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3 4	Title:	Brain size, life histories and maternal investment in eutherian mammals
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8	Authors:	Robert A. Barton ¹ & Isabella Capellini ^{1,2}
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10		^{1.} Evolutionary Anthropology Research Group,
11		Department of Anthropology, Durham University,
12		Dawson Building, South Road, Durham, DH1 3LE, U. K.
13		^{2.} Present address: School of Biological Sciences,
14		Queen's University of Belfast, 97 Lisburn Road, Belfast
15		BT7 9BL, U.K.
16		
17	Corresponding author:	Robert Barton
18		r.a.barton@durham.ac.uk
19		Evolutionary Anthropology Research Group,
20		Department of Anthropology, Durham University,
21		Dawson Building, South Road, Durham, DH1 3LE, U. K.
22		

23 Abstract

24 Brain size variation in mammals correlates with life histories: larger-brained species 25 have longer gestations, mature later and have increased lifespans. These patterns 26 have been explained in terms of both developmental costs (larger brains take longer 27 to grow) and cognitive benefits (large brains enhance survival and increase 28 lifespan). In support of the developmental cost hypothesis, we show that 29 evolutionary changes in pre- and post-natal brain growth correlate specifically with 30 duration of the relevant phases of maternal investment (gestation and lactation 31 respectively). We also find support for the hypothesis that the rate of fetal brain 32 growth is related to the energy turnover of the mother. In contrast, we find no 33 support for hypotheses proposing that costs are accommodated through direct 34 trade-offs between brain and body growth, or between brain growth and litter size. 35 Once the duration of maternal investment is taken into account, adult brain size is 36 uncorrelated with other life history traits such as lifespan. Hence, the general 37 pattern of slower life histories in large-brained species appears to be a direct 38 consequence of developmental costs.

40 \body

Brain size varies extensively between species. Many comparative studies have been 41 42 aimed at understanding how and why such variation evolved, and have identified a 43 range of factors associated with the evolution of large brains. One general factor 44 robustly correlated with brain size is life history; larger-brained species, such as 45 humans, develop slowly, have extended periods of juvenility and long lifespans, 46 effects that remain after accounting for differences in body size¹⁻⁸. These 47 associations have been interpreted in two different ways. First, life history 48 correlates could reflect the benefits of large brains in providing a "cognitive buffer" 49 against environmental unpredictability, improving survival and permitting long 50 lives^{2,6-7}. Second, selection on brain size might have secondary consequences for life 51 history because larger brains impose a developmental cost, in terms of a need for extended growth and maturation^{3-5, 8}. 52

53 Because large brains must have both benefits and costs, the two types of 54 explanation for the association between brain size and life history are not 55 necessarily incompatible³⁻⁷. They do, however, make different predictions. The 56 cognitive buffer hypothesis predicts correlations between brain size, survival and 57 lifespan⁶⁻⁷. Developmental costs hypotheses on the other hand, assume that brain 58 growth has to be traded off against aspects of production, including growth, 59 maturation time and reproductive rates, causing larger-brained species to grow and 60 mature more slowly and to have lower fertility^{4-5,8-10}. This idea overlaps with the 61 'Maternal Energy Hypothesis', which suggests that maternal investment and energy

availability constrain the development of large brains, predicting that brain size
correlates with the duration of maternal investment and with maternal basal
metabolic rate (BMR)¹¹⁻¹². Recent comparative evidence is consistent with both
cognitive buffer and developmental cost ideas; brain size variation in adult
mammals is positively correlated with lifespan⁶⁻⁷ as well as with the durations of
gestation, lactation and the juvenile period^{4-5,8,13-14}.

68 Little attempt has so far been made to distinguish between the effects of 69 these different developmental and life history traits, making individual correlations 70 with brain size difficult to interpret. In particular, it is unknown, whether maternal 71 investment and lifespan are both independently associated with brain size, or 72 whether life history correlations are driven primarily by one of these factors. 73 Furthermore, most studies focus on correlates of adult brain size, which can provide 74 only indirect evidence for developmental costs. A critical and more direct test is 75 whether brain growth during specific phases of development correlates with the 76 relevant aspects of maternal investment and maturation time. Evidence on this 77 question is limited. Some studies have demonstrated a positive correlation between 78 neonate brain size and gestation length, but these were conducted either before the 79 advent of powerful phylogenetic comparative methods⁵⁻¹⁶, or on small samples of 80 primate species^{10,17}. Studies of postnatal brain growth have also been limited to 81 small samples of primates, and do not support the critical prediction of an 82 association between postnatal brain growth and lactation^{10,17}, a finding in tension with the result that adult brain size correlates with lactation duration in a wider 83 84 range of mammals⁵. Similarly, although recent studies find that adult brain size

correlates with BMR^{8,13}, evidence that this reflects maternal metabolic constraints
on either pre- or postnatal brain growth is lacking¹⁶.

87 Indeed, it is not even clear how variability in pre- and postnatal brain growth 88 combine to influence variation in adult brain size. The relative amounts of pre- and 89 postnatal brain growth differ significantly between species¹⁷, and analysis of the 90 genetic correlates of brain size evolution suggests that the two phases of brain 91 growth are genetically dissociable¹⁸. Hence, they could in principle make 92 independent contributions to species differences in adult brain size. However, it has 93 been suggested that the relative brain sizes of neonates and adults are uncorrelated 94 in mammals^{8,10,19}, implying that pre- and postnatal brain growth are traded off. If 95 true, this would suggest that differences in prenatal maternal investment have no 96 impact on adult brain size. On the other hand, recent evidence suggests that neonate 97 and adult brain size are positively correlated in precocial species, but not in altricial 98 species^{5,20}. Thus, the question of what developmental mechanisms underpin the 99 evolution of differences in brain size requires further investigation. Given that 100 different neuro-developmental processes are concentrated in different phases²¹, and 101 that opportunities for environmental input occur principally after birth, determining 102 the developmental mechanisms of brain size evolution is likely to be important for 103 understanding its neuroanatomical and functional consequences.

Here we use phylogenetic comparative methods to examine the
developmental mechanisms underlying mammalian brain size evolution, and
comprehensively test predictions of the developmental costs hypothesis.

107 Specifically, we examine the contributions of both pre- and postnatal growth to 108 variation in adult brain size, and test the prediction that these phases correlate 109 specifically with gestation and lactation duration respectively, even after controlling 110 for other reproductive and life history variables. We also test whether costs are 111 accommodated through trade-offs between brain and body growth, or between 112 brain size and litter size, and we evaluate at which stage if any maternal metabolic 113 rate is related to brain growth. We evaluate the relative statistical power of 114 developmental costs and cognitive buffer hypotheses as explanations for 115 correlations between brain size and life history, by testing whether brain size is 116 independently associated with maternal investment and other life history variables, 117 such as lifespan. To these ends, we use phylogenetic generalized linear models 118 (PGLM) to test for correlated evolution among traits. We explore the effects of 119 specific variables on the explanatory power of the models by statistically comparing 120 models with versus without the variables in question, using the log-likelihood ratio 121 (LR) test (see Methods).

122

123 **Results**

Pre- and post-natal contributions to adult brain size. Adult and neonate brain
size are positively correlated, controlling for both adult and neonate body mass
(Figure 1 and Table 1). Additionally controlling for gestation length effectively turns
neonate brain size into a rate of relative brain growth (i.e. tests whether adult brain
size increases with the amount of prenatal brain growth relative to prenatal body

129 growth and the amount of time *in utero*); when this is done, adult brain size is 130 significantly positively correlated with neonate brain size (t_{117} =5.54, p<0.001). 131 Neonate body mass was not associated with adult brain size independently of 132 neonate brain size: adding neonate mass to the predictors did not improve the 133 model fit (model 1 versus model 2 in Table 1; $LR_1=1.8$, p>0.05), and neonate body 134 mass correlates with adult brain size only when neonate brain size is excluded from 135 the model (t_{118} =3.71, n=122, p<0.001). The addition of post-natal brain growth, 136 however, significantly improves the fit of the initial model (model 1 versus model 3 137 in Table 1; LR₁=269.9, p<0.001, increase in R^2 from 0.92 to 0.99). The effect sizes (as 138 estimated by t-values in model 3) suggest that postnatal brain growth may be a 139 stronger predictor of adult brain size, and running the initial model with postnatal 140 brain growth instead of neonate brain size yields a higher R² (0.97). Nevertheless, 141 the likelihood ratio test comparing model 3 to the same model but without neonate 142 brain size is highly significant (LR₁=70.89, p<0.001). Hence, variation in brain size at 143 birth and in the amount of brain growth postnatally have independent influences on 144 adult brain size.

145Neonate brain size and postnatal brain growth are uncorrelated, controlling146for neonate body mass and maternal mass ($t_{119}=1.30$, p>0.1), further emphasizing147the independent contributions of fetal and postnatal growth to adult brain size.148There was also no significant interaction between the effects on adult brain size of149neonatal brain size and postnatal brain growth when this interaction term was150added to model 1 (t_{118} =-0.84, p>0.1). These results therefore suggest that there is no

trade-off between pre- and postnatal brain growth, and that their effects on adultbrain size are additive rather than multiplicative.

153

154 **Correlates of neonate brain size.** Accounting for allometric effects (neonate body 155 mass and maternal body mass), neonate brain size is positively associated with 156 gestation length (Table 2, model 1). Adding litter size to the predictors in model 1 157 did not improve the model fit (LR₁=1.18, p>0.1) and litter size was not significantly 158 associated with neonate brain size (Table 2, model 2). To check that the apparent 159 effect of gestation length is not simply a side-effect of some more general growth or 160 early life-history correlate of brain size, lactation length was added as a predictor to 161 model 1 (reducing sample size to 111): neonate brain size remained significantly 162 associated with gestation length (t_{105} =6.14, p<0.001) but was unrelated to lactation 163 length (t_{105} =0.77, p>0.1), and the likelihood ratio test for models with and without 164 lactation was non-significant (LR₁=0.6, p>0.5). Because the relationship between 165 brain growth and litter size may interact with developmental state (i.e. a trade-off 166 occurs in altricial but not in precocial species⁵), we ran a model with developmental 167 state and the interaction between developmental state and litter size added as 168 predictors. The effects of neonate mass, maternal mass and gestation length 169 remained significant (neonate mass, t_{102} =5.34, p<0.001; maternal mass, t_{102} = 3.64, 170 p<0.001; gestation length, $t_{102}=4.91$, p<0.001), and in addition there was a 171 significant effect of developmental state (precociality is associated with larger brain 172 size; t_{102} = 2.49, p<0.05). However, there was still no main effect of litter size

(t₁₀₂=0.11, p>0.5), nor a significant interaction between developmental state and
litter size, (t₁₀₂=-1.81, p>0.05). Note that maternal size was positively associated
with neonate brain size in these analyses, even after controlling for other variables,
suggesting that larger females produce more encephalized offspring, reiterating the
importance of maternal investment. Note also that in all these analyses, neonate
brain size increases with neonate body size, hence showing no signs of a trade-off
between neural and somatic growth.

180 We tested for a possible association of BMR with neonatal brain size, 181 controlling for neonate body mass, maternal body mass and gestation length. 182 Gestation length remained a significant predictor of neonate brain size (t_{40} =6.41, 183 p<0.001) and BMR was also positively correlated with neonate brain size ($t_{40}=3.07$, 184 p<0.01). The model including BMR provided a significantly better fit than one 185 omitting it (LR₁=7.50, p<0.01, increase in \mathbb{R}^2 from 0.93 to 0.96). BMR remained 186 positively correlated with neonate brain size when controlling for body size using masses of individuals from which the BMR data were obtained instead of species 187 188 average female body mass (t_{40} =3.27, p<0.01). With litter size and developmental 189 state, and their interaction, added as predictors in the model, neonate brain size was 190 still significantly positively related to gestation length (t_{38} =2.94, p<0.01) and BMR 191 $(t_{39}=2.21, p<0.05)$, but unrelated to litter size $(t_{38}=-1.03, p>0.1)$, developmental state 192 $(t_{38}=-0.61, p>0.5)$, and their interaction $(t_{38}=-0.67, p>0.5)$. Gestation length and BMR 193 were uncorrelated after controlling for female body mass (t_{42} =-1.67, p>0.1). Hence, 194 these results are consistent with the hypothesis that BMR constrains neonate brain

size directly, via effects on fetal brain growth rate, rather than indirectly, through
effects on gestation length²².

197

198 **Correlates of postnatal brain growth.** The relative amount of postnatal brain 199 growth (controlling for effects of postnatal body growth) is associated with lactation 200 duration (Table 3). Litter size was not significantly related to postnatal brain 201 growth (model 2, table 3). Mirroring the analyses of neonatal brain size, gestation 202 length was added to the predictors to check that the apparent effect is specific to 203 lactation length. Postnatal brain growth remained significantly associated with 204 lactation and was unrelated to gestation length (model 3, Table 3). Similarly, the 205 effect of lactation length remains significant when either age at first reproduction or 206 juvenile period is added as a predictor (models 3 and 4, Table 3), indicating that it is 207 specifically prolongation of lactation, rather than a general slowing of postnatal 208 maturation, that is associated with increased postnatal brain growth. The test 209 comparing model 4 (including juvenile period) to model 1 is non-significant 210 $(LR_1=3.02, p>0.05)$, reinforcing the lack of an independent effect of juvenile period. 211 The addition of developmental state at birth, litter size and their interaction to the 212 predictors in model 1 (Table 3) revealed no main effects (developmental state, t₈₉=-213 0.30, p>0.5; litter size, -0.12, p>0.5) or interaction (t_{89} =-0.09, p>0.5). Hence, 214 controlling for allometry, postnatal brain growth is robustly associated with 215 lactation length and not with litter size, developmental state, or juvenile period. As 216 was the case for prenatal development, in all these analyses brain growth is

217 positively associated with body growth, hence showing no signs of a trade-off218 between neural and somatic growth.

219 Although age at first reproduction was unrelated to postnatal brain growth 220 when lactation was in the model, if lactation was removed from the predictors, age 221 at first reproduction became significant ($t_{92}=2.70$, p<0.01). This is consistent with 222 the prediction of developmental costs hypotheses that the correlation between large 223 brains and later age at first reproduction is a consequence of prolonged maternal 224 investment. The specific association between brain growth and lactation is further 225 reinforced when a similar model is run for the post-lactation juvenile period, as the 226 latter variable remains non-significant even without lactation in the model 227 (t₉₂=1.80, p>0.05).

228 There were no significant associations between postnatal brain growth and 229 milk composition (Table 4; note that the effect of lactation remained significant in 230 this smaller sample). In a smaller subset of the data (n=23) for which daily milk 231 energy intake per offspring was available, there was also no significant association 232 between this variable and postnatal brain growth (controlling for lactation and body 233 growth, t=-0.28, p>0.5). We tried running models with different combinations of 234 milk composition and intake variables, but obtained no significant results (see Table 235 S1 in supplementary information).

Adding BMR to the predictors, postnatal brain growth is significantly
positively related to both lactation (t₃₉=4.14, p<0.001), and BMR (t₃₉=2.84, p<0.05).
However, the association with BMR appears to be driven by *Homo sapiens*, which is

239 a large outlier in the regression of postnatal brain growth on body size and lactation 240 (residual approximately three standard deviations larger than the mean). When 241 humans were excluded from the analysis, there was no significant relationship 242 between postnatal brain growth and BMR (controlling for size with female body 243 mass, t_{38} =1.45, p>0.05; controlling for size using BMR sample body mass estimates, 244 t_{38} =1.10, p>0.05). In addition, even if humans were included, there was no 245 significant association between postnatal brain growth and BMR when BMR sample 246 body masses instead of mean female body mass was used to control for size 247 (t₃₉=0.92, p>0.1). Postnatal brain growth remained positively associated with 248 lactation in all models. Finally, BMR was not associated with lactation, controlling 249 for either maternal body mass (t_{41} =-0.75, p>0.5), or BMR sample body masses (t_{41} =-250 0.08, p>0.5), ruling out an indirect relationship between BMR and postnatal brain 251 growth mediated by length of lactation.

252

253 Is the association between brain size and life history independent of maternal

254 **investment?** Controlling for adult body size, adult brain size is significantly

positively associated with age at first reproduction (t_{80} =3.02, p<0.01). However,

256 inclusion of the duration of maternal investment (gestation+lactation) in the model

provides a significantly better fit (LR₁=11.52, p<0.001, increase in R² from 0.89 to

- 258 0.91). Furthermore, in this improved model, maternal investment is significantly
- associated with brain size (t_{79} =3.53, p<0.001), but age at first reproduction is not
- 260 (age at first reproduction, t_{79} =1.58, p=0.12). Juvenile period (the interval between

261	weaning and sexual maturity) is not significantly associated with brain size either
262	with or without maternal investment in the model (with, t_{79} =1.30, p>0.1; without,
263	t_{80} =1.85, p>0.05), and again the model including maternal investment provides a
264	better fit than that without (LR ₁ =11.52, p<0.001; increase in R^2 from 0.89 to 0.91).
265	Finally, controlling for body size, adult lifespan is positively correlated with brain
266	size (t ₈₀ =2.96, p<0.01, n=85), but inclusion of the duration of maternal investment in
267	the model provides a significantly better fit (LR $_1$ =12.1 , p<0.001, increase in R 2 from
268	0.89 to 0.91), and in this improved model, maternal investment is significantly
269	correlated with brain size (t_{79} =3.52, p<0.001) but adult lifespan is not (t_{79} =1.32,
270	p=0.19).

271

272 **Discussion**

273 Our results suggest that larger brains take longer to grow both pre- and 274 postnatally, resulting in prolonged maternal investment. Whilst not ruling out the 275 idea that large brains facilitate enhanced survival and slower, longer lives through a 276 generalized "cognitive buffer" effect, the specificity of the correlations between 277 brain growth and associated phases of maternal investment, together with the fact 278 that postnatal life histories are uncorrelated with adult brain size after taking 279 maternal investment into account, strongly support the argument that life history 280 correlates reflect the developmental costs of large brains⁹. Our results provide 281 support for both the maternal energy hypothesis^{11,12} and the broader "expensive 282 brain" hypothesis⁵, although, as predicted by Charnov & Berrigan⁹, some of the

283 trade-offs reported previously⁵ appear to be secondary consequences of the 284 fundamental variable of the rate at which mothers can convert energy into offspring. 285 In particular, neither litter size nor its interaction with developmental state added 286 any explanatory power to the statistical models once gestation, lactation and 287 allometry were accounted for. We conclude that brain growth is primarily related to 288 the duration and rate of maternal investment, with the apparent trade-off with litter 289 size, and differences in correlates between altricial and precocial species, being 290 secondary consequences of variability in gestation and lactation. We did however 291 find that precocial species give birth to larger-brained offspring even after 292 controlling for body size and gestation length. This indicates that the rate, as well as 293 the duration, of fetal brain growth is increased in precocial compared to altricial 294 species, and suggests that the state of the offspring at birth is not entirely 295 determined by the length of gestation.

296 We found no evidence of trade-offs between brain growth and body growth 297 either pre- or postnatally, nor between the amount of brain growth pre- versus 298 postnatally. Indeed, relative amounts of pre- and postnatal brain growth are 299 uncorrelated, consistent with independent genetic control of these two phases of 300 brain growth¹⁹ and suggesting that they have additive rather than either 301 multiplicative or mutually compensating effects on adult brain size. These findings 302 raise the important questions for future research of what structural and functional 303 implications follow from evolutionary changes in pre-versus postnatal brain 304 growth, and whether changes in the two different phases are associated with 305 different selection pressures.

306 Models of life history evolution have tended to assume that organisms vary 307 along a single "slow-fast" continuum, implying that different components of life 308 history such as growth, reproductive rate and lifespan, are tightly interlinked, and 309 thus that ratios between them are invariant across taxa²³⁻²⁴. This view has recently 310 been challenged on both theoretical and empirical grounds²⁵⁻²⁶. Empirically, 311 dissection of mammalian life history variation using phylogenetic factor analyses 312 identified two distinct dimensions²⁵. The first loads heavily on gestation length, 313 neonate size and – though less consistently - on litter size. The second factor loads 314 heavily on inter-birth interval, age at weaning and age at sexual maturity. Our 315 results suggest that brain size may be a key consideration in understanding how 316 such life history traits evolved, and we note that the two factors identified²⁵ 317 correspond broadly to pre- and postnatal influences on brain growth respectively. 318 We predict that neonatal brain size would load heavily on the first factor and 319 postnatal brain growth on the second. Although explanations of life history 320 evolution have focused on body size and environmental factors such as mortality, 321 brain size may represent an intrinsic factor whose role has so far been under 322 appreciated⁴.

Our results clarify the long-disputed relationship between brain size and metabolic rates. The maternal energy hypothesis^{11,12} suggests that basal metabolic rates constrain maternal investment in brain growth, but direct evidence linking BMR to neonate brain size has been lacking, with the only analysis of those variables finding no relationship¹⁶. Our analysis shows that neonate brain size correlates positively with BMR after taking phylogeny, allometry and gestation length into 329 account. Since the correlation is evident when controlling for gestation length, it 330 supports the hypothesis that the metabolic rate of the mother constrains the rate of 331 brain growth in the foetus¹². The finding is also consistent with the hypothesis that 332 the correlation between brain size and BMR is a placental (but not marsupial) trait 333 "related to the intimate physiological contact between mother and offspring during 334 gestation"⁸. The hypothesis that metabolic rate influences prenatal brain growth 335 through an effect on gestation length²² was however, not supported; there was no 336 significant correlation between BMR and gestation length after controlling for other 337 factors. The restriction of an effect of BMR to the prenatal period together with the 338 significant effects of other maternal investment variables operating at least partly 339 independently of one another also clarifies why the positive association between 340 BMR and adult brain size is relatively weak^{13,}.

341 Although it has been suggested that the structure of the placenta might 342 influence nutrient transfer and hence prenatal brain growth^{15,27}, recent comparative 343 studies find no evidence for a specific relationship between placental structure and 344 brain growth²⁸⁻²⁹. 'Labyrinthine' placentas, in which maternal and fetal tissues are 345 highly interdigitated, are associated with shorter gestations but no difference in 346 neonate brain or body size, suggesting that fetal growth rates are faster in species 347 with labyrinthine placentas²⁸. However, there was no difference in the relative brain 348 size of neonates, indicating that higher growth rates are not targeted specifically at 349 the brain²⁸. How higher metabolic rates are translated into additional physiological 350 support for fetal brain growth is thus an important and so far unanswered question.

One possibility is that energy turnover constrains the ability of the mother to supply
the fetus with specific nutrients, such as long-chain fatty acids²⁶.

353 Similarly, although relative rates of postnatal brain growth are likely to vary, 354 we were unable to find any relationship between brain growth and milk 355 composition, milk energy value or daily milk energy intake at peak lactation. This 356 finding agrees with the observation that convergent evolution of large brain size and 357 extended postnatal brain growth in humans and capuchin monkeys (*Cebus apella*) 358 has not resulted in convergence in milk composition³⁰. However, sample sizes were 359 relatively small in our analyses of milk composition in relation to postnatal brain 360 development, and re-analysis with larger data sets when these become available 361 would be interesting, as would analysis of specific nutrients that may play a role in 362 postnatal brain development.

363 The issue of evolutionary changes in rate versus duration of brain growth is 364 important for understanding the developmental basis of human brain evolution. 365 Most discussions of this subject assume that large relative brain size in humans was 366 developmentally achieved via an exceptional prolongation of postnatal brain 367 growth, creating enhanced opportunities for environmental input to the developing 368 brain³¹⁻³³. A re-analysis by Vinicuis³⁴ however shows that the ways in which human 369 brain and body growth patterns depart from those of other primates are more 370 complex than this, including at least three distinct mechanisms: a moderate 371 extension of postnatal brain growth, a derived developmental allometry and a 372 retardation of postnatal body growth. The first mechanism fits the general link

373 between lactation and postnatal brain growth reported here, and suggests that brain 374 size may be a better predictor of the "natural" weaning age for humans than is body 375 size. Vinicius' second mechanism³⁴ suggests a difference in the rate of brain growth 376 between humans and other anthropoids, congruent with our finding that variation 377 in brain growth rates, as well as durations, contribute to adult brain size. As we note 378 above, the physiological mechanisms that co-vary with brain growth rates remain 379 unknown. Finally, Vinicius' third mechanism³⁴ implies a trade-off between 380 postnatal brain and body growth; we found no evidence for this as a general pattern 381 among eutherian mammals, so its occurrence in humans must be presumed to be 382 evolutionarily unusual.

383 In conclusion, our results emphasize the energetic costs of brain 384 development as a driver of associations between brain size and life history in 385 mammals. Whilst large brains undoubtedly confer benefits, we found no support for 386 hypotheses predicated on specific associations between brain size and either 387 juvenile period³⁵ or adult lifespan⁶⁻⁷. It is still possible that large brains operate as 388 "cognitive buffers", since the selective advantage of slower growing, larger brains 389 may be reduced mortality mediated by cognitive capacities^{4,7}. However, the 390 cognitive buffer hypothesis as formulated assumes that such cognitive capacities are 391 'domain general', facilitating survival and long lifespans through increased 392 behavioral flexibility⁶. The lack of a significant association between brain size and 393 adult lifespan after controlling for maternal investment suggests that it is not 394 specifically lifespan and an associated need for flexibility that drives the patterns, 395 undermining the link made between life history correlations of brain size and

domain-general cognitive benefits⁶. Given that brain size evolution in mammals is
associated with a variety of specific neural systems and structures^{36-38,} domainspecific mechanisms should be given equal consideration in attempts to establish
the cognitive benefits that offset the developmental costs of large brains.

400

401 Methods

402 **Data.** We extracted data from the literature on 128 eutherian mammal species as follows. 403 (i) Brain and body masses: neonate brain and body mass, adult brain and body mass, 404 maternal (adult female) body mass (all in grams). Postnatal growth (brain or body) was 405 calculated as the difference between adult size and neonate size. (ii) Developmental, 406 maternal investment and life history variables: litter size (number of offspring per litter), 407 developmental state at birth (altricial if eyes closed at birth, versus precocial if eyes open at 408 birth), duration of gestation (days), duration of lactation (days), milk composition (as % of 409 fats, proteins and sugars) and milk energetic value (as sum of the energy provided by its 410 components, given milk composition; in KJ) both at peak lactation, daily milk energy intake 411 (milk energetic value multiplied by daily milk intake in ml/day at peak lactation; in KJ/day), 412 age at first reproduction (days), lifespan (days). (iii) Basal metabolic rate (BMR, ml 413 O_2 /hour) together with body masses for the animals from which the BMR data were taken 414 (Body mass_{BMR}, in g). We used only estimates of BMR that fulfilled the requirements of the 415 protocol described in McNab³⁹ (measurement in thermoneutral environment, on adult non-416 reproducing individuals, quietly resting and post-absorptive). Further details of data and 417 sources and the full data set are provided in supplementary information (Text S1 and 418 Dataset_S1).

419 **Statistical analysis.** We investigated the correlated evolution of brain size, body size. 420 maternal investment and life history variables using phylogenetic generalized linear models 421 (PGLM)⁴⁰, which allowed us to incorporate phylogeny into statistical models⁴⁰⁻⁴². In PGLM 422 analysis, regression parameters are found by maximum likelihood (ML) and 'weighted' by 423 the variance-covariance matrix that represents the phylogenetic relationships among the 424 species. In each regression the phylogenetic signal is estimated as the value of λ of the 425 residuals, varying between 0 (where the data have no phylogenetic structure) and 1 (where 426 the best fit to the data is provided by a 'Brownian Motion' model of trait evolution⁴³, with 427 variation at the tips proportional to the duration of common evolution^{42,44} We report λ 428 values tests for significant departure from either 0 or 1 for each analysis. The estimated ML 429 value of λ is incorporated as a parameter in the model, thus controlling for phylogenetic 430 dependence in the data. Incorporating a discrete binary trait, such as developmental state, 431 as a predictor in regression models amounts to a phylogenetic ANCOVA. In the PGLM 432 framework, more complex models can be compared to simpler models to investigate 433 whether incorporating additional variables of interest provides a better fit to the data. Our 434 tables of results indicate which variables were included in each model, significance tests for 435 these variables, and overall model parameters: values of λ , r-squared values and log-436 likelihood scores. Alternative nested models are compared using the likelihood ratio (LR) 437 test (where LR=-2[log-likelihood(better fitting model)-log-likelihood(worse fitting model)], 438 the best fitting model having the highest log-likelihood score) whose significance is 439 evaluated against a χ^2 distribution with degrees of freedom corresponding to the difference 440 in the number of parameters between the two competing models^{40,44}. All statistical tests 441 were 2-tailed with α -level of significance set at 0.05. These analyses were carried out using 442 the CAIC R package (R v.2.11.1, The R Foundation for Statistical Computing, http://ww.R-443 project.org), available from http://r-forge.r-project.org/projects/caic. The phylogeny

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445

(including branch lengths) for the species in our dataset was extracted from a published supertree of mammals⁴⁵⁻⁴⁶

446 Continuous variables were \log_{10} -transformed to improve normality, with the 447 exception of milk composition (%) data which were square-rooted and then arcsine-448 transformed⁴⁷. Because, brain mass can be a substantial proportion of overall body mass in 449 neonates analyses of these variables could potentially be influenced by autocorrelation and 450 consequent issues of collinearity. Likewise, age at first reproduction includes the period of 451 lactation and lifespan includes the period up to age at first reproduction. We therefore ran 452 analyses based on non-overlapping measures [neonate body mass with brain mass 453 subtracted, age at first reproduction with lactation subtracted (=juvenile period), and 454 lifespan with age at first reproduction subtracted (=adult lifespan]. We do include some 455 analyses in which age at first reproduction and lactation appear as predictors in the same 456 model for comparability with other studies that use this variable, whilst noting the issue of 457 autocorrelation. Bivariate plots and residuals were examined to check for violation of 458 homogeneity of variance. We checked for the effects of outliers by re-running analyses after 459 deleting data points generating large residuals (greater than the mean by three standard 460 deviations or more). However, removing outliers qualitatively affected conclusions in only 461 one case. This outlier was caused by a data point for humans. Because humans are 462 particularly large brained they potentially exert high leverage on regressions; hence we re-463 ran analyses with and without humans, but the outcome was affected in just the one case 464 mentioned above, so we report results including humans unless otherwise stated.

465

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574 Figure legends

575 Figure 1. Association between relative brain size of neonate and adult mammals.

576 Encephalization scores are the residuals from phylogenetic generalized linear

- 577 models for brain size on the appropriate body size (either neonate or adult). See
- 578 Table 1 for results of statistical analysis.

579

580 Table legends

- Table 1. PGLM analysis of pre- and postnatal contributions to adult brain size,
- 582 controlling for body size. Variables not included in a particular model are indicated
- 583 by blank entries in the Table. Significant predictors of adult brain size indicated in

bold type. Lh=maximized log-likelihood, λ =estimated ML value of lambda

- 585 (phylogenetic signal) which is included as a parameter in the models, with p-values
- for tests of statistical difference from a model with no phylogenetic signal ($p(\lambda=0)$)
- 587 and a model with $\lambda = 1$ (p($\lambda = 1$).

 588
 Table 2. PGLM analysis of neonate brain size. Significant predictors of neonate brain

 500
 Image: Significant predictors of neonate brain

- size indicated in bold type. Other details as for Table 1.
- Table 3. PGLM analysis of postnatal brain growth. Significant predictors of postnatal
- 591 brain growth indicated in bold type. Other details as for Table 1.

- Table 4. PGLM analysis of postnatal brain growth, lactation and milk composition.
- 593 Significant predictors of postnatal brain growth indicated in bold type. Other details
- as for Table 1.

Tuble I.	Tab	le 1.
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Model (n=122)	Model 1	Model 2	Model 3
Parameter	t, p-value	t, p-value	t, p-value
Intercept	-3.1, <0.01	-2.88, <0.01	4.75, <0.001
Neonatal brain size	6.82, <0.001	6.07, <0.001	17.12, <0.001
Adult body size	9.61, <0.001	8.77, <0.001	3.95, <0.001
Neonatal body mass	-	-1.54, 0.13	-
Postnatal brain growth		-	31.2, <0.001
λ	0.79	0.74	0.70
p(λ=0)	<0.001	<0.001	< 0.001
p(λ=1)	<0.001	<0.01	< 0.001
Model summary			
Lh	55.39	56.29	190.35
Adjusted R ²	0.917	0.923	0.991

Table 2.

Model (n=128)	Model 1	Model 2
Parameter	t, p-value	t, p-value
Intercept	-2.60, <0.001	-2.49, <0.001
Neonatal body mass	6.05, <0.001	6.00, <0.001
Maternal body size	3.13, <0.01	3.25, <0.01
Gestation length	7.20, <0.01	6.38, <0.001
Litter size	-	-1.14, >0.1
λ	0.90	0.90
p(λ=0)	<0.001	<0.001
p(λ=1)	<0.01	<0.01
Model summary		
Lh	58.06	58.65
Adjusted R ²	0.92	0.92

Ta	ble	3.

Model (n=96)	Model 1	Model 2	Model 3	Model 4	Model 5
Parameter	t, p-value				
Intercept	-8.41, <0.001	-8.13, <0.001	-5.56, <0.001	-8.54, <0.001	-9.64, <0.001
Postnatal body growth	17.60, <0.001	17.47, <0.001	13.89, <0.001	14.13, <0.001	14.67, <0.001
Lactation	3.83, <0.001	3.78, <0.001	3.75, <0.001	3.06, <0.01	3.80, <0.001
Litter size	-	0.18, >0.5	-	-	-
Gestation	-	-	-0.05, p>0.5	-	-
Age at first reproduction	-	-	-	1.69, >0.1	-
Juvenile period	-	-	-	-	1.82, >0.05
λ	0.67	0.67	0.67	0.60	0.57
p(λ=0)	<0.01	<0.01	<0.01	>0.05	>0.1
p(λ=1)	<0.001	<0.001	< 0.001	<0.001	< 0.001
Model summary					
Lh	1.01	1.03	1.01	2.38	2.52
Adjusted R ²	0.85	0.85	0.85	0.86	0.87

Model	Model 1	Model 2	Model 3	Model 4	Model 5
(n=48)					
Parameter	t, p-value				
Intercept	-6.94, <0.001	-8.28, <0.001	-6.25, <0.001	-7.28, <0.001	-7.42, <0.001
Postnatal body growth	13.87, <0.001	13.65, <0.001	15.02, <0.001	14.09, <0.001	14.56, <0.001
Lactation	4.20, <0.001	4.19, <0.001	4.19, <0.001	4.17, <0.001	4.31, <0.001
% dry matter	0.81, >0.1		-	-	-
% fat	-	0.71, >0.1	-	-	-
% protein	-	-	-0.24, >0.5	-	-
% sugar	-	-	-	-0.04, >0.5	-
Milk energy	-	-	-	-	0.66, >0.5
λ	0.10	0.14	0.25	0.23	0.16
p(λ=0)	>0.5	>0.5	>0.1	>0.1	>0.5
p(λ=1)	<0.001	<0.001	<0.001	<0.001	< 0.001
Model summary					
Lh	0.55	0.52	0.33	0.30	0.51
Adjusted R ²	0.92	0.92	0.90	0.91	0.91

Table 4.

