

# Package ‘ezcox’

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**Type** Package

**Title** Easily Process a Batch of Cox Models

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**Maintainer** Shixiang Wang <w\_shixiang@163.com>

**Description** A tool to operate a batch of univariate or multivariate Cox models and return tidy result.

**License** GPL-3

**URL** <https://github.com/ShixiangWang/ezcox>

**BugReports** <https://github.com/ShixiangWang/ezcox/issues>

**Depends** R (>= 3.5)

**Imports** dplyr (>= 0.8.3), forestmodel, ggplot2, magrittr (>= 1.5), methods, purrr (>= 0.3.2), rlang (>= 0.1.2), scales, survival, tibble, utf8, utils

**Suggests** covr (>= 3.2.1), furrr, future, knitr, prettydoc, rmarkdown, roxygen2 (>= 6.1.1), testthat (>= 2.1.0)

**VignetteBuilder** knitr

**Encoding** UTF-8

**RoxygenNote** 7.2.3

**NeedsCompilation** no

**Author** Shixiang Wang [aut, cre] (ORCID:  
<<https://orcid.org/0000-0001-9855-7357>>)

**Repository** CRAN

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|                 |                                         |
|-----------------|-----------------------------------------|
| clean_model_dir | <i>Clean ezcox Model File Directory</i> |
|-----------------|-----------------------------------------|

---

**Description**

Clean ezcox Model File Directory

**Usage**

```
clean_model_dir(model_dir = file.path(tempdir(), "ezcox"))
```

**Arguments**

model\_dir      a path for storing model results.

**Value**

nothing

**Examples**

```
clean_model_dir()
```

---

|       |                                       |
|-------|---------------------------------------|
| ezcox | <i>Run Cox Analysis in Batch Mode</i> |
|-------|---------------------------------------|

---

**Description**

Run Cox Analysis in Batch Mode

**Usage**

```
ezcox(
  data,
  covariates,
  controls = NULL,
  time = "time",
  status = "status",
  global_method = c("likelihood", "wald", "logrank"),
  keep_models = FALSE,
  return_models = FALSE,
  model_dir = file.path(tempdir(), "ezcox"),
  verbose = TRUE,
  ...
)
```

**Arguments**

|                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
|----------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>data</code>          | a <code>data.frame</code> containing variables, time and os status.                                                                                                                                                                                                                                                                                                                                                                                                                               |
| <code>covariates</code>    | column names specifying variables.                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| <code>controls</code>      | column names specifying controls. The names with pattern <code>"*:l()"</code> will be treated as interaction/combination term, please make sure all column names in <code>data</code> are valid R variable names.                                                                                                                                                                                                                                                                                 |
| <code>time</code>          | column name specifying time, default is <code>'time'</code> .                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| <code>status</code>        | column name specifying event status, default is <code>'status'</code> .                                                                                                                                                                                                                                                                                                                                                                                                                           |
| <code>global_method</code> | method used to obtain global p value for cox model, should be one of <code>"likelihood"</code> , <code>"wald"</code> , <code>"logrank"</code> . The likelihood-ratio test, Wald test, and score logrank statistics. These three methods are asymptotically equivalent. For large enough <code>N</code> , they will give similar results. For small <code>N</code> , they may differ somewhat. The Likelihood ratio test has better behavior for small sample sizes, so it is generally preferred. |
| <code>keep_models</code>   | If <code>TRUE</code> , keep models as local files.                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| <code>return_models</code> | default <code>FALSE</code> . If <code>TRUE</code> , return a list contains cox models.                                                                                                                                                                                                                                                                                                                                                                                                            |
| <code>model_dir</code>     | a path for storing model results.                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| <code>verbose</code>       | if <code>TRUE</code> , print extra info.                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| <code>...</code>           | other parameters passing to <code>survival::coxph()</code> .                                                                                                                                                                                                                                                                                                                                                                                                                                      |

**Value**

a `ezcox` object

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

## Examples

```
library(survival)

# Build unvariable models
t1 <- ezcox(lung, covariates = c("age", "sex", "ph.ecog"))
t1

# Build multi-variable models
# Control variable 'age'
t2 <- ezcox(lung, covariates = c("sex", "ph.ecog"), controls = "age")
t2

# Return models
t3 <- ezcox(lung,
  covariates = c("age", "sex", "ph.ecog"),
  return_models = TRUE
)
t3
t4 <- ezcox(lung,
  covariates = c("sex", "ph.ecog"), controls = "age",
  return_models = TRUE
)
t4
```

---

 ezcox\_group

*Group Cox Analysis and Visualization*


---

## Description

Group Cox Analysis and Visualization

## Usage

```
ezcox_group(
  data,
  grp_var,
  covariate,
  controls = NULL,
  time = "time",
  status = "status",
  sort = FALSE,
  decreasing = TRUE,
  add_all = FALSE,
  add_caption = TRUE,
  verbose = TRUE,
  headings = list(variable = "Group", n = "N", measure = "Hazard ratio", ci = NULL, p =
    "p"),
  ...
)
```

**Arguments**

|                          |                                                                                                                                                                                                        |
|--------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>data</code>        | a <code>data.frame</code> containing variables, time and os status.                                                                                                                                    |
| <code>grp_var</code>     | a group column.                                                                                                                                                                                        |
| <code>covariate</code>   | a covariable for cox analysis.                                                                                                                                                                         |
| <code>controls</code>    | column names specifying controls. The names with pattern " <code>*:l()</code> " will be treated as interaction/combination term, please make sure all column names in data are valid R variable names. |
| <code>time</code>        | column name specifying time, default is 'time'.                                                                                                                                                        |
| <code>status</code>      | column name specifying event status, default is 'status'.                                                                                                                                              |
| <code>sort</code>        | if TRUE, sort the models by the HR values.                                                                                                                                                             |
| <code>decreasing</code>  | logical, should the sort order be increasing or decreasing?                                                                                                                                            |
| <code>add_all</code>     | if TRUE, add a group for all data rows.                                                                                                                                                                |
| <code>add_caption</code> | if TRUE, add caption to the plot.                                                                                                                                                                      |
| <code>verbose</code>     | if TRUE, print extra info.                                                                                                                                                                             |
| <code>headings</code>    | a list for setting the heading text.                                                                                                                                                                   |
| <code>...</code>         | other arguments passing to <code>forestmodel::forest_model()</code> .                                                                                                                                  |

**Value**

a list.

**Examples**

```
library(survival)
ezcox_group(lung, grp_var = "sex", covariate = "ph.ecog")
ezcox_group(lung, grp_var = "sex", covariate = "ph.ecog", controls = "age")
p <- ezcox_group(lung,
  grp_var = "sex", covariate = "ph.ecog",
  controls = "age", add_all = TRUE
)
```

---

ezcox\_parallel

---

*Parallelly Run Cox Analysis in Batch Mode*


---

**Description**

Parallelly Run Cox Analysis in Batch Mode

**Usage**

```
ezcox_parallel(
  data,
  covariates,
  controls = NULL,
  time = "time",
  status = "status",
  batch_size = 100,
  global_method = c("likelihood", "wald", "logrank"),
  keep_models = FALSE,
  return_models = FALSE,
  model_dir = file.path(tempdir(), "ezcox"),
  parallel = TRUE,
  verbose = FALSE
)
```

**Arguments**

|                            |                                                                                                                                                                                                                                                                                                                                                                                                                             |
|----------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>data</code>          | a <code>data.frame</code> containing variables, time and os status.                                                                                                                                                                                                                                                                                                                                                         |
| <code>covariates</code>    | column names specifying variables.                                                                                                                                                                                                                                                                                                                                                                                          |
| <code>controls</code>      | column names specifying controls.                                                                                                                                                                                                                                                                                                                                                                                           |
| <code>time</code>          | column name specifying time, default is 'time'.                                                                                                                                                                                                                                                                                                                                                                             |
| <code>status</code>        | column name specifying event status, default is 'status'.                                                                                                                                                                                                                                                                                                                                                                   |
| <code>batch_size</code>    | processing size in a batch.                                                                                                                                                                                                                                                                                                                                                                                                 |
| <code>global_method</code> | method used to obtain global p value for cox model, should be one of "likelihood", "wald", "logrank". The likelihood-ratio test, Wald test, and score logrank statistics. These three methods are asymptotically equivalent. For large enough N, they will give similar results. For small N, they may differ somewhat. The Likelihood ratio test has better behavior for small sample sizes, so it is generally preferred. |
| <code>keep_models</code>   | If TRUE, keep models as local files.                                                                                                                                                                                                                                                                                                                                                                                        |
| <code>return_models</code> | default FALSE. If TRUE, return a list contains cox models.                                                                                                                                                                                                                                                                                                                                                                  |
| <code>model_dir</code>     | a path for storing model results.                                                                                                                                                                                                                                                                                                                                                                                           |
| <code>parallel</code>      | if TRUE, do parallel computation by <b>furrr</b> package.                                                                                                                                                                                                                                                                                                                                                                   |
| <code>verbose</code>       | if TRUE, print extra info. If parallel is TRUE, set verbose to FALSE may speed up.                                                                                                                                                                                                                                                                                                                                          |

**Value**

a `ezcox` object

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```
library(survival)
t <- ezcox_parallel(lung, covariates = c("sex", "ph.ecog"), controls = "age")
t
```

filter\_ezcox

*Filter ezcox***Description**

Filter ezcox

**Usage**

```
filter_ezcox(x, levels = "auto", type = c("both", "contrast", "ref"))
```

**Arguments**

|        |                                                                                                                                                                                    |
|--------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| x      | a ezcox object from <a href="#">ezcox()</a> .                                                                                                                                      |
| levels | levels to filter, default is 'auto', it will filter all control variables.                                                                                                         |
| type   | default is 'both' for filtering both contrast level and reference level. It can also be 'contrast' for filtering only contrast level and 'ref' for filtering only reference level. |

**Value**

a ezcox object

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```
library(survival)
lung$ph.ecog <- factor(lung$ph.ecog)
zz <- ezcox(lung, covariates = c("sex", "age"), controls = "ph.ecog")
zz
filter_ezcox(zz)
filter_ezcox(zz, c("0", "2"))
filter_ezcox(zz, c("0", "2"), type = "contrast")
t <- filter_ezcox(zz, c("0", "2"), type = "ref")
t
```

forester

*Create a forest plot for simple data***Description**

Create a forest plot for simple data

**Usage**

```
forester(
  data,
  display_cols = c("Variable", "HR", "lower_95", "upper_95"),
  estimate_precision = 2,
  null_line_at = 1,
  font_family = "mono",
  x_scale_linear = TRUE,
  xlim = NULL,
  xbreaks = NULL,
  point_sizes = 3,
  point_shape = 16,
  label_hjust = 0,
  label_vjust = -1,
  label_color = "blue",
  label_size = 3
)
```

**Arguments**

|                                                                                                          |                                                                                                                                                                    |
|----------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>data</code>                                                                                        | Data frame (required). The information to be displayed as the forest plot.                                                                                         |
| <code>display_cols</code>                                                                                | 4 columns stand for axis text and the forest data, default using <code>c("term", "HR", "conf.low", "conf.high")</code> .                                           |
| <code>estimate_precision</code>                                                                          | Integer. The number of decimal places on the estimate (default 2).                                                                                                 |
| <code>null_line_at</code>                                                                                | Numeric. Default 0. Change to 1 if using relative measures such as OR, RR.                                                                                         |
| <code>font_family</code>                                                                                 | String. The font to use for the ggplot. Default "mono".                                                                                                            |
| <code>x_scale_linear</code>                                                                              | Logical. Default TRUE, change to FALSE for log scale                                                                                                               |
| <code>xlim</code>                                                                                        | Vector. Manually specify limits for the x axis as a vector length 2, i.e. <code>c(low, high)</code>                                                                |
| <code>xbreaks</code>                                                                                     | Vector. X axis breaks to label. Specify limits in xlim if using this option.                                                                                       |
| <code>point_sizes</code>                                                                                 | Vector. Length should be equal to 1 or <code>nrow(left_side_data)</code> . The sizes of the points in the center plot, where 3.25 is the default.                  |
| <code>point_shape</code>                                                                                 | Vector. Length should be equal to 1 or <code>nrow(left_side_data)</code> . The shapes of the points in the center plot, where 16 (a filled circle) is the default. |
| <code>label_hjust</code> , <code>label_vjust</code> , <code>label_color</code> , <code>label_size</code> | hjust, vjust color and size for the label text.                                                                                                                    |

**Value**

a ggplot object.

**Examples**

```
library(survival)

t1 <- ezcox(lung, covariates = c(
  "age", "sex",
  "ph.karno", "pat.karno"
))
p <- forester(t1, xlim = c(0, 1.5))
p
p2 <- forester(t1, xlim = c(0.5, 1.5))
p2
```

---

get\_models

---

*Get Model List from ezcox Object*


---

**Description**

Models are renamed by the formulas.

**Usage**

```
get_models(x, variables = NULL)
```

**Arguments**

**x** a ezcox object from [ezcox\(\)](#).

**variables** a character vector representing variables to select.

**Value**

a named list with class ezcox\_models

**Examples**

```
library(survival)
zz <- ezcox(lung, covariates = c("sex", "ph.ecog"), controls = "age", return_models = TRUE)
mds <- get_models(zz)
str(mds, max.level = 1)
```

show\_forest

*Show Forest Plot***Description**

This is a wrapper of function `ezcox`, `get_models` and `show_models`. It focus on generating forest plot easily and flexibly.

**Usage**

```
show_forest(
  data,
  covariates,
  controls = NULL,
  time = "time",
  status = "status",
  merge_models = FALSE,
  model_names = NULL,
  vars_to_show = NULL,
  drop_controls = FALSE,
  add_caption = TRUE,
  point_size = 3,
  point_shape = 15,
  color = "red",
  banded = TRUE,
  headings = list(variable = "Variable", n = "N", measure = "Hazard ratio", ci = NULL, p
    = "p"),
  model_dir = file.path(tempdir(), "ezcox"),
  verbose = TRUE,
  ...
)
```

**Arguments**

|                           |                                                                                                                                                                                                                   |
|---------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>data</code>         | a <code>data.frame</code> containing variables, time and os status.                                                                                                                                               |
| <code>covariates</code>   | a character vector optionally listing the variables to include in the plot (defaults to all variables).                                                                                                           |
| <code>controls</code>     | column names specifying controls. The names with pattern <code>"*:l()"</code> will be treated as interaction/combination term, please make sure all column names in <code>data</code> are valid R variable names. |
| <code>time</code>         | column name specifying time, default is <code>'time'</code> .                                                                                                                                                     |
| <code>status</code>       | column name specifying event status, default is <code>'status'</code> .                                                                                                                                           |
| <code>merge_models</code> | if <code>'TRUE'</code> , merge all models and keep the plot tight.                                                                                                                                                |
| <code>model_names</code>  | model names to show when <code>merge_models=TRUE</code> .                                                                                                                                                         |

|               |                                                                                                                                                         |
|---------------|---------------------------------------------------------------------------------------------------------------------------------------------------------|
| vars_to_show  | default is NULL, show all variables (including controls). You can use this to choose variables to show, but remember, the models have not been changed. |
| drop_controls | works when covariates=NULL and models is a ezcox_models, if TRUE, it removes control variables automatically.                                           |
| add_caption   | if TRUE, add caption to the plot.                                                                                                                       |
| point_size    | size of point.                                                                                                                                          |
| point_shape   | shape value of point.                                                                                                                                   |
| color         | color for point and segment.                                                                                                                            |
| banded        | if TRUE (default), create banded background color.                                                                                                      |
| headings      | a list for setting the heading text.                                                                                                                    |
| model_dir     | a path for storing model results.                                                                                                                       |
| verbose       | if TRUE, print extra info.                                                                                                                              |
| ...           | other arguments passing to <code>forestmodel::forest_model()</code> .                                                                                   |

**Value**

a ggplot object

**Examples**

```
library(survival)
show_forest(lung, covariates = c("sex", "ph.ecog"), controls = "age")
show_forest(lung, covariates = c("sex", "ph.ecog"), controls = "age", merge_models = TRUE)
show_forest(lung,
  covariates = c("sex", "ph.ecog"), controls = "age", merge_models = TRUE,
  drop_controls = TRUE
)
p <- show_forest(lung,
  covariates = c("sex", "ph.ecog"), controls = "age", merge_models = TRUE,
  vars_to_show = "sex"
)
p
```

---

|             |                 |
|-------------|-----------------|
| show_models | Show Cox Models |
|-------------|-----------------|

---

**Description**

Show Cox Models

**Usage**

```
show_models(
  models,
  model_names = NULL,
  covariates = NULL,
  merge_models = FALSE,
  drop_controls = FALSE,
  headings = list(variable = "Variable", n = "N", measure = "Hazard ratio", ci = NULL, p
    = "p"),
  ...
)
```

**Arguments**

|                            |                                                                                                                                                       |
|----------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>models</code>        | a <code>ezcox_models</code> from <code>get_models()</code> or a (named) list of Cox models.                                                           |
| <code>model_names</code>   | model names to show when <code>merge_models=TRUE</code> .                                                                                             |
| <code>covariates</code>    | a character vector optionally listing the variables to include in the plot (defaults to all variables).                                               |
| <code>merge_models</code>  | if 'TRUE', merge all models and keep the plot tight.                                                                                                  |
| <code>drop_controls</code> | works when <code>covariates=NULL</code> and <code>models</code> is a <code>ezcox_models</code> , if TRUE, it removes control variables automatically. |
| <code>headings</code>      | a list for setting the heading text.                                                                                                                  |
| <code>...</code>           | other arguments passing to <code>forestmodel::forest_model()</code> .                                                                                 |

**Value**

a `ggplot` object

**Examples**

```
library(survival)
zz <- ezcox(lung, covariates = c("sex", "ph.ecog"), controls = "age", return_models = TRUE)
mds <- get_models(zz)
show_models(mds)
show_models(mds, model_names = paste0("Model ", 1:2))
show_models(mds, covariates = c("sex", "ph.ecog"))
show_models(mds, drop_controls = TRUE)
show_models(mds, merge_models = TRUE)
p <- show_models(mds, merge_models = TRUE, drop_controls = TRUE)
p
```

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