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1 RESEARCH ARTICLE (*REVISED CLEAN*)

2 **No evidence of tradeoffs in the evolution of sperm numbers and**
3 **sperm size in mammals**

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21 **Abstract**

22 Postcopulatory sexual selection, in the form sperm competition, has influenced the evolution
23 of several male reproductive traits. However, theory predicts that sperm competition would
24 lead to tradeoffs between numbers and size of spermatozoa because increased costs per cell
25 would result in a reduction of sperm number if both traits share the same energetic budget.
26 Theoretical models have proposed that, in large animals, increased sperm size would have
27 minimal fitness advantage compared with increased sperm numbers. Thus, sperm numbers
28 would evolve more rapidly than sperm size under sperm competition pressure. We tested in
29 mammals whether sperm competition maximizes sperm numbers and size, and whether there
30 is a tradeoff between these traits. Our results showed that sperm competition maximizes
31 sperm numbers in eutherian and metatherian mammals. There was no evidence of a tradeoff
32 between sperm numbers and sperm size in any of the two mammalian clades since we did not
33 observe any significant relationship between sperm numbers and sperm size once the effect of
34 sperm competition was taken into account. Maximization of both numbers and size in
35 mammals may occur because each trait is crucial at different stages in sperm's life; e.g., size-
36 determined sperm velocity is a key determinant of fertilization success. In addition, numbers
37 and size may also be influenced by diverse energetic budgets required at different stages of
38 sperm formation.

39 **Keywords:** sexual selection, sperm competition, metabolic rate, energetic constraints, sperm
40 size, sperm numbers

41

42 **Introduction**

43 One of the key predictions of sperm competition theory (Parker, 1970) is that males belonging
44 to species with high levels of sperm competition should produce more sperm in order to
45 outcompete their rivals in their quest to fertilize ova. If it is assumed that sperm competition
46 resembles a raffle, in which sperm are equivalent to fertilizing opportunity increments, more
47 sperm would increase the chances of fertilization (Parker, 1970; 1993). In line with this
48 "raffle hypothesis", high levels of sperm competition associate with increases in testes mass
49 relative to body size (Harcourt *et al.*, 1981; Gage, 1994; Birkhead & Møller, 1998; Simmons,
50 2001; Birkhead *et al.*, 2009; Soulsbury, 2010) and with high relative sperm numbers
51 (Birkhead & Møller, 1998; Parker & Pizzari, 2010; Gomez Montoto *et al.*, 2011a) in many
52 taxa.

53 Since the production of a vast amount of sperm would be highly expensive in terms of
54 energy, and larger sperm may increase the energetic cost per cell, a "direct" energetic tradeoff
55 would exist between sperm numbers and sperm size (Parker, 1982). This line of reasoning
56 assumed that total energetic budget for sperm production equaled the number of sperm
57 produced multiplied by the energetic cost of producing each sperm. Consequently, in order to
58 produce more sperm there should be a reduction in the cost of producing each individual cell
59 (i.e., via a reduction in sperm size).

60 Nevertheless, sperm size could provide an adaptive advantage. If sperm competition
61 resembles a race in which the first sperm to reach the ovum engages in fertilization, an
62 increase in sperm size (e.g., due to a longer flagellum) could provide increased sperm
63 velocity, thus affecting the outcome of sperm competition (Gomendio & Roldan, 1991). This
64 scenario adds a temporal component to the raffle hypothesis. Evidence supporting positive
65 relationships between sperm competition levels and sperm size (Gage, 1994; Briskie *et al.*,
66 1997; Tourmente *et al.*, 2009; Tourmente *et al.*, 2011b; a), sperm size and sperm velocity

67 (Gomendio & Roldan, 2008; Fitzpatrick *et al.*, 2009; Lüpold *et al.*, 2009; Gomez Montoto *et*
68 *al.*, 2011b; Tourmente *et al.*, 2011b), and sperm competition and sperm velocity (Fitzpatrick
69 *et al.*, 2009; Kleven *et al.*, 2009; Gomez Montoto *et al.*, 2011b; Tourmente *et al.*, 2011b) has
70 been found in many taxa, including mammals. Therefore, since faster sperm increase
71 fertilization success in competitive scenarios (Birkhead *et al.*, 1999; Gage *et al.*, 2004;
72 Gasparini *et al.*, 2010), males in species with high levels of sperm competition should
73 maximize both sperm numbers and sperm size to increase their fitness.

74 Subsequent theoretical models developed the concept of an "indirect tradeoff" by
75 placing the emphasis on total reproductive budget rather than just on sperm production
76 (Parker, 1993). This tradeoff definition was expanded to incorporate a new partition of the
77 total reproductive energetic budget between mate acquisition and total ejaculate investment
78 (the latter encompassing sperm numbers and sperm size) (Parker, 1993; Parker *et al.*, 2010).
79 Nevertheless, this model is still based on the idea of a direct tradeoff between sperm numbers
80 and sperm size within the ejaculate investment component, for a given value of sperm
81 competition risk. In addition, two sperm competition mechanisms have been defined on the
82 basis of the volume scale differences between the ejaculate and the female reproductive tract.
83 On the one hand, there is a raffle mechanism in which female reproductive tract volume and
84 ejaculate volume scales differ considerably and there is no space constraint for fertilization
85 (Parker, 1990). This mechanism describes conditions for externally fertilizing species and
86 many vertebrates with internal fertilization, especially those with large bodies. On the other
87 hand, a displacement mechanism, in which the volumes of the ejaculate and the female sperm
88 storage organs are similar and, thus, successive ejaculates will volumetrically displace the
89 preceding ones (Parker & Simmons, 1991). This mechanism describes the conditions
90 prevalent in insects, in which females have sperm stores that can hold a small volume of
91 ejaculate.

92 An early theoretical model on sperm size proposed that the effect of sperm size on
93 sperm competition is mainly mediated by sperm density (Parker, 1993). This model predicted
94 that, if the effect of sperm density in the female tract on sperm competitiveness is negligible,
95 there would be a single optimal size for sperm, independently of the sperm competition level.
96 In a later paper (Parker *et al.*, 2010), the probable effects of sperm density on sperm
97 competitiveness were investigated in detail. Since sperm numbers ejaculated (and hence the
98 density of competing sperm) often change with sperm competition level, this study (Parker *et al.*
99 *et al.*, 2010) examined how the balance between sperm size and number is expected to change
100 with sperm competition. Applying the indirect tradeoff hypothesis to sperm competition
101 mechanisms, theoretical studies proposed that (a) a tradeoff between sperm numbers and
102 sperm size would occur; (b) fitness advantages related to sperm size would be mediated by
103 "sperm density" (i.e., sperm numbers in relation to the space available for sperm competition);
104 and (c) in the raffle models, an increase in sperm numbers would generally yield greater
105 fitness gain than an increase in sperm size since sperm density is considered to be negligible
106 in this mechanism due to the scaling difference between ejaculate volume and female
107 reproductive tract volume (Parker *et al.*, 2010). Thus, in "raffle" models, the effect of sperm
108 size on sperm competitiveness is assumed as a weak force "reduced mainly to effects on
109 sperm motility and survival" (Immler *et al.*, 2011). A comparative study presented empirical
110 support for these predictions by comparing the relationships between sperm numbers, sperm
111 size and sperm competition risk in two taxa with different sperm competition mechanisms
112 (birds: raffle; flies: displacement) (Immler *et al.*, 2011).

113 In mammals, sperm competition is associated with increases in both sperm numbers
114 (Møller, 1989; Gomendio *et al.*, 1998; Gomez Montoto *et al.*, 2011a; Lüpold, 2013), and
115 sperm size (Tourmente *et al.*, 2011b; a). However, the existence of a positive association
116 between both sperm size and number with the risk of sperm competition does not allow any

117 predictions about the existence of a tradeoff between these two traits, since the investment in
118 one of them might be higher in relation to the investment in the other (Parker *et al.*, 2010).
119 Mammals and birds share a number of similarities regarding the reproductive traits which
120 were taken in account by the models cited above. Firstly, volume of the female reproductive
121 tract is considerably higher than the volume of the ejaculate (Brillard & Bakst, 1990; Suarez
122 & Pacey, 2006). Secondly, sperm swimming velocity is a main factor in fertility (Birkhead *et al.*
123 *et al.*, 1999; Malo *et al.*, 2005), is positively related to sperm size (Lüpold *et al.*, 2009; Gomez
124 Montoto *et al.*, 2011b; Tourmente *et al.*, 2011b) and is positively selected by sperm
125 competition (Kleven *et al.*, 2009; Gomez Montoto *et al.*, 2011b; Tourmente *et al.*, 2011b; a;
126 Tourmente *et al.*, 2013). Thirdly, immotile sperm are not transported towards the egg despite
127 passive sperm transport to the site of fertilization (Froman & Kirby, 2005; Suarez & Pacey,
128 2006). Therefore, according to current theoretical models, the existence of a tradeoff between
129 sperm numbers and size in mammals would be anticipated.

130 In this study, we analyzed the relationships between sperm numbers and sperm size in
131 mammals by comparing two sister mammalian clades, Eutheria and Metatheria. We chose to
132 compare these two groups because they present clear differences in sperm numbers, sperm
133 size, and reproductive physiology, which could strongly influence responses of sperm
134 phenotype to sperm competition. In general terms, Metatheria (i.e., marsupials) have lower
135 relative testes mass (Tourmente *et al.*, 2011a), lower sperm numbers in relation to body size
136 (Taggart *et al.*, 1998; Taggart *et al.*, 2003), longer sperm (Tourmente *et al.*, 2011a), and a
137 higher efficiency of sperm transport, with an extremely high proportion of ejaculated sperm
138 reaching the fertilization site in relation to Eutheria (i.e., the so-called placental mammals)
139 (Taggart *et al.*, 1998; Taggart *et al.*, 2003). Furthermore, in Metatheria sperm storage in the
140 female reproductive tract takes place during relatively long time periods (up to two weeks) in
141 comparison to Eutheria (with the exception of Chiroptera) (Bedford *et al.*, 1984).

142 We predicted that a tradeoff between sperm numbers and sperm size exists among
143 mammals but that, based on their reproductive physiology, there may be differences between
144 Eutheria and Metatheria with regards to the relative increase in sperm size and numbers in
145 response to sperm competition.

146

147 **Materials and Methods**

148 *Sperm numbers, sperm size, and relative testes mass*

149 Data on body mass (g), testes mass (g), total sperm length (μm), and number of sperm
150 in caudae epididymides ($\times 10^6$ sperm) were obtained from the literature for 64 species (21
151 families) of Eutheria and 14 species (7 families) of Metatheria (Table S1 for data and
152 references). Only species for which all these measures were available were included. Data
153 collection was restricted to studies on adult, healthy, reproductively mature individuals. In
154 cases of experimental studies or clinical trials, only the values of control groups were used as
155 data sources. Whenever possible, data were collected from studies in which the higher
156 number of variables analyzed were present. In those cases in which different values for the
157 same variable and species were available from different studies, averages were used to obtain
158 a representative measure. For total sperm length, only species where this parameter was
159 measured from digital images were incorporated into the dataset.

160 The measure of sperm number used in this study was the number of sperm in
161 epididymal reserves (cauda epididymides), since it is a more robust parameter of sperm
162 production than the number of sperm in the ejaculate, although both traits are strongly
163 correlated (delBarco-Trillo *et al.*, 2013). We are aware that the number of sperm in
164 epididymal reserves may not always be extremely accurate due to insufficient flushing of the
165 epididymis, seasonal or age-related variation (Johnson & Thompson, 1983), and perceived
166 risk of sperm competition during spermatogenesis (Ramm & Stockley, 2009). However, the

167 number of sperm in the ejaculate is a less direct measure of sperm production and may be
168 affected by many factors, namely number of copulations (Pierce *et al.*, 1990; Pitnick &
169 Markow, 1994; Ambriz *et al.*, 2002), number of potential partners (Sheldon, 2000), sperm
170 depletion in successive ejaculates (Preston *et al.*, 2001), differential sperm allocation through
171 perceived sperm competition risk (Dewsbury, 1982; Wedell *et al.*, 2002; delBarco-Trillo,
172 2011; Kelly & Jennions, 2011), or the effect of electrostimulation (a common procedure in
173 large mammals) in comparison to normal copulation (Mattner & Voglmayr, 1962; Aulerich *et*
174 *al.*, 1972; Schneiders *et al.*, 2004).

175 *Data analysis*

176 To test the influence of sperm competition we used relative testes size as predictor of
177 number of sperm in caudae epididymides (dependent variable). Since testes size relative to
178 body mass is a reliable indicator of investment in sperm production, this trait is considered to
179 be a very good proxy of sperm competition levels (Gage, 1994; Birkhead & Møller, 1998).
180 Experimental selection assays in insects have demonstrated that lines subjected to high sperm
181 competition environments evolve larger testes (Hosken & Ward, 2001), while lines subjected
182 to enforced monogamy evolve reduced testes size and sperm production (Pitnick *et al.*, 2001).
183 Moreover, a recent comparative study (Soulsbury, 2010) found that levels of multiple
184 paternity correlate well with relative testes size in mammals. We performed multiple
185 regression analyses introducing the following predictors in the model: body mass, testes mass,
186 a quadratic term for testes mass, and the interaction between body mass and testes mass
187 (Tomkins & Simmons, 2002; Immler *et al.*, 2011).

188 When testing the effect of sperm size on sperm numbers, we used two sets of predictor
189 variables in order to control for the effect of body size (body mass and total sperm length as
190 predictors), and sperm competition (body mass, testes mass and total sperm length as

191 predictors) on sperm numbers. All variables were \log_{10} -transformed to meet parametric
192 assumptions.

193 All regressions were performed using phylogenetic generalized least-squares analyses
194 (PGLS) (Freckleton *et al.*, 2002) since species trait values may be similar as a result of
195 phylogenetic association rather than selective evolution (Felsenstein, 1985; Harvey & Pagel,
196 1991). PGLSs incorporate phylogenetic interdependency among the data points by including
197 the phylogenetic structure within a standard linear model as a covariance matrix that assumes
198 a predetermined evolutionary model. PGLS estimates (via maximum likelihood) a
199 phylogenetic scaling parameter lambda (λ) of the tree's branch lengths that fits evolution
200 proceeding via Brownian motion. In our study, the length of all branches was set to 1. If λ
201 values are close to 0, the variables are likely to have evolved independently of phylogeny,
202 whereas λ values close to 1 indicate strong phylogenetic association of the variables.
203 Additionally, we calculated the effect size “ r ” from t -values obtained from the PGLS model
204 and the non-central confidence limits (CLs) for the z -transformed value of r (Nakagawa &
205 Cuthill, 2007). The CLs value [values?] indicate that the effect size is statistically significant
206 if 0 is not contained within the interval (Smithson, 2002).

207 All statistical analyses were performed using the CAPER v0.5 (Orme *et al.*, 2012)
208 package for R (v3.0.1; R Foundation for Statistical Computing 2013). P values were
209 considered statistically significant at $\alpha < 0.05$. Residual testes mass was calculated for each
210 group as the residual of a log-log linear regression of testes mass on body mass (Eutheria:
211 $p < 0.0001$, $R^2 = 0.87$; Metatheria: $p < 0.0001$, $R^2 = 0.89$) and used exclusively to illustrate the
212 results. The phylogenetic reconstruction used in the PGLS analyses is included in the
213 Supplementary Information (Fig. S1). This reconstruction was based on a mammalian
214 supertree (Bininda-Emonds *et al.*, 2007) for the determination of the phylogenetic position of
215 the higher groups (orders and families). Group-specific phylogenies were used to resolve the

216 within group relationships for Artiodactyla (Prothero & Foss, 2007; Agnarsson & May-
217 Collado, 2008), Carnivora (Flynn *et al.*, 2005), Soricomorpha (Dubey *et al.*, 2007),
218 Lagomorpha (Robinson & Matthee, 2005), Rodentia (Pages *et al.*, 2010; Fabre *et al.*, 2012),
219 and Metatheria (Meredith *et al.*, 2008; 2009).

220

221 **Results**

222 Data on sperm numbers, sperm size, body mass and testes mass are presented in Table
223 S1. Eutherian mammals had a mean of 8.6×10^9 spermatozoa in their sperm reserves (i.e.,
224 sperm in caudae epididymides), ranging from 2.6×10^5 spermatozoa in the Damaraland mole
225 rat (*Fukomys damarensis*) to 1.4×10^{11} spermatozoa in the sheep (*Ovis aries*) (a range of 6
226 orders of magnitude; Table S1). On the other hand, metatherian mammals showed a mean
227 value of 4.6×10^8 sperm in cauda epididymides, which ranged from 1.2×10^6 spermatozoa in
228 the fat-tailed dunnart (*Sminthopsis crassicaudata*) to 4.8×10^9 spermatozoa in the tammar
229 wallaby (*Macropus eugenii*) (a range of 2 orders of magnitude; Table S1).

230 In both Eutheria and Metatheria, sperm number exhibited a strong negative allometric
231 association with body mass (Eutheria: \log_{10} sperm numbers = $6.52 + 0.67 * \log_{10}$ body mass,
232 $F_{1,62} = 90.95$, $p < 0.0001$, $R^2 = 0.59$; Metatheria: \log_{10} sperm numbers = $5.11 + 0.81 * \log_{10}$
233 body mass, $F_{1,12} = 38.72$, $p < 0.0001$, $R^2 = 0.76$). However, the log-log equation for Metatheria
234 presents a higher slope and a lower intercept, which means that small metatherians would
235 have lower sperm numbers than their eutherian counterparts of the same body mass. To show
236 this, we calculated the "linearized mean body mass" (the antilogarithm of the mean \log_{10} -
237 transformed body mass) for each group (Eutheria= 2851.02g, Metatheria= 2299.85g) and
238 used the previously stated equations to predict the sperm numbers. Comparison of predicted
239 sperm numbers between the two clades revealed that Metatheria had roughly one order of
240 magnitude less sperm (6.66×10^7) than their Eutheria counterpart (7.02×10^8).

241 Our analyses regarding the effect of sperm competition on sperm numbers revealed
242 that in both Eutheria and Metatheria, there is a positive association between relative testes
243 size and sperm numbers in caudae epididymides (Table 1). Neither the quadratic term for
244 testes mass, nor the interaction between body mass and testes mass were significantly related
245 to sperm numbers or sperm size in any of the two clades (Table S2). We next examined
246 possible relationships between sperm size and sperm numbers. Since sperm size is negatively
247 associated with body size in both Eutheria (Gomendio *et al.*, 2011) and Metatheria
248 (Tourmente *et al.*, 2011a), as a consequence of a decrease in mass-specific metabolic rate in
249 large-bodied animals, we included body size as a controlling variable in our analyses. We
250 found that, after controlling for body size, sperm numbers in reserves and sperm size were
251 positively associated in Eutheria (Table 1, Fig. 1A) but showed no significant relationship in
252 Metatheria (Table 1, Fig. 1B).

253 According to recent theoretical models, the influence of sperm competition risk on
254 both sperm numbers and sperm size should be taken into account to identify a possible
255 tradeoff between these traits (Parker *et al.*, 2010). Thus, in order to accurately predict the
256 possible effect of one sperm trait on the other, the effect of relative testes size was included in
257 the model. When relative testes size was controlled for, no significant association between
258 sperm numbers and sperm length was observed in either Eutheria or Metatheria (Table 1, Fig.
259 1C, D).

260 Further assessments of the relationships between sperm numbers and sperm size were
261 performed by calculating two additional parameters (Parker, 1990; Parker & Simmons, 1991;
262 Parker *et al.*, 2010; Immler *et al.*, 2011): (a) total sperm investment (i.e., sperm size * sperm
263 numbers), which represents an approximation of the total energy a male spends in sperm
264 production, and (b) relative investment in sperm size (i.e., sperm size / sperm numbers),
265 which represents an approximation of the proportion between investment in sperm numbers

266 and investment in sperm size. We found that total sperm investment was positively related to
267 relative testes size in both Eutheria and Metatheria but that there were differences between
268 clades when comparing increase rates (Eutheria, slope= 4.2121, $F_{1,61}= 91.54$, $p<0.0001$;
269 Metatheria, slope= 2.9936, $F_{1,11}= 11.68$, $p=0.0057$; Table 2, Fig. 2A, B). On the other hand,
270 relative investment in sperm size showed a significant negative relationship with relative
271 testes size in Eutheria (slope= -0.0489, $F_{1,61}= 71.80$, $p<0.0001$; Table 2, Fig. 2C) but not in
272 Metatheria (slope= -0.0398, $F_{1,11}= 1.58$, $p=0.2352$, Table 2, Fig. 2D). Overall, these results
273 suggest that, in Eutheria, sperm numbers increase faster than sperm size in response to sperm
274 competition.

275

276 Discussion

277 The results of this study showed that sperm competition maximizes sperm numbers in
278 eutherian and metatherian mammals. Previous work on both clades also revealed a positive
279 relationship between sperm competition and sperm size (Tourmente *et al.*, 2011b; a). Both an
280 increase in sperm numbers and sperm size would promote a positive association between total
281 sperm investment (sperm size * sperm numbers) and sperm competition levels (Parker, 1990;
282 Parker & Simmons, 1991; Parker *et al.*, 2010; Immler *et al.*, 2011), which is supported for
283 both Eutheria and Metatheria in our analyses.

284 Theoretical models (Parker, 1990; Parker *et al.*, 2010) predict that, in large vertebrates,
285 the vast scale difference between the size of the sperm cell and that of the female reproductive
286 tract favors a "raffle" sperm competition mechanism, and that competitive advantage may be
287 gained mainly through increases in sperm numbers, which would tradeoff against sperm size.
288 Our results regarding the relationship between sperm competition and relative investment in
289 sperm size (sperm size/sperm numbers) in Eutheria suggest that, in this group, sperm numbers
290 increase more rapidly than sperm size in response to sperm competition. At first glance, these

291 trends appear to be consistent with those described for passerine birds (Immler *et al.*, 2011)
292 and seem to adjust to theoretical models (Parker *et al.*, 2010). However, we did not find any
293 significant relationship between sperm numbers and sperm size in any of the two mammalian
294 clades once the effect of sperm competition was controlled for. That is, an increase in sperm
295 size is not associated to a decrease in sperm numbers at a given level of sperm competition. In
296 addition, studies in birds showed an “inverted U-shaped” relationship between sperm
297 competition and sperm size, which led to the conclusion that little or no increase in sperm size
298 at high sperm competition risks occurred as a result of a tradeoff between sperm numbers and
299 sperm size (Parker *et al.*, 2010; Immler *et al.*, 2011). In contrast, sperm size was not
300 significantly associated with the quadratic term of testes mass, nor with the interaction term
301 between body mass and testes mass in any of the two clades analyzed. Thus, there is no
302 evidence of a deceleration in the slope of increase of sperm size in relation to sperm
303 competition in mammals. Moreover, extremes cases such as *Cricetulus griseus* (total sperm
304 length: 258 μm) and *Tarsipes rostratus* (total sperm length: 337 μm) show the longest sperm,
305 respectively, in Eutheria and Metatheria, along with extremely high relative testes size
306 (Gomendio *et al.*, 2011; Tourmente *et al.*, 2011b; a).

307 Raffle models predict that sperm numbers would have a higher rate of increment than
308 sperm size in response to sperm competition because of the differences in the relative
309 competitive advantages provided by each trait in a tradeoff scenario. Since our results found
310 no evidence of such tradeoff, we propose that a similar relationship between rates of increase
311 of these two traits could exist in mammals because their morpho-physiological reproductive
312 characteristics do not conform to certain assumptions of the general models (Parker *et al.*,
313 2010). These reproductive characteristics of mammals are discussed below.

314

315 *Sperm numbers and size may be regulated by different energetic budgets*

316 Sperm numbers and size are determined at different stages of spermatogenesis and are
317 related to separate, independent processes of sperm formation. Sperm numbers may depend
318 on testes architecture, kinetics of spermatogenesis, cell proliferation (i.e., the number of cell
319 divisions of spermatogonia) and apoptosis (i.e., cell loss) resulting in a net number of cells
320 after meiosis. These traits show considerable variations in mammals (Hess & Franca, 2008)
321 and have been shown to covary with sperm competition levels (delBarco-Trillo *et al.*, 2013).
322 Sperm size, on the other hand, depends exclusively on the post-meiotic differentiation
323 process, when spermatids give rise to sperm cells. This process also presents a high degree of
324 variability among mammals (Hess & Franca, 2008), although the possible associations
325 between sperm competition and the duration of sperm differentiation remain to be established.

326

327 *Differential constraints on sperm size and numbers*

328 At least in Eutheria, longer sperm take longer to produce, and sperm competition
329 selects for shorter sperm production times (Ramm & Stockley, 2010; delBarco-Trillo *et al.*,
330 2013). This suggests that to produce increasingly longer sperm in a competitive timeframe, an
331 enhanced mass-specific metabolic rate is required. In both Eutheria and Metatheria, sperm
332 size correlates negatively with body size, which is a consequence of differences in mass-
333 specific metabolic rates (low in large species and high in small species) (Gomendio *et al.*,
334 2011; Tourmente *et al.*, 2011a). Thus, in large mammalian species the increase on sperm size
335 in relation to high levels of sperm competition appears to be constrained by the influence of
336 mass-specific metabolic rate (Gomendio *et al.*, 2011; Tourmente *et al.*, 2011a). On the other
337 hand, an increase in sperm numbers can be attained simply by increasing the quantity of
338 sperm-producing units (i.e., increasing testes size). Absolute body mass and testes mass are
339 positively correlated with sperm numbers in sperm reserves in both Eutheria and Metatheria,
340 meaning that larger bodied species have larger testes and hence more sperm. Moreover, a

341 recent study found that sperm numbers in ejaculate, while increasing with relative testes size,
342 appear not to be associated with metabolism (Lüpold, 2013). In addition, it should be pointed
343 out that sperm size may be limited by factors other than energetic budget because size could
344 relate to hydrodynamic, biomechanical (Ishijima, 2012), and biochemical (e.g., energy
345 production, and ATP diffusion) (Takei *et al.*, 2014) constraints, beyond which motility would
346 not be efficient. Therefore, if sperm numbers and sperm size have a diverse set of energetic
347 constraints, a tradeoff between these two sperm traits is not required to explain a more
348 pronounced increase in sperm numbers associated to sperm competition.

349

350 *Variable scaling of the female reproductive tract*

351 The models described assume that the female reproductive tract is a uniform space,
352 which increases proportionally to the size of the female. In mammals, the female reproductive
353 tract generally comprises three major organs: the vagina, the uterus, and the oviducts, which
354 contain both the sperm reservoir (at the lower isthmus) and the site of fertilization (ampulla)
355 (Suarez, 2008). The size of the vagina and uterus exhibit considerable variation across
356 mammalian species and are related to adult body mass, individual offspring size, and total
357 litter mass (Austin & Short, 1985). However, the volume of the oviduct represents a small
358 fraction of the total size of the tract. The oviduct volume represents about 0.2% of the total
359 volume of the reproductive tract (estimated from (Peters & Ball, 1987; Kunhbar *et al.*, 2003))
360 and relates to body mass with an extremely low exponent (0.2809, (Gomendio & Roldan,
361 1993; Anderson *et al.*, 2006)). As an example, a cow has an oviduct 10-fold longer than a
362 mouse, while having an approximately 13,000-fold body mass (Gomendio & Roldan, 1993).
363 In this scenario, the effect of interspecific body mass variation on the scaling proportion
364 between ejaculate size and the volume of the fertilization space would be very different
365 according to the region of the female reproductive tract in which the sperm is located

366 (extremely high in the uterus and vagina; much lower in the oviduct, the actual fertilization
367 space in mammals).

368

369 *Sperm numbers at the site of fertilization and polyspermy*

370 Parker et al.'s raffle model assumes that the ejaculated sperm mass is affected only by
371 passive sperm loss in its passage through the female reproductive tract, and that the amount of
372 sperm lost is proportional to its size. In birds, this sperm loss is considerable since only 1-2%
373 of the ejaculated sperm reaches the oviductal sperm storage tubules (Brillard & Bakst, 1990;
374 Birkhead, 1992). In mammals, the sperm mass is rapidly transported (within minutes) from
375 sperm deposition sites (uterus or vagina depending on the species) to the oviduct by muscular
376 contractions. During this process, the sperm mass also sustains passive sperm loss, with a
377 reduction of sperm numbers of approximately one order of magnitude (Suarez & Pacey,
378 2006). However, at least in the Eutheria, this reduction in sperm numbers is relatively low
379 when compared to the reduction (5 to 6 orders of magnitude) that takes place when
380 spermatozoa swim through the junction between the uterus and the oviduct (utero-tubal
381 junction) (Hunter, 1993; Suarez & Pacey, 2006; Coy *et al.*, 2012). As a result, only a few
382 thousands of sperm are trapped and stored in the oviductal reservoir (Suarez, 2008; Coy *et al.*,
383 2012). It is noteworthy that, while the number of ejaculated sperm in mammals presents
384 considerable variation (a range of 5 orders of magnitude), and is related to body size [40],
385 similar sperm numbers are found in the oviductal reservoir in many species with different
386 body sizes (Harper, 1982). Following storage in the oviductal sperm reservoir, and after
387 completing the process of capacitation, sperm are released in discrete small-numbered
388 populations, which swim along the oviduct towards the site of fertilization (the oviductal
389 ampulla) (Suarez, 2008; Hunter, 2012). A final step in the reduction of sperm numbers relates
390 to the loss of cells that are not able to reach the ampulla, a process in which both thermotaxis

391 (Eisenbach & Giojalas, 2006) and chemotaxis (Guidobaldi *et al.*, 2012) are involved. Thus, as
392 a result of these complex physiological steps, the sperm:egg ratios at the time of fertilization
393 are close to unity (Hunter, 1996; Burkitt *et al.*, 2012; Coy *et al.*, 2012; Hunter, 2012). The
394 most likely explanation for such dynamics of sperm storage and release in mammals is the
395 need to prevent polyspermy (Coy & Aviles, 2010; Coy *et al.*, 2012; Hunter, 2012), which
396 generally results in the death of the embryo (Snook *et al.*, 2011). On the contrary, polyspermy
397 appears to be the rule in birds, and multiple sperm penetration in the egg's germinal disk (up
398 to 60 in some cases (Perry, 1987)) does not lead to any developmental problem for the
399 embryo (Snook *et al.*, 2011). Moreover, multiple sperm penetration in birds appears to be an
400 adaptation to assure fertility, since there is a positive correlation between the number of sperm
401 entering the ova and the ovum size (Birkhead *et al.*, 1994).

402 Taken together, differences in reproductive physiology suggest that, contrary to what
403 happens in birds, in mammals sperm velocity (which is related to sperm size) would play a
404 more essential role than sperm number in the last stage of the sperm journey to the site of
405 fertilization. Thus, this final step in sperm transport would resemble a short race, with only a
406 few participants, rather than a raffle.

407

408 *Differences between Eutheria and Metatheria*

409 In Eutheria sperm numbers tend to increase in a more pronounced way than sperm size
410 under increasing levels of sperm competition. This pattern was not observed in Metatheria. A
411 possible explanation for this difference may lay in fundamental differences that exist between
412 Eutheria and Metatheria with regards to their relative investments in sperm numbers and
413 sperm size. Metatheria have, on average, longer and fewer spermatozoa than their Eutheria
414 counterparts (mean total sperm length, Metatheria = 163 μ m; Eutheria = 91 μ m) (Tourmente *et*
415 *al.*, 2011b; a). Several species of Metatheria appear to have evolved extremely efficient

416 mechanisms to transport sperm to the fertilization site (Taggart *et al.*, 2003), measures of
417 sperm protection (such as the ability to store sperm in specialized crypts of the oviductal
418 epithelium) (Taggart *et al.*, 1998), and the formation of sperm pairs (which appear to enhance
419 velocity and/or protect the acrosomes) (Bedford *et al.*, 1984). Consequently, the proportion of
420 ejaculated sperm reaching the oviduct is up to four orders of magnitude higher than in some
421 Eutheria (Taggart *et al.*, 2003). Additionally, Metatheria have approximately 40% lower
422 testes mass in relation to their body mass than Eutheria (Tourmente *et al.*, 2011a). These
423 particular physiological characteristics (more efficient transport, high sperm survival rate, and
424 sperm storage in the female reproductive tract) may have resulted in a lower selective
425 pressure on the production of high sperm numbers in Metatheria when compared to Eutheria,
426 which may explain the comparatively lower relative testes mass and sperm numbers in the
427 former.

428

429 *Conclusions*

430 While we found some relationships that matched the prediction of the raffle model of
431 sperm competition (Parker *et al.*, 2010) regarding the relationships between sperm numbers
432 and size in mammals, we did not find evidence of a tradeoff between numbers and size when
433 controlling for sperm competition levels. These relationships between sperm traits may be
434 explained, in the absence of a tradeoff, as the product of two main features of mammalian
435 reproductive physiology: (1) The number and size of spermatozoa are regulated by different
436 molecular and cellular mechanisms underlying sperm proliferation and differentiation, which
437 are subjected to different energetic budgets and constraints. (2) The extraordinary variation in
438 the scaling between the volumes of the ejaculate and the female reproductive tract, and the
439 extreme reduction in sperm numbers at the fertilization site make increasing sperm numbers
440 and sperm size (through its influence on sperm velocity) adaptive advantages in different

441 stages of fertilization. In this way, increased sperm numbers would be more important at the
442 levels of the vagina and uterus, where sperm are likely to be diluted in a large environment,
443 while increased sperm size would be key in the oviduct, where there are only a few sperm
444 competing to reach the oocyte. These findings demand for revised or more detailed
445 hypotheses in order to elucidate the relationship between sperm traits in a sperm competition
446 context. Increased knowledge of the general and reproductive physiology of different taxa
447 should be taken into account to perform general predictions regarding the evolutionary effects
448 of sperm competition on sperm phenotype. In particular, physiological variables such as
449 mass-specific metabolic rate, timing, duration and efficiency of spermatogenesis in general
450 (and their different stages), accurate measures of the relative scaling of female reproductive
451 tract and sperm:egg ratio at the site of fertilization, and effects of sperm design and
452 physiology on sperm performance will be of foremost significance when predicting the
453 coevolution of sperm numbers and sperm size.

454

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462

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Table 1. Relation between sperm numbers, relative testes size and total sperm size in mammals. Phylogenetically controlled multiple regression analyses (PGLS). Superscripts following the λ value indicate significance levels (n.s. $p > 0.05$; * $p < 0.05$) in likelihood ratio tests against models with $\lambda = 0$ (first position) and $\lambda = 1$ (second position). Effect size r calculated from the t values and the non-central 95% confidence limits (CLs) for the z -transformed value of r are presented. Confidence intervals excluding 0 indicate statistically significant relationships. P -values and CL that indicate statistical significance are shown in bold. All variables were \log_{10} -transformed. n: number of species.

Clade	Dependent variable	Independent variable	Slope	R^2	p	t	λ	r	CL(-)	CL(+)
Eutheria (n=64)	Sperm numbers	Body mass	-0.7318	0.72	<0.0001	-5.0570	0.999 ^{*,ns}	-0.5435	-0.8601	-0.3582
		Testes mass	1.8320		<0.0001	11.1430		0.8189	0.9025	1.4044
Metatheria (n=14)	Sperm numbers	Body mass	-0.0959	0.88	0.7866	-0.2775	0.001 ^{ns,ns}	-0.0834	-0.6745	0.5074
		Testes mass	1.4944		0.0214	2.6792		0.6284	0.1478	1.3297
Eutheria (n=64)	Sperm numbers	Body mass	0.5710	0.25	0.0003	3.8675	0.988 ^{*,ns}	0.4438	0.2259	0.7279
		Total sperm length	2.5975		0.0132	2.5543		0.3108	0.0705	0.5724
Metatheria (n=14)	Sperm numbers	Body mass	0.5837	0.81	0.0640	2.0584	0.001 ^{ns,ns}	0.5273	-0.0045	1.1774
		Total sperm length	-1.4523		0.4305	-0.8183		-0.2395	-0.8353	0.3467
Eutheria (n=64)	Sperm numbers	Body mass	-0.6853	0.73	<0.0001	-4.5779	0.999 ^{*,ns}	-0.5088	-0.8120	-0.3101
		Testes mass	1.7769		<0.0001	10.4166		0.8025	0.8545	1.3564
		Total sperm length	0.7466		0.2481	1.1663		0.1489	-0.1009	0.4010
Metatheria (n=14)	Sperm numbers	Body mass	-0.4323	0.90	0.3138	-1.0606	0.001 ^{ns,ns}	-0.3180	-0.9204	0.2616
		Testes mass	1.5858		0.0146	2.9477		0.6819	0.2416	1.4235
		Total sperm length	-1.9384		0.1879	-1.4134		-0.4081	-1.0242	0.1577

Table 2. Relation between sperm investment and relative testes size in mammals. Phylogenetically controlled multiple regression analyses (PGLS). Superscripts following the λ value indicate significance levels (n.s. $p>0.05$; $*p<0.05$) in likelihood ratio tests against models with $\lambda = 0$ (first position) and $\lambda = 1$ (second position). Effect size r calculated from the t values and the non-central 95% confidence limits (CLs) for the z -transformed value of r are presented. Confidence intervals excluding 0 indicate statistically significant relationships. P -values and CL that indicate statistical significance are shown in bold. All variables were \log_{10} -transformed. n: number of species.

Clade	Dependent variable	Independent variable	Slope	R^2	p	t	λ	r	CL(-)	CL(+)
Eutheria	Total sperm investment	Body mass	-2.0393	0.63	<0.0001	-5.2897	0.980 ^{*,ns}	-0.5608	-0.8913	-0.3766
		Testes mass	4.2121		<0.0001	9.5670		0.7746	0.7745	1.2892
	Relative investment in sperm size	Body mass	0.0170	0.62	0.0014	3.3493	0.999 ^{*,ns}	0.3941	0.1593	0.6740
		Testes mass	-0.0489		<0.0001	-8.4737		-0.7353	-1.1975	-0.6828
Metatheria	Total sperm investment	Body mass	-1.2102	0.57	0.0536	-2.1612	0.999 ^{ns,ns}	-0.5459	-1.3056	0.0804
		Testes mass	2.9936		0.0057	3.4177		0.7176	0.2098	1.5957
	Relative investment in sperm size	Body mass	-0.0290	0.90	0.1669	-1.4801	0.001 ^{ns,*}	-0.4075	-1.1256	0.2603
		Testes mass	-0.0398		0.2352	-1.2557		-0.3541	-1.0631	0.3229

FIGURE LEGENDS

Figure 1. Relationships between sperm numbers and sperm size in mammals. Figure points are partial residuals estimated from multiple regression analysis using sperm numbers as dependent variable. A, B: body mass and total sperm length used as predictors. C, D: body mass, testes mass, and total sperm length used as predictors. All variables were \log_{10} -transformed. A, C: Eutheria. B, D: Metatheria.

Figure 2. Relationships between sperm investment and relative testes size in mammals. Figure points are partial residuals estimated from multiple regression analysis using total sperm investment (sperm size * sperm numbers) (A, B), or relative investment in sperm size (sperm size / numbers) (C, D) as dependent variable, and body mass and testes mass as predictors. All variables were \log_{10} -transformed. A, C: Eutheria. B, D: Metatheria.

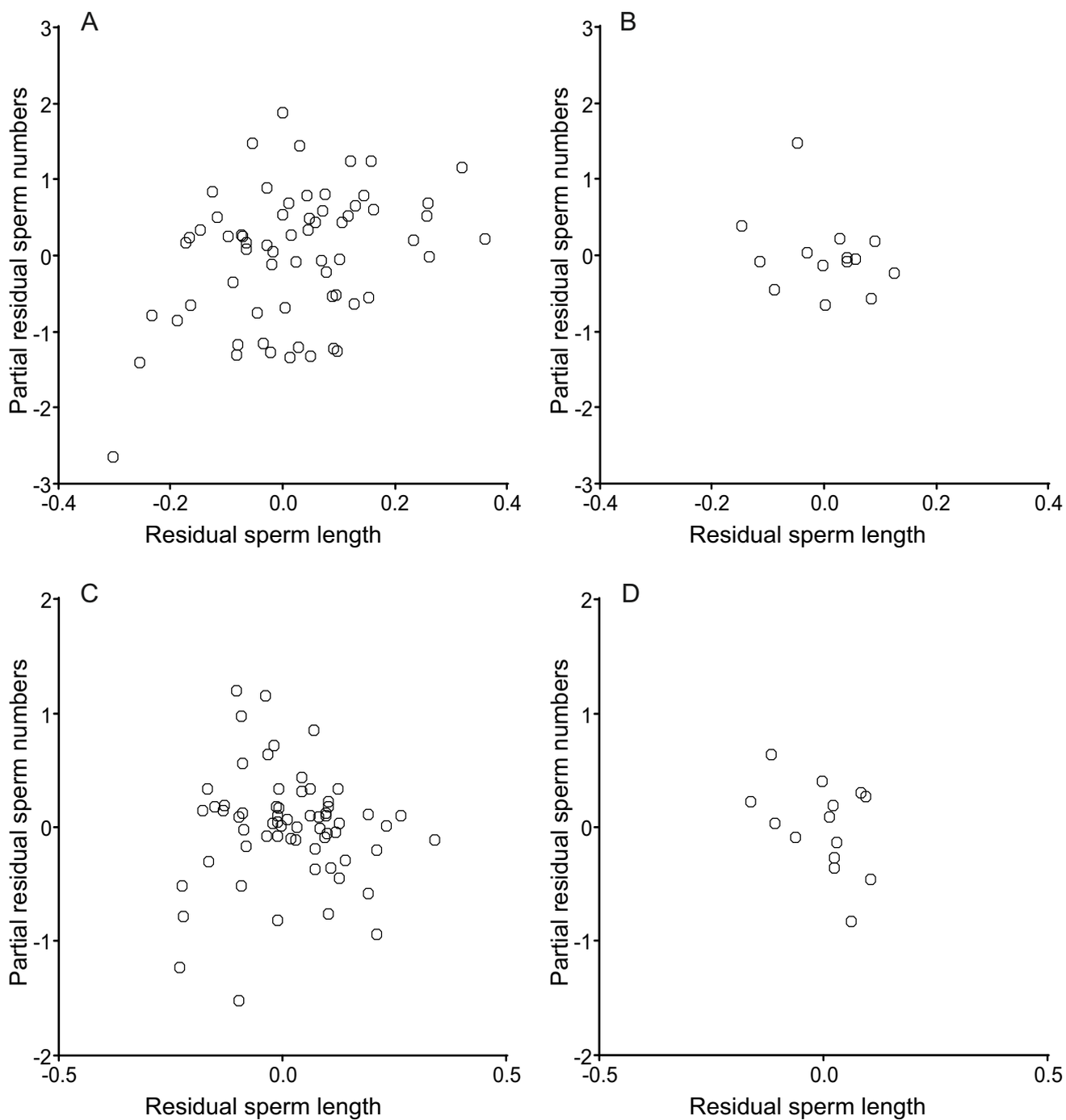


Figure 1

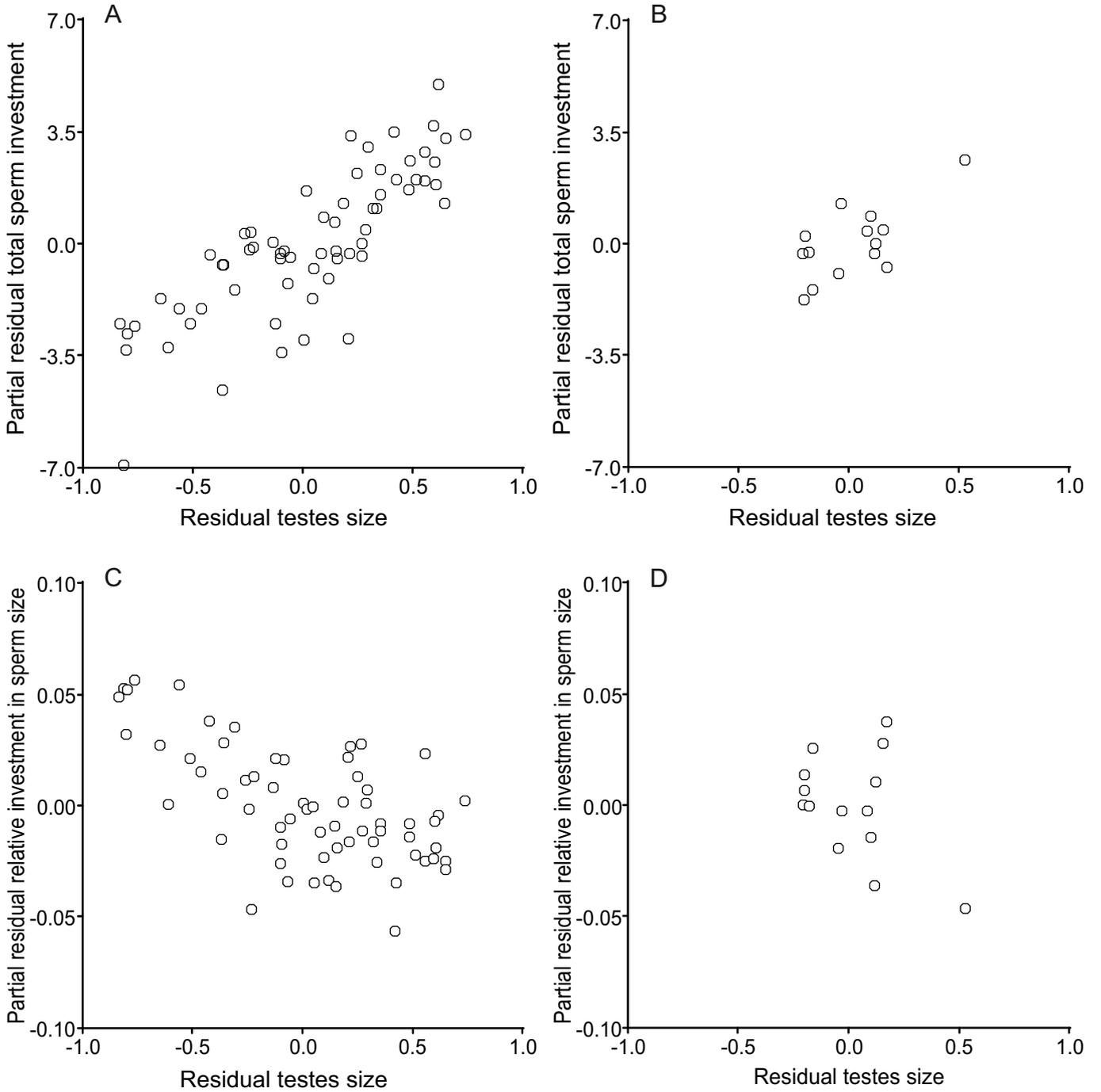


Figure 2

No evidence of tradeoffs in the evolution of sperm numbers and sperm size in mammals

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Supporting Information

Table S1. Sperm size, sperm numbers, body mass, and testes mass in 78 species of mammals.

Table S2. Alternative models for the relation between sperm numbers, sperm size and relative testes size in mammals.

Figure S1. Phylogenetic reconstruction for the 78 mammal species utilized in the PGLS analysis.

Table S1. Sperm size, sperm numbers, body mass, and testes mass in 78 species of mammals. Abbreviations: TSL: total sperm length (μm). SN: sperm numbers in caudae epididymides ($\times 10^6$). BM: body mass (g). TM: testes mass (g).

Species	Clade	Order	Family	TSL	SN	BM	TM	Ref. TSL	Ref. SN	Ref. BM & TM
<i>Bos taurus</i>	Eutheria	Artiodactyla	Bovidae	53.53	51775.00	680385.00	681.00	(1, 2)	(22, 23)	(69)
<i>Bubalus bubalis</i>	Eutheria	Artiodactyla	Bovidae	69.35	6545.45	680000.00	652.00	(3)	(3, 24)	(70)
<i>Capra hircus</i>	Eutheria	Artiodactyla	Bovidae	59.39	23680.00	25420.00	156.80	(1)	(25)	(71)
<i>Connochaetes taurinus</i>	Eutheria	Artiodactyla	Bovidae	46.77	24882.00	227000.00	306.00	(1)	(26)	(13)
<i>Ovis aries</i>	Eutheria	Artiodactyla	Bovidae	64.70	145500.00	57172.73	222.99	(1, 2)	(27, 28)	(71)
<i>Taurotragus oryx</i>	Eutheria	Artiodactyla	Bovidae	65.31	5700.00	408500.00	203.20	(4, 5)	(29)	(29)
<i>Camelus dromedarius</i>	Eutheria	Artiodactyla	Camelidae	47.21	2197.40	800000.00	121.70	(1, 2)	(30)	(71)
<i>Odocoileus virginianus</i>	Eutheria	Artiodactyla	Cervidae	52.00	8130.00	44800.00	85.50	(1, 5)	(31)	(69)
<i>Hippopotamus amphibius</i>	Eutheria	Artiodactyla	Hippopotamidae	33.49	11030.00	1600000.00	650.00	(1, 5)	(32)	(13)
<i>Sus scrofa</i>	Eutheria	Artiodactyla	Suidae	54.60	124733.33	39700.00	128.20	(1, 2)	(27, 28, 33)	(72)
<i>Canis familiaris</i>	Eutheria	Carnivora	Canidae	61.40	2119.00	21620.00	27.66	(4, 6)	(34, 35)	(35)
<i>Mustela vison</i>	Eutheria	Carnivora	Mustelidae	43.00	100.00	2387.60	5.40	(1)	(36)	(73, 74)
<i>Ursus arctos</i>	Eutheria	Carnivora	Ursidae	70.35	640.00	180900.00	68.60	(4, 2, 7)	(37)	(75)
<i>Lepus californicus</i>	Eutheria	Lagomorpha	Leporidae	50.32	1042.00	2250.00	8.48	(8)	(8)	(8)
<i>Oryctolagus cuniculus</i>	Eutheria	Lagomorpha	Leporidae	58.02	1286.00	2888.00	6.06	(1)	(8, 27, 34, 38)	(69)
<i>Romerolagus diazi</i>	Eutheria	Lagomorpha	Leporidae	54.61	310.00	486.00	1.88	(8)	(8)	(8)
<i>Elephantulus myurus</i>	Eutheria	Macroscelidea	Macroscelididae	76.60	2.59	60.00	0.08	(2)	(39, 40)	(40)
<i>Equus asinus</i>	Eutheria	Perissodactyla	Equidae	64.15	60935.00	290000.00	202.34	(5, 9)	(41)	(71)
<i>Equus caballus</i>	Eutheria	Perissodactyla	Equidae	60.60	59487.67	468000.00	416.00	(1, 2)	(27, 28, 41)	(69)
<i>Macaca mulatta</i>	Eutheria	Primates	Cercopithecidae	74.10	10358.00	10430.00	76.00	(1, 4)	(42)	(69)
<i>Homo sapiens</i>	Eutheria	Primates	Hominidae	56.90	304.64	63540.00	50.20	(4)	(34, 43)	(69)
<i>Chionomys nivalis</i>	Eutheria	Rodentia	Arvicolidae	105.23	129.40	43.64	0.85	(10)	(44)	(44)
<i>Microtus arvalis</i>	Eutheria	Rodentia	Arvicolidae	91.66	41.60	36.40	0.28	(10)	(44)	(44)
<i>Microtus cabrerai</i>	Eutheria	Rodentia	Arvicolidae	85.12	7.60	44.27	0.14	(10)	(44)	(44)
<i>Microtus duodecimcostatus</i>	Eutheria	Rodentia	Arvicolidae	62.69	4.50	29.76	0.08	(10)	(44)	(44)
<i>Microtus lusitanicus</i>	Eutheria	Rodentia	Arvicolidae	86.02	27.70	17.73	0.09	(10)	(44)	(44)
<i>Microtus ochrogaster</i>	Eutheria	Rodentia	Arvicolidae	94.40	149.89	45.40	0.52	(11)	(45)	(76)
<i>Myodes glareolus</i>	Eutheria	Rodentia	Arvicolidae	83.91	43.20	25.65	0.40	(10)	(44)	(44)
<i>Fukomys damarensis</i>	Eutheria	Rodentia	Bathyergidae	42.90	0.26	194.50	0.18	(12)	(46)	(46)
<i>Mesocricetus auratus</i>	Eutheria	Rodentia	Cricetidae	186.70	1130.00	108.00	3.17	(13)	(27, 28, 42)	(69)

<i>Peromyscus californicus</i>	Eutheria	Rodentia	Cricetidae	72.00	248.00	34.20	0.20	(1)	(47)	(77)
<i>Peromyscus leucopus</i>	Eutheria	Rodentia	Cricetidae	74.80	91.17	24.21	0.38	(1)	(48)	(77, 78)
<i>Ctenomys talarum</i>	Eutheria	Rodentia	Ctenomyidae	49.28	3.22	118.00	0.35	(11)	(49)	(49)
<i>Apodemus sylvaticus</i>	Eutheria	Rodentia	Muridae	126.15	110.10	30.43	0.96	(10)	(44)	(44)
<i>Conilurus penicillatus</i>	Eutheria	Rodentia	Muridae	124.00	1908.00	184.00	4.39	(1)	(50)	(79)
<i>Hydromys chrysogaster</i>	Eutheria	Rodentia	Muridae	115.00	1156.00	745.00	11.77	(1)	(50)	(79)
<i>Melomys littoralis</i>	Eutheria	Rodentia	Muridae	110.00	333.00	61.00	1.68	(1)	(50)	(2)
<i>Mus musculus musculus</i>	Eutheria	Rodentia	Muridae	124.64	23.20	21.13	0.13	(10)	(44)	(44)
<i>Mus pahari</i>	Eutheria	Rodentia	Muridae	137.56	9.10	30.08	0.12	(14)	(44)	(44)
<i>Mus spicilegus</i>	Eutheria	Rodentia	Muridae	103.54	99.40	14.49	0.41	(10)	(44)	(44)
<i>Mus spretus</i>	Eutheria	Rodentia	Muridae	111.49	48.00	17.01	0.29	(10)	(44)	(44)
<i>Notomys alexis</i>	Eutheria	Rodentia	Muridae	102.50	2.23	34.00	0.05	(1)	(51-53)	(69)
<i>Notomys cervinus</i>	Eutheria	Rodentia	Muridae	115.00	21.40	33.00	0.20	(1)	(53)	(69)
<i>Notomys fuscus</i>	Eutheria	Rodentia	Muridae	106.00	2.00	44.00	0.07	(1)	(53)	(69)
<i>Notomys mitchelli</i>	Eutheria	Rodentia	Muridae	98.00	1.80	39.00	0.06	(1)	(53)	(69)
<i>Pseudomys apodemoides</i>	Eutheria	Rodentia	Muridae	120.00	10.00	31.00	0.14	(1)	(53)	(69)
<i>Pseudomys australis</i>	Eutheria	Rodentia	Muridae	122.50	900.65	59.00	2.24	(1)	(51-53)	(69)
<i>Pseudomys hermannsburgensis</i>	Eutheria	Rodentia	Muridae	117.00	20.00	18.00	0.16	(1)	(53)	(69)
<i>Pseudomys nanus</i>	Eutheria	Rodentia	Muridae	127.00	387.00	78.00	1.87	(1)	(53)	(69)
<i>Pseudomys shortridgei</i>	Eutheria	Rodentia	Muridae	96.00	51.00	78.00	0.35	1, 2	(53)	(69)
<i>Rattus colletti</i>	Eutheria	Rodentia	Muridae	158.00	323.00	154.00	1.94	(1)	(50)	(79)
<i>Rattus fuscipes</i>	Eutheria	Rodentia	Muridae	162.00	387.00	110.00	4.26	(1)	(50)	(16)
<i>Rattus norvegicus</i>	Eutheria	Rodentia	Muridae	189.40	303.68	379.63	3.06	(1)	(54-58)	(57, 69, 80-82)
<i>Rattus tunneyi</i>	Eutheria	Rodentia	Muridae	155.00	130.00	243.00	4.87	(15)	(50)	(50)
<i>Rattus villosissimus</i>	Eutheria	Rodentia	Muridae	147.00	195.00	207.00	2.15	(16)	(50)	(50)
<i>Rhabdomys pumilio</i>	Eutheria	Rodentia	Muridae	117.00	161.00	70.33	1.28	(1)	(59)	(59)
<i>Uromys caudimaculatus</i>	Eutheria	Rodentia	Muridae	106.00	1342.00	820.00	7.20	(1, 2)	(50)	(2)
<i>Tupaia belangeri</i>	Eutheria	Scadentia	Tupaiaidae	73.05	6.32	141.00	1.49	(17)	(17)	(17)
<i>Blarina brevicauda</i>	Eutheria	Soricomorpha	Soricidae	70.00	60.00	24.00	0.30	(2)	(60)	(60)
<i>Crocidura russula</i>	Eutheria	Soricomorpha	Soricidae	102.70	3.89	13.40	0.04	(18)	(61)	(18)
<i>Neomys fodiens</i>	Eutheria	Soricomorpha	Soricidae	93.90	30.10	16.70	0.25	(18)	(61)	(18)
<i>Sorex araneus</i>	Eutheria	Soricomorpha	Soricidae	83.90	7.42	10.90	0.17	(18)	(61, 62)	(18)
<i>Suncus murinus</i>	Eutheria	Soricomorpha	Soricidae	113.30	3.30	66.10	0.15	(18, 19)	(61)	(69)
<i>Talpa europaea</i>	Eutheria	Soricomorpha	Talpidae	90.00	5292.00	92.31	1.80	(20)	(60)	(60)

<i>Antechinus stuartii</i>	Metatheria	Dasyuomorpha	Dasyuridae	271.10	2.07	40.00	0.64	(1)	(23, 63)	(65)
<i>Dasyuroides byrnei</i>	Metatheria	Dasyuomorpha	Dasyuridae	254.80	1.70	123.60	1.38	(1)	(23, 64)	(65)
<i>Sminthopsis crassicaudata</i>	Metatheria	Dasyuomorpha	Dasyuridae	264.90	1.22	14.60	0.17	(2)	(65)	(65)
<i>Monodelphis domestica</i>	Metatheria	Didelphiomorpha	Didelphidae	211.90	4.20	110.00	0.57	(2)	(65)	(65)
<i>Macropus eugenii</i>	Metatheria	Diprotodontia	Macropodidae	106.40	4851.00	7050.00	32.90	(1)	(66)	(83)
<i>Macropus rufus</i>	Metatheria	Diprotodontia	Macropodidae	123.90	388.00	39825.00	38.18	(1)	(64)	(63)
<i>Wallabia bicolor</i>	Metatheria	Diprotodontia	Macropodidae	109.50	482.00	31500.00	14.75	(1)	(23, 64)	(63)
<i>Trichosurus vulpecula</i>	Metatheria	Diprotodontia	Phalangeridae	94.17	218.33	3350.00	8.26	(1)	(23, 63, 64)	(63)
<i>Aepyprymnus rufescens</i>	Metatheria	Diprotodontia	Potoroidae	106.40	57.80	2400.00	4.68	(21)	(63)	(63)
<i>Potorous tridactylus</i>	Metatheria	Diprotodontia	Potoroidae	165.80	38.80	1280.00	4.38	(21)	(63)	(63)
<i>Lasiorhinus latifrons</i>	Metatheria	Diprotodontia	Vombatidae	79.50	177.33	28290.00	13.80	(1)	(23, 64, 67)	(63, 84)
<i>Vombatus ursinus</i>	Metatheria	Diprotodontia	Vombatidae	93.60	146.50	40100.00	18.42	(1)	(68)	(63)
<i>Isoodon macrourus</i>	Metatheria	Peramelemorpha	Peramelidae	171.10	101.70	2300.00	4.72	(1)	(23, 64, 63)	(63)
<i>Isoodon obesulus</i>	Metatheria	Peramelemorpha	Peramelidae	167.00	56.00	978.30	3.88	(1)	(64)	(63)

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Table S2. Alternative models for the relation between sperm numbers, sperm size and relative testes size in mammals. Phylogenetically controlled multiple regression analyses (PGLS). Superscripts following the λ value indicate significance levels (n.s. $p>0.05$; * $p<0.05$) in likelihood ratio tests against models with $\lambda = 0$ (first position) and $\lambda = 1$ (second position). Effect size r calculated from the t values and the non-central 95% confidence limits (CLs) for the z -transformed value of r are presented. Confidence intervals excluding 0 indicate statistically significant relationships. P -values and CL that indicate statistical significance are shown in bold. All variables were \log_{10} -transformed. n: number of species. Testes mass²: quadratic term of testes mass. Body mass * Testes mass: interaction between body mass and testes mass.

Clade	Dependent variable	Independent variable	Slope	R^2	p	t	λ	r	CL(-)	CL(+)
Eutheria (n=64)	Sperm numbers	Body mass	-0.7417	0.73	0.0005	-3.6646	0.999 ^{*,ns}	-0.4277	-0.7080	-0.2061
		Testes mass	2201.8000		0.2152	1.2528		0.1597	-0.0899	0.4120
		Testes mass ²	-1100.0000		0.2156	-1.2517		-0.1595	-0.4119	0.0901
		Body mass * Testes mass	-0.0003		0.9958	-0.0053		-0.0007	-0.2516	0.2503
	Sperm numbers	Body mass	-0.7066	0.72	0.0009	-3.5087	0.999 ^{*,ns}	-0.4126	-0.6897	-0.1878
		Testes mass	1.8501		<0.0001	9.5596		0.7770	0.7867	1.2886
		Body mass * Testes mass	-0.0118		0.8566	-0.1815		-0.0234	-0.2744	0.2275
	Sperm size	Body mass	-0.0432	0.09	0.2934	-1.0601	0.999 ^{*,ns}	-0.1356	-0.3874	0.1145
		Testes mass	-42.2989		0.9053	-0.1195		-0.0154	-0.2664	0.2355
		Testes mass ²	21.1929		0.9051	0.1198		0.0155	-0.2355	0.2664
		Body mass * Testes mass	-0.0088		0.5041	-0.6722		-0.0865	-0.3376	0.1643
	Sperm size	Body mass	-0.0439	0.09	0.2773	-1.0963	0.980 ^{*,ns}	-0.1401	-0.3920	0.1099
Testes mass		0.0870		0.0274	2.2617		0.2803	0.0370	0.5389	
Body mass * Testes mass		-0.0086		0.5070	-0.6675		-0.0859	-0.3370	0.1649	
Metatheria (n=14)	Sperm numbers	Body mass	-0.1076	0.88	0.8080	-0.2502	<0.001 ^{ns,*}	-0.0789	-0.6700	0.5119
		Testes mass	284.2800		0.9413	0.0757		0.0239	-0.5670	0.6149
		Testes mass ²	-141.3900		0.9416	-0.0753		-0.0238	-0.6148	0.5672
		Body mass * Testes mass	0.0073		0.9690	0.0400		0.0126	-0.5783	0.6036
	Sperm numbers	Body mass	-0.1033	0.88	0.8036	-0.2554	<0.001 ^{ns,*}	-0.0805	-0.6716	0.5103
		Testes mass	1.4800		0.0548	2.1746		0.5666	0.0516	1.2335
		Body mass * Testes mass	0.0072		0.9678	0.0414		0.0131	-0.5779	0.6041
	Sperm size	Body mass	-0.1797	0.84	0.0740	-2.0212	<0.001 ^{ns,*}	-0.5386	-1.1931	-0.0112
		Testes mass	-385.2988		0.6317	-0.4961		-0.1550	-0.7472	0.4347
		Testes mass ²	192.6532		0.6317	0.4961		0.1550	-0.4347	0.7472
		Body mass * Testes mass	0.0115		0.7691	0.3026		0.0953	-0.4954	0.6865
	Sperm size	Body mass	-0.1856	0.83	0.0533	-2.1899	<0.001 ^{ns,*}	-0.5693	-1.2375	-0.0556
Testes mass		0.0239		0.8702	0.1677		0.0530	-0.5380	0.6440	
Body mass * Testes mass		0.0116		0.7561	0.3193		0.1005	-0.4902	0.6918	

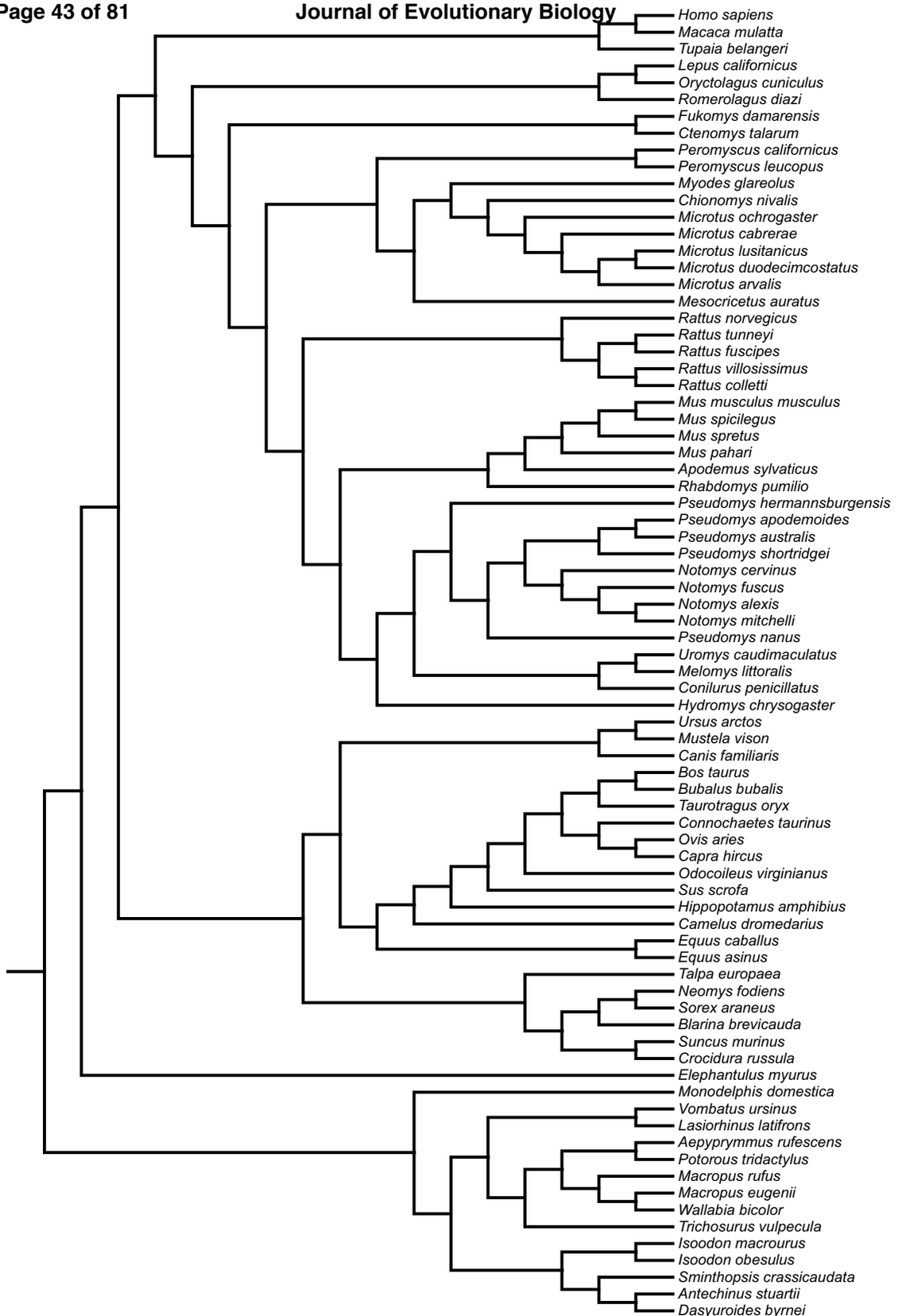


Figure S1. Phylogenetic reconstruction for the 78 mammal species utilized in the PGLS analysis.