# Package 'colocboost'

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Title Multi-Context Colocalization Analysis for QTL and GWAS Studies

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**Description** A multi-task learning approach to variable selection regression with highly correlated predictors and sparse effects,

based on frequentist statistical inference. It provides statistical evidence to identify which subsets of predictors have non-zero

effects on which subsets of response variables, motivated and designed for colocalization analysis across genome-wide association studies (GWAS) and quantitative trait loci (QTL) studies.

The ColocBoost model is described in Cao et. al. (2025) <doi:10.1101/2025.04.17.25326042>.

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## LazyData true

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URL https://github.com/StatFunGen/colocboost

BugReports https://github.com/StatFunGen/colocboost/issues

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Ambiguous\_Colocalization

A real data example includes an ambiguous colocalization between eQTL and GWAS

# Description

An example result from one of our real data applications, which shows an ambiguous colocalization between eQTL and GWAS.

# Usage

Ambiguous\_Colocalization

## Format

Ambiguous\_Colocalization:
A list with 2 elements
ColocBoost\_Results A colocboost output object
SuSiE\_Results Two susie output object for eQTL and GWAS
COLOC\_V5\_Results A coloc output object

## Source

The Ambiguous\_Colocalization dataset contains a real data example from one of our real data applications, which shows an ambiguous colocalization between eQTL and GWAS. The dataset is specifically designed for evaluating and demonstrating the capabilities of ColocBoost in real data applications. See details in tutorial vignette https://statfungen.github.io/colocboost/articles/index.html.

# See Also

Other colocboost\_data: Heterogeneous\_Effect, Ind\_5traits, Non\_Causal\_Strongest\_Marginal, Sumstat\_5traits, Weaker\_GWAS\_Effect

colocboost

*ColocBoost: A gradient boosting informed multi-omics xQTL colocalization method* 

## Description

colocboost implements a proximity adaptive smoothing gradient boosting approach for multi-trait colocalization at gene loci, accommodating multiple causal variants. This method, introduced by Cao etc. (2025), is particularly suited for scaling to large datasets involving numerous molecular quantitative traits and disease traits. In brief, this function fits a multiple linear regression model Y = XB + E in matrix form. ColocBoost can be generally used in multi-task variable selection regression problem.

## Usage

```
colocboost(
  X = NULL,
  Y = NULL,
  sumstat = NULL,
  LD = NULL,
  dict_YX = NULL,
  dict_sumstatLD = NULL,
  outcome_names = NULL,
  focal_outcome_idx = NULL,
  focal_outcome_variables = TRUE,
  overlap_variables = FALSE,
```

```
intercept = TRUE,
standardize = TRUE,
effect_est = NULL,
effect_se = NULL,
effect_n = NULL,
M = 500,
stop_thresh = 1e-06,
tau = 0.01,
learning_rate_init = 0.01,
learning_rate_decay = 1,
dynamic_learning_rate = TRUE,
prioritize_jkstar = TRUE,
func_compare = "min_max",
jk_equiv_corr = 0.8,
jk_equiv_loglik = 1,
coloc_thresh = 0.1,
lambda = 0.5,
lambda_focal_outcome = 1,
func_simplex = "LD_z2z",
func_multi_test = "lfdr",
stop_null = 1,
multi_test_max = 1,
multi_test_thresh = 1,
ash_prior = "normal",
p.adjust.methods = "fdr",
residual_correlation = NULL,
coverage = 0.95,
min_cluster_corr = 0.8,
dedup = TRUE,
overlap = TRUE,
n_{purity} = 100,
min_abs_corr = 0.5,
median_abs_corr = NULL,
median_cos_abs_corr = 0.8,
tol = 1e-09,
merge_cos = TRUE,
sec_coverage_thresh = 0.8,
weight_fudge_factor = 1.5,
check_null = 0.1,
check_null_method = "profile",
check_null_max = 0.025,
weaker_effect = TRUE,
LD_free = FALSE,
output_level = 1
```

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)

# Arguments

Х	A list of genotype matrices for different outcomes, or a single matrix if all outcomes share the same genotypes. Each matrix should have column names, if sample sizes and variables possibly differing across matrices.		
Y	A list of vectors of outcomes or an N by L matrix if it is considered for the same X and multiple outcomes.		
sumstat	A list of data.frames of summary statistics. The columns of data.frame should include either z or beta/sebeta. n is the sample size for the summary statistics, it is highly recommendation to provide. variant is required if sumstat for different outcomes do not have the same number of variables. var_y is the variance of phenotype (default is 1 meaning that the Y is in the "standardized" scale).		
LD	A list of correlation matrix indicating the LD matrix for each genotype. It also could be a single matrix if all sumstats were obtained from the same genotypes.		
dict_YX	A L by 2 matrix of dictionary for X and Y if there exist subsets of outcomes corresponding to the same X matrix. The first column should be 1:L for L outcomes. The second column should be the index of X corresponding to the outcome. The innovation: do not provide the same matrix in X to reduce the computational burden.		
dict_sumstatLD	A L by 2 matrix of dictionary for sumstat and LD if there exist subsets of outcomes corresponding to the same sumstat. The first column should be 1:L for L sumstat The second column should be the index of LD corresponding to the sumstat. The innovation: do not provide the same matrix in LD to reduce the computational burden.		
outcome_names	The names of outcomes, which has the same order for Y.		
<pre>focal_outcome_i</pre>	focal_outcome_idx		
	The index of the focal outcome if perform GWAS-xQTL ColocBoost		
focal_outcome_v	ariables		
	If focal_outcome_variables = IRUE, only consider the variables exist in the focal outcome.		
overlap_variabl	overlap_variables		
	If overlap_variables = TRUE, only perform colocalization in the overlapped region.		
intercept	If overlap_variables = TRUE, only perform colocalization in the overlapped region. If intercept = TRUE, the intercept is fitted. Setting intercept = FALSE is generally not recommended.		
intercept standardize	If overlap_variables = TRUE, only perform colocalization in the overlapped region. If intercept = TRUE, the intercept is fitted. Setting intercept = FALSE is generally not recommended. If standardize = TRUE, standardize the columns of genotype and outcomes to unit variance.		
intercept standardize effect_est	If overlap_variables = TRUE, only perform colocalization in the overlapped region. If intercept = TRUE, the intercept is fitted. Setting intercept = FALSE is gen- erally not recommended. If standardize = TRUE, standardize the columns of genotype and outcomes to unit variance. Matrix of variable regression coefficients (i.e. regression beta values) in the genomic region		
<pre>intercept standardize effect_est effect_se</pre>	If overlap_variables = TRUE, only perform colocalization in the overlapped region. If intercept = TRUE, the intercept is fitted. Setting intercept = FALSE is gen- erally not recommended. If standardize = TRUE, standardize the columns of genotype and outcomes to unit variance. Matrix of variable regression coefficients (i.e. regression beta values) in the genomic region Matrix of standard errors associated with the beta values		
<pre>intercept standardize effect_est effect_se effect_n</pre>	If overlap_variables = TRUE, only perform colocalization in the overlapped region. If intercept = TRUE, the intercept is fitted. Setting intercept = FALSE is gen- erally not recommended. If standardize = TRUE, standardize the columns of genotype and outcomes to unit variance. Matrix of variable regression coefficients (i.e. regression beta values) in the genomic region Matrix of standard errors associated with the beta values A scalar or a vector of sample sizes for estimating regression coefficients. Highly recommended!		
<pre>intercept standardize effect_est effect_se effect_n M</pre>	If overlap_variables = TRUE, only perform colocalization in the overlapped region. If intercept = TRUE, the intercept is fitted. Setting intercept = FALSE is gen- erally not recommended. If standardize = TRUE, standardize the columns of genotype and outcomes to unit variance. Matrix of variable regression coefficients (i.e. regression beta values) in the genomic region Matrix of standard errors associated with the beta values A scalar or a vector of sample sizes for estimating regression coefficients. Highly recommended! The maximum number of gradient boosting rounds for each outcome (default is 500).		

tau	The smooth parameter for proximity adaptive smoothing weights for the best update jk-star.
learning_rate_i	nit
	The minimum learning rate for updating in each iteration.
<pre>learning_rate_c</pre>	lecay
	The decayrate for learning rate. If the objective function is large at the early iterations, we need to have the higher learning rate to improve the computational efficiency.
dynamic_learnir	ng_rate
	If dynamic_learning_rate = TRUE, the dynamic learning rate based on learning_rate_init and learning_rate_decay will be used in SEC.
prioritize_jkst	ar
	When prioritize_jkstar = TRUE, the selected outcomes will prioritize best update j_k^star in SEC.
func_compare	The criterion when we update jk-star in SEC (default is "min_max").
jk_equiv_corr	The LD cutoff between overall best update jk-star and marginal best update jk-l for lth outcome
jk_equiv_loglik	
	The change of loglikelihood cutoff between overall best update jk-star and marginal best update jk-l for lth outcome
coloc_thresh	The cutoff of checking if the best update jk-star is the potential causal variable for outcome l if jk-l is not similar to jk-star (used in Delayed SEC).
lambda	The ratio [0,1] for z <sup>2</sup> and z in fun_prior simplex, default is 0.5
lambda_focal_ou	itcome
	The ratio for z <sup>2</sup> and z in fun_prior simplex for the focal outcome, default is 1
func_simplex	The data-driven local association simplex $\delta$ for smoothing the weights. Default is "LD_z2z" is the elastic net for z-score and also weighted by LD.
<pre>func_multi_test</pre>	
	The alternative method to check the stop criteria. When func_multi_test = "lfdr", boosting iterations will be stopped if the local FDR for all variables are greater than lfsr_max.
stop_null	The cutoff of nominal p-value when func_multi_test = "Z".
<pre>multi_test_max</pre>	The cutoff of the smallest FDR for stop criteria when func_multi_test = "lfdr" or func_multi_test = "lfsr".
<pre>multi_test_thre</pre>	esh
	The cutoff of the smallest FDR for pre-filtering the outcomes when func_multi_test = "lfdr" or func_multi_test = "lfsr".
ash_prior	The prior distribution for calculating lfsr when func_multi_test = "lfsr".
p.adjust.method	ls
	The adjusted pvalue method in stats:p.adj when func_multi_test = "fdr"
residual_correl	ation
	The residual correlation based on the sample overlap, it is diagonal if it is NULL.
coverage	A number between 0 and 1 specifying the "coverage" of the estimated colocal- ization confidence sets (CoS) (default is 0.95).

<pre>min_cluster_cor</pre>	r
	The small correlation for the weights distributions across different iterations to be decided having only one cluster.
dedup	If dedup = TRUE, the duplicate confidence sets will be removed in the post-processing.
overlap	If overlap = TRUE, the overlapped confidence sets will be removed in the post-processing.
n_purity	The maximum number of confidence set (CS) variables used in calculating the correlation ("purity") statistics. When the number of variables included in the CS is greater than this number, the CS variables are randomly subsampled.
min_abs_corr	Minimum absolute correlation allowed in a confidence set. The default is $0.5$ corresponding to a squared correlation of $0.25$ , which is a commonly used threshold for genotype data in genetic studies.
median_abs_corr	
	An alternative "purity" threshold for the CS. Median correlation between pairs of variables in a CS less than this threshold will be filtered out and not reported. When both min_abs_corr and median_abs_corr are set, a CS will only be removed if it fails both filters. Default set to NULL but it is recommended to set it to 0.8 in practice.
<pre>median_cos_abs_</pre>	corr
	Median absolute correlation between variants allowed to merge multiple colo- calized sets. The default is 0.8 corresponding to a stringent threshold to merge colocalized sets, which may resulting in a huge set.
tol	A small, non-negative number specifying the convergence tolerance for check- ing the overlap of the variables in different sets.
merge_cos	When merge_cos = TRUE, the sets for only one outcome will be merged if passed the median_cos_abs_corr.
<pre>sec_coverage_th</pre>	nresh
	A number between 0 and 1 specifying the weight in each SEC (default is 0.8).
<pre>weight_fudge_fa</pre>	actor
	The strength to integrate weight from different outcomes, default is 1.5
check_null	The cut off value for change conditional objective function. Default is 0.1.
check_null_meth	nod
	The metric to check the null sets. Default is "profile"
<pre>check_null_max</pre>	The smallest value of change of profile loglikelihood for each outcome.
weaker_effect	If weaker_effect = TRUE, consider the weaker single effect due to coupling effects
LD_free	When LD_free = FALSE, objective function doesn't include LD information.
output_level	When output_level = 1, return basic cos details for colocalization results When output_level = 2, return the ucos details for the single specific effects. When output_level = 3, return the entire Colocboost model to diagnostic results (more space).

## Details

The function colocboost implements the proximity smoothed gradient boosting method from Cao etc (2025). There is an additional step to help merge the confidence sets with small between\_putiry (default is 0.8) but within the same locus. This step addresses potential instabilities in linkage disequilibrium (LD) estimation that may arise from small sample sizes or discrepancies in minor allele frequencies (MAF) across different confidence sets.

# Value

A "colocboost" object with some or all of the following elements:

cos_summary	A summary table for colocalization events.
vср	The variable colocalized probability for each variable.
cos_details	A object with all information for colocalization results.
data_info	A object with detailed information from input data
model_info	A object with detailed information for colocboost model
ucos_details	A object with all information for trait-specific effects when output_level = 2.
diagnositci_det	ails
	A object with diagnostic details for ColocBoost model when output_level =
	3.

## Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ index.html

## Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 3
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5 # SNP10 affects trait 1</pre>
true_beta[10, 2] <- 0.4 # SNP10 also affects trait 2 (colocalized)</pre>
true_beta[50, 2] <- 0.3 # SNP50 only affects trait 2</pre>
true_beta[80, 3] <- 0.6 # SNP80 only affects trait 3</pre>
Y \leq matrix(0, N, L)
for (1 in 1:L) {
  Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)</pre>
}
res <- colocboost(X = X, Y = Y)</pre>
res$cos_details$cos$cos_index
```

colocboost\_plot

## Description

colocboost\_plot generates visualization plots for colocalization events from a ColocBoost analysis.

## Usage

```
colocboost_plot(
  cb_output,
  y = "log10p",
  grange = NULL,
  plot_cos_idx = NULL,
  outcome_idx = NULL,
  plot_all_outcome = FALSE,
  plot_focal_only = FALSE,
  plot_focal_cos_outcome_only = FALSE,
  points_color = "grey80",
  cos_color = NULL,
  add_vertical = FALSE,
  add_vertical_idx = NULL,
  outcome_names = NULL,
  plot_cols = 2,
  variant_coord = FALSE,
  show_top_variables = FALSE,
  show_cos_to_uncoloc = FALSE,
  show_cos_to_uncoloc_idx = NULL,
  show_cos_to_uncoloc_outcome = NULL,
  plot_ucos = FALSE,
  plot_ucos_idx = NULL,
  title_specific = NULL,
  ylim_each = TRUE,
  outcome_legend_pos = "top",
  outcome_legend_size = 1.8,
  \cos_{\log n} = c(0.05, 0.4),
  show_variable = FALSE,
  lab_style = c(2, 1),
  axis_style = c(2, 1),
  title_style = c(2.5, 2),
)
```

## Arguments

cb\_output

Output object from colocboost analysis

У	Specifies the y-axis values, default is "log10p" for -log10 transformed marginal association p-values.
grange	Optional plotting range of x-axis to zoom in to a specific region.
plot_cos_idx	Optional indices of CoS to plot
outcome_idx	Optional indices of outcomes to include in the plot. outcome_idx=NULL to plot only the outcomes having colocalization.
plot_all_outcom	ie C
	Optional to plot all outcome in the same figure.
plot_focal_only	
	Logical, if TRUE only plots colocalization with focal outcome, default is FALSE.
plot_focal_cos_	Logical, if TRUE only plots colocalization including at least on colocalized out- come with focal outcome, default is FALSE.
points_color	Background color for non-colocalized variables, default is "grey80".
cos_color	Optional custom colors for CoS.
add_vertical	Logical, if TRUE adds vertical lines at specified positions, default is FALSE
add_vertical_id	lx
	Optional indices for vertical lines.
outcome_names	Optional vector of outcomes names for the subtitle of each figure. outcome_names=NULL for the outcome name shown in data_info.
plot_cols	Number of columns in the plot grid, default is 2. If you have many colocaliza- tion. please consider increasing this.
variant_coord	Logical, if TRUE uses variant coordinates on x-axis, default is FALSE. This is required the variable names including position information.
show_top_variab	bles
	Logical, if TRUE shows top variables for each CoS, default is FALSE
show_cos_to_unc	coloc
- L	Logical, if TRUE shows colocalization to uncolocalized outcomes to diagnose, default is FALSE
snow_cos_to_unc	Ontional indices for showing CoS to all uncolocalized outcomes
show cos to unc	coloc outcome
0000_00_0	Optional outcomes for showing CoS to uncolocalized outcomes
plot_ucos	Logical, if TRUE plots also trait-specific (uncolocalized) sets, default is FALSE
plot_ucos_idx	Optional indices of trait-specific (uncolocalized) sets to plot when included
title_specific	Optional specific title to display in plot title
vlim_each	Logical, if TRUE uses separate y-axis limits for each plot, default is TRUE
outcome_legend_	pos
	Position for outcome legend, default is "top"
outcome_legend_	size
	Size for outcome legend text, default is 1.2
cos_legend_pos	Proportion of the legend from (left edge, bottom edge), default as (0.05, 0.4) at the left - median position

show_variable	Logical, if TRUE displays variant IDs, default is FALSE
lab_style	Vector of two numbers for label style (size, boldness), default is c(2, 1)
axis_style	Vector of two numbers for axis style (size, boldness), default is c(2, 1)
title_style	Vector of two numbers for title style (size, boldness), default is $c(2.5, 2)$
•••	Additional parameters passed to plot functions

## Value

Visualization plot for each colocalization event.

## Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ articles/Visualization\_ColocBoost\_Output.html

## Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 3
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5 # SNP10 affects trait 1</pre>
true_beta[10, 2] <- 0.4 # SNP10 also affects trait 2 (colocalized)</pre>
true_beta[50, 2] <- 0.3 # SNP50 only affects trait 2</pre>
true_beta[80, 3] <- 0.6 # SNP80 only affects trait 3</pre>
Y <- matrix(0, N, L)
for (1 in 1:L) {
  Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)</pre>
}
res <- colocboost(X = X, Y = Y)</pre>
colocboost_plot(res, plot_cols = 1)
colocboost_plot(res, plot_cols = 1, outcome_idx = 1:3)
```

get\_ambiguous\_colocalization

Get ambiguous colocalization events from trait-specific (uncolocalized) effects.

## Description

get\_ambiguous\_colocalization get the colocalization by discarding the weaker colocalization events or colocalized outcomes

# Usage

```
get_ambiguous_colocalization(
   cb_output,
   min_abs_corr_between_ucos = 0.5,
   median_abs_corr_between_ucos = 0.8,
   tol = 1e-09
)
```

# Arguments

cb_output	Output object from colocboost analysis
<pre>min_abs_corr_be</pre>	tween_ucos
	Minimum absolute correlation for variants across two trait-specific (uncolocal- ized) effects to be considered colocalized. The default is 0.5.
median_abs_corr	_between_ucos Median absolute correlation for variants across two trait-specific (uncolocalized) effects to be considered colocalized. The default is 0.8.
tol	A small, non-negative number specifying the convergence tolerance for check- ing the overlap of the variables in different sets.

# Value

A "colocboost" object of colocboost output with additional elements:

ambiguous\_cos If exists, a list of ambiguous trait-specific (uncolocalized) effects.

# Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ articles/Interpret\_ColocBoost\_Output.html

## See Also

Other colocboost\_inference: get\_colocboost\_summary(), get\_robust\_colocalization()

# Examples

```
data(Ambiguous_Colocalization)
test_colocboost_results <- Ambiguous_Colocalization$ColocBoost_Results
res <- get_ambiguous_colocalization(test_colocboost_results)
names(res$ambiguous_cos)</pre>
```

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get\_colocboost\_summary

Get summary tables from a ColocBoost output.

# Description

get\_colocboost\_summary get colocalization and trait-specific summary table with or without the outcomes of interest.

## Usage

```
get_colocboost_summary(
    cb_output,
    summary_level = 1,
    outcome_names = NULL,
    interest_outcome = NULL,
    region_name = NULL,
    min_abs_corr_between_ucos = 0.5,
    median_abs_corr_between_ucos = 0.8
)
```

# Arguments

cb_output	Output object from colocboost analysis	
summary_level	When summary_level = 1, return basic summary table for colocalization results. See details in get_ucos_summary function when summary_level = 2.	
outcome_names	Optional vector of names of outcomes, which has the same order as Y in the original analysis.	
interest_outcome		
	Optional vector specifying a subset of outcomes from outcome_names to focus on. When provided, only colocalization events that include at least one of these outcomes will be returned.	
region_name	Optional character string. When provided, adds a column with this gene name to the output table for easier filtering in downstream analyses.	
min_abs_corr_between_ucos		
	Minimum absolute correlation for variants across two trait-specific (uncolocal- ized) effects to be considered colocalized. The default is 0.5.	
<pre>median_abs_corr_between_ucos</pre>		
	Median absolute correlation for variants across two trait-specific (uncolocalized) effects to be considered colocalized. The default is 0.8.	

# Details

When summary\_level = 1, additional details and examples are introduced in get\_cos\_summary. When summary\_level = 2 or summary\_level = 3, additional details for trait-specific effects and ambiguous colocalization events are included. See get\_ucos\_summary for details on these tables.

## Value

A list containing results from the ColocBoost analysis:

- When summary\_level = 1 (default):
  - cos\_summary: A summary table for colocalization events with the following columns:
    - \* focal\_outcome: The focal outcome being analyzed if exists. Otherwise, it is FALSE.
    - \* colocalized\_outcomes: Colocalized outcomes for colocalization confidence set (CoS)
    - \* cos\_id: Unique identifier for colocalization confidence set (CoS)
    - \* purity: Minimum absolute correlation of variables within colocalization confidence set (CoS)
    - \* top\_variable: The variable with highest variant colocalization probability (VCP)
    - \* top\_variable\_vcp: Variant colocalization probability for the top variable
    - \* cos\_npc: Normalized probability of colocalization
    - \* min\_npc\_outcome: Minimum normalized probability of colocalized traits
    - \* n\_variables: Number of variables in colocalization confidence set (CoS)
    - \* colocalized\_index: Indices of colocalized variables
    - \* colocalized\_variables: List of colocalized variables
    - \* colocalized\_variables\_vcp: Variant colocalization probabilities for all colocalized variables
- When summary\_level = 2:
  - cos\_summary: As described above
  - ucos\_summary: A summary table for trait-specific (uncolocalized) effects
- When summary\_level = 3:
  - cos\_summary: As described above
  - ucos\_summary: A summary table for trait-specific (uncolocalized) effects
  - ambiguous\_cos\_summary: A summary table for ambiguous colocalization events from trait-specific effects

## Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ articles/Interpret\_ColocBoost\_Output.html

## See Also

Other colocboost\_inference: get\_ambiguous\_colocalization(), get\_robust\_colocalization()

## Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
```

## get\_cormat

```
X <- MASS::mvrnorm(N, rep(0, P), sigma)
colnames(X) <- paste0("SNP", 1:P)
L <- 3
true_beta <- matrix(0, P, L)
true_beta[10, 1] <- 0.5 # SNP10 affects trait 1
true_beta[10, 2] <- 0.4 # SNP10 also affects trait 2 (colocalized)
true_beta[50, 2] <- 0.3 # SNP50 only affects trait 2
true_beta[80, 3] <- 0.6 # SNP80 only affects trait 3
Y <- matrix(0, N, L)
for (l in 1:L) {
    Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)
}
res <- colocboost(X = X, Y = Y)
get_colocboost_summary(res)
```

get_cormat	A fast function to calculate correlation matrix (LD matrix) from indi-
	vidual level data

## Description

This function calculates the correlation matrix (LD matrix) from individual level data.

## Usage

```
get_cormat(X, intercepte = TRUE)
```

# Arguments

Х	A matrix of individual level data.
intercepte	A logical value indicating whether to include an intercept in the model. Default is FALSE.

# Value

A correlation matrix (LD matrix).

## See Also

Other colocboost\_utilities: get\_cos(), get\_cos\_purity(), get\_cos\_summary(), get\_hierarchical\_clusters(),
get\_ucos\_summary()

# Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))
X <- MASS::mvrnorm(N, rep(0, P), sigma)
cormat <- get_cormat(X)</pre>
```

get\_cos

Extract CoS at different coverage

# Description

get\_cos extracts colocalization confidence sets (CoS) at different coverage levels from ColocBoost results. When genotype data (X) or correlation matrix (Xcorr) is provided, it can also calculate and filter CoS based on purity statistics, ensuring that variants within each CoS are sufficiently correlated.

# Usage

```
get_cos(
   cb_output,
   coverage = 0.95,
   X = NULL,
   Xcorr = NULL,
   n_purity = 100,
   min_abs_corr = 0.5,
   median_abs_corr = NULL
)
```

# Arguments

cb_output	Output object from colocboost analysis
coverage	A number between 0 and 1 specifying the "coverage" of the estimated colocalization confidence sets (CoS) (default is $0.95$ ).
Х	Genotype matrix of values of the p variables. Used to compute correlations if Xcorr is not provided.
Xcorr	Correlation matrix of correlations between variables. Alternative to X.
n_purity	The maximum number of CoS variables used in calculating the correlation ("purity") statistics.
min_abs_corr	The minimum absolute correlation value of variants in a CoS to be considered pass ("purity") statistics.

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## get\_cos\_purity

median\_abs\_corr

The median absolute correlation value of variants in a CoS to be considered pass ("purity") statistics. When the number of variables included in the CoS is greater than this number, the CoS variables are randomly subsampled.

## Value

A list of indices of variables in each CoS.

## See Also

Other colocboost\_utilities: get\_cormat(), get\_cos\_purity(), get\_cos\_summary(), get\_hierarchical\_clusters(),
get\_ucos\_summary()

# Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 3
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5 # SNP10 affects trait 1</pre>
true_beta[10, 2] <- 0.4 # SNP10 also affects trait 2 (colocalized)</pre>
true_beta[50, 2] <- 0.3 # SNP50 only affects trait 2</pre>
true_beta[80, 3] <- 0.6 # SNP80 only affects trait 3</pre>
Y \leq matrix(0, N, L)
for (1 in 1:L) {
  Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)</pre>
}
res <- colocboost(X = X, Y = Y)
get_cos(res, coverage = 0.99, X = X)
get_cos(res, coverage = 0.99, X = X, min_abs_corr = 0.95)
```

<pre>get_cos_purity</pre>	Calculate J	ourity within a	and in-between CoS
<u> </u>			

## Description

Calculate purity statistics between all pairs of colocalization confidence sets (CoS)

## Usage

```
get_cos_purity(cos, X = NULL, Xcorr = NULL, n_purity = 100)
```

## Arguments

cos	List of variables in CoS
Х	Genotype matrix of values of the p variables. Used to compute correlations if Xcorr is not provided.
Xcorr	Correlation matrix of correlations between variables. Alternative to X.
n_purity	The maximum number of CoS variables used in calculating the correlation ("purity") statistics. When the number of variables included in the CoS is greater than this number, the CoS variables are randomly subsampled.

## Value

A list containing three matrices (min\_abs\_cor, max\_abs\_cor, median\_abs\_cor) with purity statistics for all pairs of CoS. Diagonal elements represent within-CoS purity.

# See Also

Other colocboost\_utilities: get\_cormat(), get\_cos(), get\_cos\_summary(), get\_hierarchical\_clusters(),
get\_ucos\_summary()

## Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 3
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5
true_beta[10, 2] <- 0.4
true_beta[50, 2] <- 0.3
true_beta[80, 3] <- 0.6
Y <- matrix(0, N, L)
for (1 in 1:L) {
  Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)</pre>
}
res <- colocboost(X = X, Y = Y)</pre>
cos_res <- get_cos(res, coverage = 0.8)</pre>
get_cos_purity(cos_res$cos, X = X)
```

get\_cos\_summary

# Description

get\_cos\_summary get the colocalization summary table with or without the outcomes of interest.

# Usage

```
get_cos_summary(
   cb_output,
   outcome_names = NULL,
   interest_outcome = NULL,
   region_name = NULL
)
```

# Arguments

cb_output	Output object from colocboost analysis
outcome_names	Optional vector of names of outcomes, which has the same order as Y in the original analysis.
interest_outcom	e
	Optional vector specifying a subset of outcomes from outcome_names to focus on. When provided, only colocalization events that include at least one of these outcomes will be returned.
region_name	Optional character string. When provided, adds a column with this gene name to the output table for easier filtering in downstream analyses.

# Value

A summary table for colocalization events with the following columns:

focal_outcome	The focal outcome being analyzed if exists. Otherwise, it is FALSE.	
colocalized_outcomes		
	Colocalized outcomes for colocalization confidence set (CoS)	
cos_id	Unique identifier for colocalization confidence set (CoS)	
purity	Minimum absolute correlation of variables with in colocalization confidence set (CoS)	
top_variable	The variable with highest variant colocalization probability (VCP)	
top_variable_vcp		
	Variant colocalization probability for the top variable	
cos_npc	Normalized probability of colocalization	
min_npc_outcome		
	Minimum normalized probability of colocalized traits	

n_variables	Number of variables in colocalization confidence set (CoS)
colocalized_ind	lex
	Indices of colocalized variables
colocalized_var	iables
	List of colocalized variables
colocalized_var	iables_vcp
	Variant colocalization probabilities for all colocalized variables

## Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ articles/Interpret\_ColocBoost\_Output.html

# See Also

Other colocboost\_utilities: get\_cormat(), get\_cos(), get\_cos\_purity(), get\_hierarchical\_clusters(),
get\_ucos\_summary()

# Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 3
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5 # SNP10 affects trait 1</pre>
true_beta[10, 2] <- 0.4 # SNP10 also affects trait 2 (colocalized)</pre>
true_beta[50, 2] <- 0.3 # SNP50 only affects trait 2</pre>
true_beta[80, 3] <- 0.6 # SNP80 only affects trait 3</pre>
Y \leq matrix(0, N, L)
for (1 in 1:L) {
  Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)</pre>
}
res <- colocboost(X = X, Y = Y)</pre>
get_cos_summary(res)
```

get\_hierarchical\_clusters

Perform modularity-based hierarchical clustering for a correlation matrix

# Description

This function performs a modularity-based hierarchical clustering approach to identify clusters from a correlation matrix.

## Usage

```
get_hierarchical_clusters(cormat, min_cluster_corr = 0.8)
```

# Arguments

cormat	A correlation matrix.
<pre>min_cluster_cc</pre>	orr

The small correlation for the weights distributions across different iterations to be decided having only one cluster. Default is 0.8.

# Value

A list containing:

cluster	A binary matrix indicating the cluster membership of each variable.
Q_modularity	The modularity values for the identified clusters.

# See Also

Other colocboost\_utilities: get\_cormat(), get\_cos(), get\_cos\_purity(), get\_cos\_summary(), get\_ucos\_summary()

# Examples

```
# Example usage
set.seed(1)
N <- 100
P <- 4
sigma <- matrix(0.2, nrow = P, ncol = P)
diag(sigma) <- 1
sigma[1:2, 1:2] <- 0.9
sigma[3:4, 3:4] <- 0.9
X <- MASS::mvrnorm(N, rep(0, P), sigma)
cormat <- get_cormat(X)
clusters <- get_hierarchical_clusters(cormat)
clusters$cluster
clusters$Q_modularity
```

```
get_robust_colocalization
```

Recalibrate and summarize robust colocalization events.

# Description

get\_robust\_colocalization get the colocalization by discarding the weaker colocalization events or colocalized outcomes

# Usage

```
get_robust_colocalization(
   cb_output,
   cos_npc_cutoff = 0.5,
   npc_outcome_cutoff = 0.2,
   pvalue_cutoff = NULL,
   weight_fudge_factor = 1.5,
   coverage = 0.95
)
```

# Arguments

cb_output	Output object from colocboost analysis	
cos_npc_cutoff	Minimum threshold of normalized probability of colocalization (NPC) for CoS.	
npc_outcome_cutoff		
	Minimum threshold of normalized probability of colocalized traits in each CoS.	
pvalue_cutoff	Maximum threshold of marginal p-values of colocalized variants on colocalized traits in each CoS.	
weight_fudge_factor		
	The strength to integrate weight from different outcomes, default is 1.5	
coverage	A number between 0 and 1 specifying the "coverage" of the estimated colocal- ization confidence sets (CoS) (default is 0.95).	

# Value

A "colocboost" object with some or all of the following elements:

cos_summary	A summary table for colocalization events.
vcp	The variable colocalized probability for each variable.
cos_details	A object with all information for colocalization results.
data_info	A object with detailed information from input data
model_info	A object with detailed information for colocboost model
ucos_from_cos	A object with information for trait-specific effects if exists after removing weaker signals.

## Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ articles/Interpret\_ColocBoost\_Output.html

# See Also

Other colocboost\_inference: get\_ambiguous\_colocalization(), get\_colocboost\_summary()

## Examples

```
# colocboost example
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 3
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5 # SNP10 affects trait 1</pre>
true_beta[10, 2] <- 0.4 # SNP10 also affects trait 2 (colocalized)</pre>
true_beta[50, 2] <- 0.3 # SNP50 only affects trait 2</pre>
true_beta[80, 3] <- 0.6 # SNP80 only affects trait 3</pre>
Y \leq matrix(0, N, L)
for (1 in 1:L) {
  Y[, 1] <- X %*% true_beta[, 1] + rnorm(N, 0, 1)</pre>
}
res <- colocboost(X = X, Y = Y)</pre>
res$cos_details$cos$cos_index
filter_res <- get_robust_colocalization(res, cos_npc_cutoff = 0.5, npc_outcome_cutoff = 0.2)</pre>
filter_res$cos_details$cos$cos_index
```

get\_ucos\_summary Get trait-specific summary table from a ColocBoost output.

## Description

get\_ucos\_summary produces a trait-specific summary table for uncolocalized (single-trait) associations from ColocBoost results. This is particularly useful for examining trait-specific signals or for summarizing results from single-trait FineBoost analyses.

## Usage

```
get_ucos_summary(
    cb_output,
    outcome_names = NULL,
    region_name = NULL,
```

```
ambiguous_cos = FALSE,
min_abs_corr_between_ucos = 0.5,
median_abs_corr_between_ucos = 0.8
)
```

## Arguments

cb_output	Output object from colocboost analysis	
outcome_names	Optional vector of names of outcomes, which has the same order as Y in the original analysis.	
region_name	Optional character string. When provided, adds a column with this gene name to the output table for easier filtering in downstream analyses.	
ambiguous_cos	Logical indicating whether to include ambiguous colocalization events. The default is FALSE.	
<pre>min_abs_corr_between_ucos</pre>		
	Minimum absolute correlation for variants across two trait-specific (uncolocal- ized) effects to be considered colocalized. The default is 0.5.	
<pre>median_abs_corr_between_ucos</pre>		
	Median absolute correlation for variants across two trait-specific (uncolocalized) effects to be considered colocalized. The default is 0.8.	

# Value

A list containing:

- ucos\_summary: A summary table for trait-specific, uncolocalized associations with the following columns:
  - outcomes: Outcome being analyzed
  - ucos\_id: Unique identifier for trait-specific confidence sets
  - purity: Minimum absolute correlation of variables within trait-specific confidence sets
  - top\_variable: The variable with highest variant-level probability of association (VPA)
  - top\_variable\_vpa: Variant-level probability of association (VPA) for the top variable
  - ucos\_npc: Normalized probability of causal association for the trait-specific confidence set
  - n\_variables: Number of variables in trait-specific confidence set
  - ucos\_index: Indices of variables in the trait-specific confidence set
  - ucos\_variables: List of variables in the trait-specific confidence set
  - ucos\_variables\_vpa: Variant-level probability of association (VPA) for all variables in the confidence set
  - region\_name: Region name if provided through the region\_name parameter
- ambiguous\_cos\_summary: A summary table for ambiguous colocalization events with the following columns:
  - outcomes: Outcome in the ambiguous colocalization event
  - ucos\_id: Unique identifiers for the ambiguous event
  - min\_between\_purity: Minimum absolute correlation between variables across traitspecific sets in the ambiguous event

- median\_between\_purity: Median absolute correlation between variables across traitspecific sets in the ambiguous event
- overlap\_idx: Indices of variables that overlap between ambiguous trait-specific sets
- overlap\_variables: Names of variables that overlap between ambiguous trait-specific sets
- n\_recalibrated\_variables: Number of variables in the recalibrated colocalization set from an ambiguous event
- recalibrated\_index: Indices of variables in the recalibrated colocalization set from an ambiguous event
- recalibrated\_variables: Names of variables in the recalibrated colocalization set from an ambiguous event
- recalibrated\_variables\_vcp: Variant colocalization probabilities for recalibrated variables from an ambiguous event
- region\_name: Region name if provided through the region\_name parameter

## Source

See detailed instructions in our tutorial portal: https://statfungen.github.io/colocboost/ articles/Interpret\_ColocBoost\_Output.html

## See Also

```
Other colocboost_utilities: get_cormat(), get_cos(), get_cos_purity(), get_cos_summary(),
get_hierarchical_clusters()
```

# Examples

```
# colocboost example with single trait analysis
set.seed(1)
N <- 1000
P <- 100
# Generate X with LD structure
sigma <- 0.9^abs(outer(1:P, 1:P, "-"))</pre>
X <- MASS::mvrnorm(N, rep(0, P), sigma)</pre>
colnames(X) <- paste0("SNP", 1:P)</pre>
L <- 1 # Only one trait for single-trait analysis
true_beta <- matrix(0, P, L)</pre>
true_beta[10, 1] <- 0.5 # SNP10 affects the trait</pre>
true_beta[80, 1] <- 0.2 # SNP11 also affects the trait but with lower effect
Y <- X %*% true_beta + rnorm(N, 0, 1)</pre>
res <- colocboost(X = X, Y = Y, output_level = 2)</pre>
# Get the trait-specifc effect summary
get_ucos_summary(res)
```

Heterogeneous\_Effect

## Description

An example dataset with simulated genotypes and traits for 2 traits and 2 common causal variants with heterogeneous effects

## Usage

Heterogeneous\_Effect

## Format

Heterogeneous\_Effect: A list with 3 elements X List of genotype matrices Y List of traits

variant indices of two causal variants

## Source

The Heterogeneous\_Effect dataset contains 2 simulated phenotypes alongside corresponding genotype matrices. There are two causal variants, both of which have heterogeneous effects on two traits. Due to the file size limitation of CRAN release, this is a subset of simulated data to generate Figure 2b in Cao etc. 2025. See full dataset in colocboost paper repo https://github.com/StatFunGen/ colocboost-paper.

# See Also

Other colocboost\_data: Ambiguous\_Colocalization, Ind\_5traits, Non\_Causal\_Strongest\_Marginal, Sumstat\_5traits, Weaker\_GWAS\_Effect

Ind\_5traits Individual level data for 5 traits

## Description

An example dataset with simulated genotypes and traits for 5 traits

## Usage

Ind\_5traits

# Format

Ind\_5traits:
A list with 3 elements
X List of genotype matrices
Y List of traits
true\_effect\_variants List of causal variants

## Source

The Ind\_5traits dataset contains 5 simulated phenotypes alongside corresponding genotype matrices. The dataset is specifically designed for evaluating and demonstrating the capabilities of ColocBoost in multi-trait colocalization analysis with individual-level data. See Cao etc. 2025 for details. Due to the file size limitation of CRAN release, this is a subset of simulated data. See full dataset in colocboost paper repo https://github.com/StatFunGen/colocboost-paper.

## See Also

Other colocboost\_data: Ambiguous\_Colocalization, Heterogeneous\_Effect, Non\_Causal\_Strongest\_Marginal, Sumstat\_5traits, Weaker\_GWAS\_Effect

Non\_Causal\_Strongest\_Marginal

Individual level data for 2 traits and 2 causal variants, but the strongest marginal association is not causal

## Description

An example dataset with simulated genotypes and traits for 2 traits and 2 common causal variants, but the strongest marginal association is not causal variant.

# Usage

Non\_Causal\_Strongest\_Marginal

## Format

Non\_Causal\_Strongest\_Marginal:A list with 3 elementsX List of genotype matricesY List of traitsvariant indices of two causal variants

## Source

The Non\_Causal\_Strongest\_Marginal dataset contains 2 simulated phenotypes alongside corresponding genotype matrices. There are two causal variants, but the strongest marginal association is not a causal variant. Due to the file size limitation of CRAN release, this is a subset of simulated data to generate Figure 2b in Cao etc. 2025. See full dataset in colocboost paper repo https://github.com/StatFunGen/colocboost-paper.

## See Also

Other colocboost\_data: Ambiguous\_Colocalization, Heterogeneous\_Effect, Ind\_5traits, Sumstat\_5traits, Weaker\_GWAS\_Effect

Sumstat\_Straits Summary level data for 5 traits

## Description

An example dataset with simulated statistics for 5 traits

## Usage

Sumstat\_5traits

## Format

Sumstat\_5traits:

A list with 2 elements

sumstat Summary statistics for 5 traits
true\_effect\_variants List of causal variants

## Source

The Sumstat\_5traits dataset contains 5 simulated summary statistics, where it is directly derived from the Ind\_5traits dataset using marginal association. The dataset is specifically designed for evaluating and demonstrating the capabilities of ColocBoost in multi-trait colocalization analysis with summary association data. See Cao etc. 2025 for details. Due to the file size limitation of CRAN release, this is a subset of simulated data. See full dataset in colocboost paper repo https://github.com/StatFunGen/colocboost-paper.

# See Also

Other colocboost\_data: Ambiguous\_Colocalization, Heterogeneous\_Effect, Ind\_5traits, Non\_Causal\_Strongest\_M. Weaker\_GWAS\_Effect

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Weaker\_GWAS\_Effect

Individual level data for 2 traits and 2 causal variants with weaker effects for focal trait

## Description

An example dataset with simulated genotypes and traits for 2 traits and 2 common causal variants with heterogeneous effects

## Usage

Weaker\_GWAS\_Effect

# Format

Weaker\_GWAS\_Effect:A list with 3 elementsX List of genotype matricesY List of traitsvariant indices of two causal variants

## Source

The Weaker\_GWAS\_Effect dataset contains 2 simulated phenotypes alongside corresponding genotype matrices. There are two causal variants, one of which has a weaker effect on the focal trait compared to the other trait. Due to the file size limitation of CRAN release, this is a subset of simulated data to generate Figure 2b in Cao etc. 2025. See full dataset in colocboost paper repo https://github.com/StatFunGen/colocboost-paper.

## See Also

Other colocboost\_data: Ambiguous\_Colocalization, Heterogeneous\_Effect, Ind\_5traits, Non\_Causal\_Strongest\_Massat\_5traits

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