1 Theropod dinosaurs had primate-like numbers of telencephalic neurons

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7 Abstract

8 Understanding the neuronal composition of the brains of dinosaurs and other fossil 9 amniotes would offer fundamental insight into their behavioral and cognitive capabilities, but 10 brain tissue is only rarely fossilized. However, when the bony brain case is preserved, the volume 11 and therefore mass of the brain can be estimated with computer tomography; and if the scaling 12 relationship between brain mass and numbers of neurons for the clade is known, that 13 relationship can be applied to estimate the neuronal composition of the brain. Using a recently 14 published database of numbers of neurons in the telencephalon of extant sauropsids (birds, squamates and testudines), here I show that the neuronal scaling rules that apply to these 15 16 animals can be used to infer the numbers of neurons that composed the telencephalon of 17 dinosaur, pterosaur and other fossil sauropsid species. The key to inferring numbers of 18 telencephalic neurons in these species is first using the relationship between their estimated 19 brain and body mass to determine whether bird-like (endothermic) or squamate-like 20 (ectothermic) rules apply to each fossil sauropsid species. This procedure shows that the notion 21 of "mesothermy" in dinosaurs is an artifact due to the mixing of animals with bird-like and 22 squamate-like scaling, and indicates that theropods such as Tyrannosaurus and Allosaurus were 23 endotherms with baboon- and monkey-like numbers of telencephalic neurons, respectively, 24 which would make these animals not only giant but also long-lived and endowed with flexible 25 cognition, and thus even more magnificent predators than previously thought.

26

27 Introduction

The modern mammal and bird-rich amniote fauna arose from the opportunity created by the demise, in a catastrophic astronomical event, of the giant archosaur species that dominated the Earth during the Mesozoic (Alvarez et al., 1980; Bininda-Emonds et al., 2007; Yu et al., 2021). Until then, sauropodmorphs (the long-necked, plant-eating, quadruped dinosaurs), theropods

32 (the bipedal, carnivorous dinosaurs) and ornithischians (the sometimes armored, plant-eating 33 dinosaurs; Langer et al., 2017) were the largest animals on land. They also had the largest brains 34 amongst land animals at the time, approaching 200 g (the size of a lion's brain) in 35 sauropodmorphs like Giraffatitan brancai (Brachiosaurus) and surpassing that in theropods such 36 as Tyrannosaurus rex (up to 343 g; Hurlburt, 1996; Balanoff et al., 2013; Table 1). However, the 37 fact that dinosaurs were sometimes gigantic has been used to imply that their rather large brains were actually undersized for their bodies, giving those animals very low encephalization 38 39 quotients (EQ) – the first metric to set humans apart from other species, above all others, with a 40 much larger brain than expected for our body size (Jerison, 1973). With low EQs, large dinosaurs 41 could presumably not be that cognitively competent (Jerison, 1973; Hopson, 1977; Knoll and 42 Schwarz-Wings, 2009; Rowe et al., 2011; Ksepka et al., 2020; Balanoff et al., 2013). Rather, it was 43 small theropods such as troodontids, with relatively larger brains and thus higher EQs, that were 44 considered to be "smarter" (Jerison, 1973).

45 The EQ is the ratio between the actual brain volume (or mass; these variables are 46 interchangeable, given the specific density of the brain of approximately 1.0 mg/ml) and the brain 47 volume mathematically expected for a species of a given body mass (Jerison, 1973). Calculating 48 the EQ for a species is done under the assumption that brain and body mass are universally 49 correlated across a wide range of species; interpreting the EQ as informative of cognitive 50 capacities, in turn, assumes that brain mass is a universal predictor of numbers of neurons in the 51 brain, and in the pallium (cerebral cortex, in mammals) in particular, the brain structure that most 52 decidedly confers flexibility and complexity to animal behavior (Jerison, 1973). The problem is 53 that while those were reasonable assumptions until the 2000s, we now know that they are both 54 incorrect.

Recent analyses of large datasets show that brain and body size evolve separately in both mammalian (Bertrand et al., 2022) and bird (Ksepka et al., 2020) evolution, making body size an unreliable universal predictor of brain mass. In parallel, a new line of investigation created in my lab based on an original non-stereological method to count brain cells, the isotropic fractionator (Herculano-Houzel and Lent, 2005), allowed the realization that there is no mandatory, universal relationship between body mass and numbers of brain neurons, not even in the brainstem

61 structures that operate the body (Herculano-Houzel et al., 2014, 2015a; Herculano-Houzel, 62 2017). Additionally, there is no universal relationship between the size of a brain structure and 63 its numbers of neurons; different scaling relationships apply to different clades of mammals and 64 birds (Herculano-Houzel et al., 2014; Herculano-Houzel, 2016), while scaling relationships are 65 mostly shared across reptiles (Kverkova et al., 2022; Herculano-Houzel, 2022). As a result of the lack of a universal correlation between body mass and brain composition, when numbers of 66 67 neurons (the signal processing units of circuits) should be the limiting factor that determines the computational capacity of a network (Williams and Herrup, 1980), bringing body mass into 68 69 comparisons of brain size across species of different clades is more than uninformative; it 70 muddles interpretation, by bringing a confounding factor into the mix.

71 Rather, simple, absolute numbers of neurons in the pallium (organized as a cortex in 72 mammals) are a much better proxy for cognitive abilities than brain or pallial size, whatever the 73 size of the body (Herculano-Houzel, 2017). These are the signal processing units that make 74 behavior flexible and complex and should thus constitute a primary determinant of signal 75 processing capacity (Williams and Herrup, 1980; Herculano-Houzel, 2017; Ströckens et al., 2022). 76 Absolute numbers of pallial neurons are a better predictor of flexible cognitive control than brain 77 mass or EQ across birds and primates alike (Herculano-Houzel, 2017), and, across bird species, 78 they are a great predictor of innovation rate (Sol et al., 2022). It thus follows that understanding 79 the capability for behavioral and cognitive flexibility of the extinct species that once dominated the Earth's fauna requires going beyond the veil of brain and body size and gaining direct 80 81 understanding of the numbers of neurons that composed the pallium of those animals. That 82 understanding can be reached once the scaling relationship between brain size and numbers of 83 neurons for species in a given clade is known, which makes brain size a reliable proxy of numbers 84 of neurons in that clade. In particular, numbers of pallial, or telencephalic, neurons in extant 85 species can be estimated from brain size by clade-specific allometric power functions with r² 86 values of typically 0.9, that is, which provide estimates with about 90% reliability (Herculano-87 Houzel, 2019a). Using this expedient, we have previously been able to infer the numbers of 88 neurons that composed the brain of prehistoric hominin species (Herculano-Houzel and Kaas, 89 2011) and fossil mammals (Herculano-Houzel et al., 2011).

90 How to determine the scaling relationships that applied to fossil species, when brain 91 tissue is not preserved in the fossil record? Brain mass of dinosaur species can be estimated with 92 CT or micro-CT of extant or fossilized skulls (Hulburt, 1996; Knoll et al., 1999; Witmer et al., 2003; 93 Balanoff et al., 2013). In these cases, the numbers of neurons that composed their brains, and 94 their telencephalon in particular, can be estimated if one can determine the applicable predictive 95 equations relating numbers of neurons to brain mass. Dinosaurs and pterosaurs (together with 96 turtles and crocodilians) were archosaurs, the sister clade to modern squamates, which are 97 ectothermic sauropsids; but living birds, which are endothermic sauropsids, are surviving 98 dinosaurs. Thus, the alternative hypotheses tested here are that pterosaur and dinosaur brains 99 were either composed like ancestral, and modern, ectothermic sauropsid brains, or had already 100 shifted to the composition of modern, endothermic, early-branching bird brains (Herculano-101 Houzel, 2022).

102 Here I determine whether ectothermic or endothermic sauropsid scaling rules most likely 103 applied to different dinosaur (and pterosaur) species, and use the allometric scaling equations 104 that apply to the brain structures of modern birds and non-avian sauropsids to predict the 105 numbers of neurons that composed the brains of fossil dinosaurs and pterosaurs based on their 106 brain volumes. These allometric scaling equations describe the relationship between brain 107 structure mass and numbers of neurons, determined using the isotropic fractionator (Herculano-108 Houzel and Lent, 2005). This method, which consists of turning brains or brain structures into a 109 homogeneous soup of floating cell nuclei that allows the fast, unbiased, and reproducible 110 estimation of how many neuronal and non-neuronal cells composed those structures (Herculano-111 Houzel et al., 2015b), has by now been applied to over 200 species of mammals, birds, and non-112 avian sauropsids (Kverkova et al., 2022). While the resulting dataset did not separate the pallial 113 from the subpallial structures that together compose the telencephalon of non-avian sauropsids, 114 it did establish that the vast majority of telencephalic neurons are found in the pallium of early-115 branching birds; thus, numbers of telencephalic neurons in the dataset offer a good 116 approximation for the number of pallial neurons (though the subpallium also contributes to 117 flexible behavior; Boot et al., 2017). In contrast to the recent initial analysis of the full dataset 118 which focused on relationships between numbers of neurons and body mass (Kverkova et al.,

2022), here I concentrate on the clade-specific relationships between telencephalic and brain
mass and numbers of telencephalic neurons in the different clades of living avian- and non-avian
sauropsids in search of establishing what relationships putatively applied to fossil species, which
I then use to estimate numbers of telencephalic neurons in select species with known brain
volume and mass.

- 124
- 125 Methods

126 All data on numbers of telencephalic neurons, brain and telencephalic mass, and body 127 mass for 174 extant sauropsid species (avian and non-avian) used to calculate the scaling 128 relationships in Figure 1 were taken from Kverkova et al. (2022). All power functions were 129 calculated using least-squares regression of log-transformed values using the JMP 16 software 130 package (Carey, NC). Power functions were calculated separately for different groups of birds, as 131 detailed in the results (Accipitriformes, n=4 species; Anseriformes, n= 7; Columbiformes, n=5; Falconiformes, n=3; Galliformes, n=9; Palaeognathae, n=6 species; Passeriformes, n=13; 132 133 Psittaciformes, n=11; Strigiformes, n=7); and for all non-avian sauropsids in the dataset (88 134 squamates, 19 testudines and 1 crocodilian species; Kverkova et al., 2022). Particular emphasis 135 was placed on the early-branching ("pre-K-Pg") bird species, which in this dataset belong to 136 Palaeognathae, Galliformes, Anseriformes and Collumbiformes, which share scaling relationships 137 (see Figure 1, and Herculano-Houzel, 2022). Thus, all scaling relationships reported pertain to a 138 group of species that belong to closely related clades that share a scaling relationship amongst 139 themselves. Within each group, I chose to not adjust the scaling exponent for phylogenetic 140 relatedness in the dataset because the point of calculating these scaling relationships was not to 141 study phylogeny, but to use the mathematical functions to make predictions from brain mass, in 142 which case the desired scaling relationship is the one that applies to the raw data, with no 143 modifications introduced that would distort the mathematical reality of the relationships 144 between the physical entities (structure mass) and (number of neurons).

Absolute estimated brain mass (not simply endocranial volume) and body mass values in fossil pterosaur and dinosaur species shown in Figure 2 were collected from four studies that compiled estimates from CT studies of endocranial volume (Table 10 in Hulburt, 1996; Witmer et al., 2003; Hulburt et al., 2013; Balanoff et al., 2013). Different brain-to-endocast fractions by

149 definition will impact the estimated brain mass or volume (Knoll et al., 2022); thus, care was 150 taken to use explicitly reported estimates for brain mass, whether or not that matched the 151 endocranial volume. Estimates for theropod and pterosaur species consider that theropod and 152 pterosaur brains filled the endocranial cavity (Balanoff et al., 2013; Witmer et al., 2003), whereas 153 Hurlburt's estimates for all dinosaurs except small theropods considered that brain mass was half 154 that of endocast volume (Hurlburt, 1996, p. 147). Where available, specimen numbers are 155 provided in Table 1 (they were not reported in Hurlburt's 1996 dataset). Where more than one 156 estimate was available for a species, all values are plotted so as to allow the evaluation of the 157 impact of specimen and methodological variability, as well as the range of body and brain size 158 combinations. I note that the taxonomy of several fossil species has been updated since 159 Hurlburt's data compilations that I use in Table 1. For the sake of making the present dataset 160 transparent and readily verifiable by other researchers and useful to them, and because I am not 161 a paleontologist but a neuroscientist using paleontological data, I chose to preserve in Table 1 162 the original names under which the data were originally reported by the authors.

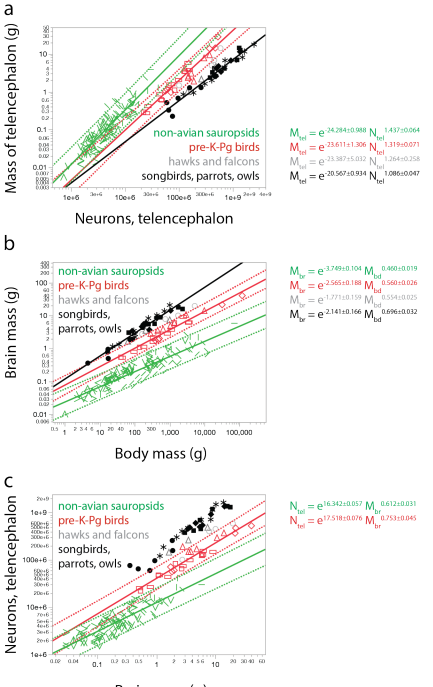
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165 Results

166 Figure 1a shows that the neuronal scaling rules that apply to the telencephalon of modern 167 birds and non-avian sauropsids are clearly distinct. For a similar telencephalic mass (which occurs 168 in the largest non-avian sauropsids and the smallest birds), early-branching bird clades in the 169 dataset (Palaeognathae, Galliformes, Anseriformes, and also Columbiformes, in red), which arose 170 before the K-Pg boundary (henceforth, pre-K-Pg birds; Brusatte et al., 2015), have ca. 5 times as 171 many telencephalic neurons as non-avian sauropsids (in green), and the post-K-Pg-branching 172 Passeriformes, Psittaciformes and Strigiformes clades have even more telencephalic neurons (in 173 black). For instance, the zebra finch has 55 million telencephalic neurons whereas the Sudan 174 plated lizard has only 14 million, though both have a telencephalon of ca. 0.3 g. Strikingly, there 175 is very little overlap in numbers of telencephalic neurons between birds and non-avian 176 sauropsids, a distinction that I have hypothesized to result from the increased oxidative rates 177 that make endothermy possible in birds compared to other sauropsids (Bennett and Ruben, 178 1979) rather than from endothermy itself (Kverkova et al., 2022; Herculano-Houzel et al., 2022). 179 Figure 1b shows that endothermic sauropsids (i.e., birds) do have larger brains compared to 180 ectothermic sauropsids of a similar body mass, but again in a clade-specific manner, such that 181 bird species belonging to post-K-Pg clades (songbirds, parrots, and owls) have even larger brain 182 mass for a similar body mass.

183 Together, the distinction between bird clades in Figures 1a and 1b shows that a shift to 184 endothermy cannot be the sole cause of increased brain mass and numbers of telencephalic 185 neurons in birds relative to body mass (Herculano-Houzel, 2022; Gillooly and McCoy, 2014), since 186 brain mass increases further relative to body mass in the post-K-Pg songbirds, parrots and owls 187 compared to the pre-K-Pg birds. Importantly, these findings establish that comparisons across species and clades for where brain scaling is involved cannot treat "birds" as a single entity, as 188 189 has been standard in the field (Balanoff et al., 2013). However, amongst non-avian sauropsid 190 clades, neuronal scaling rules are much more uniform (Herculano-Houzel, 2022), and for the 191 purposes of this study, all extant non-avian sauropsids in the dataset (88 squamates, 19 192 testudines and 1 crocodilian species) can be considered to share the scaling rules of interest, 193 which are clearly distinct from the scaling rules that apply to extant pre-K-Pg birds (Figure 1a).

194 Expressing the number of telencephalic neurons in the brain as a function of brain mass 195 shows that within a clade, brain mass has strongly predictive power to arrive at estimates of 196 numbers of telencephalic neurons in a brain of known mass, once the neuronal scaling rules that 197 presumably apply are known. Figure 1c shows that clearly different predictive scaling rules apply 198 to extant pre-K-Pg birds and to non-avian sauropsids, with non-overlapping 95% prediction 199 intervals across the entire range of bird-like brain sizes. Specifically, these distinct power laws are 200 such that over 80% of the variation in numbers of telencephalic neurons in non-avian sauropsid 201 species, and over 90% in pre-K-Pg birds, can be accounted for by the variation in brain mass, if 202 clade identity is respected.



Brain mass (g)

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Figure 1. Scaling relationships that apply to extant avian and reptile species. A, clade-specific scaling of
 telencephalic mass (M_{tel}) with numbers of telencephalic neurons (N_{tel}) distinguishes pre-K-Pg bird clades
 (Palaeognathae, red lozangles; Galliforms, red rectangles; Anseriformes, red triangles; Columbiformes,
 red squares) from post-K-Pg bird clades (Falconiformes, grey triangles; and Accipitriformes, grey circles,

210 i.e. hawks and falcons; and Passeriformes, black circles; Psittaciformes, black asterisks; and Strigiformes, 211 black squares, i.e. songbirds, parrots, and owls; and non-avian sauropsids (including Testudines and 212 Crocodilia). Power functions are indicated next to the graph for each group (non-avian sauropsids in green, 213 with different symbols for the different clades in Kverková et al. (2022), r²=0.826, p<0.0001, n=108; pre-214 K-Pg birds in red, r²=0.933, p<0.0001, n=27 species; Accipitriformes and Falconiformes, fit not plotted for 215 clarity, $r^2=0.828$, p=0.0045, n=7 species; Passeriformes, Psittaciformes and Strigiformes in black, $r^2=0.949$, 216 p<0.0001, n=31 species). B, clade-specific scaling of brain mass with body mass similarly distinguishes pre-217 K-Pg bird clades (Palaeognathae, Galliforms, Columbiformes, Anseriformes) from other, later-branching 218 bird clades (Falconiformes and Accipitriformes; and Passeriformes, Psittaciformes, and Strigiformes), and 219 non-avian sauropsids. Power functions as in A: non-avian sauropsids plotted in green, $r^2=0.842$, p<0.0001, 220 n=108 species; pre-K-Pg birds plotted in red, r²=0.948, p<0.0001, n=27 species; Accipitriformes and 221 Falconiformes, in grey, fit not plotted for clarity, $r^2=0.990$, p<0.0001, n=7 species; and Passeriformes, 222 Psittaciformes and Strigiformes plotted in black, $r^2=0.944$, p<0.0001, n=31 species. C, clade-specific 223 predictive relationships for estimating numbers of telencephalic neurons from brain mass for pre-K-Pg 224 birds and non-avian sauropsids. Power functions are indicated next to the graph for pre-K-Pg birds (in red; 225 r^2 =0.918, p<0.0001, n=27 species) and for non-avian sauropsids (in green: r^2 =0.791, p<0.0001, n=108 226 species). All data from Kverkova et al., 2022.

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228 The predictive equations shown in Figure 1c can be used to infer the numbers of 229 telencephalic neurons that composed the brains of dinosaur, pterosaur, and other fossil 230 sauropsid species provided that these species are found to conform to the scaling rules that apply 231 to either modern pre-K-Pg birds or non-avian sauropsids. Figure 2, using data compiled from the 232 literature (Table 1), shows that the scaling of brain mass with body mass can indeed provide a 233 distinguishing criterion across dinosaur species. Standard practice in the field has been to assume 234 that a single scaling relationship applies homogeneously across mixed dinosaur clades (Jerison, 235 1973; Balanoff et al., 2013; Grady et al., 2014). Figure 2a confirms that a highly significant single 236 scaling relationship can indeed be fit to the ensemble of the fossil species sampled, with a 95% 237 prediction interval that includes all but one species (Figure 2a), with an exponent of 0.455±0.028 238 (r²=0.857, p<0.0001) that is similar to the exponent that applies to extant non-avian sauropsids only (0.460±0.019, r²=0.842 in Figure 1b) but with a significantly larger intercept of e^{-2.327±0.338} 239 compared to the e^{-3.749±0.104} of modern such species. If this joint scaling relationship intermediate 240

between living non-avian sauropsids and birds truly applied across all dinosaurs, as calculated previously (Balanoff et al., 2013), and these diverse species shared a single scaling relationship between brain and body mass (for example, if they were "mesothermic", as once suggested by a similar expedient of analyzing dinosaur species together regardless of clade; Grady et al., 2014), then it would not be justified to apply the neuronal scaling rules of either modern birds or nonavian sauropsids to these fossil species.

247 In contrast, Figure 2b shows that individual dinosaur and pterosaur specimens clearly 248 conform to the brain x body scaling rules that apply to either ectothermic or pre-K-Pg 249 endothermic extant sauropsids. Both Archaeopteryx, the earliest avian species of known brain 250 and body mass, and a non-identified "protoavis" (Hulburt, 1996; filled red circles) conform to the 251 scaling relationship that applies to modern pre-K-Pg birds, which originated within Jurassic 252 theropods (Brusatte et al., 2015), with brains significantly larger than expected for a modern 253 reptile of similar body mass. Likewise, the majority of theropod dinosaur species of known brain 254 and body mass (filled pink circles) conform to the brain vs body mass relationship that applies to 255 modern pre-K-Pg birds, with the exception of *Shuvuuia deserti* (with brain mass just below the 256 prediction interval for pre-K-Pg birds) and Tsaagan mangas (with the brain mass expected for a 257 modern ectothermic sauropsid of similar body mass; unfilled pink circles). Conversely, most 258 sauropodmorph dinosaurs in the dataset had the brain mass expected for a modern ectothermic 259 sauropsid of their body mass (unfilled green circles). Ornithischian (blue circles) and pterosaur 260 species (black circles), in turn, align either with endothermic pre-K-Pg birds (filled circles) or 261 ectothermic sauropsids (unfilled circles) in their brain vs body mass relationship, depending on 262 the species (Figure 2b, Table 1). Protoceratops (filled blue circle), in particular, approached the 263 distribution of modern pre-K-Pg birds. Thus, the comparison of the brain vs body mass 264 relationships of the sampled fossil species with those of modern ectothermic and endothermic 265 pre-K-Pg sauropsids suggests that the neuronal scaling rules shared by modern ectothermic 266 sauropsid species also applied to the telencephalon of all non-theropod fossil species of 267 sauropsids, with the exception of some pterosaur and ornithischian species (filled data points in 268 Figure 2b), whereas theropods as a whole already had neuronal scaling rules similar to those of 269 modern, endothermic pre-K-Pg sauropsids, according to the cladogram in Figure 2d. Figure 3a

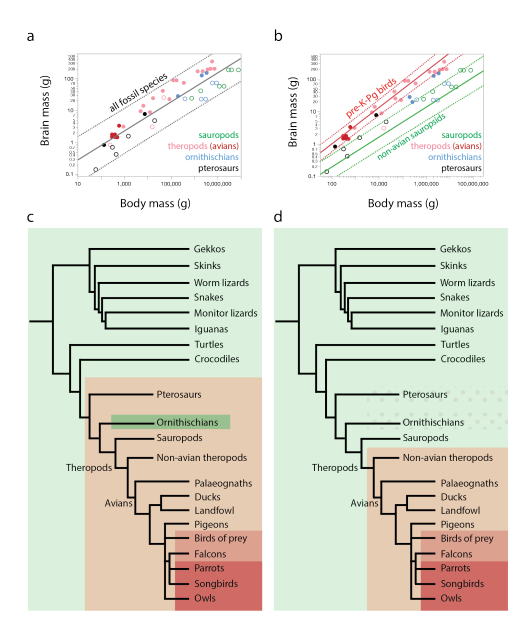
shows how the brain x body mass relationship of fossil avian and non-avian theropods matches

271 that of extant pre-K-Pg bird species (and, in comparison, primates have much larger brains for a

similar body mass). Figure 3a also shows that despite the extreme difference in body size, the

273 brain x body mass relationship of fossil sauropodmorphs, several pterosaurs and some

274 ornithischians match the relationship that applies to extant non-avian sauropsids.

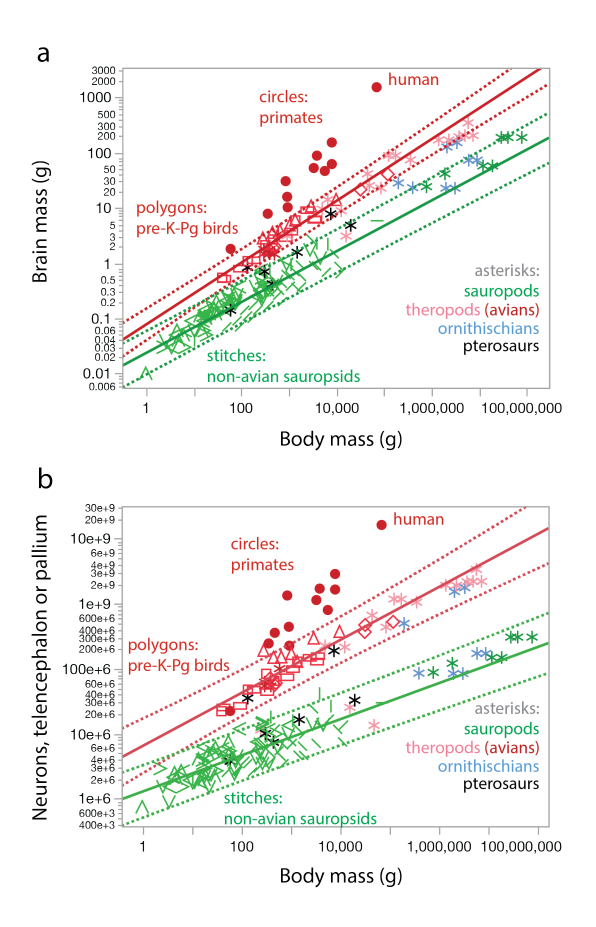


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277 Figure 2. Dinosaur and pterosaur species vary in conforming to either modern pre-K-Pg endothermic

278 (bird) or ectothermic (non-avian sauropsid) scaling relationships between brain and body mass. A, A

single power function $M_{br} = e^{-2.327 \pm 0.338} M_{bd}^{0.455 \pm 0.028}$ (r²=0.857, p<0.0001, plotted in grey) can be fit to the 279 280 relationship between brain and body mass (M_{br} and M_{bd}, respectively) across all the fossil dinosaur and 281 pterosaur species in the dataset (Table 1), color-coded as indicated in the key. However, most non-avian 282 theropod species (pink) have larger brain mass than predicted for their body mass, whereas 283 sauropodmorphs (green) and most pterosaur species (black) have smaller brain mass than predicted by 284 this joint scaling function, which is consistent with a better account of the distribution by two separate 285 functions. Filled circles correspond to species that match brain x body mass scaling in pre-K-Pg birds; 286 unfilled symbols denote species that match brain x body mass scaling in non-avian sauropsids, as in B. B, 287 Data points for fossil species in Table 1 plotted onto the fitted power functions shown in Figure 1b that 288 describe the brain x body mass relationship in modern ectothermic sauropsids and pre-K-Pg endothermic 289 sauropsids show that most non-avian theropods (pink) and early avians in the dataset have the brain mass 290 expected for a generic endothermic, pre-K-Pg modern bird that had their body mass, whereas most 291 sauropodmorphs and pterosaurs have brain mass within the range expected for a modern generic 292 ectothermic, non-avian sauropsid of their brain mass. Different ornithischian species conform to one of 293 the other scaling relationship. Power functions, plotted with 95% prediction intervals, are the same as in 294 Figure 1b. C, D, schematics of alternate proposals for the evolution of brain vs body mass relationships 295 that are characteristic of ectothermic (green) or endothermic (shades of red) modern amniotes. 296 Evolutionary trees based on Wiemann et al. (2022) and Kverkova et al. (2022). C, metabolite-based 297 analysis (Wiemann et al., 2022) predicts that a brain x body scaling relationship similar to that 298 characteristic of modern endothermic pre-K-Pg birds applied broadly across dinosaur and pterosaur 299 species, but not in ornithischians. **D**, present brain x body scaling relationships shown in **B** predicts that 300 endothermy was widespread in theropods but only occasional in pterosaurs and ornithischians.



303 Figure 3. Theropod dinosaurs overlap with modern pre-K-Pg endothermic sauropsids (birds) in their 304 brain x body mass scaling, which results in primate-like numbers of telencephalic neurons in theropods, 305 whereas most sauropodmorphs, pterosaurs and some ornithischians overlap with ectothermic non-306 avian sauropsids. A, Data points for fossil species in Table 1 (asterisks) as well as data points for extant 307 pre-K-Pg bird species (red polygons), extant non-avian sauropsids (green stitches) and primates (red 308 circles) plotted onto the fitted power functions shown in Figure 1b that describe the brain x body mass 309 relationship in modern ectothermic sauropsids and pre-K-Pg endothermic sauropsids. Power functions, 310 plotted with 95% prediction intervals, are the same as in Figure 1b. **B**, Same as in **A**, but showing measured 311 numbers of neurons in the mammalian cortex or in the telencephalon of extant sauropsids plotted 312 together with estimated numbers of neurons in the telencephalon of dinosaur and pterosaur species (bold 313 values in Table 1). Notice the similar numbers of neurons in the telencephalon of theropods and in the 314 cerebral cortex of primates despite much larger body masses in theropods.

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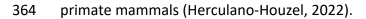
316 Given the striking distinction in brain x body scaling between extant non-avian sauropsids 317 and pre-K-Pg birds shown in Figure 1, most likely associated with the distinction between 318 ectothermy and endothermy (Kverkova et al., 2022; Herculano-Houzel, 2022), and the similar 319 segregation of fossil dinosaur and pterosaur species shown in Figure 2, here I take the approach 320 of hypothesizing that the neuronal scaling rules calculated for the telencephalon of endothermic 321 or ectothermic modern species already applied to the brains of fossil species of matching brain 322 vs body scaling relationship. Thus, considering that most fossil theropods had brains of the mass 323 expected for a modern pre-K-Pg bird of theropod-like body mass (Figure 2b), the predictive 324 neuronal scaling rule calculated for modern pre-K-Pg birds will also estimate the numbers of 325 telencephalic neurons in fossil theropod species of known brain mass.

Using the published values of brain mass estimated from CT analysis (Table 1) plotted in Figure 2, I find that theropods had primate-like numbers of telencephalic neurons (Figure 3B; Table 1), although in bodies as much as 1,000 times larger than primates of similar numbers of cortical neurons. As depicted in Figure 4, numbers of telencephalic neurons ranged from just over 1 billion telencephalic neurons in the 73 g brain of *Alioramus*, comparable to a capuchin monkey, to over 3 billion telencephalic neurons in a 343 g brain of *Tyrannosaurus rex*, which is more telencephalic neurons than found in a baboon (Herculano-Houzel et al., 2015a). In comparison, 333 scaling with ectothermic sauropsid rules, Triceratops, with a 72 g brain similar in size to 334 Alioramus, presumably had only around 172 million telencephalic neurons – fewer than the 306 335 million neurons found in the cerebral cortex of a capybara (Herculano-Houzel et al., 2015a). 336 Importantly, the use of endotherm (avian sauropsid) scaling rules to estimate numbers of 337 telencephalic neurons in theropods versus ectotherm (non-avian sauropsid) scaling rules in 338 ornithischians is supported by recent metabolite findings in these species (Wiemann et al., 2022). 339 The distinction is highly consequential: if the *Tyrannosaurus* brain scaled like a non-avian 340 ectothermic sauropsid brain, it would have an estimated 446 million telencephalic neurons – still 341 as many as in a large dog, but less than 15% of the baboon-like 3.3 billion telencephalic neurons 342 estimated if pre-K-Pg bird-like scaling rules applied (Table 1).

343 In pterosaurs, the brain x body relationship in Figure 2b supports the metabolic finding 344 consistent with endothermy in *Rhamphorhynchus muensteri* (though not in *R. gemmingi*; Table 345 1), but is consistent with ectothermy in *Pteranodon* (Wiemann et al., 2022). Assuming pre-K-Pg 346 endothermic sauropsid scaling rules, the 8 kg pterosaur Anhanguera had an estimated 189 347 million telencephalic neurons, fewer than in a marmoset, in a brain of 8 g, which is almost 4 times 348 as many telencephalic neurons as estimated in the small brain of Archaeopteryx (Figure 3). In 349 contrast, assuming ectothermic sauropsid scaling rules, a 450 g Pterodactylus animal had only an 350 estimated 7 million telencephalic neurons, fewer than found in a mouse, in its 0.4 g brain (Table 351 1).

352 Comparing the present dataset with the species analyzed in that study of fossil 353 metabolites (Wiemann et al., 2022), the only disagreement regards Diplodocus, the only 354 sauropodmorph tested in that study, suggested to have elevated oxidative metabolic rates 355 consistent with endothermy but found here to conform to the brain x body scaling relationship 356 of modern ectothermic sauropsids. Employing the latter, according to Figure 2b, 357 sauropodmorphs such as Brachiosaurus and Diplodocus had only marmoset-like numbers of 358 telencephalic neurons in their brains, even though these had mass that was similar in range to 359 the brains of theropods that had estimated monkey-like numbers of telencephalic neurons 360 (Figure 4, bottom row). Thus, until the metabolite concentrations of sauropodmorph species can 361 be systematically analyzed in more species, the present data suggest that these large quadrupeds

- 362 had telencephalons that were composed according to the neuronal scaling rules that still apply
- 363 to modern ectothermic sauropsids which, incidentally, are the same that apply to modern non-



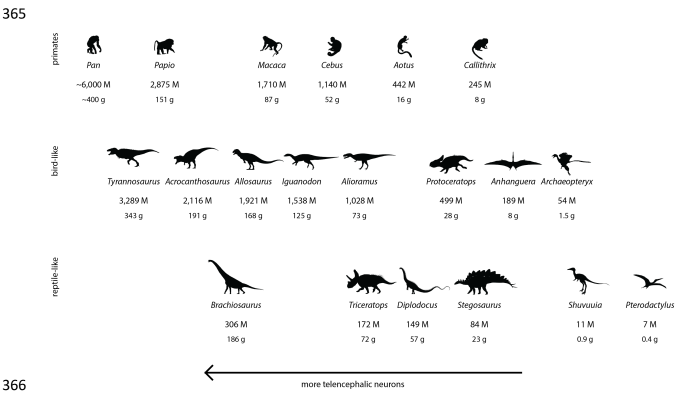


Figure 4. Estimates of numbers of telencephalic neurons in dinosaur and pterosaur species. Values below each image are numbers of telencephalic neurons (in millions, M) and brain mass (in grams). Select species from the dataset in Table 1 are depicted, ranked from left to right in decreasing order of numbers of telencephalic neurons and separated according to whether the species conforms to the brain vs body mass relationship of modern (endothermic) pre-K-Pg birds (center row) or of modern non-avian (ectothermic) sauropsid species (bottom row). For comparison, select primate species of similar numbers of pallial neurons are shown in the top row. All images from phylopic.org.

374

375 Discussion

Here I show that the scaling relationship between brain and body mass that applies to the moderately sized modern bird species originated in pre-K-Pg times still captures the scaling relationship that applied to relatively small as well as very large Mesozoic theropods, in an extreme example of how biological scaling can be very well described across several orders of

380 magnitude by power functions, which indicate the presence of scale-free, modular, self-381 organizing principles at work (Herculano-Houzel et al., 2016; West, 2017). Importantly, the 382 overlap between extinct theropods and extant pre-K-Pg birds strongly suggests that theropods 383 were endothermic. Similarly, I find that the brain x body mass scaling relationship of most species 384 of sauropodmorph and ornithischian dinosaurs, as well as pterosaurs, is captured by the scaling 385 relationship that applies to modern ectothermic sauropsids, which suggests that those species of 386 dinosaurs were ectothermic, or if they were endothermic, they had not yet expressed the 387 enlarged brain mass that is presumably affordable by the increased oxidative capacity that 388 powers endothermy (Herculano-Houzel, 2022). However, it remains possible that the values 389 presented here for sauropodmorphs and ornithischians are underestimated, since they stem 390 from a dataset that considered that the brain of these species only filled 50% of the cranial cavity 391 (Hurlburt, 1996); likewise, it is possible that the values for theropods are slightly overestimated, 392 since they assume that in these species, the brain fills the cranial cavity (Balanoff et al., 2013; 393 Witmer et al., 2003).

On a related note, I acknowledge that the nomenclature and quite possibly some of the data provided in Table 1 from the original data of Hurlburt (1996) are outdated, especially those pertaining to sauropodmorphs and ornithischians. I provide those data as initial estimates for non-Theropod species, and maintain the outdated names as in the original dataset, so that those data can be easily located by future researchers, whom I hope will carry this study forward with modern data.

400 Another methodological issue that will be disputed by some readers is that the scaling 401 relationships reported here are not "corrected" for phylogenetic relatedness, as detailed in the 402 Methods. The central issue of the present study, as in all my previous work, is the scaling 403 relationship between physical quantities such as brain structure mass and numbers of cells and 404 how one predicts the other in different ways in different clades. Whereas phylogenetic 405 comparative methods were initially introduced to alleviate issues with bias possibly introduced 406 by evolutionary relatedness in a sample. I have previously shown that any differences in the 407 scaling exponents calculated with and without accounting for phylogenetic relatedness differ by 408 only 1-2%, which is not a statistically significant difference, provided that there is no mixing in

409 the sample of species from clades already found to scale differently (Gabi et al., 2010; Herculano-410 Houzel et al., 2011). Moreover, phylogenetic relationships are constantly being updated as 411 research progresses, which would affect the predictive scaling rules calculated if relatedness 412 within each clade were taken into consideration. This is another reason why I continue to prefer 413 to report only the scaling relationships that apply to the raw data (Herculano-Houzel et al., 414 2015a). These mathematical functions will be readily useful to researchers interested in the 415 relationships between the physical entities involved; other researchers interested in the 416 phylogenetic signal possibly contained within the raw data provided are welcome to apply in their 417 own studies to further phylogenetic studies of evolution, which are not at all the focus of the 418 present work.

419 A still growing number of studies have focused on establishing whether dinosaurs were 420 ectotherms, mesotherms, or had the high metabolic rates characteristic of modern warm-421 blooded animals, and the fast-paced behavior and life history that come with it (Werner and 422 Griebeler, 2014; Grady et al., 2014; Wiemann et al., 2022). The present findings on the diverse 423 scaling of brain x body mass across dinosaur clades, which are compatible with endothermy in 424 some and ectothermy in other species, add to the still-going debate about the metabolic 425 condition of fossil dinosaurs by disputing the claim of homogeneous mesothermy across species 426 (Grady et al., 2014) in favor of much larger diversity than previously suspected, supporting the 427 finding that higher metabolic rates appeared in some but not all dinosaur clades (Wiemann et al., 2022). Specifically, while most theropods and the single sauropodmorph (Diplodocus) tested 428 429 had advanced lipoxidation end-products accumulated in quantities indicative of high metabolic 430 rates, different ornithischian and pterosaur species showed concentrations compatible with 431 either high or low metabolic rates (Wiemann et al., 2022). So much diversity amongst dinosaurs 432 and pterosaurs in both metabolism and brain x body scaling (Figure 2b) warrants discontinuation 433 of the practice of treating these species as a mixed bag in scaling studies. Instead of using all-434 encompassing scaling rules such as the power function shown in Figure 2c, clade-specific analyses 435 and scaling rules should be employed, informed by other features such as analysis of metabolites 436 (Wiemann et al., 2022), which suggests the clustering indicated by the colors in Figure 2c, or by the scaling relationship between brain and body mass, which suggests the clustering indicatedby the colors in Figure 2d.

439 As modeling techniques based on micro-CT data improve and allow the volume of other 440 brain structures to be estimated (Knoll et al., 2021), more evidence should help distinguish which 441 dinosaur and pterosaur species were ecto- or endothermic. The cerebellum, for example, is 442 decidedly larger in extant endothermic species compared to ectothermic species of similar body 443 mass (Kverkova et al., 2022); thus, the size of the cerebellum relative to the mass of the 444 telencephalon (Herculano-Houzel, 2022) and of the body may serve as a new diagnostic criterion 445 to infer the metabolic status of species of the prehistoric fauna. Importantly, this is not a 446 difference of 10-20%, but of 10-20 times in volume of the cerebellum relative to the brainstem 447 between living endothermic and ectothermic amniotes; therefore, models and simulations that 448 estimate the volume of the cerebellum in fossil animals of unknown metabolic status would 449 greatly contribute not only to understanding cerebellar evolution, but also to determining their 450 metabolic status. Still, absent volumetric analyses of the cerebellum, simply determining whether 451 the brain vs body mass relationship clusters with endothermic, pre-K-Pg birds or with 452 ectothermic sauropsids, as more fossil species have their brain and body masses estimated, 453 should already provide diagnostic evidence of the metabolic status of those species.

454 Estimating the numbers of neurons in the telencephalon, whose main component is the 455 pallium, a major contributor to behavioral flexibility, is obviously consequential for inferring the 456 cognitive capabilities of dinosaur species, whatever their body size (Herculano-Houzel, 2017). The 457 present estimates showing that apex predators such as Tyrannosaurus had the numbers of 458 telencephalic neurons found in modern medium-sized primates of impressive cognitive abilities 459 adds a new dimension to how dinosaurs are pictured; an elephant-sized but agile carnivoran 460 biped endowed with macaque- or baboon-like cognition must have been an extremely 461 competent predator indeed. But additionally, I showed recently that the number of neurons in 462 the pallium is a true and reliable predictor of age at sexual maturity and maximal longevity in 463 warm-blooded animals (Herculano-Houzel, 2019a), such that 74% of variation in these life-history 464 variables can be predicted in mammals and birds alike simply by the absolute number of neurons 465 in the cerebral cortex, whereas body mass is an irrelevant predictor once numbers of cortical

neurons are accounted for (Herculano-Houzel, 2019a). Using the reported equations $L = e^{-4.939}$ 466 $N_{cx}^{0.402}$ and S = $e^{-2.858} N_{cx}^{0.471}$ that relate maximal longevity (L) and age at female sexual maturity 467 (S), respectively, to numbers of cortical neurons (N_{cx}; Herculano-Houzel, 2019), and assuming 468 469 that most telencephalic neurons in sauropsids are pallial (Kverkova et al., 2022), I can predict that 470 a warm-blooded Tyrannosaurus of 2.2-3.3 billion telencephalic neurons would take 4-5 years to 471 reach sexual maturity, and have an estimated maximal longevity of 42-49 years, similar to 472 baboons, whereas Archaeopteryx should reach sexual maturity in ca. 8 months, and have a 473 maximal lifespan of 10 years, similar to flycatchers and other songbirds (Herculano-Houzel, 474 2019). In support of this estimate based on extant warm-blooded species, the survivorship 475 pattern of tyrannosaurs is similar to that seen in long-lived, mammals and birds (Erickson et al., 476 2006). The predicted sexual maturity of Tyrannosaurus at age 5 years, like in modern warm-477 blooded amniotes of similar numbers of cortical neurons, anticipates by a full decade the 478 previous demonstration that, at 18 years of age, this species was sexually mature (although that 479 was admittedly an upper bound; Lee and Werning, 2008). While the largest and oldest known T. 480 rex lived an estimated 28 years, well under the predicted maximal longevity, the finding that only 481 2% of the population lived long enough to attain maximal size and age for the species (Erickson 482 et al., 2006) makes the estimate of a maximal lifespan of just over 40 years compatible with the 483 oldest known fossil.

484 Through their association with delayed sexual maturity and longer lifespans, larger 485 numbers of telencephalic neurons simultaneously endow brains with the cognitive flexibility that 486 can be construed as intelligence (Herculano-Houzel, 2017) and come with increased lifetime 487 opportunities to develop that increased biological signal processing capability into abilities such 488 as using and creating tools, and devising and perpetuating problem-solving processes 489 (Herculano-Houzel, 2019b). With enough pallial neurons and a long enough lifetime that comes 490 with it, generations overlap enough that developed abilities can be transmitted and perpetuated, 491 forming a body of technology and culture that characterizes populations (Herculano-Houzel, 492 2019b). The present findings invite the speculation that theropod dinosaurs such as T. rex, with 493 even more telencephalic neurons than modern tool-using and tool-making corvids (Olkowicz et 494 al., 2016), had the biological capability to use and craft tools, and develop a culture, like modern

495	birds and primates (Beck, 1974; Whiten et al., 1999; Sapolsky and Share, 2004; von Bayern et al,
496	2018). Being able to infer what existed inside the brains of dinosaurs thus multiplies in several
497	directions our knowledge of what life was like in the pre-asteroid, Mesozoic world, and places at
498	least theropods, if not other dinosaurs as well, in the cognitive realm of tool-using and culture-
499	building modern birds and primates.
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502 503	References
504	
505	Alvarez LW, Alvarez W, Asaro F, Michel HV (1980) Extraterrestrial cause for the Cretaceous-
506	Tertiary extinction. Science 208, 1095-1108.
507	Balanoff AM, Bever GS, Rowe TS, Norell MA (2013) Evolutionary origins of the avian brain.
508	Nature 501, 93-97.
509	Beck BB (1974) Baboons, chimapnzees, and tools. J Human Evol 3, 509-516.
510	Bennett AF, Ruben JA (1979) Endothermy and activity in vertebrates. Science 206, 649-655.
511	Bertrand OC, Shelley SL, Williamson TE, Wible JR, Chester SGB, Flynn JJ, Holbrook LT, Lyson
512	TR, Meng J, Miller IM, Püschel HP, Smith T, Spaulding M, Tseng ZJ, Brusatte SL (2022) Brawn
513	before brains in placental mammals after the end-Cretaceous extinction. Science 376, 80-85.
514	Bininda-Emonds ORP, Cardillo M, Jonew KE, MacPhee RDE, Beck RMD, Grenyer R et al. (2007)
515	The delayed rise of present-day mammals. Nature 446, 507-512.
516	Boot N, Baas M, van Gaal S, Cools R, de Dreu CKW (2017) Creative cognition and dopaminergic
517	modulation of fronto-striatal networks: Integrative review and research agenda. Neurosci
518	Biobehav Rev 78, 13-23.
519	Brusatte SL, O'Connor JK, Jarvis ED (2015) The origin and diversification of birds. Curr Biol 25,
520	R888-898.

- 521 Erickson GM, Currie PJ, Inouye BD, Winn AA (2006) Tyrannosaur life tables: an example of 522 nonavian dinosaur population biology. Science 313, 213-218.
- 523 Gabi M, Collins CE, Wong P, Kaas JH, Herculano-Houzel S (2010) Cellular scaling rules for the 524 brain of an extended number of primate brains. Brain Behav Evol 76, 32-44.
- 525 Gillooly JF, McCoy MW (2014) Brain size varies with temperature in vertebrates. PeerJ 2, 526 e301.
- 527 Grady JM, Enquist BJ, Dettweiler-Robinson E, Wright NA, Smith FA (2014) Evidence for 528 mesothermy in dinosaurs. Science 344, 1268-1273.

529 Herculano-Houzel S (2016) What modern mammals teach about the cellular composition of 530 early brains and mechanisms of brain evolution. In Kaas JH, Krubitzer L eds., Evolution of 531 Nervous Systems, 2nd ed., vol. 2, 153-190.

Herculano-Houzel S (2017) Numbers of neurons as biological correlates of cognitive
capability. Curr Opin Behav Sci 16, 1-7.

534 Herculano-Houzel S (2019a) Longevity and sexual maturity vary across species with number 535 of cortical neurons, and humans are no exception. J Comp Neurol 527, 1689-1705.

- Herculano-Houzel S (2019b) Life history changes accompany increased numbers of cortical
 neurons: a new framework for understanding brain evolution. Prog Brain Res 250, 179-216.
- 538 Herculano-Houzel S (2022) Mammals, birds and non-avian reptiles have signature 539 proportions of numbers of neurons across their brain structures: Numbers of neurons 540 increased differently with endothermy in birds and mammals. bioRxiv.org, 2022/496835.
- 541 Herculano-Houzel S, Lent R (2005) Isotropic fractionator: a simple, rapid method for the 542 quantification of total cell and neuron numbers in the brain. J Neurosci 25, 2518-2521.
- 543 Herculano-Houzel S, Kaas JH (2011) Gorilla and orangutan brains conform to the primate 544 scaling rules: implications for hominin evolution. Brain Behav Evol 77, 33-44.

Herculano-Houzel S, Ribeiro P, Campos L, da Silva AV, Torres LB, Catania KC, Kaas JH (2011)
Updated neuronal scaling rules for the brains of glires (rodents/lagomorphs). Brain Behav
Evol 78, 302-314.

Herculano-Houzel S, Ribeiro PFM, Campos L, da Silva AV, Torres LB, Catania KC, Kaas JH (2011)
Updated neuronal scaling rules for the brains of Glires (rodents/lagomorphs). Brain Behav
Evol 78, 302-314.

551 Herculano-Houzel S, Manger PR, Kaas JH (2014) Brain scaling in mammalian brain evolution 552 as a consequence of concerted and mosaic changes in numbers of neurons and average 553 neuronal cell size. Front Neuroanat 8, 77.

Herculano-Houzel S, Catania K, Manger PR, Kaas JH (2015a) Mammalian brains are made of
these: a dataset on the numbers and densities of neuronal and non-neuronal cells in the brain
of glires, primates, scandentia, eulipotyphlans, afrotherians and artiodactyls, and their
relationship with body mass. Brain Behav Evol 86, 145-163.

558 Herculano-Houzel S, Kaas JH, Miller D, Von Bartheld CS (2015b) How to count cells: the 559 advantages and disadvantages of the isotropic fractionator compared with stereology. Cell 560 Tissue Res 360, 29-42.

Hopson JA (1977) Relative brain size and behavior in archosaurian reptiles. Annu Rev Ecol
Systematics 8, 429-448.

Hulburt GR (1996) Relative brain size in recent and fossil amniotes: Determination and
interpretation. PhD Thesis, University of Toronto.

565 Hulburt GR, Ridgely RC, Witmer LM (2013) Relative size of brain and cerebrum in 566 Tyrannosaurid dinosaurs: an analysis using brain-endocast quantitative relationships in 567 extant alligators. In Parrish JM, Molnar RE, Currie PJ, Koppelhus EB (eds), Tyrannosaurid 568 paleobiology. Indiana University Press, Bloomington, 134-154.

569 Jerison HJ (1973) Evolution of the brain and intelligence. New York, Academic Press.

570 Knoll F, Buffetaut E, Bülow M (1999) A theropod braincase from the Jurassic of the Vaches 571 Noires cliffs (Normandy, France): osteology and palaeoneurology. Bull Soc Geol France 170, 572 103-109.

573 Knoll F, Schwarz-Wings D (2009) Palaeoneuroanatomy of Brachiosaurus. Annales Paléontol 574 95, 165-175.

Knoll, F., S. Lautenschlager, S. Kawabe, G. Martínez, E. Espílez, L. Mampel, & L. Alcalá. 2021.
Palaeoneurology of the Early Cretaceous iguanodont *Proa valdearinnoensis* and its bearing on the
parallel developments of cognitive abilities in theropod and ornithopod dinosaurs. Journal of
Comparative Neurology, 529 (18): 3922–3945.

Knoll, F., S. Kawabe & A. Watanabe (2022) A proxy for brain-to-endocranial cavity index in nonneornithean dinosaurs and other extinct archosaurs. 10th European Conference on Comparative
Neurobiology, Abstract Book. 33. Eds: P. Němec, K. Kverková, Y. Zhang, F. Dionigi, R. Druga & G.
Pavlinkova. Praha: Univerzita Karlova.

- 583 Ksepka DT, Balanoff AM, Smith NA, Bever GS, Bhullar B-AS, Bourdon E et al. (2020) Tempo 584 and pattern of avian brain size evolution. Curr Biol 30, 2026-2036.
- 585 Kverkova K, Marhounová L, Polonyiová A, Kocourek M, Zhang Y, Olkowicz S, Stratková B,
 586 Pavelková Z, Vodicka R, Frynta D, Nemec P (2022) The evolution of brain neuron numbers in
 587 amniotes. Proc Natl Acad Sci USA 119, e2121624119.
- Lee AH, Werning S (2008) Sexual maturity in growing dinosaurs does not fit reptilian growth
 models. Proc Natl Acad Sci USA 105, 582-587.
- Olkowicz S, Kocourek M, Lucan RK, Portes M, Fitch WT, Herculano-Houzel S, Nemec P (2016)
 Birds have primate-like numbers of neurons in the telencephalon. Proc Natl Acad Sci USA 113,
 7255-7260.
- Rowe TB, Macrini TR, Luo Z-X (2011) Fossil evidence on origin of the mammalian brain.
 Science 332, 955-957.

595 Sapolsky RM, Share LJ (2004) A pacific culture among wild baboons: its emergence and 596 transmission. PLoS Biol 2, e106.

597 Sol D, Olkowicz S, Sayol F, Kocourek M, Zhang Y, Marhounová L, Osadnik C, Corssmit E, Garcia-

Porta J, Martin TE, Lefebvre L, Nemec P (2022) Neuron numbers link innovativeness with both
absolute and relative brain size in birds. Nature Ecol Evol, doi.org/10.1038/s41559-02201815-x.

Ströckens F, Neves K, Kirchem S, Schwab C, Herculano-Houzel S, Güntürkün O (2022) High
associative neuron numbers could drive cognitive performance in corvid species. J Comp
Neurol 530, 1588-1605.

Von Bayern AMP, Danel S, Auersperg AMI, Mioduszewska B, Kacelnik A (2018) Compound
tool construction by New Caledonian crows. Scientific Reports 8, 15676.

Werner J, Griebeler EM (2014) Allometries of maximum growth rate versus body mass at
 maximum growth indicate that non-avian dinosaurs had growth rates typical of fast growing
 ectothermic sauropsids. PLoS One 9, e88834.

609 West G (2017) Scale: The universal laws of growth, innovation, sustainability, and the pace of 610 life in organisms, cities, economies, and companies. Penguin Press, New York.

611 Whiten A, Goodall J, McGrew WC, Nishida T, Reynolds V, Sugiyama Y, Tutin CEG, Wrangham

612 RW, Boesch C (1999) Cultures in chimpanzees. Nature 399, 682-685.

613 Wiemann J, Menéndez I, Crawford JM, Fabbri M, Gauthier JA, Hull PM, Norell MA, Briggs DEG

614 (2022) Fossil biomolecules reveal an avian metabolism in the ancestral dinosaur. Nature,

615 htpps://doi.org/10.1038/s41586-022-04770-6.

616 Williams RW, Herrup K (1980) The control of neuron number. Annu Rev Neurosci 11, 423-617 253.

618 Witmer LM, Chatterjee S, Franzosa J, Rowe T (2003) Neuroanatomy of flying reptiles and 619 implications for flight, posture and behaviour. Nature 425, 950-953.

620	Yu Y, Zhang C, Xu X (2021) Deep time diversity and the early radiations of birds. Proc Natl
621	Acad Sci USA 118, e2019865118.

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- 630
- 631 All correspondence and material requests should be addressed to the author at the address
- 632 **above.**
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634 Table 1. Dataset and numbers of telencephalic neurons (N_{tel}) in dinosaur and pterosaur species 635 predicted from brain mass (M_{br}) estimates reported in the literature. For the sake of clarity and reproducibility, all data compiled are listed, instead of calculating averages for each species, and 636 are plotted in Figure 2a,b. For each species, estimated N_{tel} (in millions, M) are shown calculated 637 according to the scaling relationships $N_{tel} = e^{17.518} M_{br}^{0.753}$ (for endothermic, basal birds) and N_{tel} 638 = e^{16.342} M_{br}^{0.612} (for ectothermic reptiles; Figure 1c). M_{bd}, body mass in grams; M_{br}, brain mass in 639 grams, converted from estimated brain volume in cm^3 using 1 cm^3 = 1 g. Source of M_{bd} and M_{br} 640 data is indicated next to each species (a, Witmer et al., 2003; b, Hurlburt, 1996; c, Hurlburt et al., 641 642 2013; *d*, Balanoff et al., 2013). Values of N_{tel} in bold are the predictions according to Figure 2b. 643

Species	M _{bd} , g	M _{br} , g	N _{tel} if ectothermic	N _{tel} if endothermic	S (days), endothermic	L _{max} (years), endothermic
Pterosaurs						
Anhanguera piscator (AMNH 25555)ª	7,600	7.720	43.7 M	188.9 M	454	16
Pterodactylus kochi ^b	450	0.420	7.4 M	21.1 M	162	7
Pterodactylus elegans ^b	60	0.140	3.8 M	9.2 M	109	5
Pteranodon ^b	20,000	4.800	32.7 M	132 M	383	14
Rhamphorynchus gemmingi ^b	310	0.700	10.0 M	31.0 M	194	8

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Rhamphorynchus	136	0.830	11.2 M	35.2 M	206	8
muensteri (CM						
11434) ^a						
Scaphognathus	1,500	1.560	16.4 M	56.7 M	257	10
purdoni ^b						
Ornithischians						
Anatosaurus ^b	3,400,000	150.0	268.5 M	1,764 M	1,300	38
Camptosaurus ^b	400,000	23.0	85.2 M	429.9 M	668	22
Euoplocephalus ^b	1,900,000	41.0	121.4 M	664.4 M	821	26
Iguanodon ^b	2,100,000	125.0	240.2 M	1,538 M	1,219	36
<i>Kentrosaurus^b</i>	780,000	24.0	87.5 M	443.9 M	679	22
<i>Protoceratops</i> ^b	200,000	28.0	96.1 M	498.5 M	717	23
Stegosaurus ^b	3,100,000	22.5	84.1 M	422.8 M	663	22
Stegosaurus ^b	2,000,000	22.5	84.1 M	422.8 M	663	22
Triceratops ^b	9,000,000	72.2	171.7 M	1,017 M	1,003	31
Triceratops ^b	6,000,000	72.2	171.7 M	1,017 M	1,003	31
Sauropodmorphs						
Brachiosaurus ^b	78,300,000	186.0	306.3 M	2,075 M	1,403	41
Brachiosaurus ^b	40,000,000	186.0	306.3 M	2,075 M	1,403	41
Brachiosaurus ^b	29,000,000	186.0	306.3 M	2,075 M	1,403	41
Diplodocus ^b	19,000,000	57.0	148.5 M	851.4 M	922	29
Diplodocus ^b	12,000,000	57.0	148.5 M	851.4 M	922	29
Theropods	, ,					
Acrocanthosaurus	3,770,000	191.0	311.3 M	2,116 M	1,416	41
atokensis (OMNH	, ,			,	,	
10146) ^d						
Alioramus altai	359,000	73.2	173.2 M	1,028 M	1,008	31
(IGM 100/1184) ^d					,	
Allosaurus ^b	2,300,000	168.0	287.8 M	1,921 M	1,353	40
Allosaurus ^b	1,400,000	168.0	287.8 M	1,921 M	1,353	40
Bambiraptor ^c	6,582	14.0	62.9 M	295.8 M	561	19
Citipati osmolskae	85,960	22.620	84.4 M	424.5 M	665	22
(IGM 100/978) ^d						
Conchoraptor	5,020	9.440	49.4 M	219.8 M	487	17
gracilis (IGM						
100/3006) ^d						
Dromicioemimus ^b	175,000	87.85	193.6 M	1,179 M	1,075	33
Dromicioemimus ^b	125,000	87.85	193.6 M	1,179 M	1,075	33
Khaan mckennai	12,610	8.830	47.4 M	209.1 M	476	17
(IGM 100/973) ^d	ŕ					
Shuvuuia deserti	250	0.830	11.2 M	35.2 M	206	8
(IGM 100/977) ^d						
Troodon ^b	45,000	41.0	121.4 M	664.4 M	821	26

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Troodontid (IGM 100/1126) ^d	920	3.110	25.1 M	95.3 M	329	12
Tsaagan mangas (IGM 100/1015) ^d	15,950	3.070	24.9 M	94.4 M	327	12
Tyrannosaurus ^b	7,400,000	202.0	322.2 M	2,207 M	1,445	42
Tyrannosaurus ^b	5,000,000	202.0	322.2 M	2,207 M	1,445	42
Tyrannosaurus rex (AMNH 5029) ^d	5,840,000	343	445.5 M	3,289 M	1,743	49
Zanabazar junior (IGM 100/1) ^d	49,300	25.14	90.0 M	459.7 M	690	23
Aves						
Archaeopteryx ^b	400	1.470	15.8 M	54.2 M	252	10
Archaeopteryx ^b	300	1.470	15.8 M	54.2 M	252	10
Archaeopteryx ^b	400	1.760	17.7 M	62.1 M	269	10
Archaeopteryx ^b	300	1.760	17.7 M	62.1 M	269	10
Archaeopteryx lithographica (BMNH 37001) ^d	500	1.440	15.6 M	53.4 M	250	10
Unnamed protoavis ^b	600	3.320	26.1 M	100.1 M	336	13