

Package: fmrihrf (via r-universe)

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Type Package

Title Hemodynamic Response Functions for fMRI Data Analysis

Version 0.3.0

Description Creates, manipulates, and evaluates hemodynamic response functions and event-related regressors for functional magnetic resonance imaging data analysis. Supports multiple basis sets including Canonical, Gamma, Gaussian, B-spline, and Fourier bases. Features decorators for time-shifting and blocking, and efficient convolution algorithms for regressor construction. Methods are based on standard fMRI analysis techniques as described in Jezzard et al. (2001, ISBN:9780192630711).

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acquisition_onsets *Get fMRI Acquisition Onset Times*

Description

Calculate the onset time in seconds for each fMRI volume acquisition from the start of the experiment.

Usage

```

acquisition_onsets(x, ...)

## S3 method for class 'sampling_frame'
acquisition_onsets(x, ...)

```

Arguments

- x A `sampling_frame` object
- ... Additional arguments (for extensibility)

Details

Returns the temporal onset of each brain volume acquisition, accounting for TR, start_time, and run structure. This is essentially a convenience wrapper around `samples(x, global = TRUE)` that provides clearer semantic meaning for the common use case of getting acquisition times.

Note: The onset times include the start_time offset (default TR/2), so the first acquisition typically doesn't start at 0.

Value

Numeric vector of acquisition onset times in seconds

See Also

[samples](#) for more flexible timing queries

Examples

```
# Single block with default start_time (TR/2 = 1)
sf <- sampling_frame(blocklens = 100, TR = 2)
onsets <- acquisition_onsets(sf)
head(onsets) # Returns: 1, 3, 5, 7, 9, 11, ...

# Multiple blocks with same TR
sf2 <- sampling_frame(blocklens = c(100, 120), TR = 2)
onsets2 <- acquisition_onsets(sf2)
# First block: 1, 3, 5, ..., 199
# Second block: 201, 203, 205, ..., 439

# Variable TR per block
sf3 <- sampling_frame(blocklens = c(100, 100), TR = c(2, 1.5))
onsets3 <- acquisition_onsets(sf3)
# First block: 1, 3, 5, ..., 199 (TR=2)
# Second block: 200.75, 202.25, 203.75, ... (TR=1.5, start_time=0.75)

# Custom start times
sf4 <- sampling_frame(blocklens = c(50, 50), TR = 2, start_time = 0)
onsets4 <- acquisition_onsets(sf4)
head(onsets4) # Returns: 0, 2, 4, 6, 8, 10, ...
```

amplitudes

Get amplitudes from an object

Description

Generic accessor returning event amplitudes or scaling factors.

Usage

```
amplitudes(x, ...)  
  
## S3 method for class 'Reg'  
amplitudes(x, ...)
```

Arguments

```
x          Object containing amplitude information  
...       Additional arguments passed to methods
```

Value

Numeric vector of amplitudes

Examples

```
# Create a regressor with varying amplitudes  
reg <- regressor(onsets = c(1, 5, 10), hrf = HRF_SPMG1,  
                amplitude = c(1, 0.5, 2),  
                span = 20)  
amplitudes(reg)
```

block_hrf	<i>Create a Blocked HRF Object</i>
-----------	------------------------------------

Description

Creates a new HRF object representing a response to a sustained (blocked) stimulus by convolving the input HRF with a boxcar function of a given width.

Usage

```
block_hrf(  
  hrf,  
  width,  
  precision = 0.1,  
  half_life = Inf,  
  summate = TRUE,  
  normalize = FALSE  
)
```

Arguments

hrf	The HRF object (of class 'HRF') to block.
width	The width of the block in seconds.
precision	The sampling precision in seconds used for the internal convolution (default: 0.1).
half_life	The half-life of an optional exponential decay applied during the block (default: Inf, meaning no decay).
summate	Logical; if TRUE (default), responses within the block are integrated (summed). If FALSE, the integrated response is divided by the total block weight so amplitude does not grow with block width.
normalize	Logical; if TRUE, the resulting blocked HRF is scaled so that its peak value is 1 (default: FALSE).

Value

A new HRF object representing the blocked function.

See Also

Other HRF_decorator_functions: [lag_hrf\(\)](#), [normalise_hrf\(\)](#)

Examples

```
blocked_spmg1 <- block_hrf(HRF_SPMG1, width = 5)
t_vals <- seq(0, 30, by = 0.5)
plot(t_vals, HRF_SPMG1(t_vals), type = 'l', col = "blue", ylab = "Response", xlab = "Time")
lines(t_vals, blocked_spmg1(t_vals), col = "red")
legend("topright", legend = c("Original", "Blocked (width=5)"), col = c("blue", "red"), lty = 1)
```

blockids

Get block identifiers

Description

Generic accessor returning block indices for each sample or onset.

Usage

```
blockids(x, ...)

## S3 method for class 'sampling_frame'
blockids(x, ...)
```

Arguments

x	Object containing block structure
...	Additional arguments passed to methods

Value

Integer vector of block ids

Examples

```
# Get block identifiers from a sampling frame
sframe <- sampling_frame(blocklens = c(100, 120, 80), TR = 2)
blockids(sframe)
```

blocklens	<i>Get block lengths</i>
-----------	--------------------------

Description

Generic accessor returning the number of scans in each block of a sampling frame or similar object.

Usage

```
blocklens(x, ...)
```

S3 method for class 'sampling_frame'

```
blocklens(x, ...)
```

Arguments

x	Object containing block length information
...	Additional arguments passed to methods

Value

Numeric vector of block lengths

Examples

```
# Get block lengths from a sampling frame
sframe <- sampling_frame(blocklens = c(100, 120, 80), TR = 2)
blocklens(sframe)
```

`deriv`*Compute derivatives of HRF functions*

Description

Calculates the derivative of a Hemodynamic Response Function (HRF) at specified time points. This is useful for:

- Understanding HRF dynamics and rate of change
- Creating temporal derivative regressors for fMRI models
- Analyzing HRF shape characteristics
- Implementing advanced HRF basis sets

Usage

```
deriv(x, t, ...)
```

Arguments

<code>x</code>	An HRF object
<code>t</code>	Numeric vector of time points at which to evaluate the derivative
<code>...</code>	Additional arguments passed to specific methods

Details

The derivative computation method depends on the HRF type:

- Analytic derivatives are used when available (e.g., SPMG1, SPMG2, SPMG3)
- Numeric finite-difference approximation is used as fallback

The default implementation uses `numDeriv::grad` for numerical differentiation when analytic derivatives are not available.

Value

Numeric vector or matrix of derivative values at the specified time points. For multi-basis HRFs, returns a matrix with one column per basis function.

See Also

[[evaluate\(\)](#)], [[HRF_objects](#)], [[numDeriv::grad\(\)](#)]

Other hrf: [HRF_objects](#), [penalty_matrix\(\)](#)

Examples

```
# Compute derivative of SPM canonical HRF
t <- seq(0, 20, by = 0.1)
hrf_deriv <- deriv(HRF_SPMG1, t)

# Plot HRF and its derivative
hrf_vals <- evaluate(HRF_SPMG1, t)
plot(t, hrf_vals, type = "l", col = "black",
      ylab = "Response", xlab = "Time (s)")
lines(t, hrf_deriv, col = "red", lty = 2)
legend("topright", c("HRF", "Derivative"),
      col = c("black", "red"), lty = c(1, 2))

# For multi-basis HRFs, returns matrix
deriv_matrix <- deriv(HRF_SPMG3, t)
# Returns derivatives for all 3 basis functions
```

deriv.HRF

Default derivative method for HRF objects

Description

Uses numerical differentiation via `numDeriv::grad` when analytic derivatives are not available for a specific HRF type.

Uses the analytic derivative formula for the SPM canonical HRF.

Returns derivatives for both the canonical HRF and its temporal derivative. The first column contains the derivative of the canonical HRF, and the second column contains the second derivative (derivative of the temporal derivative).

Returns derivatives for the canonical HRF and its two derivatives. Since SPMG3 already includes first and second derivatives as basis functions, this method returns their derivatives (1st, 2nd, and 3rd derivatives of the original HRF).

Usage

```
## S3 method for class 'HRF'
deriv(x, t, ...)

## S3 method for class 'SPMG1_HRF'
deriv(x, t, ...)

## S3 method for class 'SPMG2_HRF'
deriv(x, t, ...)

## S3 method for class 'SPMG3_HRF'
deriv(x, t, ...)
```

Arguments

x An SPMG3_HRF object
 t Numeric vector of time points at which to evaluate the derivative
 ... Additional arguments (currently unused)

Value

Numeric vector or matrix of derivative values
 Numeric vector of derivative values
 Matrix with 2 columns of derivative values
 Matrix with 3 columns of derivative values

Examples

```
t <- seq(0, 30, by = 0.5)
d <- deriv(HRF_SPMG1, t)
```

durations

Get durations of an object

Description

Get durations of an object

Usage

```
durations(x, ...)
```

```
## S3 method for class 'Reg'
```

```
durations(x, ...)
```

Arguments

x The object to get durations from
 ... Additional arguments passed to methods

Value

A numeric vector of durations

Examples

```
# Create a regressor with event durations
reg <- regressor(onsets = c(1, 5, 10), hrf = HRF_SPMG1,
                 duration = c(2, 3, 1), span = 20)
durations(reg)
```

evaluate	<i>Evaluate a regressor object over a time grid</i>
----------	---

Description

Generic function to evaluate a regressor object over a specified time grid. Different types of regressors may have different evaluation methods.

Usage

```
evaluate(x, grid, ...)

## S3 method for class 'Reg'
evaluate(
  x,
  grid,
  precision = 0.33,
  method = c("conv", "fft", "Rconv", "loop"),
  sparse = FALSE,
  normalize = FALSE,
  ...
)
```

Arguments

x	A 'Reg' object (or an object inheriting from it, like 'regressor').
grid	Numeric vector specifying the time points (seconds) for evaluation.
...	Additional arguments passed down (e.g., to 'evaluate.HRF' in the loop method).
precision	Numeric sampling precision for internal HRF evaluation and convolution (seconds).
method	The evaluation method: conv (Default) Uses the C++ direct convolution ('evaluate_regressor_convolution'). Generally safer and more predictable. fft Uses the fast C++ FFT convolution ('evaluate_regressor_fast'). Can be faster but may fail with very fine precision or wide grids. Extremely fine 'precision' or wide 'grid' ranges may trigger an internal FFT size exceeding $\sim 1e7$, which results in an error. Rconv Uses an R-based convolution ('stats::convolve'). Requires constant event durations and a regular sampling grid. Can be faster than the R loop for many events meeting these criteria. loop Uses a pure R implementation involving looping through onsets. Can be slower, especially for many onsets.
sparse	Logical indicating whether to return a sparse matrix (from the Matrix package). Default is FALSE.

normalize Logical; if TRUE, scale evaluated regressor output to unit peak (maximum absolute value of 1). For multi-basis regressors, each basis column is normalized independently.

Value

A numeric vector or matrix containing the evaluated regressor values

See Also

[single_trial_regressor()], [regressor()]

Examples

```
# Create a regressor
reg <- regressor(onsets = c(10, 30, 50), hrf = HRF_SPMG1)

# Evaluate at specific time points
times <- seq(0, 80, by = 0.1)
response <- evaluate(reg, times)

# Plot the response
plot(times, response, type = "l", xlab = "Time (s)", ylab = "Response")
# Create a regressor
reg <- regressor(onsets = c(10, 30, 50), hrf = HRF_SPMG1)

# Evaluate with default method (conv)
times <- seq(0, 80, by = 0.5)
response <- evaluate(reg, times)

# Try different evaluation methods
response_loop <- evaluate(reg, times, method = "loop")

# With higher precision
response_precise <- evaluate(reg, times, precision = 0.1)
```

evaluate.HRF

Evaluate an HRF Object

Description

This function evaluates a hemodynamic response function (HRF) object for a given set of time points (grid) and other parameters. It handles both point evaluation (duration=0) and block evaluation (duration > 0).

Usage

```
## S3 method for class 'HRF'
evaluate(
  x,
  grid,
  amplitude = 1,
  duration = 0,
  precision = 0.2,
  summate = TRUE,
  normalize = FALSE,
  ...
)
```

Arguments

x	The HRF object (inherits from 'HRF' and 'function').
grid	A numeric vector of time points at which to evaluate the HRF.
amplitude	The scaling value for the event (default: 1).
duration	The duration of the event (seconds). If > 0, the HRF is evaluated over this duration (default: 0).
precision	The temporal resolution for evaluating responses when duration > 0 (default: 0.2).
summate	Logical; whether the HRF response should accumulate over the duration (default: TRUE). If FALSE, the convolution is averaged so the temporal profile is preserved but peak amplitude does not grow with duration.
normalize	Logical; scale output so that the peak absolute value is 1 (default: FALSE). Applied <i>after</i> amplitude scaling and duration processing.
...	Additional arguments (unused).

Value

A numeric vector or matrix of HRF values at the specified time points.

Examples

```
# Evaluate canonical HRF at specific times
times <- seq(0, 20, by = 0.5)
response <- evaluate(HRF_SPMG1, times)

# Evaluate with amplitude scaling
response_scaled <- evaluate(HRF_SPMG1, times, amplitude = 2)

# Evaluate with duration (block design)
response_block <- evaluate(HRF_SPMG1, times, duration = 5, summate = TRUE)

# Multi-basis HRF evaluation
response_multi <- evaluate(HRF_SPMG3, times) # Returns 3-column matrix
```

gen_hrf

*Construct an HRF Instance using Decorators***Description**

'gen_hrf' takes a base HRF function or object and applies optional lag, blocking, and normalization decorators based on arguments.

Usage

```
gen_hrf(
    hrf,
    lag = 0,
    width = 0,
    precision = 0.1,
    half_life = Inf,
    summate = TRUE,
    normalize = FALSE,
    name = NULL,
    span = NULL,
    ...
)
```

Arguments

hrf	A function 'f(t)' or an existing 'HRF' object.
lag	Optional lag in seconds. If non-zero, applies 'lag_hrf'.
width	Optional block width in seconds. If non-zero, applies 'block_hrf'.
precision	Sampling precision for block convolution (passed to 'block_hrf'). Default is 0.1.
half_life	Half-life decay parameter for exponential decay in seconds (passed to 'block_hrf'). Default is Inf (no decay).
summate	Passed to 'block_hrf()' when 'width > 0'. If 'TRUE' (default), block responses are integrated; if 'FALSE', the integrated response is scaled by total block weight so amplitude does not grow with block width.
normalize	If TRUE, applies 'normalise_hrf' at the end. Default is FALSE.
name	Optional name for the *final* HRF object. If NULL (default), a name is generated based on the base HRF and applied decorators.
span	Optional span for the *final* HRF object. If NULL (default), the span is determined by the base HRF and decorators.
...	Extra arguments passed to the *base* HRF function if 'hrf' is a function.

Value

A final 'HRF' object, potentially modified by decorators.

Examples

```
# Lagged SPMG1
grf_lag <- gen_hrf(HRF_SPMG1, lag=3)
# Blocked Gaussian
grf_block <- gen_hrf(hrf_gaussian, width=5, precision=0.2)
# Lagged and Blocked, then Normalized
grf_both_norm <- gen_hrf(HRF_SPMG1, lag=2, width=4, normalize=TRUE)
```

gen_hrf_blocked	<i>Generate a Blocked HRF Function</i>
-----------------	--

Description

The 'gen_hrf_blocked' function creates a blocked HRF by convolving the input HRF with a boxcar function. This can be used to model block designs in fMRI analysis.

Usage

```
gen_hrf_blocked(  
  hrf = hrf_gaussian,  
  width = 5,  
  precision = 0.1,  
  half_life = Inf,  
  summate = TRUE,  
  normalize = FALSE,  
  ...  
)
```

```
hrf_blocked(  
  hrf = hrf_gaussian,  
  width = 5,  
  precision = 0.1,  
  half_life = Inf,  
  summate = TRUE,  
  normalize = FALSE,  
  ...  
)
```

Arguments

hrf	A function representing the hemodynamic response function. Default is 'hrf_gaussian'.
width	A numeric value specifying the width of the block in seconds. Default is 5.
precision	A numeric value specifying the sampling resolution in seconds. Default is 0.1.
half_life	A numeric value specifying the half-life of the exponential decay function, used to model response attenuation. Default is 'Inf', which means no decay.

summate	Logical; if TRUE (default), responses accumulate (peak grows with duration). If FALSE, the convolution is averaged so the temporal profile is preserved but peak amplitude does not grow with duration.
normalize	A logical value indicating whether to rescale the output so that the peak of the output is 1. Default is 'FALSE'.
...	Extra arguments passed to the HRF function.

Value

A function representing the blocked HRF.

A function representing the blocked HRF.

Functions

- `hrf_blocked()`: alias for `gen_hrf_blocked`

See Also

Other `gen_hrf`: [gen_hrf_lagged\(\)](#)

Examples

```
# Deprecated: use gen_hrf(..., width = 10) or block_hrf(HRF, width = 10)
```

<code>gen_hrf_lagged</code>	<i>Generate a Lagged HRF Function</i>
-----------------------------	---------------------------------------

Description

The 'gen_hrf_lagged' function takes an HRF function and applies a specified lag to it. This can be useful for modeling time-delayed hemodynamic responses.

Usage

```
gen_hrf_lagged(hrf, lag = 2, normalize = FALSE, ...)
```

```
hrf_lagged(hrf, lag = 2, normalize = FALSE, ...)
```

Arguments

hrf	A function representing the underlying HRF to be shifted.
lag	A numeric value specifying the lag or delay in seconds to apply to the HRF. This can also be a vector of lags, in which case the function returns an HRF set.
normalize	A logical value indicating whether to rescale the output so that the maximum absolute value is 1. Defaults to 'FALSE'.
...	Extra arguments supplied to the 'hrf' function.

Value

A function representing the lagged HRF. If 'lag' is a vector of lags, the function returns an HRF set.
an lagged hrf function

Functions

- `hrf_lagged()`: alias for `gen_hrf_lagged`

See Also

Other `gen_hrf`: [gen_hrf_blocked\(\)](#)

Other `gen_hrf`: [gen_hrf_blocked\(\)](#)

Examples

```
hrf_lag5 <- gen_hrf_lagged(HRF_SPMG1, lag=5)
hrf_lag5(0:20)
```

getHRF

Get HRF by Name

Description

Retrieves an HRF by name from the registry and optionally applies decorators. This provides a unified interface for creating both pre-defined HRF objects and custom basis sets with specified parameters.

Usage

```
getHRF(  
  name = "spm1",  
  nbasis = 5,  
  span = 24,  
  lag = 0,  
  width = 0,  
  summate = TRUE,  
  normalize = FALSE,  
  ...  
)
```

Arguments

name	Character string specifying the HRF type. Options include: <ul style="list-style-type: none"> • "spm1", "spm2", "spm3" - SPM canonical HRFs • "gamma", "gaussian" - Simple parametric HRFs • "fir" - Finite Impulse Response basis • "bspline" or "bs" - B-spline basis • "fourier" - Fourier basis • "daguerre" - Daguerre spherical basis • "tent" - Tent (linear spline) basis
nbasis	Number of basis functions (for basis set types)
span	Temporal window in seconds (default: 24)
lag	Time lag in seconds to apply (default: 0)
width	Block width for block designs (default: 0)
summate	Whether to sum responses in block designs (default: TRUE)
normalize	Whether to normalize the HRF (default: FALSE)
...	Additional arguments passed to generator functions (e.g., scale for daguerre)

Details

For single HRF types (spm1, gamma, gaussian), the function returns pre-defined objects. For basis set types (fir, bspline, fourier, daguerre), it calls the appropriate generator function with the specified parameters.

Value

An HRF object

Examples

```
# Get pre-defined canonical HRF
canonical <- getHRF("spm1")

# Create custom FIR basis with 20 bins
fir20 <- getHRF("fir", nbasis = 20, span = 30)

# Create B-spline basis with lag
bs_lag <- getHRF("bspline", nbasis = 8, lag = 2)

# Create blocked Gaussian HRF
block_gauss <- getHRF("gaussian", width = 5)
```

global_onsets	<i>Convert onsets to global timing</i>
---------------	--

Description

Generic accessor for converting block-wise onsets to global onsets.

Usage

```
global_onsets(x, ...)

## S3 method for class 'sampling_frame'
global_onsets(x, onsets, blockids, ...)
```

Arguments

x	Object describing the sampling frame
...	Additional arguments passed to methods
onsets	Numeric vector of onset times within blocks
blockids	Integer vector identifying the block for each onset. Values must be whole numbers with no NAs.

Value

Numeric vector of global onset times

Examples

```
# Convert block-relative onsets to global timing
sframe <- sampling_frame(blocklens = c(100, 120), TR = 2)
global_onsets(sframe, onsets = c(10, 20), blockids = c(1, 2))
```

HRF	<i>HRF Constructor Function</i>
-----	---------------------------------

Description

The 'HRF' function creates an object representing a hemodynamic response function (HRF). It is a class constructor for HRFs.

Usage

```
HRF(fun, name, nbasis = 1, span = 24, param_names = NULL)
```

Arguments

fun	A function representing the hemodynamic response, mapping from time to BOLD response.
name	A string specifying the name of the function.
nbasis	An integer representing the number of basis functions, e.g., the columnar dimension of the HRF. Default is 1.
span	A numeric value representing the span in seconds of the HRF. Default is 24.
param_names	A character vector containing the names of the parameters for the HRF function.

Details

The package provides several pre-defined HRF types that can be used in modeling fMRI responses:

****Canonical HRFs:**** * "spm1" or 'HRF_SPMG1': SPM's canonical HRF (single basis function) * "spm2" or 'HRF_SPMG2': SPM canonical + temporal derivative (2 basis functions) * "spm3" or 'HRF_SPMG3': SPM canonical + temporal and dispersion derivatives (3 basis functions) * "gaussian" or 'HRF_GAUSSIAN': Gaussian-shaped HRF with peak around 5-6s * "gamma" or 'HRF_GAMMA': Gamma function-based HRF with longer tail

****Flexible basis sets:**** * "bspline" or "bs" or 'HRF_BSPLINE': B-spline basis for flexible HRF modeling * "tent": Tent (triangular) basis functions for flexible HRF modeling * "daguerre" or 'HRF_DAGUERRE': Daguerre basis functions

To see a complete list of available HRF types with details, use the 'list_available_hrf()' function.

Value

An HRF object with the specified properties.

Examples

```
hrf <- HRF(hrf_gamma, "gamma", nbasis=1, param_names=c("shape", "rate"))
resp <- evaluate(hrf, seq(0, 24, by=1))

# List all available HRF types
list_available_hrf(details = TRUE)
```

hrf_basis_lwu

LWU HRF Basis for Taylor Expansion

Description

Constructs the basis set for the Lag-Width-Undershoot (LWU) HRF model, intended for Taylor expansion-based fitting. The basis consists of the LWU HRF evaluated at a given expansion point θ_0 , and its partial derivatives with respect to its parameters (τ , σ , ρ).

Usage

```
hrf_basis_lwu(theta0, t, normalize_primary = "none")
```

Arguments

theta0	A numeric vector of length 3 specifying the expansion point $c(\tau_0, \sigma_0, \rho_0)$ for the LWU parameters.
t	A numeric vector of time points (in seconds) at which to evaluate the basis.
normalize_primary	Character string, one of "none" or "height". If "height", the primary HRF column ($h_0(t)$) is normalized to have a peak absolute value of 1. For Taylor expansion fitting as described in <code>Fit_LRU.md</code> , this should typically be "none" as the scaling is absorbed by the beta coefficient. Default is "none".

Value

A numeric matrix of dimension $\text{length}(t) \times 4$. The columns represent:

- h_0 : LWU HRF evaluated at θ_0
- d_τ : Partial derivative with respect to τ at θ_0
- d_σ : Partial derivative with respect to σ at θ_0
- d_ρ : Partial derivative with respect to ρ at θ_0

See Also

[hrf_lwu, grad](#)

Other `hrf_functions`: [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
t_points <- seq(0, 30, by = 0.5)
theta0_default <- c(tau = 6, sigma = 1, rho = 0.35)

# Generate the basis set
lwu_basis <- hrf_basis_lwu(theta0_default, t_points)
dim(lwu_basis) # Should be length(t_points) x 4
head(lwu_basis)

# Plot the basis functions
matplot(t_points, lwu_basis, type = "l", lty = 1,
        main = "LWU HRF Basis Functions", ylab = "Value", xlab = "Time (s)")
legend("topright", colnames(lwu_basis), col = 1:4, lty = 1, cex = 0.8)

# Example with primary HRF normalization (not typical for Taylor fitting step)
lwu_basis_norm_h0 <- hrf_basis_lwu(theta0_default, t_points, normalize_primary = "height")
plot(t_points, lwu_basis_norm_h0[,1], type="l", main="Normalized h0 in Basis")
max(abs(lwu_basis_norm_h0[,1])) # Should be 1
```

hrf_boxcar	<i>Boxcar HRF (No Hemodynamic Delay)</i>
------------	--

Description

Creates a simple boxcar (step function) HRF that is constant within a time window starting at $t=0$ and zero outside. Unlike traditional HRFs, this has no hemodynamic delay - it represents an instantaneous response.

Usage

```
hrf_boxcar(width, amplitude = 1, normalize = FALSE)
```

Arguments

width	Duration of the boxcar window in seconds.
amplitude	Height of the boxcar (default: 1).
normalize	Logical; if TRUE, the boxcar is scaled so that its integral equals 1 (i.e., amplitude = 1/width). This makes the regression coefficient interpretable as the mean signal in the window. Default is FALSE.

Details

When used in a GLM, the estimated coefficient represents a (weighted) average of the data within the specified time window. If `normalize = TRUE`, the coefficient directly estimates the mean signal in that window.

For delayed windows (not starting at $t=0$), use [lag_hrf](#) to shift the boxcar in time.

Value

An HRF object that can be used with `regressor()` and other `fmrihrf` functions.

Note on durations

The width is fixed when the HRF is created. The duration parameter in `regressor()` does **not** modify the boxcar width—it controls how long the neural input is sustained (which then gets convolved with this HRF). For trial-varying boxcar widths, use a list of HRFs:

```
widths <- c(4, 6, 8)
hrfs <- lapply(widths, function(w) hrf_boxcar(width = w, normalize = TRUE))
reg <- regressor(onsets = c(0, 20, 40), hrf = hrfs)
```

See Also

[hrf_weighted](#) for weighted/shaped boxcars, [lag_hrf](#) to shift the window in time

Other `hrf_` functions: [hrf_basis_lwu\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
# Simple boxcar of 5 seconds width
hrf1 <- hrf_boxcar(width = 5)
t <- seq(-1, 10, by = 0.1)
plot(t, evaluate(hrf1, t), type = "s", main = "Simple Boxcar HRF")

# Normalized boxcar - coefficient will estimate mean signal in window
hrf2 <- hrf_boxcar(width = 5, normalize = TRUE)
# integral is now 1, so beta estimates mean(Y[0:5])

# Use in a regressor with trial-varying widths
hrf_short <- hrf_boxcar(width = 4, normalize = TRUE)
hrf_long <- hrf_boxcar(width = 8, normalize = TRUE)
reg <- regressor(onsets = c(0, 20), hrf = list(hrf_short, hrf_long))

# For delayed windows, use lag_hrf decorator
hrf_delayed <- lag_hrf(hrf_boxcar(width = 5), lag = 10) # Window from 10-15s
```

hrf_bspline

*B-spline HRF (hemodynamic response function)***Description**

The ‘hrf_bspline’ function computes the B-spline representation of an HRF (hemodynamic response function) at given time points ‘t’.

Usage

```
hrf_bspline(t, span = 24, N = 5, degree = 3, ...)
```

Arguments

t	A vector of time points.
span	A numeric value representing the temporal window over which the basis set spans. Default value is 20.
N	An integer representing the number of basis functions. Default value is 5.
degree	An integer representing the degree of the spline. Default value is 3.
...	Additional arguments passed to ‘splines::bs’.

Value

A matrix representing the B-spline basis for the HRF at the given time points ‘t’.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
# Compute the B-spline HRF representation for time points from 0 to 20 with 0.5 increments
hrfb <- hrf_bspline(seq(0, 20, by = .5), N = 4, degree = 2)
```

hrf_bspline_generator *Create B-spline HRF Basis Set*

Description

Generates an HRF object using B-spline basis functions with custom parameters. This is the generator function that creates HRF objects with variable numbers of basis functions, unlike the pre-defined HRF_BSPLINE which has 5 functions.

Usage

```
hrf_bspline_generator(nbasis = 5, span = 24)
```

Arguments

nbasis	Number of basis functions (default: 5)
span	Temporal window in seconds (default: 24)

Value

An HRF object of class `c("BSpline_HRF", "HRF", "function")`

See Also

[HRF_objects](#) for pre-defined HRF objects, [getHRF](#) for a unified interface to create HRFs

Examples

```
# Create B-spline basis with 10 functions
custom_bs <- hrf_bspline_generator(nbasis = 10)
t <- seq(0, 24, by = 0.1)
response <- evaluate(custom_bs, t)
matplot(t, response, type = "l", main = "B-spline HRF with 10 basis functions")
```

`hrf_daguerre_generator`*Create Daguerre HRF Basis Set*

Description

Generates an HRF object using Daguerre spherical basis functions with custom parameters. These are orthogonal polynomials that naturally decay to zero.

Usage

```
hrf_daguerre_generator(nbasis = 3, scale = 4)
```

Arguments

<code>nbasis</code>	Number of basis functions (default: 3)
<code>scale</code>	Scale parameter for the time axis (default: 4)

Details

Daguerre basis functions are orthogonal polynomials on $[0, \text{Inf})$ with respect to the weight function $w(x) = x^2 * \exp(-x)$. They are particularly useful for modeling hemodynamic responses as they naturally decay to zero and can capture various response shapes with few parameters.

Value

An HRF object of class `c("Daguerre_HRF", "HRF", "function")`

See Also

[HRF_objects](#) for pre-defined HRF objects, [getHRF](#) for a unified interface to create HRFs

Examples

```
# Create Daguerre basis with 5 functions
custom_dag <- hrf_daguerre_generator(nbasis = 5, scale = 3)
t <- seq(0, 24, by = 0.1)
response <- evaluate(custom_dag, t)
matplot(t, response, type = "l", main = "Daguerre HRF with 5 basis functions")
```

hrf_fir_generator *Create FIR HRF Basis Set*

Description

Generates an HRF object using Finite Impulse Response (FIR) basis functions with custom parameters. Each basis function represents a time bin with a value of 1 in that bin and 0 elsewhere.

Usage

```
hrf_fir_generator(nbasis = 12, span = 24)
```

Arguments

nbasis	Number of time bins (default: 12)
span	Temporal window in seconds (default: 24)

Details

The FIR basis divides the time window into nbasis equal bins. Each basis function is an indicator function for its corresponding bin. This provides maximum flexibility but requires more parameters than smoother basis sets like B-splines.

Value

An HRF object of class `c("FIR_HRF", "HRF", "function")`

See Also

[HRF_objects](#) for pre-defined HRF objects, [getHRF](#) for a unified interface to create HRFs, [hrf_bspline_generator](#) for a smoother alternative

Examples

```
# Create FIR basis with 20 bins over 30 seconds
custom_fir <- hrf_fir_generator(nbasis = 20, span = 30)
t <- seq(0, 30, by = 0.1)
response <- evaluate(custom_fir, t)
matplot(t, response, type = "l", main = "FIR HRF with 20 time bins")

# Compare to default FIR with 12 bins
default_fir <- HRF_FIR
response_default <- evaluate(default_fir, t[1:241]) # 24 seconds
matplot(t[1:241], response_default, type = "l",
        main = "Default FIR HRF (12 bins over 24s)")
```

hrf_fourier	<i>Fourier basis for HRF modeling</i>
-------------	---------------------------------------

Description

Generates a set of Fourier basis functions (sine and cosine pairs) over a given span.

Usage

```
hrf_fourier(t, span = 24, nbasis = 5)
```

Arguments

t	A vector of time points.
span	The temporal window over which the basis functions span (default: 24).
nbasis	The number of basis functions (default: 5). Should be even for full sine-cosine pairs.

Value

A matrix of Fourier basis functions with nbasis columns.

Examples

```
# Create Fourier basis with 5 functions
t <- seq(0, 24, by = 0.5)
basis <- hrf_fourier(t, span = 24, nbasis = 5)
matplot(t, basis, type = "l", main = "Fourier Basis Functions")
```

hrf_fourier_generator	<i>Create Fourier HRF Basis Set</i>
-----------------------	-------------------------------------

Description

Generates an HRF object using Fourier basis functions (sine and cosine pairs) with custom parameters.

Usage

```
hrf_fourier_generator(nbasis = 5, span = 24)
```

Arguments

nbasis	Number of basis functions (default: 5). Should be even for complete sine-cosine pairs.
span	Temporal window in seconds (default: 24)

Details

The Fourier basis uses alternating sine and cosine functions with increasing frequencies. This provides a smooth, periodic basis set that can capture oscillatory components in the HRF.

Value

An HRF object of class `c("Fourier_HRF", "HRF", "function")`

See Also

[HRF_objects](#) for pre-defined HRF objects, [getHRF](#) for a unified interface to create HRFs

Examples

```
# Create Fourier basis with 8 functions
custom_fourier <- hrf_fourier_generator(nbasis = 8)
t <- seq(0, 24, by = 0.1)
response <- evaluate(custom_fourier, t)
matplot(t, response, type = "l", main = "Fourier HRF with 8 basis functions")
```

hrf_from_coefficients *Combine HRF Basis with Coefficients*

Description

Create a new HRF by linearly weighting the basis functions of an existing HRF. Useful when coefficients have been estimated for an FIR/bspline/SPMG3 basis and one wants a single functional HRF.

Usage

```
hrf_from_coefficients(hrf, h, ...)

## S3 method for class 'HRF'
hrf_from_coefficients(hrf, h, name = NULL, ...)
```

Arguments

hrf	An object of class 'HRF'.
h	Numeric vector of length 'nbasis(hrf)' giving the weights.
...	Reserved for future extensions.
name	Optional name for the resulting HRF.

Value

A new 'HRF' object with 'nbasis = 1'.

Examples

```
# Create a custom HRF from SPMG3 basis coefficients
coeffs <- c(1, 0.2, -0.1) # Main response + slight temporal shift - dispersion
custom_hrf <- hrf_from_coefficients(HRF_SPMG3, coeffs)

# Evaluate the custom HRF
t <- seq(0, 20, by = 0.1)
response <- evaluate(custom_hrf, t)

# Create from FIR basis
fir_coeffs <- c(0, 0.2, 0.5, 1, 0.8, 0.4, 0.1, 0, 0, 0, 0, 0)
custom_fir <- hrf_from_coefficients(HRF_FIR, fir_coeffs)
```

hrf_gamma	<i>Gamma HRF (hemodynamic response function)</i>
-----------	--

Description

The 'hrf_gamma' function computes the gamma density-based HRF (hemodynamic response function) at given time points 't'.

Usage

```
hrf_gamma(t, shape = 6, rate = 1)
```

Arguments

t	A vector of time points.
shape	A numeric value representing the shape parameter for the gamma probability density function. Default value is 6.
rate	A numeric value representing the rate parameter for the gamma probability density function. Default value is 1.

Value

A numeric vector representing the gamma HRF at the given time points 't'.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
# Compute the gamma HRF representation for time points from 0 to 20 with 0.5 increments
hrf_gamma_vals <- hrf_gamma(seq(0, 20, by = .5), shape = 6, rate = 1)
```

hrf_gaussian	<i>Gaussian HRF (hemodynamic response function)</i>
--------------	---

Description

The 'hrf_gaussian' function computes the Gaussian density-based HRF (hemodynamic response function) at given time points 't'.

Usage

```
hrf_gaussian(t, mean = 6, sd = 2)
```

Arguments

t	A vector of time points.
mean	A numeric value representing the mean of the Gaussian probability density function. Default value is 6.
sd	A numeric value representing the standard deviation of the Gaussian probability density function. Default value is 2.

Value

A numeric vector representing the Gaussian HRF at the given time points 't'.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
# Compute the Gaussian HRF representation for time points from 0 to 20 with 0.5 increments
hrf_gaussian_vals <- hrf_gaussian(seq(0, 20, by = .5), mean = 6, sd = 2)
```

hrf_half_cosine	<i>Half-cosine HRF</i>
-----------------	------------------------

Description

Segments: 0->f1 (h1), f1->1 (h2), 1->f2 (h3), f2->0 (h4). Negative f1 gives an initial dip; negative f2 gives an undershoot. Peak is at t = h1 + h2 (amplitude 1 by construction).

Usage

```
hrf_half_cosine(t, h1 = 1, h2 = 5, h3 = 7, h4 = 7, f1 = 0, f2 = 0)
```

Arguments

t	Numeric vector of times (s)
h1, h2, h3, h4	Segment durations (s). Must be > 0.
f1	Initial dip level (default 0), typically in [-0.2, 0]
f2	Undershoot level (default 0), typically in [-0.3, 0]

Value

Numeric vector same length as t

Examples

```
t <- seq(0, 30, by = 0.1)
y <- hrf_half_cosine(t)
```

hrf_inv_logit	<i>hrf_inv_logit</i>
---------------	----------------------

Description

A hemodynamic response function using the difference of two Inverse Logit functions.

Usage

```
hrf_inv_logit(t, mu1 = 6, s1 = 1, mu2 = 16, s2 = 1, lag = 0)
```

Arguments

t	A vector of times.
mu1	The time-to-peak for the rising phase (mean of the first logistic function).
s1	The width (slope) of the first logistic function.
mu2	The time-to-peak for the falling phase (mean of the second logistic function).
s2	The width (slope) of the second logistic function.
lag	The time delay (default: 0).

Value

A vector of the difference of two Inverse Logit HRF values.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
hrf_inv_logit_basis <- hrf_inv_logit(seq(0, 20, by = 0.5), mu1 = 6, s1 = 1, mu2 = 16, s2 = 1)
```

hrf_lwu *Lag-Width-Undershoot (LWU) HRF*

Description

Computes the Lag-Width-Undershoot (LWU) hemodynamic response function. This model uses two Gaussian components to model the main response and an optional undershoot.

Usage

```
hrf_lwu(t, tau = 6, sigma = 2.5, rho = 0.35, normalize = "none")
```

Arguments

t	A numeric vector of time points (in seconds).
tau	Lag of the main Gaussian component (time-to-peak of the positive lobe, in seconds). Default: 6.
sigma	Width (standard deviation) of the main Gaussian component (in seconds). Must be > 0.05. Default: 2.5.
rho	Amplitude of the undershoot Gaussian component, relative to the main component. Must be between 0 and 1.5. Default: 0.35.
normalize	Character string specifying normalization type. Either "none" for no normalization (default) or "height" to scale the HRF so its maximum absolute value is 1.

Details

The LWU model formula combines a positive Gaussian peak and a negative undershoot: $h(t; \tau, \sigma, \rho) = \exp(-(t-\tau)^2/(2*\sigma^2)) - \rho * \exp(-(t-\tau-2*\sigma)^2/(2*(1.6*\sigma)^2))$

Value

A numeric vector representing the LWU HRF values at the given time points 't'.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
t_points <- seq(0, 30, by = 0.1)

# Default LWU HRF
lwu_default <- hrf_lwu(t_points)
plot(t_points, lwu_default, type = "l", main = "LWU HRF (Default Params)", ylab = "Amplitude")
```

```

# LWU HRF with no undershoot
lwu_no_undershoot <- hrf_lwu(t_points, rho = 0)
lines(t_points, lwu_no_undershoot, col = "blue")

# LWU HRF with a wider main peak and larger undershoot
lwu_custom <- hrf_lwu(t_points, tau = 7, sigma = 1.5, rho = 0.5)
lines(t_points, lwu_custom, col = "red")
legend("topright", c("Default", "No Undershoot (rho=0)", "Custom (tau=7, sigma=1.5, rho=0.5)"),
      col = c("black", "blue", "red"), lty = 1, cex = 0.8)

# Height-normalized HRF
lwu_normalized <- hrf_lwu(t_points, tau = 6, sigma = 1, rho = 0.35, normalize = "height")
plot(t_points, lwu_normalized, type = "l", main = "Height-Normalized LWU HRF", ylab = "Amplitude")
abline(h = c(-1, 1), lty = 2, col = "grey") # Max absolute value should be 1

```

hrf_mexhat

*Mexican Hat HRF (hemodynamic response function)***Description**

The ‘hrf_mexhat’ function computes the Mexican hat wavelet-based HRF (hemodynamic response function) at given time points ‘t’.

Usage

```
hrf_mexhat(t, mean = 6, sd = 2)
```

Arguments

t	A vector of time points.
mean	A numeric value representing the mean of the Mexican hat wavelet. Default value is 6.
sd	A numeric value representing the standard deviation of the Mexican hat wavelet. Default value is 2.

Value

A numeric vector representing the Mexican hat wavelet-based HRF at the given time points ‘t’.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```

# Compute the Mexican hat HRF representation for time points from 0 to 20 with 0.5 increments
hrf_mexhat_vals <- hrf_mexhat(seq(0, 20, by = .5), mean = 6, sd = 2)

```

 HRF_objects

Pre-defined Hemodynamic Response Function Objects

Description

A collection of pre-defined HRF objects for common fMRI analysis scenarios. These objects can be used directly in model specifications or as templates for creating custom HRFs.

Usage

HRF_GAMMA(t)

HRF_GAUSSIAN(t)

HRF_SPMG1(t)

HRF_SPMG2(t)

HRF_SPMG3(t)

HRF_BSPLINE(t)

HRF_FIR(t)

Arguments

t Numeric vector of time points (in seconds) at which to evaluate the HRF

Value

When called as functions, return numeric vectors or matrices of HRF values. When used as objects, they are HRF objects with class `c("HRF", "function")`.

Canonical HRFs

HRF_SPMG1 SPM canonical HRF (single basis function)

HRF_SPMG2 SPM canonical HRF with temporal derivative (2 basis functions)

HRF_SPMG3 SPM canonical HRF with temporal and dispersion derivatives (3 basis functions)

HRF_GAMMA Gamma function-based HRF

HRF_GAUSSIAN Gaussian function-based HRF

Flexible Basis Sets

HRF_BSPLINE B-spline basis HRF (5 basis functions)

HRF_FIR Finite Impulse Response (FIR) basis HRF (12 basis functions)

Creating Custom Basis Sets

The pre-defined objects above have fixed numbers of basis functions. To create basis sets with custom parameters (e.g., different numbers of basis functions), use one of these approaches:

Using `getHRF()`:

- `getHRF("fir", nbasis = 20)` - FIR basis with 20 functions
- `getHRF("bspline", nbasis = 10, span = 30)` - B-spline with 10 functions
- `getHRF("fourier", nbasis = 7)` - Fourier basis with 7 functions
- `getHRF("daguerre", nbasis = 5, scale = 3)` - Daguerre basis

Using generator functions directly:

- `hrf_fir_generator(nbasis = 20, span = 30)`
- `hrf_bspline_generator(nbasis = 10, span = 30)`
- `hrf_fourier_generator(nbasis = 7, span = 24)`
- `hrf_daguerre_generator(nbasis = 5, scale = 3)`

Usage

All HRF objects can be:

- Called as functions with time argument: `HRF_SPMG1(t)`
- Used in model specifications: `hrf(condition, basis = HRF_SPMG1)`
- Evaluated with `evaluate()` method
- Combined with decorators like `lag_hrf()` or `block_hrf()`

See Also

[evaluate.HRF](#) for evaluating HRF objects, [gen_hrf](#) for creating HRFs with decorators, [list_available_hrfs](#) for listing all HRF types, [getHRF](#) for creating HRFs by name with custom parameters, [hrf_fir_generator](#), [hrf_bspline_generator](#), [hrf_fourier_generator](#), [hrf_daguerre_generator](#) for creating custom basis sets directly

Other hrf: [deriv\(\)](#), [penalty_matrix\(\)](#)

Examples

```
# Evaluate HRFs at specific time points
times <- seq(0, 20, by = 0.5)

# Single basis canonical HRF
canonical_response <- HRF_SPMG1(times)
plot(times, canonical_response, type = "l", main = "SPM Canonical HRF")

# Multi-basis HRF with derivatives
multi_response <- HRF_SPMG3(times) # Returns 3-column matrix
matplot(times, multi_response, type = "l", main = "SPM HRF with Derivatives")

# Gamma and Gaussian HRFs
```

```

gamma_response <- HRF_GAMMA(times)
gaussian_response <- HRF_GAUSSIAN(times)

# Compare different HRF shapes
plot(times, canonical_response, type = "l", col = "blue",
      main = "HRF Comparison", ylab = "Response")
lines(times, gamma_response, col = "red")
lines(times, gaussian_response, col = "green")
legend("topright", c("SPM Canonical", "Gamma", "Gaussian"),
      col = c("blue", "red", "green"), lty = 1)

# Create custom FIR basis with 20 bins
custom_fir <- getHRF("fir", nbasis = 20, span = 30)
fir_response <- evaluate(custom_fir, times)
matplot(times, fir_response, type = "l", main = "Custom FIR with 20 bins")

# Create custom B-spline basis
custom_bspline <- hrf_bspline_generator(nbasis = 8, span = 25)
bspline_response <- evaluate(custom_bspline, times)
matplot(times, bspline_response, type = "l", main = "Custom B-spline with 8 basis functions")

```

*hrf_sine**hrf_sine*

Description

A hemodynamic response function using the Sine Basis Set.

Usage

```
hrf_sine(t, span = 24, N = 5)
```

Arguments

<code>t</code>	A vector of times.
<code>span</code>	The temporal window over which the basis sets span (default: 24).
<code>N</code>	The number of basis functions (default: 5).

Value

A matrix of sine basis functions.

See Also

Other `hrf_functions`: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_spmg1\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
hrf_sine_basis <- hrf_sine(seq(0, 20, by = 0.5), N = 4)
```

hrf_spmg1	<i>hrf_spmg1</i>
-----------	------------------

Description

A hemodynamic response function based on the SPM canonical double gamma parameterization.

Usage

```
hrf_spmg1(t, P1 = 5, P2 = 15, A1 = 0.0833)
```

Arguments

t	A vector of time points.
P1	The first exponent parameter (default: 5).
P2	The second exponent parameter (default: 15).
A1	Amplitude scaling factor for the positive gamma function component; normally fixed at .0833

Details

This function models the hemodynamic response using the canonical double gamma parameterization in the SPM software. The HRF is defined by a linear combination of two gamma functions with different exponents (P1 and P2) and amplitudes (A1 and A2). It is commonly used in fMRI data analysis to estimate the BOLD (blood-oxygen-level-dependent) signal changes associated with neural activity.

Value

A vector of HRF values at the given time points.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_time\(\)](#), [hrf_weighted\(\)](#)

Examples

```
# Generate a time vector
time_points <- seq(0, 30, by=0.1)
# Compute the HRF values using the SPM canonical double gamma parameterization
hrf_values <- hrf_spmg1(time_points)
# Plot the HRF values
plot(time_points, hrf_values, type='l', main='SPM Canonical Double Gamma HRF')
```

hrf_tent_generator *Create Tent HRF Basis Set*

Description

Generates an HRF object using tent (piecewise linear) basis functions with custom parameters. This generator mirrors HRF_TENT but allows callers to control the number of basis elements and temporal span.

Usage

```
hrf_tent_generator(nbasis = 5, span = 24)
```

Arguments

nbasis	Number of tent basis functions (default: 5)
span	Temporal window in seconds (default: 24)

Value

An HRF object of class `c("Tent_HRF", "HRF", "function")`

See Also

[HRF_objects](#) for pre-defined HRF objects, [getHRF](#) for a unified interface to create HRFs, [hrf_bspline_generator](#) for a smoother alternative

Examples

```
# Create a tent basis with 6 functions over a 20 second window
custom_tent <- hrf_tent_generator(nbasis = 6, span = 20)
t <- seq(0, 20, by = 0.1)
response <- evaluate(custom_tent, t)
matplot(t, response, type = "l", main = "Tent HRF with 6 basis functions")
```

hrf_time *HRF (hemodynamic response function) as a linear function of time*

Description

The 'hrf_time' function computes the value of an HRF, which is a simple linear function of time 't', when 't' is greater than 0 and less than 'maxt'.

Usage

```
hrf_time(t, maxt = 22)
```

Arguments

t	A numeric value representing time in seconds.
maxt	A numeric value representing the maximum time point in the domain. Default value is 22.

Value

A numeric value representing the value of the HRF at the given time 't'.

See Also

Other hrf_functions: [hrf_basis_lwu\(\)](#), [hrf_boxcar\(\)](#), [hrf_bspline\(\)](#), [hrf_gamma\(\)](#), [hrf_gaussian\(\)](#), [hrf_inv_logit\(\)](#), [hrf_lwu\(\)](#), [hrf_mexhat\(\)](#), [hrf_sine\(\)](#), [hrf_spmg1\(\)](#), [hrf_weighted\(\)](#)

Examples

```
# Compute the HRF value for t = 5 seconds with the default maximum time
hrf_val <- hrf_time(5)

# Compute the HRF value for t = 5 seconds with a custom maximum time of 30 seconds
hrf_val_custom_maxt <- hrf_time(5, maxt = 30)
```

hrf_toeplitz	<i>HRF Toeplitz Matrix</i>
--------------	----------------------------

Description

Create a Toeplitz matrix for hemodynamic response function (HRF) convolution.

Usage

```
hrf_toeplitz(hrf, time, len, sparse = FALSE)
```

Arguments

hrf	The hemodynamic response function.
time	A numeric vector representing the time points.
len	The length of the output Toeplitz matrix.
sparse	Logical, if TRUE, the output Toeplitz matrix is returned as a sparse matrix (default: FALSE).

Value

A Toeplitz matrix for HRF convolution.

Examples

```
# Create HRF and time points
hrf_fun <- function(t) hrf_spmg1(t)
times <- seq(0, 30, by = 1)

# Create Toeplitz matrix
H <- hrf_toeplitz(hrf_fun, times, len = 50)

# Create sparse version
H_sparse <- hrf_toeplitz(hrf_fun, times, len = 50, sparse = TRUE)
```

hrf_weighted	<i>Weighted HRF (No Hemodynamic Delay)</i>
--------------	--

Description

Creates a flexible weighted HRF starting at t=0 with user-specified weights. Unlike traditional HRFs, this has no built-in hemodynamic delay - it directly maps weights to time points, allowing for arbitrary temporal response shapes.

Usage

```
hrf_weighted(
  weights,
  width = NULL,
  times = NULL,
  method = c("constant", "linear"),
  normalize = FALSE
)
```

Arguments

weights	Numeric vector of weights. Required.
width	Total duration of the window in seconds. If provided without times, weights are evenly spaced from 0 to width.
times	Numeric vector of time points (in seconds, relative to t=0) where weights are specified. Must be strictly increasing and start at 0 for consistency with other HRFs. If provided, width is ignored.
method	Interpolation method between time points: "constant" Step function - weight is constant until the next time point (default). Good for discrete time bins. "linear" Linear interpolation between points. Good for smooth weight transitions.
normalize	Logical; if TRUE, weights are scaled so they sum to 1 (for method = "constant") or integrate to 1 (for method = "linear"). This makes the regression coefficient interpretable as a weighted mean. Default is FALSE.

Details

This is useful for extracting weighted averages of data at specific time points. When `normalize = TRUE` and the HRF is used in a GLM, the estimated coefficient represents a weighted mean of the data at the specified times.

There are two ways to specify the temporal structure:

1. `width + weights`: Weights are evenly spaced from 0 to `width`
2. `times + weights`: Explicit time points for each weight (relative to `t=0`)

For delayed windows (not starting at `t=0`), use `lag_hrf` to shift the weighted HRF in time.

Value

An HRF object that can be used with `regressor()` and other `fmrihrf` functions.

Note on durations

The temporal structure (`width` or `times`) is fixed when the HRF is created. The `duration` parameter in `regressor()` does **not** modify the weighted HRF's structure—it controls how long the neural input is sustained (which then gets convolved with this HRF). For trial-varying weighted HRFs, use a list of HRFs:

```
hrf_early <- hrf_weighted(width = 6, weights = c(1, 1, 0, 0), normalize = TRUE)
hrf_late <- hrf_weighted(width = 6, weights = c(0, 0, 1, 1), normalize = TRUE)
reg <- regressor(onsets = c(0, 20), hrf = list(hrf_early, hrf_late))
```

See Also

`hrf_boxcar` for simple uniform boxcars, `lag_hrf` to shift the window in time, `empirical_hrf` for HRFs from measured data

Other `hrf_functions`: `hrf_basis_lwu()`, `hrf_boxcar()`, `hrf_bspline()`, `hrf_gamma()`, `hrf_gaussian()`, `hrf_inv_logit()`, `hrf_lwu()`, `hrf_mexhat()`, `hrf_sine()`, `hrf_spmg1()`, `hrf_time()`

Examples

```
# Simple: 6s window with 4 evenly-spaced weights (at 0, 2, 4, 6s)
hrf1 <- hrf_weighted(width = 6, weights = c(0.2, 0.5, 0.8, 0.3))
t <- seq(-1, 10, by = 0.1)
plot(t, evaluate(hrf1, t), type = "s", main = "Weighted HRF (width + weights)")

# Explicit times for precise control
hrf2 <- hrf_weighted(
  times = c(0, 1, 3, 5, 6),
  weights = c(0.1, 0.5, 0.8, 0.5, 0.1),
  method = "linear"
)
plot(t, evaluate(hrf2, t), type = "l", main = "Smooth Weighted HRF")

# Normalized weights - coefficient estimates weighted mean of signal
hrf3 <- hrf_weighted(
```

```
width = 8,
weights = c(1, 2, 2, 1),
normalize = TRUE
)

# Trial-varying weighted HRFs
hrf_early <- hrf_weighted(width = 6, weights = c(1, 1, 0, 0), normalize = TRUE)
hrf_late <- hrf_weighted(width = 6, weights = c(0, 0, 1, 1), normalize = TRUE)
reg <- regressor(onsets = c(0, 20), hrf = list(hrf_early, hrf_late))

# For delayed windows, use lag_hrf
hrf_delayed <- lag_hrf(hrf_weighted(width = 5, weights = c(1, 2, 1)), lag = 10)
```

lag_hrf

Lag an HRF Object

Description

Creates a new HRF object by applying a temporal lag to an existing HRF object.

Usage

```
lag_hrf(hrf, lag)
```

Arguments

hrf	The HRF object (of class 'HRF') to lag.
lag	The time lag in seconds to apply. Positive values shift the response later in time.

Value

A new HRF object representing the lagged function.

See Also

Other HRF_decorator_functions: [block_hrf\(\)](#), [normalise_hrf\(\)](#)

Examples

```
lagged_spmg1 <- lag_hrf(HRF_SPMG1, 5)
# Evaluate at time 10; equivalent to HRF_SPMG1(10 - 5)
lagged_spmg1(10)
HRF_SPMG1(5)
```

list_available_hrfs *List all available hemodynamic response functions (HRFs)*

Description

Reads the internal HRF registry to list available HRF types.

Usage

```
list_available_hrfs(details = FALSE)
```

Arguments

details Logical; if TRUE, attempt to add descriptions (basic for now).

Value

A data frame with columns: name, type (object/generator), nbasis_default.

Examples

```
# List all available HRFs
hrfs <- list_available_hrfs()
print(hrfs)

# List with details
hrfs_detailed <- list_available_hrfs(details = TRUE)
print(hrfs_detailed)
```

make_hrf *Create an HRF from a basis specification*

Description

'make_hrf' resolves a basis specification to an 'HRF' object and applies an optional temporal lag. The basis may be given as the name of a built-in HRF, as a generating function, or as an existing 'HRF' object.

Usage

```
make_hrf(basis, lag, nbasis = 1)
```

Arguments

basis	Character name of a built-in HRF, a function that generates HRF values, or an object of class 'HRF'.
lag	Numeric scalar giving the shift in seconds applied to the HRF.
nbasis	Integer specifying the number of basis functions when 'basis' is provided as a name.

Value

An object of class 'HRF' representing the lagged basis.

Examples

```
# Canonical SPM HRF delayed by 2 seconds
h <- make_hrf("spm1", lag = 2)
h(0:5)
```

nbasis	<i>Number of basis functions</i>
--------	----------------------------------

Description

Return the number of basis functions represented by an object.

Usage

```
nbasis(x, ...)
```

```
## S3 method for class 'HRF'
nbasis(x, ...)
```

```
## S3 method for class 'Reg'
nbasis(x, ...)
```

Arguments

x	Object containing HRF or regressor information.
...	Additional arguments passed to methods.

Details

This information is typically used when constructing penalty matrices or understanding the complexity of an HRF model or regressor.

Value

Integer scalar giving the number of basis functions.

Examples

```
# Number of basis functions for different HRF types
nbasis(HRF_SPMG1) # 1 basis function
nbasis(HRF_SPMG3) # 3 basis functions (canonical + 2 derivatives)
nbasis(HRF_BSPLINE) # 5 basis functions (default)

# For a regressor
reg <- regressor(onsets = c(10, 30, 50), hrf = HRF_SPMG3)
nbasis(reg) # 3 (inherits from the HRF)
```

neural_input

Generate Neural Input Function from Event Timing

Description

Converts event timing information into a neural input function representing the underlying neural activity before HRF convolution. This function is useful for:

Usage

```
neural_input(x, ...)

## S3 method for class 'Reg'
neural_input(x, start = 0, end = NULL, resolution = 0.33, ...)
```

Arguments

x	A regressor object containing event timing information
...	Additional arguments passed to methods
start	Numeric; start time of the input function
end	Numeric; end time of the input function
resolution	Numeric; temporal resolution in seconds (default: 0.33)

Details

stimulus Creating stimulus functions for fMRI analysis
modeling Modeling sustained vs. transient neural activity
inputs Generating inputs for HRF convolution
visualization Visualizing the temporal structure of experimental designs

Value

A list containing:

time Numeric vector of time points
neural_input Numeric vector of input amplitudes at each time point

See Also

[regressor](#), [evaluate.Reg](#), [HRF_SPMG1](#)

Examples

```
# Create a regressor with multiple events
reg <- regressor(
  onsets = c(10, 30, 50),
  duration = c(2, 2, 2),
  amplitude = c(1, 1.5, 0.8),
  hrf = HRF_SPMG1
)

# Generate neural input function
input <- neural_input(reg, start = 0, end = 60, resolution = 0.5)

# Plot the neural input function
plot(input$time, input$neural_input, type = "l",
      xlab = "Time (s)", ylab = "Neural Input",
      main = "Neural Input Function")

# Create regressor with varying durations
reg_sustained <- regressor(
  onsets = c(10, 30),
  duration = c(5, 10), # sustained activity
  amplitude = c(1, 1),
  hrf = HRF_SPMG1
)

# Generate and compare neural inputs
input_sustained <- neural_input(
  reg_sustained,
  start = 0,
  end = 60,
  resolution = 0.5
)
```

normalise_hrf

Normalise an HRF Object

Description

Creates a new HRF object whose output is scaled such that the maximum absolute value of the response is 1.

Usage

```
normalise_hrf(hrf)
```

Arguments

hrf The HRF object (of class 'HRF') to normalise.

Details

For multi-basis HRFs, each basis function (column) is normalised independently.

Value

A new HRF object representing the normalised function.

See Also

Other HRF_decorator_functions: [block_hrf\(\)](#), [lag_hrf\(\)](#)

Examples

```
# Create a gaussian HRF with a peak value != 1
gauss_unnorm <- as_hrf(function(t) 5 * dnorm(t, 6, 2), name="unnorm_gauss")
# Normalise it
gauss_norm <- normalise_hrf(gauss_unnorm)
t_vals <- seq(0, 20, by = 0.1)
max(gauss_unnorm(t_vals)) # Peak is > 1
max(gauss_norm(t_vals))  # Peak is 1
```

onsets

Get event onsets from an object

Description

Generic accessor returning event onset times in seconds.

Usage

```
onsets(x, ...)
```

```
## S3 method for class 'Reg'
onsets(x, ...)
```

Arguments

x Object containing onset information
... Additional arguments passed to methods

Value

Numeric vector of onsets

Examples

```
# Create a regressor with event onsets
reg <- regressor(onsets = c(1, 5, 10, 15), hrf = HRF_SPMG1, span = 20)
onsets(reg)
```

penalty_matrix	<i>Generate penalty matrix for regularization</i>
----------------	---

Description

Generate a penalty matrix for regularizing HRF basis coefficients. The penalty matrix encodes shape priors that discourage implausible or overly wiggly HRF estimates. Different HRF types use different penalty structures:

- FIR/B-spline/Tent bases: Roughness penalties based on discrete derivatives
- SPM canonical + derivatives: Differential shrinkage of derivative terms
- Fourier bases: Penalties on high-frequency components
- Daguerre bases: Increasing weights on higher-order terms
- Default: Identity matrix (ridge penalty)

Usage

```
penalty_matrix(x, ...)

## S3 method for class 'HRF'
penalty_matrix(x, order = 2, ...)

## S3 method for class 'BSpline_HRF'
penalty_matrix(x, order = 2, ...)

## S3 method for class 'Tent_HRF'
penalty_matrix(x, order = 2, ...)

## S3 method for class 'FIR_HRF'
penalty_matrix(x, order = 2, ...)

## S3 method for class 'SPMG2_HRF'
penalty_matrix(x, order = 2, shrink_deriv = 2, ...)

## S3 method for class 'SPMG3_HRF'
penalty_matrix(x, order = 2, shrink_deriv = 2, ...)

## S3 method for class 'Fourier_HRF'
penalty_matrix(x, order = 2, ...)

## S3 method for class 'Daguerre_HRF'
penalty_matrix(x, order = 2, ...)
```

Arguments

x	The HRF object or basis specification
...	Additional arguments passed to specific methods
order	Integer specifying the order of the penalty (default: 2)
shrink_deriv	Numeric; penalty weight for derivative terms in SPMG2/SPMG3 bases (default: 2)

Details

The penalty matrix R is used in regularized estimation as $\lambda * h^T R h$, where h are the basis coefficients and λ is the regularization parameter. Well-designed penalty matrices can significantly improve HRF estimation by encoding smoothness or other shape constraints.

Value

A symmetric positive definite penalty matrix of dimension $n_{\text{basis}}(x) \times n_{\text{basis}}(x)$

See Also

[nbasis()], [HRF_objects]

Other hrf: [HRF_objects](#), [deriv\(\)](#)

Examples

```
# FIR basis with smoothness penalty
fir_hrf <- HRF_FIR
R_fir <- penalty_matrix(fir_hrf)

# B-spline basis with second-order smoothness
bspline_hrf <- HRF_BSPLINE
R_bspline <- penalty_matrix(bspline_hrf, order = 2)

# SPM canonical with derivative shrinkage
spm3_hrf <- HRF_SPMG3
R_spm3 <- penalty_matrix(spm3_hrf, shrink_deriv = 4)
```

plot.HRF

Plot an HRF Object

Description

Creates a visualization of an HRF object. For single-basis HRFs, shows the response curve with peak annotation. For multi-basis HRFs (e.g., HRF_SPMG3), shows all basis functions on the same plot.

Usage

```
## S3 method for class 'HRF'
plot(x, time = NULL, normalize = FALSE, show_peak = TRUE, ...)
```

Arguments

x	An HRF object
time	Numeric vector of time points. If NULL (default), uses seq(0, span, by = 0.1) where span is the HRF's span attribute.
normalize	Logical; if TRUE, normalize responses to peak at 1. Default is FALSE.
show_peak	Logical; if TRUE (default for single-basis HRFs), annotate the peak time and amplitude on the plot.
...	Additional arguments passed to underlying plot functions.

Value

Invisibly returns a data frame with the time and response values (useful for further customization).

Examples

```
# Plot single-basis HRF
plot(HRF_SPMG1)

# Plot multi-basis HRF
plot(HRF_SPMG3)

# Plot with normalization
plot(HRF_GAMMA, normalize = TRUE)

# Custom time range
plot(HRF_SPMG1, time = seq(0, 30, by = 0.5))
```

plot.Reg

Plot a Regressor Object

Description

Creates a visualization of a regressor object showing the predicted BOLD response over time. Optionally displays event onsets as vertical lines.

Usage

```
## S3 method for class 'Reg'
plot(
  x,
  grid = NULL,
  show_onsets = TRUE,
```

```

    onset_color = "red",
    onset_alpha = 0.5,
    precision = 0.33,
    ...
)

```

Arguments

x	A 'Reg' object created by 'regressor()'.
grid	Numeric vector of time points for evaluation. If NULL (default), automatically generates a grid from 0 to max(onsets) + span with step 0.5s.
show_onsets	Logical; if TRUE (default), show vertical dashed lines at event onset times.
onset_color	Color for onset lines. Default is "red".
onset_alpha	Alpha transparency for onset lines. Default is 0.5.
precision	Numeric sampling precision for HRF evaluation. Default is 0.33.
...	Additional arguments passed to underlying plot functions.

Value

Invisibly returns a data frame with the time and response values.

Examples

```

# Create and plot a simple regressor
reg <- regressor(onsets = c(10, 30, 50), hrf = HRF_SPMG1)
plot(reg)

# Plot with custom time grid
plot(reg, grid = seq(0, 80, by = 1))

# Plot without onset markers
plot(reg, show_onsets = FALSE)
)

```

plot_hrf
Compare Multiple HRF Functions

Description

Creates a comparison plot of multiple HRF objects. This function provides a convenient way to visualize different HRFs on the same plot, with options for normalization and customization. Uses ggplot2 if available for publication-quality figures, otherwise falls back to base R graphics.

Usage

```
plot_hrfs(
  ...,
  time = NULL,
  normalize = FALSE,
  labels = NULL,
  title = NULL,
  subtitle = NULL,
  use_ggplot = TRUE
)
```

Arguments

...	HRF objects to compare. Can be passed as individual arguments or as a named list.
time	Numeric vector of time points. If NULL (default), uses <code>seq(0, max_span, by = 0.1)</code> where <code>max_span</code> is the maximum span across all HRFs.
normalize	Logical; if TRUE, normalize all HRFs to peak at 1. Useful for comparing shapes regardless of amplitude. Default is FALSE.
labels	Character vector of labels for each HRF. If NULL (default), uses the 'name' attribute of each HRF, or "HRF_1", "HRF_2", etc.
title	Character string for the plot title. If NULL (default), uses "HRF Comparison".
subtitle	Character string for the plot subtitle. If NULL (default), no subtitle is shown.
use_ggplot	Logical; if TRUE and <code>ggplot2</code> is available, use <code>ggplot2</code> for plotting. If FALSE, use base R graphics. Default is TRUE.

Value

Invisibly returns a data frame in long format with columns 'time', 'HRF', and 'response'. If `use_ggplot` is TRUE and `ggplot2` is available, also returns a `ggplot` object as an attribute 'plot'.

Examples

```
# Compare canonical HRFs
plot_hrfs(HRF_SPMG1, HRF_GAMMA, HRF_GAUSSIAN)

# Compare with custom labels
plot_hrfs(HRF_SPMG1, HRF_GAMMA,
  labels = c("SPM Canonical", "Gamma"))

# Normalize for shape comparison
plot_hrfs(HRF_SPMG1, HRF_GAMMA, HRF_GAUSSIAN,
  normalize = TRUE,
  title = "HRF Shape Comparison",
  subtitle = "All HRFs normalized to peak at 1")

# Compare blocked HRFs with different durations
hrf_1s <- block_hrf(HRF_SPMG1, width = 1)
```

```

hrf_3s <- block_hrf(HRF_SPMG1, width = 3)
hrf_5s <- block_hrf(HRF_SPMG1, width = 5)
plot_hrfs(hrf_1s, hrf_3s, hrf_5s,
          labels = c("1s duration", "3s duration", "5s duration"),
          title = "Effect of Event Duration on HRF")

# Use base R graphics instead of ggplot2
plot_hrfs(HRF_SPMG1, HRF_GAMMA, use_ggplot = FALSE)

```

plot_regressors

Compare Multiple Regressor Objects

Description

Creates a comparison plot of multiple regressor objects. This function provides a convenient way to visualize different regressors on the same plot, with options for showing event onsets and customization. Uses ggplot2 if available for publication-quality figures, otherwise falls back to base R graphics.

Usage

```

plot_regressors(
  ...,
  grid = NULL,
  labels = NULL,
  title = NULL,
  subtitle = NULL,
  show_onsets = "first",
  onset_alpha = 0.3,
  precision = 0.33,
  use_ggplot = TRUE
)

```

Arguments

...	Regressor objects to compare. Can be passed as individual arguments or as a named list.
grid	Numeric vector of time points for evaluation. If NULL (default), automatically generates a grid covering all regressors.
labels	Character vector of labels for each regressor. If NULL (default), uses "Regressor_1", "Regressor_2", etc.
title	Character string for the plot title. If NULL (default), uses "Regressor Comparison".
subtitle	Character string for the plot subtitle. If NULL, no subtitle.
show_onsets	Logical or character. If TRUE, show onset lines for all regressors. If "first", show only for the first regressor. If FALSE, hide onsets. Default is "first".

<code>onset_alpha</code>	Alpha transparency for onset lines. Default is 0.3.
<code>precision</code>	Numeric sampling precision for HRF evaluation. Default is 0.33.
<code>use_ggplot</code>	Logical; if TRUE and <code>ggplot2</code> is available, use <code>ggplot2</code> for plotting. If FALSE, use base R graphics. Default is TRUE.

Value

Invisibly returns a data frame in long format with columns `'time'`, `'Regressor'`, and `'response'`.

Examples

```
# Create regressors with different HRFs
onsets <- c(10, 30, 50)
reg1 <- regressor(onsets, HRF_SPMG1)
reg2 <- regressor(onsets, HRF_GAMMA)
reg3 <- regressor(onsets, HRF_GAUSSIAN)

# Compare regressors
plot_regressors(reg1, reg2, reg3,
                labels = c("SPM Canonical", "Gamma", "Gaussian"))

# Compare regressors with different event timings
reg_fast <- regressor(seq(0, 60, by = 10), HRF_SPMG1)
reg_slow <- regressor(seq(0, 60, by = 20), HRF_SPMG1)
plot_regressors(reg_fast, reg_slow,
                labels = c("Fast (10s ISI)", "Slow (20s ISI)"),
                title = "Effect of Inter-Stimulus Interval")

# Compare original vs shifted regressor
reg_orig <- regressor(c(10, 30, 50), HRF_SPMG1)
reg_shifted <- shift(reg_orig, 5)
plot_regressors(reg_orig, reg_shifted,
                labels = c("Original", "Shifted +5s"))
```

`print.HRF`

Print an HRF Object

Description

Displays a concise summary of an HRF object including its name, number of basis functions, temporal span, and parameters (if any).

Usage

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

Arguments

x	An HRF object
...	Additional arguments (unused)

Value

Invisibly returns the HRF object

Examples

```
# Print canonical HRF
print(HRF_SPMG1)

# Print multi-basis HRF
print(HRF_SPMG3)

# Print Gaussian HRF
print(HRF_GAUSSIAN)
```

print.Reg	<i>Print method for Reg objects</i>
-----------	-------------------------------------

Description

Provides a concise summary of the regressor object using the cli package.

Usage

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

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

Arguments

x	A 'Reg' object.
...	Not used.

Value

No return value, called for side effects (prints to console)

Examples

```
r <- regressor(onsets = c(1, 10, 20), hrf = HRF_SPMG1,
              duration = 0, amplitude = 1,
              span = 40)
print(r)
```

reconstruction_matrix *Combine HRF Basis with Coefficients*

Description

Create a new HRF by linearly weighting the basis functions of an existing HRF. This is useful for turning estimated basis coefficients into a single functional HRF.

S3 method for 'HRF' objects that returns a matrix mapping basis coefficients to sampled HRF values at the provided time grid. For single-basis HRFs, this returns a one-column matrix. For multi-basis HRFs (e.g., SPMG2/SPMG3, FIR, B-spline), this returns a matrix with one column per basis function.

Usage

```
reconstruction_matrix(hrf, sframe, ...)

## S3 method for class 'HRF'
reconstruction_matrix(hrf, sframe, ...)
```

Arguments

hrf	An object of class 'HRF'.
sframe	A numeric vector of times, or a 'sampling_frame' object from which times are extracted via 'samples()'.
...	Additional arguments passed to 'samples()' when 'sframe' is a 'sampling_frame', and to 'evaluate()' for HRF evaluation.

Details

Reconstruction matrix for an HRF basis

Returns a matrix Φ that converts basis coefficients into a sampled HRF shape.

Value

A numeric matrix with one column per basis function.

A numeric matrix of dimension 'length(times) x nbasis(hrf)'.

Examples

```
# Create reconstruction matrix for basis functions
hrf <- HRF_SPMG2 # 2-basis HRF
times <- seq(0, 20, by = 0.5)
rmat <- reconstruction_matrix(hrf, times)
dim(rmat) # Shows dimensions
```

regressor

*Construct a Regressor Object***Description**

Creates an object representing event-related regressors for fMRI modeling. This function defines event onsets and associates them with a hemodynamic response function (HRF) to generate predicted time courses.

Usage

```
regressor(
    onsets,
    hrf = HRF_SPMG1,
    duration = 0,
    amplitude = 1,
    span = 40,
    summate = TRUE
)
```

Arguments

onsets	A numeric vector of event onset times in seconds.
hrf	The hemodynamic response function (HRF) to convolve with the events. This can be: <ul style="list-style-type: none"> • A pre-defined ‘HRF’ object (e.g., ‘HRF_SPMG1’) • A custom ‘HRF’ object created with ‘as_hrf’ • A function ‘f(t)’ • A character string referring to a known HRF type (e.g., "spmgl", "gaussian") • A list of HRF objects for trial-varying HRFs (one per event, or length 1 to recycle) Defaults to ‘HRF_SPMG1’.
duration	A numeric scalar or vector specifying the duration of each event in seconds. If scalar, it’s applied to all events. Defaults to 0 (impulse events).
amplitude	A numeric scalar or vector specifying the amplitude (scaling factor) for each event. If scalar, it’s applied to all events. Defaults to 1.
span	The temporal window (in seconds) over which the HRF is defined or evaluated. This influences the length of the convolution. If not provided, it may be inferred from the ‘hrf’ object or default to 40s. For list HRFs, the maximum span across all HRFs is used. Note: Unlike some previous versions, the ‘span’ is not automatically adjusted based on ‘duration’; ensure the provided or inferred ‘span’ is sufficient for your longest event duration.

summate Logical scalar; if 'TRUE' (default), the HRF response amplitude scales with the duration of sustained events (via weighted integration). If 'FALSE', weighted integration is normalized by total block weight so amplitude does not grow with duration.

Details

This function serves as the main public interface for creating regressor objects. Internally, it utilizes the 'Reg()' constructor which performs validation and efficient storage. The resulting object can be evaluated at specific time points using the 'evaluate()' function.

Events with an amplitude of 0 are automatically filtered out.

Trial-Varying HRFs

When 'hrf' is a list of HRF objects, each event can have its own HRF. This is useful for trial-wise analyses where different events may have different temporal characteristics (e.g., different boxcar windows, different weights). The list must have either length 1 (recycled to all events) or length equal to the number of onsets.

Value

An S3 object of class 'Reg' and 'list' containing processed event information and the HRF specification. The object includes a 'filtered_all' attribute indicating whether all events were removed due to zero or 'NA' amplitudes, and an 'hrf_is_list' attribute indicating whether trial-varying HRFs are used.

Examples

```
# Create a simple regressor with 3 events
reg <- regressor(onsets = c(10, 30, 50), hrf = HRF_SPMG1)

# Regressor with durations and amplitudes
reg2 <- regressor(
  onsets = c(10, 30, 50),
  duration = c(2, 2, 2),
  amplitude = c(1, 1.5, 0.8),
  hrf = HRF_SPMG1
)

# Using different HRF types
reg_gamma <- regressor(onsets = c(10, 30), hrf = "gamma")

# Evaluate regressor at specific time points
times <- seq(0, 60, by = 0.1)
response <- evaluate(reg, times)

# Trial-varying HRFs: different boxcar windows for each event
hrf1 <- hrf_boxcar(width = 4, normalize = TRUE)
hrf2 <- hrf_boxcar(width = 6, normalize = TRUE)
reg_varying <- regressor(
  onsets = c(10, 30),
  hrf = list(hrf1, hrf2)
```

```
)
```

```
regressor_design
```

```
Build a Design Matrix from Block-wise Onsets
```

Description

'regressor_design' extends [regressor_set()] by allowing onsets to be specified relative to individual blocks and by directly returning the evaluated design matrix.

Usage

```
regressor_design(
  onsets,
  fac,
  block,
  sframe,
  hrf = HRF_SPMG1,
  duration = 0,
  amplitude = 1,
  span = 40,
  precision = 0.33,
  method = c("conv", "fft", "Rconv", "loop"),
  sparse = FALSE,
  summate = TRUE
)
```

Arguments

onsets	Numeric vector of event onset times, expressed relative to the start of their corresponding block.
fac	A factor (or object coercible to a factor) indicating the condition for each onset.
block	Integer vector identifying the block for each onset. Values must be valid block indices for 'sframe'.
sframe	A [sampling_frame] describing the temporal structure of the experiment.
hrf	Hemodynamic response function shared by all conditions.
duration	Numeric scalar or vector of event durations.
amplitude	Numeric scalar or vector of event amplitudes.
span	Numeric scalar giving the HRF span in seconds.
precision	Numeric precision used during convolution.
method	Evaluation method passed to [evaluate()].
sparse	Logical; if 'TRUE' a sparse design matrix is returned.
summate	Logical; passed to [regressor()].

Value

A numeric matrix (or sparse matrix) with one column per factor level and one row per sample defined by 'sframe'.

Examples

```
# Create a sampling frame for 2 blocks, 100 scans each, TR=2
sframe <- sampling_frame(blocklens = c(100, 100), TR = 2)

# Events in block-relative time
onsets <- c(10, 30, 50, 20, 40, 60)
conditions <- factor(c("A", "B", "A", "B", "A", "B"))
blocks <- c(1, 1, 1, 2, 2, 2)

# Build design matrix
design <- regressor_design(
  onsets = onsets,
  fac = conditions,
  block = blocks,
  sframe = sframe,
  hrf = HRF_SPMG1
)

# Design matrix has 200 rows (total scans) and 2 columns (conditions)
dim(design)
```

regressor_set

Construct a Regressor Set

Description

Creates a set of regressors, one for each level of a factor. Each condition shares the same HRF and other parameters but has distinct onsets, durations and amplitudes.

Usage

```
regressor_set(
  onsets,
  fac,
  hrf = HRF_SPMG1,
  duration = 0,
  amplitude = 1,
  span = 40,
  summate = TRUE
)

## S3 method for class 'RegSet'
evaluate(
```

```

    x,
    grid,
    precision = 0.33,
    method = c("conv", "fft", "Rconv", "loop"),
    sparse = FALSE,
    ...
)

```

Arguments

onsets	Numeric vector of event onset times.
fac	A factor (or object coercible to a factor) indicating the condition for each onset.
hrf	Hemodynamic response function used for all conditions.
duration	Numeric scalar or vector of event durations.
amplitude	Numeric scalar or vector of event amplitudes.
span	Numeric scalar giving the HRF span in seconds.
summate	Logical; passed to [regressor()].
x	A RegSet object
grid	Numeric vector of time points at which to evaluate
precision	Numeric precision for evaluation
method	Evaluation method
sparse	Logical whether to return sparse matrix
...	Additional arguments passed to evaluate

Value

An object of class ‘RegSet’ containing one ‘Reg’ per factor level.

Examples

```

# Create events for 3 conditions
onsets <- c(10, 20, 30, 40, 50, 60)
conditions <- factor(c("A", "B", "C", "A", "B", "C"))

# Create regressor set
rset <- regressor_set(onsets, conditions, hrf = HRF_SPMG1)

# With durations and amplitudes
rset2 <- regressor_set(
  onsets = onsets,
  fac = conditions,
  duration = 2,
  amplitude = c(1, 1.5, 0.8, 1, 1.5, 0.8),
  hrf = HRF_SPMG1
)

# Evaluate the regressor set

```

```
times <- seq(0, 80, by = 0.1)
design_matrix <- evaluate(rset, times)
```

samples

Get sample acquisition times

Description

Generic function retrieving sampling times from a sampling frame or related object.

Usage

```
samples(x, ...)

## S3 method for class 'sampling_frame'
samples(x, blockids = NULL, global = FALSE, ...)
```

Arguments

x	Object describing the sampling grid
...	Additional arguments passed to methods
blockids	Integer vector of block identifiers to include (default: all blocks)
global	Logical indicating whether to return global times (default: FALSE)

Value

Numeric vector of sample times

Examples

```
# Get sample times from a sampling frame
sframe <- sampling_frame(blocklens = c(100, 120), TR = 2)
samples(sframe, blockids = 1) # First block only
samples(sframe, global = TRUE) # All blocks, global timing
```

sampling_frame	<i>A sampling_frame describes the block structure and temporal sampling of an fMRI paradigm.</i>
----------------	--

Description

A `sampling_frame` describes the block structure and temporal sampling of an fMRI paradigm.

Usage

```
sampling_frame(blocklens, TR, start_time = TR/2, precision = 0.1)
```

Arguments

<code>blocklens</code>	A numeric vector representing the number of scans in each block.
<code>TR</code>	A numeric value or vector representing the repetition time in seconds (i.e., the spacing between consecutive image acquisitions). When a vector is provided, its length must be 1 or equal to the number of blocks.
<code>start_time</code>	A numeric value or vector representing the offset of the first scan of each block (default is $TR/2$). When a vector is provided, its length must be 1 or equal to the number of blocks.
<code>precision</code>	A numeric value representing the discrete sampling interval used for convolution with the hemodynamic response function (default is 0.1).

Value

A list with class "sampling_frame" describing the block structure and temporal sampling of an fMRI paradigm.

Examples

```
frame <- sampling_frame(blocklens = c(100, 100, 100), TR = 2, precision = 0.5)

# The relative time (with respect to the last block) in seconds of each sample/acquisition
sam <- samples(frame)
# The global time (with respect to the first block) of each sample/acquisition
gsam <- samples(frame, global = TRUE)

# Block identifiers for each acquisition can be retrieved using
# blockids(frame)
```

shift	<i>Shift a time series object</i>
-------	-----------------------------------

Description

Apply a temporal shift to a time series object. This function shifts the values in time while preserving the structure of the object. Common uses include:

alignment Aligning regressors with different temporal offsets

derivatives Applying temporal derivatives to time series

correction Correcting for timing differences between signals

Usage

```
shift(x, ...)
```

```
## S3 method for class 'Reg'  
shift(x, shift_amount, ...)
```

Arguments

x	An object representing a time series or a time-based data structure
...	Additional arguments passed to methods
shift_amount	Numeric; amount to shift by (positive = forward, negative = backward)

Value

An object of the same class as the input, with values shifted in time:

Values Values are moved by the specified offset

Structure Object structure and dimensions are preserved

Padding Empty regions are filled with padding value

See Also

[regressor()], [evaluate()]

Examples

```
# Create a simple time series with events  
event_data <- data.frame(  
  onsets = c(1, 10, 20, 30),  
  run = c(1, 1, 1, 1)  
)  
  
# Create regressor from events  
reg <- regressor(  

```

```

    onsets = event_data$onsets,
    hrf = HRF_SPMG1,
    duration = 0,
    amplitude = 1
  )

# Shift regressor forward by 2 seconds
reg_forward <- shift(reg, shift_amount = 2)

# Shift regressor backward by 1 second
reg_backward <- shift(reg, shift_amount = -1)

# Evaluate original and shifted regressors
times <- seq(0, 50, by = 2)
orig_values <- evaluate(reg, times)
shifted_values <- evaluate(reg_forward, times)

```

single_trial_regressor

Create a single trial regressor

Description

Creates a regressor object for modeling a single trial event in an fMRI experiment. This is particularly useful for trial-wise analyses where each trial needs to be modeled separately. The regressor represents the predicted BOLD response for a single event using a specified hemodynamic response function (HRF).

Usage

```

single_trial_regressor(
  onsets,
  hrf = HRF_SPMG1,
  duration = 0,
  amplitude = 1,
  span = 24
)

```

Arguments

onsets	the event onset in seconds, must be of length 1.
hrf	a hemodynamic response function, e.g. HRF_SPMG1
duration	duration of the event (default is 0), must be length 1.
amplitude	scaling vector (default is 1), must be length 1.
span	the temporal window of the impulse response function (default is 24).

Details

This is a convenience wrapper around ‘regressor’ that ensures inputs have length 1.

Value

A ‘Reg’ object (inheriting from ‘regressor’ and ‘list’).

See Also

[regressor](#)

Examples

```
# Create single trial regressor at 10 seconds
str1 <- single_trial_regressor(onsets = 10, hrf = HRF_SPMG1)

# Single trial with duration and custom amplitude
str2 <- single_trial_regressor(
  onsets = 15,
  duration = 3,
  amplitude = 2,
  hrf = HRF_SPMG1
)

# Evaluate the response
times <- seq(0, 40, by = 0.1)
response <- evaluate(str1, times)
```

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