

# Package ‘slendr’

December 7, 2023

**Title** A Simulation Framework for Spatiotemporal Population Genetics

**Version** 0.8.0

**Description** A framework for simulating spatially explicit genomic data which leverages real cartographic information for programmatic and visual encoding of spatiotemporal population dynamics on real geographic landscapes. Population genetic models are then automatically executed by the 'SLiM' software by Haller et al. (2019) <[doi:10.1093/molbev/msy228](https://doi.org/10.1093/molbev/msy228)> behind the scenes, using a custom built-in simulation 'SLiM' script. Additionally, fully abstract spatial models not tied to a specific geographic location are supported, and users can also simulate data from standard, non-spatial, random-mating models. These can be simulated either with the 'SLiM' built-in back-end script, or using an efficient coalescent population genetics simulator 'msprime' by Baumdicker et al. (2022) <[doi:10.1093/genetics/iyab229](https://doi.org/10.1093/genetics/iyab229)> with a custom-built 'Python' script bundled with the R package. Simulated genomic data is saved in a tree-sequence format and can be loaded, manipulated, and summarised using tree-sequence functionality via an R interface to the 'Python' module 'tskit' by Kelleher et al. (2019) <[doi:10.1038/s41588-019-0483-y](https://doi.org/10.1038/s41588-019-0483-y)>. Complete model configuration, simulation and analysis pipelines can be therefore constructed without a need to leave the R environment, eliminating friction between disparate tools for population genetic simulations and data analysis.

**Depends** R (>= 3.6.0)

**License** MIT + file LICENSE

**Encoding** UTF-8

**RoxygenNote** 7.2.3

**SystemRequirements** 'SLiM' is a forward simulation software for population genetics and evolutionary biology. See <<https://messengerlab.org/slim/>> for installation instructions and further information. The 'Python' coalescent framework 'msprime' and the 'tskit' module can be installed by following the instructions at <<https://tskit.dev/>>.

**Imports** ggplot2, dplyr, purrr, readr, magrittr, reticulate, tidyr, png, ijttiff, ape, shinyWidgets, shiny, scales

**Suggests** testthat (>= 3.0.0), sf, stars, rnaturalearth, gganimate, knitr, rmarkdown, admixr, units, magick, cowplot, forcats, rsvg

**VignetteBuilder** knitr

**URL** <https://github.com/bodkan/slendr>

**BugReports** <https://github.com/bodkan/slendr/issues>

**Config/testthat/edition** 3

**NeedsCompilation** no

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---

animate\_model

*Animate the simulated population dynamics*


---

## Description

Animate the simulated population dynamics

## Usage

```
animate_model(model, file, steps, gif = NULL, width = 800, height = 560)
```

**Arguments**

model	Compiled slendr_model model object
file	Path to the table of saved individual locations
steps	How many frames should the animation have?
gif	Path to an output GIF file (animation object returned by default)
width, height	Dimensions of the animation in pixels

**Value**

If gif = NULL, return ganimate animation object. Otherwise a GIF file is saved and no value is returned.

---

area	<i>Calculate the area covered by the given slendr object</i>
------	--

---

**Description**

Calculate the area covered by the given slendr object

**Usage**

```
area(x)
```

**Arguments**

x	Object of the class slendr
---	----------------------------

**Value**

Area covered by the input object. If a slendr\_pop was given, a table with an population range area in each time point will be returned. If a slendr\_region or slendr\_world object was specified, the total area covered by this object's spatial boundary will be returned.

**Examples**

```
region_a <- region("A", center = c(20, 50), radius = 20)
region_b <- region("B", polygon = list(c(50, 40), c(70, 40), c(70, 60), c(50, 60)))
plot_map(region_a, region_b)

# note that area won't be *exactly* equal to pi*r^2:
# https://stackoverflow.com/a/65280376
area(region_a)

area(region_b)
```

---

check_dependencies	<i>Check that the required dependencies are available for slendr to work</i>
--------------------	--

---

**Description**

Check that the required dependencies are available for slendr to work

**Usage**

```
check_dependencies(python = FALSE, slim = FALSE, quit = FALSE)
```

**Arguments**

python	Is the slendr Python environment required?
slim	Is SLiM required?
quit	Should the R interpreter quit if required slendr dependencies are missing? This option (which is not turned on by default, being set to FALSE) is used mainly in avoiding running slendr man page examples on machines which lack dependencies. If set to TRUE, a logical value is returned.

**Value**

If `quit = TRUE`, no values is returned, if `quit = FALSE`, a scalar logical value is returned indicating whether or not the dependencies are present.

---

check_env	<i>Check that the active Python environment is setup for slendr</i>
-----------	---

---

**Description**

This function inspects the Python environment which has been activated by the reticulate package and prints the versions of all slendr Python dependencies to the console.

**Usage**

```
check_env(verbose = TRUE)
```

**Arguments**

verbose	Should a log message be printed? If FALSE, only a logical value is returned (invisibly).
---------	--

**Value**

Either TRUE (slendr Python environment is present) or FALSE (slendr Python environment is not present).

**Examples**

```
init_env()
check_env()
```

---

clear_env	<i>Remove the automatically created slendr Python environment</i>
-----------	---

---

**Description**

Remove the automatically created slendr Python environment

**Usage**

```
clear_env(force = FALSE)
```

**Arguments**

force            Ask before deleting the environment?

**Value**

No return value, called for side effects

---

compile_model	<i>Compile a slendr demographic model</i>
---------------	---

---

**Description**

First, compiles the vectorized population spatial maps into a series of binary raster PNG files, which is the format that SLiM understands and uses it to define population boundaries. Then extracts the demographic model defined by the user (i.e. population divergences and gene flow events) into a series of tables which are later used by the built-in SLiM script to program the timing of simulation events.

**Usage**

```
compile_model(
  populations,
  generation_time,
  path = NULL,
  resolution = NULL,
  competition = NULL,
  mating = NULL,
  dispersal = NULL,
  gene_flow = list(),
```

```

    overwrite = FALSE,
    force = FALSE,
    simulation_length = NULL,
    direction = NULL,
    serialize = TRUE,
    slim_script = NULL,
    description = "",
    sim_length = NULL
)

```

### Arguments

populations	Object(s) of the <code>slendr_pop</code> class (multiple objects need to be specified in a list)
generation_time	Generation time (in model time units)
path	Output directory for the model configuration files which will be loaded by the backend SLiM script. If <code>NULL</code> , model configuration files will be saved to a temporary directory.
resolution	How many distance units per pixel?
competition, mating	Maximum spatial competition and mating choice distance
dispersal	Standard deviation of the normal distribution of the parent-offspring distance
gene_flow	Gene flow events generated by the <code>gene_flow</code> function (either a list of <code>data.frame</code> objects in the format defined by the <code>gene_flow</code> function, or a single <code>data.frame</code> )
overwrite	Completely delete the specified directory, in case it already exists, and create a new one?
force	Force a deletion of the model directory if it is already present? Useful for non-interactive uses. In an interactive mode, the user is asked to confirm the deletion manually.
simulation_length	Total length of the simulation (required for forward time models, optional for models specified in backward time units which by default run to "the present time")
direction	Intended direction of time. Under normal circumstances this parameter is inferred from the model and does not need to be set manually.
serialize	Should model files be serialized to disk? If not, only an R model object will be returned but no files will be created. This speeds up simulation with <code>msprime</code> but prevents using the SLiM back end.
slim_script	Path to a SLiM script to be used for executing the model (by default, a bundled backend script will be used). If <code>NULL</code> , the SLiM script bundled with <code>slendr</code> will be used.
description	Optional short description of the model
sim_length	Deprecated. Use <code>simulation_length</code> instead.

**Value**

Compiled `slendr_model` model object which encapsulates all information about the specified model (which populations are involved, when and how much gene flow should occur, what is the spatial resolution of a map, and what spatial dispersal and mating parameters should be used in a SLiM simulation, if applicable)

**Examples**

```
# spatial definitions -----

# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50)
  )
)

# population ranges can expand by a given distance in all directions
```



```

pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

distance

*Calculate the distance between a pair of spatial boundaries*


---

### Description

Calculate the distance between a pair of spatial boundaries

### Usage

```
distance(x, y, measure, time = NULL)
```

### Arguments

x, y	Objects of the class <code>slendr</code>
measure	How to measure distance? This can be either 'border' (distance between the borders of x and y) or 'center' (distance between their centroids).
time	Time closest to the spatial maps of x and y if they represent <code>slendr_pop</code> population boundaries (ignored for general <code>slendr_region</code> objects)

### Value

If the coordinate reference system was specified, a distance in projected units (i.e. meters) is returned. Otherwise the function returns a normal Euclidean distance.

**Examples**

```
# create two regions on a blank abstract landscape
region_a <- region("A", center = c(20, 50), radius = 20)
region_b <- region("B", center = c(80, 50), radius = 20)
plot_map(region_a, region_b)

# compute the distance between the centers of both population ranges
distance(region_a, region_b, measure = "center")

# compute the distance between the borders of both population ranges
distance(region_a, region_b, measure = "border")
```

---

expand\_range

*Expand the population range*


---

**Description**

Expands the spatial population range by a specified distance in a given time-window

**Usage**

```
expand_range(
  pop,
  by,
  end,
  start,
  overlap = 0.8,
  snapshots = NULL,
  polygon = NULL,
  lock = FALSE,
  verbose = TRUE
)
```

**Arguments**

pop	Object of the class <code>slendr_pop</code>
by	How many units of distance to expand by?
start, end	When does the expansion start/end?
overlap	Minimum overlap between subsequent spatial boundaries
snapshots	The number of intermediate snapshots (overrides the <code>overlap</code> parameter)
polygon	Geographic region to restrict the expansion to
lock	Maintain the same density of individuals. If <code>FALSE</code> (the default), the number of individuals in the population will not change. If <code>TRUE</code> , the number of individuals simulated will be changed (increased or decreased) appropriately, to match the new population range area.
verbose	Report on the progress of generating intermediate spatial boundaries?

## Details

Note that because slendr models have to accommodate both SLiM and msprime back ends, population sizes and times of events are rounded to the nearest integer value.

## Value

Object of the class `slendr_pop`, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

## Examples

```
# spatial definitions -----

# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
```

```

      c(600, 100), c(500, 50))
    )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

explore\_model

*Open an interactive browser of the spatial model*

---

## Description

Open an interactive browser of the spatial model

## Usage

```
explore_model(model)
```

## Arguments

model                    Compiled slendr\_model model object

## Value

No return value, called in order to start an interactive browser-based interface to explore the dynamics of a slendr model

---

gene_flow	<i>Define a gene-flow event between two populations</i>
-----------	---

---

**Description**

Define a gene-flow event between two populations

**Usage**

```
gene_flow(from, to, rate, start, end, overlap = TRUE)
```

**Arguments**

from, to	Objects of the class <code>slendr_pop</code>
rate	Scalar value in the range (0, 1] specifying the proportion of migration over given time period
start, end	Start and end of the gene-flow event
overlap	Require spatial overlap between admixing populations? (default TRUE)

**Value**

Object of the class `data.frame` containing parameters of the specified gene-flow event.

**Examples**

```
# spatial definitions -----

# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)
```

```

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                 map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

**Description**

This function attempts to activate a dedicated slendr Miniconda Python environment previously set up via `setup_env`.

**Usage**

```
init_env(quiet = FALSE)
```

**Arguments**

`quiet` Should informative messages be printed to the console? Default is FALSE.

**Value**

No return value, called for side effects

---

<code>join</code>	<i>Merge two spatial slendr objects into one</i>
-------------------	--

---

**Description**

Merge two spatial slendr objects into one

**Usage**

```
join(x, y, name = NULL)
```

**Arguments**

`x` Object of the class `slendr`

`y` Object of the class `slendr`

`name` Optional name of the resulting geographic region. If missing, name will be constructed from the function arguments.

**Value**

Object of the class `slendr_region` which encodes a standard spatial object of the class `sf` with several additional attributes (most importantly a corresponding `slendr_map` object, if applicable).

**Examples**

```
# create a blank abstract world 1000x1000 distance units in size
blank_map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# it is possible to construct custom landscapes (islands, corridors, etc.)
island1 <- region("island1", polygon = list(c(10, 30), c(50, 30), c(40, 50), c(0, 40)))
island2 <- region("island2", polygon = list(c(60, 60), c(80, 40), c(100, 60), c(80, 80)))
island3 <- region("island3", center = c(20, 80), radius = 10)
archipelago <- island1 %>% join(island2) %>% join(island3)

custom_map <- world(xrange = c(1, 100), c(1, 100), landscape = archipelago)

# real Earth landscapes can be defined using freely-available Natural Earth
# project data and with the possibility to specify an appropriate Coordinate
# Reference System, such as this example of a map of Europe

real_map <- world(xrange = c(-15, 40), yrange = c(30, 60), crs = "EPSG:3035")
```

---

 move

---

*Move the population to a new location in a given amount of time*


---

**Description**

This function defines a displacement of a population along a given trajectory in a given time frame

**Usage**

```
move(
  pop,
  trajectory,
  end,
  start,
  overlap = 0.8,
  snapshots = NULL,
  verbose = TRUE
)
```

**Arguments**

pop	Object of the class <code>slendr_pop</code>
trajectory	List of two-dimensional vectors (longitude, latitude) specifying the migration trajectory
start, end	Start/end points of the population migration
overlap	Minimum overlap between subsequent spatial boundaries
snapshots	The number of intermediate snapshots (overrides the <code>overlap</code> parameter)
verbose	Show the progress of searching through the number of sufficient snapshots?



**Value**

Object of the class `slendr_pop`, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

**Examples**

```
# spatial definitions -----

# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)
```

```

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

msprime

*Run a slendr model in msprime*


---

## Description

This function will execute a built-in msprime script and run a compiled slendr demographic model.

## Usage

```

msprime(
  model,
  sequence_length,
  recombination_rate,
  samples = NULL,
  output = NULL,
  random_seed = NULL,
  load = TRUE,
  verbose = FALSE,
  debug = FALSE,
  run = TRUE
)

```

## Arguments

model            Model object created by the compile function

sequence_length	Total length of the simulated sequence (in base-pairs)
recombination_rate	Recombination rate of the simulated sequence (in recombinations per basepair per generation)
samples	A data frame of times at which a given number of individuals should be remembered in the tree-sequence (see <code>schedule_sampling</code> for a function that can generate the sampling schedule in the correct format). If missing, only individuals present at the end of the simulation will be recorded in the tree-sequence output file.
output	Path to the output tree-sequence file. If NULL (the default), tree sequence will be saved to a temporary file.
random_seed	Random seed (if missing, SLiM's own seed will be used)
load	Should the final tree sequence be immediately loaded and returned? Default is TRUE. The alternative (FALSE) is useful when a tree-sequence file is written to a custom location to be loaded at a later point.
verbose	Write the output log to the console (default FALSE)?
debug	Write msprime's debug log to the console (default FALSE)?
run	Should the msprime engine be run? If FALSE, the command line msprime command will be printed (and returned invisibly as a character vector) but not executed.

## Value

A tree-sequence object loaded via Python-R reticulate interface function `ts_load` (internally represented by the Python object `tskit.trees.TreeSequence`). Optionally, depending on the value of the arguments `load =` or `run =`, nothing or a character vector, respectively.

## Examples

```
init_env()

# load an example model
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# afr and eur objects would normally be created before slendr model compilation,
# but here we take them out of the model object already compiled for this
# example (in a standard slendr simulation pipeline, this wouldn't be necessary)
afr <- model$populations[["AFR"]]
eur <- model$populations[["EUR"]]
chimp <- model$populations[["CH"]]

# schedule the sampling of a couple of ancient and present-day individuals
# given model at 20 ky, 10 ky, 5ky ago and at present-day (time 0)
modern_samples <- schedule_sampling(model, times = 0, list(afr, 10), list(eur, 100), list(chimp, 1))
ancient_samples <- schedule_sampling(model, times = c(40000, 30000, 20000, 10000), list(eur, 1))
```

```

# sampling schedules are just data frames and can be merged easily
samples <- rbind(modern_samples, ancient_samples)

# run a simulation using the msprime back end from a compiled slendr model object
ts <- msprime(model, sequence_length = 1e5, recombination_rate = 0, samples = samples)

# automatic loading of a simulated output can be prevented by `load = FALSE`, which can be
# useful when a custom path to a tree-sequence output is given for later downstream analyses
output_file <- tempfile(fileext = ".trees")
msprime(model, sequence_length = 1e5, recombination_rate = 0, samples = samples,
        output = output_file, load = FALSE, random_seed = 42)
# ... at a later stage:
ts <- ts_load(output_file, model)

summary(ts)

```

---

 overlap

*Generate the overlap of two slendr objects*


---

### Description

Generate the overlap of two slendr objects

### Usage

```
overlap(x, y, name = NULL)
```

### Arguments

x	Object of the class slendr
y	Object of the class slendr
name	Optional name of the resulting geographic region. If missing, name will be constructed from the function arguments.

### Value

Object of the class slendr\_region which encodes a standard spatial object of the class sf with several additional attributes (most importantly a corresponding slendr\_map object, if applicable).

---

plot\_map

*Plot slendr geographic features on a map*


---

### Description

Plots objects of the three slendr spatial classes (slendr\_map, slendr\_region, and slendr\_pop).

### Usage

```
plot_map(
  ...,
  time = NULL,
  gene_flow = FALSE,
  graticules = "original",
  intersect = TRUE,
  show_map = TRUE,
  title = NULL,
  interpolated_maps = NULL
)
```

### Arguments

...	Objects of classes slendr_map, slendr_region, or slendr_pop
time	Plot a concrete time point
gene_flow	Indicate gene flow events with an arrow
graticules	Plot graticules in the original Coordinate Reference System (such as longitude-latitude), or in the internal CRS (such as meters)?
intersect	Intersect the population boundaries against landscape and other geographic boundaries (default TRUE)?
show_map	Show the underlying world map
title	Title of the plot
interpolated_maps	Interpolated spatial boundaries for all populations in all time points (this is only used for plotting using the explore shiny app)

### Value

A ggplot2 object with the visualized slendr map

---

plot\_model

*Plot demographic history encoded in a slendr model*


---

**Description**

Plot demographic history encoded in a slendr model

**Usage**

```
plot_model(
  model,
  sizes = TRUE,
  proportions = FALSE,
  gene_flow = TRUE,
  log = FALSE,
  order = NULL,
  file = NULL,
  ...
)
```

**Arguments**

model	Compiled slendr_model model object
sizes	Should population size changes be visualized?
proportions	Should gene flow proportions be visualized (FALSE by default to prevent cluttering and overplotting)
gene_flow	Should gene-flow arrows be visualized (default TRUE).
log	Should the y-axis be plotted on a log scale? Useful for models over very long time-scales.
order	Order of the populations along the x-axis, given as a character vector of population names. If NULL (the default), the default plotting algorithm will be used, ordering populations from the most ancestral to the most recent using an in-order tree traversal.
file	Output file for a figure saved via ggsave
...	Optional argument which will be passed to ggsave

**Value**

A ggplot2 object with the visualized slendr model

**Examples**

```
init_env()
```

```
# load an example model with an already simulated tree sequence
path <- system.file("extdata/models/introgression", package = "slendr")
model <- read_model(path)

plot_model(model, sizes = FALSE, log = TRUE)
```

---

population                      *Define a population*

---

## Description

Defines the parameters of a population (non-spatial and spatial).

## Usage

```
population(
  name,
  time,
  N,
  parent = NULL,
  map = FALSE,
  center = NULL,
  radius = NULL,
  polygon = NULL,
  remove = NULL,
  intersect = TRUE,
  competition = NA,
  mating = NA,
  dispersal = NA,
  dispersal_fun = NULL,
  aquatic = FALSE
)
```

## Arguments

name	Name of the population
time	Time of the population's first appearance
N	Number of individuals at the time of first appearance
parent	Parent population object or NULL (which indicates that the population does not have an ancestor, as it is the first population in its "lineage")
map	Object of the type <code>slendr_map</code> which defines the world context (created using the <code>world</code> function). If the value <code>FALSE</code> is provided, a non-spatial model will be run.
center	Two-dimensional vector specifying the center of the circular range
radius	Radius of the circular range

polygon	List of vector pairs, defining corners of the polygon range or a geographic region of the class <code>slendr_region</code> from which the polygon coordinates will be extracted (see the <code>region()</code> function)
remove	Time at which the population should be removed
intersect	Intersect the population's boundaries with landscape features?
competition, mating	Maximum spatial competition and mating choice distance
dispersal	Standard deviation of the normal distribution of the distance that offspring disperses from its parent
dispersal_fun	Distribution function governing the dispersal of offspring. One of "normal", "uniform", "cauchy", "exponential", or "brownian" (in which vertical and horizontal displacements are drawn from a normal distribution independently).
aquatic	Is the species aquatic (FALSE by default, i.e. terrestrial species)?

### Details

There are four ways to specify a spatial boundary: i) circular range specified using a center coordinate and a radius, ii) polygon specified as a list of two-dimensional vector coordinates, iii) polygon as in ii), but defined (and named) using the `region` function, iv) with just a world map specified (circular or polygon range parameters set to the default NULL value), the population will be allowed to occupy the entire landscape.

Note that because `slendr` models have to accommodate both SLiM and `msprime` back ends, population sizes and split times are rounded to the nearest integer value.

### Value

Object of the class `slendr_pop`, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

### Examples

```
# spatial definitions -----
# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
```



```

pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
  polygon = list(c(100, 100), c(320, 30), c(500, 200),
    c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
    trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

```
print.slendr_pop
```

*Print a short summary of a slendr object*

---

### Description

All spatial objects in the slendr package are internally represented as Simple Features (sf) objects. This fact is hidden in most circumstances this, as the goal of the slendr package is to provide functionality at a much higher level (population boundaries, geographic regions, instead of individual polygons and other "low-level" geometric objects), without the users having to worry about low-level details involved in handling spatial geometries. However, the full sf object representation can be always printed by calling `x[]`.

### Usage

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

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

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

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

### Arguments

<code>x</code>	Object of a class slendr (either <code>slendr_pop</code> , <code>slendr_map</code> , <code>slendr_region</code> , or <code>slendr_table</code> )
<code>...</code>	Additional arguments passed to <code>print</code>

### Value

No return value, used only for printing

---

```
print.slendr_ts
```

*Print tskit's summary table of the Python tree-sequence object*

---

### Description

Print tskit's summary table of the Python tree-sequence object

**Usage**

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

**Arguments**

x                    Tree object of the class `slendr_phylo`  
 ...                  Additional arguments normally passed to `print` (not used in this case)

**Value**

No return value, simply prints the tskit summary table to the terminal

---

read_model	<i>Read a previously serialized model configuration</i>
------------	---

---

**Description**

Reads all configuration tables and other model data from a location where it was previously compiled to by the `compile` function.

**Usage**

```
read_model(path)
```

**Arguments**

path                  Directory with all required configuration files

**Value**

Compiled `slendr_model` model object which encapsulates all information about the specified model (which populations are involved, when and how much gene flow should occur, what is the spatial resolution of a map, and what spatial dispersal and mating parameters should be used in a SLiM simulation, if applicable)

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
path <- system.file("extdata/models/introgression", package = "slendr")
model <- read_model(path)

plot_model(model, sizes = FALSE, log = TRUE)
```

---

region	<i>Define a geographic region</i>
--------	-----------------------------------

---

### Description

Creates a geographic region (a polygon) on a given map and gives it a name. This can be used to define objects which can be reused in multiple places in a slendr script (such as region arguments of population) without having to repeatedly define polygon coordinates.

### Usage

```
region(name = NULL, map = NULL, center = NULL, radius = NULL, polygon = NULL)
```

### Arguments

name	Name of the geographic region
map	Object of the type <code>sf</code> which defines the map
center	Two-dimensional vector specifying the center of the circular range
radius	Radius of the circular range
polygon	List of vector pairs, defining corners of the polygon range or a geographic region of the class <code>slendr_region</code> from which the polygon coordinates will be extracted (see the <code>region()</code> function)

### Value

Object of the class `slendr_region` which encodes a standard spatial object of the class `sf` with several additional attributes (most importantly a corresponding `slendr_map` object, if applicable).

### Examples

```
# create a blank abstract world 1000x1000 distance units in size
blank_map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# it is possible to construct custom landscapes (islands, corridors, etc.)
island1 <- region("island1", polygon = list(c(10, 30), c(50, 30), c(40, 50), c(0, 40)))
island2 <- region("island2", polygon = list(c(60, 60), c(80, 40), c(100, 60), c(80, 80)))
island3 <- region("island3", center = c(20, 80), radius = 10)
archipelago <- island1 %>% join(island2) %>% join(island3)

custom_map <- world(xrange = c(1, 100), c(1, 100), landscape = archipelago)

# real Earth landscapes can be defined using freely-available Natural Earth
# project data and with the possibility to specify an appropriate Coordinate
# Reference System, such as this example of a map of Europe

real_map <- world(xrange = c(-15, 40), yrange = c(30, 60), crs = "EPSG:3035")
```

---

reproject	<i>Reproject coordinates between coordinate systems</i>
-----------	---

---

### Description

Converts between coordinates on a compiled raster map (i.e. pixel units) and different Geographic Coordinate Systems (CRS).

### Usage

```
reproject(
  from,
  to,
  x = NULL,
  y = NULL,
  coords = NULL,
  model = NULL,
  add = FALSE,
  input_prefix = "",
  output_prefix = "new"
)
```

### Arguments

from, to	Either a CRS code accepted by GDAL, a valid integer EPSG value, an object of class <code>crs</code> , the value "raster" (converting from/to pixel coordinates), or "world" (converting from/to whatever CRS is set for the underlying map)
x, y	Coordinates in two dimensions (if missing, coordinates are expected to be in the <code>data.frame</code> specified in the <code>coords</code> parameter as columns "x" and "y")
coords	<code>data.frame</code> -like object with coordinates in columns "x" and "y"
model	Object of the class <code>slendr_model</code>
add	Add column coordinates to the input <code>data.frame</code> <code>coords</code> (coordinates otherwise returned as a separate object)?
input_prefix, output_prefix	Input and output prefixes of data frame columns with spatial coordinates

### Value

`Data.frame` with converted two-dimensional coordinates given as input

### Examples

```
lon_lat_df <- data.frame(x = c(30, 0, 15), y = c(60, 40, 10))

reproject(
  from = "epsg:4326",
```

```

to = "epsg:3035",
coords = lon_lat_df,
add = TRUE # add converted [lon,lat] coordinates as a new column
)

```

---

resize *Change the population size*

---

### Description

Resizes the population starting from the current value of N individuals to the specified value

### Usage

```
resize(pop, N, how, time, end = NULL)
```

### Arguments

pop	Object of the class <code>slendr_pop</code>
N	Population size after the change
how	How to change the population size (options are "step" or "exponential")
time	Time of the population size change
end	End of the population size change period (used for exponential change events)

### Details

In the case of exponential size change, if the final N is larger than the current size, the population will be exponentially growing over the specified time period until it reaches N individuals. If N is smaller, the population will shrink exponentially.

Note that because `slendr` models have to accomodate both `SLiM` and `msprime` back ends, population sizes and split times are rounded to the nearest integer value.

### Value

Object of the class `slendr_pop`, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

### Examples

```

# spatial definitions -----
# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1

```

```

# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

```

```
# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)
```

---

schedule\_sampling      *Define sampling events for a given set of populations*

---

### Description

Schedule sampling events at specified times and, optionally, a given set of locations on a landscape

### Usage

```
schedule_sampling(model, times, ..., locations = NULL, strict = FALSE)
```

### Arguments

model	Object of the class <code>slendr_model</code>
times	Integer vector of times (in model time units) at which to schedule remembering of individuals in the tree-sequence
...	Lists of two elements ( <code>slendr_pop</code> population object- <code>&lt;number of individuals to sample&gt;</code> ), representing from which populations should how many individuals be remembered at times given by <code>times</code>
locations	List of vector pairs, defining two-dimensional coordinates of locations at which the closest number of individuals from given populations should be sampled. If <code>NULL</code> (the default), individuals will be sampled randomly throughout their spatial boundary.
strict	Should any occurrence of a population not being present at a given time result in an error? Default is <code>FALSE</code> , meaning that invalid sampling times for any populations will be quietly ignored.

### Details

If both times and locations are given, the the sampling will be scheduled on each specified location in each given time-point. Note that for the time-being, in the interest of simplicity, no sanity checks are performed on the locations given except the restriction that the sampling points must fall within the bounding box around the simulated world map. Other than that, `slendr` will simply instruct its SLiM backend script to sample individuals as close to the sampling points given as possible, regardless of whether those points lie within a population spatial boundary at that particular moment of time.



**Value**

Data frame with three columns: time of sampling, population to sample from, how many individuals to sample

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
path <- system.file("extdata/models/introgression", package = "slendr")
model <- read_model(path)

# afr and eur objects would normally be created before slendr model compilation,
# but here we take them out of the model object already compiled for this
# example (in a standard slendr simulation pipeline, this wouldn't be necessary)
afr <- model$populations[["AFR"]]
eur <- model$populations[["EUR"]]

# schedule the recording of 10 African and 100 European individuals from a
# given model at 20 ky, 10 ky, 5ky ago and at present-day (time 0)
schedule <- schedule_sampling(
  model, times = c(20000, 10000, 5000, 0),
  list(afr, 10), list(eur, 100)
)

# the result of `schedule_sampling` is a simple data frame (note that the locations
# of sampling locations have `NA` values because the model is non-spatial)
schedule
```

---

 setup\_env

*Setup a dedicated Python virtual environment for slendr*


---

**Description**

This function will automatically download a Python miniconda distribution dedicated to an R-Python interface. It will also create a slendr-specific Python environment with all the required Python dependencies.

**Usage**

```
setup_env(quiet = FALSE, agree = FALSE, pip = NULL)
```

**Arguments**

quiet	Should informative messages be printed to the console? Default is FALSE.
agree	Automatically agree to all questions?

pip Should pip be used instead of conda for installing slendr's Python dependencies? Note that this will still use the conda distribution to install Python itself, but will change the repository from which slendr will install its Python dependencies. Unless explicitly set to TRUE, Python dependencies will be installed from conda repositories by default, except for the case of osx-arm64 Mac architecture, for which conda dependencies are broken.

**Value**

No return value, called for side effects

---

set\_dispersal                      *Change dispersal parameters*

---

**Description**

Changes either the competition interactive distance, mating choice distance, or the dispersal of offspring from its parent

**Usage**

```
set_dispersal(
  pop,
  time,
  competition = NA,
  mating = NA,
  dispersal = NA,
  dispersal_fun = NULL
)
```

**Arguments**

pop	Object of the class slendr_pop
time	Time of the population size change
competition, mating	Maximum spatial competition and mating choice distance
dispersal	Standard deviation of the normal distribution of the distance that offspring disperses from its parent
dispersal_fun	Distribution function governing the dispersal of offspring. One of "normal", "uniform", "cauchy", "exponential", or "brownian" (in which vertical and horizontal displacements are drawn from a normal distribution independently).

**Value**

Object of the class slendr\_pop, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

**Examples**

```

# spatial definitions -----

# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

```

```

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

set\_range

*Update the population range*


---

### Description

This function allows a more manual control of spatial map changes in addition to the expand and move functions

### Usage

```

set_range(
  pop,
  time,
  center = NULL,
  radius = NULL,
  polygon = NULL,
  lock = FALSE
)

```

### Arguments

pop	Object of the class <code>slendr_pop</code>
time	Time of the change
center	Two-dimensional vector specifying the center of the circular range
radius	Radius of the circular range
polygon	List of vector pairs, defining corners of the polygon range (see also the <code>region</code> argument) or a geographic region of the class <code>slendr_region</code> from which the polygon coordinates will be extracted
lock	Maintain the same density of individuals. If <code>FALSE</code> (the default), the number of individuals in the population will not change. If <code>TRUE</code> , the number of individuals simulated will be changed (increased or decreased) appropriately, to match the new population range area.

**Value**

Object of the class `slendr_pop`, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

**Examples**

```
# spatial definitions -----

# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)
```

```

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

shrink\_range

*Shrink the population range*


---

## Description

Shrinks the spatial population range by a specified distance in a given time-window

## Usage

```

shrink_range(
  pop,
  by,
  end,
  start,
  overlap = 0.8,
  snapshots = NULL,
  lock = FALSE,
  verbose = TRUE
)

```

## Arguments

pop	Object of the class slendr_pop
by	How many units of distance to shrink by?
start, end	When does the boundary shrinking start/end?

overlap	Minimum overlap between subsequent spatial boundaries
snapshots	The number of intermediate snapshots (overrides the overlap parameter)
lock	Maintain the same density of individuals. If FALSE (the default), the number of individuals in the population will not change. If TRUE, the number of individuals simulated will be changed (increased or decreased) appropriately, to match the new population range area.
verbose	Report on the progress of generating intermediate spatial boundaries?

### Details

Note that because slendr models have to accomodate both SLiM and msprime back ends, population sizes and split times are rounded to the nearest integer value.

### Value

Object of the class `slendr_pop`, which contains population parameters such as name, time of appearance in the simulation, parent population (if any), and its spatial parameters such as map and spatial boundary.

### Examples

```
# spatial definitions -----
# create a blank abstract world 1000x1000 distance units in size
map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# create a circular population with the center of a population boundary at
# [200, 800] and a radius of 100 distance units, 1000 individuals at time 1
# occupying a map just specified
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100)

# printing a population object to a console shows a brief summary
pop1

# create another population occupying a polygon range, splitting from pop1
# at a given time point (note that specifying a map is not necessary because
# it is "inherited" from the parent)
pop2 <- population("pop2", N = 100, time = 50, parent = pop1,
                  polygon = list(c(100, 100), c(320, 30), c(500, 200),
                                c(500, 400), c(300, 450), c(100, 400)))

pop3 <- population("pop3", N = 200, time = 80, parent = pop2,
                  center = c(800, 800), radius = 200)

# move "pop1" to another location along a specified trajectory and saved the
# resulting object to the same variable (the number of intermediate spatial
# snapshots can be also determined automatically by leaving out the
# `snapshots = ` argument)
pop1_moved <- move(pop1, start = 100, end = 200, snapshots = 6,
                  trajectory = list(c(600, 820), c(800, 400), c(800, 150)))
```

```

pop1_moved

# many slendr functions are pipe-friendly, making it possible to construct
# pipelines which construct entire history of a population
pop1 <- population("pop1", N = 1000, time = 1,
                  map = map, center = c(200, 800), radius = 100) %>%
  move(start = 100, end = 200, snapshots = 6,
       trajectory = list(c(400, 800), c(600, 700), c(800, 400), c(800, 150))) %>%
  set_range(time = 300, polygon = list(
    c(400, 0), c(1000, 0), c(1000, 600), c(900, 400), c(800, 250),
    c(600, 100), c(500, 50))
  )

# population ranges can expand by a given distance in all directions
pop2 <- expand_range(pop2, by = 200, start = 50, end = 150, snapshots = 3)

# we can check the positions of all populations interactively by plotting their
# ranges together on a single map
plot_map(pop1, pop2, pop3)

# gene flow events -----

# individual gene flow events can be saved to a list
gf <- list(
  gene_flow(from = pop1, to = pop3, start = 150, end = 200, rate = 0.15),
  gene_flow(from = pop1, to = pop2, start = 300, end = 330, rate = 0.25)
)

# compilation -----

# compile model components in a serialized form to dist, returning a single
# slendr model object (in practice, the resolution should be smaller)
model <- compile_model(
  populations = list(pop1, pop2, pop3), generation_time = 1,
  resolution = 100, simulation_length = 500,
  competition = 5, mating = 5, dispersal = 1
)

```

---

slim

*Run a slendr model in SLiM*


---

## Description

This function will execute a SLiM script generated by the compile function during the compilation of a slendr demographic model.

## Usage

```

slim(
  model,

```



```

sequence_length,
recombination_rate,
samples = NULL,
output = NULL,
burnin = 0,
max_attempts = 1,
spatial = !is.null(model$world),
coalescent_only = TRUE,
method = c("batch", "gui"),
random_seed = NULL,
run = TRUE,
verbose = FALSE,
load = TRUE,
locations = NULL,
slim_path = NULL
)

```

### Arguments

model	Model object created by the compile function
sequence_length	Total length of the simulated sequence (in base-pairs)
recombination_rate	Recombination rate of the simulated sequence (in recombinations per basepair per generation)
samples	A data frame of times at which a given number of individuals should be remembered in the tree-sequence (see <code>schedule_sampling</code> for a function that can generate the sampling schedule in the correct format). If missing, only individuals present at the end of the simulation will be recorded in the tree-sequence output file.
output	Path to the output tree-sequence file. If NULL (the default), tree sequence will be saved to a temporary file.
burnin	Length of the burnin (in model's time units, i.e. years)
max_attempts	How many attempts should be made to place an offspring near one of its parents? Serves to prevent infinite loops on the SLiM backend. Default value is 1.
spatial	Should the model be executed in spatial mode? By default, if a world map was specified during model definition, simulation will proceed in a spatial mode.
coalescent_only	Should <code>initializeTreeSeq(retainCoalescentOnly = &lt;...&gt;)</code> be set to TRUE (the default) or FALSE? See "retainCoalescentOnly" in the SLiM manual for more detail.
method	How to run the script? ("gui" - open in SLiMgui, "batch" - run on the command line)
random_seed	Random seed (if missing, SLiM's own seed will be used)
run	Should the SLiM engine be run? If FALSE, the command line SLiM command will be printed (and returned invisibly as a character vector) but not executed.

verbose	Write the SLiM output log to the console (default FALSE)?
load	Should the final tree sequence be immediately loaded and returned? Default is TRUE. The alternative (FALSE) is useful when a tree-sequence file is written to a custom location to be loaded at a later point.
locations	If NULL, locations are not saved. Otherwise, the path to the file where locations of each individual throughout the simulation will be saved (most likely for use with <code>animate_model</code> ).
slim_path	Optional way to specify path to an appropriate SLiM binary (this is useful if the slim binary is not on the \$PATH).

### Value

A tree-sequence object loaded via Python-R reticulate interface function `ts_load` (internally represented by the Python object `tskit.trees.TreeSequence`). Optionally, depending on the value of the arguments `load =` or `run =`, nothing or a character vector, respectively.

### Examples

```
init_env()

# load an example model
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# afr and eur objects would normally be created before slendr model compilation,
# but here we take them out of the model object already compiled for this
# example (in a standard slendr simulation pipeline, this wouldn't be necessary)
afr <- model$populations[["AFR"]]
eur <- model$populations[["EUR"]]
chimp <- model$populations[["CH"]]

# schedule the sampling of a couple of ancient and present-day individuals
# given model at 20 ky, 10 ky, 5ky ago and at present-day (time 0)
modern_samples <- schedule_sampling(model, times = 0, list(afr, 5), list(eur, 5), list(chimp, 1))
ancient_samples <- schedule_sampling(model, times = c(30000, 20000, 10000), list(eur, 1))

# sampling schedules are just data frames and can be merged easily
samples <- rbind(modern_samples, ancient_samples)

# run a simulation using the SLiM back end from a compiled slendr model object and return
# a tree-sequence output
ts <- slim(model, sequence_length = 1e5, recombination_rate = 0, samples = samples)

# automatic loading of a simulated output can be prevented by `load = FALSE`, which can be
# useful when a custom path to a tree-sequence output is given for later downstream analyses
output_file <- tempfile(fileext = ".trees")
slim(model, sequence_length = 1e5, recombination_rate = 0, samples = samples,
      output = output_file, load = FALSE)
# ... at a later stage:
ts <- ts_load(output_file, model)
```

ts

---

subtract                      *Generate the difference between two slendr objects*

---

### Description

Generate the difference between two slendr objects

### Usage

```
subtract(x, y, name = NULL)
```

### Arguments

x	Object of the class slendr
y	Object of the class slendr
name	Optional name of the resulting geographic region. If missing, name will be constructed from the function arguments.

### Value

Object of the class slendr\_region which encodes a standard spatial object of the class sf with several additional attributes (most importantly a corresponding slendr\_map object, if applicable).

---

summary.slendr\_nodes    *Summarise the contents of a ts\_nodes result*

---

### Description

Summarise the contents of a ts\_nodes result

### Usage

```
## S3 method for class 'slendr_nodes'
summary(object, ...)
```

### Arguments

object	Data frame produced by the function ts_nodes
...	Additional formal arguments to the summary method (unused here)

### Value

Used for its output to the terminal

ts\_afs

*Compute the allele frequency spectrum (AFS)***Description**

This function computes the AFS with respect to the given set of individuals or nodes.

**Usage**

```
ts_afs(
  ts,
  sample_sets = NULL,
  mode = c("site", "branch", "node"),
  windows = NULL,
  span_normalise = FALSE,
  polarised = TRUE
)
```

**Arguments**

ts	Tree sequence object of the class <code>slendr_ts</code>
sample_sets	A list (optionally a named list) of character vectors with individual names (one vector per set). If <code>NULL</code> , allele frequency spectrum for all individuals in the tree sequence will be computed.
mode	The mode for the calculation ("sites" or "branch")
windows	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to <code>ts\$sequence_length</code> ) are added automatically
span_normalise	Argument passed to <code>tskit</code> 's <code>allele_frequency_spectrum</code> method
polarised	When <code>TRUE</code> (the default) the allele frequency spectrum will not be folded (i.e. the counts will assume knowledge of which allele is ancestral, and which is derived, which is known in a simulation)

**Details**

For more information on the format of the result and dimensions, in particular the interpretation of the first and the last element of the AFS (when `complete = TRUE`), please see the `tskit` manual at <https://tskit.dev/tskit/docs/stable/python-api.html> and the example section dedicated to AFS at [https://tskit.dev/tutorials/analysing\\_tree\\_sequences.html#allele-frequency-spectra](https://tskit.dev/tutorials/analysing_tree_sequences.html#allele-frequency-spectra).

**Value**

Allele frequency spectrum values for the given sample set. Note that the contents of the first and last elements of the AFS might surprise you. Read the links in the description for more detail on how `tskit` handles things.

**Examples**

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

samples <- ts_samples(ts) %>% .[$pop %in% c("AFR", "EUR"), ]

# compute AFS for the given set of individuals
ts_afs(ts, sample_sets = list(samples$name))

```

---

ts_ancestors	<i>Extract (spatio-)temporal ancestral history for given nodes/individuals</i>
--------------	--

---

**Description**

Extract (spatio-)temporal ancestral history for given nodes/individuals

**Usage**

```
ts_ancestors(ts, x, verbose = FALSE, complete = TRUE)
```

**Arguments**

ts	Tree sequence object of the class <code>slendr_ts</code>
x	Either an individual name or an integer node ID
verbose	Report on the progress of ancestry path generation?
complete	Does every individual in the tree sequence need to have complete metadata recorded? If TRUE, only individuals/nodes with complete metadata will be included in the reconstruction of ancestral relationships. For instance, nodes added during the coalescent recapitation phase will not be included because they don't have spatial information associated with them.

**Value**

A table of ancestral nodes of a given tree-sequence node all the way up to the root of the tree sequence

**Examples**

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# find the complete ancestry information for a given individual
ts_ancestors(ts, "EUR_1", verbose = TRUE)

```

---

ts\_coalesced

*Check that all trees in the tree sequence are fully coalesced*


---

**Description**

Check that all trees in the tree sequence are fully coalesced

**Usage**

```
ts_coalesced(ts, return_failed = FALSE)
```

**Arguments**

ts                    Tree sequence object of the class `slendr_ts`  
return\_failed       Report back which trees failed the coalescence check?

**Value**

TRUE or FALSE value if `return_failed = FALSE`, otherwise a vector of (tskit Python 0-based) indices of trees which failed the coalescence test

**Examples**

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

ts_coalesced(ts) # is the tree sequence fully coalesced? (TRUE or FALSE)

```

```
# returns a vector of tree sequence segments which are not coalesced
not_coalesced <- ts_coalesced(ts, return_failed = TRUE)
```

---

ts_descendants	<i>Extract all descendants of a given tree-sequence node</i>
----------------	--

---

## Description

Extract all descendants of a given tree-sequence node

## Usage

```
ts_descendants(ts, x, verbose = FALSE, complete = TRUE)
```

## Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
x	An integer node ID of the ancestral node
verbose	Report on the progress of ancestry path generation?
complete	Does every individual in the tree sequence need to have complete metadata recorded? If TRUE, only individuals/nodes with complete metadata will be included in the reconstruction of ancestral relationships. For instance, nodes added during the coalescent recapitation phase will not be included because they don't have spatial information associated with them.

## Value

A table of descendant nodes of a given tree-sequence node all the way down to the leaves of the tree sequence

## Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# find the complete descendency information for a given individual
ts_descendants(ts, x = 62, verbose = TRUE)
```

---

<code>ts_divergence</code>	<i>Calculate pairwise divergence between sets of individuals</i>
----------------------------	--

---

**Description**

Calculate pairwise divergence between sets of individuals

**Usage**

```
ts_divergence(
  ts,
  sample_sets,
  mode = c("site", "branch", "node"),
  windows = NULL,
  span_normalise = TRUE
)
```

**Arguments**

<code>ts</code>	Tree sequence object of the class <code>slendr_ts</code>
<code>sample_sets</code>	A list (optionally a named list) of character vectors with individual names (one vector per set)
<code>mode</code>	The mode for the calculation ("sites" or "branch")
<code>windows</code>	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to <code>ts\$sequence_length</code> ) do not have to be specified as they are added automatically.
<code>span_normalise</code>	Divide the result by the span of the window? Default TRUE, see the tskit documentation for more detail.

**Value**

For each pairwise calculation, either a single divergence value or a vector of divergence values (one for each window)

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

# collect sampled individuals from all populations in a list
```



```

sample_sets <- ts_samples(ts) %>%
  split(., .$pop) %>%
  lapply(function(pop) pop$name)

# compute the divergence between individuals from each sample set (list of
# individual names generated in the previous step)
ts_divergence(ts, sample_sets) %>% .[order(.$divergence), ]

```

---

ts\_diversity

*Calculate diversity in given sets of individuals*


---

### Description

Calculate diversity in given sets of individuals

### Usage

```

ts_diversity(
  ts,
  sample_sets,
  mode = c("site", "branch", "node"),
  windows = NULL,
  span_normalise = TRUE
)

```

### Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
sample_sets	A list (optionally a named list) of character vectors with individual names (one vector per set). If a simple vector is provided, it will be interpreted as <code>as.list(sample_sets)</code> , meaning that a given statistic will be calculated for each individual separately.
mode	The mode for the calculation ("sites" or "branch")
windows	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to <code>ts\$sequence_length</code> ) are added automatically)
span_normalise	Divide the result by the span of the window? Default TRUE, see the <code>tskit</code> documentation for more detail.

### Value

For each set of individuals either a single diversity value or a vector of diversity values (one for each window)

## Examples

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

# collect sampled individuals from all populations in a list
sample_sets <- ts_samples(ts) %>%
  split(., .$pop) %>%
  lapply(function(pop) pop$name)

# compute diversity in each population based on sample sets extracted
# in the previous step
ts_diversity(ts, sample_sets) %>% .[order(.$diversity), ]

```

---

ts\_draw

*Plot a graphical representation of a single tree*


---

## Description

This function first obtains an SVG representation of the tree by calling the `draw_svg` method of `tskit` and renders it as a bitmap image in R. All of the many optional keyword arguments of the `draw_svg` method can be provided and will be automatically passed to the method behind the scenes.

## Usage

```

ts_draw(
  x,
  width = 1000,
  height = 1000,
  labels = FALSE,
  sampled_only = TRUE,
  title = NULL,
  ...
)

```

## Arguments

x	A single tree extracted by <code>ts_tree</code>
width, height	Pixel dimensions of the rendered bitmap
labels	Label each node with the individual name?

sampled_only	Should only individuals explicitly sampled through simplification be labeled? This is relevant in situations in which sampled individuals can themselves be among the ancestral nodes.
title	Optional title for the figure
...	Keyword arguments to the tskit draw_svg function.

**Value**

No return value, called for side effects

**Examples**

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# extract the first tree in the tree sequence and draw it
tree <- ts_tree(ts, i = 1)

# ts_draw accepts various optional arguments of tskit.Tree.draw_svg
ts_draw(tree, time_scale = "rank")

```

---

ts_edges	<i>Extract spatio-temporal edge annotation table from a given tree or tree sequence</i>
----------	---

---

**Description**

Extract spatio-temporal edge annotation table from a given tree or tree sequence

**Usage**

```
ts_edges(x)
```

**Arguments**

x	Tree object generated by ts_phylo or a slendr tree sequence object produced by ts_load, ts_recapitate, ts_simplify, or ts_mutate
---	--

**Value**

Data frame of the sf type containing the times of nodes and start-end coordinates of edges across space

## Examples

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# extract an annotated table with (spatio-)temporal edge information
ts_edges(ts)

```

---

ts_eigenstrat	<i>Convert genotypes to the EIGENSTRAT file format</i>
---------------	--

---

## Description

EIGENSTRAT data produced by this function can be used by the admixr R package (<https://bodkan.net/admixr/>).

## Usage

```
ts_eigenstrat(ts, prefix, chrom = "chr1", outgroup = NULL)
```

## Arguments

ts	Tree sequence object of the class slendr_ts
prefix	EIGENSTRAT trio prefix
chrom	The name of the chromosome in the EIGENSTRAT snp file (default "chr1")
outgroup	Should a formal, artificial outgroup be added? If NULL (default), no outgroup is added. A non-NULL character name will serve as the name of the outgroup in an ind file.

## Details

In case an outgroup was not formally specified in a slendr model which generated the tree sequence data, it is possible to artificially create an outgroup sample with the name specified by the outgroup argument, which will carry all ancestral alleles (i.e. value "2" in a geno file for each position in a snp file).

## Value

Object of the class EIGENSTRAT created by the admixr package

---

`ts_f2`*Calculate the f2, f3, f4, and f4-ratio statistics*

---

**Description**

These functions present an R interface to the corresponding f-statistics methods in tskit.

**Usage**

```
ts_f2(  
  ts,  
  A,  
  B,  
  mode = c("site", "branch", "node"),  
  span_normalise = TRUE,  
  windows = NULL  
)
```

```
ts_f3(  
  ts,  
  A,  
  B,  
  C,  
  mode = c("site", "branch", "node"),  
  span_normalise = TRUE,  
  windows = NULL  
)
```

```
ts_f4(  
  ts,  
  W,  
  X,  
  Y,  
  Z,  
  mode = c("site", "branch", "node"),  
  span_normalise = TRUE,  
  windows = NULL  
)
```

```
ts_f4ratio(  
  ts,  
  X,  
  A,  
  B,  
  C,  
  O,  
  mode = c("site", "branch"),
```

```
    span_normalise = TRUE
  )
```

### Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
mode	The mode for the calculation ("sites" or "branch")
span_normalise	Divide the result by the span of the window? Default TRUE, see the tskit documentation for more detail.
windows	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to <code>ts\$sequence_length</code> ) do not have to be specified as they are added automatically.
W, X, Y, Z, A, B, C, O	Character vectors of individual names (largely following the nomenclature of Patterson 2021, but see crucial differences between tskit and ADMIXTOOLS in Details)

### Details

Note that the order of populations f3 statistic implemented in tskit (<https://tskit.dev/tskit/docs/stable/python-api.html#tskit.TreeSequence.f3>) is different from what you might expect from ADMIXTOOLS, as defined in Patterson 2012 (see <https://academic.oup.com/genetics/article/192/3/1065/5935193> under heading "The three-population test and introduction of f-statistics", as well as ADMIXTOOLS documentation at <https://github.com/DReichLab/AdmixTools/blob/master/README.3PopTest#L5>). Specifically, the widely used notation introduced by Patterson assumes the population triplet as  $f_3(C; A, B)$ , with C being the "focal" sample (i.e., either the outgroup or a sample tested for admixture). In contrast, tskit implements  $f_3(A; B, C)$ , with the "focal sample" being A.

Although this is likely to confuse many ADMIXTOOLS users, `slendr` does not have much choice in this, because its `ts_*`(`)` functions are designed to be broadly compatible with raw tskit methods.

### Value

Data frame with statistics calculated for the given sets of individuals

### Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk and add mutations to it
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

# calculate f2 for two individuals in a previously loaded tree sequence
ts_f2(ts, A = "AFR_1", B = "EUR_1")
```

```

# calculate f2 for two sets of individuals
ts_f2(ts, A = c("AFR_1", "AFR_2"), B = c("EUR_1", "EUR_3"))

# calculate f3 for two individuals in a previously loaded tree sequence
ts_f3(ts, A = "EUR_1", B = "AFR_1", C = "NEA_1")

# calculate f3 for two sets of individuals
ts_f3(ts, A = c("AFR_1", "AFR_2", "EUR_1", "EUR_2"),
      B = c("NEA_1", "NEA_2"),
      C = "CH_1")

# calculate f4 for single individuals
ts_f4(ts, W = "EUR_1", X = "AFR_1", Y = "NEA_1", Z = "CH_1")

# calculate f4 for sets of individuals
ts_f4(ts, W = c("EUR_1", "EUR_2"),
      X = c("AFR_1", "AFR_2"),
      Y = "NEA_1",
      Z = "CH_1")

# calculate f4-ratio for a given set of target individuals X
ts_f4ratio(ts, X = c("EUR_1", "EUR_2", "EUR_4", "EUR_5"),
          A = "NEA_1", B = "NEA_2", C = "AFR_1", O = "CH_1")

```

---

tsfst

*Calculate pairwise statistics between sets of individuals*


---

## Description

For a discussion on the difference between "site", "branch", and "node" options of the mode argument, please see the tskit documentation at <https://tskit.dev/tskit/docs/stable/stats.html#sec-stats-mode>.

## Usage

```

tsfst(
  ts,
  sample_sets,
  mode = c("site", "branch", "node"),
  windows = NULL,
  span_normalise = TRUE
)

```

## Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
sample_sets	A list (optionally a named list) of character vectors with individual names (one vector per set)

mode	The mode for the calculation ("sites" or "branch")
windows	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to ts\$sequence_length) do not have to be specified as they are added automatically.
span_normalise	Divide the result by the span of the window? Default TRUE, see the tskit documentation for more detail.

**Value**

For each pairwise calculation, either a single Fst value or a vector of Fst values (one for each window)

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

# compute F_st between two sets of individuals in a given tree sequence ts
tsfst(ts, sample_sets = list(afr = c("AFR_1", "AFR_2", "AFR_3"),
                             eur = c("EUR_1", "EUR_2")))
```

---

ts\_genotypes

*Extract genotype table from the tree sequence*


---

**Description**

Extract genotype table from the tree sequence

**Usage**

```
ts_genotypes(ts)
```

**Arguments**

ts Tree sequence object of the class slendr\_ts

**Value**

Data frame object of the class tibble containing genotypes of simulated individuals in columns



## Examples

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk, recapitate it, simplify it, and mutate it
ts <- ts_load(slendr_ts, model) %>%
  ts_recapitate(Ne = 10000, recombination_rate = 1e-8) %>%
  ts_simplify() %>%
  ts_mutate(mutation_rate = 1e-8)

# extract the genotype matrix (this could take a long time consume lots
# of memory!)
gts <- ts_genotypes(ts)

```

---

ts\_ibd

*Collect Identity-by-Descent (IBD) segments (EXPERIMENTAL)*


---

## Description

This function iterates over a tree sequence and returns IBD tracts between pairs of individuals or nodes

## Usage

```

ts_ibd(
  ts,
  coordinates = FALSE,
  within = NULL,
  between = NULL,
  squash = FALSE,
  minimum_length = NULL,
  maximum_time = NULL,
  sf = TRUE
)

```

## Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
coordinates	Should coordinates of all detected IBD tracts be reported? If <code>FALSE</code> (the default), only the total length of shared IBD segments and their numbers are reported. If <code>TRUE</code> , coordinates of each segment will be returned (but note that this can have a massive impact on memory usage). See details for more information.
within	A character vector with individual names or an integer vector with node IDs indicating a set of nodes within which to look for IBD segments.

between	A list of lists of character vectors with individual names or integer vectors with node IDs, indicating a set of nodes between which to look for shared IBD segments.
squash	Should adjacent IBD segments for pairs of nodes be squashed if they only differ by their 'genealogical paths' but not by their MRCA? Default is FALSE. For more context, see <a href="https://github.com/tskit-dev/tskit/issues/2459">https://github.com/tskit-dev/tskit/issues/2459</a> . This option is EXPERIMENTAL!
minimum_length	Minimum length of an IBD segment to return in results. This is useful for reducing the total amount of IBD returned (but see Details).
maximum_time	Oldest MRCA of a node to be considered as an IBD ancestor to return that IBD segment in results. This is useful for reducing the total amount of IBD returned.
sf	If IBD segments in a spatial tree sequence are being analyzed, should the returned table be a spatial sf object? Default is TRUE.

### Details

This function is considered experimental. For full control over IBD segment detection in tree-sequence data, users can (and perhaps, for the time being, should) rely on the tskit method `ibd_segments` (see [https://tskit.dev/tskit/docs/stable/python-api.html#tskit.TreeSequence.ibd\\_segments](https://tskit.dev/tskit/docs/stable/python-api.html#tskit.TreeSequence.ibd_segments)).

Internally, this function leverages the tskit `TreeSequence` method `ibd_segments`. However, note that the `ts_ibd` function always returns a data frame of IBD tracts, it does not provide an option to iterate over individual IBD segments as shown in the official tskit documentation at <https://tskit.dev/tskit/docs/stable/ibd.html>. In general, R handles heavy iteration poorly, and this function does not attempt to serve as a full wrapper to `ibd_segments`.

Unfortunately, the distinction between "squashed IBD" (what many would consider to be the expected definition of IBD) and tskit's IBD which is defined via distinct genealogical paths (see <https://github.com/tskit-dev/tskit/issues/2459> for a discussion of the topic), makes the meaning of the filtering parameter of the `ibd_segments()` method of tskit `minimum_length` somewhat unintuitive. As of this moment, this function argument filters on IBD segments on the tskit level, not the level of the squashed IBD segments!

### Value

A data frame with IBD results (either coordinates of each IBD segment shared by a pair of nodes, or summary statistics about the total IBD sharing for that pair)

### Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)
```

```
# find IBD segments between specified Neanderthals and Europeans
ts_ibd(
    ts,
    coordinates = TRUE,
    between = list(c("NEA_1", "NEA_2"), c("EUR_1", "EUR_2")),
    minimum_length = 40000
)
```

---

ts\_load

*Load a tree sequence file produced by a given model*

---

## Description

This function loads a tree sequence file simulated from a given slendr model. Optionally, the tree sequence can be recapitated and simplified.

## Usage

```
ts_load(file, model = NULL)
```

## Arguments

file	A path to the tree-sequence file (either originating from a slendr model or a standard non-slendr tree sequence).
model	Optional slendr_model object which produced the tree-sequence file. Used for adding various annotation data and metadata to the standard tskit tree-sequence object.

## Details

The loading, recapitation and simplification is performed using the Python module pylim which serves as a link between tree sequences generated by SLiM and the tskit module for manipulation of tree sequence data. All of these steps have been modelled after the official pylim tutorial and documentation available at: <https://tskit.dev/pylim/docs/latest/tutorial.html>.

The recapitation and simplification steps can also be performed individually using the functions [ts\\_recapitate](#) and [ts\\_simplify](#).

## Value

Tree-sequence object of the class slendr\_ts, which serves as an interface point for the Python module tskit using slendr functions with the ts\_ prefix.

## See Also

[ts\\_nodes](#) for extracting useful information about individuals, nodes, coalescent times and geospatial locations of nodes on a map

**Examples**

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load tree sequence generated by a given model
ts <- ts_load(slendr_ts, model)

# even tree sequences generated by non-slendr models can be
msprime_ts <- system.file("extdata/models/msprime.trees", package = "slendr")
ts <- ts_load(msprime_ts)

# load tree sequence and immediately simplify it only to sampled individuals
# (note that the example tree sequence is already simplified so this operation
# does not do anything in this case)
ts <- ts_load(slendr_ts, model = model) %>% ts_simplify(keep_input_roots = TRUE)

# load tree sequence and simplify it to a subset of sampled individuals
ts_small <- ts_simplify(ts, simplify_to = c("CH_1", "NEA_1", "NEA_2",
                                           "AFR_1", "AFR_2", "EUR_1", "EUR_2"))

# load tree sequence, recapitate it and simplify it
ts <- ts_load(slendr_ts, model) %>%
  ts_recapitate(recombination_rate = 1e-8, Ne = 10000, random_seed = 42) %>%
  ts_simplify()

# load tree sequence, recapitate it, simplify it and overlay neutral mutations
ts <- ts_load(slendr_ts, model) %>%
  ts_recapitate(recombination_rate = 1e-8, Ne = 10000, random_seed = 42) %>%
  ts_simplify() %>%
  ts_mutate(mutation_rate = 1e-8)

ts

```

---

ts\_metadata

*Extract list with tree sequence metadata saved by SLiM*


---

**Description**

Extract list with tree sequence metadata saved by SLiM

**Usage**

```
ts_metadata(ts)
```

**Arguments**

ts                    Tree sequence object of the class slendr\_ts

**Value**

List of metadata fields extracted from the tree-sequence object

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# extract the list of metadata information from the tree sequence
ts_metadata(ts)
```

---

ts\_mutate

*Add mutations to the given tree sequence*


---

**Description**

Add mutations to the given tree sequence

**Usage**

```
ts_mutate(
  ts,
  mutation_rate,
  random_seed = NULL,
  keep_existing = TRUE,
  mut_type = NULL
)
```

**Arguments**

ts                    Tree sequence object of the class slendr\_ts

mutation\_rate       Mutation rate used by msprime to simulate mutations

random\_seed         Random seed passed to msprime's mutate method

keep\_existing       Keep existing mutations?

mut\_type            Assign SLiM mutation type to neutral mutations? If NULL (default), no special mutation type will be used. If an integer number is given, mutations of the SLiM mutation type with that integer identifier will be created.

**Value**

Tree-sequence object of the class `slendr_ts`, which serves as an interface point for the Python module `tskit` using `slendr` functions with the `ts_` prefix.

**See Also**

[ts\\_nodes](#) for extracting useful information about individuals, nodes, coalescent times and geospatial locations of nodes on a map

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

ts <- ts_load(slendr_ts, model)
ts_mutate <- ts_mutate(ts, mutation_rate = 1e-8, random_seed = 42)

ts_mutate
```

---

ts\_names

*Extract names of individuals in a tree sequence*

---

**Description**

Extract names of individuals in a tree sequence

**Usage**

```
ts_names(ts, split = NULL)
```

**Arguments**

ts	Tree sequence object of the class <code>slendr_ts</code>
split	Should sample names in the tree sequence be split by a column (a population or time column)? Default is <code>NULL</code> and all names of samples will be returned as a single character vector. If set to "pop" or "time", a list of character vectors will be returned, one vector for each unique "pop" or "time" grouping.

**Value**

A vector of character sample names. If `split` is specified, a list of such vectors is returned, one element of the list per population or sampling time.

---

ts_nodes	<i>Extract combined annotated table of individuals and nodes</i>
----------	--

---

## Description

This function combines information from the table of individuals and table of nodes into a single data frame which can be used in downstream analyses.

## Usage

```
ts_nodes(x, sf = TRUE)
```

## Arguments

x	Tree sequence object of the class <code>slendr_ts</code> or a phylo object extracted by <code>ts_phylo</code>
sf	Should spatial data be returned in an sf format? If <code>FALSE</code> , spatial geometries will be returned simply as x and y columns, instead of the standard POINT data type.

## Details

The source of data (tables of individuals and nodes recorded in the tree sequence generated by SLiM) are combined into a single data frame. If the model which generated the data was spatial, coordinates of nodes (which are pixel-based by default because SLiM spatial simulations occur on a raster), the coordinates are automatically converted to an explicit spatial object of the `sf` class unless `spatial = FALSE`. See <https://r-spatial.github.io/sf/> for an extensive introduction to the `sf` package and the ways in which spatial data can be processed, analysed, and visualised.

## Value

Data frame with processed information from the tree sequence object. If the model which generated this data was spatial, result will be returned as a spatial object of the class `sf`.

## See Also

[ts\\_table](#) for accessing raw tree sequence tables without added metadata annotation. See also [ts\\_ancestors](#) to learn how to extract information about relationship between nodes in the tree sequence, and how to analysed data about distances between nodes in the spatial context.

## Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))
```

```
# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# extract an annotated table with (spatio-)temporal node information
ts_nodes(ts)
```

---

ts\_phylo

---

*Convert a tree in the tree sequence to an object of the class phylo*


---

### Description

Convert a tree in the tree sequence to an object of the class phylo

### Usage

```
ts_phylo(
  ts,
  i,
  mode = c("index", "position"),
  labels = c("tskit", "pop"),
  quiet = FALSE
)
```

### Arguments

ts	Tree sequence object of the class slendr_ts
i	Position of the tree in the tree sequence. If mode = "index", an i-th tree will be returned (in zero-based indexing as in tskit), if mode = "position", a tree covering the i-th base of the simulated genome will be returned (again, in tskit's indexing).
mode	How should the i argument be interpreted? Either "index" as an i-th tree in the sequence of genealogies, or "position" along the simulated genome.
labels	What should be stored as node labels in the final phylo object? Options are either a population name or a tskit integer node ID (which is a different thing from a phylo class node integer index).
quiet	Should ape's internal phylo validity test be printed out?

### Value

Standard phylogenetic tree object implemented by the R package ape



**Examples**

```

init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>%
  ts_recapitate(Ne = 10000, recombination_rate = 1e-8) %>%
  ts_simplify()

# extract the 1st tree from a given tree sequence, return ape object
tree <- ts_phylo(ts, i = 1, mode = "index", quiet = TRUE)
tree

# extract the tree at a 42th basepair in the given tree sequence
tree <- ts_phylo(ts, i = 42, mode = "position", quiet = TRUE)

# because the tree is a standard ape phylo object, we can plot it easily
plot(tree, use.edge.length = FALSE)
ape::nodelabels()

```

---

ts_recapitate	<i>Recapitate the tree sequence</i>
---------------	-------------------------------------

---

**Description**

Recapitate the tree sequence

**Usage**

```

ts_recapitate(
  ts,
  recombination_rate,
  Ne = NULL,
  demography = NULL,
  random_seed = NULL
)

```

**Arguments**

ts	Tree sequence object loaded by ts_load
recombination_rate	A constant value of the recombination rate
Ne	Effective population size during the recapitation process

demography	Ancestral demography to be passed internally to <code>msprime.sim_ancestry()</code> (see <code>msprime</code> 's documentation for mode detail)
random_seed	Random seed passed to <code>pyslim</code> 's <code>recapitate</code> method

**Value**

Tree-sequence object of the class `slendr_ts`, which serves as an interface point for the Python module `tskit` using `slendr` functions with the `ts_` prefix.

**See Also**

[ts\\_nodes](#) for extracting useful information about individuals, nodes, coalescent times and geospatial locations of nodes on a map

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

ts <- ts_load(slendr_ts, model) %>%
  ts_recapitate(recombination_rate = 1e-8, Ne = 10000, random_seed = 42)

ts
```

---

ts_samples	<i>Extract names and times of individuals of interest in the current tree sequence (either all sampled individuals or those that the user simplified to)</i>
------------	--

---

**Description**

Extract names and times of individuals of interest in the current tree sequence (either all sampled individuals or those that the user simplified to)

**Usage**

```
ts_samples(ts)
```

**Arguments**

ts                    Tree sequence object of the class `slendr_ts`

**Value**

Table of individuals scheduled for sampling across space and time

## Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# extract the table of individuals scheduled for simulation and sampling
ts_samples(ts)
```

---

ts_save	<i>Save a tree sequence to a file</i>
---------	---------------------------------------

---

## Description

Save a tree sequence to a file

## Usage

```
ts_save(ts, file)
```

## Arguments

ts	Tree sequence object loaded by <code>ts_load</code>
file	File to which the tree sequence should be saved

## Value

No return value, called for side effects

## Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree sequence
ts <- ts_load(slendr_ts, model)

# save the tree-sequence object to a different location
another_file <- paste(tempfile(), ".trees")
ts_save(ts, another_file)
```

---

ts_segregating	<i>Calculate the density of segregating sites for the given sets of individuals</i>
----------------	---

---

### Description

Calculate the density of segregating sites for the given sets of individuals

### Usage

```
ts_segregating(
  ts,
  sample_sets,
  mode = c("site", "branch", "node"),
  windows = NULL,
  span_normalise = FALSE
)
```

### Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
sample_sets	A list (optionally a named list) of character vectors with individual names (one vector per set). If a simple vector is provided, it will be interpreted as <code>as.list(sample_sets)</code> , meaning that a given statistic will be calculated for each individual separately.
mode	The mode for the calculation ("sites" or "branch")
windows	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to <code>ts\$sequence_length</code> ) are added automatically
span_normalise	Divide the result by the span of the window? Default TRUE, see the <code>tskit</code> documentation for more detail.

### Value

For each set of individuals either a single diversity value or a vector of diversity values (one for each window)

### Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)
```

```
# collect sampled individuals from all populations in a list
sample_sets <- ts_samples(ts) %>%
  split(., .$pop) %>%
  lapply(function(pop) pop$name)

ts_segregating(ts, sample_sets)
```

---

ts\_simplify

*Simplify the tree sequence down to a given set of individuals*


---

### Description

This function is a convenience wrapper around the `simplify` method implemented in `tskit`, designed to work on tree sequence data simulated by SLiM using the **slendr** R package.

### Usage

```
ts_simplify(
  ts,
  simplify_to = NULL,
  keep_input_roots = FALSE,
  keep_unary = FALSE,
  keep_unary_in_individuals = FALSE,
  filter_nodes = TRUE
)
```

### Arguments

<code>ts</code>	Tree sequence object of the class <code>slendr_ts</code>
<code>simplify_to</code>	A character vector of individual names. If <code>NULL</code> , all explicitly remembered individuals (i.e. those specified via the <a href="#">schedule_sampling</a> function) will be left in the tree sequence after the simplification.
<code>keep_input_roots</code>	Should the history ancestral to the MRCA of all samples be retained in the tree sequence? Default is <code>FALSE</code> .
<code>keep_unary</code>	Should unary nodes be preserved through simplification? Default is <code>FALSE</code> .
<code>keep_unary_in_individuals</code>	Should unary nodes be preserved through simplification if they are associated with an individual recorded in the table of individuals? Default is <code>FALSE</code> . Cannot be set to <code>TRUE</code> if <code>keep_unary</code> is also <code>TRUE</code>
<code>filter_nodes</code>	Should nodes be reindexed after simplification? Default is <code>TRUE</code> . See <code>tskit</code> 's documentation for the Python method <code>simplify()</code>

**Details**

The simplification process is used to remove redundant information from the tree sequence and retains only information necessary to describe the genealogical history of a set of samples.

For more information on how simplification works in pyslim and tskit, see the official documentation at <https://tskit.dev/tskit/docs/stable/python-api.html#tskit.TreeSequence.simplify> and <https://tskit.dev/pyslim/docs/latest/tutorial.html#simplification>.

A very clear description of the difference between remembering and retaining and how to use these techniques to implement historical individuals (i.e. ancient DNA samples) is in the pyslim documentation at <https://tskit.dev/pyslim/docs/latest/tutorial.html#historical-individuals>.

**Value**

Tree-sequence object of the class `slendr_ts`, which serves as an interface point for the Python module `tskit` using `slendr` functions with the `ts_` prefix.

**See Also**

[ts\\_nodes](#) for extracting useful information about individuals, nodes, coalescent times and geospatial locations of nodes on a map

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

ts <- ts_load(slendr_ts, model)
ts

# simplify tree sequence to sampled individuals
ts_simplified <- ts_simplify(ts)

# simplify to a subset of sampled individuals
ts_small <- ts_simplify(ts, simplify_to = c("CH_1", "NEA_1", "NEA_2", "AFR_1",
                                           "AFR_2", "EUR_1", "EUR_2"))

ts_small
```

---

ts\_table

*Get the table of individuals/nodes/edges/mutations from the tree sequence*

---

**Description**

This function extracts data from a given tree sequence table. All times are converted to model-specific time units from tskit's "generations backwards" time direction.

**Usage**

```
ts_table(ts, table = c("individuals", "edges", "nodes", "mutations"))
```

**Arguments**

ts	Tree sequence object of the class <code>slendr_ts</code>
table	Which tree sequence table to return

**Details**

For further processing and analyses, the output of the function [ts\\_nodes](#) might be more useful, as it merges the information in node and individual tables into one table and further annotates it with useful information from the model configuration data.

**Value**

Data frame with the information from the give tree-sequence table (can be either a table of individuals, edges, nodes, or mutations).

**See Also**

[ts\\_nodes](#) and [ts\\_edges](#) for accessing an annotated, more user-friendly and analysis-friendly tree-sequence table data

**Examples**

```
# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk and add mutations to it
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

# get the 'raw' tskit table of individuals
ts_table(ts, "individuals")

# get the 'raw' tskit table of edges
ts_table(ts, "edges")

# get the 'raw' tskit table of nodes
ts_table(ts, "nodes")

# get the 'raw' tskit table of mutations
ts_table(ts, "mutations")
```

ts\_tajima

*Calculate Tajima's D for given sets of individuals***Description**

For a discussion on the difference between "site" and "branch" options of the mode argument, please see the tskit documentation at <https://tskit.dev/tskit/docs/stable/stats.html#sec-stats-mode>

**Usage**

```
ts_tajima(ts, sample_sets, mode = c("site", "branch", "node"), windows = NULL)
```

**Arguments**

ts	Tree sequence object of the class <code>slendr_ts</code>
sample_sets	A list (optionally a named list) of character vectors with individual names (one vector per set). If a simple vector is provided, it will be interpreted as <code>as.list(sample_sets)</code> , meaning that a given statistic will be calculated for each individual separately.
mode	The mode for the calculation ("sites" or "branch")
windows	Coordinates of breakpoints between windows. The first coordinate (0) and the last coordinate (equal to <code>ts\$sequence_length</code> ) are added automatically)

**Value**

For each set of individuals either a single Tajima's D value or a vector of Tajima's D values (one for each window)

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model) %>% ts_mutate(mutation_rate = 1e-8, random_seed = 42)

# calculate Tajima's D for given sets of individuals in a tree sequence ts
ts_tajima(ts, list(eur = c("EUR_1", "EUR_2", "EUR_3", "EUR_4", "EUR_5"),
                  nea = c("NEA_1", "NEA_2")))
```



---

ts_tracts	<i>Extract ancestry tracts from a tree sequence (EXPERIMENTAL)</i>
-----------	--

---

### Description

Extract a data frame with coordinates of ancestry tracts from a given tree sequence.

### Usage

```
ts_tracts(ts, census, squashed = TRUE, source = NULL, target = NULL)
```

### Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
census	Census time. See the documentation linked in the Details for more information. If a <code>slendr</code> -specific tree sequence was provided as <code>ts</code> , the census time is expected to be given in <code>slendr</code> model-specific time units, and must correspond to some gene-flow event encoded by the model.
squashed	Should ancestry tracts be squashed (i.e., should continuous tracts that can be traced to different ancestral nodes be merged)? Default is <code>TRUE</code> . If <code>FALSE</code> , these effectively continuous ancestry tracts will be split into individual segments, each assigned to a specific ancestral node ID (recorded in a column <code>ancestor_id</code> ).
source	From which source population to extract tracts for? if <code>NULL</code> (the default), ancestry tracts for all populations contributing gene flow at the census time will be reported. Otherwise, ancestry tracts from only specified source populations will be extracted. Note that this option is ignored for non- <code>slendr</code> tree sequences!
target	Similar purpose as <code>source</code> above, except that it filters for tracts discovered in the target population(s)

### Details

This function implements an R-friendly interface to an algorithm for extracting ancestry tracts provided by the Python module `tspop` <https://tspop.readthedocs.io/en/latest/> and described by Georgia Tsambos. Please make sure to cite their paper which describes the algorithm in detail: <https://academic.oup.com/bioinformaticsadvances/article/3/1/vbad163/7429395>. For more technical details, see also the tutorial at: <https://tspop.readthedocs.io/en/latest/basicusage.html>.

In general, when using this function on a `slendr`-generated tree sequence, please be aware that the output changes slightly to what you would get by running the pure `tspop.get_pop_ancestry()` in Python. First, `ts_tracts()` populates the output data frame with additional metadata (such as names of individuals or populations). Additionally, for `slendr` models, it is specifically designed to only return ancestry tracts originating to an ancestral population which contributed its ancestry during a gene-flow event which started at a specific time (i.e., scheduled in a model via the `gene_flow()` function). It does not return every single ancestry tract present in the tree sequence for every single sample node (and every single potential ancestry population) as does the `tspop.get_pop_ancestry()` Python method.

That said, when run on a tree sequence which does not originate from a slendr simulation, the behavior of `ts_tracts()` is identical to that of the underlying `tspop.get_pop_ancestry()`.

As of the current version of slendr, `ts_tracts()` only works for slendr/msprime sequences but not on slendr/SLiM tree sequences. Support for slendr-generated SLiM tree sequences is in development. Tree sequences from non-slendr msprime and SLiM tree sequences is not restricted in any way and, as mentioned in the previous paragraph, `ts_tracts()` on such tree sequences effectively reduces to the standard `tspop.get_pop_ancestry()` call.

## Value

A data frame containing coordinates of ancestry tracts

## Examples

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_msprime.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk and add mutations to it
ts <- ts_load(slendr_ts, model)

# extract Neanderthal ancestry tracts (i.e. those corresponding to the
# census event at the gene-flow time at 55000 kya)
nea_tracts <- ts_tracts(ts, census = 55000, source = "NEA")
nea_tracts
```

---

ts\_tree

*Get a tree from a given tree sequence*

---

## Description

For more information about optional keyword arguments see tskit documentation: <https://tskit.dev/tskit/docs/stable/python-api.html#the-treesequences-class>

## Usage

```
ts_tree(ts, i, mode = c("index", "position"), ...)
```

## Arguments

ts	Tree sequence object of the class <code>slendr_ts</code>
i	Position of the tree in the tree sequence. If <code>mode = "index"</code> , an <i>i</i> -th tree will be returned (in zero-based indexing as in tskit), if <code>mode = "position"</code> , a tree covering the <i>i</i> -th base of the simulated genome will be returned (again, in tskit's indexing).

mode	How should the <code>i</code> argument be interpreted? Either "index" as an <code>i</code> -th tree in the sequence of genealogies, or "position" along the simulated genome.
...	Additional keyword arguments accepted by <code>tskit.TreeSequence.at</code> and <code>tskit.TreeSequence.at_in</code> methods

**Value**

Python-reticulate-based object of the class `tskit.trees.Tree`

**Examples**

```
init_env()

# load an example model with an already simulated tree sequence
slendr_ts <- system.file("extdata/models/introgression_slim.trees", package = "slendr")
model <- read_model(path = system.file("extdata/models/introgression", package = "slendr"))

# load the tree-sequence object from disk
ts <- ts_load(slendr_ts, model)

# extract the zero-th tree in the tree sequence
tree <- ts_tree(ts, i = 0)

# extract the tree at a position in the tree sequence
tree <- ts_tree(ts, i = 100000, mode = "position")
```

---

ts_vcf	<i>Save genotypes from the tree sequence as a VCF file</i>
--------	--

---

**Description**

Save genotypes from the tree sequence as a VCF file

**Usage**

```
ts_vcf(ts, path, chrom = NULL, individuals = NULL)
```

**Arguments**

ts	Tree sequence object of the class <code>slendr_ts</code>
path	Path to a VCF file
chrom	Chromosome name to be written in the <code>CHROM</code> column of the VCF
individuals	A character vector of individuals in the tree sequence. If missing, all individuals present in the tree sequence will be saved.

**Value**

No return value, called for side effects

---

 world

*Define a world map for all spatial operations*


---

### Description

Defines either an abstract geographic landscape (blank or containing user-defined landscape) or using a real Earth cartographic data from the Natural Earth project (<https://www.naturalearthdata.com>).

### Usage

```
world(
  xrange,
  yrange,
  landscape = "naturalearth",
  crs = NULL,
  scale = c("small", "medium", "large")
)
```

### Arguments

xrange	Two-dimensional vector specifying minimum and maximum horizontal range ("longitude" if using real Earth cartographic data)
yrange	Two-dimensional vector specifying minimum and maximum vertical range ("latitude" if using real Earth cartographic data)
landscape	Either "blank" (for blank abstract geography), "naturalearth" (for real Earth geography) or an object of the class <i>sf</i> defining abstract geographic features of the world
crs	EPSG code of a coordinate reference system to use for spatial operations. No CRS is assumed by default (NULL), implying an abstract landscape not tied to any real-world geographic region (when <code>landscape = "blank"</code> or when <code>landscape</code> is a custom-defined geographic landscape), or implying WGS-84 (EPSG 4326) coordinate system when a real Earth landscape was defined ( <code>landscape = "naturalearth"</code> ).
scale	If Natural Earth geographic data is used (i.e. <code>landscape = "naturalearth"</code> ), this parameter determines the resolution of the data used. The value "small" corresponds to 1:110m data and is provided with the package, values "medium" and "large" correspond to 1:50m and 1:10m respectively and will be downloaded from the internet. Default value is "small".

### Value

Object of the class `slendr_map`, which encodes a standard spatial object of the class `sf` with additional `slendr`-specific attributes such as requested x-range and y-range.

**Examples**

```
# create a blank abstract world 1000x1000 distance units in size
blank_map <- world(xrange = c(0, 1000), yrange = c(0, 1000), landscape = "blank")

# it is possible to construct custom landscapes (islands, corridors, etc.)
island1 <- region("island1", polygon = list(c(10, 30), c(50, 30), c(40, 50), c(0, 40)))
island2 <- region("island2", polygon = list(c(60, 60), c(80, 40), c(100, 60), c(80, 80)))
island3 <- region("island3", center = c(20, 80), radius = 10)
archipelago <- island1 %>% join(island2) %>% join(island3)

custom_map <- world(xrange = c(1, 100), c(1, 100), landscape = archipelago)

# real Earth landscapes can be defined using freely-available Natural Earth
# project data and with the possibility to specify an appropriate Coordinate
# Reference System, such as this example of a map of Europe

real_map <- world(xrange = c(-15, 40), yrange = c(30, 60), crs = "EPSG:3035")
```

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