Package ‘EpiNow2’

September 26, 2023

Type Package

Title Estimate Real-Time Case Counts and Time-Varying Epidemiological Parameters

Version 1.4.0

Description Estimates the time-varying reproduction number, rate of spread, and doubling time using a range of open-source tools (Abbott et al. (2020) <doi:10.12688/wellcomeopenres.16006.1>), and current best practices (Gostic et al. (2020) <doi:10.1101/2020.06.18.20134858>). It aims to help users avoid some of the limitations of naive implementations in a framework that is informed by community feedback and is actively supported.

License MIT + file LICENSE


BugReports https://github.com/epiforecasts/EpiNow2/issues

Depends R (>= 3.5.0)

Imports data.table, futile.logger (>= 1.4), future, future.apply, ggplot2, lifecycle, lubridate, methods, patchwork, progressr, purrr, R.utils (>= 2.0.0), Rcpp (>= 0.12.0), rlang (>= 0.4.7), rstan (>= 2.26.0), rstantools (>= 2.2.0), runner, scales, stats, truncnorm, utils

Suggests covr, dplyr, here, kableExtra, knitr, magrittr, precommit, rmarkdown, spelling, styler, testthat, tidyr, withr

LinkingTo BH (>= 1.66.0), Rcpp (>= 0.12.0), RcppEigen (>= 0.3.3.3.0), RcppParallel (>= 5.0.1), rstan (>= 2.26.0), StanHeaders (>= 2.26.0)

Biarch true

Config/testthat/edition 3

Encoding UTF-8
R topics documented:

Language en-GB
LazyData true
RoxygenNote 7.2.3
NeedsCompilation yes
SystemRequirements GNU make C++17
VignetteBuilder knitr
Author Sam Abbott [aut, cre] (<https://orcid.org/0000-0001-8057-8037>),
Joel Hellewell [aut] (<https://orcid.org/0000-0003-2683-0849>),
Katharine Sherratt [aut],
Katelyn Gostic [aut],
Joe Hickson [aut],
Hamada S. Badr [aut] (<https://orcid.org/0000-0002-9808-2344>),
Michael DeWitt [aut] (<https://orcid.org/0000-0001-8940-1967>),
Robin Thompson [ctb],
Sophie Meakin [ctb],
James Munday [ctb],
Nikos Bosse [ctb],
Paul Mee [ctb],
Peter Ellis [ctb],
Pietro Monticone [ctb],
Lloyd Chapman [ctb],
James M. Azam [ctb] (<https://orcid.org/0000-0001-5782-7330>),
Andrew Johnson [ctb],
EpiForecasts [aut],
Sebastian Funk [aut] (<https://orcid.org/0000-0002-2842-3406>)
Maintainer Sam Abbott <sam.abbott@lshtm.ac.uk>
Repository CRAN
Date/Publication 2023-09-26 12:00:02 UTC

R topics documented:

+ dist_spec .................................................. 5
add_day_of_week ........................................... 6
adjust_infection_to_report .................................. 6
allocate_delays ............................................. 8
allocate_empty ............................................. 9
backcalc_opts ............................................... 10
bootstrapped_dist_fit ...................................... 11
c.dist_spec .................................................. 12
calc_CrI ..................................................... 12
calc_CrIs ................................................... 13
calc_summary_measures ..................................... 14
calc_summary_stats ......................................... 14
clean_nowcasts ............................................. 15
clean_regions ............................................... 16
<table>
<thead>
<tr>
<th>R topics documented:</th>
</tr>
</thead>
<tbody>
<tr>
<td>construct_output</td>
</tr>
<tr>
<td>convert_to_logmean</td>
</tr>
<tr>
<td>convert_to_logsd</td>
</tr>
<tr>
<td>copy_results_to_latest</td>
</tr>
<tr>
<td>create_backcalc_data</td>
</tr>
<tr>
<td>create_clean_reported_cases</td>
</tr>
<tr>
<td>create_future_rt</td>
</tr>
<tr>
<td>create_gp_data</td>
</tr>
<tr>
<td>create_initial_conditions</td>
</tr>
<tr>
<td>create_obs_model</td>
</tr>
<tr>
<td>create_rt_data</td>
</tr>
<tr>
<td>create_shifted_cases</td>
</tr>
<tr>
<td>create_stan_args</td>
</tr>
<tr>
<td>create_stan_data</td>
</tr>
<tr>
<td>create_stan_delays</td>
</tr>
<tr>
<td>delay_opts</td>
</tr>
<tr>
<td>dist_fit</td>
</tr>
<tr>
<td>dist_skel</td>
</tr>
<tr>
<td>dist_spec</td>
</tr>
<tr>
<td>dist_spec_plus</td>
</tr>
<tr>
<td>epinow</td>
</tr>
<tr>
<td>estimates_by_report_date</td>
</tr>
<tr>
<td>estimate_delay</td>
</tr>
<tr>
<td>estimate_infections</td>
</tr>
<tr>
<td>estimate_secondary</td>
</tr>
<tr>
<td>estimate_truncation</td>
</tr>
<tr>
<td>example_confirmed</td>
</tr>
<tr>
<td>expose_stan_fns</td>
</tr>
<tr>
<td>extract_CrIs</td>
</tr>
<tr>
<td>extract_inits</td>
</tr>
<tr>
<td>extract_parameter</td>
</tr>
<tr>
<td>extract_parameter_samples</td>
</tr>
<tr>
<td>extract_stan_param</td>
</tr>
<tr>
<td>extract_static_parameter</td>
</tr>
<tr>
<td>filter_opts</td>
</tr>
<tr>
<td>fit_model_with_nuts</td>
</tr>
<tr>
<td>fit_model_with_vb</td>
</tr>
<tr>
<td>forecast_secondary</td>
</tr>
<tr>
<td>format_fit</td>
</tr>
<tr>
<td>gamma_dist_def</td>
</tr>
<tr>
<td>generation_times</td>
</tr>
<tr>
<td>generation_time_opts</td>
</tr>
<tr>
<td>get_dist</td>
</tr>
<tr>
<td>get_generation_time</td>
</tr>
<tr>
<td>get_incubation_period</td>
</tr>
<tr>
<td>get_raw_result</td>
</tr>
<tr>
<td>get_regional_results</td>
</tr>
<tr>
<td>get_regions</td>
</tr>
<tr>
<td>R topics documented</td>
</tr>
<tr>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>get_regions_with_most_reports</td>
</tr>
<tr>
<td>get_seeding_time</td>
</tr>
<tr>
<td>gp_opts</td>
</tr>
<tr>
<td>growth_to_R</td>
</tr>
<tr>
<td>incubation_periods</td>
</tr>
<tr>
<td>init_cumulative_fit</td>
</tr>
<tr>
<td>lognorm_dist_def</td>
</tr>
<tr>
<td>make_conf</td>
</tr>
<tr>
<td>map_prob_change</td>
</tr>
<tr>
<td>match_output_arguments</td>
</tr>
<tr>
<td>mean.dist_spec</td>
</tr>
<tr>
<td>obs_opts</td>
</tr>
<tr>
<td>opts_list</td>
</tr>
<tr>
<td>plot.dist_spec</td>
</tr>
<tr>
<td>plot.epinow</td>
</tr>
<tr>
<td>plot.estimate_infections</td>
</tr>
<tr>
<td>plot.estimate_secondary</td>
</tr>
<tr>
<td>plot.estimate_truncation</td>
</tr>
<tr>
<td>plot_CrIs</td>
</tr>
<tr>
<td>plot_estimates</td>
</tr>
<tr>
<td>plot_summary</td>
</tr>
<tr>
<td>print.dist_spec</td>
</tr>
<tr>
<td>process_region</td>
</tr>
<tr>
<td>process_regions</td>
</tr>
<tr>
<td>regional_epinow</td>
</tr>
<tr>
<td>regional_runtimes</td>
</tr>
<tr>
<td>regional_summary</td>
</tr>
<tr>
<td>report_cases</td>
</tr>
<tr>
<td>report_plots</td>
</tr>
<tr>
<td>report_summary</td>
</tr>
<tr>
<td>rstan_opts</td>
</tr>
<tr>
<td>rstan_sampling_OPTS</td>
</tr>
<tr>
<td>rstan_vb_opts</td>
</tr>
<tr>
<td>rt_opts</td>
</tr>
<tr>
<td>run_region</td>
</tr>
<tr>
<td>R_to_growth</td>
</tr>
<tr>
<td>sample_approx_dist</td>
</tr>
<tr>
<td>save_estimate_infections</td>
</tr>
<tr>
<td>save_input</td>
</tr>
<tr>
<td>secondary_opts</td>
</tr>
<tr>
<td>setup_default_logging</td>
</tr>
<tr>
<td>setup_dt</td>
</tr>
<tr>
<td>setup_future</td>
</tr>
<tr>
<td>setup_logging</td>
</tr>
<tr>
<td>setup_target_folder</td>
</tr>
<tr>
<td>simulate_infections</td>
</tr>
<tr>
<td>simulate_secondary</td>
</tr>
<tr>
<td>stan_opts</td>
</tr>
</tbody>
</table>
+dist_spec

DESCRIPTION

This is done via convolution with stats::convolve(). Nonparametric delays that can be combined are processed together, and their cumulative distribution function is truncated at a specified tolerance level, ensuring numeric stability.

USAGE

## S3 method for class 'dist_spec'
e1 + e2

ARGUMENTS

e1
  The first delay distribution (from a call to dist_spec()) to combine.
e2
  The second delay distribution (from a call to dist_spec()) to combine.

VALUE

A delay distribution representing the sum of the two delays (with class dist_spec())

AUTHOR(S)

Sebastian Funk

EXAMPLES

# A fixed lognormal distribution with mean 5 and sd 1.
lognormal <- dist_spec(
  mean = 1.6, sd = 1, max = 20, distribution = "lognormal"
)
lognormal + lognormal

# An uncertain gamma distribution with mean 3 and sd 2
gamma <- dist_spec(
  mean = 3, sd = 2, mean_sd = 0.5, sd_sd = 0.5, max = 20,
adjust_infection_to_report

Adjust from Case Counts by Infection Date to Date of Report

Description

[Stable] Maps from cases by date of infection to date of report via date of onset.

distribution = "gamma"
}
lognormal + gamma

# Using tolerance parameter
EpiNow2:::dist_spec_plus(lognormal, lognormal, tolerance = 0.5)

add_day_of_week

Add a day of the week vector

Description

Adds a day of the week vector

Usage

add_day_of_week(dates, week_effect = 7)

Arguments

dates Vector of dates

week_effect Numeric from 1 to 7 defaults to 7

Value

A numeric vector containing the period day of the week index

Examples

dates <- seq(as.Date("2020-03-15"), by = "days", length.out = 15)
# Add date based day of week
add_day_of_week(dates, 7)

# Add shorter week
add_day_of_week(dates, 4)
Usage

adjust_infection_to_report(
  infections,
  delay_defs,
  reporting_model,
  reporting_effect,
  type = "sample",
  truncate_future = TRUE
)

Arguments

infections    data.table containing a date variable and a numeric cases variable.
delay_defs    A list of single row data.tables that each defines a delay distribution (model, parameters and maximum delay for each model). See lognorm_dist_def for an example of the structure.
reporting_model    A function that takes a single numeric vector as an argument and returns a single numeric vector. Can be used to apply stochastic reporting effects. See the examples for details.
reporting_effect    A numeric vector of length 7 that allows the scaling of reported cases by the day on which they report (1 = Monday, 7 = Sunday). By default no scaling occurs.
type    Character string indicating the method to use to transform counts. Supports either "sample" which approximates sampling or "median" would shift by the median of the distribution.
truncate_future    Logical, should cases be truncated if they occur after the first date reported in the data. Defaults to TRUE.

Value

A data.table containing a date variable (date of report) and a cases variable. If return_onset = TRUE there will be a third variable reference which indicates what the date variable refers to.

Author(s)

Sam Abbott

Examples

# define example cases
cases <- data.table::copy(example_confirmed)[, cases := as.integer(confirm)]

# define a single report delay distribution
delay_def <- lognorm_dist_def(
  mean = 5, mean_sd = 1, sd = 3, sd_sd = 1,
allocate_delays

Allocate Delays into Required Stan Format

Description

[Stable] Allocate delays for stan. Used in delay_opts().

```r
# define a single incubation period
incubation_def <- lognorm_dist_def(
  mean = incubation_periods[1,]$mean,
  mean_sd = incubation_periods[1,]$mean_sd,
  sd = incubation_periods[1,]$sd,
  sd_sd = incubation_periods[1,]$sd_sd,
  max_value = 30, samples = 1
)

# simple mapping
report <- adjust_infection_to_report(
  cases, delay_defs = list(incubation_def, delay_def)
)
print(report)

# mapping with a weekly reporting effect
report_weekly <- adjust_infection_to_report(
  cases, delay_defs = list(incubation_def, delay_def),
  reporting_effect = c(1.1, rep(1, 4), 0.95, 0.95)
)
print(report_weekly)

# map using a deterministic median shift for both delays
report_median <- adjust_infection_to_report(cases,
  delay_defs = list(incubation_def, delay_def),
  type = "median")
print(report_median)

# map with a weekly reporting effect and stochastic reporting model
report_stochastic <- adjust_infection_to_report(
  cases, delay_defs = list(incubation_def, delay_def),
  reporting_effect = c(1.1, rep(1, 4), 0.95, 0.95),
  reporting_model = function(n) {
    out <- suppressWarnings(rbinom(length(n), as.integer(n), 0.5))
    out <- ifelse(is.na(out), 0, out)
  }
)
print(report_stochastic)
```
**allocate_empty**

**Usage**

```r
allocate_delays(delay_var, no_delays)
```

**Arguments**

- `delay_var` List of numeric delays
- `no_delays` Numeric, number of delays

**Value**

A numeric array

---

**allocate_empty**  
*Allocate Empty Parameters to a List*

**Description**

[Stable] Allocate missing parameters to be empty two dimensional arrays. Used internally by `simulate_infections`.

**Usage**

```r
allocate_empty(data, params, n = 0)
```

**Arguments**

- `data` A list of parameters
- `params` A character vector of parameters to allocate to empty if missing.
- `n` Numeric, number of samples to assign an empty array

**Value**

A list of parameters some allocated to be empty
backcalc_opts  

Back Calculation Options

Description

[Stable] Defines a list specifying the optional arguments for the back calculation of cases. Only used if \( rt = \text{NULL} \).

Usage

\[
\text{backcalc_opts(prior = "reports", prior_window = 14, rt_window = 1)}
\]

Arguments

- **prior**: A character string defaulting to "reports". Defines the prior to use when de-convolving. Currently implemented options are to use smoothed mean delay shifted reported cases ("reports"), to use the estimated infections from the previous time step seeded for the first time step using mean shifted reported cases ("infections"), or no prior ("none"). Using no prior will result in poor real time performance. No prior and using infections are only supported when a Gaussian process is present. If observed data is not reliable then it a sensible first step is to explore increasing the \( \text{prior_window} \) with a sensible second step being to no longer use reported cases as a prior (i.e set \( \text{prior} = \text{"none"} \)).

- **prior_window**: Integer, defaults to 14 days. The mean centred smoothing window to apply to mean shifted reports (used as a prior during back calculation). 7 days is minimum recommended settings as this smooths day of the week effects but depending on the quality of the data and the amount of information users wish to use as a prior (higher values equalling a less informative prior).

- **rt_window**: Integer, defaults to 1. The size of the centred rolling average to use when estimating \( R_t \). This must be odd so that the central estimate is included.

Value

A list of back calculation settings.

Author(s)

Sam Abbott

Examples

# default settings
backcalc_opts()
Fit a Subsampled Bootstrap to Integer Values and Summarise Distribution Parameters

Description

[Stable] Fits an integer adjusted distribution to a subsampled bootstrap of data and then integrates the posterior samples into a single set of summary statistics. Can be used to generate a robust reporting delay that accounts for the fact the underlying delay likely varies over time or that the size of the available reporting delay sample may not be representative of the current case load.

Usage

```r
bootstrapped_dist_fit(
  values,
  dist = "lognormal",
  samples = 2000,
  bootstraps = 10,
  bootstrap_samples = 250,
  max_value,
  verbose = FALSE
)
```

Arguments

- `values`: Integer vector of values.
- `dist`: Character string, which distribution to fit. Defaults to lognormal ("lognormal") but gamma ("gamma") is also supported.
- `samples`: Numeric, number of samples to take overall from the bootstrapped posteriors.
- `bootstraps`: Numeric, defaults to 1. The number of bootstrap samples (with replacement) of the delay distribution to take.
- `bootstrap_samples`: Numeric, defaults to 100. The number of samples to take in each bootstrap. When the sample size of the supplied delay distribution is less than 100 this is used instead.
- `max_value`: Numeric, defaults to the maximum value in the observed data. Maximum delay to allow (added to output but does impact fitting).
- `verbose`: Logical, defaults to FALSE. Should progress messages be printed.

Value

A `dist_spec` object summarising the bootstrapped distribution

Author(s)

Sam Abbott
Examples

```r
# lognormal
delays <- rlnorm(500, log(5), 1)
out <- bootstrapped_dist_fit(delays,
samples = 1000, bootstraps = 10,
dist = "lognormal"
)
out
```

---

**c.dist_spec**  
*Combines multiple delay distributions for further processing*

---

**Description**

This combines the parameters so that they can be fed as multiple delay distributions to `epinow()` or `estimate_infections()`.

**Usage**

```r
## S3 method for class 'dist_spec'
c(...)
```

**Arguments**

- ...  
The delay distributions (from calls to `dist_spec()`) to combine

**Value**

Combined delay distributions (with class `dist_spec()`)

---

**Author(s)**

Sebastian Funk

---

**calc_CrI**  
*Calculate Credible Interval*

---

**Description**

[Stable] Adds symmetric a credible interval based on quantiles.

**Usage**

```r
calc_CrI(samples, summarise_by = NULL, CrI = 0.9)
```
calc_CrI

Arguments

samples A data.table containing at least a value variable
summarise_by A character vector of variables to group by.
CrI Numeric between 0 and 1. The credible interval for which to return values. Defaults to 0.9.

Value

A data.table containing the upper and lower bounds for the specified credible interval.

Examples

samples <- data.frame(value = 1:10, type = "car")
# add 90% credible interval
calc_CrI(samples)
# add 90% credible interval grouped by type
calc_CrI(samples, summarise_by = "type")

calc_CrIs

Calculate Credible Intervals

Description

[Stable] Adds symmetric credible intervals based on quantiles.

Usage

calc_CrIs(samples, summarise_by = NULL, CrIs = c(0.2, 0.5, 0.9))

Arguments

samples A data.table containing at least a value variable
summarise_by A character vector of variables to group by.
CrIs Numeric vector of credible intervals to calculate.

Value

A data.table containing the summarise_by variables and the specified lower and upper credible intervals.

Examples

samples <- data.frame(value = 1:10, type = "car")
# add credible intervals
calc_CrIIs(samples)
# add 90% credible interval grouped by type
calc_CrIIs(samples, summarise_by = "type")
calc_summary_measures  Calculate All Summary Measures

Description

[Stable] Calculate summary statistics and credible intervals from a data frame by group.

Usage

```r
calc_summary_measures(
  samples,
  summarise_by = NULL,
  order_by = NULL,
  CrIs = c(0.2, 0.5, 0.9)
)
```

Arguments

- `samples`: A data.table containing at least a value variable
- `summarise_by`: A character vector of variables to group by.
- `order_by`: A character vector of parameters to order by, defaults to all `summarise_by` variables.
- `CrIs`: Numeric vector of credible intervals to calculate.

Value

A data.table containing summary statistics by group.

Examples

```r
samples <- data.frame(value = 1:10, type = "car")
# default
calc_summary_measures(samples)
# by type
calc_summary_measures(samples, summarise_by = "type")
```

calc_summary_stats  Calculate Summary Statistics

Description

[Stable] Calculate summary statistics from a data frame by group. Currently supports the mean, median and standard deviation.
calc_summary_stats(samples, summarise_by = NULL)

Arguments

samples A data.table containing at least a value variable
summarise_by A character vector of variables to group by.

Value

A data.table containing the upper and lower bounds for the specified credible interval

Examples

samples <- data.frame(value = 1:10, type = "car")
# default
calc_summary_stats(samples)
# by type
calc_summary_stats(samples, summarise_by = "type")

clean_nowcasts

Clean Nowcasts for a Supplied Date

Description

[Stable] This function removes nowcasts in the format produced by EpiNow2 from a target directory for the date supplied.

Usage

clean_nowcasts(date = NULL, nowcast_dir = ".")

Arguments

date Date object. Defaults to today's date
nowcast_dir Character string giving the filepath to the nowcast results directory. Defaults to the current directory.

Value

No return value, called for side effects
clean_regions  

Clean Regions

Description

[Stable] Removes regions with insufficient time points, and provides logging information on the input.

Usage

clean_regions(reported_cases, non_zero_points)

Arguments

reported_cases  A data frame of confirmed cases (confirm) by date (date), and region (region).
non_zero_points  Numeric, the minimum number of time points with non-zero cases in a region required for that region to be evaluated. Defaults to 7.

Value

A dataframe of cleaned regional data

See Also

regional_epinow

construct_output  

Construct Output

Description

[Stable] Combines the output produced internally by epinow into a single list.

Usage

construct_output(  
estimates,  
estimated_reported_cases,  
plots = NULL,  
summary = NULL,  
samples = TRUE  
)
**convert_to_logmean**

**Arguments**

- **estimates** List of data frames as output by `estimate_infections`
- **estimated_reported_cases**
  A list of dataframes as produced by `estimates_by_report_date`.
- **plots** A list of plots as produced by `report_plots`.
- **summary** A list of summary output as produced by `report_summary`.
- **samples** Logical, defaults to TRUE. Should samples be saved

**Value**

A list of output as returned by `epinow`

**Author(s)**

Sam Abbott

---

**convert_to_logmean**  
*Convert mean and sd to log mean for a log normal distribution*

**Description**

[Stable] Convert from mean and standard deviation to the log mean of the lognormal distribution. Useful for defining distributions supported by `estimate_infections`, `epinow`, and `regional_epinow`.

**Usage**

```r
convert_to_logmean(mean, sd)
```

**Arguments**

- **mean** Numeric, mean of a distribution
- **sd** Numeric, standard deviation of a distribution

**Value**

The log mean of a lognormal distribution

**Examples**

```r
convert_to_logmean(2, 1)
```
**convert_to_logsd**  
*Convert mean and sd to log standard deviation for a log normal distribution*

**Description**

[Stable] Convert from mean and standard deviation to the log standard deviation of the lognormal distribution. Useful for defining distributions supported by `estimate_infections`, `epinow`, and `regional_epinow`.

**Usage**

```r
close_to_logsd(mean, sd)
```

**Arguments**

- `mean`  
  Numeric, mean of a distribution

- `sd`  
  Numeric, standard deviation of a distribution

**Value**

The log standard deviation of a lognormal distribution

**Examples**

```r
close_to_logsd(2, 1)
```

**copy_results_to_latest**  
*Copy Results From Dated Folder to Latest*

**Description**

[Questioning] Copies output from the dated folder to a latest folder. May be undergo changes in later releases.

**Usage**

```r
copy_results_to_latest(target_folder = NULL, latest_folder = NULL)
```

**Arguments**

- `target_folder`  
  Character string specifying where to save results (will create if not present).

- `latest_folder`  
  Character string containing the path to the latest target folder. As produced by `setup_target_folder`. 
create_backcalc_data

Value

No return value, called for side effects

Author(s)

Sam Abbott

create_backcalc_data  Create Back Calculation Data

Description

[Stable] Takes the output of backcalc_opts() and converts it into a list understood by stan.

Usage

create_backcalc_data(backcalc = backcalc_opts())

Arguments

backcalc  A list of options as generated by backcalc_opts() to define the back calculation. Defaults to backcalc_opts().

Value

A list of settings defining the Gaussian process

Author(s)

Sam Abbott

See Also

backcalc_opts

Examples

create_backcalc_data(backcalc = backcalc_opts())
create_clean_reported_cases

Create Clean Reported Cases

Description

[Stable] Cleans a data frame of reported cases by replacing missing dates with 0 cases and applies an optional threshold at which point 0 cases are replaced with a moving average of observed cases. See zero_threshold for details.

Usage

create_clean_reported_cases(
    reported_cases,
    horizon,
    filter_leading_zeros = TRUE,
    zero_threshold = Inf
)

Arguments

reported_cases A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.

horizon Numeric, defaults to 7. Number of days into the future to forecast.

filter_leading_zeros Logical, defaults to TRUE. Should zeros at the start of the time series be filtered out.

zero_threshold [Experimental] Numeric defaults to Inf. Indicates if detected zero cases are meaningful by using a threshold number of cases based on the 7 day average. If the average is above this threshold then the zero is replaced with the backwards looking rolling average. If set to infinity then no changes are made.

Value

A cleaned data frame of reported cases

Author(s)

Sam Abbott

Lloyd Chapman
**create_future_rt**

*Construct the Required Future Rt assumption*

**Description**

[Stable] Converts the future argument from `rt_opts()` into arguments that can be passed to `stan`.

**Usage**

```
create_future_rt(future = "latest", delay = 0)
```

**Arguments**

- `future` (A character string or integer. This argument indicates how to set future Rt values. Supported options are to project using the Rt model ("project"), to use the latest estimate based on partial data ("latest"), to use the latest estimate based on data that is over 50% complete ("estimate"). If an integer is supplied then the Rt estimate from this many days into the future (or past if negative) past will be used forwards in time.)
- `delay` (Numeric mean delay)

**Value**

A list containing a logical called fixed and an integer called from

**Author(s)**

Sam Abbott

---

**create_gp_data**

*Create Gaussian Process Data*

**Description**

[Stable] Takes the output of `gp_opts()` and converts it into a list understood by `stan`.

**Usage**

```
create_gp_data(gp = gp_opts(), data)
```

**Arguments**

- `gp` (A list of options as generated by `gp_opts()` to define the Gaussian process. Defaults to `gp_opts()`. Set to NULL to disable the Gaussian process.)
- `data` (A list containing the following numeric values: t, seeding_time, horizon.)
create_initial_conditions

Value
A list of settings defining the Gaussian process

Author(s)
Sam Abbott

See Also
gp_opts

Examples

# define input data required
data <- list(
  t = 30,
  seeding_time = 7,
  horizon = 7
)

# default gaussian process data
create_gp_data(data = data)

# settings when no gaussian process is desired
create_gp_data(NULL, data)

# custom lengths
create_gp_data(gp_opts(ls_mean = 14), data)

create_initial_conditions

Create Initial Conditions Generating Function

Description

[Stable] Uses the output of create_stan_data to create a function which can be used to sample from the prior distributions (or as close as possible) for parameters. Used in order to initialise each stan chain within a range of plausible values.

Usage

create_initial_conditions(data)

Arguments

data A list of data as produced by create_stan_data.

Value
An initial condition generating function
**create_obs_model**

Create Observation Model Settings

Description

[Stable] Takes the output of `obs_opts()` and converts it into a list understood by `stan`.

Usage

```r
create_obs_model(obs = obs_opts(), dates)
```

Arguments

- **obs**: A list of options as generated by `obs_opts()` defining the observation model. Defaults to `obs_opts()`.
- **dates**: A vector of dates used to calculate the day of the week.

Value

A list of settings ready to be passed to `stan` defining the Observation Model

Author(s)

Sam Abbott

See Also

`obs_opts`

Examples

```r
dates <- seq(as.Date("2020-03-15"), by = "days", length.out = 15)
# default observation model data
create_obs_model(dates = dates)

# Poisson observation model
create_obs_model(obs_opts(family = "poisson"), dates = dates)

# Applying a observation scaling to the data
create_obs_model(
  obs_opts(scale = list(mean = 0.4, sd = 0.01)), dates = dates
)

# Apply a custom week week length
create_obs_model(obs_opts(week_length = 3), dates = dates)
```
create_rt_data Create Time-varying Reproduction Number Data

Description

[Stable] Takes the output from rt_opts() and converts it into a list understood by stan.

Usage

create_rt_data(rt = rt_opts(), breakpoints = NULL, delay = 0, horizon = 0)

Arguments

rt A list of options as generated by rt_opts() defining Rt estimation. Defaults to rt_opts(). Set to NULL to switch to using back calculation rather than generating infections using Rt.
breakpoints An integer vector (binary) indicating the location of breakpoints.
delay Numeric mean delay
horizon Numeric, forecast horizon.

Value

A list of settings defining the time-varying reproduction number

Author(s)

Sam Abbott

See Also

rt_settings

Examples

# default Rt data
create_rt_data()

# settings when no Rt is desired
create_rt_data(rt = NULL)

# using breakpoints
create_rt_data(rt_opts(use_breakpoints = TRUE), breakpoints = rep(1, 10))
create_shifted_cases

Create Delay Shifted Cases

Description

[Stable]

This function creates a data frame of reported cases that has been smoothed using a centred partial rolling average (with a period set by `smoothing_window`) and shifted back in time by some delay. It is used by `estimate_infections` to generate the mean shifted prior on which the back calculation method (see `backcalc_opts()`) is based.

Usage

```r
create_shifted_cases(reported_cases, shift, smoothing_window, horizon)
```

Arguments

- `reported_cases`: A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.
- `shift`: Numeric, mean delay shift to apply.
- `smoothing_window`: Numeric, the rolling average smoothing window to apply. Must be odd in order to be defined as a centred average.
- `horizon`: Numeric, defaults to 7. Number of days into the future to forecast.

Value

A data frame for shifted reported cases

Author(s)

Sam Abbott

Examples

```r
create_shifted_cases(example_confirmed, 7, 14, 7)
```
create_stan_args  Create a List of Stan Arguments

Description

[Stable] Generates a list of arguments as required by `rstan::sampling` or `rstan::vb` by combining the required options, with data, and type of initialisation. Initialisation defaults to random but it is expected that `create_initial_conditions` will be used.

Usage

```r
create_stan_args(
  stan = stan_opts(),
  data = NULL,
  init = "random",
  verbose = FALSE
)
```

Arguments

- **stan**: A list of stan options as generated by `stan_opts()`. Defaults to `stan_opts()`. Can be used to override `data`, `init`, and `verbose` settings if desired.
- **data**: A list of stan data as created by `create_stan_data`
- **init**: Initial conditions passed to `rstan`. Defaults to "random" but can also be a function (as supplied by `create_initial_conditions`).
- **verbose**: Logical, defaults to `FALSE`. Should verbose progress messages be returned.

Value

A list of stan arguments

Author(s)

Sam Abbott

Examples

```r
# default settings
create_stan_args()

# increasing warmup
create_stan_args(stan = stan_opts(warmup = 1000))
```
create_stan_data  
Create Stan Data Required for estimate_infections

Description

[Stable] Takes the output of `stan_opts()` and converts it into a list understood by `stan`. Internally calls the other create_ family of functions to construct a single list for input into `stan` with all data required present.

Usage

```r
create_stan_data(
    reported_cases,
    seeding_time,
    rt,
    gp,
    obs,
    horizon,
    backcalc,
    shifted_cases
)
```

Arguments

- `reported_cases`: A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.
- `seeding_time`: Integer; seeding time, usually obtained using `get_seeding_time()`.
- `rt`: A list of options as generated by `rt_opts()` defining Rt estimation. Defaults to `rt_opts()`. Set to NULL to switch to using back calculation rather than generating infections using Rt.
- `gp`: A list of options as generated by `gp_opts()` to define the Gaussian process. Defaults to `gp_opts()`. Set to NULL to disable the Gaussian process.
- `obs`: A list of options as generated by `obs_opts()` defining the observation model. Defaults to `obs_opts()`.
- `horizon`: Numeric, forecast horizon.
- `backcalc`: A list of options as generated by `backcalc_opts()` to define the back calculation. Defaults to `backcalc_opts()`.
- `shifted_cases`: A dataframe of delay shifted cases

Value

A list of stan data
create_stan_delays  Create delay variables for stan

Description

Create delay variables for stan

Usage

create_stan_delays(..., weight = 1)

Arguments

...  Named delay distributions specified using dist_spec(). The names are assigned to IDs
weight  Numeric, weight associated with delay priors; default: 1

Value

A list of variables as expected by the stan model

Author(s)

Sebastian Funk

delay_opts  Delay Distribution Options

Description

[Stable] Returns delay distributions formatted for usage by downstream functions.

Usage

delay_opts(dist = dist_spec(), ..., fixed = FALSE)

Arguments

dist  A delay distribution or series of delay distributions generated using dist_spec(). Default is an empty call to dist_spec(), i.e. no delay
...  deprecated; use dist instead
fixed  deprecated; use dist instead
Value

A list summarising the input delay distributions.

Author(s)

Sam Abbott
Sebastian Funk

See Also

convert_to_logmean convert_to_logsd bootstrapped_dist_fit dist_spec

Examples

# no delays
delay_opts()

# A single delay that has uncertainty
delay <- dist_spec(mean = 1, mean_sd = 0.2, sd = 0.5, sd_sd = 0.1, max = 15)
delay_opts(delay)

# A single delay without uncertainty
delay <- dist_spec(mean = 1, sd = 0.5, max = 15)
delay_opts(delay)

# Multiple delays (in this case twice the same)
delay_opts(delay + delay)

---

**dist_fit**

*Fit an Integer Adjusted Exponential, Gamma or Lognormal distributions*

---

**Description**

[Stable] Fits an integer adjusted exponential, gamma or lognormal distribution using `stan`.

**Usage**

dist_fit(
    values = NULL,
    samples = 1000,
    cores = 1,
    chains = 2,
    dist = "exp",
    verbose = FALSE
)
Arguments

values  Numeric vector of values
samples Numeric, number of samples to take. Must be >= 1000. Defaults to 1000.
cores  Numeric, defaults to 1. Number of CPU cores to use (no effect if greater than
        the number of chains).
chains Numeric, defaults to 2. Number of MCMC chains to use. More is better with
        the minimum being two.
dist  Character string, which distribution to fit. Defaults to exponential ("exp") but
        gamma ("gamma") and lognormal ("lognormal") are also supported.
verbose Logical, defaults to FALSE. Should verbose progress messages be printed.

Value

A stan fit of an interval censored distribution

Author(s)

Sam Abbott

Examples

# integer adjusted exponential model
dist_fit(rexp(1:100, 2),
    samples = 1000, dist = "exp",
    cores = ifelse(interactive(), 4, 1), verbose = TRUE
)

# integer adjusted gamma model
dist_fit(rgamma(1:100, 5, 5),
    samples = 1000, dist = "gamma",
    cores = ifelse(interactive(), 4, 1), verbose = TRUE
)

# integer adjusted lognormal model
dist_fit(rlnorm(1:100, log(5), 0.2),
    samples = 1000, dist = "lognormal",
    cores = ifelse(interactive(), 4, 1), verbose = TRUE
)
**dist_skel**

**Distribution Skeleton**

**Description**

[Questioning] This function acts as a skeleton for a truncated distribution defined by model type, maximum value and model parameters. It is designed to be used with the output from get_dist.

**Usage**

```r
dist_skel(
  n,
  dist = FALSE,
  cum = TRUE,
  model,
  discrete = FALSE,
  params,
  max_value = 120
)
```

**Arguments**

- **n**
  Numeric vector, number of samples to take (or days for the probability density).

- **dist**
  Logical, defaults to `FALSE`. Should the probability density be returned rather than a number of samples.

- **cum**
  Logical, defaults to `TRUE`. If `dist = TRUE` should the returned distribution be cumulative.

- **model**
  Character string, defining the model to be used. Supported options are exponential ("exp"), gamma ("gamma"), and log normal ("lognormal")

- **discrete**
  Logical, defaults to `FALSE`. Should the probability distribution be discretised. In this case each entry of the probability mass function corresponds to the 1-length interval ending at the entry, i.e. the probability mass function is a vector where the first entry corresponds to the integral over the (0,1] interval of the continuous distribution, the second entry corresponds to the (1,2] interval etc.

- **params**
  A list of parameters values (by name) required for each model. For the exponential model this is a rate parameter and for the gamma model this is alpha and beta.

- **max_value**
  Numeric, the maximum value to allow. Defaults to 120. Samples outside of this range are resampled.

**Value**

A vector of samples or a probability distribution.
Author(s)
Sam Abbott
Sebastian Funk

Examples

```r
## Exponential model
# sample
dist_skel(10, model = "exp", params = list(rate = 1))

# cumulative prob density
dist_skel(1:10, model = "exp", dist = TRUE, params = list(rate = 1))

# probability density
dist_skel(1:10,
  model = "exp", dist = TRUE,
  cum = FALSE, params = list(rate = 1))
)

## Gamma model
# sample
dist_skel(10, model = "gamma", params = list(shape = 1, scale = 2))

# cumulative prob density
dist_skel(0:10,
  model = "gamma", dist = TRUE,
  params = list(shape = 1, scale = 2))
)

# probability density
dist_skel(0:10,
  model = "gamma", dist = TRUE,
  cum = FALSE, params = list(shape = 2, scale = 2))
)

## Log normal model
# sample
dist_skel(10, model = "lognormal", params = list(mean = log(5), sd = log(2)))

# cumulative prob density
dist_skel(0:10,
  model = "lognormal", dist = TRUE,
  params = list(mean = log(5), sd = log(2)))
)

# probability density
dist_skel(0:10,
  model = "lognormal", dist = TRUE, cum = FALSE,
  params = list(mean = log(5), sd = log(2)))
)
```
**dist_spec**

Specify a distribution.

**Description**

**[Stable]** Defines the parameters of a supported distribution for use in onward modelling. Multiple distribution families are supported - see the documentation for `family` for details. Alternatively, a nonparametric distribution can be specified using the `pmf` argument. This function provides distribution functionality in `delay_opts()`, `generation_time_opts()`, and `trunc_opts()`.

**Usage**

```r
dist_spec(
  mean,
  sd = 0,
  mean_sd = 0,
  sd_sd = 0,
  distribution = c("lognormal", "gamma"),
  max,
  pmf = numeric(0),
  fixed = FALSE
)
```

**Arguments**

- `mean` Numeric. If the only non-zero summary parameter then this is the fixed interval of the distribution. If the `sd` is non-zero then this is the mean of the distribution given by `dist`. If this is not given a vector of empty vectors is returned.
- `sd` Numeric, defaults to 0. Sets the standard deviation of the distribution.
- `mean_sd` Numeric, defaults to 0. Sets the standard deviation of the uncertainty around the mean of the distribution assuming a normal prior.
- `sd_sd` Numeric, defaults to 0. Sets the standard deviation of the uncertainty around the `sd` of the distribution assuming a normal prior.
- `distribution` Character, defaults to "lognormal". The (discretised) distribution to be used. If `sd == 0` then the distribution is fixed and a delta function is used. If `sd > 0` then the distribution is discretised and truncated.

The following distributions are currently supported:

- "lognormal" - a lognormal distribution. For this distribution `mean` is the mean of the natural logarithm of the delay (on the log scale) and `sd` is the standard deviation of the natural logarithm of the delay.
- "gamma" - a gamma distribution. For this distribution `mean` is the mean of the delay and `sd` is the standard deviation of the delay. During model fitting these are then transformed to the shape and scale of the gamma distribution.

When `distribution` is the default lognormal distribution the other function arguments have the following definition:
dist_spec_plus

- mean is the mean of the natural logarithm of the delay (on the log scale).
- sd is the standard deviation of the natural logarithm of the delay.

max Numeric, maximum value of the distribution. The distribution will be truncated at this value.

pmf Numeric, a vector of values that represent the (nonparametric) probability mass function of the delay (starting with 0); defaults to an empty vector corresponding to a parametric specification of the distribution (using mean, sd and corresponding uncertainties)

fixed Logical, defaults to FALSE. Should delays be treated as coming from fixed (vs uncertain) distributions. Overrides any values assigned to mean_sd and sd_sd by setting them to zero. reduces compute requirement but may produce spuriously precise estimates.

Value

A list of distribution options.

Author(s)

Sebastian Funk
Sam Abbott

Examples

# A fixed lognormal distribution with mean 5 and sd 1.
dist_spec(mean = 5, sd = 1, max = 20, distribution = "lognormal")

# An uncertain gamma distribution with mean 3 and sd 2
dist_spec(
    mean = 3, sd = 2, mean_sd = 0.5, sd_sd = 0.5, max = 20,
    distribution = "gamma"
)

dist_spec_plus Creates a delay distribution as the sum of two other delay distributions

Description

This is done via convolution with stats::convolve(). Nonparametric delays that can be combined are processed together, and their cumulative distribution function is truncated at a specified tolerance level, ensuring numeric stability.

Usage

dist_spec_plus(e1, e2, tolerance = 0.001)
Arguments

- **e1**
  The first delay distribution (from a call to `dist_spec()`) to combine.

- **e2**
  The second delay distribution (from a call to `dist_spec()`) to combine.

- **tolerance**
  A numeric value that sets the cumulative probability to retain when truncating the cumulative distribution function of the combined nonparametric delays. The default value is 0.001 with this retaining 0.999 of the cumulative probability. Note that using a larger tolerance may result in a smaller number of points in the combined nonparametric delay but may also impact the accuracy of the combined delay (i.e., change the mean and standard deviation).

Value

A delay distribution representing the sum of the two delays (with class `dist_spec()`)

Author(s)

- Sebastian Funk
- Sam Abbott

---

Description

[Maturing] This function wraps the functionality of `estimate_infections()` and `forecast_infections()` in order to estimate Rt and cases by date of infection, forecast into these infections into the future. It also contains additional functionality to convert forecasts to date of report and produce summary output useful for reporting results and interpreting them. See [here](#) for an example of using `epinow` to estimate Rt for Covid-19 in a country from the ECDC data source.

Usage

```r
epinow(
  reported_cases,
  generation_time = NULL,
  delays = delay_opts(),
  truncation = trunc_opts(),
  rt = rt_opts(),
  backcalc = backcalc_opts(),
  gp = gp_opts(),
  obs = obs_opts(),
  stan = stan_opts(),
  horizon = 7,
  CrIs = c(0.2, 0.5, 0.9),
  filter_leading_zeros = TRUE,
  zero_threshold = Inf,
)```
Arguments

reported_cases A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.

generation_time A call to generation_time_opts() defining the generation time distribution used. For backwards compatibility a list of summary parameters can also be passed.

delays A call to delay_opts() defining delay distributions and options. See the documentation of delay_opts() and the examples below for details.

truncation A call to trunc_opts() defining the truncation of observed data. Defaults to trunc_opts(). See estimate_truncation() for an approach to estimating truncation from data.

rt A list of options as generated by rt_opts() defining Rt estimation. Defaults to rt_opts(). Set to NULL to switch to using back calculation rather than generating infections using Rt.

backcalc A list of options as generated by backcalc_opts() to define the back calculation. Defaults to backcalc_opts().

gp A list of options as generated by gp_opts() to define the Gaussian process. Defaults to gp_opts(). Set to NULL to disable the Gaussian process.

obs A list of options as generated by obs_opts() defining the observation model. Defaults to obs_opts().

stan A list of stan options as generated by stan_opts(). Defaults to stan_opts(). Can be used to override data, init, and verbose settings if desired.

horizon Numeric, defaults to 7. Number of days into the future to forecast.

CrIs Numeric vector of credible intervals to calculate.

filter_leading_zeros Logical, defaults to TRUE. Should zeros at the start of the time series be filtered out.

zero_threshold [Experimental] Numeric defaults to Inf. Indicates if detected zero cases are meaningful by using a threshold number of cases based on the 7 day average. If the average is above this threshold then the zero is replaced with the backwards looking rolling average. If set to infinity then no changes are made.

return_output Logical, defaults to FALSE. Should output be returned, this automatically updates to TRUE if no directory for saving is specified.
output A character vector of optional output to return. Supported options are samples ("samples"), plots ("plots"), the run time ("timing"), copying the dated folder into a latest folder (if target_folder is not null, set using "latest"), and the stan fit ("fit"). The default is to return all options.

plot_args A list of optional arguments passed to plot.epinow().

target_folder Character string specifying where to save results (will create if not present).

target_date Date, defaults to maximum found in the data if not specified.

logs Character path indicating the target folder in which to store log information. Defaults to the temporary directory if not specified. Default logging can be disabled if logs is set to NULL. If specifying a custom logging setup then the code for setup_default_logging and the setup_logging function are a sensible place to start.

id A character string used to assign logging information on error. Used by regional_epinow to assign errors to regions. Alter the default to run with error catching.

verbose Logical, defaults to TRUE when used interactively and otherwise FALSE. Should verbose debug progress messages be printed. Corresponds to the "DEBUG" level from futile.logger. See setup_logging for more detailed logging options.

Value

A list of output from estimate_infections, forecast_infections, report_cases, and report_summary.

Author(s)

Sam Abbott

See Also

estimate_infections simulate_infections forecast_infections regional_epinow

Examples

# set number of cores to use
old_opts <- options()
options(mc.cores = ifelse(interactive(), 4, 1))
# construct example distributions
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- dist_spec(
  mean = convert_to_logmean(2, 1),
  mean_sd = 0.1,

estimates_by_report_date

Estimate Cases by Report Date

Description

[Questioning] Either extracts or converts reported cases from an input data table. For output from estimate_infections this is a simple filtering step.

Usage

estimates_by_report_date(
  estimates,
  CrIs = c(0.2, 0.5, 0.9),
  target_folder = NULL,
  samples = TRUE
)

Arguments

estimates List of data frames as output by estimate_infections
CrIs Numeric vector of credible intervals to calculate.
target_folder Character string specifying where to save results (will create if not present).
samples Logical, defaults to TRUE. Should samples be saved
Value

A list of samples and summarised estimates of estimated cases by date of report.

Author(s)

Sam Abbott

estimate_delay  Estimate a Delay Distribution

Description

[Maturing] Estimate a log normal delay distribution from a vector of integer delays. Currently this function is a simple wrapper for bootstrapped_dist_fit.

Usage

estimate_delay(delays, ...)

Arguments

delays  Integer vector of delays
...
Arguments to pass to internal methods.

Value

A dist_spec summarising the bootstrapped distribution

Author(s)

Sam Abbott

See Also

bootstrapped_dist_fit

Examples

delays <- rlnorm(500, log(5), 1)
estimate_delay(delays, samples = 1000, bootstraps = 10)
estimate_infections

Estimate Infections, the Time-Varying Reproduction Number and the Rate of Growth

Description

[Maturing] Uses a non-parametric approach to reconstruct cases by date of infection from reported cases. It uses either a generative Rt model or non-parametric back calculation to estimate underlying latent infections and then maps these infections to observed cases via uncertain reporting delays and a flexible observation model. See the examples and function arguments for the details of all options. The default settings may not be sufficient for your use case so the number of warmup samples (stan_args = list(warmup)) may need to be increased as may the overall number of samples. Follow the links provided by any warnings messages to diagnose issues with the MCMC fit. It is recommended to explore several of the Rt estimation approaches supported as not all of them may be suited to users own use cases. See here for an example of using estimate_infections within the epinow wrapper to estimate Rt for Covid-19 in a country from the ECDC data source.

Usage

```r
estimate_infections(
  reported_cases,
  generation_time = generation_time_opts(),
  delays = delay_opts(),
  truncation = trunc_opts(),
  rt = rt_opts(),
  backcalc = backcalc_opts(),
  gp = gp_opts(),
  obs = obs_opts(),
  stan = stan_opts(),
  horizon = 7,
  CrIs = c(0.2, 0.5, 0.9),
  filter_leading_zeros = TRUE,
  zero_threshold = Inf,
  weigh_delay_priors = TRUE,
  id = "estimate_infections",
  verbose = interactive()
)
```

Arguments

- `reported_cases` A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.
- `generation_time` A call to generation_time_opts() defining the generation time distribution used. For backwards compatibility a list of summary parameters can also be passed.
A call to `delay_opts()` defining delay distributions and options. See the documentation of `delay_opts()` and the examples below for details.

A call to `trunc_opts()` defining the truncation of observed data. Defaults to `trunc_opts()`. See `estimate_truncation()` for an approach to estimating truncation from data.

A list of options as generated by `rt_opts()` defining Rt estimation. Defaults to `rt_opts()`. Set to `NULL` to switch to using back calculation rather than generating infections using Rt.

A list of options as generated by `backcalc_opts()` to define the back calculation. Defaults to `backcalc_opts()`.

A list of options as generated by `gp_opts()` to define the Gaussian process. Defaults to `gp_opts()`. Set to `NULL` to disable the Gaussian process.

A list of options as generated by `obs_opts()` defining the observation model. Defaults to `obs_opts()`.

A list of stan options as generated by `stan_opts()`. Defaults to `stan_opts()`. Can be used to override data, init, and verbose settings if desired.

Numeric, defaults to 7. Number of days into the future to forecast.

Numeric vector of credible intervals to calculate.

Logical, defaults to `TRUE`. Should zeros at the start of the time series be filtered out.

Numeric defaults to `Inf`. Indicates if detected zero cases are meaningful by using a threshold number of cases based on the 7 day average. If the average is above this threshold then the zero is replaced with the backwards looking rolling average. If set to infinity then no changes are made.

Logical. If `TRUE` (default), all delay distribution priors will be weighted by the number of observation data points, in doing so approximately placing an independent prior at each time step and usually preventing the posteriors from shifting. If `FALSE`, no weight will be applied, i.e. delay distributions will be treated as a single parameters.

A character string used to assign logging information on error. Used by `regional_epinow` to assign errors to regions. Alter the default to run with error catching.

Logical, defaults to `TRUE` when used interactively and otherwise `FALSE`. Should verbose debug progress messages be printed. Corresponds to the "DEBUG" level from `futile.logger`. See `setup_logging` for more detailed logging options.

A list of output including: posterior samples, summarised posterior samples, data used to fit the model, and the fit object itself.

Sam Abbott
See Also

epinow regional_epinow forecast_infections simulate_infections

Examples

# set number of cores to use
old_opts <- options()
options(mc.cores = ifelse(interactive(), 4, 1))

# get example case counts
reported_cases <- example_confirmed[1:60]

# set up example generation time
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani", fixed = TRUE
)

# set delays between infection and case report
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer", fixed = TRUE
)

# set delays between infection and case report, with uncertainty
incubation_period_uncertain <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- dist_spec(
  mean = convert_to_logmean(2, 1), mean_sd = 0,
  sd = convert_to_logsd(2, 1), sd_sd = 0, max = 10
)

# default settings but assuming that delays are fixed rather than uncertain
def <- estimate_infections(reported_cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  rt = rt_opts(prior = list(mean = 2, sd = 0.1)),
  stan = stan_opts(control = list(adapt_delta = 0.95))
)

# real time estimates
summary(def)

# summary plot
plot(def)

# decreasing the accuracy of the approximate Gaussian to speed up
# computation.
# These settings are an area of active research. See ?gp_opts for details.
agp <- estimate_infections(reported_cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  rt = rt_opts(prior = list(mean = 2, sd = 0.1)),
  gp = gp_opts(ls_min = 10, basis_prop = 0.1),
  stan = stan_opts(control = list(adapt_delta = 0.95))
)
estimate_infections

summary(agp)
plot(agp)

# Adjusting for future susceptible depletion
dep <- estimate_infections(reported_cases,
generation_time = generation_time_opts(generation_time),
delays = delay_opts(incubation_period + reporting_delay),
rt = rt_opts(
  prior = list(mean = 2, sd = 0.1),
  pop = 1000000, future = "latest"),
gp = gp_opts(ls_min = 10, basis_prop = 0.1), horizon = 21,
stan = stan_opts(control = list(adapt_delta = 0.95))
)
plot(dep)

# Adjusting for truncation of the most recent data
# See estimate_truncation for an approach to estimating this from data
trunc_dist <- dist_spec(
  mean = convert_to_logmean(0.5, 0.5), mean_sd = 0.1,
  sd = convert_to_logsd(0.5, 0.5), sd_sd = 0.1,
  max = 3
)
trunc <- estimate_infections(reported_cases,
generation_time = generation_time_opts(generation_time),
delays = delay_opts(incubation_period + reporting_delay),
truncation = trunc_opts(trunc_dist),
rt = rt_opts(prior = list(mean = 2, sd = 0.1)),
gp = gp_opts(ls_min = 10, basis_prop = 0.1),
stan = stan_opts(control = list(adapt_delta = 0.95))
)
plot(trunc)

# using back calculation (combined here with under reporting)
# this model is in the order of 10 ~ 100 faster than the gaussian process
# method
# it is likely robust for retrospective Rt but less reliable for real time
# estimates
# the width of the prior window controls the reliance on observed data and
# can be optionally switched off using backcalc_opts(prior = "none"),
# see ?backcalc_opts for other options
backcalc <- estimate_infections(reported_cases,
generation_time = generation_time_opts(generation_time),
delays = delay_opts(incubation_period + reporting_delay),
rt = NULL, backcalc = backcalc_opts(),
obs = obs_opts(scale = list(mean = 0.4, sd = 0.05)),
horizon = 0
)
plot(backcalc)

# Rt projected into the future using the Gaussian process
project_rt <- estimate_infections(reported_cases,
generation_time = generation_time_opts(generation_time),
rt = rt_opts(prior = list(mean = 2, sd = 0.1)),
gp = gp_opts(ls_min = 10, basis_prop = 0.1), horizon = 21,
stan = stan_opts(control = list(adapt_delta = 0.95))
)
delays = delay_opts(incubation_period + reporting_delay),
rt = rt_opts(
    prior = list(mean = 2, sd = 0.1),
    future = "project"
)
)
plot(project_rt)

# default settings on a later snapshot of data
snapshot_cases <- example_confirmed[80:130]
snapshot <- estimate_infections(snapshot_cases,
    generation_time = generation_time_opts(generation_time),
    delays = delay_opts(incubation_period + reporting_delay),
    rt = rt_opts(prior = list(mean = 1, sd = 0.1))
)
plot(snapshot)

# stationary Rt assumption (likely to provide biased real-time estimates)
# with uncertain reporting delays
stat <- estimate_infections(reported_cases,
    generation_time = generation_time_opts(generation_time),
    delays = delay_opts(incubation_period_uncertain + reporting_delay),
    rt = rt_opts(prior = list(mean = 2, sd = 0.1), gp_on = "R0")
)
plot(stat)

# no gaussian process (i.e fixed Rt assuming no breakpoints)
# with uncertain reporting delays
fixed <- estimate_infections(reported_cases,
    generation_time = generation_time_opts(generation_time),
    delays = delay_opts(incubation_period_uncertain + reporting_delay),
    gp = NULL
)
plot(fixed)

# no delays
no_delay <- estimate_infections(
    reported_cases,
    generation_time = generation_time_opts(generation_time)
)
plot(no_delay)

# break point but otherwise static Rt
# with uncertain reporting delays
bp_cases <- data.table::copy(reported_cases)
bp_cases[, 
    breakpoint := ifelse(date == as.Date("2020-03-16"), 1, 0)
]
bkp <- estimate_infections(bp_cases,
    generation_time = generation_time_opts(generation_time),
    delays = delay_opts(incubation_period_uncertain + reporting_delay),
    rt = rt_opts(prior = list(mean = 2, sd = 0.1)),
    gp = NULL
```r
# break point effect
summary(bkp, type = "parameters", params = "breakpoints")
plot(bkp)

# weekly random walk
# with uncertain reporting delays
rw <- estimate_infections(reported_cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period_uncertain + reporting_delay),
  rt = rt_opts(prior = list(mean = 2, sd = 0.1), rw = 7),
  gp = NULL
)

# random walk effects
summary(rw, type = "parameters", params = "breakpoints")
plot(rw)

options(old_opts)
```

---

**estimate_secondary**  
*Estimate a Secondary Observation from a Primary Observation*

**Description**

[Stable] Estimates the relationship between a primary and secondary observation, for example hospital admissions and deaths or hospital admissions and bed occupancy. See `secondary_opts()` for model structure options. See parameter documentation for model defaults and options. See the examples for case studies using synthetic data and here for an example of forecasting Covid-19 deaths from Covid-19 cases. See here for a prototype function that may be used to estimate and forecast a secondary observation from a primary across multiple regions and here # nolint for an application forecasting Covid-19 deaths in Germany and Poland.

**Usage**

```r
estimate_secondary(
  reports,
  secondary = secondary_opts(),
  delays = delay_opts(dist_spec(mean = 2.5, mean_sd = 0.5, sd = 0.47, sd_sd = 0.25, max = 30)),
  truncation = trunc_opts(),
  obs = obs_opts(),
  burn_in = 14,
  CrIs = c(0.2, 0.5, 0.9),
  priors = NULL,
  model = NULL,
  weigh_delay_priors = FALSE,
  verbose = interactive(),
)```
Arguments

- **reports**
  A data frame containing the date of report and both primary and secondary reports.

- **secondary**
  A call to `secondary_opts()` or a list containing the following binary variables:
  - cumulative
  - historic
  - primary_hist_additive
  - current
  - primary_current_additive.
  These parameters control the structure of the secondary model, see `secondary_opts()` for details.

- **delays**
  A call to `delay_opts()` defining delay distributions between primary and secondary observations. See the documentation of `delay_opts()` for details. By default a diffuse prior is assumed with a mean of 14 days and standard deviation of 7 days (with a standard deviation of 0.5 and 0.25 respectively on the log scale).

- **truncation**
  A call to `trunc_opts()` defining the truncation of observed data. Defaults to `trunc_opts()`. See `estimate_truncation()` for an approach to estimating truncation from data.

- **obs**
  A list of options as generated by `obs_opts()` defining the observation model. Defaults to `obs_opts()`.

- **burn_in**
  Integer, defaults to 14 days. The number of data points to use for estimation but not to fit to at the beginning of the time series. This must be less than the number of observations.

- **CrIs**
  Numeric vector of credible intervals to calculate.

- **priors**
  A data.frame of named priors to be used in model fitting rather than the defaults supplied from other arguments. This is typically useful if wanting to inform an estimate from the posterior of another model fit.

- **model**
  A compiled stan model to override the default model. May be useful for package developers or those developing extensions.

- **weigh_delay_priors**
  Logical. If TRUE, all delay distribution priors will be weighted by the number of observation data points, in doing so approximately placing an independent prior at each time step and usually preventing the posteriors from shifting. If FALSE (default), no weight will be applied, i.e. delay distributions will be treated as a single parameters.

- **verbose**
  Logical, should model fitting progress be returned. Defaults to `interactive()`.

- **...**
  Additional parameters to pass to `rstan::sampling`.

Value

A list containing:
- **predictions** (a data frame ordered by date with the primary, and secondary observations, and a summary of the model estimated secondary observations),
- **posterior** (which contains a summary of the entire model posterior),
- **data** (a list of data used to fit the model),
- **fit** (the stanfit object).
Author(s)
Sam Abbott

Examples

```r
# set number of cores to use
old_opts <- options()
options(mc.cores = ifelse(interactive(), 4, 1))

# load data.table for manipulation
library(data.table)

#### Incidence data example ####
# make some example secondary incidence data
cases <- example_confirmed
cases <- as.data.table(cases)[, primary := confirm]
# Assume that only 40 percent of cases are reported
cases[, scaling := 0.4]
# Parameters of the assumed log normal delay distribution
cases[, meanlog := 1.8][, sdlog := 0.5]

# Simulate secondary cases
cases <- simulate_secondary(cases, type = "incidence")

# fit model to example data specifying a weak prior for fraction reported
# with a secondary case
inc <- estimate_secondary(cases[1:60],
    obs = obs_opts(scale = list(mean = 0.2, sd = 0.2), week_effect = FALSE)
)
plot(inc, primary = TRUE)

# forecast future secondary cases from primary
inc_preds <- forecast_secondary(inc, cases[seq(61, .N)][, value := primary])
plot(inc_preds, new_obs = cases, from = "2020-05-01")

#### Prevalence data example ####

# make some example prevalence data
cases <- example_confirmed
cases <- as.data.table(cases)[, primary := confirm]
# Assume that only 30 percent of cases are reported
cases[, scaling := 0.3]
# Parameters of the assumed log normal delay distribution
cases[, meanlog := 1.6][, sdlog := 0.8]

# Simulate secondary cases
cases <- simulate_secondary(cases, type = "prevalence")
```

# fit model to example prevalence data
prev <- estimate_secondary(cases[1:100],
    secondary = secondary_opts(type = "prevalence"),
    obs = obs_opts(
        week_effect = FALSE,
        scale = list(mean = 0.4, sd = 0.1)
    )
)
plot(prev, primary = TRUE)

# forecast future secondary cases from primary
prev_preds <- forecast_secondary(prev, cases[seq(101, .N)][, value := primary]
)
plot(prev_preds, new_obs = cases, from = "2020-06-01")

options(old_opts)

---

**estimate_truncation**  
*Estimate Truncation of Observed Data*

**Description**

[Stable] Estimates a truncation distribution from multiple snapshots of the same data source over time. This distribution can then be used in `regional_epinow`, `epinow`, and `estimate_infections` to adjust for truncated data. See [here](#) for an example of using this approach on Covid-19 data in England. The functionality offered by this function is now available in a more principled manner in the `epinowcast` R package.

The model of truncation is as follows:

1. The truncation distribution is assumed to be discretised log normal with a mean and standard deviation that is informed by the data.
2. The data set with the latest observations is adjusted for truncation using the truncation distribution.
3. Earlier data sets are recreated by applying the truncation distribution to the adjusted latest observations in the time period of the earlier data set. These data sets are then compared to the earlier observations assuming a negative binomial observation model with an additive noise term to deal with zero observations.

This model is then fit using `stan` with standard normal, or half normal, prior for the mean, standard deviation, 1 over the square root of the overdispersion and additive noise term.

This approach assumes that:

- Current truncation is related to past truncation.
- Truncation is a multiplicative scaling of underlying reported cases.
- Truncation is log normally distributed.
Usage

```r
estimate_truncation(
  obs,
  max_truncation,
  trunc_max = 10,
  trunc_dist = "lognormal",
  truncation = dist_spec(mean = 0, sd = 0, mean_sd = 1, sd_sd = 1, max = 10),
  model = NULL,
  CrIs = c(0.2, 0.5, 0.9),
  weigh_delay_priors = FALSE,
  verbose = TRUE,
  ...
)
```

Arguments

- **obs**: A list of data frames each containing a date variable and a confirm (integer) variable. Each data set should be a snapshot of the reported data over time. All data sets must contain a complete vector of dates.
- **max_truncation**: Deprecated; use truncation instead.
- **trunc_max**: Deprecated; use truncation instead.
- **trunc_dist**: Deprecated; use truncation instead.
- **truncation**: A call to trunc_opts() defining the truncation of observed data. Defaults to trunc_opts(). See estimate_truncation() for an approach to estimating truncation from data.
- **model**: A compiled stan model to override the default model. May be useful for package developers or those developing extensions.
- **CrIs**: Numeric vector of credible intervals to calculate.
- **weigh_delay_priors**: Logical. If TRUE, all delay distribution priors will be weighted by the number of observation data points, in doing so approximately placing an independent prior at each time step and usually preventing the posteriors from shifting. If FALSE (default), no weight will be applied, i.e. delay distributions will be treated as a single parameters.
- **verbose**: Logical, should model fitting progress be returned.
- **...**: Additional parameters to pass to rstan::sampling.

Value

A list containing: the summary parameters of the truncation distribution (dist), the estimated CMF of the truncation distribution (cmf, can be used to adjusted new data), a data frame containing the observed truncated data, latest observed data and the adjusted for truncation observations (obs), a data frame containing the last observed data (last_obs, useful for plotting and validation), the data used for fitting (data) and the fit object (fit).
Author(s)

Sam Abbott
Sebastian Funk

Examples

# set number of cores to use
old_opts <- options()
options(mc.cores = ifelse(interactive(), 4, 1))

# get example case counts
reported_cases <- example_confirmed[1:60]

# define example truncation distribution (note not integer adjusted)
trunc <- dist_spec(
    mean = convert_to_logmean(3, 2),
    mean_sd = 0.1,
    sd = convert_to_logsd(3, 2),
    sd_sd = 0.1,
    max = 10
)

# apply truncation to example data
construct_truncation <- function(index, cases, dist) {
    set.seed(index)
    if (dist$dist == 0) {
        dfunc <- dlnorm
    } else {
        dfunc <- dgamma
    }
    cmf <- cumsum(
        dfunc(
            1:(dist$max + 1),
            rnorm(1, dist$mean_mean, dist$mean_sd),
            rnorm(1, dist$sd_mean, dist$sd_sd)
        )
    )
    cmf <- cmf / cmf[dist$max + 1]
    cmf <- rev(cmf)[-1]
    trunc_cases <- data.table::copy(cases)[1:(.N - index)]
    trunc_cases[
        (.N - length(cmf) + 1):.N, confirm := as.integer(confirm * cmf)
    ]
    return(trunc_cases)
}

example_data <- purrr::map(c(20, 15, 10, 0),
    construct_truncation,
    cases = reported_cases,
    dist = trunc
)

# fit model to example data
example_confirmed

example_confirmed <- estimate_truncation(example_data,
   verbose = interactive(),
   chains = 2, iter = 2000
)

# summary of the distribution
est$dist
# summary of the estimated truncation cmf (can be applied to new data)
print(est$cmf)
# observations linked to truncation adjusted estimates
print(est$obs)
# validation plot of observations vs estimates
plot(est)

options(oldOpts)

example_confirmed  Example Confirmed Case Data Set

Description

[Stable] An example data frame of observed cases

Usage

example_confirmed

Format

A data frame containing cases reported on each date.

expose_stan_fns  Expose internal package stan functions in R

Description

[Stable] his function exposes internal stan functions in R from a user supplied list of target files. Allows for testing of stan functions in R and potentially user use in R code.

Usage

expose_stan_fns(files, target_dir, ...)

Arguments

files  A character vector indicating the target files.
target_dir  A character string indicating the target directory for the file.
...  Additional arguments passed to rstan::expose_stan_functions.
extract_CrIs

Extract Credible Intervals Present

Description

[Stable] Helper function to extract the credible intervals present in a data frame.

Usage

extract_CrIs(summarised)

Arguments

summarised A data frame as processed by calc_CrIs

Value

A numeric vector of credible intervals detected in the data frame.

Examples

samples <- data.frame(value = 1:10, type = "car")
summarised <- calc_CrIs(samples,
summarise_by = "type",
CrIs = c(seq(0.05, 0.95, 0.05))
)
extract_CrIs(summarised)

event_inits

Generate initial conditions from a Stan fit

Description

[Experimental] Extracts posterior samples to use to initialise a full model fit. This may be useful for certain data sets where the sampler gets stuck or cannot easily be initialised. In estimate_infections(), epinow() and regional_epinow() this option can be engaged by setting stan_opts(init_fit = <stanfit>). This implementation is based on the approach taken in epidemia authored by James Scott.

Usage

extract_inits(fit, current_inits, exclude_list = NULL, samples = 50)
Arguments

fit A stanfit object.

current_inits A function that returns a list of initial conditions (such as `create_initial_conditions()`). Only used in `exclude_list` is specified.

exclude_list A character vector of parameters to not initialise from the fit object, defaulting to NULL.

samples Numeric, defaults to 50. Number of posterior samples.

Value

A function that when called returns a set of initial conditions as a named list.

Author(s)

Sam Abbott

---

**extract_parameter** Extract Samples for a Parameter from a Stan model

**Description**

[Stable] Extracts a single from a list of stan output and returns it as a data.table.

**Usage**

`extract_parameter(param, samples, dates)`

**Arguments**

`param` Character string indicating the parameter to extract

`samples` Extracted stan model (using `rstan::extract`)

`dates` A vector identifying the dimensionality of the parameter to extract. Generally this will be a date.

**Value**

A data frame containing the parameter name, date, sample id and sample value.

**Author(s)**

Sam Abbott
extract_parameter_samples

Extract Parameter Samples from a Stan Model

Description

[Stable] Extracts a custom set of parameters from a stan object and adds stratification and dates where appropriate.

Usage

extract_parameter_samples(
  stan_fit,
  data,
  reported_dates,
  reported_inf_dates,
  drop_length_1 = FALSE,
  merge = FALSE
)

Arguments

  stan_fit  A fit Stan model as returned by rstan::sampling.
  data      A list of the data supplied to the rstan::sampling call.
  reported_dates  A vector of dates to report estimates for.
  reported_inf_dates  A vector of dates to report infection estimates for.
  drop_length_1   Logical; whether the first dimension should be dropped if it is off length 1; this is necessary when processing simulation results.
  merge     if TRUE, merge samples and data so that parameters can be extracted from data.

Value

  A list of dataframes each containing the posterior of a parameter

Author(s)

  Sam Abbott
Extract a Parameter Summary from a Stan Object

Description

[Stable] Extracts summarised parameter posteriors from a stanfit object using rstan::summary in a format consistent with other summary functions in EpiNow2.

Usage

```r
extract_stan_param(
  fit, 
  params = NULL, 
  CrIs = c(0.2, 0.5, 0.9), 
  var_names = FALSE 
)
```

Arguments

- `fit` A stanfit objec.
- `params` A character vector of parameters to extract. Defaults to all parameters.
- `CrIs` Numeric vector of credible intervals to calculate.
- `var_names` Logical defaults to FALSE. Should variables be named. Automatically set to TRUE if multiple parameters are to be extracted.

Value

A data.table summarising parameter posteriors. Contains a following variables: variable, mean, mean_se, sd, median, and lower_, upper_ followed by credible interval labels indicating the credible intervals present.

Author(s)

Sam Abbott

Extract Samples from a Parameter with a Single Dimension

Description

Extract Samples from a Parameter with a Single Dimension
Usage

```
extract_static_parameter(param, samples)
```

Arguments

- `param`: Character string indicating the parameter to extract
- `samples`: Extracted stan model (using rstan::extract)

Value

A data frame containing the parameter name, sample id and sample value

Author(s)

Sam Abbott

---

**filter_OPTS**

Filter Options for a Target Region

Description

[Maturing] A helper function that allows the selection of region specific settings if present and otherwise applies the overarching settings.

Usage

```
filter_opts(opts, region)
```

Arguments

- `opts`: Either a list of calls to an _opts() function or a single call to an _opts() function.
- `region`: A character string indicating a region of interest.

Value

A list of options

Author(s)

Sam Abbott
fit_model_with_nuts  

Fit a Stan Model using the NUTs sampler

Description

[Maturing] Fits a stan model using rstan::sampling. Provides the optional ability to run chains using future with error catching, timeouts and merging of completed chains.

Usage

fit_model_with_nuts(
  args,
  future = FALSE,
  max_execution_time = Inf,
  id = "stan"
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>args</td>
<td>List of stan arguments.</td>
</tr>
<tr>
<td>future</td>
<td>Logical, defaults to FALSE. Should future be used to run stan chains in parallel.</td>
</tr>
<tr>
<td>max_execution_time</td>
<td>Numeric, defaults to Inf. What is the maximum execution time per chain in seconds. Results will still be returned as long as at least 2 chains complete successfully within the timelimit.</td>
</tr>
<tr>
<td>id</td>
<td>A character string used to assign logging information on error. Used by regional_epinow to assign errors to regions. Alter the default to run with error catching.</td>
</tr>
</tbody>
</table>

Value

A stan model object

Author(s)

Sam Abbott

fit_model_with_vb  

Fit a Stan Model using Variational Inference

Description


Usage

fit_model_with_vb(args, future = FALSE, id = "stan")
forecast_secondary

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>args</td>
<td>List of stan arguments.</td>
</tr>
<tr>
<td>future</td>
<td>Logical, defaults to FALSE. Should future be used to run stan chains in parallel.</td>
</tr>
<tr>
<td>id</td>
<td>A character string used to assign logging information on error. Used by regional_epinow to assign errors to regions. Alter the default to run with error catching.</td>
</tr>
</tbody>
</table>

Value

A stan model object

Author(s)

Sam Abbott

Description

[Experimental] This function forecasts secondary observations using the output of estimate_secondary() and either observed primary data or a forecast of primary observations. See the examples of estimate_secondary() for one use case. It can also be combined with estimate_infections() to produce a forecast for a secondary observation from a forecast of a primary observation. See the examples of estimate_secondary() for example use cases on synthetic data. See here for an example of forecasting Covid-19 deaths from Covid-19 cases.

Usage

```r
forecast_secondary(
  estimate,
  primary,
  primary_variable = "reported_cases",
  model = NULL,
  samples = NULL,
  all_dates = FALSE,
  CrIs = c(0.2, 0.5, 0.9)
)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>estimate</td>
<td>An object of class &quot;estimate_secondary&quot; as produced by estimate_secondary().</td>
</tr>
<tr>
<td>primary</td>
<td>A data.frame containing at least date and value (integer) variables and optionally sample. Used as the primary observation used to forecast the secondary observations. Alternatively, this may be an object of class &quot;estimate_infections&quot; as produced by estimate_infections(). If primary is of class &quot;estimate_infections&quot; then the internal samples will be filtered to have a minimum date ahead of those observed in the estimate object.</td>
</tr>
</tbody>
</table>
`format_fit` 59

**primary_variable**
A character string indicating the primary variable, defaulting to "reported\_cases". Only used when primary is of class "estimate\_infections".

**model**
A compiled stan model as returned by `rstan::stan_model`.

**samples**
Numeric, number of posterior samples to simulate from. The default is to use all samples in the primary input when present. If not present the default is to use 1000 samples.

**all_dates**
Logical, defaults to FALSE. Should a forecast for all dates and not just those in the forecast horizon be returned.

**CrIs**
Numeric vector of credible intervals to calculate.

**Value**
A list containing: predictions (a data frame ordered by date with the primary, and secondary observations, and a summary of the forecast secondary observations. For primary observations in the forecast horizon when uncertainty is present the median is used), samples a data frame of forecast secondary observation posterior samples, and forecast a summary of the forecast secondary observation posterior.

**Author(s)**
Sam Abbott

**See Also**

`estimate_secondary`

---

**format_fit**  Format Posterior Samples

**Description**

[Stable] Summaries posterior samples and adds additional custom variables.

**Usage**

`format_fit(posterior_samples, horizon, shift, burn_in, start_date, CrIs)`

**Arguments**

- **posterior_samples**
  A list of posterior samples as returned by `extract\_parameter\_samples`.
- **horizon**
  Numeric, forecast horizon.
- **shift**
  Numeric, the shift to apply to estimates.
- **burn_in**
  Numeric, number of days to discard estimates for.
- **start_date**
  Date, earliest date with data.
- **CrIs**
  Numeric vector of credible intervals to calculate.
Value

A list of samples and summarised posterior parameter estimates.

Author(s)

Sam Abbott

---

**gamma_dist_def**

*Generate a Gamma Distribution Definition Based on Parameter Estimates*

Description

*[Soft-deprecated]* Generates a distribution definition when only parameter estimates are available for gamma distributed parameters. See `rgamma` for distribution information.

Usage

```r
gamma_dist_def(
  shape, shape_sd, scale, scale_sd, mean, mean_sd, sd, sd_sd, max_value, samples
)
```

Arguments

- `shape`: Numeric, shape parameter of the gamma distribution.
- `shape_sd`: Numeric, standard deviation of the shape parameter.
- `scale`: Numeric, scale parameter of the gamma distribution.
- `scale_sd`: Numeric, standard deviation of the scale parameter.
- `mean`: Numeric, log mean parameter of the gamma distribution.
- `mean_sd`: Numeric, standard deviation of the log mean parameter.
- `sd`: Numeric, log sd parameter of the gamma distribution.
- `sd_sd`: Numeric, standard deviation of the log sd parameter.
- `max_value`: Numeric, the maximum value to allow. Defaults to 120. Samples outside of this range are resampled.
- `samples`: Numeric, number of sample distributions to generate.
Value

A data.table defining the distribution as used by dist_skel

Author(s)

Sam Abbott

Examples

```r
# using estimated shape and scale
def <- gamma_dist_def(
  shape = 5.807, shape_sd = 0.2,
  scale = 0.9, scale_sd = 0.05,
  max_value = 20, samples = 10
)
print(def)
def$params[[1]]

# using mean and sd
def <- gamma_dist_def(
  mean = 3, mean_sd = 0.5,
  sd = 3, sd_sd = 0.1,
  max_value = 20, samples = 10
)
print(def)
def$params[[1]]
```

Description


Usage

generation_times

Format

A data.table of summarising the distribution
**generation_time_opts**  
*Generation Time Distribution Options*

**Description**

[Stable] Returns generation time parameters in a format for lower level model use. The generation time can either be given as a disease and source to be passed to `get_generation_time`, or as parameters of a distribution to be passed to `dist_spec`.

**Usage**

```r
generation_time_opts(
  dist = dist_spec(mean = 1),
  ..., 
  disease, 
  source, 
  max = 15L, 
  fixed = FALSE, 
  prior_weight 
)
```

**Arguments**

- **dist**: A delay distribution or series of delay distributions generated using `dist_spec()` or `get_generation_time()`. If no distribution is given a fixed generation time of 1 will be assumed.
- **...**: deprecated; use `dist` instead
- **disease**: deprecated; use `dist` instead
- **source**: deprecated; use `dist` instead
- **max**: deprecated; use `dist` instead
- **fixed**: deprecated; use `dist` instead
- **prior_weight**: deprecated; prior weights are now specified as a model option. Use the `weigh_delay_priors` argument of `estimate_infections` instead.

**Value**

A list summarising the input delay distributions.

**Author(s)**

Sebastian Funk  
Sam Abbott

**See Also**

`convert_to_logmean` `convert_to_logsd` `bootstrapped_dist_fit` `dist_spec`
get_dist

Examples

# default settings with a fixed generation time of 1
generation_time_opts()

# A fixed gamma distributed generation time
generation_time_opts(dist_spec(mean = 3, sd = 2, max = 15))

# An uncertain gamma distributed generation time
generation_time_opts(
  dist_spec(mean = 3, sd = 2, mean_sd = 1, sd_sd = 0.5, max = 15)
)

# A generation time sourced from the literature
dist <- get_generation_time(disease = "SARS-CoV-2", source = "ganyani")
generation_time_opts(dist)

get_dist(data, disease, source, max_value = 15, fixed = FALSE)

Arguments

data A data.table in the format of generation_times.
disease A character string indicating the disease of interest.
source A character string indicating the source of interest.
max_value Numeric, the maximum value to allow. Defaults to 15 days.
fixed Logical, defaults to FALSE. Should distributions be supplied as fixed values (vs with uncertainty)?

Value

A list defining a distribution

Author(s)

Sam Abbott
Examples

```r
get_dist(  
  EpiNow2::generation_times, disease = "SARS-CoV-2", source = "ganyani"
)
```

---

get_generation_time  
*Get a Literature Distribution for the Generation Time*

Description

[Stable] Extracts a literature distribution from `generation_times`.

Usage

```r
get_generation_time(disease, source, max_value = 15, fixed = FALSE)
```

Arguments

- **disease**: A character string indicating the disease of interest.
- **source**: A character string indicating the source of interest.
- **max_value**: Numeric, the maximum value to allow. Defaults to 15 days.
- **fixed**: Logical, defaults to `FALSE`. Should distributions be supplied as fixed values (vs with uncertainty)?

Value

A list defining a distribution

Author(s)

Sam Abbott

Examples

```r
get_generation_time(disease = "SARS-CoV-2", source = "ganyani")
```
get_incubation_period  Get a Literature Distribution for the Incubation Period

Description

[Stable] Extracts a literature distribution from incubation_periods.

Usage

get_incubation_period(disease, source, max_value = 15, fixed = FALSE)

Arguments

disease  A character string indicating the disease of interest.
source   A character string indicating the source of interest.
max_value Numeric, the maximum value to allow. Defaults to 15 days.
fixed    Logical, defaults to FALSE. Should distributions be supplied as fixed values (vs with uncertainty)?

Value

A list defining a distribution

Author(s)

Sam Abbott

Examples

get_incubation_period(disease = "SARS-CoV-2", source = "lauer")

get_raw_result  Get a Single Raw Result

Description

[Stable]

Usage

get_raw_result(file, region, date, result_dir)
get_regional_results

Arguments

file Character string giving the result files name.
region Character string giving the region of interest.
date Target date (in the format "yyyy-mm-dd").
result_dir Character string giving the location of the target directory.

Value

An R object read in from the targeted .rds file

Author(s)

Sam Abbott

get_regional_results  Get Combined Regional Results

Description

[Stable] Summarises results across regions either from input or from disk. See the examples for details.

Usage

get_regional_results(
  regional_output,  # A list of output as produced by regional_epinow and stored in the regional list.
  results_dir,  # A character string indicating the folder containing the EpiNow2 results to extract.
  date,  # A Character string (in the format "yyyy-mm-dd") indicating the date to extract data for. Defaults to "latest" which finds the latest results available.
  samples = TRUE,  # Logical, defaults to TRUE. Should samples be returned.
  forecast = FALSE  # Logical, defaults to FALSE. Should forecast results be returned.
)

Arguments

regional_output
results_dir
date
samples
forecast

Value

A list of estimates, forecasts and estimated cases by date of report.
get_regions

Author(s)

Sam Abbott

Examples

# construct example distributions
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- estimate_delay(rlnorm(100, log(6), 1), max_value = 10)

# example case vector
cases <- example_confirmed[1:30]
cases <- data.table::rbindlist(list(
  data.table::copy(cases)[, region := "testland"],
  cases[, region := "realland"]
))

# save results to tmp folder
dir <- file.path(tempdir(check = TRUE), "results")
# run multiregion estimates
regional_out <- regional_epinow(
  reported_cases = cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  rt = rt_opts(rw = 7), gp = NULL,
  output = c("regions", "latest"),
  target_folder = dir,
  return_output = TRUE
)
# from output
results <- get_regional_results(regional_out$regional, samples = FALSE)
names(results)

# from a folder
folder_results <- get_regional_results(results_dir = dir, samples = FALSE)
names(folder_results)

---

get_regions

Get Folders with Results

Description

[Stable]
get_regions_with_most_reports

Get Regions with Most Reported Cases

Description

[Stable] Extract a vector of regions with the most reported cases in a set time window.

Usage

get_regions_with_most_reports(reported_cases, time_window = 7, no_regions = 6)

Arguments

reported_cases A data frame of confirmed cases (confirm) by date (date), and region (region).
time_window Numeric, number of days to include from latest date in data. Defaults to 7 days.
no_regions Numeric, number of regions to return. Defaults to 6.

Value

A character vector of regions with the highest reported cases

Author(s)

Sam Abbott
get_seeding_time

Estimate seeding time from delays and generation time

Description

The seeding time is set to the mean of the specified delays, constrained to be at least the maximum generation time.

Usage

get_seeding_time(delays, generation_time)

Arguments

delays Delays as specified using dist_spec

generation_time Generation time as specified using dist_spec

Value

An integer seeding time

Author(s)

Sebastian Funk

gp_opts

Approximate Gaussian Process Settings

Description

[Stable] Defines a list specifying the structure of the approximate Gaussian process. Custom settings can be supplied which override the defaults.

Usage

gp_opts(
    basis_prop = 0.2,
    boundary_scale = 1.5,
    ls_mean = 21,
    ls_sd = 7,
    ls_min = 0,
    ls_max = 60,
    alpha_sd = 0.05,
    kernel = "matern",
    matern_type = 3/2
    )
Arguments

- **basis_prop**
  - Numeric, proportion of time points to use as basis functions. Defaults to 0.2. Decreasing this value results in a decrease in accuracy but a faster compute time (with increasing it having the first effect). In general smaller posterior length scales require a higher proportion of basis functions. See (Riutort-Mayol et al. 2020 https://arxiv.org/abs/2004.11408) for advice on updating this default.

- **boundary_scale**

- **ls_mean**
  - Numeric, defaults to 21 days. The mean of the lognormal length scale.

- **ls_sd**
  - Numeric, defaults to 7 days. The standard deviation of the log normal length scale. If \( ls\_sd = 0 \), inverse-gamma prior on Gaussian process length scale will be used with recommended parameters \( \text{inv\_gamma}(1.499007, 0.057277 * \text{ls\_max}) \).

- **ls_min**
  - Numeric, defaults to 0. The minimum value of the length scale.

- **ls_max**
  - Numeric, defaults to 60. The maximum value of the length scale. Updated in create_gp_data to be the length of the input data if this is smaller.

- **alpha_sd**
  - Numeric, defaults to 0.05. The standard deviation of the magnitude parameter o the Gaussian process kernel. Should be approximately the expected standard deviation of the logged Rt.

- **kernel**
  - Character string, the type of kernel required. Currently supporting the squared exponential kernel ("se") and the 3 over 2 Matern kernel ("matern", with \( \text{matern\_type} = 3/2 \)). Defaulting to the Matern 3 over 2 kernel as discontinuities are expected in Rt and infections.

- **matern_type**
  - Numeric, defaults to 3/2. Type of Matern Kernel to use. Currently only the Matern 3/2 kernel is supported.

Value

A list of settings defining the Gaussian process

Author(s)

Sam Abbott

Examples

```r
# default settings
gp_opts()

# add a custom length scale
gp_opts(ls_mean = 4)
```
growth_to_R

Convert Growth Rates to Reproduction numbers.

Description

[Questioning] See here # nolint for justification. Now handled internally by stan so may be removed in future updates if no user demand.

Usage

growth_to_R(r, gamma_mean, gamma_sd)

Arguments

r Numeric, rate of growth estimates.

gamma_mean Numeric, mean of the gamma distribution

gamma_sd Numeric, standard deviation of the gamma distribution.

Value

Numeric vector of reproduction number estimates

Examples

growth_to_R(0.2, 4, 1)

incubation_periods

Literature Estimates of Incubation Periods

Description


Usage

incubation_periods

Format

A data.table of summarising the distribution
init_cumulative_fit  Generate initial conditions by fitting to cumulative cases

Description

[Experimental] Fits a model to cumulative cases. This may be a useful approach to initialising a full model fit for certain data sets where the sampler gets stuck or cannot easily be initialised as fitting to cumulative cases changes the shape of the posterior distribution. In `estimate_infections()`, `epinow()` and `regional_epinow()` this option can be engaged by setting `stan_opts(init_fit = "cumulative")`.

This implementation is based on the approach taken in `epidemia` authored by James Scott.

Usage

```r
init_cumulative_fit(
  args,
  samples = 50,
  warmup = 50,
  id = "init",
  verbose = FALSE
)
```

Arguments

- **args**: List of stan arguments.
- **samples**: Numeric, defaults to 50. Number of posterior samples.
- **warmup**: Numeric, defaults to 50. Number of warmup samples.
- **id**: A character string used to assign logging information on error. Used by `regional_epinow` to assign errors to regions. Alter the default to run with error catching.
- **verbose**: Logical, should fitting progress be returned. Defaults to FALSE.

Value

A stanfit object

Author(s)

Sam Abbott
Generate a Log Normal Distribution Definition Based on Parameter Estimates

Description

[Soft-deprecated] Generates a distribution definition when only parameter estimates are available for log normal distributed parameters. See rlnorm for distribution information.

Usage

lognorm_dist_def(mean, mean_sd, sd, sd_sd, max_value, samples, to_log = FALSE)

Arguments

- **mean**: Numeric, log mean parameter of the gamma distribution.
- **mean_sd**: Numeric, standard deviation of the log mean parameter.
- **sd**: Numeric, log sd parameter of the gamma distribution.
- **sd_sd**: Numeric, standard deviation of the log sd parameter.
- **max_value**: Numeric, the maximum value to allow. Defaults to 120. Samples outside of this range are resampled.
- **samples**: Numeric, number of sample distributions to generate.
- **to_log**: Logical, should parameters be logged before use.

Value

A data.table defining the distribution as used by dist_skel

Author(s)

Sam Abbott

Examples

```r
# Example 1
def1 <- lognorm_dist_def(
  mean = 1.621, mean_sd = 0.0640,
  sd = 0.418, sd_sd = 0.0691,
  max_value = 20, samples = 10
)
print(def1)
def1$params[[1]]

# Example 2
def2 <- lognorm_dist_def(
  mean = 5, mean_sd = 1,
  sd = 3, sd_sd = 1,
  max_value = 20, samples = 10,
  to_log = TRUE
)```
map_prob_change

make_conf

Format Credible Intervals

Description

[Stable] Combines a list of values into formatted credible intervals.

Usage

make_conf(value, CrI = 90, reverse = FALSE)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>value</td>
<td>List of value to map into a string. Requires, point, lower, and upper.</td>
</tr>
<tr>
<td>CrI</td>
<td>Numeric, credible interval to report. Defaults to 90.</td>
</tr>
<tr>
<td>reverse</td>
<td>Logical, defaults to FALSE. Should the reported credible interval be switched.</td>
</tr>
</tbody>
</table>

Value

A character vector formatted for reporting

Examples

```r
value <- list(median = 2, lower_90 = 1, upper_90 = 3)
make_conf(value)
```

map_prob_change

Categorise the Probability of Change for Rt

Description

[Stable] Categorises a numeric variable into "Increasing" (< 0.05), "Likely increasing" (< 0.4), "Stable" (< 0.6), "Likely decreasing" (< 0.95), "Decreasing" (<= 1)

Usage

map_prob_change(var)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>var</td>
<td>Numeric variable to be categorised</td>
</tr>
</tbody>
</table>
**match_output_arguments**

**Value**

A character variable.

**Examples**

```r
var <- seq(0.01, 1, 0.01)
var
map_prob_change(var)
```

**Description**

[Stable] Match user supplied arguments with supported options and return a logical list for internal usage.

**Usage**

```r
match_output_arguments(
  input_args = NULL,
  supported_args = NULL,
  logger = NULL,
  level = "info"
)
```

**Arguments**

- `input_args` A character vector of input arguments (can be partial).
- `supported_args` A character vector of supported output arguments.
- `logger` A character vector indicating the logger to target messages at. Defaults to no logging.
- `level` Character string defaulting to "info". Logging level see documentation of futile.logger for details. Supported options are "info" and "debug".

**Value**

A logical vector of named output arguments
mean.dist_spec

Returns the mean of one or more delay distribution

Description

This works out the mean of all the (parametric / nonparametric) delay distributions combined in the passed dist_spec().

Usage

## S3 method for class 'dist_spec'
mean(x, ...)

Arguments

x The dist_spec() to use
...
Not used

Value

A vector of means.

Author(s)

Sebastian Funk

Examples

# A fixed lognormal distribution with mean 5 and sd 1.
lognormal <- dist_spec(
  mean = 5, sd = 1, max = 20, distribution = "lognormal"
)
mean(lognormal)

# An uncertain gamma distribution with mean 3 and sd 2
gamma <- dist_spec(
  mean = 3, sd = 2, mean_sd = 0.5, sd_sd = 0.5, max = 20,
  distribution = "gamma"
)
mean(gamma)

# The mean of the sum of two distributions
mean(lognormal + gamma)
**obs_opts**

**Observation Model Options**

**Description**

[Stable] Defines a list specifying the structure of the observation model. Custom settings can be supplied which override the defaults.

**Usage**

```r
obs_opts(
  family = "negbin",
  phi = c(0, 1),
  weight = 1,
  week_effect = TRUE,
  week_length = 7,
  scale = list(),
  likelihood = TRUE,
  return_likelihood = FALSE
)
```

**Arguments**

- **family** Character string defining the observation model. Options are Negative binomial ("negbin"), the default, and Poisson.
- **phi** A numeric vector of length 2, defaults to 0, 1. Indicates the mean and standard deviation of the normal prior used for the observation process.
- **weight** Numeric, defaults to 1. Weight to give the observed data in the log density.
- **week_effect** Logical defaulting to TRUE. Should a day of the week effect be used in the observation model.
- **week_length** Numeric assumed length of the week in days, defaulting to 7 days. This can be modified if data aggregated over a period other than a week or if data has a non-weekly periodicity.
- **scale** List, defaulting to an empty list. Should an scaling factor be applied to map latent infections (convolved to date of report). If none empty a mean (mean) and standard deviation (sd) needs to be supplied defining the normally distributed scaling factor.
- **likelihood** Logical, defaults to TRUE. Should the likelihood be included in the model.
- **return_likelihood** Logical, defaults to FALSE. Should the likelihood be returned by the model.

**Value**

A list of observation model settings.
Examples

# default settings
obs_opts()

# Turn off day of the week effect
obs_opts(week_effect = TRUE)

# Scale reported data
obs_opts(scale = list(mean = 0.2, sd = 0.02))

Description

[Maturing] Define a list of _opts() to pass to regional_epinow _opts() accepting arguments. This is useful when different settings are needed between regions within a single regional_epinow call. Using opts_list the defaults can be applied to all regions present with an override passed to regions as necessary (either within opts_list or externally).

Usage

opts_list(opts, reported_cases, ...)

Arguments

- **opts**: An _opts() function call such as rt_opts().
- **reported_cases**: A data frame containing a region variable indicating the target regions.
- **...**: Optional override for region defaults. See the examples for use case.

Value

A named list of options per region which can be passed to the _opt accepting arguments of regional_epinow.

Author(s)

Sam Abbott

See Also

regional_epinow rt_opts
Examples

# uses example case vector
cases <- example_confirmed[1:40]
cases <- data.table::rbindlist(list(
    data.table::copy(cases)[, region := "testland"],
    cases[, region := "realland"]
))

# default settings
opts_list(rt_opts(), cases)

# add a weekly random walk in realland
opts_list(rt_opts(), cases, realland = rt_opts(rw = 7))

# add a weekly random walk externally
rt <- opts_list(rt_opts(), cases)
rt$realland$rw <- 7
rt

plot.dist_spec  

Plot PMF and CDF for a dist_spec object

Description

This function takes a dist_spec object and plots its probability mass function (PMF) and cumulative distribution function (CDF) using ggplot2. Note that currently uncertainty in distributions is not plot.

Usage

## S3 method for class 'dist_spec'
plot(x, ...)

Arguments

x  
A dist_spec object

...  
Additional arguments to pass to ggplot

Author(s)

Sam Abbott

Examples

#' # A fixed lognormal distribution with mean 5 and sd 1.
lognormal <- dist_spec(
  mean = 1.6, sd = 0.5, max = 20, distribution = "lognormal"
)
plot(lognormal)
# An uncertain gamma distribution with mean 3 and sd 2
gamma <- dist_spec(
    mean = 3, sd = 2, mean_sd = 0.5, sd_sd = 0.5, max = 20,
    distribution = "gamma"
)
plot(gamma)

# Multiple distributions
plot(lognormal + gamma + lognormal)

# A combination of the two fixed distributions
plot(lognormal + lognormal)

plot.epinow  
Plot method for epinow

Description

[Maturing] plot method for class "epinow".

Usage

## S3 method for class 'epinow'
plot(x, type = "summary", ...)

Arguments

x  A list of output as produced by epinow

type  A character vector indicating the name of plots to return. Defaults to "summary" with supported options being "infections", "reports", "R", "growth_rate", "summary", "all".

...  Pass additional arguments to report_plots

Value

List of plots as produced by report_plots

See Also

plot plot.epinow plot.estimate_infections report_plots estimate_infections
plot.estimate_infections

Plot method for estimate_infections

Description

[Maturing] plot method for class "estimate_infections".

Usage

## S3 method for class 'estimate_infections'
plot(x, type = "summary", ...)

Arguments

x A list of output as produced by estimate_infections

type A character vector indicating the name of plots to return. Defaults to "summary" with supported options being "infections", "reports", "R", "growth_rate", "summary", "all".

... Pass additional arguments to report_plots

Value

List of plots as produced by report_plots

See Also

plot report_plots estimate_infections

plot.estimate_secondary

Plot method for estimate_secondary

Description

[Experimental] plot method for class "estimate_secondary".

Usage

## S3 method for class 'estimate secondary'
plot(x, primary = FALSE, from = NULL, to = NULL, new_obs = NULL, ...)


#### Arguments

- **x**: A list of output as produced by `estimate_secondary`
- **primary**: Logical, defaults to `FALSE`. Should primary reports also be plot?
- **from**: Date object indicating when to plot from.
- **to**: Date object indicating when to plot up to.
- **new_obs**: A data.frame containing the columns `date` and `secondary` which replace the secondary observations stored in the `estimate_secondary` output.
- **...**: Pass additional arguments to plot function. Not currently in use.

#### Value

A `ggplot` object.

#### Author(s)

Sam Abbott

#### See Also

`plot estimate_secondary`

---

**plot.estimate_truncation**

*Plot method for estimate_truncation*

#### Description

[Experimental] plot method for class "estimate_truncation". Returns a plot faceted over each dataset used in fitting with the latest observations as columns, the data observed at the time (and so truncated) as dots and the truncation adjusted estimates as a ribbon.

#### Usage

```r
## S3 method for class 'estimate_truncation'
plot(x, ...)  
```

#### Arguments

- **x**: A list of output as produced by `estimate_truncation`
- **...**: Pass additional arguments to plot function. Not currently in use.

#### Value

`ggplot2` object
**plot_CrIs**

**Author(s)**
Sam Abbott

**See Also**
plot estimate_truncation

---

**plot_CrIs**  
*Plot EpiNow2 Credible Intervals*

**Description**

[Stable] Adds lineranges for user specified credible intervals

**Usage**

```r
plot_CrIs(plot, CrIs, alpha, linewidth)
```

**Arguments**

- `plot`: A ggplot2 plot
- `CrIs`: Numeric list of credible intervals present in the data. As produced by `extract_CrIs`
- `alpha`: Numeric, overall alpha of the target line range
- `linewidth`: Numeric, line width of the default line range.

**Value**

A ggplot2 plot.

---

**plot_estimates**  
*Plot Estimates*

**Description**

[Questioning] Allows users to plot the output from `estimate_infections` easily. In future releases it may be depreciated in favour of increasing the functionality of the S3 plot methods.

**Usage**

```r
plot_estimates(
    estimate,  
    reported,  
    ylab = "Cases",  
    hline,  
    obs_as_col = TRUE,  
    max_plot = 10,  
    estimate_type = NULL
)
```
Arguments

- **estimate**: A data.table of estimates containing the following variables: date, type (must contain "estimate", "estimate based on partial data" and optionally "forecast").
- **reported**: A data.table of reported cases with the following variables: date, confirm.
- **ylab**: Character string, defaulting to "Cases". Title for the plot y axis.
- **hline**: Numeric, if supplied gives the horizontal intercept for a indicator line.
- **obs_as_col**: Logical, defaults to TRUE. Should observed data, if supplied, be plotted using columns or as points (linked using a line).
- **max_plot**: Numeric, defaults to 10. A multiplicative upper bound on the number of cases shown on the plot. Based on the maximum number of reported cases.
- **estimate_type**: Character vector indicating the type of data to plot. Default to all types with supported options being: "Estimate", "Estimate based on partial data", and "Forecast".

Value

A ggplot2 object

Examples

```r
# define example cases
cases <- example_confirmed[1:40]

# set up example delays
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- estimate_delay(rlnorm(100, log(6), 1), max_value = 10)

# run model
out <- estimate_infections(cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay)
)

# plot infections
plot_estimates(
  estimate = out$summarised[variable == "infections"],
  reported = cases,
  ylab = "Cases", max_plot = 2
) + ggplot2::facet_wrap(~type, scales = "free_y")

# plot reported cases estimated via Rt
plot_estimates(
  estimate = out$summarised[variable == "reported_cases"],
  reported = cases,
```
plot_summary

Plot a Summary of the Latest Results

Description

[Questioning] Used to return a summary plot across regions (using results generated by summarise_results). May be depreciated in later releases in favour of enhanced S3 methods.

Usage

plot_summary(summary_results, x_lab = "Region", log_cases = FALSE, max_cases)

Arguments

summary_results
  A data.table as returned by summarise_results (the data object).

x_lab
  A character string giving the label for the x axis, defaults to region.

log_cases
  Logical, should cases be shown on a logged scale. Defaults to FALSE.

max_cases
  Numeric, no default. The maximum number of cases to plot.

Value

A ggplot2 object
print.dist_spec  

Prints the parameters of one or more delay distributions

Description

This displays the parameters of the uncertain and probability mass functions of fixed delay distributions combined in the passed dist_spec().

Usage

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

Arguments

- `x`  
  The dist_spec() to use
- `...`  
  Not used

Value

invisible

Author(s)

Sebastian Funk

Examples

```r
#' # A fixed lognormal distribution with mean 5 and sd 1.
lognormal <- dist_spec(
  mean = 1.5, sd = 0.5, max = 20, distribution = "lognormal"
)
print(lognormal)

#' # An uncertain gamma distribution with mean 3 and sd 2
gamma <- dist_spec(
  mean = 3, sd = 2, mean_sd = 0.5, sd_sd = 0.5, max = 20,
  distribution = "gamma"
)
print(gamma)
```
process_region

**Process regional estimate**

**Description**

**[Maturing]** Internal function that removes output that is not required, and returns logging information.

**Usage**

```r
code
process_region(
  out,
  target_region,
  timing,
  return_output = TRUE,
  return_timing = TRUE,
  complete_logger = "EpiNow2.epinow"
)
```

**Arguments**

- `out`: List of output returned by `epinow`
- `target_region`: Character string indicating the region being evaluated
- `timing`: Output from `Sys.time`
- `return_output`: Logical, defaults to `FALSE`. Should output be returned, this automatically updates to `TRUE` if no directory for saving is specified.
- `return_timing`: Logical, should runtime be returned
- `complete_logger`: Character string indicating the logger to output the completion of estimation to.

**Value**

A list of processed output

**See Also**

`regional_epinow`
process_regions Process all Region Estimates

Description

[Stable] Internal function that processes the output from multiple epinow runs, adds summary logging information.

Usage

process_regions(regional_out, regions)

Arguments

regional_out A list of output from multiple runs of regional_epinow
regions A character vector identifying the regions that have been run

Value

A list of all regional estimates and successful regional estimates

See Also

regional_epinow epinow

region_epinow Real-time Rt Estimation, Forecasting and Reporting by Region

Description

[Maturing] Efficiently runs epinow() across multiple regions in an efficient manner and conducts basic data checks and cleaning such as removing regions with fewer than non_zero_points as these are unlikely to produce reasonable results whilst consuming significant resources. See the documentation for epinow for further information.

By default all arguments supporting input from _opts() functions are shared across regions (including delays, truncation, Rt settings, stan settings, and gaussian process settings). Region specific settings are supported by passing a named list of _opts() calls (with an entry per region) to the relevant argument. A helper function (opts_list) is available to facilitate building this list.

Regions can be estimated in parallel using the {future} package (see setup_future). The progress of producing estimates across multiple regions is tracked using the progressr package. Modify this behaviour using progressr::handlers and enable it in batch by setting R_PROGRESSR_ENABLE=TRUE as an environment variable.
Usage

regional_epinow(
  reported_cases,
  generation_time,
  delays = delay_opts(),
  truncation = trunc_opts(),
  rt = rt_opts(),
  backcalc = backcalc_opts(),
  gp = gp_opts(),
  obs = obs_opts(),
  stan = stan_opts(),
  horizon = 7,
  CrIs = c(0.2, 0.5, 0.9),
  target_folder = NULL,
  target_date,
  non_zero_points = 2,
  output = c("regions", "summary", "samples", "plots", "latest"),
  return_output = FALSE,
  summary_args = list(),
  verbose = FALSE,
  logs = tempdir(check = TRUE),
  ...
)

Arguments

reported_cases A data frame of confirmed cases (confirm) by date (date), and region (region).

generation_time A call to generation_time_opts() defining the generation time distribution used. For backwards compatibility a list of summary parameters can also be passed.

delays A call to delay_opts() defining delay distributions and options. See the documentation of delay_opts() and the examples below for details.

truncation A call to trunc_opts() defining the truncation of observed data. Defaults to trunc_opts(). See estimate_truncation() for an approach to estimating truncation from data.

rt A list of options as generated by rt_opts() defining Rt estimation. Defaults to rt_opts(). Set to NULL to switch to using back calculation rather than generating infections using Rt.

backcalc A list of options as generated by backcalc_opts() to define the back calculation. Defaults to backcalc_opts().

gp A list of options as generated by gp_opts() to define the Gaussian process. Defaults to gp_opts(). Set to NULL to disable the Gaussian process.

obs A list of options as generated by obs_opts() defining the observation model. Defaults to obs_opts().
stan A list of stan options as generated by `stan_opts()`. Defaults to `stan_opts()`. Can be used to override data, `init`, and `verbose` settings if desired.

horizon Numeric, defaults to 7. Number of days into the future to forecast.

cris Numeric vector of credible intervals to calculate.

target_folder Character string specifying where to save results (will create if not present).

target_date Date, defaults to maximum found in the data if not specified.

non_zero_points Numeric, the minimum number of time points with non-zero cases in a region required for that region to be evaluated. Defaults to 7.

output A character vector of optional output to return. Supported options are the individual regional estimates ("regions"), samples ("samples"), plots ("plots"), copying the individual region dated folder into a latest folder (if `target_folder` is not null, set using "latest"), the stan fit of the underlying model ("fit"), and an overall summary across regions ("summary"). The default is to return samples and plots alongside summarised estimates and summary statistics. If `target_folder` is not NULL then the default is also to copy all results into a latest folder.

return_output Logical, defaults to FALSE. Should output be returned, this automatically updates to TRUE if no directory for saving is specified.

summary_args A list of arguments passed to `regional_summary`. See the `regional_summary` documentation for details.

verbose Logical defaults to FALSE. Outputs verbose progress messages to the console from `epinow`.

logs Character path indicating the target folder in which to store log information. Defaults to the temporary directory if not specified. Default logging can be disabled if `logs` is set to NULL. If specifying a custom logging setup then the code for `setup_default_logging` and the `setup_logging` function are a sensible place to start.

... Pass additional arguments to `epinow`. See the documentation for `epinow` for details.

Value A list of output stratified at the top level into regional output and across region output summary output

See Also `epinow`, `estimate_infections`, `forecast_infections`, `setup_future`, `regional_summary`

Examples

```r
# set number of cores to use
old_opts <- options()
options(mc.cores = ifelse(interactive(), 4, 1))
```
# construct example distributions

```r
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
```

```r
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
```

```r
reporting_delay <- dist_spec(
  mean = convert_to_logmean(2, 1),
  mean_sd = 0.1,
  sd = convert_to_logsd(2, 1),
  sd_sd = 0.1, max = 15
)
```

# uses example case vector

```r
cases <- example_confirmed[1:60]
cases <- data.table::rbindlist(list(
  data.table::copy(cases)[, region := "testland"],
  cases[, region := "realland"
))
```

# run epinow across multiple regions and generate summaries
# samples and warmup have been reduced for this example

def <- regional_epinow(
  reported_cases = cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  rt = rt_opts(prior = list(mean = 2, sd = 0.2)),
  stan = stan_opts(
    samples = 100, warmup = 200,
    control = list(adapt_delta = 0.95)
  ),
  verbose = interactive()
)

# apply a different rt method per region
# (here a gaussian process and a weekly random walk)

gp <- opts_list(gp_opts(), cases)
gp <- update_list(gp, list(realland = NULL))
rt <- opts_list(rt_opts(), cases, realland = rt_opts(rw = 7))
region_rt <- regional_epinow(
  reported_cases = cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  rt = rt, gp = gp,
  stan = stan_opts(
    samples = 100, warmup = 200,
    control = list(adapt_delta = 0.95)
  ),
  verbose = interactive()
)
options(old_opts)

regional_runtimes  

*Summarise Regional Runtimes*

---

**Description**

*[Maturing]* Used internally by `regional_epinow` to summarise region run times.

**Usage**

```r
regional_runtimes(
  regional_output = NULL,
  target_folder = NULL,
  target_date = NULL,
  return_output = FALSE
)
```

**Arguments**

- `regional_output`: A list of output as produced by `regional_epinow` and stored in the `regional` list.
- `target_folder`: Character string specifying where to save results (will create if not present).
- `target_date`: A character string giving the target date for which to extract results (in the format "yyyy-mm-dd"). Defaults to latest available estimates.
- `return_output`: Logical, defaults to FALSE. Should output be returned, this automatically updates to TRUE if no directory for saving is specified.

**Value**

A data.table of region run times

**See Also**

`regional_summary` `regional_epinow`

**Examples**

```r
# example delays
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
```
regional_summary <- regional_epinow(
  reported_cases = cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  stan = stan_opts(samples = 100, warmup = 100),
  output = c("region", "timing")
)

regional_runtimes(regional_output = regional_out$regional)

---

### regional_summary

#### Regional Summary Output

**Description**

[Maturing] Used to produce summary output either internally in `regional_epinow` or externally.

**Usage**

```r
regional_summary(
  regional_output = NULL,
  reported_cases,
  results_dir = NULL,
  summary_dir = NULL,
  target_date = NULL,
  region_scale = "Region",
  all_regions = TRUE,
  return_output = FALSE,
  plot = TRUE,
  max_plot = 10,
  ...
)
```

**Arguments**

- **regional_output**
  A list of output as produced by `regional_epinow` and stored in the `regional` list.

- **reported_cases**
  A data frame of confirmed cases (confirm) by date (date), and region (region).
**regional_summary**

- **results_dir**: An optional character string indicating the location of the results directory to extract results from.
- **summary_dir**: A character string giving the directory in which to store summary of results.
- **target_date**: A character string giving the target date for which to extract results (in the format "yyyy-mm-dd"). Defaults to latest available estimates.
- **region_scale**: A character string indicating the name to give the regions being summarised.
- **all_regions**: Logical, defaults to TRUE. Should summary plots for all regions be returned rather than just regions of interest.
- **return_output**: Logical, defaults to FALSE. Should output be returned, this automatically updates to TRUE if no directory for saving is specified.
- **plot**: Logical, defaults to TRUE. Should regional summary plots be produced.
- **max_plot**: Numeric, defaults to 10. A multiplicative upper bound on the number of cases shown on the plot. Based on the maximum number of reported cases.
- **...**: Additional arguments passed to `report_plots`.

**Value**

A list of summary measures and plots

**See Also**

`regional_epinow`

**Examples**

```r
# example delays
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- estimate_delay(rlnorm(100, log(6), 1), max_value = 30)

# example case vector from EpiSoon
cases <- example_confirmed[1:30]
cases <- data.table::rbindlist(list(
  data.table::copy(cases)[, region := "testland"],
  cases[, region := "realland"]
))

# run basic nowcasting pipeline
out <- regional_epinow(
  reported_cases = cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
  output = "region",
  rt = NULL
)
```
```r
report_cases

regional_summary(
  regional_output = out$regional,
  reported_cases = cases
)
```

---

**Description**

[Soft-deprecated] Convolves latent infections to reported cases via an observation model. Likely to be removed/replaced in later releases by functionality drawing on the `stan` implementation.

**Usage**

```r
report_cases(
  case_estimates,
  case_forecast = NULL,
  delays,
  type = "sample",
  reporting_effect,
  CrIs = c(0.2, 0.5, 0.9)
)
```

**Arguments**

- `case_estimates` A data.table of case estimates with the following variables: date, sample, cases
- `case_forecast` A data.table of case forecasts with the following variables: date, sample, cases. If not supplied the default is not to incorporate forecasts.
- `delays` A call to `delay_opts()` defining delay distributions and options. See the documentation of `delay_opts()` and the examples below for details.
- `type` Character string indicating the method to use to transform counts. Supports either "sample" which approximates sampling or "median" would shift by the median of the distribution.
- `reporting_effect` A data.table giving the weekly reporting effect with the following variables: sample (must be the same as in nowcast), effect (numeric scaling factor for each weekday), day (numeric 1 - 7 (1 = Monday and 7 = Sunday)). If not supplied then no weekly reporting effect is assumed.
- `CrIs` Numeric vector of credible intervals to calculate.

**Value**

A list of data.tables. The first entry contains the following variables sample, date and cases with the second being summarised across samples.
Examples

```r
# define example cases
cases <- example_confirmed[1:40]

# set up example delays
generation_time <- get_generation_time(
  disease = "SARS-CoV-2", source = "ganyani"
)
incubation_period <- get_incubation_period(
  disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- dist_spec(
  mean = convert_to_logmean(2, 1), mean_sd = 0.1,
  sd = convert_to_logsd(2, 1), sd_sd = 0.1, max = 10
)

# Instead of running them model we use example
data for speed in this example.
cases <- cases[, cases := as.integer(confirm)]
cases <- cases[, confirm := NULL][, sample := 1]

reported_cases <- report_cases(
  case_estimates = cases,
  delays = delay_opts(incubation_period + reporting_delay),
  type = "sample"
)
print(reported_cases)
```

**Description**

[Questioning] Returns key summary plots for estimates. May be depreciated in later releases as current S3 methods are enhanced.

**Usage**

```r
report_plots(summarised_estimates, reported, target_folder = NULL, ...)
```

**Arguments**

- **summarised_estimates**
  
  A data.table of summarised estimates containing the following variables: variable, median, bottom, and top.
  
  It should also contain the following estimates: R, infections, reported_cases_rt, and r (rate of growth).
report_plots

reported A data.table of reported cases with the following variables: date, confirm.
target_folder Character string specifying where to save results (will create if not present).
...
Additional arguments passed to plot_estimates().

Value

A named list of ggplot2 objects, list(infections, reports, R, growth_rate, summary), which correspond to a summary combination (last item) and for the leading items.

See Also

plot_estimates() of summarised_estimates[variable == "infections"], summarised_estimates[variable == "reported_cases"], summarised_estimates[variable == "R"], and summarised_estimates[variable == "growth_rate"], respectively.

Examples

# define example cases
cases <- example_confirmed[1:40]

# set up example delays
generation_time <- get_generation_time(disease = "SARS-CoV-2", source = "ganyani")
incubation_period <- get_incubation_period(disease = "SARS-CoV-2", source = "lauer")
reporting_delay <- bootstrapped_dist_fit(rlnorm(100, log(6), 1), max_value = 30)

# run model
out <- estimate_infections(cases, stan = stan_opts(samples = 500), generation_time = generation_time_opts(generation_time), delays = delay_opts(incubation_period + reporting_delay), rt = NULL)

# plot infections
plots <- report_plots(summarised_estimates = out$summarised, reported = cases)
plots
**rstan_opts**

---

### report_summary

*Provide Summary Statistics for Estimated Infections and Rt*

#### Description

**[Questioning]** Creates a snapshot summary of estimates. May be removed in later releases as S3 methods are enhanced.

#### Usage

```r
report_summary(
    summarised_estimates, rt_samples, target_folder = NULL, return_numeric = FALSE
)
```

#### Arguments

- **summarised_estimates**: A data.table of summarised estimates containing the following variables: variable, median, bottom, and top. It should contain the following estimates: R, infections, and r (rate of growth).
- **rt_samples**: A data.table containing Rt samples with the following variables: sample and value.
- **target_folder**: Character string specifying where to save results (will create if not present).
- **return_numeric**: Should numeric summary information be returned.

#### Value

A data.table containing formatted and numeric summary measures

---

### rstan_opts

*Rstan Options*

#### Description

**[Stable]** Defines a list specifying the arguments passed to underlying rstan functions via `rstan_sampling_opts()` and `rstan_vb_opts()`. Custom settings can be supplied which override the defaults.

#### Usage

```r
rstan_opts(object = NULL, samples = 2000, method = "sampling", ...)
```
Arguments

- **object**: Stan model object. By default uses the compiled package default.
- **samples**: Numeric, default 2000. Overall number of posterior samples. When using multiple chains iterations per chain is `samples / chains`.
- **method**: A character string, defaulting to sampling. Currently supports `rstan::sampling` ("sampling") or `rstan::vb" ("vb")
- **...**: Additional parameters to pass underlying option functions.

Value

A list of arguments to pass to the appropriate rstan functions.

Author(s)

Sam Abbott

See Also

`rstan_sampling_opts` `rstan_vb_opts`

Examples

```r
rstan_opts(samples = 1000)

# using vb
rstan_opts(method = "vb")
```

Description

[Stable] Defines a list specifying the arguments passed to `rstan::sampling`. Custom settings can be supplied which override the defaults.

Usage

```r
rstan_sampling_opts(
  cores = getOption("mc.cores", 1L),
  warmup = 250,
  samples = 2000,
  chains = 4,
  control = list(),
  save_warmup = FALSE,
  seed = as.integer(runif(1, 1, 1e+08)),
  future = FALSE,
  max_execution_time = Inf,
  ...
)
```
Arguments

- **cores**
  - Number of cores to use when executing the chains in parallel, which defaults to 1 but it is recommended to set the mc.cores option to be as many processors as the hardware and RAM allow (up to the number of chains).

- **warmup**
  - Numeric, defaults to 250. Number of warmup samples per chain.

- **samples**
  - Numeric, default 2000. Overall number of posterior samples. When using multiple chains iterations per chain is samples / chains.

- **chains**
  - Numeric, defaults to 4. Number of MCMC chains to use.

- **control**
  - List, defaults to empty. control parameters to pass to underlying rstan function. By default adapt_delta = 0.95 and max_treedepth = 15 though these settings can be overwritten.

- **save_warmup**
  - Logical, defaults to FALSE. Should warmup progress be saved.

- **seed**
  - Numeric, defaults uniform random number between 1 and 1e8. Seed of sampling process.

- **future**
  - Logical, defaults to FALSE. Should stan chains be run in parallel using future. This allows users to have chains fail gracefully (i.e. when combined with max_execution_time). Should be combined with a call to future::plan.

- **max_execution_time**
  - Numeric, defaults to Inf (seconds). If set will kill off processing of each chain if not finished within the specified timeout. When more than 2 chains finish successfully estimates will still be returned. If less than 2 chains return within the allowed time then estimation will fail with an informative error.

... Additional parameters to pass to rstan::sampling.

Value

A list of arguments to pass to rstan::sampling.

Author(s)

Sam Abbott

Examples

rstan_sampling_opts(samples = 2000)
Usage

rt_opts(samples = 2000, trials = 10, iter = 10000, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>samples</td>
<td>Numeric, default 2000. Overall number of approximate posterior samples.</td>
</tr>
<tr>
<td>trials</td>
<td>Numeric, defaults to 10. Number of attempts to use rstan::vb before failing.</td>
</tr>
<tr>
<td>iter</td>
<td>Numeric, defaulting to 10000. Number of iterations to use in rstan::vb.</td>
</tr>
<tr>
<td>...</td>
<td>Additional parameters to pass to rstan::vb.</td>
</tr>
</tbody>
</table>

Value

A list of arguments to pass to rstan::vb.

Author(s)

Sam Abbott

Examples

rt_opts(samples = 1000)
Arguments

prior  List containing named numeric elements "mean" and "sd". The mean and standard deviation of the log normal Rt prior. Defaults to mean of 1 and standard deviation of 1.

use_rt Logical, defaults to TRUE. Should Rt be used to generate infections and hence reported cases.

rw Numeric step size of the random walk, defaults to 0. To specify a weekly random walk set \( rw = 7 \). For more custom break point settings consider passing in a breakpoints variable as outlined in the next section.

use_breakpoints Logical, defaults to TRUE. Should break points be used if present as a breakpoint variable in the input data. Break points should be defined as 1 if present and otherwise 0. By default breakpoints are fit jointly with a global non-parametric effect and so represent a conservative estimate of break point changes (alter this by setting gp = NULL).

future A character string or integer. This argument indicates how to set future Rt values. Supported options are to project using the Rt model ("project"), to use the latest estimate based on partial data ("latest"), to use the latest estimate based on data that is over 50% complete ("estimate"). If an integer is supplied then the Rt estimate from this many days into the future (or past if negative) past will be used forwards in time.

gp_on Character string, defaulting to "R_t-1". Indicates how the Gaussian process, if in use, should be applied to Rt. Currently supported options are applying the Gaussian process to the last estimated Rt (i.e Rt = Rt-1 * GP), and applying the Gaussian process to a global mean (i.e Rt = R0 * GP). Both should produced comparable results when data is not sparse but the method relying on a global mean will revert to this for real time estimates, which may not be desirable.

pop Integer, defaults to 0. Susceptible population initially present. Used to adjust Rt estimates when otherwise fixed based on the proportion of the population that is susceptible. When set to 0 no population adjustment is done.

Value

A list of settings defining the time-varying reproduction number.

Author(s)

Sam Abbott

Examples

# default settings
rt_opts()

# add a custom length scale
rt_opts(prior = list(mean = 2, sd = 1))
# add a weekly random walk
rt_opts(rw = 7)

---

**run_region**

*Run epinow with Regional Processing Code*

**Description**

[Maturing] Internal function that handles calling `epinow`. Future work will extend this function to better handle `stan` logs and allow the user to modify settings between regions.

**Usage**

```r
run_region(
  target_region,
  generation_time,
  delays,
  truncation,
  rt,
  backcalc,
  gp,
  obs,
  stan,
  horizon,
  CrIs,
  reported_cases,
  target_folder,
  target_date,
  return_output,
  output,
  complete_logger,
  verbose,
  progress_fn,
  ...
)
```

**Arguments**

- `target_region`: Character string indicating the region being evaluated
- `generation_time`: A call to `generation_time_opts()` defining the generation time distribution used. For backwards compatibility a list of summary parameters can also be passed.
- `delays`: A call to `delay_opts()` defining delay distributions and options. See the documentation of `delay_opts()` and the examples below for details.
truncation A call to `trunc_opts()` defining the truncation of observed data. Defaults to `trunc_opts()`. See `estimate_truncation()` for an approach to estimating truncation from data.

rt A list of options as generated by `rt_opts()` defining Rt estimation. Defaults to `rt_opts()`. Set to NULL to switch to using back calculation rather than generating infections using Rt.

backcalc A list of options as generated by `backcalc_opts()` to define the back calculation. Defaults to `backcalc_opts()`.

gp A list of options as generated by `gp_opts()` to define the Gaussian process. Defaults to `gp_opts()`. Set to NULL to disable the Gaussian process.

obs A list of options as generated by `obs_opts()` defining the observation model. Defaults to `obs_opts()`.

stan A list of stan options as generated by `stan_opts()`. Defaults to `stan_opts()`. Can be used to override data, init, and verbose settings if desired.

horizon Numeric, defaults to 7. Number of days into the future to forecast.

CrIs Numeric vector of credible intervals to calculate.

reported_cases A data frame of confirmed cases (confirm) by date (date), and region (region).

target_folder Character string specifying where to save results (will create if not present).

target_date Date, defaults to maximum found in the data if not specified.

return_output Logical, defaults to FALSE. Should output be returned, this automatically updates to TRUE if no directory for saving is specified.

output A character vector of optional output to return. Supported options are the individual regional estimates ("regions"), samples ("samples"), plots ("plots"), copying the individual region dated folder into a latest folder (if `target_folder` is not null, set using "latest"), the stan fit of the underlying model ("fit"), and an overall summary across regions ("summary"). The default is to return samples and plots alongside summarised estimates and summary statistics. If `target_folder` is not NULL then the default is also to copy all results into a latest folder.

complete_logger Character string indicating the logger to output the completion of estimation to.

verbose Logical defaults to FALSE. Outputs verbose progress messages to the console from `epinow`.

progress_fn Function as returned by `progressr::progressor`. Allows the use of a progress bar.

... Pass additional arguments to `epinow`. See the documentation for `epinow` for details.

Value

A list of processed output as produced by `process_region`

See Also

`regional_epinow`
R_to_growth

Convert Reproduction Numbers to Growth Rates

Description

[Questioning] See here # nolint for justification. Now handled internally by stan so may be removed in future updates if no user demand.

Usage

R_to_growth(R, gamma_mean, gamma_sd)

Arguments

R Numeric, Reproduction number estimates

gamma_mean Numeric, mean of the gamma distribution

gamma_sd Numeric, standard deviation of the gamma distribution.

Value

Numeric vector of reproduction number estimates

Examples

R_to_growth(2.18, 4, 1)

sample_approx_dist

Approximate Sampling a Distribution using Counts

Description

[Soft-deprecated] Convolves cases by a PMF function. This function will soon be removed or replaced with a more robust stan implementation.

Usage

sample_approx_dist(
  cases = NULL,
  dist_fn = NULL,
  max_value = 120,
  earliest_allowed_mapped = NULL,
  direction = "backwards",
  type = "sample",
  truncate_future = TRUE
)
Arguments

cases A dataframe of cases (in date order) with the following variables: date and cases.
dist_fn Function that takes two arguments with the first being numeric and the second being logical (and defined as dist). Should return the probability density or a sample from the defined distribution. See the examples for more.
max_value Numeric, maximum value to allow. Defaults to 120 days
earliest_allowed_mapped A character string representing a date ("2020-01-01"). Indicates the earliest allowed mapped value.
direction Character string, defato "backwards". Direction in which to map cases. Supports either "backwards" or "forwards".
type Character string indicating the method to use to transform counts. Supports either "sample" which approximates sampling or "median" would shift by the median of the distribution.
truncate_future Logical, should cases be truncated if they occur after the first date reported in the data. Defaults to TRUE.

Value

A data.table of cases by date of onset

Examples

cases <- example_confirmed
cases <- cases[, cases := as.integer(confirm)]
print(cases)

# total cases
sum(cases$cases)

delay_fn <- function(n, dist, cum) {
  if (dist) {
    pgamma(n + 0.9999, 2, 1) - pgamma(n - 1e-5, 2, 1)
  } else {
    as.integer(rgamma(n, 2, 1))
  }
}

onsets <- sample_approx_dist(
cases = cases,
dist_fn = delay_fn
)

# estimated onset distribution
print(onsets)
# check that sum is equal to reported cases
total_onsets <- median(
  purrr::map_dbl(
    1:10,
    ~ sum(sample_approx_dist(
          cases = cases,
          dist_fn = delay_fn
        )$cases)
  )
)
total_onsets

# map from onset cases to reported
reports <- sample_approx_dist(
  cases = cases,
  dist_fn = delay_fn,
  direction = "forwards"
)

# map from onset cases to reported using a mean shift
reports <- sample_approx_dist(
  cases = cases,
  dist_fn = delay_fn,
  direction = "forwards",
  type = "median"
)

---

**save_estimate_infections**

*Save Estimated Infections*

**Description**

[Stable] Saves output from estimate_infections to a target directory.

**Usage**

```r
save_estimate_infections(
  estimates,
  target_folder = NULL,
  samples = TRUE,
  return_fit = TRUE
)
```
save_input

Arguments

  estimates  List of data frames as output by estimate_infections
  target_folder  Character string specifying where to save results (will create if not present).
  samples  Logical, defaults to TRUE. Should samples be saved
  return_fit  Logical, defaults to TRUE. Should the fit stan object be returned.

Value

  No return value, called for side effects

Author(s)

  Sam Abbott

See Also

  estimate_infections

Description

  [Stable] Saves observed data to a target location if given.

Usage

  save_input(reported_cases, target_folder)

Arguments

  reported_cases  A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.
  target_folder  Character string specifying where to save results (will create if not present).

Value

  No return value, called for side effects

Author(s)

  Sam Abbott
**secondary_opts**

---

## Secondary Reports Options

### Description

**[Stable]** Returns a list of options defining the secondary model used in `estimate_secondary()`. This model is a combination of a convolution of previously observed primary reports combined with current primary reports (either additive or subtractive). It can optionally be cumulative. See the documentation of `type` for sensible options to cover most use cases and the returned values of `secondary_opts()` for all currently supported options.

### Usage

```r
ger`secondary_opts(type = "incidence", ...)
```

### Arguments

- `type` A character string indicating the type of observation the secondary reports are. Options include:
  - "incidence": Assumes that secondary reports equal a convolution of previously observed primary reported cases. An example application is deaths from an infectious disease predicted by reported cases of that disease (or estimated infections).
  - "prevalence": Assumes that secondary reports are cumulative and are defined by currently observed primary reports minus a convolution of secondary reports. An example application is hospital bed usage predicted by hospital admissions.

- `...` Overwrite options defined by `type`. See the returned values for all options that can be passed.

### Value

A list of binary options summarising secondary model used in `estimate_secondary()`. Options returned are cumulative (should the secondary report be cumulative), historic (should a convolution of primary reported cases be used to predict secondary reported cases), `primary_hist_additive` (should the historic convolution of primary reported cases be additive or subtractive), `current` (should currently observed primary reported cases contribute to current secondary reported cases), `primary_current_additive` (should current primary reported cases be additive or subtractive).

### Author(s)

Sam Abbott

### See Also

`estimate_secondary`
**Examples**

```r
# incidence model
secondary_opts("incidence")

# prevalence model
secondary_opts("prevalence")
```

---

**setup_default_logging  Setup Default Logging**

**Description**

[Questioning] Sets up default logging. Usage of logging is currently being explored as the current setup cannot log stan errors or progress.

**Usage**

```r
setup_default_logging(
  logs = tempdir(check = TRUE),
  mirror_epinow = FALSE,
  target_date = NULL
)
```

**Arguments**

- **logs**: Character path indicating the target folder in which to store log information. Defaults to the temporary directory if not specified. Default logging can be disabled if `logs` is set to `NULL`. If specifying a custom logging setup then the code for `setup_default_logging` and the `setup_logging` function are a sensible place to start.

- **mirror_epinow**: Logical, defaults to `FALSE`. Should internal logging be returned from `epinow` to the console.

- **target_date**: Date, defaults to maximum found in the data if not specified.

**Value**

No return value, called for side effects

**Examples**

```r
setup_default_logging()
```
**setup_dt**  
*Convert to Data Table*

**Description**

[Stable] Convenience function that sets the number of data.table cores to 1 and maps input to be a data.table.

**Usage**

```r
setup_dt(reported_cases)
```

**Arguments**

- `reported_cases` A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.

**Value**

A data table

---

**setup_future**  
*Set up Future Backend*

**Description**

[Stable] A utility function that aims to streamline the set up of the required future backend with sensible defaults for most users of regional_epinow. More advanced users are recommended to setup their own future backend based on their available resources.

**Usage**

```r
setup_future(
  reported_cases,
  strategies = c("multisession", "multisession"),
  min_cores_per_worker = 4
)
```

**Arguments**

- `reported_cases` A data frame of confirmed cases (confirm) by date (date), and region (region).
- `strategies` A vector length 1 to 2 of strategies to pass to `future::plan`. Nesting of parallelisation is from the top level down. The default is to set up nesting parallelisation with both using `future::multisession (future::multicore will likely be a faster option on supported platforms). For single level parallelisation use a single strategy or `future::plan` directly. See `?future::plan` for options.
min_cores_per_worker

Numeric, the minimum number of cores per worker. Defaults to 4 which assumes 4 MCMC chains are in use per region.

Value

Numeric number of cores to use per worker. If greater than 1 pass to stan_args = list(cores = "output from setup future") or use future = TRUE. If only a single strategy is used then nothing is returned.

setup_logging

Description

[Questioning] Sets up futile.logger logging, which is integrated into EpiNow2. See the documentation for futile.logger for full details. By default EpiNow2 prints all logs at the "INFO" level and returns them to the console. Usage of logging is currently being explored as the current setup cannot log stan errors or progress.

Usage

setup_logging(
  threshold = "INFO",
  file = NULL,
  mirror_to_console = FALSE,
  name = "EpiNow2"
)

Arguments

threshold Character string indicating the logging level see (?futile.logger for details of the available options). Defaults to "INFO".

file Character string indicating the path to save logs to. By default logs will be written to the console.

mirror_to_console Logical, defaults to FALSE. If saving logs to a file should they also be duplicated in the console.

name Character string defaulting to EpiNow2. This indicates the name of the logger to setup. The default logger for EpiNow2 is called EpiNow2. Nested options include: Epinow2.epinow which controls all logging for epinow and nested functions, EpiNow2.epinow.estimate_infections (logging in estimate_infections), and EpiNow2.epinow.estimate_infections.fit (logging in fitting functions).

Value

Nothing
**setup_target_folder**  
*Setup Target Folder for Saving*

**Description**

**[Stable]** Sets up a folders for saving results

**Usage**

```r
setup_target_folder(target_folder = NULL, target_date)
```

**Arguments**

- **target_folder**  
  Character string specifying where to save results (will create if not present).
- **target_date**  
  Date, defaults to maximum found in the data if not specified.

**Value**

A list containing the path to the dated folder and the latest folder

---

**simulate_infections**  
*Simulate infections using a given trajectory of the time-varying reproduction number*

**Description**

**[Stable]** This function simulates infections using an existing fit to observed cases but with a modified time-varying reproduction number. This can be used to explore forecast models or past counterfactuals. Simulations can be run in parallel using `future::plan`.

**Usage**

```r
simulate_infections(
    estimates,
    R = NULL,
    model = NULL,
    samples = NULL,
    batch_size = 10,
    verbose = interactive()
)
```
simulate_infections

Arguments

estimates The estimates element of an epinow run that has been done with output = "fit", or the result of estimate_infections with return_fit set to TRUE.

R A numeric vector of reproduction numbers; these will overwrite the reproduction numbers contained in estimates, except elements set to NA. Alternatively accepts a data.frame containing at least date and value (integer) variables and optionally sample. More (or fewer) days than in the original fit can be simulated.

model A compiled stan model as returned by rstan::stan_model.

samples Numeric, number of posterior samples to simulate from. The default is to use all samples in the estimates input.

batch_size Numeric, defaults to 10. Size of batches in which to simulate. May decrease run times due to reduced IO costs but this is still being evaluated. If set to NULL then all simulations are done at once.

verbose Logical defaults to interactive(). Should a progress bar (from progressr) be shown.

Value

A list of output as returned by estimate_infections() but based on results from the specified scenario rather than fitting.

Examples

```r
# set number of cores to use
old_opts <- options()
options(mc.cores = ifelse(interactive(), 4, 1))

# get example case counts
reported_cases <- example_confirmed[1:50]

# set up example generation time
generation_time <- get_generation_time(  
disease = "SARS-CoV-2", source = "ganyani"
)

# set delays between infection and case report
incubation_period <- get_incubation_period(  
disease = "SARS-CoV-2", source = "lauer"
)
reporting_delay <- dist_spec(  
  mean = convert_to_logmean(2, 1), mean_sd = 0.1,
  sd = convert_to_logsd(2, 1), sd_sd = 0.1, max = 15
)

# fit model to data to recover Rt estimates
est <- estimate_infections(reported_cases,
  generation_time = generation_time_opts(generation_time),
  delays = delay_opts(incubation_period + reporting_delay),
```
\begin{verbatim}
rt = rt_opts(prior = list(mean = 2, sd = 0.1), rw = 7),
stan = stan_opts(control = list(adapt_delta = 0.9)),
obs = obs_opts(scale = list(mean = 0.1, sd = 0.01)),
gp = NULL, horizon = 0
)

# update Rt trajectory and simulate new infections using it
R <- c(rep(NA_real_, 26), rep(0.5, 10), rep(0.8, 7))
sims <- simulate_infections(est, R)
plot(sims)

# with a data.frame input of samples
R_dt <- data.frame(
  date = seq(
    min(summary(est, type = "parameters", param = "R")$date),
    by = "day", length.out = length(R)
  ),
  value = R
)
sims <- simulate_infections(est, R_dt)
plot(sims)

#' # with a data.frame input of samples
R_samples <- summary(est, type = "samples", param = "R")
R_samples <- R_samples[,.
  (date, sample, value)][sample <= 1000][date <= "2020-04-10"
]
R_samples <- R_samples[date >= "2020-04-01", value := 1.1]
sims <- simulate_infections(est, R_samples)
plot(sims)

options(old_opts)
\end{verbatim}

### simulate_secondary

**Simulate a secondary observation**

#### Description

Simulate a secondary observation

#### Usage

\begin{verbatim}
simulate_secondary(
  data,
  type = "incidence",
  family = "poisson",
  delay_max = 30,
  ...
)
\end{verbatim}
Arguments

data  A data frame containing the date of report and primary cases as a numeric vector.

type  A character string indicating the type of observation the secondary reports are. Options include:

- "incidence": Assumes that secondary reports equal a convolution of previously observed primary reported cases. An example application is deaths from an infectious disease predicted by reported cases of that disease (or estimated infections).
- "prevalence": Assumes that secondary reports are cumulative and are defined by currently observed primary reports minus a convolution of secondary reports. An example application is hospital bed usage predicted by hospital admissions.

cfamily  Character string defining the observation model. Options are Negative binomial ("negbin"), the default, Poisson ("poisson"), and "none" meaning the expectation is returned.

delay_max  Integer, defaulting to 30 days. The maximum delay used in the convolution model.

... Additional parameters to pass to the observation model (i.e. rbinom or rpois).

Value

A data frame containing simulated data in the format required by `estimate_secondary()`.

Author(s)

Sam Abbott
Sebastian Funk

See Also

`estimate_secondary`

Examples

# load data.table for manipulation
library(data.table)

#### Incidence data example ####

# make some example secondary incidence data
cases <- example_confirmed
cases <- as.data.table(cases)[, primary := confirm]

# Assume that only 40 percent of cases are reported
cases[, scaling := 0.4]

# Parameters of the assumed log normal delay distribution
cases[, meanlog := 1.8][, sdlog := 0.5]

# Simulate secondary cases
cases <- simulate_secondary(cases, type = "incidence")
cases

### Prevalence data example ###

# make some example prevalence data
cases <- example_confirmed
cases <- as.data.table(cases)[, primary := confirm]

# Assume that only 30 percent of cases are reported
cases[, scaling := 0.3]

# Parameters of the assumed log normal delay distribution
cases[, meanlog := 1.6][, sdlog := 0.8]

# Simulate secondary cases
cases <- simulate_secondary(cases, type = "prevalence")
cases

---

**stan_opts**

*Stan Options*

**Description**

**[Stable]** Defines a list specifying the arguments passed to underlying stan backend functions via `rstan_sampling_opts()` and `rstan_vb_opts()`. Custom settings can be supplied which override the defaults.

**Usage**

```r
stan_opts(
    samples = 2000,
    backend = "rstan",
    init_fit = NULL,
    return_fit = TRUE,
    ...
)
```

**Arguments**

<table>
<thead>
<tr>
<th>samples</th>
<th>Numeric, default 2000. Overall number of posterior samples. When using multiple chains iterations per chain is samples / chains.</th>
</tr>
</thead>
<tbody>
<tr>
<td>backend</td>
<td>Character string indicating the backend to use for fitting stan models. Currently only &quot;rstan&quot; is supported.</td>
</tr>
</tbody>
</table>
summarise_key_measures

**init_fit**  
[Experimental] Character string or stanfit object, defaults to NULL. Should an initial fit be used to initialise the full fit. An example scenario would be using a national level fit to parametrise regional level fits. Optionally a character string can be passed with the currently supported option being "cumulative". This fits the model to cumulative cases and may be useful for certain data sets where the sampler gets stuck or struggles to initialise. See `init_cumulative_fit()` for details.

This implementation is based on the approach taken in `epidemia` authored by James Scott.

**return_fit**  
Logical, defaults to TRUE. Should the fit stan model be returned.

**...**  
Additional parameters to pass underlying option functions.

**Value**  
A list of arguments to pass to the appropriate rstan functions.

**Author(s)**  
Sam Abbott

**See Also**  
rstan_opts

**Examples**

```r
# using default of rstan::.sampling
stan_opts(samples = 1000)

# using vb
stan_opts(method = "vb")
```

---

### summarise_key_measures

_Summarise rt and cases_

**Description**

[Maturing] Produces summarised data frames of output across regions. Used internally by `regional_summary`.

**Usage**

```r
summarise_key_measures(
    regional_results = NULL,
    results_dir = NULL,
    summary_dir = NULL,
    type = "region",
    date = "latest"
)
```
summarise_results

Arguments

- `regional_results`  
  A list of dataframes as produced by `get_regional_results`
- `results_dir`  
  Character string indicating the directory from which to extract results.
- `summary_dir`  
  Character string the directory into which to save results as a csv.
- `type`  
  Character string, the region identifier to apply (defaults to region).
- `date`  
  A Character string (in the format "yyyy-mm-dd") indicating the date to extract data for. Defaults to "latest" which finds the latest results available.

Value

A list of summarised Rt, cases by date of infection and cases by date of report

See Also

- `regional_summary`

**summarise_results**  
*Summarise Real-time Results*

Description

[Questioning] Used internally by `regional_summary` to produce a summary table of results. May be streamlined in later releases.

Usage

```r
define_summarise_results(  
  regions,  
  summaries = NULL,  
  results_dir = NULL,  
  target_date = NULL,  
  region_scale = "Region"  
)
```

Arguments

- `regions`  
  An character string containing the list of regions to extract results for (must all have results for the same target date).
- `summaries`  
  A list of summary data frames as output by `epinow`
- `results_dir`  
  An optional character string indicating the location of the results directory to extract results from.
- `target_date`  
  A character string indicating the target date to extract results for. All regions must have results for this date.
- `region_scale`  
  A character string indicating the name to give the regions being summarised.
**summary.epinow**  
*Summary output from epinow*

**Description**

[Stable] summary method for class "epinow".

**Usage**

```r
## S3 method for class 'epinow'
summary(object, output = "estimates", date = NULL, params = NULL, ...)
```

**Arguments**

- `object`: A list of output as produced by "epinow".
- `output`: A character string of output to summarise. Defaults to "estimates" but also supports "forecast", and "estimated_reported_cases".
- `date`: A date in the form "yyy-mm-dd" to inspect estimates for.
- `params`: A character vector of parameters to filter for.
- `...`: Pass additional summary arguments to lower level methods

**Value**

Returns a data frame of summary output

**See Also**

summary.estimate_infections epinow

---

**summary.estimate_infections**  
*Summary output from estimate_infections*

**Description**

[Stable] summary method for class "estimate_infections".

**Usage**

```r
## S3 method for class 'estimate_infections'
summary(object, type = "snapshot", date = NULL, params = NULL, ...)
```

**Value**

A list of summary data

**See Also**

summary.estimate_infections epinow
trunc_opts (Truncation Distribution Options)

Description

[Stable] Returns a truncation distribution formatted for usage by downstream functions. See estimate_truncation() for an approach to estimate these distributions.

Usage

trunc_opts(dist = dist_spec())

Arguments

dist A delay distribution or series of delay distributions reflecting the truncation generated using dist_spec() or estimate_truncation(). Default is an empty call to dist_spec(), i.e. no truncation

Value

A list summarising the input truncation distribution.

Author(s)

Sam Abbott
Sebastian Funk
update_horizon

See Also

convert_to_logmean convert_to_logsd bootstrapped_dist_fit dist_spec

Examples

# no truncation
trunc_opts()

# truncation dist
trunc_opts(dist = dist_spec(mean = 3, sd = 2, max = 10))

update_horizon 

Updates Forecast Horizon Based on Input Data and Target

Description

[Stable] Makes sure that a forecast is returned for the user specified time period beyond the target date.

Usage

update_horizon(horizon, target_date, reported_cases)

Arguments

horizon Numeric, defaults to 7. Number of days into the future to forecast.

target_date Date, defaults to maximum found in the data if not specified.

reported_cases A data frame of confirmed cases (confirm) by date (date). confirm must be integer and date must be in date format.

Value

Numeric forecast horizon adjusted for the users intention

Author(s)

Sam Abbott
update_list  Update a List

**Description**

[Stable] Used to handle updating settings in a list. For example when making changes to opts_list output.

**Usage**

```r
update_list(defaults = list(), optional = list())
```

**Arguments**

- **defaults**  A list of default settings
- **optional**  A list of optional settings to override defaults

**Value**

A list

---

update_secondary_args  Update estimate_secondary default priors

**Description**

[Stable] This functions allows the user to more easily specify data driven or model based priors for estimate_secondary() from example from previous model fits using a data.frame to overwrite other default settings. Note that default settings are still required.

**Usage**

```r
update_secondary_args(data, priors, verbose = TRUE)
```

**Arguments**

- **data**  A list of data and arguments as returned by create_stan_data().
- **priors**  A data.frame of named priors to be used in model fitting rather than the defaults supplied from other arguments. This is typically useful if wanting to inform a estimate from the posterior of another model fit. Priors that are currently use to update the defaults are the scaling fraction ("frac_obs"), the mean delay ("delay_mean"), and standard deviation of the delay ("delay_sd"). The data.frame should have the following variables: variable, mean, and sd.
- **verbose**  Logical, defaults to FALSE. Should verbose progress messages be returned.
Value

A list as produced by create_stan_data().

Author(s)

Sam Abbott

Examples

```r
priors <- data.frame(variable = "frac_obs", mean = 3, sd = 1)
data <- list(obs_scale_mean = 4, obs_scale_sd = 3)
update_secondary_args(data, priors)
```
Index

* datasets
  example_confirmed, 51
  generation_times, 61
  incubation_periods, 71
  +.dist_spec, 5

  add_day_of_week, 6
  adjust_infection_to_report, 6
  allocate_delays, 8
  allocate_empty, 9

  backcalc_opts, 10
  bootstrapped_dist_fit, 11

  c.dist_spec, 12
  calc_CrI, 12
  calc_CrIs, 13
  calc_summary_measures, 14
  calc_summary_stats, 14
  clean_nowcasts, 15
  clean_regions, 16
  construct_output, 16
  convert_to_logmean, 17
  convert_to_logsd, 18
  copy_results_to_latest, 18
  create_backcalc_data, 19
  create_clean_reported_cases, 20
  create_future_rt, 21
  create_gp_data, 21
  create_initial_conditions, 22
  create_obs_model, 23
  create_rt_data, 24
  create_shifted_cases, 25
  create_stan_args, 26
  create_stan_data, 27
  create_stan_delays, 28

  delay_opts, 28
  delay_opts(), 33, 46
  dist_fit, 29

  dist_skel, 31
  dist_spec, 33, 62, 79
  dist_spec(), 5, 12, 28, 35, 62, 76, 86, 121
  dist_spec_plus, 34

  epinow, 35
  epinow(), 12
  estimate_delay, 39
  estimate_infections, 40
  estimate_infections(), 12, 114
  estimate_secondary, 45
  estimate_secondary(), 116
  estimate_truncation, 48
  estimate_truncation(), 121
  estimates_by_report_date, 38
  example_confirmed, 51
  expose_stan_fns, 51
  extract_CrIs, 52
  extract_inits, 52
  extract_parameter, 53
  extract_parameter_samples, 54
  extract_stan_param, 55
  extract_static_parameter, 55

  filter_opts, 56
  fit_model_with_nuts, 57
  fit_model_with_vb, 57
  forecast_secondary, 58
  format_fit, 59

  gamma_dist_def, 60
  generation_time_opts, 62
  generation_time_opts(), 33
  generation_times, 61
  get_dist, 63
  get_generation_time, 62, 64
  get_generation_time(), 62
  get_incubation_period, 65
  get_raw_result, 65
  get_regional_results, 66
get_regions, 67
get_regions_with_most_reports, 68
get_seeding_time, 69
ggplot2, 79
gp_opts, 69
growth_to_R, 71

incubation_periods, 71
init_cumulative_fit, 72

lognorm_dist_def, 73

make_conf, 74
map_prob_change, 74
match_output_arguments, 75
mean.dist_spec, 76

obs_opts, 77
opts_list, 78

plot (plot.estimate_infections), 81
plot.dist_spec, 79
plot.epinow, 80
plot.estimate_infections, 81
plot.estimate_secondary, 81
plot.estimate_truncation, 82
plot_CrIs, 83
plot.estimates, 83
plot.estimates(), 97
plot_summary, 85
print.dist_spec, 86
process_region, 87
process_regions, 88

R_to_growth, 105
regional_epinow, 88
regional_runtimes, 92
regional_summary, 93
report_cases, 95
report_plots, 96
report_summary, 98
rstan_opts, 98
rstan_sampling_opts, 99
rstan_vb_opts, 100
rt_opts, 101
run_region, 103

sample_approx_dist, 105
save_estimate_infections, 107
save_input, 108

secondary_opts, 109
setup_default_logging, 110
setup_dt, 111
setup_future, 111
setup_logging, 112
setup_target_folder, 113
simulate_infections, 113
simulate_secondary, 115
stan_opts, 117
summarise_key_measures, 118
summarise_results, 119
summary (summary.epinow), 120
summary.epinow, 120
summary.estimate_infections, 120

trunc_opts, 121
trunc_opts(), 33

update_horizon, 122
update_list, 123
update_secondary_args, 123