egssimtools: an R package for the analysis of a model of speciation

1 Introduction

The ExplicitGenomeSpeciation program is a simulation of a speciation event with explicit genetics and genotype-phenotype map (see the main page for details). This vignette introduces egssimtools, an R package that comes with the simulation program, containing a series of tools to read and analyze the simulation data within R. Here we will show how to use it with a few use cases. We assume that the program has been run and that simulation data have already been saved. Throughout the vignette we will use example simulation data from the data folder.

The functions in egssimtools provide an interface between the data saved by the simulation, which consist in binary files (see details on the main page), and the R environment. Specifically, these functions try as much as possible to produce data frames allowing to process, plot and analyze the many types of data that can be retrieved from the simulations in multiple ways, using the tidyverse workflow. As such, the functions make heavy use of the tidyverse packages and their outputs are tailored to eing use in tidyverse pipelines, especially plotting with ggplot2. We recommend the user to be familiar with the tidyverse and some of its extensions, such as patchwork, which we will use throughout this vignette. We refer the reader to the ggplot2 documentation to customize the plots as needed, as this is out of the scope of this vignette.

Because of the diversity of the simulation data, and the large number of ways they can be viewed, this package avoids providing ready-made functions to plot specific results directly from the simulation folders. Instead, we provide functions such that pretty much any plot can be produced in a few chunks of code only, with a common flow. We will go through examples here and explain the usage of the functions as we go.

In a first part we will take a tour of the main functions in the package, and in a second part we will cover several use-cases with short snippets of code showing how to read, process, and plot the data.

2 Installation

As this package comes as part of the ExplicitGenomeSpeciation repository, it cannot be installed from GitHub using devtools::install_github. Instead, you can install it by running devtools::install() from within the egssimtools folder, or by opening the project egssimtools.Rproj in RStudio and clicking on "Install and Restart", in the "Build" menu.

3 Use cases

Here we show how to use the package through a series of examples, with increasing complexity. You can find all examples in the scripts/examples.R script. For more specific information about each function, please refer to the documentation. Pretty much all use cases go through the same repeated phases of (1) reading, (2) processing and (3) plotting the data.

We start by loading the packages we will need.

```
library(egssimtools)
library(tidyverse)
```

library(patchwork)

```
library(tidygraph)
library(ggraph)
```

Next, we set up the path to one of our simulations.

```
root <- "../data/example_1"</pre>
```

3.1 Simulation-wise data

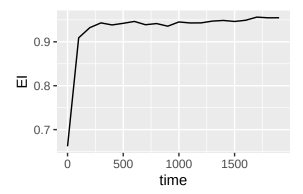
Data saved as a single value per time point, such as the degree of ecological divergence, are the easiest to read and plot. We can read these simulation-wise variables using read_sim:

```
data <- read_sim(root, "EI")</pre>
data
# # A tibble: 20 x 2
#
      time EI
#
    <dbl> <dbl>
#
        0 0.662
#
  2
      100 0.909
#
  3
      200 0.932
#
      300 0.943
 4
  5
#
      400 0.938
#
  6
      500 0.942
  7
      600 0.946
#
 8
      700 0.939
#
  9
      800 0.942
# 10
      900 0.936
# 11 1000 0.945
# 12 1100 0.943
# 13 1200 0.943
# 14 1300 0.947
# 15 1400 0.949
# 16 1500 0.946
# 17 1600 0.949
# 18 1700 0.956
# 19 1800 0.955
# 20 1900 0.955
```

Here, the function read_sim reads the files time.dat (by default) and EI.dat, which have the same dimensions. Note that most reading functions read time.dat by default.

We can then use the regular ggplot2 workflow to plot the data:

```
ggplot(data, aes(x = time, y = EI)) +
geom_line()
```



3.2 Pivotting the data

It is possible to read multiple variables, for example:

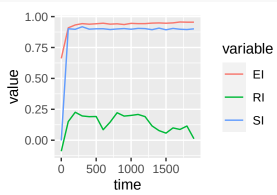
```
data <- read_sim(root, c("EI", "RI", "SI"))</pre>
data
# # A tibble: 20 x 4
#
                       RI
      time
               EI
                                 SI
#
     <dbl> <dbl>
                    <db1>
                              <db1>
#
         0 0.662
                  -0.0892
   1
                           -0.00245
#
   2
       100 0.909
                   0.151
                            0.901
#
   3
       200 0.932
                   0.226
                            0.897
#
   4
       300 0.943
                   0.196
                            0.918
#
   5
       400 0.938
                   0.190
                            0.898
#
   6
       500 0.942
                   0.192
                            0.901
   7
#
       600 0.946
                   0.0847
                            0.902
#
   8
       700 0.939
                   0.147
                            0.895
#
   9
       800 0.942
                   0.222
                            0.900
# 10
       900 0.936
                   0.195
                            0.903
# 11
      1000 0.945
                   0.200
                            0.898
# 12
      1100 0.943
                   0.208
                            0.904
# 13
      1200 0.943
                   0.191
                            0.903
# 14
      1300 0.947
                   0.116
                            0.894
# 15
      1400 0.949
                   0.0775
                            0.907
                   0.0571
# 16
      1500 0.946
                            0.893
# 17
      1600 0.949
                   0.0994
                            0.904
# 18
      1700 0.956
                   0.0870
                            0.898
# 19
      1800 0.955
                   0.115
                            0.895
     1900 0.955
                   0.0116 0.901
```

We could plot the variables indepdently from this data frame, but for practical purposes it is often handy to reshape such "wide" data frame into its "long" counterpart (according to the nomenclature of the tidyverse), where several columns are gathered in a single one. We use pivot_data (which internally calls pivot_longer from tidyr) for this:

```
3
          0 SI
                      -0.00245
#
       100 EI
                       0.909
   4
#
   5
       100 RI
                       0.151
#
   6
       100 SI
                       0.901
#
   7
       200 EI
                       0.932
#
   8
       200 RI
                       0.226
#
   9
       200 SI
                       0.897
#
 10
       300 EI
                       0.943
        with 50 more rows
```

where we specify that the EI, RI and SI columns must be gathered in a single one. We can then use this long data frame to plot the variables, for example, in different colors:

```
ggplot(data, aes(x = time, y = value, color = variable)) +
  geom_line()
```



3.3 Splitting variables into several columns

Some files contain data that may have to be splitted into multiple columns in order to be arranged with other variables in a single data frame with a common unit of observation. For example, the file Fst contains genome-wide Fst values for each trait, therefore consisting of three values for each time point. To read this file into a simulation-wise data frame (and combine it with the time column), we must split it into three columns, one for each trait. The by argument of read_sim does exactly that:

```
data <- read_sim(root, "Fst", by = 3)</pre>
data
# # A tibble: 20 x 4
#
      time
               Fst1
                         Fst2
                                  Fst3
                        <db1>
#
     <db1>
              <db1>
                                 <db1>
#
         0 0.00158 0.000659 0.000451
   1
#
   2
       100 0.0371
                    0.0167
                              0.0165
#
   3
       200 0.0454
                    0.0176
                              0.0228
#
   4
       300 0.0509
                    0.0232
                              0.0247
#
   5
       400 0.0517
                    0.0257
                              0.0254
#
   6
       500 0.0568
                    0.0280
                              0.0304
#
   7
       600 0.0608
                    0.0268
                              0.0328
#
   8
       700 0.0617
                    0.0351
                              0.0338
   9
       800 0.0646
                    0.0411
                              0.0379
#
 10
       900 0.0646
                    0.0472
                              0.0440
      1000 0.0697
                              0.0481
# 11
                    0.0509
                              0.0529
# 12
      1100 0.0678
                    0.0535
      1200 0.0722 0.0540
# 13
                              0.0511
```

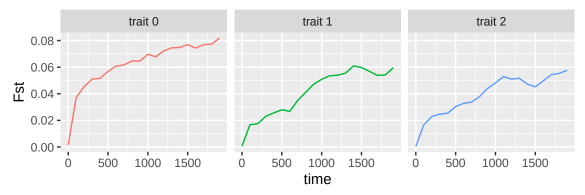
```
# 14 1300 0.0745
                   0.0555
                            0.0516
                            0.0473
# 15
    1400 0.0749
                   0.0610
# 16
     1500 0.0769
                            0.0452
                   0.0597
# 17 1600 0.0744
                   0.0569
                            0.0497
# 18
     1700 0.0769
                   0.0538
                            0.0543
# 19
     1800 0.0773
                   0.0542
                            0.0553
# 20
     1900 0.0819 0.0596
                            0.0576
```

Again, we may want to pivot this table to the long format to facilitate plotting:

```
data <- pivot_data(data, paste0("Fst", 1:3), newnames = paste0("trait ", 0:2))</pre>
data <- data %>% rename(trait = "variable", Fst = "value")
data
# # A tibble: 60 x 3
#
      time trait
                         Fst
     <dbl> <fct>
#
                       <db1>
#
   1
         0 trait 0 0.00158
#
   2
         0 trait 1 0.000659
         0 trait 2 0.000451
#
   3
#
   4
       100 trait 0 0.0371
   5
#
       100 trait 1 0.0167
       100 trait 2 0.0165
   7
#
       200 trait 0 0.0454
       200 trait 1 0.0176
#
  9
       200 trait 2 0.0228
# 10
       300 trait 0 0.0509
# # ... with 50 more rows
```

Here, we used the newnames argument of pivot_data to replace the labels Fst1, Fst2 and Fst3 by the mentions trait 0, trait 1 and trait 2, and we used the rename function from dplyr to rename the variable column to trait. This produces a nicer graph:

```
ggplot(data, aes(x = time, y = Fst, color = trait)) +
  geom_line() +
  facet_grid(. ~ trait) +
  theme(legend.position = "none")
```



3.4 Individual-wise data

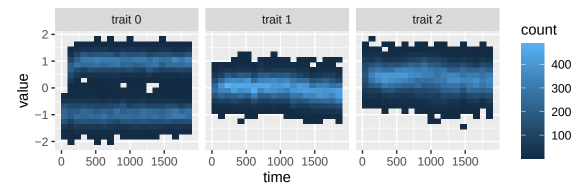
While variables such as time, EI or Fst are (or can easily be converted to) simulation-wise variables, some files contain data with other units of observation, such as the individual organism, thus consisting of one value per individual per time point. To read individual-wise data, use read_pop:

```
data <- read_pop(root, "individual_trait", by = 3)</pre>
data
# # A tibble: 33,953 x 4
#
      time\ individual\_trait1\ individual\_trait2\ individual\_trait3
#
     <db1>
                         <dbl>
                                              <db1>
                                                                  <db1>
#
   1
          0
                        -1.20
                                            0.0673
                                                                0.958
#
   2
          0
                        -0.885
                                            0.138
                                                                -0.0359
#
   3
          0
                        -0.493
                                            -0.0451
                                                                0.585
#
   4
                                                                0.898
          0
                        -1.02
                                            0.128
#
   5
          0
                        -1.11
                                           -0.237
                                                                0.430
#
   6
          0
                        -1.00
                                           -0.131
                                                                -0.0151
#
   7
          0
                        -1.30
                                           -0.415
                                                                0.255
   8
#
          0
                                                                0.698
                        -0.803
                                            0.190
   9
#
          0
                        -0.743
                                           -0.0497
                                                                -0.0427
# 10
          0
                        -0.596
                                            0.384
                                                                0.822
        with 33,943 more rows
```

Here, individual_trait consists of three trait values per individual and must be splitted into three columns to yield an individual-wise data frame.

We pivot the data again to allow plotting the three traits as different facets, except that now we show a density map of individual traits through time:

```
newnames <- paste0("trait ", 0:2)
data <- pivot_data(data, paste0("individual_trait", 1:3), newnames = newnames)
data <- data %>% rename(trait = "variable")
ggplot(data, aes(x = time, y = value)) +
   geom_bin2d(bins = 20) +
   facet_grid(. ~ trait)
```



The function plot_trait_values can be used to produce such density map too, if provided with the dataset with traits as different columns.

3.5 Locus-wise data

Yet other variables are recorded for every locus at every time point. Use read_genome to read these data in a locus-wise data frame:

```
data <- read_genome(root, "genome_Fst")
data
# # A tibble: 6,000 x 3
# time genome_Fst locus
# <dbl> <dbl> <int>
```

```
0.0000528
#
   2
           0.000304
                           2
         0
#
   3
                           3
         0
#
         0 0.00129
  4
                           4
#
  5
         0
           0
                           5
#
   6
         0
            0
                           6
#
   7
         0 0.0000659
                           7
#
   8
         0 0.00161
                           8
#
  9
         0 0.00188
                           9
            0.000534
# 10
         0
                          10
# # ... with 5,990 more rows
```

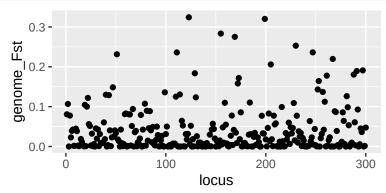
Note that a given set of data may be read in different formats. Here, genome_Fst is read on a locus-wise basis, but we could have read it on a simulation-wise basis, with 300 variables recorded at every time point:

```
read_sim(root, "genome_Fst", by = 300)
# # A tibble: 20 x 301
#
      time genome_Fst1 genome_Fst2 genome_Fst3 genome_Fst4 genome_Fst5 genome_Fst6
#
                 <dbl>
                              <dbl>
                                          <dbl>
                                                       <dbl>
                                                                   <db1>
#
  1
         0
             0.0000528
                           0.000304
                                      0
                                                   0.00129
                                                                0
                                                                            0
#
   2
       100
             0.0604
                           0.0348
                                      0
                                                   0.0000345
                                                                0
                                                                            0
#
   3
       200
             0.0713
                           0.0240
                                      0
                                                   0.00711
                                                                0.00340
                                                                            0
#
       300
             0.105
                           0.0987
                                      0.000275
                                                   0.0218
                                                                0.00165
                                                                            0
  4
#
                           0.146
                                      0.00375
                                                   0.0107
                                                                0.000104
                                                                            0.000266
   5
       400
             0.134
#
   6
       500
             0.185
                           0.123
                                      0
                                                   0.0234
                                                                0.00789
                                                                            0.000724
#
  7
       600
             0.179
                           0.146
                                      0
                                                   0.0128
                                                                0.00426
                                                                            0.000275
#
   8
       700
             0.208
                           0.159
                                      0
                                                   0.0441
                                                                0.000273
                                                                            0.000317
#
  9
       800
             0.180
                           0.167
                                                   0.0343
                                                                0.0166
                                                                            0.0000747
                                      0
# 10
                                                   0.0216
       900
             0.151
                           0.187
                                      0.000268
                                                                0.00756
                                                                            0.00111
# 11
     1000
             0.189
                           0.184
                                      0.000864
                                                   0.0585
                                                                0.0149
                                                                            0.00389
# 12
     1100
             0.140
                           0.142
                                      0.000293
                                                   0.0520
                                                                0.0197
                                                                            0.00591
# 13
      1200
             0.103
                           0.130
                                      0.000924
                                                   0.0219
                                                                0.0504
                                                                            0.0125
# 14
     1300
             0.166
                           0.134
                                      0
                                                   0.0658
                                                                0.0582
                                                                            0.00485
# 15
     1400
             0.128
                           0.171
                                      0.000275
                                                   0.0698
                                                                0.0655
                                                                            0.0111
# 16
     1500
             0.107
                           0.129
                                                                0.0681
                                      0
                                                   0.0954
                                                                            0.0201
# 17
      1600
             0.0541
                           0.0819
                                      0.000523
                                                   0.0898
                                                                0.0590
                                                                            0.0352
# 18
     1700
             0.0577
                           0.125
                                      0
                                                   0.0780
                                                                0.0403
                                                                            0.0228
# 19
     1800
             0.0671
                           0.102
                                      0.00154
                                                   0.0951
                                                                0.0317
                                                                            0.0163
# 20
     1900
             0.0807
                           0.107
                                      0.0000157
                                                   0.0776
                                                                0.0228
                                                                            0.0393
# # ... with 294 more variables: genome_Fst7 <dbl>, genome_Fst8 <dbl>,
# #
      genome_Fst9 <dbl>, genome_Fst10 <dbl>, genome_Fst11 <dbl>,
      genome Fst12 <dbl>, genome Fst13 <dbl>, genome Fst14 <dbl>,
# #
# #
      genome Fst15 <dbl>, genome Fst16 <dbl>, genome Fst17 <dbl>,
# #
      genome_Fst18 <dbl>, genome_Fst19 <dbl>, genome_Fst20 <dbl>,
# #
      genome_Fst21 <dbl>, genome_Fst22 <dbl>, genome_Fst23 <dbl>,
# #
      qenome_Fst24 <dbl>, qenome_Fst25 <dbl>, qenome_Fst26 <dbl>,
# #
      genome_Fst27 <dbl>, genome_Fst28 <dbl>, genome_Fst29 <dbl>,
# #
      genome_Fst30 <dbl>, genome_Fst31 <dbl>, genome_Fst32 <dbl>,
# #
      qenome_Fst33 <dbl>, qenome_Fst34 <dbl>, qenome_Fst35 <dbl>,
# #
      genome_Fst36 <dbl>, genome_Fst37 <dbl>, genome_Fst38 <dbl>,
# #
      qenome_Fst39 <dbl>, qenome_Fst40 <dbl>, qenome_Fst41 <dbl>,
# #
      genome_Fst42 <dbl>, genome_Fst43 <dbl>, genome_Fst44 <dbl>,
# #
      qenome_Fst45 <dbl>, qenome_Fst46 <dbl>, qenome_Fst47 <dbl>,
# #
      genome_Fst48 <dbl>, genome_Fst49 <dbl>, genome_Fst50 <dbl>,
```

```
# #
      genome_Fst51 <dbl>, genome_Fst52 <dbl>, genome_Fst53 <dbl>,
# #
      genome_Fst54 <dbl>, genome_Fst55 <dbl>, genome_Fst56 <dbl>,
# #
      genome Fst57 <dbl>, genome Fst58 <dbl>, genome Fst59 <dbl>,
      genome_Fst60 <dbl>, genome_Fst61 <dbl>, genome_Fst62 <dbl>,
# #
# #
      genome Fst63 <dbl>, genome Fst64 <dbl>, genome Fst65 <dbl>,
      genome_Fst66 <dbl>, genome_Fst67 <dbl>, genome_Fst68 <dbl>,
# #
# #
      genome_Fst69 <dbl>, genome_Fst70 <dbl>, genome_Fst71 <dbl>,
# #
      genome_Fst72 <dbl>, genome_Fst73 <dbl>, genome_Fst74 <dbl>,
# #
      genome Fst75 <dbl>, genome Fst76 <dbl>, genome Fst77 <dbl>,
# #
      qenome_Fst78 <dbl>, qenome_Fst79 <dbl>, qenome_Fst80 <dbl>,
# #
      genome_Fst81 <dbl>, genome_Fst82 <dbl>, genome_Fst83 <dbl>,
# #
      qenome_Fst84 <dbl>, qenome_Fst85 <dbl>, qenome_Fst86 <dbl>,
# #
      genome_Fst87 <dbl>, genome_Fst88 <dbl>, genome_Fst89 <dbl>,
# #
      qenome_Fst90 <dbl>, qenome_Fst91 <dbl>, qenome_Fst92 <dbl>,
# #
      qenome_Fst93 <dbl>, qenome_Fst94 <dbl>, qenome_Fst95 <dbl>,
# #
      qenome_Fst96 <dbl>, qenome_Fst97 <dbl>, qenome_Fst98 <dbl>,
# #
      qenome_Fst99 <dbl>, qenome_Fst100 <dbl>, qenome_Fst101 <dbl>,
# #
      genome_Fst102 <dbl>, genome_Fst103 <dbl>, genome_Fst104 <dbl>,
      genome_Fst105 <dbl>, genome_Fst106 <dbl>, ...
# #
```

From our long, locus-wise data frame we can plot the Fst scan along the genome at the last generation with some help from dplyr's filter function:

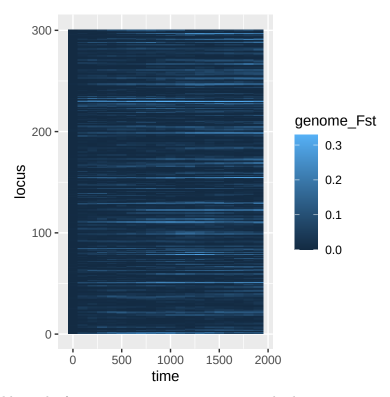
```
data <- data %>% filter(time == last(time))
ggplot(data, aes(x = locus, y = genome_Fst)) +
  geom_point()
```



For ready-made genome scan plots the function plot_genome_scan can be used. This function expects a dataset on a per-locus basis, and has multiple options such as displaying the timepoint, showing bars or lollipops instead of points, or even showing an animation through time. Also note that the function plot_genome_scatter is an extension of plot_genome_scan where the x-axis does not necessarily correspond to locus positions, but can be any locus-specific variable, thus resulting in a 2D scatter plot.

We could also plot Fst through time using a heatmap across the genome:

```
data <- read_genome(root, "genome_Fst")
ggplot(data, aes(x = time, y = locus, fill = genome_Fst)) +
    geom_tile()</pre>
```



Alternatively we could use the function plot_genome_heatmap to do the same.

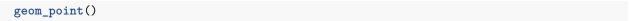
3.6 Reading the genetic architecture

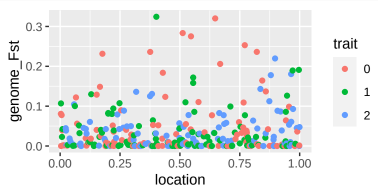
Locus-wise data are best analyzed in the light of the underlying genetic architecture of the loci, which can be read from the corresponding architecture file. Use the architecture argument of read_genome to append locus-wise architecture information to the data:

```
data <- read_genome(root, "genome_Fst", architecture = TRUE)</pre>
# Joining, by = c("time", "locus")
data
# # A tibble: 6,000 x 9
      time genome_Fst locus location trait effect dominance chromosome degree
#
                 <dbl> <int>
                                 <dbl> <fct>
                                               <db1>
                                                           <dbl>
                                                                       \langle int \rangle
                                                                              <db1>
#
   1
         0 0.0000528
                           1 0.00309 0
                                              -0.0793
                                                         0.0245
                                                                           1
                                                                                 19
#
   2
         0
            0.000304
                           2
                              0.00339 1
                                               0.103
                                                         0.0223
                                                                           1
                                                                                  14
#
   3
         0 0
                            3
                              0.00404 0
                                               0.0940
                                                         0.0240
                                                                           1
                                                                                 15
#
         0
            0.00129
                               0.00622 0
                                              -0.0632
                                                         0.0168
                                                                           1
   4
                           4
                                                                                 12
#
                                              -0.0933
   5
         0
            0
                            5
                               0.00758 1
                                                         0.143
                                                                           1
                                                                                 11
#
   6
         0
            0
                            6
                               0.00867 0
                                               0.0703
                                                         0.00695
                                                                           1
                                                                                 13
#
   7
         0
            0.0000659
                            7
                               0.00968 1
                                               0.0778
                                                         0.0379
                                                                           1
                                                                                  5
#
   8
         0
            0.00161
                            8
                               0.0121 0
                                              -0.107
                                                         0.0900
                                                                           1
                                                                                 14
   9
#
         0
            0.00188
                           9
                                              -0.0734
                                                                           1
                               0.0159
                                       0
                                                         0.105
                                                                                 18
# 10
         0 0.000534
                          10
                              0.0233 2
                                              -0.0219
                                                         0.0370
                                                                           1
                                                                                 16
# # ... with 5,990 more rows
```

Now that we have more information about each locus, we can refine our plot, for example by adding colors based on the encoded trait of each locus:

```
data <- data %>% filter(time == last(time))
ggplot(data, aes(x = location, y = genome_Fst, color = trait)) +
```



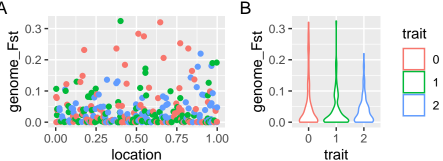


3.7 Combining plots

It may be handy to pool different plots into the same figure (as in, not different facets from the same plot). We can use the patchwork package to do this (no new egssimtools function here):

```
data <- read_genome(root, "genome_Fst", architecture = TRUE)
# Joining, by = c("time", "locus")
data <- data %>% filter(time == last(time))
p1 <- ggplot(data, aes(x = location, y = genome_Fst, color = trait)) +
    geom_point() +
    theme(legend.position = "none")
p2 <- ggplot(data, aes(x = trait, y = genome_Fst, color = trait)) +
    geom_violin()

# From patchwork
wrap_plots(p1, p2, widths = c(2, 1)) + plot_annotation(tag_levels = "A")</pre>
```

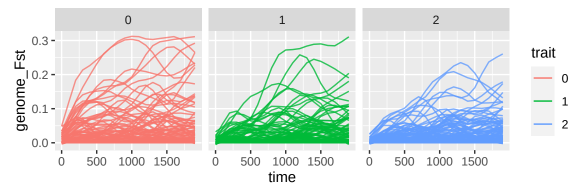


3.8 More complex plots

We can use the full breadths of tricks from ggplot2 to make our plots look their best. For example, it is difficult to plot many lines on the same plot (without a lot of copy-and-paste), yet it is often needed in simulation studies. One way to go around that is to plot each line with a different transparency, using the alpha aesthetics. One can then constrain the possible alpha values to a very narrow range so differences in transparency between the lines are not noticeable. Note that this is what the gglineplot function from the ggsim package does, but here we will stick to base ggplot2.

Let us apply this trick to plotting Fst through time on a per-locus basis. We also use the smoothen_data (from egssimtools) to smoothen our Fst curves prior to plotting.

```
data <- read_genome(root, "genome_Fst", architecture = TRUE)
# Joining, by = c("time", "locus")
data <- smoothen_data(data, "time", "genome_Fst", span = 0.3, line = "locus")
ggplot(data, aes(x = time, y = genome_Fst, alpha = factor(locus), color = trait)) +
    geom_line() +
    facet_grid(. ~ trait) +
    scale_alpha_manual(values = runif(length(unique(data$locus)), 0.79, 0.81)) +
    guides(alpha = FALSE)</pre>
```



Trajectories of multipe loci through time can also be plotted using the plot_genome_trajectories function.

3.9 Edge-wise data

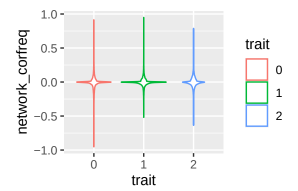
Our simulation implements a genotype-phenotype map involving a gene regulatory network, and some variables are specific to edges in the network, such as the correlation in allele frequencies between interacting loci. To read them into an edge-wise data frame, use read_network:

```
data <- read_network(root, "network_corfreq", architecture = TRUE)</pre>
# Joining, by = c("time", "edge")
data
# # A tibble: 60,000 x 7
#
      time network_corfreq
                             edge
                                    from
                                             to
                                                    weight trait
#
     <db1>
                      <dbl> <int> <int> <int>
                                                     <dbl> <fct>
#
   1
         0
                     0.0528
                                 1
                                       62
                                            247 -0.0722
#
   2
         0
                                 2
                                       62
                                            161 -0.00219
                     0.0412
                                                           0
   3
                                 3
#
         0
                    -0.0350
                                      247
                                            161
                                                  0.0449
#
         0
                     0.0725
                                 4
                                      247
                                            230
                                                  0.00321
                                                           0
#
   5
         0
                     0.0648
                                 5
                                      161
                                            230
                                                  0.00181
#
   6
         0
                    -0.0229
                                 6
                                            230
                                                  0.000663 0
                                       62
   7
#
         0
                    -0.0293
                                 7
                                       62
                                            118
                                                  0.0644
#
   8
         0
                    -0.0296
                                 8
                                      230
                                            118 -0.00207
                                                           0
#
   9
         0
                    -0.0348
                                 9
                                      247
                                            118
                                                  0.0124
                                                            0
# 10
         0
                    -0.0445
                                10
                                      161
                                            118
                                                  0.0118
# # ... with 59,990 more rows
```

Here, the architecture argument tells the function to read edge-wise parameters from the genetic architecture, such as the indices of the partner genes or the interaction weights, and append them to the data.

From this dataset, we can for example plot distributions of edge-wise variables across traits,

```
ggplot(data, aes(x = trait, y = network_corfreq, color = trait)) +
  geom_violin()
```



3.10 Reading data: what happens backstage?

So far we have used the functions read_sim, read_pop, read_genome and read_network to read the data. These functions are all wrappers around one same function, read_data. The difference between them is in the preformatting of the arguments to be passed to read_data in order to produce a simulation-wise, individual-wise, locus-wise or edge-wise data frame, respectively. But what happens within read_data?

read_data is a flexible function that can be used to read data into a variety of formats. It can be used instead of its simplified, preformatted counterparts, but this will typically result in longer snippets of code and should be reserved to cases when the desired output cannot be achieved using the simplified versions. For example, the following code:

```
read_data(
 root,
  c("time", "genome_Fst", "genome_Cst"),
  by = c(1, 1, 1),
  dupl = c(300, 1, 1)
)
#
  # A tibble: 6,000 x 3
#
      time genome_Fst genome_Cst
#
     <db1>
                 <db1>
                             <db1>
#
   1
         0
            0.0000528 0.0149
   2
#
            0.000304
                        0.000567
         0
   3
#
         0
            0
                        0.00277
#
   4
         0
            0.00129
                        0.00562
#
   5
         0
            0
                        0.00278
#
   6
         0
            0
                        0.00503
#
   7
         0
            0.0000659
                        0.00155
            0.00161
                        0.00163
         0
   9
#
         0
            0.00188
                        0.0488
# 10
         0 0.000534
                        0.0000189
       with 5,990 more rows
```

is equivalent to

```
read_genome(root, c("genome_Fst", "genome_Cst"))
# # A tibble: 6,000 x 4
#
      time genome_Fst genome_Cst locus
#
     <db1>
                <db1>
                           <dbl> <int>
           0.0000528
  1
         0
                      0.0149
                                      1
  2
         0
                                      2
           0.000304
                       0.000567
         0 0
                       0.00277
```

```
#
            0.00129
                        0.00562
   4
#
   5
         0
            0
                         0.00278
                                        5
#
   6
         0
            0
                         0.00503
                                        6
#
   7
                                        7
         0
            0.0000659
                        0.00155
#
   8
         0
            0.00161
                         0.00163
                                        8
#
  9
         0
            0.00188
                         0.0488
                                        9
# 10
         0 0.000534
                         0.0000189
                                       10
# # ... with 5,990 more rows
```

In read_data, the file time.dat is not read in by default and must be explicitly provided. This means that the by argument needs to account for the presence of time in the variables argument, i.e., by must be provided for all the variables (unless all of them take a value of one, in this case leave unspecified). The dupl argument is being taken care of for you in the simplified versions of read_data, while it must be provided here. This argument determines how many times each variable should be duplicated (with copies stacked on top of each other). This matters when, even after splitting some variables into multiple columns, some are still much shorter than the rest. Here, for example, we need to duplicate the time column 300 times to get a viable data frame, because we have 300 loci recorded for each time point in genome Fst and genome Cst.

Sometimes, different values of a too short column may need to be duplicated different numbers of times. This is the case for time in individual-wise data, for example, because the number of individuals changes from one generation to the next, so each time point has to be duplicated by the number of individuals in that specific time point. For this reason, dupl accepts as some of its values the name of a variable in which to look up the number of times the corresponding column should be duplicated. For example,

```
read data(
  root,
  c("time", "individual trait", "individual ecotype"),
  by = c(1, 3, 1),
  dupl = list("population_size", 1, 1)
)
#
  # A tibble: 33,953 x 5
#
      time individual_trait1 individual_trait2 individual_trait3 individual_ecoty~
#
     <db1>
                         <dbl>
                                             <db1>
                                                                 <db1>
                                                                                    <db1>
#
   1
         0
                        -1.20
                                            0.0673
                                                               0.958
                                                                                         0
#
   2
         0
                        -0.885
                                           0.138
                                                               -0.0359
                                                                                         1
   3
#
         0
                        -0.493
                                                               0.585
                                                                                         1
                                           -0.0451
#
   4
         0
                        -1.02
                                            0.128
                                                               0.898
                                                                                         0
#
                                                                                         0
   5
         0
                        -1.11
                                           -0.237
                                                               0.430
#
   6
         0
                        -1.00
                                           -0.131
                                                               -0.0151
                                                                                         0
#
   7
         0
                        -1.30
                                                               0.255
                                                                                         0
                                           -0.415
#
   8
         0
                        -0.803
                                           0.190
                                                               0.698
                                                                                         1
#
   9
         0
                        -0.743
                                           -0.0497
                                                               -0.0427
                                                                                         1
# 10
                        -0.596
                                                                                         1
         0
                                            0.384
                                                               0.822
# # ... with 33,943 more rows
```

looks up in population_size.dat the number of times each time point in the time column should be duplicated. Note the use of a list instead of a vector here, because "population_size" is not a number. Also note that here we splitted individual_trait into three columns but individual_ecotype remained a single column.

The above snippet is equivalent to:

```
read_pop(root, paste0("individual_", c("trait", "ecotype")), by = c(3, 1))
# # A tibble: 33,953 x 5
# time individual_trait1 individual_trait2 individual_trait3 individual_ecoty~
# <dbl> <dbl> <dbl> <dbl>
```

#	1	0	-1.20	0.0673	0.958	0
#	2	0	-0.885	0.138	-0.0359	1
#	3	0	-0.493	-0.0451	0.585	1
#	4	0	-1.02	0.128	0.898	0
#	5	0	-1.11	-0.237	0.430	0
#	6	0	-1.00	-0.131	-0.0151	0
#	7	0	-1.30	-0.415	0.255	0
#	8	0	-0.803	0.190	0.698	1
#	9	0	-0.743	-0.0497	-0.0427	1
#	10	0	-0.596	0.384	0.822	1
# # with 33,943 more rows						

3.11 Plot a gene network

We previously saw how to read edge-wise data, but the data frame resulting from read_network is not enough in order to plot the actual gene network. Networks are graphs, and graphs are complicated plots to make because they do not fit into a "tidy" data frame representation, where one of the criteria for tidyness (sensu tidyverse) means that the data should have one unit of observation. This is because a graph typically has two equally valid units of observation: nodes and edges, both of which are needed to produce a plot. The packages tidygraph and ggraph solved this problem by introducing the tbl_graph object. This object contains a representation of two tibbles (data frames): one for nodes, the other one for edges. This sort-of-tidy object can readily be interpreted by graph-plotting functions from ggraph.

In egssimtools, the function read_arch_network reads the content of a genetic architecture file and returns a tbl_graph (unless its as_list argument is TRUE, which is the case e.g. when it is called from within read_network) containing locus-wise as well as edge-wise data.

So, to plot a gene regulatory network, we first read the genetic architecture as a tbl_graph:

```
arch <- read_arch_network(root)</pre>
arch
# # A tbl_graph: 300 nodes and 3000 edges
# #
# # A directed acyclic simple graph with 3 components
# #
# # Node Data: 300 x 7 (active)
    locus location trait effect dominance chromosome degree
#
#
    \langle int \rangle
              <dbl> <fct>
                              <db1>
                                         <db1>
                                                     \langle int \rangle
# 1
        1 0.00309 0
                            -0.0793
                                       0.0245
                                                         1
                                                                19
# 2
        2
            0.00339 1
                             0.103
                                       0.0223
                                                          1
                                                                14
# 3
        3
            0.00404 0
                             0.0940
                                                          1
                                                                15
                                       0.0240
# 4
            0.00622 0
                            -0.0632
                                       0.0168
        4
                                                         1
                                                                12
# 5
        5
           0.00758 1
                            -0.0933
                                       0.143
                                                         1
                                                                11
# 6
           0.00867 0
                             0.0703
                                      0.00695
                                                          1
                                                                13
# # ... with 294 more rows
# #
# # Edge Data: 3,000 x 5
     from
#
              to
                   weight trait
                                   edge
#
    <int> <int>
                      <dbl> < fct> < int> 
# 1
       62
             247 -0.0722 0
                                       1
                                       2
# 2
       62
             161 -0.00219 0
                  0.0449 0
                                       3
# 3
      247
             161
# # ... with 2,997 more rows
```

This is already enough for plotting. But let us say that we want to map some locus-specific data onto the nodes of the network, for example, the locus-specific Fst at the final generation. To do that we must read these locus-specific data:

```
data_n <- read_genome(root, "genome_Fst", architecture = TRUE)
data_n <- data_n %>% filter(time == last(time))
```

and then attach them to the tbl_graph:

```
arch <- arch %>% activate(nodes) %>% right_join(data_n)
arch
# # A tbl_graph: 300 nodes and 3000 edges
# #
# # A directed acyclic simple graph with 3 components
# #
# # Node Data: 300 x 9 (active)
#
    locus location trait effect dominance chromosome degree time genome_Fst
#
    \langle int \rangle
           <\!db\,l> <\!fc\,t> <\!db\,l> <\!db\,l>
                                                 <int> <dbl> <dbl>
# 1
      1 0.00309 0
                         -0.0793
                                   0.0245
                                                     1
                                                           19 1900 0.0807
# 2
       2 0.00339 1
                          0.103
                                   0.0223
                                                     1
                                                           14 1900 0.107
# 3
       3 0.00404 0
                          0.0940 0.0240
                                                     1
                                                           15
                                                               1900 0.0000157
# 4
       4 0.00622 0
                         -0.0632 0.0168
                                                     1
                                                           12 1900 0.0776
# 5
       5 0.00758 1
                         -0.0933 0.143
                                                           11 1900 0.0228
                                                     1
       6 0.00867 0
# 6
                          0.0703 0.00695
                                                     1
                                                           13 1900 0.0393
# # ... with 294 more rows
# #
# # Edge Data: 3,000 x 5
            to weight trait edge
#
     from
    \langle int \rangle \langle int \rangle
                   < dbl > < fct > < int > 
# 1
            247 -0.0722 0
       62
                                   1
# 2
       62
          161 -0.00219 0
                                    2
      247 161 0.0449 0
                                    3
# # ... with 2,997 more rows
```

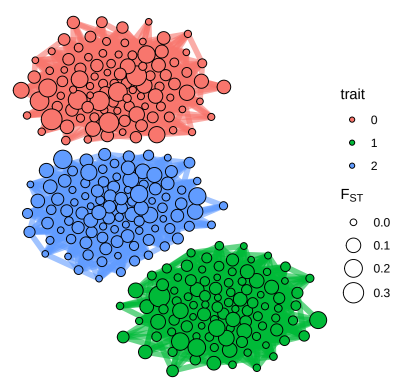
where right_join attaches our data_n tibble to the "top-facing" tibble in the arch tbl_graph, which we set to nodes using the activate function from tidygraph. The alternative activation of nodes and edges allows tbl_graph objects to be treated as a single tibble (the activated one) by other functions, which makes their incorporation smooth inside tidyverse pipelines.

We could also map edge-specific variables onto the network, following the same logic:

```
data_e <- read_network(root, "network_corfreq", architecture = TRUE)
data_e <- data_e %>% filter(time == last(time))
arch <- arch %>% activate(edges) %>% right_join(data_e)
```

We can now plot the network using the ggraph package, which produces a ggplot object:

```
# This may take a while
ggraph(arch, layout = "graphopt", charge = 0.1, mass = 30, niter = 100000) +
geom_edge_link(aes(color = trait), width = 2, alpha = 0.6) +
geom_node_point(aes(fill = trait, size = genome_Fst), shape = 21) +
scale_size_continuous(range = c(2, 7)) +
scale_alpha(range = c(0.6, 1)) +
labs(size = parse(text = "F[ST]"), fill = "trait") +
theme_void() +
guides(edge_color = FALSE)
```



Note that the function plot_genome_network produces a similar output, with multiple options, including animations!

Of course, one can play with the graphical parameters as needed. Visit the ggraph website for more details on how to use its functions!

3.12 Combining simulations

So far we have worked with single simulations. Some use cases may require to work with multiple simulations, however, such as looking at the effect of some parameter value. To combine data from multiple simulations, we use the combine_data function, which is essentially a wrapper around the read_* functions we previously used for single simulations:

```
# First we reset the working directory to where multiple simulations are
root <- "../data"
data <- combine_data(</pre>
 root, pattern = "example", level = 1, type = "genome",
  variables = "genome_Fst", architecture = TRUE,
  parnames = c("ecosel", "hsymmetry")
)
data
# # A tibble: 18,000 x 12
#
            time genome_Fst hsymmetry ecosel locus location trait
                                                                     effect
     <chr> <dbl>
                      <dbl> <chr>
                                                                      <db1>
#
  1 1
               0 0.0000528 0
                                       1
                                                  1 0.00309 0
                                                                    -0.0793
  2 1
               0
                  0.000304
                            0
                                       1
                                                  2 0.00339 1
                                                                     0.103
  3 1
               0
                  0
                            0
                                                  3 0.00404 0
                                       1
                                                                     0.0940
  4 1
               0
                  0.00129
                            0
                                       1
                                                  4
                                                     0.00622 0
                                                                    -0.0632
#
  5 1
               0
                  0
                            0
                                       1
                                                  5
                                                     0.00758 1
                                                                    -0.0933
  6 1
                                                     0.00867 0
                                                                     0.0703
```

```
0.0000659 0
                                                     0.00968 1
#
  8 1
                 0.00161
                                       1
                                                     0.0121
                                                                    -0.107
                            0
                                                  8
  9 1
                  0.00188
                            0
                                       1
                                                  9
                                                     0.0159
                                                                    -0.0734
                                                             0
# 10 1
               0
                 0.000534 0
                                       1
                                                 10 0.0233 2
                                                                    -0.0219
# # ... with 17,990 more rows, and 3 more variables: dominance <dbl>,
      chromosome <int>, degree <dbl>
```

Here, we are combining the results of three simulations called example_1, example_2 and example_3 into one data frame. To do this, we have to give the function:

- a place to look for the simulations, here root
- the level of recursion at which the function has to search, here one means that the simulation folders are *directly* within root, two would mean than they are in one or multiple folders within root, and so on. A value of zero would mean that root is actually a vector of simulation folders (in our case it would be c(".../data/example_1", ".../data/example_2", ".../data/example3"))
- a pattern to match to identify simulation folder names (if level is nonzero)

The function internally calls fetch_dirs to search for simulation folders, which comes in handy when the folder containing the simulation has some substructure (e.g. multiple batches of simulations stored in different subfolders).

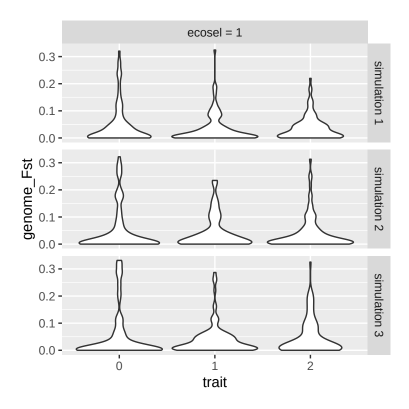
In addition, the type argument tells combine_data what version of read_data to use to read the multiple simulations. Here, we are internally calling read_genome because we are reading locus-wise Fst values from multiple simulations, and we attach to them their respective architecture data.

All other parameters passed to combine_data are passed to the read_* function you have chosen, so make sure those parameters are relevant by checking the documentation of the different functions. For example, it would not make sense to use architecture = TRUE if type is pop, because read_pop does not have an architecture argument.

Among the arguments passed to the read_* function are arguments regarding the reading of simulation parameters, which are read from a parameter file. Here, for example, we read the ecosel and the hsymmetry parameter values for each simulation. The function being called internally for that is read_param. All read_* functions can take arguments to pass on to read_param. We did not use them before because each simulation is generated from a single set of parameter values, but parameters start to matter when we combine data from multiple simulations together.

Once we have a data frame summing up multiple simulations, we can manipulate it and plot it just like the tibbles we get from single simulations:

```
data <- data %>% filter(time == last(time))
data <- data %>% mutate(sim = str_replace(sim, "^", "simulation "))
data <- data %>% mutate(ecosel = str_replace(ecosel, "^", "ecosel = "))
ggplot(data, aes(x = trait, y = genome_Fst)) +
    geom_violin() +
    facet_grid(sim ~ ecosel)
```



3.13 The split-apply-combine routine

We may want to produce a different plot for different subsets of a data frame, for example, for different simulations. This can be done using the split-apply-combine routine from the tidyverse. We first read the data from multiple simulations:

```
data <- combine_data(
  root, pattern = "example", level = 1, type = "genome",
  variables = "genome_Fst", architecture = TRUE
)</pre>
```

Then, we nest our tibble by simulation using some tidyverse functions:

```
data <- data %>%
  group_by(sim) %>%
  nest()
data
# # A tibble: 3 x 2
# # Groups:
               sim [3]
    sim
          data
#
    <chr> <chr> ist>
#
          <tibble [6,000 x 9]>
 1 1
# 2 2
          <tibble [6,000 x 9]>
          <tibble [6,000 x 9]>
# 3 3
```

This returns a nested tibble, where the data have been split between the different simulations and stored in a new list-column called data. The data has not disappeared, instead it is now located in a list of tibbles, with one tibble per simulation. Looping through this list-column will allow us to produce one plot per simulation. But before that, we must define a function to apply to each simulation, that will do the plotting. This function must take a tibble as argument, because it will be called on each tibble within the data list-column.

One example is:

```
ggplot(
  data,
  aes(x = time, y = genome_Fst, color = trait, alpha = factor(locus))
) +
  geom_line() +
  facet_grid(. ~ trait) +
  guides(alpha = FALSE)
}
```

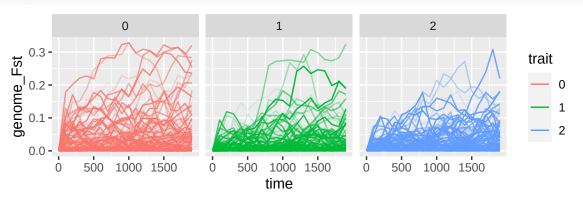
This function will plot Fst values through time for each locus, facetted by trait. But in practice, any plotting routine that can work with a tibble containing data for one simulation will do.

We now use the map function from the purrrr package to iterate through the data list-column within the data tibble, and apply our custom plot_this function to each of its tibbles:

This has added a new column called fig to our tibble. fig is also a list-column, but instead of being a list of tibbles, it is a list of ggplot objects returned by map (map always returns a list), one for each simulation.

We check out the plots individually:

data\$fig[[1]]



But we can also perform other operations, such as saving them separately in their own respective image files, using:

```
data <- data %>% mutate(figname = sprintf("sim%s.png", sim))
data
# # A tibble: 3 x 4
# # Groups: sim [3]
# sim data fig figname
# <chr> tist> <chr>
```

Here, we have created a new column called figname, which contains the names of the PNG files where each plot should be saved. We then used the walk2 function from purrr to save each plot in fig to its respective file figname. Check out the purrr documentation for more details on how map, walk and their extensions work.

3.14 Looping through simulation folders

So far we have dealt with a situation where the data from multiple simulations fits into a single data frame. However, we may want to save the same kind of figure (e.g. a genome scan of Fst values through time) for thousands of replicate simulations, which may be too large to store in a single data frame using combine_data. For such cases, it may be better to loop through the different simulation folders, read the data for each of them using a read_* function, plot it and store the output plot in a list for later manipulation or save it directly from within the loop (using e.g. ggsave).

We would also run into this problem if we wanted to plot a gene network for multiple simulations, because the data for a single network does not fit within a standard tibble, and therefore it may be very clunky to try to combine the data from multiple networks into a single object.

We can get a vector of simulation folders using the fetch_dirs function:

```
roots <- fetch_dirs("../data", pattern = "example_", level = 1)
roots
# [1] "../data/example_1" "../data/example_2" "../data/example_3"</pre>
```

Note that the find_extant function can be used on this vector of simulation folder names to find out and retain the ones that did not go extinct or did not crash.

To loop through these folders and plot the data, we can e.g. refine our plot_this function so it operates at the level of the simulation folder, and not on a tibble. This means that plot_this would have to read the data too. For example, we could have:

```
plot_this <- function(root) {

# This is the only line added
data <- read_genome(root, "genome_Fst", architecture = TRUE)

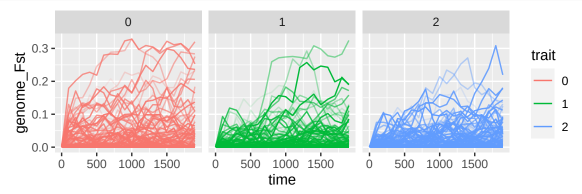
ggplot(
   data,
   aes(x = time, y = genome_Fst, color = trait, alpha = factor(locus))
) +
   geom_line() +
   facet_grid(. ~ trait) +
   guides(alpha = FALSE)
}</pre>
```

which we then would use within a loop through the simulation folders:

```
plots <- map(roots, plot_this)</pre>
```

This has produced a list of ggplot objects:

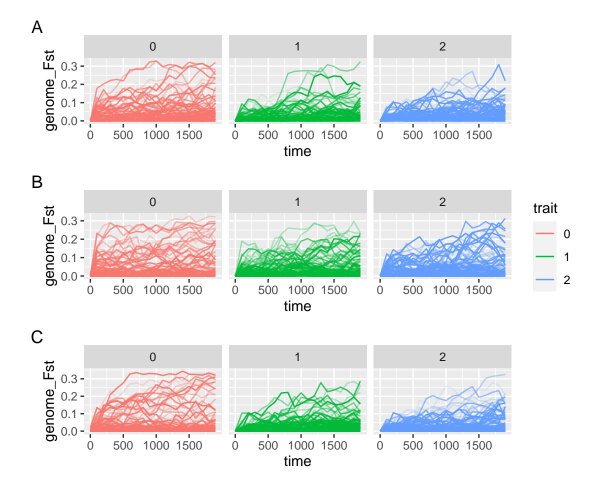
plots[[1]]



which we can then manipulate as needed, for example, by combining them into a single figure using patchwork:

```
# We remove the legend of all plots but one
for (i in 2:length(plots)) {
   plots[[i]] <- plots[[i]] + theme(legend.position = "none")
}

# Then we assemble the plots with a common legend
wrap_plots(plots) +
   plot_layout(guides = 'collect', nrow = length(plots)) +
   plot_annotation(tag_levels = 'A')</pre>
```



4 To sum up

The complexity and diversity of the data saved by the ExplicitGenomeSpeciation simulation require an analysis toolkit that is both flexible and very standardized. The tools of the tidyverse allow for exactly that purpose with the R computing environment. As often in R, the same result can be obtained in multiple ways. For these reasons we strongly recommend its use when analyzing the data from our simulaitons, but of course other computing languages can be used (e.g. Python, MATLAB or C++), as long as they can read the binary files saved by EGS.