Optimal Semelparity

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Abstract

Semelparous organisms have a simple life cycle characterized by immediate death after reproduction. We assume that semelparous life histories can be separated into a juvenile non-reproductive period followed by an adult period during which reproduction is possible. We derive formulae for the optimal age and size at reproduction and for the optimal size of the offspring (e.g., seeds). Our main contribution is to determine the conditions under which the optimal size of the offspring does not depend on the optimal size at reproduction and vice versa.

The life cycle of semelparous species can be viewed as a two-phase process, driven by different mechanisms. Stage 1 is a juvenile non-reproductive period, in which some individuals survive to become adults. Adults can reproduce and, when they do, they die. Hence stage 2 is the period of life in which individuals seek to maximize their reproduction by weighing at each instant the benefits of delaying reproduction further against the risk of death associated with this delay. We assume that stage 1 is the milestone between the two phases. Without loss of generality, we can further assume that size 1 corresponds to adult age 0. If the organism survives to size 1, it is larger than the smallest size that can reproduce at age 0 (see Fig. 1). Our first question is what determines the size of the smallest size at which reproduction is possible. Second, what is the optimal number of offspring and what is the optimal size of each offspring? Third and most importantly, does the optimal number of offspring depend on the optimal size of each offspring? Our first question is whether the optimal size of an organism at reproduction determines the optimal offspring size. If it does, they die. Hence stage 2 is the period of life in which individuals seek to maximize their reproduction by weighing at each instant the benefits of delaying reproduction further against the risk of death associated with this delay. We assume that size 1 is the milestone between the two phases.
Our third and most important question concerns the relationship between $\xi_0$ and $i_1$. The assumptions we made about the separation of the two stages imply that $\xi_0$ and $i_0$ are independent and, similarly, $\xi_1$ and $i_1$ are independent. The question of interest is whether $\xi_0$ and $i_1$ are independent. This formulation has not been clearly developed in previous studies [12] and is a key contribution.

**Semelparous Strategies: Models and Results**

**Optimal Age and Size at Reproduction**

Stage 2, which starts once seed size no longer affects the risk of dying, is the stage of adult growth during which reproduction is possible. If reproduction occurs only at age $\bar{a}$ in stage 2, if the chance $R$ of surviving to $\bar{a}$ is constant over time and across environments, and if $m(a)$ and $\delta$ are similarly constant, then the net reproduction rate $R$ for such semelparous species can be expressed as

$$R = e^{-\delta \bar{a}} l(\bar{a})m(\bar{a}),$$

(1)

where $r$ is the rate of population growth, and $m(\bar{a})$ measures reproduction at age $\bar{a}$; $m(a)$ at any age $a$ other than $\bar{a}$ is zero. This implies that

$$e^{-\delta \bar{a}} l(\bar{a})m(\bar{a}) = 1$$

(2)

[13, p189], an expression that follows directly from the Lotka equation,

$$\int_0^\infty e^{-ra} l(a)m(a) da = 1.$$  

(3)

Proof that $r$ represents the growth rate in the Lotka equation is not straightforward and depends on the assumption of stable populations (see [14]), but (2) for semelparous species is true by definition. The simplicity of (2) facilitates analytical insights into optimal age at reproduction and optimal offspring size.

Solving (2) for $r$ yields (see [13], p.189)

$$r = \frac{\ln(l(\bar{a})m(\bar{a}))}{\bar{a}}.$$  

(4)

The value of $\bar{a}$ that maximizes $r$ is the optimal age at reproduction, $\bar{a}$. It satisfies the condition

$$\frac{dr}{d\bar{a}}|_{\bar{a}=\bar{a}} = 0.$$  

(5)

Inserting the expression for $r$ from (4) into (5), using the equation for the derivative to solve for $\bar{a}$, and rearranging terms yields the requirement that the optimal age at reproduction, denoted by $\bar{a}$, must satisfy:

$$m'(\bar{a}) - \mu(\bar{a}) = \frac{\ln(l(\bar{a})m(\bar{a}))}{\bar{a}}.$$  

(6)

where $m'(\bar{a}) = \frac{[m'(\bar{a})]}{[m(\bar{a})]} / [m(\bar{a})]$ and $\mu(\bar{a}) = [-d\bar{a}(l(\bar{a})/l(\bar{a})) / l(\bar{a})].$ Note that $m'(\bar{a})$ is the relative rate of improvement in reproductive capacity at age $\bar{a}$, and $\mu(\bar{a})$ is the hazard of death (force of mortality) at age $\bar{a}$. Substituting (4) into (6) shows that

$$m'(\bar{a}) - \mu(\bar{a}) = r(\bar{a}).$$  

(7)

At equilibrium, $r=0$ and the optimal age at reproduction is defined by a balance between the rate of growth in reproductive capacity and the force of mortality,

$$m'(\bar{a}) = \mu(\bar{a}).$$  

(8)

Note that in reality populations, especially semelparous populations, might not be always at equilibrium. We will, nevertheless, assume they are in order to illustrate the trade-off mechanism in determining the optimal timing of reproduction. From (8), reproduction should be delayed as long as the reproductive benefits of further growth outweigh the risk of mortality occasioned by delaying. The optimal age at reproduction is the age at which the benefits of further growth are exactly offset by the risk of dying. Note that $\delta$, the duration of stage 1, does not appear in (8) and does not affect the optimal age (in stage 2) of reproduction. If the population were growing or shrinking, then $\delta$ would matter, as it would affect time to reproduction; with earlier times being favored in growing populations (see [15,16]); and later
times in shrinking populations (see [17]). In the rest of this article we focus on the equilibrium case when \( r = 0 \) and we will use “age” to refer to age in stage 2.

The optimal size at reproduction \( \hat{z} = \hat{z}(\hat{a}) \) is the size of the semelparous organism at the optimal age at reproduction. We assume semelparous organisms grow until they reproduce, i.e. \( \hat{z}(a) \) is an increasing function of age (this might not always be the case as shown in [18,19]). As a result, this optimal size can be determined by

\[
m'(\hat{z}(\hat{a})) \frac{d\hat{z}}{da} = \mu(\hat{z}),
\]

where the parameter \( k \) captures how the growth rate increases with size, and \( \kappa \) modulates the increase so that eventually size reaches an asymptote. For illustration, we use the exponent 0.75, following predictions from the fractal model of scaling (see [23]). However, using a different exponent would not alter the main conclusions of the article. This equation provides a fairly general description of asymptotic growth. If size at age 0 is 1, we have

\[
\hat{z}(a) = \left( \frac{k}{\kappa} - \left( \frac{k}{\kappa} - 1 \right) e^{-\frac{k}{4} a} \right)^4,
\]

which reduces to

\[
(\eta k + c) \hat{z} - \eta k \hat{z}^{0.75} + b = 0
\]

The latter is a quartic equation for \( \hat{z}^{0.25} \) and its analytic solution is given by Ferrari’s formula. Denoting

\[
A = \frac{\eta k}{\eta k + c}, \quad B = \frac{b}{\eta k + c},
\]

we can express the positive root of the quartic equation (14) in the following manner

\[
\hat{z} = -\frac{A}{4} + \frac{1}{2} \sqrt{\frac{A^2}{4} + 2C + \frac{2B}{3C}} + \frac{1}{4} \sqrt{\frac{7A^2}{4} + 2C + \frac{2B}{3C} + \frac{A^3}{4C}}.
\]

As a result, \( \hat{z} \) increases with \( A \) and decreases with \( B \) (see Fig. 2). Therefore, the optimal size of reproduction \( \hat{z} \) will increase with positive changes in the reproduction scale parameter \( \eta \) or the determinants of asymptotic size \( \frac{k}{\kappa} \) as well as negative changes in mortality parameters \( b \) or \( c \).

These mathematical results aid biological insight. Because optimal size does not depend on the parameter \( \phi \), species suffering proportional reduction in offspring production will, ceteris paribus, not vary in flowering size (see [24]). An example of this
might be density dependence of seed establishment [see 20]. Furthermore, if species’ relative ranking with respect to asymptotic size \( k/k_i \), scaling of reproductive output with size \( g \), and mortality parameters, \( b \) and \( c \), are known, relative ranking in terms of flowering size could be predicted.

Optimal Seed Size and Number

Let \( p(i) \) be the probability that a seed germinates and grows until initial size no longer influences mortality, i.e. to \( x = 0 \) and size taken as \( \xi = 1 \). Generally \( p(i) \) increases with seed size \( i \). Let reproductive output i.e., number of seeds produced, be denoted by \( n(x,i) \) which is an increasing function of plant size (and age), and a decreasing function of seed size. The net reproductive rate is then

\[
R = p(i)l(x)n(x,i). \quad (16)
\]

If the population is in equilibrium, maximizing \( r \) is generally equivalent to maximizing \( R \) [see 25]. Further, in [24] it has been shown that maximizing \( R \) provides the evolutionary stable strategy if population regulation operates on offspring establishment. Such density dependence characterizes many semelparous species [see 20]. The optimal life history is therefore defined by the derivative or relative derivative of \( R \) being equal to zero. Hence, the optimal age at reproduction can be specified by

\[
\frac{dR}{dx} = 0 = n'(x,i) - \mu(\xi) \quad (17)
\]

where \( n'(x,i) = (dn(x,i)/dx)/n(x,i) \) defines the rate of change in the number of offspring produced at age \( x \). Equation (17) implies \( n'(x,i) = \mu(\xi) \), which is similar to the result obtained in (8). Note that optimal time at reproduction depends only on \( x \) in stage 2 and does not depend on time taken by a seed to grow to \( \xi = 1 \) (see [12]). The optimal offspring size is specified by

\[
\frac{dR}{di} = 0 = n'(x,i) + p'(i) \quad (18)
\]

where \( p'(i) = (dp(i)/di)/p(i) \) and \( n'(x,i) = (dn(x,i)/di)/n(x,i) \). This implies \( p'(i) = -n'(x,i) \). At equilibrium, optimal offspring size is the size at which the benefits accrued through investing less in each offspring and thereby producing more offspring are offset by the risk of mortality for an offspring of that size.

Optimal Seed Size in a Specific Model for Stage 1

Specific functional forms can be used to deepen understanding. The number of seeds \( n \) of size \( i \) produced at age \( x \) can be determined by
where parameter $\beta$ captures both saturating and accelerating functional forms of producing larger offspring. The probability of reaching size $\xi = 1$ can be specified by a concave function

$$p(t) = \left(\frac{t - t_0}{1 - t_0}\right)^\gamma, \quad 0 < \gamma < 1,$$  

where $t_0$ is the minimal possible seed size and $\gamma$ accounts for the speed of reaching reference size $\xi = 1$. As a result, the optimal offspring size $t$ will be the solution of (18) i.e.

$$i = \frac{\beta t_0}{\beta - \gamma}.$$  

Discussion

When is Optimal Seed Size $i$ Independent of Optimal Adult Size at Reproduction $\xi$?

Eq. (21) implies that the optimal seed size $t$ does not depend on the optimal plant size at reproduction $\xi$. Using (18), it can be similarly shown that optimal plant size at reproduction does not depend on the optimal size of the seeds produced. This mutual independence holds in general if the number of seeds of size $t$ produced at age $z$ is proportional to the product of a function of adult size and a function of seed size, i.e.

$$n(z,t) = CF(t)G(z), \quad C \equiv \text{const}$$  

where $C$ is a scaling factor. In this case

$$n'_i = F'_i$$  

does not depend on $\xi$ and neither does $p'(t)$. This is also true for

$$n'_\xi = F'_\xi$$  

Eq. (22) is a necessary and sufficient condition, in our framework, for the independence of the parent’s optimal size at reproduction from the optimal seed size of its offspring. The condition is not implausible, but it is also not trivial. For instance, in (19) $\beta$ might be a function of $\xi$: bigger plants might be more efficient at producing large seeds than smaller plants are. Also in (19), $\gamma$ might be a function of $t$: the relationship between plant size and reproductive capacity may be modulated by seed size.

Note that the assumptions about a juvenile vs. an adult stage imply that $\xi_\phi$ is independent of $t_0$ and $\xi_1$ is independent of $i_1$ (see Fig. 1). To prove independence of optimal seed size and optimal size at maturity, it is also necessary to show that $\xi_\phi$ and $i_1$ are independent. Eq. (22) gives the condition for this.

The independence of two characteristics means that the optimal value of either of them does not depend on the value of the other characteristic. This causal independence is different from lack of empirical correlation. For instance, suppose a species grows in two environments, one unfavorable (perhaps because of poor soil or lack of sunlight) and the other favorable. Then, the time it takes a plant to grow from seed to adult size, and $z$, the time it takes for the plant to grow from adult size to size at reproduction and death, might be correlated across the two environments: e.g., both times might be long in the unfavorable environment and short in the favorable one. The long time to develop, however, does not cause the long time to mature: the unfavorable environment causes both and the correlation is merely a statistical association. As explained above, the duration $\delta$ is irrelevant to the optimization problems we addressed.

Conclusion

The simplicity of the semelparous life cycle aids formulating general mathematical models that predict key features of life histories. The analytical framework presented here unifies predictions of timing of reproduction and offspring size. This framework provides insights into how basic demographic features shape the diversity of age trajectories across species and plasticity within species in response to environmental cues. This permits separation of these patterns from complications such as variation in growth, both across individuals (see [20]) and through time (see [26]). Variants of the models may also be relevant for other life-history switches such as metamorphosis (see [27]).

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