

# Package ‘SVEMnet’

June 3, 2026

**Type** Package

**Title** Self-Validated Ensemble Models with Lasso and Relaxed Elastic Net Regression

**Version** 3.2.1

**Date** 2026-06-02

**Description** Tools for fitting self-validated ensemble models (SVEM; Lemkus et al. (2021) <[doi:10.1016/j.chemolab.2021.104439](https://doi.org/10.1016/j.chemolab.2021.104439)>) in small-sample design-of-experiments and related workflows, using elastic net and relaxed elastic net regression via 'glmnet' (Friedman et al. (2010) <[doi:10.18637/jss.v033.i01](https://doi.org/10.18637/jss.v033.i01)>). Fractional random-weight bootstraps with anti-correlated validation copies are used to tune penalty paths by validation-weighted AIC/BIC. Supports Gaussian and binomial responses, deterministic expansion helpers for shared factor spaces, prediction with bootstrap uncertainty, and a random-search optimizer that respects mixture constraints and combines multiple responses via desirability functions. Also includes a permutation-based whole-model test for Gaussian SVEM fits (Karl (2024) <[doi:10.1016/j.chemolab.2024.105122](https://doi.org/10.1016/j.chemolab.2024.105122)>). Package code was drafted with assistance from generative AI tools.

**URL** <https://arxiv.org/abs/2511.20968>

**Depends** R (>= 4.0.0)

**Imports** glmnet (>= 4.1-6), stats, cluster, ggplot2, lhs, foreach, doParallel, parallel, gamlss, gamlss.dist, utils

**Suggests** covr, knitr, rmarkdown, testthat (>= 3.0.0), withr, vdiff, RhpcBLASctl

**License** GPL-2 | GPL-3

**Encoding** UTF-8

**Config/testthat/edition** 3

**LazyData** true

**Config/roxygen2/version** 8.0.0

**NeedsCompilation** no

**Author** Andrew T. Karl [cre, aut] (ORCID: <<https://orcid.org/0000-0002-5933-8706>>)

**Maintainer** Andrew T. Karl <akar1@asu.edu>

**Repository** CRAN

**Date/Publication** 2026-06-02 22:20:02 UTC

## Contents

SVEMnet-package . . . . .	2
bigexp_formula . . . . .	6
bigexp_prepare . . . . .	7
bigexp_terms . . . . .	9
bigexp_train . . . . .	14
coef.svem_model . . . . .	15
glmnet_with_cv . . . . .	16
lipid_screen . . . . .	22
plot.svem_binomial . . . . .	28
plot.svem_model . . . . .	30
plot.svem_significance_test . . . . .	31
predict.svem_model . . . . .	32
predict_cv . . . . .	35
print.bigexp_spec . . . . .	37
print.svem_significance_test . . . . .	39
SVEMnet . . . . .	39
svem_export_candidates_csv . . . . .	48
svem_nonzero . . . . .	51
svem_random_table_multi . . . . .	53
svem_score_random . . . . .	56
svem_select_from_score_table . . . . .	64
svem_significance_test_parallel . . . . .	65
svem_wmt_multi . . . . .	71
with_bigexp_contrasts . . . . .	73
<b>Index</b>	<b>75</b>

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SVEMnet-package	<i>SVEMnet: Self-Validated Ensemble Models with Relaxed Lasso and Elastic-Net Regression</i>
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## Description

The SVEMnet package implements Self-Validated Ensemble Models (SVEM) using Elastic Net (including lasso and ridge) regression via `glmnet`. SVEM averages predictions from multiple models fitted to fractionally weighted bootstraps of the data, tuned with anti-correlated validation weights. The package supports multi-response optimization with uncertainty-aware candidate generation for iterative formulation and process development.

## Details

A typical workflow is:

1. Build a wide, deterministic factor expansion (optionally via `bigexp_terms`) and reuse it across responses with `bigexp_formula`.
2. Fit one or more SVEM models with `SVEMnet`.
3. Optionally run whole-model testing via `svem_significance_test_parallel` (and `svem_wmt_multi`) to assess factor relationships or reweight response goals.
4. Call `svem_score_random` to draw random points in the factor space, compute multi-response Derringer–Suich scores, optional WMT-reweighted scores, and an uncertainty measure; then use `svem_select_from_score_table` to pick a single "best" row and diverse medoid candidates, and `svem_export_candidates_csv` to export candidate tables for the next experimental round.
5. Run new experiments at the suggested candidates, append the data, refit the models, and repeat as needed (closed-loop optimization).

## Core modeling and summaries

`SVEMnet` Fit an SVEMnet model using Elastic Net regression (including relaxed elastic net) on fractionally weighted bootstraps.

`predict.svem_model` Predict method for SVEM models (ensemble-mean aggregation by default, optional debiasing, and percentile prediction intervals when available).

`coef.svem_model` Averaged (optionally debiased) coefficients from an SVEM model.

`svem_nonzero` Bootstrap nonzero percentages for each coefficient, with an optional quick plot.

`plot.svem_model` Quick actual-versus-predicted plot for a fitted model (with optional group colorings).

## Deterministic wide expansions (bigexp helpers)

The `bigexp_*` helpers build and reuse a locked polynomial/interaction expansion across multiple responses and datasets:

`bigexp_terms` Build a deterministic expanded RHS (polynomials, interactions, optional partial-cubic terms) with locked factor levels and numeric ranges.

`bigexp_prepare` Coerce new data to match a stored `bigexp_spec`, including factor levels and numeric types.

`bigexp_formula` Reuse a locked expansion for another response to ensure an identical factor space across models.

`with_bigexp_contrasts` Temporarily restore the contrast options used when a `bigexp_spec` was built.

`bigexp_train` Convenience wrapper that builds a `bigexp_spec` and prepares training data in one call.

### Random tables, optimization, and candidate generation

- `svem_random_table_multi` Generate one shared random predictor table (with optional mixture constraints) from cached factor-space information and obtain predictions from multiple SVEM models at those points. Supports both Gaussian and binomial models; binomial predictions are returned on the probability scale. This is the lower-level sampler used by `svem_score_random`.
- `svem_score_random` Random-search scoring for multiple responses with Derringer–Suich desirabilities, user weights, optional whole-model-test (WMT) reweighting, percentile CI-based uncertainty, and (optionally) scoring of existing experimental data. Returns a scored random-search table and, when data is supplied, an augmented copy of the original data with `<resp>_pred`, desirabilities, scores, and an `uncertainty_measure`.
- `svem_select_from_score_table` Given a scored table (typically `svem_score_random()`\$`score_table`), select one "best" row under a chosen objective and a small, diverse set of medoid candidates via PAM clustering on predictors.
- `svem_export_candidates_csv` Concatenate one or more selection objects from `svem_select_from_score_table` and export candidate tables (with metadata, predictions, and optional design-only trimming) to CSV or return them in-memory for inspection.

### Whole-model testing and plotting

- `svem_significance_test_parallel` Parallel whole-model significance test (using `foreach + doParallel`) with support for mixture-constrained sampling and reuse of a locked `bigexp_spec`. Designed for continuous (Gaussian) responses.
- `svem_wmt_multi` Helper to run `svem_significance_test_parallel` across multiple responses and construct whole-model p-values and reweighting multipliers for use in `svem_score_random`.
- `plot.svem_significance_test` Plot helper for visualizing multiple significance-test outputs (observed vs permutation distances, fitted null, and p-values).

### Auxiliary utilities and data

- `glmnet_with_cv` Convenience wrapper around repeated `cv.glmnet()` selection for robust lambda (and optional alpha) choice.
- `lipid_screen` Example dataset for multi-response modeling, whole-model testing, and mixture-constrained optimization demonstrations.

### Families

SVEMnet currently supports:

- Gaussian responses (`family = "gaussian"`) with identity link and optional debiasing / percentile prediction intervals.
- Binomial responses (`family = "binomial"`) with logit link. The response must be 0/1 numeric or a two-level factor (first level treated as 0). Use `predict(..., type = "response")` for event probabilities or `type = "class"` for 0/1 labels (threshold = 0.5 by default).

Some higher-level utilities place additional constraints:

- `svem_significance_test_parallel` is designed and interpreted for continuous (Gaussian) responses.

- `svem_score_random` supports mixed Gaussian + binomial response sets, treating binomial predictions and CIs on the probability scale, but WMT-based goal reweighting (via `svem_wmt_multi` and the `wmt` argument) is only allowed when all responses are Gaussian.

## Acknowledgments

OpenAI's GPT models (o1-preview through GPT-5 Pro) were used to assist with coding and roxygen documentation; all content was reviewed and finalized by the author.

## Author(s)

**Maintainer:** Andrew T. Karl <akar1@asu.edu> (ORCID)

Authors:

- Andrew T. Karl <akar1@asu.edu> (ORCID)

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### See Also

Useful links:

- <https://arxiv.org/abs/2511.20968>

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bigexp_formula	<i>Construct a formula for a new response using a bigexp_spec</i>
----------------	---

---

### Description

bigexp\_formula() lets you reuse an existing expansion spec for multiple responses. It keeps the right hand side locked but changes the response variable on the left hand side.

### Usage

```
bigexp_formula(spec, response)
```

### Arguments

spec	A "bigexp_spec" object created by bigexp_terms().
response	Character scalar giving the name of the new response column in your data. If omitted, the original formula is returned unchanged.

### Details

This is useful when you want to fit separate models for several responses on the same factor space while guaranteeing that they all use exactly the same design columns and coding.

### Value

A formula of the form response ~ rhs, where the right-hand side is taken from the locked expansion stored in spec.

## Examples

```
set.seed(1)
df2 <- data.frame(
  y1 = rnorm(10),
  y2 = rnorm(10),
  X1 = rnorm(10),
  X2 = rnorm(10)
)

spec2 <- bigexp_terms(
  y1 ~ X1 + X2,
  data      = df2,
  factorial_order = 2,
  polynomial_order = 2
)

f2 <- bigexp_formula(spec2, "y2")
f2
```

---

bigexp\_prepare

*Prepare data to match a bigexp\_spec*

---

## Description

`bigexp_prepare()` coerces a new data frame so that it matches a previously built `bigexp_terms` spec. It:

- applies the locked factor levels for categorical predictors,
- enforces that continuous variables remain numeric (and errors if they are not), and
- optionally warns about or errors on unseen factor levels.

## Usage

```
bigexp_prepare(spec, data, unseen = c("warn_na", "error"))
```

## Arguments

spec	Object returned by <code>bigexp_terms</code> .
data	New data frame (for example, training, test, or future batches).
unseen	How to handle unseen factor levels in data: "warn_na" (default) maps unseen levels to NA and issues a warning, or "error" stops with an error if any unseen levels are encountered.

## Details

Columns that are not listed in `spec$vars` (for example, the response or extra metadata columns) are left unchanged.

The goal is that `model.matrix(spec$formula, data)` will produce the same set of columns in the same order across all datasets prepared with the same spec, even if some levels are missing in a particular batch.

## Value

A list with two elements:

- `formula`: the expanded formula stored in the spec (same as `spec$formula`).
- `data`: a copy of the input data with predictor columns coerced to match the spec (types and levels), suitable for `model.frame()` / `model.matrix()`.

## See Also

[bigexp\\_terms](#)

## Examples

```
set.seed(1)
train <- data.frame(
  y = rnorm(10),
  X1 = rnorm(10),
  X2 = rnorm(10),
  G = factor(sample(c("A", "B"), 10, replace = TRUE))
)

spec <- bigexp_terms(
  y ~ X1 + X2 + G,
  data = train,
  factorial_order = 2,
  polynomial_order = 2
)

newdata <- data.frame(
  y = rnorm(5),
  X1 = rnorm(5),
  X2 = rnorm(5),
  G = factor(sample(c("A", "B"), 5, replace = TRUE))
)

prep <- bigexp_prepare(spec, newdata)
str(prepare$data)
```

---

bigexp_terms	<i>Create a deterministic expansion spec for wide polynomial and interaction models</i>
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---

## Description

bigexp\_terms() builds a specification object that:

- decides which predictors are treated as continuous or categorical,
- optionally treats selected variables as blocking factors that enter the model only additively and never in interactions or polynomials,
- locks factor levels from the supplied data,
- records the contrast settings used when the model matrix is first built, and
- constructs a reusable right-hand side (RHS) expression string for a large expansion that can be shared across multiple responses and datasets.

## Usage

```
bigexp_terms(
  formula,
  data,
  factorial_order = 3L,
  polynomial_order = 3L,
  include_pc_2way = TRUE,
  include_pc_3way = FALSE,
  intercept = TRUE,
  blocking = NULL,
  discrete_numeric = NULL,
  audit = c("warn", "error", "none"),
  audit_numeric_rate = 0.9,
  audit_unique_ratio = 0.8,
  audit_min_n = 12L,
  report = TRUE
)
```

## Arguments

formula	Main-effects formula of the form $y \sim X1 + X2 + G$ or $y \sim ..$ . The right-hand side should contain main effects only; do not include : (interactions), ^ (factorial shortcuts), I() powers, or inline polynomial expansions. The helper will generate interactions and polynomial terms automatically.
data	Data frame used to decide types and lock factor levels.
factorial_order	Integer $\geq 1$ . Maximum order of factorial interactions among the non-blocking main effects. For example, 1 gives main effects only, 2 gives up to two-way interactions, 3 gives up to three-way interactions, and so on.

polynomial_order	Integer $\geq 1$ . Maximum polynomial degree for continuous non-blocking predictors. A value of 1 means only linear terms; 2 adds squares $I(X^2)$ ; 3 adds cubes $I(X^3)$ ; in general, all powers $I(X^k)$ for $k$ from 2 up to <code>polynomial_order</code> are added.
include_pc_2way	Logical. If TRUE (default) and <code>polynomial_order</code> $\geq 2$ , include partial-cubic two-way terms of the form $Z:I(X^2)$ where $X$ is continuous and $Z$ is another non-blocking predictor.
include_pc_3way	Logical. If TRUE and <code>polynomial_order</code> $\geq 2$ , include partial-cubic three-way terms $I(X^2):Z:W$ among non-blocking predictors.
intercept	Logical. If TRUE (default), include an intercept in the expansion; if FALSE, the generated RHS drops the intercept.
blocking	Optional character vector of column names in data to treat as blocking factors. These variables are included in the spec and typed like other predictors (categorical vs continuous), but they enter the model only as additive main effects and never appear in interactions, polynomials, or partial-cubic terms. <b>Important:</b> when using $y \sim .$ , blocking variables are automatically excluded from the "non-blocking" predictor set so they do not trigger a conflict error. When using an explicit RHS (for example $y \sim X1 + X2$ ), blocking variables must not also be explicitly listed on the right-hand side.
discrete_numeric	Optional specification of "discrete numeric" predictors for downstream sampling (for example in <code>svem_random_table_multi()</code> ). These predictors are still treated as numeric for modeling and expansion (that is, they remain continuous in the design matrix and may participate in polynomial and interaction terms). This option only records a finite set of preferred numeric levels to be used when randomly generating recipes. Supply either: <ul style="list-style-type: none"> <li>• a character vector of predictor names, in which case the allowed levels are inferred as the sorted unique finite values observed in data; or</li> <li>• a named list mapping predictor names to numeric vectors of allowed levels. If an entry is NULL or length zero, levels are inferred from data for that predictor.</li> </ul>
audit	How to handle suspicious typing / high-cardinality issues when building the spec. One of "warn" (default), "error", or "none". Audits cover numeric-like character/factor columns (including percent strings like "25%"), and very high-cardinality categorical predictors that are likely IDs or mis-typed numerics.
audit_numeric_rate	Numeric in (0,1). If at least this fraction of non-missing values parse as numeric (after stripping commas and an optional trailing %), the column is flagged as numeric-like when stored as character/factor.
audit_unique_ratio	Numeric in (0, 1). For categorical predictors, warn/error if <code>unique(non-missing) / n_nonmissing</code> $\geq$ <code>audit_unique_ratio</code> .
audit_min_n	Integer $\geq 1$ . Minimum number of non-missing values required before audits are applied.

report Logical. If TRUE (default), print a compact summary of the inferred predictor types and settings (via `print.bigexp_spec`) when `bigexp_terms()` returns.

## Details

The expansion for non-blocking predictors can include:

- full factorial interactions among the listed main effects, up to a chosen order;
- polynomial terms  $I(X^k)$  for continuous predictors up to a chosen degree; and
- optional partial-cubic interactions of the form  $Z: I(X^2)$  and  $I(X^2): Z: W$ .

Predictor types are inferred from data:

- factors, characters, and logicals are treated as categorical;
- all other numeric predictors are treated as continuous, and their observed ranges are stored.

Variables listed in `blocking` are included in the `spec` and are typed using the same rules as other predictors (for example, a numeric blocking variable with many distinct values is treated as continuous). However, blocking variables enter the model only as additive main effects, without interactions or polynomial terms, regardless of `factorial_order` or `polynomial_order`.

Once built, a "bigexp\_spec" can be reused to create consistent expansions for new datasets via `bigexp_prepare` and `bigexp_formula`. The RHS and contrast settings are locked, so the same spec applied to different data produces design matrices with the same columns in the same order (up to missing levels for specific batches).

## Value

An object of class "bigexp\_spec" with components:

- `formula`: expanded formula of the form  $y \sim \langle \text{big expansion} \rangle$ , using the response from the input formula.
- `rhs`: right-hand side expansion string (reusable for any response).
- `vars`: character vector of predictor names (including blocking variables) in the order discovered from the formula and data.
- `is_cat`: named logical vector indicating which predictors are treated as categorical (TRUE) versus continuous (FALSE).
- `levels`: list of locked factor levels for categorical predictors.
- `num_range`:  $2 \times p$  numeric matrix of ranges for continuous variables (rows `c("min", "max")`).
- `settings`: list of expansion settings, including `factorial_order`, `polynomial_order`, `include_pc_2way`, `include_pc_3way`, `intercept`, `blocking`, and stored contrast information.

## Typical workflow

In a typical multi-response workflow you:

1. Call `bigexp_terms()` once on your training data to build and lock the expansion (types, levels, contrasts, RHS).

2. Fit models using `spec$formula` and the original data (for example, `SVMnet(spec$formula, data, ...)` or `lm(spec$formula, data)`).
3. For new batches, call `bigexp_prepare` with the same `spec` so that design matrices have exactly the same columns and coding.
4. For additional responses on the same factor space, use `bigexp_formula` to swap the left-hand side while reusing the locked expansion.

### See Also

[bigexp\\_prepare](#), [bigexp\\_formula](#), [bigexp\\_train](#)

### Examples

```
## Example 1: small design with one factor
set.seed(1)
df <- data.frame(
  y = rnorm(20),
  X1 = rnorm(20),
  X2 = rnorm(20),
  G = factor(sample(c("A", "B"), 20, replace = TRUE))
)

## Two-way interactions and up to cubic terms in X1 and X2
spec <- bigexp_terms(
  y ~ X1 + X2 + G,
  data = df,
  factorial_order = 2,
  polynomial_order = 3
)

print(spec)

## Example 2: pure main effects (no interactions, no polynomial terms)
spec_main <- bigexp_terms(
  y ~ X1 + X2 + G,
  data = df,
  factorial_order = 1, # main effects only
  polynomial_order = 1 # no I(X^2) or higher
)

## Example 3: blocking factors (categorical and continuous)
set.seed(2)
df_block <- data.frame(
  y = rnorm(30),
  X1 = rnorm(30),
  X2 = rnorm(30),
  G = factor(sample(c("A", "B"), 30, replace = TRUE)),
  Operator = factor(sample(paste0("Op", 1:3), 30, replace = TRUE)),
  AmbientTemp = rnorm(30, mean = 22, sd = 2) # continuous blocking covariate
)

## Here Operator (categorical) and AmbientTemp (continuous) are blocking factors:
```

```

## they enter additively, but do not appear in interactions or polynomials.
spec_block <- bigexp_terms(
  y ~ X1 + X2 + G,
  data      = df_block,
  factorial_order = 2,
  polynomial_order = 3,
  blocking   = c("Operator", "AmbientTemp")
)

print(spec_block)
spec_block$rhs

## Example 4: discrete numeric predictors (finite numeric support)
## A common case is a numeric process setting that only takes a small set
## of allowed values (e.g., 0.5, 1, 2, 4). Use `discrete_numeric` in
## bigexp_terms() so downstream sampling respects those levels automatically.

set.seed(3)
D_allowed <- c(0.5, 1, 2, 4)
df_disc <- data.frame(
  y = rnorm(60),
  D = sample(D_allowed, 60, replace = TRUE), # numeric with discrete support
  X1 = rnorm(60),
  G = factor(sample(c("A", "B"), 60, replace = TRUE))
)

# Record that D should be treated as "discrete numeric" for downstream sampling.
# Levels are inferred automatically from the training data.
spec_disc <- bigexp_terms(
  y ~ D + X1 + G,
  data      = df_disc,
  factorial_order = 2,
  polynomial_order = 2,
  discrete_numeric = "D"
)

# Fit. The discrete support is expected to propagate into fit$sampling_schema
# (assuming the updated SVEMnet implementation that stores sampling_schema).
fit_disc <- SVEMnet(spec_disc, df_disc, nBoot = 20)

# Score random candidates; sampled D values stay in D_allowed
scored <- svem_score_random(
  objects      = list(y = fit_disc),
  goals       = list(y = list(goal = "max", weight = 1)),
  n           = 2000,
  numeric_sampler = "random",
  verbose     = FALSE
)

table(scored$score_table$D)
stopifnot(all(scored$score_table$D %in% D_allowed))

```

---

bigexp\_train

*Build a spec and prepare training data in one call*


---

### Description

bigexp\_train() is a convenience wrapper around `bigexp_terms` and `bigexp_prepare`. It:

- builds a deterministic expansion spec from the training data; and
- immediately prepares that same data to match the locked types and levels.

### Usage

```
bigexp_train(formula, data, ...)
```

### Arguments

formula	Main-effects formula such as $y \sim X1 + X2 + G$ or $y \sim ..$ . Only main effects should appear on the right hand side.
data	Training data frame used to lock types and levels.
...	Additional arguments forwarded to <code>bigexp_terms()</code> , such as <code>factorial_order</code> , <code>polynomial_order</code> , <code>include_pc_2way</code> , <code>include_pc_3way</code> , and <code>intercept</code> .

### Details

This is handy when you want a single object that contains both the spec and the training data in a form that is ready to pass into a modeling function. For more control, you can call `bigexp_terms()` and `bigexp_prepare()` explicitly instead.

### Value

An object of class "bigexp\_train" which is a list with components:

- `spec`: the "bigexp\_spec" object returned by `bigexp_terms()`.
- `formula`: the expanded formula `spec$formula`.
- `data`: the prepared training data (predictors coerced to match spec), suitable for passing directly to modeling functions such as `lm()`, `glm()`, or `SVMnet()`.

### Examples

```
set.seed(1)
df5 <- data.frame(
  y = rnorm(20),
  X1 = rnorm(20),
  X2 = rnorm(20)
)

tr <- bigexp_train(
```

```

y ~ X1 + X2,
data          = df5,
factorial_order = 2,
polynomial_order = 3
)

## Prepared training data and expanded formula:
str(tr$data)
tr$formula

## Example: fit a model using the expanded formula
fit_lm <- lm(tr$formula, data = tr$data)
summary(fit_lm)

```

---

coef.svem_model	<i>Coefficients for SVEM Models</i>
-----------------	-------------------------------------

---

## Description

Extracts averaged coefficients from an `svem_model` fitted by [SVEMnet](#).

## Usage

```
## S3 method for class 'svem_model'
coef(object, debiased = FALSE, ...)
```

## Arguments

<code>object</code>	An object of class <code>svem_model</code> , typically returned by <a href="#">SVEMnet</a> .
<code>debiased</code>	Logical; if TRUE and debiased coefficients are available (Gaussian fits with <code>parms_debiased</code> ), return those instead of <code>parms</code> . Default is FALSE.
<code>...</code>	Unused; present for S3 method compatibility.

## Details

For Gaussian fits, you can optionally request debiased coefficients (if they were computed and stored) via `debiased = TRUE`. In that case, the function returns `object$parms_debiased`. If debiased coefficients are not available, or if `debiased = FALSE`, the function returns `object$parms`, which are the ensemble-averaged coefficients across bootstrap members.

For Binomial models, `debiased` is ignored and the averaged coefficients in `object$parms` are returned.

This is a lightweight accessor around the stored components of an `svem_model`:

- `parms`: ensemble-averaged coefficients over bootstrap members, on the model's link scale;
- `parms_debiased`: optional debiased coefficients (Gaussian only), if requested at fit time.

Passing `debiased = TRUE` has no effect if `parms_debiased` is NULL.

**Value**

A named numeric vector of coefficients (including the intercept).

**See Also**

[svem\\_nonzero](#) for bootstrap nonzero percentages and a quick stability plot.

**Examples**

```

set.seed(1)
n <- 200
x1 <- rnorm(n)
x2 <- rnorm(n)
eps <- rnorm(n, sd = 0.3)
y_g <- 1 + 2*x1 - 0.5*x2 + eps
dat_g <- data.frame(y_g, x1, x2)

# Small nBoot to keep runtime light in examples
fit_g <- SVEMnet(y_g ~ x1 + x2, data = dat_g, nBoot = 30, relaxed = TRUE)

# Ensemble-averaged coefficients
cc <- coef(fit_g)
head(cc)

# Debiased (only if available for Gaussian fits)
ccd <- coef(fit_g, debiased = TRUE)
head(ccd)

# Binomial example (0/1 outcome)
set.seed(2)
n <- 250
x1 <- rnorm(n)
x2 <- rnorm(n)
eta <- -0.4 + 1.1*x1 - 0.7*x2
p <- 1/(1+exp(-eta))
y_b <- rbinom(n, 1, p)
dat_b <- data.frame(y_b, x1, x2)

fit_b <- SVEMnet(y_b ~ x1 + x2, data = dat_b,
                 family = "binomial", nBoot = 30, relaxed = TRUE)

# Averaged coefficients (binomial; debiased is ignored)
coef(fit_b)

```

**Description**

Repeated K-fold cross-validation over a per-alpha lambda path, with a combined 1-SE rule across repeats. Preserves fields expected by `predict.svem_model()` and internal prediction helpers. Optionally uses glmnet's built-in relaxed elastic net for both the warm-start path and each CV fit. When `relaxed = TRUE`, the final coefficients are taken from a `cv.glmnet()` object at the chosen lambda so that the returned model reflects the relaxed solution (including its chosen gamma).

**Usage**

```
glmnet_with_cv(
  formula,
  data,
  glmnet_alpha = c(0.5, 1),
  standardize = TRUE,
  nfolds = 10,
  repeats = 5,
  choose_rule = c("min", "1se"),
  seed = NULL,
  exclude = NULL,
  relaxed = FALSE,
  relax_gamma = NULL,
  family = c("gaussian", "binomial"),
  ...
)
```

**Arguments**

<code>formula</code>	Model formula.
<code>data</code>	Data frame containing the variables in the model.
<code>glmnet_alpha</code>	Numeric vector of Elastic Net mixing parameters (alphas) in $[0, 1]$ ; default <code>c(0.5, 1)</code> . When <code>relaxed = TRUE</code> , any <code>alpha = 0</code> (ridge) is dropped with a warning.
<code>standardize</code>	Logical passed to <code>glmnet()</code> and <code>cv.glmnet()</code> (default <code>TRUE</code> ).
<code>nfolds</code>	Requested number of CV folds (default 10). Internally constrained so that there are at least about 3 observations per fold and at least 5 folds when possible.
<code>repeats</code>	Number of independent CV repeats (default 5). Each repeat reuses the same folds across all alphas for paired comparisons.
<code>choose_rule</code>	Character; how to choose lambda within each alpha: <ul style="list-style-type: none"> <li>• <code>"min"</code>: lambda minimizing the cross-validated criterion.</li> <li>• <code>"1se"</code>: largest lambda within 1 combined SE of the minimum, where the SE includes both within- and between-repeat variability.</li> </ul> Default is <code>"min"</code> . In small-mixture simulations, the 1-SE rule tended to increase RMSE on held-out data, so <code>"min"</code> is used as the default here.
<code>seed</code>	Optional integer seed for reproducible fold IDs (and the ridge fallback, if used).
<code>exclude</code>	Optional vector or function for glmnet's <code>exclude=</code> argument. If a function, <code>cv.glmnet()</code> applies it inside each training fold (requires glmnet $\geq$ 4.1-2).

relaxed	Logical; if TRUE, call <code>glmnet()</code> and <code>cv.glmnet()</code> with <code>relax = TRUE</code> and optionally a gamma path (default FALSE). If <code>cv.glmnet(relax = TRUE)</code> fails for a particular repeat/alpha, the function retries that fit without relaxation; the number of such fallbacks is recorded in <code>meta\$relax_cv_fallbacks</code> .
relax_gamma	Optional numeric vector passed as <code>gamma=</code> to <code>glmnet()</code> and <code>cv.glmnet()</code> when <code>relaxed = TRUE</code> . If NULL, <code>glmnet</code> 's internal default gamma grid is used.
family	Model family: either "gaussian" or "binomial", or the corresponding <code>stats::gaussian()</code> or <code>stats::binomial()</code> family objects with canonical links. For Gaussian, <code>y</code> must be numeric. For binomial, <code>y</code> must be 0/1 numeric, logical, or a factor with exactly 2 levels (the second level is treated as 1). Non-canonical links are not supported.
...	Additional arguments forwarded to both <code>cv.glmnet()</code> and <code>glmnet()</code> , for example: <code>weights</code> , <code>parallel</code> , <code>type.measure</code> , <code>intercept</code> , <code>lower.limits</code> , <code>upper.limits</code> , <code>penalty.factor</code> , <code>offset</code> , <code>standardize.response</code> , <code>keep</code> , and so on. <code>Glmnet</code> algorithm-control values such as <code>maxit</code> , <code>thresh</code> , <code>dfmax</code> , and <code>pmax</code> may be supplied directly for compatibility or via <code>control = list(...)</code> with <code>glmnet</code> versions that support it; they are routed in a version-compatible way when possible. If <code>family</code> is supplied here, it is ignored in favor of the explicit <code>family</code> argument.

## Details

This function is a convenience wrapper around `glmnet()` and `cv.glmnet()` that returns an object in the same structural format as `SVEMnet()` (class "svem\_model"). It is intended for:

- direct comparison of standard cross-validated `glmnet` fits to `SVEMnet` models using the same prediction and schema tools, or
- users who want a repeated-`cv.glmnet()` workflow without any `SVEM` weighting or bootstrap ensembling.

It is not called internally by the `SVEM` bootstrap routines.

The basic workflow is:

1. For each alpha in `glmnet_alpha`, generate a set of CV fold IDs (shared across alphas and repeats).
2. For that alpha, run repeats independent `cv.glmnet()` fits, align the lambda paths, and aggregate the CV curves.
3. At each lambda, compute a combined SE that accounts for both within-repeat and between-repeat variability.
4. Apply `choose_rule` ("min" or "1se") to select lambda for that alpha, then choose the best alpha by comparing these per-alpha scores.

Special cases and fallbacks:

- If there are no predictors after `model.matrix()` (an intercept-only model), the function returns an intercept-only fit without calling `glmnet()`, along with a minimal schema for safe prediction.

- If all `cv.glmnet()` attempts fail for every alpha (a rare edge case), the function falls back to a manual ridge ( $\alpha = 0$ ) CV search over a fixed lambda grid and returns the best ridge solution. For Gaussian models this search uses a mean-squared-error criterion; for binomial models it uses a negative log-likelihood (deviance-equivalent) criterion.

Family-specific behavior:

- For the Gaussian family, an optional calibration  $\text{lm}(y \sim y_{\text{pred}})$  is fit on the training data (when there is sufficient variation), and both `y_pred` and `y_pred_debiased` are stored.
- For the binomial family, `y_pred` is always on the probability (response) scale and debiasing is not applied. Both the primary cross-validation and any ridge fallback use deviance-style criteria (binomial negative log-likelihood) rather than squared error.

Design-matrix schema and contrasts:

- The training terms are stored with environment set to `baseenv()`.
- Factor and character levels are recorded in `xlevels` for safe prediction.
- Per-factor contrasts are stored in `contrasts`, normalized so that any contrasts recorded as character names are converted back to contrast functions at prediction time.

The returned object inherits classes `"svem_cv"` and `"svem_model"` and is designed to be compatible with SVEMnet prediction and schema utilities. It is a standalone, standard glmnet CV workflow that does not use SVEM-style bootstrap weighting or ensembling.

## Value

A list of class `c("svem_cv", "svem_model")` with elements:

- `parms` Named numeric vector of coefficients (including `"(Intercept)"`).
- `glmnet_alpha` Numeric vector of alphas searched.
- `best_alpha` Numeric; winning alpha.
- `best_lambda` Numeric; winning lambda.
- `y_pred` In-sample predictions from the returned coefficients (fitted values for Gaussian; probabilities for binomial).
- `debias_fit` For Gaussian, an optional  $\text{lm}(y \sim y_{\text{pred}})$  calibration model; NULL otherwise.
- `y_pred_debiased` If `debias_fit` exists, its fitted values; otherwise NULL.
- `cv_summary` Named list (one element per alpha) of data frames with columns `lambda`, `mean_cvm`, `sd_cvm`, `se_combined`, `n_repeats`, `idx_min`, `idx_1se`.
- `formula` Original modeling formula.
- `terms` Training terms object with environment set to `baseenv()`.
- `training_X` Training design matrix (without intercept column).
- `actual_y` Training response vector used for glmnet: numeric `y` for Gaussian, or 0/1 numeric `y` for binomial.
- `xlevels` Factor and character levels seen during training (for safe prediction).
- `contrasts` Contrasts used for factor predictors during training.

- `schema` List `list(feature_names, terms_str, xlevels, contrasts, terms_hash)` for deterministic prediction.
- `note` Character vector of notes (for example, dropped rows, intercept-only path, ridge fallback, relaxed-coefficient source).
- `meta` List with fields such as `nfolds`, `repeats`, `rule`, `family`, `relaxed`, `relax_cv_fallbacks`, and `cv_object` (the final `cv.glmnet()` object when `relaxed = TRUE` and `keep = TRUE`, otherwise `NULL`).
- `diagnostics` List of simple diagnostics for the selected model, currently including:
  - `k_final`: number of coefficients estimated as nonzero *including* the intercept.
  - `k_final_no_intercept`: number of nonzero slope coefficients (excludes the intercept).
- `family` Character scalar giving the resolved family ("gaussian" or "binomial"), mirroring `meta$family`.

## Acknowledgments

OpenAI's GPT models (o1-preview through GPT-5 Pro) were used to assist with coding and roxy-gen documentation; all content was reviewed and finalized by the author.

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## Examples

```
set.seed(123)
n <- 100; p <- 10
X <- matrix(rnorm(n * p), n, p)
beta <- c(1, -1, rep(0, p - 2))
y <- as.numeric(X %*% beta + rnorm(n))
df_ex <- data.frame(y = y, X)
colnames(df_ex) <- c("y", paste0("x", 1:p))

# Gaussian example, v1-like behavior: choose_rule = "min"
fit_min <- glmnet_with_cv(
  y ~ ., df_ex,
  glmnet_alpha = 1,
  nfolds = 5,
  repeats = 1,
  choose_rule = "min",
  seed = 42,
  family = "gaussian"
)

# Gaussian example, relaxed path with gamma search
fit_relax <- glmnet_with_cv(
  y ~ ., df_ex,
  glmnet_alpha = 1,
  nfolds = 5,
  repeats = 1,
  relaxed = TRUE,
  seed = 42,
  family = "gaussian"
)

# Binomial example (numeric 0/1 response)
set.seed(456)
n2 <- 150; p2 <- 8
X2 <- matrix(rnorm(n2 * p2), n2, p2)
beta2 <- c(1.0, -1.5, rep(0, p2 - 2))
```

```

linpred <- as.numeric(X2 %*% beta2)
prob <- plogis(linpred)
y_bin <- rbinom(n2, size = 1, prob = prob)
df_bin <- data.frame(y = y_bin, X2)
colnames(df_bin) <- c("y", paste0("x", 1:p2))

fit_bin <- glmnet_with_cv(
  y ~ ., df_bin,
  glmnet_alpha = c(0.5, 1),
  nfolds = 5,
  repeats = 2,
  seed = 99,
  family = "binomial"
)

```

lipid\_screen

*Lipid formulation screening data***Description**

An example dataset for modeling Potency, Size, and PDI as functions of formulation and process settings. Percent composition columns are stored as proportions in  $[0, 1]$  (for example, 4.19 percent is 0.0419). This table is intended for demonstration of SVEMnet multi-response modeling, desirability-based random-search optimization, and probabilistic design-space construction.

**Usage**

```
data(lipid_screen)
```

**Format**

A data frame with one row per experimental run and the following columns:

**RunID** character. Optional identifier for each run.

**PEG** numeric. Proportion (0-1).

**Helper** numeric. Proportion (0-1).

**Ionizable** numeric. Proportion (0-1).

**Cholesterol** numeric. Proportion (0-1).

**Ionizable\_Lipid\_Type** factor. Categorical identity of the ionizable lipid.

**N\_P\_ratio** numeric. Molar or mass  $N : P$  ratio (unitless).

**flow\_rate** numeric. Process flow rate (arbitrary units).

**Operator** factor. Categorical blocking factor.

**Potency** numeric. Response (for example, normalized activity).

**Size** numeric. Response (for example, particle size in nm).

**PDI** numeric. Response (polydispersity index).

**Notes** character. Optional free-text notes.

## Details

The four composition columns PEG, Helper, Ionizable, and Cholesterol are stored as proportions in  $[0, 1]$ , and in many rows they sum (approximately) to 1, making them natural candidates for mixture constraints in optimization and design-space examples.

This dataset accompanies examples showing:

- fitting three SVEM models (Potency, Size, PDI) on a shared expanded factor space via `bigexp_terms` and `bigexp_formula`,
- random design generation using SVEM random-table helpers (for use with multi-response optimization),
- multi-response scoring and candidate selection with `svem_score_random` (Derringer–Suich desirabilities, weights, uncertainty) and `svem_select_from_score_table` (optimal and high-uncertainty medoid candidates),
- returning both high-score optimal candidates and high-uncertainty exploration candidates from the same scored table by changing the target column (for example score vs uncertainty\_measure or wmt\_score),
- optional whole-model reweighting (WMT) of response weights via `svem_wmt_multi` (for p-values and multipliers) together with `svem_score_random` (via its `wmt` argument),
- constructing a probabilistic design space in one step by passing process-mean specifications via the `specs` argument of `svem_score_random` (internally using `svem_append_design_space_cols` when needed).

## Acknowledgments

OpenAI’s GPT models (o1-preview through GPT-5 Pro) were used to assist with coding and roxygen documentation; all content was reviewed and finalized by the author.

## Source

Simulated screening table supplied by the package author.

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Lumley, T. and Scott, A. (2015). AIC and BIC for modelling with complex survey data. *Journal of Survey Statistics and Methodology*, 3(1), 1–18.

## Examples

```
# 1) Load the bundled dataset
data(lipid_screen)
str(lipid_screen)

#2) Build a deterministic expansion using bigexp_terms()
# Provide main effects only on the right-hand side; expansion width
# is controlled via arguments. Here Operator is treated as a blocking
# factor: additive only, no interactions or polynomial terms.
spec <- bigexp_terms(
  Potency ~ PEG + Helper + Ionizable + Cholesterol +
    Ionizable_Lipid_Type + N_P_ratio + flow_rate,
  data      = lipid_screen,
  factorial_order = 3, # up to 3-way interactions
  polynomial_order = 3, # include up to cubic terms I(X^2), I(X^3)
  include_pc_2way = TRUE, # partial-cubic two-way terms Z:I(X^2)
  include_pc_3way = FALSE, # no partial-cubic three-way terms I(X^2):Z
  blocking      = "Operator",
  discrete_numeric = c("N_P_ratio", "flow_rate")
)

# 3) Reuse the same locked expansion for other responses
```

```

form_pot <- bigexp_formula(spec, "Potency")
form_siz <- bigexp_formula(spec, "Size")
form_pdi <- bigexp_formula(spec, "PDI")

# 4) Fit SVEM models with the shared factor space and expansion
set.seed(1)
fit_pot <- SVEMnet(form_pot, lipid_screen)
fit_siz <- SVEMnet(form_siz, lipid_screen)
fit_pdi <- SVEMnet(form_pdi, lipid_screen)

# 5) Collect models in a named list by response
objs <- list(Potency = fit_pot, Size = fit_siz, PDI = fit_pdi)

# 6) Define multi-response goals and weights (DS desirabilities under the hood)
#   Maximize Potency (0.6), minimize Size (0.3), minimize PDI (0.1)
goals <- list(
  Potency = list(goal = "max", weight = 0.6),
  Size     = list(goal = "min", weight = 0.3),
  PDI     = list(goal = "min", weight = 0.1)
)

# Mixture constraints: components sum to total, with bounds
mix <- list(list(
  vars = c("PEG", "Helper", "Ionizable", "Cholesterol"),
  lower = c(0.01, 0.10, 0.10, 0.10),
  upper = c(0.05, 0.60, 0.60, 0.60),
  total = 1.0
))

# 7) Optional: whole-model tests and WMT multipliers via svem_wmt_multi()
#   This wrapper runs svem_significance_test_parallel() for each response,
#   plots the distance distributions, and prints p-values and multipliers.

set.seed(123)
wmt_out <- svem_wmt_multi(
  formulas      = list(Potency = form_pot,
                      Size     = form_siz,
                      PDI     = form_pdi),
  data          = lipid_screen,
  mixture_groups = mix,
  wmt_control   = list(seed = 123),
  plot          = TRUE
)

# Inspect WMT p-values and multipliers (also printed by the wrapper)
wmt_out$p_values
wmt_out$multipliers

# 8) Optional: define process-mean specifications for a joint design space.
#   Potency at least 78, Size no more than 100, PDI less than 0.25.
#   Here we only specify the bounded side; the unbounded side defaults to
#   lower = -Inf or upper = Inf inside svem_score_random().
specs_ds <- list(

```

```

Potency = list(lower = 78),
Size    = list(upper = 100),
PDI     = list(upper = 0.25)
)

# 9) Random-search scoring in one step via svem_score_random()
# This draws a random candidate table, computes DS desirabilities,
# a combined multi-response score, WMT-adjusted wmt_score (if `wmt`
# is supplied), CI-based uncertainty, and (when `specs` is supplied)
# appends mean-level design-space columns.
#
# The `wmt` and `specs` arguments are optional:
# - Omit `wmt` for no whole-model reweighting.
# - Omit `specs` if you do not need design-space probabilities.

set.seed(3)
scored <- svem_score_random(
  objects      = objs,
  goals       = goals,
  data        = lipid_screen, # scored and returned as original_data_scored
  n           = 25000,
  mixture_groups = mix,
  level       = 0.95,
  combine     = "geom",
  numeric_sampler = "random",
  wmt         = wmt_out, # optional: NULL for no WMT
  specs      = specs_ds, # optional: NULL for no design-space columns
  verbose    = TRUE
)

# 10) Select optimal and exploration sets from the same scored table

# Optimal medoid candidates (maximizing DS score)
opt_sel <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "score", # score column is maximized
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.1,
  label       = "round1_score_optimal"
)

# Optimal medoid candidates (maximizing WMT-adjusted wmt_score)
opt_sel_wmt <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "wmt_score", # wmt_score column is maximized
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.1,
  label       = "round1_wmt_optimal"
)

```

```

# Exploration medoid candidates (highest uncertainty_measure)
explore_sel <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "uncertainty_measure", # uncertainty_measure column is maximized
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.1,
  label       = "round1_explore"
)

# In-spec medoid candidates (highest joint mean-level assurance)
inspec_sel <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "p_joint_mean",       # p_joint_mean column is maximized
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.10,
  label       = "round1_inspec"
)

# Single best by score, including per-response CIs
opt_sel$best

# Single best by WMT-adjusted score, including per-response CIs
opt_sel_wmt$best

# Diverse high-score candidates (medoids)
head(opt_sel_wmt$candidates)

# Highest-uncertainty setting and its medoid candidates
explore_sel$best
head(explore_sel$candidates)

# Highest probability mean-in-spec setting and its medoid candidates
inspec_sel$best
head(inspec_sel$candidates)

# 11) Scored original data (predictions, desirabilities, score, wmt_score, uncertainty)
head(scored$original_data_scored)

# 12) Example: combine new candidate settings with the best existing run
#       and (optionally) export a CSV for the next experimental round.

# Best existing run from the original scored data (no new medoids; k = 0)
best_existing <- svem_select_from_score_table(
  score_table = scored$original_data_scored,
  target      = "score",
  direction   = "max",
  k           = 0, # k <= 0 => only the best row, no medoids
  top_type    = "frac",

```

```

    top      = 1.0,
    label    = "round1_existing_best"
  )

# 13) Prepare candidate export tables for the next experimental round.
#   svem_export_candidates_csv() accepts individual selection objects.
#   Calls are commented out so examples/tests do not write files.

# Export a design-style candidate table (factors + responses + predictions)
# out.df <- svem_export_candidates_csv(
#   opt_sel,
#   opt_sel_wmt,
#   explore_sel,
#   inspec_sel,
#   best_existing,
#   file      = "lipid_screen_round1_candidates.csv",
#   overwrite = TRUE
# )
# head(out.df)

# Export all columns including desirabilities, CI widths, and design-space columns
# out.df2 <- svem_export_candidates_csv(
#   opt_sel,
#   opt_sel_wmt,
#   explore_sel,
#   inspec_sel,
#   best_existing,
#   file      = "lipid_screen_round1_candidates_all.csv",
#   overwrite = TRUE
# )
# head(out.df2)

```

---

plot.svem\_binomial      *Plot Method for SVEM Binomial Models*

---

### Description

Diagnostics for svem\_binomial fits from SVEMnet(..., family = "binomial"). Produces one of:

- type = "calibration": Reliability curve (binned average predicted probability vs observed rate), with jittered raw points for context.
- type = "roc": ROC curve with trapezoidal AUC in the title.
- type = "pr": Precision–Recall curve with step-wise Average Precision (AP).

### Usage

```

## S3 method for class 'svem_binomial'
plot(

```

```

  x,
  type = c("calibration", "roc", "pr"),
  bins = 10,
  jitter_width = 0.05,
  ...
)

```

### Arguments

x	An object of class <code>svem_binomial</code> .
type	One of "calibration", "roc", or "pr" (default "calibration").
bins	Integer number of equal-frequency bins for calibration (default 10).
jitter_width	Vertical jitter amplitude for raw points in calibration (default 0.05).
...	Additional aesthetics passed to <code>ggplot2::geom_line()</code> or <code>ggplot2::geom_point()</code> .

### Details

For ROC/PR, simple one-class guards are used (returns a diagonal ROC and trivial PR). The function assumes binomial models store `x$y_pred` on the *probability* scale.

### Value

A `ggplot2` object.

### Examples

```

## Not run:
## --- Binomial example: simulate, fit, and plot -----
set.seed(2025)
n <- 600
x1 <- rnorm(n); x2 <- rnorm(n); x3 <- rnorm(n)
eta <- -0.4 + 1.1*x1 - 0.8*x2 + 0.5*x3
p_true <- plogis(eta)
y <- rbinom(n, 1, p_true)
dat_b <- data.frame(y, x1, x2, x3)

fit_b <- SVEMnet(
  y ~ x1 + x2 + x3 + I(x1^2) + (x1 + x2 + x3)^2,
  data = dat_b,
  family = "binomial",
  glmnet_alpha = c(1, 0.5),
  nBoot = 60,
  objective = "auto",
  weight_scheme = "SVEM",
  relaxed = TRUE
)

# Calibration / ROC / PR
plot(fit_b, type = "calibration", bins = 12)
plot(fit_b, type = "roc")

```

```
plot(fit_b, type = "pr")

## End(Not run)
```

---

plot.svem_model	<i>Plot Method for SVEM Models (Gaussian / Generic)</i>
-----------------	---

---

### Description

Plots actual versus predicted values for an `svem_model`. This is the default plot for models fit with `SVEMnet(..., family = "gaussian")` and any other non-binomial models that share the `svem_model` class.

### Usage

```
## S3 method for class 'svem_model'
plot(x, plot_debiased = FALSE, ...)
```

### Arguments

<code>x</code>	An object of class <code>svem_model</code> .
<code>plot_debiased</code>	Logical; if TRUE, include debiased predictions (when available) as an additional series. Default FALSE.
<code>...</code>	Additional aesthetics passed to <code>ggplot2::geom_point()</code> .

### Details

Points show fitted values against observed responses; the dashed line is the 45-degree identity. If available and requested, debiased predictions are included as a second series.

This method assumes the fitted object stores the training response in `$actual_y` and in-sample predictions in `$y_pred`, as produced by `SVEMnet()` and `glmnet_with_cv()`.

### Value

A `ggplot2` object.

### Examples

```
## Not run:
## --- Gaussian example: simulate, fit, and plot -----
set.seed(2026)
n <- 300
X1 <- rnorm(n); X2 <- rnorm(n); X3 <- rnorm(n)
eps <- rnorm(n, sd = 0.4)
y_g <- 1.2 + 2*X1 - 0.7*X2 + 0.3*X3 + 1.1*(X1*X2) + 0.8*(X1^2) + eps
dat_g <- data.frame(y_g, X1, X2, X3)
```

```

fit_g <- SVMnet(
  y_g ~ (X1 + X2 + X3)^2 + I(X1^2) + I(X2^2),
  data      = dat_g,
  family    = "gaussian",
  glmnet_alpha = c(1, 0.5),
  nBoot     = 60,
  objective  = "auto",
  weight_scheme = "SVEM",
  relaxed   = TRUE
)

# Actual vs predicted (with and without debias overlay)
plot(fit_g, plot_debiased = FALSE)
plot(fit_g, plot_debiased = TRUE)

## End(Not run)

```

---

```
plot.svem_significance_test
```

*Plot SVEM significance test results for one or more responses*

---

## Description

Plots the Mahalanobis-like distances for original and permuted data from one or more SVEM significance test results returned by `svem_significance_test_parallel()`.

## Usage

```
## S3 method for class 'svem_significance_test'
plot(x, ..., labels = NULL)
```

## Arguments

<code>x</code>	An object of class <code>svem_significance_test</code> .
<code>...</code>	Optional additional <code>svem_significance_test</code> objects to include in the same plot.
<code>labels</code>	Optional character vector of labels for the responses. If not provided, the function uses inferred response names (from <code>data_d\$Response</code> or <code>x\$response</code> ) and ensures uniqueness.

## Details

If additional `svem_significance_test` objects are provided via `...`, their distance tables (`$data_d`) are stacked and plotted together using a shared x-axis grouping of "Response / Source" and a fill aesthetic indicating "Original" vs "Permutation".

**Value**

A ggplot2 object showing the distributions of distances for original vs. permuted data, grouped by response.

---

predict.svem\_model      *Predict Method for SVEM Models (Gaussian and Binomial)*

---

**Description**

Generate predictions from a fitted SVEM model (Gaussian or binomial), with optional bootstrap uncertainty and family-appropriate output scales.

**Usage**

```
## S3 method for class 'svem_model'
predict(
  object,
  newdata,
  type = c("response", "link", "class"),
  debias = FALSE,
  se.fit = FALSE,
  interval = FALSE,
  level = 0.95,
  ...
)
```

**Arguments**

object	A fitted SVEM model (class svem_model; binomial models typically also inherit class svem_binomial). Created by SVEMnet().
newdata	A data frame of new predictor values.
type	(Binomial only) One of: <ul style="list-style-type: none"> <li>• "response" (default): predicted probabilities in [0, 1].</li> <li>• "link": linear predictor (log-odds).</li> <li>• "class": 0/1 class labels (threshold 0.5). Uncertainty summaries are not available for this type.</li> </ul> Ignored for Gaussian models.
debias	(Gaussian only) Logical; default FALSE. If TRUE, apply the linear calibration fit $\text{lm}(y \sim y_{\text{pred}})$ learned at training when available. Ignored (and internally set to FALSE) for binomial models.
se.fit	Logical; if TRUE, return bootstrap standard errors computed from member predictions (requires coef_matrix). Not available for type = "class". For Gaussian models, this forces use of bootstrap member predictions instead of aggregate coefficients.

interval	Logical; if TRUE, return percentile confidence limits from member predictions (requires <code>coef_matrix</code> ). Not available for <code>type = "class"</code> . For Gaussian models, this forces use of bootstrap member predictions instead of aggregate coefficients.
level	Confidence level for percentile intervals. Default 0.95.
...	Currently unused.

### Details

This method dispatches on `object$family`:

- **Gaussian**: returns predictions on the response (identity) scale. Optional linear calibration ("debias") learned at training may be applied.
- **Binomial**: supports glmnet-style `type = "link", "response",` or `"class"` predictions. No debiasing is applied; `type = "response"` returns probabilities in  $[0, 1]$ .

Uncertainty summaries (`se.fit`, `interval`) and all binomial predictions are based on per-bootstrap member predictions obtained from the coefficient matrix stored in `object$coef_matrix`. If `coef_matrix` is NULL, these options are not available (and binomial prediction will fail). For Gaussian models with `se.fit = FALSE` and `interval = FALSE`, predictions are computed directly from the aggregated coefficients.

### Value

If `se.fit = FALSE` and `interval = FALSE`:

- **Gaussian**: a numeric vector of predictions on the response (identity) scale.
- **Binomial**: a numeric vector for `type = "response"` (probabilities) or `type = "link"` (log-odds), or an integer vector of 0/1 labels for `type = "class"`.

If `se.fit` and/or `interval` are TRUE (and `type != "class"`), a list with components:

- `fit`: predictions on the requested scale.
- `se.fit`: bootstrap standard errors (when `se.fit = TRUE`).
- `lwr`, `upr`: percentile confidence limits (when `interval = TRUE`).

Rows containing unseen or missing factor levels produce NA predictions (and NA SEs/intervals), with a warning.

### Design-matrix reconstruction

The function rebuilds the design matrix for `newdata` to match the training design:

- Uses the training terms (with environment set to `baseenv()`).
- Harmonizes factor and character predictors to the training `xlevels`.
- Reuses stored per-factor contrasts when available; otherwise falls back to saved global contrast options.
- Zero-fills any columns present at training but absent in `newdata`, and reorders columns to match the training order.

Rows containing unseen factor levels yield NA predictions (with a warning).

## Aggregation and debiasing

For Gaussian SVEM models:

**Point predictions** When `se.fit = FALSE` and `interval = FALSE`, predictions are computed from the aggregated coefficients saved at fit time (`object$params`; or `object$params_debiased` when `debias = TRUE`). This is algebraically equivalent to averaging member predictions when the coefficients were formed as bootstrap means.

**Bootstrap-based summaries** When `se.fit = TRUE` and/or `interval = TRUE`, predictions are computed from per-bootstrap member predictions using `object$coef_matrix`. For `debias = TRUE`, the linear calibration is applied to member predictions before summarizing.

For binomial SVEM models, predictions are always aggregated from member predictions on the requested scale (probability or link) using `coef_matrix`; the stored coefficient averages (`parms`, `parms_debiased`) are retained for diagnostics but are not used in prediction. The `debias` argument is ignored and treated as `FALSE` for binomial models.

For Gaussian fits, if `debias = TRUE` and a calibration model `lm(y ~ y_pred)` was learned at training, predictions (and, when applicable, member predictions) are transformed by that calibration. This debiasing is never applied for binomial fits.

## Uncertainty

When `se.fit = TRUE`, standard errors are computed as the row-wise standard deviations of member predictions on the requested scale. When `interval = TRUE`, percentile intervals are computed from member predictions on the requested scale, using the requested level. Both require a non-null `coef_matrix`. For `type = "class"` (binomial), uncertainty summaries are not available.

## See Also

[SVEMnet](#)

## Examples

```
## ---- Gaussian example -----
set.seed(1)
n <- 60
X1 <- rnorm(n); X2 <- rnorm(n); X3 <- rnorm(n)
y <- 1 + 0.8 * X1 - 0.6 * X2 + 0.2 * X3 + rnorm(n, 0, 0.4)
dat <- data.frame(y, X1, X2, X3)

fit_g <- SVEMnet(
  y ~ (X1 + X2 + X3)^2, dat,
  nBoot = 40, glmnet_alpha = c(1, 0.5),
  relaxed = TRUE, family = "gaussian"
)

## Aggregate-coefficient predictions (with and without debiasing)
p_g <- predict(fit_g, dat) # debias = FALSE (default)
p_gd <- predict(fit_g, dat, debias = TRUE) # apply calibration, if available

## Bootstrap-based uncertainty (requires coef_matrix)
```

```

out_g <- predict(
  fit_g, dat,
  debias = TRUE,
  se.fit = TRUE,
  interval = TRUE,
  level = 0.90
)
str(out_g)

## ---- Binomial example -----
set.seed(2)
n <- 120
X1 <- rnorm(n); X2 <- rnorm(n); X3 <- rnorm(n)
eta <- -0.3 + 1.1 * X1 - 0.8 * X2 + 0.5 * X1 * X3
p <- plogis(eta)
yb <- rbinom(n, 1, p)
db <- data.frame(yb = yb, X1 = X1, X2 = X2, X3 = X3)

fit_b <- SVMnet(
  yb ~ (X1 + X2 + X3)^2, db,
  nBoot = 50, glmnet_alpha = c(1, 0.5),
  relaxed = TRUE, family = "binomial"
)

## Probabilities, link, and classes
p_resp <- predict(fit_b, db, type = "response")
p_link <- predict(fit_b, db, type = "link")
y_hat <- predict(fit_b, db, type = "class") # 0/1 labels (no SE or interval)

## Bootstrap-based uncertainty on the probability scale
out_b <- predict(
  fit_b, db,
  type = "response",
  se.fit = TRUE,
  interval = TRUE,
  level = 0.90
)
str(out_b)

```

---

predict\_cv

*Predict from glmnet\_with\_cv Fits (svem\_cv Objects)*


---

### Description

Generate predictions from a fitted object returned by `glmnet_with_cv()` (class "svem\_cv").

**Usage**

```
predict_cv(object, newdata, debias = FALSE, strict = FALSE, ...)

## S3 method for class 'svem_cv'
predict(object, newdata, debias = FALSE, strict = FALSE, ...)
```

**Arguments**

<code>object</code>	A fitted object returned by <code>glmnet_with_cv()</code> (class "svem_cv").
<code>newdata</code>	A data frame of new predictor values.
<code>debias</code>	Logical; if TRUE and a debiasing fit is available, apply it. Has an effect only for Gaussian models where <code>debias_fit</code> is present.
<code>strict</code>	Logical; if TRUE, enforce a strict column-name match between the aligned design for <code>newdata</code> and the training design (including the intercept position). Default FALSE.
<code>...</code>	Additional arguments (currently unused).

**Details**

The design matrix for `newdata` is rebuilt using the stored training terms (with environment set to `baseenv()`), together with the saved factor `xlevels` and contrasts (cached in `object$schema`). Columns are then aligned back to the training design in a robust way:

- Any training columns that `model.matrix()` drops for `newdata` (for example, a factor collapsing to a single level) are added back as zero columns.
- Columns are reordered to exactly match the training order.
- Rows containing unseen factor/character levels are warned about and their predictions are set to NA.

For Gaussian fits (`family = "gaussian"`), the returned values are on the original response (identity-link) scale. For binomial fits (`family = "binomial"`), the returned values are probabilities in  $[0, 1]$  (logit-link inverted via `plogis()`).

If `debias = TRUE` and a calibration model `lm(y ~ y_pred)` is present with a finite slope, predictions are adjusted via  $a + b * \text{pred}$ . Debiasing is only fitted and used for Gaussian models; for binomial models the `debias` argument has no effect.

`predict_cv()` is a small convenience wrapper that simply calls the underlying S3 method `predict.svem_cv()`, keeping a single code path for prediction from `glmnet_with_cv()` objects.

**Value**

A numeric vector of predictions on the response scale: numeric fitted values for Gaussian models; probabilities in  $[0, 1]$  for binomial models. Rows with unseen factor/character levels return NA.

**Examples**

```

set.seed(1)
n <- 50; p <- 5
X <- matrix(rnorm(n * p), n, p)
y <- X[, 1] - 0.5 * X[, 2] + rnorm(n)
df_ex <- data.frame(y = as.numeric(y), X)
colnames(df_ex) <- c("y", paste0("x", 1:p))

fit <- glmnet_with_cv(
  y ~ ., df_ex,
  glmnet_alpha = 1,
  nfolds = 5,
  repeats = 2,
  seed = 9,
  family = "gaussian"
)

preds_raw <- predict_cv(fit, df_ex)
preds_db <- predict_cv(fit, df_ex, debias = TRUE)
cor(preds_raw, df_ex$y)

# Binomial example (probability predictions on [0,1] scale)
set.seed(2)
n2 <- 80; p2 <- 4
X2 <- matrix(rnorm(n2 * p2), n2, p2)
eta2 <- X2[, 1] - 0.8 * X2[, 2]
pr2 <- plogis(eta2)
y2 <- rbinom(n2, size = 1, prob = pr2)
df_bin <- data.frame(y = y2, X2)
colnames(df_bin) <- c("y", paste0("x", 1:p2))

fit_bin <- glmnet_with_cv(
  y ~ ., df_bin,
  glmnet_alpha = c(0.5, 1),
  nfolds = 5,
  repeats = 2,
  seed = 11,
  family = "binomial"
)

prob_hat <- predict_cv(fit_bin, df_bin)
summary(prob_hat)

```

---

print.bigexp\_spec

*Print method for bigexp\_spec objects*


---

**Description**

This print method shows a compact summary of the expansion settings and the predictors that are treated as continuous or categorical. It also reports any variables that were designated as blocking

factors and therefore enter the model only additively (no interactions, no polynomials).

### Usage

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

### Arguments

```
x          A "bigexp_spec" object.  
...        Unused.
```

### Examples

```
set.seed(1)  
df4 <- data.frame(  
  y = rnorm(10),  
  X1 = rnorm(10),  
  G = factor(sample(c("A", "B"), 10, replace = TRUE))  
)  
  
spec4 <- bigexp_terms(  
  y ~ X1 + G,  
  data = df4,  
  factorial_order = 2,  
  polynomial_order = 2  
)  
  
print(spec4)  
  
## Example with a blocking factor:  
set.seed(2)  
df_block2 <- data.frame(  
  y = rnorm(12),  
  X1 = rnorm(12),  
  G = factor(sample(c("A", "B"), 12, replace = TRUE)),  
  Operator = factor(sample(letters[1:3], 12, replace = TRUE)),  
  AmbientTemp = rnorm(12, mean = 22, sd = 1.5)  
)  
  
spec_block2 <- bigexp_terms(  
  y ~ X1 + G,  
  data = df_block2,  
  factorial_order = 2,  
  polynomial_order = 3,  
  blocking = c("Operator", "AmbientTemp")  
)  
  
print(spec_block2)
```

---

```
print.svem_significance_test
      Print Method for SVEM Significance Test
```

---

**Description**

Prints the median p-value from an object of class `svem_significance_test`.

**Usage**

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

**Arguments**

<code>x</code>	An object of class <code>svem_significance_test</code> .
<code>...</code>	Additional arguments (unused).

---

<code>SVEMnet</code>	<i>Fit an SVEMnet model (Self-Validated Ensemble Elastic Net)</i>
----------------------	---

---

**Description**

Fit a Self-Validated Ensemble Model (SVEM) with elastic net or relaxed elastic net base learners using `glmnet`. Fractional random-weight (FRW) train/validation weights are drawn on each bootstrap replicate, a validation-weighted information criterion (wAIC, wBIC, or wSSE) is minimized to select the penalty, and predictions are ensembled across replicates. Gaussian and binomial responses are supported.

**Usage**

```
SVEMnet(
  formula,
  data,
  nBoot = 200,
  glmnet_alpha = c(0.5, 1),
  weight_scheme = c("SVEM", "FRW_plain", "Identity"),
  objective = c("auto", "wAIC", "wBIC", "wSSE"),
  relaxed = "auto",
  response = NULL,
  unseen = c("warn_na", "error"),
  family = c("gaussian", "binomial"),
  ...
)
```

**Arguments**

<code>formula</code>	A formula specifying the model to be fitted, or a <code>bigexp_spec</code> created by <code>bigexp_terms()</code> .
<code>data</code>	A data frame containing the variables in the model.
<code>nBoot</code>	Integer. Number of bootstrap replicates (default 200). Each replicate draws FRW weights and fits a <code>glmnet</code> path.
<code>glmnet_alpha</code>	Numeric vector of elastic net mixing parameters $\alpha$ in $[0, 1]$ . $\alpha = 1$ is lasso, $\alpha = 0$ is ridge. Defaults to <code>c(0.5, 1)</code> . When <code>relaxed = TRUE</code> , $\alpha = 0$ is automatically dropped (ridge + relaxed is not used).
<code>weight_scheme</code>	Character. Weighting scheme for train/validation copies. One of: <ul style="list-style-type: none"> <li>"SVEM" (default): Self-Validated Ensemble Model weights. For each replicate and row, a shared uniform draw <math>U_i \sim \text{Unif}(0, 1)</math> is converted to anti-correlated FRW weights <math>w_{\text{train}_i} = -\log(U_i)</math> and <math>w_{\text{valid}_i} = -\log(1 - U_i)</math>. Each weight vector is then rescaled to have mean 1 (sum <math>n</math>).</li> <li>"FRW_plain": Fractional random-weight regression without a separate validation copy. A single FRW vector <math>w_i = -\log(U_i)</math> is used for both training and validation and rescaled to have mean 1 (sum <math>n</math>). This reproduces the FRW bootstrap regression of Xu et al. (2020) and related work.</li> <li>"Identity": Uses unit weights for both training and validation (no resampling). In combination with <code>nBoot = 1</code> this wraps a single <code>glmnet</code> fit and selects the penalty by the chosen information criterion, while still using SVEMnet's expansion and diagnostics.</li> </ul>
<code>objective</code>	Character. One of "auto", "wAIC", "wBIC", or "wSSE". <ul style="list-style-type: none"> <li>"wAIC": Gaussian AIC-like criterion based on the weighted SSE.</li> <li>"wBIC": Gaussian BIC-like criterion based on the weighted SSE.</li> </ul> <p>See Details for the exact criteria in the Gaussian and binomial cases.</p>
<code>relaxed</code>	Logical or character. Default "auto". If TRUE, use <code>glmnet</code> 's relaxed elastic-net path and select both the penalty $\lambda$ and the relaxed refit parameter $\gamma$ on each bootstrap. If FALSE, fit the standard <code>glmnet</code> path without the relaxed step. If "auto" (default), SVEMnet uses <code>relaxed = TRUE</code> for <code>family = "gaussian"</code> and <code>relaxed = FALSE</code> for <code>family = "binomial"</code> .
<code>response</code>	Optional character. When <code>formula</code> is a <code>bigexp_spec</code> , this names the response column to use on the left-hand side. Defaults to the response stored in the spec.
<code>unseen</code>	How to treat factor levels not seen in the original <code>bigexp_spec</code> when <code>formula</code> is a <code>bigexp_spec</code> . One of "warn_na" (default; convert unseen levels to NA with a warning) or "error" (stop with an error).
<code>family</code>	Character. One of "gaussian" (default) or "binomial". For Gaussian models SVEMnet uses the identity link; for binomial it uses the canonical logit link. The binomial response must be numeric 0/1, logical, or a factor with exactly two levels (the second level is treated as 1).
<code>...</code>	Additional arguments passed to <code>glmnet()</code> , such as <code>penalty.factor</code> , <code>lower.limits</code> , <code>upper.limits</code> , <code>offset</code> , or <code>standardize.response</code> . Any user-supplied weights are ignored (SVEMnet supplies its own bootstrap weights). Any user-supplied <code>standardize</code> is ignored; SVEMnet always calls <code>glmnet</code> with <code>standardize =</code>

TRUE. Glmnet algorithm-control values supplied directly or through `control = list(...)` are routed in a version-compatible way when possible.

## Details

You can pass either:

- a standard model formula, e.g.,  $y \sim X1 + X2 + X3 + I(X1^2) + (X1 + X2 + X3)^2$ , or
- a `bigexp_spec` created by `bigexp_terms()`, in which case SVEMnet will build the design matrix deterministically (locked types, levels, and contrasts) and, if requested, swap the response to fit multiple independent responses over the same expansion.

SVEMnet implements Self-Validated Ensemble Models using elastic net and relaxed elastic net base learners from `glmnet`. Each bootstrap replicate draws fractional random weights, builds a train and validation copy, fits a path over  $\lambda$  (and optionally over  $\alpha$  and relaxed  $\gamma$ ), and selects a path point by minimizing a validation-weighted criterion. Final predictions are obtained by averaging replicate predictions on the chosen scale.

By default, `relaxed = "auto"` resolves to TRUE for Gaussian fits and FALSE for binomial fits.

The function is typically used in small-n design-of-experiments (DOE) workflows where classical train/validation splits and cross-validation can be unstable. A common pattern is: (1) build a deterministic expansion with `bigexp_terms()`, (2) fit SVEM models via `SVEMnet()`, (3) perform whole-model significance testing, and (4) call `svem_score_random()` for constrained multi-response optimization.

Weighting schemes:

- With `weight_scheme = "SVEM"`, SVEMnet uses a pair of anti-correlated FRW vectors for train and validation. All rows appear in every replicate, but train and validation contributions are separated through the shared uniform draws.
- With `weight_scheme = "FRW_plain"`, a single FRW vector is used for both train and validation, which reproduces FRW regression without a self-validation split. This is mainly provided for method comparison and teaching.
- With `weight_scheme = "Identity"`, both train and validation weights are 1. Setting `nBoot = 1` in this mode yields a single `glmnet` fit whose penalty is chosen by the selected information criterion, without any bootstrap variation.

Selection criteria (Gaussian): For `family = "gaussian"`, the validation loss is a weighted sum of squared errors on the validation copy. Let  $SSE_w = \sum_i w_i^{valid} r_i^2$  denote the weighted SSE. The criteria are:

- "wSSE": loss-only selector that minimizes the weighted SSE  $SSE_w$ ,
- "wAIC": Gaussian AIC analog  $C(\lambda) = n \log\{SSE_w(\lambda)/n\} + 2k$ ,
- "wBIC": Gaussian BIC analog  $C(\lambda) = n \log\{SSE_w(\lambda)/n\} + \log(n_{eff} k)$ .

The FRW validation weights are rescaled to have mean one, so that their sum is always  $\sum_i w_i^{valid} = n$ . The AIC/BIC analogs therefore use  $n \log(SSE_w/n)$  as the Gaussian loss term, while the "wSSE" selector uses  $SSE_w$  directly.

The effective validation size is computed from the FRW weights using Kish's effective sample size  $n_{eff} = (\sum_i w_i^{valid})^2 / \sum_i (w_i^{valid})^2$  and then truncated to lie between 2 and  $n$  to form  $n_{eff}$ .

The AIC-style selector uses a  $2k$  penalty; the BIC-style selector uses a  $\log(n_{eff} f_a dm)k$  penalty, so that the loss term is scaled by total validation weight while the complexity penalty reflects the effective amount of information under unequal weights. For "wAIC" and "wBIC", path points with more than  $n_{eff} f_a dm$  non-intercept coefficients are treated as inadmissible when evaluating the criterion.

Because the FRW validation weights are random rather than fixed design weights, these information-criterion scores are used heuristically for relative model comparison within each FRW replicate, rather than as exact AIC/BIC values.

For diagnostics, SVEMnet reports the raw Kish effective sizes across bootstraps (see `diagnostics$n_eff_summary`), while  $n_{eff} f_a dm$  is used internally in the penalty and model-size guardrail. Near-interpolating path points are screened out via a simple model size guardrail before minimization. When `objective = "auto"`, SVEMnet uses "wAIC".

This structure (pseudo-likelihood using total weight and BIC penalty using a Kish-type effective sample size) parallels survey-weighted information criteria as in Lumley and Scott (2015) and Kish (1965).

Selection criteria (binomial): For `family = "binomial"`, the validation loss is the weighted negative log-likelihood on the FRW validation copy (equivalently, proportional to the binomial deviance up to a constant factor). Let NLL denote the weighted negative log-likelihood. The same labels are used:

- "wSSE": loss-only selector based on NLL (the name is retained for backward compatibility),
- "wAIC": deviance-style criterion  $C(\lambda) = 2 \text{NLL}(\lambda) + 2k$ ,
- "wBIC": deviance-style criterion  $C(\lambda) = 2 \text{NLL}(\lambda) + \log(n_{eff} f_a dm)k$ .

The effective validation size  $n_{eff} f_a dm$  and the model size guardrail are handled as in the Gaussian case: for "wAIC" and "wBIC" we compute a Kish effective size from the FRW validation weights, truncate it to lie between 2 and `n`, and require the number of nonzero coefficients (excluding the intercept) to be less than this effective size when evaluating the criterion.

Auto rule: When `objective = "auto"`, SVEMnet selects the criterion by family:

- `family = "gaussian" -> "wAIC"`
- `family = "binomial" -> "wBIC"`

Relaxed elastic net: When `relaxed = TRUE`, SVEMnet calls `glmnet` with `relax = TRUE` and traverses a small grid of relaxed refit values (`gamma`). For each `alpha` and `gamma`, SVEMnet evaluates all lambda path points on the validation copy and records the combination that minimizes the selected criterion. Model size is always defined as the number of nonzero coefficients including the intercept, so standard and relaxed paths are scored on the same scale.

Gaussian debiasing: For Gaussian models, SVEMnet optionally performs a simple linear calibration of ensemble predictions on the training data. When there is sufficient variation in the fitted values and `nBoot` is at least 10, the function fits `lm(y ~ y_pred)` and uses the coefficients to construct debiased coefficients and debiased fitted values. Binomial fits do not use debiasing; predictions are ensembled on the probability or link scale directly.

Implementation notes:

- Predictors are always standardized internally via `glmnet(..., standardize = TRUE)`.
- The `terms` object is stored with its environment set to `baseenv()` so that prediction does not accidentally capture objects from the calling environment.

- A compact schema (feature names, factor levels, contrasts, and a simple hash) is stored to allow `predict()` and companion functions to rebuild model matrices deterministically, even when the original data frame is not available.
- A separate sampling schema stores raw predictor ranges and factor levels for use in random candidate generation for optimization.

## Value

An object of class "svem\_model" (and "svem\_binomial" when `family = "binomial"`) with components:

- `parms`: Vector of ensemble-averaged coefficients, including the intercept.
- `parms_debiased`: Vector of coefficients after optional debiasing (see Details; Gaussian only).
- `debias_fit`: If debiasing was performed, the calibration model  $\text{lm}(y \sim y_{\text{pred}})$ ; otherwise NULL.
- `coef_matrix`: Matrix of per-bootstrap coefficients (rows = bootstraps, columns = intercept and predictors).
- `nBoot`: Number of bootstrap replicates actually used.
- `glmnet_alpha`: Vector of alpha values considered.
- `best_alphas`: Per-bootstrap alpha selected by the criterion.
- `best_lambdas`: Per-bootstrap lambda selected by the criterion.
- `best_relax_gammas`: Per-bootstrap relaxed gamma selected when `relaxed = TRUE`; NA otherwise.
- `weight_scheme`: The weighting scheme that was used.
- `relaxed`: Logical flag indicating whether relaxed paths were used.
- `relaxed_input`: The user-supplied value for `relaxed` (one of TRUE, FALSE, or "auto"). The resolved flag actually used is reported in `relaxed`.
- `dropped_alpha0_for_relaxed`: Logical; TRUE if  $\alpha = 0$  was dropped because `relaxed = TRUE`.
- `objective_input`: The objective requested by the user.
- `objective_used`: The objective actually used after applying the "auto" rule (for example "wAIC" or "wBIC").
- `objective`: Same as `objective_used` (for convenience).
- `auto_used`: Logical; TRUE if `objective = "auto"`.
- `auto_decision`: The objective selected by the auto rule (wAIC or wBIC) when `auto_used = TRUE`.
- `diagnostics`: List with summary information, including:
  - `k_summary`: Median and IQR of selected model size (number of nonzero coefficients including intercept).
  - `fallback_rate`: Proportion of bootstraps that fell back to an intercept-only fit.
  - `n_eff_summary`: Summary of raw Kish effective validation sizes  $n_{eff} = (\sum_i w_i^{valid})^2 / \sum_i (w_i^{valid})^2$  across bootstraps (before truncation to form  $n_{eff_{adm}}$ ).

- alpha\_freq: Relative frequency of selected alpha values (if any).
- relax\_gamma\_freq: Relative frequency of selected relaxed gamma values (if relaxed = TRUE and any were selected).
- actual\_y: Numeric response vector used in fitting (0/1 for binomial).
- training\_X: Numeric model matrix without the intercept column used for training.
- y\_pred: Fitted values from the ensemble on the training data. For Gaussian this is on the response scale; for binomial it is on the probability scale.
- y\_pred\_debiased: Debiased fitted values on the training data (Gaussian only); NULL otherwise.
- nobs: Number of observations used in fitting.
- nparam: Number of parameters in the full expansion (intercept plus predictors).
- formula: The formula used for fitting (possibly derived from a bigexp\_spec).
- terms: terms object used for building the design matrix, with environment set to baseenv() for safety.
- xlevels: Factor levels recorded at training time.
- contrasts: Contrasts used for building the design matrix.
- schema: Compact description for safe prediction, including feature\_names, terms\_str, xlevels, contrasts, contrasts\_options, and a simple hash.
- sampling\_schema: Schema used to generate random candidate tables, including predictor names, variable classes, numeric ranges, and factor levels.
- used\_bigexp\_spec: Logical flag indicating whether a bigexp\_spec was used.
- family: The fitted family ("gaussian" or "binomial").

## Acknowledgments

OpenAI's GPT models (o1-preview through GPT-5 Pro) were used to assist with coding and roxygen documentation; all content was reviewed and finalized by the author.

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## Examples

```
set.seed(42)

n <- 30
X1 <- rnorm(n)
X2 <- rnorm(n)
X3 <- rnorm(n)
eps <- rnorm(n, sd = 0.5)
y <- 1 + 2 * X1 - 1.5 * X2 + 0.5 * X3 + 1.2 * (X1 * X2) +
  0.8 * (X1^2) + eps
dat <- data.frame(y, X1, X2, X3)

# Minimal hand-written expansion
mod_relax <- SVEMnet(
  y ~ (X1 + X2 + X3)^2 + I(X1^2) + I(X2^2),
  data      = dat,
  glmnet_alpha = c(1, 0.5),
  nBoot     = 75,
  objective  = "auto",
  weight_scheme = "SVEM",
  relaxed   = FALSE
)

pred_in_raw <- predict(mod_relax, dat, debias = FALSE)
```

```

pred_in_db <- predict(mod_relax, dat, debias = TRUE)

# -----
# Big expansion (full factorial + polynomial surface + partial-cubic crosses)
# Build once, reuse for one or more responses
# -----
spec <- bigexp_terms(
  y ~ X1 + X2 + X3,
  data      = dat,
  factorial_order = 3,
  polynomial_order = 3,
  include_pc_3way = FALSE
)

# Fit using the spec (auto-prepares data)
fit_y <- SVEMnet(
  spec, dat,
  glmnet_alpha = c(1, 0.5),
  nBoot        = 50,
  objective    = "auto",
  weight_scheme = "SVEM"
)

# A second, independent response over the same expansion
set.seed(99)
dat$y2 <- 0.5 + 1.4 * X1 - 0.6 * X2 + 0.2 * X3 + rnorm(n, 0, 0.4)
fit_y2 <- SVEMnet(
  spec, dat, response = "y2",
  glmnet_alpha = c(1, 0.5),
  nBoot        = 50,
  objective    = "auto",
  weight_scheme = "SVEM"
)

svem_nonzero(fit_y2)

p1 <- predict(fit_y, dat)
p2 <- predict(fit_y2, dat, debias = TRUE)

# Show that a new batch expands identically under the same spec
newdat <- data.frame(
  y = y,
  X1 = X1 + rnorm(n, 0, 0.05),
  X2 = X2 + rnorm(n, 0, 0.05),
  X3 = X3 + rnorm(n, 0, 0.05)
)

prep_new <- bigexp_prepare(spec, newdat)
stopifnot(identical(
  colnames(model.matrix(spec$formula, bigexp_prepare(spec, dat)$data)),
  colnames(model.matrix(spec$formula, prep_new$data))
))
preds_new <- predict(fit_y, prep_new$data)

```

```

## Binomial example
set.seed(2)
n <- 120
X1 <- rnorm(n); X2 <- rnorm(n); X3 <- rnorm(n)
eta <- -0.3 + 1.1 * X1 - 0.8 * X2 + 0.5 * X1 * X3
p <- plogis(eta)
yb <- rbinom(n, 1, p)
db <- data.frame(yb = yb, X1 = X1, X2 = X2, X3 = X3)

fit_b <- SVEMnet(
  yb ~ (X1 + X2 + X3)^2, db,
  nBoot = 50,
  glmnet_alpha = c(1, 0.5),
  family = "binomial"
)

## Probabilities, link, and classes
p_resp <- predict(fit_b, db, type = "response")
p_link <- predict(fit_b, db, type = "link")
y_hat <- predict(fit_b, db, type = "class") # 0/1 labels

## Mean aggregation with uncertainty on probability scale
out_b <- predict(
  fit_b, db,
  type = "response",
  se.fit = TRUE,
  interval = TRUE,
  level = 0.9
)
str(out_b)

#' ## Example with blocking (requires SVEMnet to store sampling_schema$blocking)
set.seed(2)
df_block <- data.frame(
  y1 = rnorm(40),
  y2 = rnorm(40),
  X1 = runif(40),
  X2 = runif(40),
  Operator = factor(sample(paste0("Op", 1:3), 40, TRUE)),
  AmbientTmp = rnorm(40, mean = 22, sd = 2)
)

spec_block <- bigexp_terms(
  y1 ~ X1 + X2,
  data = df_block,
  factorial_order = 2,
  polynomial_order = 2,
  blocking = c("Operator", "AmbientTmp")
)

fit_b1 <- SVEMnet(spec_block, df_block, response = "y1", nBoot = 30)
fit_b2 <- SVEMnet(spec_block, df_block, response = "y2", nBoot = 30)

```

```
tab_block <- svem_random_table_multi(list(fit_b1, fit_b2), n = 500)
```

---

```
svem_export_candidates_csv
```

*Export SVEM candidate sets to CSV*

---

## Description

Given one or more selection objects returned by [svem\\_select\\_from\\_score\\_table](#), concatenate their \$best rows and \$candidates and export a CSV suitable for planning new experimental runs.

Each row is tagged with:

- candidate\_type: "best" or "medoid".
- selection\_label: derived from the label argument used in [svem\\_select\\_from\\_score\\_table\(\)](#) when available.

The function does not modify any response or prediction columns (for example, Potency, Potency\_pred); it simply harmonizes columns across inputs (adding NA-filled columns where necessary), concatenates rows, and reorders a few metadata columns for readability.

Any columns named candidate\_type, selection\_label, or Notes\_from\_SVEMnet that are present in the final data frame are moved to the leftmost positions in that order.

## Usage

```
svem_export_candidates_csv(
  ...,
  file = NULL,
  overwrite = FALSE,
  write_file = TRUE
)
```

## Arguments

...	One or more objects returned by <a href="#">svem_select_from_score_table</a> . You may also pass a single list of such objects.
file	Character scalar; path to the CSV file to be written. Required only when write_file = TRUE.
overwrite	Logical; if FALSE (default) and file already exists, an error is thrown. If TRUE, any existing file at file is overwritten. Only used when write_file = TRUE.
write_file	Logical; if TRUE (default), write the combined table to file as CSV and print the full path. If FALSE, no file is written and file may be NULL; the concatenated data.frame is still returned (invisibly).

**Value**

Invisibly, the data.frame that was written to CSV (or would be written, when write\_file = FALSE).

**Examples**

```
# 1) Load example data
data(lipid_screen)

# 2) Build a deterministic expansion using bigexp_terms()
spec <- bigexp_terms(
  Potency ~ PEG + Helper + Ionizable + Cholesterol +
    Ionizable_Lipid_Type + N_P_ratio + flow_rate,
  data      = lipid_screen,
  factorial_order = 3, # up to 3-way interactions
  polynomial_order = 3, # include up to cubic terms I(X^2), I(X^3)
  include_pc_2way = TRUE,
  include_pc_3way = FALSE
)

# 3) Shared deterministic expansion for all three responses
form_pot <- bigexp_formula(spec, "Potency")
form_siz <- bigexp_formula(spec, "Size")
form_pdi <- bigexp_formula(spec, "PDI")

# 4) Fit SVEM models
set.seed(1)
fit_pot <- SVEMnet(form_pot, lipid_screen)
fit_siz <- SVEMnet(form_siz, lipid_screen)
fit_pdi <- SVEMnet(form_pdi, lipid_screen)

objs <- list(Potency = fit_pot, Size = fit_siz, PDI = fit_pdi)

# 5) Multi-response goals (DS desirabilities under the hood)
goals <- list(
  Potency = list(goal = "max", weight = 0.6),
  Size    = list(goal = "min", weight = 0.3),
  PDI     = list(goal = "min", weight = 0.1)
)

# 6) Mixture constraints on the four lipid components
mix <- list(list(
  vars = c("PEG", "Helper", "Ionizable", "Cholesterol"),
  lower = c(0.01, 0.10, 0.10, 0.10),
  upper = c(0.05, 0.60, 0.60, 0.60),
  total = 1.0
))

# 7) Optional process-mean specifications for a design-space example
specs_ds <- list(
  Potency = list(lower = 78),
  Size    = list(upper = 100),
```

```
PDI      = list(upper = 0.25)
)

# 8) Random-search scoring (predictions stored in *_pred columns)
set.seed(3)
scored <- svem_score_random(
  objects      = objs,
  goals        = goals,
  data         = lipid_screen,
  n            = 2500,
  mixture_groups = mix,
  level        = 0.95,
  combine      = "geom",
  numeric_sampler = "random",
  specs        = specs_ds,
  verbose      = FALSE
)

# 9) Build several selection objects from the scored table

# High-score optimal medoids (user-weighted score)
opt_sel <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "score",
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.02,
  label       = "round1_score_optimal"
)

# High-uncertainty exploration medoids
explore_sel <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "uncertainty_measure",
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.05,
  label       = "round1_explore"
)

# High joint mean-in-spec medoids (design-space view)
inspec_sel <- svem_select_from_score_table(
  score_table = scored$score_table,
  target      = "p_joint_mean",
  direction   = "max",
  k           = 5,
  top_type    = "frac",
  top         = 0.10,
  label       = "round1_inspec"
)
```

```

# Best existing screened run (from original_data_scored; k <= 0 -> no medoids)
best_existing <- svem_select_from_score_table(
  score_table = scored$original_data_scored,
  target      = "score",
  direction   = "max",
  k           = 0,
  top_type    = "frac",
  top         = 1.0,
  label       = "round1_existing_best"
)

# 10) Combine all selection objects in a single list
candidate_sels <- list(
  opt_sel,
  explore_sel,
  inspec_sel,
  best_existing
)

# 11a) Export all candidates to CSV for the next experimental round
# svem_export_candidates_csv(
#   candidate_sels,
#   file          = "lipid_screen_round1_candidates.csv",
#   overwrite     = FALSE,
#   write_file    = TRUE
# )

# 11b) Or inspect the combined table in-memory without writing a file
cand_tbl <- svem_export_candidates_csv(
  candidate_sels,
  write_file = FALSE
)
head(cand_tbl)

# 11c) Alternatively, pass selection objects directly as separate arguments
cand_tbl2 <- svem_export_candidates_csv(
  opt_sel,
  explore_sel,
  inspec_sel,
  best_existing,
  write_file = FALSE
)
head(cand_tbl2)

```

## Description

Summarizes variable-selection stability across SVEM bootstrap refits by computing the percentage of bootstrap iterations in which each coefficient (excluding the intercept) is nonzero, using a small tolerance. Optionally produces a quick **ggplot2** summary and/or prints a compact table.

## Usage

```
svem_nonzero(object, tol = 1e-07, plot = TRUE, print_table = TRUE, ...)
```

## Arguments

object	An object of class <code>svem_model</code> with a non-empty <code>\$coef_matrix</code> component. <code>svem_nonzero()</code> is not defined for <code>svem_cv</code> objects.
tol	Numeric tolerance for "nonzero". Coefficients with $ \text{beta}  > \text{tol}$ are counted as nonzero. Default is $1e-7$ .
plot	Logical; if TRUE, draws a quick <b>ggplot2</b> summary plot of the nonzero percentages (default TRUE).
print_table	Logical; if TRUE, prints a compact table of nonzero percentages to the console (default TRUE).
...	Unused; included for future extension.

## Details

This function expects `object$coef_matrix` to contain the per-bootstrap coefficients (including an intercept column), typically created by [SVEMnet](#) when `save_boot = TRUE` (or similar) is enabled. Rows correspond to bootstrap fits; columns correspond to coefficients.

Internally, `svem_nonzero()`:

- checks for and drops rows of `coef_matrix` that contain any non-finite values, to keep summaries stable;
- drops an "(Intercept)" column if present;
- computes  $100 * \text{mean}(|\text{beta}_j| > \text{tol})$  across bootstrap rows for each remaining coefficient.

The plot is a simple line + point chart with labels, ordered by decreasing nonzero percentage. It is intended as a quick diagnostic; for publication graphics, you may want to customize the output data frame with your own plotting code.

## Value

Invisibly returns a data frame with columns:

- Variable: coefficient name (excluding the intercept).
- Percent of Bootstraps Nonzero: percentage (0–100) of bootstrap fits in which  $|\text{beta}| > \text{tol}$ .

If no non-intercept coefficients are found (for example, if only the intercept is present), an empty data frame is returned and a message is issued.

**See Also**

[coef.svem\\_model](#) for ensemble-averaged (optionally debiased) coefficients.

**Examples**

```
## ----- Gaussian demo -----
set.seed(10)
n <- 220
x1 <- rnorm(n)
x2 <- rnorm(n)
x3 <- rnorm(n)
eps <- rnorm(n, sd = 0.4)
y <- 0.7 + 1.5*x1 - 0.8*x2 + 0.05*x3 + eps
dat <- data.frame(y, x1, x2, x3)

fit <- SVEMnet(y ~ (x1 + x2 + x3)^2, data = dat,
              nBoot = 40, relaxed = TRUE)

# Table + plot of bootstrap nonzero percentages
nz <- svem_nonzero(fit, tol = 1e-7, plot = TRUE, print_table = TRUE)
head(nz)

## ----- Binomial demo -----
set.seed(11)
n <- 260
x1 <- rnorm(n)
x2 <- rnorm(n)
x3 <- rnorm(n)
lp <- -0.3 + 0.9*x1 - 0.6*x2 + 0.2*x3
p <- 1/(1+exp(-lp))
y <- rbinom(n, 1, p)
dat_b <- data.frame(y, x1, x2, x3)

fit_b <- SVEMnet(y ~ x1 + x2 + x3, data = dat_b,
                family = "binomial", nBoot = 40, relaxed = TRUE)

# Still summarizes bootstrap selection frequencies for binomial fits
svem_nonzero(fit_b, plot = TRUE, print_table = TRUE)
```

---

svem\_random\_table\_multi

*Generate a Random Prediction Table from Multiple SVEMnet Models  
(no refit)*

---

**Description**

Samples the original predictor factor space cached in fitted `svem_model` objects and computes predictions from each model at the same random points. This is intended for multiple responses

built over the same factor space and a deterministic factor expansion (for example via a shared `bigexp_terms`), so that a shared sampling schema is available.

### Usage

```
svem_random_table_multi(
  objects,
  n = 1000,
  mixture_groups = NULL,
  debias = FALSE,
  range_tol = 1e-08,
  numeric_sampler = c("random", "uniform")
)
```

### Arguments

<code>objects</code>	A list of fitted <code>svem_model</code> objects returned by <code>SVEMnet()</code> . Each object must contain a valid <code>\$sampling_schema</code> produced by the updated <code>SVEMnet()</code> implementation. A single model is also accepted and treated as a length-one list.
<code>n</code>	Number of random points to generate (rows in the output tables). Default is 1000.
<code>mixture_groups</code>	Optional list of mixture constraint groups. Each group is a list with elements <code>vars</code> , <code>lower</code> , <code>upper</code> , <code>total</code> (see <i>Notes on mixtures</i> ). Mixture variables must be numeric-like and must also appear in the models' <code>predictor_vars</code> (that is, they must be used as predictors in all models).
<code>debias</code>	Logical; if TRUE, apply each model's calibration during prediction when available (for Gaussian fits). This is passed to <code>predict.svem_model()</code> . Default is FALSE.
<code>range_tol</code>	Numeric tolerance for comparing numeric ranges across models (used when checking that all <code>\$sampling_schema\$num_ranges</code> agree). Default is 1e-8.
<code>numeric_sampler</code>	Sampler for non-mixture numeric predictors: "random" (default), or "uniform". <ul style="list-style-type: none"> <li>"random": random Latin hypercube when the <b>lhs</b> package is available; otherwise independent uniforms via <code>runif()</code>.</li> <li>"uniform": independent uniform draws within numeric ranges (fastest; no <b>lhs</b> dependency).</li> </ul>

### Details

No refitting is performed. Predictions are obtained by averaging per-bootstrap member predictions on the requested scale.

All models must share an identical predictor schema. Specifically, their `$sampling_schema` entries must agree on:

- The same `predictor_vars` in the same order.
- The same `var_classes` for each predictor.
- Identical factor levels and level order for all categorical predictors.

- Numeric num\_ranges that match within range\_tol for all continuous predictors.
- When present, the same blocking set (up to order).

The function stops with an informative error message if any of these checks fail.

**Discrete numeric predictors (automatic).** If any supplied model stores discrete-numeric sampling information in its \$sampling\_schema, this function will automatically respect it (no separate user argument).

In the updated SVEMnet() implementation this information is stored as:

- \$sampling\_schema\$discrete\_numeric: a character vector of discrete numeric variable names; and
- \$sampling\_schema\$discrete\_levels: a named list mapping those names to allowed numeric values.

(Older objects may use \$sampling\_schema\$discrete\_values instead of discrete\_levels; this function accepts both for backward compatibility.)

Discrete numeric variables are sampled independently (uniform over allowed values) and are excluded from Latin hypercube sampling; LHS (when used) is applied only to the remaining continuous numeric predictors. Discrete numeric variables are not allowed to be mixture variables.

Models may be Gaussian or binomial. For binomial fits, predictions are returned on the probability scale (that is, on the response scale) by default, consistent with the default behaviour of predict.svem\_model().

## Value

A list with three data frames:

- data: the sampled predictor settings, one row per random point.
- pred: one column per response, aligned to data rows.
- all: cbind(data, pred) for convenience.

Each prediction column is named by the model's response (left-hand side) with a "\_pred" suffix (for example, "y1\_pred"). If that name would collide with a predictor name or with another prediction column, the function stops with an error and asks the user to rename the response or predictor.

## Typical workflow

1. Build a deterministic expansion (for example with `bigexp_terms`) and fit several SVEMnet() models for different responses on the same factor space, using the same expansion / sampling settings.
2. Ensure that the fitted models were created with a version of SVEMnet() that populates \$sampling\_schema.
3. Collect the fitted models in a list and pass them to svem\_random\_table\_multi().
4. Use \$data (predictors), \$pred (response columns), or \$all (cbind(data, pred)) for downstream plotting, summarization, or cross-response comparison.

### Blocking variables

If the models were fit using a `bigexp_spec` that included blocking variables (for example `blocking = c("Operator", "Plate_ID")`) and `SVEMnet()` stored these in `$sampling_schema$blocking`, then `svem_random_table_multi()` will:

- treat those variables as blocking factors; and
- hold them fixed at a single value across the sampled table.

Specifically:

- For blocking numeric variables, the function uses the midpoint of the recorded numeric range,  $(\min + \max) / 2$ , for all rows. If the variable also has stored discrete support, the midpoint is snapped deterministically to the nearest allowed discrete value.
- For blocking categorical variables, the function uses a single reference level equal to the most frequent observed level (mode) in the training data, with ties broken deterministically; if the mode is unavailable, it falls back to the first stored level.

Blocking variables are not allowed to appear in `mixture_groups`. If any mixture group tries to use a blocking variable, the function stops with an error.

When no blocking information is present in `$sampling_schema` (for example for models fit without a `bigexp_spec` or without `blocking`), the behavior is unchanged from earlier versions: all predictors are sampled according to the rules described under "Sampling strategy".

---

svem_score_random	<i>Random-search scoring for SVEM models</i>
-------------------	--

---

### Description

Draw random points from the SVEM sampling schema, compute multi-response desirability scores and (optionally) whole-model-test (WMT) reweighted scores, and attach a scalar uncertainty measure based on percentile CI widths. This function does *not* choose candidates; see [svem\\_select\\_from\\_score\\_table](#) for selection and clustering.

Predictions used inside this scorer are always generated with debiasing disabled (i.e., `debias = FALSE`) regardless of whether the underlying SVEM fits support calibration.

When `specs` is supplied, the function also attempts to append mean-level "in spec" probabilities and related joint indicators using the SVEM bootstrap ensemble via `svem_append_design_space_cols`. These quantities reflect uncertainty on the *process mean* at each sampled setting under the fitted SVEM models, not unit-level predictive probabilities. If any error occurs in this spec-limit augmentation, it is caught; a message may be issued when `verbose = TRUE`, and the affected table(s) are returned without the spec-related columns.

**Usage**

```
svem_score_random(
  objects,
  goals,
  data = NULL,
  n = 50000,
  mixture_groups = NULL,
  level = 0.95,
  combine = c("geom", "mean"),
  numeric_sampler = c("random", "uniform"),
  wmt = NULL,
  verbose = TRUE,
  specs = NULL
)
```

**Arguments**

- |         |   |
|---------|---|
| objects | List of svem_model objects (from <a href="#">SVEMnet</a> ). When unnamed, svem_score_random() attempts to infer response names from the left-hand sides of the model formulas. Names (when present) are treated as response identifiers and should typically match the model response names. All models must share a common sampling schema (predictor set, factor levels, numeric ranges) compatible with <a href="#">svem_random_table_multi</a> .  |
| goals   | <p>List of per-response goal specifications. Either:</p> <ul style="list-style-type: none"> <li>• a named list, where names may be either names(objects) or the left-hand-side response names from the fitted models; or</li> <li>• an unnamed list with the same length as objects, in which case entries are matched to models by position.</li> </ul> <p>Each goals[[response]] must be a list with at least:</p> <ul style="list-style-type: none"> <li>• goal: one of "max", "min", "target";</li> <li>• weight: nonnegative numeric weight.</li> </ul> <p>For goal = "target", also provide target. Optional Derringer–Suich controls:</p> <ul style="list-style-type: none"> <li>• For "max" or "min": lower_acceptable, upper_acceptable, shape.</li> <li>• For "target": tol (symmetric), or tol_left / tol_right, and shape_left / shape_right.</li> </ul> <p>When anchors/tolerances are not supplied, robust defaults are inferred from the sampled table using the q0.02–q0.98 span.</p> |
| data    | Optional data frame. When supplied (regardless of whether wmt is used), it is scored and returned as original_data_scored, with predictions (in <resp>_pred columns), per-response desirabilities, score (and wmt_score if applicable), and uncertainty_measure appended. When specs is supplied and the spec-limit augmentation succeeds, the same mean-level spec columns as in score_table (per-response <resp>_p_in_spec_mean, <resp>_in_spec_point, and joint_p_joint_mean, joint_in_spec_point) are appended as well.   |
| n       | Number of random samples to draw in the predictor space. This is the number of rows in the sampled table used for scoring.  |

mixture_groups	Optional mixture and simplex constraints passed to <code>svem_random_table_multi</code> . Each group typically specifies mixture variable names, bounds, and a total.
level	Confidence level for percentile intervals used in the CI width and uncertainty calculations. Default 0.95.
combine	How to combine per-response desirabilities into a scalar score. One of: <ul style="list-style-type: none"> <li>• "geom": weighted geometric mean (default);</li> <li>• "mean": weighted arithmetic mean.</li> </ul>
numeric_sampler	Character string controlling how numeric predictors are sampled inside <code>svem_random_table_multi</code> . One of: <ul style="list-style-type: none"> <li>• "random": Latin hypercube sampling when <code>lhs</code> is available, otherwise independent uniforms;</li> <li>• "uniform": independent uniforms over stored numeric ranges.</li> </ul>
wmt	Optional object returned by <code>svem_wmt_multi</code> . When non-NULL, its multipliers (and <code>p_values</code> , if present) are aligned to <code>names(objects)</code> and used to define WMT weights, <code>wmt_score</code> . When NULL, only user weights are used and no WMT reweighting is applied.
verbose	Logical; if TRUE, print a compact summary of the run (and any WMT diagnostics from upstream) to the console.
specs	Optional named list of specification objects, one per response in <code>objects</code> for which you want to define a mean-level spec constraint. Each entry should be either NULL (no specs for that response) or a list with components: <ul style="list-style-type: none"> <li>• lower: numeric lower limit (may be <code>-Inf</code>, <code>NA</code>, or <code>NULL</code> for a one-sided upper spec);</li> <li>• upper: numeric upper limit (may be <code>Inf</code>, <code>NA</code>, or <code>NULL</code> for a one-sided lower spec).</li> </ul> Names of specs, when provided, should be a subset of <code>names(objects)</code> or of the model response names (left-hand sides). The specification structure matches that used by <code>svem_append_design_space_cols</code> .

## Details

**Typical workflow:** A common pattern is:

1. Fit one or more `SVEMnet()` models for the responses of interest.
2. Call `svem_score_random()` to:
  - draw candidate settings in factor space,
  - compute Derringer–Suich (DS) desirabilities and a combined multi-response score, and
  - attach a scalar uncertainty measure derived from percentile CI widths.
3. Optionally provide specs to append mean-level "in spec" probabilities and joint indicators based on the SVEM bootstrap ensemble (process-mean assurance).
4. Use `svem_select_from_score_table` to:
  - select one "best" row (e.g., maximizing score or `wmt_score`), and
  - pick a small, diverse set of medoid candidates for optimality or exploration (e.g. high uncertainty measure).

5. Run selected candidates, append the new data, refit the SVEM models, and repeat as needed.

**Multi-response desirability scoring:** Each response is mapped to a Derringer–Suich desirability  $d_r \in [0, 1]$  according to its goal:

- goal = "max": larger values are better;
- goal = "min": smaller values are better;
- goal = "target": values near a target are best.

Per-response anchors (acceptable lower/upper limits or target-band tolerances) can be supplied in goals; when not provided, robust defaults are inferred from the sampled responses using the q0.02–q0.98 span.

Per-response desirabilities are combined into a single scalar score using either:

- a weighted arithmetic mean (combine = "mean"), or
- a weighted geometric mean (combine = "geom"), with a small floor applied inside the log to avoid  $\log(0)$ .

User-provided weights in goals[[resp]]\$weight are normalized to sum to one and always define weights\_original and the user-weighted score.

**Whole-model reweighting (WMT):** When a WMT object from svem\_wmt\_multi is supplied via the wmt argument, each response receives a multiplier derived from its whole-model p-value. Final WMT weights are proportional to the product of the user weight and the multiplier, then renormalized to sum to one:

$$w_r^{(\text{final})} \propto w_r^{(\text{user})} \times m_r,$$

where  $m_r$  comes from wmt\$multipliers. The user weights always define score; the WMT-adjusted weights define wmt\_score. The uncertainty measure is always weighted using the user weights, even when WMT is supplied.

**Binomial responses.** If any responses are fitted with family = "binomial", supplying a non-NULL wmt object is not allowed and the function stops with a clear error. Predictions and CI bounds for binomial responses are interpreted on the probability (response) scale and clamped to  $[0, 1]$  before desirability and uncertainty calculations.

**Uncertainty measure:** The uncertainty\_measure is a weighted sum of robustly normalized percentile CI widths across responses. For each response, we compute the bootstrap percentile CI width  $\text{CIwidth}_r(x) = u_r(x) - \ell_r(x)$  and then map it to the unit interval using an affine rescaling based on the empirical q0.02 and q0.98 quantiles of the CI widths for that response (computed from the table being scored):

$$\tilde{W}_r(x) = \frac{\min\{\max(\text{CIwidth}_r(x), q_{0.02}(r)), q_{0.98}(r)\} - q_{0.02}(r)}{q_{0.98}(r) - q_{0.02}(r)}.$$

The scalar uncertainty\_measure is then

$$\text{uncertainty}(x) = \sum_r w_r \tilde{W}_r(x),$$

where  $w_r$  are the user-normalized response weights derived from goals[[resp]]\$weight. Larger values of uncertainty\_measure indicate settings where the ensemble CI is relatively wide compared to the response's typical scale and are natural targets for exploration.

**Spec-limit mean-level probabilities:** If `specs` is provided, `svem_score_random()` attempts to pass the scored table and models to `svem_append_design_space_cols` to compute, for each response with an active spec:

- `<resp>p_in_spec_mean`: estimated probability (under the SVEM bootstrap ensemble) that the process mean at a setting lies within the specified interval;
- `<resp>_in_spec_point`: 0/1 indicator that the point prediction lies within the same interval.

and joint quantities:

- `p_joint_mean`: product of per-response mean-level probabilities over responses with active specs;
- `joint_in_spec_point`: 0/1 indicator that all point predictions are in spec across responses with active specs.

Names in `specs` may refer either to `names(objects)` or to the model response names; they are automatically aligned to the fitted models.

These probabilities are defined on the *conditional means* at each sampled setting, not on individual units or lots, and are best interpreted as ensemble-based assurance measures under the SVEM + FRW pipeline. If the augmentation step fails for any reason (for example, missing predictor columns or incompatible models), the error is caught; a message may be issued when `verbose = TRUE`, and `score_table` and/or `original_data_scored` are returned without the spec-related columns.

## Value

A list with components:

`score_table` Data frame with predictors, predicted responses (columns `<resp>_pred` for each `resp` in `names(objects)`), per-response desirabilities, score, optional `wmt_score`, and `uncertainty_measure`. For each response `r` in `names(objects)`, additional columns `r_lwr`, `r_upr` (percentile CI bounds at level `level`) and `r_ciw_w` (weighted, normalized CI width contribution to `uncertainty_measure`) are appended. When `specs` is supplied and the spec-limit augmentation succeeds, additional columns `<resp>p_in_spec_mean`, `<resp>_in_spec_point`, `p_joint_mean`, and `joint_in_spec_point` are appended.

`original_data_scored` If data is supplied, that data augmented with prediction columns `<resp>_pred`, per-response desirabilities, score, optional `wmt_score`, and `uncertainty_measure`; otherwise NULL. When `specs` is supplied and the spec-limit augmentation succeeds, the same mean-level spec columns as in `score_table` are appended to `original_data_scored` as well.

`weights_original` User-normalized response weights.

`weights_final` Final weights after WMT, if `wmt` is supplied; otherwise equal to `weights_original`.

`wmt_p_values` Named vector of per-response whole-model p-values when `wmt` is supplied and contains `p_values`; otherwise NULL.

`wmt_multipliers` Named vector of per-response WMT multipliers when `wmt` is supplied; otherwise NULL.

## See Also

[SVEMnet](#), [svem\\_random\\_table\\_multi](#), [svem\\_select\\_from\\_score\\_table](#), [svem\\_append\\_design\\_space\\_cols\(\)](#), [svem\\_wmt\\_multi](#)

**Examples**

```

## -----
## Multi-response SVEM scoring with Derringer–Suich desirabilities
## -----

data(lipid_screen)

# Build a deterministic expansion once and reuse for all responses
spec <- bigexp_terms(
  Potency ~ PEG + Helper + Ionizable + Cholesterol +
  Ionizable_Lipid_Type + N_P_ratio + flow_rate,
  data      = lipid_screen,
  factorial_order = 3,
  polynomial_order = 3,
  include_pc_2way = TRUE,
  include_pc_3way = FALSE
)

form_pot <- bigexp_formula(spec, "Potency")
form_siz <- bigexp_formula(spec, "Size")
form_pdi <- bigexp_formula(spec, "PDI")

set.seed(1)
fit_pot <- SVEMnet(form_pot, lipid_screen)
fit_siz <- SVEMnet(form_siz, lipid_screen)
fit_pdi <- SVEMnet(form_pdi, lipid_screen)

# Collect SVEM models in a named list by response
objs <- list(Potency = fit_pot, Size = fit_siz, PDI = fit_pdi)

# Targets and user weights for Derringer–Suich desirabilities
goals <- list(
  Potency = list(goal = "max", weight = 0.6),
  Size    = list(goal = "min", weight = 0.3),
  PDI     = list(goal = "min", weight = 0.1)
)

# Optional mixture constraints (composition columns sum to 1)
mix <- list(list(
  vars = c("PEG", "Helper", "Ionizable", "Cholesterol"),
  lower = c(0.01, 0.10, 0.10, 0.10),
  upper = c(0.05, 0.60, 0.60, 0.60),
  total = 1.0
))

# Basic random-search scoring without WMT or design-space specs
set.seed(3)
scored_basic <- svem_score_random(
  objects      = objs,
  goals        = goals,
  n            = 10000,          # number of random candidates
  mixture_groups = mix,

```

```

combine      = "geom",
numeric_sampler = "random",
verbose      = FALSE
)

# Scored candidate table: predictors, <resp>_pred, <resp>_des, score, uncertainty
names(scored_basic$score_table)
head(scored_basic$score_table)

# Scored original data (if 'data' is supplied)
# scored_basic$original_data_scored contains predictions + desirabilities

## -----
## With whole-model tests (WMT) and process-mean specifications
## -----

set.seed(123)
wmt_out <- svem_wmt_multi(
  formulas      = list(Potency = form_pot,
                      Size     = form_siz,
                      PDI      = form_pdi),
  data          = lipid_screen,
  mixture_groups = mix,
  wmt_control   = list(seed = 123),
  plot         = FALSE,
  verbose      = FALSE
)

# Simple process-mean specs for a joint design space:
# Potency >= 78, Size <= 100, PDI <= 0.25
specs_ds <- list(
  Potency = list(lower = 78),
  Size    = list(upper = 100),
  PDI     = list(upper = 0.25)
)

set.seed(4)
scored_full <- svem_score_random(
  objects      = objs,
  goals        = goals,
  data         = lipid_screen, # score the original runs as well
  n            = 25000,
  mixture_groups = mix,
  level        = 0.95,
  combine      = "geom",
  numeric_sampler = "random",
  wmt          = wmt_out,      # optional: WMT reweighting
  specs        = specs_ds,    # optional: design-space columns
  verbose      = TRUE
)

# The scored table now includes:
# * score, wmt_score, uncertainty_measure

```

```

# * per-response CIs: <resp>_lwr, <resp>_upr
# * design-space columns, e.g. Potency_p_in_spec_mean, p_joint_mean
names(scored_full$score_table)

## -----
## Positional (unnamed) goals matched to objects by position
## -----

data(lipid_screen)

# Build a deterministic expansion once and reuse for all responses
spec <- bigexp_terms(
  Potency ~ PEG + Helper + Ionizable + Cholesterol +
    Ionizable_Lipid_Type + N_P_ratio + flow_rate,
  data      = lipid_screen,
  factorial_order = 3,
  polynomial_order = 3,
  include_pc_2way = TRUE,
  include_pc_3way = FALSE
)

form_pot <- bigexp_formula(spec, "Potency")
form_siz <- bigexp_formula(spec, "Size")
form_pdi <- bigexp_formula(spec, "PDI")

set.seed(1)
fit_pot <- SVEMnet(form_pot, lipid_screen)
fit_siz <- SVEMnet(form_siz, lipid_screen)
fit_pdi <- SVEMnet(form_pdi, lipid_screen)

# Collect SVEM models in a list.
# Here goals will be matched by position: Potency, Size, PDI.
objs <- list(fit_pot, fit_siz, fit_pdi)

# Positional goals (unnamed list): must have same length as 'objects'
goals_positional <- list(
  list(goal = "max", weight = 0.6), # for Potency (objs[[1]])
  list(goal = "min", weight = 0.3), # for Size   (objs[[2]])
  list(goal = "min", weight = 0.1)  # for PDI    (objs[[3]])
)

set.seed(5)
scored_pos <- svem_score_random(
  objects      = objs,
  goals        = goals_positional,
  n            = 5000,
  numeric_sampler = "random",
  verbose      = FALSE
)

names(scored_pos$score_table)

```

---

svem\_select\_from\_score\_table

*Select best row and diverse candidates from an SVEM score table*


---

## Description

Given a scored random-search table (e.g. from `svem_score_random()`), pick a single "best" row under a chosen objective column and sample a small, diverse set of medoid candidates from the top of that ranking. Any per-response CI columns (e.g. `*_lwr / *_upr`) present in `score_table` are carried through unchanged.

Optionally, a string label can be supplied to annotate the returned best row and candidates by appending that label to a "Notes\_from\_SVEMnet" column. If "Notes\_from\_SVEMnet" is missing, it is created. If it exists and is nonempty, the label is appended with "; " as a separator.

## Usage

```
svem_select_from_score_table(
  score_table,
  target = "score",
  direction = c("max", "min"),
  k = 5,
  top_type = c("frac", "n"),
  top = 0.1,
  predictor_cols = NULL,
  label = NULL
)
```

## Arguments

<code>score_table</code>	Data frame with predictors, responses, scores, and <code>uncertainty_measure</code> , typically <code>scored\$score_table</code> from <code>svem_score_random</code> . When medoids are requested ( $k > 0$ ), the predictor columns used for clustering are taken from the "svem_predictor_cols" attribute by default. If that attribute is missing, a heuristic is used. If you accidentally pass the full scored list, a helpful error is thrown reminding you to use <code>scored\$score_table</code> .
<code>target</code>	Character scalar naming the column in <code>score_table</code> to optimize (e.g. "score", "wmt_score", "uncertainty_measure").
<code>direction</code>	Either "max" or "min" indicating whether larger or smaller values of target are preferred.
<code>k</code>	Integer; desired number of medoid candidates to return. If $k \leq 0$ , only the best row is returned and no clustering is performed.
<code>top_type</code>	Either "frac" or "n" specifying whether top is a fraction of rows or an integer count.

top	Value for the top set: a fraction in (0,1] if top_type = "frac", or an integer >= 1 if top_type = "n".
predictor_cols	Optional character vector of predictor column names used to measure diversity in the PAM step when $k > 0$ . When NULL (default), the function first tries <code>attr(score_table, "svem_predictor_cols")</code> . If that is unavailable, it falls back to a heuristic that prefers non-derived predictor columns (excluding e.g. *_pred, *_des, *_lwr, *_upr, *_ciw_w, *_p_in_spec_mean, *_in_spec_point, score, wmt_score, uncertainty_measure, p_joint_mean, joint_in_spec_point, candidate_type, selection_label, Notes_from_SVEMnet). If no usable predictor columns can be inferred, a warning is issued and only best is returned.
label	Optional character scalar. When non-NULL, this label is appended into a "Notes_from_SVEMnet" column for the returned best row and candidates. If "Notes_from_SVEMnet" is missing, it is created; if present and nonempty, the label is appended using ";" as separator.

### Value

A list with components:

**best** One-row data frame at the optimum of target under the specified direction, including any columns present in score\_table (e.g. \*\_lwr / \*\_upr).

**candidates** Data frame of medoid candidates (possibly empty or NULL) drawn from the top top of the ranking on target, with all columns carried through from score\_table.

**call** The matched call, including all arguments used to create this selection object.

### See Also

[svem\\_score\\_random](#), [svem\\_select\\_candidates\(\)](#)

---

svem\_significance\_test\_parallel

*SVEM whole-model significance test with mixture support (parallel)*

---

### Description

Perform a permutation-based whole-model significance test for a continuous (Gaussian) SVEM fit, with optional mixture-factor groups and parallel SVEM refits.

### Usage

```
svem_significance_test_parallel(
  formula,
  data,
  mixture_groups = NULL,
  nPoint = 2000,
  nSVEM = 10,
```

```

nPerm = 150,
percent = 90,
nBoot = 100,
glmnet_alpha = c(1),
weight_scheme = c("SVEM"),
objective = c("wAIC", "wBIC", "wSSE", "auto"),
relaxed = FALSE,
verbose = TRUE,
nCore = parallel::detectCores() - 2,
seed = NULL,
spec = NULL,
response = NULL,
use_spec_contrasts = TRUE,
...
)

```

### Arguments

formula	A model formula. If spec is provided, the right-hand side is ignored and replaced by the locked expansion in spec.
data	A data frame containing the variables in the model.
mixture_groups	Optional list describing one or more mixture-factor groups. Each element should be a list with components: <ul style="list-style-type: none"> <li>vars: character vector of column names;</li> <li>lower: numeric vector of lower bounds (same length as vars);</li> <li>upper: numeric vector of upper bounds (same length as vars);</li> <li>total: scalar specifying the sum of the mixture variables.</li> </ul> All mixture variables must appear in exactly one group. Defaults to NULL.
nPoint	Number of random evaluation points in the factor space (default 2000).
nSVEM	Number of SVEM fits on the original (unpermuted) data used to summarize the observed surface (default 10).
nPerm	Number of SVEM fits on permuted responses used to build the null reference distribution (default 150).
percent	Percentage of variance to capture in the SVD of the permutation surfaces (default 90).
nBoot	Number of bootstrap iterations within each inner SVEM fit (default 100).
glmnet_alpha	Numeric vector of glmnet alpha values (default c(1)).
weight_scheme	Weighting scheme for SVEM (default "SVEM"). Passed to SVEMnet().
objective	Objective used inside SVEMnet() to pick the bootstrap path solution. One of "wAIC", "wBIC", or "wSSE" (default "wAIC").
relaxed	Logical; default FALSE. When TRUE, inner SVEMnet() fits use glmnet's relaxed elastic-net path and select both lambda and relaxed gamma on each bootstrap. When FALSE, the standard glmnet path is used. If relaxed = TRUE and glmnet_alpha includes 0, ridge (alpha = 0) is dropped by SVEMnet() for relaxed fits.

verbose	Logical; if TRUE, display progress messages (default TRUE).
nCore	Number of CPU cores for parallel processing. Default is <code>parallel::detectCores() - 2</code> , with a floor of 1.
seed	Optional integer seed for reproducible RNG (default NULL). When supplied, the master RNG kind is set to "L'Ecuyer-CMRG" (with <code>sample.kind = "Rounding"</code> when supported), and deterministic per-iteration seeds are generated on the master and applied inside each parallel <code>%dopar%</code> iteration via <code>set.seed()</code> . This yields reproducibility regardless of parallel scheduling and core count.
spec	Optional <code>bigexp_spec</code> created by <code>bigexp_terms()</code> . If provided, the test reuses its locked expansion. The working formula becomes <code>bigexp_formula(spec, response_name)</code> , where <code>response_name</code> is taken from <code>response</code> if supplied, otherwise from the left-hand side of formula. Categorical sampling uses <code>spec\$levels</code> , and numeric sampling prefers <code>spec\$num_range</code> when available. Discrete numeric predictors recorded by <code>bigexp_terms()</code> ( <code>spec\$settings\$discrete_numeric + spec\$settings\$discrete_levels</code> ) are sampled only from their recorded allowed levels when building the evaluation grid.
response	Optional character name for the response variable to use when <code>spec</code> is supplied. If omitted, the response is taken from the left-hand side of formula.
use_spec_contrasts	Logical; default TRUE. When <code>spec</code> is supplied and <code>use_spec_contrasts = TRUE</code> , the function replays <code>spec\$settings\$contrasts_options</code> on the parallel workers for deterministic factor coding.
...	Additional arguments passed to <code>SVEMnet()</code> and then to <code>glmnet()</code> (for example: <code>penalty.factor</code> , <code>offset</code> , <code>lower.limits</code> , <code>upper.limits</code> , <code>standardize.response</code> , etc.). The relaxed setting is controlled by the <code>relaxed</code> argument of this function and any relaxed value passed via <code>...</code> is ignored with a warning.

## Details

The procedure follows Karl (2024): it generates a space-filling evaluation grid in the factor space, fits multiple SVEM models on the original data and on permuted responses, standardizes grid predictions, reduces them via an SVD-based low-rank representation, and summarizes each fit by a Mahalanobis-type distance in the reduced space. A flexible SHASHo distribution is then fit to the permutation distances and used to obtain a whole-model  $p$ -value for the observed surface.

Because the test is based on a finite number of permutations and a fitted null distribution, the reported  $p$ -values are approximate and are intended as a diagnostic measure of global factor signal, not as exact hypothesis tests.

All SVEM refits (for the original and permuted responses) are run in parallel using `foreach + doParallel`.

Reproducible parallel RNG (Windows/macOS/Linux): when `seed` is supplied, the function sets the master RNG to `RNGkind("L'Ecuyer-CMRG", sample.kind = "Rounding")` (falling back to "L'Ecuyer-CMRG" on older R), and generates a deterministic, per-iteration seed schedule on the master. Each parallel `foreach` iteration then calls `set.seed()` with its assigned seed before performing any random draws (including permutations and the bootstrap randomness inside `SVEMnet()`). This makes results reproducible regardless of worker scheduling (including `preschedule = FALSE`) and independent of the number of cores.

The function can optionally reuse a deterministic, locked expansion built with `bigexp_terms()`. Supply `spec` (and optionally `response`) to ensure that categorical levels, contrasts, and the polynomial/interaction structure are identical across repeated calls and across multiple responses sharing the same factor space.

Although the implementation calls `SVEMnet()` internally and will technically run for any supported family, the significance test is *designed* for continuous (Gaussian) responses and should be interpreted in that setting.

## Value

An object of class "svem\_significance\_test", a list with components:

- `p_value`: median whole-model  $p$ -value across the `nSVEM` original SVEM fits.
- `p_values`: numeric vector of length `nSVEM` with the per-fit  $p$ -values.
- `d_Y`: numeric vector of distances for the original SVEM fits.
- `d_pi_Y`: numeric vector of distances for the permutation fits.
- `distribution_fit`: fitted SHASHo distribution object.
- `data_d`: data frame of distances and source labels (original vs permutation), suitable for plotting.

## Acknowledgments

OpenAI's GPT models (o1-preview through GPT-5 Pro) were used to assist with coding and roxygen documentation; all content was reviewed and finalized by the author.

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### See Also

[SVEMnet](#), [bigexp\\_terms](#), [bigexp\\_formula](#)

### Examples

```
set.seed(1)

# Small toy data with a 3-component mixture A, B, C
n <- 40
sample_trunc_dirichlet <- function(n, lower, upper, total) {
  k <- length(lower)
  stopifnot(length(upper) == k, total >= sum(lower), total <= sum(upper))
  avail <- total - sum(lower)
  if (avail <= 0) return(matrix(rep(lower, each = n), nrow = n))
  out <- matrix(NA_real_, n, k)
  i <- 1L
  while (i <= n) {
    g <- rgamma(k, 1, 1)
    w <- g / sum(g)
    x <- lower + avail * w
    if (all(x <= upper + 1e-12)) { out[i, ] <- x; i <- i + 1L }
  }
  out
}

lower <- c(0.10, 0.20, 0.05)
upper <- c(0.60, 0.70, 0.50)
total <- 1.0
ABC <- sample_trunc_dirichlet(n, lower, upper, total)
```

```

A <- ABC[, 1]; B <- ABC[, 2]; C <- ABC[, 3]
X <- runif(n)
F <- factor(sample(c("red", "blue"), n, replace = TRUE))
y <- 2 + 3*A + 1.5*B + 1.2*C + 0.5*X + 1*(F == "red") + rnorm(n, sd = 0.3)
dat <- data.frame(y = y, A = A, B = B, C = C, X = X, F = F)

mix_spec <- list(list(
  vars = c("A", "B", "C"),
  lower = lower,
  upper = upper,
  total = total
))

## Example 1: direct formula interface (no locked expansion spec)
res1 <- svem_significance_test_parallel(
  y ~ A + B + C + X + F,
  data = dat,
  mixture_groups = mix_spec,
  glmnet_alpha = 1,
  weight_scheme = "SVEM",
  objective = "auto",
  relaxed = FALSE, # default, shown for clarity
  nCore = 2,
  seed = 123,
  verbose = FALSE
)
res1$p_value

## Example 2: using a deterministic bigexp expansion spec
## Build a wide expansion once and reuse it via `spec`
spec <- bigexp_terms(
  y ~ A + B + C + X + F,
  data = dat,
  factorial_order = 2, # up to 2-way interactions
  polynomial_order = 2 # up to quadratic terms in continuous vars
)

## Run the same significance test, but with the locked expansion:
## - `formula` is still required, but its RHS is ignored when `spec` is given
## - `response` tells the helper which LHS to use with `spec`
res2 <- svem_significance_test_parallel(
  y ~ A + B + C + X + F,
  data = dat,
  mixture_groups = mix_spec,
  glmnet_alpha = 1,
  weight_scheme = "SVEM",
  objective = "auto",
  relaxed = FALSE,
  nCore = 2,
  seed = 123,
  spec = spec,
  response = "y",
  use_spec_contrasts = TRUE,

```

```

    verbose          = FALSE
  )
  res2$p_value

```

---

svem_wmt_multi	<i>Whole-model tests for multiple SVEM responses (WMT wrapper)</i>
----------------	--

---

## Description

Convenience wrapper around [svem\\_significance\\_test\\_parallel](#) for running whole-model tests (WMT) on multiple responses that share the same dataset and mixture constraints. This helper:

- takes a formula or a list of formulas and a single data frame,
- calls `svem_significance_test_parallel()` for each response,
- extracts per-response p-values and converts them to WMT multipliers via a chosen transform, and
- optionally plots the WMT objects together using `plot.svem_significance_test` and prints a compact summary of p-values and multipliers.

The resulting multipliers vector is designed to be passed directly to downstream scoring functions (for example, as an optional WMT argument to `svem_score_random()`), with response names matched by `names()`.

## Usage

```

svem_wmt_multi(
  formulas,
  data,
  mixture_groups = NULL,
  wmt_transform = c("neglog10", "one_minus_p"),
  wmt_control = list(seed = 123),
  plot = TRUE,
  verbose = TRUE
)

```

## Arguments

- |                             |   |
|-----------------------------|---|
| <code>formulas</code>       | A single formula or a (preferably named) list of formulas, one per response. If unnamed, response names are inferred from the left-hand side of each formula; non-unique names are made unique.         |
| <code>data</code>           | Data frame containing the predictors and responses referenced in formulas.  |
| <code>mixture_groups</code> | Optional mixture and simplex constraints passed to <a href="#">svem_significance_test_parallel</a> .  |
| <code>wmt_transform</code>  | Character; transformation used to convert WMT p-values into multipliers. One of: <ul style="list-style-type: none"> <li>• "neglog10": <math>f(p) = [-\log_{10}(p)]^{\text{strength}}</math>,</li> </ul> |

- "one\_minus\_p":  $f(p) = (1 - p)^{\text{strength}}$ .

Currently, strength = 1 is used internally.

wmt_control	Optional list of extra arguments passed directly to <a href="#">svem_significance_test_parallel</a> . By default this is <code>list(seed = 123)</code> so that WMT calls are reproducible; you may override or extend this (e.g. <code>list(seed = 999, nPerm = 300)</code> ). Any entries not recognized by <code>svem_significance_test_parallel</code> are ignored by that function.
plot	Logical; if TRUE (default), attempt to plot all successfully computed WMT objects together via <a href="#">plot.svem_significance_test</a> .
verbose	Logical; if TRUE (default), print progress and a compact summary of p-values and multipliers.

### Value

A list of class "svem\_wmt\_multi" with components:

wmt_objects	Named list of WMT objects (one per response), as returned by <code>svem_significance_test_parallel()</code> . Entries are NULL where a WMT call failed.
p_values	Named numeric vector of per-response p-values (bounded away from 0/1), or NA when unavailable.
multipliers	Named numeric vector of per-response WMT multipliers derived from the p-values using <code>wmt_transform</code> .
wmt_transform	The transformation used.
wmt_control	The list of arguments passed through to <code>svem_significance_test_parallel()</code> .

### See Also

[svem\\_significance\\_test\\_parallel](#), [plot.svem\\_significance\\_test](#)

### Examples

```
data(lipid_screen)

spec <- bigexp_terms(
  Potency ~ PEG + Helper + Ionizable + Cholesterol +
    Ionizable_Lipid_Type + N_P_ratio + flow_rate,
  data      = lipid_screen,
  factorial_order = 3,
  polynomial_order = 3,
  include_pc_2way = TRUE,
  include_pc_3way = FALSE
)

form_pot <- bigexp_formula(spec, "Potency")
form_siz <- bigexp_formula(spec, "Size")
form_pdi <- bigexp_formula(spec, "PDI")

mix <- list(list(
```

```

vars = c("PEG", "Helper", "Ionizable", "Cholesterol"),
lower = c(0.01, 0.10, 0.10, 0.10),
upper = c(0.05, 0.60, 0.60, 0.60),
total = 1.0
))

set.seed(123)
wmt_out <- svem_wmt_multi(
  formulas      = list(Potency = form_pot,
                      Size    = form_siz,
                      PDI     = form_pdi),
  data          = lipid_screen,
  mixture_groups = mix,
  wmt_transform = "neglog10",
  wmt_control   = list(seed = 123),
  plot          = TRUE
)

wmt_out$p_values
wmt_out$multipliers

## later: pass wmt_out$multipliers into svem_score_random()

```

---

with\_bigexp\_contrasts *Evaluate code with the spec's recorded contrast options*

---

## Description

with\_bigexp\_contrasts() temporarily restores the contrasts options that were active when the spec was built, runs a block of code, and then restores the original options. This is useful when a modeling function uses the global options("contrasts") to decide how to encode factors (for example, lm(), glm(), or other modeling functions that call model.matrix() internally).

## Usage

```
with_bigexp_contrasts(spec, code)
```

## Arguments

spec	A "bigexp_spec" object with stored contrasts_options in settings.
code	Code to evaluate with temporarily restored options.

## Examples

```

set.seed(1)
df4 <- data.frame(
  y = rnorm(10),

```

```
X1 = rnorm(10),
G = factor(sample(c("A", "B"), 10, replace = TRUE))
)

spec4 <- bigexp_terms(
  y ~ X1 + G,
  data = df4,
  factorial_order = 2,
  polynomial_order = 2
)

with_bigexp_contrasts(spec4, {
  mm4 <- model.matrix(spec4$formula, df4)
  head(mm4)
})
```

# Index

- \* **SVEM methods**
  - predict.svem\_model, 32
- \* **datasets**
  - lipid\_screen, 22
- \* **package**
  - SVEMnet-package, 2
  
- bigexp\_formula, 3, 6, 11, 12, 23, 69
- bigexp\_prepare, 3, 7, 11, 12, 14
- bigexp\_terms, 3, 7, 8, 9, 14, 23, 54, 55, 69
- bigexp\_train, 3, 12, 14
  
- coef.svem\_model, 3, 15, 53
  
- glmnet\_with\_cv, 4, 16
  
- lipid\_screen, 4, 22
  
- plot.svem\_binomial, 28
- plot.svem\_model, 3, 30
- plot.svem\_significance\_test, 4, 31, 71, 72
- predict.svem\_cv (predict\_cv), 35
- predict.svem\_model, 3, 32
- predict\_cv, 35
- print.bigexp\_spec, 37
- print.svem\_significance\_test, 39
  
- svem\_export\_candidates\_csv, 3, 4, 48
- svem\_nonzero, 3, 16, 51
- svem\_random\_table\_multi, 4, 53, 57, 58, 60
- svem\_score\_random, 3–5, 23, 56, 64, 65
- svem\_select\_from\_score\_table, 3, 4, 23, 48, 56, 58, 60, 64
- svem\_significance\_test\_parallel, 3, 4, 65, 71, 72
- svem\_wmt\_multi, 3–5, 23, 58–60, 71
- SVEMnet, 3, 15, 34, 39, 52, 57, 60, 69
- SVEMnet-package, 2
  
- with\_bigexp\_contrasts, 3, 73