

slimr: An R package for integrating data and tailor-made population genomic simulations over space and time

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Running headline: Integrating SLiM and R with slimr

Abstract

1) Software for realistically simulating complex population genomic processes is revolutionizing our understanding of evolutionary processes, and providing novel opportunities for integrating empirical with simulated data. However, the integration between simulation software and software designed for working with empirical data is currently not well developed. In particular, SLiM 3.0, which is one of the most powerful population genomic simulation frameworks for linking evolutionary dynamics with ecological patterns and processes, has a unique scripting language which limits its availability to potential users. Here we present slimr, an R package designed to

create a seamless link between SLiM 3.0 and the R development environment, with its powerful data manipulation and analysis tools.

- 2) We show how *slimr*, in combination with SLiM, facilitates smooth integration between genetic data, ecological data and simulation in a single environment. The package enables pipelines that begin with data reading, cleaning, and manipulation, proceed to constructing empirically-based parameters and initial conditions for simulations, then to running numerical simulations, and finally to retrieving simulation results in a format suitable for comparisons with empirical data – aided by advanced analysis and visualization tools provided by R (such as ABC and deep learning).
- 3) We demonstrate the use of *slimr* with an example from our own work on the landscape population genomics of desert mammals, highlighting the advantage of having a single integrated tool for both data analysis and simulation.
- 4) *slimr* makes the powerful simulation ability of SLiM 3.0 directly accessible to R users, allowing integrated simulation projects that incorporate empirical data without the need to switch between software environments. This should provide more opportunities for evolutionary biologists and ecologists to use realistic simulations to better understand the interplay between ecological and evolutionary processes. *slimr* is available at <https://rdinnager.github.io/slimr/>.

Keywords: population genomics; simulation; landscape genomics; evolution; ecology; evolutionary ecology; application; software

Introduction

Mathematical modelling and simulation are critical cornerstones of population genetic practice. At a fundamental level, empirical datasets demand analytical tool-kits that can accommodate their high complexity, and recent developments in sophisticated simulation software have the potential to provide mechanistic insight into increasingly complex evolutionary scenarios (Strand 2002; Carvajal-Rodríguez

2010; Yuan *et al.* 2012; Messer 2013; Hoban 2014; Kelleher *et al.* 2016; Haller & Messer 2019).
However, utilising flexible simulations requires exploration of large parameter space, which often
generates large amounts of data that need sophisticated computational tools to unpack, interrogate and
synthesize. Likewise, using simulations to model empirical data is an emerging field because it allows
researchers to deal with complex situations where it is difficult to obtain a closed likelihood (Beaumont *et al.* 2002; Marjoram *et al.* 2003; Sisson 2018; Torada *et al.* 2019; Brehmer *et al.* 2020; Wang *et al.* 2020; Cranmer *et al.* 2020). To facilitate more rapid and seamless interrogation and synthesis between
empirical data and population genetics simulation, we present `slimr` (<https://rdinnager.github.io/slimr/>).
`slimr` is an R package designed to link the very large and widely used ecosystem of analysis and
visualization tools in the R statistical language to the SLiM scripting language (Haller & Messer 2019), a
popular, powerful and flexible population genetics simulation tool. The package creates a smooth fusion
between the computational power and flexible model specification of SLiM with the advanced statistical
analysis, visualisation, and metaprogramming tools of R.

Package Description

`slimr` is an R package that interfaces with SLiM 3.0 software for forward population genetics simulations
(see Messer 2013; Haller & Messer 2019 for full details on SLiM, as well as the website at
<https://messerlab.org/slim/>).

`slimr` implements a Domain Specific Language (DSL) that mimics the syntax of SLiM, allowing users to
write and run SLiM scripts and capture resulting simulation output, all within the R environment. Much of
the syntax is identical to SLiM, but `slimr` offers additional R functions that allow users to manipulate
SLiM scripts (“`slimr` verbs”) by inserting them directly into any SLiM code block. This enables R users to
create SLiM scripts that explore large numbers of different parameters and also automatically produce
output from SLiM for powerful downstream analysis within R.

74 The features of slimr fall into three categories: 1) SLiM script integrated development, 2) data
75 input/output, and 3) SLiM script metaprogramming. The first set of features is designed to make it easy to
76 develop SLiM scripts in an R development environment such as Rstudio, and mostly recapitulates
77 features that SLiM users already have access to in the form of SLiMgui and QtSLiM
78 (<https://messengerlab.org/slim/>). The second and third features are implemented using what we call “slimr
79 verbs”, allowing SLiM and R features to be combined in advanced ways. The integration between R and
80 SLiM provided by slimr compensates knowledgeable users of R for a lack of knowledge of SLiM, helping
81 to lower the barrier to learning and using SLiM.

82 Each of the 3 categories has subcategories of features as follows:

83 1) Integrated Development

- 84 a) Autocomplete and Documentation (within R) for SLiM code
- 85 b) Code highlighting and pretty printing of SLiM code
- 86 c) Rstudio addins
- 87 d) Run code in SLiM from R

88 2) Data Input/Output

- 89 a) Automatic output generation and extraction from SLiM to R (`slimr_output()`)
- 90 b) Insert arbitrary R objects into SLiM scripts through inlining (`slimr_inline()`)

91 3) Metaprogramming

- 92 a) Code templating for SLiM scripts (`slimr_template()`)
- 93 b) Flexible general metaprogramming tools (support for `rlang`'s `!!` and `!!!` forcing
94 operators)

95 In the next section we describe each of these features in greater detail, showing examples through
96 screenshots and code snippets.

97 Integrated Development

98 `slimr` allows the user to write SLiM code from within an R integrated development environment (IDE).
99 `slimr` is designed to work well with Rstudio, but can be used in any R IDE. The syntax used to write SLiM
100 code is very similar to the native SLiM syntax, with a few modifications to make it work with the R
101 interpreter. As an example, here is a minimal SLiM program, and its counterpart written in `slimr`.

102 **SLiM code:**

```
103 initialize()  
104 {  
105   initializeMutationRate(1e-7);  
106   initializeMutationType("m1", 0.5, "f", 0.0);  
107   initializeGenomicElementType("g1", m1, 1.0);  
108   initializeGenomicElement(g1, 0, 99999);  
109   initializeRecombinationRate(1e-8);  
110 }  
111 1  
112 {  
113   sim.addSubpop("p1", 500);  
114 }  
115 10000  
116 {  
117   sim.simulationFinished();  
118 }
```

119 **slimr code:**

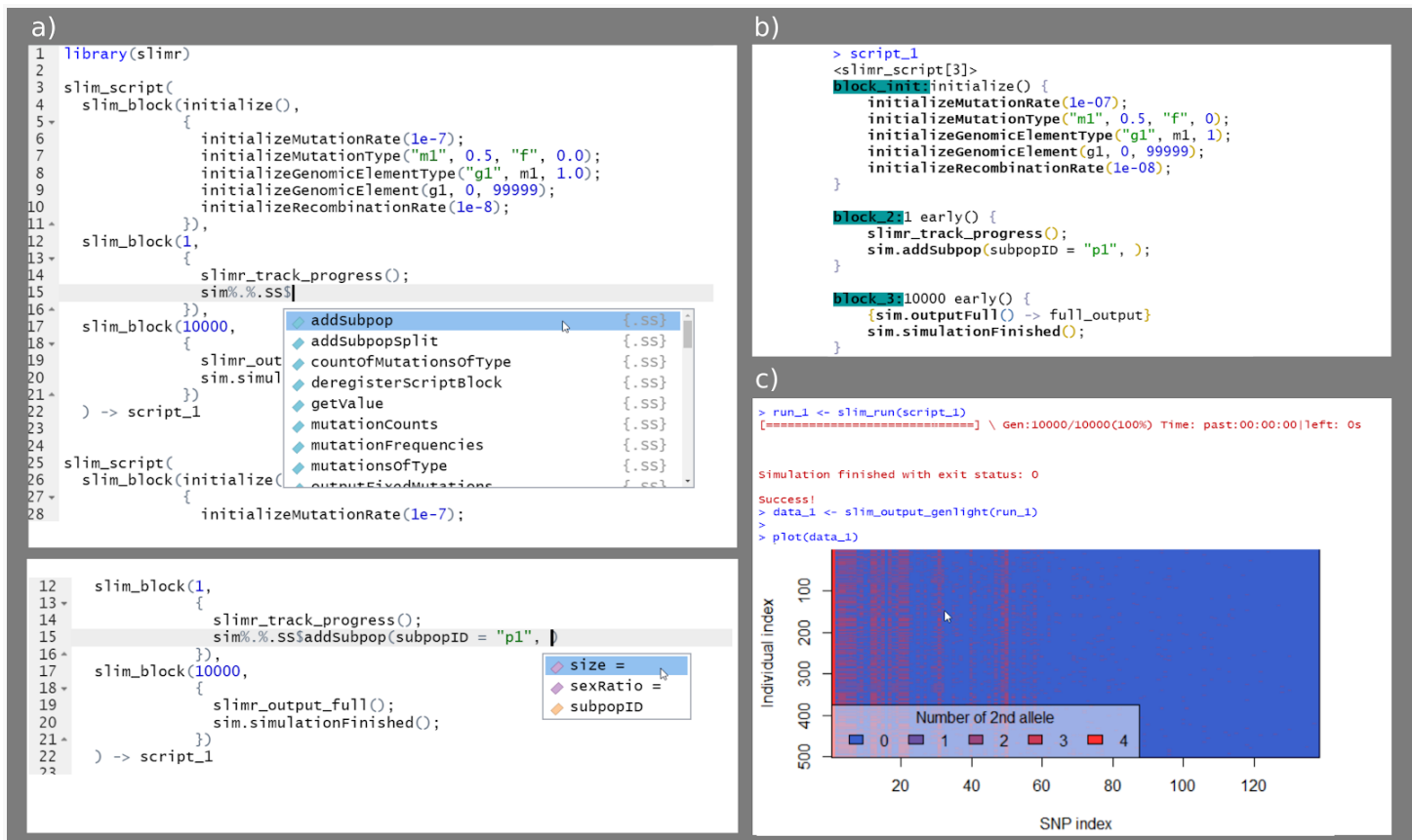
```
120 slim_script(  
121   slim_block(initialize(),  
122     {  
123       initializeMutationRate(1e-7);  
124       initializeMutationType("m1", 0.5, "f", 0.0);  
125       initializeGenomicElementType("g1", m1, 1.0);  
126       initializeGenomicElement(g1, 0, 99999);  
127       initializeRecombinationRate(1e-8);  
128     }  
129   ),  
130   slim_block(1,  
131     {  
132       sim.addSubpop("p1", 500);  
133     }  
134   )  
135 )
```

```

132     }},
133     slim_block(10000,
134     {
135         slimr_output_full();
136         sim.simulationFinished();
137     })
138 ) -> script_1

```

139 The above code assigns the script to an R object `script_1`, which can then be further manipulated,
 140 printed prettily, and sent to SLiM to run. See Fig. 1 to see what the above script looks like in the Rstudio
 141 IDE, and examples of things you can do with it. A script is specified using the `slimr_script` function,
 142 within which you create `slimr` coding blocks, using the `slimr_block` function. The user can create as
 143 many `slimr` code blocks as desired within a `slimr_script`. We've added a `slimr` "verb"
 144 (`slimr_output_full`), which tells SLiM to output the full state of the simulation and return it to R during
 145 the execution of the block. We will discuss `slimr` verbs in more detail in the next section.



146 **Figure 1.** Screenshots of working with `slimr` in Rstudio. An example of autocomplete for SLiM code (a),
 147 an example of pretty printing of a `slimr_script` object (b), and an example of running a script,
 148 converting its output to a standard R format for genetic data (adeigenet package's `genlight`), and plotting
 149 it.

150 `slimr` makes it easy to write SLiM code in R after the user learns a few differences between SLiM and
151 `slimr`. This means users can learn how to write complex SLiM simulations by reading SLiM 3.0
152 documentation and the examples found within it (<https://messerlab.org/slim/>). To make this process
153 easier for `slimr` users, the entire reference documentation for functions in SLiM 3.0 and Eidos scripting
154 language (on which SLiM is based) is included in `slimr` (with the original author's permission). Hence, not
155 only can R users look up relevant SLiM functions in their R session, but the R IDE can perform
156 autocompletion.

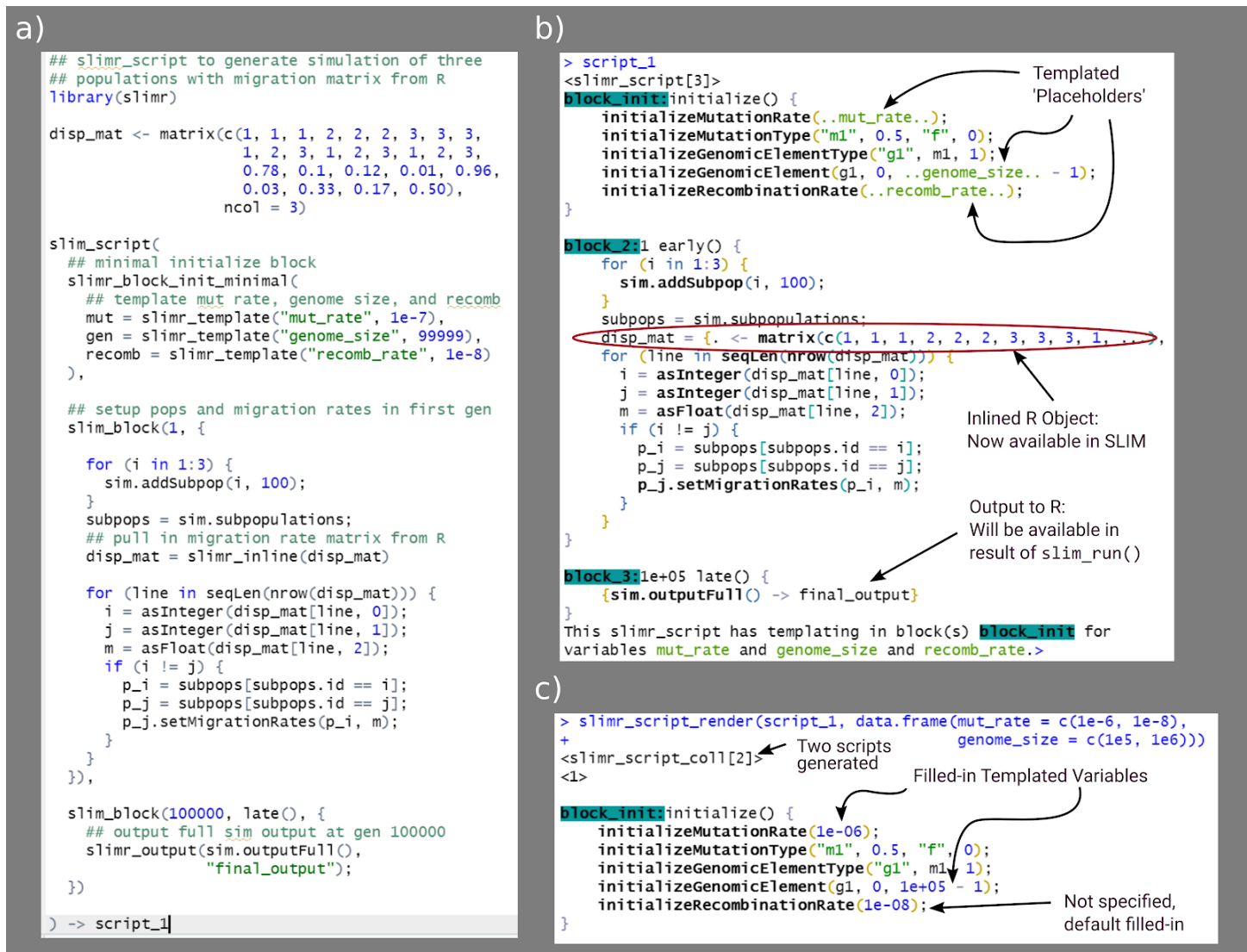
157 `slimr` also provides several Rstudio addins to make common tasks simple. These include an addin that
158 converts SLiM code to `slimr` code automatically by pasting from the clipboard, and an addin to easily
159 send `slimr_script` calls to be run in SLiM. Converting code from `slimr` to SLiM is also supported,
160 including the ability to open the converted code in SLiMGUI or QtSLiM if installed.

161 `slimr_script` objects (and `slimr_script_coll`, which contain lists of `slimr_script` objects) can be
162 run in SLiM, and their results collected and returned using the `slim_run` function.

163 Data Input/Output

164 The input/output and metaprogramming features of `slimr` are achieved using special `slimr` “verbs” that
165 can be inserted directly into `slimr` coding blocks (Fig. 2). These verbs are pure R functions that modify
166 how the SLiM script will be generated and run in SLiM. They are not passed directly to SLiM, but make it
167 easy for R to interact with SLiM. In this way, `slimr` code appears to be a hybrid between SLiM and R
168 code. `slimr` verbs allow all setup and logic required to use SLiM with R to occur inside the coding blocks
169 comprising the `slimr_script` object, thus requiring fewer arguments to be set in preparation for
170 downstream analysis (for example `slim_run` does not require many complex arguments because most

171 of what it needs to know is embedded in the `slimr_script` object). In our experience, this leads to a
 172 very smooth experience using `slimr` by reducing the frequency of switches between different mental
 173 modes. By convention, all `slimr` verbs have the prefix `slimr_`, and are meant to be used only inside
 174 `slim_script` calls. All other `slimr` functions are prefixed with `slim_`, which means they are to be used
 175 on `slimr_script` objects, and not inside a `slim_script` call. The following are the main `slimr` verbs
 176 supported:



177 **Figure 2.** Example of a single script using the main `slimr` verbs (`slimr_template`, `slimr_output`, and
 178 `slimr_inline`). A) Code to specify the `slimr_script`. B) Pretty printing of the script, showing special
 179 `slimr` syntax. C) Example of running `slim_script_render` on the `slimr_script` object, demonstrating
 180 how placeholder variables specified in `slimr_template` are replaced with provided values. All code from
 181 the above example can be accessed as a package vignette
 182 (https://rdinnager.github.io/slimr/articles/simple_example.html).

183 `slimr_inline(object, delay = FALSE)`

184 `slimr_inline` allows `slimr` users to embed (or “inline”) an R object directly into a SLiM script so that it
185 can be accessed from within a SLiM simulation. This is a powerful way to use empirical data that has
186 been generated, loaded, and / or cleaned from within R within a simulation. `slimr_inline` automatically
187 detects the type of R object and attempts to coerce it into a format compatible with SLiM. Currently
188 supported types are all atomic vectors, matrices, arrays, and Raster* objects from the raster package,
189 which will allow users to insert maps for use in spatial simulations.

190 `slimr_output(slimr_expr, name, do_every = 1, callbacks = NULL)`

191 `slimr_output` makes it simple to output data from the simulation by wrapping a SLiM expression. Where
192 it is called in the `slimr_script`, it will produce SLiM code to take the output of the expression and send
193 it to R. The output will be available after running `slim_run` in the returned object as a `data.frame`. Output
194 can even be accessed live during the simulation run via the use of callback functions. The `do_every`
195 argument tells `slimr_output` not to output every time it is called, but rather only after every `do_every`
196 generations.

197 `slimr` includes several functions to create different commonly desired outputs and visualizations, which
198 use the `slimr_output_prefix` (e.g. `slimr_output_nucleotides()`, which outputs DNA sequences data
199 for nucleotide-based simulations).

200 Metaprogramming

201 Metaprogramming is programming that generates or manipulates programming code itself. `slimr` has
202 facilities for manipulating SLiM programming code and generating scripts. The main `slimr` verb for doing
203 this is `slimr_template`. `slimr` also supports the metaprogramming operators for forcing (!!) and splicing

204 (!!!), as used in the `{rlang}` R package. Here we briefly describe `slimr_template`, designed to help
205 users easily generate many versions of a `slimr_script` with different parameters.

206 `slimr_template(var_name, default = NULL)`

207 `slimr_template` allows the user to insert “templated” variables into a `slimr_script`; the call to
208 `slimr_template` will be replaced in the SLiM script with a placeholder with the name `var_name`. This
209 placeholder can be replaced with values of the user’s choice by calling
210 `slim_script_render(slm_script, template = tmplt)`, and providing a template – a list or
211 `data.frame` containing values with names matching `var_name`. This action can be performed on multiple
212 `slimr_template` variables simultaneously, as well as producing multiple replicate scripts with different
213 combinations of replacements. This feature can create a swathe of parameter values to be run
214 (automatically) in parallel to explore parameter space, conduct sensitivity analyses, or fit data to
215 simulation output using methods requiring many simulation runs, such as Approximate Bayesian
216 Computation (ABC). Users can provide a default value for each templated variable, which will be used if
217 the user does not specify a replacement for that variable.

218 These features together make `slimr` far more than a simple wrapper for SLiM – its goal is to enhance
219 and complement SLiM by creating a hybrid domain specific language for R. We plan to continue to
220 increase integration of our package with SLiM, and to continuously update it as new SLiM versions are
221 released in the future.

222 `slim_run(slimr_ob, slim_path, parallel = FALSE)`

223 Once a `slimr_script` or `slimr_script_coll` object has been created, with all SLiM simulation logic
224 and `slimr` verbs for interacting with R, it can be sent to the SLiM software to be run using the `slim_run`
225 function. To access this functionality, users must install SLiM on their computers. This is facilitated by the

226 `slim_setup()` function, which will attempt to automatically install a platform appropriate version of SLiM
227 on the user's system and set it up to work with `slimr`.

228 Calling `slim_run` will run the simulation. While the simulation is running, `slimr_run` produces progress
229 updates if requested, as well as any output generated by calls to `slimr_output` with custom callbacks. If
230 called on a `slimr_script_coll` containing multiple `slimr_script` objects, each `slimr_script` object
231 will be run, optionally in parallel, and the result returned in a list.

232 Once finished, `slim_run` will return a `slimr_results` object, which contains information about the
233 simulation run, such as whether it succeeded or failed, any error messages produced, all output
234 generated from `slimr_output` calls, and any file names where additional data from the run are stored. This
235 can then be used for any downstream analysis the user desires.

236 Example: Investigating population genomics of small 237 mammals in a periodic environment

238 In this section we provide a brief description of a full example analysis using simulation (Fig. 3, Fig. 4).
239 Full code for the example can be found in the supplementary material.

240 The context for this example is a long-term ecological study in the Simpson Desert in central Australia.
241 Several authors of this paper have studied the population dynamics of small mammals and reptiles in this
242 desert for more than 30 years (Dickman *et al.* 2014; Greenville *et al.* 2016, 2017). Recently, we have
243 begun sequencing tissue samples taken from animals captured during the past 15 years, and obtained
244 single nucleotide polymorphism (SNP) data using DArT technology. Here, we use SNP data from 167
245 individuals of a common native rodent species, the sandy inland mouse *Pseudomys hermannsburgensis*,
246 sampled at 7 sites over three years (2006-2008), and subsequently aggregated to 3 subpopulations for

247 analysis. The three sample years span periods before and after a major rainfall event at the end of 2006;
248 big rains occur infrequently in the study region (every 8-12 years) (Greenville *et al.* 2012) but drive major
249 population eruptions.

250 We used the SNP data to calculate pairwise F_{st} values among the three subpopulations in each year,
251 revealing that pairwise F_{st} values dropped rapidly to nearly zero immediately after the rainfall event from
252 a high recorded just prior to the event when the populations were more genetically differentiated. We
253 interpreted this result to mean that the rainfall event, which caused the sandy inland mouse population to
254 rapidly increase, also allowed animals to move out of spatially scattered refuge patches to which they
255 had been confined during the preceding dry period (Dickman *et al.* 2011). This movement allowed the
256 subpopulations to mix, leading to a decrease in population genetic structure as measured by F_{st} .

257 In the example, we use simulations to evaluate our interpretation regarding the processes driving
258 changes in F_{st} values. We found that our initial hypothesis, that the rainfall event led to the mixing of
259 previously unconnected populations in refuge patches, provided a good match to the data when we
260 simulated the population and genomic processes. However, we also identified several other processes
261 that could generate similar outcomes, which raises the question as to what data or analyses would be
262 required to distinguish among these competing processes.

263 The results from these preliminary simulations will thus be invaluable in guiding which individuals and
264 time periods we should focus sequencing on to maximize the chances of distinguishing among
265 competing hypotheses that might explain the combined population and genetic patterns in the data.
266 Ultimately, we aim to use this approach to understand how future climate change could alter the
267 population and genetic structure of desert animals, highlighting the value of *slimr* in a scientific workflow.

a)

```

pop_sim <- slim_script(
  slim_block(initialize(), {
    initializeMutationRate(slimr_template("mut_rate", 1e-6));
    initializeMutationType("m1", 0.5, "n", 0, slimr_template("selection_strength", 0.1));
    initializeGenomicElementType("g1", m1, 1.0);
    initializeGenomicElement(g1, 0, slimr_template("genome_size", 50000) - 1);
    initializeRecombinationRate(slimr_template("recomb_rate", 1e-8));
    initializeSex("A");
    defineConstant("abund", slimr_inline(pop_abunds, delay = TRUE));
    defineConstant("sample_these", slimr_inline(sample_these, delay = TRUE));
  }),
  slim_block(1, {
    init_pop = slimr_inline(init_popsiz, delay = TRUE)

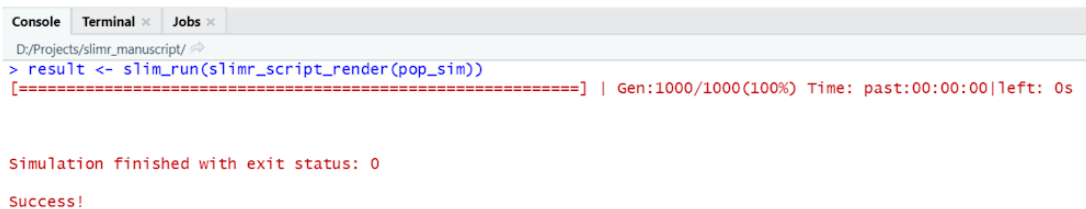
    ## set populations to initial size
    sim.addSubpop("p1", asInteger(init_pop[0]));
    sim.addSubpop("p2", asInteger(init_pop[1]));
    sim.addSubpop("p3", asInteger(init_pop[2]));
  }),
  slim_block(1, late(), {
    ## get starting population from a file which we will fill-in later
    sim.readFromPopulationFile(slimr_inline(starting_pop, delay = TRUE));
    ## migration on or off flags for pops 1-3 (using tag)
    p1.tag = 0;
    p2.tag = 0;
    p3.tag = 0;
  }),
  slim_block(1, 1000, late(), {
    ## update generation number
    gen = sim.generation %% 50
    if(gen == 0) {
      ## set population size to observed levels
      p1.setSubpopulationSize(asInteger(ceil(abund[0, gen - 1] * slimr_template("popsiz_scaling", 100))));
      p2.setSubpopulationSize(asInteger(ceil(abund[1, gen - 1] * ..popsiz_scaling..)));
      p3.setSubpopulationSize(asInteger(ceil(abund[2, gen - 1] * ..popsiz_scaling..)));

      ## increase migration when above abundance threshold
      if(p1.tag == 0 & abund[0, gen - 1] > slimr_template("abund_threshold", 5)) {
        p2.setMigrationRates(p1, slimr_template("migration_rate", 0))
        p3.setMigrationRates(p1, ..migration_rate..)
        p1.tag = 1;
      }
      if(p1.tag == 1 & abund[0, gen - 1] <= ..abund_threshold..) {
        p2.setMigrationRates(p1, 0)
        p3.setMigrationRates(p1, 0)
        p1.tag = 0;
      }
      if(p2.tag == 0 & abund[1, gen - 1] > ..abund_threshold..) {
        p3.setMigrationRates(p2, ..migration_rate..)
        p2.tag = 1;
      }
      if(p2.tag == 1 & abund[1, gen - 1] <= ..abund_threshold..) {
        p3.setMigrationRates(p2, 0)
        p2.tag = 0;
      }
      if(p3.tag == 0 & abund[2, gen - 1] > ..abund_threshold..) {
        p1.setMigrationRates(p3, ..migration_rate..)
        p1.tag = 1;
      }
      if(p3.tag == 1 & abund[2, gen - 1] <= ..abund_threshold..) {
        p1.setMigrationRates(p3, 0)
        p3.tag = 0;
      }

      if(any(match(sample_these, sim.generation) >= 0)) {
        ind_sample = sample(sim.subpopulations.individuals, 50)
        slimr_output(ind_sample.genomes.output(), "pop_sample", do_every = 1);
      }
    }
  }),
  slim_block(1000, late(), {
  })
)

```

b)



```

D:/Projects/slimr_manuscript/
> result <- slimr_run(slimr_script_render(pop_sim))
[=====] | Gen:1000/1000(100%) Time: past:00:00:00|left: 0s

Simulation finished with exit status: 0
Success!

```

268 **Figure 3.** Screenshots of the main `slimr` example from this manuscript. **a)** Rstudio screenshot showing
 269 completed `slimr` script for specifying the model. **Note:** some code sections have been collapsed for
 270 brevity. **b)** Screen shot of the Rstudio console after running the example using default values for
 271 templated variables. This shows how fast SLiM can run – this example took less than 1 second!

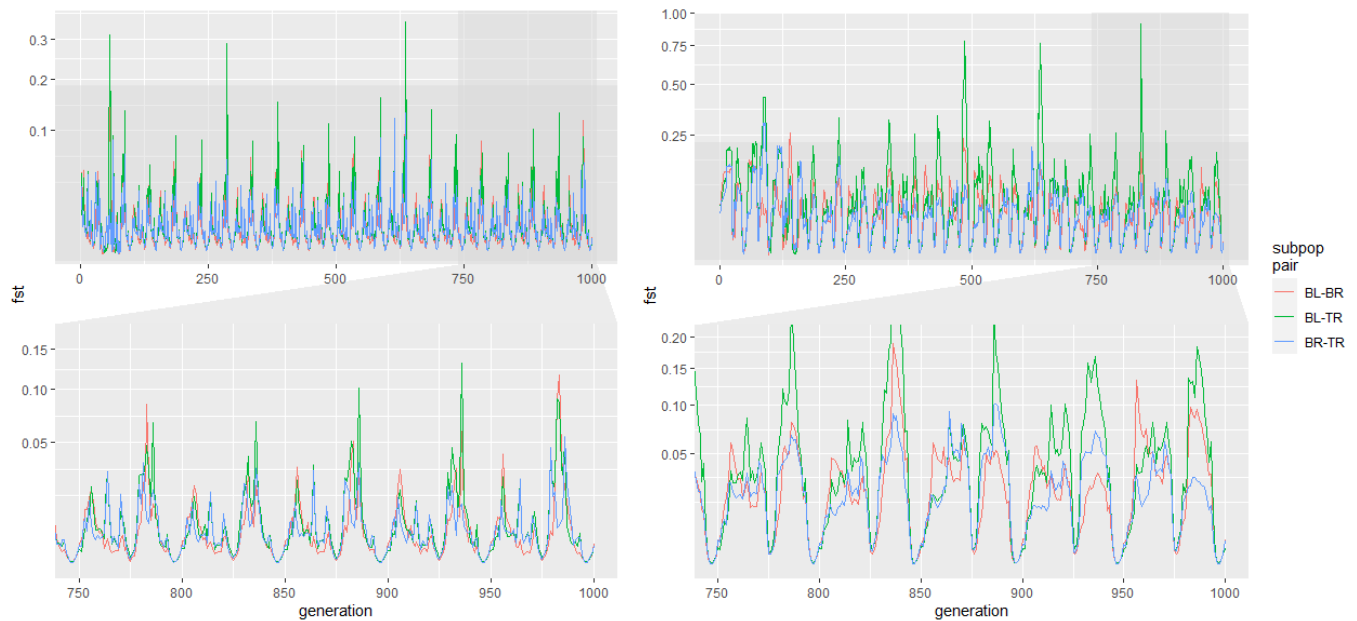


Figure 4. Two very different evolutionary scenarios that produce similar patterns of fluctuations in pairwise F_{st} between three subpopulations when simulated using *slimr*. **Left Panel:** Maximum migration rate between subpopulations is at its maximum value (panmictic), selection is relatively high, and migration is “on” all the time. **Right Panel:** Maximum migration rate is 0.2 (20% population exchange), there is no selection, and migration is only “on” when total population size is high (e.g. during and just after rainfall events), otherwise there is no migration.

Acknowledgments

We thank Benjamin C. Haller and Phillip W. Messer for permission to reproduce the documentation and examples of SLiM in *slimr*, and for valuable feedback on the package (from B.C.H.). Thanks to Emily Stringer for providing additional test data to help develop the methods used in this manuscript.

Author Contributions

RD, BG, SS, and RPD developed the concept for the package. RD coded the package and wrote the manuscript draft. CD, GW, and AG contributed data for testing of the package, and BG helped test the package as a user. All authors contributed critically to manuscript drafts and gave final approval for publication.

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