

# Package ‘epiomics’

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**Title** Analysis of Omics Data in Observational Studies

**Version** 1.1.0

**Description** A collection of fast and flexible functions for analyzing omics data in observational studies. Multiple different approaches for integrating multiple environmental/genetic factors, omics data, and/or phenotype data are implemented. This includes functions for performing omics wide association studies with one or more variables of interest as the exposure or outcome; a function for performing a meet in the middle analysis for linking exposures, omics, and outcomes (as described by Chadeau-Hyam et al., (2010) [doi:10.3109/1354750X.2010.533285](https://doi.org/10.3109/1354750X.2010.533285)); and a function for performing a mixtures analysis across all omics features using quantile-based g-Computation (as described by Keil et al., (2019) [doi:10.1289/EHP5838](https://doi.org/10.1289/EHP5838)).

**License** GPL (>= 3)

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coef_plot_from_owas	<i>Create volcano plot using results from owas</i>
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## Description

Creates a coefficient plot based on ggplot using the results from the owas function.

## Usage

```
coef_plot_from_owas(
  df,
  main_cat_var = NULL,
  order_effects = TRUE,
  highlight_adj_p = TRUE,
  highlight_adj_p_threshold = 0.05,
  effect_ratio = FALSE,
  flip_axis = FALSE,
  filter_p_less_than = 1
)
```

## Arguments

df	output from owas function call, using conf_int = TRUE.
main_cat_var	Which variable should be the primary categorical variable? Should be either var_name or feature_name. Only relevant if both var_name and feature_name have more than one level. Default is NULL, and the y-axis is chosen as the variable that has more levels.
order_effects	Should features be ordered by the mean effect estimate? Default is TRUE.
highlight_adj_p	Should features which meet a specific adjusted p-value threshold be highlighted? Default is TRUE.
highlight_adj_p_threshold	If highlight_adj_p = TRUE, can set annotation_adj_p_threshold to change the adjusted p-value threshold for which features will be highlighted. Defaults to 0.05.
effect_ratio	Are the effect estimates on the ratio scale (ie, should the null effect line be centered at 1)? Defaults to FALSE.

`flip_axis` Flip the x and y axis? Default is FALSE, and the y-axis is plotted with the features or variable names.

`filter_p_less_than` P-value threshold for which features/variables will be included in the plot. Default is 1, and all features will be included.

**Value**

A ggplot figure

**Examples**

```
data("example_data")

# Get names of omics
colnames_omic_fts <- colnames(example_data)[
  grep("feature_",
       colnames(example_data))][1:5]

# Run function with continuous exposure as the variable of interest
owas_out <- owas(df = example_data,
                 var = "exposure1",
                 omics = colnames_omic_fts,
                 covars = c("age", "sex"),
                 var_exposure_or_outcome = "exposure",
                 family = "gaussian",
                 conf_int = TRUE)

coef_plot_from_owas(owas_out)
```

---

example\_data

*Example data with multiple exposures, multiple outcomes,*

---

**Description**

Example data with multiple exposures, multiple outcomes,

**Usage**

```
data(example_data)
```

**Format**

An dataframe with multiple exposures, outcomes, and omics features.

**Examples**

```
data(example_data)
```

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meet_in_middle	<i>Perform 'omics wide association study</i>
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### Description

Implements a meet in the middle analysis for identifying omics associated with both exposures and outcomes, as described by Chadeau-Hyam et al., 2010.

### Usage

```
meet_in_middle(
  df,
  exposure,
  outcome,
  omics,
  covars = NULL,
  outcome_family = "gaussian",
  confidence_level = 0.95,
  conf_int = FALSE,
  ref_group_exposure = NULL,
  ref_group_outcome = NULL
)
```

### Arguments

df	Dataframe
exposure	Name of the exposure of interest. Can be either continuous or dichotomous. Currently, only a single exposure is supported.
outcome	Name of the outcome of interest. Can be either continuous or dichotomous. For dichotomous variables, must set outcome_family to "logistic", and values must be either 0/1 or a factor with the first level representing the reference group. Currently, only a single outcome is supported.
omics	Names of all omics features in the dataset
covars	Names of covariates (can be NULL)
outcome_family	"gaussian" for linear models (via lm) or "binomial" for logistic (via glm)
confidence_level	Confidence level for marginal significance (defaults to 0.95)
conf_int	Should Confidence intervals be generated for the estimates? Default is FALSE. Setting to TRUE will take longer. For logistic models, calculates Wald confidence intervals via conf_int.default.
ref_group_exposure	Reference category if the exposure is a character or factor. If not, can leave empty.
ref_group_outcome	Reference category if the outcome is a character or factor. If not, can leave empty.

**Value**

A list of three dataframes, containing:

1. Results from the Exposure-Omics Wide Association Study
2. Results from the Omics-Outcome Wide Association Study
3. Overlapping significant features from 1 and 2. For each omics wide association, results are provided in a data frame with 6 columns: `feature_name`: name of the omics feature estimate: the model estimate for the feature. For linear models, this is the beta: for logistic models, this is the log odds. `se`: Standard error of the estimate `p_value`: p-value for the estimate `adjusted_pval`: FDR adjusted p-value threshold: Marginal significance, based on unadjusted p-values

**Examples**

```
# Load Example Data
data("example_data")

# Get names of omics
colnames_omic_fts <- colnames(example_data)[grep("feature_",
                                                colnames(example_data))][1:10]

# Meet in the middle with a dichotomous outcome
res <- meet_in_middle(df = example_data,
                     exposure = "exposure1",
                     outcome = "disease1",
                     omics = colnames_omic_fts,
                     covars = c("age", "sex"),
                     outcome_family = "binomial")

# Meet in the middle with a continuous outcome
res <- meet_in_middle(df = example_data,
                     exposure = "exposure1",
                     outcome = "weight",
                     omics = colnames_omic_fts,
                     covars = c("age", "sex"),
                     outcome_family = "gaussian")

# Meet in the middle with a continuous outcome and no covariates
res <- meet_in_middle(df = example_data,
                     exposure = "exposure1",
                     outcome = "weight",
                     omics = colnames_omic_fts,
                     outcome_family = "gaussian")
```

## Description

Implements an omics wide association study with the option of using the 'omics data as either the dependent variable (i.e., for performing an exposure → 'omics analysis) or using the 'omics as the independent variable (i.e., for performing an 'omics → outcome analysis). Allows for either continuous or dichotomous outcomes, and provides the option to adjust for covariates.

## Usage

```
owas(
  df,
  var,
  omics,
  covars = NULL,
  var_exposure_or_outcome,
  family = "gaussian",
  confidence_level = 0.95,
  conf_int = FALSE,
  ref_group = NULL
)
```

## Arguments

df	Dataset
var	Name of the variable or variables of interest- this is usually either an exposure variable or an outcome variable. Can be either continuous or dichotomous. For dichotomous variables, must set family to "binomial", and values must be either 0/1 or a factor with the first level representing the reference group. Can handle multiple variables, but they must all be of the same family.
omics	Names of all omics features in the dataset
covars	Names of covariates (can be NULL)
var_exposure_or_outcome	Is the variable of interest an exposure (independent variable) or outcome (dependent variable)? Must be either "exposure" or "outcome"
family	"gaussian" (default) for linear models (via lm) or "binomial" for logistic (via glm)
confidence_level	Confidence level for marginal significance (defaults to 0.95, or an alpha of 0.05)
conf_int	Should Confidence intervals be generated for the estimates? Default is FALSE. Setting to TRUE will take longer. For logistic models, calculates Wald confidence intervals via <code>confint.default</code> .
ref_group	Reference category if the variable of interest is a character or factor. If not, can leave empty.

## Value

A data frame with 6 columns: `feature_name`: name of the omics feature `estimate`: the model estimate for the feature. For linear models, this is the beta; for logistic models, this is the log odds. `se`:

Standard error of the estimate test\_statistic: t-value p\_value: p-value for the estimate adjusted\_pval:  
FDR adjusted p-value threshold: Marginal significance, based on unadjusted p-values

### Examples

```
# Load Example Data
data("example_data")

# Get names of omics
colnames_omic_fts <- colnames(example_data)[grep("feature_",
                                                colnames(example_data))][1:10]

# Get names of exposures
expnms = c("exposure1", "exposure2", "exposure3")

# Run function with one continuous exposure as the variable of interest
owas(df = example_data,
     var = "exposure1",
     omics = colnames_omic_fts,
     covars = c("age", "sex"),
     var_exposure_or_outcome = "exposure",
     family = "gaussian")

# Run function with multiple continuous exposures as the variable of interest
owas(df = example_data,
     var = expnms,
     omics = colnames_omic_fts,
     covars = c("age", "sex"),
     var_exposure_or_outcome = "exposure",
     family = "gaussian")

# Run function with dichotomous outcome as the variable of interest
owas(df = example_data,
     var = "disease1",
     omics = colnames_omic_fts,
     covars = c("age", "sex"),
     var_exposure_or_outcome = "outcome",
     family = "binomial")
```

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owas\_clogit

---

*Perform 'omics wide association study for matched case control studies*


---

### Description

Implements an omics wide association study for matched case control studies using conditional logistic regression. For this function, the variable of of interest should be a dichotomous outcome, and the strata is the variable indicating the matching.

**Usage**

```
owas_clogit(
  df,
  cc_status,
  cc_set,
  omics,
  covars = NULL,
  confidence_level = 0.95,
  conf_int = FALSE,
  method = "efron"
)
```

**Arguments**

df	Dataset
cc_status	Name of the variable indicating case control status. Must be either 0/1 or a factor with the first level representing the reference group.
cc_set	Name of the variable indicating the case control set.
omics	Names of all omics features in the dataset reference group.
covars	Names of covariates (can be NULL)
confidence_level	Confidence level for marginal significance (defaults to 0.95, or an alpha of 0.05)
conf_int	Should Confidence intervals be generated for the estimates? Default is FALSE. Setting to TRUE will take longer. For logistic models, calculates Wald confidence intervals via <code>confint.default</code> .
method	method used the correct (exact) calculation in the conditional likelihood or one of the approximations. Default is "efron". Passed to <code>clogit</code> .

**Value**

A data frame with 6 columns: `feature_name`: name of the omics feature `estimate`: the model estimate for the feature. For linear models, this is the beta; for logistic models, this is the log odds. `se`: Standard error of the estimate `t_value`: t-value `p_value`: p-value for the estimate `adjusted_pval`: FDR adjusted p-value threshold: Marginal significance, based on unadjusted p-values

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owas\_qgcomp

---

*Perform omics wide association study using qgcomp*


---

**Description**

Omics wide association study using quantile-based g-Computation (as described by Keil et al., (2019) [doi:10.1289/EHP5838](https://doi.org/10.1289/EHP5838)) to examine associations of exposure mixtures with each individual 'omics feature as an outcome 'omics data as either the dependent variable. Allows for either continuous or dichotomous outcomes, and provides the option to adjust for covariates.



**Usage**

```
owas_qgcomp(
  df,
  expnms,
  omics,
  covars = NULL,
  q = 4,
  confidence_level = 0.95,
  family = "gaussian",
  rr = TRUE,
  run.qgcomp.boot = TRUE
)
```

**Arguments**

df	Dataset
expnms	Name of the exposures. Can be either continuous or dichotomous. For dichotomous variables, must set q to "NULL", and values must be either 0/1.
omics	Names of all omics features in the dataset
covars	Names of covariates (can be NULL)
q	NULL or number of quantiles used to create quantile indicator variables representing the exposure variables. Defaults to 4. If NULL, then qgcomp proceeds with un-transformed version of exposures in the input datasets (useful if data are already transformed, or for performing standard g-computation).
confidence_level	Confidence level for marginal significance (defaults to 0.95, or an alpha of 0.05)
family	Currently only "gaussian" (default) for linear models (via lm) or "binomial" for logistic. Default is "gaussian".
rr	see qgcomp()
run.qgcomp.boot	Should the model be fit with qgcomp.boot? See package <a href="#">qgcomp.boot</a> for details. Default is TRUE. Setting to FALSE decreases computational time.

**Value**

A data frame with the following columns: feature: name of the omics feature psi: the model estimate for the feature. For linear models, this is the beta; for logistic models, this is the log odds. lcl\_psi: the lower confidence interval. ucl\_psi: the upper confidence interval. p\_value: p-value for the estimate test\_statistic: t-statistic for psi coefficient adjusted\_pval: FDR adjusted p-value threshold: Marginal significance, based on unadjusted p-values covariates: the names of covariates in the model, if any coef\_exposure: the individual coefficient of each exposure

**Examples**

```
# Load Example Data
data("example_data")
```

```
# Get names of omics
colnames_omic_fts <- colnames(example_data)[grep("feature_",
                                                colnames(example_data))][1:5]

# Names of exposures in mixture
exposure_names = c("exposure1", "exposure2", "exposure3")

# Run function without covariates
out <- owas_qgcomp(df = example_data,
                  expnms = exposure_names,
                  omics = colnames_omic_fts,
                  q = 4,
                  confidence_level = 0.95)

# Run analysis with covariates
out <- owas_qgcomp(df = example_data,
                  expnms = c("exposure1", "exposure2", "exposure3"),
                  covars = c("weight", "age", "sex"),
                  omics = colnames_omic_fts,
                  q = 4,
                  confidence_level = 0.95)
```

---

volcano\_owas

*Create volcano plot using results from owas*

---

## Description

Creates a volcano plot based on ggplot using the results from the owas function.

## Usage

```
volcano_owas(
  df,
  annotate_ftrs = TRUE,
  annotation_p_threshold = 0.05,
  highlight_adj_p = TRUE,
  highlight_adj_p_threshold = 0.05,
  horizontal_line_p_value = 0.05
)
```

## Arguments

**df** output from owas function call

**annotate\_ftrs** Should features be annotated with the feature name? Default is TRUE. If necessary can change the p\_value\_threshold as well.

`annotation_p_threshold`  
If `annotate_fts = TRUE`, can set `annotation_p_threshold` to change the p-value threshold for which features will be annotated. Defaults to 0.05.

`highlight_adj_p`  
Should features which meet a specific adjusted p-value threshold be highlighted? Default is `TRUE`.

`highlight_adj_p_threshold`  
If `highlight_adj_p = TRUE`, can set `annotation_adj_p_threshold` to change the adjusted p-value threshold for which features will be highlighted. Defaults to 0.05.

`horizontal_line_p_value`  
Set the p-value for the horizontal line for the threshold of significance.

**Value**

A ggplot figure

**Examples**

```
data("example_data")

# Get names of omics
colnames_omic_fts <- colnames(example_data)[
  grep("feature_",
       colnames(example_data))][1:5]

# Run function with continuous exposure as the variable of interest
owas_out <- owas(df = example_data,
                 var = "exposure1",
                 omics = colnames_omic_fts,
                 covars = c("age", "sex"),
                 var_exposure_or_outcome = "exposure",
                 family = "gaussian")

vp <- volcano_owas(owas_out)
```

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