

Estimation of diplodocid length for fragmentary fossils with a ‘suspension bridge’ (zero-torque) model

PHILIP L. KAHN

ABSTRACT—By using simple geometrical arguments, I predict the total length of *Diplodocus carnegii* using a least-squares fit on the cervical series, total dorsal series length, and the height of the neural spine on vertebra D10. This is done by using a simple, torque-based ‘suspension bridge’, or zero-torque, model after using a geometric model to compute neck and tail component masses. The successful results of this model to *Diplodocus* CM84 and to a recent specimen of *Apatosaurus ajax* suggest that this model is general among diplodocid sauropod dinosaurs. This model therefore can be used to provide length estimates for fragmentary diplodocid fossils.

INTRODUCTION

Dinosaur biomechanics can provide insights into the various problems that life encounters when it reaches massive sizes. Sauropod dinosaurs were the largest known land animals ever to live, peaking in size around the late Jurassic. Sauropod gigantism begs several questions, including ones related to energetics and biomechanical / structural concerns. Work on the BMR of sauropod dinosaurs (CITE RECENT READS) indicates metabolic rates that are comparable to modern mammals and birds. Though recent work suggests that raising the head above approximately 30° would be prohibitively expensive metabolically, the costs of holding the head at a level height are not commonly accounted for. Evaluating the energetics costs of this will demonstrate that for energetic efficiency, long-necked sauropods of the diplodocid body type must be in an energetically neutral position with its axial body elements held parallel to the ground. This assumption yields biomechanical results that permit a predictive model that allows total animal length to be estimated with few initial parameters.

Sauropods, particularly those of the low-slung, diplodocid body type have no modern analog. While reconstruction of brachiosaurids suggest comparisons with animals like the modern giraffe, diplodocids have a long neck that mammalian skulls are generally poorly suited to permit (CITE NEW), and long, suspended tails are generally absent in extant endotherms. This makes baseline comparisons difficult to implement. However, a brief discussion of the giraffe implementation of long necks is worthwhile.

A brief consideration of the energetics of sauropod necks, then, will provide a basis for consideration of a model which uses no net energy expenditure as its basic assumption. A *Diplodocus carnegii* stood X m at the shoulder with a neck length of L m. To estimate the torque that this exerts on the base of the neck, the cervical vertebrae, with a steadily reducing radius toward the head, is easily approximated as a conical frustrum.

Thus, to actively maintain the head at shoulder level, a constant torque of T would have to be exerted at the base of the neck.

University of California Museum of Paleontology, 1101 Valley Life Sciences Building, Berkeley, CA 94720-4780, pl.kahn@gmail.com

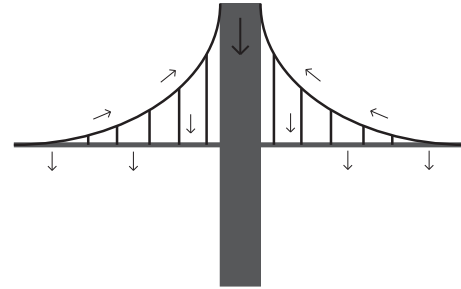


FIGURE 1. Caption about a bridge here. Are torque arrows needed? Loading transferred to support column. This basic unit is repeated along the length of a bridge.

Given this energy requirement for a simple neutral position, it is reasonable to assume that derived sauropods would have evolved some mechanism to passively maintain its head — an tail — against the pull of gravity. To this end, I endeavor to demonstrate that the previously proposed “suspension bridge” model (CITE) can be applied to a biological system and thus be used to predict the length of the animal with minimal information.

To establish the meaning of a “suspension bridge” in this context, consider a bridge like the Golden Gate in San Francisco. The weight of the road is supported by cables affixed to the road at regular intervals. This larger cable then attached to a large central support structure. The entire thing is kept balanced by an equal torque exerted on the opposite side, by the weight of that segment of the road, and a significant portion of the weight being loaded directly on the support (see Figure 1). The diplodocid-type sauropods can be modeled after this style, then, as shown in Figure 2. To follow a suspension-bridge approach, an anchor point is required, which I postulate to be the neural spines above the acetabulum. I chose this point for generally being the tallest neural spines in a diplodocid skeleton and for the structural support available from the pelvic girdle.

In order to avoid the colloquial term “suspension-bridge” model, despite being established before in the literature (Alexander 2006), I will refer to the approach as a “zero-torque” (ZT) approach, referring to the net zero torque on the support structure used. Thus, the corresponding mathematical model is one that has the primary load-bearing on connective elements. Such force-balance approaches have been used successfully before (Alexander 1989).

In an attempt to model these animals, I examine *Diplodocus carnegii* (Carnegie Museum 84) and construct a simple mathematical model to predict diplodocid sizes with fragmentary information. This modeling technique abstracts the form of the dinosaur to simple geometrical shapes and uses simple physical approximations to predict the length and mass

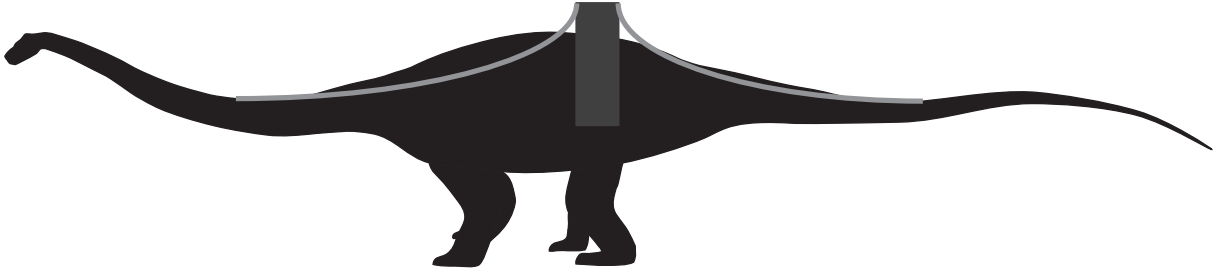


FIGURE 2. Analogous to Figure 1, in the proposed model the acetabulum of diplodocid dinosaurs act as the primary support structure for a ZT model. This is both close to the center of mass and is in-line with the rear legs of the animal for support. Here a simplified suspension bridge is superimposed on a silhouette of *D. carnegii* for clarity.

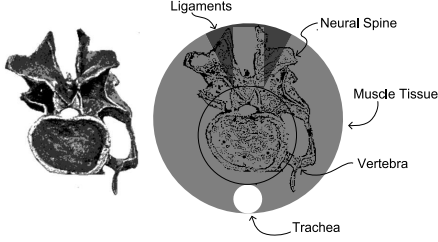


FIGURE 3. A cross-sectional schematic of the model neck construction. The elements identified here are subscripted in equations by b for the bone, l for the ligaments, m for muscle mass, ns for the neural spines, t for tissue mass, and tr for the trachea. The capital subscript T is added when these variables refer to the tail. For a side-schematic view, see Figure 5. Vertebrae taken from Hatcher (1901), plate VI.

of sauropod necks and tails that conform to this suspension bridge model, such as diplodocids.

To accomplish this, several parameters are needed. The ranges of neck motion have been modeled for CM 84 by Parrish and Stevens (1999), and those parameters are taken as canonical in this analysis (Stevens 2002; Stevens and Parrish 2005a). Despite possible problems with the result as discussed by Upchurch *et al.* (2000), the model constraints provided by the Parrish and Stevens (1999) analysis are important to this analysis, and the problems are generally limited to those that should not affect a rigid-body solution. However, the primary reason CM84 was chosen as the test case is because the animal has comprehensive measurements (Hatcher 1901) for the entire vertebral series.

This approach relies on the value of finding a least-squares linear fit to describe the radius of cervical vertebrae as a function of distance from the neck. By calculating the mass of bony elements, the musculature required to move the neck can be calculated, and the mass of the entire neck follows. The torque that the neck would then apply on dorsal spines can be equated to a torque that the tail must apply, yielding a quadratic that provides an accompanying tail length.

METHODS

Length and Mass Estimates of a *Diplodocus* Neck

By modeling a diplodocid neck as a series of reducing cylinders (of length l and density ρ) and idealizing this to a frustum of a cone, the total mass M of the neck can be calculated. This model is shown graphically in Figures 3 and

5. Pneumatic aspects of the skeleton are important to consider for sauropods, because they are relevant to thermoregulation problems, mass, and oxygen supply, and are well established in many sauropod lineages (Schwarz and Fritsch 2006; Wedel 2003, 2007). Because the pneumatic features are “virtually identical to those of birds” for all Neosauropoda, pneumaticity is assumed and denoted as k_p (Wedel 2005). For a description of subscript convention describing body parts, see Figure 3. All variables are used as is standard in physics notation, such as m for mass, ρ for volume density, r for radius, l for length, h for height, I for moment, and τ for torque. Capital letters M and L are used to represent total mass and length of a part, respectively. Given these assumptions, the total mass of the neck can be described by

$$M = m_t + m_b$$

$$m_b = k_p \int_l^0 \int_0^{r_b} 2\pi\rho_b r(l) dr dl + m_{ns} \quad (1)$$

$$m_{ns} = 2\pi \left(\frac{2}{3} r_b(l) \right)^2 h(l)$$

$$m_t = \int_l^0 \int_{\frac{\pi}{4}}^{\frac{7}{4}\pi} \int_{r_b}^r \rho_t r(l) dr d\theta dl - \int_0^{r_{tr}} 2\pi\rho_{tr} r dr + \int_0^{\frac{\pi}{8} r_t} \int_0^l 2\pi\rho_l r dl dr \quad (2)$$

This construction accounts for the basic morphology of the skeleton, the primary tissue mass, and an empty column for the trachea. The muscle is treated as leaving a 45° gap at the top of the cervical vertebrae for ligamentary attachment, which are added separately (Dzemski *et al.* 2007; Organ 2006; Schwarz *et al.* 2007b; Wedel 2002). Based on data from Wedel (2005), $k_p = 0.3$ is taken to be accurate to a first approximation, and assumed to apply to the entire neck. Figure 4 shows that the heights of the neural spines have a very strong linear fit ($p < 10^{-4}$) with the distance from the base of the neck, and this least-squares result can be used as the solution to $h(l)$. These fits were then taken at the locations of the vertebrae, assumed to have a radius of $\frac{2}{3}r_b(l)$, and computed as a cylinder. This result was then doubled to account for the bifurcated neural spines on *Diplodocus*.

Figure 4 also shows the radius of the cervical vertebrae falls off linearly with length to a good approximation. Table 1 shows the variation in the linear fitting coefficients for different

	C1 — C15	C1 — C14	C6 — C8	C1, C7, & C15	C2 — C4
Constant k_1	-1.74×10^{-2}	-1.92×10^{-2}	-2.27×10^{-2}	-1.42×10^{-2}	-2.64×10^{-2}
Offset k_2	1.55×10^{-1}	1.66×10^{-1}	1.74×10^{-1}	1.31×10^{-1}	1.84×10^{-1}
Uncertainty	7.62%	5.72%	40.51%	1.49%	37.67%
$ r $	0.97	0.98	0.93	1.00	0.94

TABLE 1. Taper coefficients and constant offset for various methods of measurement. Uncertainty is the maximum fractional uncertainty, $|r|$ is the correlation coefficient.

selections of vertebrae, for possible representative fragmentary finds. The resulting coefficient k_1 and constant k_2 are for equations of the form $r(l) = k_1 l + k_2$. The measurements used to calculate these fits were taken from Hatcher (1901). C1 — C14 are used for the rest of this analysis due to a high $|r|$. The fit using C1, C7 and C15 is left for completeness, but will generally not be practical in the field, because it would require knowing the distance between the elements before solving. Nevertheless, the table shows that the fit constant is relatively stable among various choices of vertebrae, reaching a mean of -2.00×10^{-2} with a standard deviation of $\sigma = 4.70 \times 10^{-3}$. Larger fractional uncertainties occur for fewer vertebrae closer to each other, but the results are nevertheless consistent among all fit types, with the smallest linear coefficient as 0.93. At four significant figures, the fit for the first two columns and the fit using C1, C7, and C15 are highly significant ($p < 0.01$). The other two fits fail this test when applied rigorously, with a two-tailed p -value of 24.5% and 22.9%. However, this is to be expected with a very small sample size, and the strong other fits suggest that this degree of correlation should be acceptable in practice. In particular, the fits with a correlation above 0.95 validate the continuous tapering model as used in Figure 4.

There are several possible methods by which the length of the neck can either be empirically determined or calculated, which is required to calculate the bony core mass and the moment of inertia. For CM84, the sum of the length of the cervical vertebrae comes to 7.29 m; however, when the vertebrae are connected and overlap of the centra is taken into account (neglecting intervertebral discs), a length of 6.7 m is obtained for the cervical series — average for a diplodocid (Hatcher 1901; Stevens and Parrish 2005b; Wedel 2005). With the constants given in Table 1, it is possible to calculate the length of the neck without actually knowing the entire cervical series. For example, by choosing a 3 cm final vertebral radius and using the fit from the second column, the predicted neck length is 7.05 m, which is intermediate between the two methods detailed above. The choice of a 3 cm minimum radius is based on the radius of the C2 vertebra in CM84 and an extrapolation from an *Apatosaurus ajax* described by Upchurch *et al.* (2004b). The worst fit, by using two adjacent vertebrae, under-predicts the neck length by nearly a meter, but still provides a guideline that is not entirely arbitrary. This method enables only partial cervical series to be of use in calculating the entire length of the animal, but is not as accurate as direct measurement from a complete or nearly-complete cervical series. For the purposes of this treatment, the calculated length $L = 7.05$ m is used.

Using a bone density of 3.16 g cm^{-3} (treating homogeneous dense bone as $\text{Ca}(\text{PO}_4)_3\text{OH}$), and a total neural spine mass of $1.40 \times 10^2 \text{ kg}$:

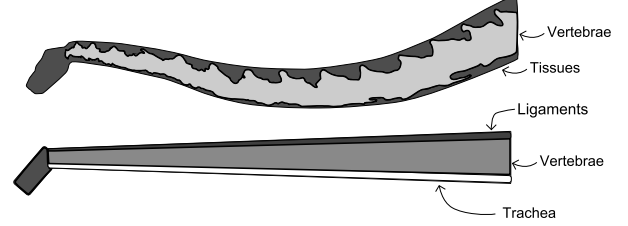


FIGURE 5. Side schematic view of model, reflecting neck tapering in model. The top image is a silhouette CM84, and the bottom image the proposed model.

$$\begin{aligned}
 m_b &= \rho_b \pi k_b \left(1.233 \times 10^{-4} l^3 - 1.592 \times 10^{-3} l^2 \right. \\
 &\quad \left. + 2.742 \times 10^{-2} l \right) \Big|_{l=7.05} + m_{ns} \\
 &\approx 6.09 \times 10^2 \text{ kg}
 \end{aligned}$$

If biomechanical limits on neck movement are established, the amount of musculature required to manipulate the neck on either side can be approximated. The muscle can then be extended as a column that reduces in radial size as it reaches the cranium.

The moment of inertia of the system can be calculated by looking at the torque of the neck on the lumbar vertebrae. Starting from the definition $I = \int r^2 dm$,

$$\begin{aligned}
 dm &= \pi r(b)^2 \rho_b dl \\
 &= k_p \rho_b \pi [-0.019l + .166]^2 dl \rightarrow
 \end{aligned}$$

$$\begin{aligned}
 I &= k_p \rho_b \pi \int_0^l l^2 [-0.019l + 0.166]^2 dl \quad (3) \\
 I &= \rho_b \pi k_p \left(7.40 \times 10^{-5} l^5 \right. \\
 &\quad \left. - 7.96 \times 10^{-4} l^4 + 9.14 \times 10^{-3} l^3 \right) \Big|_{l=7.05} \\
 &\approx 7.52 \times 10^3 \text{ kg}\cdot\text{m}^2 \quad (4)
 \end{aligned}$$

This result is accurate to a first approximation, assuming that each cross-sectional piece is of the same density. Given the assumption that majority of the moment of inertia is supplied by the bone, this approximation is justified because the neck tissues are generally isotropic and homogeneous.

To proceed, the extremes of neck flexion are needed. The movement axis can be rewritten in terms of spherical coordinates, with $(r, \theta, \phi) = (0, 0, 0)$ at the base of the neck, following mathematical rather than physics conventions (where ϕ is longitudinal). By transforming the Cartesian coordinates of Parrish and Stevens (1999) to spherical coordinates, one obtains a maximum ϕ range of approximately 34.9° (0.607 rad) and a maximum θ of 43.1° (0.752 rad) from

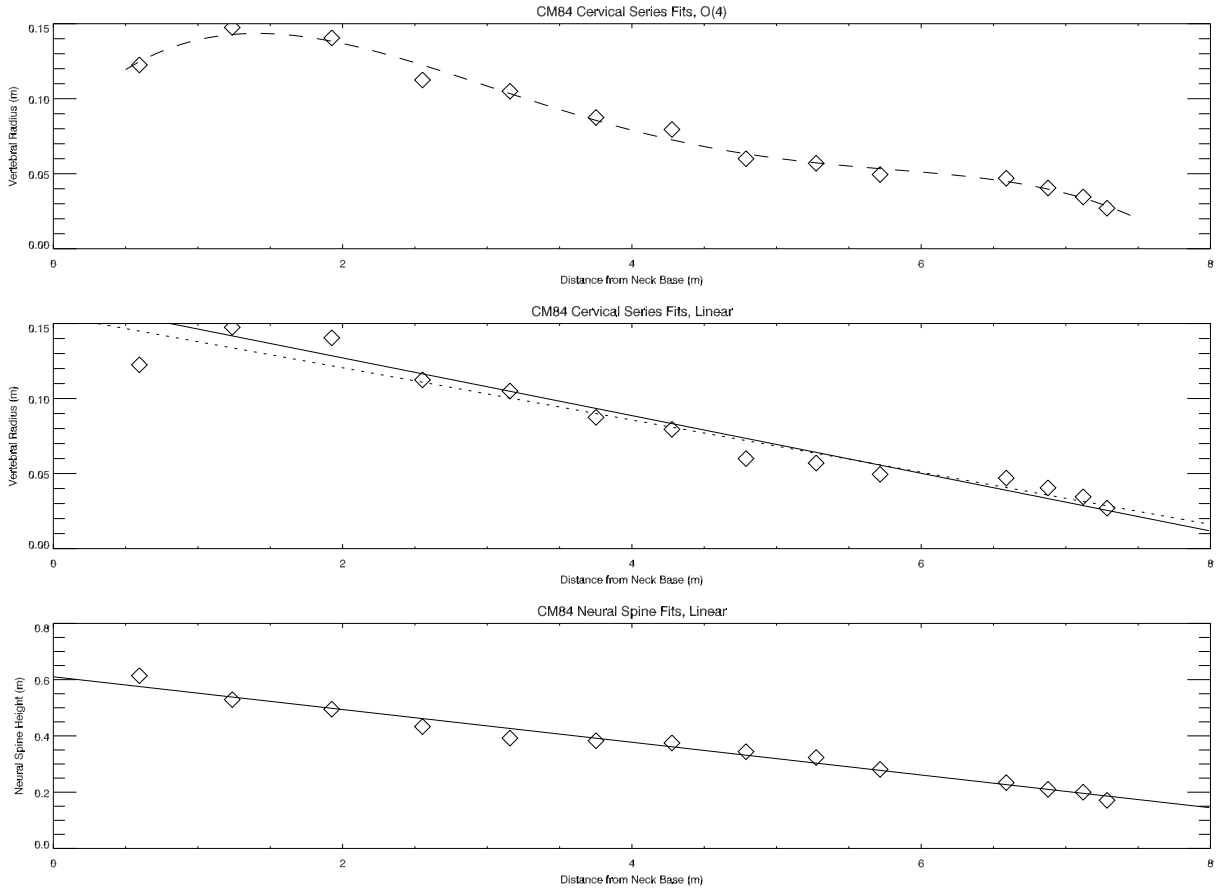


FIGURE 4. Plots of $O(4)$ and linear fits for the cervical series, and a linear fit for the neural spine heights. The dotted line is the complete least squares fit, the solid line the one neglecting the base cervical. It is this fit that is used in the text. The neural spine fit has $h(l) = -5.82 \times 10^{-2}l + 0.61$, a maximum fractional uncertainty of 4.21%, and $r = 0.99$.

the center, or a total range of $\Delta\theta \approx 86.17^\circ$ (1.50 rad). This limited vertical range implies the ligament support as modeled in Figure 5 has merit. Using result (4) and the angles through which the neck moves, the approximate amount of musculature for the neck can be calculated.

Here, I take the shoulder girdles to be approximately 30° ($\pi/6$ rad) off the long axis of the animal as in Schwarz *et al.* (2007a). The measurement of the coefficient is based on *Diplodocus* reconstructions but the value of r_a should not vary much between animals. The muscle attachment is approximated as occurring over the first meter of the neck, with the posterior end attached at the scapula and shoulder girdles. The resulting geometry gives a radial distance r_a from the vertebrae, which can be used to calculate the torque exerted on the neck by the muscles. Using this value for a projection reduces the cross product into scalar form.

$$r_a \approx 1.240 \cos(\pi/6) + r_b \approx 1.221 \text{ m}$$

To complete a calculation of the muscle mass, it is necessary to find the maximum angular acceleration that the *Diplodocus* head and brain could withstand. If the blood pressure system is treated as similar to *Giraffa camelopardalis*, it can be feasibly postulated to be capable of handling up to double standard mammalian blood pressure levels (Hargans *et al.* 1987). Thus a reasonable solution should be found for a model that treats

the maximum acceleration at the head of the dinosaur to be approximately $a = 3g$ (Seymour and Lillywhite 2000). Using the fact that the torque $\tau = a|I|$,

$$\begin{aligned} \tau &= 3g|I| \\ &= F \int_0^1 r \sin\left(\arctan\left(\frac{r_a}{r}\right)\right) dr \\ &= F r_a r \sqrt{\frac{r_a^2}{r^2} + 1} \Bigg|_{r=1} \rightarrow \end{aligned} \quad (5)$$

$$\begin{aligned} F &= \frac{3g|I|}{r_a \sqrt{r_a^2 + 1}} \\ &\approx 1.06 \times 10^5 \text{ N} \end{aligned} \quad (6)$$

Taking this as the maximum force required to swing the base of the neck, the resulting arc can be extended over a 43.1° lateral rotation as by Parrish and Stevens (1999) resulting in a work output of:

$$(1.06 \times 10^5 \text{ N})(1 \text{ m})(0.752 \text{ rad}) = 7.97 \times 10^4 \text{ J}$$

Given that the mean power output of muscle is approximately $1 \text{ kW} \cdot \text{kg}^{-1}$, this then yields $67.9 \text{ kg} \cdot \text{s}^{-1}$ of muscle required to accelerate the neck for any length of time t (Alexander 2002). It is simple to show the maximal rotational velocity occurs in

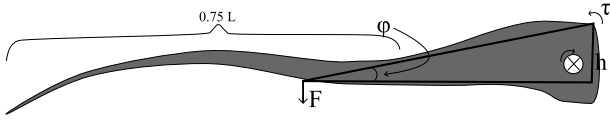


FIGURE 6. Schematic of force setup on the tail as used to begin the torque-balance equations. The setup is analogous for the neck, with the neutral point of torque chosen to be large neural spines above the acetabulum.

$t = \sqrt{2L\theta/a} \approx .607$ seconds, resulting in approximately 49.9 kg of muscle required to swing the neck in a given direction.

To complete the analysis, this base muscle-mass result must be multiplied by two to account for both sides of the neck. This calculation can be repeated for the top and bottom of the neck, substituting the appropriate angle terms. Substituting in a density of $\rho_m \approx 1050 \text{ kg}\cdot\text{m}^{-3}$, this yields $m_m \approx 1.47 \times 10^2$ kg (Alexander 2002).

Using a mean ligament density of $\langle \rho_l \rangle \approx 1 \text{ g}\cdot\text{cm}^{-3} = 1000 \text{ kg}\cdot\text{m}^{-3}$, and tracheal radius r_{tr} , the total neck tissue mass can be represented by

$$m_t \approx m_m - \pi r_{tr}^2 l + \int_0^l \frac{\pi^2}{8} r_t l \rho_l dl \quad (7)$$

Stretching the tissue across the neck in a partial cylindrical shell (as Figure 3),

$$m_m = \rho_t l \frac{7}{8} \pi (r_t(l) - r_b(l))^2 \rightarrow$$

$$r_t(l) = \sqrt{\frac{8m_m}{7\pi}} - r_b(l) \quad (8)$$

Which can be substituted into the final integral term in Equation 7 to calculate the ligamentary mass:

$$m_l = \frac{\pi^2 \rho_l}{8} \int_0^l l \sqrt{\frac{8m_m}{7\pi}} - l r_b(l) dl \quad (9)$$

$$\approx 2.22 \times 10^2 \text{ kg} \quad (10)$$

Finally, it is assumed that the trachea has a radius about equal to that of the C2 vertebra, or 0.027 m (Hatcher 1901), giving:

$$m_t \approx m_m - \pi r_{tr}^2 l + m_l$$

$$\approx 3.33 \times 10^2 \text{ kg}$$

For a total neck mass of $M = 5.85 \times 10^2 + 3.04 \times 10^2 = 9.77 \times 10^2$ kg for a *Diplodocus* neck — an upper bound of 8.1% of its total mass (Henderson 2003). This result neglects the mass of the head, which would have to weigh over 40 kg to result in a 5% correction.

Estimating Sauropod Tail Length

Estimation of *Diplodocus*' tail length can be accomplished by using a 'suspension bridge' design to the lumbar neural spines and by applying a similar mass analysis to the one just performed on the neck. Solving for a net zero torque on thoracic and lumbar neural spines can then yield a predicted tail length. This is an oversimplification of

ligamentary attachment along the vertebrae, but accurate to a first approximation, and such a force-balance approach has precedent, and prior work has shown elastic ligaments would be strong enough to support the neck (Alexander 1989, 2006; Schwarz *et al.* 2007b).

For the sake of simplicity, it is assumed that a single ligament connects the highest neural spine over the acetabulum to the middle of its neck, with that neural spine supporting the entire torque of the neck. The neck and tail is supported at its center of mass, $L/4$. This is then equated to the torque placed on the same neural spine by the tail; that is, $\tau_{neck} - \tau_{tail} = 0$. It is this equality that prompted the mass of the neck to be calculated previously.

The tail radius as a function of length is

$$r_T(l) = \frac{R_f - R_i}{L_T} l + R_i$$

Where R_i is the initial radius at the base of the tail, R_f the final radius at its extremum, L its total length, and l the position of an element along its length. For convenience, $\eta \equiv R_f - R_i$. In these calculations, the values are known from Hatcher (1901); however, even for incomplete specimens these values should not be difficult to extrapolate.

Rewriting Equations 1 and 3,

$$m_{bT} = \rho_b k_{pt} \pi \int_0^L r_T(l)^2 dl$$

$$I_T = \rho_b k_{pt} \pi \int_0^L r_T(l)^4 dl$$

Leading to

$$m_{bT} = \rho_b k_{pt} \pi \left(\frac{1}{2} \eta^2 + \eta R_i + R_i^2 \right) \quad (11)$$

$$I_T = \rho_b k_{pt} \pi \left(\frac{1}{4} \eta^4 + \eta^3 R_i + 2\eta^2 R_i^2 + 2\eta R_i^3 + R_i^4 \right) \quad (12)$$

Since current tail reconstructions suggest an only partially pneumatic tail (such as Wedel (2005)), here I assume that approximately 75% of the tail is fully pneumatic, and the rest of the tail is apneumatic, instead of the reducing pneumaticity toward the posterior end of the tail. This results in a mean pneumaticity of $k_{pt} \approx 0.53$. This results in a tail bone mass of $m_b \approx 2.97 \times 10^2$ kg and a moment of inertia of 2.87×10^2 N·m.

To proceed, the assumption is made that the extremum of the tail can break the speed of sound ($c_s = 343 \text{ m}\cdot\text{s}^{-1}$ at 293 K) as suggested by Mhyrvold and Currie (1997). Assuming the tail reaches 110% c_s ($v_{ss} = 377.3 \text{ m}\cdot\text{s}^{-1}$) over a quarter rotation of the available range,

$$\tau_w = I\alpha$$

$$= 2I_T v t^{-1}$$

$$= \frac{2I_T v^2}{L_T \sin \pi/8}$$

This can then be substituted into a force-torque equation, as in Equation 5. The joule-kilograms expended is $F\phi_{ss}$, where ϕ_{ss} is the angle through which the tail rotates in this time frame ($\pi/8$ rad, or 27.5°).

The product of the energy, elapsed time, and the mean power output of muscle $4Et\langle P_m \rangle$ results in

$$\begin{aligned} m_{mT} &= \left(\frac{4L \sin(\pi/8)}{10^3 v_{ss}} \right) \left(\frac{2|I_T|v_{ss}^2}{L_T \sin(\pi/8)} \right) \phi_{ss} \\ &\quad \left(R_i \sqrt{R_i^2 + 1} \right)^{-1} \\ &= \frac{|I_T|v_{ss}}{125R_i \sqrt{R_i^2 + 1}} \phi_{ss} \\ &\approx 9.03 \times 10^2 \text{ kg} \\ M_T &= m_{bT} + m_{mT} + m_l \frac{m_{bT} + m_{mT}}{M} \end{aligned} \quad (13)$$

The final term in the total mass is a correction for the ligaments holding the tail, where they are assumed to be scaled linearly with the mass ratio of the rest of the tail components to the neck. The forces and torques can be set up as illustrated in Figure 6. With the ligament modeled to be connected to the middle of the tail, the model results in $\tan \phi = 4h/L_p$, and $F_y = mg \sin \phi$. The hypotenuse and overall force are

$$\begin{aligned} r &= \sqrt{(L_d + L_p/4)^2 + h^2} \\ F_y &= Mg \sin \left(\arctan \left(\frac{4h}{L_p} \right) \right) \end{aligned}$$

Where L_d is the dorsal vertebra distance to the acetabulum, $h = \Delta h + h_{ns*}$ the vertical offset to the neutral position as shown in Figure 6. L_p is used to show that this can be used for either the neck or the tail, and the correct value can be input / calculated for as required by context. D_n as a variable will also represent the length of the n th dorsal vertebra. In context, $L_d = D_{11}$ for the tail, and the sum of dorsals 1-9 for the neck.

The torque of the neck can be explicitly solved with its calculated values of M and L , model value $\Delta h = 0$, the estimated or known height of the tallest neural spine h_{ns*} , and $L_d = \sum_{n=1}^9 D_n$. The resulting τ_n can then be equated with the same equation from the tail:

$$\begin{aligned} \tau_n &= 2.22 \times 10^4 \text{ N} \cdot \text{m} \\ &= M_T g \left((L_d + L_T/4)^2 + h_T^2 \right)^{1/2} \sin \left(\arctan \left(\frac{4h_T}{L_T} \right) \right) \end{aligned}$$

Squaring, taking a small angle approximation, and expanding,

$$(L_d^2 + L_T L_d + h_T^2) \left(\frac{4h_T}{L_T} \right)^2 + h_T^2 \approx \left(\frac{\tau_n}{M_T g} \right)^2$$

Reducing to the quadratic in L :

$$\left(h_T^2 - \left(\frac{\tau_n}{M_T g} \right)^2 \right) L_T^2 + 8h_T^2 L_d L_T + 16(L_d^2 h_T^2 + h_T^4) = 0 \quad (14)$$

Applying the quadratic formula, and using $L_d = D_{11} = 0.270$ m and approximating $\Delta h_T = 1.25$ m (Hatcher 1901), the physical solution to the equation is then a tail length of 16.7 m and mass of 1.47×10^3 kg. Summing this up with the other known lengths of the body parts, the model obtains a total animal length of 28.6 m, or within about 4.3% removed from the length of the current restoration at 27.4 m.

Summary

By finding a least-squares linear fit for the neck ($r(l)$), extrapolating to a minimum radius, the following combined equations from the Methods section can be solved in order to calculate the length and the masses of both the neck and tail of a diplodocid.

$$I = k_p \rho_b \pi \int_0^l l^2 r_b(l)^2 dl \quad (15)$$

$$m_m = \frac{|I|}{r_a} \sqrt{\frac{6gL}{r_a^2 + 1}} \left(\theta_{horiz}^{3/2} + \theta_{vert}^{3/2} \right) \quad (16)$$

$$\begin{aligned} M &= m_m - \pi r_{tr}^2 l + \\ &\quad \frac{\pi^2 \rho_l}{8} \int_0^l l \sqrt{\frac{8m_m}{7\pi}} - l r_b(l) dl \end{aligned} \quad (17)$$

$$\begin{aligned} \tau_n &= Mg \sin \left(\arctan \left(\frac{4h_{ns*}}{L} \right) \right) \\ &\quad \left(\sqrt{(L_d + L/4)^2 + h_{ns*}^2} \right) \end{aligned} \quad (18)$$

$$m_{bT} = \rho_b k_{pt} \pi \left(\frac{1}{2} \eta^2 + \eta R_i + R_i^2 \right) \quad (19)$$

$$\begin{aligned} I_T &= \rho_b k_{pt} \pi \left(\frac{1}{4} \eta^4 + \eta^3 R_i + 2\eta^2 R_i^2 + \right. \\ &\quad \left. 2\eta R_i^3 + R_i^4 \right) \end{aligned} \quad (20)$$

$$\begin{aligned} M_T &= \frac{\pi |I_T| v_{ss}}{10^3 R_i \sqrt{R_i^2 + 1}} + m_l \frac{m_{bT} + m_{mT}}{M} \\ &\quad + m_{bT} \end{aligned} \quad (21)$$

$$\begin{aligned} 0 &= \left(h_T^2 - \left(\frac{\tau_n}{M_T g} \right)^2 \right) L_T^2 - 8h_T^2 L_d L_T \\ &\quad - 16(L_d^2 h_T^2 + h_T^4) \end{aligned} \quad (22)$$

This model then requires in addition only the length of the dorsal series, scapular girdle, initial tail radius, and vertical tail offset.

DISCUSSION

Based on the results generated by the physical model, his method of modeling length can be generalized to various sauropods. The accuracy of the results can always be increased; for the mass of the skeleton, the neck can be better modeled by using a function that would give a better approximation to the shape of the cervical vertebrae, and the tapering of the neck can always be better approximated by a nonlinear expression. The tissue mass can be better approximated by breaking it into the sums of integrals over

particular muscle groups (instead of a single group over the first meter), and the ligaments can be better treated by having a separate set of expressions for them, and modeling where and how the ligaments connect in a more rigorous fashion. Practically, however, the results given are more than sufficient in accuracy. The most influential factors in the result are the pneumaticity factors and the precise fit used. Whereas any cervical vertebra series may be used here, the longest continuous series starting close to the base of the neck will generally return the best fits.

The tendency of this solution is to over-estimate the mass; insufficient modeling of the vertebrae gives them more volume than they actually had. This over-estimation of mass will tend to decrease animal size if biased toward the head, and increase animal size if biased toward the tail. Finally, calculating musculature based on a maximum head and tail velocity might be problematic, but was the most limiting case available.

Both the neck and the tail are taken to be θ independent, and the calculations only have neural spine data added at the end, which are treated separately. The bifurcated cervical spines are treated as a single element for simplicity. The k_p factor that appears is 1-ASP or a variation on the “Airspace Proportion” proposed by Wedel (2005). This treatment also assumes that the density of the tracheal space is negligible; that is, $\rho_{air} \approx 1.3 \text{ mg/cc} \rightarrow 0$.

The modeling of the tail could be improved by using more than an ‘average’ ligamentary connection, and providing proper connective chains of the ligaments rather than a rigid-model approximation. Additionally, the ligaments could be separately accounted for. However, this runs contrary to the goal of this paper in constructing a low-order polynomial solution that can be quickly done on a calculator. The k_p for the tail used is in line with approximate values that can be obtained from the Wedel’s (2005) analysis. There, Wedel calculates an approximate mass savings of 300 kg for the caudal series, which, for a 600 kg apneumatic tail, represents a tail $k_{pT} \approx 0.5$, in line with the ‘75%’ estimate used here.

The pneumaticity of the axial skeleton is interesting to note here. The combined mass of the tail and the neck — which comprises the majority of the length of the animal — comprises at most 22% the mass of the total body as calculated by Henderson (2003). Additionally, the fact that these measurements of pneumaticity are required to predict sizes of the animal provides an additional method by which one can obtain pneumaticity estimates. ‘Bogus’ estimates of pneumaticity result only in nonphysical solutions to the quadratic.

To ensure the result was not a coincidence, the analysis was performed on the incomplete neck of a “medium-sized” *Apatosaurus ajax* (NSMT-PV 20375) described by Upchurch *et al.* (2004b). Δh_T was obtained by adding the last dorsal and sacral vertebra’s radii, CM84’s neural spine mass was used, the tail and the fit obtained with C3—C9. k_p was based on cervical midcentrum measurements and k_{pe} used the ‘75%’ estimate with respect to the cervical condyle (Wedel 2005). This resulted in a very typical total length of 22.0 m, suggesting that the applicability to *Diplodocus* is unlikely to be a chance artifact.

CONCLUSION

The approximations and type of solutions done in this paper only apply to sauropods whose biomechanics can be described as similar to a suspension bridge, such as diplodocids. Currently, I propose no method by which the length of the dorsal series can be estimated with only fragmentary finds, and the requirement of this as the variable L_d makes this the strongest requirement of specimen completeness. In a broader context, the results of this study would suggest that because a zero-torque model predicts diplodocid length with high-precision, it is likely this model which was the primary enabling feature that allowed sauropods to reach their massive sizes.

Future work would include brachiosaurids, juvenile diplodocids, titanosaurids, and oddly-proportioned members of Sauropoda that should otherwise conform to this model. Many of these would require fairly significant changes to the model, and sauropods that are not based on a zero-torque model, such as *Brachiosaurus*, will fail to have reasonable predictions. Smaller animals such as juveniles would not obey the assumptions made about speed of the tail and neck. There are interesting candidates that would test the assertion made here that the massive head should not be significant (such as *Dicraeosaurus*) and the predictive abilities of this method on oddly-proportioned individuals such as *Mamenchisaurus*. Finally, titanosaurids are more robust and not traditionally thought of as ‘living suspension bridges’. However, these are diverse enough in methods and significant enough changes to warrant a separate treatment. The changes required are validated by a direct application of this theory to a juvenile diplodocid (SMA 0009) described in Schwarz *et al.* (2007c), in which constants assumed for adult animals yield nonphysical ligamentary masses.

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REFERENCES

- Alexander, R. M. 1989. Mechanics of fossil vertebrates: William Smith lecture. *Journal of the Geological Society*, London 146:1–52. doi:10.1144/gsjgs.146.1.0041.
- Alexander, R. M. 2002. Tendon elasticity and muscle function. *Comparative Biochemical Physiology, A* 133:1001–1011. doi:10.1016/S1095-6433(02)00143-5.
- Alexander, R. M. 2006. Dinosaur Biomechanics. *Proceedings of the Royal Society B* 273:1849–1855. doi: 10.1098/rspb.2006.3532.
- Dzemski, G., and A. Christian. 2007. Flexibility along the neck of the ostrich (*Struthio camelus*) and consequences

- for the reconstruction of dinosaurs with extreme neck length. *Journal of Morphology* 268:701–714. doi:10.1002/jmor.10542.
- Hargans, A. R., R. W. Millard, K. Pettersson, and K. Johansen. 1987. Gravitational haemodynamics and oedema prevention in the giraffe. *Nature* 329:59–60. doi:10.1038/329059a0.
- Hatcher, J. B. 1901. *Diplodocus: Its osteology, taxonomy, and probable habits, with restoration of its skeleton.* The Carnegie Institute, Washington, DC.
- Henderson, D. M. 2003. Tippy punters: Sauropod dinosaur pneumaticity, buoyancy, and aquatic habits. *Proceedings of the Royal Society B* 271:180–183. doi:10.1098/rsbl.2003.0136.
- Mhyrvold, N., and P. Currie. 1997. Supersonic sauropods? Tail dynamics in the diplodocids. *Paleobiology* 23:393–409.
- Organ, C. L. 2006. Thoracic epaxial muscles in living archosaurs and ornithomimid dinosaurs. *The Anatomical Record Part A: Discoveries in Molecular, Cellular, and Evolutionary Biology* 288:782–793. doi:10.1002/ar.a.20341.
- Parrish, J. M., and K. A. Stevens. 1999. Neck posture and feeding habits of two Jurassic sauropod dinosaurs. *Science* 284:98–800. doi:10.1126/science.284.5415.798.
- Schwarz, D., E. D. Frey, and C. A. Meyer. 2007a. Novel reconstruction of the orientation of the pectoral girdle in sauropods. *Anatomical Record B* 290:32–47. doi:10.1002/ar.20405.
- Schwarz, D., E. D. Frey, and C. A. Meyer. 2007b. Pneumaticity and soft-tissue reconstructions in the neck of diplodocid and dicraeosaurid sauropods. *Acta Palaeontologica Polonica* 52:167–188.
- Schwarz, D., and G. Fritsch. 2006. Pneumatic structures in the cervical vertebrae of the late Jurassic Tendaguru sauropods *Brachiosaurus branchai* and *Dicraeosaurus*. *Ecologiae Geologicae Helvetica* 99:65–78. doi:10.1007/s00015-006-1177-x.
- Schwarz, D., T. Ikejiri, B. H. Breithaupt, and P. M. Sander. 2007c. A nearly complete skeleton of an early juvenile diplodocid (Dinosauria: Sauropoda) from the Lower Morrison Formation (Late Jurassic) of north central Wyoming and its implications for early ontogeny and pneumaticity in sauropods. *Historical Biology* 19:225–253. doi:10.1080/08912960601118651.
- Seymour, R. S., and H. B. Lillywhite. 2000. Hearts, neck posture, and metabolic intensity of sauropod dinosaurs. *Proceedings of the Royal Society B* 267:1883–1887. doi:10.1098/rspb.2000.1225.
- Stevens, K. A. 2002. Dinomorph: Parametric modeling of skeletal structures. *Senckenbergiana Lethaea* 82:23–34.
- Stevens, K. A., and J. M. Parrish. 2005a. Digital Reconstructions of Sauropod Dinosaurs and Implications for Feeding; pp178–200 in K. C. Rogers (ed.) and J. Wilson (ed.), *The Sauropods: Evolution and Paleobiology.* University of California Press, Berkeley and Los Angeles.
- Stevens, K. A., and J. M. Parrish. 2005b. Neck posture, dentition, and feeding strategies in Jurassic sauropod dinosaurs; pp212–232 in V. Tidwell (ed.) and K. Carpenter (ed.), *Thunder-lizards: The Sauropodomorph Dinosaurs.* Indiana University Press, Bloomington.
- Upchurch, P., K. A. Stevens, J. M. Parrish. 2000. Technical comment: Neck posture of sauropod dinosaurs. *Science* 287:547. doi:10.1126/science.287.5453.547b.
- Upchurch, P., P. M. Barrett, and P. Dodson. 2004. Sauropoda; pp316–318 in D. B. Weishampel (ed.), P. Dodson (ed.), and H. Osmólska (ed.), *The Dinosauria*, 2nd edn. University of California Press, Berkeley and Los Angeles.
- Upchurch, P., Y. Tomida, and P. M. Barrett. 2004. A new specimen of *Apatosaurus ajax* from the Morrison formation (Upper Jurassic) of Wyoming, USA. *National Science Museum monographs* 26:pp. i–118.
- Wedel, M. J. 2003. Vertebral pneumaticity, air sacs, and the physiology of sauropod dinosaurs. *Paleobiology* 29:243–255. doi:10.1666/0094-8373(2003)029<0243:VPASAT>2.0.CO;2.
- Wedel, M. J. 2005. Postcranial skeletal pneumaticity in sauropods and its implications for mass estimates. pp201–228 in K. C. Rogers (ed.) and J. Wilson (ed.), *The Sauropods: Evolution and Paleobiology.* University of California Press, Berkeley and Los Angeles.
- Wedel, M. J. 2007. Postcranial pneumaticity in dinosaurs and the origin of the avian lung. Ph.D. dissertation, University of California, Berkeley, California, 304pp.
- Wedel, M. J., and R. K. Sanders. 2002. Osteological correlates of cervical musculature in aves and sauropoda (dinosauria: saurischia), with comments on the cervical ribs of *Apatosaurus*. *PaleoBios* 22:1–6.
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