

# Package ‘auditor’

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**Title** Model Audit - Verification, Validation, and Error Analysis

**Version** 1.3.0

**Description** Provides an easy to use unified interface for creating validation plots for any model. The 'auditor' helps to avoid repetitive work consisting of writing code needed to create residual plots. This visualizations allow to asses and compare the goodness of fit, performance, and similarity of models.

**Depends** R (>= 3.5.0)

**License** GPL

**Encoding** UTF-8

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**Imports** DALEX, ggplot2, ggrepel, grid, gridExtra, hnp, scales

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**VignetteBuilder** knitr

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audit	<i>Deprecated</i>
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---

## Description

The `audit()` function is deprecated, use [explain](#) from the DALEX package instead.

## Usage

```
audit(
  object,
  data = NULL,
  y = NULL,
  predict.function = NULL,
  residual.function = NULL,
  label = NULL,
```

```

    predict_function = NULL,
    residual_function = NULL
  )

```

### Arguments

<code>object</code>	An object containing a model or object of class explainer (see <a href="#">explain</a> ).
<code>data</code>	Data.frame or matrix - data that will be used by further validation functions. If not provided, will be extracted from the model.
<code>y</code>	Response vector that will be used by further validation functions. Some functions may require an integer vector containing binary labels with values 0,1. If not provided, will be extracted from the model.
<code>predict.function</code>	Function that takes two arguments: model and data. It should return a numeric vector with predictions.
<code>residual.function</code>	Function that takes three arguments: model, data and response vector. It should return a numeric vector with model residuals for given data. If not provided, response residuals ( $y - \hat{y}$ ) are calculated.
<code>label</code>	Character - the name of the model. By default it's extracted from the 'class' attribute of the model.
<code>predict_function</code>	Function that takes two arguments: model and data. It should return a numeric vector with predictions.
<code>residual_function</code>	Function that takes three arguments: model, data and response vector. It should return a numeric vector with model residuals for given data. If not provided, response residuals ( $y - \hat{y}$ ) are calculated.

### Value

An object of class explainer.

### Examples

```

data(titanic_imputed, package = "DALEX")

model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)
audit_glm <- audit(model_glm,
  data = titanic_imputed,
  y = titanic_imputed$survived)

p_fun <- function(model, data) { predict(model, data, response = "link") }
audit_glm_newpred <- audit(model_glm,
  data = titanic_imputed,
  y = titanic_imputed$survived,
  predict.function = p_fun)

```

```
library(randomForest)
model_rf <- randomForest(Species ~ ., data=iris)
audit_rf <- audit(model_rf)
```

---

auditorData	<i>auditorData</i>
-------------	--------------------

---

### Description

The auditor Data is an artificial data set. It consists of 2000 observations. First four of simulated variables are treated as continuous while the fifth one is categorical.

### Examples

```
data("auditorData", package = "auditor")
head(auditorData)
```

---

check_residuals	<i>Automated tests for model residuals</i>
-----------------	--

---

### Description

Currently three tests are performed - for outliers in residuals - for autocorrelation in target variable or in residuals - for trend in residuals as a function of target variable (detection of bias)

### Usage

```
check_residuals(object, ...)
```

### Arguments

object	An object of class 'explainer' created with function <a href="#">explain</a> from the DALEX package.
...	other parameters that will be passed to further functions.

### Value

list with statistics for particular checks

## Examples

```
dragons <- DALEX::dragons[1:100, ]
lm_model <- lm(life_length ~ ., data = dragons)
lm_audit <- audit(lm_model, data = dragons, y = dragons$life_length)
check_residuals(lm_audit)
## Not run:
library("randomForest")
rf_model <- randomForest(life_length ~ ., data = dragons)
rf_audit <- audit(rf_model, data = dragons, y = dragons$life_length)
check_residuals(rf_audit)

## End(Not run)
```

---

check\_residuals\_autocorrelation

*Checks for autocorrelation in target variable or in residuals*

---

## Description

Checks for autocorrelation in target variable or in residuals

## Usage

```
check_residuals_autocorrelation(object, method = "pearson")
```

## Arguments

object	An object of class 'explainer' created with function <a href="#">explain</a> from the DALEX package.
method	will be passed to the cor.test functions

## Value

autocorrelation between target variable and between residuals

## Examples

```
dragons <- DALEX::dragons[1:100, ]
lm_model <- lm(life_length ~ ., data = dragons)
lm_audit <- audit(lm_model, data = dragons, y = dragons$life_length)
check_residuals_autocorrelation(lm_audit)
```

---

check\_residuals\_outliers  
*Checks for outliers*

---

**Description**

Checks for outliers

**Usage**

```
check_residuals_outliers(object, n = 5)
```

**Arguments**

object	An object of class 'explainer' created with function <a href="#">explain</a> from the DALEX package.
n	number of lowest and highest standardized residuals to be presented

**Value**

indexes of lowest and highest standardized residuals

**Examples**

```
dragons <- DALEX::dragons[1:100, ]  
lm_model <- lm(life_length ~ ., data = dragons)  
lm_audit <- audit(lm_model, data = dragons, y = dragons$life_length)  
check_residuals_outliers(lm_audit)
```

---

check\_residuals\_trend *Checks for trend in residuals*

---

**Description**

Calculates loess fit for residuals and then extracts statistics that shows how far is this fit from one without trend

**Usage**

```
check_residuals_trend(object, B = 20)
```

**Arguments**

object	An object of class 'explainer' created with function <a href="#">explain</a> from the DALEX package.
B	number fo samplings

**Value**

standardized loess fit for residuals

**Examples**

```
library(DALEX)
dragons <- DALEX::dragons[1:100, ]
lm_model <- lm(life_length ~ ., data = dragons)
lm_exp <- explain(lm_model, data = dragons, y = dragons$life_length)
library(auditor)
check_residuals_trend(lm_exp)
```

---

model\_cooksdistance    *Cook's distances*

---

**Description**

Calculates Cook's distances for each observation. Please, note that it will work only for functions with specified update method.

**Usage**

```
model_cooksdistance(object)

observationInfluence(object)
```

**Arguments**

object            An object of class explainer created with function [explain](#) from the DALEX package.

**Value**

An object of the class auditor\_model\_cooksdistance.

**References**

Cook, R. Dennis (1977). "Detection of Influential Observations in Linear Regression". doi:10.2307/1268249.

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# use DALEX package to wrap up a model into explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
```



```
      y = titanic_imputed$survived)

# validate a model with auditor
mc <- model_cooksdistance(glm_audit)
mc

plot(mc)
```

---

model_evaluation	<i>Create model evaluation explanation</i>
------------------	--

---

### Description

Creates explanation of classification model.

Returns, among others, true positive rate (tpr), false positive rate (fpr), rate of positive prediction (rpp), and true positives (tp).

Created object of class `auditor_model_evaluation` can be used to plot Receiver Operating Characteristic (ROC) curve (plot `plot_roc`) and LIFT curve (plot `plot_lift`).

### Usage

```
model_evaluation(object)
```

```
modelEvaluation(object)
```

### Arguments

`object` An object of class `explainer` created with function `explain` from the DALEX package.

### Value

An object of the class `auditor_model_evaluation`.

### Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data= titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
me <- model_evaluation(glm_audit)
```

```
me  
plot(me)
```

---

model_halfnormal	<i>Create Halfnormal Explanation</i>
------------------	--------------------------------------

---

## Description

Creates `auditor_model_halfnormal` object that can be used for plotting halfnormal plot.

## Usage

```
model_halfnormal(object, quant = FALSE, ...)  
modelFit(object, quant = FALSE, ...)
```

## Arguments

<code>object</code>	An object of class <code>explainer</code> created with function <code>explain</code> from the DALEX package.
<code>quant</code>	if TRUE values on axis are on quantile scale.
<code>...</code>	other parameters passed do <code>hnp</code> function.

## Value

An object of the class `auditor_model_halfnormal`.

## References

Moral, R., Hinde, J., & Demétrio, C. (2017). Half-Normal Plots and Overdispersed Models in R: The `hnp` Package.  
doi:<http://dx.doi.org/10.18637/jss.v081.i10>

## Examples

```
data(titanic_imputed, package = "DALEX")  
  
# fit a model  
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)  
  
glm_audit <- audit(model_glm,  
                  data = titanic_imputed,  
                  y = titanic_imputed$survived)  
  
# validate a model with auditor  
mh <- model_halfnormal(glm_audit)  
mh
```

```
plot(mh)
```

---

```
model_performance      Create Model Performance Explanation
```

---

## Description

Creates `auditor_model_performance` object that can be used to plot radar with ranking of models.

## Usage

```
model_performance(
  object,
  score = c("mae", "mse", "rec", "rroc"),
  new_score = NULL,
  data = NULL,
  ...
)
```

```
modelPerformance(
  object,
  score = c("mae", "mse", "rec", "rroc"),
  new_score = NULL
)
```

## Arguments

<code>object</code>	An object of class <code>explainer</code> created with function <code>explain</code> from the DALEX package.
<code>score</code>	Vector of score names to be calculated. Possible values: <code>acc</code> , <code>auc</code> , <code>cookdistance</code> , <code>dw</code> , <code>f1</code> , <code>gini</code> , <code>halfnormal</code> , <code>mae</code> , <code>mse</code> , <code>peak</code> , <code>precision</code> , <code>r2</code> , <code>rec</code> , <code>recall</code> , <code>rmse</code> , <code>rroc</code> , <code>runs</code> , <code>specificity</code> , <code>one_minus_acc</code> , <code>one_minus_auc</code> , <code>one_minus_f1</code> , <code>one_minus_gini</code> , <code>one_minus_precision</code> , <code>one_minus_recall</code> , <code>one_minus_specificity</code> (for detailed description see functions in <code>see also</code> section). Pass <code>NULL</code> if you want to use only custom scores by <code>new_score</code> parameter.
<code>new_score</code>	A named list of functions that take one argument: object of class <code>'explainer'</code> and return a numeric value. The measure calculated by the function should have the property that lower score value indicates better model.
<code>data</code>	New data that will be used to calculate scores. Pass <code>NULL</code> if you want to use data from <code>object</code> .
<code>...</code>	Other arguments dependent on the score list.

## Value

An object of the class `auditor_model_performance`.

**See Also**

[score\\_acc](#), [score\\_auc](#), [score\\_cooksdistance](#), [score\\_dw](#), [score\\_f1](#), [score\\_gini](#), [score\\_halfnormal](#), [score\\_mae](#), [score\\_mse](#), [score\\_peak](#), [score\\_precision](#), [score\\_r2](#), [score\\_rec](#), [score\\_recall](#), [score\\_rmse](#), [score\\_rroc](#), [score\\_runs](#), [score\\_specificity](#), [score\\_one\\_minus\\_acc](#), [score\\_one\\_minus\\_auc](#), [score\\_one\\_minus\\_f1](#), [score\\_one\\_minus\\_precision](#), [score\\_one\\_minus\\_gini](#), [score\\_one\\_minus\\_recall](#), [score\\_one\\_minus\\_specificity](#)

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# use DALEX package to wrap up a model into explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
library(auditor)
mp <- model_performance(glm_audit)
mp

plot(mp)
```

---

model\_residual

*Create Model Residuals Explanation*


---

**Description**

Creates `auditor_model_residual` that contains sorted residuals. An object can be further used to generate plots. For the list of possible plots see [see also](#) section.

**Usage**

```
model_residual(object, ...)
```

```
modelResiduals(object, ...)
```

**Arguments**

<code>object</code>	An object of class <code>explainer</code> created with function <a href="#">explain</a> from the DALEX package.
<code>...</code>	other parameters

**Value**

An object of the class `auditor_model_residual`.

**See Also**

[plot\\_acf](#), [plot\\_autocorrelation](#), [plot\\_residual](#), [plot\\_residual\\_boxplot](#), [plot\\_pca](#), [plot\\_correlation](#), [plot\\_pr](#)

**Examples**

```
library(DALEX)

# fit a model
model_glm <- glm(m2.price ~ ., data = apartments)

glm_audit <- explain(model_glm,
                    data = apartments,
                    y = apartments$m2.price)

# validate a model with auditor
mr <- model_residual(glm_audit)
mr

plot(mr)
```

---

plotD3

*Model Diagnostic Plots in D3 with r2d3 package.*

---

**Description**

This function provides several diagnostic plots for regression and classification models. Provide object created with one of auditor's computational functions, [model\\_residual](#), [model\\_cooksdistance](#), [model\\_evaluation](#), [model\\_performance](#), [model\\_evaluation](#).

**Usage**

```
plotD3(x, ...)

plotD3_auditor(x, ..., type = "residual")

## S3 method for class 'auditor_model_residual'
plotD3(x, ..., type = "residual")

## S3 method for class 'auditor_model_halfnormal'
plotD3(x, ..., type = "residual")

## S3 method for class 'auditor_model_evaluation'
plotD3(x, ..., type = "residual")
```

```
## S3 method for class 'auditor_model_cooksdistance'
plotD3(x, ..., type = "residual")
```

### Arguments

x	object of class <code>auditor_model_residual</code> (created with <a href="#">model_residual</a> function), <code>auditor_model_performance</code> (created with <a href="#">model_performance</a> function), <code>auditor_model_evaluation</code> (created with <a href="#">model_evaluation</a> function), <code>auditor_model_cooksdistance</code> (created with <a href="#">model_cooksdistance</a> function), or <code>auditor_model_halfnormal</code> (created with <a href="#">model_halfnormal</a> function).
...	other arguments dependent on the type of plot or additional objects of classes <code>'auditor_model_residual'</code> , <code>'auditor_model_performance'</code> , <code>'auditor_model_evaluation'</code> , <code>'auditor_model_cooksdistance'</code> , or <code>'auditor_model_halfnormal'</code> .
type	the type of plot. Single character. Possible values: <code>'acf'</code> , <code>'autocorrelation'</code> , <code>'cooksdistance'</code> , <code>'halfnormal'</code> , <code>'prediction'</code> , <code>'residual'</code> , <code>'performance'</code> , <code>'evaluation'</code> , <code>'cooksdistance'</code> , <code>'halfnormal'</code> (for detailed description see corresponding functions in <a href="#">see also</a> section).

### See Also

[plotD3\\_acf](#), [plotD3\\_autocorrelation](#), [plotD3\\_cooksdistance](#), [plotD3\\_halfnormal](#), [plotD3\\_residual](#), [plotD3\\_prediction](#), [plotD3\\_performance](#), [plotD3\\_evaluation](#), [plotD3\\_cooksdistance](#), [plotD3\\_halfnormal](#)

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plotD3(mr_lm)
plotD3(mr_lm, type = "prediction")

hn_lm <- model_halfnormal(lm_audit)
plotD3(hn_lm)
```

---

plotD3\_acf

*Plot Autocorrelation Function in D3 with r2d3 package.*

---

### Description

Plot Autocorrelation Function of models' residuals.

**Usage**

```
plotD3_acf(object, ..., variable = NULL, alpha = 0.95, scale_plot = FALSE)
```

```
plotD3ACF(object, ..., variable = NULL, alpha = 0.95, scale_plot = FALSE)
```

**Arguments**

object	An object of class 'auditor_model_residual' created with <a href="#">model_residual</a> function.
...	Other 'auditor_model_residual' objects to be plotted together.
variable	Name of variable to order residuals on a plot. If variable="_y_", the data is ordered by a vector of actual response (y parameter passed to the <a href="#">explain</a> function). If variable = "_y_hat_" the data on the plot will be ordered by predicted response. If variable = NULL, unordered observations are presented.
alpha	Confidence level of the interval.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.

**Value**

a 'r2d3' object.

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plotD3_acf(mr_lm)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plotD3_acf(mr_lm, mr_rf)
```

---

 plotD3\_autocorrelation

*Autocorrelation Plot in D3 with r2d3 package.*


---

### Description

Plot of i-th residual vs i+1-th residual.

### Usage

```
plotD3_autocorrelation(
  object,
  ...,
  variable = NULL,
  points = TRUE,
  smooth = FALSE,
  point_count = NULL,
  single_plot = TRUE,
  scale_plot = FALSE,
  background = FALSE
)
```

```
plotD3Autocorrelation(
  object,
  ...,
  variable = NULL,
  points = TRUE,
  smooth = FALSE,
  point_count = NULL,
  single_plot = TRUE,
  scale_plot = FALSE,
  background = FALSE
)
```

### Arguments

object	An object of class 'auditor_model_residual' created with <a href="#">model_residual</a> function.
...	Other 'auditor_model_residual' objects to be plotted together.
variable	Name of variable to order residuals on a plot. If variable="_y_", the data is ordered by a vector of actual response (y parameter passed to the <a href="#">explain</a> function).
points	Logical, indicates whenever observations should be added as points. By default it's TRUE.
smooth	Logical, indicates whenever smoothed lines should be added. By default it's FALSE.



point_count	Number of points to be plotted per model. Points will be chosen randomly. By default plot all of them.
single_plot	Logical, indicates whenever single or facets should be plotted. By default it's TRUE.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.
background	Logical, available only if single_plot = FALSE. Indicates whenever background plots should be plotted. By default it's FALSE.

**Value**

a r2d3 object

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plotD3_autocorrelation(mr_lm)
plotD3_autocorrelation(mr_lm, smooth = TRUE)
```

---

plotD3\_cooksdistance *Influence of observations Plot in D3 with r2d3 package.*

---

**Description**

Plot of Cook's distances used for estimate the influence of an single observation.

**Usage**

```
plotD3_cooksdistance(  
  object,  
  ...,  
  nlabel = 3,  
  single_plot = FALSE,  
  scale_plot = FALSE,  
  background = FALSE  
)
```

```
plotD3CooksDistance(  
  object,  
  ...,  
  nlabel = 3,  
  single_plot = FALSE,  
  scale_plot = FALSE,  
  background = FALSE  
)
```

### Arguments

object	An object of class 'auditor_model_cooksdistance' created with <a href="#">model_cooksdistance</a> function.
...	Other objects of class 'auditor_model_cooksdistance'.
nlabel	Number of observations with the biggest Cook's distances to be labeled.
single_plot	Logical, indicates whenever single or facets should be plotted. By default it's FALSE.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.
background	Logical, available only if single_plot = FALSE. Indicates whenever background plots should be plotted. By default it's FALSE.

### Details

Cook's distance is a tool for identifying observations that may negatively affect the model. They may be also used for indicating regions of the design space where it would be good to obtain more observations. Data points indicated by Cook's distances are worth checking for validity.

Cook's Distances are calculated by removing the  $i$ -th observation from the data and recalculating the model. It shows how much all the values in the model change when the  $i$ -th observation is removed.

For model classes other than `lm` and `glm` the distances are computed directly from the definition.

### Value

a `r2d3` object

### References

Cook, R. Dennis (1977). "Detection of Influential Observations in Linear Regression". doi:10.2307/1268249.

### See Also

[plot\\_cooksdistance](#)

**Examples**

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
cd_lm <- model_cooksdistance(lm_audit)

# plot results
plotD3_cooksdistance(cd_lm, nlabel = 5)

```

---

plotD3\_halfnormal      *Plot Half-Normal in D3 with r2d3 package.*

---

**Description**

The half-normal plot is one of the tools designed to evaluate the goodness of fit of a statistical models. It is a graphical method for comparing two probability distributions by plotting their quantiles against each other. Points on the plot correspond to ordered absolute values of model diagnostic (i.e. standardized residuals) plotted against theoretical order statistics from a half-normal distribution.

**Usage**

```

plotD3_halfnormal(object, ..., quantiles = FALSE, sim = 99, scale_plot = FALSE)

plotD3HalfNormal(object, ..., quantiles = FALSE, sim = 99, scale_plot = FALSE)

```

**Arguments**

object	An object of class 'auditor_model_halfnormal' created with <a href="#">model_halfnormal</a> function.
...	Other 'auditor_model_halfnormal' objects.
quantiles	If TRUE values on axis are on quantile scale.
sim	Number of residuals to simulate.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.

**Value**

a r2d3 object

**See Also**

[model\\_halfnormal](#)  
[score\\_halfnormal](#), [plot\\_halfnormal](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
hn_lm <- model_halfnormal(lm_audit)

# plot results
plotD3_halfnormal(hn_lm)
```

---

plotD3\_lift

*Plot LIFT in D3 with r2d3 package.*


---

**Description**

LIFT is a plot of the rate of positive prediction against true positive rate for the different thresholds. It is useful for measuring and comparing the accuracy of the classifiers.

**Usage**

```
plotD3_lift(object, ..., scale_plot = FALSE, zeros = TRUE)

plotD3LIFT(object, ..., scale_plot = FALSE)
```

**Arguments**

object	An object of class 'auditor_model_evaluation' created with <a href="#">model_evaluation</a> function.
...	Other 'auditor_model_evaluation' objects to be plotted together.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.
zeros	Logical. It makes the lines start from the $(0, 0)$ point. By default it's TRUE.

**Value**

a r2d3 object

**See Also**[plot\\_lift](#)**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
eva_glm <- model_evaluation(glm_audit)

# plot results
plot_roc(eva_glm)
plot(eva_glm)

#add second model
model_glm_2 <- glm(survived ~ .-age, family = binomial, data = titanic_imputed)
glm_audit_2 <- audit(model_glm_2,
                   data = titanic_imputed,
                   y = titanic_imputed$survived,
                   label = "glm2")
eva_glm_2 <- model_evaluation(glm_audit_2)

plotD3_lift(eva_glm, eva_glm_2)
```

---

plotD3\_prediction      *Plot Prediction vs Target, Observed or Variable Values in D3 with r2d3 package.*

---

**Description**

Function plotD3\_prediction plots predicted values observed or variable values in the model.

**Usage**

```
plotD3_prediction(
  object,
  ...,
  variable = "_y_",
  points = TRUE,
  smooth = FALSE,
  abline = FALSE,
```

```

    point_count = NULL,
    single_plot = TRUE,
    scale_plot = FALSE,
    background = FALSE
  )

plotD3Prediction(
  object,
  ...,
  variable = NULL,
  points = TRUE,
  smooth = FALSE,
  abline = FALSE,
  point_count = NULL,
  single_plot = TRUE,
  scale_plot = FALSE,
  background = FALSE
)

```

### Arguments

object	An object of class 'auditor_model_residual'.
...	Other modelAudit or modelResiduals objects to be plotted together.
variable	Name of variable to order residuals on a plot. If variable="_y_", the data is ordered by a vector of actual response (y parameter passed to the <a href="#">explain</a> function). If variable = "_y_hat_" the data on the plot will be ordered by predicted response. If variable = NULL, unordered observations are presented.
points	Logical, indicates whenever observations should be added as points. By default it's TRUE.
smooth	Logical, indicates whenever smoothed lines should be added. By default it's FALSE.
abline	Logical, indicates whenever function $y = x$ should be added. Works only with variable = NULL which is a default option.
point_count	Number of points to be plotted per model. Points will be chosen randomly. By default plot all of them.
single_plot	Logical, indicates whenever single or facets should be plotted. By default it's TRUE.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.
background	Logical, available only if single_plot = FALSE. Indicates whenever background plots should be plotted. By default it's FALSE.

### Value

a r2d3 object

**See Also**[plot\\_prediction](#)**Examples**

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plotD3_prediction(mr_lm, abline = TRUE)
plotD3_prediction(mr_lm, variable = "height", smooth = TRUE)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plotD3_prediction(mr_lm, mr_rf, variable = "weight", smooth = TRUE)

```

plotD3\_rec

*Regression Error Characteristic Curves (REC) in D3 with r2d3 package.*

**Description**

Error Characteristic curves are a generalization of ROC curves. On the x axis of the plot there is an error tolerance and on the y axis there is a percentage of observations predicted within the given tolerance.

**Usage**

```
plotD3_rec(object, ..., scale_plot = FALSE)
```

```
plotD3REC(object, ..., scale_plot = FALSE)
```

**Arguments**

object	An object of class 'auditor_model_residual' created with <a href="#">model_residual</a> function.
...	Other 'auditor_model_residual' objects to be plotted together.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.

**Details**

REC curve estimates the Cumulative Distribution Function (CDF) of the error  
Area Over the REC Curve (REC) is a biased estimate of the expected error

**Value**

a r2d3 object

**References**

Bi J., Bennett K.P. (2003). Regression error characteristic curves, in: Twentieth International Conference on Machine Learning (ICML-2003), Washington, DC.

**See Also**

[plot\\_rec](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)
plotD3_rec(mr_lm)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plotD3_rec(mr_lm, mr_rf)
```

---

plotD3\_residual

*Plot Residuals vs Observed, Fitted or Variable Values in D3 with r2d3 package.*

---

**Description**

Function plotD3\_residual plots residual values vs fitted, observed or variable values in the model.



**Usage**

```
plotD3_residual(
  object,
  ...,
  variable = "_y_",
  points = TRUE,
  smooth = FALSE,
  std_residuals = FALSE,
  nlabel = 0,
  point_count = NULL,
  single_plot = TRUE,
  scale_plot = FALSE,
  background = FALSE
)
```

```
plotD3Residual(
  object,
  ...,
  variable = NULL,
  points = TRUE,
  smooth = FALSE,
  std_residuals = FALSE,
  point_count = NULL,
  single_plot = TRUE,
  scale_plot = FALSE,
  background = FALSE
)
```

**Arguments**

object	An object of class 'auditor_model_residual' created with <code>model_residual</code> function.
...	Other 'auditor_model_residual' objects to be plotted together.
variable	Name of variable to order residuals on a plot. If <code>variable="_y_"</code> , the data is ordered by a vector of actual response (y parameter passed to the <code>explain</code> function). If <code>variable="_y_hat_"</code> the data on the plot will be ordered by predicted response. If <code>variable = NULL</code> , unordered observations are presented.
points	Logical, indicates whenever observations should be added as points. By default it's TRUE.
smooth	Logical, indicates whenever smoothed lines should be added. By default it's FALSE.
std_residuals	Logical, indicates whenever standardized residuals should be used. By default it's FALSE.
nlabel	Number of observations with the biggest residuals to be labeled.
point_count	Number of points to be plotted per model. Points will be chosen randomly. By default plot all of them.

single_plot	Logical, indicates whenever single or facets should be plotted. By default it's TRUE.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.
background	Logical, available only if single_plot = FALSE. Indicates whenever background plots should be plotted. By default it's FALSE.

**Value**

a r2d3 object

**See Also**

[plot\\_residual](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# use DALEX package to wrap up a model into explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plotD3_residual(mr_lm)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plotD3_residual(mr_lm, mr_rf)
```

---

plotD3\_roc

*Receiver Operating Characteristic (ROC) in D3 with r2d3 package.*

---

**Description**

Receiver Operating Characteristic Curve is a plot of the true positive rate (TPR) against the false positive rate (FPR) for the different thresholds. It is useful for measuring and comparing the accuracy of the classifiers.

**Usage**

```
plotD3_roc(object, ..., nlabel = NULL, scale_plot = FALSE)
```

**Arguments**

object	An object of class <code>auditor_model_evaluation</code> created with <code>model_evaluation</code> function.
...	Other <code>auditor_model_evaluation</code> objects to be plotted together.
nlabel	Number of cutoff points to show on the plot. Default is <code>NULL</code> .
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's <code>FALSE</code> .

**Value**

a `r2d3` object

**See Also**

[plot\\_roc](#)

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# use DALEX package to wrap up a model into explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
eva_glm <- model_evaluation(glm_audit)

# plot results
plot_roc(eva_glm)
plot(eva_glm)

#add second model
model_glm_2 <- glm(survived ~ .-age, family = binomial, data = titanic_imputed)
glm_audit_2 <- audit(model_glm_2,
                   data = titanic_imputed,
                   y = titanic_imputed$survived,
                   label = "glm2")
eva_glm_2 <- model_evaluation(glm_audit_2)

plotD3_roc(eva_glm, eva_glm_2)
```

---

plotD3_rroc	<i>Regression Receiver Operating Characteristic (RROC) in D3 with r2d3 package.</i>
-------------	---

---

### Description

The basic idea of the ROC curves for regression is to show model asymmetry. The RROC is a plot where on the x-axis we depict total over-estimation and on the y-axis total under-estimation.

### Usage

```
plotD3_rroc(object, ..., scale_plot = FALSE)
```

### Arguments

object	An object of class 'auditor_model_residual' created with <a href="#">model_residual</a> function.
...	Other 'auditor_model_residual' objects to be plotted together.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.

### Details

For RROC curves we use a shift, which is an equivalent to the threshold for ROC curves. For each observation we calculate new prediction:  $\hat{y}' = \hat{y} + s$  where  $s$  is the shift. Therefore, there are different error values for each shift:  $e_i = \hat{y}'_i - y_i$

Over-estimation is calculated as:  $OVER = \sum(e_i | e_i > 0)$ .

Under-estimation is calculated as:  $UNDER = \sum(e_i | e_i < 0)$ .

The shift equals 0 is represented by a dot.

The Area Over the RROC Curve (AOC) equals to the variance of the errors multiplied by  $frac{n^2$ .

### Value

a 'r2d3' object

### References

Hernández-Orallo, José. 2013. 'ROC Curves for Regression'. Pattern Recognition 46 (12): 3395–3411.

### See Also

[plotD3\\_rroc](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# use DALEX package to wrap up a model into explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plotD3_rroc(mr_lm)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plotD3_rroc(mr_lm, mr_rf)
```

---

plotD3\_scalelocation *Scale Location Plot in D3 with r2d3 package.*

---

**Description**

Function plotD3\_scalelocation plots square root of the absolute value of the residuals vs target, observed or variable values in the model. A vertical line corresponds to median.

**Usage**

```
plotD3_scalelocation(
  object,
  ...,
  variable = NULL,
  smooth = FALSE,
  peaks = FALSE,
  point_count = NULL,
  single_plot = TRUE,
  scale_plot = FALSE,
  background = FALSE
)

plotD3ScaleLocation(
  object,
  ...,
  variable = NULL,
```

```

smooth = FALSE,
peaks = FALSE,
point_count = NULL,
single_plot = TRUE,
scale_plot = FALSE,
background = FALSE
)

```

### Arguments

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
variable	Name of variable to order residuals on a plot. If <code>variable = "_y_"</code> , the data is ordered by a vector of actual response (y parameter passed to the <code>explain</code> function). If <code>variable = "_y_hat_"</code> the data on the plot will be ordered by predicted response. If <code>variable = NULL</code> , unordered observations are presented.
smooth	Logical, indicates whenever smoothed lines should be added. By default it's FALSE.
peaks	Logical, indicates whenever peak observations should be highlighted. By default it's FALSE.
point_count	Number of points to be plotted per model. Points will be chosen randomly. By default plot all of them.
single_plot	Logical, indicates whenever single or facets should be plotted. By default it's TRUE.
scale_plot	Logical, indicates whenever the plot should scale with height. By default it's FALSE.
background	Logical, available only if <code>single_plot = FALSE</code> . Indicates whenever background plots should be plotted. By default it's FALSE.

### Value

a r2d3 object

### See Also

[plot\\_scalelocation](#)

### Examples

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor

```

```

mr_lm <- model_residual(lm_audit)

# plot results
plotD3_scalelocation(mr_lm, peaks = TRUE)

```

---

plot\_acf

*Autocorrelation Function Plot*


---

## Description

Plot Autocorrelation Function of models' residuals.

## Usage

```

plot_acf(object, ..., variable = NULL, alpha = 0.95)

plotACF(object, ..., variable = NULL, alpha = 0.95)

```

## Arguments

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
variable	Name of variable to order residuals on a plot. If <code>variable="_y_"</code> , the data is ordered by a vector of actual response ( <code>y</code> parameter passed to the <code>explain</code> function). If <code>variable="_y_hat_"</code> the data on the plot will be ordered by predicted response. If <code>variable = NULL</code> , unordered observations are presented.
alpha	Confidence level of the interval.

## Value

A `ggplot` object.

## Examples

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot(mr_lm, type = "acf")

```

```
plot_acf(mr_lm)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_acf(mr_lm, mr_rf)
plot(mr_lm, mr_rf, type="acf")
```

---

plot\_auditor

*Model Diagnostic Plots*


---

## Description

This function provides several diagnostic plots for regression and classification models. Provide object created with one of auditor's computational functions, [model\\_residual](#), [model\\_cooksdistance](#), [model\\_evaluation](#), [model\\_performance](#), [model\\_evaluation](#).

## Usage

```
plot_auditor(x, ..., type = "residual", ask = TRUE, grid = TRUE)

## S3 method for class 'auditor_model_residual'
plot(x, ..., type = "residual", ask = TRUE, grid = TRUE)

## S3 method for class 'auditor_model_performance'
plot(x, ..., type = "residual", ask = TRUE, grid = TRUE)

## S3 method for class 'auditor_model_halfnormal'
plot(x, ..., type = "residual", ask = TRUE, grid = TRUE)

## S3 method for class 'auditor_model_evaluation'
plot(x, ..., type = "residual", ask = TRUE, grid = TRUE)

## S3 method for class 'auditor_model_cooksdistance'
plot(x, ..., type = "residual", ask = TRUE, grid = TRUE)
```

## Arguments

x object of class `auditor_model_residual` (created with [model\\_residual](#) function), `auditor_model_performance` (created with [model\\_performance](#) function), `auditor_model_evaluation` (created with [model\\_evaluation](#) function), `auditor_model_cooksdistance` (created with [model\\_cooksdistance](#) function), or `auditor_model_halfnormal` (created with [model\\_halfnormal](#) function).



...	other arguments dependent on the type of plot or additional objects of classes 'auditor_model_residual', 'auditor_model_performance', 'auditor_model_evaluation', 'auditor_model_evaluation', 'auditor_model_evaluation', 'auditor_model_evaluation'
type	the type of plot. Character or vector of characters. Possible values: 'acf', 'autocorrelation', 'cooksdistance', 'halfnormal', 'lift', 'prediction', 'residual_boxplot'. (for detailed description see corresponding functions in see also section).
ask	logical; if TRUE, the user is asked before each plot, see <a href="#">par(ask=)</a> .
grid	logical; if TRUE plots will be plotted on the grid.

**Value**

A ggplot object.

**See Also**

[plot\\_acf](#), [plot\\_autocorrelation](#), [plot\\_cooksdistance](#), [plot\\_halfnormal](#), [plot\\_residual\\_boxplot](#), [plot\\_lift](#), [plot\\_prediction](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot(mr_lm)
plot(mr_lm, type = "prediction")

hn_lm <- model_halfnormal(lm_audit)
plot(hn_lm)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)

mp_rf <- model_performance(rf_audit)
mp_lm <- model_performance(lm_audit)
plot(mp_lm, mp_rf)
```

---

plot\_autocorrelation *Autocorrelation of Residuals Plot*

---

**Description**

Plot of  $i$ -th residual vs  $i+1$ -th residual.

**Usage**

```
plot_autocorrelation(object, ..., variable = "_y_hat_", smooth = FALSE)
```

```
plotAutocorrelation(object, ..., variable, smooth = FALSE)
```

**Arguments**

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
variable	Name of variable to order residuals on a plot. If <code>variable="_y_"</code> , the data is ordered by a vector of actual response (y parameter passed to the <code>explain</code> function).
smooth	Logical, if TRUE smooth line will be added.

**Value**

A ggplot object.

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot_autocorrelation(mr_lm)
plot(mr_lm, type = "autocorrelation")
plot_autocorrelation(mr_lm, smooth = TRUE)
plot(mr_lm, type = "autocorrelation", smooth = TRUE)
```

---

plot\_cooksdistance      *Influence of Observations Plot*

---

**Description**

Plot of Cook's distances used for estimate the influence of an single observation.

**Usage**

```
plot_cooksdistance(object, ..., nlabel = 3)
```

```
plotCooksDistance(object, ..., nlabel = 3)
```

**Arguments**

object	An object of class <code>auditor_model_cooksdistance</code> created with <code>model_cooksdistance</code> function.
...	Other objects of class <code>auditor_model_cooksdistance</code> .
nlabel	Number of observations with the biggest Cook's distances to be labeled.

**Details**

Cook's distance is a tool for identifying observations that may negatively affect the model. They may be also used for indicating regions of the design space where it would be good to obtain more observations. Data points indicated by Cook's distances are worth checking for validity.

Cook's Distances are calculated by removing the *i*-th observation from the data and recalculating the model. It shows how much all the values in the model change when the *i*-th observation is removed.

For model classes other than `lm` and `glm` the distances are computed directly from the definition.

**Value**

A `ggplot` object.

**References**

Cook, R. Dennis (1977). "Detection of Influential Observations in Linear Regression". doi:10.2307/1268249.

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
library(auditor)
cd_lm <- model_cooksdistance(lm_audit)

# plot results
plot_cooksdistance(cd_lm)
plot(cd_lm, type = "cooksdistance")
```

---

plot_correlation	<i>Correlation of Model's Residuals Plot</i>
------------------	--

---

### Description

Matrix of plots. Left-down triangle consists of plots of fitted values (alternatively residuals), on the diagonal there are density plots of fitted values (alternatively residuals), in the right-top triangle there are correlations between fitted values (alternatively residuals).

### Usage

```
plot_correlation(object, ..., values = "fit")  
  
plotModelCorrelation(object, ..., values = "fit")
```

### Arguments

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
values	"fit" for model fitted values or "res" for residual values.

### Value

Invisibly returns a `gtable` object.

### Examples

```
dragons <- DALEX::dragons[1:100, ]  
  
# fit a model  
model_lm <- lm(life_length ~ ., data = dragons)  
  
# use DALEX package to wrap up a model into explainer  
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)  
  
# validate a model with auditor  
mr_lm <- model_residual(lm_audit)  
  
library(randomForest)  
model_rf <- randomForest(life_length~., data = dragons)  
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)  
mr_rf <- model_residual(rf_audit)  
  
# plot results  
plot_correlation(mr_lm, mr_rf)  
plot(mr_lm, mr_rf, type = "correlation")
```

---

plot_halfnormal	<i>Half-Normal plot</i>
-----------------	-------------------------

---

### Description

The half-normal plot is one of the tools designed to evaluate the goodness of fit of a statistical models. It is a graphical method for comparing two probability distributions by plotting their quantiles against each other. Points on the plot correspond to ordered absolute values of model diagnostic (i.e. standardized residuals) plotted against theoretical order statistics from a half-normal distribution.

### Usage

```
plot_halfnormal(object, ..., quantiles = FALSE, sim = 99)
```

```
plotHalfNormal(object, ..., quantiles = FALSE, sim = 99)
```

### Arguments

object	An object of class <code>auditor_model_halfnormal</code> created with <a href="#">model_halfnormal</a> function.
...	Other <code>auditor_model_halfnormal</code> objects.
quantiles	If TRUE values on axis are on quantile scale.
sim	Number of residuals to simulate.

### Value

A ggplot object.

### See Also

[model\\_halfnormal](#)  
[score\\_halfnormal](#)

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
hn_lm <- model_halfnormal(lm_audit)

# plot results
plot_halfnormal(hn_lm)
```

```
plot(hn_lm)
```

---

plot\_lift

*LIFT Chart*

---

### Description

LIFT is a plot of the rate of positive prediction against true positive rate for the different thresholds. It is useful for measuring and comparing the accuracy of the classifiers.

### Usage

```
plot_lift(object, ..., zeros = TRUE)
```

```
plotLIFT(object, ...)
```

### Arguments

object	An object of class <code>auditor_model_evaluation</code> created with <code>model_evaluation</code> function.
...	Other <code>auditor_model_evaluation</code> objects to be plotted together.
zeros	Logical. It makes the lines start from the $(0,0)$ point. By default it's TRUE.

### Value

A ggplot object.

### See Also

[model\\_evaluation](#)

### Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
eva_glm <- model_evaluation(glm_audit)

# plot results
plot_lift(eva_glm)
plot(eva_glm, type = "lift")
```

```

model_glm_2 <- glm(survived ~ .-age, family = binomial, data = titanic_imputed)
glm_audit_2 <- audit(model_glm_2,
                    data = titanic_imputed,
                    y = titanic_imputed$survived,
                    label = "glm2")
eva_glm_2 <- model_evaluation(glm_audit_2)

plot_lift(eva_glm, eva_glm_2)
plot(eva_glm, eva_glm_2, type = "lift")

```

---

plot\_pca

*Principal Component Analysis of models*


---

### Description

Principal Component Analysis of models residuals. PCA can be used to assess the similarity of the models.

### Usage

```

plot_pca(object, ..., scale = TRUE, arrow_size = 2)

plotModelPCA(object, ..., scale = TRUE)

```

### Arguments

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
scale	A logical value indicating whether the models residuals should be scaled before the analysis.
arrow_size	Width of the arrows.

### Value

A ggplot object.

### Examples

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

```

```
# validate a model with auditor
mr_lm <- model_residual(lm_audit)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)

# plot results
plot_pca(mr_lm, mr_rf)
```

---

plot\_prc

*Precision-Recall Curve (PRC)*


---

### Description

Precision-Recall Curve summarize the trade-off between the true positive rate and the positive predictive value for a model. It is useful for measuring performance and comparing classifiers.

Receiver Operating Characteristic Curve is a plot of the true positive rate (TPR) against the false positive rate (FPR) for the different thresholds. It is useful for measuring and comparing the accuracy of the classifiers.

### Usage

```
plot_prc(object, ..., nlabel = NULL)
```

```
plot_roc(object, ..., nlabel = NULL)
```

```
plotROC(object, ..., nlabel = NULL)
```

### Arguments

object	An object of class <code>auditor_model_evaluation</code> created with <code>model_evaluation</code> function.
...	Other <code>auditor_model_evaluation</code> objects to be plotted together.
nlabel	Number of cutoff points to show on the plot. Default is <code>NULL</code> .

### Value

A `ggplot` object.

A `ggplot` object.

### See Also

[plot\\_rrroc](#), [plot\\_rec](#)



**Examples**

```
library(DALEX)

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
eva_glm <- model_evaluation(glm_audit)

# plot results
plot_prc(eva_glm)
plot(eva_glm)

#add second model
model_glm_2 <- glm(survived ~ .-age, family = binomial, data = titanic_imputed)
glm_audit_2 <- audit(model_glm_2,
                   data = titanic_imputed,
                   y = titanic_imputed$survived,
                   label = "glm2")
eva_glm_2 <- model_evaluation(glm_audit_2)

plot_prc(eva_glm, eva_glm_2)
plot(eva_glm, eva_glm_2)

data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# use DALEX package to wrap up a model into explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
eva_glm <- model_evaluation(glm_audit)

# plot results
plot_roc(eva_glm)
plot(eva_glm)

#add second model
model_glm_2 <- glm(survived ~ .-age, family = binomial, data = titanic_imputed)
glm_audit_2 <- audit(model_glm_2,
                   data = titanic_imputed,
                   y = titanic_imputed$survived,
                   label = "glm2")
eva_glm_2 <- model_evaluation(glm_audit_2)
```

```
plot_roc(eva_glm, eva_glm_2)
plot(eva_glm, eva_glm_2)
```

---

plot_prediction	<i>Predicted response vs Observed or Variable Values</i>
-----------------	--

---

### Description

Plot of predicted response vs observed or variable Values.

### Usage

```
plot_prediction(object, ..., variable = "_y_", smooth = FALSE, abline = FALSE)
plotPrediction(object, ..., variable = NULL, smooth = FALSE, abline = FALSE)
```

### Arguments

object	An object of class auditor_model_residual.
...	Other auditor_model_residual objects to be plotted together.
variable	Name of variable to order residuals on a plot. If variable="y", the data is ordered by a vector of actual response (y parameter passed to the <a href="#">explain</a> function). If variable = "_y_hat_" the data on the plot will be ordered by predicted response. If variable = NULL, unordered observations are presented.
smooth	Logical, indicates whenever smooth line should be added.
abline	Logical, indicates whenever function $y = x$ should be added. Works only with variable = "_y_" (which is a default option) or when variable equals actual response variable.

### Value

A ggplot2 object.

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)
```

```

# plot results
plot_prediction(mr_lm, abline = TRUE)
plot_prediction(mr_lm, variable = "height", smooth = TRUE)
plot(mr_lm, type = "prediction", abline = TRUE)

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_prediction(mr_lm, mr_rf, variable = "height", smooth = TRUE)

```

---

plot\_radar

*Model Ranking Plot*


---

### Description

Radar plot with model score. score are scaled to  $[0, 1]$ , each score is inversed and divided by maximum score value.

### Usage

```

plot_radar(object, ..., verbose = TRUE)

plotModelRanking(object, ..., verbose = TRUE)

```

### Arguments

object	An object of class <code>auditor_model_performance</code> created with <code>model_performance</code> function.
...	Other <code>auditor_model_performance</code> objects to be plotted together.
verbose	Logical, indicates whether values of scores should be printed.

### Value

A ggplot object.

### Examples

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mp_lm <- model_performance(lm_audit)

```

```

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mp_rf <- model_performance(rf_audit)

# plot results
plot_radar(mp_lm, mp_rf)

```

---

plot\_rec

*Regression Error Characteristic Curves (REC)*


---

### Description

Error Characteristic curves are a generalization of ROC curves. On the x axis of the plot there is an error tolerance and on the y axis there is a percentage of observations predicted within the given tolerance.

### Usage

```
plot_rec(object, ...)
```

```
plotREC(object, ...)
```

### Arguments

`object` An object of class `auditor_model_residual` created with `model_residual` function.

`...` Other `auditor_model_residual` objects to be plotted together.

### Details

REC curve estimates the Cumulative Distribution Function (CDF) of the error

Area Over the REC Curve (REC) is a biased estimate of the expected error

### Value

A `ggplot` object.

### References

Bi J., Bennett K.P. (2003). Regression error characteristic curves, in: Twentieth International Conference on Machine Learning (ICML-2003), Washington, DC.

### See Also

[plot\\_roc](#), [plot\\_rroc](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# use DALEX package to wrap up a model into explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)
plot_rec(mr_lm)
plot(mr_lm, type = "rec")

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_rec(mr_lm, mr_rf)
plot(mr_lm, mr_rf, type = "rec")
```

---

plot\_residual

*Plot Residuals vs Observed, Fitted or Variable Values*

---

**Description**

A plot of residuals against fitted values, observed values or any variable.

**Usage**

```
plot_residual(  
  object,  
  ...,  
  variable = "_y_",  
  smooth = FALSE,  
  std_residuals = FALSE,  
  nlabel = 0  
)
```

```
plotResidual(  
  object,  
  ...,  
  variable = NULL,  
  smooth = FALSE,  
  std_residuals = FALSE,  
  nlabel = 0  
)
```

**Arguments**

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
variable	Name of variable to order residuals on a plot. If <code>variable="_y_"</code> , the data is ordered by a vector of actual response ( <code>y</code> parameter passed to the <code>explain</code> function). If <code>variable="_y_hat_"</code> the data on the plot will be ordered by predicted response. If <code>variable = NULL</code> , unordered observations are presented.
smooth	Logical, indicates whenever smoothed lines should be added. By default it's <code>FALSE</code> .
std_residuals	Logical, indicates whenever standardized residuals should be used.
nlabel	Number of observations with the biggest absolute values of residuals to be labeled.

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot_residual(mr_lm)
plot(mr_lm, type = "residual")

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_residual(mr_lm, mr_rf)
plot(mr_rf, mr_rf, type = "residual")
```

---

plot\_residual\_boxplot *Plot Boxplots of Residuals*

---

**Description**

A boxplot of residuals.

**Usage**

```
plot_residual_boxplot(object, ...)
```

```
plotResidualBoxplot(object, ...)
```

**Arguments**

**object** An object of class `auditor_model_residual` created with `model_residual` function.

**...** Other `auditor_model_residual` objects to be plotted together.

**Value**

A ggplot object.

**See Also**

[plot\\_residual](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot_residual_boxplot(mr_lm)
plot(mr_lm, type = "residual_boxplot")

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_residual_boxplot(mr_lm, mr_rf)
plot(mr_lm, mr_rf)
```

---

 plot\_residual\_density *Residual Density Plot*


---

**Description**

Density of model residuals.

**Usage**

```
plot_residual_density(object, ..., variable = "", show_rugs = TRUE)
plotResidualDensity(object, ..., variable = NULL)
```

**Arguments**

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
variable	Split plot by variable's factor level or median. If <code>variable = "_y_"</code> , the plot will be splitted by actual response (y parameter passed to the <code>explain</code> function). If <code>variable = "_y_hat_"</code> the plot will be splitted by predicted response. If <code>variable = NULL</code> , the plot will be splitted by observation index. If <code>variable = ""</code> plot is not splitted (default option).
show_rugs	Adds rugs layer to the plot. By default it's TRUE

**Value**

A ggplot object.

**See Also**

[plot\\_residual](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot_residual_density(mr_lm)
plot(mr_lm, type = "residual_density")
```



```

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_residual_density(mr_lm, mr_rf)
plot(mr_lm, mr_rf, type = "residual_density")

```

---

plot\_rroc

*Regression Receiver Operating Characteristic (RROC)*


---

### Description

The basic idea of the ROC curves for regression is to show model asymmetry. The RROC is a plot where on the x-axis we depict total over-estimation and on the y-axis total under-estimation.

### Usage

```
plot_rroc(object, ...)
```

```
plotRROC(object, ...)
```

### Arguments

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.

### Details

For RROC curves we use a shift, which is an equivalent to the threshold for ROC curves. For each observation we calculate new prediction:  $\hat{y}' = \hat{y} + s$  where  $s$  is the shift. Therefore, there are different error values for each shift:  $e_i = \hat{y}'_i - y_i$

Over-estimation is calculated as:  $OVER = \sum(e_i | e_i > 0)$ .

Under-estimation is calculated as:  $UNDER = \sum(e_i | e_i < 0)$ .

The shift equals 0 is represented by a dot.

The Area Over the RROC Curve (AOC) equals to the variance of the errors multiplied by  $frac{n^2}{2}$ .

### Value

A ggplot object.

### References

Hernández-Orallo, José. 2013. 'ROC Curves for Regression'. *Pattern Recognition* 46 (12): 3395–3411.

**See Also**

[plot\\_roc](#), [plot\\_rec](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot_rroc(mr_lm)
plot(mr_lm, type = "rroc")

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_rroc(mr_lm, mr_rf)
plot(mr_lm, mr_rf, type="rroc")
```

---

plot\_scalelocation      *Scale location plot*

---

**Description**

Variable values vs square root of the absolute value of the residuals. A vertical line corresponds to median.

**Usage**

```
plot_scalelocation(
  object,
  ...,
  variable = "_y_",
  smooth = FALSE,
  peaks = FALSE
)

plotScaleLocation(object, ..., variable = NULL, smooth = FALSE, peaks = FALSE)
```

**Arguments**

object	An object of class <code>auditor_model_residual</code> created with <code>model_residual</code> function.
...	Other <code>auditor_model_residual</code> objects to be plotted together.
variable	Name of variable to order residuals on a plot. If <code>variable="_y_"</code> , the data is ordered by a vector of actual response (y parameter passed to the <code>explain</code> function). If <code>variable = "_y_hat_"</code> the data on the plot will be ordered by predicted response. If <code>variable = NULL</code> , unordered observations are presented.
smooth	Logical, indicates whenever smoothed lines should be added. By default it's FALSE.
peaks	A logical value. If TRUE peaks are marked on plot by black dots.

**Value**

A ggplot object.

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)

# plot results
plot_scalelocation(mr_lm)
plot(mr_lm, type = "scalelocation")
```

---

plot\_tsecdf

*Two-sided Cumulative Distribution Function*

---

**Description**

Cumulative Distribution Function for positive and negative residuals.

**Usage**

```
plot_tsecdf(
  object,
  ...,
  scale_error = TRUE,
  outliers = NA,
```

```

    residuals = TRUE,
    reverse_y = FALSE
  )

plotTwoSidedECDF(
  object,
  ...,
  scale_error = TRUE,
  outliers = NA,
  residuals = TRUE,
  reverse_y = FALSE
)

```

### Arguments

object	An object of class 'auditor_model_residual' created with <code>model_residual</code> function.
...	Other modelAudit objects to be plotted together.
scale_error	A logical value indicating whether ECDF should be scaled by proportions of positive and negative proportions.
outliers	Number of outliers to be marked.
residuals	A logical value indicating whether residuals should be marked.
reverse_y	A logical value indicating whether values on y axis should be reversed.

### Value

A ggplot object.

### Examples

```

dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# validate a model with auditor
mr_lm <- model_residual(lm_audit)
plot_tsecdf(mr_lm)
plot(mr_lm, type="tsecdf")

library(randomForest)
model_rf <- randomForest(life_length~., data = dragons)
rf_audit <- audit(model_rf, data = dragons, y = dragons$life_length)
mr_rf <- model_residual(rf_audit)
plot_tsecdf(mr_lm, mr_rf, reverse_y = TRUE)

```

---

```
print.auditor_model_cooksdistance
      Prints Model Cook's Distances Summary
```

---

**Description**

Prints Model Cook's Distances Summary

**Usage**

```
## S3 method for class 'auditor_model_cooksdistance'
print(x, ...)
```

**Arguments**

x	an object auditor_model_cooksdistance created with <a href="#">model_cooksdistance</a> function.
...	other parameters

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
model_cooksdistance(lm_audit)
```

---

```
print.auditor_model_evaluation
      Prints Model Evaluation Summary
```

---

**Description**

Prints Model Evaluation Summary

**Usage**

```
## S3 method for class 'auditor_model_evaluation'
print(x, ...)
```

**Arguments**

x                    an object `auditor_model_evaluation` created with `model_evaluation` function.  
...                   other parameters

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data= titanic_imputed,
                  y = titanic_imputed$survived)

# validate a model with auditor
model_evaluation(glm_audit)
```

---

```
print.auditor_model_halfnormal
```

*Prints Model Halfnormal Summary*

---

**Description**

Prints Model Halfnormal Summary

**Usage**

```
## S3 method for class 'auditor_model_halfnormal'
print(x, ...)
```

**Arguments**

x                    an object `auditor_model_halfnormal` created with `model_halfnormal` function.  
...                   other parameters

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
```

```
      data = titanic_imputed,  
      y = titanic_imputed$survived)  
  
# validate a model with auditor  
model_halfnormal(glm_audit)
```

---

```
print.auditor_model_performance  
      Prints Model Performance Summary
```

---

### **Description**

Prints Model Performance Summary

### **Usage**

```
## S3 method for class 'auditor_model_performance'  
print(x, ...)
```

### **Arguments**

x	an object auditor_model_performance created with <a href="#">model_performance</a> function.
...	other parameters

### **Examples**

```
data(titanic_imputed, package = "DALEX")  
  
# fit a model  
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)  
  
glm_audit <- audit(model_glm,  
                  data = titanic_imputed,  
                  y = titanic_imputed$survived)  
  
# validate a model with auditor  
model_performance(glm_audit)
```

```
print.auditor_model_residual  
Prints Model Residual Summary
```

---

**Description**

Prints Model Residual Summary

**Usage**

```
## S3 method for class 'auditor_model_residual'  
print(x, ...)
```

**Arguments**

x                    an object auditor\_model\_residual created with [model\\_residual](#) function.  
...                   other parameters

**Examples**

```
data(titanic_imputed, package = "DALEX")  
  
# fit a model  
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)  
  
glm_audit <- audit(model_glm,  
                  data = titanic_imputed,  
                  y = titanic_imputed$survived)  
  
# validate a model with auditor  
model_residual(glm_audit)
```

---

```
print.auditor_score    Prints of Models Scores
```

---

**Description**

Prints of Models Scores

**Usage**

```
## S3 method for class 'auditor_score'  
print(x, ...)
```



**Arguments**

x                    an object `auditor_score` created with `score` function.  
 ...                   other parameters

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                   data = titanic_imputed,
                   y = titanic_imputed$survived)

# calculate score
score(glm_audit, type = "auc")
```

---

score                                    *Model Scores computations*

---

**Description**

This function provides several scores for model validation and performance assessment. Scores can be also used to compare models.

**Usage**

```
score(object, type = "mse", data = NULL, ...)
```

**Arguments**

object                An object of class `explainer` created with function `explain` from the DALEX package.

type                   The score to be calculated. Possible values: `acc`, `auc`, `cookdistance`, `dw`, `f1`, `gini`, `halfnormal`, `mae`, `mse`, `peak`, `precision`, `r2`, `rec`, `recall`, `rmse`, `rroc`, `runs`, `specificity`, `one_minus_acc`, `one_minus_auc`, `one_minus_f1`, `one_minus_gini`, `one_minus_precision`, `one_minus_recall`, `one_minus_specificity` (for detailed description see functions in `see also` section).

data                   New data that will be used to calculate the score. Pass `NULL` if you want to use data from object.

...                    Other arguments dependent on the type of score.

**Value**

An object of class `auditor_score`, except Cooks distance, where numeric vector is returned.

**See Also**

score\_acc, score\_auc, score\_cooksdistance, score\_dw, score\_f1, score\_gini, score\_halfnormal, score\_mae, score\_mse, score\_peak, score\_precision, score\_r2, score\_rec, score\_recall, score\_rmse, score\_rroc, score\_runs, score\_specificity, score\_one\_minus\_acc, score\_one\_minus\_auc, score\_one\_minus\_f1, score\_one\_minus\_gini, score\_one\_minus\_precision, score\_one\_minus\_recall, score\_one\_minus\_specificity

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score(lm_audit, type = 'mae')
```

---

score\_acc

*Accuracy*

---

**Description**

Accuracy

**Usage**

```
score_acc(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $y_{\hat{}}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class `auditor_score`.

**Examples**

```

data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                   data = titanic_imputed,
                   y = titanic_imputed$survived)

# calculate score
score_acc(glm_audit)

```

---

score_auc	<i>Area Under ROC Curve (AUC)</i>
-----------	-----------------------------------

---

**Description**

Area Under Curve (AUC) for Receiver Operating Characteristic.

**Usage**

```

score_auc(object, data = NULL, y = NULL, ...)

scoreROC(object)

```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**See Also**

[plot\\_roc](#)

**Examples**

```

data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_auc(glm_audit)

```

---

score_auprc	<i>Area under precision-recall curve</i>
-------------	--

---

**Description**

Area under precision-recall curve

**Usage**

```
score_auprc(object, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**Examples**

```

data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,

```

```

y = titanic_imputed$survived)

# calculate score
score_auprc(glm_audit)

```

---

score\_cooksdistance    *Score based on Cooks Distance*

---

### Description

Cook's distance are used for estimate of the influence of an single observation.

### Usage

```

score_cooksdistance(object, verbose = TRUE, ...)

scoreCooksDistance(object, verbose = TRUE)

```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
verbose	If TRUE progress is printed.
...	Other arguments dependent on the type of score.

### Details

Cook's distance is a tool for identifying observations that may negatively affect the model. They may be also used for indicating regions of the design space where it would be good to obtain more observations. Data points indicated by Cook's distances are worth checking for validity.

Cook's Distances are calculated by removing the  $i$ -th observation from the data and recalculating the model. It shows how much all the values in the model change when the  $i$ -th observation is removed.

Models of classes other than `lm` and `glm` the distances are computed directly from the definition, so this may take a while.

### Value

A vector of Cook's distances for each observation.  
numeric vector

### See Also

[score](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_cooksdistance(lm_audit)
```

---

score_dw	<i>Durbin-Watson Score</i>
----------	----------------------------

---

**Description**

Score based on Durbin-Watson test statistic. The score value is helpful in comparing models. It is worth pointing out that results of tests like p-value makes sense only when the test assumptions are satisfied. Otherwise test statistic may be considered as a score.

**Usage**

```
score_dw(object, variable = NULL, data = NULL, y = NULL, ...)

scoreDW(object, variable = NULL)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
variable	Name of model variable to order residuals.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class `auditor_score`.

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_dw(lm_audit)
```

---

score_f1	<i>F1 Score</i>
----------	-----------------

---

**Description**

F1 Score

**Usage**

```
score_f1(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $\hat{y}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
```

```

y = titanic_imputed$survived)

# calculate score
score_f1(glm_audit)

```

---

score_gini	<i>Gini Coefficient</i>
------------	-------------------------

---

### Description

The Gini coefficient measures the inequality among values of a frequency distribution. A Gini coefficient equals 0 means perfect equality, where all values are the same. A Gini coefficient equals 100

### Usage

```
score_gini(object, data = NULL, y = NULL, ...)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

### Value

An object of class auditor\_score.

### See Also

[plot\\_roc](#)

### Examples

```

library(DALEX)

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
exp_glm <- explain(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

```



```
# calculate score
score_gini(exp_glm)
```

---

score_halfnormal	<i>Half-Normal Score</i>
------------------	--------------------------

---

### Description

Score is approximately:  $\sum \#[res_i \leq simres_{i,j}] - n$  with the distinction that each element of sum is also scaled to take values from [0,1].

$res_i$  is a residual for i-th observation,  $simres_{i,j}$  is the residual of j-th simulation for i-th observation, and  $n$  is the number of simulations for each observation. Scores are calculated on the basis of simulated data, so they may differ between function calls.

### Usage

```
score_halfnormal(object, ...)
scoreHalfNormal(object, ...)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
...	...

### Value

An object of class auditor\_score.

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_halfnormal(lm_audit)
```

---

score_mae	<i>Mean Absolute Error</i>
-----------	----------------------------

---

**Description**

Mean Absolute Error.

**Usage**

```
score_mae(object, data = NULL, y = NULL, ...)
```

```
scoreMAE(object)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**See Also**

[score](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_mae(lm_audit)
```

---

score_mse	<i>Mean Square Error</i>
-----------	--------------------------

---

**Description**

Mean Square Error.

**Usage**

```
score_mse(object, data = NULL, y = NULL, ...)
```

```
scoreMSE(object)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**See Also**

[score](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_mse(lm_audit)
```

---

score\_one\_minus\_acc     *One minus accuracy*

---

## Description

One minus accuracy

## Usage

```
score_one_minus_acc(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

## Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

## Value

An object of class auditor\_score.

## Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_acc(glm_audit)
```

---

score_one_minus_auc	<i>One minus Area Under ROC Curve (AUC)</i>
---------------------	---

---

### Description

One minus Area Under Curve (AUC) for Receiver Operating Characteristic.

### Usage

```
score_one_minus_auc(object, data = NULL, y = NULL, ...)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

### Value

An object of class auditor\_score.

### Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_auc(glm_audit)
```

---

score\_one\_minusauprc *One Minus area under precision-recall curve*

---

## Description

One Minus area under precision-recall curve

## Usage

```
score_one_minusauprc(object, data = NULL, y = NULL, ...)
```

## Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

## Value

An object of class auditor\_score.

## Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minusauprc(glm_audit)
```

---

score_one_minus_f1	<i>One Minus F1 Score</i>
--------------------	---------------------------

---

**Description**

One Minus F1 Score

**Usage**

```
score_one_minus_f1(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $\hat{y}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_f1(glm_audit)
```

---

score\_one\_minus\_gini *One minus Gini Coefficient*

---

### Description

One minus Gini COefficient 100 0 expresses maximal inequality of values.

### Usage

```
score_one_minus_gini(object, data = NULL, y = NULL, ...)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

### Value

An object of class auditor\_score.

### Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_gini(glm_audit)
```



---

`score_one_minus_precision`*One Minus Precision*

---

**Description**

One Minus Precision

**Usage**`score_one_minus_precision(object, cutoff = 0.5, data = NULL, y = NULL, ...)`**Arguments**

<code>object</code>	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
<code>cutoff</code>	Threshold value, which divides model predicted values ( <code>y_hat</code> ) to calculate confusion matrix. By default it's 0.5.
<code>data</code>	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
<code>y</code>	New y parameter will be used to calculate score.
<code>...</code>	Other arguments dependent on the type of score.

**Value**An object of class `auditor_score`.**Examples**

```
library(DALEX)

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
exp_glm <- explain(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_precision(exp_glm)
```

---

`score_one_minus_recall`*One minus recall*

---

**Description**

One minus recall

**Usage**

```
score_one_minus_recall(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

<code>object</code>	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
<code>cutoff</code>	Threshold value, which divides model predicted values ( <code>y_hat</code> ) to calculate confusion matrix. By default it's 0.5.
<code>data</code>	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
<code>y</code>	New y parameter will be used to calculate score.
<code>...</code>	Other arguments dependent on the type of score.

**Value**

An object of class `auditor_score`.

**Examples**

```
library(DALEX)

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
exp_glm <- explain(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_recall(exp_glm)
```

---

score\_one\_minus\_specificity  
*One minus specificity*

---

## Description

One minus specificity

## Usage

```
score_one_minus_specificity(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

## Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $y_{\hat{}}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

## Value

An object of class auditor\_score.

## Examples

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_one_minus_specificity(glm_audit)
```

---

score_peak	<i>Peak Score</i>
------------	-------------------

---

### Description

This score is calculated on the basis of Peak test, which is used for checking for homoscedasticity of residuals in regression analyses.

### Usage

```
score_peak(object, variable = NULL, data = NULL, y = NULL, ...)  
  
scorePeak(object)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
variable	Name of model variable to order residuals.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

### Value

An object of class auditor\_score.

### Examples

```
dragons <- DALEX::dragons[1:100, ]  
  
# fit a model  
model_lm <- lm(life_length ~ ., data = dragons)  
  
# create an explainer  
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)  
  
# calculate score  
score_peak(lm_audit)
```

---

score_precision	<i>Precision</i>
-----------------	------------------

---

**Description**

Precision

**Usage**

```
score_precision(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $y_{\hat{}}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_precision(glm_audit)
```

---

`score_r2`*R-squared*

---

### Description

The R2 is the coefficient of determination, An R2 coefficient equals 0 means that model explains none of the variability of the response. An R2 coefficient equals 1 means that model explains all the variability of the response.

### Usage

```
score_r2(object, data = NULL, y = NULL, ...)
```

### Arguments

<code>object</code>	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
<code>data</code>	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
<code>y</code>	New y parameter will be used to calculate score.
<code>...</code>	Other arguments dependent on the type of score.

### Value

An object of class `auditor_score`.

### See Also

[score](#)

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# use DALEX package to wrap up a model into explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score with auditor
score_r2(lm_audit)
```

---

score_rec	<i>Area Over the Curve for REC Curves</i>
-----------	---

---

**Description**

The area over the Regression Error Characteristic curve is a measure of the expected error for the regression model.

**Usage**

```
score_rec(object, data = NULL, y = NULL, ...)
scoreREC(object)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**References**

J. Bi, and K. P. Bennet, "Regression error characteristic curves," in Proc. 20th Int. Conf. Machine Learning, Washington DC, 2003, pp. 43-50

**See Also**

[plot\\_rec](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
lm_model <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(lm_model, data = dragons, y = dragons$life_length)

# calculate score
score_rec(lm_audit)
```

---

score_recall	<i>Recall</i>
--------------	---------------

---

**Description**

Recall

**Usage**

```
score_recall(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $\hat{y}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class `auditor_score`.

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

# create an explainer
glm_audit <- audit(model_glm,
                  data = titanic_imputed,
                  y = titanic_imputed$survived)

# calculate score
score_recall(glm_audit)
```



---

score_rmse	<i>Root Mean Square Error</i>
------------	-------------------------------

---

**Description**

Root Mean Square Error.

**Usage**

```
score_rmse(object, data = NULL, y = NULL, ...)  
  
scoreRMSE(object)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**See Also**

[score](#)

**Examples**

```
dragons <- DALEX::dragons[1:100, ]  
  
# fit a model  
model_lm <- lm(life_length ~ ., data = dragons)  
  
# create an explainer  
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)  
  
# calculate score  
score_rmse(lm_audit)
```

---

score_rroc	<i>Area Over the Curve for RROC Curves</i>
------------	--

---

### Description

The area over the Regression Receiver Operating Characteristic.

### Usage

```
score_rroc(object, data = NULL, y = NULL, ...)
```

```
scoreRROC(object)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

### Value

An object of class `auditor_score`.

### References

Hernández-Orallo, José. 2013. 'ROC Curves for Regression'. *Pattern Recognition* 46 (12): 3395–3411.

### See Also

[plot\\_rroc](#)

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_rroc(lm_audit)
```

---

 score\_runs

*Runs Score*


---

### Description

Score based on Runs test statistic. Note that this test is not very strong. It utilizes only signs of the residuals. The score value is helpful in comparing models. It is worth pointing out that results of tests like p-value makes sense only when the test assumptions are satisfied. Otherwise test statistic may be considered as a score.

### Usage

```
score_runs(object, variable = NULL, data = NULL, y = NULL, ...)
```

```
scoreRuns(object, variable = NULL)
```

### Arguments

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
variable	name of model variable to order residuals.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

### Value

An object of class `auditor_score`.

### Examples

```
dragons <- DALEX::dragons[1:100, ]

# fit a model
model_lm <- lm(life_length ~ ., data = dragons)

# create an explainer
lm_audit <- audit(model_lm, data = dragons, y = dragons$life_length)

# calculate score
score_runs(lm_audit)
```

---

score\_specificity      *Specificity*

---

**Description**

Specificity

**Usage**

```
score_specificity(object, cutoff = 0.5, data = NULL, y = NULL, ...)
```

**Arguments**

object	An object of class explainer created with function <a href="#">explain</a> from the DALEX package.
cutoff	Threshold value, which divides model predicted values ( $y_{\hat{}}$ ) to calculate confusion matrix. By default it's 0.5.
data	New data that will be used to calculate the score. Pass NULL if you want to use data from object.
y	New y parameter will be used to calculate score.
...	Other arguments dependent on the type of score.

**Value**

An object of class auditor\_score.

**Examples**

```
data(titanic_imputed, package = "DALEX")

# fit a model
model_glm <- glm(survived ~ ., family = binomial, data = titanic_imputed)

exp_glm <- audit(model_glm,
                data = titanic_imputed,
                y = titanic_imputed$survived)

# calculate score
score_specificity(exp_glm)
```

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