

# Package ‘math3190package’

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**Title** Package for MATH 3190 - Fundamentals of Data Science at SUU

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**Description** This package contains functions and data files needed for MATH 3190 - Fundamentals of Data Science taught at Southern Utah University.

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bootstrapping	<i>Outputs and restyles an object from the boot() function from the boot package.</i>
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## Description

This function relies on the `boot` function. It generates R bootstrap replicates of a statistic applied to data. Both parametric and nonparametric resampling are possible. For the nonparametric bootstrap, possible resampling methods are the ordinary bootstrap, the balanced bootstrap, antithetic resampling, and permutation. For nonparametric multi-sample problems stratified resampling is used: this is specified by including a vector of strata in the call to `boot`. Importance resampling weights may be specified.

## Usage

```
bootstrapping(data, statistic, R, ...)
```

## Arguments

<code>data</code>	The data as a vector, matrix or data frame. If it is a matrix or data frame then each row is considered as one multivariate observation.
<code>statistic</code>	A function which when applied to data returns a vector containing the statistic(s) of interest. When <code>sim = "parametric"</code> , the first argument to <code>statistic</code> must be the data. For each replicate a simulated dataset returned by <code>ran.gen</code> will be passed. In all other cases <code>statistic</code> must take at least two arguments. The first argument passed will always be the original data. The second will be a vector of indices, frequencies or weights which define the bootstrap sample. Further, if predictions are required, then a third argument is required which would be a vector of the random indices used to generate the bootstrap predictions. Any further arguments can be passed to <code>statistic</code> through the <code>...</code> argument.

R	The number of bootstrap replicates. Usually this will be a single positive integer. For importance resampling, some resamples may use one set of weights and others use a different set of weights. In this case R would be a vector of integers where each component gives the number of resamples from each of the rows of weights.
...	See more documentation in the <a href="#">boot</a> function.

**Value**

This function returns an object of class boot3150 that works much like an object of class boot from the boot package.

**Examples**

```
med <- function(x, i){median(x[i])}
sample_data <- rnorm(100)
boot_out <- bootstrapping(sample_data, med, 1000)
boot_out
```

**boot\_ci**

*Outputs and restyles an object from the boot.ci() function from the boot package.*

**Description**

This function relies on the [boot.ci](#) function. This function generates 5 different types of equi-tailed two-sided nonparametric confidence intervals. These are the first order normal approximation, the basic bootstrap interval, the studentized bootstrap interval, the bootstrap percentile interval, and the adjusted bootstrap percentile (BCa) interval. All or a subset of these intervals can be generated.

**Usage**

```
boot_ci(boot_out, conf = 0.95, type = "all", ...)
```

**Arguments**

boot_out	An object of class "boot" or "boot3150" containing the output of a bootstrap calculation.
conf	A scalar or vector containing the confidence level(s) of the required interval(s).
type	A vector of character strings representing the type of intervals required. The value should be any subset of the values c("norm", "basic", "stud", "perc", "bca") or simply "all" which will compute all five types of intervals.
...	See more documentation in the <a href="#">boot.ci</a> function.

**Value**

This function returns an object of class boot3150ci that works much like an object of class bootci from the boot package.

## Examples

```
med <- function(x, i){median(x[i])}
sample_data <- rnorm(100)
boot_out <- bootstrapping(sample_data, med, 1000)
bootci_out <- boot_ci(boot_out)
```

cars99

*cars99 Data Set*

## Description

This data set contains information on 109 vehicles from 1999.

## Usage

cars99

## Format

A tibble with 109 rows and 11 variables:

**Model** The vehicle model name.

**CityMPG** The miles per gallon (MPG) for the vehicle in the city.

**HwyMPG** The miles per gallon (MPG) for the vehicle on the highway

**FuelCap** The fuel capacity of the vehicle in gallons.

**Weight** The weight of the vehicle in lbs.

**FrontWt** The front weight of the vehicle.

**Accel0\_30** The time it takes, in seconds, for the vehicle to accelerate from 0 to 30 mph.

**Accel0\_60** The time it takes, in seconds, for the vehicle to accelerate from 0 to 60 mph.

**QtrMile** The time it takes, in seconds, for the vehicle to travel a quarter of a mile.

classifier\_plot

*Plots the Points and Predicted Region for a Classifier Model*

## Description

This function plots the points and predicted region for a classifier model of class "naive\_bayes", "lda", or "svm". This function can support up to three predictor variables: two of which can be quantitative and one can be categorical.

## Usage

```
classifier_plot(classifier_model, data, res = 200, sv_diamonds = TRUE)
```

**Arguments**

<code>classifier_model</code>	The classifier model of class "naive_bayes", "lda", or "svm".
<code>data</code>	The dataset that should be plotted. This need not be the dataset on which the classifier was trained.
<code>res</code>	The resolution of the plot. Larger values will make the lines look straighter and make the grid more fine, but take longer.
<code>sv_diamonds</code>	A boolean indicating whether the support vectors should be plotted with diamonds around them when plotting a support vector machine classifier.

**Value**

This function returns a ggplot object plot. It can be added onto like any ggplot object.

**Examples**

```
data(mtcars)
mtcars2 <- mtcars |> mutate(vs = factor(vs), am = factor(am))
classifier <- MASS::lda(vs ~ mpg + disp + am, data = mtcars2)
classifier_plot(classifier, data = mtcars2)
```

**ESL.mixture***ESL Mixture Data Set***Description**

These are the data used in Figures 2.1-2.3, and elsewhere through the Elements of Statistical Learning book.

**Usage**

`ESL.mixture`

**Format**

A list with 8 components The components are

- x** 200 x 2 matrix of training predictors.
- y** class variable; logical vector of TRUES and FALSES - 100 of each.
- xnew** matrix 6831 x 2 of lattice points in predictor space.
- prob** vector of 6831 probabilities (of class TRUE) at each lattice point.
- marginal** marginal probability at each lattice point.
- px1** 69 lattice coordinates for x.1.
- px2** 99 lattice values for x.2 (69\*99=6831).
- means** 20 x 2 matrix of the mixture centers, first ten for one class, next ten for the other

**ilogit***Compute the Inverse Logit of a Value***Description**

A function that computes the inverse logit (the inverse log odds) of a value.

**Usage**

```
ilogit(x)
```

**Arguments**

**x** A value for which that we want to find the inverse.

**Examples**

```
data(mtcars)
mod <- glm(vs ~ mpg, data = mtcars, family = binomial())
ilogit(predict(mod, data.frame(mpg = 20)))
```

**influence\_plots***Creates many diagnostic plots***Description**

This function creates many diagnostic plots for a given model. These plots include residual plots, a QQ plot, a leverage plot, a Cook's distance plot, a DFFits plot, and DFBetas plots for the intercept and all slopes.

**Usage**

```
influence_plots(model, missing_group = NULL)
```

**Arguments**

**model** The model for which we would like these plots. This can be of class "lm" or "glm" with a binomial family.  
**missing\_group** Used for multinomial regression to indicate which group is not being plotted.

**Value**

This function returns plots. The user must press "enter" or "return" to view subsequent plots.

**Examples**

```
mod <- lm(mpg ~ disp, data = mtcars)
influence_plots(mod)
```

---

**interaction\_plot**      *Two-way Interaction Plot with ggplot2*

---

**Description**

This function is similar to [interaction.plot](#), but plots in ggplot2 instead of base R. It plots the mean (or other summary) of the response for two-way combinations of factors, thereby illustrating possible interactions.

**Usage**

```
interaction_plot(  
  df,  
  response,  
  x_factor,  
  group_factor,  
  linewidth = 1,  
  point_size = 2,  
  .f = mean,  
  ...  
)
```

**Arguments**

<code>df</code>	The data frame that contains the response and factor variables.
<code>response</code>	The name of the numeric variable giving the response. Can be entered with or without quotes.
<code>x_factor</code>	The name of a factor whose levels will form the x axis. Can be entered with or without quotes.
<code>group_factor</code>	The name of a factor whose levels will split up the other variables. Can be entered with or without quotes.
<code>linewidth</code>	The linewidth of the lines plotted. Enter 0 if the lines should be omitted.
<code>point_size</code>	The size of the points plotted. Enter 0 if the points should be omitted.
<code>...</code>	Other parameters that can be passed to <a href="#">geom_point</a> .

**Value**

This function returns a two-way interaction plot as a ggplot object. All the ggplot add-on functions can be added onto the object returned for more customization.

**Examples**

```
data(warpbreaks)  
interaction_plot(warpbreaks, breaks, wool, tension)  
interaction_plot(warpbreaks, "breaks", "wool", "tension")
```

<code>logistic_plots</code>	<i>Creates many diagnostic plots for logistic or multinomial models</i>
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## Description

This function feeds into the `influence_plots()` function to create many diagnostic plots for a given logistic or multinomial model. These plots include residual plots, a leverage plot, a Cook's distance plot, a DFFits plot, and DFBetas plots for the intercept and all slopes.

## Usage

```
logistic_plots(model)
```

## Arguments

- |                            |   |
|----------------------------|---|
| <code>model</code>         | The model for which we would like these plots. This can be of class "glm" with a binomial family or of class "multinom" from the <code>nnet</code> library. |
| <code>missing_group</code> | Used for multinomial regression to indicate which group is not being plotted.   |

## Value

This function returns plots. The user must press "enter" or "return" to view subsequent plots.

## Examples

```
mod <- glm(vs ~ disp, data = mtcars, family = "binomial")
logistic_plots(mod)
```

<code>multinom_summary</code>	<i>Outputs estimates, standard errors, test statistic, and p-values for a multinomial logistic regression object from the nnet library.</i>
-------------------------------	---

## Description

When given an `nnet` `multinom` object, this function will compute the test statistic and p-value for each term in the models and return a tibble with the coefficients, standard errors, test statistic, and p-values in a list with one list element per model.

## Usage

```
multinom_summary(model)
```

## Arguments

- |                    |                                     |
|--------------------|-------------------------------------|
| <code>model</code> | An object of type "multinom" "nnet" |
|--------------------|-------------------------------------|

### Value

This function returns a list of tibbles with the coefficient estimates, the standard errors, the test statistics, and the p-values for the models comparing each category to the baseline. The AIC and deviance are also given as the last elements in the list.

### Examples

```
mod <- mtcars |>
  mutate(carb = factor(carb)) |>
  multinom(carb ~ mpg + disp + hp, data = _)
multinom_summary(mod)
```

plot.boot3150

*Plots a boot3150 object*

### Description

When given a boot3150 object, this function plots the samples with both histograms and QQ plots.

### Usage

```
## S3 method for class 'boot3150'
plot(x)
```

### Arguments

x An object of class boot3150

### Value

This function plots the boot3150 object.

plot\_tukey

*Plots the Tukey HSD Confidence Intervals*

### Description

This function plots the Tukey HSD confidence intervals when given a tibble outputted by the [tukeyhsd](#) function. If more than one variable, all will be plotted by default, but that can be adjusted with the which option.

### Usage

```
plot_tukey(t, which = "all")
```

### Arguments

t	The Tukey HSD tibble outputted by the <a href="#">tukeyhsd</a> function.
which	Indicates which variable for which the graph should be created. The default is "all", but a variable name used in the model can also be entered (in quotes).

**Value**

This function returns plots. The user must press "enter" or "return" to view subsequent plots if there are any.

**Examples**

```
data(warpbreaks)
fm1 <- aov(breaks ~ wool + tension, data = warpbreaks)
t <- tukeyhsd(fm1, "tension", ordered = TRUE)
plot_tukey(t)
```

**print.boot3150**      *Prints a boot3150 object*

**Description**

When given a boot3150 object, this function ensures it prints correctly.

**Usage**

```
## S3 method for class 'boot3150'
print(x)
```

**Arguments**

x      An object of class boot3150

**Value**

This function prints the output for a boot3150 object.

**print.boot3150ci**      *Prints a boot3150ci object*

**Description**

When given a boot3150ci object, this function ensures it prints correctly.

**Usage**

```
## S3 method for class 'boot3150ci'
print(x)
```

**Arguments**

x      An object of class boot3150ci

**Value**

This function prints the output for a boot3150ci object.

---

`print.innerboot3150ci` *Prints a innerboot3150ci object*

---

**Description**

When given a innerboot3150ci object, this function ensures it prints correctly.

**Usage**

```
## S3 method for class 'innerboot3150ci'
print(x)
```

**Arguments**

<code>x</code>	An object of class innerboot3150ci
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**Value**

This function prints the output for a innerboot3150ci object.

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<code>residual_plots</code>	<i>Creates residual and QQ plots</i>
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---

**Description**

This function creates a residual and a QQ plot for a given model.

**Usage**

```
residual_plots(model, resid_type = "jackknife", smoother = TRUE, se = T)
```

**Arguments**

<code>model</code>	The model for which we would like these plots. This can be of class "lm" or "glm" with a binomial family.
<code>resid_type</code>	The type of residuals that should be used in the plots. The options are "jackknife", "raw", "standard", and "pearson". Jackknife (externally studentized) residuals are used by default for "lm" objects. Standardized Pearson residuals are used for "glm" objects regardless of what is entered here.
<code>smoother</code>	Logical indicating whether a loess smoother should be added to the residual plot.
<code>se</code>	Logical indicating whether the standard error bands should be added to the smoother.

**Value**

This function returns plots. The user must press "enter" or "return" to view subsequent plots.

**Examples**

```
mod <- lm(mpg ~ disp, data = mtcars)
residual_plots(mod)
```

---

**umap\_plot***Create a 1D or 2D UMAP Plot*

---

**Description**

This function plots the UMAP layouts in either 1D or 2D and can optionally color them using a variable from a given dataset.

**Usage**

```
umap_plot(umap_object, data, color_umap = NULL, ...)
```

**Arguments**

umap_object	An object fit by the <code>umap()</code> function in the <code>umap</code> library.
data	The dataset that has the variable to be used for <code>color_umap</code> .
...	Extra arguments for the <code>geom_point()</code> function when plotting in 2D or for the <code>geom_histogram()</code> function when plotting in 1D.
res	The variable that should be used to change the color of the points on the graph.

**Value**

This function returns a ggplot object plot. It can be added onto like any ggplot object.

**Examples**

```
data(iris)
iris_umap <- iris[,1:4] |> umap(random_state = 1)
umap_plot(iris_umap, data = iris, color_umap = "Species")
```

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