

THE EVOLUTION OF INTERTEMPORAL PREFERENCES

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The Evolution of Intertemporal Preferences

*Arthur Robson and Larry Samuelson**

Where do preferences come from? What determines their properties? Though traditionally reluctant to ask such questions, economists have recently turned to evolutionary models for answers.

We focus on intertemporal preferences here, arising out of the evolutionary implications of different reproductive strategies or *life histories*. An agent's life history specifies the agent's number and timing (and, in a richer model, quality) of offspring. Evolution will select the life history that maximizes the growth rate of the associated group of individuals.

We begin with the simplest possible biological life history—that of a *semelparous* agent that, if it survives a fixed number of years, reproduces and then dies. We show the evolutionary criterion for success in this case entails hyperbolic time discounting of the log of the number of offspring produced. However, the rate of time preference is a function of age, not of time relative to the present, and hence there are no preference reversals in the sense of behavioral economics. At the same time, the *optimal* strategy maximizes the exponentially discounted number of offspring, provided we discount at the sum of the death rate and the *maximal* growth rate. Conventional discounting thus suffices to induce optimal choices from the agent.

More generally, if the animal is *iteroparous*, and so may have a non-degenerate profile of offspring, we show the evolutionary indifference curves over offspring of various ages are hyperplanes that are not parallel, but tilt to reflect greater impatience as the growth rate increases. There is no additively separable function of the age profile of expected offspring that is globally equivalent to this basic biological growth-rate criterion, even if arbitrary age dependence is allowed. Further, the rate of time discounting of offspring falls with age, if there is a positive discounting to begin with. Evolution's preferences thus exhibit a

present bias, though this still cannot imply preference reversals. Again, the *optimal* strategy maximizes the appropriately exponentially discounted number of offspring.

1 Semelparous Life Histories

We begin with the simplest case, that of a *semelparous* life history, in which an organism reproduces at a fixed, single age (if it survives that long) and then dies.

Let time be discrete, given by $t = 0, 1, 2, \dots$. An agent is characterized by a pair (x, τ) identifying the agent's expected number of offspring x , to be produced at age τ , conditional on the agent's surviving until age τ .

In keeping with our focus on preferences over reproduction, we represent the strategy (x, τ) simply as a number of offspring and a time, though it presumably reflects a collection of activities, such as where to forage, what food to eat, when to mate, and so on, that determine reproduction. The strategy (x, τ) is heritable, so that offspring are characterized by the same strategy as their parents. We are interested in which such strategy will be favored by evolution.

While waiting to reproduce, the agent faces an instantaneous death risk of $\delta > 0$. An agent choosing (x, τ) thus survives for τ periods with probability $e^{-\delta\tau}$. If and only if the agent survives, the x offspring appear, at which point the agent dies. We work with a continuum of agents and without aggregate uncertainty.

Consider a population characterized by strategy (x, τ) . Let $N^T(t)$ be a (row) vector $(N_1(t), N_2(t), \dots, N_\tau(t))$ describing the measures of agents of each age in the population, at the beginning of time t . During time t , the τ -period-old agents produce an average of x offspring apiece, and then die. A proportion $e^{-\delta}$ of the remaining agents then survive until the next period, each becoming one period older in the process. The basic difference

equation governing the population is

$$N^T(t) = N^T(t-1) \begin{bmatrix} 0 & e^{-\delta} & 0 & \dots & 0 & 0 \\ 0 & 0 & e^{-\delta} & \dots & 0 & 0 \\ \vdots & \vdots & \vdots & & \vdots & \vdots \\ 0 & 0 & 0 & \dots & 0 & e^{-\delta} \\ e^{-\delta}x & 0 & 0 & \dots & 0 & 0 \end{bmatrix}.$$

In this $\tau \times \tau$ matrix, each row $n \in \{1, \dots, \tau - 1\}$ applies to the agents who enter period t at age n . The single term $e^{-\delta}$ that appears in row n identifies the proportion of these agents who survive until the beginning of the next period, at which point they are one period older. The final row corresponds to agents who enter at age τ . These agents have x offspring each, with a fraction $e^{-\delta}$ of these offspring surviving to become next period's 1-period-olds and with the parents disappearing from the population. This is referred to as a *Leslie matrix* (Brian Charlesworth (1994)).

Asymptotically, the growth factor $\tilde{\lambda}$ for this population is the unique positive real root of the characteristic equation of the Leslie matrix.¹ The characteristic equation is equivalent to $\lambda^\tau = e^{-\delta\tau}x$, so the growth factor is $\tilde{\lambda} = e^{-\delta}(x)^{\frac{1}{\tau}}$.

We now consider a population of agents characterized by a variety of life histories (x, τ) . Evolution will select for the value (x, τ) that maximizes $e^{-\delta}(x)^{\frac{1}{\tau}}$ or, equivalently, that maximizes

$$\frac{\ln x}{\tau}.$$

Hence, evolution evaluates births according to the function $\ln(\cdot)$ and discounts them hyperbolically—evolution exhibits a present bias. The equilibrium population will grow geometrically at rate $\tilde{\lambda}$, or (equivalently) exponentially at the growth rate $r = \ln \tilde{\lambda}$.

Evolution can induce agents to behave optimally via conventional exponential discounting. Let (x_1, τ_1) be the optimal life history, giving rise to exponential growth at rate

$r_1 = \ln \left(e^{-\delta} x_1^{\frac{1}{\tau_1}} \right) = -\delta + \frac{1}{\tau_1} \ln x_1$. Let the alternative strategy (x_2, τ_2) give growth rate $r_2 = \ln \left(e^{-\delta} x_2^{\frac{1}{\tau_2}} \right) = -\delta + \frac{1}{\tau_2} \ln x_2 < r_1$. Suppose that agents are designed to evaluate births linearly and discount exponentially at rate $-(\delta + r_1)$, so that (x, τ) is evaluated as $e^{-(\delta+r_1)\tau} x$. This choice of discount rates is intuitive. There are two costs of delaying reproduction. One of these is simply that death occurs at rate δ . The other is that a given number of offspring will comprise a smaller fraction of a population growing at rate r_1 . The sum of these two rates is the rate at which delaying births causes an agent to fall behind the population. The comparison between two reproductive strategies is then given by (using $e^{-(\delta+r_1)\tau_1} x_1 = 1 = e^{-(\delta+r_2)\tau_2} x_2$)

$$\begin{aligned}
e^{-(\delta+r_1)\tau_1} x_1 > e^{-(\delta+r_1)\tau_2} x_2 &\Leftrightarrow e^{-(\delta+r_1)\tau_1} x_1 > e^{-r_1\tau_2} e^{r_2\tau_2} e^{-(\delta+r_2)\tau_2} x_2 \\
&\Leftrightarrow 1 > e^{-r_1\tau_2} e^{r_2\tau_2} \\
&\Leftrightarrow r_1 > r_2.
\end{aligned}$$

Exponential discounting, at the sum of the death and optimal growth rates, thus identifies the optimal strategy (x, τ) . More rapid population growth induces a higher discount rate. (See Ingemar Hansson and Charles Stuart (1990) and Alan R. Rogers (1994) for similar results.) A population that shrinks rapidly enough will induce negative discounting, in which case reproduction is better deferred. If the population growth rate is zero, agents will discount at the death rate δ . This will be the case if Malthusian considerations constrain the set of feasible (x, τ) values to those featuring nonpositive growth.

2 Iteroparous Life Histories

We now generalize the analysis to *iteroparous* life histories, in which an individual may have offspring at more than one age.

2.1 Asymptotic Growth Rate

Suppose that agents live for ℓ periods, producing x_i offspring in each period $i = 1, \dots, \ell$. A life history is then a collection $(x_1, x_2, \dots, x_\ell)$, where some of these entries may be zero. The Leslie matrix is

$$\begin{bmatrix} e^{-\delta}x_1 & e^{-\delta} & 0 & \dots & 0 & 0 \\ e^{-\delta}x_2 & 0 & e^{-\delta} & \dots & 0 & 0 \\ \vdots & \vdots & \vdots & & \vdots & \vdots \\ e^{-\delta}x_{\ell-1} & 0 & 0 & \dots & 0 & e^{-\delta} \\ e^{-\delta}x_\ell & 0 & 0 & \dots & 0 & 0 \end{bmatrix}.$$

The population's asymptotic growth factor is again the largest real root $\tilde{\lambda}$ of the characteristic equation of this matrix. This equation can be written as

$$(1) \quad 1 = \frac{x_1}{e^\delta \lambda} + \frac{x_2}{(e^\delta \lambda)^2} + \dots + \frac{x_\ell}{(e^\delta \lambda)^\ell}.$$

Evolution will select for the life history that maximizes this growth factor. Along any indifference surface for evolution, $e^\delta \tilde{\lambda}$ is constant and hence we have a linear function of the values x_1, \dots, x_ℓ . Evolution's indifference surfaces are thus hyperplanes in a space of ℓ dimensions. Figure 1 illustrates.

It is immediate from (1) that evolution is indifferent over two semelparous strategies (x_1, τ_1) and (x_2, τ_2) if and only if $x_1^{\frac{1}{\tau_1}} = x_2^{\frac{1}{\tau_2}}$. This confirms that the semelparous analysis is a special case of this more general model. Preferences over the remaining iteroparous strategies are captured by connecting indifferent semelparous strategies with linear indifference surfaces.

2.2 No Time Separable Representation

The basic biological criterion for the population growth rate is a complex function of the fertility profile. Let $e^\delta \lambda = \theta(x_1, x_2, \dots)$ be the function implicitly defined by (1). Then the

marginal rate of substitution between x_t and x_{t+1} is θ itself, which is a strictly increasing function of each x_s for $s \neq t, t+1$. It is then immediate that there then can be no additively separable representation of preferences, even if an arbitrary age dependence is permitted.

2.3 Induced Preferences Over Consumption

Economists are typically interested in preferences over consumption rather than births. The simplest transition from preferences over births to preferences over consumption is made by assuming that births are a function of consumption, where preferences over consumption are those induced by the underlying preferences over births.

Consider for simplicity the case in which age- τ births depend only on age- τ consumption. Let $f_\tau(c_\tau)$ give age- τ births as a function of period- τ consumption c_τ . Suppose that all the f_τ are strictly increasing and concave.

For any consumption vector c , an indifference curve is defined by (from (1), replacing $e^\delta \lambda$ with θ)

$$(2) \quad 1 = \frac{f_1(c_1)}{\theta} + \dots + \frac{f_\tau(c_\tau)}{\theta^\tau} + \dots + \frac{f(c_{\ell-1})}{\theta^{\ell-1}} + \frac{f_\ell(c_\ell)}{\theta^\ell},$$

where $\theta > 0$ is constant on a particular indifference surface. A higher value of θ indicates a higher indifference curve, so that consumption plan (c'_1, \dots, c'_ℓ) is preferred to (c_1, \dots, c_ℓ) if and only if

$$1 = \frac{f_1(c_1)}{\theta} + \dots + \frac{f_\ell(c_\ell)}{\theta^\ell} < \frac{f_1(c'_1)}{\theta} + \dots + \frac{f_\ell(c'_\ell)}{\theta^\ell}.$$

It follows readily that evolution's indifference surfaces over consumption bundles (c_1, \dots, c_ℓ) have the usual shape, in the sense that evolution's preferences can be described by a utility function $\theta(c_1, \dots, c_\ell)$ that is strictly increasing and quasi-concave.

2.4 Present Bias

As long as the evolution has a preference for early births, evolution's criterion generates a decreasing rate of impatience with age. We illustrate this by considering a type that lives to age three with fertility/utility f_1 , f_2 and f_3 in periods 1, 2 and 3. Consider the possibilities of augmenting fertility/utility by η_1 in the first period and augmenting it by η_2 in the second period. Suppose there is indifference between these alternatives, so that, for some $\theta > 1$ (cf. (2)),

$$1 = \frac{f_1 + \eta_1}{\theta} + \frac{f_2}{\theta^2} + \frac{f_3}{\theta^3} = \frac{f_1}{\theta} + \frac{f_2 + \eta_2}{\theta^2} + \frac{f_3}{\theta^3}$$

and hence $\theta\eta_1 = \eta_2$. Now consider the possibilities of augmenting fertility/utility by η_1 in the second period and augmenting it by η_2 in the third. Let the first option lie on an indifference surface characterized by θ' , so that

$$1 = \frac{f_1}{\theta'} + \frac{f_2 + \eta_1}{(\theta')^2} + \frac{f_3}{(\theta')^3}.$$

As long as $\eta_1 > 0$, we have $\theta' < \theta$ (i.e., a pure postponement of births is disadvantageous when the population is growing). Hence, we have $\theta'\eta_1 < \eta_2$, implying

$$1 = \frac{f_1}{\theta'} + \frac{f_2 + \eta_1}{(\theta')^2} + \frac{f_3}{(\theta')^3} < \frac{f_1}{\theta} + \frac{f_2}{\theta^2} + \frac{f_3 + \eta_2}{\theta^3}.$$

Evolution is thus indifferent when deferring the increment in fertility/utility from age 1 to 2, but strictly prefers to defer from period 2 to 3, i.e., evolution's preferences exhibit a present bias. In essence, deferring increments in fertility/utility lowers the associated growth rate, leading to less impatience.

Evolution's present bias does not lead to preference reversals here, in contrast to the models of present bias that form the heart of behavioral economics. Preferences are defined over age, rather than over time relative to the present. Suppose, for example, that

$$1 = \frac{f_1}{\theta'} + \frac{f_2 + \eta_1}{(\theta')^2} + \frac{f_3}{(\theta')^3} < \frac{f_1}{\theta} + \frac{f_2}{\theta^2} + \frac{f_3 + \eta_2}{\theta^3}$$

as before, but we break the indifference in the first choice to give

$$1 = \frac{f_1 + \eta_1}{\theta} + \frac{f_2}{\theta^2} + \frac{f_3}{\theta^3} > \frac{f_1}{\theta} + \frac{f_2 + \eta_2}{\theta^2} + \frac{f_3}{\theta^3}.$$

Hence the individual would choose $(f_1, f_2, f_3 + \eta_2)$ over $(f_1, f_2 + \eta_1, f_3)$ if offered this choice at age 0. If again offered this choice at age 1, having already obtained utility/fertility f_1 , the individual will reiterate her original choice. That is, the growth rate that is generated by $(f_1, f_2, f_3 + \eta_2)$ exceeds that generated by $(f_1, f_2 + \eta_1, f_3)$ even if this calculation is redone at age 1.

2.5 Exponential Discounting

Agents can once again be induced to make optimal choices via exponentially discounting offspring at the optimal growth rate. Letting (f_1, \dots, f_ℓ) be the optimal fertility/utility profile and $\theta = e^{\delta \tilde{\lambda}}$, where $\tilde{\lambda}$ is the leading eigenvalue of the corresponding Leslie matrix, (2) gives

$$1 = \frac{f_1}{\theta} + \frac{f_2}{\theta^2} + \dots + \frac{f_\ell}{\theta^\ell}.$$

Now suppose an alternative fertility/utility profile (f'_1, \dots, f'_ℓ) gives a smaller growth rate.

Then

$$\frac{f_1}{\theta} + \frac{f_2}{\theta^2} + \dots + \frac{f_\ell}{\theta^\ell} = 1 > \frac{f'_1}{\theta} + \frac{f'_2}{\theta^2} + \dots + \frac{f'_\ell}{\theta^\ell}.$$

The agent can thus be induced to make the optimal decision by exponentially discounted preferences. In a particular richer model, in which different agents would “choose” their life histories from random feasible sets determined idiosyncratically, exponential discounting would be the only parsimonious way to describe their preferences (cf. Emil Iantchev and Balazs Szentes (2005)).

2.6 Discussion

The evolutionary criterion for maximizing a population's growth rate entails a present bias. If we are interested in which of a collection of possible life histories will dominate a population or why one mutant is likely to be more successful than another, then we must take this present bias into account. At the same time, once we have identified the optimal life history and the attendant growth rate, the behavior induced by this life history can be described by exponential discounting at the sum of that growth rate and the death rate.

Peter Sozou (1998) and Partha Dasgupta and Eric Maskin (2005) show that if future consumption may disappear before it can be realized, and if this possibility is not explicitly recognized by the agent, then it may be evolutionarily advantageous to compensate with a present bias in the agent's discounting. More generally, if evolution embeds aspects of the feasible set into our preferences, there is little reason to expect consistent discounting.

We have implicitly maintained throughout the standard assumption that risks are independent across agents. Our continuing research explores the implications of environmental fluctuations that induce correlated risks.

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Footnotes:

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¹ More precisely, the Perron-Frobenius Theorem implies that the population vector $N(t)$ satisfies $N^T(t)/\tilde{\lambda}^t \rightarrow N^T(0)wv^T$, as $t \rightarrow \infty$, where v^T and w are the strictly positive left (row) and right (column) eigenvectors of the Leslie matrix associated with $\tilde{\lambda} > 0$, and $v^T w = 1$. If $\sum_k v_k = 1$, then v describes the limiting proportion of each age and w gives the reproductive value of an individual of each age in the initial population.

Figure 1:

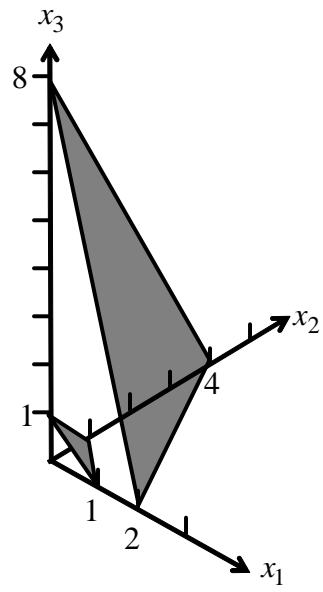


Figure 1 caption:

Illustration of evolution's indifference surface when $\ell = 3$. Evolution is indifferent over the three semelparous strategies $(x, 1)$, $(x^2, 2)$ and $(x^3, 3)$. Indifference surfaces are hyperplanes containing such equivalent semelparous strategies.