

# Package: partition (via r-universe)

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**Type** Package

**Title** Agglomerative Partitioning Framework for Dimension Reduction

**Version** 0.2.2

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**Description** A fast and flexible framework for agglomerative partitioning. 'partition' uses an approach called Direct-Measure-Reduce to create new variables that maintain the user-specified minimum level of information. Each reduced variable is also interpretable: the original variables map to one and only one variable in the reduced data set. 'partition' is flexible, as well: how variables are selected to reduce, how information loss is measured, and the way data is reduced can all be customized. 'partition' is based on the Partition framework discussed in Millstein et al. (2020) <doi:10.1093/bioinformatics/btz661>.

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**URL** <https://uscbiostats.github.io/partition/>,  
<https://github.com/USCbiostats/partition>

**BugReports** <https://github.com/USCbiostats/partition/issues>

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<i>as_director</i>	<i>Create a custom director</i>
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---

**Description**

Directors are functions that tell the partition algorithm what to try to reduce. `as_director()` is a helper function to create new directors to be used in partitioners. partitioners can be created with `as_partitioner()`.

**Usage**

```
as_director(.pairs, .target, ...)
```

**Arguments**

- `.pairs` a function that returns a matrix of targets (e.g. a distance matrix of variables)
- `.target` a function that returns a vector of targets (e.g. the minimum pair)
- `...` Extra arguments passed to `.f`.

**Value**

a function to use in `as_partitioner()`

**See Also**

Other directors: `direct_distance()`, `direct_k_cluster()`

**Examples**

```
# use euclidean distance to calculate distances
euc_dist <- function(.data) as.matrix(dist(t(.data)))

# find the pair with the minimum distance
min_dist <- function(.x) {
  indices <- arrayInd(which.min(.x), dim(as.matrix(.x)))

  # get variable names with minimum distance
  c(
    colnames(.x)[indices[1]],
```

```

      colnames(.x)[indices[2]]
    )
  }

  as_director(euc_dist, min_dist)

```

---

as\_measure

*Create a custom metric*


---

## Description

Metrics are functions that tell how much information would be lost for a given reduction in the data. reduce. [as\\_measure\(\)](#) is a helper function to create new metrics to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

## Usage

```
as_measure(.f, ...)
```

## Arguments

```
.f          a function that returns either a numeric vector or a data.frame
...         Extra arguments passed to .f.
```

## Value

a function to use in [as\\_partitioner\(\)](#)

## See Also

Other metrics: [measure\\_icc\(\)](#), [measure\\_min\\_icc\(\)](#), [measure\\_min\\_r2\(\)](#), [measure\\_std\\_mutualinfo\(\)](#), [measure\\_variance\\_explained\(\)](#)

Other metrics: [measure\\_icc\(\)](#), [measure\\_min\\_icc\(\)](#), [measure\\_min\\_r2\(\)](#), [measure\\_std\\_mutualinfo\(\)](#), [measure\\_variance\\_explained\(\)](#)

## Examples

```

inter_item_reliability <- function(mat) {
  corrs <- corr(mat)
  corrs[lower.tri(corrs, diag = TRUE)] <- NA

  corrs %>%
    colMeans(na.rm = TRUE) %>%
    mean(na.rm = TRUE)
}

measure_iir <- as_measure(inter_item_reliability)
measure_iir

```

---

`as_partitioner`*Create a partitioner*

---

## Description

Partitioners are functions that tell the partition algorithm 1) what to try to reduce 2) how to measure how much information is lost from the reduction and 3) how to reduce the data. In partition, functions that handle 1) are called directors, functions that handle 2) are called metrics, and functions that handle 3) are called reducers. `partition` has a number of pre-specified partitioners for agglomerative data reduction. Custom partitioners can be created with `as_partitioner()`.

Pass partitioner objects to the `partitioner` argument of `partition()`.

## Usage

```
as_partitioner(direct, measure, reduce)
```

## Arguments

<code>direct</code>	a function that directs, possibly created by <code>as_director()</code>
<code>measure</code>	a function that measures, possibly created by <code>as_measure()</code>
<code>reduce</code>	a function that reduces, possibly created by <code>as_reducer()</code>

## Value

a partitioner

## See Also

Other partitioners: `part_icc()`, `part_kmeans()`, `part_minr2()`, `part_pc1()`, `part_stdmi()`, `replace_partitioner()`

## Examples

```
as_partitioner(  
  direct = direct_distance_pearson,  
  measure = measure_icc,  
  reduce = reduce_scaled_mean  
)
```

---

as\_partition\_step      *Create a partition object from a data frame*

---

### Description

as\_partition\_step() creates a partition\_step object. partition\_steps are used while iterating through the partition algorithm: it stores necessary information about how to proceed in the partitioning, such as the information threshold. as\_partition\_step() is primarily called internally by partition() but can be helpful while developing partitioners.

### Usage

```
as_partition_step(
  .x,
  threshold = NA,
  reduced_data = NA,
  target = NA,
  metric = NA,
  tolerance = 0.01,
  var_prefix = NA,
  partitioner = NA,
  ...
)
```

### Arguments

.x	a data.frame or partition_step object
threshold	The minimum information loss allowable
reduced_data	A data set with reduced variables
target	A character or integer vector: the variables to reduce
metric	A measure of information
tolerance	A tolerance around the threshold to accept a reduction
var_prefix	Variable name for reduced variables
partitioner	A partitioner, a part_*() function or one created with <a href="#">as_partitioner()</a> .
...	Other objects to store during the partition step

### Value

a partition\_step object

### Examples

```
.df <- data.frame(x = rnorm(100), y = rnorm(100))
as_partition_step(.df, threshold = .6)
```

---

as_reducer	<i>Create a custom reducer</i>
------------	--------------------------------

---

## Description

Reducers are functions that tell the partition algorithm how to reduce the data. `as_reducer()` is a helper function to create new reducers to be used in partitioners. partitioners can be created with `as_partitioner()`.

## Usage

```
as_reducer(.f, ..., returns_vector = TRUE, first_match = NULL)
```

## Arguments

<code>.f</code>	a function that returns either a numeric vector or a <code>data.frame</code>
<code>...</code>	Extra arguments passed to <code>.f</code> .
<code>returns_vector</code>	logical. Does <code>.f</code> return a vector? TRUE by default. If FALSE, assumes that <code>.f</code> returns a <code>data.frame</code> .
<code>first_match</code>	logical. Should the partition algorithm stop when it finds a reduction that is equal to the threshold? Default is TRUE for reducers that return a <code>data.frame</code> and FALSE for reducers that return a vector

## Value

a function to use in `as_partitioner()`

## See Also

Other reducers: `reduce_first_component()`, `reduce_kmeans()`, `reduce_scaled_mean()`

Other reducers: `reduce_first_component()`, `reduce_kmeans()`, `reduce_scaled_mean()`

## Examples

```
reduce_row_means <- as_reducer(rowMeans)
reduce_row_means
```

---

`baxter_data`*Microbiome data*

---

## Description

Clinical and microbiome data derived from "Microbiota-based model improves the sensitivity of fecal immunochemical test for detecting colonic lesions" by Baxter et al. (2016). These data represent a subset of 172 health participants. `baxter_clinical` contains 8 clinical variables for each of the participants: `sample_name`, `id`, `age`, `bmi`, `gender`, `height`, `total_reads`, and `disease_state` (all H for healthy). `baxter_otu` has 1,234 columns, where each column represents an Operational Taxonomic Unit (OTU). OTUs are species-like relationships among bacteria determined by analyzing their RNA. The cells are logged counts for how often the OTU was detected in a participant's stool sample. Each column name is a shorthand name, e.g. `otu1`; you can find the true name of the OTU mapped in `baxter_data_dictionary`. `baxter_family` and `baxter_genus` are also logged counts but instead group OTUs at the family and genus level, respectively, a common approach to reducing microbiome data. Likewise, the column names are shorthands, which you can find mapped in `baxter_data_dictionary`.

## Usage

`baxter_clinical``baxter_otu``baxter_family``baxter_genus``baxter_data_dictionary`

## Format

5 data frames

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 172 rows and 1234 columns.

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 172 rows and 35 columns.

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 172 rows and 82 columns.

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 1351 rows and 3 columns.

## Source

Baxter et al. (2016) [doi:10.1186/s1307301602903](https://doi.org/10.1186/s1307301602903)



---

corr *Efficiently fit correlation coefficient for matrix or two vectors*

---

### Description

Efficiently fit correlation coefficient for matrix or two vectors

### Usage

```
corr(x, y = NULL, spearman = FALSE)
```

### Arguments

x	a matrix or vector
y	a vector. Optional.
spearman	Logical. Use Spearman's correlation?

### Value

a numeric vector, the correlation coefficient

### Examples

```
library(dplyr)
# fit for entire data set
iris %>%
  select_if(is.numeric) %>%
  corr()

# just fit for two vectors
corr(iris$Sepal.Length, iris$Sepal.Width)
```

---

direct\_distance *Target based on minimum distance matrix*

---

### Description

Directors are functions that tell the partition algorithm what to try to reduce. `as_director()` is a helper function to create new directors to be used in partitioners. partitioners can be created with `as_partitioner()`.

`direct_distance()` fits a distance matrix using either Pearson's or Spearman's correlation and finds the pair with the smallest distance to target. If the distance matrix already exists, `direct_distance()` only fits the distances for any new reduced variables. `direct_distance_pearson()` and `direct_distance_spearman()` are convenience functions that directly call the type of distance matrix.

**Usage**

```
direct_distance(.partition_step, spearman = FALSE)

direct_distance_pearson(.partition_step)

direct_distance_spearman(.partition_step)
```

**Arguments**

```
.partition_step      a partition_step object
spearman             Logical. Use Spearman's correlation?
```

**Value**

a partition\_step object

**See Also**

Other directors: [as\\_director\(\)](#), [direct\\_k\\_cluster\(\)](#)

---

direct_k_cluster	<i>Target based on K-means clustering</i>
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---

**Description**

Directors are functions that tell the partition algorithm what to try to reduce. [as\\_director\(\)](#) is a helper function to create new directors to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`direct_k_cluster()` assigns each variable to a cluster using K-means. As the partition looks for the best reduction, `direct_k_cluster()` iterates through values of k to assign clusters. This search is handled by the binary search method by default and thus does not necessarily need to fit every value of k.

**Usage**

```
direct_k_cluster(
  .partition_step,
  algorithm = c("armadillo", "Hartigan-Wong", "Lloyd", "Forgy", "MacQueen"),
  search = c("binary", "linear"),
  init_k = NULL,
  seed = 1L
)
```

**Arguments**

<code>.partition_step</code>	a <code>partition_step</code> object
<code>algorithm</code>	The K-Means algorithm to use. The default is a fast version of the LLoyd algorithm written in <code>armadillo</code> . The rest are options in <code>kmeans()</code> . In general, <code>armadillo</code> is fastest, but the other algorithms can be faster in high dimensions.
<code>search</code>	The search method. Binary search is generally more efficient but linear search can be faster in very low dimensions.
<code>init_k</code>	The initial k to test. If <code>NULL</code> , then the initial k is the threshold times the number of variables.
<code>seed</code>	The seed to set for reproducibility

**Value**

a `partition_step` object

**See Also**

Other directors: `as_director()`, `direct_distance()`

---

<code>filter_reduced</code>	<i>Filter the reduced mappings</i>
-----------------------------	------------------------------------

---

**Description**

`filter_reduced()` and `unnest_reduced()` are convenience functions to quickly retrieve the mappings for only the reduced variables. `filter_reduced()` returns a nested tibble while `unnest_reduced()` unnests it.

**Usage**

```
filter_reduced(.partition)
```

```
unnest_reduced(.partition)
```

**Arguments**

<code>.partition</code>	a <code>partition</code> object
-------------------------	---------------------------------

**Value**

a tibble with mapping key

## Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)
# fit partition
prt <- partition(df, threshold = .6)

# A tibble: 3 x 4
filter_reduced(prt)

# A tibble: 9 x 4
unnest_reduced(prt)
```

---

icc

*Calculate the intraclass correlation coefficient*

---

## Description

icc() efficiently calculates the ICC for a numeric data set.

## Usage

```
icc(.x, method = c("r", "c"))
```

## Arguments

.x	a data set
method	The method source: both the pure R and C++ versions are efficient

## Value

a numeric vector of length 1

## Examples

```
library(dplyr)
iris %>%
  select_if(is.numeric) %>%
  icc()
```

---

*is\_partition*                      *Is this object a partition?*

---

**Description**

Is this object a partition?

**Usage**

`is_partition(x)`

**Arguments**

`x`                      an object to be tested

**Value**

logical: TRUE or FALSE

---

*is\_partitioner*                      *Is this object a partitioner?*

---

**Description**

Is this object a partitioner?

**Usage**

`is_partitioner(x)`

**Arguments**

`x`                      an object to be tested

**Value**

logical: TRUE or FALSE

---

is_partition_step	<i>Is this object a partition_step?</i>
-------------------	---

---

**Description**

Is this object a partition\_step?

**Usage**

```
is_partition_step(x)
```

**Arguments**

x                    an object to be tested

**Value**

logical: TRUE or FALSE

---

mapping_key	<i>Return partition mapping key</i>
-------------	-------------------------------------

---

**Description**

mapping\_key() returns a data frame with each reduced variable and its mapping and information loss; the mapping and indices are represented as list-cols (so there is one row per variable in the reduced data set). unnest\_mappings() unnests the list columns to return a tidy data frame. mapping\_groups() returns a list of mappings (either the variable names or their column position).

**Usage**

```
mapping_key(.partition)
```

```
unnest_mappings(.partition)
```

```
mapping_groups(.partition, indices = FALSE)
```

**Arguments**

.partition        a partition object

indices           logical. Return just the indices instead of the names? Default is FALSE.

**Value**

a tibble

## Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)
# fit partition
prt <- partition(df, threshold = .6)

# tibble: 6 x 4
mapping_key(prt)

# tibble: 12 x 4
unnest_mappings(prt)

# list: length 6
mapping_groups(prt)
```

---

map\_partition

*Map a partition across a range of minimum information*

---

## Description

map\_partition() fits partition() across a range of minimum information values, specified in the information argument. The output is a tibble with a row for each value of information, a summary of the partition, and a list-col containing the partition object.

## Usage

```
map_partition(
  .data,
  partitioner = part_icc(),
  ...,
  information = seq(0.1, 0.5, by = 0.1)
)
```

## Arguments

.data	a data set to partition
partitioner	the partitioner to use. The default is <a href="#">part_icc()</a> .
...	arguments passed to <a href="#">partition()</a>
information	a vector of minimum information to fit in <a href="#">partition()</a>

## Value

a tibble

## Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

map_partition(df, partitioner = part_pc1())
```

---

measure_icc	<i>Measure the information loss of reduction using intraclass correlation coefficient</i>
-------------	---

---

## Description

Metrics are functions that tell how much information would be lost for a given reduction in the data. `reduce.as_measure()` is a helper function to create new metrics to be used in partitioners. partitioners can be created with `as_partitioner()`.

`measure_icc()` assesses information loss by calculating the intraclass correlation coefficient for the target variables.

## Usage

```
measure_icc(.partition_step)
```

## Arguments

```
.partition_step  
  a partition_step object
```

## Value

```
a partition_step object
```

## See Also

Other metrics: `as_measure()`, `measure_min_icc()`, `measure_min_r2()`, `measure_std_mutualinfo()`, `measure_variance_explained()`



---

measure_min_icc	<i>Measure the information loss of reduction using the minimum intra-class correlation coefficient</i>
-----------------	--

---

### Description

Metrics are functions that tell how much information would be lost for a given reduction in the data. reduce. [as\\_measure\(\)](#) is a helper function to create new metrics to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`measure_min_icc()` assesses information loss by calculating the intraclass correlation coefficient for each set of the target variables and finding their minimum.

### Usage

```
measure_min_icc(.partition_step, search_method = c("binary", "linear"))
```

### Arguments

`.partition_step` a `partition_step` object

`search_method` The search method. Binary search is generally more efficient but linear search can be faster in very low dimensions.

### Value

a `partition_step` object

### See Also

Other metrics: [as\\_measure\(\)](#), [measure\\_icc\(\)](#), [measure\\_min\\_r2\(\)](#), [measure\\_std\\_mutualinfo\(\)](#), [measure\\_variance\\_explained\(\)](#)

---

measure_min_r2	<i>Measure the information loss of reduction using minimum R-squared</i>
----------------	--

---

### Description

Metrics are functions that tell how much information would be lost for a given reduction in the data. reduce. [as\\_measure\(\)](#) is a helper function to create new metrics to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`measure_min_r2()` assesses information loss by calculating the minimum R-squared for the target variables.

### Usage

```
measure_min_r2(.partition_step)
```

**Arguments**

`.partition_step`  
a `partition_step` object

**Value**

a `partition_step` object

**See Also**

Other metrics: [as\\_measure\(\)](#), [measure\\_icc\(\)](#), [measure\\_min\\_icc\(\)](#), [measure\\_std\\_mutualinfo\(\)](#), [measure\\_variance\\_explained\(\)](#)

---

`measure_std_mutualinfo`

*Measure the information loss of reduction using standardized mutual information*

---

**Description**

Metrics are functions that tell how much information would be lost for a given reduction in the data. reduce. [as\\_measure\(\)](#) is a helper function to create new metrics to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`measure_std_mutualinfo()` assesses information loss by calculating the standardized mutual information for the target variables. See [mutual\\_information\(\)](#).

**Usage**

```
measure_std_mutualinfo(.partition_step)
```

**Arguments**

`.partition_step`  
a `partition_step` object

**Value**

a `partition_step` object

**See Also**

Other metrics: [as\\_measure\(\)](#), [measure\\_icc\(\)](#), [measure\\_min\\_icc\(\)](#), [measure\\_min\\_r2\(\)](#), [measure\\_variance\\_explained\(\)](#)

---

measure\_variance\_explained

*Measure the information loss of reduction using the variance explained.*

---

### Description

Metrics are functions that tell how much information would be lost for a given reduction in the data. reduce. [as\\_measure\(\)](#) is a helper function to create new metrics to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

measure\_variance\_explained() assesses information loss by calculating the variance explained by the first component of a principal components analysis. Because the PCA calculates the components and the variance explained at the same time, if the reducer is reduce\_first\_component(), then measure\_variance\_explained() will store the first component for later use to avoid recalculation.

### Usage

```
measure_variance_explained(.partition_step)
```

### Arguments

.partition\_step  
a partition\_step object

### Value

a partition\_step object

### See Also

Other metrics: [as\\_measure\(\)](#), [measure\\_icc\(\)](#), [measure\\_min\\_icc\(\)](#), [measure\\_min\\_r2\(\)](#), [measure\\_std\\_mutualinfo\(\)](#)

---

mutual\_information

*Calculate the standardized mutual information of a data set*

---

### Description

mutual\_information calculate the standardized mutual information of a data set using the infotheo package.

### Usage

```
mutual_information(.data)
```

**Arguments**

`.data` a dataframe of numeric values

**Value**

a list containing the standardized MI and the scaled row means

**Examples**

```
library(dplyr)
iris %>%
  select_if(is.numeric) %>%
  mutual_information()
```

---

partition

*Agglomerative partitioning*

---

**Description**

`partition()` reduces data while minimizing information loss using an agglomerative partitioning algorithm. The partition algorithm is fast and flexible: at every iteration, `partition()` uses an approach called Direct-Measure-Reduce (see Details) to create new variables that maintain the user-specified minimum level of information. Each reduced variable is also interpretable: the original variables map to one and only one variable in the reduced data set.

**Usage**

```
partition(
  .data,
  threshold,
  partitioner = part_icc(),
  tolerance = 1e-04,
  niter = NULL,
  x = "reduced_var",
  .sep = "_"
)
```

**Arguments**

`.data` a data.frame to partition

`threshold` the minimum proportion of information explained by a reduced variable; `threshold` sets a boundary for information loss because each reduced variable must explain at least as much as `threshold` as measured by the metric.

`partitioner` a partitioner. See the `part_*`() functions and `as_partitioner()`.

`tolerance` a small tolerance within the threshold; if a reduction is within the threshold plus/minus the tolerance, it will reduce.

niter	the number of iterations. By default, it is calculated as 20% of the number of variables or 10, whichever is larger.
x	the prefix of the new variable names
.sep	a character vector that separates x from the number (e.g. "reduced_var_1").

### Details

`partition()` uses an approach called Direct-Measure-Reduce. Directors tell the partition algorithm what to reduce, metrics tell it whether or not there will be enough information left after the reduction, and reducers tell it how to reduce the data. Together these are called a partitioner. The default partitioner for `partition()` is `part_icc()`: it finds pairs of variables to reduce by finding the pair with the minimum distance between them, it measures information loss through ICC, and it reduces data using scaled row means. There are several other partitioners available (`part_*` functions), and you can create custom partitioners with `as_partitioner()` and `replace_partitioner()`.

### Value

a partition object

### References

Millstein, Joshua, Francesca Battaglin, Malcolm Barrett, Shu Cao, Wu Zhang, Sebastian Stintzing, Volker Heinemann, and Heinz-Josef Lenz. 2020. "Partition: A Surjective Mapping Approach for Dimensionality Reduction." *Bioinformatics* 36 (3): <https://doi.org/676-81.10.1093/bioinformatics/btz661>.

Barrett, Malcolm and Joshua Millstein (2020). `partition`: A fast and flexible framework for data reduction in R. *Journal of Open Source Software*, 5(47), 1991, <https://doi.org/10.21105/joss.01991>

### See Also

`part_icc()`, `part_kmeans()`, `part_minr2()`, `part_pc1()`, `part_stdmi()`, `as_partitioner()`, `replace_partitioner()`

### Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

# don't accept reductions where information < .6
prt <- partition(df, threshold = .6)
prt

# return reduced data
partition_scores(prt)

# access mapping keys
mapping_key(prt)
unnest_mappings(prt)

# use a lower threshold of information loss
```

```
partition(df, threshold = .5, partitioner = part_kmeans())

# use a custom partitioner
part_icc_rowmeans <- replace_partitioner(part_icc, reduce = as_reducer(rowMeans))
partition(df, threshold = .6, partitioner = part_icc_rowmeans)
```

---

partition_scores	<i>Return the reduced data from a partition</i>
------------------	---

---

## Description

The reduced data is stored as `reduced_data` in the partition object and can thus be returned by subsetting `object$reduced_data`. Alternatively, the functions `partition_score()` and `fitted()` also return the reduced data.

## Usage

```
partition_scores(object, ...)

## S3 method for class 'partition'
fitted(object, ...)
```

## Arguments

<code>object</code>	a partition object
<code>...</code>	not currently used (for S3 consistency with <code>fitted()</code> )

## Value

a tibble containing the reduced data for the partition

## Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)
# fit partition
prt <- partition(df, threshold = .6)

# three ways to retrieve reduced data
partition_scores(prt)
fitted(prt)
prt$reduced_data
```

---

`part_icc`*Partitioner: distance, ICC, scaled means*

---

## Description

Partitioners are functions that tell the partition algorithm 1) what to try to reduce 2) how to measure how much information is lost from the reduction and 3) how to reduce the data. In partition, functions that handle 1) are called directors, functions that handle 2) are called metrics, and functions that handle 3) are called reducers. `partition` has a number of pre-specified partitioners for agglomerative data reduction. Custom partitioners can be created with `as_partitioner()`.

Pass partitioner objects to the `partitioner` argument of `partition()`.

`part_icc()` uses the following direct-measure-reduce approach:

- **direct:** `direct_distance()`, Minimum Distance
- **measure:** `measure_icc()`, Intraclass Correlation
- **reduce:** `reduce_scaled_mean()`, Scaled Row Means

## Usage

```
part_icc(spearman = FALSE)
```

## Arguments

`spearman`            logical. Use Spearman's correlation for distance matrix?

## Value

a partitioner

## See Also

Other partitioners: `as_partitioner()`, `part_kmeans()`, `part_minr2()`, `part_pc1()`, `part_stdmi()`, `replace_partitioner()`

## Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

# fit partition using part_icc()
partition(df, threshold = .6, partitioner = part_icc())
```

---

part_kmeans	<i>Partitioner: K-means, ICC, scaled means</i>
-------------	--

---

### Description

Partitioners are functions that tell the partition algorithm 1) what to try to reduce 2) how to measure how much information is lost from the reduction and 3) how to reduce the data. In partition, functions that handle 1) are called directors, functions that handle 2) are called metrics, and functions that handle 3) are called reducers. `partition` has a number of pre-specified partitioners for agglomerative data reduction. Custom partitioners can be created with `as_partitioner()`.

Pass partitioner objects to the partitioner argument of `partition()`.

`part_kmeans()` uses the following direct-measure-reduce approach:

- **direct:** `direct_k_cluster()`, K-Means Clusters
- **measure:** `measure_min_icc()`, Minimum Intraclass Correlation
- **reduce:** `reduce_kmeans()`, Scaled Row Means

### Usage

```
part_kmeans(
  algorithm = c("armadillo", "Hartigan-Wong", "Lloyd", "Forgy", "MacQueen"),
  search = c("binary", "linear"),
  init_k = NULL,
  n_hits = 4
)
```

### Arguments

algorithm	The K-Means algorithm to use. The default is a fast version of the LLoyd algorithm written in <code>armadillo</code> . The rest are options in <code>kmeans()</code> . In general, <code>armadillo</code> is fastest, but the other algorithms can be faster in high dimensions.
search	The search method. Binary search is generally more efficient but linear search can be faster in very low dimensions.
init_k	The initial k to test. If NULL, then the initial k is the threshold times the number of variables.
n_hits	In linear search method, the number of iterations that should be under the threshold before reducing; useful for preventing false positives.

### Value

a partitioner

### See Also

Other partitioners: `as_partitioner()`, `part_icc()`, `part_minr2()`, `part_pc1()`, `part_stdmi()`, `replace_partitioner()`



**Examples**

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

# fit partition using part_kmeans()
partition(df, threshold = .6, partitioner = part_kmeans())
```

---

part_minr2	<i>Partitioner: distance, minimum R-squared, scaled means</i>
------------	---

---

**Description**

Partitioners are functions that tell the partition algorithm 1) what to try to reduce 2) how to measure how much information is lost from the reduction and 3) how to reduce the data. In partition, functions that handle 1) are called directors, functions that handle 2) are called metrics, and functions that handle 3) are called reducers. `partition` has a number of pre-specified partitioners for agglomerative data reduction. Custom partitioners can be created with [as\\_partitioner\(\)](#).

Pass partitioner objects to the partitioner argument of [partition\(\)](#).

`part_minr2()` uses the following direct-measure-reduce approach:

- **direct:** `direct_distance()`, Minimum Distance
- **measure:** `measure_min_r2()`, Minimum R-Squared
- **reduce:** `reduce_scaled_mean()`, Scaled Row Means

**Usage**

```
part_minr2(spearman = FALSE)
```

**Arguments**

`spearman`            logical. Use Spearman's correlation for distance matrix?

**Value**

a partitioner

**See Also**

Other partitioners: [as\\_partitioner\(\)](#), [part\\_icc\(\)](#), [part\\_kmeans\(\)](#), [part\\_pc1\(\)](#), [part\\_stdmi\(\)](#), [replace\\_partitioner\(\)](#)

**Examples**

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

# fit partition using part_minr2()
partition(df, threshold = .6, partitioner = part_minr2())
```

---

part\_pc1

*Partitioner: distance, first principal component, scaled means*


---

**Description**

Partitioners are functions that tell the partition algorithm 1) what to try to reduce 2) how to measure how much information is lost from the reduction and 3) how to reduce the data. In partition, functions that handle 1) are called directors, functions that handle 2) are called metrics, and functions that handle 3) are called reducers. `partition` has a number of pre-specified partitioners for agglomerative data reduction. Custom partitioners can be created with `as_partitioner()`.

Pass partitioner objects to the partitioner argument of `partition()`.

`part_pc1()` uses the following direct-measure-reduce approach:

- **direct:** `direct_distance()`, Minimum Distance
- **measure:** `measure_variance_explained()`, Variance Explained (PCA)
- **reduce:** `reduce_first_component()`, First Principal Component

**Usage**

```
part_pc1(spearman = FALSE)
```

**Arguments**

`spearman`            logical. Use Spearman's correlation for distance matrix?

**Value**

a partitioner

**See Also**

Other partitioners: `as_partitioner()`, `part_icc()`, `part_kmeans()`, `part_minr2()`, `part_stdmi()`, `replace_partitioner()`

## Examples

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

# fit partition using part_pc1()
partition(df, threshold = .6, partitioner = part_pc1())
```

---

part_stdmi	<i>Partitioner: distance, mutual information, scaled means</i>
------------	--

---

## Description

Partitioners are functions that tell the partition algorithm 1) what to try to reduce 2) how to measure how much information is lost from the reduction and 3) how to reduce the data. In partition, functions that handle 1) are called directors, functions that handle 2) are called metrics, and functions that handle 3) are called reducers. `partition` has a number of pre-specified partitioners for agglomerative data reduction. Custom partitioners can be created with [as\\_partitioner\(\)](#).

Pass partitioner objects to the partitioner argument of [partition\(\)](#).

`part_stdmi()` uses the following direct-measure-reduce approach:

- **direct:** `direct_distance()`, Minimum Distance
- **measure:** `measure_std_mutualinfo()`, Standardized Mutual Information
- **reduce:** `reduce_scaled_mean()`, Scaled Row Means

## Usage

```
part_stdmi(spearman = FALSE)
```

## Arguments

`spearman`            logical. Use Spearman's correlation for distance matrix?

## Value

a partitioner

## See Also

Other partitioners: [as\\_partitioner\(\)](#), [part\\_icc\(\)](#), [part\\_kmeans\(\)](#), [part\\_minr2\(\)](#), [part\\_pc1\(\)](#), [replace\\_partitioner\(\)](#)

**Examples**

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

# fit partition using part_stdmi()
partition(df, threshold = .6, partitioner = part_stdmi())
```

---

permute\_df                      *Permute a data set*

---

**Description**

permute\_df() permutes a data set: it randomizes the order within each variable, which breaks any association between them. Permutation is useful for testing against null statistics.

**Usage**

```
permute_df(.data)
```

**Arguments**

.data                      a data.frame

**Value**

a permuted data.frame

**Examples**

```
permute_df(iris)
```

---

plot\_area\_clusters              *Plot partitions*

---

**Description**

plot\_stacked\_area\_clusters() and plot\_area\_clusters() plot the partition against a permuted partition. plot\_ncluster() plots the number of variables per cluster. If .partition is the result of map\_partition() or test\_permutation(), plot\_ncluster() facets the plot by each partition. plot\_information() plots a histogram or density plot of the information of each variable in the partition. If .partition is the result of map\_partition() or test\_permutation(), plot\_information() plots a scatterplot of the targeted vs. observed information with a 45 degree line indicating perfect alignment.

**Usage**

```

plot_area_clusters(
  .data,
  partitioner = part_icc(),
  information = seq(0.1, 0.5, length.out = 25),
  ...,
  obs_color = "#E69F00",
  perm_color = "#56B4E9"
)

plot_stacked_area_clusters(
  .data,
  partitioner = part_icc(),
  information = seq(0.1, 0.5, length.out = 25),
  ...,
  stack_colors = c("#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2", "#D55E00")
)

plot_ncluster(
  .partition,
  show_n = 100,
  fill = "#0172B1",
  color = NA,
  labeller = "target information:"
)

plot_information(
  .partition,
  fill = "#0172B1",
  color = NA,
  geom = ggplot2::geom_density
)

```

**Arguments**

.data	a data.frame to partition
partitioner	a partitioner. See the <code>part_*()</code> functions and <code>as_partitioner()</code> .
information	a vector of minimum information to fit in <code>partition()</code>
...	arguments passed to <code>partition()</code>
obs_color	the color of the observed partition
perm_color	the color of the permuted partition
stack_colors	the colors of the cluster sizes
.partition	either a partition or a tibble, the result of <code>map_partition()</code> or <code>test_permutation()</code>
show_n	the number of reduced variables to plot
fill	the color of the fill for geom

color	the color of the geom
labeller	the facet label
geom	the geom to use. The default is geom_density.

**Value**

a ggplot

**Examples**

```
set.seed(123)
df <- simulate_block_data(c(3, 4, 5), lower_corr = .4, upper_corr = .6, n = 100)

df %>%
  partition(.6, partitioner = part_pc1()) %>%
  plot_ncluster()
```

---

plot_permutation	<i>Plot permutation tests</i>
------------------	-------------------------------

---

**Description**

plot\_permutation() takes the results of [test\\_permutation\(\)](#) and plots the distribution of permuted partitions compared to the observed partition.

**Usage**

```
plot_permutation(
  permutations,
  .plot = c("information", "nclusters", "nreduced"),
  labeller = "target information:",
  perm_color = "#56B4EA",
  obs_color = "#CC78A8",
  geom = ggplot2::geom_density
)
```

**Arguments**

permutations	a tibble, the result of <a href="#">test_permutation()</a>
.plot	the variable to plot: observed information, the number of clusters created, or the number of observed variables reduced
labeller	the facet label
perm_color	the color of the permutation fill
obs_color	the color of the observed statistic line
geom	the geom to use. The default is geom_density.

**Value**

a ggplot

---

reduce_cluster	<i>Reduce a target</i>
----------------	------------------------

---

**Description**

reduce\_cluster() and map\_cluster() apply the data reduction to the targets found in the director step. They only do so if the metric is above the threshold, however. reduce\_cluster() is for functions that return vectors while map\_cluster() is for functions that return data.frames. If you're using [as\\_reducer\(\)](#), there's no need to call these functions directly.

**Usage**

```
reduce_cluster(.partition_step, .f, first_match = FALSE)
```

```
map_cluster(.partition_step, .f, rewind = FALSE, first_match = FALSE)
```

**Arguments**

.partition_step	a partition_step object
.f	a function to reduce the data to either a vector or a data.frame
first_match	logical. Should the partition algorithm stop when it finds a reduction that is equal to the threshold? Default is TRUE for reducers that return a data.frame and FALSE for reducers that return a vector
rewind	logical. Should the last target be used instead of the current target?

**Value**

a partition\_step object

**Examples**

```
reduce_row_means <- function(.partition_step, .data) {
  reduce_cluster(.partition_step, rowMeans)
}

replace_partitioner(
  part_icc,
  reduce = reduce_row_means
)
```

---

 reduce\_first\_component

*Reduce selected variables to first principal component*


---

### Description

Reducers are functions that tell the partition algorithm how to reduce the data. [as\\_reducer\(\)](#) is a helper function to create new reducers to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`reduce_first_component()` returns the first component from the principal components analysis of the target variables. Because the PCA calculates the components and the variance explained at the same time, if the metric is `measure_variance_explained()`, that function will store the first component for use in `reduce_first_component()` to avoid recalculation. If the partitioner uses a different metric, the first component will be calculated by `reduce_first_component()`.

### Usage

```
reduce_first_component(.partition_step)
```

### Arguments

```
.partition_step
      a partition_step object
```

### Value

a partition\_step object

### See Also

Other reducers: [as\\_reducer\(\)](#), [reduce\\_kmeans\(\)](#), [reduce\\_scaled\\_mean\(\)](#)

---

 reduce\_kmeans

*Reduce selected variables to scaled means*


---

### Description

Reducers are functions that tell the partition algorithm how to reduce the data. [as\\_reducer\(\)](#) is a helper function to create new reducers to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`reduce_kmeans()` is efficient in that it doesn't reduce until the closest k to the information threshold is found.

### Usage

```
reduce_kmeans(.partition_step, search = c("binary", "linear"), n_hits = 4)
```



**Arguments**

<code>.partition_step</code>	a <code>partition_step</code> object
<code>search</code>	The search method. Binary search is generally more efficient but linear search can be faster in very low dimensions.
<code>n_hits</code>	In linear search method, the number of iterations that should be under the threshold before reducing; useful for preventing false positives.

**Value**

a `partition_step` object

**See Also**

Other reducers: [as\\_reducer\(\)](#), [reduce\\_first\\_component\(\)](#), [reduce\\_scaled\\_mean\(\)](#)

---

<code>reduce_scaled_mean</code>	<i>Reduce selected variables to scaled means</i>
---------------------------------	--

---

**Description**

Reducers are functions that tell the partition algorithm how to reduce the data. [as\\_reducer\(\)](#) is a helper function to create new reducers to be used in partitioners. partitioners can be created with [as\\_partitioner\(\)](#).

`reduce_scaled_mean()` returns the scaled row means of the target variables to reduce.

**Usage**

```
reduce_scaled_mean(.partition_step)
```

**Arguments**

<code>.partition_step</code>	a <code>partition_step</code> object
------------------------------	--------------------------------------

**Value**

a `partition_step` object

**See Also**

Other reducers: [as\\_reducer\(\)](#), [reduce\\_first\\_component\(\)](#), [reduce\\_kmeans\(\)](#)

---

replace\_partitioner     *Replace the director, metric, or reducer for a partitioner*

---

### Description

Replace the director, metric, or reducer for a partitioner

### Usage

```
replace_partitioner(partitioner, direct = NULL, measure = NULL, reduce = NULL)
```

### Arguments

partitioner	a partitioner
direct	a function that directs, possibly created by <a href="#">as_director()</a>
measure	a function that measures, possibly created by <a href="#">as_measure()</a>
reduce	a function that reduces, possibly created by <a href="#">as_reducer()</a>

### Value

a partitioner

### See Also

Other partitioners: [as\\_partitioner\(\)](#), [part\\_icc\(\)](#), [part\\_kmeans\(\)](#), [part\\_minr2\(\)](#), [part\\_pc1\(\)](#), [part\\_stdmi\(\)](#)

### Examples

```
replace_partitioner(
  part_icc,
  reduce = as_reducer(rowMeans)
)
```

---

scaled\_mean     *Average and scale rows in a data.frame*

---

### Description

scaled\_mean() calculates scaled row means for a dataframe.

### Usage

```
scaled_mean(.x, method = c("r", "c"))
```

**Arguments**

.x                    a data.frame  
 method              The method source: both the pure R and C++ versions are efficient

**Value**

a numeric vector

**Examples**

```
library(dplyr)
iris %>%
  select_if(is.numeric) %>%
  scaled_mean()
```

---

simulate\_block\_data    *Simulate correlated blocks of variables*

---

**Description**

simulate\_block\_data() creates a dataset of blocks of data where variables within each block are correlated. The correlation for each pair of variables is sampled uniformly from lower\_corr to upper\_corr, and the values of each are sampled using [MASS: :mvrnorm\(\)](#).

**Usage**

```
simulate_block_data(
  block_sizes,
  lower_corr,
  upper_corr,
  n,
  block_name = "block",
  sep = "_",
  var_name = "x"
)
```

**Arguments**

block\_sizes        a vector of block sizes. The size of each block is the number of variables within it.  
 lower\_corr        the lower bound of the correlation within each block  
 upper\_corr        the upper bound of the correlation within each block  
 n                  the number of observations or rows  
 block\_name        description prepended to the variable to indicate the block it belongs to  
 sep                a character, what to separate the variable names with  
 var\_name          the name of the variable within the block

**Value**

a tibble with `sum(block_sizes)` columns and `n` rows.

**Examples**

```
# create a 100 x 15 data set with 3 blocks
simulate_block_data(
  block_sizes = rep(5, 3),
  lower_corr = .4,
  upper_corr = .6,
  n = 100
)
```

---

super_partition	<i>super_partition</i>
-----------------	------------------------

---

**Description**

`super_partition` implements the agglomerative, data reduction method `Partition` for datasets with large numbers of features by first 'super-partitioning' the data into smaller clusters to `Partition`.

**Usage**

```
super_partition(
  full_data,
  threshold = 0.5,
  cluster_size = 4000,
  partitioner = part_icc(),
  tolerance = 1e-04,
  niter = NULL,
  x = "reduced_var",
  .sep = "_",
  verbose = TRUE,
  progress_bar = TRUE
)
```

**Arguments**

<code>full_data</code>	sample by feature data frame or matrix
<code>threshold</code>	the minimum proportion of information explained by a reduced variable; <code>threshold</code> sets a boundary for information loss because each reduced variable must explain at least as much as <code>threshold</code> as measured by the metric.
<code>cluster_size</code>	maximum size of any single cluster; default is 4000
<code>partitioner</code>	a partitioner. See the <code>part_*</code> () functions and <code>as_partitioner()</code> .
<code>tolerance</code>	a small tolerance within the threshold; if a reduction is within the threshold plus/minus the tolerance, it will reduce.

niter	the number of iterations. By default, it is calculated as 20% of the number of variables or 10, whichever is larger.
x	the prefix of the new variable names; must not be contained in any existing data names
.sep	a character vector that separates x from the number (e.g. "reduced_var_1").
verbose	logical for whether or not to display information about super partition step; default is TRUE
progress_bar	logical for whether or not to show progress bar; default is TRUE

### Details

super\_partition scales up partition with an approximation, using Genie, a fast, hierarchical clustering algorithm with similar qualities of those to Partition, to first super-partition the data into  $\text{ceiling}(N/c)$  clusters, where  $N$  is the number of features in the full dataset and  $c$  is the user-defined maximum cluster size (default value = 4,000). Then, if any cluster from the super-partition has a size greater than  $c$ , use Genie again on that cluster until all cluster sizes are less than  $c$ . Finally, apply the Partition algorithm to each of the super-partitions.

It may be the case that large super-partitions cannot be easily broken with Genie due to high similarity between features. In this case, we use k-means to break the cluster.

### Value

Partition object

### Author(s)

Katelyn Queen, <kjqueen@usc.edu>

### References

- Barrett, Malcolm and Joshua Millstein (2020). partition: A fast and flexible framework for data reduction in R. *Journal of Open Source Software*, 5(47), 1991, <https://doi.org/10.21105/joss.01991> Millstein J, Battaglin F, Barrett M, Cao S, Zhang W, Stintzing S, et al. Partition: a surjective mapping approach for dimensionality reduction. *Bioinformatics* 36 (2019) 676–681. doi:10.1093/bioinformatics/btz661.
- Gagolewski, Marek, Maciej Bartoszuk, and Anna Cena. "Genie: A new, fast, and outlier-resistant hierarchical clustering algorithm." *Information Sciences* 363 (2016): 8-23.
- Millstein, Joshua, Francesca Battaglin, Malcolm Barrett, Shu Cao, Wu Zhang, Sebastian Stintzing, Volker Heinemann, and Heinz-Josef Lenz. 2020. "Partition: A Surjective Mapping Approach for Dimensionality Reduction." *Bioinformatics* 36 (3): <https://doi.org/676–81.10.1093/bioinformatics/btz661>.

### See Also

[partition\(\)](#)

**Examples**

```
set.seed(123)
df <- simulate_block_data(c(15, 20, 10), lower_corr = .4, upper_corr = .6, n = 100)

# don't accept reductions where information < .6
prt <- super_partition(df, threshold = .6, cluster_size = 30)
prt
```

---

test_permutation	<i>Permute partitions</i>
------------------	---------------------------

---

**Description**

`test_permutation()` permutes data and partitions the results to generate a distribution of null statistics for observed information, number of clusters, and number of observed variables reduced to clusters. The result is a tibble with a summary of the observed data results and the averages of the permuted results. The partitions and and permutations are also available in `list-cols`. `test_permutation()` tests across a range of target information values, as specified in the `information` argument.

**Usage**

```
test_permutation(
  .data,
  information = seq(0.1, 0.6, by = 0.1),
  partitioner = part_icc(),
  ...,
  nperm = 100
)
```

**Arguments**

<code>.data</code>	a data set to partition
<code>information</code>	a vector of minimum information to fit in <code>partition()</code>
<code>partitioner</code>	the partitioner to use. The default is <code>part_icc()</code> .
<code>...</code>	arguments passed to <code>partition()</code>
<code>nperm</code>	Number of permuted data sets to test. Default is 100.

**Value**

a tibble with summaries on observed and permuted data (the means of the permuted summaries), as well as `list-cols` containing them

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