

Package: lgspline (via r-universe)

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Type Package

Title Lagrangian Multiplier Smoothing Splines for Smooth Function Estimation

Version 1.1.0

Description Implements Lagrangian multiplier smoothing splines for flexible nonparametric regression and function estimation. Provides tools for fitting, prediction, and inference using a constrained optimization approach to enforce smoothness. Supports generalized linear models, Weibull accelerated failure time (AFT) models, Cox proportional hazards models, quadratic programming constraints, and customizable working-correlation structures, with options for parallel fitting. The core spline construction builds on Ezhov et al. (2018) <doi:10.1515/jag-2017-0029>. Quadratic-programming and SQP details follow Goldfarb & Idnani (1983) <doi:10.1007/BF02591962> and Nocedal & Wright (2006) <doi:10.1007/978-0-387-40065-5>. For smoothing spline and penalized spline background, see Wahba (1990) <doi:10.1137/1.9781611970128> and Wood (2017) <doi:10.1201/9781315370279>. For variance-component and correlation-parameter estimation, see Searle et al. (2006) <ISBN:978-0470009598>. The default multivariate partitioning step uses k-means clustering as in MacQueen (1967).

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coef.lgspline	<i>Extract Coefficients from a Fitted lgspline</i>
---------------	--

Description

Returns the per-partition polynomial coefficient lists from a fitted lgspline model.

Usage

```
## S3 method for class 'lgspline'
coef(object, ...)
```

Arguments

object	A fitted lgspline model object.
...	Not used.

Details

Coefficient names reflect the polynomial expansion terms, e.g.:

- intercept
- v: linear term for predictor v
- v^2: quadratic term
- v^3: cubic term
- _v_x_w_: two-way interaction

Column/variable names replace numeric indices when available.

To get all coefficients as a single matrix: `Reduce('cbind', coef(model_fit))`.

Value

A list of per-partition coefficient vectors. Returns NULL with a warning if `object$B` is not found.

See Also

[lgspline](#)

Examples

```
set.seed(1234)
t <- runif(1000, -10, 10)
y <- 2*sin(t) + -0.06*t^2 + rnorm(length(t))
model_fit <- lgspline(t, y)

coefficients <- coef(model_fit)
print(coefficients[[1]])
print(Reduce('cbind', coefficients))
```

`coef.wald_lgspline` *Extract Coefficients from a wald_lgspline Object*

Description

Extract Coefficients from a wald_lgspline Object

Usage

```
## S3 method for class 'wald_lgspline'
coef(object, ...)
```

Arguments

<code>object</code>	A "wald_lgspline" object.
<code>...</code>	Not used.

Value

Named numeric vector of coefficient estimates, or NULL if no estimate column is available.

confint.lgspline	<i>Confidence Intervals for lgspline Coefficients</i>
------------------	---

Description

Wald-based confidence intervals for regression coefficients and, when available, correlation parameters (on the working scale).

Usage

```
## S3 method for class 'lgspline'
confint(object, parm, level = 0.95, ...)
```

Arguments

object	A fitted lgspline object with return_varcovmat = TRUE.
parm	Optional vector of parameter indices or names. Default returns all regression parameters; working-scale correlation parameters are appended when available.
level	Confidence level. Default 0.95.
...	Additional arguments passed to wald_univariate .

Details

For Gaussian identity-link models, t-distribution quantiles are used with effective degrees of freedom $N - \text{trace}(\mathbf{XUGX}^\top)$. All other families use normal quantiles.

Correlation parameter intervals (if `VhalfInv_params_estimates` and `VhalfInv_params_vcov` are present) are computed on the unbounded working scale via a Wald interval.

Value

A matrix with columns giving lower and upper confidence limits, named e.g. 2.5% and 97.5% for 95% intervals. When available, rows for working-scale correlation parameters are appended after the regression coefficients.

confint.wald_lgspline *Extract Confidence Intervals from a wald_lgspline Object*

Description

Extract Confidence Intervals from a wald_lgspline Object

Usage

```
## S3 method for class 'wald_lgspline'
confint(object, parm = NULL, level = NULL, ...)
```

Arguments

object	A "wald_lgspline" object.
parm	Parameter specification (ignored; all returned).
level	Confidence level (ignored; uses the object's critical value).
...	Not used.

Value

Matrix with columns lower and upper, or NULL if confidence limits are not available.

cox_dispersion_function
Cox PH Dispersion Function

Description

Returns 1 unconditionally. Cox PH has no dispersion parameter; this function exists solely for interface compatibility with lgspline's dispersion_function argument.

Usage

```
cox_dispersion_function(
  mu,
  y,
  order_indices,
  family,
  observation_weights,
  VhalfInv,
  ...
)
```

Arguments

mu	Predicted values.
y	Observed survival times.
order_indices	Observation indices.
family	Family object.
observation_weights	Observation weights.
VhalfInv	Inverse square root of correlation matrix.
...	Additional arguments (including status).

Value

Scalar 1.

cox_family	<i>Cox Proportional Hazards Family for lgspline</i>
------------	---

Description

Creates a family-like object for Cox PH models. The link function is log (the linear predictor is log-relative-hazard), but unlike standard GLM families there is no dispersion parameter and no closed-form mean-variance relationship.

The family provides `$loglik` and `$aic` methods compatible with [logLik.lgspline](#).

Usage

```
cox_family()
```

Details

Cox PH is semiparametric: the baseline hazard is unspecified. The partial log-likelihood depends only on the order of event times and the linear predictor $\eta = \mathbf{X}\beta$. Consequently:

- No dispersion parameter is estimated (`sigma_sq_tilde` is fixed at 1).
- `dev.resids` returns martingale-style residuals for GCV tuning compatibility.
- The response variable is survival time (positive), and the link is log.

Value

A list with family components used by `lgspline`.

Examples

```
fam <- cox_family()
fam$family
fam$link
```

 cox_glm_weight_function

Cox PH GLM Weight Function

Description

Computes working weights for the Cox PH information matrix, used by lgspline when updating **G** after obtaining constrained estimates. The weights are a diagonal approximation built from the Breslow tied-event information contributions.

Usage

```
cox_glm_weight_function(
  mu,
  y,
  order_indices,
  family,
  dispersion,
  observation_weights,
  status
)
```

Arguments

mu	Predicted values (exp(eta), i.e., relative hazard).
y	Observed survival times.
order_indices	Observation indices in partition order.
family	Cox family object (unused, for interface compatibility).
dispersion	Dispersion parameter (fixed at 1 for Cox PH).
observation_weights	Observation weights.
status	Event indicator (1 = event, 0 = censored).

Details

For a tied event-time block g , the diagonal approximation uses

$$W_{jj}^{(g)} = d_g^{(w)} \frac{h_j}{S_g} \left(1 - \frac{h_j}{S_g}\right), \quad j \in R_g$$

where $h_j = w_j \exp(\eta_j)$, $S_g = \sum_{k \in R_g} h_k$, and $d_g^{(w)} = \sum_{i \in D_g} w_i$.

When the natural weights are degenerate (all zero or non-finite), falls back to a vector of ones.

Value

Numeric vector of working weights, length N.

Examples

```
## Used internally by lgspline; see cox_helpers examples below.
```

cox_helpers	<i>Cox Proportional Hazards Helpers for lgspline</i>
-------------	--

Description

Functions for fitting Cox proportional hazards regression models within the lgspline framework. Analogous to the Weibull AFT helpers, these provide the partial log-likelihood, score, information, and all interface functions needed by lgspline's unconstrained fitting, penalty tuning, and inference machinery.

cox_qp_score_function	<i>Cox PH Score Function for Quadratic Programming and Blockfit</i>
-----------------------	---

Description

Computes the score (gradient of partial log-likelihood) in the format expected by lgspline's qp_score_function interface. The block-diagonal design matrix X and response y are in partition order; this function internally sorts by event time, computes the Cox score using the Breslow approximation for tied event times, and returns the result in the original partition order.

Usage

```
cox_qp_score_function(  
  X,  
  y,  
  mu,  
  order_list,  
  dispersion,  
  VhalfInv,  
  observation_weights,  
  status  
)
```

Arguments

X	Block-diagonal design matrix ($N \times P$).
y	Response vector (survival times, $N \times 1$).
μ	Predicted values ($N \times 1$), same order as X and y .
order_list	List of observation indices per partition.
dispersion	Dispersion (fixed at 1).

VhalfInv Inverse square root correlation matrix (NULL for independent observations).
 observation_weights Observation weights.
 status Event indicator (1 = event, 0 = censored).

Value

Numeric column vector of length P (gradient w.r.t. coefficients).

cox_schur_correction *Cox PH Schur Correction*

Description

Returns zero corrections for all partitions. Cox PH has no nuisance dispersion parameter, so no Schur complement correction to the information matrix is needed. This function exists for interface compatibility with lgspline's schur_correction_function.

Usage

```
cox_schur_correction(  
  X,  
  y,  
  B,  
  dispersion,  
  order_list,  
  K,  
  family,  
  observation_weights,  
  ...  
)
```

Arguments

X List of partition design matrices.
 y List of partition response vectors.
 B List of partition coefficient vectors.
 dispersion Dispersion (fixed at 1).
 order_list List of observation indices per partition.
 K Number of knots.
 family Family object.
 observation_weights Observation weights.
 ... Additional arguments.

Value

List of K+1 zeros.

create_onehot	<i>Create One-Hot Encoded Matrix</i>
---------------	--------------------------------------

Description

Converts a categorical vector into a one-hot encoded matrix where each unique value becomes a binary column.

Usage

```
create_onehot(x, drop_first = FALSE)
```

Arguments

x	A vector containing categorical values (factors, character, etc.)
drop_first	Logical; if TRUE and more than one dummy column is created, drop the first column.

Details

The function creates dummy variables for each unique value in the input vector using `model.matrix()` with dummy-intercept coding. Column names are cleaned by removing the 'x' prefix added by `model.matrix()`.

Value

A data frame containing the one-hot encoded binary columns with cleaned column names

Examples

```
## lgspline will not accept this format of "catvar", because inputting data
# this way can cause difficult-to-diagnose issues in formula parsing
# all variables must be numeric
df <- data.frame(numvar = rnorm(100),
                 catvar = rep(LETTERS[1:4],
                             25))

print(head(df))

## Instead, replace with dummy-intercept coding by
# 1) applying one-hot encoding
# 2) dropping the first column
# 3) appending to our data

dummy_intercept_coding <- create_onehot(df$catvar)[,-1]
```

```
df$catvar <- NULL
df <- cbind(df, dummy_intercept_coding)
print(head(df))
```

Description

This document gives mathematical and implementation details for Lagrangian Multiplier Smoothing Splines (LMSS) as implemented in **lgspline**.

LMSS fits smoothing splines directly in a monomial basis. Smoothness is imposed by external equality constraints rather than by embedding knots in a specialized spline basis. This keeps the fitted model interpretable at the coefficient level, but moves some complexity into constraint construction and constrained estimation.

B-spline fits can be converted to monomial form in principle, but multivariate tensor-product conversions can be high-dimensional, numerically unstable, and unavailable in standard software.

Statistical Problem Formulation

Consider an $N \times q$ matrix of predictors $\mathbf{T} = (\mathbf{t}_1, \dots, \mathbf{t}_N)^\top$ and an $N \times 1$ response vector $\mathbf{y} = (y_1, \dots, y_N)^\top$. We assume the relationship follows a generalized linear model with unknown smooth function f :

$$y_i \sim \mathcal{D}(g^{-1}(f(\mathbf{t}_i)), \sigma^2)$$

where \mathcal{D} is a distribution (e.g. exponential family or related) with mean $\mu_i = g^{-1}(f(\mathbf{t}_i))$, link function $g(\cdot)$, and dispersion parameter σ^2 . For Gaussian response with identity link, observations are independently distributed as $y_i | \mathbf{t}_i, \sigma^2 \sim \mathcal{N}(f(\mathbf{t}_i), \sigma^2)$.

The objective is to estimate f by:

1. Partitioning the predictor space into $K + 1$ mutually exclusive regions.
2. Fitting local polynomial models within each partition.
3. Enforcing smoothness at partition boundaries via Lagrangian multipliers.
4. Penalizing the integrated squared second derivative to discourage roughness.

Unlike other smoothing spline formulations, no post-fitting algebraic rearrangement or disentanglement of a spline basis is needed to obtain interpretable models. The polynomial expansions are homogeneous across partitions, and the relationship between predictor and response is explicit at the coefficient level.

To anchor the notation, in the single-predictor cubic case one would write

$$\hat{f}(t_i) = \hat{\beta}_{(0)} + \hat{\beta}_{(1)}t_i + \hat{\beta}_{(2)}t_i^2 + \hat{\beta}_{(3)}t_i^3 = \mathbf{x}_i^\top \hat{\boldsymbol{\beta}},$$

where $\mathbf{x}_i = (1, t_i, t_i^2, t_i^3)^\top$. The LMSS formulation preserves exactly this kind of polynomial representation, but now does so within each partition and then forces neighboring pieces to agree in the smoothness conditions described below.

Core notation used throughout:

- $\mathbf{y}_{(N \times 1)}$: Response vector.
- $\mathbf{T}_{(N \times q)}$: Matrix of predictors.
- $\mathbf{X}_{(N \times P)}$: Block-diagonal matrix of polynomial expansions, with diagonal blocks \mathbf{X}_k of dimension $n_k \times p$.
- $\mathbf{\Lambda}_{(P \times P)}$: Block-diagonal penalty matrix, with blocks $\mathbf{\Lambda}_k$ of dimension $p \times p$.
- $\hat{\boldsymbol{\beta}}_{(P \times 1)}$: Unconstrained penalized estimate.
- $\tilde{\boldsymbol{\beta}}_{(P \times 1)}$: Constrained coefficient estimates.
- $\mathbf{G}_{(P \times P)}$: Block-diagonal matrix with blocks $\mathbf{G}_k = (\mathbf{X}_k^\top \mathbf{W}_k \mathbf{D}_k \mathbf{X}_k + \mathbf{\Lambda}_k)^{-1}$, where \mathbf{W}_k and \mathbf{D}_k are defined below.
- $\mathbf{A}_{(P \times r)}$: Constraint matrix encoding smoothness conditions. Reduced to linearly independent columns via pivoted QR decomposition by default, or by the `parallel_qr` reduced-system route when that option is active.
- $\mathbf{U}_{(P \times P)}$: $\mathbf{I} - \mathbf{G}\mathbf{A}(\mathbf{A}^\top \mathbf{G}\mathbf{A})^{-1} \mathbf{A}^\top$.
- $\mathbf{D}_{(N \times N)}$: Diagonal matrix of user-supplied observation weights (`observation_weights` or `weights`). Defaults to the identity. These play the role of prior precision on individual observations: a weight of 2 is equivalent to seeing that observation twice.
- $\mathbf{W}_{(N \times N)}$: Diagonal matrix of GLM working weights. In the implementation these diagonal entries are whatever is returned by `glm_weight_function`; by default this is the GLM working weight `family$mu.eta(eta)^2 / family$variance(mu)`, optionally multiplied by user-supplied observation weights. For Gaussian response with identity link, $\mathbf{W} = \mathbf{I}$. For other families, \mathbf{W} depends on the current fitted values and is updated at each Newton-Raphson iteration. For the common canonical families used by default, this matches the familiar Fisher-scoring weighting role.
- $\mathbf{V}_{(N \times N)}$: Correlation matrix of errors. When no correlation structure is specified, $\mathbf{V} = \mathbf{I}$. Otherwise supplied via `VhalfInv` or estimated through `VhalfInv_fxn`.

In the Gaussian identity case with unit weights and no correlation, $\mathbf{G}_k = (\mathbf{X}_k^\top \mathbf{X}_k + \mathbf{\Lambda}_k)^{-1}$ and most formulas simplify accordingly. When \mathbf{D} or \mathbf{W} appear in a formula, the product $\mathbf{W}\mathbf{D}$ means “GLM working weights times observation weights”; whenever one of them is the identity it drops out.

Before these quantities reach the main fitting stage, the user-facing inputs are parsed, standardized, and organized by `process_input`. When the formula interface is used and `auto_encode_factors = TRUE`, that preprocessing step also relies on helpers such as `create_onehot` to encode factor levels before the design reaches `lgspline.fit()`. The notation in the remainder of this document therefore refers to the internal objects that actually enter `lgspline.fit()`, not necessarily the raw objects originally supplied by the user.

Most user-facing controls can also be supplied through grouped lists (`penalty_args`, `tuning_args`, `expansion_args`, `constraint_args`, `qp_args`, `parallel_args`, `covariance_args`, `return_args`, and `glm_args`); these are unpacked before entering the same fitting pipeline. Setting `dummy_fit = TRUE` runs preprocessing, partitioning, expansion, and penalty construction without estimating nonzero coefficients, which is useful for inspecting X , A , partitions, and penalties before fitting.

Model Formulation and Estimation

Piecewise Polynomial Structure: For K knots (one predictor) or $K + 1$ partitions (multiple predictors) there are $K + 1$ mutually exclusive partitions $\mathcal{P}_0, \dots, \mathcal{P}_K$. Each observation i belongs to exactly one partition. Within partition k , the function is represented as a polynomial of degree $p - 1$ in each predictor:

$$\hat{f}_k(\mathbf{t}) = \mathbf{x}^\top \tilde{\boldsymbol{\beta}}_k$$

where \mathbf{x} collects the polynomial basis terms (intercept, linear, quadratic, cubic, and optionally quartic and interaction terms) and $\tilde{\boldsymbol{\beta}}_k$ are the corresponding coefficients. In one predictor, the same idea can be written more explicitly as

$$\hat{f}(t_i) = \sum_{k=0}^K \mathbf{x}_{ik}^\top \hat{\boldsymbol{\beta}}_k \mathbf{1}(t_i \in \mathcal{P}_k),$$

which highlights that the unconstrained problem is just a collection of local polynomial regressions. The expansions are homogeneous across partitions, so coefficients are directly comparable. This is implemented via [get_polynomial_expansions](#).

The exact contents of \mathbf{x} are controlled by the basis-expansion arguments documented in [lgspline](#): `include_quadratic_terms`, `include_cubic_terms`, `include_quartic_terms`, `include_2way_interactions`, `include_3way_interactions`, `include_quadratic_interactions`, `exclude_interactions_for`, `exclude_these_expansions`, and `custom_basis_fxn`. Likewise, `just_linear_with_interactions` and `just_linear_without_interactions` determine which predictors remain structurally linear even though they still participate in the same partition-wise polynomial bookkeeping described here.

Letting p denote the number of basis terms per partition, $P = p(K + 1)$ is the total number of coefficients. The full design matrix \mathbf{X} and penalty matrix $\boldsymbol{\Lambda}$ are block-diagonal with $K + 1$ blocks, so unconstrained estimation reduces to $K + 1$ independent penalized regressions, which appears as follows for the identity link case:

$$\hat{\boldsymbol{\beta}}_k = \mathbf{G}_k \mathbf{X}_k^\top \mathbf{W}_k \mathbf{D}_k \mathbf{y}_k, \quad \mathbf{G}_k = (\mathbf{X}_k^\top \mathbf{W}_k \mathbf{D}_k \mathbf{X}_k + \boldsymbol{\Lambda}_k)^{-1}.$$

For Gaussian identity with unit weights this reduces to the familiar $\mathbf{G}_k = (\mathbf{X}_k^\top \mathbf{X}_k + \boldsymbol{\Lambda}_k)^{-1}$. The block-diagonal structure means these can be computed in parallel across partitions. In the user-facing interface this is realized by supplying a cluster through `cl`, controlling work splitting with `chunk_size`, and enabling stages such as `parallel_eigen`, `parallel_unconstrained`, `parallel_penalty`, `parallel_make_constraint`, and `parallel_qr`. The `parallel_qr` option does not change the estimator; it replaces one QR decomposition of a tall transformed system with parallel cross-product accumulation and a smaller dense solve, with fallback to base linear algebra when needed. The eigendecomposition and matrix square roots of each \mathbf{G}_k are computed by [compute_G_eigen](#) and can be returned as `G` and `Ghalf`.

Fitted values for the canonical Gaussian case appear as $\tilde{\mathbf{y}} = \mathbf{X}\tilde{\boldsymbol{\beta}} = \mathbf{H}\mathbf{y}$ for $\mathbf{H} = \mathbf{X}\mathbf{U}\mathbf{G}\mathbf{X}^\top$.

Smoothing Constraints and the Constraint Matrix: Without further intervention the piecewise polynomial will be discontinuous. The central idea of LMSS is that smoothness is not hidden inside a special basis, but instead imposed directly where neighboring partitions meet. At each knot $t_{k,k+1}$ between neighboring partitions k and $k + 1$, up to three smoothing constraints are imposed:

1. Continuity: $\mathbf{x}_{k,k+1}^\top \boldsymbol{\beta}_k = \mathbf{x}_{k,k+1}^\top \boldsymbol{\beta}_{k+1}$.

2. First-derivative continuity: $\mathbf{x}'_{k,k+1}\boldsymbol{\beta}_k = \mathbf{x}'_{k,k+1}\boldsymbol{\beta}_{k+1}$.
3. Second-derivative continuity: $\mathbf{x}''_{k,k+1}\boldsymbol{\beta}_k = \mathbf{x}''_{k,k+1}\boldsymbol{\beta}_{k+1}$.

where \mathbf{x}' and \mathbf{x}'' are elementwise first and second derivatives of the basis with respect to \mathbf{t} . For the familiar cubic single-predictor basis $\mathbf{x} = (1, t, t^2, t^3)^\top$, these derivative vectors are

$$\mathbf{x}' = (0, 1, 2t, 3t^2)^\top, \quad \mathbf{x}'' = (0, 0, 2, 6t)^\top.$$

With K knots this yields up to $3K$ scalar constraints for one predictor. With multiple spline-expanded predictors, the default combines the corresponding first- and second-derivative rows before adding them to the equality matrix. With one spline-expanded predictor and additional non-spline effects, derivative constraints are kept separate by default. The flag `add_first_and_second_derivative_constraints` can force either construction. The constraints are collected as linear equations $\mathbf{A}^\top \boldsymbol{\beta} = \mathbf{0}$ in a $P \times r$ matrix \mathbf{A} . The constraint matrix is built by `make_constraint_matrix` and returned in the fitted object as `A`.

In higher dimensions or with many partitions, separate derivative constraints can over-specify the fit and push it toward a single global polynomial. Combining first- and second-derivative rows reduces redundant equality constraints while retaining a smooth join condition. The constraint level can be controlled via `include_constrain_fitted`, `include_constrain_first_deriv`, `include_constrain_second_deriv`, and `add_first_and_second_derivative_constraints`. The companion flag `include_constrain_interactions` determines whether the analogous mixed-partial constraints are imposed for interaction terms, and `no_intercept` adds the special homogeneous equality constraint that fixes the intercept at zero (the same behavior triggered by using `0 +` in the formula interface).

Before computing the projection \mathbf{U} , the constraint matrix is optionally reduced to a linearly independent subset of columns. When `qr_pivot_smoothing_constraints = TRUE` (the default), this uses pivoted QR decomposition; when `parallel_qr = TRUE` and a cluster is supplied, the code instead partitions the rows of \mathbf{A} into blocks $\mathbf{A}_{(1)}, \dots, \mathbf{A}_{(B)}$ and computes

$$\mathbf{A}^\top \mathbf{A} = \sum_{b=1}^B \mathbf{A}_{(b)}^\top \mathbf{A}_{(b)}$$

in parallel. The linearly independent constraint basis is then recovered from that reduced system, so the package avoids running one full serial QR on a tall redundant matrix whenever the parallel route is well-conditioned. If the reduced system appears unstable, the code falls back to pivoted QR. This avoids numerical instability from redundant constraints and ensures $\mathbf{A}^\top \mathbf{G} \mathbf{A}$ is invertible. When `qr_pivot_smoothing_constraints = FALSE`, the original equality columns are carried forward unchanged.

Lagrangian Projection: The constrained estimate is derived via Lagrangian multipliers. Define the $P \times P$ projection matrix:

$$\mathbf{U} = \mathbf{I} - \mathbf{G} \mathbf{A} (\mathbf{A}^\top \mathbf{G} \mathbf{A})^{-1} \mathbf{A}^\top.$$

Then the constrained estimate is:

$$\tilde{\boldsymbol{\beta}} = \mathbf{U} \hat{\boldsymbol{\beta}}.$$

The matrix \mathbf{U} has the property that $\mathbf{U} \mathbf{G} \mathbf{U}^\top = \mathbf{U} \mathbf{G}$, which is used extensively in variance estimation and posterior draws. In words, the unconstrained penalized estimate is projected back into the coefficient space that satisfies the smoothness restrictions, and all subsequent uncertainty

calculations inherit that same projected geometry. The projection is computed via `get_U` and, when requested, returned in the fitted object as `U` through `return_U = TRUE`.

When the constraints are inhomogeneous ($\mathbf{A}^\top \boldsymbol{\beta} = \mathbf{c}$ with $\mathbf{c} \neq \mathbf{0}$), a particular solution $\boldsymbol{\beta}_0$ satisfying $\mathbf{A}^\top \boldsymbol{\beta}_0 = \mathbf{c}$ is added back after projection, yielding the full Lagrangian solution $\mathbf{U}\hat{\boldsymbol{\beta}} + (\mathbf{I} - \mathbf{U})\boldsymbol{\beta}_0$. In `lgspline` and `lgspline.fit`, users realize this by supplying extra equality columns in `constraint_vectors` together with matching right-hand sides in `constraint_values`; `null_constraint` provides the alternate shorthand documented in `lgspline` when `constraint_vectors` is supplied and `constraint_values` is left empty.

In practice \mathbf{U} is never explicitly formed during fitting. The constrained estimate is obtained from a transformed OLS residual problem (the $\mathbf{G}^{1/2}\mathbf{r}^*$ trick) in four steps:

1. Obtain the unconstrained partition-wise unconstrained estimate $\hat{\boldsymbol{\beta}}$.
2. Set $\mathbf{y}^* = \mathbf{G}^{-1/2}\hat{\boldsymbol{\beta}}$ and $\mathbf{X}^* = \mathbf{G}^{1/2}\mathbf{A}$.
3. Fit the linear model $\mathbb{E}[\mathbf{y}^*] = \mathbf{X}^*\boldsymbol{\gamma}$ by OLS using QR decomposition, or with the `parallel_qr` cross-product route when that option is enabled.
4. Compute the residuals $\mathbf{r}^* = \mathbf{y}^* - \mathbf{X}^*(\mathbf{X}^{*\top}\mathbf{X}^*)^{-1}\mathbf{X}^{*\top}\mathbf{y}^*$ from that transformed OLS fit and recover the constrained estimate by $\tilde{\boldsymbol{\beta}} = \mathbf{G}^{1/2}\mathbf{r}^*$.

A scaling factor $1/\sqrt{K+1}$ is applied to both \mathbf{X}^* and \mathbf{y}^* prior to the OLS call and divided out afterward, keeping the absolute scale moderate when the constraint matrix has many rows.

This is where `parallel_qr` enters most directly. Write the rows of the transformed system as $\mathbf{X}_{(1)}^*, \dots, \mathbf{X}_{(B)}^*$ and $\mathbf{y}_{(1)}^*, \dots, \mathbf{y}_{(B)}^*$. The default route fits Step 3 with a standard QR decomposition of the full tall matrix \mathbf{X}^* . When `parallel_qr = TRUE`, the package instead forms

$$\mathbf{M} = \mathbf{X}^{*\top}\mathbf{X}^* = \sum_{b=1}^B \mathbf{X}_{(b)}^{*\top}\mathbf{X}_{(b)}^*, \quad \mathbf{u} = \mathbf{X}^{*\top}\mathbf{y}^* = \sum_{b=1}^B \mathbf{X}_{(b)}^{*\top}\mathbf{y}_{(b)}^*$$

in parallel, solves the reduced dense system $\mathbf{M}\hat{\boldsymbol{\gamma}} = \mathbf{u}$, and then computes $\mathbf{r}^* = \mathbf{y}^* - \mathbf{X}^*\hat{\boldsymbol{\gamma}}$. So the costly operation being replaced is not the Lagrangian projection itself, but the single tall QR decomposition inside the transformed OLS solve. Algebraically the fit is the same projection; computationally it is a parallel reduction followed by a much smaller dense solve. If that reduced system is judged too unstable, the code falls back to the serial `.lm.fit()` or `qr()` route.

The same idea reappears in other equality-only solves used during tuning, active-set refinement, and constraint reduction. In each case the package replaces one tall QR or least-squares call with a chunked accumulation of cross-products, while keeping the surrounding notation and estimator unchanged.

The most expensive object in this approach remains the $P \times r$ transformed matrix $\mathbf{X}^* = \mathbf{G}^{1/2}\mathbf{A}$, but either route is far cheaper than working with the full $P \times P$ system directly. Without correlation or SQP constraints, \mathbf{G} is stored and operated upon as a list of $K+1$ small $p \times p$ matrices rather than the full $P \times P$ block-diagonal, saving substantial memory when K is large and allowing for parallelism.

When correlation is present, $\mathbf{X}^\top\mathbf{V}^{-1}\mathbf{X}$ is no longer block-diagonal, so the equality-only solve must use either the full correlated system or, when available, the Woodbury reduction (see `.woodbury_decompose_V`). When additional inequality constraints are present, the same active-set controller is still called first. Block structure now determines only how each equality-only re-solve is carried out: partition-wise when available, Woodbury on the low-rank correlated path, or full-system otherwise. Dense `solve.QP` and damped SQP are retained as fallback paths only when the active-set route is unavailable or does not converge.

GLM Extension and Iterative Updates

Working Quantities: For GLM responses with mean $\mu_i = g^{-1}(\eta_i)$ and working weights $w_i = [V(\mu_i)\{g'(\mu_i)\}^2]^{-1}$, the penalized information becomes $\mathbf{G}_k^{-1} = \mathbf{X}_k^\top \mathbf{W}_k \mathbf{X}_k + \mathbf{\Lambda}_k$ with $\mathbf{W}_k = \text{diag}(w_i : i \in \mathcal{P}_k)$. Because \mathbf{W} is diagonal, $\mathbf{X}^\top \mathbf{W} \mathbf{X}$ remains block-diagonal and the four-step procedure carries through with $\mathbf{G}_k = (\mathbf{X}_k^\top \mathbf{W}_k \mathbf{X}_k + \mathbf{\Lambda}_k)^{-1}$ in place of $(\mathbf{X}_k^\top \mathbf{X}_k + \mathbf{\Lambda}_k)^{-1}$. Under the default `glm_weight_function`, the diagonal entries are `family$mu.eta(eta)^2 / family$variance(mu)`, optionally scaled by observation weights.

The partition-wise unconstrained estimates are obtained by `unconstrained_fit_fxn`, by default `unconstrained_fit_default`, which initializes via an augmented ridge trick (appending $\mathbf{\Lambda}^{1/2}$ as pseudo-observations to `glm.fit`) then refines using `damped_newton_r` with `nr_iterate`. For non-canonical log-link gamma regression, the `keep_weighted_Lambda = TRUE` option correctly returns the augmented ridge estimate directly.

Three Fitting Paths:

Path 1: Correlation structure present. When a working correlation \mathbf{V} is present, such as for marginal means models or generalized estimating equation (GEE)-like models, \mathbf{V}^{-1} couples observations across partitions, so partition-wise fitting is not directly available. Two sub-paths handle this.

Path 1a (Gaussian identity + GEE) solves the whitened system directly via $\tilde{\mathbf{G}} = (\tilde{\mathbf{X}}^\top \tilde{\mathbf{X}} + \mathbf{\Lambda}_{\text{block}})^{-1}$ where $\tilde{\mathbf{X}} = \mathbf{V}^{-1/2} \mathbf{X}$, then applies the Lagrangian projection in the full P -space. See `.get_B_gee_gaussian`.

Path 1b (non-Gaussian GEE) uses a damped SQP iteration on the whitened system. Each quadratic subproblem first attempts the same active-set refinement used elsewhere in the package; the dense `solve.QP` bridge is used only as fallback. See `.get_B_gee_glm`.

Both sub-paths have Woodbury-accelerated variants described below.

Path 2: Gaussian identity, no correlation. The constrained estimate is obtained by a single Lagrangian projection from the per-partition penalized least-squares cross-products (the four steps in closed form). No outer iteration is needed. When $K = 0$ and there are no additional constraints, this reduces to the ordinary penalized closed form $\hat{\boldsymbol{\beta}} = \mathbf{G} \mathbf{X}^\top \mathbf{y}$. Implemented in `.get_B_gaussian_nocorr`.

Path 3: Non-Gaussian GLM, no correlation. Unconstrained estimates are obtained separately within each partition, then projected onto the constraint space. When the link is non-identity, the information \mathbf{G} depends on the current fitted values through the working weights, so the projection must be iterated. The unconstrained anchor $\hat{\boldsymbol{\beta}}$ is held fixed while \mathbf{G} is recomputed at each iterate's constrained estimate:

$$\tilde{\boldsymbol{\beta}}^{(s+1)} = \mathbf{U}^{(s)} \hat{\boldsymbol{\beta}}, \quad \mathbf{U}^{(s)} = \mathbf{I} - \mathbf{G}^{(s)} \mathbf{A} (\mathbf{A}^\top \mathbf{G}^{(s)} \mathbf{A})^{-1} \mathbf{A}^\top.$$

Iteration stops when the mean absolute coefficient change falls below `tol`, or when the update begins increasing (in which case the previous iterate is restored). The recomputation of weighted Gram matrices, Schur corrections, and square-root information factors at each step is handled by `.solver_recompute_G_at_estimate`. Implemented in `.get_B_glm_nocorr`.

Woodbury Acceleration for Structured Correlation: For structured correlation, write the penalized information as

$$\mathbf{G}^{-1} = \mathbf{G}_{\text{on}}^{-1} + \mathbf{G}_{\text{off}}^{-1},$$

where $\mathbf{G}_{\text{on}}^{-1}$ contains the $K + 1$ block-diagonal $p \times p$ pieces of $\mathbf{X}^\top \mathbf{V}^{-1} \mathbf{X} + \mathbf{\Lambda}$, and $\mathbf{G}_{\text{off}}^{-1}$ contains the cross-partition part. Without correlation, $\mathbf{G}_{\text{off}}^{-1} = \mathbf{0}$.

When the cross-partition part is low rank, the code factors it as

$$\mathbf{G}_{\text{off}}^{-1} = \mathbf{E}\mathbf{J}\mathbf{E}^\top,$$

with \mathbf{E} of size $P \times r$ and \mathbf{J} diagonal with entries ± 1 . Then

$$\mathbf{G} = \mathbf{G}_{\text{on}} - \mathbf{G}_{\text{on}}\mathbf{E}(\mathbf{J}^{-1} + \mathbf{E}^\top\mathbf{G}_{\text{on}}\mathbf{E})^{-1}\mathbf{E}^\top\mathbf{G}_{\text{on}}.$$

The only dense inverse is the small $r \times r$ matrix $(\mathbf{J}^{-1} + \mathbf{E}^\top\mathbf{G}_{\text{on}}\mathbf{E})^{-1}$.

Define $\mathbf{N} = \mathbf{G}_{\text{on}}^{1/2}\mathbf{E}$ and a thin QR decomposition $\mathbf{N} = \mathbf{Q}\mathbf{R}$. With $\mathbf{P} = \mathbf{Q}\mathbf{Q}^\top$ and $\mathbf{S} = \mathbf{I}_r + \mathbf{R}\mathbf{J}\mathbf{R}^\top$,

$$\mathbf{F} = (\mathbf{I}_P + \mathbf{N}\mathbf{J}\mathbf{N}^\top)^{-1} = \mathbf{I}_P - \mathbf{P} + \mathbf{Q}\mathbf{S}^{-1}\mathbf{Q}^\top,$$

so that a convenient, generally non-symmetric factor $\mathbf{L}\mathbf{L}^\top = \mathbf{G}$ is

$$\mathbf{L} = \mathbf{G}_{\text{on}}^{1/2}\mathbf{F}^{1/2}, \quad \mathbf{L}^{-1} = \mathbf{F}^{-1/2}\mathbf{G}_{\text{on}}^{-1/2},$$

with

$$\mathbf{F}^{1/2} = (\mathbf{I}_P - \mathbf{P}) + \mathbf{Q}\mathbf{S}^{-1/2}\mathbf{Q}^\top, \quad \mathbf{F}^{-1/2} = (\mathbf{I}_P - \mathbf{P}) + \mathbf{Q}\mathbf{S}^{1/2}\mathbf{Q}^\top.$$

This preserves the same transformed OLS projection used in the independent case, using the transpose orientation for the transformed response and constraint design:

$$\mathbf{y}^* = \mathbf{L}^\top \mathbf{w}, \quad \mathbf{X}^* = \mathbf{L}^\top \mathbf{A}, \quad \tilde{\boldsymbol{\beta}} = \mathbf{L}\mathbf{r}^*,$$

where $\mathbf{w} = \mathbf{X}^\top \mathbf{V}^{-1}\mathbf{y}$. The full-space operations remain the QR on \mathbf{X}^* and a rank- r correction through the manuscript basis \mathbf{Q} .

Implementation map. The code uses the same algebra but stores a numerically convenient representation. The correspondence is:

- manuscript $\mathbf{E} \leftrightarrow$ helper output \mathbf{E} ;
- manuscript diagonal sign matrix $\mathbf{J} \leftrightarrow$ helper output \mathbf{J} ;
- manuscript inverse $(\mathbf{J}^{-1} + \mathbf{E}^\top\mathbf{G}_{\text{on}}\mathbf{E})^{-1} \leftrightarrow$ helper output `inner_inv`;
- manuscript projector basis $\mathbf{Q} \leftrightarrow$ helper output \mathbf{Q} ;
- manuscript projector $\mathbf{P} = \mathbf{Q}\mathbf{Q}^\top$ is not stored explicitly, but is represented through rank- r multiplies involving \mathbf{Q} ;
- manuscript factors $\mathbf{F}^{1/2}$ and $\mathbf{F}^{-1/2}$ are stored internally as $\mathbf{I} - \mathbf{Q}\mathbf{C}\mathbf{Q}^\top$ and $\mathbf{I} + \mathbf{Q}\mathbf{C}_{\text{inv}}\mathbf{Q}^\top$, where \mathbf{C} and \mathbf{C}_{inv} are diagonal.

For the non-Gaussian Woodbury path (`.get_B_gee_glm_woodbury`), the fixed correlation perturbation $\mathbf{V}^{-1} - \mathbf{I}$ and its product with \mathbf{X} are precomputed once and held fixed across Newton iterations. At each step, the weighted perturbation $\mathbf{X}^\top \mathbf{W}(\mathbf{V}^{-1} - \mathbf{I})\mathbf{X}$ is formed using the precomputed product, then split and re-factored via `.woodbury_redecompose_weighted`.

The Woodbury path is used when $r < 2P/3$; otherwise the code falls back to the dense whitened solve. If \mathbf{F} is not positive definite, the dense path is also used. See `.get_B_gee_woodbury` (Gaussian) and `.get_B_gee_glm_woodbury` (non-Gaussian).

Step Control: Different paths use different step-control strategies.

In the GEE paths (Paths 1a and 1b), the coefficient update is damped:

$$\boldsymbol{\beta}^{(s+1)} = (1 - \alpha_s)\boldsymbol{\beta}^{(s)} + \alpha_s\boldsymbol{\beta}_{\text{cand}}^{(s)},$$

with $\alpha_s = 2^{-d_s}$ and d_s increased whenever the candidate gives non-finite or larger deviance. The loop terminates after 10 consecutive rejections or 100 total iterations, and exits early when both coefficient change and deviance improvement fall below `tol` after a burn-in period.

In Path 3 (non-Gaussian, no correlation), the outer projection loop has no line search. The algorithm recomputes \mathbf{G} at the current constrained estimate and applies a fresh projection. If the mean absolute coefficient change begins increasing, the previous iterate is restored and the loop stops. The partition-wise unconstrained estimates themselves are obtained by damped Newton-Raphson inside `unconstrained_fit_default`, where `damped_newton_r` computes a Newton direction once per iteration and halves the step size until the penalized log-likelihood improves.

Accommodating Correlation Structures

Parametric Correlation Structures: Suppose $\text{Cov}(\mathbf{y}) = \sigma^2 \mathbf{V}(\boldsymbol{\theta})$ for a known parametric family indexed by $\boldsymbol{\theta}$ (e.g., AR(1) with $\theta = \rho$, Matern with $\boldsymbol{\theta} = (\ell, \nu)$, exchangeable with $\theta = \rho$). The penalized generalized least-squares problem becomes

$$\min_{\boldsymbol{\beta}} (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})^\top \mathbf{V}^{-1} (\mathbf{y} - \mathbf{X}\boldsymbol{\beta}) + \boldsymbol{\beta}^\top \boldsymbol{\Lambda} \boldsymbol{\beta} \quad \text{s.t. } \mathbf{A}^\top \boldsymbol{\beta} = \mathbf{0}.$$

The correlation matrix \mathbf{V} is supplied through the fitted-object components `Vhalf` and `VhalfInv`, either directly or via user functions `Vhalf_fxn` and `VhalfInv_fxn`. When both are non-NULL, `get_B` dispatches to the GEE paths (Path 1a or 1b). For built-in correlation structures, the required square-root matrices are assembled numerically using `matsqrt`, `matinvsqrt`, and `invert`.

Whitening and Permutation: Because the data are stored in partition ordering (all observations from partition 0, then partition 1, etc.) while \mathbf{V} is in the original observation ordering, a permutation is applied internally: $\mathbf{V}_{\text{perm}}^{-1/2} = \mathbf{V}^{-1/2}[\boldsymbol{\pi}, \boldsymbol{\pi}]$, where $\boldsymbol{\pi} = \text{unlist}(\text{order_list})$ maps original indices to partition-ordered indices. The whitened design and response are $\tilde{\mathbf{X}} = \mathbf{V}_{\text{perm}}^{-1/2} \mathbf{X}_{\text{block}}$ and $\tilde{\mathbf{y}} = \mathbf{V}_{\text{perm}}^{-1/2} \mathbf{y}$.

The \mathbf{X} and \mathbf{y} inputs to `lgspline.fit` are preserved in their unwhitened form. Whitening is applied inside `get_B` and `blockfit_solve` where the full $N \times P$ block-diagonal design is available, since applying $\mathbf{V}^{-1/2}$ to only the diagonal blocks of the partitioned design would silently discard cross-partition contributions and corrupt the Gram matrix.

Loss of Block-Diagonal Structure: Unlike the independent-errors case, $\mathbf{X}^\top \mathbf{V}^{-1} \mathbf{X}$ is not block-diagonal because \mathbf{V}^{-1} introduces cross-partition coupling. The unconstrained estimator

$$\hat{\boldsymbol{\beta}} = (\mathbf{X}^\top \mathbf{V}^{-1} \mathbf{X} + \boldsymbol{\Lambda})^{-1} \mathbf{X}^\top \mathbf{V}^{-1} \mathbf{y}$$

requires a full $P \times P$ solve, and the constraint projection proceeds with $\mathbf{G} = (\mathbf{X}^\top \mathbf{V}^{-1} \mathbf{X} + \boldsymbol{\Lambda})^{-1}$. For structured correlation matrices (AR(1), exchangeable, banded), the perturbation $\mathbf{V}^{-1} - \mathbf{I}$ is sparse and the cross-partition coupling has low effective rank. In these cases the Woodbury-accelerated paths (`.get_B_gee_woodbury`, `.get_B_gee_glm_woodbury`) recover the partition-wise computational structure by decomposing the coupling into a block-diagonal correction plus a low-rank remainder, as described in the GLM Extension section. For dense or high-rank \mathbf{V}^{-1} , the code falls back to the full whitened system (`.get_B_gee_gaussian`, `.get_B_gee_glm`).

GEE Deviance Monitoring: For non-Gaussian models with correlation, the deviance used for convergence monitoring is computed in the whitened space by `.bf_gee_deviance`. When the

family supplies `custom_dev.resids`, the raw deviance residuals $r_i = \text{sign}(d_i)\sqrt{|d_i|}$ are divided by $\sqrt{w_i}$ and pre-multiplied by $\mathbf{V}_{\text{perm}}^{-1/2}$ before squaring and averaging:

$$D_{\text{GEE}} = \frac{1}{N} \left\| \mathbf{V}_{\text{perm}}^{-1/2} \mathbf{w}^{-1/2} \mathbf{r} \right\|^2,$$

where \mathbf{w} is the vector of working weights at the current iterate, clamped below at $\sqrt{\varepsilon_{\text{mach}}}$. When only `dev.resids` is available, the function falls back to the standard mean deviance; otherwise it uses mean squared error.

REML Estimation of Correlation Parameters: Correlation parameters θ are estimated by minimizing a negative restricted log-likelihood (REML) objective. The criterion implemented in `lgspline` is a central-limit-theorem-based working approximation to a Laplace-style marginal likelihood criterion, applied here solely to correlation structure estimation rather than penalty parameter selection. Let $\mathbf{D} = \text{diag}(d_i)$ be the observation weight matrix, $\tilde{\mathbf{W}} = \text{diag}(\tilde{w}_i)$ the GLM working weight matrix at the current fitted values, \mathbf{V} the correlation matrix parameterized by ρ (a vector on the unconstrained real line), and $\tilde{\sigma}^2$ the dispersion profiled at its restricted maximum likelihood estimate. The negative REML objective implemented in `lgspline`, scaled by $1/N$, is

$$-\ell_R(\rho) = \frac{1}{N} \left[-\log |\mathbf{V}^{-1/2}| + \frac{N}{2} \log \tilde{\sigma}^2 + \frac{1}{2\tilde{\sigma}^2} \|\mathbf{V}^{-1/2} \mathbf{D}^{1/2} \tilde{\mathbf{W}}^{-1/2} (\mathbf{y} - \boldsymbol{\mu})\|^2 + \frac{1}{2} \log |(\tilde{\sigma}^2 \mathbf{U} \mathbf{G} \mathbf{U}^\top)^{-1}|^+ \right],$$

where $|\cdot|^+$ denotes the generalized determinant (product of nonzero eigenvalues), and $\boldsymbol{\mu} = g^{-1}(\mathbf{X}\boldsymbol{\beta})$ are the fitted values on the response scale.

Gradients with respect to correlation parameters are available in closed form for all built-in structures except Matern, which uses finite-difference approximation due to the complexity of differentiating the modified Bessel function K_ν with respect to ν . See `reml_grad_from_dV` for the full gradient derivation and notation. Custom analytic gradients can be supplied through `REML_grad`, and a fully custom criterion can replace REML through `custom_vhalfInv_loss`. The Toeplitz example in `lgspline` demonstrates how to supply custom correlation structures with user-defined gradient functions. Optimization over these working correlation parameters is then carried out by the same quasi-Newton engine used elsewhere in the package, with a finite-difference fallback when needed. In the user-facing interface, this machinery is activated through `correlation_structure`, `correlation_id`, `spacetime`, `VhalfInv`, `Vhalf`, `VhalfInv_fxn`, `Vhalf_fxn`, `VhalfInv_par_init`, `REML_grad`, `custom_vhalfInv_loss`, and `VhalfInv_logdet`.

The gradient of the negative REML has three terms per parameter:

1. $\frac{1}{2} \text{tr}(\mathbf{V}^{-1} \partial \mathbf{V} / \partial \theta_j)$: the log-determinant contribution.
2. $-\frac{1}{2\tilde{\sigma}^2} \mathbf{r}^\top (\partial \mathbf{V} / \partial \theta_j) \mathbf{r}$: the residual quadratic form contribution, where $\mathbf{r} = \text{diag}(\sqrt{d_i}/\sqrt{\tilde{w}_i}) \mathbf{V}^{-1/2} (\mathbf{y} - \boldsymbol{\mu})$.
3. $-\frac{1}{2} \text{tr}(\mathbf{M}^+ \mathbf{X}_*^\top \mathbf{V}^{-1} (\partial \mathbf{V} / \partial \theta_j) \mathbf{V}^{-1} \tilde{\mathbf{W}} \mathbf{D} \mathbf{X}_*)$: the REML correction, where $\mathbf{X}_* = \mathbf{X} \mathbf{U}$ is the constrained design and $\mathbf{M} = \mathbf{X}_*^\top \mathbf{V}^{-1} \tilde{\mathbf{W}} \mathbf{D} \mathbf{X}_* + \mathbf{U}^\top \boldsymbol{\Lambda} \mathbf{U}$ is the projected penalized information.

For each supported correlation family, the derivatives $\partial \mathbf{V} / \partial \theta_j$ are available in closed form, enabling analytic gradient computation for use with the quasi-Newton optimizer.

Connection to Standard Mixed Model REML: The quadratic penalty $\boldsymbol{\Lambda}$ acts as the inverse prior covariance of a Gaussian random effect on the spline coefficients, with the smoothing parameter satisfying $\lambda = \tau^2 / \sigma^2$; this is the same mixed model representation used by `mgcv`, and Monte

Carlo draws under the resulting Laplace-style approximation are available via `generate_posterior`. For Gaussian responses with identity link and no penalization, the implemented criterion coincides exactly with classical REML. For non-Gaussian responses, the criterion substitutes the Fisher information for the full penalized log-likelihood Hessian, exploiting the CLT approximation that $\tilde{\mathbf{W}}^{-1/2}(\mathbf{y} - \boldsymbol{\mu})$ is approximately Gaussian; this yields a method-of-moments style estimator for the correlation and variance parameters that depends only on mean-variance relationships through \mathbf{W} , and therefore generalizes naturally to quasi-likelihood and other settings where a fully specified log-likelihood is unavailable. The REML correction term $\log |(\tilde{\sigma}^2 \mathbf{U} \mathbf{G} \mathbf{U}^\top)^{-1}|$ uses a generalized log-determinant so that only nonzero eigenvalues contribute when rank deficiency arises from smoothness constraints or identifiability conditions in \mathbf{A} . When \mathbf{A} is full rank, the criterion coincides with the exact marginal likelihood from integrating out $\boldsymbol{\beta}$ under its Gaussian prior; when \mathbf{A} is rank-deficient, unpenalized coefficients are projected out in the REML sense while penalized coefficients are integrated through their prior. During penalty tuning, the block-diagonal approximation is retained for GCV criteria and gradients; since GCV is rotation-invariant the practical effect on automatic selection is expected to be negligible, though this is not confirmed and the tuned penalties can always be overridden.

Built-In Correlation Structures: The package provides several built-in correlation structures for modeling spatial and temporal dependence. These are specified via `correlation_structure` with group membership in `correlation_id` and spatial or temporal coordinates in `spacetime` (an N -row matrix). Exchangeable correlation does not require `spacetime`.

All positive scale parameters are estimated on the log scale, with back-transform $\exp(\cdot)$. Parameters constrained to $(0, 1)$ use a double-exponential back-transform of the form $\exp(-\exp(\eta))$, so optimization still occurs on the unconstrained real line while the correlation remains bounded.

Exchangeable Aliases: 'exchangeable', 'cs', 'CS', 'compoundsymmetric', 'compound-symmetric'.

A constant correlation ν between any two observations within the same cluster. Parameterization: $\nu = \exp(-\exp(\rho))$, so $\nu \in (0, 1)$. Only positive within-cluster correlation is supported under this parameterization.

Spatial Exponential Aliases: 'spatial-exponential', 'spatialexponential', 'exp', 'exponential'.

Correlation decays exponentially with distance: $\exp(-\omega d)$ where d is Euclidean distance and $\omega > 0$. Parameterization: $\omega = \exp(\rho)$. Mathematically equivalent to the power correlation θ^d with $\theta = e^{-\omega}$, but with better numerical properties during optimization.

AR(1) Aliases: 'ar1', 'ar(1)', 'AR(1)', 'AR1'.

Correlation depends on rank difference between observations: ν^r where r is the rank difference within cluster. Parameterization: $\nu = \exp(-\exp(\rho))$, so $\nu \in (0, 1)$. Only positive autocorrelation is supported.

Gaussian / Squared Exponential Aliases: 'gaussian', 'rbf', 'squared-exponential'.

Smooth decay with squared distance: $\exp(-d^2/(2\ell^2))$ where ℓ is the length scale. Parameterization: $\ell = \exp(\rho)$.

Spherical Aliases: 'spherical', 'Spherical', 'cubic', 'sphere'.

Polynomial decay with a hard cutoff at range r : $1 - 1.5(d/r) + 0.5(d/r)^3$ for $d \leq r$, and 0 otherwise. Parameterization: $r = \exp(\rho)$.

Matern Aliases: 'matern', 'Matern'.

Flexible correlation with adjustable smoothness: $(2^{1-\nu}/\Gamma(\nu))(\sqrt{2\nu} d/\ell)^\nu K_\nu(\sqrt{2\nu} d/\ell)$. Two parameters: length scale $\ell = \exp(\rho_1)$ and smoothness $\nu = \exp(\rho_2)$. No analytical gradient is available for ν due to the difficulty of differentiating the modified Bessel function K_ν with respect to its order, so finite differences are used; this makes Matern slower and potentially less stable than other structures.

Gamma-Cosine Aliases: 'gamma-cosine', 'gammacosine', 'GammaCosine'.

Oscillatory dependence: $(d^{\alpha-1}e^{-\gamma d})/(\Gamma(\alpha)/\gamma^\alpha) \cdot \cos(\omega d)$. Three parameters: shape $\alpha = \exp(\rho_1)$, rate $\gamma = \exp(\rho_2)$, frequency $\omega = \exp(\rho_3)$. Reduces to exponential when $\alpha = 1$ and $\omega \approx 0$.

Gaussian-Cosine Aliases: 'gaussian-cosine', 'gaussiancosine', 'GaussianCosine'.

Smooth oscillatory correlation: $\exp(-d^2/(2\ell^2)) \cdot \cos(\omega d)$. Two parameters: length scale $\ell = \exp(\rho_1)$ and frequency $\omega = \exp(\rho_2)$. Reduces to Gaussian when $\omega \approx 0$.

Interpreting Estimated Correlation Parameters: Correlation parameters are estimated on transformed scales; they must be back-transformed for interpretation. When `confint.lgspline` is called and the inverse Hessian from BFGS is available, confidence intervals are returned on the untransformed (working) scale and should be back-transformed as described in the examples for `lgspline`.

Custom Correlation Structures: Custom correlation structures can be specified through:

- `VhalfInv_fxn`: Creates $\mathbf{V}^{-1/2}$.
- `Vhalf_fxn`: Creates $\mathbf{V}^{1/2}$. When omitted, the code computes it by explicit inversion of `VhalfInv`.
- `REML_grad`: Provides the analytical gradient of the REML objective.
- `VhalfInv_logdet`: Efficient log-determinant computation.
- `custom_VhalfInv_loss`: Replaces the REML objective entirely.

These functions enter `lgspline` through `correlation_structure`, `VhalfInv_fxn`, `Vhalf_fxn`, `REML_grad`, and `custom_VhalfInv_loss`, and the fitted object retains the resulting correlation machinery in components such as `VhalfInv_fxn`, `Vhalf_fxn`, and `VhalfInv_params_estimates`. When `VhalfInv` is supplied but `Vhalf` is not, `Vhalf` is computed unconditionally as the inverse of `VhalfInv` for all family/link combinations, since both `get_B` and `blockfit_solve` require it for GEE estimation.

Variance and Dispersion Estimation

Once the constrained estimate has been obtained, the next questions are how much flexibility the fitted model effectively used and how uncertainty should be propagated through the same constrained geometry. The quantities in this section are therefore all built on the projected information matrices from the previous sections.

Effective Degrees of Freedom and Dispersion: The effective degrees of freedom is the trace of the hat matrix. In the Gaussian identity case with observation weights and correlation, the fitted linear operator is built from the dense GLS analogue $\mathbf{G}_{\text{correct}}$ and can be written schematically as

$$\mathbf{H} = \mathbf{V}^{-1/2}(\mathbf{WD})^{1/2}\mathbf{XUG}_{\text{correct}}\mathbf{X}^\top(\mathbf{WD})^{1/2}\mathbf{V}^{-1/2},$$

where for Gaussian identity $\mathbf{W} = \mathbf{I}$. In the no-correlation Gaussian case this reduces to the familiar $\mathbf{H} = \mathbf{XUGX}^\top\mathbf{D}$.

For Gaussian identity fits, the dispersion estimate is computed as a weighted mean squared residual, optionally scaled by $N/(N - \text{tr}(\mathbf{H}))$ when `unbias_dispersion = TRUE`:

$$\tilde{\sigma}^2 = \frac{1}{N - \text{tr}(\mathbf{H})} \|\mathbf{y} - \tilde{\mathbf{y}}_i\|^2.$$

More generally with weights, a correlation structure and non-linear link function:

$$\tilde{\sigma}^2 = \frac{1}{N - \text{tr}(\mathbf{H})} \|\mathbf{V}^{-1/2} \mathbf{W}^{-1/2} \mathbf{D}^{1/2} (\mathbf{y} - \tilde{\mathbf{y}})\|^2.$$

This estimated dispersion is returned as `sigmasq_tilde`, and the corresponding effective degrees of freedom trace is returned as `trace_XUGX`. For non-Gaussian families, the fitting code delegates dispersion estimation to `dispersion_function`; thus the package does not assume a single closed-form Pearson-style formula outside the Gaussian identity setting. The hat-matrix trace itself is assembled by `compute_trace_H` in the dense correlation-aware case and by the same blockwise products summarized by `trace_XUGX` in the simpler no-correlation paths. A concrete built-in non-Gaussian example is the Weibull AFT path, which pairs `weibull_family` with `weibull_dispersion_function`, `weibull_glm_weight_function`, and `weibull_schur_correction`. Users who want these quantities available for downstream inference should keep `estimate_dispersion = TRUE` and `return_varcovmat = TRUE` (the defaults), since `wald_univariate`, `confint.lgspline`, and the prediction-standard-error path in `predict.lgspline` all rely on the post-fit dispersion and covariance components documented here.

Variance-Covariance Matrix: The variance-covariance matrix of $\tilde{\beta}$ is estimated as:

$$\text{Var}(\tilde{\beta}) = \tilde{\sigma}^2 (\mathbf{U} \mathbf{G}^{1/2}) (\mathbf{U} \mathbf{G}^{1/2})^\top$$

using the outer-product form for numerical stability. The result is returned as `varcovmat` when `return_varcovmat = TRUE`. The algebraically equivalent expression $\tilde{\sigma}^2 \mathbf{U} \mathbf{G}$ is not used because \mathbf{G} is only positive semi-definite when the penalty matrix $\mathbf{\Lambda}$ has zero eigenvalues (e.g., the intercept and linear terms under the smoothing spline penalty when `flat_ridge_penalty = 0`), which can introduce negative diagonal entries in finite precision arithmetic. The outer-product form also guarantees symmetry.

This is the Bayesian posterior covariance, treating the penalty as a Gaussian prior on the coefficients. When `exact_varcovmat = TRUE`, a frequentist correction is additionally computed:

$$\text{Var}_{\text{exact}}(\tilde{\beta}) = \tilde{\sigma}^2 \mathbf{U} \mathbf{G}^{1/2} (\mathbf{X}^\top \mathbf{W} \mathbf{D} \mathbf{V}^{-1} \mathbf{X}) \mathbf{G}^{1/2} \mathbf{U}^\top = \tilde{\sigma}^2 \mathbf{U} \mathbf{G} \mathbf{U}^\top - \tilde{\sigma}^2 \mathbf{U} \mathbf{G} \mathbf{\Lambda} \mathbf{G} \mathbf{U}^\top.$$

The first term is the Bayesian posterior covariance; the second is a frequentist correction such that for Gaussian identity link (with or without correlation), this is the exact variance-covariance matrix of the constrained estimator.

When a correlation structure is present (`VhalfInv` non-NULL), the block-diagonal \mathbf{G} is replaced by the full weighted GLS analogue

$$\mathbf{G}_{\text{correct}} = (\mathbf{X}^\top \mathbf{W} \mathbf{D} \mathbf{V}^{-1} \mathbf{X} + \mathbf{\Lambda})^{-1},$$

where $\mathbf{W} = \mathbf{I}$ in the Gaussian identity case. This dense matrix is what enters the correlation-aware \mathbf{U} , $\text{Var}(\tilde{\beta})$, and $\text{Var}_{\text{exact}}(\tilde{\beta})$ computations.

In user-facing terms, `return_varcovmat` controls whether this matrix is stored at all, while `exact_varcovmat` switches between the default posterior/Laplace approximation and the exact frequentist correction in the Gaussian-identity setting. The stored covariance is what powers `wald_univariate`, `confint.lgspline`, and `se.fit = TRUE` in `predict.lgspline`; the `critical_value` argument supplied at fit time is carried forward as the default cutoff for those interval-producing helpers.

Recomputation of \mathbf{G} at Convergence: At the final iterate, \mathbf{G} is recomputed to reflect the converged working weights and Schur corrections. The implementation computes the weighted design $\mathbf{X}_w^{(k)} = \mathbf{X}_k \cdot \text{diag}(\sqrt{\mathbf{w}_k})$, forms the weighted Gram matrix $\mathbf{X}_w^{(k)\top} \mathbf{X}_w^{(k)}$, adds the Schur correction, and performs eigendecomposition via `compute_G_eigen` to obtain \mathbf{G}_k , $\mathbf{G}_k^{1/2}$, and $\mathbf{G}_k^{-1/2}$. The relationship $\mathbf{G}_k = \mathbf{G}_k^{1/2} (\mathbf{G}_k^{1/2})^\top$ is enforced exactly by construction, and the fitted object can retain these as `G` and `Ghalf` when `return_G = TRUE` and `return_Ghalf = TRUE`. The square-root factors are numerically stabilized through the helper routines `matsqrt` and `matinvsqrt`, which are also used elsewhere in the package when dense analogues of $\mathbf{G}^{1/2}$ or $\mathbf{G}^{-1/2}$ are required.

Bayesian Interpretation

The penalty has a natural Gaussian-prior interpretation, so once the constrained estimator and its covariance are available, Bayesian-style posterior simulation follows almost immediately. This section records the interpretation that is already implicit in the fitted object and in the package's posterior simulation helpers.

A Bayesian interpretation follows from viewing the penalty as a Gaussian prior on the coefficients. Conditional on the fitted smoothing parameters, the code samples on the coefficient scale from

$$\boldsymbol{\beta}^{(m)} = \tilde{\boldsymbol{\beta}} + \sqrt{\hat{\sigma}^2} \mathbf{U} \mathbf{G}^{1/2} \mathbf{z}^{(m)}, \quad \mathbf{z}^{(m)} \sim \mathcal{N}(\mathbf{0}, \mathbf{I}).$$

The coefficients are then back-transformed to the original response and predictor scales.

When inequality constraints are absent, these draws are i.i.d. Gaussian posterior draws around the fitted mode. The underlying coefficient-draw closure also contains an elliptical slice sampling route for active inequality constraints, using the same covariance factor to keep retained draws in the feasible region.

At the implementation level, standard Gaussian posterior draws may place positive mass on the infeasible region, so the constrained-draw closure instead targets the truncated posterior

$$\pi(\boldsymbol{\beta} \mid \mathbf{y}) \propto \exp\left(-\frac{1}{2}(\boldsymbol{\beta} - \tilde{\boldsymbol{\beta}})^\top \mathbf{G}^{-1}(\boldsymbol{\beta} - \tilde{\boldsymbol{\beta}})\right) \mathbf{1}(\mathbf{C}^\top \boldsymbol{\beta} \succeq \mathbf{c}),$$

yielding credible intervals that respect the constraint boundaries. The public `generate_posterior` wrapper forwards `enforce_qp_constraints` to the stored constrained-draw closure, so constrained draws can be requested directly from the user-facing interface. When a working correlation structure is present, the companion helper `generate_posterior_correlation` extends this idea by propagating uncertainty in the fitted correlation parameters through the same `VhalfInv_fn/Vhalf_fn` machinery described in the correlation section, rather than conditioning only on fixed covariance parameters. Correlation parameters are drawn from a multivariate normal distribution centered about their estimates with the inverse approxiamte BFGS Hessian of the REML optimization problem `VhalfInv_params_vcov` used by default (or a custom alternative as supplied to the argument `correlation_param_vcov_sc`).

Inequality Constraints via Sequential Quadratic Programming

Overview: Inequality constraints of the form $\mathbf{C}^\top \boldsymbol{\beta} \succeq \mathbf{c}$ handle shape restrictions such as monotonicity, convexity, or boundedness. In the monomial basis, these are linear in $\boldsymbol{\beta}$. Monotonicity

at a grid of points requires the first derivative polynomial to be non-negative there; convexity requires the second derivative to be non-negative; range constraints bound the fitted values directly. All of these translate to linear inequality constraints on the coefficient vector.

The inequality pieces are assembled by `process_qp`, which returns the `qp_Amat`, `qp_bvec`, and `qp_meq` objects passed to `solve.QP`, together with a `quadprog` flag indicating whether any inequality constraints are active. For derivative-sign constraints, `process_qp` calls `.build_deriv_qp`, which uses `make_derivative_matrix` on the expansion-standardized design and maps the derivative rows into the full P -dimensional coefficient space partition by partition.

Partition-Wise Active-Set Method: The sparsity pattern of \mathbf{C} is inspected automatically by `.detect_qp_global` (equivalently `.solver_detect_qp_global`). When every column of \mathbf{C} has nonzero entries in only a single partition block, the constraint system is block-separable and an active-set method can replace the dense QP.

At each iteration, the active set \mathcal{A} (constraints satisfied at equality) is appended to \mathbf{A} as additional equality constraints:

$$\mathbf{A}_{\text{aug}} = [\mathbf{A} \mid \mathbf{C}_{\mathcal{A}}].$$

The constrained estimate is obtained by the same OLS projection with \mathbf{A}_{aug} in place of \mathbf{A} , and since \mathbf{A}_{aug} retains block-diagonal compatibility, all operations remain partition-wise.

The active set is updated by checking primal feasibility (adding the most-violated inactive constraint) and dual feasibility (dropping the active constraint with the most negative Lagrange multiplier) until the KKT conditions are satisfied. Lagrange multipliers for active inequalities are recovered from the OLS fit used in the Lagrangian projection: the fitted coefficients on $\mathbf{X}^* = \mathbf{G}^{1/2}\mathbf{A}_{\text{aug}}$ give the multipliers up to sign. Implemented in `.active_set_refine` and `.check_kkt_partitionwise`.

During penalty tuning, the active inequality set from the last accepted tuning evaluation is used as the next initial working set. This only changes where the add/drop loop starts; the KKT check still determines the final active set for each penalty value.

The method adds or drops one constraint at a time. It is usually fast when the final active set is small and well separated, but can slow down when many derivative constraints are nearly binding, redundant, or nearly collinear. The implementation tracks visited active sets, filters rank deficient augmented systems, and falls back to dense SQP if the active-set route does not converge within its iteration budget.

When any column of \mathbf{C} spans multiple partition blocks (for example, cross-knot monotonicity constraints), the equality re-solve switches to the full-system bridge rather than the partition-wise bridge. The selection is automatic; dense SQP is retained as the fallback.

Dense SQP Iteration: The dense SQP approach, implemented in `.qp_refine`, solves a sequence of quadratic subproblems approximating the penalized log-likelihood. At each iteration s :

1. Compute the information matrix $\mathbf{M}^{(s)} = \mathbf{X}^\top \mathbf{W}^{(s)} \mathbf{X} + \mathbf{\Lambda}_{\text{block}} + \mathbf{S}^{(s)}$, where $\mathbf{S}^{(s)}$ is the Schur complement correction.
2. Compute the score vector via `qp_score_function`.
3. Solve the QP with `solve.QP`, where the combined constraint matrix is $[\mathbf{A} \mid \mathbf{C}]$ with the first R columns treated as equalities.
4. Apply a damped update $\boldsymbol{\beta}^{(s+1)} = (1 - \alpha)\boldsymbol{\beta}^{(s)} + \alpha\boldsymbol{\beta}_{\text{QP}}$, with $\alpha = 2^{-d}$ and d incremented upon deviance increase.

A rescaling factor $\text{sc} = \sqrt{\text{mean}(|\mathbf{M}^{(s)}|)}$ is applied to the Hessian and linear term before calling the QP solver. The equality-constrained estimate from the Lagrangian projection serves as a

warm start. The `qp_score_function` defaults to the GLM score using `family$mu.eta(eta) / family$variance(mu)`; for custom models a different score can be supplied. With `denseVhalfInv`, the same score weight is applied after whitening.

Active Set and Lagrange Multipliers: The active set at the solution identifies binding inequality constraints. The Lagrange multipliers quantify the cost of each: a multiplier of zero means the constraint is not binding. The implementation stores active constraint indices, the corresponding submatrix of the constraint matrix, and the multiplier vector in the `qp_info` list returned alongside coefficient estimates, with components `lagrangian`, `iact`, and `Amat_active`. When `return_lagrange_multipliers = TRUE`, the fitted object also stores the final multiplier vector directly as `lagrange_multipliers`.

The `qp_info$method` string records which solver path produced the final inequality-constrained estimate: `"active_set"` = partition-wise active-set on the standard block-diagonal path; `"active_set_full"` = active-set with full-system equality re-solves; `"active_set_woodbury"` = active-set with Woodbury equality re-solves on the correlated low-rank path; `"dense_qp_gee_gaussian"` = dense QP fallback for correlated Gaussian fits; `"dense_qp_gee_glm"` = dense SQP / dense QP-subproblem fallback for correlated non-Gaussian fits.

The original assembled inequality data are retained in the fitted object's `quadprog_list` component (containing `qp_Amat`, `qp_bvec`, and `qp_meq` from `process_qp`) so the final active set can be interpreted relative to the full specification. The final \mathbf{U} used in constructing the posterior variance-covariance matrix is built from both the equality constraints and the active inequality constraints at the solution.

Built-In Constraints: Built-in inequality constraints include:

- **Monotonicity:** `qp_monotonic_increase`, `qp_monotonic_decrease`. Enforced by requiring consecutive fitted values to be non-decreasing (or non-increasing): $(\mathbf{x}_i - \mathbf{x}_{i-1})^\top \boldsymbol{\beta} \geq 0$. These are constructed by `process_qp` from the partition-stacked block design reordered to observation order.
- **Derivative sign:** `qp_positive_derivative`, `qp_negative_derivative`. Enforced through the first-derivative design matrix from `make_derivative_matrix`. May be TRUE/FALSE or a character/integer vector selecting specific predictors.
- **Second-derivative sign:** `qp_positive_2ndderivative`, `qp_negative_2ndderivative`. Same construction using the second-derivative design matrix.
- **Response range:** `qp_range_lower`, `qp_range_upper`. Bounds on the linear predictor; for non-identity links, the bounds are transformed to the link scale inside `process_qp`.
- **Custom constraints** via `qp_Amat_fxn`, `qp_bvec_fxn`, and `qp_meq_fxn`, which receive the design matrix structure and return the constraint matrix, bound vector, and number of equalities. These are commonly paired with a custom `qp_score_function` when the quadratic approximation uses a non-default likelihood. The low-level objects `qp_Amat`, `qp_bvec`, and `qp_meq` remain documented in `lgspline` but in the current implementation serve as activation markers rather than being merged into the constraint set assembled by `process_qp`.

Built-in constraints can be thinned via `qp_observations`. A single numeric vector applies the same subset to every active built-in constraint, while a named list keyed by `"var:qp_<type>"` or bare `"qp_<type>"` lets different constraint types use different row subsets. For range and monotonicity, matching keyed entries are unioned; derivative entries dispatch per variable. Unknown keys are ignored with a warning when `include_warnings = TRUE`.

After assembly, `process_qp` can also stabilize the partition-local inequality blocks by QR pivoting them before the final QP solve. When `qr_pivot_inequality_constraints = TRUE`, each

partition-local block of \mathbf{C} is scanned for linearly dependent columns and only the pivot columns are carried forward, while the leading equality columns are left untouched. If `parallel_qr_qp = TRUE`, those partition-wise QR pivots are farmed out across workers in the same style as the package’s other parallel QR reductions. Built-in range and monotonicity constraints are intentionally excluded from this reduction. For range bounds, keeping the full block preserves the direct user-facing lower/upper-value meaning; for monotonicity, the consecutive differences are defined in observation order and can bridge neighboring partitions even when some individual difference columns happen to sit inside one partition block.

Blockfit Backfitting for Linear Non-Interactive Effects

Motivation: When a model contains both spline terms (receiving $K + 1$ partition-specific coefficient vectors constrained to smoothness) and non-interactive linear terms (“flat” terms, specified via `just_linear_without_interactions`, receiving a single shared coefficient vector \mathbf{v} across all partitions), the standard solver carries $K + 1$ copies of \mathbf{v} linked by equality constraints. Backfitting avoids this inflation by solving a lower-dimensional problem at each step. Write the partition- k design as $\mathbf{X}_k = [\mathbf{Z}_k \mid \mathbf{X}_{\text{flat}}^{(k)}]$, where \mathbf{Z}_k contains the spline columns and $\mathbf{X}_{\text{flat}}^{(k)}$ the flat columns. This is invoked when `blockfit = TRUE`, flat columns are non-empty, and $K > 0$.

The design, penalty, and constraint matrices are split into spline and flat components by `.bf_split_components`, which extracts spline rows from \mathbf{A} , drops null columns, rank-reduces via QR, and detects mixed constraints (columns of \mathbf{A} with nonzero entries on both spline and flat rows).

Block-Coordinate Descent: Spline step. Holding \mathbf{v} fixed, the code forms the adjusted response $\mathbf{y}_k - \mathbf{X}_{\text{flat}}^{(k)}\mathbf{v}$ and applies the spline-only Lagrangian projection via `.bf_lagrangian_project`:

$$\tilde{\boldsymbol{\beta}}_{\text{spline}}^{(k)} = \mathbf{U}_{\text{spline}} \mathbf{G}_{\text{spline}} \mathbf{Z}_k^\top (\mathbf{y}_k - \mathbf{X}_{\text{flat}}^{(k)}\mathbf{v}).$$

Flat step. Holding spline coefficients fixed, the shared flat vector is updated by pooled penalized regression on residuals:

$$\mathbf{v} = \left(\sum_{k=0}^K \mathbf{X}_{\text{flat}}^{(k)\top} \mathbf{X}_{\text{flat}}^{(k)} + \boldsymbol{\Lambda}_{\text{flat}} \right)^{-1} \sum_{k=0}^K \mathbf{X}_{\text{flat}}^{(k)\top} (\mathbf{y}_k - \mathbf{Z}_k \tilde{\boldsymbol{\beta}}_{\text{spline}}^{(k)}).$$

When the constraint matrix \mathbf{A} has mixed columns (nonzero entries on both spline and flat rows), the flat update instead solves a KKT system enforcing the residual equality constraint $\mathbf{A}_{\text{flat}}^\top \mathbf{v} = \mathbf{c} - \mathbf{A}_{\text{spline}}^\top \tilde{\boldsymbol{\beta}}_{\text{spline}}$ via `.bf_constrained_flat_update`.

Convergence is checked using the maximum absolute change across both spline and flat coefficients. In the weighted inner loop used by the GLM solvers, the flat-block change alone determines stopping.

Four Estimation Cases: `blockfit_solve` dispatches to one of four paths.

Case (a): Gaussian identity + GEE (`.bf_case_gauss_gee`). Whitening destroys block-diagonal structure, so the code skips backfitting and performs the same full-system Gaussian GEE projection used by `get_B` Path 1a. The result is split back into spline and flat components for downstream assembly.

Case (b): Gaussian identity, no correlation (`.bf_case_gauss_no_corr`). Standard block-coordinate descent as described above. The spline-only $\mathbf{G}_{\text{spline}}$ factors are precomputed once, and the pooled flat penalized inverse is reused across iterations.

Case (c): GLM + GEE (`.bf_case_glm_gee`). Two stages. Stage 1 forms a warm start by running damped Newton-Raphson on the unwhitened working response: each outer iteration computes working responses and weights at the current linear predictor, then calls `.bf_inner_weighted` for the inner backfitting loop. Stage 2 refines the warm start on the full whitened system using the damped SQP loop (`.bf_sqp_loop`), replicating the approach used by `.get_B_gee_glm`.

Case (d): GLM without GEE (`.bf_case_glm_no_corr`). A damped Newton-Raphson outer loop updates working responses and weights, while each inner iteration alternates between weighted spline and weighted flat updates via `.bf_inner_weighted`. Deviance is monitored across outer iterations for convergence and damping.

Inequality Constraints and Reassembly: Because flat coefficients are shared by construction, the corresponding equality constraints are satisfied exactly. Smoothness constraints on the spline block are enforced by the spline-only Lagrangian projection. After backfitting convergence, inequality constraints use the same active-set-first refinement used by `get_B`. Block-separable constraints use partition-wise equality re-solves through `.active_set_refine`; cross-partition constraints use the full-system bridge; and GEE paths use the whitened system, with Woodbury equality re-solves when the low-rank gate succeeds. Dense SQP through `.bf_sqp_loop` remains the fallback when active-set refinement does not converge.

After convergence, the shared flat vector \mathbf{v} is copied into each partition's coefficient vector, yielding $\beta_k = [\tilde{\beta}_{\text{spline}}^{(k)\top}, \mathbf{v}^\top]^\top$ for compatibility with downstream inference. If `blockfit_solve` throws an error, a warning is issued and the code falls back to `get_B`.

Knot Selection and Partitioning

The topic of knot selection is not the main focus of the package, but the partition structure is central because every later design matrix, penalty, and smoothness constraint depends on it. The defaults in `lgspline` are therefore meant to be practical and transparent rather than theoretically final.

Univariate Case: For a single predictor, the default partitioning is now handled by `make_partitions` in the same k -means framework used more generally: $K + 1$ centers are fit on an internally standardized copy of the predictor, controlled by `standardize_predictors_for_knots`, and then returned on the raw scale. Custom knots can still be supplied via `custom_knots`, in which case partition assignment is built directly from those raw-scale breakpoints. The default number of knots K is chosen adaptively based on N , p , q , and the GLM family. For multivariate fits, the resulting partition metadata are returned as `make_partition_list` and can be re-used in later calls to `lgspline`. This is particularly useful when one wants to hold the partition geometry fixed across repeated fits, for example while varying penalties, families, or correlation structures.

Multivariate Case: For multiple predictors, $K + 1$ cluster centers are identified by k -means on an internally standardized predictor matrix via `make_partitions`. This is the partitioning mechanism used to determine the multivariate spline regions; see MacQueen (1967) for the classical clustering formulation and Kisi et al. (2025) for a recent applied example of k -means-driven partitioning in a nonlinear prediction setting. Midpoints between neighboring centers (those whose midpoint does not fall into a third cluster) serve as knot locations. Observations are assigned to the nearest cluster center using `get.knnx`, and the returned centers and knots are on the original predictor scale. The resulting partition structure is a type of Voronoi diagram and is stored in the fitted object as `make_partition_list`. The `do_not_cluster_on_these` argument can exclude certain predictors from clustering (e.g., a treatment indicator that should not drive partitioning). The lower-level clustering behavior can be further controlled by `cluster_args` and

neighbor_tolerance, while cluster_on_indicators determines whether binary predictors are allowed to influence the partition geometry at all.

Standardizing Predictors: Higher-order polynomial terms can dramatically inflate or deflate the magnitude of basis expansions, introducing numerical instability. All polynomial basis expansions are scaled by $q_{0.69} - q_{0.31}$, where q_ζ is the ζ -th quantile of the expansion. For a standard normal distribution this quantity is approximately 1, so the scaling is close to one standard deviation for symmetric distributions. This fitting-stage rescaling is controlled by `standardize_expansions_for_fitting`, while knot construction is controlled separately by `standardize_predictors_for_knots`. The same scaling is applied to the constraint matrix to maintain smoothness, and coefficients are back-transformed to the original scale after fitting.

Smoothing Spline Penalty

Penalty Construction: The penalty matrix Λ_s penalizes the integrated squared total curvature of the fitted function over the observed predictor ranges. This is the step that makes the piecewise polynomial fit genuinely behave like a smoothing spline rather than merely a constrained regression spline. The package computes this penalty directly from the monomial structure of the basis rather than by appealing to a pre-tabulated spline basis. For a single partition k with basis expansion $\mathbf{x}_k = (\phi_1(\mathbf{t}), \dots, \phi_p(\mathbf{t}))^\top$ where each $\phi_i(\mathbf{t}) = \prod_{j=1}^q t_j^{\alpha_{ij}}$ is a multivariate monomial:

$$\beta_k^\top \Lambda_s \beta_k = \int_{\mathbf{a}}^{\mathbf{b}} \|\tilde{f}_k''(\mathbf{t})\|^2 dt,$$

where \mathbf{a} and \mathbf{b} are the observed predictor minimums and maximums (computed globally from the data, not partition-specific), and $\tilde{f}_k(\mathbf{t}) = \mathbf{x}_k^\top \beta_k$ is the fitted function for partition \mathcal{P}_k .

Total curvature operator. The integrated squared second derivative decomposes into q curvature operators, one per predictor. For predictor v , the curvature operator D_v is defined as

$$D_v = \frac{\partial^2}{\partial t_v^2} + \sum_{s \neq v} \frac{\partial^2}{\partial t_v \partial t_s}.$$

That is, D_v captures both the pure second derivative with respect to t_v and all mixed second partial derivatives involving t_v . The penalty matrix entries are then

$$[\Lambda_s]_{ij} = \sum_{v=1}^q \int_{\mathbf{a}}^{\mathbf{b}} D_v(\phi_i) D_v(\phi_j) dt.$$

Monomial derivative rule. For a monomial $\phi(\mathbf{t}) = \prod_j t_j^{\alpha_j}$, the derivatives entering D_v have closed forms. The pure second derivative is

$$\frac{\partial^2}{\partial t_v^2} \prod_j t_j^{\alpha_j} = \alpha_v(\alpha_v - 1) t_v^{\alpha_v - 2} \prod_{j \neq v} t_j^{\alpha_j},$$

which is zero when $\alpha_v < 2$. The mixed second derivative is

$$\frac{\partial^2}{\partial t_v \partial t_s} \prod_j t_j^{\alpha_j} = \alpha_v \alpha_s t_v^{\alpha_v - 1} t_s^{\alpha_s - 1} \prod_{j \neq v, s} t_j^{\alpha_j},$$

which is zero when $\alpha_v < 1$ or $\alpha_s < 1$. Applying D_v to a monomial ϕ_i produces a sum of monomials with known coefficients and exponent vectors.

Factorized integration. Because every $D_v(\phi_i)$ is polynomial, the product $D_v(\phi_i) D_v(\phi_j)$ is also polynomial and the multivariate integral factorizes over predictors:

$$\int_{\mathbf{a}}^{\mathbf{b}} \prod_{j=1}^q t_j^{e_j} dt = \prod_{j=1}^q \frac{b_j^{e_j+1} - a_j^{e_j+1}}{e_j + 1}.$$

The integral runs over all q predictor ranges, including predictors absent from the integrand, for which $e_j = 0$ and the factor is $b_j - a_j$.

Single-predictor verification. For $q = 1$ with expansion $\mathbf{x} = (1, t, t^2, t^3)^\top$ on $[a, b]$, the curvature operator reduces to $D_1 = \partial^2 / \partial t^2$ (no mixed partials exist), and the penalty matrix reduces to

$$\mathbf{\Lambda}_s = \int_a^b \mathbf{x}'' \mathbf{x}''^\top dt = \begin{pmatrix} 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 4(b-a) & 6(b^2-a^2) \\ 0 & 0 & 6(b^2-a^2) & 12(b^3-a^3) \end{pmatrix},$$

equivalent to the classical cubic smoothing spline penalty originally proposed by Reinsch.

Handling of non-spline predictors. Predictors specified via `just_linear_without_interactions` or `just_linear_with_interactions` do not receive higher-order polynomial expansions in the design matrix. To ensure their curvature contributions are still correctly computed (particularly through interaction terms), the implementation temporarily appends phantom higher-order columns (with zero data) for these predictors, computes the full curvature penalty on the augmented basis, and then subsets the result back to the original $p \times p$ dimensions. This ensures that interaction terms involving non-spline predictors receive appropriate penalty contributions without affecting the rest of the estimation pipeline.

Parallel computation. Because the total penalty is an additive sum over predictors ($\mathbf{\Lambda}_s = \sum_{v=1}^q \mathbf{\Lambda}_{s,v}$), the computation can be parallelized by distributing the per-predictor curvature matrices across workers via `parallel::parLapply` and summing the results. This is controlled by the `parallel_penalty` argument and is beneficial when q is large.

The penalty is computed by `get_2ndDerivPenalty` (single predictor or subset) and `get_2ndDerivPenalty_wrapper` (full assembly with optional parallelism and non-spline handling).

Because the smoothing penalty has zero eigenvalues for the intercept and linear terms (whose second derivatives vanish), an optional ridge penalty on lower-order terms is added for computational stability.

The full penalty block for partition k is:

$$\mathbf{\Lambda}_k = \lambda_w \left(\mathbf{\Lambda}_s + \lambda_r \mathbf{\Lambda}_r + \sum_{l=1}^L \lambda_{l,k} \mathbf{\Lambda}_{l,k} \right),$$

and the full penalty matrix is

$$\mathbf{\Lambda} = \text{blockdiag}(\mathbf{\Lambda}_0, \dots, \mathbf{\Lambda}_K).$$

Here $\mathbf{\Lambda}_s$ is the integrated curvature penalty, $\mathbf{\Lambda}_r$ is the ridge penalty on the null space, and $\mathbf{\Lambda}_{l,k}$ denotes any additional penalty matrix attached to partition k . The tuning scalars are λ_w (`wiggle_penalty`), λ_r (`flat_ridge_penalty`), and $\lambda_{l,k}$ (represented through `predictor_penalties`

and `partition_penalties` in the current interface). Internally, the package stores these components with implementation labels such as `L1`, `L2`, `L_predictor_list`, and `L_partition_list`. This assembly is handled by `compute_Lambda`.

The penalty matrix Λ is stored as a list of $K + 1$ $p \times p$ square, symmetric, positive semi-definite matrices.

Penalty Optimization via Leave-One-Out or Generalized Cross-Validation: After the structural pieces of the model are fixed, the main remaining question is how much smoothing to apply. In **lgspline**, that tuning is performed with exact leave-one-out by default, or with generalized cross-validation when `tuning_criterion = "gcv"`. In either case, the criterion is evaluated using the same constrained estimator that will be used in the final model fit. Penalty parameters are estimated on the log scale via exponential parameterization ($\lambda = \exp(\theta)$, $\theta \in \mathbb{R}$), ensuring positivity. The chain rule factor $\partial \exp(\theta) / \partial \theta = \exp(\theta) = \lambda$ is applied throughout. User-facing arguments (`initial_wiggle`, `initial_flat`, `predictor_penalties`, `partition_penalties`) accept values on the raw, natural scale; conversion to log scale is handled internally. The final tuned values and assembled penalty pieces are returned in the fitted object's `penalties` component.

With `tuning_criterion = "loo"`, the criterion is

$$\text{LOO} = \frac{1}{N} \sum_{i=1}^N \left(\frac{r_i}{1 - h_{ii}} \right)^2,$$

where h_{ii} is the diagonal of the tuning hat matrix. For Gaussian identity-link tuning with fixed transformed design, this is exact. With correlation structure and/or non-identity links, **lgspline** applies the same transformed or linearized tuning problem already used by GCV, so the leave-one-out calculation is carried out on that working scale. In empirical diagnostics, the criterion itself behaves stably, while the fully analytic derivative of the observation-wise leverage term can be numerically delicate in some problems; when that sensitivity matters in practice, the finite-difference optimizer path (`use_custom_bfgs = FALSE`) remains available. This exact observation-wise calculation is also the main reason LOO tuning becomes more expensive than GCV as N grows; in routine use, `tuning_criterion = "gcv"` is often the more practical choice once the sample size is above about 250,000, although this varies by data set.

With `tuning_criterion = "gcv"`, the package instead uses

$$\text{GCV}_{u,\gamma} = \frac{\sum_{i=1}^N D_{ii} r_i^2}{N(1 - \gamma \bar{W})^2}$$

where $\bar{W} = \text{tr}(\mathbf{H})/N$ is the mean of the hat-matrix diagonal. The symbol γ is set by `gcv_gamma` and controls optional inflation of the effective-degrees-of-freedom term.

For identity link, $r_i = y_i - \hat{\eta}_i$. For non-identity links, the residuals are $r_i = g((y_i + \delta)/(1 + 2\delta)) - (\hat{\eta}_i + \delta)/(1 + 2\delta)$, where $\delta \geq 0$ is a pseudocount that stabilizes the link transformation, automatically tuned within `tune_Lambda` if not supplied to `delta`. Non-Gaussian families with observation weights ω_i have their residuals scaled by ω_i . When the family provides a custom deviance residual function, that function is used in place of the link-scale residuals.

Pseudocount selection. The pseudocount δ is chosen to make the transformed response distribution most closely approximate a t -distribution with $N - 1$ degrees of freedom, in the sense of minimizing the (optionally weighted) mean absolute deviation between the sorted standardized transformed responses and the corresponding t -quantiles. This is solved via Brent's method over

$[10^{-64}, 1]$. When the link is identity, or when the response naturally lies in the domain of the link function, $\delta = 0$. This behavior is exposed through the `delta` argument in `lgspline`: supplying a fixed numeric value bypasses the internal search, while leaving it `NULL` allows the tuning code to choose the stabilizing pseudocount automatically when needed.

Meta-penalty regularization. A regularization term pulls the predictor- and partition-specific penalty parameters toward 1 on the raw scale:

$$P_{\text{meta}}(\lambda_w, \lambda_{l,k}) = \frac{1}{2} c_{\text{meta}} \sum_k \sum_l (\lambda_{l,k} - 1)^2 + \frac{1}{2} \cdot 10^{-32} (\lambda_w - 1)^2$$

where c_{meta} is a user-specified coefficient (`meta_penalty`). The gradient of P_{meta} on the log scale, incorporating the exp chain rule, is $\partial P_{\text{meta}} / \partial \theta_{l,k} = c_{\text{meta}} (\lambda_{l,k} - 1) \lambda_{l,k}$ and $\partial P_{\text{meta}} / \partial \theta_1 = 10^{-32} (\lambda_w - 1) \lambda_w$. The total objective is the selected tuning criterion plus P_{meta} .

Closed-Form Gradients for Tuning Criteria: Let ℓ denote the selected tuning criterion and

$$\mathbf{\Lambda}_k = \lambda_w (\mathbf{\Lambda}_s + \lambda_r \mathbf{\Lambda}_r + \sum_l \lambda_{l,k} \mathbf{\Lambda}_{l,k}).$$

The analytic gradient differentiates ℓ through \mathbf{G} , \mathbf{U} , and \mathbf{H} once, giving the partition-level derivative $\mathbf{M}_k = \partial \ell / \partial \mathbf{\Lambda}_k$. The penalty derivatives then follow directly:

$$\frac{\partial \ell}{\partial \lambda_w} = \frac{1}{\lambda_w} \sum_{k=0}^K \text{tr}(\mathbf{M}_k \mathbf{\Lambda}_k), \quad \frac{\partial \ell}{\partial \lambda_r} = \lambda_w \sum_{k=0}^K \text{tr}(\mathbf{M}_k \mathbf{\Lambda}_r),$$

and

$$\frac{\partial \ell}{\partial \lambda_{l,k}} = \lambda_w \text{tr}(\mathbf{M}_k \mathbf{\Lambda}_{l,k}).$$

Shared predictor penalties use the corresponding sum over partitions. Since optimization is on $\theta_j = \log \lambda_j$, each derivative is multiplied by λ_j . Thus, after \mathbf{M}_k is available, each additional penalty gradient costs only a blockwise trace product.

For exact leave-one-out, ℓ is the mean squared PRESS residual on the transformed problem, using blockwise constrained leverages rather than forming the full hat matrix. For GCV, ℓ uses the usual trace-based denominator, optionally inflated by `gcv_gamma`.

Optimization Procedure: Grid search initialization. The selected tuning criterion is evaluated over a grid of candidate values for (λ_w, λ_r) on the log scale. All combinations of user-supplied candidate vectors (`initial_wiggle` and `initial_flat`) are formed, and the combination yielding the smallest finite criterion value is selected as the starting point for BFGS optimization. Grid points producing non-finite objective values are discarded. If all grid points fail, an error is raised advising the user to check the data or adjust the grid. When `parallel_grideval = TRUE` and a cluster is supplied, these candidate fits are spread across workers. If more than six workers are available, additional raw-scale penalty pairs are drawn from $[\min / 10, \max \times 10]$ in each direction before the starting point is chosen.

Damped BFGS optimizer. A custom damped BFGS quasi-Newton optimizer, implemented in `tune_Lambda` through `.damped_bfgs()`, minimizes the selected tuning criterion plus P_{meta} . When analytic gradients are not preferred, or when the exact LOO leverage derivative appears too noisy for a given problem, a base-R `optim` call with method "BFGS" provides the finite-difference fallback. When `parallel_bfgs = TRUE` and a cluster is supplied, the damping trial values are evaluated as a batch across workers, and the best improving candidate is retained.

Iterations 1-2: steepest descent. The first two iterations use steepest descent with a damping factor α : $\phi^{(t+1)} = \phi^{(t)} - \alpha \nabla \phi$.

Iterations 3+: BFGS. From iteration 3, an inverse Hessian approximation $\mathbf{K}^{(t)}$ is maintained via the standard secant update. Let $\mathbf{s}^{(t)} = \phi^{(t)} - \phi^{(t-1)}$ and $\mathbf{v}^{(t)} = \nabla^{(t)} - \nabla^{(t-1)}$. The BFGS update is:

$$\mathbf{K}^{(t+1)} = (\mathbf{I} - \mathbf{u}\mathbf{v}^\top)\mathbf{K}^{(t)}(\mathbf{I} - \mathbf{u}\mathbf{v}^\top) + \mathbf{u}\mathbf{s}^\top, \quad \mathbf{u} = (\mathbf{v}^\top\mathbf{s})^{-1}.$$

When $|\mathbf{v}^\top\mathbf{s}| < 10^{-64}$, the approximation is reset to \mathbf{I} and the iteration is flagged for restart. The search direction is $\mathbf{d}^{(t)} = -\mathbf{K}^{(t)}\nabla^{(t)}$.

Step acceptance. A step is accepted if the new tuning criterion value is no larger than the old one. On rejection, α is halved. If $\alpha < 2^{-10}$ (early iterations) or $\alpha < 2^{-12}$ (later iterations), the optimizer terminates with the best solution found. With inequality constraints, the active set from the last accepted tuning evaluation initializes the next coefficient solve, avoiding repeated reconstruction of the same working set for nearby penalty values.

Convergence. After at least 10 iterations, the optimizer terminates when the absolute change in the tuning criterion is less than ϵ , when $\|\phi^{(t)} - \phi^{(t-1)}\|_\infty < \epsilon$, or when accepted criterion changes remain below a small scale-aware plateau threshold for several iterations. The plateau threshold is $\max\{\epsilon, \min(10^{-5}, 10^{-5}|\ell_{\text{best}}|)\}$.

Alternative. A base-R `optim` call with method "BFGS" and finite-difference gradients can be used instead via `use_custom_bfgs = FALSE`.

Post-optimization sample-size adjustment. After optimization, the tuned penalty parameters are divided by $(N + 1)/(N - 1)$ (equivalently multiplied by $(N - 1)/(N + 1)$) for both GCV- and LOO-based tuning. This decreases the final penalties at small sample sizes.

The tuning loop is implemented in `tune_Lambda`.

Incorporating Non-Spline Effects

Fixed effects and spline effects share the same partition-wise polynomial bookkeeping. Predictors included only linearly via `just_linear_without_interactions` or `just_linear_with_interactions` remain structurally linear. The first-derivative constraint then keeps their slope constant across partitions, while spline-expanded predictors remain free to vary nonlinearly.

When `blockfit = TRUE` is specified alongside `just_linear_without_interactions`, the flat-block path provides an alternative enforcement mechanism: flat coefficients are pooled across partitions during backfitting. The point estimate agrees with constraint projection, but uncertainty quantification differs; see the Blockfit section.

Integration

Because each partition retains an explicit polynomial representation, numerical integration can be carried out directly.

The user-facing method `integrate.lgspline` applies Gauss-Legendre quadrature to predictions from `predict.lgspline`.

Implementation: For a rectangular domain, `integrate.lgspline` constructs a tensor-product grid of Gauss-Legendre nodes, evaluates the fitted model, and forms the weighted sum. The fitted partition structure is handled internally.

The `vars` argument selects which predictors are integrated over. Predictors not listed in `vars` are held fixed at `initial_values` when supplied, or otherwise at the midpoint of their observed

training range. The optional `B_predict` argument makes it possible to integrate posterior draws or other alternate coefficient sets, and `n_quad` controls the number of Gauss-Legendre nodes used per integrated dimension.

Integration is performed on the response scale by default. Setting `link_scale = TRUE` instead integrates the linear predictor $\eta = f(\mathbf{t})$. For identity-link Gaussian models the two scales coincide.

Lagrange Multipliers

When `return_lagrange_multipliers = TRUE`, the multiplier vector

$$\boldsymbol{\lambda} = (\mathbf{A}^\top \mathbf{G} \mathbf{A})^{-1} \mathbf{A}^\top \hat{\boldsymbol{\beta}}$$

is returned. These quantify the sensitivity of the penalized objective to relaxing each smoothness or user-supplied equality constraint. When constraint target values are nonzero ($\mathbf{A}^\top \boldsymbol{\beta}_0 \neq \mathbf{0}$), the modified formulation is used:

$$\boldsymbol{\lambda} = (\mathbf{A}^\top \mathbf{G} \mathbf{A})^{-1} \mathbf{A}^\top (\hat{\boldsymbol{\beta}} - \boldsymbol{\beta}_0)$$

where $\mathbf{A}^\top \boldsymbol{\beta}_0$ is the vector of constraint target values. Multipliers are `NULL` when no constraints are active (\mathbf{A} is `NULL` or $K = 0$).

For inequality constraints, multipliers are returned as computed by `solve.QP`. The Lagrange multipliers for active inequality constraints can be used diagnostically to identify which shape constraints are most costly in terms of goodness of fit.

S3 Methods

Standard S3 methods are provided for objects of class `lgspline`:

- `print.lgspline` and `summary.lgspline`: Provide concise model summaries, with `print.summary.lgspline` formatting coefficient tables in a familiar regression-style layout.
- `logLik.lgspline`: Returns a standard `logLik` object. For Gaussian responses with identity link, the exact log-likelihood is computed. When a correlation structure is present via `VhalfInv`, the log-likelihood includes the $\log |\mathbf{V}^{-1/2}|$ adjustment and the corresponding whitened quadratic form. For other families, the method falls back to `family$aic` or a deviance-based approximation. An `include_prior` argument (default `TRUE`) optionally adds the Gaussian prior penalty interpretation of the smoothing spline penalty $-\frac{1}{2\sigma^2} \tilde{\boldsymbol{\beta}}^\top \boldsymbol{\Lambda} \tilde{\boldsymbol{\beta}}$ to obtain a penalized MAP log-likelihood. Alternate coefficient lists and fixed dispersions can be supplied through `B_predict` and `sigmasq_predict`.
- `predict.lgspline`: Produces fitted values and related quantities (e.g., derivatives and standard errors through `se.fit = TRUE`), lets `new_predictors` override `newdata`, accepts alternate coefficient lists through `B_predict`, and supports prediction on new predictor matrices consistent with the original spline expansions.
- `coef.lgspline`: Extracts partition-specific coefficient vectors.
- `confint.lgspline`: Extracts confidence intervals. When the inverse Hessian from BFGS optimization is available for correlation parameters, intervals for those correlation parameters are returned on the working (transformed) scale and should be back-transformed as described in the correlation section.

- `plot.lgspline`: For one-dimensional fits, produces base R graphics showing the fitted function (with optional partition-wise formulas) and supports overlay via `add = TRUE`. For two or more predictors, an interactive **plotly**-based visualization is returned. Specific predictors may be selected via `vars`.
- `integrate.lgspline`: Computes definite integrals of the fitted surface over rectangular domains by Gauss-Legendre quadrature.

Additional user-facing helpers include `wald_univariate` for coefficient-wise Wald inference, `generate_posterior` for posterior and posterior-predictive sampling, `generate_posterior_correlation` for correlation-aware posterior simulation, `equation` for closed-form display of the fitted partition formulas, and `find_extremum` for optimizing the fitted surface or a custom acquisition function built from it.

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 equation

Print Closed-Form Fitted Equation from lgspline Model

Description

Displays the closed-form polynomial equation for each partition of a fitted lgspline model, along with partition boundary or cluster center information. Optionally prints the first derivative, second derivative, or antiderivative of the fitted equation with respect to a single specified variable.

Usage

```
equation(object, ...)

## S3 method for class 'lgspline'
equation(
  object,
  digits = 4,
  scientific = FALSE,
  show_bounds = TRUE,
  predictor_names = NULL,
  response_name = NULL,
  collapse_zero = TRUE,
  first_derivative = NULL,
  second_derivative = NULL,
  antiderivative = NULL,
  ...
)

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

## S3 method for class 'lgspline'
equation(
  object,
  digits = 4,
  scientific = FALSE,
  show_bounds = TRUE,
```

```

    predictor_names = NULL,
    response_name = NULL,
    collapse_zero = TRUE,
    first_derivative = NULL,
    second_derivative = NULL,
    antiderivative = NULL,
    ...
)

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

Arguments

object	A fitted lgspline model object.
...	Not used.
digits	Integer; decimal places for coefficient display. Default 4.
scientific	Logical; use scientific notation for coefficients with absolute value $< 1e-3$ or $> 1e4$. Default FALSE.
show_bounds	Logical; display partition bounds (1D) or knot midpoint boundaries (multi-D). Default TRUE.
predictor_names	Character vector; custom names for predictor variables. If NULL (default), uses original column names or "_j_" labels.
response_name	Character; label for response. If NULL (default), uses "y" for identity link Gaussian, or "link(E[y])" otherwise.
collapse_zero	Logical; omit terms with coefficient exactly 0. Default TRUE.
first_derivative	Default: NULL. Character name or integer index of the predictor variable with respect to which the first derivative is printed. Only one variable at a time is supported. When non-NULL, the printed equations show df/dx_j for each partition.
second_derivative	Default: NULL. Character name or integer index of the predictor variable with respect to which the second derivative is printed. Only one variable at a time is supported. When non-NULL, the printed equations show d^2f/dx_j^2 for each partition. Ignored if first_derivative is also non-NULL.
antiderivative	Default: NULL. Character name or integer index of the predictor variable with respect to which the antiderivative (indefinite integral) is printed. Only one variable at a time is supported. When non-NULL, the printed equations show $\int f dx_j$ for each partition, with an unspecified constant of integration C . Ignored if first_derivative or second_derivative is also non-NULL.
x	An object returned by equation() for printing.

Details

For 1D models with K knots, partition boundaries are displayed as intervals on the predictor scale. For multi-predictor models, partition boundaries are computed as the midpoints between adjacent

cluster centers along each predictor dimension. When the model's `make_partition_list` contains knots (midpoint boundaries between clusters), those are used directly. Otherwise, cluster centers are displayed.

Coefficients are displayed on the original (unstandardized) predictor scale. For GLMs with non-identity link, the left-hand side shows the link function applied to the expected response.

Derivative and antiderivative modes. Only one of `first_derivative`, `second_derivative`, or `antiderivative` may be non-NULL. If more than one is supplied, the priority order is: first derivative, second derivative, antiderivative.

Derivatives and antiderivatives are computed symbolically from the polynomial coefficients. For a term ax^n , the first derivative is nax^{n-1} , the second derivative is $n(n-1)ax^{n-2}$, and the antiderivative is $ax^{n+1}/(n+1)$. Cross-terms (interactions) involving the target variable are differentiated or integrated with respect to that variable only, treating all other variables as constants.

A warning is emitted if the user attempts to differentiate or integrate with respect to more than one variable simultaneously. Multi-variable calculus operations should be performed one variable at a time by calling `equation()` repeatedly.

Value

Invisibly returns a list with components:

formulas Character vector of equation strings per partition.

bounds Matrix or list of partition boundary information.

link Character; link function name.

mode Character; one of "equation", "first_derivative", "second_derivative", or "antiderivative".

variable Character; the variable name for the calculus operation, or NULL if mode is "equation".

See Also

[lgspline](#), [plot.lgspline](#), [coef.lgspline](#)

Examples

```
## 1D example
set.seed(1234)
t <- runif(500, -5, 5)
y <- 2*sin(t) + 0.1*t^2 + rnorm(length(t), 0, 0.5)
fit <- lgspline(t, y, K = 2)
equation(fit)
equation(fit, digits = 2, predictor_names = "time")

## First derivative with respect to predictor
equation(fit, first_derivative = 1)

## Second derivative
equation(fit, second_derivative = 1)

## Antiderivative
equation(fit, antiderivative = 1)
```

```

## 2D example with named predictors
x1 <- runif(300, 0, 10)
x2 <- runif(300, 0, 10)
y <- x1 + 0.5*x2 + 0.1*x1*x2 + rnorm(300)
fit2d <- lgspline(cbind(x1, x2), y, K = 3)
equation(fit2d, predictor_names = c("Length", "Width"))

## Derivative w.r.t. first variable only
equation(fit2d, first_derivative = "Length",
         predictor_names = c("Length", "Width"))

## GLM example
y_bin <- rbinom(500, 1, plogis(0.5*t))
fit_glm <- lgspline(t, y_bin, family = binomial(), K = 1)
equation(fit_glm)

```

find_extremum

Find the Extremum of a Fitted lgspline

Description

Finds the global maximum or minimum of a fitted lgspline using L-BFGS-B, with options for partition-based heuristics, stochastic exploration, and custom objective functions (e.g., acquisition functions for Bayesian optimization).

Usage

```

find_extremum(
  object,
  vars = NULL,
  quick_heuristic = TRUE,
  initial = NULL,
  B_predict = NULL,
  minimize = FALSE,
  stochastic = FALSE,
  stochastic_draw = function(mu, sigma, ...) {
    N <- length(mu)
    rnorm(N, mu,
          sigma)
  },
  sigmasq_predict = object$sigmasq_tilde,
  custom_objective_function = NULL,
  custom_objective_derivative = NULL,
  ...
)

```

Arguments

object	A fitted lgspline model object.
vars	Integer or character vector; indices or names of predictors to optimize over. Default NULL optimizes all predictors.
quick_heuristic	Logical; if TRUE (default) searches only the best-performing partition. If FALSE, initiates searches from all partition local maxima.
initial	Numeric vector; optional starting values. Useful for fixing binary predictors. Default NULL.
B_predict	List; optional coefficient list for prediction, e.g. from generate_posterior . Default NULL uses object\$B.
minimize	Logical; find minimum instead of maximum. Default FALSE.
stochastic	Logical; add noise during optimization for exploration. Default FALSE.
stochastic_draw	Function; generates noise for stochastic optimization. Takes mu, sigma, and Default <code>rnorm(length(mu), mu, sigma)</code> .
sigmasq_predict	Numeric; variance for stochastic draws. Default <code>object\$sigmasq_tilde</code> .
custom_objective_function	Function; optional custom objective. Takes mu, sigma, y_best, Default NULL.
custom_objective_derivative	Function; optional gradient of custom_objective_function. Takes mu, sigma, y_best, d_mu, Default NULL.
...	Additional arguments passed to internal optimization routines.

Value

A list with elements:

- t** Numeric vector; predictor values at the extremum.
- y** Numeric; objective value at the extremum.

See Also

[lgspline](#), [generate_posterior](#)

Examples

```
set.seed(1234)
t <- runif(1000, -10, 10)
y <- 2*sin(t) + -0.06*t^2 + rnorm(length(t))
model_fit <- lgspline(t, y)
plot(model_fit)

max_point <- find_extremum(model_fit)
min_point <- find_extremum(model_fit, minimize = TRUE)
```

```

abline(v = max_point$t, col = 'blue')
abline(v = min_point$t, col = 'red')

## Expected improvement acquisition function
ei_obj <- function(mu, sigma, y_best, ...) {
  d <- y_best - mu
  d * pnorm(d/sigma) + sigma * dnorm(d/sigma)
}
ei_deriv <- function(mu, sigma, y_best, d_mu, ...) {
  d <- y_best - mu
  z <- d/sigma
  d_z <- -d_mu/sigma
  pnorm(z)*d_mu - d*dnorm(z)*d_z + sigma*z*dnorm(z)*d_z
}

post_draw <- generate_posterior(model_fit)
acq <- find_extremum(model_fit,
                    stochastic = TRUE,
                    B_predict = post_draw$post_draw_coefficients,
                    sigmasq_predict = post_draw$post_draw_sigmasq,
                    custom_objective_function = ei_obj,
                    custom_objective_derivative = ei_deriv)
abline(v = acq$t, col = 'green')

```

generate_posterior *Generate Posterior Samples from a Fitted lgspline*

Description

Draws from the posterior distribution of model coefficients, with optional dispersion sampling, posterior predictive draws, and propagation of uncertainty in estimated correlation parameters.

Usage

```

generate_posterior(
  object,
  new_sigmasq_tilde = object$sigmasq_tilde,
  new_predictors = NULL,
  theta_1 = 0,
  theta_2 = 0,
  posterior_predictive_draw = function(N, mean, sqrt_dispersion, ...) {
    rnorm(N,
          mean, sqrt_dispersion)
  },
  draw_dispersion = TRUE,
  include_posterior_predictive = FALSE,
  num_draws = 1,
  enforce_qp_constraints = TRUE,

```

```

draw_correlation = FALSE,
correlation_param_mean = NULL,
correlation_param_vcov = NULL,
correlation_VhalfInv_fxn = NULL,
correlation_Vhalf_fxn = NULL,
correlation_param_vcov_scale = NULL,
include_warnings = TRUE,
...
)

```

Arguments

object A fitted lgspline model object.

new_sigmasq_tilde Numeric; dispersion $\tilde{\sigma}^2$ used as the point estimate when `draw_dispersion = FALSE`. Default `object$sigmasq_tilde`.

new_predictors Matrix; predictor matrix for posterior predictive sampling. Default uses in-sample predictors.

theta_1 Numeric; shape increment for the inverse-gamma prior on σ^2 . Default 0.

theta_2 Numeric; rate increment for the inverse-gamma prior. Default 0.

posterior_predictive_draw Function; sampler for posterior predictive realisations. Must accept `N`, `mean`, `sqrt_dispersion`, ... Default `rnorm`.

draw_dispersion Logical; sample σ^2 from its posterior. Default `TRUE`.

include_posterior_predictive Logical; generate posterior predictive draws at `new_predictors`. Default `FALSE`.

num_draws Positive integer; number of draws. Default 1.

enforce_qp_constraints Logical; if `TRUE`, enforce active QP inequality constraints during posterior sampling via the stored elliptical-slice constrained sampler. Default `TRUE`.

draw_correlation Logical; propagate correlation parameter uncertainty. Requires `VhalfInv_fxn` and `VhalfInv_params_estimates` in the fitted object. Default `FALSE`.

correlation_param_mean Numeric vector; mean of the approximate normal posterior for correlation parameters on the unbounded (working) scale. Default: `object$VhalfInv_params_estimates`.

correlation_param_vcov Matrix; variance-covariance for correlation parameter draws. Default: inverse Hessian from BFGS (`object$VhalfInv_params_vcov`).

correlation_VhalfInv_fxn Function; maps correlation parameter vector to $\mathbf{V}^{-1/2}$. Default `object$VhalfInv_fxn`.

correlation_Vhalf_fxn Function or `NULL`; maps to $\mathbf{V}^{1/2}$. Passed through to `generate_posterior_correlation`; the current correlation-aware posterior path only requires `correlation_VhalfInv_fxn`.

correlation_param_vcov_scale	NULL or numeric; if supplied, divides a user-supplied correlation_param_vcov by this value before passing it to <code>generate_posterior_correlation</code> . When NULL, no additional scaling is applied.
include_warnings	Logical; emit warnings for degenerate draws, constraint violations, etc. Default TRUE.
...	Additional arguments forwarded to the GLM weight function, dispersion function, and posterior_predictive_draw.

Details

Uses a Laplace approximation centred at the MAP estimate for non-Gaussian responses.

Dispersion posterior. When `draw_dispersion = TRUE`, σ^2 is drawn from

$$\sigma^2 \mid \mathbf{y} \sim \text{InvGamma}(\alpha_1, \alpha_2),$$

where

$$\alpha_1 = \theta_1 + \frac{1}{2}(N - s \cdot \text{tr}(\mathbf{H})), \quad \alpha_2 = \theta_2 + \frac{1}{2}(N - s \cdot \text{tr}(\mathbf{H}))\tilde{\sigma}^2,$$

$\mathbf{H} = \mathbf{XUGX}^\top$ is the hat matrix, $s = 1$ when `unbias_dispersion = TRUE` (else $s = 0$), and $\theta_1 = -1, \theta_2 = 0$ recovers an improper uniform prior on σ^2 , while $\theta_1 = 0, \theta_2 = 0$ gives the usual scale-invariant improper prior $p(\sigma^2) \propto 1/\sigma^2$.

Correlation parameter posterior. When `draw_correlation = TRUE` and the fitted model contains an estimated correlation structure, each draw first samples $\boldsymbol{\rho}$ from

$$\boldsymbol{\rho}^{(m)} \sim \mathcal{N}(\hat{\boldsymbol{\rho}}_{\text{REML}}, \mathbf{H}_{\text{BFGS}}^{-1}),$$

rebuilds the posterior covariance under the drawn correlation structure (reusing all pre-computed design matrices, constraints, and penalty matrices) and then draws coefficients from the updated posterior. Knot placement, partitioning, coefficient re-estimation, and GCV tuning are skipped entirely. Draws producing non-positive-definite correlation matrices are rejected and redrawn (up to 50 attempts).

When `draw_correlation = FALSE` (default), correlation parameters are fixed at their estimated values.

Inequality constraints. Active QP inequalities can be enforced during posterior sampling via elliptical slice sampling, producing draws from the corresponding truncated multivariate normal posterior on the coefficient scale. The public `enforce_qp_constraints` argument is forwarded to the stored sampler for both the standard and correlation-aware posterior paths.

Value

When `num_draws = 1`, a named list:

- post_draw_coefficients** List of length $K+1$; per-partition coefficient vectors on the original scale.
- post_draw_sigmasq** Drawn (or fixed) dispersion.
- post_pred_draw** Posterior predictive vector (only when `include_posterior_predictive = TRUE`).
- post_draw_correlation_params** Drawn correlation parameters on the working scale (only when `draw_correlation = TRUE`).

When `num_draws > 1`, each element becomes a list of length `num_draws`, and `post_pred_draw` (if requested) is an $N_{\text{new}} \times M$ matrix, where $M = \text{num_draws}$.

See Also

[lgspline](#), [generate_posterior_correlation](#), [wald_univariate](#)

Examples

```
set.seed(1234)
n_blocks <- 100; block_size <- 5; N <- n_blocks * block_size
rho_true <- 0.3
t <- seq(-5, 5, length.out = N)
true_mean <- sin(t)
errors <- Reduce("rbind",
  lapply(1:n_blocks, function(i) {
    sigma <- diag(block_size) + rho_true *
      (matrix(1, block_size, block_size) - diag(block_size))
    matsqrt(sigma) %*% rnorm(block_size)
  })
)
y <- true_mean + errors * 0.5

model_fit <- lgspline(t, y,
  K = 3,
  correlation_id = rep(1:n_blocks, each = block_size),
  correlation_structure = "exchangeable",
  include_warnings = FALSE
)

## Propagate correlation uncertainty across 50 draws
post <- generate_posterior(model_fit,
  draw_correlation = TRUE, num_draws = 50,
  include_warnings = FALSE
)

## Fixed correlation parameters for comparison
post_fixed <- generate_posterior(model_fit, num_draws = 50)

corr_draws <- unlist(post$post_draw_correlation_params)
rho_draws <- exp(-exp(corr_draws))
print(summary(rho_draws))
```

generate_posterior_correlation

Generate Posterior Samples Propagating Correlation Parameter Uncertainty

Description

Called internally by `generate_posterior` when `draw_correlation = TRUE`, but can be used directly for finer control. For each draw, samples the correlation parameter vector from its approximate normal posterior, rebuilds the posterior covariance under that drawn correlation structure without re-solving for a new coefficient mode, then draws coefficients from the updated posterior.

Usage

```
generate_posterior_correlation(
  object,
  new_sigmasq_tilde = object$sigmasq_tilde,
  new_predictors = NULL,
  theta_1 = 0,
  theta_2 = 0,
  posterior_predictive_draw = function(N, mean, sqrt_dispersion, ...) {
    rnorm(N,
          mean, sqrt_dispersion)
  },
  draw_dispersion = TRUE,
  include_posterior_predictive = FALSE,
  num_draws = 1,
  enforce_qp_constraints = TRUE,
  correlation_param_mean = NULL,
  correlation_param_vcov_sc = NULL,
  correlation_VhalfInv_fxn = NULL,
  correlation_Vhalf_fxn = NULL,
  include_warnings = TRUE,
  ...
)
```

Arguments

<code>object</code>	A fitted <code>lgspline</code> object with a correlation structure (i.e., <code>VhalfInv_fxn</code> and <code>VhalfInv_params_estimates</code> present, or supplied via override arguments).
<code>new_sigmasq_tilde</code>	Numeric; dispersion starting value when <code>draw_dispersion = FALSE</code> . Default <code>object\$sigmasq_tilde</code> .
<code>new_predictors</code>	Matrix or <code>NULL</code> ; predictor matrix for posterior predictive sampling. Default uses in-sample predictors.
<code>theta_1</code>	Numeric; shape increment for the inverse-gamma prior. Default 0.
<code>theta_2</code>	Numeric; rate increment for the inverse-gamma prior. Default 0.
<code>posterior_predictive_draw</code>	Function; sampler for posterior predictive realisations. Default <code>rnorm</code> .
<code>draw_dispersion</code>	Logical; sample σ^2 within each draw. Default <code>TRUE</code> .
<code>include_posterior_predictive</code>	Logical; generate posterior predictive draws. Default <code>FALSE</code> .

num_draws	Positive integer; number of draws (each requires one correlation parameter sample and one covariance rebuild). Default 1.
enforce_qp_constraints	Logical; if TRUE, enforce active QP inequality constraints during each coefficient draw via the stored elliptical-slice constrained sampler. Default TRUE.
correlation_param_mean	Numeric vector or NULL; mean of the approximate normal posterior on the working scale. Default: object\$VhalfInv_params_estimates. Supplying this allows correlation draws for models fit with a fixed (non-optimised) correlation structure.
correlation_param_vcov_sc	Matrix or NULL; variance-covariance on the working scale. Default: object\$VhalfInv_params_vcov. No further scaling is applied within this function.
correlation_VhalfInv_fxn	Function or NULL; maps parameter vector to $\mathbf{V}^{-1/2}$. Default object\$VhalfInv_fxn.
correlation_Vhalf_fxn	Function or NULL; maps to $\mathbf{V}^{1/2}$. Not consumed in the current method body but resolved for interface consistency and potential future use.
include_warnings	Logical; emit warnings. Default TRUE.
...	Additional arguments forwarded to the GLM weight function, dispersion function, and posterior_predictive_draw.

Details

Each draw proceeds in three steps:

1. **Draw correlation parameters.** $\boldsymbol{\rho}^{(m)} \sim \mathcal{N}(\hat{\boldsymbol{\rho}}_{\text{REML}}, \mathbf{H}_{\text{BFGS}}^{-1})$ on the unbounded working scale. Draws producing a non-PD correlation matrix are rejected and redrawn (up to 50 attempts); if all fail, the point estimate is used with a warning.
2. **Rebuild posterior covariance.** Using the already-expanded \mathbf{X}_k , \mathbf{A} , and $\boldsymbol{\Lambda}$ from the original fit, recompute only the covariance-side quantities implied by the drawn correlation structure:

$$\mathbf{G}_{\text{correct}}^{(m)} = \left(\mathbf{X}^\top \mathbf{W} \mathbf{D} \mathbf{V}^{-1} (\boldsymbol{\rho}^{(m)}) \mathbf{X} + \boldsymbol{\Lambda} \right)^{-1},$$

and from this the updated constraint projection $\mathbf{U}^{(m)}$ and effective degrees of freedom $\text{trace}(\mathbf{H}^{(m)})$.

The coefficient mode $\hat{\boldsymbol{\beta}}_{\text{raw}}$ and fitted mean $\tilde{\mathbf{y}}$ are held fixed at the original fit values. Knot placement, partitioning, polynomial expansion, penalty tuning, and coefficient re-estimation are all skipped entirely.

3. **Draw coefficients.** Updated quantities (\mathbf{U} , $\mathbf{G}_{\text{half_correct}}$, $\mathbf{V}_{\text{halfInv}}$, sigmasq_tilde , trace_XUGX) are passed to the stored closure via `override_*` arguments so that the draw is centred at the original mode but uses the covariance implied by the drawn correlation structure. The stored mode `object$B_raw` is passed as `override_B_raw` so it is not recomputed.

Why the mode is held fixed. Re-solving for a new MAP estimate under each drawn correlation structure is expensive, requires iterative solvers, and risks convergence failures on draws far from the REML estimate. The posterior draw is centred at the original mode, which remains a reasonable

approximation when the REML surface is not sharply peaked. The covariance update captures the primary effect of correlation uncertainty on posterior width and shape.

BFGS inverse Hessian caveat. The BFGS inverse Hessian approximation for the correlation parameter covariance is asymptotically valid but may be poor for small samples, near-boundary estimates, or multimodal REML surfaces. It is not guaranteed to converge to the observed information matrix. Users should inspect `object$VhalfInv_params_vcov` before relying on these draws.

Value

When `num_draws = 1`, a named list:

post_draw_coefficients List of length $K+1$; per-partition coefficient vectors on the original scale.

post_draw_sigmasq Drawn dispersion.

post_pred_draw Posterior predictive vector (only when `include_posterior_predictive = TRUE`).

post_draw_correlation_params Drawn correlation parameters on the working scale.

When `num_draws > 1`:

post_draw_coefficients List of `num_draws` lists of $K+1$ coefficient vectors.

post_draw_sigmasq List of `num_draws` scalars.

post_pred_draw $N_{\text{new}} \times M$ matrix, where $M = \text{num_draws}$ (only when `include_posterior_predictive = TRUE`).

post_draw_correlation_params List of `num_draws` vectors.

See Also

[generate_posterior](#), [lgspline](#), [lgspline.fit](#)

Examples

```
## See ?generate_posterior for a complete worked example.
```

get_B_verification_examples

Verification Examples for get_B

Description

Simple, self-contained examples that reviewers can run to verify that `get_B` produces correct output. These exercise Path 2 (Gaussian, no correlation) and Path 3 (binomial GLM).

Examples

```

## Not run:
## Example 1: Path 2 - Gaussian identity, with knots
set.seed(1234)
t <- runif(200, -5, 5)
y <- sin(t) + rnorm(200, 0, 0.5)
fit1 <- lgspline(t, y, K = 3, opt = FALSE, wiggle_penalty = 1e-4)
stopifnot(inherits(fit1, "lgspline"))
stopifnot(length(fit1$B) == 4) # K+1 = 4 partitions
cat("Example 1 passed: Gaussian identity, K=3\n")

preds1 <- predict(fit1, new_predictors = rnorm(10))
stopifnot(all(is.finite(preds1)))
cat(" Predictions finite: OK\n")

## Example 2: Path 2 - Gaussian identity, K=0 (no constraints)
fit2 <- lgspline(t, y, K = 0, opt = FALSE, wiggle_penalty = 1e-4)
stopifnot(inherits(fit2, "lgspline"))
stopifnot(length(fit2$B) == 1)
preds2 <- predict(fit2, new_predictors = rnorm(10))
stopifnot(all(is.finite(preds2)))
cat("Example 2 passed: Gaussian identity, K=0\n")

## Example 3: Path 3 - Binomial GLM
y_bin <- rbinom(200, 1, plogis(sin(t)))
fit3 <- lgspline(t, y_bin, K = 2, family = binomial(),
               opt = FALSE, wiggle_penalty = 1e-3)
stopifnot(inherits(fit3, "lgspline"))
preds3 <- predict(fit3, new_predictors = rnorm(10))
stopifnot(all(preds3 >= 0 & preds3 <= 1))
cat("Example 3 passed: Binomial GLM, K=2\n")

## Example 4: Path 2 with QP constraints (monotonic increase)
t_sorted <- sort(runif(100, -3, 3))
y_mono <- t_sorted + 0.5 * sin(t_sorted) + rnorm(100, 0, 0.3)
fit4 <- lgspline(t_sorted, y_mono, K = 2,
               qp_monotonic_increase = TRUE,
               opt = FALSE, wiggle_penalty = 1e-4)
preds4 <- predict(fit4, new_predictors = cbind(sort(rnorm(50))))
stopifnot(all(diff(preds4) >= -sqrt(.Machine$double.eps)))
cat("Example 4 passed: Monotonic increase QP constraint\n")

## Example 5: Path 2 with range constraints
fit5 <- lgspline(t, y, K = 3,
               qp_range_lower = -2, qp_range_upper = 2,
               opt = FALSE, wiggle_penalty = 1e-4)
stopifnot(all(fit5$ytilde >= -2 - 0.01))
stopifnot(all(fit5$ytilde <= 2 + 0.01))
cat("Example 5 passed: Range constraints\n")

## Example 6: Multi-predictor
set.seed(1234)

```

```

x1 <- rnorm(300)
x2 <- rnorm(300)
y6 <- sin(x1) + cos(x2) + rnorm(300, 0, 0.5)
fit6 <- lgspline(cbind(x1, x2), y6, K = 4,
                opt = FALSE, wiggle_penalty = 1e-5)
stopifnot(inherits(fit6, "lgspline"))
stopifnot(fit6$q_predictors == 2)
cat("Example 6 passed: 2D predictor, K=4\n")

## Example 7: Coefficient consistency (determinism)
set.seed(1234)
t7 <- runif(150, -4, 4)
y7 <- 2 * cos(t7) + rnorm(150, 0, 0.4)
fit7a <- lgspline(t7, y7, K = 2, opt = FALSE, wiggle_penalty = 1e-5)
set.seed(999)
fit7b <- lgspline(t7, y7, K = 2, opt = FALSE, wiggle_penalty = 1e-5)
max_diff <- max(abs(unlist(fit7a$B) - unlist(fit7b$B)))
stopifnot(max_diff < 1e-12)
cat("Example 7 passed: Deterministic coefficient reproduction\n")

cat("\nAll verification examples passed.\n")

## End(Not run)

```

integrate

Generic for Numerical Integration

Description

S3 generic that dispatches to `integrate.lgspline` for fitted `lgspline` objects and falls back to [integrate](#) for ordinary functions.

Usage

```
integrate(f, ...)
```

```
## Default S3 method:
integrate(f, ...)
```

Arguments

`f` A fitted model object or a function.
`...` Arguments passed to methods.

integrate.lgspline *Definite Integral of a Fitted lgspline*

Description

Given a fitted lgspline object, computes the definite integral of the fitted surface over a rectangular domain using Gauss–Legendre quadrature on predict().

Usage

```
## S3 method for class 'lgspline'
integrate(
  f,
  lower,
  upper,
  vars = NULL,
  initial_values = NULL,
  B_predict = NULL,
  link_scale = FALSE,
  n_quad = 50L,
  ...
)
```

Arguments

f	A fitted lgspline object.
lower	Numeric vector of lower bounds, one per integration variable. Scalar values are recycled.
upper	Numeric vector of upper bounds, one per integration variable. Scalar values are recycled.
vars	Default: NULL. Character or integer vector identifying which predictor(s) to integrate over. When NULL all numeric predictors are integrated simultaneously.
initial_values	Default: NULL. Numeric vector of length q supplying fixed values for predictors not among the integration variables. When NULL, non-integration predictors are held at the midpoint of their training range.
B_predict	Default: NULL. Optional list of coefficient vectors, one per partition. When NULL the fitted coefficients are used.
link_scale	Default: FALSE. Logical; when TRUE the integral is computed on the link (linear predictor) scale η rather than the response scale μ .
n_quad	Default: 50. Number of Gauss–Legendre nodes per integration dimension.
...	Additional arguments (currently unused; present for S3 method compatibility).

Value

A numeric scalar: the estimated definite integral.

Method

The integration domain is discretised into a tensor-product grid of Gauss–Legendre quadrature nodes. Predicted values at each node come from the model’s `predict()` method, which correctly handles partition assignment and piecewise polynomial evaluation. The integral is the weighted sum

$$\int_{a_1}^{b_1} \cdots \int_{a_d}^{b_d} \hat{f}(\mathbf{t}) dt_1 \cdots dt_d \approx \sum_{i=1}^M w_i \hat{f}(\mathbf{t}_i)$$

where $M = n_{\text{quad}}^d$ and each weight incorporates the Jacobian $(b_j - a_j)/2$ for the affine map from $[-1, 1]$ to $[a_j, b_j]$. Nodes and weights on $[-1, 1]$ are computed via the Golub–Welsch algorithm (eigenvalues of the symmetric tridiagonal Jacobi matrix).

For smooth polynomials, 30–50 nodes per dimension is typically sufficient; highly partitioned models (large K) may benefit from more. Total evaluation points scale as n_{quad}^d , so problems with $d \geq 4$ may require reducing `n_quad`.

Integration scale

By default (`link_scale = FALSE`), integration is on the response scale $\mu = g^{-1}(\eta)$. Setting `link_scale = TRUE` integrates the linear predictor $\eta = \mathbf{x}^\top \boldsymbol{\beta}$ directly, which is useful when the quantity of interest is the area under the link-transformed surface rather than the response. For the identity link the two coincide.

Examples

```
## 1-D: integral of fitted sin(t) over [-pi, pi] should be near 0
set.seed(1234)
t <- seq(-pi, pi, length.out = 1000)
y <- sin(t) + rnorm(length(t), 0, 0.01)
fit <- lgspline(t, y, K = 4, opt = FALSE)
integrate(fit, lower = -pi, upper = pi)

## Base R integrate still works as expected
integrate(sin, lower = -pi, upper = pi)

## 2-D: volume under fitted volcano surface
data(volcano)
vlong <- cbind(
  rep(seq_len(nrow(volcano)), ncol(volcano)),
  rep(seq_len(ncol(volcano)), each = nrow(volcano)),
  as.vector(volcano)
)
colnames(vlong) <- c("Length", "Width", "Height")
fit_v <- lgspline(vlong[, 1:2], vlong[, 3], K = 18,
  include_quadratic_interactions = TRUE, opt = FALSE)
integrate(fit_v, lower = c(1, 1), upper = c(87, 61))
```

leave_one_out	<i>Compute Leave-One-Out Cross-Validated Predictions for Gaussian Response/Identity Link under Constraint</i>
---------------	---

Description

Computes the leave-one-out cross-validated predictions from a model fit, assuming Gaussian-distributed response with identity link.

The LOO closed-formula for observation i is $\hat{y}_{(-i)} = y_i - \frac{1}{1-H_{ii}}(y_i - \hat{y}_i)$ where \mathbf{H} is the effective hat matrix under smoothing constraints, adjusted for weights and correlation structure if present.

Observations with leverage at or above `leverage_threshold` are flagged in a warning, since extreme hat values can make the shortcut numerically unreliable. The default `leverage_threshold = 100` is intentionally permissive, so users who want diagnostic warnings for large H_{ii} should set a smaller threshold explicitly.

For related discussion of prediction-sum-of-squares calculations under linear restrictions, see Tarpey (2000), who studies the PRESS statistic for restricted least squares. That setting is closely related to the constraint-adjusted hat-matrix shortcut used here.

Usage

```
leave_one_out(model_fit, leverage_threshold = 100)
```

Arguments

`model_fit` A fitted lgspline model object.

`leverage_threshold`
 Numeric scalar. Observations with $H_{ii} \geq \text{leverage_threshold}$ are treated as high-leverage for the warning below. Default 100.

Value

A vector of leave-one-out cross-validated predictions

References

Tarpey, T. (2000). A note on the prediction sum of squares statistic for restricted least squares. *The American Statistician*, 54(2), 116–118. doi:10.2307/2686028

Examples

```
## Basic usage with Gaussian response, computing PRESS
set.seed(1234)
t <- rnorm(50)
y <- sin(t) + rnorm(50, 0, .25)
model_fit <- lgspline(t, y)
loo <- leave_one_out(model_fit)
press <- mean((y - loo)^2, na.rm = TRUE)
```

```
plot(loo, y,
     main = "LOO Cross-Validation Prediction vs. Observed Response",
     xlab = 'Prediction', ylab = 'Response')
abline(0, 1)
```

lgspline

Fit Lagrangian Multiplier Smoothing Splines

Description

`lgspline` fits penalized piecewise-polynomial regression splines in a monomial basis, with smoothness imposed directly as linear constraints at partition boundaries rather than absorbed into a special spline basis.

The estimator is obtained with Lagrangian multipliers and enforces the usual smoothing restrictions:

- Equivalent fitted values at knots
- Equivalent first derivatives at knots, with respect to predictors
- Equivalent second derivatives at knots, with respect to predictors

The coefficients are penalized by the closed-form cubic smoothing-spline penalty, with optional predictor- and partition-specific modifications. The same framework extends to GLMs, shape constraints, and correlated responses, while keeping the fitted partition-wise polynomials explicit and interpretable.

Usage

```
lgspline(
  predictors = NULL,
  y = NULL,
  formula = NULL,
  response = NULL,
  standardize_response = TRUE,
  standardize_predictors_for_knots = TRUE,
  standardize_expansions_for_fitting = TRUE,
  family = gaussian(),
  glm_weight_function = default_glm_weight_function,
  schur_correction_function = function(X, y, B, dispersion, order_list, K, family,
  observation_weights, ...) {
  lapply(1:(K + 1), function(k) 0)
},
  need_dispersion_for_estimation = FALSE,
  dispersion_function = function(mu, y, order_indices, family, observation_weights,
  VhalfInv, ...) {
  if (!is.null(VhalfInv)) {
    VhalfInv <-
```

```

VhalfInv[order_indices, order_indices]
  c(mean((tcrossprod(VhalfInv, t(y -
mu)))^2/family$variance(mu)))
  }
  else {
    c(mean((y -
mu)^2/family$variance(mu)))
  }
},
K = NULL,
custom_knots = NULL,
cluster_on_indicators = FALSE,
make_partition_list = NULL,
previously_tuned_penalties = NULL,
smoothing_spline_penalty = NULL,
opt = TRUE,
use_custom_bfgs = TRUE,
delta = NULL,
tol = 10 * sqrt(.Machine$double.eps),
tuning_criterion = "loo",
gcv_gamma = 1.4,
initial_wiggle = c(2e-12, 2e-07, 2e-04, 0.2),
initial_flat = c(0.5, 5),
wiggle_penalty = 2e-07,
flat_ridge_penalty = 0.5,
unique_penalty_per_partition = TRUE,
unique_penalty_per_predictor = TRUE,
meta_penalty = 1e-08,
predictor_penalties = NULL,
partition_penalties = NULL,
include_quadratic_terms = TRUE,
include_cubic_terms = TRUE,
include_quartic_terms = NULL,
include_2way_interactions = TRUE,
include_3way_interactions = TRUE,
include_quadratic_interactions = FALSE,
offset = c(),
just_linear_with_interactions = NULL,
just_linear_without_interactions = NULL,
exclude_interactions_for = NULL,
exclude_these_expansions = NULL,
custom_basis_fxn = NULL,
include_constrain_fitted = TRUE,
include_constrain_first_deriv = TRUE,
include_constrain_second_deriv = TRUE,
include_constrain_interactions = TRUE,
add_first_and_second_derivative_constraints = NULL,
qr_pivot_smoothing_constraints = TRUE,

```

```
cl = NULL,
chunk_size = NULL,
parallel_eigen = TRUE,
parallel_trace = FALSE,
parallel_aga = FALSE,
parallel_matmult = FALSE,
parallel_qr = FALSE,
parallel_bfgs = FALSE,
parallel_grideval = TRUE,
parallel_qr_qp = FALSE,
parallel_unconstrained = TRUE,
parallel_find_neighbors = TRUE,
parallel_penalty = FALSE,
parallel_make_constraint = TRUE,
unconstrained_fit_fxn = unconstrained_fit_default,
keep_weighted_Lambda = FALSE,
iterate_tune = TRUE,
iterate_final_fit = TRUE,
blockfit = TRUE,
qp_score_function = function(X, y, mu, order_list, dispersion, VhalfInv,
  observation_weights, ...) {
  default_qp_score_function(X, y, mu, order_list,
    dispersion, VhalfInv, observation_weights, family, ...)
},
qp_observations = NULL,
qp_Amat = NULL,
qp_bvec = NULL,
qp_meq = 0,
qp_positive_derivative = FALSE,
qp_negative_derivative = FALSE,
qp_positive_2ndderivative = FALSE,
qp_negative_2ndderivative = FALSE,
qp_monotonic_increase = FALSE,
qp_monotonic_decrease = FALSE,
qp_range_upper = NULL,
qp_range_lower = NULL,
qr_pivot_inequality_constraints = FALSE,
qp_Amat_fxn = NULL,
qp_bvec_fxn = NULL,
qp_meq_fxn = NULL,
constraint_values = cbind(),
constraint_vectors = cbind(),
return_G = TRUE,
return_Ghalf = TRUE,
return_U = TRUE,
estimate_dispersion = TRUE,
unbias_dispersion = NULL,
return_varcovmat = TRUE,
```

```

exact_varcovmat = FALSE,
return_lagrange_multipliers = FALSE,
custom_penalty_mat = NULL,
cluster_args = c(custom_centers = NA, nstart = 10),
dummy_divisor = 1.2345672152894e-22,
dummy_adder = 2.234567210529e-18,
verbose = FALSE,
verbose_tune = FALSE,
dummy_fit = FALSE,
auto_encode_factors = TRUE,
observation_weights = NULL,
do_not_cluster_on_these = c(),
neighbor_tolerance = 1 + 1e-08,
null_constraint = NULL,
critical_value = qnorm(1 - 0.05/2),
data = NULL,
weights = NULL,
no_intercept = FALSE,
correlation_id = NULL,
spacetime = NULL,
correlation_structure = NULL,
VhalfInv = NULL,
Vhalf = NULL,
VhalfInv_fxn = NULL,
Vhalf_fxn = NULL,
VhalfInv_par_init = c(),
REML_grad = NULL,
custom_VhalfInv_loss = NULL,
VhalfInv_logdet = NULL,
include_warnings = TRUE,
penalty_args = NULL,
tuning_args = NULL,
expansion_args = NULL,
constraint_args = NULL,
qp_args = NULL,
parallel_args = NULL,
covariance_args = NULL,
return_args = NULL,
glm_args = NULL,
...
)

```

Arguments

predictors	Default: NULL. Numeric matrix or data frame of predictor variables, or a formula when using the formula interface.
y	Default: NULL. Numeric response variable vector.
formula	Default: NULL. Optional statistical formula for model specification, supporting

	spl() (and the alias s()) for spline terms.
response	Default: NULL. Alternative name for response variable.
standardize_response	Default: TRUE. Logical indicator controlling whether the response variable should be centered and scaled before model fitting. Only offered for identity link functions.
standardize_predictors_for_knots	Default: TRUE. Logical flag controlling whether predictors are internally standardized for partitioning / knot placement. The exact transformation is handled inside <code>make_partitions</code> and depends on the effective clustering dimension.
standardize_expansions_for_fitting	Default: TRUE. Logical switch to standardize polynomial basis expansions during model fitting. Design matrices, variance-covariance matrices, and coefficients are backtransformed after fitting. U and G remain on the transformed scale; B_raw corresponds to coefficients on the expansion-standardized scale.
family	Default: <code>gaussian()</code> . GLM family specifying the error distribution and link function. Minimally requires: family name, link name, linkfun, linkinv, variance.
glm_weight_function	Default: GLM working weight $\text{family}\$\mu.\text{eta}(\text{eta})^2 / \text{family}\$\text{variance}(\mu)$, optionally multiplied by <code>observation_weights</code> .
schur_correction_function	Default: function returning list of zeros. Computes Schur complements S added to G : $\mathbf{G}^* = (\mathbf{G}^{-1} + \mathbf{S})^{-1}$.
need_dispersion_for_estimation	Default: FALSE. Logical indicator specifying whether a dispersion parameter is required for coefficient estimation (e.g. Weibull AFT).
dispersion_function	Default: function returning mean squared residuals. Custom function for estimating the exponential dispersion parameter.
K	Default: NULL. Integer specifying the number of knot locations. Intuitively, total partitions minus 1.
custom_knots	Default: NULL. Optional matrix providing user-specified knot locations in 1-D.
cluster_on_indicators	Default: FALSE. Logical flag for whether indicator variables should be used for clustering knot locations.
make_partition_list	Default: NULL. Optional list allowing direct specification of custom partition assignments. The <code>make_partition_list</code> returned by one model can be supplied here to reuse knot locations.
previously_tuned_penalties	Default: NULL. Optional list of pre-computed penalty components from a previous model fit.
smoothing_spline_penalty	Default: NULL. Optional custom smoothing spline penalty matrix.

<code>opt</code>	Default: TRUE. Logical switch controlling automatic penalty optimization.
<code>use_custom_bfgs</code>	Default: TRUE. Selects between a native damped-BFGS implementation with closed-form gradients or base R's BFGS with finite-difference gradients. The native path is usually faster, while the finite-difference fallback can be preferable when tuning under exact LOO and the leverage derivative is numerically noisy.
<code>delta</code>	Default: NULL. Numeric pseudocount for stabilizing optimization in non-identity link function scenarios.
<code>tol</code>	Default: $10 \times \sqrt{.Machine\$double.eps}$. Numeric convergence tolerance.
<code>tuning_criterion</code>	Default: "loo". Character scalar selecting the tuning criterion. Use "loo" for exact leave-one-out on the transformed tuning problem, or "gcv" for the generalized cross-validation criterion. The LOO path computes the needed hat-matrix diagonal exactly from blockwise constrained-G quantities, without explicitly forming the full projection matrix or full hat matrix. In empirical diagnostics, the observation-wise derivative of the LOO leverage term can be numerically delicate even when the overall tuning criterion and fitted penalties remain well behaved; users who prefer a more conservative optimization path can set <code>use_custom_bfgs = FALSE</code> . Penalty gradients reuse the partition penalty-matrix derivatives $\partial \ell / \partial \Lambda_k$, so additional tuned penalty directions require only trace products. For very large samples, generalized cross-validation is often the more practical choice; as a rough guideline, "gcv" is recommended once the sample size is above about 250,000.
<code>gcv_gamma</code>	Default: 1.4. Numeric scalar, at least 1, used only when <code>tuning_criterion = "gcv"</code> . Multiplies the effective degrees of freedom in the GCV denominator during automatic penalty tuning. It is accepted but ignored when <code>tuning_criterion = "loo"</code> .
<code>initial_wiggle</code>	Default: <code>c(2e-12, 2e-7, 2e-4, 0.2)</code> . Numeric vector of initial grid points for wiggle penalty optimization, on the raw (non-negative) scale.
<code>initial_flat</code>	Default: <code>c(0.5, 5)</code> . Numeric vector of initial grid points for ridge penalty optimization, on the raw scale (ratio of ridge to wiggle).
<code>wiggle_penalty</code>	Default: $2e-7$. Numeric penalty on the integrated squared second derivative, governing function smoothness.
<code>flat_ridge_penalty</code>	Default: 0.5. Numeric flat ridge penalty for intercepts and linear terms only. Multiplied by <code>wiggle_penalty</code> to obtain total ridge penalty.
<code>unique_penalty_per_partition</code>	Default: TRUE. Logical flag allowing penalty magnitude to differ across partitions.
<code>unique_penalty_per_predictor</code>	Default: TRUE. Logical flag allowing penalty magnitude to differ between predictors.
<code>meta_penalty</code>	Default: $1e-8$. Numeric regularization coefficient for predictor- and partition-specific penalties during tuning. On the raw scale, the implemented meta-penalty shrinks these penalty multipliers toward 1; the wiggle penalty receives only a tiny stabilizing penalty by default.

`predictor_penalties`
Default: NULL. Optional vector of custom penalties per predictor, on the raw (positive) scale.

`partition_penalties`
Default: NULL. Optional vector of custom penalties per partition, on the raw (positive) scale.

`include_quadratic_terms`
Default: TRUE. Logical switch to include squared predictor terms.

`include_cubic_terms`
Default: TRUE. Logical switch to include cubic predictor terms.

`include_quartic_terms`
Default: NULL. Includes quartic terms; when NULL, set to FALSE for single predictor and TRUE otherwise. Highly recommended for multi-predictor models to avoid over-specified constraints.

`include_2way_interactions`
Default: TRUE. Logical switch for linear two-way interactions.

`include_3way_interactions`
Default: TRUE. Logical switch for three-way interactions.

`include_quadratic_interactions`
Default: FALSE. Logical switch for linear-quadratic interaction terms.

`offset`
Default: Empty vector. Column indices/names to include as offsets. Coefficients for offset terms are automatically constrained to 1.

`just_linear_with_interactions`
Default: NULL. Integer or character vector specifying predictors to retain as linear terms while still allowing interactions.

`just_linear_without_interactions`
Default: NULL. Integer or character vector specifying predictors to retain only as linear terms without interactions. Eligible for blockfitting.

`exclude_interactions_for`
Default: NULL. Integer or character vector of predictors to exclude from all interaction terms.

`exclude_these_expansions`
Default: NULL. Character vector of basis expansions to exclude. Named columns of data, or in the form "_1_", "_2_", "_1_x_2_", "_2_^2" etc.

`custom_basis_fxn`
Default: NULL. Optional user-defined function for custom basis expansions. See [get_polynomial_expansions](#).

`include_constrain_fitted`
Default: TRUE. Logical switch to constrain fitted values at knot points.

`include_constrain_first_deriv`
Default: TRUE. Logical switch to constrain first derivatives at knot points.

`include_constrain_second_deriv`
Default: TRUE. Logical switch to constrain second derivatives at knot points.

`include_constrain_interactions`
Default: TRUE. Logical switch to constrain interaction terms at knot points.

<code>add_first_and_second_derivative_constraints</code>	Default: NULL. Logical switch controlling how first- and second-derivative smoothness constraints are assembled. If TRUE, the corresponding first- and second-derivative rows are added before entering the equality constraint matrix. If FALSE, they are kept as separate equality constraints. If NULL, they are combined only when more than one predictor is spline-expanded, so a model with one spline effect and any remaining non-spline effects keeps the derivative constraints separate.
<code>qr_pivot_smoothing_constraints</code>	Default: TRUE. Logical switch to reduce the smoothness/equality constraint matrix to a linearly independent set before fitting. Disabling this keeps the original equality columns.
<code>cl</code>	Default: NULL. Parallel processing cluster object (use <code>parallel::makeCluster()</code>).
<code>chunk_size</code>	Default: NULL. Integer specifying custom chunk size for parallel processing.
<code>parallel_eigen</code>	Default: TRUE. Logical flag for parallel eigenvalue decomposition. Ignored inside tuning fits when <code>parallel_grideval</code> or <code>parallel_bfgs</code> is using the cluster.
<code>parallel_trace</code>	Default: FALSE. Logical flag for parallel trace computation.
<code>parallel_aga</code>	Default: FALSE. Logical flag for parallel G and A matrix operations.
<code>parallel_matmult</code>	Default: FALSE. Logical flag for parallel block-diagonal matrix multiplication.
<code>parallel_qr</code>	Default: FALSE. Logical flag for the tall-skinny least-squares and rank-reduction steps that arise in transformed constraint solves. When active and a cluster is supplied, these steps use row-chunked cross-products with small dense fallback solves instead of relying entirely on base QR; unstable cases fall back automatically to <code>.lm.fit()</code> or <code>qr()</code> .
<code>parallel_bfgs</code>	Default: False. Logical flag for parallel evaluation of damped BFGS step candidates during penalty tuning. When active and a cluster is supplied, multiple damping factors are evaluated across workers and inner parallel flags are ignored for those fits.
<code>parallel_grideval</code>	Default: TRUE. Logical flag for parallel evaluation of the initial tuning grid. When active and a cluster is supplied, grid points are distributed across workers and inner parallel flags are ignored for those fits.
<code>parallel_qr_qp</code>	Default: FALSE. Logical flag for parallel QR pivot reduction of partition-local inequality constraint columns. When active and a cluster is supplied, each partition's reducible QP block is handled independently across workers before the final QP matrix is assembled.
<code>parallel_unconstrained</code>	Default: TRUE. Logical flag for parallel unconstrained MLE for non-identity-link-Gaussian models.
<code>parallel_find_neighbors</code>	Default: TRUE. Logical flag for parallel neighbor identification.
<code>parallel_penalty</code>	Default: FALSE. Logical flag for parallel penalty matrix construction.

<code>parallel_make_constraint</code>	Default: TRUE. Logical flag for parallel constraint matrix generation.
<code>unconstrained_fit_fxn</code>	Default: <code>unconstrained_fit_default</code> . Custom function for fitting unconstrained models per partition.
<code>keep_weighted_Lambda</code>	Default: FALSE. Logical flag to retain GLM weights in penalty constraints using Tikhonov parameterization. Advised for non-canonical GLMs.
<code>iterate_tune</code>	Default: TRUE. Logical switch for iterative optimization during penalty tuning.
<code>iterate_final_fit</code>	Default: TRUE. Logical switch for iterative optimization in final model fitting.
<code>blockfit</code>	Default: TRUE. Logical switch for backfitting with mixed spline and non-interactive linear terms. When the blockfit conditions are met, both tuning and the final fit use <code>blockfit_solve</code> ; otherwise the code uses <code>get_B</code> . Any failure falls back to <code>get_B</code> .
<code>qp_score_function</code>	Default: GLM score using <code>family\$mu.eta(eta) / family\$variance(mu)</code> . Used for quadratic programming, blockfit, and GEE formulations. With dense <code>VhalfInv</code> , the same score weight is applied after whitening. Accepts arguments "X, y, mu, order_list, dispersion, VhalfInv, observation_weights, ...".
<code>qp_observations</code>	Default: NULL. Either a numeric vector of observation indices at which every active built-in QP constraint is evaluated, or a named list keyed by "var: qp_<type>" (or bare "qp_<type>") giving different built-in constraints different observation subsets. The known types are <code>qp_range_lower</code> , <code>qp_range_upper</code> , <code>qp_positive_derivative</code> , <code>qp_negative_derivative</code> , <code>qp_positive_2ndderivative</code> , <code>qp_negative_2ndderivative</code> , <code>qp_monotonic_increase</code> , and <code>qp_monotonic_decrease</code> . For range and monotonicity the canonical keys are the bare forms such as "qp_range_lower" and "qp_monotonic_increase", because those constraints are not tied to a specific variable; prefixed entries are still accepted and unioned. Derivative entries dispatch per variable, so different variables and constraint types may use different subsets. Unknown keys are ignored with a warning when <code>include_warnings = TRUE</code> .
<code>qp_Amat</code>	Default: NULL. Optional pre-built QP constraint matrix. In the current pipeline its presence marks QP handling as active, but the built-in constructor does not merge it into the assembled constraint set; use <code>qp_Amat_fxn</code> for custom assembled constraints.
<code>qp_bvec</code>	Default: NULL. Optional pre-built QP right-hand side paired with <code>qp_Amat</code> . Like <code>qp_Amat</code> , it is currently treated as an advanced placeholder rather than merged into the built-in constructor.
<code>qp_meq</code>	Default: 0. Optional number of equality constraints paired with <code>qp_Amat</code> . Like <code>qp_Amat</code> , it is currently treated as an advanced placeholder rather than merged into the built-in constructor.
<code>qp_positive_derivative</code>	Default: FALSE. Require nonnegative first derivatives. Accepts FALSE (inactive), TRUE (all predictors), or a character / integer vector selecting the predictor variables to constrain.

`qp_negative_derivative`
 Default: FALSE. Require nonpositive first derivatives. Same input types as `qp_positive_derivative`; may be used simultaneously on different predictors.

`qp_positive_2ndderivative`
 Default: FALSE. Require nonnegative second derivatives (convexity). Same input types as `qp_positive_derivative`.

`qp_negative_2ndderivative`
 Default: FALSE. Require nonpositive second derivatives (concavity). Same input types as `qp_positive_derivative`.

`qp_monotonic_increase`
 Default: FALSE. Logical only. Require fitted values to be nondecreasing in observation order.

`qp_monotonic_decrease`
 Default: FALSE. Logical only. Require fitted values to be nonincreasing in observation order.

`qp_range_upper` Default: NULL. Optional upper bound on constrained fitted values.

`qp_range_lower` Default: NULL. Optional lower bound on constrained fitted values.

`qr_pivot_inequality_constraints`
 Default: FALSE. Logical switch to reduce partition-local inequality constraint columns to QR pivot columns before solving. Built-in range and monotonicity constraints are left unchanged, and more generally any inequality columns spanning multiple partitions are also left unchanged.

`qp_Amat_fxn` Default: NULL. Custom function generating `Amat`.

`qp_bvec_fxn` Default: NULL. Custom function generating `bvec`.

`qp_meq_fxn` Default: NULL. Custom function generating `meq`.

`constraint_values`
 Default: `cbind()`. Optional matrix encoding nonzero equality targets paired with `constraint_vectors`. When left empty, added equality constraints are treated as homogeneous.

`constraint_vectors`
 Default: `cbind()`. Optional matrix of user-supplied equality-constraint vectors, appended to the internally generated smoothness constraints.

`return_G` Default: TRUE. Logical switch to return the unscaled unconstrained variance-covariance matrix \mathbf{G} .

`return_Ghalf` Default: TRUE. Logical switch to return $\mathbf{G}^{1/2}$.

`return_U` Default: TRUE. Logical switch to return the constraint projection matrix \mathbf{U} .

`estimate_dispersion`
 Default: TRUE. Logical flag to estimate dispersion after fitting.

`unbias_dispersion`
 Default: NULL. Logical switch to multiply dispersion by $N/(N - \text{trace}(\mathbf{H}))$. When NULL, set to TRUE for Gaussian identity link and FALSE otherwise.

`return_varcovmat`
 Default: TRUE. Logical switch to return the variance-covariance matrix of estimated coefficients. Needed for Wald inference.

<code>null_constraint</code>	Default: NULL. Alternative parameterization for a nonzero equality target when <code>constraint_vectors</code> is supplied and <code>constraint_values</code> is left empty.
<code>critical_value</code>	Default: <code>qnorm(1-0.05/2)</code> . Numeric critical value for Wald confidence intervals.
<code>data</code>	Default: NULL. Optional data frame for formula-based model specification.
<code>weights</code>	Default: NULL. Alias for <code>observation_weights</code> .
<code>no_intercept</code>	Default: FALSE. Logical flag to constrain intercept to 0. Formulas with " <code>0+</code> " set this to TRUE automatically.
<code>correlation_id, spacetime</code>	Default: NULL. N-length vector and N-row matrix of cluster ids and longitudinal/spatial variables, respectively.
<code>correlation_structure</code>	Default: NULL. Native implementations: " <code>exchangeable</code> ", " <code>spatial-exponential</code> ", " <code>squared-exponential</code> ", " <code>ar(1)</code> ", " <code>spherical</code> ", " <code>gaussian-cosine</code> ", " <code>gamma-cosine</code> ", " <code>matern</code> ", and aliases. Estimated via REML.
<code>VhalfInv</code>	Default: NULL. Fixed custom $N \times N$ square-root-inverse covariance matrix $\mathbf{V}^{-1/2}$. Triggers GLS with known covariance. Post-fit inference recomputed from whitened Gram matrices.
<code>Vhalf</code>	Default: NULL. Fixed custom $N \times N$ square-root covariance $\mathbf{V}^{1/2}$. Computed as inverse of <code>VhalfInv</code> if not supplied.
<code>VhalfInv_fxn</code>	Default: NULL. Parametric function for $\mathbf{V}^{-1/2}$; takes single numeric vector " <code>par</code> ", returns $N \times N$ matrix. If <code>VhalfInv</code> is not supplied, correlation parameters are optimized from this function; when <code>VhalfInv_par_init</code> is omitted, the optimizer starts at <code>1e-2</code> . Helper functions that also use <code>correlation_id</code> , <code>spacetime</code> , or matching extra arguments are supported.
<code>Vhalf_fxn</code>	Default: NULL. Optional function for efficient computation of $\mathbf{V}^{1/2}$ from the same parameter vector used by <code>VhalfInv_fxn</code> . When omitted, <code>Vhalf</code> is obtained by explicit matrix inversion of <code>VhalfInv</code> .
<code>VhalfInv_par_init</code>	Default: <code>c()</code> . Initial parameter values for <code>VhalfInv_fxn</code> optimization, on unbounded transformed scale. If left empty while <code>VhalfInv_fxn</code> is supplied and <code>VhalfInv</code> is NULL, it is set internally to <code>1e-2</code> .
<code>REML_grad</code>	Default: NULL. Function for the gradient of the negative REML (or custom loss) with respect to the parameters of <code>VhalfInv_fxn</code> . Takes " <code>par</code> ", " <code>model_fit</code> ", and " <code>...</code> ".
<code>custom_VhalfInv_loss</code>	Default: NULL. Alternative to negative REML for the correlation parameter objective function. Takes " <code>par</code> ", " <code>model_fit</code> ", and " <code>...</code> ".
<code>VhalfInv_logdet</code>	Default: NULL. Function for efficient $\log \mathbf{V}^{-1/2} $ computation. Takes same " <code>par</code> " as <code>VhalfInv_fxn</code> .
<code>include_warnings</code>	Default: TRUE. Logical switch to control display of warnings.

penalty_args	Default: NULL. Optional named list grouping penalty-related arguments. See section "Grouped Argument Lists".
tuning_args	Default: NULL. Optional named list grouping tuning-related arguments.
expansion_args	Default: NULL. Optional named list grouping basis expansion arguments.
constraint_args	Default: NULL. Optional named list grouping constraint arguments.
qp_args	Default: NULL. Optional named list grouping quadratic programming arguments.
parallel_args	Default: NULL. Optional named list grouping parallel processing arguments.
covariance_args	Default: NULL. Optional named list grouping correlation structure arguments.
return_args	Default: NULL. Optional named list grouping return-control arguments.
glm_args	Default: NULL. Optional named list grouping GLM customization arguments.
...	Additional arguments passed to the unconstrained model fitting function.

Details

Main features include:

- Multiple predictors and interaction terms
- Various GLM families and link functions
- Correlation structures for longitudinal/clustered data
- Shape constraints via quadratic programming
- Parallel computation for large datasets
- Comprehensive inference tools

Value

A list of class "lgspline" containing model components:

y Original response vector.

ytilde Fitted/predicted values on the scale of the response.

X List of design matrices \mathbf{X}_k for each partition k , containing basis expansions including intercept, linear, quadratic, cubic, and interaction terms as specified. Returned on the unstandardized scale.

A Constraint matrix **A** encoding smoothness constraints at knot points and any user-specified linear constraints. When `qr_pivot_smoothing_constraints = TRUE`, only a linearly independent subset of columns is retained via pivoted QR decomposition; otherwise the original equality columns are kept.

B List of fitted coefficients β_k for each partition k on the original, unstandardized scale of the predictors and response.

B_raw List of fitted coefficients for each partition on the predictor-and-response standardized scale.

K Number of interior knots with one predictor (number of partitions minus 1 with > 1 predictor).

p Number of basis expansions of predictors per partition.

q Number of predictor variables.

P Total number of coefficients ($p \times (K + 1)$).

N Number of observations.

penalties List containing optimized penalty matrices and components:

- **Lambda**: Baseline per-partition penalty matrix corresponding to $\lambda_w(\mathbf{\Lambda}_s + \lambda_r \mathbf{\Lambda}_r + \sum_l \lambda_{l,k} \mathbf{\Lambda}_{l,k})$ before any partition-specific block is added.
- **L1**: Implementation label for the curvature penalty matrix $\mathbf{\Lambda}_s$.
- **L2**: Implementation label for the ridge penalty matrix $\mathbf{\Lambda}_r$.
- **L_predictor_list**: Implementation lists for additional penalty matrices $\mathbf{\Lambda}_{l,k}$ associated with predictor-specific tuning directions.
- **L_partition_list**: Implementation lists for additional penalty matrices $\mathbf{\Lambda}_{l,k}$ associated with partition-specific tuning directions.

knot_scale_transf Function for transforming predictors to standardized scale used for knot placement.

knot_scale_inv_transf Function for transforming standardized predictors back to original scale.

knots Matrix of knot locations on original unstandardized predictor scale for one predictor.

partition_codes Vector assigning observations to partitions.

partition_bounds Vector or matrix specifying the boundaries between partitions.

knot_expand_function Internal function for expanding data according to partition structure.

predict Function for generating predictions on new data. For multi-predictor models, `take_first_derivatives = TRUE`, `take_second_derivatives` returns derivatives as a named list of components per predictor variable, rather than a concatenated vector. When `new_predictors` contains columns not present in the data, extraneous columns are silently dropped before prediction.

assign_partition Function for assigning new observations to partitions.

family GLM family object specifying the error distribution and link function.

estimate_dispersion Logical indicating whether dispersion parameter was estimated.

unbias_dispersion Logical indicating whether dispersion estimates should be unbiased.

backtransform_coefficients Function for converting standardized coefficients to original scale.

forwtransform_coefficients Function for converting coefficients to standardized scale.

mean_y, sd_y Mean and standard deviation of response if standardized.

og_order Original ordering of observations before partitioning.

order_list List containing observation indices for each partition.

constraint_values, constraint_vectors Matrices specifying linear equality constraints if provided.

make_partition_list List containing partition information for > 1-D cases.

expansion_scales Vector of scaling factors used for standardizing basis expansions.

take_derivative, take_interaction_2ndderivative Functions for computing derivatives of basis expansions.

get_all_derivatives_insample Function for computing all derivatives on training data.

numerics Indices of numeric predictors used in basis expansions.

- power1_cols, power2_cols, power3_cols, power4_cols** Column indices for linear through quartic terms.
- quad_cols** Column indices for all quadratic terms (including interactions).
- interaction_single_cols, interaction_quad_cols** Column indices for linear-linear and linear-quadratic interactions.
- triplet_cols** Column indices for three-way interactions.
- nonspline_cols** Column indices for terms excluded from spline expansion.
- return_varcovmat** Logical indicating whether variance-covariance matrix was computed.
- raw_expansion_names** Names of basis expansion terms.
- std_X, unstd_X** Functions for standardizing/unstandardizing design matrices.
- parallel_cluster_supplied** Logical indicating whether a parallel cluster was supplied.
- weights** Original observation weights on the data scale. When no weights were supplied, this is a vector of ones.
- G** List of unscaled partition-wise information inverses \mathbf{G}_k if `return_G = TRUE`. These are the blockwise quantities stored on the fitting scale; correlation-aware trace, posterior, and variance calculations additionally use dense GLS analogues internally when needed.
- Ghalf** List of $\mathbf{G}_k^{1/2}$ matrices if `return_Ghalf = TRUE`. As with **G**, dense GLS square-root factors may also be constructed internally for correlation-aware post-fit calculations.
- U** Constraint projection matrix \mathbf{U} if `return_U = TRUE`. For $K=0$ and no constraints, returns identity. Otherwise, returns $\mathbf{U} = \mathbf{I} - \mathbf{G}\mathbf{A}(\mathbf{A}^\top\mathbf{G}\mathbf{A})^{-1}\mathbf{A}^\top$. Used for computing the variance-covariance matrix $\sigma^2\mathbf{U}\mathbf{G}$.
- sigmasq_tilde** Estimated (or fixed) dispersion parameter $\tilde{\sigma}^2$. For Gaussian identity fits without correlation, this is the weighted mean squared residual with optional bias correction. When `VhalfInv` is non-NULL, Gaussian-identity residuals are whitened before this calculation.
- trace_XUGX** Effective degrees of freedom ($\text{trace}(\mathbf{X}\mathbf{U}\mathbf{G}\mathbf{X}^\top)$), where $\mathbf{X}\mathbf{U}\mathbf{G}\mathbf{X}^\top$ serves as the "hat" matrix. When `VhalfInv` is non-NULL, computed as $\|\mathbf{V}^{-1/2}\mathbf{X}\mathbf{U}\mathbf{G}_{\text{correct}}^{1/2}\|_F^2$ using the full penalized GLS information.
- varcovmat** Variance-covariance matrix of coefficient estimates if `return_varcovmat = TRUE`. Computed as $\sigma^2(\mathbf{U}\mathbf{G}^{1/2})(\mathbf{U}\mathbf{G}^{1/2})^\top$ for numerical stability. When `VhalfInv` is non-NULL, uses the full $\mathbf{G}_{\text{correct}}^{1/2}$ in place of the block-diagonal $\mathbf{G}^{1/2}$.
- lagrange_multipliers** Vector of Lagrangian multipliers if `return_lagrange_multipliers = TRUE`. For equality-only fits these correspond to the active columns of \mathbf{A} ; when quadratic-programming constraints are active they are taken directly from `solve.QP` and therefore refer to the combined equality/inequality constraint system. NULL if no constraints are active (\mathbf{A} is NULL or $K == \emptyset$).
- VhalfInv** The $\mathbf{V}^{-1/2}$ matrix used for implementing correlation structures, if specified.
- VhalfInv_fxn, Vhalf_fxn, VhalfInv_logdet, REML_grad** Functions for generating $\mathbf{V}^{-1/2}$, $\mathbf{V}^{1/2}$, $\log|\mathbf{V}^{-1/2}|$, and gradient of REML if provided.
- VhalfInv_params_estimates** Vector of estimated correlation parameters when using `VhalfInv_fxn`.
- VhalfInv_params_vcov** Approximate variance-covariance matrix of estimated correlation parameters from BFGS optimization.

wald_univariate Function for computing univariate Wald statistics and confidence intervals. Returns an S3 object of class "wald_lgspline" with dedicated print, summary, plot, coef, and confint methods. The print method uses printCoefmat() for standard R coefficient table formatting with significance stars.

critical_value Critical value used for confidence interval construction.

generate_posterior Function for drawing from the posterior distribution of coefficients. When VhalfInv is non-NULL, draws are from the correct joint posterior $\mathbf{UG}_{\text{correct}}^{1/2} \mathbf{z}$ using the full penalized GLS information, reflecting cross-partition posterior covariance induced by off-diagonal blocks of $\mathbf{V}^{-1/2}$.

find_extremum Function for optimizing the fitted function. Accepts both numeric column indices and character column names for vars. When select_vars_fl = TRUE, L-BFGS-B bounds are correctly subsetted to the optimized variables.

plot Function for visualizing fitted curves.

qp_info Metadata for inequality-constrained solves, or NULL. Includes active constraints, multipliers, and a method string describing the solver path: "active_set" = partition-wise active-set on the standard block-diagonal path; "active_set_full" = active-set with full-system equality re-solves; "active_set_woodbury" = active-set with Woodbury equality re-solves on the correlated low-rank path; "dense_qp_gee_gaussian" = dense QP fallback for correlated Gaussian fits; "dense_qp_gee_glm" = dense SQP / dense QP-subproblem fallback for correlated non-Gaussian fits.

quadprog_list List containing the assembled QP objects qp_Amat, qp_bvec, qp_meq, and when applicable a copy of qp_info.

.fit_call_args List containing the arguments passed to lgspline.

The returned object has class "lgspline" and provides comprehensive tools for model interpretation, inference, prediction, and visualization. All coefficients and predictions can be transformed between standardized and original scales using the provided transformation functions. The object includes both frequentist and Bayesian inference capabilities through Wald statistics and posterior sampling. S3 methods logLik.lgspline and confint.lgspline are available for standard log-likelihood extraction and confidence interval computation, respectively. Advanced customization options are available for analyzing arbitrarily complex study designs.

Response and Predictor Setup

These arguments control the primary data inputs and the initial standardization steps applied before knot placement and fitting.

GLM Customization

These options let you override the default GLM working-weight, dispersion, and partition-wise unconstrained fitting behavior.

Knots and Partitioning

These arguments determine how the predictor space is partitioned and how knot locations are chosen or reused.

Penalty

These arguments control the per-partition penalty $\Lambda_k = \lambda_w(\Lambda_s + \lambda_r\Lambda_r + \sum_{l=1}^L \lambda_{l,k}\Lambda_{l,k})$, and hence the full block-diagonal penalty $\Lambda = \text{blockdiag}(\Lambda_0, \dots, \Lambda_K)$. By default, tuning uses exact leave-one-out on the transformed problem; modified GCV is also available.

Basis Expansions

These arguments control which polynomial and interaction terms are included in the partition-specific design matrices.

Constraints

These arguments govern the smoothness equalities at partition boundaries and any additional user-supplied linear equality constraints. Shape restrictions such as monotonicity, convexity, or range bounds are handled separately in the quadratic-programming stage below.

Quadratic Programming

These arguments activate built-in or custom linear inequality constraints on fitted values or derivatives. Internally they are assembled in the form $\mathbf{C}^\top \boldsymbol{\beta} \geq \mathbf{c}$ and passed to the constrained solver only when requested.

Parallel Processing

These arguments control which computational subroutines may run in parallel and how work is chunked across cluster workers.

Tuning Control

These options control iterative updates during penalty tuning and the final constrained fit.

Return Control

These arguments determine which intermediate matrices and inferential quantities are retained in the returned fit object.

Correlation Structures

These arguments enable built-in or custom working-correlation structures for longitudinal, clustered, or spatially indexed responses. In the notation of [Details](#), the correlated penalized information is written as $\mathbf{G}^{-1} = \mathbf{G}_{\text{on}}^{-1} + \mathbf{G}_{\text{off}}^{-1}$. When the cross-partition part is low rank, the Woodbury path factors $\mathbf{G}_{\text{off}}^{-1} = \mathbf{E}\mathbf{J}\mathbf{E}^\top$, define $\mathbf{N} = \mathbf{G}_{\text{on}}^{1/2}\mathbf{E}$, and works through $\mathbf{G}^{1/2} = \mathbf{G}_{\text{on}}^{1/2}\mathbf{F}^{1/2}$ without changing the final estimator. If that low-rank path is unavailable or fails its numerical checks, the code falls back to the dense correlated solve.

Grouped Argument Lists

For convenience, related arguments can be bundled into named lists. When a grouped argument is non-NULL, its entries overwrite the corresponding individual arguments. Individual arguments remain available for backward compatibility.

`penalty_args` Groups: `wiggle_penalty`, `flat_ridge_penalty`, `unique_penalty_per_partition`, `unique_penalty_per_predictor`, `meta_penalty`, `predictor_penalties`, `partition_penalties`, `custom_penalty_mat`, `previously_tuned_penalties`, `smoothing_spline_penalty`.

`tuning_args` Groups: `opt`, `use_custom_bfgs`, `delta`, `tol`, `tuning_criterion`, `gcv_gamma`, `initial_wiggle`, `initial_flat`, `iterate_tune`, `iterate_final_fit`, `blockfit`.

`expansion_args` Groups: `include_quadratic_terms`, `include_cubic_terms`, `include_quartic_terms`, `include_2way_interactions`, `include_3way_interactions`, `include_quadratic_interactions`, `just_linear_with_interactions`, `just_linear_without_interactions`, `exclude_interactions_for`, `exclude_these_expansions`, `custom_basis_fxn`, `offset`.

`constraint_args` Groups: `include_constrain_fitted`, `include_constrain_first_deriv`, `include_constrain_second_deriv`, `include_constrain_interactions`, `add_first_and_second_derivative_constraints`, `qr_pivot_smoothing_constraints`, `constraint_values`, `constraint_vectors`, `null_constraint`, `no_intercept`.

`qp_args` Groups: `qp_score_function`, `qp_observations`, `qp_Amat`, `qp_bvec`, `qp_meq`, `qp_positive_derivative`, `qp_negative_derivative`, `qp_positive_2ndderivative`, `qp_negative_2ndderivative`, `qp_monotonic_increase`, `qp_monotonic_decrease`, `qp_range_upper`, `qp_range_lower`, `qr_pivot_inequality_constraints`, `qp_Amat_fxn`, `qp_bvec_fxn`, `qp_meq_fxn`.

`parallel_args` Groups: `cl`, `chunk_size`, `parallel_eigen`, `parallel_trace`, `parallel_aga`, `parallel_matmult`, `parallel_qr`, `parallel_bfgs`, `parallel_grideval`, `parallel_qr_qp`, `parallel_unconstrained`, `parallel_find_neighbors`, `parallel_penalty`, `parallel_make_constraint`.

`covariance_args` Groups: `correlation_id`, `spacetime`, `correlation_structure`, `VhalfInv`, `Vhalf`, `VhalfInv_fxn`, `Vhalf_fxn`, `VhalfInv_par_init`, `REML_grad`, `custom_VhalfInv_loss`, `VhalfInv_logdet`.

`return_args` Groups: `return_G`, `return_Ghalf`, `return_U`, `estimate_dispersion`, `unbias_dispersion`, `return_varcovmat`, `exact_varcovmat`, `return_lagrange_multipliers`.

`glm_args` Groups: `glm_weight_function`, `schur_correction_function`, `need_dispersion_for_estimation`, `dispersion_function`, `unconstrained_fit_fxn`, `keep_weighted_Lambda`.

Miscellaneous

These remaining arguments affect inference defaults, numerical safeguards, verbosity, and developer-oriented diagnostics.

See Also

- [lgspline.fit](#) for the low-level fitting interface
- [logLik.lgspline](#) for log-likelihood extraction
- [confint.lgspline](#) for confidence interval extraction
- [leave_one_out](#) for leave-one-out cross-validated predictions
- [blockfit_solve](#) for the standalone backfitting solver


```

## Composite function
fxn <- function(x)(slinky(t) +
                    coil(t) +
                    exponential_log(t) +
                    scaled_abs_gamma(t))

## Bind together with random noise
dat <- cbind(t, fxn(t) + rnorm(length(t), 0, 50))
colnames(dat) <- c('t', 'y')
t <- dat[, 't']
y <- dat[, 'y']

## Fit Model, 4 equivalent ways are shown below
model_fit <- lgspline(t, y, opt = FALSE)
model_fit <- lgspline(y ~ spl(t), as.data.frame(dat), opt = FALSE)
model_fit <- lgspline(response = y, predictors = t, opt = FALSE)
model_fit <- lgspline(data = as.data.frame(dat), formula = y ~ ., opt = FALSE)

# This is not valid: lgspline(y ~ ., t)
# This is not valid: lgspline(y, data = as.data.frame(dat))
# Do not put operations in formulas, not valid: lgspline(y ~ log(t) + spl(t))

## Basic Functionality
predict(model_fit, new_predictors = rnorm(1)) # make prediction on new data
loo_vals <- suppressWarnings(head(leave_one_out(model_fit)))
loo_vals # may contain NA when leverage is too high
coef(model_fit) # extract coefficients
summary(model_fit) # model information and Wald inference
generate_posterior(model_fit) # generate draws of parameters from posterior distribution
find_extremum(model_fit, minimize = TRUE) # find the minimum of the fitted function

## Incorporate range constraints, custom knots, keep penalization identical
# across partitions and predictors
model_fit <- lgspline(y ~ spl(t),
                     unique_penalty_per_partition = FALSE,
                     unique_penalty_per_predictor = FALSE,
                     custom_knots = cbind(c(-2, -1, 0, 1, 2)),
                     data = data.frame(t = t, y = y),
                     qp_range_lower = -150,
                     qp_range_upper = 150,
                     qp_observations = sample(1:length(t), 50),
                     opt = FALSE)

## Plotting the constraints and knots
plot(model_fit,
      custom_title = 'Fitted Function Constrained to Lie Between (-150, 150)',
      cex.main = 0.75)
# knot locations
abline(v = model_fit$knots)
# lower bound from quadratic program
abline(h = -150, lty = 2)
# upper bound from quadratic program
abline(h = 150, lty = 2)

```



```

    l1_bound) {

  ## Combine matrices
  l1_bvec <- rep(-l1_bound, ncol(qp_Amat)) * c(1, scales)

  return(l1_bvec)
}

## Fit model, using predictor-response formulation, assuming
# Gamma-distributed response and custom quadratic-programming constraints
# as well as quartic terms, keeping the effect of height constant across
# partitions, and 3 partitions in total. The default GLM score and weights
# handle the non-canonical log link.
# You can modify this code for performing l1-regularization in general.
model_fit <- lgspline(
  Volume ~ spl(Girth) + Height*Girth,
  data = with(trees, cbind(Girth, Height, Volume)),
  family = Gamma(link = 'log'),
  keep_weighted_Lambda = TRUE,
  unbiased_dispersion = TRUE, # multiply dispersion by N/(N-trace(XUGX^T))
  K = 2, # 3 partitions
  opt = FALSE, # keep penalties fixed
  unique_penalty_per_partition = FALSE,
  unique_penalty_per_predictor = FALSE,
  flat_ridge_penalty = 1e-64,
  wiggle_penalty = 1e-64,
  qp_Amat_fxn = function(N, p, K, X, colnm, scales, deriv_fxn, ...) {
    l1_constraint_matrix(p, K)
  },
  qp_bvec_fxn = function(qp_Amat, N, p, K, X, colnm, scales, deriv_fxn, ...) {
    l1_bound_vector(qp_Amat, scales, 25)
  },
  qp_meq_fxn = function(qp_Amat, N, p, K, X, colnm, scales, deriv_fxn, ...) 0
)

## Notice, interaction effect is constant across partitions as is the effect
# of Height alone
Reduce('cbind', coef(model_fit))

## Many constraints, many coefficients, and small sample size makes inference
# using asymptotic variance-covariance matrix untrustworthy.
# Confidence intervals are often too wide or narrow, even for "good" fit.
# Consider bootstrapping or alternative.
print(summary(model_fit))

## Plot results
plot(model_fit, custom_predictor_lab1 = 'Girth',
      custom_predictor_lab2 = 'Height',
      custom_response_lab = 'Volume',
      custom_title = 'Girth and Height Predicting Volume of Trees',
      show_formulas = TRUE)

## Verify magnitude of unstandardized coefficients does not exceed bound (25)

```

```

print(max(abs(unlist(model_fit$B))))

## Find height and girth where tree volume is closest to 42
# Uses custom objective that minimizes MSE discrepancy between predicted
# value and 42.
# The vanilla find_extremum function can be thought of as
# using "function(mu)mu" aka the identity function as the
# objective, where mu = "f(t)", our estimated function. The derivative is then
# d_mu = "df/dt" with respect to predictors t.
# But with more creative objectives, and since we have machinery for
# df/dt already available, we can compute gradients for (and optimize)
# arbitrary differentiable functions of our predictors too.
# For any objective, differentiate w.r.t. to mu, then multiply by d_mu to
# satisfy chain rule.
# Here, we have objective function: 0.5*(mu-42)^2
# and gradient : (mu-42)*d_mu
# and L-BFGS-B will be used to find the height and girth that most closely
# yields a prediction of 42 within the bounds of the observed data.
# The d_mu also takes into account link function transforms automatically
# for most common link functions, and will return warning + instructions
# on how to program the link-function derivatives otherwise.

## Custom acquisition functions for Bayesian optimization could be coded here.
find_extremum(
  model_fit,
  minimize = TRUE,
  custom_objective_function = function(mu, sigma, ybest, ...){
    0.5*(mu - 42)^2
  },
  custom_objective_derivative = function(mu, sigma, ybest, d_mu, ...){
    (mu - 42) * d_mu
  }
)

## ## ## ## How to Use Formulas in lgspline ## ## ## ## ## ## ## ## ## ## ## ## ## ## ## ##
## Demonstrates splines with multiple mixed predictors and interactions

## Generate data
n <- 2500
t <- rnorm(n)
u <- rnorm(n)
z <- sin(t)*mean(abs(u))/2

## Categorical predictors
cat1 <- rbinom(n, 1, 0.5)
cat2 <- rbinom(n, 1, 0.5)
cat3 <- rbinom(n, 1, 0.5)

## Response with mix of effects
response <- u + z + 0.1*(2*cat1 - 1)

## Continuous predictors re-named
continuous1 <- t

```

```

continuous2 <- z

## Combine data
dat <- data.frame(
  response = response,
  continuous1 = continuous1,
  continuous2 = continuous2,
  cat1 = cat1,
  cat2 = cat2,
  cat3 = cat3
)

## Example 1: Basic Model with Default Terms, No Intercept
# standardize_response = FALSE often needed when constraining intercepts to 0
fit1 <- lgspline(
  formula = response ~ 0 + spl(continuous1, continuous2) +
    cat1*cat2*continuous1 + cat3,
  K = 2,
  standardize_response = FALSE,
  data = dat
)
## Examine coefficients included
rownames(fit1$B$partition1)
## Verify intercept term is near 0 up to some numeric tolerance
abs(fit1$B[[1]][1]) < 1e-8

## Example 2: Similar Model with Intercept, Other Terms Excluded
fit2 <- lgspline(
  formula = response ~ spl(continuous1, continuous2) +
    cat1*cat2*continuous1 + cat3,
  K = 1,
  standardize_response = FALSE,
  include_cubic_terms = FALSE,
  exclude_these_expansions = c( # Not all need to actually be present
    '_batman_x_robin_',
    '_3_x_4_', # no cat1 x cat2 interaction, coded using column indices
    'continuous1xcontinuous2', # no continuous1 x continuous2 interaction
    'thejoker'
  ),
  data = dat
)
## Examine coefficients included
rownames(Reduce('cbind',coef(fit2)))
# Intercept will probably be present and non-0 now
abs(fit2$B[[1]][1]) < 1e-8

## ## ## ## Compare Inference to survreg for Weibull AFT Model Validation ##
# Only linear predictors, no knots, no penalties, using Weibull AFT Model
# The goal here is to ensure that for the special case of no spline effects
# and no knots, this implementation will be consistent with other model
# implementations.
# Also note, that when using models (like Weibull AFT) where dispersion is
# being estimated and is required for estimating beta coefficients,

```



```

      lapply(1:n_blocks,
            function(i){
              sigma <- diag(block_size) + rho_true *
                (matrix(1, block_size, block_size) -
                 diag(block_size))
              matsqrt(sigma) %*% rnorm(block_size)
            })

## Generate response with correlated errors
y <- true_mean + errors * 0.5

## Fit model with correlation structure
# include_warnings = FALSE is a good idea here, since many proposed
# correlations will not work
model_fit <- lgspline(t,
                     y,
                     K = 4,
                     correlation_id = rep(1:n_blocks, each = block_size),
                     correlation_structure = 'exchangeable',
                     include_warnings = FALSE
                    )

## Assess overall fit
plot(t, y, main = 'Sinusoidal Fit Under Correlation Structure')
plot(model_fit, add = TRUE, show_formulas = TRUE, custom_predictor_lab = 't')

## Compare estimated vs true correlation
# Built-in exchangeable uses rho = exp(-exp(par)), so par in (-Inf, Inf)
# maps to rho in (0, 1). Only positive correlation is supported.
rho_est <- exp(-exp(model_fit$VhalfInv_params_estimates))
print(c("True correlation:" = rho_true,
       "Estimated correlation:" = rho_est))

## Also check SD (should be close to 0.5)
print(sqrt(model_fit$sigma_sq_tilde))

## Toeplitz Simulation Setup, with demonstration of custom functions
# and boilerplate. Toep is not implemented by default, because it makes
# strong assumptions on the study design and missingness that are rarely met,
# with non-obvious workarounds.
# If a GLM was to-be-fit, you would also submit a function "Vhalf_fxn" analogous
# to VhalfInv_fxn with same argument (par) and an output of an N x N matrix
# that yields the inverse of VhalfInv_fxn output.
n_blocks <- 250 # Number of correlation_ids
block_size <- 8 # Observations per correlation_id
N <- n_blocks * block_size # total sample size
sigma_true <- 2 # Marginal standard deviation

## True Toeplitz components
# This example uses a convex combination of two geometric lag kernels:
# corr(h) = mix * rho_fast^h + (1 - mix) * rho_slow^h
# which is Toeplitz and positive definite for mix in (0, 1) and
# rho_fast, rho_slow in (0, 1).

```

```

rho_fast_true <- 0.25
rho_slow_true <- 0.75
mix_true <- 0.40

## Create time and correlation_id variables
time_var <- rep(1:block_size, n_blocks)
correlation_id_var <- rep(1:n_blocks, each = block_size)

## Create nonlinear predictor-response relationship
# Not sinusoidal and not polynomial.
t_base <- seq(-2, 2, length.out = block_size)
t <- rep(t_base, n_blocks) + rnorm(N, sd = 0.10)
f_true <- function(t) {
  1.4 + 0.9 * atan(1.8 * t) + 0.8 * exp(-1.2 * (t - 0.4)^2)
}

## Generate mean structure
mu_true <- f_true(t)

## Toeplitz correlation helper
corr_from_components <- function(rho_fast, rho_slow, mix) {
  corr <- matrix(0, block_size, block_size)
  for(i in 1:block_size) {
    for(j in 1:block_size) {
      lag <- abs(i - j)
      if(lag == 0) {
        corr[i, j] <- 1
      } else {
        corr[i, j] <- mix * rho_fast^lag + (1 - mix) * rho_slow^lag
      }
    }
  }
  corr
}

## Toeplitz correlation function
# Custom functions can use any parameterization. Here we map:
# par[1] -> rho_fast = exp(-exp(par[1]))
# par[2] -> rho_slow = exp(-exp(par[2]))
# par[3] -> mix = plogis(par[3])
# so the parameter space is unconstrained, while the resulting Toeplitz
# correlation matrix remains valid.
corr_from_par <- function(par) {
  rho_fast <- exp(-exp(par[1]))
  rho_slow <- exp(-exp(par[2]))
  mix <- plogis(par[3])
  corr_from_components(rho_fast, rho_slow, mix)
}

## Create block Toeplitz errors from the same family we will fit
corr_true <- corr_from_components(rho_fast_true, rho_slow_true, mix_true)
errors <- Reduce('c',
  lapply(1:n_blocks, function(i) {

```

```

        c(matsqrt(corr_true) %**% rnorm(block_size))
      )))

## Generate response with correlated errors and nonlinear covariate effect
y <- mu_true + sigma_true * errors

VhalfInv_fxn <- function(par) {
  corr <- corr_from_par(par)
  kronecker(diag(n_blocks), matinvsqrt(corr))
}

Vhalf_fxn <- function(par) {
  corr <- corr_from_par(par)
  kronecker(diag(n_blocks), matsqrt(corr))
}

## Determinant function (for efficiency)
# This avoids taking determinant of N by N matrix
VhalfInv_logdet <- function(par) {
  corr <- corr_from_par(par)
  log_det_invsqrt_corr <- -0.5 * determinant(corr, logarithm = TRUE)$modulus[1]
  n_blocks * log_det_invsqrt_corr
}

## GLM weights for REML gradient helper
# For Gaussian identity, these are all 1.
glm_weight_function <- function(mu, y, order_indices, family,
                                dispersion, observation_weights, ...) {
  rep(1, length(mu))
}

## REML gradient function
# The helper reml_grad_from_dV computes the three REML terms once dV / dpar
# is supplied. For this parameterization, dV / dpar has closed form.
REML_grad <- function(par, model_fit, ...) {
  rho_fast <- exp(-exp(par[1]))
  rho_slow <- exp(-exp(par[2]))
  mix <- plogis(par[3])

  dV1_block <- matrix(0, block_size, block_size)
  dV2_block <- matrix(0, block_size, block_size)
  dV3_block <- matrix(0, block_size, block_size)

  for(i in 1:block_size) {
    for(j in 1:block_size) {
      lag <- abs(i - j)
      if(lag > 0) {
        ## d/dpar[1] through rho_fast = exp(-exp(par[1]))
        dV1_block[i, j] <- -mix * lag * exp(par[1]) * rho_fast^lag
        ## d/dpar[2] through rho_slow = exp(-exp(par[2]))
        dV2_block[i, j] <- -(1 - mix) * lag * exp(par[2]) * rho_slow^lag
        ## d/dpar[3] through mix = plogis(par[3])
        dV3_block[i, j] <- mix * (1 - mix) * (rho_fast^lag - rho_slow^lag)
      }
    }
  }
}

```

```

    }
  }
}

dV1 <- kronecker(diag(n_blocks), dV1_block)
dV2 <- kronecker(diag(n_blocks), dV2_block)
dV3 <- kronecker(diag(n_blocks), dV3_block)

gradient <- numeric(3)
gradient[1] <- lgspline::reml_grad_from_dV(dV1, model_fit,
                                          glm_weight_function, ...)
gradient[2] <- reml_grad_from_dV(dV2, model_fit,
                                  glm_weight_function, ...)
gradient[3] <- reml_grad_from_dV(dV3, model_fit,
                                  glm_weight_function, ...)
gradient
}

## Visualization
plot(t, y, col = correlation_id_var,
     main = 'Simulated Data with Toeplitz Correlation')

## Fit model with custom Toeplitz
model_fit <- lgspline(
  response = y,
  predictors = t,
  K = 4,
  standardize_response = FALSE,
  VhalfInv_fxn = VhalfInv_fxn,
  Vhalf_fxn = Vhalf_fxn,
  VhalfInv_logdet = VhalfInv_logdet,
  REML_grad = REML_grad,
  VhalfInv_par_init = c(0, -1, 0),
  include_warnings = FALSE
)

## Print comparison of true and estimated correlations
lag_values <- 1:(block_size - 1)
corr_true_by_lag <- sapply(lag_values, function(h) {
  mix_true * rho_fast_true^h + (1 - mix_true) * rho_slow_true^h
})
rho_fast_est <- exp(-exp(model_fit$VhalfInv_params_estimates[1]))
rho_slow_est <- exp(-exp(model_fit$VhalfInv_params_estimates[2]))
mix_est <- plogis(model_fit$VhalfInv_params_estimates[3])
corr_est_by_lag <- sapply(lag_values, function(h) {
  mix_est * rho_fast_est^h + (1 - mix_est) * rho_slow_est^h
})
cat('Toeplitz Correlation Estimates by Lag:\n')
print(data.frame(
  Lag = lag_values,
  True.Correlation = round(corr_true_by_lag, 4),
  Estimated.Correlation = round(corr_est_by_lag, 4)
))

```



```

c <- rnorm(500000)
d <- rpois(500000, 1)
y <- sin(a) + cos(b) - 0.2*sqrt(a^2 + b^2) +
  abs(a) + b +
  0.5*(a^2 + b^2) +
  (1/6)*(a^3 + b^3) +
  a*b*c -
  c +
  d +
  rnorm(500000, 0, 5)

## Set up cores
cl <- parallel::makeCluster(1)
on.exit(try(parallel::stopCluster(cl), silent = TRUE), add = TRUE)

## This example shows some options for what operations can be parallelized
# By default, only parallel_eigen and parallel_unconstrained are TRUE
# parallel_unconstrained is only for GLMs, for identity link Gaussian
# response, use parallel_matmult=TRUE to ensure parallel fitting across
# partitions.
# G, G^{-1/2}, and G^{1/2} are computed in parallel across each of the
# K+1 partitions.
# However, parallel_unconstrained only affects GLMs without corr. components
# - it does not affect fitting here
system.time({
  parfit <- lgspline(y ~ spl(a, b) + a*b*c + d,
    data = data.frame(y = y,
                      a = a,
                      b = b,
                      c = c,
                      d = d),
    cl = cl,
    K = 1,
    parallel_eigen = TRUE,
    parallel_unconstrained = TRUE,
    parallel_aga = FALSE,
    parallel_find_neighbors = TRUE,
    parallel_trace = FALSE,
    parallel_matmult = TRUE,
    parallel_qr = FALSE,
    parallel_make_constraint = TRUE,
    parallel_penalty = FALSE)
})
print(summary(parfit))
}

```

Description

Convenience wrapper that calls `lgspline` with the correct family, weight, dispersion, score, and unconstrained-fit functions for Cox proportional hazards regression. All standard `lgspline` arguments (knots, penalties, constraints, parallel, etc.) are passed through.

Usage

```
lgspline_cox(formula, data, status, ...)
```

Arguments

<code>formula</code>	Formula specifying the model. The response should be survival time; <code>status</code> is passed separately.
<code>data</code>	Data frame.
<code>status</code>	Integer vector of event indicators (1 = event, 0 = censored), same length as the number of rows in <code>data</code> .
<code>...</code>	Additional arguments passed to <code>lgspline</code> .

Details

Internally sets:

- `family = cox_family()`
- `unconstrained_fit_fxn = unconstrained_fit_cox`
- `glm_weight_function = cox_glm_weight_function`
- `qp_score_function = cox_qp_score_function`
- `dispersion_function = cox_dispersion_function`
- `schur_correction_function = cox_schur_correction`
- `need_dispersion_for_estimation = FALSE`
- `estimate_dispersion = FALSE`
- `standardize_response = FALSE`

A formula interface is needed, e.g. `lgspline_cox(t, y, ...)` won't work, unlike for ordinary [lgspline](#).

Value

An object of class "lgspline".

Examples

```
## Cox PH with a nonlinear age effect on lung cancer survival
if(requireNamespace("survival", quietly = TRUE)) {
  library(survival)
  set.seed(1234)
  lung <- na.omit(lung[, c("time", "status", "age")])
  lung$age_std <- std(lung$age)
```

```

## survival codes status as 1 = censored, 2 = dead
event <- as.integer(lung$status == 2)

## Spline on age
fit <- lgspline_cox(
  time ~ spl(age_std),
  data = lung,
  status = event,
  K = 1
)
print(summary(fit))
plot(fit,
     show_formulas = TRUE,
     custom_response_lab = 'HR',
     custom_predictor_lab = 'Standardized Age',
     ylim = c(0, 5))
}

```

lgspline_negbin

Fit Negative Binomial Model via lgspline

Description

Convenience wrapper that calls `lgspline` with the correct family, weight, dispersion, score, and unconstrained-fit functions for NB2 regression. All standard `lgspline` arguments (knots, penalties, constraints, parallel, correlation structures, etc.) are passed through.

Usage

```
lgspline_negbin(formula, data, ...)
```

Arguments

formula	Formula specifying the model. The response should be non-negative integer counts.
data	Data frame.
...	Additional arguments passed to <code>lgspline</code> , including <code>Vhalf</code> and <code>VhalfInv</code> for correlation structures.

Details

Internally sets:

- `family = negbin_family()`
- `unconstrained_fit_fxn = unconstrained_fit_negbin`

- `glm_weight_function = negbin_glm_weight_function`
- `qp_score_function = negbin_qp_score_function`
- `dispersion_function = negbin_dispersion_function`
- `schur_correction_function = negbin_schur_correction`
- `need_dispersion_for_estimation = TRUE`
- `estimate_dispersion = TRUE`
- `standardize_response = FALSE`

A formula interface is needed, e.g. `lgspline_negbin(t, y)` won't work, unlike for ordinary [lgspline](#).

When a correlation structure is supplied via `Vhalf/VhalfInv`, the model is fitted through the GEE Path 1b machinery in `get_B`. The dispersion function uses `VhalfInv` to whiten Pearson residuals for a better moment-based initialization of θ , which stabilizes the profile MLE under moderate to strong correlation. The score function handles the whitened design consistently with the Weibull AFT GEE convention.

Value

An object of class "lgspline".

See Also

[lgspline_cox](#) for Cox PH, [lgspline_weibull](#) for Weibull AFT, [negbin_family](#), [unconstrained_fit_negbin](#)

Examples

```
set.seed(1234)
N <- 300
t <- rnorm(N)
mu <- exp(1 + 0.5 * sin(2 * t))
y <- rbinom(N, size = 3, mu = mu)
df <- data.frame(response = y, predictor = t)

fit <- lgspline_negbin(
  response ~ spl(predictor),
  data = df,
  K = 2,
  opt = FALSE,
  wiggle_penalty = 1e-2
)
print(summary(fit))
plot(fit, show_formulas = TRUE,
      custom_response_lab = 'Count')
points(t, mu, col = 'grey', cex=0.67)
```

lgspline_weibull *Fit Weibull Accelerated Failure Time Model via lgspline*

Description

Convenience wrapper that calls `lgspline` with the correct family, weight, dispersion, score, and unconstrained-fit functions for Weibull accelerated failure time regression. All standard `lgspline` arguments (knots, penalties, constraints, parallel, etc.) are passed through.

Usage

```
lgspline_weibull(formula, data, status, ...)
```

Arguments

<code>formula</code>	Formula specifying the model. The response should be survival time; status is passed separately.
<code>data</code>	Data frame.
<code>status</code>	Integer vector of event indicators (1 = event, 0 = censored), same length as the number of rows in data.
<code>...</code>	Additional arguments passed to <code>lgspline</code> .

Details

Internally sets:

- `family = weibull_family()`
- `unconstrained_fit_fxn = unconstrained_fit_weibull`
- `glm_weight_function = weibull_glm_weight_function`
- `qp_score_function = weibull_qp_score_function`
- `dispersion_function = weibull_dispersion_function`
- `schur_correction_function = weibull_schur_correction`
- `need_dispersion_for_estimation = TRUE`
- `estimate_dispersion = TRUE`
- `standardize_response = FALSE`

A formula interface is needed, e.g. `lgspline_weibull(t, y, ...)` won't work, unlike for ordinary [lgspline](#).

Value

An object of class "lgspline".

See Also

[lgspline_cox](#) for Cox proportional hazards, [weibull_family](#), [unconstrained_fit_weibull](#)

Examples

```
## Weibull AFT with a nonlinear age effect on lung cancer survival
if(requireNamespace("survival", quietly = TRUE)) {
  library(survival)
  set.seed(1234)
  lung <- na.omit(lung[, c("time", "status", "age")])
  lung$age_std <- std(lung$age)

  ## survival codes status as 1 = censored, 2 = dead
  event <- as.integer(lung$status == 2)

  ## Spline on age
  fit <- lgspline_weibull(
    time ~ spl(age_std),
    data = lung,
    status = event,
    K = 1,
    opt = FALSE,
    wiggle_penalty = 1e-4,
    flat_ridge_penalty = 1
  )
  print(summary(fit))
  plot(fit,
    show_formulas = TRUE,
    custom_response_lab = 'Survival Time',
    custom_predictor_lab = 'Standardized Age')
}
```

logLik.lgspline

Extract Log-Likelihood from a Fitted lgspline

Description

Returns the log-likelihood as a "logLik" object for use with [AIC](#), [BIC](#), and other model comparison tools.

Usage

```
## S3 method for class 'lgspline'
logLik(
  object,
  include_prior = TRUE,
  new_weights = NULL,
  B_predict = NULL,
  sigmasq_predict = NULL,
  ...
)
```

Arguments

object	A fitted lgspline model object.
include_prior	Logical; add the log-prior penalty term. Default TRUE.
new_weights	Numeric scalar or N-vector; optional observation weights overriding object\$weights.
B_predict	List; optional coefficient list at which to evaluate the likelihood. Default NULL uses object\$B.
sigmasq_predict	Numeric scalar; optional dispersion at which to evaluate the likelihood. Default NULL uses object\$sigmasq_tilde.
...	Not used.

Details**Gaussian identity, no correlation.**

$$\ell = -\frac{N}{2} \log(2\pi\tilde{\sigma}^2) - \frac{1}{2\tilde{\sigma}^2} \sum_i w_i (y_i - \hat{y}_i)^2 + \frac{1}{2} \sum_i \log w_i$$

Gaussian identity, with correlation. GLS log-likelihood:

$$\ell = -\frac{N}{2} \log(2\pi\tilde{\sigma}^2) + \log |\mathbf{V}^{-1/2}| - \frac{1}{2\tilde{\sigma}^2} \sum_i w_i [\mathbf{V}^{-1/2}(\mathbf{y} - \hat{\mathbf{y}})]_i^2 + \frac{1}{2} \sum_i \log w_i$$

$\log |\mathbf{V}^{-1/2}|$ is obtained from `VhalfInv_logdet` when available, or computed directly from `VhalfInv`.

Prior contribution. When `include_prior = TRUE` (default), the log-prior

$$-\frac{1}{2\tilde{\sigma}^2} \sum_k \beta_k^\top \Lambda_k \beta_k$$

is added, giving the penalised MAP log-likelihood coherent with the smoothing spline objective. Set `include_prior = FALSE` for the unpenalised marginal likelihood, which is more appropriate when comparing models with different penalty structures or numbers of knots.

Other GLM families. Uses `family$aic()` when available. For correlated models the whitened residuals and fitted values are passed. When `family$aic()` is unavailable, a deviance-based approximation is used (valid for relative comparisons; a warning is emitted).

This function returns the marginal (full) GLS log-likelihood, not the REML log-likelihood. This is consistent with `REML = FALSE` in `lme` and `gls`, and is the conventional choice for AIC/BIC comparisons of fixed-effects structure.

The `df` attribute is set to $N - \text{trace}(\mathbf{XUGX}^\top)$.

Value

A "logLik" object with attributes `df` (effective degrees of freedom) and `nobs` (number of observations).

See Also

[lgspline](#), [prior_loglik](#), [logLik](#), [AIC](#), [BIC](#)

Examples

```

set.seed(1234)
t <- runif(1000, -10, 10)
y <- 2*sin(t) + -0.06*t^2 + rnorm(length(t))
model_fit <- lgspline(t, y)

logLik(model_fit)
logLik(model_fit, include_prior = FALSE)
logLik(model_fit, B_predict = model_fit$B,
        sigmasq_predict = model_fit$sigmasq_tilde)

AIC(model_fit)
BIC(model_fit)

## Compare models with different K using unpenalized likelihood
fit_k3 <- lgspline(t, y, K = 3)
fit_k7 <- lgspline(t, y, K = 7)
AIC(fit_k3, fit_k7)

```

loglik_cox

Compute Cox Partial Log-Likelihood

Description

Evaluates the Cox partial log-likelihood for a given coefficient vector, using the Breslow approximation for tied event times.

Observations must be sorted in ascending order of survival time before calling this function. The internal helpers handle sorting automatically; this function is exposed for diagnostics and testing.

Usage

```
loglik_cox(eta, status, y = NULL, weights = 1)
```

Arguments

eta	Numeric vector of linear predictors $\mathbf{X}\beta$, length N, sorted by ascending event time.
status	Integer/logical vector of event indicators (1 = event, 0 = censored), same length and order as eta.
y	Optional numeric vector of observed event/censor times, same length and order as eta. When supplied, tied event times are handled by the Breslow approximation using a common risk-set denominator within each tied event-time block. When omitted, the function assumes there are no ties (or that ties have already been expanded appropriately).
weights	Optional numeric vector of observation weights (default 1).

Details

The partial log-likelihood (Breslow) is

$$\ell(\boldsymbol{\beta}) = \sum_g \left[\sum_{i \in D_g} w_i \eta_i - d_g^{(w)} \log \left(\sum_{j \in R_g} w_j \exp(\eta_j) \right) \right]$$

where D_g is the event set at tied event time t_g , $R_g = \{j : t_j \geq t_g\}$ is the corresponding risk set, and $d_g^{(w)} = \sum_{i \in D_g} w_i$.

Value

Scalar partial log-likelihood value.

Examples

```
set.seed(1234)
eta <- rnorm(50)
status <- rbinom(50, 1, 0.6)
y <- rexp(50)
loglik_cox(eta, status, y)
```

loglik_negbin

Compute Negative Binomial Log-Likelihood

Description

Evaluates the NB2 log-likelihood for given mean vector and shape parameter.

Usage

```
loglik_negbin(y, mu, theta, weights = 1)
```

Arguments

y	Non-negative integer response vector.
mu	Positive mean vector, same length as y.
theta	Positive scalar shape parameter.
weights	Optional observation weights (default 1).

Details

The log-likelihood is

$$\ell(\mu, \theta) = \sum_i w_i [\log \Gamma(y_i + \theta) - \log \Gamma(\theta) - \log \Gamma(y_i + 1) + \theta \log \theta - \theta \log(\mu_i + \theta) + y_i \log \mu_i - y_i \log(\mu_i + \theta)]$$

Value

Scalar log-likelihood value.

Examples

```
set.seed(1234)
mu <- exp(rnorm(50))
y <- rpois(50, mu)
loglik_negbin(y, mu, theta = 5)
```

loglik_weibull	<i>Compute Log-Likelihood for Weibull Accelerated Failure Time Model</i>
----------------	--

Description

Calculates the log-likelihood for a Weibull accelerated failure time (AFT) survival model, supporting right-censored survival data.

Usage

```
loglik_weibull(log_y, log_mu, status, scale, weights = 1)
```

Arguments

log_y	Numeric vector of logarithmic response/survival times
log_mu	Numeric vector of logarithmic predicted survival times
status	Numeric vector of censoring indicators (1 = event, 0 = censored) Indicates whether an event of interest occurred (1) or the observation was right-censored (0). In survival analysis, right-censoring occurs when the full survival time is unknown, typically because the study ended or the subject was lost to follow-up before the event of interest occurred.
scale	Numeric scalar representing the Weibull scale parameter (sigma), equivalent to <code>survreg\$scale</code> . This is the square root of the dispersion stored in <code>lgspline\$sigmasq_tilde</code> .
weights	Optional numeric vector of observation weights (default = 1)

Details

The function computes log-likelihood contributions for a Weibull AFT model, explicitly accounting for right-censored observations. It supports optional observation weighting to accommodate complex sampling designs.

This both provides a tool for actually fitting Weibull AFT models, and boilerplate code for users who wish to incorporate Lagrangian multiplier smoothing splines into their own custom models.

Value

A numeric scalar representing the weighted total log-likelihood of the model

Examples

```
## Minimal example of fitting a Weibull Accelerated Failure Time model
# Simulating survival data with right-censoring
set.seed(1234)
x1 <- rnorm(1000)
x2 <- rbinom(1000, 1, 0.5)
yraw <- rexp(exp(0.01*x1 + 0.01*x2))
# status: 1 = event occurred, 0 = right-censored
status <- rbinom(1000, 1, 0.25)
yobs <- ifelse(status, runif(length(yraw), 0, yraw), yraw)
df <- data.frame(
  y = yobs,
  x1 = x1,
  x2 = x2
)

## Fit model using lgspline with Weibull AFT specifics
model_fit <- lgspline(y ~ spl(x1) + x2,
  df,
  unconstrained_fit_fxn = unconstrained_fit_weibull,
  family = weibull_family(),
  need_dispersion_for_estimation = TRUE,
  dispersion_function = weibull_dispersion_function,
  glm_weight_function = weibull_glm_weight_function,
  schur_correction_function = weibull_schur_correction,
  status = status,
  opt = FALSE,
  K = 1)

loglik_weibull(log(model_fit$y), log(model_fit$ytilde), status,
  sqrt(model_fit$sigmasq_tilde))
```

matinvsqrt

Calculate Matrix Inverse Square Root for Symmetric Matrices

Description

Calculate Matrix Inverse Square Root for Symmetric Matrices

Usage

```
matinvsqrt(mat)
```

Arguments

mat A symmetric matrix **M**

Details

Uses an eigenvalue-decomposition-based approach.

Non-positive eigenvalues are set to 0 before taking inverse fourth roots.

This implementation is particularly useful for whitening procedures in GLMs with correlation structures and for computing variance-covariance matrices under constraints.

You can use this to help construct a custom `VhalfInv_fxn` for `lgspline`. When only `VhalfInv` is supplied there, the corresponding `Vhalf` is reconstructed internally by inversion for the GEE code paths.

Value

A matrix **B** such that **BB** equals the Moore-Penrose-style inverse on the positive-eigenvalue subspace, with non-positive components truncated to 0.

Examples

```
## Identity matrix
m1 <- diag(2)
matinvsqrt(m1) # Returns identity matrix

## Compound symmetry correlation matrix
rho <- 0.5
m2 <- matrix(rho, 3, 3) + diag(1-rho, 3)
B <- matinvsqrt(m2)
# Verify: B %**% B approximately equals solve(m2)
all.equal(B %**% B, solve(m2))

## Example for GLM correlation structure
n_blocks <- 2 # Number of subjects
block_size <- 3 # Measurements per subject
rho <- 0.7 # Within-subject correlation
# Correlation matrix for one subject
R <- matrix(rho, block_size, block_size) +
  diag(1-rho, block_size)
## Full correlation matrix for all subjects
V <- kronecker(diag(n_blocks), R)
## Create whitening matrix
VhalfInv <- matinvsqrt(V)

# Example construction of VhalfInv_fxn for lgspline
VhalfInv_fxn <- function(par) {
  rho <- tanh(par) # Transform parameter to (-1, 1)
  R <- matrix(rho, block_size, block_size) +
    diag(1-rho, block_size)
  kronecker(diag(n_blocks), matinvsqrt(R))
}
```

matsqrt

Calculate Matrix Square Root for Symmetric Matrices

Description

Calculate Matrix Square Root for Symmetric Matrices

Usage

```
matsqrt(mat)
```

Arguments

mat A symmetric matrix **M**

Details

Uses an eigenvalue-decomposition-based approach.

Non-positive eigenvalues are set to 0 before taking fourth roots.

This implementation is particularly useful for whitening procedures in GLMs with correlation structures and for computing variance-covariance matrices under constraints.

You can use this to help construct a custom `Vhalf_fxn`, or more directly to build the $\mathbf{V}^{1/2}$ input supplied to [lgspline](#) for correlation-aware fits.

Value

A matrix **B** such that **BB** equals **M** on the positive-eigenvalue subspace, with non-positive components truncated to 0.

Examples

```
## Identity matrix
m1 <- diag(2)
matsqrt(m1) # Returns identity matrix

## Compound symmetry correlation matrix
rho <- 0.5
m2 <- matrix(rho, 3, 3) + diag(1-rho, 3)
B <- matsqrt(m2)
# Verify: B %**% B approximately equals m2
all.equal(B %**% B, m2)

## Example for correlation structure
n_blocks <- 2 # Number of subjects
block_size <- 3 # Measurements per subject
rho <- 0.7 # Within-subject correlation
# Correlation matrix for one subject
R <- matrix(rho, block_size, block_size) +
```

```

      diag(1-rho, block_size)
# Full correlation matrix for all subjects
V <- kronecker(diag(n_blocks), R)
Vhalf <- matsqrt(V)

```

negbin_dispersion_function
NB Dispersion Function

Description

Estimates the shape parameter θ from current fitted values. When a correlation structure is present (`VhalfInv` is non-NULL), the Pearson residuals are whitened before computing the moment-based initial value, giving a better starting point for the profile MLE under correlated data. The final estimate is always the profile MLE over θ .

Usage

```

negbin_dispersion_function(
  mu,
  y,
  order_indices,
  family,
  observation_weights,
  VhalfInv
)

```

Arguments

<code>mu</code>	Predicted values.
<code>y</code>	Observed counts.
<code>order_indices</code>	Observation indices.
<code>family</code>	NB family object.
<code>observation_weights</code>	Observation weights.
<code>VhalfInv</code>	Inverse square root of the correlation matrix, or NULL for independent observations. When non-NULL, used to whiten residuals for the moment-based initialization of θ .

Details

The profile MLE maximizes $\ell(\theta | \mu)$ via Brent's method. When `VhalfInv` is provided, the Pearson residuals $r_i = (y_i - \mu_i) / \sqrt{V(\mu_i)}$ are pre-whitened as $\tilde{r} = V^{-1/2}r$ before computing the moment estimator used for initialization. This accounts for the correlation structure in the variance

decomposition and produces a more stable starting point for the optimizer, particularly when the correlation inflates the marginal variance beyond what the NB model alone would predict.

The profile MLE itself does not use `VhalfInv` because the NB log-likelihood is a marginal quantity; the correlation structure affects estimation only through the mean model (handled by the GEE paths in `get_B`).

Value

Scalar θ estimate (stored as `sigmasq_tilde`).

<code>negbin_family</code>	<i>Negative Binomial Family for lgspline</i>
----------------------------	--

Description

Creates a family-like object for NB2 regression. The link is log, the variance function is $V(\mu) = \mu + \mu^2/\theta$, and the dispersion stored by `lgspline` (`sigmasq_tilde`) is the shape parameter θ .

Usage

```
negbin_family()
```

Details

The NB2 model has a nuisance shape parameter θ analogous to the Weibull scale parameter. It is estimated jointly with β and its uncertainty is propagated via the Schur complement correction.

Value

A list with family components used by `lgspline`.

Examples

```
fam <- negbin_family()
fam$family
fam$link
```

`negbin_glm_weight_function`*NB GLM Weight Function*

Description

Computes working weights for the NB2 information matrix used by lgspline when updating \mathbf{G} after obtaining constrained estimates.

Usage

```
negbin_glm_weight_function(  
  mu,  
  y,  
  order_indices,  
  family,  
  dispersion,  
  observation_weights  
)
```

Arguments

<code>mu</code>	Predicted values $\exp(\eta)$.
<code>y</code>	Observed counts.
<code>order_indices</code>	Observation indices in partition order.
<code>family</code>	NB family object (unused, for interface compatibility).
<code>dispersion</code>	Shape parameter θ .
<code>observation_weights</code>	Observation weights.

Details

The IRLS weight for NB2 with log link is

$$W_i = w_i \mu_i \theta / (\theta + \mu_i)$$

Falls back to observation weights when natural weights are degenerate.

Value

Numeric vector of working weights, length N.

VhalfInv Inverse square root of correlation matrix; when non-NULL the score is computed on the whitened scale as $\tilde{\mathbf{X}}^\top \tilde{\mathbf{r}}$ where $\tilde{\mathbf{X}} = \mathbf{V}^{-1/2} \mathbf{X}$ and the residual vector accounts for the correlation.

observation_weights Observation weights.

Details

Without correlation (VhalfInv = NULL), the score is $\mathbf{X}^\top \mathbf{w} \odot (y - \mu)\theta / (\theta + \mu)$.

With correlation, the GEE score is $\tilde{\mathbf{X}}^\top \text{diag}(\mathbf{W}) \mathbf{V}^{-1} (\mathbf{y} - \boldsymbol{\mu})$ where \mathbf{W} contains the NB working weights. The whitening is absorbed by pre-multiplying both \mathbf{X} and the residual by $\mathbf{V}^{-1/2}$.

Value

Numeric column vector of length P.

negbin_schur_correction

NB Schur Correction

Description

Computes the Schur complement correction to account for uncertainty in estimating θ . Structure is identical to [weibull_schur_correction](#): the joint information is partitioned into $(\boldsymbol{\beta}, \theta)$ blocks and the correction is $-\mathbf{I}_{\boldsymbol{\beta}\theta} \mathbf{I}_{\theta\theta}^{-1} \mathbf{I}_{\boldsymbol{\beta}\theta}^\top$.

Usage

```
negbin_schur_correction(
  X,
  y,
  B,
  dispersion,
  order_list,
  K,
  family,
  observation_weights
)
```

Arguments

X List of partition design matrices.

y List of partition response vectors.

B List of partition coefficient vectors.

dispersion Scalar shape parameter θ .

order_list List of observation indices per partition.

K	Number of knots.
family	Family object.
observation_weights	Observation weights.

Details

The cross-derivative (score of η_i w.r.t. θ) is

$$\frac{\partial^2 \ell}{\partial \eta_i \partial \theta} = \frac{(y_i - \mu_i) \mu_i}{(\theta + \mu_i)^2}$$

The second derivative of the log-likelihood w.r.t. θ is

$$I_{\theta\theta} = - \sum_i w_i \left[\psi'(y_i + \theta) - \psi'(\theta) + \frac{1}{\theta} - \frac{2}{\mu_i + \theta} + \frac{y_i + \theta}{(\mu_i + \theta)^2} \right]$$

where ψ' is the trigamma function.

Value

List of K+1 correction matrices, with \emptyset for empty partitions.

negbin_theta	<i>Estimate Negative Binomial Shape Parameter</i>
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Description

Computes the profile MLE of the shape parameter θ given current mean estimates μ .

Usage

```
negbin_theta(y, mu, weights = 1, init = NULL)
```

Arguments

y	Response vector.
mu	Mean vector.
weights	Observation weights (default 1).
init	Optional initial value for θ . If NULL, uses a moment-based estimate.

Details

Maximizes the profile log-likelihood over θ via Brent's method on $[10^{-4}, 10^4]$.

Value

Scalar MLE of θ .

Examples

```
set.seed(1234)
mu <- rep(5, 200)
y <- rnbino(200, size = 3, mu = 5)
negbin_theta(y, mu)
```

plot.lgspline

Plot Method for lgspline Objects

Description

Wrapper for the internal lgspline plot function. Produces a 1D line plot (base R) or interactive 3D surface plot (plotly) depending on the number of predictors, with optional formula overlays per partition. When plotting a subset of variables via vars, non-plotted predictors are automatically set to zero (or a user-specified value via fixed_values).

Usage

```
## S3 method for class 'lgspline'
plot(
  x,
  show_formulas = FALSE,
  include_all_terms_in_formulas = FALSE,
  digits = 4,
  legend_pos = "topright",
  custom_response_lab = "y",
  custom_predictor_lab = NULL,
  custom_predictor_lab1 = NULL,
  custom_predictor_lab2 = NULL,
  custom_formula_lab = NULL,
  custom_title = "Fitted Function",
  text_size_formula = NULL,
  legend_args = list(),
  new_predictors = NULL,
  xlim = NULL,
  ylim = NULL,
  color_function = NULL,
  add = FALSE,
  vars = c(),
  legend_order = NULL,
  se.fit = FALSE,
  cv = 1,
  band_col = "grey80",
  band_border = NA,
  fixed_values = NULL,
  n_grid = 200,
```

```
    ...
  )
```

Arguments

<code>x</code>	A fitted lgspline model object.
<code>show_formulas</code>	Logical; display partition-level polynomial formulas. Default FALSE.
<code>include_all_terms_in_formulas</code>	Logical; when <code>show_formulas = TRUE</code> and plotting only a subset of predictors via <code>vars</code> , include all fitted terms in the displayed formulas rather than only the terms involving the plotted predictor(s). Default FALSE retains the current marginal-only formula display.
<code>digits</code>	Integer; decimal places for formula coefficients. Default 4.
<code>legend_pos</code>	Character; legend position for 1D plots. Default "topright".
<code>custom_response_lab</code>	Character; response axis label. Default "y".
<code>custom_predictor_lab</code>	Character; predictor axis label (1D). Default NULL uses the column name.
<code>custom_predictor_lab1</code>	Character; first predictor axis label (2D). Default NULL.
<code>custom_predictor_lab2</code>	Character; second predictor axis label (2D). Default NULL.
<code>custom_formula_lab</code>	Character; fitted response label on the link scale. Default NULL.
<code>custom_title</code>	Character; plot title. Default "Fitted Function".
<code>text_size_formula</code>	Numeric; formula text size. Passed to <code>cex</code> (1D) or <code>hover font size</code> (2D). Default NULL (0.8 for 1D, 8 for 2D).
<code>legend_args</code>	List; additional arguments passed to <code>legend()</code> (1D only).
<code>new_predictors</code>	Matrix; optional predictor values for prediction. Default NULL. When <code>vars</code> is specified and <code>new_predictors</code> is NULL, a grid is automatically generated with non-plotted variables set to zero (or values from <code>fixed_values</code>).
<code>xlim</code>	Numeric vector; x-axis limits (1D only). Default NULL.
<code>ylim</code>	Numeric vector; y-axis limits (1D only). Default NULL.
<code>color_function</code>	Function; returns K+1 colors, one per partition. Default NULL uses <code>grDevices::rainbow(K+1)</code> for 1D and a Spectral palette for 2D.
<code>add</code>	Logical; add to an existing plot (1D only). Default FALSE.
<code>vars</code>	Numeric or character vector; predictor indices or names to plot. Default <code>c()</code> plots all.
<code>legend_order</code>	Numeric; re-ordered partition indices for the legend.
<code>se.fit</code>	Logical; if TRUE, plot pointwise confidence bands. Default FALSE.
<code>cv</code>	Numeric; critical value for confidence bands. Default 1.
<code>band_col</code>	Character; color for confidence band fill. Default "grey80".

band_border	Character or NA; border color for confidence band polygon. Default NA (no border).
fixed_values	Named list; fixed values for non-plotted predictors when vars is specified. Names should match predictor names. Default NULL sets non-plotted predictors to zero.
n_grid	Integer; number of grid points for automatic grid generation when vars is specified and new_predictors is NULL. Default 200.
...	Additional arguments passed to plot (1D) or plot_ly (2D).

Details

Partition boundaries are indicated by color changes. For 1D models, observation points can be overlaid. For 2D models, `plotly` is used.

When using `vars` to plot a subset of predictors, the non-plotted predictors are automatically set to zero. This can be overridden by passing a named list to `fixed_values` (e.g., `fixed_values = list(Height = 75)`). The automatic zeroing replaces the previous behavior where the user had to manually construct `new_predictors` with non-plotted variables set to fixed values.

When `se.fit = TRUE`, pointwise confidence bands are drawn around the fitted function. These are Wald-type intervals constructed on the link scale and back-transformed to the response scale, using `cv` as the critical value to multiply `se.fit` by (default 1 for actual `se`).

The function extracts predictor positions from linear expansion terms. If linear terms are excluded (e.g., via `exclude_these_expansions`), plotting will fail. As a workaround, constrain those terms to zero via `constraint_vectors / constraint_values` so they remain in the expansion but are zeroed out.

Value

For 1D models: invisibly returns NULL (base R plot drawn to device). For 2D models: returns a `plotly` object.

See Also

[lgspline](#), [plot](#), [plot_ly](#)

Examples

```
set.seed(1234)
t_data <- runif(1000, -10, 10)
y_data <- 2*sin(t_data) + -0.06*t_data^2 + rnorm(length(t_data))
model_fit <- lgspline(t_data, y_data, K = 9)

## Basic plot
plot(model_fit)

## Plot with confidence bands
plot(model_fit,
      se.fit = TRUE,
      cv = 1.96,
      custom_title = 'Fitted Function with 95% CI')
```

```
## Multi-predictor: automatically zeros non-plotted variables
# plot(model_fit_2d, vars = 'x1', se.fit = TRUE)
```

plot.wald_lgspline *Plot Method for wald_lgspline Objects*

Description

Forest-style plot of coefficient estimates with confidence intervals.

Usage

```
## S3 method for class 'wald_lgspline'
plot(
  x,
  parm = NULL,
  which = NULL,
  main = "Coefficient Estimates and CIs",
  xlab = "Estimate",
  ...
)
```

Arguments

x	A "wald_lgspline" object.
parm	Integer vector of coefficient indices or character vector of names to plot. Default NULL plots all.
which	Integer vector of coefficient indices to plot (alternative to parm). Default NULL.
main	Plot title. Default "Coefficient Estimates and CIs".
xlab	x-axis label. Default "Estimate".
...	Additional arguments passed to plot .

Value

Invisibly returns NULL.

See Also

[wald_univariate](#), [confint.lgspline](#)

predict.lgspline

Predict Method for lgspline Objects

Description

Generates predictions, derivatives, and basis expansions from a fitted lgspline model. Wrapper for the internal predict closure stored in the object.

Usage

```
## S3 method for class 'lgspline'
predict(
  object,
  newdata = NULL,
  parallel = FALSE,
  cl = NULL,
  chunk_size = NULL,
  num_chunks = NULL,
  rem_chunks = NULL,
  B_predict = NULL,
  take_first_derivatives = FALSE,
  take_second_derivatives = FALSE,
  expansions_only = FALSE,
  new_predictors = NULL,
  ...
)
```

Arguments

object	A fitted lgspline model object.
newdata	Matrix or data.frame; new predictor values. Default NULL.
parallel	Logical; use parallel processing (experimental). Default FALSE.
cl	Cluster object for parallel processing. Default NULL.
chunk_size	Integer; chunk size for parallel processing. Default NULL.
num_chunks	Integer; number of chunks. Default NULL.
rem_chunks	Integer; remainder chunks. Default NULL.
B_predict	List; per-partition coefficient list for prediction, e.g. from generate_posterior . Default NULL uses object\$B.
take_first_derivatives	Logical; compute first derivatives. Default FALSE.
take_second_derivatives	Logical; compute second derivatives. Default FALSE.
expansions_only	Logical; return basis expansion matrix only. Default FALSE.
new_predictors	Matrix or data.frame; overrides newdata.
...	Additional arguments passed to the internal predict method.

Details

`new_predictors` takes priority over `newdata` when both are supplied. When both are NULL, the training data is used.

Fitted values are also accessible directly as `model_fit$ytilde` or via `model_fit$predict()`.

The parallel processing feature is experimental.

Additional arguments passed through `...` include `se.fit` and `cv` for pointwise interval summaries.

Predictor input should use the original predictor columns. Named extra columns are dropped when they can be identified as irrelevant to the fitted expansions.

Value

A numeric vector of predictions, or a list when derivatives or interval summaries are requested:

preds Numeric vector of predictions when derivatives are requested.

fit Numeric vector of predictions when `se.fit = TRUE` and no derivatives are requested.

first_deriv Numeric vector or named list of first derivatives (if requested).

second_deriv Numeric vector or named list of second derivatives (if requested).

se.fit Pointwise standard errors on the link scale (if requested).

lower Pointwise lower interval bound (if requested).

upper Pointwise upper interval bound (if requested).

cv Critical value returned when `se.fit = TRUE` without derivative requests.

If `expansions_only = TRUE`, returns a list of basis expansions.

See Also

[lgspline](#), [plot.lgspline](#)

Examples

```
set.seed(1234)
t <- runif(1000, -10, 10)
y <- 2*sin(t) + -0.06*t^2 + rnorm(length(t))
model_fit <- lgspline(t, y)

newdata <- matrix(sort(rnorm(10000)), ncol = 1)
preds <- predict(model_fit, newdata)

deriv1_res <- predict(model_fit, newdata, take_first_derivatives = TRUE)
deriv2_res <- predict(model_fit, newdata, take_second_derivatives = TRUE)

oldpar <- par(no.readonly = TRUE)
layout(matrix(c(1,1,2,2,3,3), byrow = TRUE, ncol = 2))

plot(newdata[,1], preds, main = 'Fitted Function',
      xlab = 't', ylab = "f(t)", type = 'l')
plot(newdata[,1], deriv1_res$first_deriv, main = 'First Derivative',
```

```

      xlab = 't', ylab = "f'(t)", type = 'l')
plot(newdata[,1], deriv2_res$second_deriv, main = 'Second Derivative',
      xlab = 't', ylab = "f''(t)", type = 'l')

par(oldpar)

```

`print.lgspline` *Print Method for lgspline Objects*

Description

Prints a concise summary of the fitted model to the console.

Usage

```

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

```

Arguments

`x` An lgspline model object.
`...` Not used.

Value

Invisibly returns `x`.

`print.summary.lgspline` *Print Method for lgspline Summaries*

Description

Displays a formatted model summary using `printCoefmat` for the coefficient table.

Usage

```

## S3 method for class 'summary.lgspline'
print(x, ...)

```

Arguments

`x` A `summary.lgspline` object.
`...` Not used.

Value

Invisibly returns x.

See Also

[printCoefmat](#)

`print.wald_lgspline` *Print Method for wald_lgspline Objects*

Description

Prints the coefficient table using [printCoefmat](#) with significance stars.

Usage

```
## S3 method for class 'wald_lgspline'
print(
  x,
  digits = max(3, getOption("digits") - 3),
  signif.stars = getOption("show.signif.stars"),
  ...
)
```

Arguments

<code>x</code>	A "wald_lgspline" object from wald_univariate .
<code>digits</code>	Number of significant digits.
<code>signif.stars</code>	Logical; show significance stars.
<code>...</code>	Additional arguments passed to printCoefmat .

Value

Invisibly returns x.

See Also

[wald_univariate](#), [printCoefmat](#)

prior_loglik *Log-Prior Distribution Evaluation for lgspline Models*

Description

Evaluates the log-prior on the spline coefficients conditional on the dispersion and penalty matrices.

Usage

```
prior_loglik(
  model_fit,
  B_predict = NULL,
  sigmasq_predict = NULL,
  include_constant = TRUE,
  ...
)
```

Arguments

model_fit	An lgspline model object.
B_predict	Optional list of coefficient vectors at which to evaluate the prior. Default NULL uses the fitted coefficients.
sigmasq_predict	Numeric scalar dispersion parameter. If NULL, model_fit\$sigmasq_tilde is used. Legacy sigmasq calls are still accepted.
include_constant	Logical; if TRUE (default), include the multivariate normal normalizing constant.
...	Optional legacy arguments.

Details

Returns the quadratic form of $\beta^T \Lambda \beta$ evaluated at the tuned or fixed penalties, scaled by negative one-half inverse dispersion.

Assuming fixed penalties, the prior on β is taken to be

$$\beta | \sigma^2 \sim \mathcal{N}(\mathbf{0}, \sigma^2 \Lambda^{-1})$$

so that, up to a normalizing constant C with respect to β ,

$$\implies \log P(\beta | \sigma^2) = C - \frac{1}{2\sigma^2} \beta^T \Lambda \beta$$

The value of C is included when include_constant = TRUE, and omitted when FALSE.

This is useful for computing joint penalized log-likelihoods and related MAP-style diagnostics for a fitted lgspline object.

Value

A numeric scalar representing the prior log-likelihood.

See Also

[lgspline](#)

Examples

```
## Data
t <- sort(runif(100, -5, 5))
y <- sin(t) - 0.1*t^2 + rnorm(100)

## Model keeping penalties fixed
model_fit <- lgspline(t, y, opt = FALSE)

## Full joint log-likelihood, conditional upon known sigma^2 = 1
jntloglik <- sum(dnorm(model_fit$y,
                      model_fit$ytilde,
                      1,
                      log = TRUE)) +
  prior_loglik(model_fit, sigmasq_predict = 1)
print(jntloglik)
```

 process_qp

Prepare Quadratic Programming Constraints for lgspline

Description

Builds the linear inequality system used for shape-restricted fitting. The helper collects built-in range, derivative-sign, second-derivative, and monotonicity restrictions, together with any user-supplied custom constraint functions, and returns the resulting $\mathbf{C}^T \boldsymbol{\beta} \geq \mathbf{c}$ objects.

This logic was refactored out of [lgspline.fit](#) so the constraint construction can be reviewed and tested on its own. Existing calls that pass TRUE/FALSE for derivative flags remain backward-compatible.

Usage

```
process_qp(
  X,
  K,
  p_expansions,
  order_list,
  colnm_expansions,
  expansion_scales,
```

```

power1_cols,
power2_cols,
nonspline_cols,
interaction_single_cols,
interaction_quad_cols,
triplet_cols,
include_2way_interactions,
include_3way_interactions,
include_quadratic_interactions,
family,
mean_y,
sd_y,
N_obs,
qp_observations = NULL,
qp_positive_derivative = FALSE,
qp_negative_derivative = FALSE,
qp_positive_2ndderivative = FALSE,
qp_negative_2ndderivative = FALSE,
qp_monotonic_increase = FALSE,
qp_monotonic_decrease = FALSE,
qp_range_upper = NULL,
qp_range_lower = NULL,
qp_Amat_fxn = NULL,
qp_bvec_fxn = NULL,
qp_meq_fxn = NULL,
qp_Amat = NULL,
qp_bvec = NULL,
qp_meq = 0,
qr_pivot_inequality_constraints = FALSE,
parallel_qr_qp = FALSE,
all_derivatives_fxn = NULL,
og_cols = NULL,
cl = NULL,
include_warnings = TRUE,
...
)

```

Arguments

X	List of per-partition design matrices.
K	Integer. Number of interior knots.
p_expansions	Integer. Number of basis expansions per partition.
order_list	List of per-partition observation index vectors.
colnm_expansions	Character vector of expansion column names.
expansion_scales	Numeric vector of expansion standardization scales.

power1_cols	Integer vector of linear-term column indices.
power2_cols	Integer vector of quadratic-term column indices.
nonspline_cols	Integer vector of non-spline linear column indices.
interaction_single_cols	Integer vector of linear-by-linear interaction column indices.
interaction_quad_cols	Integer vector of linear-by-quadratic interaction column indices.
triplet_cols	Integer vector of three-way interaction column indices.
include_2way_interactions	Logical switch controlling whether two-way interactions are included in derivative construction.
include_3way_interactions	Logical switch controlling whether three-way interactions are included in derivative construction.
include_quadratic_interactions	Logical switch controlling whether quadratic interactions are included in derivative construction.
family	GLM family object.
mean_y, sd_y	Numeric scalars for response standardization.
N_obs	Integer. Total sample size.
qp_observations	Optional observation subset. Supply either a single integer vector to thin every active built-in QP constraint the same way, or a named list keyed by "var: qp_<type>" (or bare "qp_<type>") to give different constraints different subsets. Known types are qp_range_lower, qp_range_upper, qp_positive_derivative, qp_negative_derivative, qp_positive_2ndderivative, qp_negative_2ndderivative, qp_monotonic_increase, and qp_monotonic_decrease. For range and monotonicity, use the bare keys such as "qp_range_lower" and "qp_monotonic_increase" because those constraints are variable-independent; prefixed versions are still accepted and unioned internally. Unknown keyed entries are ignored with a warning when include_warnings = TRUE.
qp_positive_derivative, qp_negative_derivative	Logical scalar, character vector, or integer vector. See section <i>Per-Variable Derivative Constraints</i> .
qp_positive_2ndderivative, qp_negative_2ndderivative	Same as above but for second derivatives.
qp_monotonic_increase, qp_monotonic_decrease	Logical. Constrain fitted values to be monotonic in observation order.
qp_range_upper, qp_range_lower	Optional numeric upper/lower bounds for fitted values.
qp_Amat_fxn, qp_bvec_fxn, qp_meq_fxn	Optional user-supplied constraint-generating functions.
qp_Amat, qp_bvec, qp_meq	Optional pre-built QP objects. Their presence marks QP handling as active, but this helper does not append them to the built-in constraints it constructs; they are expected to be handled outside this constructor.

qr_pivot_inequality_constraints	Logical. If TRUE, partition-local inequality columns are thinned to QR pivot columns before the final solve. QP objects are stacked. Leading equality columns are left untouched; built-in range and monotonicity blocks are also left untouched; and columns spanning multiple partitions are left untouched.
parallel_qr_qp	Logical. If TRUE and a cluster is supplied in <code>cl</code> , the partition-local QR pivot steps used by <code>qr_pivot_inequality_constraints</code> are parallelized across partitions.
all_derivatives_fxn	Function to compute derivatives from expansion matrices (the <code>all_derivatives</code> closure from <code>lgspline.fit</code>).
og_cols	Optional character vector of original predictor column names.
cl	Optional parallel cluster object used only when <code>parallel_qr_qp = TRUE</code> .
include_warnings	Logical. Whether to issue warnings.
...	Additional arguments forwarded to custom constraint functions.

Value

A list with components:

- qp_Amat** $P \times M$ combined constraint matrix.
- qp_bvec** Numeric vector of length M .
- qp_meq** Integer. Number of leading equality constraints.
- quadprog** Logical. TRUE if any QP constraints are active.

Per-Variable Derivative Constraints

The arguments `qp_positive_derivative`, `qp_negative_derivative`, `qp_positive_2ndderivative`, and `qp_negative_2ndderivative` now accept three forms:

FALSE No constraint (default).

TRUE Constrain **all** predictor variables (backward compatible with previous behavior).

Character or integer vector Constrain only the specified predictor variables. Character entries are resolved via `og_cols`; integer entries refer to column positions in the predictor matrix.

This allows, for example, enforcing a nonnegative first derivative for "Dose" and a nonpositive first derivative for "Time" simultaneously:

```
lgspline(...,
  qp_positive_derivative = "Dose",
  qp_negative_derivative = "Time")
```

The arguments `qp_monotonic_increase` and `qp_monotonic_decrease` remain TRUE/FALSE only, because they constrain fitted values in observation order (not per-variable).

Examples

```

## Not run:
## Standalone verification: simple 1-D monotonic increase
set.seed(1234)
t <- seq(-5, 5, length.out = 200)
y <- 3 * sin(t) + t + rnorm(200, 0, 0.5)

## Fit with positive first-derivative constraint on all variables
fit1 <- lgspline(t, y, K = 3,
                qp_positive_derivative = TRUE)

## Verify: first derivative should be >= 0 everywhere
derivs1 <- predict(fit1, new_predictors = sort(t),
                  take_first_derivatives = TRUE)
stopifnot(all(derivs1$first_deriv >= -1e-8))

## Fit with monotonic increase (observation-order)
fit2 <- lgspline(t, y, K = 3,
                qp_monotonic_increase = TRUE)
preds2 <- predict(fit2, new_predictors = sort(t))
stopifnot(all(diff(preds2) >= -1e-8))

## Per-variable constraints: 2-D example
t1 <- runif(500, -5, 5)
t2 <- runif(500, -5, 5)
y2 <- t1 + sin(t2) + rnorm(500, 0, 0.5)
dat2 <- data.frame(t1 = t1, t2 = t2, y = y2)

## Constrain t1 to have positive derivative, t2 to have negative
fit3 <- lgspline(y ~ spl(t1, t2), data = dat2, K = 2,
                qp_positive_derivative = "t1",
                qp_negative_derivative = "t2")

## Verify per-variable derivatives
newdat <- expand.grid(t1 = seq(-4, 4, length.out = 50),
                    t2 = seq(-4, 4, length.out = 50))
d3 <- predict(fit3, new_predictors = newdat,
              take_first_derivatives = TRUE)

## t1 derivative should be >= 0
stopifnot(all(unlist(d3$first_deriv[["_1_"]] ) >= -1e-6))
## t2 derivative should be <= 0
stopifnot(all(unlist(d3$first_deriv[["_2_"]] ) <= 1e-6))

## Numeric column indices work identically
fit4 <- lgspline(y ~ spl(t1, t2), data = dat2, K = 2,
                qp_positive_derivative = 1,
                qp_negative_derivative = 2)

## Range + derivative constraints simultaneously
fit5 <- lgspline(t, y, K = 3,
                qp_positive_derivative = TRUE,

```

```

qp_range_lower = -5,
qp_range_upper = 15)
preds5 <- predict(fit5)
stopifnot(all(preds5 >= -5 - 1e-6))
stopifnot(all(preds5 <= 15 + 1e-6))

## End(Not run)

```

reml_grad_from_dV *Evaluate the REML gradient with respect to a single correlation parameter*

Description

Computes the derivative of the negative REML objective with respect to a scalar correlation parameter, given the matrix derivative $\partial\mathbf{V}/\partial\rho$.

Usage

```
reml_grad_from_dV(dV, model_fit, glm_weight_function, ...)
```

Arguments

dV $N \times N$ numeric matrix giving $\partial\mathbf{V}/\partial\rho$ for one scalar parameter ρ , with the chain rule through any reparameterization already applied.

model_fit A fitted lgspline object; see [lgspline](#).

glm_weight_function The GLM weight function used during model fitting; see the `glm_weight_function` argument of [lgspline](#).

... Additional arguments forwarded to `glm_weight_function`.

Details

Notation: $\mathbf{D} = \text{diag}(d_1, \dots, d_N)$ is the diagonal matrix of observation weights (`observation_weights`), and $\mathbf{W} = \text{diag}(w_1, \dots, w_N)$ is the diagonal matrix of GLM working weights evaluated at the current fitted values via `glm_weight_function`, with the observation-weight contribution carried separately by \mathbf{D} . For canonical GLM families, w_i is the usual IRWLS/Fisher-scoring weight on the mean–variance scale; for example, logistic regression gives $w_i = \mu_i(1 - \mu_i)$. The combined weighting entering the information matrix is \mathbf{WD} . In Gaussian identity models both reduce to scalar multiples of the identity.

\mathbf{V} is the $N \times N$ correlation matrix implied by `VhalfInv`, with $\mathbf{V}^{-1} = (\mathbf{V}^{-1/2})^\top \mathbf{V}^{-1/2}$.

The penalized observed information at the current iterate is

$$\mathbf{M} = (\mathbf{X}^*)^\top \mathbf{V}^{-1} \mathbf{W} \mathbf{D} \mathbf{X}^* + \mathbf{U}^\top \boldsymbol{\Lambda} \mathbf{U},$$

where $\mathbf{X}^* = \mathbf{X}\mathbf{U}$ is the constrained design ($N \times P$) and the first term is the quadratic approximation to the penalized log-likelihood Hessian at the current $\boldsymbol{\mu}$. For non-Gaussian families this is a local approximation (the IRWLS/Fisher scoring Hessian), not an exact GLS information matrix. The constraint projection is $\mathbf{U} = \mathbf{I} - \mathbf{G}\mathbf{A}(\mathbf{A}^\top\mathbf{G}\mathbf{A})^{-1}\mathbf{A}^\top$ with $\mathbf{G} = \mathbf{M}^{-1}$. \mathbf{U} is idempotent ($\mathbf{U}^2 = \mathbf{U}$) but *not* symmetric, so $\mathbf{U}^\top\boldsymbol{\Lambda}\mathbf{U} \neq \mathbf{U}\boldsymbol{\Lambda}\mathbf{U}^\top$.

The \mathbf{U} stored in `model_fit$U` is on the expansion- and response-standardised scale. The rescaled version used here is

$$\tilde{\mathbf{U}} = \mathbf{U} \cdot \text{diag}(1, s_1, \dots, s_{p-1}, 1, s_1, \dots) / \hat{\sigma}_y,$$

where s_j are the expansion scales and $\hat{\sigma}_y$ standardises the response. All quantities below use $\tilde{\mathbf{U}}$ in place of \mathbf{U} .

REML objective and its gradient: The REML objective is constructed by integrating out the fixed effects from the penalized log-likelihood, using a Laplace approximation to the marginal likelihood for non-Gaussian families. This approximation is exact for Gaussian identity models and is the standard extension used in restricted maximum likelihood estimation for GLMMs.

The REML correction term is $-\frac{1}{2} \log |\mathbf{M}|$, where \mathbf{M} is the penalized observed information defined above. Differentiating with respect to ρ (noting only \mathbf{V} depends on ρ) gives the REML correction gradient

$$-\frac{1}{2} \text{tr} \left(\mathbf{M}^{-1} (\mathbf{X}^*)^\top \mathbf{V}^{-1} \frac{\partial \mathbf{V}}{\partial \rho} \mathbf{V}^{-1} \mathbf{W} \mathbf{D} \mathbf{X}^* \right).$$

Full gradient:

$$\frac{\partial(-\ell_R)}{\partial \rho} = \frac{1}{N} \left[\underbrace{\frac{1}{2} \text{tr} \left(\mathbf{V}^{-1} \frac{\partial \mathbf{V}}{\partial \rho} \right)}_{\text{(i) log-det of } \mathbf{V}} \quad \underbrace{-\frac{1}{2\hat{\sigma}^2} \mathbf{r}^\top \frac{\partial \mathbf{V}}{\partial \rho} \mathbf{r}}_{\text{(ii) residual quadratic form}} \quad \underbrace{-\frac{1}{2} \text{tr} \left(\mathbf{M}^{-1} (\mathbf{X}^*)^\top \mathbf{V}^{-1} \frac{\partial \mathbf{V}}{\partial \rho} \mathbf{V}^{-1} \mathbf{W} \mathbf{D} \mathbf{X}^* \right)}_{\text{(iii) REML correction}} \right],$$

where the whitened residual is

$$\mathbf{r} = \text{diag} \left(\sqrt{d_i} / \sqrt{w_i} \right) \mathbf{V}^{-1/2} (\mathbf{y} - \boldsymbol{\mu}),$$

and $r_i = [\mathbf{V}^{-1/2} (\mathbf{y} - \boldsymbol{\mu})]_i \sqrt{d_i} / \sqrt{w_i}$.

Term (i) does not involve \mathbf{D} or \mathbf{W} ; the log-determinant of \mathbf{V} depends only on the correlation structure. For non-Gaussian families terms (ii) and (iii) are evaluated at the current IRWLS iterate and constitute a local approximation.

Value

A scalar: the gradient of the negative REML objective with respect to ρ , divided by N .

See Also

[lgspline](#), [lgspline.fit](#)

std	<i>Standardize Vector to Z-Scores</i>
-----	---------------------------------------

Description

Centers a vector by its sample mean, then scales it by its sample standard deviation $(x - \text{mean}(x)) / \text{sd}(x)$.

Usage

```
std(x)
```

Arguments

x	Numeric vector to standardize
---	-------------------------------

Value

Standardized vector with sample mean 0 and standard deviation 1

Examples

```
v <- c(1, 2, 3, 4, 5)
std(v)
print(mean(v))
print(sd(v))
```

summary.lgspline	<i>Summary Method for lgspline Objects</i>
------------------	--

Description

Summary Method for lgspline Objects

Usage

```
## S3 method for class 'lgspline'
summary(object, ...)
```

Arguments

object	An lgspline model object.
...	Not used.

Value

An object of class `summary.lgspline`, a list containing:

model_family The `family` object.

observations Number of observations N .

predictors Number of predictor variables q .

knots Number of knots K .

basis_functions Basis functions per partition p .

estimate_dispersion "Yes" or "No".

cv Critical value used for confidence intervals.

coefficients Coefficient matrix from `wald_univariate`, or a single-column estimate matrix if `return_varcovmat = FALSE`.

sigmasq_tilde Estimated dispersion $\tilde{\sigma}^2$.

trace_XUGX Trace of the hat matrix $\text{trace}(\mathbf{XUGX}^T)$.

N Number of observations.

`summary.wald_lgspline` *Summary Method for wald_lgspline Objects*

Description

Prints a header with model info then delegates to `print.wald_lgspline`.

Usage

```
## S3 method for class 'wald_lgspline'
summary(object, ...)
```

Arguments

`object` A "wald_lgspline" object.
`...` Passed to `print.wald_lgspline`.

Value

Invisibly returns object.

See Also

`wald_univariate`, `print.wald_lgspline`

wald_univariate	<i>Univariate Wald Tests and Confidence Intervals for lgspline Coefficients</i>
-----------------	---

Description

Computes per-coefficient Wald tests and confidence intervals from a fitted lgspline. For Gaussian identity-link models, t-statistics and t-intervals are used; otherwise z-statistics.

Usage

```
wald_univariate(object, scale_vcovmat_by = 1, cv, ...)
```

Arguments

object	A fitted lgspline object. Must have been fit with <code>return_varcovmat = TRUE</code> .
scale_vcovmat_by	Numeric; scaling factor for the variance-covariance matrix. Default 1.
cv	Numeric; critical value for confidence intervals. If missing, defaults to <code>object\$critical_value</code> or <code>qnorm(0.975)</code> .
...	Additional arguments passed to the internal <code>wald_univariate</code> method.

Value

An object of class "wald_lgspline", a list with:

coefficients Matrix with columns: Estimate, Std. Error, t value or z value, Pr(>|t|) or Pr(>|z|), CI LB, CI UB.

critical_value Critical value used.

family GLM family from the fitted model.

N Number of observations.

trace_XUGX Effective df trace term.

statistic_name "t value" or "z value".

p_value_name "Pr(>|t|)" or "Pr(>|z|)".

df.residual Residual degrees of freedom when supplied by the internal Wald method.

Print, summary, and plot methods are available; see [print.wald_lgspline](#), [summary.wald_lgspline](#), [plot.wald_lgspline](#).

See Also

[lgspline](#), [confint.lgspline](#), [print.wald_lgspline](#), [summary.wald_lgspline](#), [plot.wald_lgspline](#)

Examples

```

set.seed(1234)
t <- runif(1000, -10, 10)
y <- 2*sin(t) + -0.06*t^2 + rnorm(length(t))
model_fit <- lgspline(t, y, return_varcovmat = TRUE)

wald_default <- wald_univariate(model_fit)
print(wald_default)

## t-distribution critical value
eff_df <- model_fit$N - model_fit$trace_XUGX
wald_t <- wald_univariate(model_fit, cv = qt(0.975, eff_df))
print(wald_t)

coef_table <- wald_default$coefficients
plot(wald_default)

```

weibull_dispersion_function

Estimate Weibull Dispersion for Accelerated Failure Time Model

Description

Computes the dispersion parameter ($\sigma^2 = \text{scale}^2$) for a Weibull accelerated failure time (AFT) model, supporting right-censored survival data. The returned value is σ^2 , where σ is the Weibull scale parameter matching `survreg$scale`.

This both provides a tool for actually fitting Weibull AFT Models, and boilerplate code for users who wish to incorporate Lagrangian multiplier smoothing splines into their own custom models.

Usage

```

weibull_dispersion_function(
  mu,
  y,
  order_indices,
  family,
  observation_weights,
  VhalfInv,
  status
)

```

Arguments

<code>mu</code>	Predicted survival times
<code>y</code>	Observed response/survival times
<code>order_indices</code>	Indices to align status with response

family	Weibull AFT model family specification; unused here and retained for interface compatibility.
observation_weights	Optional observation weights
VhalfInv	Inverse square root of the correlation matrix; unused here and retained for interface compatibility.
status	Censoring indicator (1 = event, 0 = censored) Indicates whether an event of interest occurred (1) or the observation was right-censored (0). In survival analysis, right-censoring occurs when the full survival time is unknown, typically because the study ended or the subject was lost to follow-up before the event of interest occurred.

Value

Dispersion estimate (σ^2) for the Weibull AFT model, i.e., the squared scale parameter. The Weibull scale (σ) matching `survreg$scale` is `sqrt()` of this value.

See Also

[weibull_scale](#) for the underlying scale estimation function

Examples

```
## Simulate survival data with covariates
set.seed(1234)
n <- 1000
t1 <- rnorm(n)
t2 <- rbinom(n, 1, 0.5)

## Generate survival times with Weibull-like structure
lambda <- exp(0.5 * t1 + 0.3 * t2)
yraw <- rexp(n, rate = 1/lambda)

## Introduce right-censoring
status <- rbinom(n, 1, 0.75)
y <- ifelse(status, yraw, runif(length(yraw), 0, yraw))

## Example of using dispersion function
mu <- mean(y)
order_indices <- seq_along(y)
weights <- rep(1, n)

## Estimate dispersion (= scale^2 = sigma^2)
dispersion_est <- weibull_dispersion_function(
  mu = mu,
  y = y,
  order_indices = order_indices,
  family = weibull_family(),
  observation_weights = weights,
  VhalfInv = NULL,
  status = status
```

```

)

print(dispersion_est)      # sigma^2
print(sqrt(dispersion_est)) # sigma (comparable to survreg$scale)

```

weibull_family *Weibull Family for Survival Model Specification*

Description

Creates a family-like object for Weibull accelerated failure time (AFT) models, including custom log-likelihood, AIC, and deviance helpers.

This both provides a tool for actually fitting Weibull AFT Models, and boilerplate code for users who wish to incorporate Lagrangian multiplier smoothing splines into their own custom models.

Usage

```
weibull_family()
```

Details

Provides a comprehensive family specification for Weibull AFT models, including family name, link function, inverse link function, custom loss function for model tuning, and methods for AIC and log-likelihood computation compatible with `logLik.lgspline`.

Supports right-censored survival data with flexible parameter estimation.

Note on scale vs. dispersion: throughout this package, the `lgspline` object stores `sigmasq_tilde` which equals σ^2 (dispersion), where σ is the Weibull scale parameter matching `survreg$scale`. Functions that accept a dispersion argument receive σ^2 ; functions that accept a scale argument receive σ .

Value

A family-like list containing the link functions and Weibull-specific methods used by `lgspline`.

Examples

```

## Simulate survival data with covariates
set.seed(1234)
n <- 1000
t1 <- rnorm(n)
t2 <- rbinom(n, 1, 0.5)

## Generate survival times with Weibull-like structure
lambda <- exp(0.5 * t1 + 0.3 * t2)
yraw <- rexp(n, rate = 1/lambda)

## Introduce right-censoring

```

```

status <- rbinom(n, 1, 0.75)
y <- ifelse(status, yraw, runif(length(yraw), 0, yraw))

## Prepare data
df <- data.frame(y = y, t1 = t1, t2 = t2, status = status)

## Fit model using custom Weibull family
model_fit <- lgspline(y ~ spl(t1) + t2,
  df,
  unconstrained_fit_fxn = unconstrained_fit_weibull,
  family = weibull_family(),
  need_dispersion_for_estimation = TRUE,
  dispersion_function = weibull_dispersion_function,
  glm_weight_function = weibull_glm_weight_function,
  schur_correction_function = weibull_schur_correction,
  status = status,
  opt = FALSE,
  K = 1)

summary(model_fit)

## Log-likelihood now works via logLik.lgspline:
# logLik(model_fit)

```

```
weibull_glm_weight_function
```

Weibull GLM Weight Function for Constructing Information Matrix

Description

Computes the working weights used in the Weibull AFT information matrix, including the observation-weight contribution returned on the vector scale.

Usage

```

weibull_glm_weight_function(
  mu,
  y,
  order_indices,
  family,
  dispersion,
  observation_weights,
  status
)

```

Arguments

mu	Predicted survival times
y	Observed response/survival times
order_indices	Order of observations when partitioned to match "status" to "response"
family	Weibull AFT family; unused here and retained for interface compatibility.
dispersion	Estimated dispersion parameter ($\sigma^2 = \text{scale}^2$). The lgspline framework stores and passes dispersion (σ^2); the Weibull scale (σ) matching <code>survreg\$scale</code> is $\sqrt{\text{dispersion}}$.
observation_weights	Weights of observations submitted to function
status	Censoring indicator (1 = event, 0 = censored) Indicates whether an event of interest occurred (1) or the observation was right-censored (0). In survival analysis, right-censoring occurs when the full survival time is unknown, typically because the study ended or the subject was lost to follow-up before the event of interest occurred.

Details

This function generates weights used in constructing the information matrix after unconstrained estimates have been found. Specifically, it is used in the construction of the **U** and **G** matrices following initial unconstrained parameter estimation.

These weights are analogous to the variance terms in generalized linear models (GLMs). Like logistic regression uses $\mu(1 - \mu)$, Poisson regression uses e^μ , and Linear regression uses constant weights, Weibull AFT models use $\exp((\log y - \log \mu)/\sigma)$ where $\sigma = \sqrt{\text{dispersion}}$ is the scale parameter.

Value

Numeric vector of working weights for the information matrix, including observation weights when finite and a fallback of 1s when the natural Weibull weights are non-finite.

Examples

```
## Demonstration of glm weight function in constrained model estimation
set.seed(1234)
n <- 1000
t1 <- rnorm(n)
t2 <- rbinom(n, 1, 0.5)

## Generate survival times
lambda <- exp(0.5 * t1 + 0.3 * t2)
yraw <- rexp(n, rate = 1/lambda)

## Introduce right-censoring
status <- rbinom(n, 1, 0.75)
y <- ifelse(status, yraw, runif(length(yraw), 0, yraw))

## Fit model demonstrating use of custom glm weight function
```

```

model_fit <- lgspline(y ~ spl(t1) + t2,
  data.frame(y = y, t1 = t1, t2 = t2),
  unconstrained_fit_fxn = unconstrained_fit_weibull,
  family = weibull_family(),
  need_dispersion_for_estimation = TRUE,
  dispersion_function = weibull_dispersion_function,
  glm_weight_function = weibull_glm_weight_function,
  schur_correction_function = weibull_schur_correction,
  status = status,
  opt = FALSE,
  K = 1)

print(summary(model_fit))

```

weibull_qp_score_function

Compute Gradient of Log-Likelihood of Weibull Accelerated Failure Model

Description

Calculates the gradient of log-likelihood for a Weibull accelerated failure time (AFT) survival model, supporting right-censored survival data.

Usage

```

weibull_qp_score_function(
  X,
  y,
  mu,
  order_list,
  dispersion,
  VhalfInv,
  observation_weights,
  status
)

```

Arguments

X	Design matrix
y	Response vector
mu	Predicted mean vector
order_list	List of observation indices per partition
dispersion	Dispersion parameter ($\sigma^2 = \text{scale}^2$). The lgspline framework stores and passes dispersion (σ^2); the Weibull scale (σ) matching <code>survreg\$scale</code> is <code>sqrt(dispersion)</code> .

VhalfInv	Inverse square root of correlation matrix; unused here and retained for interface compatibility.
observation_weights	Observation weights
status	Censoring indicator (1 = event, 0 = censored)

Details

Needed if using "blockfit", correlation structures, or quadratic programming with Weibull AFT models.

The gradient is computed on a scale that omits the $1/\sigma$ prefactor. Specifically, the true score is $(1/\sigma) * X^T \text{diag}(w) (\exp(z) - \text{status})$, but both this function and the corresponding information matrix used internally omit $1/\sigma$ and $1/\sigma^2$ respectively, so the Newton-Raphson step G^*u remains correct. This matches the convention in [unconstrained_fit_weibull](#).

Value

Numeric column vector representing the gradient with respect to coefficients.

Examples

```
set.seed(1234)
t1 <- rnorm(1000)
t2 <- rbinom(1000, 1, 0.5)
yraw <- rexp(exp(0.01*t1 + 0.01*t2))
status <- rbinom(1000, 1, 0.25)
yobs <- ifelse(status, runif(length(yraw), 0, yraw), yraw)
df <- data.frame(
  y = yobs,
  t1 = t1,
  t2 = t2
)

## Example using blockfit for t2 as a linear term - output does not look
# different, but internal methods used for fitting change
model_fit <- lgspline(y ~ spl(t1) + t2,
  df,
  unconstrained_fit_fxn = unconstrained_fit_weibull,
  family = weibull_family(),
  need_dispersion_for_estimation = TRUE,
  qp_score_function = weibull_qp_score_function,
  dispersion_function = weibull_dispersion_function,
  glm_weight_function = weibull_glm_weight_function,
  schur_correction_function = weibull_schur_correction,
  K = 1,
  blockfit = TRUE,
  opt = FALSE,
  status = status,
  verbose = TRUE)

print(summary(model_fit))
```

weibull_scale *Estimate Scale for Weibull Accelerated Failure Time Model*

Description

Computes maximum log-likelihood scale estimate (σ) for a Weibull accelerated failure time (AFT) survival model.

This both provides a tool for actually fitting Weibull AFT Models, and boilerplate code for users who wish to incorporate Lagrangian multiplier smoothing splines into their own custom models.

Usage

```
weibull_scale(log_y, log_mu, status, weights = 1)
```

Arguments

log_y	Logarithm of response/survival times
log_mu	Logarithm of predicted survival times
status	Censoring indicator (1 = event, 0 = censored) Indicates whether an event of interest occurred (1) or the observation was right-censored (0). In survival analysis, right-censoring occurs when the full survival time is unknown, typically because the study ended or the subject was lost to follow-up before the event of interest occurred.
weights	Optional observation weights (default = 1)

Details

Calculates maximum log-likelihood estimate of scale (σ) for Weibull AFT model accounting for right-censored observations using Brent's method for optimization.

Value

Scalar representing the estimated Weibull scale (σ), equivalent to `survreg$scale`. The dispersion (as stored in `lgspline$sigma_sq_tilde`) is σ^2 .

Examples

```
## Simulate exponential data with censoring
set.seed(1234)
mu <- 2 # mean of exponential distribution
n <- 500
y <- rexp(n, rate = 1/mu)

## Introduce censoring (25% of observations)
status <- rbinom(n, 1, 0.75)
```

```

y_obs <- ifelse(status, y, NA)

## Compute scale estimate
scale_est <- weibull_scale(
  log_y = log(y_obs[!is.na(y_obs)]),
  log_mu = log(mu),
  status = status[!is.na(y_obs)]
)

print(scale_est)

```

```
weibull_schur_correction
```

Correction for the Variance-Covariance Matrix for Uncertainty in Scale

Description

Computes the Schur complement \mathbf{S} such that $\mathbf{G}^* = (\mathbf{G}^{-1} + \mathbf{S})^{-1}$ properly accounts for uncertainty in estimating the Weibull scale parameter when estimating the variance-covariance matrix. Otherwise, the variance-covariance matrix is optimistic and assumes the scale is known, when it was in fact estimated. Note that the parameterization adds the output of this function elementwise (not subtract) so for most cases, the output of this function will be negative or a negative definite/semi-definite matrix.

Usage

```

weibull_schur_correction(
  X,
  y,
  B,
  dispersion,
  order_list,
  K,
  family,
  observation_weights,
  status
)

```

```

weibull_shur_correction(
  X,
  y,
  B,
  dispersion,
  order_list,
  K,

```

```

    family,
    observation_weights,
    status
  )

```

Arguments

<code>x</code>	Block-diagonal matrices of spline expansions
<code>y</code>	Block-vector of response
<code>B</code>	Block-vector of coefficient estimates
<code>dispersion</code>	Scalar, estimate of dispersion ($\sigma^2 = \text{scale}^2$). The <code>lgspline</code> framework stores and passes dispersion (σ^2); the Weibull scale (σ) matching <code>survreg\$scale</code> is <code>sqrt(dispersion)</code> .
<code>order_list</code>	List of partition orders
<code>K</code>	Number of partitions minus 1 (K)
<code>family</code>	Distribution family
<code>observation_weights</code>	Optional observation weights (default = 1)
<code>status</code>	Censoring indicator (1 = event, 0 = censored) Indicates whether an event of interest occurred (1) or the observation was right-censored (0). In survival analysis, right-censoring occurs when the full survival time is unknown, typically because the study ended or the subject was lost to follow-up before the event of interest occurred.

Details

Adjusts the variance-covariance matrix unscaled for coefficients to account for uncertainty in estimating the Weibull scale parameter, that otherwise would be lost if simply using $\mathbf{G} = (\mathbf{X}^T \mathbf{W} \mathbf{X} + \mathbf{L})^{-1}$. This is accomplished using a correction based on the Schur complement so we avoid having to construct the entire variance-covariance matrix, or modifying the procedure for `lgspline` substantially. For any model with nuisance parameters that must have uncertainty accounted for, this tool will be helpful.

This both provides a tool for actually fitting Weibull accelerated failure time (AFT) models, and boilerplate code for users who wish to incorporate Lagrangian multiplier smoothing splines into their own custom models.

Value

List of blockwise Schur-complement corrections \mathbf{S}_k to be elementwise added to each block of the information matrix before inversion, with \emptyset returned for empty partitions.

Examples

```

## Minimal example of fitting a Weibull Accelerated Failure Time model
# Simulating survival data with right-censoring
set.seed(1234)
t1 <- rnorm(1000)

```

```
t2 <- rbinom(1000, 1, 0.5)
yraw <- rexp(exp(0.01*t1 + 0.01*t2))
# status: 1 = event occurred, 0 = right-censored
status <- rbinom(1000, 1, 0.25)
yobs <- ifelse(status, runif(length(yraw), 0, yraw), yraw)
df <- data.frame(
  y = yobs,
  t1 = t1,
  t2 = t2
)

## Fit model using lgspline with Weibull Schur correction
model_fit <- lgspline(y ~ spl(t1) + t2,
  df,
  unconstrained_fit_fxn = unconstrained_fit_weibull,
  family = weibull_family(),
  need_dispersion_for_estimation = TRUE,
  dispersion_function = weibull_dispersion_function,
  glm_weight_function = weibull_glm_weight_function,
  schur_correction_function = weibull_schur_correction,
  status = status,
  opt = FALSE,
  K = 1)

print(summary(model_fit))

## Fit model using lgspline without Weibull Schur correction
naive_fit <- lgspline(y ~ spl(t1) + t2,
  df,
  unconstrained_fit_fxn = unconstrained_fit_weibull,
  family = weibull_family(),
  need_dispersion_for_estimation = TRUE,
  dispersion_function = weibull_dispersion_function,
  glm_weight_function = weibull_glm_weight_function,
  status = status,
  opt = FALSE,
  K = 1)

print(summary(naive_fit))
```

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