Simple workflow example for a population model

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In this vignette we present a simple example of the *poems* workflow using a fictitious population model. The purpose of this example is to demonstrate how the components of the package are used to build an ensemble of viable models that best match known or desired patterns. Although the package is designed to facilitate building complex models and running multitudes of sample simulations, the scale and complexity of this demonstration model is deliberately minimal so as to easily examine the outputs at every stage of the workflow.

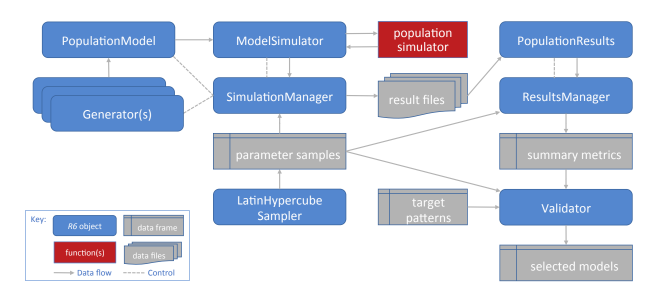
Setup

We begin by loading the *poems* package and setting our output directory.

```
library(poems)
OUTPUT_DIR <- tempdir()</pre>
```

Workflow

The *poems* workflow, which implements a pattern-oriented modeling (POM) approach (Grimm et al., 2005), is achieved via a framework of interoperable components:



The workflow is summarized by the following six steps:

- 1. Build the population model for the study region.
- 2. Build generators for dynamically generating model parameters.
- 3. Sample model and generator parameters for each simulation.

- 4. Build a simulation manager to run each simulation.
- 5. Build a results manager to generate summary results (metrics).
- 6. Build a validator to select a model ensemble.

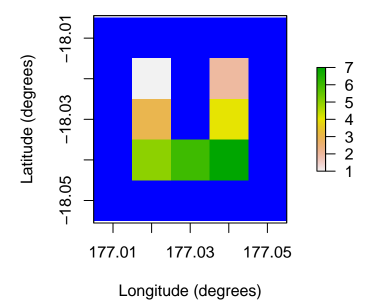
Step 1: Build the population model for the study region

Create a model template using the *PopulationModel* class. If the model is spatially explicit, then define the study region via the *Region* class. All fixed model inputs, such as environmental correlations, as well as any user-defined functions for processes such as harvesting, should be set at this stage.

We could create these components in any order and set model parameters separately, but let's setup our study region, generate environmental correlations, and define a harvest function prior to initializing the template model with all the fixed parameters.

Study region First, we'll define our study region (denoted U Island) with some longitude-latitude coordinates.

Example region (cell indices)



Environmental correlation Next, we'll define a distance-based spatial correlation for applying environmental stochasticity within our model. The generated correlation data is compacted for computational efficiency (with large-scale models).

```
# Distance-based environmental correlation (via a compacted Cholesky decomposition)
env_corr <- SpatialCorrelation$new(region = region, amplitude = 0.4, breadth = 500)
correlation <- env_corr$get_compact_decomposition(decimals = 2)</pre>
correlation # examine
#> $matrix
     [,1] [,2] [,3] [,4] [,5] [,6] [,7]
#> [1,] 1 0.01 0.04 0.04 0.04 0.02 0.04
#> [2,] 0 1.00 1.00 0.01 1.00 0.02 0.01
#> [3,] 0 0.00 0.00 1.00 0.00 0.05 0.05
#> [4,] 0 0.00 0.00 0.00 0.00 1.00 1.00
#>
#> $map
     [,1] [,2] [,3] [,4] [,5] [,6] [,7]
#> [1,] 1 1 1 2 3 3 4
            2 3 3 5
#> [2,] NA
                                 4
                                     5
#> [3,] NA NA NA 4 NA
                                5
                                     6
#> [4,] NA NA NA NA
                           NA
```

Harvest function Let's now define a simple harvest function, which is optionally list-nested with a harvest rate parameter. We'll also define an alias to the harvest rate so we can sample this parameter later.

Template model Finally, we can build our template model with these and other fixed parameters.

```
# Population (simulation) model template for fixed parameters
stage_matrix <- matrix(c(0, 2.5, # Leslie/Lefkovitch matrix</pre>
                         0.8, 0.5), nrow = 2, ncol = 2, byrow = TRUE,
                       dimnames = list(c("juv", "adult"), c("juv", "adult")))
stage_matrix # examine
#> juv adult
#> juv 0.0 2.5
#> adult 0.8 0.5
model_template <- PopulationModel$new(region = region,</pre>
                                      time steps = 10, # years
                                      populations = region$region_cells, # 7
                                      stages = 2,
                                      stage_matrix = stage_matrix,
                                      demographic_stochasticity = TRUE,
                                      standard deviation = 0.05,
                                      correlation = correlation,
                                      density_dependence = "logistic",
                                      harvest = harvest,
                                      results_selection = c("abundance", "harvested"),
                                      attribute_aliases = harvest_rate_alias)
```

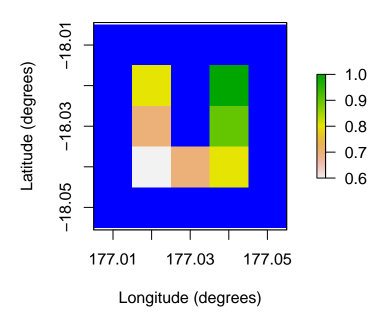
Step 2: Build generators for dynamically generating model parameters

Some model parameters are single values, whilst others are represented as arrays and other multi-value data structures. Usually, we don't wish to sample each individual value within these multi-value parameters (in step 3), but to

generate them dynamically via one or more intermediate sampled parameters. Here we build generators for model initial abundance, carrying capacity, and dispersal.

Habitat suitability Firstly, our initial abundance and carrying capacity generator utilizes an example (mock) habitat suitability for our defined study region.

Example habitat suitability



Initial abundance and carrying capacity generator The generator utilizes generic template functionality for user-defined custom functions. It uses sampled input parameters (initial total abundance and maximum cell density), along with habitat suitability, to generate the desired output model parameters (initial abundance and carrying capacity) via these user-defined functions. Generators can also be configured to read values from files or generate values via probabilistic distributions.

Dispersal generator Our dispersal generator uses default functionality for generating dispersal rates between cells. Its sampled inputs parameterize the distance-based dispersal function (proportion dispersing and breadth of dispersal). The generator can be configured with a dispersal friction helper class object, which calculates equivalent dispersal distances for frictional landscapes and coastlines. Here it is used to ensure dispersal is not performed directly across the "water" in our example U-shaped island. The generated dispersal data is calculated via precalculated distance data and compacted for computational efficiency (with large-scale models).

```
# Distance-based dispersal generator
dispersal gen <- DispersalGenerator$new(region = region,
                                        dispersal max distance = 3000, # in m
                                        dispersal friction = DispersalFriction$new(),
                                        inputs = c("dispersal_p", "dispersal_b"),
                                        decimals = 5)
dispersal_gen$calculate_distance_data() # pre-calculate
test_dispersal <- dispersal_gen$generate(input_values = list(dispersal_p = 0.5,
                                                              dispersal_b = 700))
head(test_dispersal$dispersal_data[[1]])
     target_pop source_pop emigrant_row immigrant_row dispersal_rate
#> 1
              3
                       1
                                      1
                                                     1
                                                              0.10284
              5
                                      2
#> 2
                         1
                                                     1
                                                              0.02115
#> 3
              6
                                      3
                         1
                                                     1
                                                              0.01501
                         2
              4
                                       1
                                                     1
                                                              0.10284
#> 5
              6
                         2
                                       2
                                                     2
                                                              0.01501
#> 6
              7
                         2
                                       3
                                                              0.02115
```

Note that there is no dispersal rate between cells 1 and 2 as there is "water" between those cells, and consequently dispersal between those cells must travel around the U-shaped island, which can't be achieved in one simulation time-step since the "round" distance between those cells is greater than 3000 m.

Step 3: Sample model and generator parameters for each simulation

In order to explore the model parameter space to find the best models, we generate Latin hypercube samples of model and generator parameters to be simulated, using the *LatinHypercubeSampler* class. This class has functionality for generating sample parameters via Uniform, Normal, Lognormal, Beta, and Triangular distributions. For our example we only generate 12 samples. We encourage the user to generate hundreds, or thousands, of samples.

```
# Generate sampled values for variable model parameters via LHS
lhs_gen <- LatinHypercubeSampler$new()
lhs_gen$set_uniform_parameter("growth_rate_max", lower = 0.4, upper = 0.6, decimals = 2)</pre>
```

```
lhs_gen$set_uniform_parameter("harvest_rate", lower = 0.05, upper = 0.15, decimals = 2)
lhs_gen$set_uniform_parameter("initial_n", lower = 400, upper = 600, decimals = 0)
lhs_gen$set_uniform_parameter("density_max", lower = 80, upper = 120, decimals = 0)
lhs_gen$set_uniform_parameter("dispersal_p", lower = 0.2, upper = 0.5, decimals = 2)
lhs_gen$set_uniform_parameter("dispersal_b", lower = 400, upper = 1000, decimals = 0)
sample_data <- lhs_gen$generate_samples(number = 12, random_seed = 123)</pre>
sample data # examine
#>
      qrowth_rate_max harvest_rate initial_n density_max dispersal_p dispersal_b
#> 1
                  0.50
                               0.09
                                           575
                                                        111
                                                                   0.26
#> 2
                  0.41
                               0.07
                                           487
                                                         96
                                                                   0.34
                                                                                 589
#> 3
                  0.44
                               0.09
                                           483
                                                        110
                                                                   0.42
                                                                                 509
#> 4
                               0.11
                                           589
                                                                   0.38
                                                                                 859
                  0.59
                                                         86
                  0.56
                               0.06
                                                         92
                                                                   0.21
                                                                                 667
#> 5
                                           441
#> 6
                  0.51
                                                                   0.50
                                                                                 980
                               0.14
                                           458
                                                        116
#> 7
                  0.54
                               0.05
                                           416
                                                         88
                                                                   0.28
                                                                                 447
#> 8
                  0.43
                               0.13
                                           542
                                                        107
                                                                   0.36
                                                                                 920
#> 9
                               0.12
                                                                                 849
                  0.46
                                           422
                                                        101
                                                                   0.43
#> 10
                  0.52
                               0.08
                                           552
                                                         99
                                                                   0.32
                                                                                 745
#> 11
                  0.48
                               0.11
                                           525
                                                        119
                                                                   0.25
                                                                                 789
#> 12
                  0.58
                               0.15
                                           505
                                                         81
                                                                   0.45
                                                                                 609
```

Step 4: Build a simulation manager to run each simulation

We now wish to run a simulation for each set (or row) of sampled parameters. The *SimulationManager* class manages the generation of parameters (via the generators), the running the model simulations, and writing simulation results to disk. It also maintains a log of each simulation's success and any errors or warnings encountered.

```
# Create a simulation manager and run the sampled model simulations
sim_manager <- SimulationManager$new(sample_data = sample_data,</pre>
                                      model_template = model_template,
                                      generators = list(capacity_gen, dispersal_gen),
                                      parallel_cores = 2,
                                      results_dir = OUTPUT_DIR)
run output <- sim manager$run()</pre>
run_output$summary
#> [1] "12 of 12 sample models ran and saved results successfully"
dir(OUTPUT_DIR, "*.RData") # includes 12 result files
    [1] "sample_10_results.RData" "sample_11_results.RData"
   [3] "sample_12_results.RData" "sample_1_results.RData"
#>
   [5] "sample 2 results.RData" "sample 3 results.RData"
   [7] "sample_4_results.RData"
                                   "sample_5_results.RData"
                                   "sample\_7\_results.RData"
   [9] "sample_6_results.RData"
#> [11] "sample_8_results.RData"
                                   "sample_9_results.RData"
dir(OUTPUT_DIR, "*.txt") # plus simulation log
#> [1] "simulation_log.txt"
```

Note that the output directory contains a R-data result files for each sample simulation and a simulation log file.

Step 5: Build a results manager to generate summary results (metrics)

We now wish to collate summary results for each of our simulations via the *ResultsManager* class. This manager loads the results from each sample simulation into an intermediate *PopulationResults* class object, which dynamically generates further results. We need to define functions for calculating summary metrics, as well as any matrices (one row of values per simulation) that we may be interested in examining. Each metric (or matrix) is associated with

a user-defined function that utilizes results object attributes, or alternatively direct access to an attribute may be defined via a string. Once generated, the result metrics (a data frame) and/or matrices (a list) can be accessed via the manager. We may utilize the collated results in a variety of ways. However, with the objective of selecting the best models, we wish to compare (or validate) these result metrics to (with) known or desired target patterns (in step 6).

```
results_manager <- ResultsManager$new(simulation_manager = sim_manager,
                                        simulation_results = PopulationResults$new(),
                                        summary_metrics = c("trend_n", "total_h"),
                                        summary_matrices = c("n", "h"),
                                        summary_functions = list(
                                          trend n = function(results) {
                                            round(results$all$abundance_trend, 2)
                                          },
                                          total_h = function(results) {
                                            sum(results$harvested)
                                          },
                                          n = "all$abundance", # string
                                          h = "all$harvested"),
                                        parallel_cores = 2)
gen_output <- results_manager$generate()</pre>
gen_output$summary
#> [1] "12 of 12 summary metrics/matrices generated from sample results successfully"
dir(OUTPUT_DIR, "*.txt") # plus generation log
#> [1] "generation_log.txt" "simulation_log.txt"
results_manager$summary_metric_data
#>
      index trend_n total_h
#> 1
          1
               0.67
                         517
#> 2
          2
              -3.50
                         348
              -6.33
#> 3
          3
                         452
             -13.20
                         478
#> 4
          4
#> 5
          5
               2.00
                         283
          6
               5.00
                         691
#> 6
#> 7
          7
                         223
              -0.40
#> 8
          8
              -5.00
                         591
#> 9
          9
               4.14
                         548
#> 10
         10
               -5.12
                         414
#> 11
               0.57
                         588
         11
#> 12
         12
               -6.50
                         533
results_manager$summary_matrix_list
#> $n
#>
         [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
#>
    [1,] 542
               557
                    480
                         505
                               519
                                    516
                                         563
                                               525
                                                    529
                                                           509
          455
               471
                    481
                          480
                               463
                                    457
                                          442
                                               440
                                                    449
                                                           444
          481
               490
   [3,]
                    479
                          470
                               470
#>
                                    460
                                          431
                                               492
                                                    424
                                                           437
          462
               414
                     422
                          393
                               383
                                    368
                                          332
                                               352
                                                    327
                                                           366
                                               400
#>
   [5,]
          421
               405
                    464
                          466
                               455
                                    433
                                          459
                                                    492
                                                           434
   [6,]
          411
               413
                    428
                          428
                               409
                                    418
                                         403
                                               480
                                                    452
                                                           457
#>
   [7,]
          421
                               387
                                    457
                                               435
               438
                    467
                          440
                                          436
                                                    408
                                                           458
    [8,]
                               371
                                    366
                                          375
                                                    380
          474
               395
                     416
                          387
                                               360
                                                           388
                                          409
                    407
#>
   [9,]
               370
                          405
                               395
                                    410
                                               421
                                                    401
                                                           433
          392
#> [10,]
          480
               497
                    531
                          494
                               497
                                    448
                                          491
                                               451
                                                    455
                                                           456
#> [11,]
          509
               455
                     478
                          467
                               452
                                    482
                                          491
                                               513
                                                    474
                                                           461
#> [12,]
          383
               330
                    313
                          257
                               294
                                    267
                                         281
                                               316
                                                    306
                                                           293
#>
#> $h
```

```
#>
          [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
#>
    [1,]
                  56
                                               55
            52
                        46
                             49
                                   52
                                         53
                                                    51
                                                          52
                                                                 51
    [2,]
#>
            34
                  37
                        37
                             38
                                   34
                                         33
                                               33
                                                    34
                                                          35
                                                                 33
   [3,]
                             46
#>
            48
                  48
                        47
                                   46
                                         46
                                               42
                                                    47
                                                          41
                                                                 41
                                         47
#>
    [4,]
            59
                  50
                        54
                             48
                                   46
                                               42
                                                    43
                                                          42
                                                                 47
#>
    [5,]
            27
                  26
                        29
                             30
                                   27
                                         27
                                               30
                                                    27
                                                          32
                                                                 28
#>
   [6,]
            67
                  67
                        70
                             68
                                   66
                                         67
                                               65
                                                    75
                                                          72
                                                                 74
#>
   [7,]
            20
                  22
                        24
                             24
                                   19
                                         24
                                               23
                                                    23
                                                          20
                                                                 24
    [8,]
#>
            71
                  59
                        63
                             57
                                   56
                                         56
                                               58
                                                    54
                                                          58
                                                                 59
#>
   [9,]
            54
                  50
                        55
                             55
                                   54
                                         56
                                               55
                                                    58
                                                          53
                                                                 58
#> [10,]
                  42
                        46
                             43
                                   42
                                         39
                                               46
                                                    38
                                                          38
                                                                 39
            41
#> [11,]
                  55
                             57
                                   56
                                         58
                                               60
                                                    63
                                                          59
                                                                 58
                        58
            64
#> [12,]
                  56
                                               51
                                                    56
                                                          52
                             46
                                   49
                                         48
                                                                 54
```

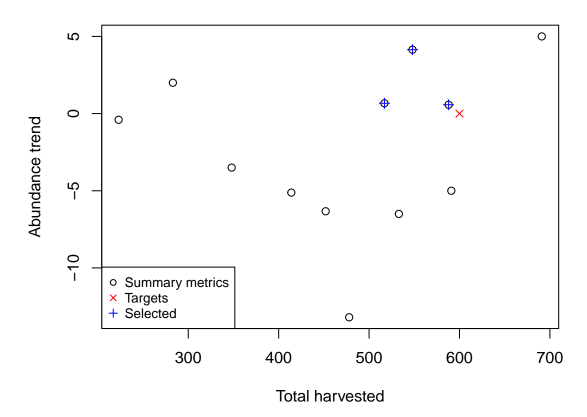
Step 6: Build a validator to select a model ensemble

We now select and analyze our 'best' models via a Validator class object, which by default utilizes an approximate Bayesian computation (ABC) approach (Beaumont, Zhang, & Balding, 2002) provided by the abc library (Csillery et al., 2015). The validator can be configured appropriately for a problem domain (see the abc documentation). Here we use the default configuration to select the best 3 models, along with a weight value, which is indicative of the congruence between each model's summary metrics and the corresponding target patterns. Also provided (with the default settings) is a diagnostic output (PDF) file, containing statistical information for analyzing the contribution of model parameters in the selection/validation process (see the abc documentation). For our simple demonstration the metrics and corresponding targets are relatively trivial, having the aim of producing stable population abundances with high harvest. However, the package facilitates the use more complex spatio-temporal metrics and targets (demonstrated in more advanced vignettes).

```
# Create a validator for selecting the 'best' example models
validator <- Validator $\text{simulation parameters} = \text{sample data},
                            simulation_summary_metrics =
                              results_manager$summary_metric_data[-1],
                            observed_metric_targets = c(trend_n = 0, total_h = 600),
                            output dir = OUTPUT DIR)
suppressWarnings(validator$run(tolerance = 0.25, output_diagnostics = TRUE))
#> 12345678910
#> 12345678910
dir(OUTPUT_DIR, "*.pdf") # plus validation diagnostics (see abc library documentation)
#> [1] "validation_diagnostics.pdf"
validator$selected_simulations # top 3 models (stable abundance and high harvest)
              weight
#>
     index
#> 1
         1 0.4469105
#> 2
         9 0.0000000
        11 0.9738769
```

We encourage the user to examine the generated diagnostics (PDF) output file, and to become acquainted with the analysis that this information facilitates (see the *abc* documentation). As our simple example only uses two metrics/targets, we can visualize the congruence of the selected models with the targets via a simple plot.

Example model validation



Summary

This demonstration has provided an overview of the *poems* workflow and modules via a simple population model example. We hope it has given you the foundation to progress to our more advanced Tasmanian *Thylacine* vignette, and towards utilizing the package for your own modeling projects.

Thank you:-)

References

Beaumont, M. A., Zhang, W., & Balding, D. J. (2002). 'Approximate Bayesian computation in population genetics'. Genetics, vol. 162, no. 4, pp, 2025–2035.

Csillery, K., Lemaire L., Francois O., & Blum M. (2015). 'abc: Tools for Approximate Bayesian Computation (ABC)'. R package version 2.1. Retrieved from https://CRAN.R-project.org/package=abc

Grimm, V., Revilla, E., Berger, U., Jeltsch, F., Mooij, W. M., Railsback, S. F., Thulke, H. H., Weiner, J., Wiegand, T., DeAngelis, D. L., (2005). 'Pattern-Oriented Modeling of Agent-Based Complex Systems: Lessons from Ecology'. *Science* vol. 310, no. 5750, pp. 987–991.