Case Study 2: Ellner, Childs & Rees 2016

A more complicated, age and size structured model

Many life cycles cannot be described by a single, continuous state variable. For example, some plants are best modeled using height or diameter at breast height (DBH) as a state variable, and may also form a seed bank. Seeds in the seed bank can't have a value for height or DBH, but may lose their viability as they age. Thus, we require a discrete state to capture the dynamics of the seed bank. Models that include discrete states and/or multiple continuous state variables are general IPMs.

Many species experience some form of senesence, where survival increases and fecundity decreases after a certain age is attained. Age may interact with a measure of size, for example body mass, resulting in neither single variable reliably predicting demography on its own. Data sets where individuals are cross-classified by age and size represent an interesting opportunity for demographic research. Unfortunately, they can be a bit more complicated to model. This case study will use an age and size structured population to illustrate how to implement general IPMs in ipmr.

The model itself can be written on paper as:

1.
$$n_0(z', t+1) = \sum_{a=0}^{M} \int_{L}^{U} F_a(z', z) n_a(z, t) dz$$

2.
$$n_a(z',t+1) = \int_L^U P_{a-1}(z',z) n_{a-1}(z,t) dz$$
 a = 1,2,..., M-1

In this case, there is also a maximum age class M defined as a > 20. We need to define one more equation to indicate the number of individuals in that age group.

3.
$$n_M(z',t+1) = \int_L^U [P_M(z',z)n_M(z,t) + P_{M-1}(z',z)n_{M-1}(z,t)]dz$$

The sub-kernel $P_a(z',z)$ is comprised of the following functions:

4.
$$P_a(z',z) = S(z,a) * G(z',z,a)$$

5.
$$Logit^{-1}(S(z,a)) = \alpha_s + \beta_s^z * z + \beta_s^a * a$$

6.
$$G(z', z, a) \sim Norm(\mu_q(z, a), \sigma_q)$$

7.
$$\mu_g(z) = \alpha_g + \beta_g^z * z + \beta_g^a * a$$

and the sub-kernel \$F_a(z',z) is comprised of the following functions:

8.
$$F_0 = 0$$

9.
$$F_a = S(z, a) * p_b(z, a) * p_r(a) * f_d(z', z) * 0.5$$

The F_a kernel is multiplied by 0.5 because this model only models the population dynamics of females.

10.
$$Logit^{-1}(p_b(z,a)) = \alpha_{p_b} + \beta_{p_b}^z * z + \beta_{p_b}^a * a$$

11.
$$Logit^{-1}(p_r(a)) = \alpha_{p_r} + \beta_{p_r}^a * a$$

12.
$$f_d(z',z) \sim Norm(\mu_{f_d}(z), \sigma_{f_d})$$

13.
$$\mu_{f_d}(z) = \alpha_{f_d} + \beta_{f_d}^z * z$$

Equations 5, 7, 10, 11, and 13 are parameterized from regression models, and equations 6 and 12 are constant parameters derived from observed data. This example will skip the step of model parameterization, and go straight into the IPM.

First, we will define the model parameters and function for the F_a kernels.

```
# library(ipmr)
devtools::load_all()
## Loading ipmr
##
## Attaching package: 'rlang'
## The following object is masked from 'package:ipmr':
##
##
       flatten
## The following objects are masked from 'package:testthat':
##
       is_false, is_null, is_true
# Set parameter values and names
param_list <- list(</pre>
  ## Survival
  surv_int = -1.70e+1,
  surv_z = 6.68e+0,
         = -3.34e-1,
  surv_a
  ## growth
  grow_int = 1.27e+0,
  grow_z = 6.12e-1,
  grow_a
         = -7.24e-3,
  grow_sd = 7.87e-2,
  ## reproduce or not
  repr int = -7.88e+0,
           = 3.11e+0,
  repr_z
          = -7.80e-2
  repr_a
  ## recruit or not
 recr_int = 1.11e+0,
          = 1.84e-1,
  recr_a
  ## recruit size
 rcsz_int = 3.62e-1,
 rcsz_z = 7.09e-1,
  rcsz_sd = 1.59e-1
# define a custom function to handle the F kernels. We could write a rather
# verbose if (age == 0) {0} else {other_math} in the define_kernel(), but that
# might look ugly. Note that we CANNOT use ifelse(), as its output is the same
# same length as its input (in this case, it would return 1 value, not 10000 values).
f_fun <- function(age, s_age, pb_age, pr_age, recr) {</pre>
  if(age == 0) return(0)
  s_age * pb_age * pr_age * recr * 0.5
}
```

Next, we set up the P_a kernels (Equations 4-7 above). Because this is a general, deterministic IPM, we set the model class to "general_di_det", and set has_age = TRUE to indicate that our model has age structure as well as size structure. There are 3 k ey things to note:

- 1. the use of the suffix _age appended to the names of the "P" kernel and the mu_g_age variable
- 2. the value age used in the vital rate expressions
- 3. the list in the levels_ages argument

The values in the levels_ages list will automatically get substituted in for "age" each time it appears in the vital rate expressions and kernels. This single call to define_kernel() will result in 22 actual kernels, one for each value of age from 0-21.

The plogis function is part of the stats package in R, and performs the inverse logit transformation.

```
age_size_ipm <- init_ipm("general_di_det", has_age = TRUE) %>%
  define_kernel(
                  = "P_age",
    name
                  = "CC",
    family
                  = s_age * g_age * d_z,
    formula
                  = plogis(surv_int + surv_z * z_1 + surv_a * age),
    s_age
                 = dnorm(z_2, mu_g_age, grow_sd),
    g_age
              = grow_int +
= param_list,
                 = grow_int + grow_z * z_1 + grow_a * age,
    mu_g_age
    data_list
                 = list(c("z")),
    states
    has_hier_effs = FALSE,
    levels_ages = list(age = c(0:20), max_age = 21),
                  = FALSE
    evict_cor
  )
```

The F_a kernel (equations 8-13) will follow a similar pattern - we append a suffix to the name paramter, and then make sure that our functions also include _age suffixes and age values where they need to appear.

```
define_kernel(
age_size_ipm <-
  proto_ipm
                = age_size_ipm,
                = "F_age",
  name
                = "CC",
  family
  formula
                = f_fun(age, s_age, pb_age, pr_age, recr) * d_z,
                = plogis(surv_int + surv_z * z_1 + surv_a * age),
  s_age
                = plogis(repr_int + repr_z * z_1 + repr_a * age),
  pb_age
               = plogis(recr_int + recr_a * age),
  pr_age
                = dnorm(z_2, rcsz_mu, rcsz_sd),
  recr
               = rcsz_int + rcsz_z * z_1,
  rcsz_mu
  data_list
                = param_list,
                = list(c("z")),
  states
  has_hier_effs = FALSE,
  levels_ages = list(age = c(0:20), max_age = 21),
  evict_cor
                = FALSE
)
```

Once we've defined the P_a and F_a kernels, we need to define how they interact with age-specific size distributions to produce size distributions at the next time step. This corresponds to equations 1-3 above. There a couple things to note here:

- 1. We do not add "_age" to the name here. This is because we do not want to expand this into 22 separate iteration procedures we only want 1 that iterates the 22 separate P_a and F_a kernels together.
- 2. the n_z_0_t_1 = sum(F_age %*% n_z_age_t). This takes the place of the $\sum_{a=0}^{M}$... in Eq. 1 above.

We don't need to supply a=0 and M as those are inferred in the levels_ages list.

- 3. The insertion of age_minus_1 in the n_z_age_t_1 expression. ipmr automatically generates the correct indices using these markers, so there is no need to worry about indexing correctly. Tragically, we could not implement a literal age 1. _plus_ is also available for models with different formats. The amount to add/subtract is not limited to 1, though it is hard to think of examples where we would want to index by larger increments (perhaps with censuses every 2 or 3 years instead annual?).
- 4. the max_age in n_z_max_age_t_1. This corresponds to equation 3. We can adjust the maximum age value in the model by changing the max_age slot in the levels_ages list.

```
age_size_ipm <- define_k(</pre>
  proto_ipm
                  = age_size_ipm,
                  = "K",
  name
                  = "IPM",
  family
                  = sum(F_age %*% n_z_age_t),
  n_z_0_t_1
                  = P_age_minus_1 %*% n_z_age_minus_1_t,
  n_z_age_t_1
  n_z_max_age_t_1 = P_max_age %*% n_z_max_age_t +
                    P_max_age_minus_1 %*% n_z_max_age_minus_1_t,
  data_list
                  = param_list,
                  = list (c("z")),
  states
  has_hier_effs
                  = FALSE,
                  = list(age = c(0:20), max_age = 21),
  levels_ages
  evict_cor
                  = FALSE
  )
```

The rest of the model definition code looks like the first case study. We define the implementation arguments using define_impl(), the domains using define_domains(), and the initial population state using define_pop_state(). We can compute the populations asymptotic growth rate, and stable size distributions for each age using the lambda() and right_ev() functions.

```
age size ipm <- define impl(
 proto_ipm = age_size_ipm,
  make impl args list(
    kernel_names = c("P_age", "F_age", "K"),
               = rep("midpoint", 3),
    int rule
                 = rep("z", 3),
    dom_start
                 = rep("z", 3)
    dom end
  )
) %>%
  define_domains(
    z = c(1.6, 3.7, 100)
  ) %>%
  define_pop_state(
    pop_vectors = list(
      n_z_{age} = runif(100)
  ) %>%
  make_ipm(
    usr_funs = list(f_fun = f_fun),
    iterate = TRUE,
    iterations = 100
  )
             <- lambda(age size ipm)
stable_dists <- right_ev(age_size_ipm)</pre>
```

We can plot the stable distributions using an lapply() on the stable_dists object.

```
plot(stable_dists[[1]], type = 'l')
cap <- lapply(stable_dists[2:22], function(x) lines(x))</pre>
```

