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Intertemporal choice in lemurs

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Abstract

Different species vary in their ability to wait for delayed rewards in intertemporal choice tasks. Models of rate maximization account for part of this variation, but other factors such as social structure and feeding ecology seem to underly some species differences. Though studies have evaluated intertemporal choice in several primate species, including Old World monkeys, New World monkeys, and apes, prosimians have not been tested. This study investigated intertemporal choices in three species of lemur (black-and-white ruffed lemur, \textit{Varecia variegata}, red ruffed lemur, \textit{Varecia rubra}, and black lemur, \textit{Eulemur macaco}) to assess how they compare to other primate species and whether their choices are consistent with rate maximization. We offered lemurs a choice between two food items available immediately and six food items available after a delay. We found that by adjusting the delay to the larger reward, the lemurs were indifferent between the two options at a mean delay of 17 s, ranging from 9-25 s. These data are comparable to data collected from common marmosets (\textit{Callithrix jacchus}). The lemur data were not consistent with models of rate maximization. The addition of lemurs to the list of species tested in these tasks will help uncover the role of life history and socio-ecological factors influencing intertemporal choices.

Keywords: intertemporal choice, primate, prosimian, rate maximization, temporal discounting

1. Introduction

The black-and-white ruffed lemur dangles by her hind legs at the top of the leafy canopy, plucking figs from the branch below. At some point she has consumed many of the figs from the branch, leaving a few small fruits hidden beneath the leaves. The lemur now faces a choice: should she continue to search for the remaining figs or move on to another branch full of fruit? These questions of when to leave a patch (Charnov, 1976; Stephens and Krebs, 1986) are an example of an intertemporal choice—a choice between options whose benefits are available at different points in time. Historically, most experimental studies of intertemporal choices in animals tested primarily pigeons and rats (McDiarmid and Rilling, 1965; Richards et al., 1997). More recently, however, researchers have tested many more species, including insects, fish, birds, mice, and primates (Bateson and Kacelnik, 1996; Tobin et al., 1996; Brunner and Hen, 1997; Cheng et al., 2002; Stevens et al., 2005a; Rosati et al., 2007; Pearson et al., 2010; Vick et al., 2010; Mülhoff et al., 2011).

Many studies of intertemporal choice in animals assume that they temporally discount the options (Mazur, 1987; Green and Myerson, 2004). Temporal discounting occurs when an individual assigns a subjective present value to a future reward, and that value decreases as the time delay to receiving that reward increases. In the hyperbolic discounting model (Mazur, 1987), value decreases with the reciprocal of time:

\[ V = \frac{A}{1 + kd} \]  

where \( A \) represents the amount of the food reward, \( d \) represents the time delay to receive the reward, and \( k \) represents a discounting parameter. This hyperbolic
model describes the intertemporal choices of animals quite well (Green and Myerson, 2004). Most of the animal studies, however, offer subjects repeated instances of the same choice. An alternative to the discounting account proposes that, rather than generating a subjective value, subjects simply use reinforcement rate (Skinner, 1938) or food intake rate (Stephens and Krebs, 1986) to make these decisions (Kacelnik, 2003; Stevens and Stephens, 2009). Two different types of rates are possible, depending on which temporal components individuals use: short-term or long-term rates. Short-term rates are defined as

\[ R_{ST} = \frac{A}{d + h} \]

where \( h \) represents the handling time required to process and consume the reward. Blue jays (Cyanocitta cristata) and cotton-top tamarins (Saguinus oedipus) may use short-term rates to make intertemporal choices (Stephens and Anderson, 2001; Stevens et al., 2005a). For the long-term rate

\[ R_{LT} = \frac{A}{d + h + ITI} \]

the time intervals after choice and consumption (here referring to the intertrial interval or \( ITI \)) are also included in the rate calculation. Though early work in pigeons suggested that the intertrial interval played little role in their choices (Mazur, 1994), rhesus macaques (Macaca mulatta) use post-choice information when it is made more salient to them (Pearson et al., 2010). Moreover, intertemporal choices in bonobos (Pan paniscus) are consistent with using long-term rates (Rosati et al., 2007). Once individuals estimate a rate for each option, they must then apply a decision rule such as matching (distributing responses to options in proportion to their relative rates—e.g., an option with twice the reward rate will receive twice as many responses, Herrnstein, 1961) or maximizing (always choosing the option that offers the highest rate, Stephens and Krebs, 1986). Combining the valuation or rate with a decision rule results in a choice.

Researchers have investigated intertemporal choice in several primate species using a variety of methodologies (Beran et al., 1999; Ramseyer et al., 2006; Pearson et al., 2010). Tobin et al. (1996), Stevens et al. (2005a), Rosati et al. (2007), and Addessi et al. (2011) have used the same adjusting-delay ‘self-control’ paradigm to test intertemporal choices in Old World monkeys, New World monkeys, and apes. In this task, subjects make binary choices between two immediately available food items and six food items available after a time delay. By adjusting the delay to the larger reward until subjects choose equally between the two options, researchers can establish an indifference point for each subject. To our knowledge, however, prosimian species have not been tested in an intertemporal choice task. The aim of this study was to test lemurs in this task, thereby adding the more phylogenetically basal prosimian group to the comparative data. Lemurs are good species to test because we already have data on their numerical discrimination (Lewis et al., 2005; Santos et al., 2005), inhibitory control (Genty et al., 2004), and risk sensitivity (MacLean et al., 2011). Therefore, this study will contribute to our emerging understanding of prosimian cognition and decision making. Our study examined two research questions:

1. How does lemur intertemporal choice compare to other species?
2. Do lemurs maximize their intake rate?

To explore these questions, we tested several species of lemurs in an adjusting-delay intertemporal choice procedure.

2. Methods

2.1. Subjects

In this experiment, we tested twelve lemurs of four different species at the Tierpark Berlin: four black-and-white ruffed lemurs (Varecia variegata), two red ruffed lemurs (Varecia rubra), four red-bellied lemurs (Eulemur rubriventer) and two black lemurs (Eulemur macaco). Only five lemurs (three black-and-white ruffed, one red ruffed, and one black lemur) completed the experiment. The remaining subjects failed to pass the number discrimination phase (\( N = 2 \)), stopped responding to the choices in the titration phase (\( N = 4 \)), or were moved into inaccessible facilities (\( N = 1 \)). We tested the ruffed lemurs from Nov 2007 to Dec 2008 and the black lemur from May to July 2009.

None of the lemurs had prior experience with experiments. Zookeepers fed them in the early morning approximately three hours prior to our testing and in the afternoon approximately one hour after our testing. The lemurs had constant access to water.

2.2. Materials

For testing, we separated pair-housed individuals. The testing apparatus (Figure 1) attached to openings of the cages that were used for feeding. The apparatus consisted of a platform that was affixed horizontally to the grating outside the cage. An opaque box covered...
the top and sides of the platform. The box was open on the experimenter’s side but covered by a transparent Plexiglas barrier with two holes on the subject’s side. In addition, the experimenter inserted two opaque barriers into the box near the Plexiglas barrier to prevent physical and visual access to the contents of the box. Rewards were placed on two slides on either side of the box. To choose a reward, subjects reached through one of the holes. The experimenter dispensed the chosen rewards by pushing the respective slide towards the subject. We used apple pieces (about 1.5 g) as rewards because they were a preferred food item and did not interfere with the subjects’ diet. The experimenter used a stopwatch to measure the time intervals and videotaped all trials for behavioral analysis.

Figure 1: Experimental apparatus (experimenter view). The experimental platform attached to the subject’s home cage and consisted of an opaque cover (grey), two opaque barriers (black), and two slides. The barriers prevented visual access to the rewards during the intertrial interval. Once the experimenter removed the barriers, the subject could view the options and reach through one of the two holes to make a choice. After the appropriate delay, the experimenter slid the chosen reward into the subject’s reach and removed the unchosen reward.

2.3. Experimental phases

All tests occurred between 1100 and 1300 h about five days a week, with one session per subject per day. Subjects experienced three experimental phases: training, number discrimination, and titration. In each phase, subjects experienced a daily session of 14 trials. Within a session, intertrial intervals (ITIs) of 30 seconds separated trials. During the ITI, the placement of the opaque barriers prevented the subject from seeing inside the box. The reward slides remained pulled back towards the experimenter. The experimenter stood beside the cage with her back to the cage during this period. After 20 s, the experimenter placed the rewards for the upcoming trial on the two reward slides out of sight of the subject. Reward side assignments were randomized across trials. Once the ITI elapsed, the experimenter pushed the slides forward and removed both opaque barriers simultaneously. Subjects had 15 s to examine the options and choose by reaching into one of the two holes. If they failed to choose an option within 15 s, the experimenter recorded the trial as invalid and replaced the barriers. Once subjects made a choice, the experimenter removed the unchosen reward and pushed the chosen reward slide forward into the subject’s reach after the programmed delay elapsed. When the subject had removed all food pieces from the slide (food dropped on the floor was regarded as consumed), the next ITI began.

2.3.1. Training phase

The training phase familiarized the subjects to the general testing procedure and trained them to examine their options and make a choice. In the training phase, subjects experienced only forced-choice trials—that is, subjects chose between either zero versus two rewards or zero versus six rewards, with counterbalanced order of trial types and side assignments. In this phase, subjects received the rewards immediately after choosing an option, and thereby learned about the different reward quantities available. Subjects had to make a choice in 12 of the 14 trials for the session to be considered valid, and they had to repeat invalid sessions. To enter the next experimental phase, subjects needed to choose correctly in 12 of 14 trials for three consecutive sessions. In this phase, we scored a correct choice as reaching only into the chamber that contained the reward (either two or six food pieces).

2.3.2. Number discrimination phase

In the number discrimination phase, we wanted to ensure that the subjects discriminated between the two reward amounts. We offered them the choice between the smaller reward (two apple pieces) and the larger reward (six apple pieces), with side assignments counterbalanced within a session. Subjects received the rewards immediately after choosing an option. The two versus six comparison is within the lemurs’ numerical discrimination abilities (Lewis et al., 2005). As in the training phase, subjects needed to choose correctly in 12 of 14 trials for three consecutive sessions to enter the next experimental phase. Subjects chose correctly when reaching for the larger reward.

2.3.3. Titration phase

In the titration phase, we investigated how long the lemurs would wait for the larger option. We based the
subjects became indifferent between the two options when the mean delay of the last five sessions did not differ from the mean delay of the preceding five sessions by more than 10% (Stevens et al., 2005a; Rosati et al., 2007). Subjects stopped the experiment after achieving indifference. We used the mean delay of the last five sessions for each subject as the primary dependent variable, the mean adjusted delay to indifference.

In addition to choice behavior, we measured temporal variables of choice, including the latency to retrieve the food and the handling time. We measured retrieval latency from the time the experimenter pushed the chosen slide forward until the subject placed the first piece of food in his/her mouth. We measured handling time as the time between the subject placing the first and last piece of food into his/her mouth. We randomly selected six sessions for each subject and measured the retrieval latency and handling time in all four forced-choice trials in each session. If these data were not available because the subject did not choose on the trial or we could not assess the appropriate start and stop times from the video, we randomly selected other sessions until we had temporal data in 24 trials (12 for the smaller option and 12 for the larger option).

For a comparative analysis, we used the mean adjusted delay from 12 animal species. Seven species were tested in a manner similar to this experiment. For these species, the subjects chose between two food items available immediately and six food items after an adjusting delay. In addition to the lemur, this included cotton-top tamarins (*Saguinus oedipus*) and common marmosets (*Callithrix jacchus*, Stevens et al., 2005a), long-tailed macaques (*Macaca fascicularis*, Tobin et al., 1996), brown capuchins (*Cebus apella*, Addessi et al., 2011), and bonobos (*Pan paniscus*) and chimpanzees (*Pan troglodytes*, Rosati et al., 2007). In addition, seven species were tested with slightly different methodologies. Amici et al. (2008) tested brown capuchins (some of the same subjects as Addessi et al., 2011), black-handed spider monkeys (*Ateles geoffroyi*), long-tailed macaques, lowland gorillas (*Gorilla gorilla*), and orangutans (*Pongo pygmaeus*), using the same techniques as used here except the subjects chose between one and three food items. Perry et al. (2004) tested *Rattus norvegicus* also with one versus three food items. Finally, Green et al. (2007) tested White Carneau pigeons (*Columba livia*) with an adjusting-delay procedure in which the subjects chose between 5 immediate versus 30 delayed food items, and the adjustments occurred within a session. We estimated the pigeons’ mean adjusted delay to the larger reward by substituting each subject’s discounting parameter $k$ into a set of hyperbolic equations (equation 1) and solving for the delay to larger reward. Body mass for each species was taken from Kangas and Branch (2006) for pigeons, Perry et al. (2004) for rats, and Isler et al. (2008) for all other primate species.

We analyzed the data using R statistical software version 2.13.2 (R Development Core Team, 2011) and the ape version 2.7-3 (Paradis et al., 2004), beetswarm version 0.1.1 (Eklund, 2010), and xtable version 1.5-6 (Dahl, 2009) packages. Data and R code are available in the Supplementary Materials (Data S1-S3), and data are available at the Dryad data repository (doi:10.5061/dryad.fb778jf2). The original Sweave-embedded R code (Leisch, 2002) to allow re-
production of analyses (de Leeuw, 2001), is available from JRS.

3. Results

Subjects required between 6-38 sessions (mean = 14.8, median = 9) to pass the number discrimination phase. In the titration phase, the mean adjusted delay at indifference observed for the lemurs tested in this study ranged from 9-25 s, with a mean of 17 s. Table 1 shows the mean adjusted delay for the individual subjects. There do not appear to be strong differences between lemur species (the minimum and maximum delays both come from the black-and-white ruffed lemurs), but we tested too few subjects to make proper between-species comparisons. Thus, we pool all of the lemur species in the subsequent comparative analyses.

Previous studies with cotton-top tamarins and bonobos suggest that their intertemporal choice patterns are consistent with short-term and long-term rate maximization, respectively (Stevens et al., 2005a; Rosati et al., 2007). To investigate whether lemur intertemporal choice is consistent with maximizing intake rate, we measured individual response latencies and handling times for both the smaller and larger rewards (Table 1). The inclusion or exclusion of retrieval latency did not influence the results, so all analyses added retrieval latency to the handling time. If lemurs maximize their short-term intake rate, they should choose the option with the higher rate (Equation 2). We can solve the equation \( \frac{A_i}{d + h} = \frac{A_l}{d + h} \) (where the subscripts s and l refer to the smaller and larger option, respectively) for \( d_l \) to generate a predicted mean adjusted delay at indifference for each subject. The mean short-term rate maximization prediction for the adjusted is -4.9 s (range -13.09 to 0.24 s), clearly not matching the observed mean of 17.0 s. The long-term rate (Equation 3) includes the intertrial interval (ITI) of 30 s. When solving \( \frac{d_i}{d + h + ITI} = \frac{d_l}{d + h + ITI} \), the mean long-term rate maximization prediction for the adjusted delay is 55.1 s (range 46.91 to 60.24 s), again not matching the observed adjusted delay.

4. Discussion

Using an adjusting-delay procedure in three species of lemur, we measured the mean adjusted delay to indifference such that subjects were indifferent between two food items received immediately and six food items received after a delay. The observed mean adjusted delays ranged from 9-25 s. Relative to other species tested under similar circumstances (Figure 2A), these adjusted delays are most similar to those of common marmosets (Stevens et al., 2005a).

Across primates, indifference point times vary by more than an order of magnitude from cotton-top tamarins to chimpanzees. What can account for this variation across species? One possibility is that animals simply maximize their rate of food intake. There are two ways in which individuals or species may have different indifference points when rate maximizing. First, they may vary in the time it takes to handle and consume food. Second, they may include different temporal components in the rate estimation (Mazur, 1994; Rosati et al., 2006; Stevens and Stephens, 2009; Pearson et al., 2010).

Optimal foraging theory (Stephens and Krebs, 1986) assumes that animals maximize their long-term rate, so they should include all temporal components in their rate. In our study, that includes the delay, handling time, and intertrial interval. Animals may, however, ignore the intertrial interval and focus on temporal components that occur only between choice and consumption, resulting in short-term rate maximization (Stephens and Anderson, 2001). Though consistent

![Figure 2: Intertemporal choice across animal species. (A) Seven species have been tested using the same adjusting-delay procedure with two versus six food items: lemurs (this study), cotton-top tamarins and common marmosets (Stevens et al., 2005a), black capuchins (Addessi et al., 2011), long-tailed macaques (Tobin et al., 1996), and bonobos and chimpanzees (Rosati et al., 2007). The mean adjusted delay at indifference for lemurs matched that found in common marmosets. (B) Other studies used similar but not identical adjusting-delay procedures for seven species: White Carneau pigeons (Green et al., 2007), Wistar rats (Perry et al., 2004), and brown capuchins, black-handed spider monkeys, long-tailed macaques, lowland gorillas, and orangutans (Amici et al., 2008). The capuchins and macaques are the same species in both panels, and some of the capuchin data are from the same individuals in both panels. In this figure, circles represent data points for individual subjects, triangles represent the mean, lines represent the median, boxes represent the interquartile range (25-75%), and whiskers represent the range.](image-url)
with intertemporal choices in other primates (Stevens et al., 2005a; Rosati et al., 2007), our analysis here indicates that neither short- nor long-term rate maximization models accounted for the intertemporal choice pattern observed in lemurs.

A number of other possible factors may influence intertemporal choice, ranging from metabolic rate and life expectancy to social structure and foraging ecology (reviewed in Stevens and Stephens, 2009; Addessi et al., 2011). Metabolic rate is relevant for intertemporal choice for food because the rewards directly feed into metabolism. Therefore, species with higher metabolisms need food more quickly and may opt for sooner rewards than those with lower metabolisms (Tobin and Logue, 1994). Life expectancy is also relevant because shorter life spans should make delayed rewards less beneficial, so shorter-lived species should choose sooner rewards (Stevens and Stephens, 2009). Unfortunately, we do not have good direct measures of metabolic rate and life expectancy for all of the primates analyzed here. Body size, however, offers a reasonable correlate for both metabolic rate and life expectancy: in mammals, larger species tend to have lower metabolisms and longer life expectancies (Speakman, 2010) and the pigeon and rat branch lengths calculated in their methodologies: Addessi et al. uses two versus six food items, whereas Amici et al. (2008) uses one versus three. Though the absolute magnitude of the rewards influences human intertemporal choices (e.g., 100 vs. 500 dollars is not the same as 1000 vs. 5000 dollars; Green et al., 1999), they have not been shown to influence choices in animals (Kirby and Maraković, 1996; Green et al., 2004). Moreover, Stevens et al. (2005b) directly compared one versus three and two versus six food items in a spatial discounting task with tamarins and marmosets and found no effect of the reward magnitude. Thus, we pooled these data and observe that intertemporal choice correlates with body mass.

Lemurs fall below the regression line in this analysis, suggesting that they wait for shorter times than expected by their body size. Capuchins provide an interesting comparison because, though they share similar body sizes, capuchins wait on average three times longer (almost five times longer using only the data from Addessi et al., 2011) than lemurs. Though making comparisons across phylogenetically distantly related taxa warrants caution, socio-ecological characters such as social complexity, tool use, and extractive foraging for the lemur species tested here may not be as developed as those of capuchins (Jolly, 1998; Vasey, 2003). Without these pressures, lemurs may not have evolved the ability to wait like other primate species.

Though body size correlates with metabolic rate and life expectancy, it also correlates with many other factors such as diet and territory size. Therefore, rather than using body size as a proxy for other variables, we must measure those specific variables to assess their influence on temporal preferences. Moreover, in our data, body size also correlates with phylogenetic relationship. To control for phylogenetic relationship between species (Figure S1), we conducted phylogenetically independent contrasts with weighted branch lengths (Felsenstein, 1985) based on a primate phylogeny created from 10kTrees version 3 (Arnold et al., 2010) and the pigeon and rat branch lengths calculated using TimeTree (Hedges et al., 2006). Independent contrasts suggest that body size remains an influence on intertemporal choice even after accounting for phylogeny ($F_{1,10} = 10.55, p = 0.01$, Figure S2).

Though large-scale relationships between intertemporal choice and body size and phylogeny exist across species, other factors such as social structure and for-

<table>
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<th>Subject</th>
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<th>Sex</th>
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<th>Small amount</th>
<th>Large amount</th>
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<th>Small latency</th>
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*For species, R is red ruffed lemur, BW is black-and-white ruffed lemur, and BL is black lemur.
aging ecology may also influence intertemporal choice for more closely related species. Amici et al. (2008) proposed that social structure shapes choice via fission-fusion dynamics. In species with high fission-fusion dynamics, group members enter and leave the group frequently, resulting in a very fluid social structure. Amici et al. suggested that this turnover may select for more inhibitory control to suppress “prepotent but ineffectual responses in a changing social environment” (p. 1415). That is, fission-fusion species require flexibility in inhibiting their social behavior and in turn may have more control over inhibitory processes in the food domain as well. When they tested the primate species in Figure 2B, Amici et al. reported that fission-fusion species tend to wait longer for food rewards, thereby confirming their predictions.

Foraging ecology offers another factor relevant to intertemporal choice. Species that frequently face delays in their natural foraging ecology (such as guminvores, stalking and sit-and-wait hunters, and extractive foragers) likely face selection pressure for the ability to wait in intertemporal choices (Stevens and Stephens, 2009). The ability to wait (or not) may be “ecologically rational” (Todd and Gigerenzer, 2007) depending on the temporal structure of the environment. Foraging ecology may act as an important factor explaining why common marmosets wait longer than tamarins and chimpanzees wait longer than bonobos (Stevens et al., 2005a; Rosati et al., 2007). Moreover, Addessi et al. (2011) proposed that the extractive foraging used by capuchins explains their long wait times in intertemporal choice tasks compared to other New World monkeys. Thus, the social and foraging environments likely exert strong selection pressure for intertemporal choices. Other, more internal, factors such as differential valuation of rewards and time perception may also influence choice. The factors mentioned here are not mutually exclusive, however, and likely combine to shape intertemporal choice.

4.1. Conclusions

In summary, lemurs’ intertemporal choices match those of the callitrichid primates tested with similar methodologies. They do not appear to maximize their food intake rate in this task. A comparative analysis with 12 species suggests that intertemporal choices correlate with body size, even when accounting for phylogeny.

In this study, we tested too few species to make general claims about lemurs broadly because lemur species differ greatly in their social and ecological characters. For instance, in a comparative analysis of 19 species, MacLean et al. (2009) explored how pairbonding, group size, diet, and activity pattern relate to brain size in lemurs. Though the social factors did not relate to brain size in their analysis, the ecological factors did: frugivores had larger brains that foliovores and cathemeral species (those active both during day and night) had larger brains than diurnal species. The lemur species tested here are primarily frugivorous and diurnal. Comparisons with foliovorous and cathemeral species might result in interesting differences in waiting times. Despite the overall lack of an influence of group size on brain size in MacLean et al. (2009), studies of specific cognitive tasks do show differences in performance. For instance, the more highly social ring-tailed lemurs (Lemur catta) outperformed less social species such as mongoose lemurs (Eulemur mongoz), black lemurs, and ruffed lemurs in transitive inference and social cognition tasks (MacLean et al., 2008; Sandel et al., 2011). Lemur species vary widely in life history, ecology, and social structure, ranging from rather solitary, specialized extractive foragers such as aye-ayes (Daubentonia madagascariensis) to the fission-fusion and frugivorous ruffed lemurs (Morland, 1991; Vasey, 2006). This broad diversity among species makes lemurs a particularly interesting group for testing comparative hypotheses regarding factors influencing intertemporal choice.
Acknowledgments

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References


Supplementary Material

**Movie S1.** Intertemporal choice test. Video recording of black lemur free-choice trials. In the first clip, the subject chose six apple pieces after 13 s and in the second clip she chose two apple pieces available immediately.

**Data sheet S1.** Lemur intertemporal choice data.

**Data sheet S2.** Comparative intertemporal choice data. Individual subject intertemporal choice data for 12 animal species.

**Data sheet S3.** R code for data analysis.

**Figure S1.** Phylogeny of species in comparative analysis.

**Figure S2.** Independent contrasts of body mass and intertemporal choice.

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**Figure S1:** Phylogeny of species in comparative analysis. We used 10kTrees version 3 (http://10ktrees.fas.harvard.edu/index.html, Arnold et al., 2010) to construct the weighted branch lengths of the primate phylogeny and TreeTime (http://www.timetree.org/, Hedges et al., 2006) for the weighted branch lengths of pigeons and rats.

**Figure S2:** Independent contrasts of log body mass and intertemporal choice. When accounting for phylogeny, the relationship between intertemporal choice and log body size is maintained ($y = 0.35x$, $F_{1,10} = 10.55$, $p = 0.01$, adjusted $R^2 = 0.46$).