will be preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::filter(feature_1 > 50)
```

---

<code>ggplot.flowFrame</code>	<i>Create a new ggplot.</i>
-------------------------------	-----------------------------

---

**Description**

Create a new ggplot.

**Usage**

```
## S3 method for class 'flowFrame'
ggplot(
  data = NULL,
  mapping = ggplot2::aes(),
  ...,
  environment = parent.frame()
)
```

**Arguments**

<code>data</code>	Default dataset to use for plot in the form of a <a href="#">flowFrame</a> . If not specified, must be supplied in each layer added to the plot.
<code>mapping</code>	Default list of aesthetic mappings to use for plot. If not specified, must be supplied in each layer added to the plot. Note that variable names used for aesthetic mappings come from the <a href="#">featureNames</a> of the input <a href="#">flowFrame</a> .
<code>...</code>	Other arguments passed on to methods. Not currently used.
<code>environment</code>	Deprecated. Used prior to tidy evaluation.

**Value**

A [ggplot](#)

**Examples**

```
simulations <- simulate_cytometry_data()
test_flowframe <- simulations$flowframe

flowframe_plot <-
  test_flowframe |>
  ggplot2::ggplot(ggplot2::aes(x = feature_1, y = feature_2)) +
  ggplot2::geom_point()
```

---

<code>ggplot.flowSet</code>	<i>Create a new ggplot.</i>
-----------------------------	-----------------------------

---

**Description**

Create a new `ggplot`.

**Usage**

```
## S3 method for class 'flowSet'
ggplot(
  data = NULL,
  mapping = ggplot2::aes(),
  ...,
  environment = parent.frame()
)
```

**Arguments**

<code>data</code>	Default dataset to use for plot in the form of a <a href="#">flowSet</a> . If not specified, must be supplied in each layer added to the plot. Note that any metadata stored in <a href="#">pData</a> will be merged into the underlying <code>flowCore</code> -tibble abstraction and will thus be available for plotting.
<code>mapping</code>	Default list of aesthetic mappings to use for plot. If not specified, must be supplied in each layer added to the plot. Note that variable names used for aesthetic mappings come from the <a href="#">featureNames</a> of the input <code>flowSet</code> 's constituent <a href="#">flowFrames</a> .
<code>...</code>	Other arguments passed on to methods. Not currently used.
<code>environment</code>	Deprecated. Used prior to tidy evaluation.

**Value**

A [ggplot](#)

**Examples**

```

simulations <- simulate_cytometry_data()
test_flowset <- simulations$flowset

flowset_plot <-
  test_flowset |>
  ggplot2::ggplot(ggplot2::aes(x = feature_1, y = feature_2)) +
  ggplot2::geom_point()

flowset_plot_with_metadata <-
  test_flowset |>
  # note that `patient` below comes from the flowSet's metadata (pData)
  ggplot2::ggplot(ggplot2::aes(x = feature_1, y = feature_2, color = patient)) +
  ggplot2::geom_point()

```

---

group\_by.flowFrame      *Group a flowFrame into a flowSet using one or more variables.*

---

**Description**

Group a flowFrame into a flowSet using one or more variables.

**Usage**

```

## S3 method for class 'flowFrame'
group_by(.data, ..., .add = FALSE, .drop = dplyr::group_by_drop_default(.data))

```

**Arguments**

.data	A <a href="#">flowFrame</a>
...	Unquoted variables or columns to group by according to .data's <a href="#">featureNames</a> .
.add	Unused.
.drop	Unused.

**Value**

A [flowSet](#) containing one [flowFrame](#) for each of the unique combinations of columns selected in .... Metadata about grouping columns will be stored in the output [flowSet](#)'s [pData](#).

**Examples**

```

my_flowframe <-
  simulate_cytometry_data()$flowframe |>
  dplyr::mutate(
    random_group =
      sample(
        c("a", "b"),
        size = nrow(simulate_cytometry_data()$flowframe),
        replace = TRUE
      )
  )

```

```
my_flowframe |>
  dplyr::group_by(random_group)
```

---

```
make_flowcore_annotated_data_frame
```

*Make the AnnotatedDataFrame needed for the flowFrame class*

---

### Description

Make the AnnotatedDataFrame needed for the flowFrame class

### Usage

```
make_flowcore_annotated_data_frame(maxes_and_mins)
```

### Arguments

`maxes_and_mins` a data.frame containing information about the max and min values of each channel to be saved in the flowFrame.

### Value

An AnnotatedDataFrame.

### Examples

```
NULL
```

---

```
metal_masterlist
```

*A character vector of CyTOF metal name patterns supported by tidyFlowCore*

---

### Description

A character vector used by 'tof\_find\_panel\_info' to detect and parse which CyTOF metals correspond to each channel in an input .fcs file.

### Usage

```
data(metal_masterlist)
```

### Format

A character vector in which each entry is a pattern that tidyFlowCore searches for in every CyTOF channel in input .fcs files. These patterns are an amalgamate of example .fcs files sampled from the studies linked below.

### Value

None

**Source**

<https://github.com/kara-davis-lab/DDPR> <https://cytobank.org/nolanlab/reports/Levine2015.html> <https://cytobank.org/nolanlab/reports/Spitzer2015.html> <https://cytobank.org/nolanlab/reports/Spitzer2017.html> <https://community.cytobank.org/cytobank/projects/609>

---

mutate.flowFrame	<i>Create, modify, and delete columns.</i>
------------------	--

---

**Description**

Create, modify, and delete columns.

**Usage**

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

**Arguments**

.data	A <a href="#">flowFrame</a>
...	Name-value pairs. The name (the left side of the equals sign) gives the name of the column in the output. The right side of the equation performs computations using the names of each channel according to <a href="#">featureNames</a> . Supports tidyselection.

**Value**

A [flowFrame](#). 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.

**Examples**

```
my_flowframe <-
  simulate_cytometry_data()$flowframe |>
  dplyr::mutate(
    random_group =
      sample(
        c("a", "b"),
        size = nrow(simulate_cytometry_data()$flowframe),
        replace = TRUE
      )
  )

my_flowframe |>
  dplyr::mutate(new_feature = feature_1 + feature_2)
```

---

mutate.flowSet	<i>Create, modify, and delete columns.</i>
----------------	--

---

**Description**

Create, modify, and delete columns.

**Usage**

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

**Arguments**

.data	A <a href="#">flowSet</a>
...	Name-value pairs. The name (the left side of the equals sign) gives the name of the column in the output. The right side of the equation performs computations using the names of each channel according to <a href="#">featureNames</a> . Supports tidyselection.

**Value**

A [flowSet](#). 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.

**Examples**

```
my_flowset <-
  simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::mutate(new_feature = feature_1 + feature_2)
```

---

nest.flowFrame	<i>Nest a <a href="#">flowFrame</a> into a <a href="#">flowSet</a></i>
----------------	--

---

**Description**

Nest a [flowFrame](#) into a [flowSet](#)

**Usage**

```
## S3 method for class 'flowFrame'
nest(.data, ..., .by = NULL, .key = NULL, .names_sep = NULL)
```



**Arguments**

<code>.data</code>	A <code>flowFrame</code>
<code>...</code>	Columns to nest; these will appear in the inner <code>flowFrames</code> comprising the output <code>flowSet</code> . Specified using name-variable pairs of the form <code>new_col = c(col1, col2, col3)</code> . The right hand side can be any valid tidyselct expression. If not supplied, then <code>...</code> is derived as all columns not selected by <code>.by</code> .
<code>.by</code>	Columns to nest by; these will be stored in the <code>pData</code> of the output <code>flowSet</code> . <code>.by</code> can be used in place of or in conjunction with columns supplied through <code>....</code> . If not supplied, then <code>.by</code> is derived as all columns not selected by <code>....</code>
<code>.key</code>	Unused.
<code>.names_sep</code>	Unused.

**Value**

A `flowSet` wherein cells are grouped into constituent `flowFrames` based on which columns are used to nest.

**Examples**

```
my_flowframe <-
  simulate_cytometry_data()$flowframe |>
  dplyr::mutate(
    random_group =
      sample(
        c("a", "b"),
        size = nrow(simulate_cytometry_data()$flowframe),
        replace = TRUE
      )
  )
my_flowframe |>
  tidyr::nest(.by = random_group)
```

---

`new_tof_tibble`      *Constructor for a tof\_tibble.*

---

**Description**

Constructor for a `tof_tibble`.

**Usage**

```
new_tof_tibble(x = dplyr::tibble(), panel = dplyr::tibble())
```

**Arguments**

<code>x</code>	A <code>data.frame</code> or <code>tibble</code> containing single-cell mass cytometry data such that rows are cells and columns are CyTOF measurements.
<code>panel</code>	A <code>data.frame</code> or <code>tibble</code> containing information about the panel for the mass cytometry data in <code>x</code> .

**Value**

A 'tof\_tbl', a tibble extension that tracks a few other attributes that are useful for CyTOF data analysis.

**See Also**

Other tof\_tbl utilities: [tof\\_get\\_panel\(\)](#), [tof\\_set\\_panel\(\)](#)

---

pull.flowFrame	<i>Extract a single column.</i>
----------------	---------------------------------

---

**Description**

pull() is similar to \$. It's mostly useful because it looks a little nicer in pipes.

**Usage**

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

**Arguments**

.data	A <a href="#">flowFrame</a> .
var	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.
name	An optional parameter that specifies the column to be used as names for a named vector. Specified in a similar manner as var.
...	For use by methods.

**Value**

A vector the same size as .data.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::pull(feature_1)
```

---

pull.flowSet	<i>Extract a single column.</i>
--------------	---------------------------------

---

**Description**

pull() is similar to \$. It's mostly useful because it looks a little nicer in pipes.

**Usage**

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

**Arguments**

.data	A <a href="#">flowSet</a> .
var	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.
name	An optional parameter that specifies the column to be used as names for a named vector. Specified in a similar manner as var.
...	For use by methods.

**Value**

A vector the same size as .data.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::pull(feature_1)
```

---

reexports	<i>Objects exported from other packages</i>
-----------	---

---

**Description**

These objects are imported from other packages. Follow the links below to see their documentation.

**rlang** :=, [.data](#)

**Value**

See documentation in each object's original package.

**Examples**

```
# See examples in each object's original package
NULL
```

---

rename.flowFrame	<i>Rename columns in a flowFrame</i>
------------------	--------------------------------------

---

**Description**

Rename columns in a [flowFrame](#)

**Usage**

```
## S3 method for class 'flowFrame'
rename(.data, ...)
```

**Arguments**

.data	A <a href="#">flowFrame</a>
...	Unquoted name-value pairs (as specified by <a href="#">featureNames</a> ). Use new_name = old_name to rename selected columns

**Value**

An object of the same type as .data. The output has the following properties: \* Rows are not affected. \* Column names are changed; column order is preserved. \* The [flowFrame](#)'s [identifier](#) will be preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::rename(new_feature = feature_1)
```

---

rename.flowSet	<i>Rename columns in a flowSet</i>
----------------	------------------------------------

---

**Description**

Rename columns in a [flowSet](#)

**Usage**

```
## S3 method for class 'flowSet'
rename(.data, ...)
```

**Arguments**

.data	A <a href="#">flowSet</a>
...	Unquoted name-value pairs (as specified by the <a href="#">featureNames</a> of the <a href="#">flowFrames</a> making up the <a href="#">flowSet</a> ). Use new_name = old_name to rename selected columns

**Value**

An object of the same type as `.data`. The output has the following properties: \* Rows are not affected. \* Column names are changed; column order is preserved. \* The `flowSet`'s `pData` will be preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::rename(new_feature = feature_1)
```

---

rename\_with.flowFrame *Rename columns in a flowFrame*

---

**Description**

Rename columns in a `flowFrame`

**Usage**

```
## S3 method for class 'flowFrame'
rename_with(.data, .fn, .cols = dplyr::everything(), ...)
```

**Arguments**

<code>.data</code>	A <code>flowFrame</code>
<code>.fn</code>	A function used to transform the selected <code>.cols</code> . Should return a character vector the same length as the input.
<code>.cols</code>	Unquoted column names indicating which columns to rename (as specified by <code>featureNames</code> ).
<code>...</code>	Additional arguments passed onto <code>.fn</code> .

**Value**

An object of the same type as `.data`. The output has the following properties: \* Rows are not affected. \* Column names are changed; column order is preserved. \* The `flowFrame`'s `identifier` will be preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::rename_with(.fn = toupper)
```

---

```
rename_with.flowSet Rename columns in a flowSet
```

---

**Description**

Rename columns in a `flowSet`

**Usage**

```
## S3 method for class 'flowSet'
rename_with(.data, .fn, .cols = dplyr::everything(), ...)
```

**Arguments**

<code>.data</code>	A <code>flowSet</code>
<code>.fn</code>	A function used to transform the selected <code>.cols</code> . Should return a character vector the same length as the input.
<code>.cols</code>	Unquoted column names indicating which columns to rename (as specified by the <code>featureNames</code> of the <code>flowFrames</code> making up the <code>flowSet</code> ).
<code>...</code>	Additional arguments passed onto <code>.fn</code> .

**Value**

An object of the same type as `.data`. The output has the following properties: \* Rows are not affected. \* Column names are changed; column order is preserved. \* The `flowSet`'s `pData` will be preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::rename_with(fn = toupper)
```

---

```
select.flowFrame Keep or drop columns using their names and types.
```

---

**Description**

Keep or drop columns using their names and types.

**Usage**

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

**Arguments**

`.data` A `flowFrame`

`...` One or more unquoted expressions separated by commas. Variables names (as specified by `featureNames`) can be used as if they were positions in the `flowFrame`). Supports tidyselection.

**Value**

A `flowFrame`. 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. \* The `flowFrame`'s `identifier` will be preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe
my_flowframe |>
  dplyr::select(feature_1)
```

---

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

---

**Description**

Keep or drop columns using their names and types.

**Usage**

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

**Arguments**

`.data` A `flowSet`

`...` One or more unquoted expressions separated by commas. Variables names (as specified by the `featureNames` of the component `flowFrames` that make up the `flowSet`) can be used as if they were positions in the `flowSet`). Supports tidyselection.

**Value**

A `flowSet`. 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. \* The `flowSet`'s `pData` will be preserved.

**Examples**

```
my_flowset <-
  simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::select(feature_1)
```

---

```
simulate_cytometry_data
```

*Simulate Cytometry Data for FlowSet and FlowFrame Analysis*

---

**Description**

Simulate Cytometry Data for FlowSet and FlowFrame Analysis

**Usage**

```
simulate_cytometry_data(num_cells = 100, num_features = 10, num_flowframes = 5)
```

**Arguments**

`num_cells` An integer indicating the number of cells to simulate.  
`num_features` An integer indicating how many features to simulate.  
`num_flowframes` An integer indicating how many flowFrames to simulate for the simulated flowSet.

**Value**

A list containing two entries: a flowFrame and a flowSet.

**Examples**

```
simulate_cytometry_data()
```

---

```
slice.flowFrame
```

*Subset rows using their positions*

---

**Description**

Subset rows using their positions

**Usage**

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



**Arguments**

.data	A <a href="#">flowFrame</a>
...	Integer row values (to keep).
.by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to <code>group_by</code> .
.preserve	Currently unused.

**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. \* A [flowSet](#)'s `pData` is preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::slice(1)
```

---

slice.flowSet	<i>Subset rows using their positions</i>
---------------	--

---

**Description**

Subset rows using their positions

**Usage**

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

**Arguments**

.data	A <a href="#">flowSet</a>
...	Integer row values (to keep).
.by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to <code>group_by</code> .
.preserve	Currently unused.

**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. \* A [flowSet](#)'s `pData` is preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::slice(1)
```

---

slice\_head.flowFrame *Subset rows at the head of a data structure.*

---

**Description**

Subset rows at the head of a data structure.

**Usage**

```
## S3 method for class 'flowFrame'
slice_head(.data, ..., n, prop, by = NULL)
```

**Arguments**

.data	A <a href="#">flowFrame</a>
...	Unused.
n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows.  A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.

**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. \* A [flowFrame](#)'s [identifier](#) is preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::slice_head(n = 5)
```

---

slice\_head.flowSet      *Subset rows at the head of a data structure.*

---

## Description

Subset rows at the head of a data structure.

## Usage

```
## S3 method for class 'flowSet'  
slice_head(.data, ..., n, prop, by = NULL)
```

## Arguments

.data	A <a href="#">flowSet</a>
...	Unused.
n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows.  A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.

## 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. \* A [flowSet](#)'s [pData](#) is preserved.

## Examples

```
my_flowset <- simulate_cytometry_data()$flowset  
  
my_flowset |>  
  dplyr::slice_head(n = 10)
```

---

slice\_max.flowFrame     *Subset rows of a data structure in order.*

---

### Description

Subset rows of a data structure in order.

### Usage

```
## S3 method for class 'flowFrame'
slice_max(
  .data,
  order_by,
  ...,
  n,
  prop,
  by = NULL,
  with_ties = TRUE,
  na_rm = FALSE
)
```

### Arguments

.data	A <a href="#">flowFrame</a>
order_by	Variable or function of variables to order by. To order by multiple variables, wrap them in a data frame or tibble.
...	Unused.
n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows. A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.
with_ties	Should ties be kept together? The default, TRUE, may return more rows than you request. Use FALSE to ignore ties, and return the first n rows.
na_rm	Should missing values in order_by be removed from the result? If FALSE, NA values are sorted to the end so they will only be included if there are insufficient non-missing values to reach n/prop.

### 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. \* A [flowFrame](#)'s `identifier` is preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::slice_max(order_by = feature_1, n = 5)
```

---

slice\_max.flowSet      *Subset rows of a data structure in order.*

---

**Description**

Subset rows of a data structure in order.

**Usage**

```
## S3 method for class 'flowSet'
slice_max(
  .data,
  order_by,
  ...,
  n,
  prop,
  by = NULL,
  with_ties = TRUE,
  na_rm = FALSE
)
```

**Arguments**

.data	A <a href="#">flowSet</a>
order_by	Variable or function of variables to order by. To order by multiple variables, wrap them in a data frame or tibble.
...	Unused.
n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows. A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.
with_ties	Should ties be kept together? The default, TRUE, may return more rows than you request. Use FALSE to ignore ties, and return the first n rows.
na_rm	Should missing values in order_by be removed from the result? If FALSE, NA values are sorted to the end so they will only be included if there are insufficient non-missing values to reach n/prop.

**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. \* A `flowSet`'s `pData` is preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::slice_max(order_by = feature_1, n = 10)
```

---

`slice_min.flowFrame`     *Subset rows of a data structure in order.*

---

**Description**

Subset rows of a data structure in order.

**Usage**

```
## S3 method for class 'flowFrame'
slice_min(
  .data,
  order_by,
  ...,
  n,
  prop,
  by = NULL,
  with_ties = TRUE,
  na_rm = FALSE
)
```

**Arguments**

<code>.data</code>	A <code>flowFrame</code>
<code>order_by</code>	Variable or function of variables to order by. To order by multiple variables, wrap them in a data frame or tibble.
<code>...</code>	Unused.
<code>n, prop</code>	Provide either <code>n</code> , the number of rows, or <code>prop</code> , the proportion of rows to select. If neither are supplied, <code>n = 1</code> will be used. If <code>n</code> is greater than the number of rows in the group (or <code>prop &gt; 1</code> ), the result will be silently truncated to the group size. <code>prop</code> will be rounded towards zero to generate an integer number of rows. A negative value of <code>n</code> or <code>prop</code> will be subtracted from the group size. For example, <code>n = -2</code> with a group of 5 rows will select $5 - 2 = 3$ rows; <code>prop = -0.25</code> with 8 rows will select $8 * (1 - 0.25) = 6$ rows.
<code>by</code>	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to <code>group_by</code> .

with_ties	Should ties be kept together? The default, TRUE, may return more rows than you request. Use FALSE to ignore ties, and return the first n rows.
na_rm	Should missing values in order_by be removed from the result? If FALSE, NA values are sorted to the end so they will only be included if there are insufficient non-missing values to reach n/prop.

### 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. \* A `flowFrame`'s `identifier` is preserved.

### Examples

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::slice_min(order_by = feature_1, n = 5)
```

---

slice\_min.flowSet      *Subset rows of a data structure in order.*

---

### Description

Subset rows of a data structure in order.

### Usage

```
## S3 method for class 'flowSet'
slice_min(
  .data,
  order_by,
  ...,
  n,
  prop,
  by = NULL,
  with_ties = TRUE,
  na_rm = FALSE
)
```

### Arguments

.data	A <code>flowSet</code>
order_by	Variable or function of variables to order by. To order by multiple variables, wrap them in a data frame or tibble.
...	Unused.

n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows. A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.
with_ties	Should ties be kept together? The default, TRUE, may return more rows than you request. Use FALSE to ignore ties, and return the first n rows.
na_rm	Should missing values in order_by be removed from the result? If FALSE, NA values are sorted to the end so they will only be included if there are insufficient non-missing values to reach n/prop.

**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. \* A flowSet's pData is preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::slice_max(order_by = feature_1, n = 10)
```

---

slice\_sample.flowFrame

*Subset rows randomly*

---

**Description**

Subset rows randomly

**Usage**

```
## S3 method for class 'flowFrame'
slice_sample(.data, ..., n, prop, by = NULL, weight_by = NULL, replace = FALSE)
```

**Arguments**

.data	A flowFrame
...	Unused.



n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows. A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.
weight_by	Sampling weights. This must evaluate to a vector of non-negative numbers the same length as the input. Weights are automatically standardized to sum to 1.
replace	Should sampling be performed with (TRUE) or without (FALSE, the default) replacement.

**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. \* A `flowFrame`'s `identifier` is preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::slice_sample(n = 5)
```

---

slice\_sample.flowSet    *Subset rows randomly*

---

**Description**

Subset rows randomly

**Usage**

```
## S3 method for class 'flowSet'
slice_sample(.data, ..., n, prop, by = NULL, weight_by = NULL, replace = FALSE)
```

**Arguments**

.data	A <code>flowSet</code>
...	Unused.
n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows. A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.

by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.
weight_by	Sampling weights. This must evaluate to a vector of non-negative numbers the same length as the input. Weights are automatically standardized to sum to 1.
replace	Should sampling be performed with (TRUE) or without (FALSE, the default) replacement.

### 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. \* A flowSet's pData is preserved.

### Examples

```
my_flowset <- simulate_cytometry_data()$flowset
```

```
my_flowset |>
  dplyr::slice_sample(n = 10)
```

---

slice\_tail.flowFrame *Subset rows at the tail of a data structure.*

---

### Description

Subset rows at the tail of a data structure.

### Usage

```
## S3 method for class 'flowFrame'
slice_tail(.data, ..., n, prop, by = NULL)
```

### Arguments

.data	A flowFrame
...	Unused.
n, prop	Provide either n, the number of rows, or prop, the proportion of rows to select. If neither are supplied, n = 1 will be used. If n is greater than the number of rows in the group (or prop > 1), the result will be silently truncated to the group size. prop will be rounded towards zero to generate an integer number of rows. A negative value of n or prop will be subtracted from the group size. For example, n = -2 with a group of 5 rows will select 5 - 2 = 3 rows; prop = -0.25 with 8 rows will select 8 * (1 - 0.25) = 6 rows.
by	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to group_by.

**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. \* A `flowFrame`'s `identifier` is preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::slice_tail(n = 5)
```

---

`slice_tail.flowSet`      *Subset rows at the tail of a data structure.*

---

**Description**

Subset rows at the tail of a data structure.

**Usage**

```
## S3 method for class 'flowSet'
slice_tail(.data, ..., n, prop, by = NULL)
```

**Arguments**

<code>.data</code>	A <code>flowSet</code>
<code>...</code>	Unused.
<code>n, prop</code>	Provide either <code>n</code> , the number of rows, or <code>prop</code> , the proportion of rows to select. If neither are supplied, <code>n = 1</code> will be used. If <code>n</code> is greater than the number of rows in the group (or <code>prop &gt; 1</code> ), the result will be silently truncated to the group size. <code>prop</code> will be rounded towards zero to generate an integer number of rows. A negative value of <code>n</code> or <code>prop</code> will be subtracted from the group size. For example, <code>n = -2</code> with a group of 5 rows will select $5 - 2 = 3$ rows; <code>prop = -0.25</code> with 8 rows will select $8 * (1 - 0.25) = 6$ rows.
<code>by</code>	Optionally, an unquoted selection of columns to group by for just this operation. An alternative to <code>group_by</code> .

**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. \* A `flowSet`'s `pData` is preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::slice_tail(n = 10)
```

---

summarise.flowFrame     *Summarize a flowFrame.*

---

### Description

Summarize a flowFrame.

### Usage

```
## S3 method for class 'flowFrame'
summarise(.data, ..., .by = NULL, .groups = NULL)
```

### Arguments

<code>.data</code>	<code>.data</code> A <a href="#">flowFrame</a>
<code>...</code>	Name-value pairs of summary functions. The name will be the name of the variable in the result.
<code>.by</code>	Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> .
<code>.groups</code>	Grouping structure of the result. * "drop_last": dropping the last level of grouping. * "drop": All levels of grouping are dropped. * "keep": Same grouping structure as <code>.data</code> . * "rowwise": Each row is its own group.

### Value

A data.frame containing the summarized result.

### Examples

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::summarise(feature_1_mean = mean(feature_1))
```

---

summarise.flowSet     *Summarize a flowSet.*

---

### Description

Summarize a flowSet.

### Usage

```
## S3 method for class 'flowSet'
summarise(.data, ..., .by = NULL, .groups = NULL)
```

**Arguments**

<code>.data</code>	<code>.data</code> A <a href="#">flowSet</a>
<code>...</code>	Name-value pairs of summary functions. The name will be the name of the variable in the result.
<code>.by</code>	Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> .
<code>.groups</code>	Grouping structure of the result. * <code>"drop_last"</code> : dropping the last level of grouping. * <code>"drop"</code> : All levels of grouping are dropped. * <code>"keep"</code> : Same grouping structure as <code>.data</code> . * <code>"rowwise"</code> : Each row is its own group.

**Value**

A data.frame containing the summarized result.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::summarise(feature_1_mean = mean(feature_1))
```

---

`summarize.flowFrame`     *Summarize a flowFrame.*

---

**Description**

Summarize a flowFrame.

**Usage**

```
## S3 method for class 'flowFrame'
summarize(.data, ..., .by = NULL, .groups = NULL)
```

**Arguments**

<code>.data</code>	<code>.data</code> A <a href="#">flowFrame</a>
<code>...</code>	Name-value pairs of summary functions. The name will be the name of the variable in the result.
<code>.by</code>	Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> .
<code>.groups</code>	Grouping structure of the result. * <code>"drop_last"</code> : dropping the last level of grouping. * <code>"drop"</code> : All levels of grouping are dropped. * <code>"keep"</code> : Same grouping structure as <code>.data</code> . * <code>"rowwise"</code> : Each row is its own group.

**Value**

A data.frame containing the summarized result.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::summarize(feature_1_mean = mean(feature_1))
```

---

summarize.flowSet      *Summarize a flowSet.*

---

**Description**

Summarize a flowSet.

**Usage**

```
## S3 method for class 'flowSet'
summarize(.data, ..., .by = NULL, .groups = NULL)
```

**Arguments**

<code>.data</code>	A <a href="#">flowSet</a>
<code>...</code>	Name-value pairs of summary functions. The name will be the name of the variable in the result.
<code>.by</code>	Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> .
<code>.groups</code>	Grouping structure of the result. * "drop_last": dropping the last level of grouping. * "drop": All levels of grouping are dropped. * "keep": Same grouping structure as <code>.data</code> . * "rowwise": Each row is its own group.

**Value**

A data.frame containing the summarized result.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::summarize(feature_1_mean = mean(feature_1))
```

---

tof_find_panel_info	<i>Use tidyFlowCore's opinionated heuristic for extracting a high-dimensional cytometry panel's channel-antigen pairs from a flowFrame (read from a .fcs file.)</i>
---------------------	---

---

### Description

Using the character vectors obtained from the 'name' and 'desc' columns of the parameters of the data of a flowFrame, infer the cytometry panel used to collect the data and return it as a tidy tibble.

### Usage

```
tof_find_panel_info(input_flowFrame)
```

### Arguments

input\_flowFrame

A flowFrame (just read from an .fcs file) from which a high-dimensional cytometry panel should be extracted

### Value

A tibble with 4 columns ('channels', 'antigens', '.flowCore\_featureNames' and '.flowCore\_colnames'). The first two columns correspond to the channels and antigens of the high-dimensional cytometry panel used during data acquisition, respectively. The last two channels represent the featureNames and colnames attributes used to represent each channel in the input flowFrame.

---

tof_get_panel	<i>Get panel information from a tof_tibble</i>
---------------	--

---

### Description

Get panel information from a tof\_tibble

### Usage

```
tof_get_panel(tof_tibble)
```

### Arguments

tof\_tibble      A 'tof\_tbl'.

### Value

A tibble containing information about the CyTOF panel that was used during data acquisition for the data contained in 'tof\_tibble'.

### See Also

Other tof\_tbl utilities: [new\\_tof\\_tibble\(\)](#), [tof\\_set\\_panel\(\)](#)

**Examples**

NULL

---

tof_set_panel	<i>Set panel information from a tof_tbl</i>
---------------	---

---

**Description**

Set panel information from a tof\_tbl

**Usage**

```
tof_set_panel(tof_tibble, panel)
```

**Arguments**

tof_tibble	A 'tof_tbl'.
panel	A data.frame containing two columns ('channels' and 'antigens') representing the information about a panel

**Value**

A 'tof\_tbl' containing information about the CyTOF panel that was used during data acquisition for the data contained in the input 'tof\_tibble'. Two columns are required: "metals" and "antigens".

**See Also**

Other tof\_tbl utilities: [new\\_tof\\_tibble\(\)](#), [tof\\_get\\_panel\(\)](#)

**Examples**

NULL

---

transmute.flowFrame	<i>Create, modify, and delete columns.</i>
---------------------	--

---

**Description**

Create, modify, and delete columns.

**Usage**

```
## S3 method for class 'flowFrame'
transmute(.data, ...)
```



**Arguments**

`.data` A [flowFrame](#)

`...` Name-value pairs. The name (the left side of the equals sign) gives the name of the column in the output. The right side of the equation performs computations using the names of each channel according to [featureNames](#). Supports tidyselection.

**Value**

A [flowFrame](#). The output has the following properties: \* Columns created or modified through ... will be returned in the order specified by ... \* The number of rows is not affected. \* Columns given the value NULL will be removed. \* The [flowFrame](#)'s [identifier](#) will be preserved.

**Examples**

```
my_flowframe <- simulate_cytometry_data()$flowframe

my_flowframe |>
  dplyr::transmute(new_feature = feature_1 + feature_2)
```

---

`transmute.flowSet`      *Create, modify, and delete columns.*

---

**Description**

Create, modify, and delete columns.

**Usage**

```
## S3 method for class 'flowSet'
transmute(.data, ...)
```

**Arguments**

`.data` A [flowSet](#)

`...` Name-value pairs. The name (the left side of the equals sign) gives the name of the column in the output. The right side of the equation performs computations using the names of each channel according to the [featureNames](#) of `.data`'s constituent [flowFrames](#). Supports tidyselection.

**Value**

A [flowSet](#). The output has the following properties: \* Columns created or modified through ... will be returned in the order specified by ... \* The number of rows is not affected. \* Columns given the value NULL will be removed. \* The [flowSet](#)'s [pData](#) will be preserved.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::transmute(new_feature = feature_1 + feature_2)
```

---

ungroup.flowSet	<i>Ungroup a flowSet</i>
-----------------	--------------------------

---

**Description**

Ungroup a flowSet

**Usage**

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

**Arguments**

x	A <a href="#">flowSet</a>
...	Variables/columns in <a href="#">pData</a> to remove from the grouping. Note that the "name" field in a flowSet's <a href="#">pData</a> is special in <a href="#">flowCore</a> , so requesting an ungrouping by name will result in a copied column called ".tidyFlowCore_name" in the result. Also note that the column ".tidyof_unique_identifier" is used internally and will not have any effect on the ungrouping.

**Value**

A [flowFrame](#) or [flowSet](#) depending on the degree of ungrouping. Note that unnest-ing and ungrouping a [flowSet](#) are equivalent.

**Examples**

```
my_flowset <- simulate_cytometry_data()$flowset

my_flowset |>
  dplyr::ungroup()
```

---

unnest.flowSet	<i>Unnest a <a href="#">flowSet</a> into a single <a href="#">flowFrame</a></i>
----------------	---

---

## Description

Unnest a [flowSet](#) into a single [flowFrame](#)

## Usage

```
## S3 method for class 'flowSet'  
unnest(  
  data,  
  cols,  
  ...,  
  keep_empty = FALSE,  
  ptype = NULL,  
  names_sep = NULL,  
  names_repair = "check_unique"  
)
```

## Arguments

data	A <a href="#">flowSet</a>
cols	Columns in <a href="#">pData</a> to unnest.
...	Unused.
keep_empty	Unused.
ptype	Unused.
names_sep	Unused.
names_repair	Unused.

## Value

A [flowFrame](#) or [flowSet](#) depending on the degree of unnest-ing. Note that unnest-ing and ungrouping a [flowSet](#) are equivalent.

## Examples

```
my_flowset <- simulate_cytometry_data()$flowset  
  
my_flowset |>  
  tidyr::unnest(cols = c(patient, cell_type))  
  
my_flowset |>  
  tidyr::unnest(cols = patient)
```

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