

# Package: healthyR.ai (via r-universe)

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**Title** The Machine Learning and AI Modeling Companion to 'healthyR'

**Version** 0.1.0.9000

**Description** Hospital machine learning and ai data analysis workflow tools, modeling, and automations. This library provides many useful tools to review common administrative hospital data. Some of these include predicting length of stay, and readmits. The aim is to provide a simple and consistent verb framework that takes the guesswork out of everything.

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**Encoding** UTF-8

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**URL** <https://www.spsanderson.com/healthyR.ai/>,  
<https://github.com/spsanderson/healthyR.ai>

**BugReports** <https://github.com/spsanderson/healthyR.ai/issues>

**Imports** magrittr, rlang (>= 0.1.2), yardstick (>= 0.0.8), utils, broom, ggrepel, tibble, dplyr, ggplot2, tidyr, forcats, recipes (>= 1.0.0), purrr, h2o, stats, dials, parsnip, tune, workflows, modeltime

**Suggests** rmarkdown, knitr, healthyR.data, scales, tidymodels, janitor, timetk, plotly, rsample, kknn, hardhat, uwot, stringr

**VignetteBuilder** knitr

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**Repository** <https://spsanderson.r-universe.dev>

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---

color_blind	<i>Provide Colorblind Compliant Colors</i>
-------------	--

---

**Description**

8 Hex RGB color definitions suitable for charts for colorblind people.

**Usage**

```
color_blind()
```

**Details**

This function is used in others in order to help render plots for those that are color blind.

**Value**

A vector of 8 Hex RGB definitions.

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Color\_Blind: [hai\\_scale\\_color\\_colorblind\(\)](#), [hai\\_scale\\_fill\\_colorblind\(\)](#)

**Examples**

```
color_blind()
```

---

generate\_mesh\_data      *Generate Mesh Data*

---

**Description**

This function creates a square mesh by sampling nodes uniformly on a square and then connecting these nodes with edges. The nodes are distributed based on the provided side length and number of segments. Horizontal, vertical, and diagonal edges are generated to fully connect the mesh. The function returns a list containing the nodes and edges, along with data frames and a ggplot object for visualization.

**Usage**

```
generate_mesh_data(.side_length = 1, .n_seg = 1)
```

**Arguments**

`.side_length`      A single numeric value representing the side length of the square.

`.n_seg`              A positive integer representing the number of segments along each side of the square.

**Details**

This function generates a mesh of nodes and edges based on the provided side length and number of segments.

This function creates a square mesh of nodes and edges, where the nodes are sampled uniformly on a square. The edges are generated to connect the nodes horizontally, vertically, and diagonally.

**Value**

A list containing:

**nodes** A matrix with coordinates of the nodes.

**edges** A list of edges connecting the nodes.

**nodes\_df** A data frame of nodes for ggplot.

**edges\_df** A data frame of edges for ggplot.

**plot** A ggplot object visualizing the nodes and edges.

Additionally, the list contains attributes:

**side\_length** The side length used to generate the mesh.

**n\_seg** The number of segments used to generate the mesh.

**nodes\_df\_dim** Dimensions of the nodes data frame.

**edges\_df\_dim** Dimensions of the edges data frame.

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Data Generation: [get\\_juiced\\_data\(\)](#)

**Examples**

```
generate_mesh_data(1, 1)
generate_mesh_data(1, 2)
```

---

get\_juiced\_data

*Get the Juiced Data*

---

**Description**

This is a simple function that will get the juiced data from a recipe.

**Usage**

```
get_juiced_data(.recipe_object)
```

**Arguments**

`.recipe_object` The recipe object you want to pass.

**Details**

Instead of typing out something like: `recipe_object %>% prep() %>% juice() %>% glimpse()`

**Value**

A tibble of the prepped and juiced data from the given recipe

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Data Generation: [generate\\_mesh\\_data\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(timetk))
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(purrr))
suppressPackageStartupMessages(library(healthyR.data))
suppressPackageStartupMessages(library(rsample))
suppressPackageStartupMessages(library(recipes))

data_tbl <- healthyR_data %>%
  select(visit_end_date_time) %>%
  summarise_by_time(
    .date_var = visit_end_date_time,
    .by       = "month",
    value     = n()
  ) %>%
  set_names("date_col", "value") %>%
  filter_by_time(
    .date_var = date_col,
    .start_date = "2013",
    .end_date = "2020"
  )

splits <- initial_split(data = data_tbl, prop = 0.8)

rec_obj <- recipe(value ~ ., training(splits))

get_juiced_data(rec_obj)
```

---

 hai\_auto\_c50 *Boilerplate Workflow*


---

**Description**

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

**Usage**

```
hai_auto_c50(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_c50_data_prepper()</code> an automatic <code>recipe_object</code> .
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "f_meas". You can choose a metric depending on the <code>model_type</code> used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
<code>.model_type</code>	Default is classification, can also be regression.

**Details**

This uses the `parsnip::boost_tree()` with the engine set to `C5.0`

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Boiler\_Plate: [hai\\_auto\\_cubist\(\)](#), [hai\\_auto\\_earth\(\)](#), [hai\\_auto\\_glmnet\(\)](#), [hai\\_auto\\_knn\(\)](#), [hai\\_auto\\_ranger\(\)](#), [hai\\_auto\\_svm\\_poly\(\)](#), [hai\\_auto\\_svm\\_rbf\(\)](#), [hai\\_auto\\_wflw\\_metrics\(\)](#), [hai\\_auto\\_xgboost\(\)](#)

Other C5.0: [hai\\_c50\\_data\\_prepper\(\)](#)

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_c50_data_prepper(data, Species ~ .)

auto_c50 <- hai_auto_c50(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas",
  .model_type = "classification"
)

auto_c50$recipe_info

## End(Not run)
```

---

hai\_auto\_cubist

*Boilerplate Workflow*

---

**Description**

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)



**Usage**

```
hai_auto_cubist(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "rmse"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_cubist_data_prepper()</code> an automatic <code>recipe_object</code> .
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "rmse". The only <code>.model_type</code> you can use with Cubist is regression so use <code>hai_default_regression_metric_set()</code> to get the available metrics. Because of this the <code>.model_type</code> parameter is omitted from this function.

**Details**

This uses the `parsnip::cubist_rules()` with the engine set to `cubist`

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Boiler\_Plate: [hai\\_auto\\_c50\(\)](#), [hai\\_auto\\_earth\(\)](#), [hai\\_auto\\_glmnet\(\)](#), [hai\\_auto\\_knn\(\)](#), [hai\\_auto\\_ranger\(\)](#), [hai\\_auto\\_svm\\_poly\(\)](#), [hai\\_auto\\_svm\\_rbf\(\)](#), [hai\\_auto\\_wflw\\_metrics\(\)](#), [hai\\_auto\\_xgboost\(\)](#)

Other cubist: [hai\\_cubist\\_data\\_prepper\(\)](#)

## Examples

```
## Not run:
data <- mtcars

rec_obj <- hai_cubist_data_prepper(data, mpg ~ .)

auto_cube <- hai_auto_cubist(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "rmse"
)

auto_cube$recipe_info

## End(Not run)
```

---

hai\_auto\_earth

*Boilerplate Workflow*

---

## Description

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

## Usage

```
hai_auto_earth(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_earth_data_prepper()</code> an automatic recipe_object.
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "f_meas". You can choose a metric depending on the model_type used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
<code>.model_type</code>	Default is classification, can also be regression.

**Details**

This uses the `parsnip::mars()` with the engine set to earth

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<http://uc-r.github.io/mars>

Other Boiler\_Plate: `hai_auto_c50()`, `hai_auto_cubist()`, `hai_auto_glmnet()`, `hai_auto_knn()`, `hai_auto_ranger()`, `hai_auto_svm_poly()`, `hai_auto_svm_rbf()`, `hai_auto_wflw_metrics()`, `hai_auto_xgboost()`

Other Earth: `hai_earth_data_prepper()`

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_earth_data_prepper(data, Species ~ .)

auto_earth <- hai_auto_earth(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas",
  .model_type = "classification"
```

```

)
auto_earth$recipe_info

## End(Not run)

```

---

 hai\_auto\_glmnet

*Boilerplate Workflow*


---

## Description

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

## Usage

```

hai_auto_glmnet(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)

```

## Arguments

.data	The data being passed to the function. The time-series object.
.rec_obj	This is the recipe object you want to use. You can use <code>hai_glmnet_data_prepper()</code> an automatic <code>recipe_object</code> .
.splits_obj	NULL is the default, when NULL then one will be created.
.rsamp_obj	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
.tune	Default is TRUE, this will create a tuning grid and tuned workflow
.grid_size	Default is 10
.num_cores	Default is 1
.best_metric	Default is "f_meas". You can choose a metric depending on the <code>model_type</code> used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
.model_type	Default is classification, can also be regression.

**Details**

This uses the `parsnip::multinom_reg()` with the engine set to `glmnet`

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Boiler\_Plate: [hai\\_auto\\_c50\(\)](#), [hai\\_auto\\_cubist\(\)](#), [hai\\_auto\\_earth\(\)](#), [hai\\_auto\\_knn\(\)](#), [hai\\_auto\\_ranger\(\)](#), [hai\\_auto\\_svm\\_poly\(\)](#), [hai\\_auto\\_svm\\_rbf\(\)](#), [hai\\_auto\\_wflw\\_metrics\(\)](#), [hai\\_auto\\_xgboost\(\)](#)

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_glmnet_data_prepper(data, Species ~ .)

auto_glm <- hai_auto_glmnet(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas",
  .model_type = "classification"
)

auto_glm$recipe_info

## End(Not run)
```

---

hai\_auto\_knn

*Boilerplate Workflow*

---

**Description**

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

**Usage**

```
hai_auto_knn(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "rmse",
  .model_type = "regression"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_knn_data_prepper()</code> an automatic <code>recipe_object</code> .
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "rmse". You can choose a metric depending on the <code>model_type</code> used. If regression then see <a href="#">hai_default_regression_metric_set()</a> , if classification then see <a href="#">hai_default_classification_metric_set()</a> .
<code>.model_type</code>	Default is regression, can also be classification.

**Details**

This uses the `parsnip::nearest_neighbor()` with the engine set to `kkn`

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Boiler\_Plate: [hai\\_auto\\_c50\(\)](#), [hai\\_auto\\_cubist\(\)](#), [hai\\_auto\\_earth\(\)](#), [hai\\_auto\\_glmnet\(\)](#), [hai\\_auto\\_ranger\(\)](#), [hai\\_auto\\_svm\\_poly\(\)](#), [hai\\_auto\\_svm\\_rbf\(\)](#), [hai\\_auto\\_wflw\\_metrics\(\)](#), [hai\\_auto\\_xgboost\(\)](#)

**Examples**

```
## Not run:
library(dplyr)

data <- iris

rec_obj <- hai_knn_data_prepper(data, Species ~ .)

auto_knn <- hai_auto_knn(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas",
  .model_type = "classification"
)

auto_knn$recipe_info

## End(Not run)
```

---

hai\_auto\_ranger

*Boilerplate Workflow*

---

**Description**

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

**Usage**

```
hai_auto_ranger(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_ranger_data_prepper()</code> an automatic recipe_object.
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "f_meas". You can choose a metric depending on the model_type used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
<code>.model_type</code>	Default is classification, can also be regression.

**Details**

This uses the `parsnip::rand_forest()` with the engine set to kernlab

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/rand\\_forest.html](https://parsnip.tidymodels.org/reference/rand_forest.html)

Other Boiler\_Plate: `hai_auto_c50()`, `hai_auto_cubist()`, `hai_auto_earth()`, `hai_auto_glmnet()`, `hai_auto_knn()`, `hai_auto_svm_poly()`, `hai_auto_svm_rbf()`, `hai_auto_wflw_metrics()`, `hai_auto_xgboost()`

Other Ranger: `hai_ranger_data_prepper()`

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_ranger_data_prepper(data, Species ~ .)

auto_ranger <- hai_auto_ranger(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas"
)
```



```

auto_ranger$recipe_info

## End(Not run)

```

---

```

hai_auto_svm_poly      Boilerplate Workflow

```

---

## Description

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

## Usage

```

hai_auto_svm_poly(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)

```

## Arguments

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_svm_poly_data_prepper()</code> an automatic recipe object.
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "f_meas". You can choose a metric depending on the <code>model_type</code> used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
<code>.model_type</code>	Default is classification, can also be regression.

**Details**

This uses the `parsnip::svm_poly()` with the engine set to `kernlab`

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/svm\\_poly.html](https://parsnip.tidymodels.org/reference/svm_poly.html)

Other Boiler\_Plate: [hai\\_auto\\_c50\(\)](#), [hai\\_auto\\_cubist\(\)](#), [hai\\_auto\\_earth\(\)](#), [hai\\_auto\\_glmnet\(\)](#), [hai\\_auto\\_knn\(\)](#), [hai\\_auto\\_ranger\(\)](#), [hai\\_auto\\_svm\\_rbf\(\)](#), [hai\\_auto\\_wflw\\_metrics\(\)](#), [hai\\_auto\\_xgboost\(\)](#)

Other SVM\_Poly: [hai\\_svm\\_poly\\_data\\_prepper\(\)](#)

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_svm_poly_data_prepper(data, Species ~ .)

auto_svm_poly <- hai_auto_svm_poly(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas"
)

auto_svm_poly$recipe_info

## End(Not run)
```

---

hai\_auto\_svm\_rbf      *Boilerplate Workflow*

---

**Description**

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

**Usage**

```
hai_auto_svm_rbf(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_svm_rbf_data_prepper()</code> an automatic <code>recipe_object</code> .
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "f_meas". You can choose a metric depending on the <code>model_type</code> used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
<code>.model_type</code>	Default is classification, can also be regression.

**Details**

This uses the `parsnip::svm_rbf()` with the engine set to `kernlab`

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/svm\\_rbf.html](https://parsnip.tidymodels.org/reference/svm_rbf.html)

Other Boiler\_Plate: `hai_auto_c50()`, `hai_auto_cubist()`, `hai_auto_earth()`, `hai_auto_glmnet()`, `hai_auto_knn()`, `hai_auto_ranger()`, `hai_auto_svm_poly()`, `hai_auto_wflw_metrics()`, `hai_auto_xgboost()`

Other SVM\_RBF: `hai_svm_rbf_data_prepper()`

## Examples

```
## Not run:
data <- iris

rec_obj <- hai_svm_rbf_data_prepper(data, Species ~ .)

auto_rbf <- hai_auto_svm_rbf(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas"
)

auto_rbf$recipe_info

## End(Not run)
```

---

hai\_auto\_wflw\_metrics *Collect Metrics from Boilerplat Workflows*

---

## Description

This function will extract the metrics from the hai\_auto\_ boilerplate functions.

## Usage

```
hai_auto_wflw_metrics(.data)
```

## Arguments

.data            The output of the hai\_auto\_ boilerplate function in it's entirety.

## Details

This function will extract the metrics from the hai\_auto\_ boilerplate functions. This function looks for a specific attribute from the hai\_auto\_ functions so that it will extract the tuned\_results from the tuning process if it has indeed been tuned.

## Value

A tibble

## Author(s)

Steven P. Sanderson II, MPH

## See Also

Other Boiler\_Plate: [hai\\_auto\\_c50\(\)](#), [hai\\_auto\\_cubist\(\)](#), [hai\\_auto\\_earth\(\)](#), [hai\\_auto\\_glmnet\(\)](#), [hai\\_auto\\_knn\(\)](#), [hai\\_auto\\_ranger\(\)](#), [hai\\_auto\\_svm\\_poly\(\)](#), [hai\\_auto\\_svm\\_rbf\(\)](#), [hai\\_auto\\_xgboost\(\)](#)

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_knn_data_prepper(data, Species ~ .)

auto_knn <- hai_auto_knn(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas",
  .model_type = "classification",
  .grid_size = 2,
  .num_cores = 4
)

hai_auto_wflw_metrics(auto_knn)

## End(Not run)
```

---

hai\_auto\_xgboost

*Boilerplate Workflow*

---

**Description**

This is a boilerplate function to create automatically the following:

- recipe
- model specification
- workflow
- tuned model (grid ect)

**Usage**

```
hai_auto_xgboost(
  .data,
  .rec_obj,
  .splits_obj = NULL,
  .rsamp_obj = NULL,
  .tune = TRUE,
  .grid_size = 10,
  .num_cores = 1,
  .best_metric = "f_meas",
  .model_type = "classification"
)
```

**Arguments**

<code>.data</code>	The data being passed to the function. The time-series object.
<code>.rec_obj</code>	This is the recipe object you want to use. You can use <code>hai_xgboost_data_prepper()</code> an automatic recipe_object.
<code>.splits_obj</code>	NULL is the default, when NULL then one will be created.
<code>.rsamp_obj</code>	NULL is the default, when NULL then one will be created. It will default to creating an <code>rsample::mc_cv()</code> object.
<code>.tune</code>	Default is TRUE, this will create a tuning grid and tuned workflow
<code>.grid_size</code>	Default is 10
<code>.num_cores</code>	Default is 1
<code>.best_metric</code>	Default is "f_meas". You can choose a metric depending on the model_type used. If regression then see <code>hai_default_regression_metric_set()</code> , if classification then see <code>hai_default_classification_metric_set()</code> .
<code>.model_type</code>	Default is classification, can also be regression.

**Details**

This uses the `parsnip::boost_tree()` with the engine set to xgboost

**Value**

A list

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/details\\_boost\\_tree\\_xgboost.html](https://parsnip.tidymodels.org/reference/details_boost_tree_xgboost.html)

Other Boiler\_Plate: `hai_auto_c50()`, `hai_auto_cubist()`, `hai_auto_earth()`, `hai_auto_glmnet()`, `hai_auto_knn()`, `hai_auto_ranger()`, `hai_auto_svm_poly()`, `hai_auto_svm_rbf()`, `hai_auto_wflw_metrics()`

**Examples**

```
## Not run:
data <- iris

rec_obj <- hai_xgboost_data_prepper(data, Species ~ .)

auto_xgb <- hai_auto_xgboost(
  .data = data,
  .rec_obj = rec_obj,
  .best_metric = "f_meas"
)

auto_xgb$recipe_info
```

```
## End(Not run)
```

---

hai\_c50\_data\_prepper *Prep Data for C5.0 - Recipe*

---

## Description

Automatically prep a data.frame/tibble for use in the C5.0 algorithm.

## Usage

```
hai_c50_data_prepper(.data, .recipe_formula)
```

## Arguments

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::recipe()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the iris data then the formula would most likely be something like `Species ~ .`

## Details

This function will automatically prep your data.frame/tibble for use in the C5.0 algorithm. The C5.0 algorithm is a lazy learning classification algorithm. It expects data to be presented in a certain fashion.

This function will output a recipe specification.

## Value

A recipe object

## Author(s)

Steven P. Sanderson II, MPH

## See Also

<https://www.rulequest.com/see5-unix.html>

Other Preprocessor: [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

Other C5.0: [hai\\_auto\\_c50\(\)](#)

**Examples**

```
library(ggplot2)

hai_c50_data_prepper(.data = Titanic, .recipe_formula = Survived ~ .)
rec_obj <- hai_c50_data_prepper(Titanic, Survived ~ .)
get_juiced_data(rec_obj)
```

---

hai_control_chart	<i>Create a control chart</i>
-------------------	-------------------------------

---

**Description**

Create a control chart, aka Shewhart chart: [https://en.wikipedia.org/wiki/Control\\_chart](https://en.wikipedia.org/wiki/Control_chart).

**Usage**

```
hai_control_chart(
  .data,
  .value_col,
  .x_col,
  .center_line = mean,
  .std_dev = 3,
  .plt_title = NULL,
  .plt_catpion = NULL,
  .plt_font_size = 11,
  .print_plot = TRUE
)
```

**Arguments**

.data	data frame or a path to a csv file that will be read in
.value_col	variable of interest mapped to y-axis (quoted, ie as a string)
.x_col	variable to go on the x-axis, often a time variable. If unspecified row indices will be used (quoted)
.center_line	Function used to calculate central tendency. Defaults to mean
.std_dev	Number of standard deviations above and below the central tendency to call a point influenced by "special cause variation." Defaults to 3
.plt_title	Plot title
.plt_catpion	Plot caption
.plt_font_size	Font size; text elements will be scaled to this
.print_plot	Print the plot? Default = TRUE. Set to FALSE if you want to assign the plot to a variable for further modification, as in the last example.



**Details**

Control charts, also known as Shewhart charts (after Walter A. Shewhart) or process-behavior charts, are a statistical process control tool used to determine if a manufacturing or business process is in a state of control. It is more appropriate to say that the control charts are the graphical device for Statistical Process Monitoring (SPM). Traditional control charts are mostly designed to monitor process parameters when underlying form of the process distributions are known. However, more advanced techniques are available in the 21st century where incoming data streaming can-be monitored even without any knowledge of the underlying process distributions. Distribution-free control charts are becoming increasingly popular.

**Value**

Generally called for the side effect of printing the control chart. Invisibly, returns a ggplot object for further customization.

**Author(s)**

Steven P. Sanderson II, MPH

**Examples**

```
data_tbl <- tibble::tibble(
  day = sample(
    c("Monday", "Tuesday", "Wednesday", "Thursday", "Friday"),
    100, TRUE
  ),
  person = sample(c("Tom", "Jane", "Alex"), 100, TRUE),
  count = rbinom(100, 20, ifelse(day == "Friday", .5, .2)),
  date = Sys.Date() - sample.int(100)
)

hai_control_chart(.data = data_tbl, .value_col = count, .x_col = date)

# In addition to printing or writing the plot to file, hai_control_chart
# returns the plot as a ggplot2 object, which you can then further customize

library(ggplot2)
my_chart <- hai_control_chart(data_tbl, count, date)
my_chart +
  ylab("Number of Adverse Events") +
  scale_x_date(name = "Week of ... ", date_breaks = "week") +
  theme(axis.text.x = element_text(angle = -90, vjust = 0.5, hjust = 1))
```

**Description**

Automatically prep a data.frame/tibble for use in the cubist algorithm.

**Usage**

```
hai_cubist_data_prepper(.data, .recipe_formula)
```

**Arguments**

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::recip()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the diamonds data then the formula would most likely be something like `price ~ .`

**Details**

This function will automatically prep your data.frame/tibble for use in the cubist algorithm. The cubist algorithm is for regression only.

This function will output a recipe specification.

**Value**

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<https://rulequest.com/cubist-info.html>

Other Preprocessor: `hai_c50_data_prepper()`, `hai_data_impute()`, `hai_data_poly()`, `hai_data_scale()`, `hai_data_transform()`, `hai_data_trig()`, `hai_earth_data_prepper()`, `hai_glmnet_data_prepper()`, `hai_knn_data_prepper()`, `hai_ranger_data_prepper()`, `hai_svm_poly_data_prepper()`, `hai_svm_rbf_data_prepper()`, `hai_xgboost_data_prepper()`

Other cubist: `hai_auto_cubist()`

**Examples**

```
library(ggplot2)

hai_cubist_data_prepper(.data = diamonds, .recipe_formula = price ~ .)
rec_obj <- hai_cubist_data_prepper(diamonds, price ~ .)
get_juiced_data(rec_obj)
```

---

hai_data_impute	<i>Data Preprocessor - Imputation</i>
-----------------	---------------------------------------

---

### Description

Takes in a recipe and will impute missing values using a selected recipe. To call the recipe use a quoted argument like "median" or "bagged".

### Usage

```
hai_data_impute(
  .recipe_object = NULL,
  ...,
  .seed_value = 123,
  .type_of_imputation = "mean",
  .number_of_trees = 25,
  .neighbors = 5,
  .mean_trim = 0,
  .roll_statistic,
  .roll_window = 5
)
```

### Arguments

<code>.recipe_object</code>	The data that you want to process
<code>...</code>	One or more selector functions to choose variables to be imputed. When used with <code>imp_vars</code> , these dots indicate which variables are used to predict the missing data in each variable. See <code>selections()</code> for more details
<code>.seed_value</code>	To make results reproducible, set the seed.
<code>.type_of_imputation</code>	This is a quoted argument and can be one of the following: <ul style="list-style-type: none"> <li>• "bagged"</li> <li>• "knn"</li> <li>• "linear"</li> <li>• "lower"</li> <li>• "mean"</li> <li>• "median"</li> <li>• "mode"</li> <li>• "roll"</li> </ul>
<code>.number_of_trees</code>	This is used for the <code>recipes::step_impute_bag()</code> <code>trees</code> parameter. This should be an integer.
<code>.neighbors</code>	This should be filled in with an integer value if <code>.type_of_imputation</code> selected is "knn".

<code>.mean_trim</code>	This should be filled in with a fraction if <code>.type_of_imputation</code> selected is "mean".
<code>.roll_statistic</code>	This should be filled in with a single unquoted function that takes with it a single argument such as mean. This should be filled in if <code>.type_of_imputation</code> selected is "roll".
<code>.roll_window</code>	This should be filled in with an integer value if <code>.type_of_imputation</code> selected is "roll".

### Details

This function will get your data ready for processing with many types of ml/ai models.

This is intended to be used inside of the data processor and therefore is an internal function. This documentation exists to explain the process and help the user understand the parameters that can be set in the pre-processor function.

### Value

A list object

### Author(s)

Steven P. Sanderson II, MPH

### See Also

<https://recipes.tidymodels.org/reference/index.html#section-step-functions-imputation/>

`step_impute_bag`

`recipes::step_impute_bag()`

[https://recipes.tidymodels.org/reference/step\\_impute\\_bag.html](https://recipes.tidymodels.org/reference/step_impute_bag.html)

`step_impute_knn`

`recipes::step_impute_knn()`

[https://recipes.tidymodels.org/reference/step\\_impute\\_knn.html](https://recipes.tidymodels.org/reference/step_impute_knn.html)

`step_impute_linear`

`recipes::step_impute_linear()`

[https://recipes.tidymodels.org/reference/step\\_impute\\_linear.html](https://recipes.tidymodels.org/reference/step_impute_linear.html)

`step_impute_lower`

`recipes::step_impute_lower()`

[https://recipes.tidymodels.org/reference/step\\_impute\\_lower.html](https://recipes.tidymodels.org/reference/step_impute_lower.html)

`step_impute_mean`

`recipes::step_impute_mean()`

[https://recipes.tidymodels.org/reference/step\\_impute\\_mean.html](https://recipes.tidymodels.org/reference/step_impute_mean.html)

`step_impute_median`

```

recipes::step_impute_median()
https://recipes.tidymodels.org/reference/step\_impute\_median.html
step_impute_mode
recipes::step_impute_mode()
https://recipes.tidymodels.org/reference/step\_impute\_mode.html
step_impute_roll
recipes::step_impute_roll()
https://recipes.tidymodels.org/reference/step\_impute\_roll.html
Other Data Recipes: hai_data_poly(), hai_data_scale(), hai_data_transform(), hai_data_trig(),
pca_your_recipe()
Other Preprocessor: hai_c50_data_prepper(), hai_cubist_data_prepper(), hai_data_poly(),
hai_data_scale(), hai_data_transform(), hai_data_trig(), hai_earth_data_prepper(),
hai_glmnet_data_prepper(), hai_knn_data_prepper(), hai_ranger_data_prepper(), hai_svm_poly_data_prepper(),
hai_svm_rbf_data_prepper(), hai_xgboost_data_prepper()

```

## Examples

```

suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

date_seq <- seq.Date(from = as.Date("2013-01-01"), length.out = 100, by = "month")
val_seq <- rep(c(rnorm(9), NA), times = 10)
df_tbl <- tibble(
  date_col = date_seq,
  value    = val_seq
)

rec_obj <- recipe(value ~ ., df_tbl)

hai_data_impute(
  .recipe_object = rec_obj,
  value,
  .type_of_imputation = "roll",
  .roll_statistic = median
)$impute_rec_obj %>%
  get_juiced_data()

```

---

hai\_data\_poly

*Data Preprocessor - Polynomial Function*

---

## Description

Takes in a recipe and will scale values using a selected recipe.

**Usage**

```
hai_data_poly(.recipe_object = NULL, ..., .p_degree = 2)
```

**Arguments**

`.recipe_object` The data that you want to process

`...` One or more selector functions to choose variables to be imputed. When used with `imp_vars`, these dots indicate which variables are used to predict the missing data in each variable. See `selections()` for more details

`.p_degree` The polynomial degree, an integer.

**Details**

This function will get your data ready for processing with many types of ml/ai models.

This is intended to be used inside of the data processor and therefore is an internal function. This documentation exists to explain the process and help the user understand the parameters that can be set in the pre-processor function.

[recipes::step\\_poly\(\)](#)

**Value**

A list object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://recipes.tidymodels.org/reference/step\\_poly.html](https://recipes.tidymodels.org/reference/step_poly.html)

Other Data Recipes: [hai\\_data\\_impute\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [pca\\_your\\_recipe\(\)](#)

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

date_seq <- seq.Date(from = as.Date("2013-01-01"), length.out = 100, by = "month")
val_seq <- rep(rnorm(10, mean = 6, sd = 2), times = 10)
df_tbl <- tibble(
  date_col = date_seq,
  value    = val_seq
)
```

```

rec_obj <- recipe(value ~ ., df_tbl)

hai_data_poly(
  .recipe_object = rec_obj,
  value
)$scale_rec_obj %>%
  get_juiced_data()

```

---

 hai\_data\_scale

*Data Preprocessor - Scale/Normalize*


---

### Description

Takes in a recipe and will scale values using a selected recipe. To call the recipe use a quoted argument like "scale" or "normalize".

### Usage

```

hai_data_scale(
  .recipe_object = NULL,
  ...,
  .type_of_scale = "center",
  .range_min = 0,
  .range_max = 1,
  .scale_factor = 1
)

```

### Arguments

- `.recipe_object` The data that you want to process
- `...` One or more selector functions to choose variables to be imputed. When used with `imp_vars`, these dots indicate which variables are used to predict the missing data in each variable. See `selections()` for more details
- `.type_of_scale` This is a quoted argument and can be one of the following:
  - "center"
  - "normalize"
  - "range"
  - "scale"
- `.range_min` A single numeric value for the smallest value in the range. This defaults to 0.
- `.range_max` A single numeric value for the largest value in the range. This defaults to 1.
- `.scale_factor` A numeric value of either 1 or 2 that scales the numeric inputs by one or two standard deviations. By dividing by two standard deviations, the coefficients attached to continuous predictors can be interpreted the same way as with binary inputs. Defaults to 1. More in reference below.

## Details

This function will get your data ready for processing with many types of ml/ai models.

This is intended to be used inside of the data processor and therefore is an internal function. This documentation exists to explain the process and help the user understand the parameters that can be set in the pre-processor function.

## Value

A list object

## Author(s)

Steven P. Sanderson II, MPH

## References

Gelman, A. (2007) "Scaling regression inputs by dividing by two standard deviations." Unpublished. Source: <http://www.stat.columbia.edu/~gelman/research/unpublished/standardizing.pdf>.

## See Also

<https://recipes.tidymodels.org/reference/index.html#section-step-functions-normalization>  
step\_center

[recipes::step\\_center\(\)](#)

[https://recipes.tidymodels.org/reference/step\\_center.html](https://recipes.tidymodels.org/reference/step_center.html)

step\_normalize

[recipes::step\\_normalize\(\)](#)

[https://recipes.tidymodels.org/reference/step\\_normalize.html](https://recipes.tidymodels.org/reference/step_normalize.html)

step\_range

[recipes::step\\_range\(\)](#)

[https://recipes.tidymodels.org/reference/step\\_range.html](https://recipes.tidymodels.org/reference/step_range.html)

step\_scale

[recipes::step\\_scale\(\)](#)

[https://recipes.tidymodels.org/reference/step\\_scale.html](https://recipes.tidymodels.org/reference/step_scale.html)

Other Data Recipes: [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [pca\\_your\\_recipe\(\)](#)

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)



## Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

date_seq <- seq.Date(from = as.Date("2013-01-01"), length.out = 100, by = "month")
val_seq <- rep(rnorm(10, mean = 6, sd = 2), times = 10)
df_tbl <- tibble(
  date_col = date_seq,
  value    = val_seq
)

rec_obj <- recipe(value ~ ., df_tbl)

hai_data_scale(
  .recipe_object = rec_obj,
  value,
  .type_of_scale = "center"
)$scale_rec_obj %>%
get_juiced_data()
```

---

hai\_data\_transform      *Data Preprocessor - Transformation Functions*

---

## Description

Takes in a recipe and will perform the desired transformation on the selected variable(s) using a selected recipe. To call the desired transformation recipe use a quoted argument like "boxcox", "bs" etc.

## Usage

```
hai_data_transform(
  .recipe_object = NULL,
  ...,
  .type_of_scale = "log",
  .bc_limits = c(-5, 5),
  .bc_num_unique = 5,
  .bs_deg_free = NULL,
  .bs_degree = 3,
  .log_base = exp(1),
  .log_offset = 0,
  .logit_offset = 0,
  .ns_deg_free = 2,
  .rel_shift = 0,
  .rel_reverse = FALSE,
  .rel_smooth = FALSE,
  .yj_limits = c(-5, 5),
```

```
.yj_num_unique = 5
)
```

### Arguments

`.recipe_object` The data that you want to process

`...` One or more selector functions to choose variables to be imputed. When used with `imp_vars`, these dots indicate which variables are used to predict the missing data in each variable. See `selections()` for more details

`.type_of_scale` This is a quoted argument and can be one of the following:

- "boxcox"
- "bs"
- "log"
- "logit"
- "ns"
- "relu"
- "sqrt"
- "yeojohnson"

`.bc_limits` A length 2 numeric vector defining the range to compute the transformation parameter lambda.

`.bc_num_unique` An integer to specify minimum required unique values to evaluate for a transformation

`.bs_deg_free` The degrees of freedom for the spline. As the degrees of freedom for a spline increase, more flexible and complex curves can be generated. When a single degree of freedom is used, the result is a rescaled version of the original data.

`.bs_degree` Degree of polynomial spline (integer).

`.log_base` A numeric value for the base.

`.log_offset` An optional value to add to the data prior to logging (to avoid  $\log(0)$ )

`.logit_offset` A numeric value to modify values of the columns that are either one or zero. They are modified to be  $x - \text{offset}$  or  $\text{offset}$  respectively.

`.ns_deg_free` The degrees of freedom for the natural spline. As the degrees of freedom for a natural spline increase, more flexible and complex curves can be generated. When a single degree of freedom is used, the result is a rescaled version of the original data.

`.rel_shift` A numeric value dictating a translation to apply to the data.

`.rel_reverse` A logical to indicate if the left hinge should be used as opposed to the right hinge.

`.rel_smooth` A logical indicating if the softplus function, a smooth approximation to the rectified linear transformation, should be used.

`.yj_limits` A length 2 numeric vector defining the range to compute the transformation parameter lambda.

`.yj_num_unique` An integer where data that have less possible values will not be evaluated for a transformation.

## Details

This function will get your data ready for processing with many types of ml/ai models.

This is intended to be used inside of the data processor and therefore is an internal function. This documentation exists to explain the process and help the user understand the parameters that can be set in the pre-processor function.

`recipes::step_BoxCox()`

## Value

A list object

## Author(s)

Steven P. Sanderson II, MPH

## See Also

[https://recipes.tidymodels.org/reference/step\\_BoxCox.html](https://recipes.tidymodels.org/reference/step_BoxCox.html)

`recipes::step_bs()`

[https://recipes.tidymodels.org/reference/step\\_bs.html](https://recipes.tidymodels.org/reference/step_bs.html)

`recipes::step_log()`

[https://recipes.tidymodels.org/reference/step\\_log.html](https://recipes.tidymodels.org/reference/step_log.html)

`recipes::step_logit()`

[https://recipes.tidymodels.org/reference/step\\_logit.html](https://recipes.tidymodels.org/reference/step_logit.html)

`recipes::step_ns()`

[https://recipes.tidymodels.org/reference/step\\_ns.html](https://recipes.tidymodels.org/reference/step_ns.html)

`recipes::step_relu()`

[https://recipes.tidymodels.org/reference/step\\_relu.html](https://recipes.tidymodels.org/reference/step_relu.html)

`recipes::step_sqrt()`

[https://recipes.tidymodels.org/reference/step\\_sqrt.html](https://recipes.tidymodels.org/reference/step_sqrt.html)

`recipes::step_YeoJohnson()`

[https://recipes.tidymodels.org/reference/step\\_YeoJohnson.html](https://recipes.tidymodels.org/reference/step_YeoJohnson.html)

Other Data Recipes: `hai_data_impute()`, `hai_data_poly()`, `hai_data_scale()`, `hai_data_trig()`, `pca_your_recipe()`

Other Preprocessor: `hai_c50_data_prepper()`, `hai_cubist_data_prepper()`, `hai_data_impute()`, `hai_data_poly()`, `hai_data_scale()`, `hai_data_trig()`, `hai_earth_data_prepper()`, `hai_glmnet_data_prepper()`, `hai_knn_data_prepper()`, `hai_ranger_data_prepper()`, `hai_svm_poly_data_prepper()`, `hai_svm_rbf_data_prepper()`, `hai_xgboost_data_prepper()`

**Examples**

```

suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

date_seq <- seq.Date(from = as.Date("2013-01-01"), length.out = 100, by = "month")
val_seq <- rep(rnorm(10, mean = 6, sd = 2), times = 10)
df_tbl <- tibble(
  date_col = date_seq,
  value    = val_seq
)

rec_obj <- recipe(value ~ ., df_tbl)

hai_data_transform(
  .recipe_object = rec_obj,
  value,
  .type_of_scale = "log"
)$scale_rec_obj %>%
  get_juiced_data()

```

---

 hai\_data\_trig

*Data Preprocessor - Trigonometric Functions*


---

**Description**

Takes in a recipe and will scale values using a selected recipe. To call the recipe use a quoted argument like "sinh", "cosh" or "tanh".

**Usage**

```

hai_data_trig(
  .recipe_object = NULL,
  ...,
  .type_of_scale = "sinh",
  .inverse = FALSE
)

```

**Arguments**

`.recipe_object` The data that you want to process

`...` One or more selector functions to choose variables to be imputed. When used with `imp_vars`, these dots indicate which variables are used to predict the missing data in each variable. See `selections()` for more details

`.type_of_scale` This is a quoted argument and can be one of the following:

- "sinh"
- "cosh"
- "tanh"

`.inverse` A logical: should the inverse function be used? Default is FALSE

## Details

This function will get your data ready for processing with many types of ml/ai models.

This is intended to be used inside of the data processor and therefore is an internal function. This documentation exists to explain the process and help the user understand the parameters that can be set in the pre-processor function.

`recipes::step_hyperbolic()`

## Value

A list object

## Author(s)

Steven P. Sanderson II, MPH

## See Also

[https://recipes.tidymodels.org/reference/step\\_hyperbolic.html](https://recipes.tidymodels.org/reference/step_hyperbolic.html)

Other Data Recipes: `hai_data_impute()`, `hai_data_poly()`, `hai_data_scale()`, `hai_data_transform()`, `pca_your_recipe()`

Other Preprocessor: `hai_c50_data_prepper()`, `hai_cubist_data_prepper()`, `hai_data_impute()`, `hai_data_poly()`, `hai_data_scale()`, `hai_data_transform()`, `hai_earth_data_prepper()`, `hai_glmnet_data_prepper()`, `hai_knn_data_prepper()`, `hai_ranger_data_prepper()`, `hai_svm_poly_data_prepper()`, `hai_svm_rbf_data_prepper()`, `hai_xgboost_data_prepper()`

## Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

date_seq <- seq.Date(from = as.Date("2013-01-01"), length.out = 100, by = "month")
val_seq <- rep(rnorm(10, mean = 6, sd = 2), times = 10)
df_tbl <- tibble(
  date_col = date_seq,
  value    = val_seq
)

rec_obj <- recipe(value ~ ., df_tbl)

hai_data_trig(
  .recipe_object = rec_obj,
  value,
  .type_of_scale = "sinh"
)$scale_rec_obj %>%
  get_juiced_data()
```

---

hai\_default\_classification\_metric\_set  
*Metric Set*

---

**Description**

Default classification metric sets from yardstick

**Usage**

```
hai_default_classification_metric_set()
```

**Details**

Default classification metric sets from yardstick

**Value**

A yardstick metric set tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Default Metric Sets: [hai\\_default\\_regression\\_metric\\_set\(\)](#)

**Examples**

```
hai_default_classification_metric_set()
```

---

hai\_default\_regression\_metric\_set  
*Metric Set*

---

**Description**

Default regression metric sets from yardstick

**Usage**

```
hai_default_regression_metric_set()
```

**Details**

Default regression metric sets from yardstick

**Value**

A yardstick metric set tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Default Metric Sets: [hai\\_default\\_classification\\_metric\\_set\(\)](#)

**Examples**

```
hai_default_regression_metric_set()
```

---

hai\_density\_hist\_plot *Density Histogram Plot*

---

**Description**

this will produce a ggplot2 or plotly histogram plot of the density information provided from the hai\_get\_density\_data\_tbl function.

**Usage**

```
hai_density_hist_plot(  
  .data,  
  .dist_name_col = distribution,  
  .value_col = dist_data,  
  .alpha = 0.382,  
  .interactive = FALSE  
)
```

**Arguments**

.data	The data that is produced from using hai_get_density_data_tbl
.dist_name_col	The column that has the distribution name, should be distribution and that is set as the default.
.value_col	The column that contains the x values that comes from the hai_get_density_data_tbl function.
.alpha	The alpha parameter for ggplot
.interactive	This is a Boolean fo TRUE/FALSE and is defaulted to FALSE. TRUE will produce a plotly plot.

**Details**

This will produce a histogram of the density information that is produced from the function `hai_get_density_data_tbl`. It will look for an attribute from the `.data` param to ensure the function was used.

**Value**

A plot, either `ggplot2` or `plotly`

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Distribution Plots: [hai\\_density\\_plot\(\)](#), [hai\\_density\\_qq\\_plot\(\)](#)

**Examples**

```
library(dplyr)

df <- hai_scale_zero_one_vec(.x = mtcars$mpg) %>%
  hai_distribution_comparison_tbl()

dist_data_tbl <- hai_get_dist_data_tbl(df)

hai_density_hist_plot(
  .data = dist_data_tbl,
  .dist_name_col = distribution,
  .value_col = dist_data,
  .alpha = 0.5,
  .interactive = FALSE
)
```

---

`hai_density_plot`*Density Histogram Plot*

---

**Description**

this will produce a `ggplot2` or `plotly` histogram plot of the density information provided from the `hai_get_density_data_tbl` function.

**Usage**

```
hai_density_plot(
  .data,
  .dist_name_col,
  .x_col,
```



```

    .y_col,
    .size = 1,
    .alpha = 0.382,
    .interactive = FALSE
  )

```

### Arguments

<code>.data</code>	The data that is produced from using <code>hai_get_density_data_tbl</code>
<code>.dist_name_col</code>	The column that has the distribution name, should be <code>distribution</code> and that is set as the default.
<code>.x_col</code>	The x value from the tidied density object.
<code>.y_col</code>	The y value from the tidied density object.
<code>.size</code>	The size parameter for <code>ggplot</code> .
<code>.alpha</code>	The alpha parameter for <code>ggplot</code> .
<code>.interactive</code>	This is a Boolean for TRUE/FALSE and is defaulted to FALSE. TRUE will produce a <code>plotly</code> plot.

### Details

This will produce a density plot of the density information that is produced from the function `hai_get_density_data_tbl`. It will look for an attribute from the `.data` param to ensure the function was used.

### Value

A plot, either `ggplot2` or `plotly`

### Author(s)

Steven P. Sanderson II, MPH

### See Also

Other Distribution Plots: [hai\\_density\\_hist\\_plot\(\)](#), [hai\\_density\\_qq\\_plot\(\)](#)

### Examples

```

library(dplyr)

df <- hai_scale_zero_one_vec(.x = mtcars$mpg) %>%
  hai_distribution_comparison_tbl()

tidy_density_tbl <- hai_get_density_data_tbl(df)

hai_density_plot(
  .data = tidy_density_tbl,
  .dist_name_col = distribution,
  .x_col = x,

```

```

.y_col = y,
.alpha = 0.5,
.interactive = FALSE
)

```

---

hai\_density\_qq\_plot    *Density QQ Plot*

---

### Description

this will produce a ggplot2 or plotly histogram plot of the density information provided from the hai\_get\_density\_data\_tbl function.

### Usage

```

hai_density_qq_plot(
  .data,
  .dist_name_col = distribution,
  .x_col = x,
  .y_col = y,
  .size = 1,
  .alpha = 0.382,
  .interactive = FALSE
)

```

### Arguments

.data	The data that is produced from using hai_get_density_data_tbl
.dist_name_col	The column that has the distribution name, should be distribution and that is set as the default.
.x_col	The column that contains the x values that comes from the hai_get_density_data_tbl function.
.y_col	The column that contains the y values that comes from the hai_get_density_data_tbl function.
.size	The size parameter for ggplot
.alpha	The alpha parameter for ggplot
.interactive	This is a Boolean fo TRUE/FALSE and is defaulted to FALSE. TRUE will produce a plotly plot.

### Details

This will produce a qq plot of the density information that is produced from the function hai\_get\_density\_data\_tbl. It will look for an attribute from the .data param to ensure the function was used.

**Value**

A plot, either ggplot2 or plotly

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Distribution Plots: [hai\\_density\\_hist\\_plot\(\)](#), [hai\\_density\\_plot\(\)](#)

**Examples**

```
library(dplyr)

df <- hai_scale_zero_one_vec(.x = mtcars$mpg) %>%
  hai_distribution_comparison_tbl()

tidy_density_tbl <- hai_get_density_data_tbl(df)

hai_density_qq_plot(
  .data = tidy_density_tbl,
  .dist_name_col = distribution,
  .x_col = x,
  .y_col = y,
  .size = 1,
  .alpha = 0.5,
  .interactive = FALSE
)
```

---

hai\_distribution\_comparison\_tbl

*Compare Data Against Distributions*

---

**Description**

This function will attempt to get some key information on the data you pass to it. It will also automatically normalize the data from 0 to 1. This will not change the distribution just it's scale in order to make sure that many different types of distributions can be fit to the data, which should help identify what the distribution of the passed data could be.

The resulting output has attributes added to it that get used in other functions that are meant to compliment each other.

This function will automatically pass the `.x` parameter to [hai\\_skewness\\_vec\(\)](#) and [hai\\_kurtosis\\_vec\(\)](#) in order to help create the random data from the distributions.

The distributions that can be chosen from are:

Distribution	R stats::dist
normal	rnorm
uniform	runif
exponential	rexp
logistic	rlogis
beta	rbeta
lognormal	rlnorm
gamma	rgamma
weibull	weibull
chisquare	rchisq
cauchy	rcauchy
hypergeometric	rhyper
f	rf
poisson	rpois

### Usage

```
hai_distribution_comparison_tbl(
  .x,
  .distributions = c("gamma", "beta"),
  .normalize = TRUE
)
```

### Arguments

`.x` The numeric vector to analyze.

`.distributions` A character vector of distributions to check. For example, `c("gamma", "beta")`

`.normalize` A boolean value of TRUE/FALSE, the default is TRUE. This will normalize the data using the `hai_scale_zero_one_vec` function.

### Details

Get information on the empirical distribution of your data along with generated densities of other distributions. This information is in the resulting tibble that is generated. Three columns will generate, `Distribution`, from the param `.distributions`, `dist_data` which is a list vector of density values passed to the underlying stats r distribution function, and `density_data`, which is the `dist_data` column passed to `list(stats::density(unlist(dist_data)))`

This has the effect of giving you the desired vector that can be used in resultant plots (`dist_data`) or you can interact with the density object itself.

If the skewness of the distribution is negative, then for the gamma and beta distributions the skew is set equal to the kurtosis and the kurtosis is set equal to  $\sqrt{(\text{skew})^2}$

### Value

A tibble.

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Distribution Functions: [hai\\_get\\_density\\_data\\_tbl\(\)](#), [hai\\_get\\_dist\\_data\\_tbl\(\)](#)

**Examples**

```
x_vec <- hai_scale_zero_one_vec(mtcars$mpg)
df <- hai_distribution_comparison_tbl(
  .x = x_vec,
  .distributions = c("beta", "gamma")
)
df
```

---

hai\_earth\_data\_prepper

*Prep Data for Earth - Recipe*

---

**Description**

Automatically prep a data.frame/tibble for use in the Earth algorithm.

**Usage**

```
hai_earth_data_prepper(.data, .recipe_formula)
```

**Arguments**

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::reciep()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the diamonds data then the formula would most likely be something like `price ~ .`

**Details**

This function will automatically prep your data.frame/tibble for use in the Earth algorithm. The Earth algorithm is for classification and regression.

This function will output a recipe specification.

**Value**

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<http://uc-r.github.io/mars>

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

Other Earth: [hai\\_auto\\_earth\(\)](#)

**Examples**

```
library(ggplot2)

# Regression
hai_earth_data_prepper(.data = diamonds, .recipe_formula = price ~ .)
reg_obj <- hai_earth_data_prepper(diamonds, price ~ .)
get_juiced_data(reg_obj)

# Classification
hai_earth_data_prepper(Titanic, Survived ~ .)
cla_obj <- hai_earth_data_prepper(Titanic, Survived ~ .)
get_juiced_data(cla_obj)
```

---

hai\_fourier\_augment    *Augment Function Fourier*

---

**Description**

Takes a numeric vector(s) or date and will return a tibble of one of the following:

- "sin"
- "cos"
- "sincos"
- c("sin", "cos", "sincos")

**Usage**

```
hai_fourier_augment(
  .data,
  .value,
  .period,
  .order,
  .names = "auto",
  .scale_type = c("sin", "cos", "sincos")
)
```

## Arguments

.data	The data being passed that will be augmented by the function.
.value	This is passed <code>rlang::enquo()</code> to capture the vectors you want to augment.
.period	The number of observations that complete a cycle
.order	The fourier term order
.names	The default is "auto"
.scale_type	A character of one of the following: "sin", "cos", or "sincos" All can be passed by setting the param equal to <code>c("sin", "cos", "sincos")</code>

## Details

Takes a numeric vector or date and will return a vector of one of the following:

- "sin"
- "cos"
- "sincos"
- `c("sin", "cos", "sincos")`

This function is intended to be used on its own in order to add columns to a tibble.

## Value

A augmented tibble

## Author(s)

Steven P. Sanderson II, MPH

## See Also

Other Augment Function: [hai\\_fourier\\_discrete\\_augment\(\)](#), [hai\\_hyperbolic\\_augment\(\)](#), [hai\\_polynomial\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_winsorized\\_move\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

## Examples

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)
```

```
hai_fourier_augment(data_tbl, b, .period = 12, .order = 1, .scale_type = "sin")
hai_fourier_augment(data_tbl, b, .period = 12, .order = 1, .scale_type = "cos")
```

---

```
hai_fourier_discrete_augment
```

*Augment Function Fourier Discrete*

---

## Description

Takes a numeric vector(s) or date and will return a tibble of one of the following:

- "sin"
- "cos"
- "sincos"
- c("sin", "cos", "sincos") When either of these values falls below zero, then zero else one

## Usage

```
hai_fourier_discrete_augment(
  .data,
  .value,
  .period,
  .order,
  .names = "auto",
  .scale_type = c("sin", "cos", "sincos")
)
```

## Arguments

.data	The data being passed that will be augmented by the function.
.value	This is passed <code>rlang::enquo()</code> to capture the vectors you want to augment.
.period	The number of observations that complete a cycle
.order	The fourier term order
.names	The default is "auto"
.scale_type	A character of one of the following: "sin", "cos", or "sincos" All can be passed by setting the param equal to c("sin", "cos", "sincos")

## Details

Takes a numeric vector or a date and will return a vector of one of the following:

- "sin"
- "cos"
- "sincos"
- c("sin", "cos", "sincos")

This function is intended to be used on its own in order to add columns to a tibble.



**Value**

A augmented tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Augment Function: [hai\\_fourier\\_augment\(\)](#), [hai\\_hyperbolic\\_augment\(\)](#), [hai\\_polynomial\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_winsorized\\_move\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 24
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

hai_fourier_discrete_augment(data_tbl, b, .period = 2 * 12, .order = 1, .scale_type = "sin")
hai_fourier_discrete_augment(data_tbl, b, .period = 2 * 12, .order = 1, .scale_type = "cos")
```

---

hai\_fourier\_discrete\_vec

*Vector Function Discrete Fourier*

---

**Description**

Takes a numeric vector or date and will return a vector of one of the following:

- "sin"
- "cos"
- "sincos" This will do value =  $\sin(x) * \cos(x)$  When either of these values falls below zero, then zero else one

**Usage**

```
hai_fourier_discrete_vec(
  .x,
  .period,
  .order,
  .scale_type = c("sin", "cos", "sincos")
)
```

**Arguments**

<code>.x</code>	A numeric vector
<code>.period</code>	The number of observations that complete a cycle
<code>.order</code>	The fourier term order
<code>.scale_type</code>	A character of one of the following: "sin","cos","sincos"

**Details**

Takes a numeric vector or date and will return a vector of one of the following:

- "sin"
- "cos"
- "sincos"

The internal calculation is straightforward:

- $\sin = \sin(2 * \pi * h * x)$ , where  $h = .order/.period$
- $\cos = \cos(2 * \pi * h * x)$ , where  $h = .order/.period$
- $\text{sincos} = \sin(2 * \pi * h * x) * \cos(2 * \pi * h * x)$  where  $h = .order/.period$

This function can be used on its own. It is also the basis for the function [hai\\_fourier\\_discrete\\_augment\(\)](#).

**Value**

A numeric vector of 1's and 0's

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Vector Function: [hai\\_fourier\\_vec\(\)](#), [hai\\_hyperbolic\\_vec\(\)](#), [hai\\_kurtosis\\_vec\(\)](#), [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [hai\\_skewness\\_vec\(\)](#), [hai\\_winsorized\\_move\\_vec\(\)](#), [hai\\_winsorized\\_truncate\\_vec\(\)](#)

**Examples**

```

suppressPackageStartupMessages(library(dplyr))

len_out <- 24
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

vec_1 <- hai_fourier_discrete_vec(data_tbl$a, .period = 12, .order = 1, .scale_type = "sin")
vec_2 <- hai_fourier_discrete_vec(data_tbl$a, .period = 12, .order = 1, .scale_type = "cos")
vec_3 <- hai_fourier_discrete_vec(data_tbl$a, .period = 12, .order = 1, .scale_type = "sincos")

plot(data_tbl$b)
lines(vec_1, col = "blue")
lines(vec_2, col = "red")
lines(vec_3, col = "green")

```

---

hai_fourier_vec	<i>Vector Function Fourier</i>
-----------------	--------------------------------

---

**Description**

Takes a numeric vector and will return a vector of one of the following:

- "sin"
- "cos"
- "sincos" This will do value =  $\sin(x) * \cos(x)$

**Usage**

```
hai_fourier_vec(.x, .period, .order, .scale_type = c("sin", "cos", "sincos"))
```

**Arguments**

.x	A numeric vector
.period	The number of observations that complete a cycle
.order	The fourier term order
.scale_type	A character of one of the following: "sin","cos","sincos"

**Details**

Takes a numeric vector and will return a vector of one of the following:

- "sin"
- "cos"
- "sincos"

The internal calculation is straightforward:

- $\sin = \sin(2 * \pi * h * x)$ , where  $h = .order / .period$
- $\cos = \cos(2 * \pi * h * x)$ , where  $h = .order / .period$
- $\text{sincos} = \sin(2 * \pi * h * x) * \cos(2 * \pi * h * x)$  where  $h = .order / .period$

This function can be used on it's own. It is also the basis for the function [hai\\_fourier\\_augment\(\)](#).

**Value**

A numeric vector

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Vector Function: [hai\\_fourier\\_discrete\\_vec\(\)](#), [hai\\_hyperbolic\\_vec\(\)](#), [hai\\_kurtosis\\_vec\(\)](#), [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [hai\\_skewness\\_vec\(\)](#), [hai\\_winsorized\\_move\\_vec\(\)](#), [hai\\_winsorized\\_truncate\\_vec\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 25
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

vec_1 <- hai_fourier_vec(data_tbl$b, .period = 12, .order = 1, .scale_type = "sin")
vec_2 <- hai_fourier_vec(data_tbl$b, .period = 12, .order = 1, .scale_type = "cos")
vec_3 <- hai_fourier_vec(data_tbl$date_col, .period = 12, .order = 1, .scale_type = "sincos")

plot(data_tbl$b)
lines(vec_1, col = "blue")
lines(vec_2, col = "red")
lines(vec_3, col = "green")
```

---

`hai_get_density_data_tbl`*Get Density Data Helper*

---

**Description**

This function will return a tibble that can either be nested/unnested, and grouped or un-grouped. The `.data` argument must be the output of the `hai_distribution_comparison_tbl()` function.

**Usage**

```
hai_get_density_data_tbl(.data, .unnest = TRUE, .group_data = TRUE)
```

**Arguments**

<code>.data</code>	The data from the <code>hai_distribution_comparison_tbl()</code> function as this function looks for an attribute of <code>hai_dist_compare_tbl</code>
<code>.unnest</code>	Should the resulting tibble be un-nested, a Boolean value TRUE/FALSE. The default is TRUE
<code>.group_data</code>	Should the resulting tibble be grouped, a Boolean value TRUE/FALSE. The default is FALSE

**Details**

This function expects to take the output of the `hai_distribution_comparison_tbl()` function. It returns a tibble of the tidy density data.

**Value**

A tibble.

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Distribution Functions: [hai\\_distribution\\_comparison\\_tbl\(\)](#), [hai\\_get\\_dist\\_data\\_tbl\(\)](#)

**Examples**

```
library(dplyr)

df <- hai_scale_zero_one_vec(.x = mtcars$mpg) %>%
  hai_distribution_comparison_tbl()
hai_get_density_data_tbl(df)
```

---

hai\_get\_dist\_data\_tbl *Get Distribution Data Helper*

---

### Description

This function will return a tibble that can either be nested/unnested, and grouped or ungrouped. The `.data` argument must be the output of the `hai_distribution_comparison_tbl()` function.

### Usage

```
hai_get_dist_data_tbl(.data, .unnest = TRUE, .group_data = FALSE)
```

### Arguments

<code>.data</code>	The data from the <code>hai_distribution_comparison_tbl()</code> function as this function looks for a class of 'hai_dist_data'
<code>.unnest</code>	Should the resulting tibble be unnested, a boolean value TRUE/FALSE. The default is TRUE
<code>.group_data</code>	Should the resulting tibble be grouped, a boolean value TRUE/FALSE. The default is FALSE

### Details

This function expects to take the output of the `hai_distribution_comparison_tbl()` function. It returns a tibble of the distribution and the randomly generated data produced from the associated stats r function like `rnorm`

### Value

A tibble.

### Author(s)

Steven P. Sanderson II, MPH

### See Also

Other Distribution Functions: [hai\\_distribution\\_comparison\\_tbl\(\)](#), [hai\\_get\\_density\\_data\\_tbl\(\)](#)

### Examples

```
library(dplyr)

df <- hai_scale_zero_one_vec(.x = mtcars$mpg) %>%
  hai_distribution_comparison_tbl()
hai_get_dist_data_tbl(df)
```

---

`hai_glmnet_data_prepper`*Prep Data for glmnet - Recipe*

---

## Description

Automatically prep a `data.frame/tibble` for use in the `glmnet` algorithm.

## Usage

```
hai_glmnet_data_prepper(.data, .recipe_formula)
```

## Arguments

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the `data` parameter of the `recipes::recip()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the `iris` data then the formula would most likely be something like `Species ~ .`

## Details

This function will automatically prep your `data.frame/tibble` for use in the `glmnet` algorithm. It expects data to be presented in a certain fashion.

This function will output a recipe specification.

## Value

A recipe object

## Author(s)

Steven P. Sanderson II, MPH

## See Also

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

Other knn: [hai\\_knn\\_data\\_prepper\(\)](#)

**Examples**

```
library(ggplot2)

hai_glmnet_data_prepper(.data = Titanic, .recipe_formula = Survived ~ .)
rec_obj <- hai_glmnet_data_prepper(Titanic, Survived ~ .)
get_juiced_data(rec_obj)
```

---

```
hai_histogram_facet_plot
Histogram Facet Plot
```

---

**Description**

This function expects a data.frame/tibble and will return a faceted histogram.

**Usage**

```
hai_histogram_facet_plot(
  .data,
  .bins = 10,
  .scale_data = FALSE,
  .ncol = 5,
  .fct_reorder = FALSE,
  .fct_rev = FALSE,
  .fill = "steelblue",
  .color = "white",
  .scale = "free",
  .interactive = FALSE
)
```

**Arguments**

.data	The data you want to pass to the function.
.bins	The number of bins for the histograms.
.scale_data	This is a boolean set to FALSE. TRUE will use hai_scale_zero_one_vec() to [0, 1] scale the data.
.ncol	The number of columns for the facet_warp argument.
.fct_reorder	Should the factor column be reordered? TRUE/FALSE, default of FALSE
.fct_rev	Should the factor column be reversed? TRUE/FALSE, default of FALSE
.fill	Default is steelblue
.color	Default is 'white'
.scale	Default is 'free'
.interactive	Default is FALSE, TRUE will produce a plotly plot.



**Details**

Takes in a data.frame/tibble and returns a faceted histogram.

**Value**

A ggplot or plotly plot

**Author(s)**

Steven P. Sanderson II, MPH

**Examples**

```
hai_histogram_facet_plot(.data = iris)
hai_histogram_facet_plot(.data = iris, .scale_data = TRUE)
```

---

hai\_hyperbolic\_augment

*Augment Function Hyperbolic*

---

**Description**

Takes a numeric vector(s) or date and will return a tibble of one of the following:

- "sin"
- "cos"
- "tan"
- "sincos"
- c("sin", "cos", "tan", "sincos")

**Usage**

```
hai_hyperbolic_augment(  
  .data,  
  .value,  
  .names = "auto",  
  .scale_type = c("sin", "cos", "tan", "sincos")  
)
```

**Arguments**

.data	The data being passed that will be augmented by the function.
.value	This is passed <code>rlang::enquo()</code> to capture the vectors you want to augment.
.names	The default is "auto"
.scale_type	A character of one of the following: "sin", "cos", "tan", "sincos" All can be passed by setting the param equal to c("sin", "cos", "tan", "sincos")

**Details**

Takes a numeric vector or date and will return a vector of one of the following:

- "sin"
- "cos"
- "tan"
- "sincos"
- c("sin", "cos", "tan", "sincos")

This function is intended to be used on its own in order to add columns to a tibble.

**Value**

A augmented tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Augment Function: [hai\\_fourier\\_augment\(\)](#), [hai\\_fourier\\_discrete\\_augment\(\)](#), [hai\\_polynomial\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_winsorized\\_move\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

hai_hyperbolic_augment(data_tbl, b, .scale_type = "sin")
hai_hyperbolic_augment(data_tbl, b, .scale_type = "tan")
```

---

hai\_hyperbolic\_vec      *Vector Function Hyperbolic*

---

### Description

Takes a numeric vector and will return a vector of one of the following:

- "sin"
- "cos"
- "tan"
- "sincos" This will do value =  $\sin(x) * \cos(x)$

### Usage

```
hai_hyperbolic_vec(.x, .scale_type = c("sin", "cos", "tan", "sincos"))
```

### Arguments

.x                    A numeric vector  
.scale\_type        A character of one of the following: "sin","cos","tan","sincos"

### Details

Takes a numeric vector and will return a vector of one of the following:

- "sin"
- "cos"
- "tan"
- "sincos"

This function can be used on it's own. It is also the basis for the function [hai\\_hyperbolic\\_augment\(\)](#).

### Value

A numeric vector

### Author(s)

Steven P. Sanderson II, MPH

### See Also

Other Vector Function: [hai\\_fourier\\_discrete\\_vec\(\)](#), [hai\\_fourier\\_vec\(\)](#), [hai\\_kurtosis\\_vec\(\)](#), [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [hai\\_skewness\\_vec\(\)](#), [hai\\_winsorized\\_move\\_vec\(\)](#), [hai\\_winsorized\\_truncate\\_vec\(\)](#)

## Examples

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 25
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

vec_1 <- hai_hyperbolic_vec(data_tbl$b, .scale_type = "sin")
vec_2 <- hai_hyperbolic_vec(data_tbl$b, .scale_type = "cos")
vec_3 <- hai_hyperbolic_vec(data_tbl$b, .scale_type = "sincos")

plot(data_tbl$b)
lines(vec_1, col = "blue")
lines(vec_2, col = "red")
lines(vec_3, col = "green")
```

---

hai\_kmeans\_automl      *Automatic K-Means H2O*

---

## Description

This is a wrapper around the `h2o:h2o.kmeans()` function that will return a list object with a lot of useful and easy to use tidy style information.

## Usage

```
hai_kmeans_automl(
  .data,
  .split_ratio = 0.8,
  .seed = 1234,
  .centers = 10,
  .standardize = TRUE,
  .print_model_summary = TRUE,
  .predictors,
  .categorical_encoding = "auto",
  .initialization_mode = "Furthest",
  .max_iterations = 100
)
```

**Arguments**

- `.data` The data that is to be passed for clustering.
- `.split_ratio` The ratio for training and testing splits.
- `.seed` The default is 1234, but can be set to any integer.
- `.centers` The default is 1. Specify the number of clusters (groups of data) in a data set.
- `.standardize` The default is set to TRUE. When TRUE all numeric columns will be set to zero mean and unit variance.
- `.print_model_summary`  
This is a boolean and controls if the model summary is printed to the console. The default is TRUE.
- `.predictors` This must be in the form of `c("column_1", "column_2", ... "column_n")`
- `.categorical_encoding`  
Can be one of the following:
- "auto"
  - "enum"
  - "one\_hot\_explicit"
  - "binary"
  - "eigen"
  - "label\_encoder"
  - "sort\_by\_response"
  - "enum\_limited"
- `.initialization_mode`  
This can be one of the following:
- "Random"
  - "Furthest (default)"
  - "PlusPlus"
- `.max_iterations`  
The default is 100. This specifies the number of training iterations

**Value**

A list object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Kmeans: [hai\\_kmeans\\_automl\\_predict\(\)](#), [hai\\_kmeans\\_mapped\\_tbl\(\)](#), [hai\\_kmeans\\_obj\(\)](#), [hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_scree\\_plt\(\)](#), [hai\\_kmeans\\_tidy\\_tbl\(\)](#), [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)

## Examples

```
## Not run:
h2o.init()
output <- hai_kmeans_automl(
  .data = iris,
  .predictors = c("Sepal.Width", "Sepal.Length", "Petal.Width", "Petal.Length"),
  .standardize = FALSE
)
h2o.shutdown()

## End(Not run)
```

---

hai\_kmeans\_automl\_predict  
*Automatic K-Means H2O*

---

## Description

This is a wrapper around the `h2o::h2o.predict()` function that will return a list object with a lot of useful and easy to use tidy style information.

## Usage

```
hai_kmeans_automl_predict(.input)
```

## Arguments

`.input` This is the output of the `hai_kmeans_automl()` function.

## Details

This function will internally take in the output assigned from the `hai_kmeans_automl()` function only and return a list of useful information. The items that are returned are as follows:

1. `prediction` - The h2o dataframe of predictions
2. `prediction_tbl` - The h2o predictions in tibble format
3. `valid_tbl` - The validation data in tibble format
4. `pred_full_tbl` - The entire validation set with the predictions attached using `base::cbind()`. The predictions are in a column called `predicted_cluster` and are in the format of a factor using `forcats::as_factor()`

## Value

A list object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Kmeans: [hai\\_kmeans\\_automl\(\)](#), [hai\\_kmeans\\_mapped\\_tbl\(\)](#), [hai\\_kmeans\\_obj\(\)](#), [hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_scree\\_plt\(\)](#), [hai\\_kmeans\\_tidy\\_tbl\(\)](#), [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)

**Examples**

```
## Not run:
h2o.init()

output <- hai_kmeans_automl(
  .data = iris,
  .predictors = c("Sepal.Width", "Sepal.Length", "Petal.Width", "Petal.Length"),
  .standardize = FALSE
)

pred <- hai_kmeans_automl_predict(output)

h2o.shutdown()

## End(Not run)
```

---

hai\_kmeans\_mapped\_tbl *K-Means Mapping Function*

---

**Description**

Create a tibble that maps the [hai\\_kmeans\\_obj\(\)](#) using `purrr::map()` to create a nested data.frame/tibble that holds n centers. This tibble will be used to help create a scree plot.

**Usage**

```
hai_kmeans_mapped_tbl(.data, .centers = 15)
```

```
kmeans_mapped_tbl(.data, .centers = 15)
```

**Arguments**

<code>.data</code>	You must have a tibble in the working environment from the <a href="#">hai_kmeans_user_item_tbl()</a>
<code>.centers</code>	How many different centers do you want to try

**Details**

Takes in a single parameter of `.centers`. This is used to create the tibble and map the [hai\\_kmeans\\_obj\(\)](#) function down the list creating a nested tibble.

**Value**

A nested tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://en.wikipedia.org/wiki/Scree\\_plot](https://en.wikipedia.org/wiki/Scree_plot)

Other Kmeans: [hai\\_kmeans\\_automl\(\)](#), [hai\\_kmeans\\_automl\\_predict\(\)](#), [hai\\_kmeans\\_obj\(\)](#),  
[hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_scree\\_plt\(\)](#), [hai\\_kmeans\\_tidy\\_tbl\(\)](#), [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)

**Examples**

```
library(healthyR.data)
library(dplyr)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

ui_tbl <- hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
)

hai_kmeans_mapped_tbl(ui_tbl)
```

---

hai\_kmeans\_obj

*K-Means Object*

---

**Description**

Takes the output of the [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#) function and applies the k-means algorithm to it using `stats::kmeans()`

**Usage**

```
hai_kmeans_obj(.data, .centers = 5)
```

```
kmeans_obj(.data, .centers = 5)
```



**Arguments**

.data            The data that gets passed from [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)  
.centers        How many initial centers to start with

**Details**

Uses the [stats::kmeans\(\)](#) function and creates a wrapper around it.

**Value**

A stats k-means object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Kmeans: [hai\\_kmeans\\_automl\(\)](#), [hai\\_kmeans\\_automl\\_predict\(\)](#), [hai\\_kmeans\\_mapped\\_tbl\(\)](#),  
[hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_scree\\_plt\(\)](#), [hai\\_kmeans\\_tidy\\_tbl\(\)](#), [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)

**Examples**

```
library(healthyR.data)
library(dplyr)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
) %>%
  hai_kmeans_obj()
```

---

`hai_kmeans_scee_data_tbl`*K-Means Scree Plot Data Table*

---

**Description**

Take data from the `hai_kmeans_mapped_tbl()` and unnest it into a tibble for inspection and for use in the `hai_kmeans_scee_plt()` function.

**Usage**

```
hai_kmeans_scee_data_tbl(.data)
```

```
kmeans_scee_data_tbl(.data)
```

**Arguments**

`.data` You must have a tibble in the working environment from the `hai_kmeans_mapped_tbl()`

**Details**

Takes in a single parameter of `.data` from `hai_kmeans_mapped_tbl()` and transforms it into a tibble that is used for `hai_kmeans_scee_plt()`. It will show the values (tot.withinss) at each center.

**Value**

A nested tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Kmeans: `hai_kmeans_automl()`, `hai_kmeans_automl_predict()`, `hai_kmeans_mapped_tbl()`, `hai_kmeans_obj()`, `hai_kmeans_scee_plt()`, `hai_kmeans_tidy_tbl()`, `hai_kmeans_user_item_tbl()`

**Examples**

```
library(healthyR.data)
library(dplyr)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()
```

```
ui_tbl <- hai_kmeans_user_item_tbl(  
  .data = data_tbl,  
  .row_input = service_line,  
  .col_input = payer_grouping,  
  .record_input = record  
)  
  
kmm_tbl <- hai_kmeans_mapped_tbl(ui_tbl)  
  
hai_kmeans_scree_data_tbl(kmm_tbl)
```

---

hai\_kmeans\_scree\_plt *K-Means Scree Plot*

---

## Description

Create a scree-plot from the [hai\\_kmeans\\_mapped\\_tbl\(\)](#) function.

## Usage

```
hai_kmeans_scree_plt(.data)  
  
kmeans_scree_plt(.data)  
  
hai_kmeans_scree_plot(.data)
```

## Arguments

.data            The data from the [hai\\_kmeans\\_mapped\\_tbl\(\)](#) function

## Details

Outputs a scree-plot

## Value

A ggplot2 plot

## Author(s)

Steven P. Sanderson II, MPH

## See Also

[https://en.wikipedia.org/wiki/Scree\\_plot](https://en.wikipedia.org/wiki/Scree_plot)

Other Kmeans: [hai\\_kmeans\\_automl\(\)](#), [hai\\_kmeans\\_automl\\_predict\(\)](#), [hai\\_kmeans\\_mapped\\_tbl\(\)](#), [hai\\_kmeans\\_obj\(\)](#), [hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_tidy\\_tbl\(\)](#), [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)

**Examples**

```

library(healthyR.data)
library(dplyr)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

ui_tbl <- hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
)

kmm_tbl <- hai_kmeans_mapped_tbl(ui_tbl)

hai_kmeans_scree_plt(.data = kmm_tbl)

```

---

hai\_kmeans\_tidy\_tbl    *K-Means Object Tidy Functions*

---

**Description**

K-Means tidy functions

**Usage**

```
hai_kmeans_tidy_tbl(.kmeans_obj, .data, .tidy_type = "tidy")
```

```
kmeans_tidy_tbl(.kmeans_obj, .data, .tidy_type = "tidy")
```

**Arguments**

.kmeans_obj	A <code>stats::kmeans()</code> object
.data	The user item tibble created from <code>hai_kmeans_user_item_tbl()</code>
.tidy_type	"tidy", "glance", or "augment"

**Details**

Takes in a k-means object and its associated user item tibble and then returns one of the items asked for. Either: `broom::tidy()`, `broom::glance()` or `broom::augment()`. The function defaults to `broom::tidy()`.

**Value**

A tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Kmeans: [hai\\_kmeans\\_automl\(\)](#), [hai\\_kmeans\\_automl\\_predict\(\)](#), [hai\\_kmeans\\_mapped\\_tbl\(\)](#), [hai\\_kmeans\\_obj\(\)](#), [hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_scree\\_plt\(\)](#), [hai\\_kmeans\\_user\\_item\\_tbl\(\)](#)

**Examples**

```
library(healthyR.data)
library(dplyr)
library(broom)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

uit_tbl <- hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
)

km_obj <- hai_kmeans_obj(uit_tbl)

hai_kmeans_tidy_tbl(
  .kmeans_obj = km_obj,
  .data = uit_tbl,
  .tidy_type = "augment"
)

hai_kmeans_tidy_tbl(
  .kmeans_obj = km_obj,
  .data = uit_tbl,
  .tidy_type = "glance"
)

hai_kmeans_tidy_tbl(
  .kmeans_obj = km_obj,
  .data = uit_tbl,
  .tidy_type = "tidy"
) %>%
```

```
glimpse()
```

---

```
hai_kmeans_user_item_tbl
```

*K-Means User Item Tibble*

---

## Description

Takes in a data.frame/tibble and transforms it into an aggregated/normalized user-item tibble of proportions. The user will need to input the parameters for the rows/user and the columns/items.

## Usage

```
hai_kmeans_user_item_tbl(.data, .row_input, .col_input, .record_input)
```

```
kmeans_user_item_tbl(.data, .row_input, .col_input, .record_input)
```

## Arguments

<code>.data</code>	The data that you want to transform
<code>.row_input</code>	The column that is going to be the row (user)
<code>.col_input</code>	The column that is going to be the column (item)
<code>.record_input</code>	The column that is going to be summed up for the aggregation and normalization process.

## Details

This function should be used before using a k-mean model. This is commonly referred to as a user-item matrix because "users" tend to be on the rows and "items" (e.g. orders) on the columns. You must supply a column that can be summed for the aggregation and normalization process to occur.

## Value

A aggregated/normalized user item tibble

## Author(s)

Steven P. Sanderson II, MPH

## See Also

Other Kmeans: [hai\\_kmeans\\_automl\(\)](#), [hai\\_kmeans\\_automl\\_predict\(\)](#), [hai\\_kmeans\\_mapped\\_tbl\(\)](#), [hai\\_kmeans\\_obj\(\)](#), [hai\\_kmeans\\_scree\\_data\\_tbl\(\)](#), [hai\\_kmeans\\_scree\\_plt\(\)](#), [hai\\_kmeans\\_tidy\\_tbl\(\)](#)

## Examples

```
library(healthyR.data)
library(dplyr)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
)
```

---

hai\_knn\_data\_prepper *Prep Data for k-NN - Recipe*

---

## Description

Automatically prep a data.frame/tibble for use in the k-NN algorithm.

## Usage

```
hai_knn_data_prepper(.data, .recipe_formula)
```

## Arguments

**.data** The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::recip()` function.

**.recipe\_formula** The formula that is going to be passed. For example if you are using the iris data then the formula would most likely be something like `Species ~ .`

## Details

This function will automatically prep your data.frame/tibble for use in the k-NN algorithm. The k-NN algorithm is a lazy learning classification algorithm. It expects data to be presented in a certain fashion.

This function will output a recipe specification.

## Value

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

Other knn: [hai\\_glmnet\\_data\\_prepper\(\)](#)

**Examples**

```
library(ggplot2)

hai_knn_data_prepper(.data = Titanic, .recipe_formula = Survived ~ .)
rec_obj <- hai_knn_data_prepper(iris, Species ~ .)
get_juiced_data(rec_obj)
```

---

hai\_kurtosis\_vec

*Compute Kurtosis of a Vector*

---

**Description**

This function takes in a vector as it's input and will return the kurtosis of that vector. The length of this vector must be at least four numbers. The kurtosis explains the sharpness of the peak of a distribution of data.

$$\left(\frac{1}{n} * \sum(x - \mu)^4\right) / \left(\left(\frac{1}{n} * \sum(x - \mu)^2\right)^2\right)$$
**Usage**

```
hai_kurtosis_vec(.x)
```

**Arguments**

`.x` A numeric vector of length four or more.

**Details**

A function to return the kurtosis of a vector.

**Value**

The kurtosis of a vector



**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<https://en.wikipedia.org/wiki/Kurtosis>

Other Vector Function: `hai_fourier_discrete_vec()`, `hai_fourier_vec()`, `hai_hyperbolic_vec()`, `hai_scale_zero_one_vec()`, `hai_scale_zscore_vec()`, `hai_skewness_vec()`, `hai_winsorized_move_vec()`, `hai_winsorized_truncate_vec()`

**Examples**

```
hai_kurtosis_vec(rnorm(100, 3, 2))
```

---

hai\_polynomial\_augment  
*Augment Polynomial Features*

---

**Description**

This function takes in a data table and a predictor column. A user can either create their own formula using the `.formula` parameter or, if they leave the default of `NULL` then the user must enter a `.degree` **AND** `.pred_col` column.

**Usage**

```
hai_polynomial_augment(  
  .data,  
  .formula = NULL,  
  .pred_col = NULL,  
  .degree = 1,  
  .new_col_prefix = "nt_"  
)
```

**Arguments**

<code>.data</code>	The data being passed that will be augmented by the function.
<code>.formula</code>	This should be a valid formula like <code>'y ~ .^2'</code> or <code>NULL</code> .
<code>.pred_col</code>	This is passed <code>rlang::enquo()</code> to capture the vector that you designate as the 'y' column.
<code>.degree</code>	This should be an integer and is used to set the degree in the poly function. The degree must be less than the unique data points or it will error out.
<code>.new_col_prefix</code>	The default is <code>"nt_"</code> which stands for <code>"new_term"</code> . You can set this to whatever you like, as long as it is a quoted string.

## Details

A valid `data.frame/tibble` must be passed to this function. It is required that a user either enter a `.formula` or a `.degree` **AND** `.pred_col` otherwise this function will stop and error out.

Under the hood this function will create a `stats::poly()` function if the `.formula` is left as `NULL`. For example:

- `.formula = A ~ .^2`
- OR `.degree = 2, .pred_col = A`

There is also a parameter `.new_col_prefix` which will add a character string to the column names so that they are easily identified further down the line. The default is `'nt_'`

## Value

An augmented tibble

## Author(s)

Steven P. Sanderson II, MPH

## See Also

Other Augment Function: [hai\\_fourier\\_augment\(\)](#), [hai\\_fourier\\_discrete\\_augment\(\)](#), [hai\\_hyperbolic\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_winsorized\\_move\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

## Examples

```
suppressPackageStartupMessages(library(dplyr))
data_tbl <- data.frame(
  A = c(0, 2, 4),
  B = c(1, 3, 5),
  C = c(2, 4, 6)
)

hai_polynomial_augment(.data = data_tbl, .pred_col = A, .degree = 2, .new_col_prefix = "n")
hai_polynomial_augment(.data = data_tbl, .formula = A ~ .^2, .degree = 1)
```

---

hai\_ranger\_data\_prepper

*Prep Data for Ranger - Recipe*

---

## Description

Automatically prep a `data.frame/tibble` for use in the Ranger algorithm.

**Usage**

```
hai_ranger_data_prepper(.data, .recipe_formula)
```

**Arguments**

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::recipe()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the diamonds data then the formula would most likely be something like `price ~ .`

**Details**

This function will automatically prep your `data.frame/tibble` for use in the Ranger algorithm.

This function will output a recipe specification.

**Value**

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/rand\\_forest.html](https://parsnip.tidymodels.org/reference/rand_forest.html)

Other Preprocessor: `hai_c50_data_prepper()`, `hai_cubist_data_prepper()`, `hai_data_impute()`, `hai_data_poly()`, `hai_data_scale()`, `hai_data_transform()`, `hai_data_trig()`, `hai_earth_data_prepper()`, `hai_glmnet_data_prepper()`, `hai_knn_data_prepper()`, `hai_svm_poly_data_prepper()`, `hai_svm_rbf_data_prepper()`, `hai_xgboost_data_prepper()`

Other Ranger: `hai_auto_ranger()`

**Examples**

```
library(ggplot2)

# Regression
hai_ranger_data_prepper(.data = diamonds, .recipe_formula = price ~ .)
reg_obj <- hai_ranger_data_prepper(diamonds, price ~ .)
get_juiced_data(reg_obj)

# Classification
hai_ranger_data_prepper(Titanic, Survived ~ .)
cla_obj <- hai_ranger_data_prepper(Titanic, Survived ~ .)
get_juiced_data(cla_obj)
```

hai\_range\_statistic *Get the range statistic*

---

**Description**

Takes in a numeric vector and returns back the range of that vector

**Usage**

```
hai_range_statistic(.x)
```

**Arguments**

.x                    A numeric vector

**Details**

Takes in a numeric vector and returns the range of that vector using the diff and range functions.

**Value**

A single number, the range statistic

**Author(s)**

Steven P. Sandeson II, MPH

**Examples**

```
hai_range_statistic(seq(1:10))
```

---

hai\_scale\_color\_colorblind  
*Provide Colorblind Compliant Colors*

---

**Description**

8 Hex RGB color definitions suitable for charts for colorblind people.

**Usage**

```
hai_scale_color_colorblind(..., theme = "hai")
```

**Arguments**

... Data passed in from a ggplot object  
theme Right now this is hai only. Anything else will render an error.

**Details**

This function is used in others in order to help render plots for those that are color blind.

**Value**

A ggplot layer

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Color\_Blind: [color\\_blind\(\)](#), [hai\\_scale\\_fill\\_colorblind\(\)](#)

---

hai\_scale\_fill\_colorblind

*Provide Colorblind Compliant Colors*

---

**Description**

8 Hex RGB color definitions suitable for charts for colorblind people.

**Usage**

```
hai_scale_fill_colorblind(..., theme = "hai")
```

**Arguments**

... Data passed in from a ggplot object  
theme Right now this is hai only. Anything else will render an error.

**Details**

This function is used in others in order to help render plots for those that are color blind.

**Value**

A ggplot layer

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Color\_Blind: [color\\_blind\(\)](#), [hai\\_scale\\_color\\_colorblind\(\)](#)

---

hai\_scale\_zero\_one\_augment

*Augment Function Scale Zero One*

---

**Description**

Takes a numeric vector and will return a vector that has been scaled from  $[0, 1]$

**Usage**

```
hai_scale_zero_one_augment(.data, .value, .names = "auto")
```

**Arguments**

<code>.data</code>	The data being passed that will be augmented by the function.
<code>.value</code>	This is passed <a href="#">rlang::enquo()</a> to capture the vectors you want to augment.
<code>.names</code>	This is set to 'auto' by default but can be a user supplied character string.

**Details**

Takes a numeric vector and will return a vector that has been scaled from  $[0, 1]$  The input vector must be numeric. The computation is fairly straightforward. This may be helpful when trying to compare the distributions of data where a distribution like beta from the `fitdistrplus` package which requires data to be between 0 and 1

$$y[h] = (x - \min(x)) / (\max(x) - \min(x))$$

This function is intended to be used on its own in order to add columns to a tibble.

**Value**

An augmented tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Augment Function: [hai\\_fourier\\_augment\(\)](#), [hai\\_fourier\\_discrete\\_augment\(\)](#), [hai\\_hyperbolic\\_augment\(\)](#), [hai\\_polynomial\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_winsorized\\_move\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

Other Scale: [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#)

**Examples**

```
df <- data.frame(x = rnorm(100, 2, 1))
hai_scale_zero_one_augment(df, x)
```

---

hai\_scale\_zero\_one\_vec

*Vector Function Scale to Zero and One*

---

**Description**

Takes a numeric vector and will return a vector that has been scaled from  $[0, 1]$

**Usage**

```
hai_scale_zero_one_vec(.x)
```

**Arguments**

.x                    A numeric vector to be scaled from  $[0, 1]$  inclusive.

**Details**

Takes a numeric vector and will return a vector that has been scaled from  $[0, 1]$  The input vector must be numeric. The computation is fairly straightforward. This may be helpful when trying to compare the distributions of data where a distribution like beta from the `fitdistrplus` package which requires data to be between 0 and 1

$$y[h] = (x - \min(x)) / (\max(x) - \min(x))$$

This function can be used on it's own. It is also the basis for the function `hai_scale_zero_one_augment()`.

**Value**

A numeric vector

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Vector Function: [hai\\_fourier\\_discrete\\_vec\(\)](#), [hai\\_fourier\\_vec\(\)](#), [hai\\_hyperbolic\\_vec\(\)](#), [hai\\_kurtosis\\_vec\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [hai\\_skewness\\_vec\(\)](#), [hai\\_winsorized\\_move\\_vec\(\)](#), [hai\\_winsorized\\_truncate\\_vec\(\)](#)

Other Scale: [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#)

**Examples**

```
vec_1 <- rnorm(100, 2, 1)
vec_2 <- hai_scale_zero_one_vec(vec_1)

dens_1 <- density(vec_1)
dens_2 <- density(vec_2)
max_x <- max(dens_1$x, dens_2$x)
max_y <- max(dens_1$y, dens_2$y)
plot(dens_1,
     asp = max_y / max_x, main = "Density vec_1 (Red) and vec_2 (Blue)",
     col = "red", xlab = "", ylab = "Density of Vec 1 and Vec 2"
)
lines(dens_2, col = "blue")
```

---

```
hai_scale_zscore_augment
```

*Augment Function Scale Zero One*

---

**Description**

Takes a numeric vector and will return a vector that has been scaled by mean and standard deviation

**Usage**

```
hai_scale_zscore_augment(.data, .value, .names = "auto")
```

**Arguments**

.data	The data being passed that will be augmented by the function.
.value	This is passed <code>rlang::enquo()</code> to capture the vectors you want to augment.
.names	This is set to 'auto' by default but can be a user supplied character string.

**Details**

Takes a numeric vector and will return a vector that has been scaled by mean and standard deviation.

The input vector must be numeric. The computation is fairly straightforward. This may be helpful when trying to compare the distributions of data where a distribution like beta from the `fitdistrplus` package which requires data to be between 0 and 1

$$y[h] = (x - \text{mean}(x)) / \text{sd}(x)$$

This function is intended to be used on its own in order to add columns to a tibble.

**Value**

An augmented tibble



**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Augment Function: [hai\\_fourier\\_augment\(\)](#), [hai\\_fourier\\_discrete\\_augment\(\)](#), [hai\\_hyperbolic\\_augment\(\)](#), [hai\\_polynomial\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_winsorized\\_move\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

Other Scale: [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_scale\\_zscore\\_vec\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#)

**Examples**

```
df <- data.frame(x = mtcars$mpg)
hai_scale_zscore_augment(df, x)
```

---

hai\_scale\_zscore\_vec *Vector Function Scale to Zero and One*

---

**Description**

Takes a numeric vector and will return a vector that has been scaled from by mean and standard deviation

**Usage**

```
hai_scale_zscore_vec(.x)
```

**Arguments**

`.x` A numeric vector to be scaled by mean and standard deviation inclusive.

**Details**

Takes a numeric vector and will return a vector that has been scaled from mean and standard deviation.

The input vector must be numeric. The computation is fairly straightforward. This may be helpful when trying to compare the distributions of data where a distribution like beta from the `fitdistrplus` package which requires data to be between 0 and 1

$$y[h] = (x - \text{mean}(x)) / \text{sd}(x)$$

This function can be used on it's own. It is also the basis for the function [hai\\_scale\\_zscore\\_augment\(\)](#).

**Value**

A numeric vector

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

Other Vector Function: [hai\\_fourier\\_discrete\\_vec\(\)](#), [hai\\_fourier\\_vec\(\)](#), [hai\\_hyperbolic\\_vec\(\)](#), [hai\\_kurtosis\\_vec\(\)](#), [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_skewness\\_vec\(\)](#), [hai\\_winsorized\\_move\\_vec\(\)](#), [hai\\_winsorized\\_truncate\\_vec\(\)](#)

Other Scale: [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_vec\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#)

**Examples**

```
vec_1 <- mtcars$mpg
vec_2 <- hai_scale_zscore_vec(vec_1)

ax <- pretty(min(vec_1, vec_2):max(vec_1, vec_2), n = 12)

hist(vec_1, breaks = ax, col = "blue")
hist(vec_2, breaks = ax, col = "red", add = TRUE)
```

---

hai\_skewed\_features    *Get Skewed Feature Columns*

---

**Description**

Takes in a data.frame/tibble and returns a vector of names of the columns that are skewed.

**Usage**

```
hai_skewed_features(.data, .threshold = 0.6, .drop_keys = NULL)
```

**Arguments**

<code>.data</code>	The data.frame/tibble you are passing in.
<code>.threshold</code>	A level of skewness that indicates where you feel a column should be considered skewed.
<code>.drop_keys</code>	A c() character vector of columns you do not want passed to the function.

**Details**

Takes in a data.frame/tibble and returns a vector of names of the skewed columns. There are two other parameters. The first is the `.threshold` parameter that is set to the level of skewness you want in order to consider the column too skewed. The second is `.drop_keys`, these are columns you don't want to be considered for whatever reason in the skewness calculation.

**Value**

A character vector of column names that are skewed.

**Author(s)**

Steven P. Sanderson II, MPH

**Examples**

```
hai_skewed_features(mtcars)
hai_skewed_features(mtcars, .drop_keys = c("mpg", "hp"))
hai_skewed_features(mtcars, .drop_keys = "hp")
```

---

hai_skewness_vec	<i>Compute Skewness of a Vector</i>
------------------	-------------------------------------

---

**Description**

This function takes in a vector as it's input and will return the skewness of that vector. The length of this vector must be at least four numbers. The skewness explains the 'tailedness' of the distribution of data.

$$\left(\frac{1}{n} * \sum(x - \mu)^3\right) / \left(\left(\frac{1}{n} * \sum(x - \mu)^2\right)^{3/2}\right)$$

**Usage**

```
hai_skewness_vec(.x)
```

**Arguments**

.x                    A numeric vector of length four or more.

**Details**

A function to return the skewness of a vector.

**Value**

The skewness of a vector

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<https://en.wikipedia.org/wiki/Skewness>

Other Vector Function: `hai_fourier_discrete_vec()`, `hai_fourier_vec()`, `hai_hyperbolic_vec()`, `hai_kurtosis_vec()`, `hai_scale_zero_one_vec()`, `hai_scale_zscore_vec()`, `hai_winsorized_move_vec()`, `hai_winsorized_truncate_vec()`

**Examples**

```
hai_skewness_vec(rnorm(100, 3, 2))
```

---

```
hai_svm_poly_data_prepper  
  Prep Data for SVM_Poly - Recipe
```

---

**Description**

Automatically prep a data.frame/tibble for use in the SVM\_Poly algorithm.

**Usage**

```
hai_svm_poly_data_prepper(.data, .recipe_formula)
```

**Arguments**

<code>.data</code>	The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the <code>recipes::recip()</code> function.
<code>.recipe_formula</code>	The formula that is going to be passed. For example if you are using the diamonds data then the formula would most likely be something like <code>price ~ .</code>

**Details**

This function will automatically prep your data.frame/tibble for use in the SVM\_Poly algorithm. The SVM\_Poly algorithm is for regression only.

This function will output a recipe specification.

**Value**

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/svm\\_poly.html](https://parsnip.tidymodels.org/reference/svm_poly.html)

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

Other SVM\_Poly: [hai\\_auto\\_svm\\_poly\(\)](#)

**Examples**

```
library(ggplot2)

# Regression
hai_svm_poly_data_prepper(.data = diamonds, .recipe_formula = price ~ .)
reg_obj <- hai_svm_poly_data_prepper(diamonds, price ~ .)
get_juiced_data(reg_obj)

# Classification
hai_svm_poly_data_prepper(Titanic, Survived ~ .)
cla_obj <- hai_svm_poly_data_prepper(Titanic, Survived ~ .)
get_juiced_data(cla_obj)
```

---

hai\_svm\_rbf\_data\_prepper  
*Prep Data for SVM\_RBF - Recipe*

---

**Description**

Automatically prep a data.frame/tibble for use in the SVM\_RBF algorithm.

**Usage**

```
hai_svm_rbf_data_prepper(.data, .recipe_formula)
```

**Arguments**

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::reciep()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the diamonds data then the formula would most likely be something like `price ~ .`

**Details**

This function will automatically prep your data.frame/tibble for use in the SVM\_RBF algorithm. The SVM\_RBF algorithm is for regression only.

This function will output a recipe specification.

**Value**

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/svm\\_rbf.html](https://parsnip.tidymodels.org/reference/svm_rbf.html)

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_xgboost\\_data\\_prepper\(\)](#)

Other SVM\_RBF: [hai\\_auto\\_svm\\_rbf\(\)](#)

**Examples**

```
library(ggplot2)

# Regression
hai_svm_rbf_data_prepper(.data = diamonds, .recipe_formula = price ~ .)
reg_obj <- hai_svm_rbf_data_prepper(diamonds, price ~ .)
get_juiced_data(reg_obj)

# Classification
hai_svm_rbf_data_prepper(Titanic, Survived ~ .)
cla_obj <- hai_svm_rbf_data_prepper(Titanic, Survived ~ .)
get_juiced_data(cla_obj)
```

---

hai\_umap\_list

*UMAP Projection*

---

**Description**

Create a umap object from the `uwot::umap()` function.

**Usage**

```
hai_umap_list(.data, .kmeans_map_tbl, .k_cluster = 5)
```

```
umap_list(.data, .kmeans_map_tbl, .k_cluster = 5)
```

## Arguments

- `.data`            The data from the `hai_kmeans_user_item_tbl()` function.
- `.kmeans_map_tbl`            The data from the `hai_kmeans_mapped_tbl()`.
- `.k_cluster`        Pick the desired amount of clusters from your analysis of the scree plot.

## Details

This takes in the user item table/matix that is produced by `hai_kmeans_user_item_tbl()` function. This function uses the defaults of `uwot::umap()`.

## Value

A list of tibbles and the umap object

## Author(s)

Steven P. Sanderson II, MPH

## See Also

- <https://cran.r-project.org/package=uwot> (CRAN)
- <https://github.com/jlmelville/uwot> (GitHub)
- <https://github.com/jlmelville/uwot> (arXiv paper)

Other UMAP: `hai_umap_plot()`

## Examples

```
library(healthyR.data)
library(dplyr)
library(broom)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

uit_tbl <- hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
)

kmm_tbl <- hai_kmeans_mapped_tbl(uit_tbl)
```

```
umap_list(.data = uit_tbl, kmm_tbl, 3)
```

---

hai\_umap\_plot

*UMAP and K-Means Cluster Visualization*

---

## Description

Create a UMAP Projection plot.

## Usage

```
hai_umap_plot(.data, .point_size = 2, .label = TRUE)
```

```
umap_plt(.data, .point_size = 2, .label = TRUE)
```

## Arguments

<code>.data</code>	The data from the <code>umap_list()</code> function.
<code>.point_size</code>	The desired size for the points of the plot.
<code>.label</code>	Should <code>ggrepel::geom_label_repel()</code> be used to display cluster user labels.

## Details

This takes in `umap_kmeans_cluster_results_tbl` from the `umap_list()` function output.

## Value

A ggplot2 UMAP Projection with clusters represented by colors.

## Author(s)

Steven P. Sanderson II, MPH

## See Also

- <https://cran.r-project.org/package=uwot> (CRAN)
- <https://github.com/jlmelville/uwot> (GitHub)
- <https://github.com/jlmelville/uwot> (arXiv paper)

Other UMAP: `hai_umap_list()`



**Examples**

```

library(healthyR.data)
library(dplyr)
library(broom)
library(ggplot2)

data_tbl <- healthyR_data %>%
  filter(ip_op_flag == "I") %>%
  filter(payer_grouping != "Medicare B") %>%
  filter(payer_grouping != "?") %>%
  select(service_line, payer_grouping) %>%
  mutate(record = 1) %>%
  as_tibble()

uit_tbl <- hai_kmeans_user_item_tbl(
  .data = data_tbl,
  .row_input = service_line,
  .col_input = payer_grouping,
  .record_input = record
)

kmm_tbl <- hai_kmeans_mapped_tbl(uit_tbl)

ump_lst <- hai_umap_list(.data = uit_tbl, kmm_tbl, 3)

hai_umap_plot(.data = ump_lst, .point_size = 3)

```

---

```
hai_winsorized_move_augment
```

*Augment Function Winsorize Move*

---

**Description**

Takes a numeric vector and will return a tibble with the winsorized values.

**Usage**

```
hai_winsorized_move_augment(.data, .value, .multiple, .names = "auto")
```

**Arguments**

.data	The data being passed that will be augmented by the function.
.value	This is passed <code>rlang::enquo()</code> to capture the vectors you want to augment.
.multiple	A positive number indicating how many times the the zero center mean absolute deviation should be multiplied by for the scaling parameter.
.names	The default is "auto"

**Details**

Takes a numeric vector and will return a winsorized vector of values that have been moved some multiple from the mean absolute deviation zero center of some vector. The intent of winsorization is to limit the effect of extreme values.

**Value**

An augmented tibble

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<https://en.wikipedia.org/wiki/Winsorizing>

Other Augment Function: [hai\\_fourier\\_augment\(\)](#), [hai\\_fourier\\_discrete\\_augment\(\)](#), [hai\\_hyperbolic\\_augment\(\)](#), [hai\\_polynomial\\_augment\(\)](#), [hai\\_scale\\_zero\\_one\\_augment\(\)](#), [hai\\_scale\\_zscore\\_augment\(\)](#), [hai\\_winsorized\\_truncate\\_augment\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 24
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

hai_winsorized_move_augment(data_tbl, a, .multiple = 3)
```

---

hai\_winsorized\_move\_vec

*Vector Function Winsorize Move*

---

**Description**

Takes a numeric vector and will return a vector of winsorized values.

**Usage**

```
hai_winsorized_move_vec(.x, .multiple = 3)
```

## Arguments

- `.x` A numeric vector
- `.multiple` A positive number indicating how many times the the zero center mean absolute deviation should be multiplied by for the scaling parameter.

## Details

Takes a numeric vector and will return a winsorized vector of values that have been moved some multiple from the mean absolute deviation zero center of some vector. The intent of winsorization is to limit the effect of extreme values.

## Value

A numeric vector

## Author(s)

Steven P. Sanderson II, MPH

## See Also

<https://en.wikipedia.org/wiki/Winsorizing>

This function can be used on it's own. It is also the basis for the function `hai_winsorized_move_augment()`.

Other Vector Function: `hai_fourier_discrete_vec()`, `hai_fourier_vec()`, `hai_hyperbolic_vec()`, `hai_kurtosis_vec()`, `hai_scale_zero_one_vec()`, `hai_scale_zscore_vec()`, `hai_skewness_vec()`, `hai_winsorized_truncate_vec()`

## Examples

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 25
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

vec_1 <- hai_winsorized_move_vec(data_tbl$a, .multiple = 1)

plot(data_tbl$a)
lines(data_tbl$a)
lines(vec_1, col = "blue")
```

---

`hai_winsorized_truncate_augment`*Augment Function Winsorize Truncate*

---

## Description

Takes a numeric vector and will return a tibble with the winsorized values.

## Usage

```
hai_winsorized_truncate_augment(.data, .value, .fraction, .names = "auto")
```

## Arguments

<code>.data</code>	The data being passed that will be augmented by the function.
<code>.value</code>	This is passed <code>rlang::enquo()</code> to capture the vectors you want to augment.
<code>.fraction</code>	A positive fractional between 0 and 0.5 that is passed to the <code>stats::quantile</code> parameter of <code>probs</code> .
<code>.names</code>	The default is "auto"

## Details

Takes a numeric vector and will return a winsorized vector of values that have been truncated if they are less than or greater than some defined fraction of a quantile. The intent of winsorization is to limit the effect of extreme values.

## Value

An augmented tibble

## Author(s)

Steven P. Sanderson II, MPH

## See Also

<https://en.wikipedia.org/wiki/Winsorizing>

Other Augment Function: `hai_fourier_augment()`, `hai_fourier_discrete_augment()`, `hai_hyperbolic_augment()`, `hai_polynomial_augment()`, `hai_scale_zero_one_augment()`, `hai_scale_zscore_augment()`, `hai_winsorized_move_augment()`

**Examples**

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 24
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

hai_winsorized_truncate_augment(data_tbl, a, .fraction = 0.05)
```

---

hai\_winsorized\_truncate\_vec

*Vector Function Winsorize Truncate*

---

**Description**

Takes a numeric vector and will return a vector of winsorized values.

**Usage**

```
hai_winsorized_truncate_vec(.x, .fraction = 0.05)
```

**Arguments**

<code>.x</code>	A numeric vector
<code>.fraction</code>	A positive fractional between 0 and 0.5 that is passed to the <code>stats::quantile</code> parameter of <code>probs</code> .

**Details**

Takes a numeric vector and will return a winsorized vector of values that have been truncated if they are less than or greater than some defined fraction of a quantile. The intent of winsorization is to limit the effect of extreme values.

**Value**

A numeric vector

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

<https://en.wikipedia.org/wiki/Winsorizing>

This function can be used on it's own. It is also the basis for the function `hai_winsorized_truncate_augment()`.

Other Vector Function: `hai_fourier_discrete_vec()`, `hai_fourier_vec()`, `hai_hyperbolic_vec()`, `hai_kurtosis_vec()`, `hai_scale_zero_one_vec()`, `hai_scale_zscore_vec()`, `hai_skewness_vec()`, `hai_winsorized_move_vec()`

**Examples**

```
suppressPackageStartupMessages(library(dplyr))

len_out <- 25
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

vec_1 <- hai_winsorized_truncate_vec(data_tbl$a, .fraction = 0.05)

plot(data_tbl$a)
lines(data_tbl$a)
lines(vec_1, col = "blue")
```

---

hai\_xgboost\_data\_prepper

*Prep Data for XGBoost - Recipe*

---

**Description**

Automatically prep a data.frame/tibble for use in the xgboost algorithm.

**Usage**

```
hai_xgboost_data_prepper(.data, .recipe_formula)
```

**Arguments**

`.data` The data that you are passing to the function. Can be any type of data that is accepted by the data parameter of the `recipes::reciep()` function.

`.recipe_formula` The formula that is going to be passed. For example if you are using the diamonds data then the formula would most likely be something like `price ~ .`

**Details**

This function will automatically prep your data.frame/tibble for use in the XGBoost algorithm.

This function will output a recipe specification.

**Value**

A recipe object

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://parsnip.tidymodels.org/reference/details\\_boost\\_tree\\_xgboost.html](https://parsnip.tidymodels.org/reference/details_boost_tree_xgboost.html)

Other Preprocessor: [hai\\_c50\\_data\\_prepper\(\)](#), [hai\\_cubist\\_data\\_prepper\(\)](#), [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#), [hai\\_earth\\_data\\_prepper\(\)](#), [hai\\_glmnet\\_data\\_prepper\(\)](#), [hai\\_knn\\_data\\_prepper\(\)](#), [hai\\_ranger\\_data\\_prepper\(\)](#), [hai\\_svm\\_poly\\_data\\_prepper\(\)](#), [hai\\_svm\\_rbf\\_data\\_prepper\(\)](#)

**Examples**

```
library(ggplot2)

# Regression
hai_xgboost_data_prepper(.data = diamonds, .recipe_formula = price ~ .)
reg_obj <- hai_xgboost_data_prepper(diamonds, price ~ .)
get_juiced_data(reg_obj)

# Classification
hai_xgboost_data_prepper(Titanic, Survived ~ .)
cla_obj <- hai_xgboost_data_prepper(Titanic, Survived ~ .)
get_juiced_data(cla_obj)
```

---

pca\_your\_recipe

*Perform PCA*

---

**Description**

This is a simple function that will perform PCA analysis on a passed recipe.

**Usage**

```
pca_your_recipe(.recipe_object, .data, .threshold = 0.75, .top_n = 5)
```

**Arguments**

- `.recipe_object` The recipe object you want to pass.
- `.data` The full data set that is used in the original recipe object passed into `.recipe_object` in order to obtain the baked data of the transform.
- `.threshold` A number between 0 and 1. A fraction of the total variance that should be covered by the components.
- `.top_n` How many variables loadings should be returned per PC

**Details**

This is a simple wrapper around some recipes functions to perform a PCA on a given recipe. This function will output a list and return it invisible. All of the components of the analysis will be returned in a list as their own object that can be selected individually. A scree plot is also included. The items that get returned are:

1. `pca_transform` - This is the pca recipe.
2. `variable_loadings`
3. `variable_variance`
4. `pca_estimates`
5. `pca_juiced_estimates`
6. `pca_baked_data`
7. `pca_variance_df`
8. `pca_rotattion_df`
9. `pca_variance_scree_plt`
10. `pca_loadings_plt`
11. `pca_loadings_plotly`
12. `pca_top_n_loadings_plt`
13. `pca_top_n_plotly`

**Value**

A list object with several components.

**Author(s)**

Steven P. Sanderson II, MPH

**See Also**

[https://recipes.tidymodels.org/reference/step\\_pca.html](https://recipes.tidymodels.org/reference/step_pca.html)

Other Data Recipes: [hai\\_data\\_impute\(\)](#), [hai\\_data\\_poly\(\)](#), [hai\\_data\\_scale\(\)](#), [hai\\_data\\_transform\(\)](#), [hai\\_data\\_trig\(\)](#)



**Examples**

```

suppressPackageStartupMessages(library(timetk))
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(purrr))
suppressPackageStartupMessages(library(healthyR.data))
suppressPackageStartupMessages(library(rsample))
suppressPackageStartupMessages(library(recipes))
suppressPackageStartupMessages(library(ggplot2))
suppressPackageStartupMessages(library(plotly))

data_tbl <- healthyR_data %>%
  select(visit_end_date_time) %>%
  summarise_by_time(
    .date_var = visit_end_date_time,
    .by       = "month",
    value     = n()
  ) %>%
  set_names("date_col", "value") %>%
  filter_by_time(
    .date_var = date_col,
    .start_date = "2013",
    .end_date = "2020"
  ) %>%
  mutate(date_col = as.Date(date_col))

splits <- initial_split(data = data_tbl, prop = 0.8)

rec_obj <- recipe(value ~ ., training(splits)) %>%
  step_timeseries_signature(date_col) %>%
  step_rm(matches("(iso$)|(xts$)|(hour)|(min)|(sec)|(am.pm)"))

output_list <- pca_your_recipe(rec_obj, .data = data_tbl)
output_list$pca_variance_scee_plt
output_list$pca_loadings_plt
output_list$pca_top_n_loadings_plt

```

---

step\_hai\_fourier

*Recipes Step Fourier Generator*


---

**Description**

step\_hai\_fourier creates a *specification* of a recipe step that will convert numeric data into either a 'sin', 'cos', or 'sincos' feature that can aid in machine learning.

**Usage**

```

step_hai_fourier(
  recipe,

```

```

    ...,
    role = "predictor",
    trained = FALSE,
    columns = NULL,
    scale_type = c("sin", "cos", "sincos"),
    period = 1,
    order = 1,
    skip = FALSE,
    id = rand_id("hai_fourier")
  )

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class <code>numeric</code>
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
scale_type	A character string of a scaling type, one of "sin", "cos", or "sincos"
period	The number of observations that complete a cycle
order	The fourier term order
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

### Details

**Numeric Variables** Unlike other steps, `step_hai_fourier` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

### Value

For `step_hai_fourier`, an updated version of recipe with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

## See Also

Other Recipes: [step\\_hai\\_fourier\\_discrete\(\)](#), [step\\_hai\\_hyperbolic\(\)](#), [step\\_hai\\_scale\\_zero\\_one\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#), [step\\_hai\\_winsorized\\_move\(\)](#), [step\\_hai\\_winsorized\\_truncate\(\)](#)

## Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

# Create a recipe object
rec_obj <- recipe(a ~ ., data = data_tbl) %>%
  step_hai_fourier(b, scale_type = "sin") %>%
  step_hai_fourier(b, scale_type = "cos") %>%
  step_hai_fourier(b, scale_type = "sincos")

# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prepare(rec_obj), data_tbl)

rec_obj %>% get_juiced_data()
```

---

step\_hai\_fourier\_discrete

*Recipes Step Fourier Discrete Generator*

---

## Description

`step_hai_fourier_discrete` creates a *specification* of a recipe step that will convert numeric data into either a 'sin', 'cos', or 'sincos' feature that can aid in machine learning.

**Usage**

```
step_hai_fourier_discrete(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  columns = NULL,
  scale_type = c("sin", "cos", "sincos"),
  period = 1,
  order = 1,
  skip = FALSE,
  id = rand_id("hai_fourier_discrete")
)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class numeric or date,POSIXct
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
scale_type	A character string of a scaling type, one of "sin","cos", or "sincos"
period	The number of observations that complete a cycle
order	The fourier term order
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

**Details**

**Numeric Variables** Unlike other steps, `step_hai_fourier_discrete` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

**Value**

For `step_hai_fourier_discrete`, an updated version of recipe with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

### See Also

Other Recipes: [step\\_hai\\_fourier\(\)](#), [step\\_hai\\_hyperbolic\(\)](#), [step\\_hai\\_scale\\_zero\\_one\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#), [step\\_hai\\_winsorized\\_move\(\)](#), [step\\_hai\\_winsorized\\_truncate\(\)](#)

### Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

# Create a recipe object
rec_obj <- recipe(a ~ ., data = data_tbl) %>%
  step_hai_fourier_discrete(b, scale_type = "sin") %>%
  step_hai_fourier_discrete(b, scale_type = "cos") %>%
  step_hai_fourier_discrete(b, scale_type = "sincos")

# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prep(rec_obj), data_tbl)

rec_obj %>% get_juiced_data()
```

---

`step_hai_hyperbolic`     *Recipes Step Hyperbolic Generator*

---

### Description

`step_hai_hyperbolic` creates a *specification* of a recipe step that will convert numeric data into either a 'sin', 'cos', or 'tan' feature that can aid in machine learning.

**Usage**

```
step_hai_hyperbolic(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  columns = NULL,
  scale_type = c("sin", "cos", "tan", "sincos"),
  skip = FALSE,
  id = rand_id("hai_hyperbolic")
)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class <code>numeric</code>
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
scale_type	A character string of a scaling type, one of "sin", "cos", "tan" or "sincos"
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

**Details**

**Numeric Variables** Unlike other steps, `step_hai_hyperbolic` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

**Value**

For `step_hai_hyperbolic`, an updated version of `recipe` with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

**See Also**

Other Recipes: [step\\_hai\\_fourier\(\)](#), [step\\_hai\\_fourier\\_discrete\(\)](#), [step\\_hai\\_scale\\_zero\\_one\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#), [step\\_hai\\_winsorized\\_move\(\)](#), [step\\_hai\\_winsorized\\_truncate\(\)](#)

**Examples**

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

# Create a recipe object
rec_obj <- recipe(a ~ ., data = data_tbl) %>%
  step_hai_hyperbolic(b, scale_type = "sin") %>%
  step_hai_hyperbolic(b, scale_type = "cos")

# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prep(rec_obj), data_tbl)

rec_obj %>% get_juiced_data()
```

---

```
step_hai_scale_zero_one
```

*Recipes Data Scale to Zero and One*

---

**Description**

`step_hai_scale_zero_one` creates a *specification* of a recipe step that will convert numeric data into from a time series into its velocity.

**Usage**

```
step_hai_scale_zero_one(
  recipe,
```

```

    ...,
    role = "predictor",
    trained = FALSE,
    columns = NULL,
    skip = FALSE,
    id = rand_id("hai_scale_zero_one")
  )

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class <code>numeric</code>
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

### Details

**Numeric Variables** Unlike other steps, `step_hai_scale_zero_one` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

### Value

For `step_hai_scale_zero_one`, an updated version of `recipe` with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

### See Also

Other Recipes: `step_hai_fourier()`, `step_hai_fourier_discrete()`, `step_hai_hyperbolic()`, `step_hai_scale_zscore()`, `step_hai_winsorized_move()`, `step_hai_winsorized_truncate()`



## Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

data_tbl <- data.frame(a = rnorm(200, 3, 1), b = rnorm(200, 2, 2))

# Create a recipe object
rec_obj <- recipe(a ~ ., data = data_tbl) %>%
  step_hai_scale_zero_one(b)

# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prepare(rec_obj), data_tbl)

rec_obj %>%
  prep() %>%
  juice()
```

---

step\_hai\_scale\_zscore *Recipes Data Scale by Z-Score*

---

## Description

step\_hai\_scale\_zscore creates a *specification* of a recipe step that will convert numeric data into from a time series into its velocity.

## Usage

```
step_hai_scale_zscore(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("hai_scale_zscore")
)
```

## Arguments

recipe            A recipe object. The step will be added to the sequence of operations for this recipe.

...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class <code>numeric</code>
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

### Details

**Numeric Variables** Unlike other steps, `step_hai_scale_zscore` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

### Value

For `step_hai_scale_zscore`, an updated version of recipe with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

### See Also

Other Recipes: `step_hai_fourier()`, `step_hai_fourier_discrete()`, `step_hai_hyperbolic()`, `step_hai_scale_zero_one()`, `step_hai_winsorized_move()`, `step_hai_winsorized_truncate()`

Other Scale: `hai_scale_zero_one_augment()`, `hai_scale_zero_one_vec()`, `hai_scale_zscore_augment()`, `hai_scale_zscore_vec()`

### Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

data_tbl <- data.frame(
  a = mtcars$mpg,
  b = AirPassengers %>% as.vector() %>% head(32)
)

# Create a recipe object
```

```

rec_obj <- recipe(a ~ ., data = data_tbl) %>%
  step_hai_scale_zscore(b)

# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prepare(rec_obj), data_tbl)

rec_obj %>%
  prep() %>%
  juice()

```

---

step\_hai\_winsorized\_move

*Recipes Step Winsorized Move Generator*


---

## Description

step\_hai\_winsorized\_move creates a *specification* of a recipe step that will winsorize numeric data.

## Usage

```

step_hai_winsorized_move(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  columns = NULL,
  multiple = 3,
  skip = FALSE,
  id = rand_id("hai_winsorized_move")
)

```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class <code>numeric</code>
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.

trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
multiple	A positive number indicating how many times the the zero center mean absolute deviation should be multiplied by for the scaling parameter.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

### Details

**Numeric Variables** Unlike other steps, `step_hai_winsorize_move` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

### Value

For `step_hai_winsorize_move`, an updated version of recipe with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

### See Also

Other Recipes: [step\\_hai\\_fourier\(\)](#), [step\\_hai\\_fourier\\_discrete\(\)](#), [step\\_hai\\_hyperbolic\(\)](#), [step\\_hai\\_scale\\_zero\\_one\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#), [step\\_hai\\_winsorized\\_truncate\(\)](#)

### Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

# Create a recipe object
rec_obj <- recipe(b ~ ., data = data_tbl) %>%
```

```

step_hai_winsorized_move(a, multiple = 3)

# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prepare(rec_obj), data_tbl)

rec_obj %>% get_juiced_data()

```

---

step\_hai\_winsorized\_truncate

*Recipes Step Winsorized Truncate Generator*


---

### Description

step\_hai\_winsorized\_truncate creates a *specification* of a recipe step that will winsorize numeric data.

### Usage

```

step_hai_winsorized_truncate(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  columns = NULL,
  fraction = 0.05,
  skip = FALSE,
  id = rand_id("hai_winsorized_truncate")
)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class <code>numeric</code>
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.

columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <code>recipes::prep()</code> is used.
fraction	A positive fractional between 0 and 0.5 that is passed to the <code>stats::quantile</code> parameter of <code>probs</code> .
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

### Details

**Numeric Variables** Unlike other steps, `step_hai_winsorize_truncate` does *not* remove the original numeric variables. `recipes::step_rm()` can be used for this purpose.

### Value

For `step_hai_winsorize_truncate`, an updated version of recipe with the new step added to the sequence of existing steps (if any).

Main Recipe Functions:

- `recipes::recipe()`
- `recipes::prep()`
- `recipes::bake()`

### See Also

Other Recipes: [step\\_hai\\_fourier\(\)](#), [step\\_hai\\_fourier\\_discrete\(\)](#), [step\\_hai\\_hyperbolic\(\)](#), [step\\_hai\\_scale\\_zero\\_one\(\)](#), [step\\_hai\\_scale\\_zscore\(\)](#), [step\\_hai\\_winsorized\\_move\(\)](#)

### Examples

```
suppressPackageStartupMessages(library(dplyr))
suppressPackageStartupMessages(library(recipes))

len_out <- 10
by_unit <- "month"
start_date <- as.Date("2021-01-01")

data_tbl <- tibble(
  date_col = seq.Date(from = start_date, length.out = len_out, by = by_unit),
  a = rnorm(len_out),
  b = runif(len_out)
)

# Create a recipe object
rec_obj <- recipe(b ~ ., data = data_tbl) %>%
  step_hai_winsorized_truncate(a, fraction = 0.05)
```

```
# View the recipe object
rec_obj

# Prepare the recipe object
prep(rec_obj)

# Bake the recipe object - Adds the Time Series Signature
bake(prepare(rec_obj), data_tbl)

rec_obj %>% get_juiced_data()
```

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