

Management frequency and extinction risk

GMSE: an R package for generalised management strategy evaluation (Supporting Information 6)

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The individual-based approach of default GMSE submodels

The default submodels of GMSE (`resource`, `observation`, `manager`, `user`) are individual-based (also called ‘agent-based’), meaning that they model discrete individuals (resources or agents), which in GMSE are represented by individual table rows (as in `RESOURCES`, `AGENTS`, and `OBSERVATION`) or layers of three-dimensional arrays (as in `COST` and `ACTION`). Individual-based models (IBMs) have been a useful approach in ecology for decades ([Uchmański and Grimm, 1996](#); [Grimm, 1999](#)), providing both a pragmatic tool for the mechanistic modelling of complex populations and a powerful technique for theoretical investigation. A key advantage of the individual-based modelling approach is the discrete nature of individuals, which allows for detailed trait variation and complex interactions among individuals. In GMSE, some of the most important traits for resources include types, ages, demographic parameter values, locations, etc., and for agents (manager and users), traits include different types, utilities, budgets, etc. The traits that resources and managers have can potentially affect their interactions, and default GMSE submodels take advantage of this by simulating interactions explicitly on a landscape (see [SI7](#) for an introduction to GMSE default data structures).

Replicate simulations as a tool for model inference

Mechanistically modelling complex interactions among discrete individuals typically causes some degree of stochasticity in IBMs (in the code, this is caused by the sampling of random values, which determine probabilistically whether or not events such as birth or death occur for individuals), reflecting the uncertainty that is inherent to complex systems. We can see a simple example of this by calling `gmse_apply` under the same default conditions twice.

```
rand_eg_1 <- gmse_apply();
print(rand_eg_1);

## $resource_results
## [1] 1097
##
## $observation_results
## [1] 1247.166
##
## $manager_results
##      resource_type scaring culling castration feeding help_offspring
## policy_1          1      NA      65         NA         NA         NA
##
## $user_results
##      resource_type scaring culling castration feeding help_offspring
```

```
## Manager      1      NA      0      NA      NA      NA
## user_1       1      NA     15      NA      NA      NA
## user_2       1      NA     15      NA      NA      NA
## user_3       1      NA     15      NA      NA      NA
## user_4       1      NA     15      NA      NA      NA
##      tend_crops kill_crops
## Manager      NA      NA
## user_1       NA      NA
## user_2       NA      NA
## user_3       NA      NA
## user_4       NA      NA
```

Although a second call of `gmse_apply` has identical initial conditions, because resource demographics (e.g., birth and death) and agent decision making (e.g., policy generation and user actions) is not deterministic, a slightly different result is obtained below.

```
rand_eg_2 <- gmse_apply();
print(rand_eg_2);

## $resource_results
## [1] 1070
##
## $observation_results
## [1] 929.7052
##
## $manager_results
##      resource_type scaring culling castration feeding help_offspring
## policy_1          1      NA     62          NA      NA          NA
##
## $user_results
##      resource_type scaring culling castration feeding help_offspring
## Manager          1      NA      0          NA      NA          NA
## user_1           1      NA     16          NA      NA          NA
## user_2           1      NA     16          NA      NA          NA
## user_3           1      NA     16          NA      NA          NA
## user_4           1      NA     16          NA      NA          NA
##      tend_crops kill_crops
## Manager      NA      NA
## user_1       NA      NA
## user_2       NA      NA
## user_3       NA      NA
## user_4       NA      NA
```

To make meaningful model inferences, it is often necessary to replicate simulations under the same initial conditions to understand the range of predicted outcomes for a particular set of parameter values. This can be computationally intense, but it can also lead to a more robust understanding of the range of dynamics that might be expected within a system. Additionally, when parameter values are unknown but believed to be important, replicate simulations can be applied across a range of values to understand how a particular parameter might affect system dynamics. Below, we show how to use the `gmse_replicates` function to simulate a simple example of a managed population that is hunted by users. This function calls `gmse` multiple times and aggregates the results from replicate simulations into a single table.

For a single simulation, the `gmse_table` function prints out key information from a `gmse` simulation result. The example provided in the GMSE documentation is below.

```
gmse_sim <- gmse(time_max = 10, plotting = FALSE);
```

```
## [1] "Initialising simulations ... "
```

```
sim_table <- gmse_table(gmse_sim = gmse_sim);
print(sim_table)
```

```
##      time_step resources estimate cost_culling cost_unused act_culling
## [1,]         1      1118 1065.7596          52          58          76
## [2,]         2      1180  997.7324         109           1          36
## [3,]         3      1269 1383.2200          10         100         312
## [4,]         4      1084 1337.8685          10         100         400
## [5,]         5       821 1292.5170          10         100         400
## [6,]         6       552  408.1633         110           0          36
## [7,]         7       605  589.5692         110           0          36
## [8,]         8       666  680.2721         110           0          36
## [9,]         9       749  884.3537         110           0          36
## [10,]        10       877 1065.7596          45          65          88
##      act_unused harvested
## [1,]          2         76
## [2,]          2         36
## [3,]         88        312
## [4,]          0        400
## [5,]          0        400
## [6,]          3         36
## [7,]          4         36
## [8,]          3         36
## [9,]          3         36
## [10,]         0         88
```

The above table can be saved as a CSV file using the `write.csv` function.

```
write.csv(x= sim_table, file = "file_path/gmse_table");
```

Instead of recording all time steps in the simulation, we can instead record only the last time step in `gmse_table` using the `all_time` argument.

```
sim_table_last <- gmse_table(gmse_sim = gmse_sim, all_time = FALSE);
print(sim_table_last)
```

```
##      time_step      resources      estimate cost_culling cost_unused
##          10.00          877.00         1065.76          45.00          65.00
## act_culling act_unused harvested
##          88.00           0.00          88.00
```

The `gmse_replicates` function replicates multiple simulations `replicates` times under the same initial conditions, then returns a table showing the values of all simulations. This can be useful, for example, for testing how frequently a population is expected to go to extinction or carrying capacity under a given set of parameter values. First, we demonstrate the `gmse_replicates` function for simulations of up to 20 time steps. The `gmse_replicates` function accepts all arguments used in `gmse`, and also all arguments of `gmse_table` (`all_time` and `hide_unused_options`) to summarise multiple `gmse` results. Here we use default `gmse` values in replicate simulations, except `plotting`, which we set to `FALSE` to avoid plotting each simulation result. We run 10 replicates below.

```
gmse_reps1 <- gmse_replicates(replicates = 10, time_max = 20, plotting = FALSE);
print(gmse_reps1);
```

```
##      time_step resources estimate cost_culling cost_unused act_culling
```

```
## [1,]      20      1006 1111.1111      15      95      264
## [2,]      20       941  839.0023     110       0       36
## [3,]      20      1270 1496.5986      10     100      400
## [4,]      20      1204 1269.8413      10     100      400
## [5,]      20      1143 1088.4354      31      79      128
## [6,]      20      1091 1043.0839      53      57       72
## [7,]      20      1290 1746.0317      10     100      400
## [8,]      20      1205 1020.4082     108       2       36
## [9,]      20      1012 1111.1111      23      87      172
## [10,]     20      1097 1383.2200      10     100      306
##      act_unused harvested
## [1,]          1       264
## [2,]          4        36
## [3,]          0       400
## [4,]          0       400
## [5,]          0       128
## [6,]          4        72
## [7,]          0       400
## [8,]          4        36
## [9,]          3       172
## [10,]         94       306
```

Note from the results above that resources in all simulations persisted for 20 time steps, which means that extinction never occurred. We can also see that the population in all simulations never terminated at a density near the default carrying capacity of `res_death_K = 2000`, and was instead consistently near the target population size of `manage_target = 1000`. If we wish to define management success as having a population density near target levels after 20 time steps (perhaps interpreted as 20 years), then we might assess this population as successfully managed under the conditions of the simulation. We can then see what happens if managers only respond to changes in the social-ecological system with a change in policy once every two years, perhaps as a consequence of reduced funding for management or increasing demands for management attention elsewhere. This can be done by changing the default `manage_freq = 1` to `manage_freq = 2`.

```
gmse_reps2 <- gmse_replicates(replicates = 10, time_max = 20, plotting = FALSE,
                             manage_freq = 2);
print(gmse_reps2);
```

```
##      time_step resources estimate cost_culling cost_unused act_culling
## [1,]      20      985 1043.0839      73      37       52
## [2,]      20     1905 1632.6531      10     100      400
## [3,]      20     1670 1632.6531      10     100      296
## [4,]      20      903  748.2993     110       0       36
## [5,]      20     1536 1405.8957      10     100      400
## [6,]      20     1179  997.7324     110       0       36
## [7,]      20      975  929.7052     110       0       36
## [8,]      20      965  748.2993     110       0       36
## [9,]      20      811  634.9206     110       0       36
## [10,]     20      686  725.6236     110       0       36
##      act_unused harvested
## [1,]          9        52
## [2,]          0       400
## [3,]        104       296
## [4,]          2        36
## [5,]          0       400
## [6,]          1        36
## [7,]          2        36
```

```
## [8,]      1      36
## [9,]      1      36
## [10,]     1      36
```

Note that while extinction still does not occur in these simulations, when populations are managed less frequently, they tend to be less close to the target size of 1000 after 20 generations. The median population size of `gmse_reps1` (management in every time step) was 1120, with a maximum of 1290 and minimum of 941. The median population size of the newly simulated `gmse_reps2` (management every two time steps) is 980, with a maximum of 1905 and minimum of 686. We can now see what happens when management occurs only once in every three time steps.

```
gmse_reps3 <- gmse_replicates(replicates = 10, time_max = 20, plotting = FALSE,
                             manage_freq = 3);
print(gmse_reps3);
```

```
##      time_step resources    estimate cost_culling cost_unused act_culling
## [1,]         20     1055 1315.19274          10        100          400
## [2,]         20     1043 1428.57143          10        100          400
## [3,]         20      986  907.02948         108           2           36
## [4,]         20      956  748.29932         110           0           36
## [5,]         20       78  136.05442         110           0           36
## [6,]         20     1062 1541.95011          10        100          400
## [7,]         20      805  634.92063         110           0           36
## [8,]          16        0   68.02721         110           0           36
## [9,]         20      392  294.78458         110           0           36
## [10,]        20     1025  680.27211         110           0           36
##      act_unused harvested
## [1,]           0        400
## [2,]           0        400
## [3,]           2         36
## [4,]           1         36
## [5,]           3         36
## [6,]           0        400
## [7,]           3         36
## [8,]           3          0
## [9,]           2         36
## [10,]          2         36
```

Given a management frequency of once every three time steps, the median population size of `gmse_reps3` (management in every time step) is 971, with a maximum of 1062 and minimum of 0. The number of extinctions observed in these replicate populations was 1. Below we change the management frequency to once every four time steps.

```
gmse_reps4 <- gmse_replicates(replicates = 10, time_max = 20, plotting = FALSE,
                             manage_freq = 4);
print(gmse_reps4);
```

```
##      time_step resources    estimate cost_culling cost_unused act_culling
## [1,]         20      330  226.75737          110           0           36
## [2,]         12        0   0.00000          110           0           36
## [3,]         20      653  589.56916          110           0           36
## [4,]         20     1713 1451.24717          10        100          367
## [5,]         20      705  839.00227          110           0           36
## [6,]         20      553  544.21769          110           0           36
## [7,]         12        0   0.00000          110           0           36
## [8,]         20      868  725.62358          110           0           36
```

```
## [9,]      8      0 0.00000      110      0      36
## [10,]     18      5 22.67574      110      0      36
##      act_unused harvested
## [1,]         1      36
## [2,]         3       0
## [3,]         2      36
## [4,]        33     367
## [5,]         2      36
## [6,]         3      36
## [7,]         3       0
## [8,]         2      36
## [9,]         3       0
## [10,]        2       5
```

Now note from the first column of `gmse_reps4` above that 4 populations did not persist to the 20th time step; i.e., 4 populations went to extinction (note that GMSE has a minimum resource population size of 5). This has occurred because managers cannot respond quickly enough to changes in the population density, and therefore cannot increase the cost of culling to maintain target resource levels if population size starts to decrease. We can see the extinction risk increase even further if management only occurs once every 5 time steps.

```
gmse_reps5 <- gmse_replicates(replicates = 10, time_max = 20, plotting = FALSE,
                             manage_freq = 5);
print(gmse_reps5);
```

```
##      time_step resources estimate cost_culling cost_unused act_culling
## [1,]         5       0       0       110         0         36
## [2,]         5       0       0       108         2         36
## [3,]         5       0       0       110         0         36
## [4,]         5       0       0       110         0         36
## [5,]         5       0       0       110         0         36
## [6,]         5       0       0       110         0         36
## [7,]         5       0       0       110         0         36
## [8,]         5       0       0       109         1         36
## [9,]         5       0       0       110         0         36
## [10,]        5       0       0       110         0         36
##      act_unused harvested
## [1,]         4       0
## [2,]         4       0
## [3,]         4       0
## [4,]         4       0
## [5,]         3       0
## [6,]         2       0
## [7,]         3       0
## [8,]         2       0
## [9,]         2       0
## [10,]        2       0
```

When a manager can only make policy decisions once every five time steps, extinction occurs in 10 out of 10 simulated populations before year 20. If we wanted to summarise these results, we could plot how extinction risk changes with increasing `manage_freq`.

```
ext_risk1 <- sum(gmse_reps1[,2] < 20);
ext_risk2 <- sum(gmse_reps2[,2] < 20);
ext_risk3 <- sum(gmse_reps3[,2] < 20);
ext_risk4 <- sum(gmse_reps4[,2] < 20);
```

```

ext_risk5 <- sum(gmse_reps5[,2] < 20);
y_var     <- c(ext_risk1, ext_risk2, ext_risk3, ext_risk4, ext_risk5);
x_var     <- 1:5;
plot(x = x_var, y = y_var, type = "b", pch = 20, lwd = 2, cex = 1.5,
     xlab = "Management every N time steps (manage_freq)",
     ylab = "Freq. of population extinction", cex.lab = 1.25)

```

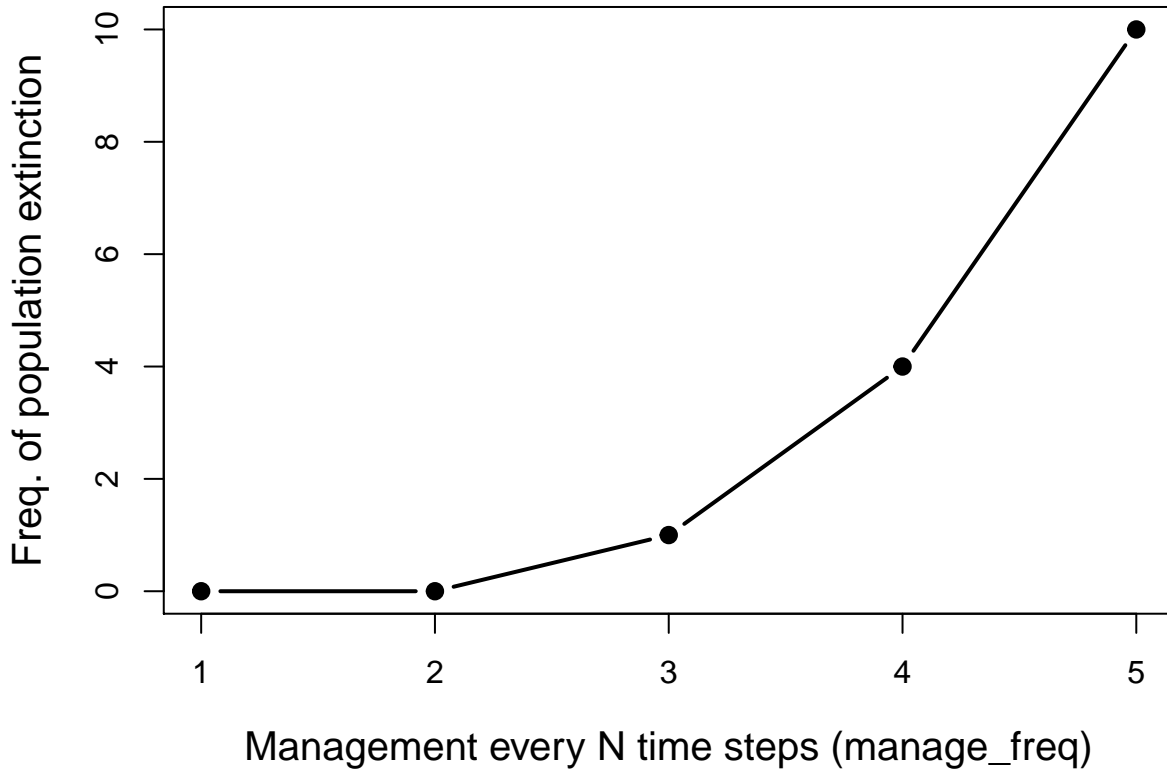


Figure 1: Extinction risk given an increasing number of time steps between updating policy decisions for culling costs in a simulated population. Higher values on the x-axis correspond to more time passing before a new policy is set. For each point, a total of 10 replicate simulations were run.

The above plot and the simulations from which it was derived illustrates a greatly simplified example of how GMSE might be used to assess the risk of extinction in a managed population. A comprehensive analysis would need more than 10 replicate simulations to accurately infer extinction risk, and would require careful parameterisation of all sub-models and a sensitivity analysis where such parameters are unknown. A benefit of this approach is that it allows for the simulation of multiple different scenarios under conditions of uncertainty and stochasticity, modelling the range of outcomes that might occur within and among scenarios and facilitating the development of social-ecological theory. Future expansion on the complexity of individual-based default submodels of GMSE will further increase the realism of targeted case studies.

References

- Grimm, V. (1999). Ten years of individual-based modelling in ecology: what have we learned and what could we learn in the future? *Ecological Modelling*, 115(2-3):129–148.
- Uchmański, J. and Grimm, V. (1996). Individual-based modelling in ecology: what makes the difference? *Trends in Ecology & Evolution*, 11(10):437–441.