New Insights into Non-Avian Dinosaur Reproduction and Their Evolutionary and Ecological Implications: Linking Fossil Evidence to Allometries of Extant Close Relatives

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Abstract

It has been hypothesized that a high reproductive output contributes to the unique gigantism in large dinosaur taxa. In order to infer more information on dinosaur reproduction, we established allometries between body mass and different reproductive traits (egg mass, clutch mass, annual clutch mass) for extant phylogenetic brackets (birds, crocodiles and tortoises) of extinct non-avian dinosaurs. Allometries were applied to nine non-avian dinosaur taxa (theropods, hadrosaurs, and sauropodomorphs) for which fossil estimates on relevant traits are currently available. We found that the reproductive traits of most dinosaurs conformed to similar-sized or scaled-up extant reptiles or birds. The reproductive traits of theropods, which are considered more bird-like, were indeed consistent with birds, while the traits of sauropodomorphs conformed better to reptiles. Reproductive traits of hadrosaurs corresponded to both reptiles and birds. Excluding Massospondylus carinatus, all dinosaurs studied had an intermediary egg to body mass relationship to reptiles and birds. In contrast, dinosaur clutch masses fitted with either the masses predicted from allometries of birds (theropods) or to the masses of reptiles (all other taxa). Theropods studied had probably one clutch per year. For sauropodomorphs and hadrosaurs, more than one clutch per year was predicted. Contrary to current hypotheses, large dinosaurs did not have exceptionally high annual egg numbers (AEN). Independent of the extant model, the estimated dinosaur AEN did not exceed 850 eggs (75,000 kg sauropod) for any of the taxa studied. This estimated maximum is probably an overestimation due to unrealistic assumptions. According to most AEN estimations, the dinosaurs studied laid less than 200 eggs per year. Only some AEN estimates obtained for medium to large sized sauropods were higher (200-400 eggs). Our results provide new (testable) hypotheses, especially for reproductive traits that are insufficiently documented or lacking from the fossil record. This contributes to the understanding of their evolution.

Introduction

The discovery of the gigantic sauropods and other large dinosaurs has stimulated scientists to understand the biology of dinosaurs. Several researchers [1–3] recently argued that the reproductive strategy of producing many small offspring contributed to the exceptional gigantism seen in the sauropods, a hypothesis introduced by Janis and Carrano [4] and recently corroborated by Werner and Griebeler [5]. In contrast to any living species, our information on dinosaurs and their reproduction is limited to fossils. Unfortunately, fossils do not allow for the complete reconstruction of an organisms’ traits (e.g. of the life history). Traits are often inaccurately preserved or simply absent from the fossil record (e.g. clutches can be incomplete and breeding frequency is simply not documented in the fossil record). This hampers our understanding of the reproductive strategies employed by dinosaurs.

Equations linking body mass to other traits derived from extant taxa are commonly used to estimate these traits for extinct species, including those dealing with reproduction (e.g. [6–8]). While the correlations between clutch/litter size or annual offspring number and body mass differ between different extant amniotic taxa [4,5,9,10], mass specific reproductive traits such as egg mass, clutch mass and annual reproductive mass (clutch mass × number of clutches per year) do significantly, positively correlate with body mass [9,11–17]. The relationships between traits (Y) and body mass (BM) follow a power function $Y = c \times BM^b$, where $c$ is a normalization constant and $b$ is an exponent. These so-called allometric
(allometric because $b$ usually differs from one) functions are usually log-log plotted, whereby the normalization constant $c$ is the intercept and the exponent $b$ is the slope of a straight line. The exponent $b$ differs between different amniotic groups when egg mass is plotted against body mass \cite{9,10}. Conversely, $b$ is often very similar between different amniotic groups when clutch mass or annual reproductive mass is plotted against body mass \cite{9,10,16}. While the slopes $b$ are often statistically indistinguishable for annual clutch/litter mass, the normalization constants vary significantly among taxa \cite{16,18}. These normalization constants are frequently similar among species/taxa with similar lifestyles \cite{18}. We assume from these observations that analogous allometries on body mass and reproductive traits exist within close extant phylogenetic relatives of non-avian dinosaurs. We expect that these allometries are applicable to extinct non-avian dinosaurs as well.

To derive information on dinosaur reproduction, the procedure presented by Bryant and Russell \cite{19} and Witmer \cite{20} was employed, specifically a combination of phylogenetic inference and extrapolatory analysis. As an extant phylogenetic bracket (EPB, \cite{20}) for non-avian dinosaurs, we chose phylogenetically close relatives with reproductive characteristics similar to those assumed for non-avian dinosaurs. Reproductive traits included, among others, ground breeding, and calcified, rigid-shelled eggs (for the distribution of egg types over the phylogeny of early amniotes, see Sander \cite{21}). For the extant bird and reptile species which meet these specifications, we established allometries between body mass and different reproductive traits (egg mass, clutch mass, annual clutch mass). Allometries were then applied to nine non-avian dinosaur taxa for which information on body mass, egg mass and clutch mass is currently available from the fossil record. We expected, (i) reproductive trait estimates of non-avian dinosaurs, which are considered to be more “bird-like” in their reproductive mode (e.g. theropods \cite{22–24}), to conform to those seen in extant birds. Similarly, those dinosaurs expressing traits probably more “reptile-like” in their reproductive mode (e.g. sauropods \cite{24}) were expected to fit to those of extant reptiles.

Our established allometries between body mass and clutch mass might also provide further support for the hypothesis of Seymour \cite{25}, who argued that the buried clutch mass of large sauropods is limited due to physiological constrains imposed on the clutch. Assuming a sea turtle model, Seymour \cite{26} showed that unfavorable respiratory gas pressures can occur inside large buried clutches. If clutches are too large, buried eggs do not receive enough oxygen through the soil \cite{25}. Because of this physiological limitation of clutch mass, Seymour \cite{25} suggested that fully buried sauropod eggs were distributed over several small clutches, each clutch containing not more than 13 eggs. Sander et al. \cite{27} also hypothesized that the clutch mass of buried sauropod clutches is smaller than expected given their body mass, suggesting that these sauropods produced several clutches per year. According to these hypotheses, we expected (ii) the clutch masses of buried clutches of large dinosaurs, e.g. of the sauropods producing Megaloolithus mammillare eggs \cite{27}, to be smaller than those predicted from any extant species studied. Assuming that the annual clutch mass allometries derived from extant taxa are also valid for dinosaurs, (iii) we estimated annual breeding frequencies of dinosaurs from their fossil clutch mass. Finally, (iv) we estimated the total number of eggs laid per year for each dinosaur taxon, calculated from the annual clutch mass allometries of extant taxa using the respective fossil egg masses.

**Material and Methods**

**Phylogenetic framework and EPB**

For our analyses, we selected three extant taxa (birds N=217, crocodiles N=22 and tortoises N=20; Table S2), each of which is phylogenetically close to non-avian dinosaurs and/or has reproductive characteristics similar to non-avian dinosaurs. We chose the extant phylogenetic bracket of non-avian dinosaurs, i.e. birds and crocodiles, as the closest phylogenetic relatives of ancient dinosaurs \cite{28–30}. Given that dinosaurs were most probably terrestrial and ground breeding \cite{27,31,32}, we also aimed to restrict our dataset to ground breeding and terrestrial bird species. We initially focused on avian orders presumed to have less derived reproductive characteristics (e.g. ground breeding and precocial). According to traditional taxonomy, these include Paleognathae with the orders Struthioniformes and Tinamiformes. Since the sample sizes obtained for Paleognathae were too small, we also included the orders Galliformes and Anseriformes in our analysis; both are phylogenetically closely related to Paleognathae and are ground breeding and precocial. For Struthioniformes, data from only seven species (two kiwi species and five other ratites) were available, with the two kiwi species (~1.3 kg, cave breeding) strongly differing from the other ratites (~20–90 kg, open breeding) in terms of body masses and reproductive strategies \cite{33,34}. For Tinamiformes, we found no single species for which information on body mass and all studied reproductive traits was available. Our allometries were finally based on 60 galliforme, 150 anseriforme and 7 ratite species. The allometries obtained from these avian species are hereafter referred to as bird model.

For the crocodile model, we chose all extant crocodilian species (N = 22). Since crocodiles are non-terrestrial, we also included tortoises (N = 20) in our allometric analyses. Molecular data suggest that turtles are more closely related to archosaurs than to lepidosaurs \cite{35–37} and, similar to crocodiles and birds, and tortoises have calcified, rigid-shelled eggs. For both the crocodile and tortoises model, all extant species for which information on reproductive traits and body mass was available were included in the model. For each of the three extant taxa, we established allometries linking reproductive traits of species to their body mass.

**Dinosaurs**

We applied established allometries to all dinosaur taxa for which body mass estimates and assignments of fossil eggs or clutches to taxa are currently available (Table S1). These assignments only exist for four theropods (Troodon formosus, Oviraptor philoceratops, Clipitati osmolskai, and
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Lourinhanosaurus antunesi), two hadrosaurs (Maiasaura peeblesorum, lambeosaurine dinosaur), two sauropod oospecies (Megaloolithus patagonicus, Megaloolithus mammillare) and one prosauropod (Massospondylus carinatus). It should be noted that Megaloolithus siruguei is considered a junior synonym of Megaloolithus mammillare.

Life-history traits

For all extant species, we gathered data on adult body mass (BM), egg mass (EM), clutch size (CS) and number of clutches per year (CY). When more than one trait value was found in the literature for the same species, the mean value was calculated (Table S2). For body mass estimates, data on females was preferentially used because mass is more strongly linked to reproductive traits in females than in males. In some cases, however, it was not possible to distinguish between male and female body masses because only the averages of both sexes were available or the sex was not denoted in the source or was unknown (especially for all dinosaurs). To maximize our sample size while minimizing any bias introduced by male body masses, we used female body masses wherever possible and otherwise averaged body masses.

For dinosaurs (for details, see Table S1, supporting information), we used the average species body masses for our estimations when eggs were assigned to a specific dinosaur species (Troodon formosus, Oviraptor philoceratops, Citipati osmolskai, Lourinhanosaurus antunesi, Maiasaura peeblesorum, Massospondylus carinatus). When eggs were assigned instead to a taxonomic group, we used averages of the specific taxonomic group (Megaloolithus patagonicus, Megaloolithus mammillare, lambeosaurine dinosaur). Dinosaur egg masses were calculated from an egg’s volume, assuming an egg density of 1.13 g/cm³ (birds [38]; mean egg density of the six bird orders from Table 3 in this reference; crocodiles [39]; Table 3 in this reference). Egg volume was calculated from fossil egg dimensions (mean) using either the equation V = 0.51L² D² (asymmetrical, bird-like, theropods) or V = 0.524L² D² (elliptical/globular, crocodile-like, sauropodomorphs and hadrosaurs), where V is the egg volume, L the egg length and D the egg diameter [39,40].

Given that all studied extant birds and reptiles lay at least one clutch per year, an initial conservative estimate of the unknown annual breeding frequency was assumed to be one clutch per year for all non-avian dinosaurs. Clutch mass is egg mass multiplied by clutch size. Annual clutch mass is clutch mass multiplied by the number of clutches per year.

Statistical analyses

Establishment of allometries for birds, crocodiles and tortoises. We began by separately analysing the relationship between body mass and reproductive traits for extant birds, crocodiles and tortoises. For each taxa and each of the three reproductive traits, we calculated regression slopes and normalization constants using ordinary least square regressions (OLS) on log-log-transformed data (Table S3).

In these regression analyses, we did not control for phylogenetic effects on reproductive traits. In general, phylogenetic comparative methods perform best when the phylogeny itself and branch lengths are correct [41–43]. However, to the best of our knowledge, no phylogenetic trees resolving to the species level are currently available for birds, crocodiles and tortoises. More notably, a phylogenetic tree containing all studied species with reliable branch lengths is not available. Furthermore, the purpose of taking phylogeny into account is to reduce the variance in the estimated regression or correlation coefficients. However, the estimates of regression coefficients are unbiased and independent of phylogeny; estimates of correlation coefficients are only slightly biased [44]. Because we were mainly interested in the mean coefficient values, and we obtained highly significant coefficients, controlling for phylogeny would not have improved our analyses.

Next, we tested the homogeneity of the regression lines obtained for the three taxonomic groups within each reproductive trait (Table S4, S5). Therein, the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable). When these overall analyses indicated that the calculated regression slopes and the normalization constants of all taxonomic groups of one reproductive trait were compared (analyses of covariance ANCOVA, groups as categorical variable).
The previous analyses could either reveal no statistical differences in regression slopes between at least two taxonomic groups or a significant difference between groups for a reproductive trait. If slopes were statistically homogeneous but intercepts differed between taxonomic groups, we calculated a new OLS regression function with a common regression slope for each group. We used this common regression slope (average value of taxa) as a fixed parameter in these regression models and only estimated the normalization constant for each group. If both slopes and intercepts of taxonomic groups were statistically homogeneous, we determined a common regression function (with the average slope and average intercept of taxa) for these taxa. If a slope of a single taxonomic group statistically differed from all other groups, the initially found OLS regression function was used as allometry for the respective taxon and reproductive trait (Table S6). This statistical procedure led to the development of three allometries/models for egg mass and clutch mass (birds, crocodiles, tortoises, Figures 1 and 2, Table S6) and two allometries for annual clutch mass (birds, reptiles = crocodiles + tortoises, see results, Figure 3, Table S6).

Finally, as a measure of variability in residuals and the deviation of single species from the expected average, we calculated 95% prediction intervals for each regression line (Figures 1-3).

**Application of allometries to non-avian dinosaurs.** Each of the trait pairs (body mass and reproductive trait) of non-avian dinosaurs were compared to the regression lines and to the respective 95% prediction intervals for the bird, the crocodile, and the tortoise model (Figures 1-3). Due to uncertainties in dinosaur body masses and clutch masses, we considered not only the mean fossil values but also their known variability (determined by reported minimum and maximum values taken from literature). For egg masses however, only the mean values were used. We assume that errors in egg mass are negligible in comparison to errors in the estimates of body mass and clutch size and mass.

**Estimation of clutch sizes, annual egg numbers and clutches per year.** Clutch size for non-avian dinosaurs was calculated from fossil egg mass and the clutch mass estimates from the regression lines derived for birds, crocodiles and tortoises. Analogously, the total number of eggs per year (AEN) for dinosaurs was calculated from annual clutch mass (ACM). Since the regression lines for crocodiles and tortoises did not differ statistically (see results), the ACM was estimated from uncertainties in dinosaur body masses and clutch masses, we only estimated the normalization constant for each group. If both slopes and intercepts of taxonomic groups were statistically homogeneous, we determined a common regression function (with the average slope and average intercept of taxa) for these taxa. If a slope of a single taxonomic group statistically differed from all other groups, the initially found OLS regression function was used as allometry for the respective taxon and reproductive trait (Table S6).

The ACM of the prosauropod *M. carinatus* and the sauropod *M. patagonicus*, all dinosaur EMs fell outside the 95% prediction interval of all extant models (Figure 1, B–D). *Massospondylus carinatus* fitted very well to the crocodile and the tortoise model (Figure 1, C and D). The EM of *M. patagonicus* fitted the tortoise model, but only when we assume that its BM is equal to or higher than the mean BM of 22,399 kg, as given in literature (Figure 1D). None of the other dinosaur EMs fitted to the EM estimates of similar-sized or scaled-up extant species. The two models on ACM versus BM were applied to estimate AEN and CY for dinosaurs.

**Application of allometries to dinosaurs**

**Egg mass.** Except for the prosauropod *M. carinatus* and the sauropod *M. patagonicus*, all dinosaur EMs fell outside the 95% prediction interval of all extant models (Figure 1, B–D). *Massospondylus carinatus* fitted very well to the crocodile and the tortoise model (Figure 1, C and D). The EM of *M. patagonicus* fitted the tortoise model, but only when we assume that its BM is equal to or higher than the mean BM of 22,399 kg, as given in literature (Figure 1D). None of the other dinosaur EMs fitted to the EM estimates of similar-sized or scaled-up birds or reptiles. Instead, they were immediately located between these two models (Figure 1). In particular, the theropods (T. formosus, O. philoceratops, C. osmolskae, and L. antunesi) had EMs somewhat closer to those of birds than to reptiles, whereas the EMs of the hadrosaur *M. peeblesorum* and of the sauropod *M. patagonicus* were closer to those of reptiles (Figure 1). EMs of the lambeosaurine dinosaur (hadrosaur) and of the sauropod *M. mammiliare* ranged between the predicted EM in the bird and the two reptile models (Figure 1).

**Clutch mass.** All fossil dinosaur CMs fell within the 95% prediction interval of at least one of the extant models (Figure 2, B–D). All non-theropod CMs matched at least one of the two reptile models (crocodiles, tortoises, Figure 2, C and D), while all theropod CMs matched the bird model (Figure 2B). However, the extant model best suited for the CM of a specific dinosaur differed between dinosaur taxa. Theropod CMs and the CM of the lambeosaurine dinosaur clearly conformed best to the bird model (Figure 2B), but the lambeosaurine dinosaur was in the 95% prediction interval of the crocodile model, too (Figure 2C). Sauropod CMs and the CM of the hadrosaur *M. peeblesorum* matched the tortoise model best, but were still realistic under the crocodile model (Figure 2, C and D). The CM of the prosauropod *M. carinatus* corresponded well to both the crocodile and the tortoise model (Figure 2, C and D). The CM of the sauropod *M. mammiliare* fell within the 95% prediction interval of the reptile models (Figure 2C).
Figure 1. Allometries between body mass (BM) and egg mass (EM) of birds, crocodiles, and tortoises and their comparison to non-avian dinosaurs. (A) General comparison of dinosaur EMs to the EM allometry of birds (grey squares/line), crocodiles (green triangles/line) and tortoises (dark green circles/line). (B) Detailed comparison of dinosaur EMs to the EM allometry of birds. Grey continuous line = regression line of birds. Grey scattered lines = 95% prediction interval of the bird regression. Dinosaurs in the graphs = theropods (yellow squares, from left to right): *Troodon formosus*, *Oviraptor philoceratops*, *Clitiphi osmolskae*, *Lourinhanosaurus antunesi*; hadrosaurs (light blue triangles, from bottom to top): *Maiasaura peeblesorum*, lambeosaurine dinosaur; sauropod oospecies (red circles, from bottom to top): *Megaloolithus patagonicus*, *Megaloolithus mammilare*; prosauropod (orange circle): *Massospondylus carinatus*. Black error bars = possible value ranges for a non-avian dinosaur taxon (left/lower bar = minimum value derived from the fossil record, right/upper bar = maximum value). (C) Analogous to (B), but for crocodiles (green triangles/lines) and dinosaurs. (D) Analogous to (B), but for tortoises (dark green circles/lines) and dinosaurs.

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Figure 2. Allometries between body mass (BM) and clutch mass (CM) of birds, crocodiles, and tortoises and their comparison to non-avian dinosaurs. (A) General comparison of dinosaur CMs to the CM allometry of birds (grey squares/line), crocodiles (green triangles/line) and tortoises (dark green circles/line). (B) Detailed comparison of dinosaur CMs to the CM allometry of birds. Grey continuous line = regression line of birds. Grey scattered lines = 95% prediction interval of the bird regression. Dinosaurs in the graph = theropods (yellow squares, from left to right): Troodon formosus, Oviraptor philoceratops, Citipati osmolskae, Lourinhanosaurus antunesi; hadrosaurs (light blue triangles, from bottom to top): Maiasaura peeblesorum, lambeosaurine dinosaur; sauropod oospecies (red circles, from bottom to top): Megaloolithus patagonicus, Megaloolithus mammiliare; prosauropod (orange circle): Massospondylus carinatus. Black error bars/scattered rectangles = possible value ranges for a non-avian dinosaur taxon (left/lower bar/edge = minimum value derived from the fossil record, right/upper bar/edge = maximum value). (C) Analogous to (B), but for crocodiles (green triangles/lines) and dinosaurs. (D) Analogous to (B), but for tortoises (dark green circles/lines) and dinosaurs.

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Figure 3. Allometries between body mass (BM) and clutch mass (ACM) of birds, crocodiles, and tortoises and their comparison to non-avian dinosaurs. (A) General comparison of dinosaur ACMs to the ACM allometry of birds (grey squares/line), crocodiles (green triangles/line) and tortoises (dark green circles/line). (B) Detailed comparison of dinosaur ACMs to the ACM allometry of birds. Grey continuous line = regression line of birds. Grey scattered lines = 95% prediction interval of the bird regression. Dinosaurs in the graph = theropods (yellow squares, from left to right): Troodon formosus, Oviraptor philoceratops, Citipati osmolskae, Lourinhanosaurus antunesi; hadrosaurs (light blue triangles, from bottom to top): Maiasaura peeblesorum, lambeosaurine dinosaur; sauropod oospecies (red circles, from bottom to top): Megaloolithus patagonicus, Megaloolithus mammillare; prosauropod (orange circle): Massospondylus carinatus. Black error bars/scattered rectangles = possible value ranges for a non-avian dinosaur taxon (left/lower bar/edge = minimum value derived from the fossil record, right/upper bar/edge = maximum value). (C) Analogous to (B), but for reptiles (crocodiles + tortoises, green triangles/lines) and dinosaurs.

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Table 1. Reproductive characteristics of sauropodomorphs as documented in the fossil record and estimated by the reptile (crocodiles, tortoises) and the bird model.

| Taxon                      | Fossil | Reptile model | Bird model |
|----------------------------|--------|---------------|------------|
|                            | BM (kg) | EM (kg) | CS | CS¹ | CS² | AEN | CY | CS² | AEN | CY |
| Massospondylus carinatus   | 107     | 0.128    | 34 | 14.0 | 39.7 | 40.2 | 1.2 | 93.1 | 100.0 | 2.9 |
| Massospondylus carinatus   | 175     | 0.128    | 34 | 19.9 | 56.5 | 57.3 | 1.7 | 132.4 | 142.9 | 4.2 |
| Massospondylus carinatus   | 280     | 0.128    | 34 | 27.8 | 79.1 | 80.6 | 2.4 | 185.3 | 200.8 | 5.9 |
| Megaloolithus mammilare    | 5000    | 5.211    | 19 [9, 28] | 5.4 | 15.3 | 16.0 | 0.8 [0.6, 1.8] | 35.8 | 38.8 | 2.1 [1.4, 4.4] |
| Megaloolithus mammilare    | 22399   | 5.211    | 19 [9, 28] | 15.7 | 44.7 | 47.3 | 2.5 [1.7, 5.3] | 104.7 | 118.0 | 6.2 [4.2, 13.1] |
| Megaloolithus mammilare    | 75000   | 5.211    | 19 [9, 28] | 37.3 | 106.1 | 113.6 | 6.0 [4.1, 12.6] | 248.6 | 283.1 | 14.9 [10.1, 31.5] |
| Megaloolithus patagonicus  | 5000    | 1.741    | 28 [15, 40] | 16.1 | 45.7 | 47.8 | 1.7 [1.2, 3.2] | 107.2 | 119.1 | 4.3 [3.0, 7.9] |
| Megaloolithus patagonicus  | 22399   | 1.741    | 28 [15, 40] | 47.1 | 133.8 | 141.7 | 5.1 [3.5, 9.4] | 313.4 | 353.0 | 12.6 [8.8, 23.5] |
| Megaloolithus patagonicus  | 75000   | 1.741    | 28 [15, 40] | 111.8 | 317.6 | 340.1 | 12.1 [8.5, 22.7] | 744.0 | 847.3 | 30.3 [21.2, 56.5] |

BM: minimum, mean and maximum fossil body mass of a taxon taken from literature in kilograms; CS: mean clutch size observed in fossil record, with minimum and maximum fossil values given in square brackets or CS calculated from an allometric clutch mass (CM) model (CS¹ = tortoise model, CS² = crocodile model, CS² = bird model) using the fossil egg mass (EM, for calculating of fossil egg mass see Material and Method), CS = CM divided by EM. AEN (annual egg number): total number of eggs laid per year, calculated from an allometric ACM (annual clutch mass) model using the fossil egg mass, AEN = ACM divided by EM. CY: number of clutches per year, calculated from an allometric ACM model using the fossil CS and estimated AEN, CY = AEN divided by CS. Minima and maxima are given in brackets. References for fossil data are given in Table S3. Equations for the reptile and the bird model are given in Table S6. Note: For Massospondylus carinatus only one CS estimate was available.

Prediction interval of the crocodile model (Figure 2C). The CM of the sauropod M. patagonicus and of the hadrosaur M. peeblesorum also conform to the crocodile model (Figure 2C) when assuming a lower BM for both taxa (M. patagonicus: 5,000-10,000 kg, M. peeblesorum: minimum reported BM of 1,500 kg), or in the case of M. patagonicus when the maximum CS reported in literature was used for predicting CM (40 eggs).

Annual clutch mass. When assuming one clutch per year, all estimated ACMs of non-avian dinosaurs fell within the 95% prediction interval of at least one of the extant models, with the exception of the sauropod M. patagonicus and the hadrosaur M. peeblesorum (Figure 3). All ACMs of theropods conformed well to the bird model (Figure 3B). In contrast, with the exception of the lambeosaurine dinosaur, none of the non-theropod dinosaur ACMs coincided with the bird model when assuming one clutch per year (Figure 3B). The ACM of the lambeosaurine dinosaur was intermediary when referring to the mass expected under the bird model and the reptile model (Figure 3). For the sauropod M. patagonicus and the hadrosaur M. peeblesorum, the ACMs derived from the assumption of one clutch per year coincided with the reptile model only when assuming body masses lower than the mean body masses derived from literature (Figure 3C). Combining M. patagonicus EMs and sauropod BMs with the maximum CS of 40 eggs reported in literature also leads to an ACM that conforms to the reptile model. ACMs of very large sauropods (75,000 kg and more) conformed neither to the bird nor to the reptile model when assuming the CS and EM reported in literature (for M. patagonicus or M. mammilare) and only one clutch per year (Figure 3).

Estimates of annual egg numbers and clutches per year from allometries

Independent of the extant model used, the estimated dinosaur AEN did not exceed 850 eggs (75,000 kg sauropod) for any of the taxa studied. According to most estimations, dinosaurs lay less than 200 eggs per year and only some estimates obtained for medium to large sized sauropods were higher (Table 1). Assuming the bird model, for example, the AENs of theropods were comparable to fossil clutch sizes. This suggests that theropods had one clutch per year (Table 2). By contrast, hadrosaurs and sauropodomorphs probably had more than one clutch per year. This is supported by AEN estimates of nearly all BM and CM combinations (minimum, maximum and average values for BM and CM are considered). For all hadrosaurs and sauropodomorphs, these AEN estimates exceeded the egg number of fossil clutches, independent of the extant model assumed. The AENs of the lambeosaurine dinosaur calculated under the reptile and bird model indicated that this taxon probably had one or a maximum of two clutches per year. The M. peeblesorum had at least two clutches per year under the reptile model, and up to 11 clutches under the bird model (Table 3). Depending on the extant models applied for M. carinatus this taxon might have had one up to two (reptile model) or three up to six (bird model) clutches per year (Table 1). Small / young sauropods (BM ~ 5,000 kg), likely producers of oospecies M. mammilare or M. patagonicus eggs, might have had one (reptile model) up to two (reptile model) or two (reptile model) up to four (bird model) clutches per year (Table 1). Medium sized sauropods (BM ~ 22,399 kg) might have laid two or three (M. mammilare, reptile model), but up to six (M. mammilare, bird model) clutches per year, depending on the model employed. For M. patagonicus, the number of clutches per year ranged from five (reptile model) to a maximum of 13 (M. patagonicus, bird model). The AEN estimate for very large sauropods (BM ~ 75,000 kg) was also variable depending on the species and model used. Estimates range from six (M. mammilare, reptile model) up to 15 (M. mammilare, bird model) or even 13 (M. patagonicus, reptile model) up to 30 (M. patagonicus, bird model) clutches per year (Table 1).
Reptiles (Figure 1). In contrast, large reptiles have a larger hypothesis (i), the reproductive traits of dinosaurs that are dinosaur reproductive biology. First, corroborating our birds [9,15]. This results in less distinction between clutch masses/annual clutch masses of large birds and reptiles than the size difference between a dinosaur egg and the egg-laying mass) are highly correlated in extant reptiles and birds in egg masses (Figures 1, 2 and 3).

Reproductive investment in dinosaurs

In summary, our results revealed four important insights into dinosaur reproductive biology. First, corroborating our hypothesis (i), the reproductive traits of dinosaurs that are considered to be more bird-like (theropods) did indeed coincide with reproductive traits of birds. Similarly, those traits of dinosaurs that were probably more reptile-like (prosauropods, sauropods) coincided with those of reptiles. Second, although the size difference between a dinosaur egg and the egg-laying female is very impressive, for all dinosaurs studied the egg to body mass relationship was similar to similar-sized or scaled-up reptiles (in M. carinatus) or even higher (in all other dinosaurs). However, it was lower than in similar-sized or scaled-up birds. Third, contrary to our hypothesis (ii) clutch masses of all dinosaurs and even of sauropods matched at least one of the extant models. We thus did not find any evidence that sauropods clutch sizes are small in comparison to their body mass. This in turn questions the idea that a physiological limitation imposed on the clutch [25] leads to the “small” clutch size of fully buried sauropod clutches. Under such a limitation, the predicted CM to BM relationship would be too high in large dinosaurs, regardless of the extant model used. Fourth, annual clutch mass estimates (iii) suggest that theropods had only one clutch per year, whereas all other studied dinosaurs had probably several clutches per year (except for the lambeosaurine hadrosaur, for which one clutch per year is also realistic). This is especially true for the large sauropods. However, contrary to our expectation (iv), most of the dinosaurs studied probably laid no more than 200 eggs per year (Tables 1, 2 and 3). Even large sauropods (75,000 kg) probably had less than 400 eggs per year (Table 1), which is a smaller annual egg number than extant sea turtles (up to 513 eggs [47]).

**Egg mass.** Our results suggest that the egg masses of most dinosaurs match neither the egg masses of similar-sized or scaled-up birds nor those of reptiles, but were in fact in-between (Figure 1). This could reflect the reproductive strategy differences of most dinosaurs compared to the reproductive strategy seen in extant birds or reptiles [1] and suggests that their reproductive strategy was intermediary [24]. The great variability in egg mass to body mass relations found in dinosaurs (Figure 1) could indicate that different reproductive strategies existed in dinosaurs. The suggested variability in reproduction strategies is corroborated by the variability seen in extant birds or reptiles [1] and suggests that the reptile reproductive model is pliosomorph and the bird model is phylogenetically derived, none of the studied dinosaurs with egg masses close to the reptile model belong to the theropods (Figure 1). Furthermore, the egg mass of the

### Table 2. Reproductive characteristics of theropods as documented in the fossil record and estimated by the reptile (crocodiles, tortoises) and the bird model.

| Taxon                  | Fossil        | Reptile model | Bird model |
|------------------------|---------------|---------------|------------|
|                        | BM (kg)       | CS            | CS [a] CS [b] | AEN | CY | CS [c] | AEN | CY |
| *Troodon formosus*     | 34            | 23 [22, 24]   | 2.4 6.5    | 6.6 | 0.3 [0.3, 0.3] | 15.9 | 17.0 | 0.7 [0.7, 0.8] |
| *Troodon formosus*     | 44            | 23 [22, 24]   | 2.9 8.2    | 8.2 | 0.4 [0.3, 0.4] | 19.2 | 20.4 | 0.9 [0.9, 0.9] |
| *Troodon formosus*     | 51            | 23 [22, 24]   | 3.2 9.1    | 9.1 | 0.4 [0.4, 0.4] | 21.3 | 22.8 | 1.0 [0.9, 1.0] |
| *Oviraptor philoceratops* | 33        | 24 [20, 30]   | 2.9 8.4    | 8.4 | 0.3 [0.3, 0.4] | 19.6 | 20.8 | 0.9 [0.7, 1.0] |
| *Oviraptor philoceratops* | 37       | 24 [20, 30]   | 3.2 9.1    | 9.1 | 0.4 [0.3, 0.5] | 21.3 | 22.6 | 0.9 [0.8, 1.1] |
| *Citipati osmolskae*   | 40            | 24 [20, 30]   | 3.4 9.6    | 9.6 | 0.4 [0.3, 0.5] | 22.5 | 24.0 | 1.0 [0.8, 1.2] |
| *Lourinhanosaurus antunesi* | 79      | 22 [15, 30]   | 3.0 8.7    | 8.7 | 0.4 [0.3, 0.6] | 20.3 | 21.7 | 1.0 [0.7, 1.4] |
| *Lourinhanosaurus atunesi* | 176     | 63 [25, 100]  | 4.2 12.1   | 12.2 | 0.2 [0.1, 0.1] | 26.3 | 30.5 | 0.5 [0.3, 1.2] |

BM: minimum, mean and maximum fossil body mass of a taxon taken from literature in kilograms; CS: mean clutch size observed in fossil record, with minimum and maximum fossil values given in square brackets or CS calculated from an allometric clutch mass (CM) model (CS [a] = tortoise model, CS [b] = crocodile model, CS [c] = bird model) using the fossil egg mass (EM, for calculating of fossil egg mass see Material and Method), CS = CM divided by EM. AEN (annual egg number): total number of eggs laid per year, calculated from an allometric ACM (annual clutch mass) model using the fossil egg mass, AEN = ACM divided by EM. CY: number of clutches per year, calculated from an allometric ACM model using the fossil CS and estimated AEN. CY = AEN divided by CS. Minima and maxima are given in brackets. References for fossil data are given in Table S3 (supplementary). Equations for the reptile and the bird model are given in Table S6. Note: For *Citipati osmolskae* and *Lourinhanosaurus antunesi* only one BM estimate was available.

**Discussion**

The allometries of body mass and reproductive investment in extant amniotes

Our results corroborate that body mass and reproductive investment (in terms of egg mass, clutch mass or annual clutch mass) are highly correlated in extant reptiles and birds [9,11–17]. In amniotes, the relative reproductive investment generally declines with body mass, whereas the absolute reproductive investment increases ([16,46], Figures 1–3, Table S3). Our analysis provides additional evidence [9,15] that the egg mass of large birds is higher compared to similar-sized reptiles (Figure 1). In contrast, large reptiles have a larger number of eggs per clutch and/or per year than similar-sized birds [9,15]. This results in less distinction between clutch masses/annual clutch masses of large birds and reptiles than in egg masses (Figures 1, 2 and 3).
most basal sauropodomorph (M. carinatus) matched both the crocodile and tortoise model well. Thus our results corroborate our hypothesis (i).

**Clutch mass/size.** In contrast to the egg masses, all dinosaur (mean) clutch masses matched the masses of similar-sized or scaled-up birds or reptiles.

Theropods. As expected under our initial hypothesis, the bird model was the best model for theropods. Fossils indicate that at least some avian reproductive characteristics, such as adult brooding [22,50–52], asymmetrical eggs [22,50,53,54], unornamented eggshell surface and complex eggshell ultrastructure, existed in non-avian theropods [55]. Thus, our results provide further evidence of a bird-like reproduction mode in theropods. Furthermore, our results on theropods suggest accurate body mass and egg mass estimates and the completeness of fossil clutches. The stronger deviation of L. antunesi from the bird model could possibly be explained by a higher inaccuracy in the estimates of its body mass (a not fully grown sub-adult individual is the holotype of this taxon, Mateus et al. [56]) and fossil clutch size (eggs of the clutch could come from different females [57]) than in the other theropods.

Varricchio et al. [23] assumed that some theropods received/ provided paternal care, because clutch-associated adults lack the maternal and reproductively associated histological feature common to extant archosaurs, including the medullary bone. Furthermore, theropods have relatively large clutch volumes. However, our analyses revealed no large clutch masses relative to body masses for theropods when compared to the studied extant birds showing biparental or maternal care. Thus, our data provide no evidence for the postulation presented in Varricchio et al. [23], that the theropods “sitting” on eggs were really males. The discrepancy in the results could have been caused by different sample compositions. We focused only on preadult birds in our analyses and used female body masses, as far as possible. We did not take into consideration the different parental care strategies of species. In contrast, Varricchio et al. [23] mixed different development modes of birds and used body mass averages without accounting for differences between sexes, but did allow for different parental care strategies. However, a recent study [58] corroborates our conclusion, showing that the development mode is a better predictor of the parental care strategy than clutch mass.

Sauropodomorpha. Contrary to our initial hypothesis (ii) clutch masses of sauropods were consistent with an extant species model, the tortoise model (Figure 2D). Several authors have argued that the clutch sizes of buried clutches in sauropods are bounded by physiological constrains [25,27,59], resulting in lower clutch mass to body mass relations compared to smaller taxa. Our analysis showed that the mean clutch masses for all studied dinosaurs matched the 95% prediction interval of at least one of the extant species models (birds, crocodiles or tortoises). Hence, they could be still consistent with the extant variability. For the two analyzed sauropods, the largest dinosaurs in our dataset, body mass and clutch size (particularly for M. mamillare) [27,49, but see 60] is uncertain. However, even when assuming large errors in the body mass and clutch mass/size estimates for these two sauropods (Figure 2, scattered rectangles), the clutch mass to body mass relations did not conflict with those seen in scaled-up recent taxa (Figure 2D, the rectangles are completely located within the 95% prediction interval of the tortoise model). Additionally, the clutch mass of the prosauropod M. carinatus is also well described by the tortoise model (Figure 2D). All these observations suggest that the tortoise model might be appropriate for sauropodomorphs in general. Thus, our results provided no evidence that the “small” clutch sizes of M. mamillare are caused by physiological limits imposed on the clutch [25,27,59]. We think that the use of a sea turtle model, as Seymour [25] did, to determine physiological limits on a large buried sauropod clutch is problematic. Sea turtles bury their clutches much deeper than most other reptiles [61]. In crocodilian clutches, for example, the respiratory gas pressure is closer to the atmospheric level than in sea turtle clutches [61]. In a buried clutch of the turtle species Chelodina expansa the respiratory gas pressure is also similar to the atmospheric pressure [62]. Thus, oxygen availability plays a stronger role in sea turtle clutch size than in other egg-burying reptiles, and is presumably not such a limiting factor in sauropods.

Hadrosaurids. The applicability of allometric models for clutch mass differed between the two hadrosaurids. For the lambeosaurine hadrosaur, the bird model was best, but the crocodile model was also applicable. For M. peeblesorum, the tortoise model was best; the crocodile model was also applicable, but only when assuming the lowest body mass estimates for that species. This discrepancy could indicate that reproduction strategies differed in hadrosaurs, as already suggested by Horner [63]. However, our results could be biased by an incomplete M. peeblesorum fossil clutch count. This would lead to a low assumed clutch mass. Horner [63] noted that counting individual eggs in a M. peeblesorum clutch was very difficult and for this reason assumed that one clutch consisted of at least 16 eggs.

**Annual clutch mass/clutch per year/annual egg number.** A reliable estimate of the number of clutches per year and the annual egg number for dinosaurs is uncertain because of the high variability observed in traits of extant species. Furthermore, there is a high inaccuracy in body mass and clutch mass estimates of dinosaurs, making it difficult to completely rule out any of the extant models for dinosaur taxa. Irrespective of all these limitations, we are able to provide qualitative estimates for the number of clutches per year and the annual egg number laid by dinosaurs. By using further information from the fossil record, we were able to identify the most likely extant model for a taxon.

Theropods. Fossil egg masses and clutch masses of theropods are consistent with the bird model, which suggests a general applicability of allometries on reproductive traits of birds to theropods. Assuming one clutch per year for theropods, the theropod ACMs match the bird model (Figure 3B). Furthermore, the low annual breeding frequency of theropods might provide further evidence for parental care in this taxon [23].

Sauropodomorpha. ACMs of sauropodomorphs were lower than all “average” extant species studied when assuming the studied sauropodomorphs (Figure 3) to have one clutch per year and mean values for BM and CM/CS. However, with the
exception of *M. patagonicus*, ACMs still fall within the 95% prediction interval of the reptile model (Figure 3C). Due to this observation and the finding that sauropods/sauropodomorphs were more "reptile-like" in their reproductive mode [24], we think that the reptile model is more appropriate for Sauropodomorpha than the bird model. The bird model provided high numbers of clutches per year, which could be flawed. These high estimates could be the result of incomplete fossil clutches (estimated clutch size is too small) or an assumed body mass too large for the producer. It is also possible that the bird model is not applicable for sauropods in general because of the more "reptile-like" reproduction strategy in sauropods.

Under the reptile model, an "average" *M. carinatus* could have had one or a maximum of two clutches, and laid between 40 and 81 eggs per year. Our estimates of the number of clutches per year for sauropods are more imprecise than those for the theropods and prosauropod. The two studied oospecies were assigned to the taxon *Titanosauria*, which covers a wide range of body masses for the potential egg producer. However, even if the body mass of a fully grown adult would be known, a wide range of body masses for the egg producer is still possible because sauropods probably became sexually mature well before they were fully-grown [64,65]. Irrespective of all these uncertainties, the reptile model revealed reliable estimates of the number of eggs and clutches per year for Sauropodomorpha. A sauropod weighing 75,000 kg could have laid the eggs of the *M. patagonicus* oospecies and is predicted to have had twelve clutches per year. This would result in an annual egg number of 340. However, the small eggs of the *M. patagonicus* oospecies imply that they were likely produced by a small to medium sized sauropod. Based on the smaller clutch size found in *M. mammiliare* compared to *M. patagonicus*, Sander et al. [27] argued that *M. mammiliare* had several clutches per year, whereas *M. patagonicus* had only one clutch per year. Contrary to these authors, our results indicate that both sauropods had multiple clutches per year, assuming that the egg laying individuals were not very small sauropods (BM < 5,000 kg, Table 1). Based on the reptile model, we suggest that *M. patagonicus* had between two and five clutches per year resulting in an annual egg production between 48 and 142, whereas *M. mammiliare* could have laid one up to six clutches per year resulting in around 16 but up to 114 eggs per year.

Altogether, our results imply that sauropods probably had several clutches per year, resulting in more than one hundred eggs per year. However, sauropod clutch and egg numbers probably did not exceed the numbers found in some recent reptile species (e.g. sea turtles [66]). Nevertheless, the high annual egg numbers estimated for Sauropodomorpha in comparison to other non-avian dinosaurs, recent birds or mammals could indicate a high predation rate of hatchlings and little parental care in this dinosaurian lineage.

Hadrosaurs. We could not clearly identify the most likely extant model for the ACM of hadrosaurs. The ACM to BM relation of the lambeosaurine hadrosaur was intermediary to both the bird and reptile model; for *M. peeblesorum* its relation was lower than observed in any extant model. Nevertheless, our results suggest that hadrosaurs probably had several clutches per year. For the lambeosaurine hadrosaur, one (reptile model) but up to two clutches (bird model) per year is estimated (Table 3). *Maiasaura peeblesorum* probably had more clutches per year than the lambeosaurine hadrosaur. The reptile model estimates two, up to four clutches per year and the bird model five up to eleven (Table 3). However, as discussed before, the assumed clutch size and mass of *M. peeblesorum* could be too low. The resulting number of clutches per year estimated could therefore be much closer to those of the lambeosaurine hadrosaur. In any case, the annual egg numbers of hadrosaurs studied were less than 200 (Table 3).

**Ecological implications**

Why should some dinosaurs have several clutches per year and others not? As seen in extant species, they could have lived in different environments, each favoring different reproductive strategies. Producing several small clutches within a breeding season could reflect a bet-hedging strategy [67–69]. If few of many clutches are lost to predation or other unfavorable environmental conditions cause considerable egg mortality, the eggs/hatchlings from other clutches may survive by chance. Such a strategy is favorable in environments with a long breeding season and when the time intervals between clutches are relatively long. This is also true in environments with a short breeding season, when clutches are laid more or less simultaneously. Dinosaurs which had one clutch per year could have lived in environments with a short breeding season. Under such environmental conditions putting all eggs into a single clutch/reproductive event (producing all eggs in a specific time period) is the only option, even at high rates of egg mortality, because the length of the breeding season limits reproduction [70]. Irrespective of the length of the breeding season, one clutch per year can be sufficient if offspring mortality due to environmental conditions is low.

**Evolutionary implications**

Since birds are dinosaurs, there must have been an evolutionary shift in the reproductive mode from the basal reptilian/non-avian dinosaur mode to that currently observed in birds. This shift might be observable in the studied dinosaurs. As expected, all studied reproductive traits of Sauropodomorpha were more reptile-like, whereas traits of studied theropods conform well to those of recent birds. Furthermore, it is likely that within the dinosaur lineage (including birds), an increase in egg size was linked to a decrease in egg numbers per clutch/year and vice versa. The prosauropod had many small eggs for its body mass. The two sauropods had in fact larger eggs than the prosauropod, but they still had many eggs in comparison to other dinosaurs because of their large size (Figure 1, Tables 1, 2 and 3). The four theropods had larger but fewer eggs than the sauropodomorphs (Figure 1, Tables 1 and 2), whereas recent birds have the largest eggs in comparison to their body mass (Figure 1) and also the fewest egg numbers (2.2-4.5 eggs per clutch, geometric mean of 5290 bird species [33]). This evolutionary change in egg size and number observed in the dinosaurian clade probably coincides with other changes in life
history traits. High mortality during the egg and juvenile phase could have led to the evolution of and selection for parental care; in birds, this would have sustained viable populations over evolutionary time. A similar shift in life history traits might have also occurred in the hadrosaurian lineage (Ornithischia). Interestingly, other researchers have also assumed extended parental care for species of this lineage based on fossil clutches with “altricial hatchlings” (e.g. Maiasaura peeblesorum [71,72], but see 48,73). However, our results imply that extended parental care was more likely in the studied lambeosaurine hadrosaur than in M. peeblesorum. This is due to the increased size observed in eggs and estimated fewer egg/clutch numbers in the former taxon.

Conclusion

From our study we conclude i) that allometric regression functions are a suitable approach to describe the relation between body mass and the studied reproductive traits in birds or reptiles. It is ii) appropriate to transfer these established allometries to specific taxa of extinct non-avian dinosaurs. Although we found a high variability in reproductive traits around the (average) allometric regression lines in extant species, we think that the results provide new testable hypotheses about dinosaur reproduction, its evolution and their ecological implications, especially for reproductive traits that are insufficiently documented or lacking in the fossil record.

Supporting Information

Table S1. Average body mass and reproductive characteristics of dinosaurs as documented in the fossil record. The Megaloolithus patagonicus oospecies is assigned to a titanosaurian sauropod based on embryonic remains in the eggs, and the Megaloolithus siruguei/mammillare egg type (with a highly porous shell) is commonly assigned to titanosaurian sauropod dinosaurs, because titanosaur bones had been found in the same horizon or formation as the eggs. It should be noted that Megaloolithus siruguei is considered as a junior synonym of Megaloolithus mammillare. Megaloolithus eggs have also been assigned to titanosaurs because of the find of a hatchling in a nest of Megaloolithus eggs from India. Although taxonomic identification of the eggs and their producers is problematic in sauropods, species with a mass of at least 5000 kg were assigned to both Megaloolithus oospecies. BM = body mass. ES = egg size, expressed in length (L) and diameter (D). EM = egg mass. CS = clutch size, number of eggs per clutch. CM/ACM = clutch mass (CM; CM = EM x CS) respectively annual clutch mass (ACM). ACM equals CM because as a first approximation we assumed one clutch per year for all dinosaurs studied. Values in brackets are minimum and maximum values, where no maxima are given only one value was available. Note: the mean BMs for both titanosaur taxa was established from body masses of sauropods larger than 5000 kg (because sauropod species with a BM of at least 5000 kg were assigned to both Megaloolithus oospecies) and for which the body mass estimation method was given in the source. The exact body mass range is 6853 kg to 72936 kg, however we assumed, as a somewhat more conservative measure, minimum and maximum body masses of 5000 kg and of 75000 kg.

Table S2. Data used in the allometric analyses of extant species. G = Galliformes, A = Anseriformes, S = Struthioniformes, C = Crocodylia, T = Testudinidae, BM = body mass, EM = egg mass, CM = clutch mass, ACM = annual clutch mass.

Table S3. Results of the OLS regression analyses carried out for body mass against several reproductive traits for tortoises, crocodiles and birds. Allometric functions follow c×BM^b (c+b×BM, log-log-plot), where BM is the body mass in kilograms, c is the intercept and b the slope of the line in a log-log-plot. N = sample size. 95% CI = 95% confidence interval of c or b. EM = egg mass. CM = clutch mass. ACM = annual clutch mass.

Table S4. Summary of results obtained in ANCOVAs. With these analyses we tested for differences in regression slopes ((a); for EM, CM, ACM) and intercepts ((b); CM, ACM) obtained for birds, crocodiles and tortoises under ordinary least squares regression analysis. EM = egg mass, CM = clutch mass, ACM = annual clutch mass, group = categorical variable, coding whether the species is a bird, crocodile or tortoises, log = logarithm to the base 10. n.a. = not applicable. For sample sizes refer to Table S3.

Table S5. Summary of results obtained in ANCOVAs. With these analyses we tested for differences in regression slopes ((a); for EM, CM, ACM) and intercepts ((b); CM, ACM) obtained for crocodiles and tortoises under ordinary least squares regression analysis. EM = egg mass, CM = clutch mass, ACM = annual clutch mass, group = categorical variable, coding whether the species is a crocodile or tortoises, log = logarithm to the base 10. n.a. = not applicable. For sample sizes refer to Table S3.

Table S6. Allometric models used to estimate reproductive traits of non-avian dinosaurs. EM = egg mass, CM = clutch mass, ACM = annual clutch mass; c = intercept and b = slope of the respective allometry.

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**Author Contributions**

Conceived and designed the experiments: JW EMG. Performed the experiments: JW. Analyzed the data: JW.

**References**

1. Sander PM, Claus M (2008) Sauropod gigantism. Science 322: 200-201. doi:10.1126/science.1158094. PubMed: 18845734.

2. Griebeler EM, Werner J (2011) The life-cycle of sauropods. In: N KleinC RemesCT GeePM Sander. Biology of the saurapod dinosaurs: understanding the life of giants. Bloomington: Indiana University Press. pp. 263-275.

3. Szykman RA (1977) Respiratory exchange of sea turtle nests. Respir Physiol 31: 19-38. doi:10.1016/0034-5887(77)90062-7. PubMed: 918411.

4. Sander PM, Peitz C, Jackson FD, Chiappe LM (2008) Upper Cretaceous titanosaur nesting sites and their implications for sauropod reproductive biology. Palaeontogr Abt A 284: 69-107.

5. Padian K, de Ricqlès AJ, Horner JR (2001) Dinosaurian growth rates and bird origins. Nature 412: 405-408. doi:10.1038/35086500. PubMed: 11473307.

6. Klein C, Remes CT, Gee PM, Sander PM, Ackerman RA (1977) Avian paternal care had dinosaur origin. Science 322: 1826-1828. doi:10.1126/science.1905938.

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12. Testudines). Herpetol Monogr 6: 25-42. doi:10.2307/1466960.

17. Peters RH (1993) The ecological implications of body size. Cambridge: Cambridge University Press. 344pp.

22. Peters RH (1983) The ecological implications of body size. Cambridge: Cambridge University Press.

27. Hendriks AJ, Mulder C (2008) Scaling of offspring number and mass to plant and animal size: model and meta-analysis. Oecologia 155: 705-716. doi:10.1007/s00442-007-0952-3. PubMed: 18196279.

32. Sander PM, Peitz C, Jackson FD, Chiappe LM (2008) Upper Cretaceous titanosaur nesting sites and their implications for sauropod reproductive biology. Palaeontogr Abt A 284: 69-107.

37. Weishampel DB, Dodson P, Osmolska H, editors (2004) The Dinosauria: University of California Press.

42. Turner AH, Pol D, Clarke JA, Erickson GM, Norell MA (2007) A basal dromaeosaurid and size evolution preceding avian flight. Science 317: 1378-1381. doi:10.1126/science.1144066. PubMed: 17823350.

47. Sander PM, Peitz C, Jackson FD, Chiappe LM (2008) Upper Cretaceous titanosaur nesting sites and their implications for sauropod reproductive biology. Palaeontogr Abt A 284: 69-107.

52. Paul GS (1994) Dinosaur reproduction in the fast lane: implications for size, success, and extinction. In: K CarpenterKF HirschJR Horner. Dinosaur eggs and babies. New York: Cambridge University Press. pp. 244-255.

57. Horner JR (2000) Dinosaur reproduction and parenting. Annu Rev Earth Planet Sc 28: 19-45.

62. Peters RH (1993) The ecological implications of body size. Cambridge: Cambridge University Press. 344pp.

67. Sander PM, Peitz C, Jackson FD, Chiappe LM (2008) Upper Cretaceous titanosaur nesting sites and their implications for sauropod reproductive biology. Palaeontogr Abt A 284: 69-107.

72. Padian K, de Ricqlès AJ, Horner JR (2001) Dinosaurian growth rates and bird origins. Nature 412: 405-408. doi:10.1038/35086500. PubMed: 11473307.

77. Martin TE, Li P (1992) Life history traits of open- vs. cavity-nesting birds. Ecology 73: 579-592. doi:10.2307/1940764.

82. Zardoya R, Meyer A (2001) The evolutionary position of turtles revised. Naturwissenschaften 88: 193-200. doi:10.1007/s001140100228. PubMed: 11482432.

87. Iwabe N, Hara Y, Kumazawa Y, Shibamoto K, Saito Y et al. (2005) Sister group relationship of turtles to the bird-crocodilian clade revealed by nuclear DNA-coded proteins. Mol Biol Evol 22: 810-813. doi:10.1093/molbev/msi075. PubMed: 15625185.

92. Chacartegui J, Funes A, Serna J, Sánchez E (1997) Hatching success and hatching rate in gecko (Gekkonidae) eggs from different habitats. J Zool 246: 49-55. doi:10.1111/j.1469-7998.1997.tb02692.x.

97. Webster SM, Leventhal P, Fowles J, Wynn-Jones J (1996) Temperature controls on hatching success and hatching rate in newts. J Zool 240: 49-55. doi:10.1111/j.1469-7998.1996.tb02692.x.

102. Iverson JB (1992) Correlates of reproductive output in turtles (Order Testudines). Herpetol Monogr 6: 25-42. doi:10.2307/1466960.

107. Rahn H, Paganeli CV, Ar A (1975) Relation of avian egg weight to hatching success. Auk 92: 750-765. doi:10.2307/4084786.

112. Peters RH (1983) The ecological implications of body size. Cambridge: Cambridge University Press.

117. Rahn H, Paganeli CV (1975) Relation of avian egg weight to hatching success. Auk 92: 750-765. doi:10.2307/4084786.

122. Martin TE, Li P (1992) Life history traits of open- vs. cavity-nesting birds. Ecology 73: 579-592. doi:10.2307/1940764.

127. Rahn H, Paganeli CV (1975) Relation of avian egg weight to hatching success. Auk 92: 750-765. doi:10.2307/4084786.
46. Reiss MJ (1985) The allometry of reproduction - why larger species invest relatively less in their offspring. J Theor Biol 113: 529-544. doi: 10.1016/S0022-5193(85)80037-0.

47. Hirth HF (1980) Some aspects of the nesting behavior and reproductive biology of sea turtles. Am Zool 20: 507-523.

48. Carpenter K (1999) Eggs, nests, and baby dinosaurs: a look at dinosaur reproduction. Bloomington: Indiana University Press.

49. Jackson FD, Varricchio DJ, Jackson RA, Vila B, Chiappe LM (2008) Comparison of water vapor conductance in a titanosaurs egg from the Upper Cretaceous of Argentina and a Megaloolithus siruguei egg from Spain. Paleobiology 34: 229-246. doi: 10.1666/0094-8373(2008)034[0229:COAVCN]2.0.CO;2.

50. Norell MA, Clark JM, Chiappe LM, Dashzeveg D (1995) A nesting dinosaur. Nature 378: 774-776. doi:10.1038/378774a0.

51. Dong ZM, Currie PJ (1996) On the discovery of an oviraptorid skeleton on a nest of eggs at Bayan Mandahu, Inner Mongolia, People's Republic of China. Can J Earth Sci 33: 631-636. doi:11.1093/e96-046.

52. Varricchio DJ, Jackson F, Trueman CN (1999) A nesting trace with eggs for the Cretaceous theropod dinosaur Troodon formosus. J Verteb Paleontol 19: 91-100. doi:10.1080/02724634.1999.10011125.

53. Horner JR, Weishampel DB (1988) A comparative embryological study of two ornithischian dinosaurs. Nature 332: 256-257. doi: 10.1038/332256a0.

54. Horner JR, Weishampel DB (1996) A comparative embryological study of two ornithischian dinosaurs (vol 332, pg 256, 1988). Nature 383: 10.1038/332256a0.

55. Zelenitsky DK, Therrien F (2008a) Phylogenetic analysis of reproductive traits of maniraptoran theropods and its implications for egg parataxyonomy. Palaeontology 51: 807-816. doi:11.1011/j.1475-4963.2008.00770.x.

56. Mateus O, Walen A, Antunes MT (2006) The large theropod fauna of the Lourinhã Formation (Portugal) and its similarity to that of the Morrison Formation, with a description of a new species of Allosaurus. In: JR Foster SG Lucas. Paleontology and geology of the Upper Jurassic Morrison Formation. New Mexico: Museum of Natural History & Science Bulletin 36. pp. 123-130

57. Mateus I, Mateus H, Antunes MT, Mateus O, Taquet P et al. (1998) Upper Jurassic theropod dinosaur embryos from Lourinhã (Portugal) and comparison with those of Megalosaurus from Oxford. Biol Lett 9: 37-40. doi:10.1098/rspb.1999.0500. PubMed: 10.1038/282296a0.

58. Birchard GF, Ruta M, Deeming DC (2013) Evolution of parental incubation behaviour in dinosaurs cannot be inferred from clutch mass in birds. Biol Lett 9: 20130036. PubMed: 23676654.

59. Seymour RS, Ackerman RA (1980) Adaptations to underground nesting in birds and reptiles. Am Zool 20: 437-447.

60. Vila B, Jackson FD, Fortuny J, Sellés AG, Galobart À (2010) 3-D modelling of megaloolithid clutches: insights about nest construction and dinosaur behaviour. PLOS ONE 5: e10362. doi:10.1371/journal.pone.0010362. PubMed: 20403853.

61. Booth DT, Thompson MB (1991) A comparison of reptilian eggs with those of megapode birds. In: DC Deeming MWJ Ferguson. Egg incubation: its effects on embryonic development in birds and reptiles. Cambridge, UK: Cambridge University Press. pp. 117-146.

62. Booth DT (1998) Nest temperature and respiratory gases during natural incubation in the broad-shelled river turtle, Chelodina expansa (Testudinata : Chelidae). Aust J Zool 46: 183-191. doi:10.1071/ZO98003.

63. Horner JR (1999) Egg clutches and embryos of two hadrosaurian dinosaurs. J Vertebr Paleontol 19: 607-611. doi: 10.1080/02724634.1999.10011174.

64. Sander PM (2000) Long bone histology of the Tendaguru sauropods: implications for growth and biology. Paleobiology 26: 466-488. doi: 10.1666/0094-8373(2000)026.2000.

65. Griebeler EM, Klein N, Sander PM (2013) Aging, maturation and growth of sauropodomorph dinosaurs as deduced from growth curves using long bone histological data: an assessment of methodological constraints and solutions. PLOS ONE 8: e67012. doi:10.1371/journal.pone.0067012. PubMed: 23840575.

66. SWOT (2012) The state of the world’s sea turtles. Available: http://sawturlestatus.org/learn/meet-the-turtles. Accessed 2012 November 13.

67. Slatkin M (1974) Hedging one's evolutionary bets. Nature 250: 704-705. doi:10.1038/250704b0.

68. Olofsson H, Ripa J, Jonzén N (2009) Bet-hedging as an evolutionary game: the trade-off between egg size and number. Proc R Soc Lond B Biol Sci 276: 2963-2969. doi:10.1098/rspb.2009.0500. PubMed: 19474039.

69. Beaumont HUE, Gallie J, Kost C, Ferguson GC, Rainey PB (2009) Experimental evolution of bet hedging. Nature 462: 90-93. doi:10.1038/nature08504. PubMed: 19890329.

70. Griebeler EM, Caprano T, Böhning-Gaese K (2010) Evolution of avian clutch size along latitudinal gradients: do seasonality, nest predation or breeding season length matter? J Evol Biol 23: 888-901. doi:10.1111/j.1420-9101.2010.01958.x. PubMed: 20210827.

71. Horner JR, Makela R (1979) Nest of juveniles provides evidence of colonial nesting among ornithischian dinosaurs. Nature 282: 296-298. doi: 10.1038/282296a0.

72. Horner JR (1982) Evidence of colonial nesting and site fidelity among ornithischian dinosaurs. Nature 297: 675-676. doi:10.1038/297675a0.

73. Geist NR, Jones TD (1996) Juvenile skeletal structure and the reproductive habits of dinosaurs. Science 272: 712-714. doi:10.1126/science.272.5262.712. PubMed: 8662567.