Feeding height stratification among the herbivorous dinosaurs from the Dinosaur Park Formation (upper Campanian) of Alberta, Canada

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Abstract

**Background:** Herbivore coexistence on the Late Cretaceous island continent of Laramidia has been a topic of great interest, stemming from the paradoxically high diversity and biomass of these animals in relation to the relatively small landmass available to them. Various hypotheses have been advanced to account for these facts, of which niche partitioning is among the most frequently invoked. However, despite its wide acceptance, this hypothesis has not been rigorously tested. This study uses the fossil assemblage from the Dinosaur Park Formation of Alberta as a model to investigate whether niche partitioning facilitated herbivorous dinosaur coexistence on Laramidia. Specifically, the question of feeding height stratification is examined in light of the role it plays in facilitating modern ungulate coexistence.

**Results:** Most herbivorous dinosaur species from the Dinosaur Park Formation were restricted to feeding no higher than approximately 1 m above the ground. There is minimal evidence for feeding height partitioning at this level, with ceratopsids capable of feeding slightly higher than ankylosaurs, but the ecological significance of this is ambiguous. Hadrosaurids were uniquely capable of feeding up to 2 m quadrupedally, or up to 5 m bipedally. There is no evidence for either feeding height stratification within any of these clades, or for change in these ecological relationships through the approximately 1.5 Ma record of the Dinosaur Park Formation.

**Conclusions:** Although we cannot reject the possibility, we find no good evidence that feeding height stratification, as revealed by reconstructed maximum feeding heights, played an important role in facilitating niche partitioning among the herbivorous dinosaurs of Laramidia. Most browsing pressure was concentrated in the herb layer, although hadrosaurids were capable of reaching shrubs and low-growing trees that were out of reach from ceratopsids, ankylosaurs, and other small herbivores, effectively dividing the herbivores in terms of relative abundance. Sympatric hadrosaurids may have avoided competing with one another by feeding differentially using bipedal and quadrupedal postures. These ecological relationships evidently proved to be evolutionarily stable because they characterize the herbivore assemblage of the Dinosaur Park Formation through time. If niche partitioning served to facilitate the rich diversity of these animals, it may have been achieved by other means in addition to feeding height stratification. Consideration of other feeding height proxies, including dental microwear and skull morphology, may help to alleviate problems of underdetermination identified here.

**Keywords:** Dinosaur Park Formation, Feeding height stratification, Niche partitioning, Evolutionary palaeoecology, Ankylosauria, Ceratopsidae, Hadrosauridae

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Background

During the Late Cretaceous, a shallow inland sea divided North America into two longitudinally-arrayed landmasses. The eastern landmass—called Appalachia [1]—supported an enigmatic fauna known only from scant fossil remains [2]. The western landmass—called Laramidia [1]—supported a rich diversity of herbivorous dinosaurs. Included among these were such small forms (< 100 kg) as hypsilophodontids, pachycephalosaurids, and leptoceratopsids. Putatively facultative herbivores (or omnivores) included troodontids, oviraptorosaurs, ornithomimids, and therizinosaurids [3]. The megaherbivorous dinosaurs—primarily represented by ankylosaurs, ceratopsids, and hadrosaurids—form the most significant component of this fauna [4–6], in terms of both population (75–82% of total dinosaur fauna; [7]) and body size.

This high diversity of contemporaneous herbivores in Laramidia has long puzzled investigators. Laramidia had a total estimated area between just 4 million km² [8] and 7.7 million km² [5], and dinosaur distribution was segregated into distinct northern and southern faunal provinces that presumably reflect strict habitat preferences linked to a palaeoclimatic gradient [4–6,9]. By contrast, the area of sub-Saharan Africa, where living vertebrate herbivores are most diverse, is approximately 23.6 million km², with the highest diversity occurring in the East African savannah [10]. The diverse herbivorous dinosaur fauna of Laramidia therefore appears to have been spatially restricted compared to modern analogs, and this likely would have increased the potential for resource competition. The problem of the coexistence of these dinosaurs is further exaggerated when considering the presumably large nutritional requirements of the megaherbivorous forms [11–14], as well as their high population densities [15–22], both of which would have placed increased pressure on the resource base.

Two main hypotheses have been proposed to account for the coexistence of so many herbivores in Laramidia. The first hypothesis contends that plant resources were simply not limiting, a scenario that could be explained by numerous causal factors. For example, it may have been that, in spite of the fact that the nutritional requirements of the megaherbivorous dinosaurs were absolutely high, the inferred bradymetabolic thermoregulatory systems of these animals imparted relatively low nutritional requirements compared to their mammalian counterparts, minimizing pressure on the resource base [5,14,23]. Alternatively, plant resources may not have been limiting because primary productivity was elevated during the Late Cretaceous [23,24]. This suggestion was also advanced to account for the increased abundance of browsers during the Miocene [25,26]. Increased primary productivity may have resulted in part from the prevalence of marginal coastal environments during the Late Cretaceous [27]; however, palaeoclimatological modeling [28,29] and experimental evidence [30] also suggest that elevated atmospheric pressure and CO₂ during the Late Cretaceous would have yielded similar effects. A third option might be that predation pressure from tyrannosaurusids and dromaeosaurids was sufficiently high during the Late Cretaceous to suppress herbivore population densities, leading to reduced pressure on the resource base as in the first scenario. A similar mechanism is thought to help shape the structure of modern African ungulate communities [31].

The second hypothesis maintains that plant resources were limiting, and that herbivorous dinosaur coexistence was facilitated by dietary niche partitioning [6,13,32]. If so, these animals should have been adapted in such a way as to avoid competing with one another, perhaps varying in morphology so as to specialize on different plant types [7,8,33–42]. Numerous ecomorphological correlates among the herbivores have been hypothesized to reflect niche partitioning, including differences in snout shape [7,33,34,36–41,43,44], jaw mechanics and tooth shape [36,40,45,46], and feeding height stratification [12,13,33,34].

The late Campanian-aged Dinosaur Park Formation (DPF) of Alberta preserves the most diverse dinosaur assemblage currently known, including the most diverse assemblage of large-bodied herbivores from Laramidia [47]. Currently, approximately 33 herbivorous and omnivorous dinosaur species are recognized [48]. Biostratigraphic work shows that these taxa are distributed heterogeneously throughout the formation, such that different species are restricted to different horizons within the formation [47,49–51]. Although not all 33 taxa co-existed, over half likely co-existed in certain time intervals, including at least eight megaherbivorous taxa (Megaherbivore Assemblage Zone 1 [51]). Therefore, this assemblage provides a unique opportunity to evaluate possible dietary niche partitioning among Late Cretaceous herbivorous dinosaurs.

Niche partitioning via feeding height stratification is thought to have facilitated sauropod diversity during the Late Jurassic [33,34,44,52–58]. Likewise, some [12,13,33] have suggested that the rich diversity of herbivorous dinosaurs in the Late Cretaceous of Western Interior North America was fostered, in part, by a similar mechanism, but rigorous quantification of these patterns and formal tests of specific hypotheses are lacking. The aim of this paper is to test the hypothesis that the long-term coexistence of these animals was facilitated by dietary niche partitioning, more specifically via feeding height stratification.

Methods

We examined the question of feeding height stratification by estimating the maximum feeding heights (MFHs)
of the dinosaurs from the DPF. It is prudent to note from the outset that MFH does not necessarily reflect habitual feeding height. Habitual feeding height is very difficult to determine in fossil forms, particularly in light of the fact that amniotes rarely maintain the cervical column in an osteologically neutral pose [59]. Regardless, most discussions to date regarding the feeding habits of dinosaurs from the DPF have focused on MFH, which is our primary concern. See Discussion below for further treatment of maximal and habitual feeding heights.

The data used in this study are listed in Additional file 1. We excluded juvenile specimens, identified by their small size and/or under-developed cranial ornamentation, because we were specifically interested in estimating the MFHs of fully-grown individuals. Presumably, juveniles fed at intermediate heights and their inclusion in this study would have no effect on the calculation of MFHs. We also excluded the facultatively herbivorous/omnivorous thero- pods because they formed only a small component of the DPF fauna [7,12,60,61], and were therefore unlikely to have consumed an appreciable amount of plant matter. Many of these animals are also known only from fragmentary remains, and their MFHs can be estimated only with a great deal of uncertainty.

We studied all fossil specimens with permission from the following institutions: ACM, Beneski Museum of Natural History, Amherst College, Amherst; AMNH, American Museum of Natural History, New York; CMN, Canadian Museum of Nature, Ottawa; MCSNM, Museo Civico di Storia Naturale di Milano, Milan; NSM PV, National Museum of Nature and Science, Tokyo; ROM, Royal Ontario Museum, Toronto; TMP, Royal Tyrrell Museum of Palaeontology, Drumheller; UALVP, University of Alberta Laboratory of Vertebrate Palaeontology, Edmonton; YPM, Yale Peabody Museum, New Haven.

Maximum feeding height estimation
When estimating the MFHs of the herbivorous dinosaurs, it was important to devise some metric that allowed for the use of both articulated and disarticulated skeletons, thereby maximizing sample size. MFH can often be measured directly from mounted specimens, but many reasonably complete skeletons remain disarticulated in museum collections. For this reason, we calculated MFH as follows: for the quadrupedal forms, we estimated MFH from shoulder height, calculated by adding the lengths of the humerus, radius, and metacarpal III. Observations of mounted skeletons reveal that this is an appropriate proxy for MFH because the mouth is approximately level with the glenoid of the shoulder (Figure 1A, B). In the case of ceratopsids, the mouth could not be brought much higher than this because the extension of the head at the atlanto-occipital joint would eventually cause the huge parietosquamosal frill to abut the dorsal surface of the shoulders (Figure 1B). For hadrosaurids, quadrupedal MFH was estimated from hip height, calculated by summing the lengths of the femur, tibia/fibula, and metatarsal III (the fibula was used if the tibia was not available). Again, observations of mounted hadrosaurid skeletons confirm that this is an appropriate proxy for quadrupedal MFH (Figure 1C).

For bipedal forms (including the facultatively bipedal hadrosaurids and leptoceratopsid), bipedal MFH was calculated using trigonometry: assuming that the animal in question could rear up on its hind legs only so far as the distal tip of its tail would touch the ground (a reasonable assumption, given that the tails of many of these animals were relatively inflexible due the presence of either tendon trellises or ossified myorhabdoi [62] that extended much of their lengths), a right triangle is formed (Figure 2) with the vertical side equal to hip height and...
the hypotenuse equal to tail length. The intersection of the hypotenuse and the horizontal side of the triangle forms angle \( \theta \), which can be calculated as \( \theta = \sin^{-1}(\text{hip height}/\text{tail length}) \). The triangle can then be extended by adding the combined length of the trunk and neck to the hypotenuse to give the full body length. The product of \( \sin \theta \) and body length yields an estimate for bipedal MFH.

We took measurements from the literature when they could not be made directly. Examples of complete, articulated vertebral columns from the DPF are nonexistent. For this reason, we assumed that mounts that included restored vertebrae were reasonably accurate and were included in this study; however, entirely missing or restored sections of the vertebral column (e.g., cervicals, thoracics, caudals) or limb elements were not used. We estimated these missing elements from more complete specimens using reduced major axis regression (RMA; Additional file 2). RMA is preferable to ordinary least squares (OLS) regression because it does not assume that the independent variable is measured without error. There is some debate, however, about whether RMA is preferable to OLS when extrapolating beyond the dataset [63], as we occasionally did here. For species lacking postcranial material, we modeled MFHs after closely related taxa of similar size (e.g., *Unescoceratops koppelhusae* was modeled after *Leptoceratops gracilis* [CMN 8889], “*Stegoceras* ("Prenocephale") breve and the unnamed pachycephalosaur after *S. validum* [UALVP 2], and cf. *Orodromeus* after *Parkosaurus warreni* [ROM 802]).

The various sources of error associated with the reconstruction of MFHs must be noted. For instance, habitual limb flexion in any of the taxa considered here would result in overestimated MFHs. Ceratopsids, in particular, are generally thought to have held their forelimbs in a semi-sprawled posture [64-68], which would result in smaller MFHs. Conversely, our inability to account for the influence of epiphyseal cartilage on feeding height calculations would result in underestimated MFHs. Holliday et al. [69] recently showed that, among archosaurs, epiphyseal cartilage significantly increases limb length relative to that calculated from the bones alone, the results differing by up to 10%. By accounting for missing cartilage in multi-tonned dinosaurs using various “cartilage correction factors” (CCFs), Holliday et al. [69] found that the heights of these animals may be consistently underestimated by up to 0.5 m. Unfortunately, it is not possible to determine exactly how much epiphyseal cartilage is missing from every individual, and to apply different CCFs to separate taxa in the reconstruction of maximum feeding heights would tell us more about the influence of CCFs in producing (or reducing) statistical differences, and less about how the fossil bones themselves differ. On the other hand, it is unlikely that the same CCF should apply across all taxa. In the best case scenario, the confounding effects of limb flexion and epiphyseal cartilage on the calculation of maximum feeding heights would cancel each other out. Overall, the error in reconstructing MFH is likely on the decimetre scale. Although this may affect the results in some cases (e.g., discerning between low-level browsers), we believe that this level of error should permit the major patterns to be discerned.

### Statistical comparisons

We compared MFHs at coarse (family/suborder), medium (subfamily/family), and fine (genus) taxonomic scales. We did not consider the species level because sample size was generally too low at this resolution to permit meaningful statistical comparisons. We used the non-parametric Kruskal-Wallis test because samples were typically quite small (\( n < 30 \)) and non-normal. This test lacks the power of parametric tests, and is more prone to committing Type II errors (reporting false negatives), but is more robust to committing Type I errors (reporting false positives). We conducted post hoc pairwise comparisons using Mann-Whitney U tests with Bonferroni correction. Bonferroni correction was designed to counteract the problem of multiple comparisons, wherein the probability of committing a Type I error increases with the number of simultaneous comparisons being made [70]. This problem is rectified by multiplying the \( p \)-value by the number of pairwise comparisons, effectively lowering the significance level. However, because Bonferroni correction provides little power and is probably too conservative [70,71], we also report uncorrected probabilities for the purpose of interpretation. We performed all statistical procedures using the software program PAST 2.12 [72].

Because the DPF does not represent a single assemblage of contemporaneous organisms, time-averaging is
an issue. This has the effect of masking palaeoecological patterns that are otherwise distinguishable only at fine temporal resolutions [73]. For this reason, we minimized the effects of time-averaging by making the above comparisons within each of the two most inclusive Megaherbivore Assemblage Zones (MAZs) identified by Mallon et al. [51]. To summarize, MAZ-1 encompasses the lower 28 m of the DPF, and MAZ-2 encompasses intervals from 29–52 m. Although this time-constrained approach theoretically increases the probability of recovering differences that would otherwise be masked by the effects of time-averaging, there is a trade-off in that sample size (and hence statistical power) is reduced considerably. Also, this approach does not completely remove the effects of time-averaging because the MAZs are themselves time-averaged over a period of approximately 600 Ka [51]. Although we took steps to minimize the amount of time-averaging in this study, the remaining time bias is likely too large to effectively capture true, interacting palaeocommunities. It is possible to reduce the time bias further by dividing the MAZs into approximately 300 Ka sub-zones [51], more closely approximating true palaeocommunities, but the sample size (and resulting statistical power) per sub-zone becomes drastically reduced in doing so.

**Results**

**Time-averaged approach**

The MFHs for the time-averaged analysis are depicted in Figure 3A. The most striking aspect of this diagram is that, irrespective of body mass, most species were restricted to feeding below approximately 1 m from the ground. The leptoceratopsid *Unescoceratops* appears to have been particularly limited in its feeding range because it was unable to reach above 0.5 m while in a quadrupedal posture. However, this animal probably could have reared bipedally to reach somewhat beyond 1 m in a manner similar to other coexisting, small ornithischians (cf. *Orodromeus* and pachycephalosaurids).

With respect to the megaherbivorous forms, the average quadrupedal MFHs of ankylosaurs generally fall slightly below 1 m, whereas those of ceratopsids plot slightly above 1 m. Hadrosaurids clearly have the most distinct MFHs, which consistently plot above 2 m in a quadrupedal posture, or 4 m in a bipedal posture. Hadrosaurines (*Gryposaurus* and *Prosaurolophus*) have slightly taller average bipedal MFHs than lambeosaurines, but the uncertainty intervals for all hadrosaurids in this posture are quite large and overlap substantially. This may reflect variation in reconstructed portions of the spinal column and estimated missing elements.

The Kruskal-Wallis test reveals highly significant differences among all higher-level taxa with adequate representation (N = 62, H = 55.82, p < 0.001); Ankylosauria, Ceratopsidae, and Hadrosauridae each differ significantly from one another in quadrupedal MFH (Table 1). As expected, the differences are exacerbated when bipedal hadrosaurids are considered, which differ significantly from all quadrupedal postures. No further significant differences are recovered with increasingly finer taxonomic resolution, even between the apparently different bipedal hadrosaurines and lambeosaurines (Tables 2–3).
Time-constrained approach
The results of the time-constrained analyses (MAZ-1 and –2) largely mirror those of the time-averaged analysis; the same MFH relationships are maintained (Figure 3B, C). Note that some taxa, while known from the DPF, are not represented in either MAZ-1 or –2 due either to a lack of appropriate material or imprecise provenance data (e.g., ankylosaurs in MAZ-2, Chasmosaurus in MAZ-1). Probabilities are reduced in the time-constrained analyses, often below significant levels, due to correspondingly reduced sample sizes (Tables 4, 5, 6, 7, 8 and 9).

Discussion
Feeding height stratification in extant ecosystems
Ecological separation of coexisting species is achieved along the axes of food, time, and space [74,75]. For example, the African savannah biome supports 31 species of large (> 5 kg), herbivorous mammals [76], and competition among its members is typically alleviated via the selection of different food types, the occupation of the same habitat at different times, or the occupation of different habitats at the same time [77-79]. The ecological separation of these ungulates may also be achieved along a vertical gradient, with different species feeding at different heights within the canopy or grass cover [77-83]. For example, Bell [78] demonstrated that, among grazers, zebra (Equus burchelli) tend to ingest the tallest, most fibrous portions of the herb layer, wildebeest (Connochaetes taurinus) and topi (Damaliscus korrigum) select the more nutritious middle layer, and Thomson’s gazelle (Gazella thomsoni) take in fruits from the ground. With the exception of the topi, this grazing succession is reflected by the decreasing body size of the animals (but see [83]). Similarly, feeding height stratification has been said to operate among browsing ungulates [77,79,81]. This hypothesis was tested explicitly on a subset of African browsers by du Toit [82], who found that giraffe (Giraffa camelopardalis), kudu (Tragelaphus strepsiceros), impala (Aepyceros melampus), and steenbok (Raphicerus campesteris) are stratified by mean feeding-height. On average, giraffe feed at heights between 2 and 3 m, kudu at heights near 1 m, and impala and steenbok at heights below 0.5 m. Although there was significant overlap between the browsing heights of some of these species, du Toit [82] observed that feeding height stratification is more pronounced during the dry season, when the use of woody browse is increased to compensate for the reduced availability of green forage in the herb layer. The long neck of the giraffe has been cited as an example of an adaptation to escape competition occurring at lower browsing heights [84].

Feeding height stratification in the Dinosaur Park Formation assemblage
As documented in extant taxa, numerous studies have commented on the importance of feeding height stratification as a mechanism for niche partitioning among herbivorous dinosaurs. For example, within the Upper Jurassic Morrison Formation of the western United States, up to five sauropod genera are thought to have lived in sympathy [85]. Feeding height stratification has been repeatedly invoked as a means of facilitating their coexistence, with different sauropods using their long necks to feed at different heights within the environment. Evidence for this interpretation includes reconstructed neck morphology [33,34,56,57], tooth wear analysis [44,53-55,58,86], and jaw mechanics [52,54].

Feeding height stratification has likewise been invoked to account for the diverse herbivorous dinosaur fauna of the DPF [12,13,33], but no formal test of this hypothesis has been conducted to date. Notably, most treatments of

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### Table 1 Coarse-scale Mann-Whitney pairwise comparisons of the time-averaged analysis

|                  | Ankylosauria | Ceratopsidae | Hadrosauridae (Q) | Hadrosauridae (B) |
|------------------|--------------|--------------|-------------------|-------------------|
| Ankylosauria     | 0.00         | 0.00         | 0.00              | 0.00              |
| Ceratopsidae     | 0.00         | 0.00         | 0.00              | 0.00              |
| Hadrosauridae (Q)| 0.00         | 0.00         | 0.00              | 0.00              |
| Hadrosauridae (B)| 0.00         | 0.00         | 0.00              | 0.00              |

N = 62, H = 55.82, p = 4.554 × 10^-12.
Bonferroni corrected p-values shown in lower left triangle; uncorrected pairwise comparison p-values shown in upper right triangle. Significant results reported in bold. Abbreviations: Q, quadrupedal posture; B, bipedal posture.

### Table 2 Medium-scale Mann-Whitney pairwise comparisons of the time-averaged analysis

|      | An | No | Ce | Ch | Ha (Q) | Ha (B) | La (Q) | La (B) |
|------|----|----|----|----|--------|--------|--------|--------|
| An   | 0.51| 0.00| 0.01| 0.00| 0.00   | 0.00   | 0.00   | 0.00   |
| No   | 0.11| 0.19| 0.01| 0.01| 0.00   | 0.00   | 0.00   | 0.00   |
| Ce   | 0.04| 1   | 0.67| 0.00| 0.00   | 0.00   | 0.00   | 0.00   |
| Ch   | 0.30| 0.00| 0.01| 0.01| 0.00   | 0.00   | 0.00   | 0.00   |
| Ha (Q)| 0.10| 0.40| 0.40| 0.40| 0.00   | 0.36   | 0.00   | 0.00   |
| Ha (B)| 0.06| 0.30| 0.30| 0.10| 0.00   | 0.30   | 0.00   | 0.00   |
| La (Q)| 0.01| 0.11| 0.01| 0.11| 0.01   | 0.01   | 0.00   | 0.00   |
| La (B)| 0.01| 0.11| 0.01| 0.11| 0.02   | 0.01   | 1      | 0.00   |

N = 62, H = 56.17, p = 8.721 × 10^-10.
Bonferroni corrected p-values shown in lower left triangle; uncorrected pairwise comparison p-values shown in upper right triangle. Significant results reported in bold. Taxonomic abbreviations: An, Ankylosauridae; Ce, Centrosaurinae; Ch, Chasmosaurinae; Ha, Hadrosauridae; La, Lambeosaurinae; No, Nodosauridae. Other abbreviations as in Table 1.
feeding height stratification make little or no mention of the small ornithischians that inhabited the Late Cretaceous landscape of Laramidia. This is probably because these forms are both poorly known and relatively rare in the fossil record. However, Brown et al. [48] recently demonstrated a substantial taphonomic size bias in the record of the DPF, and it is therefore likely that small ornithischians comprised a larger proportion of the herbivore fauna than previously assumed. The MFHs of these animals were probably restricted to less than approximately 1 m, which may have placed them in competition with the megaherbivores discussed below.

**Ankylosaurs**

Beland and Russell [12] and Coe et al. [13] suggested that ankylosaurs from the DPF exhibited different MFHs, with *Euoplocephalus* feeding on herbs below 0.5 m and *Panoplosaurus* feeding on woody vegetation up to 1 m. However, the current data do not support this hypothesis. The mean MFH of *Euoplocephalus* is closer to 0.8 m, whereas that of *Panoplosaurus* approximates 0.9 m. There is also considerable overlap of MFHs between these taxa, with some specimens of *Euoplocephalus* reaching as high as 0.88 m, and some specimens of *Panoplosaurus* only reaching 0.77 m. Apart from the reduced ankylosaur sample (see below), this moderate overlap of MFHs likely accounts for the fact that *Euoplocephalus* and *Panoplosaurus* do not differ significantly from one another. The contention of Weishampel and Norman [87] that ankylosaurs were generally restricted to browsing between 1 and 2 m is not supported here. By extending the forelimb proportions of *Euoplocephalus* (AMNH 5403) to the 542 mm-long humerus of *Ankylosaurus* (AMNH 5214), the largest known ankylosaur, the total forelimb length is estimated to be only 1.1 m.

**Ceratopsids**

Several authors [12,13,87,88] have likewise suggested that ceratopsids were capable of reaching heights of 2 m; however, no evidence was provided for this value.

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### Table 3: Fine-scale Mann–Whitney pairwise comparisons of the time-averaged analysis

|      | Ankylosauria | Ceratopsidae | Hadrosauridae (Q) | Hadrosauridae (B) |
|------|--------------|--------------|-------------------|-------------------|
| Ankylosauria | 0.59         | 0.01         | 0.07              | 0.03              |
| Ceratopsidae | 0.03         | 0.03         | 0.38              | 0.03              |
| Hadrosauridae (Q) | 0.01         | 0.01         | 0.03              | 0.03              |
| Hadrosauridae (B) | 0.01         | 0.01         | 0.03              | 0.01              |

### Table 4: Coarse-scale Mann–Whitney pairwise comparisons of the MAZ-1 analysis

|      | Ankylosauria | Ceratopsidae | Hadrosauridae (Q) | Hadrosauridae (B) |
|------|--------------|--------------|-------------------|-------------------|
| Ankylosauria | 0.24         | 0.00         | 0.00              | 0.00              |
| Ceratopsidae | 0.00         | 0.00         | 0.00              | 0.00              |
| Hadrosauridae (Q) | 0.00         | 0.02         | 0.00              | 0.00              |
| Hadrosauridae (B) | 0.00         | 0.02         | 0.00              | 0.00              |

N = 59, H = 53.52, p = 1.553 × 10⁻⁶. Bonferroni corrected p-values shown in lower left triangle; uncorrected pairwise comparison p-values shown in upper right triangle. Significant results reported in bold. Taxonomic abbreviations: Cen, *Centrosaurus*; Cha, *Chasmosaurus*; Cor, *Corythosaurus*; Euo, *Euoplocephalus*; Gry, *Gryposaurus*; Lam, *Lambeosaurus*; Pan, *Panoplosaurus*; Par, *Parasaurolophus*; Pro, *Prosaurolophus*; Sty, *Styracosaurus*. Other abbreviations as in Table 1.
Instead, our results indicate that ceratopsids more likely browsed no higher than approximately 1 m, as suggested by Dodson [7]. One well-preserved specimen of Triceratops (NSM PV 20379), among the largest ceratopsids that ever existed, could not reach above 1.2 m, either [68]. Hadrosaurids

The common claim that hadrosaurids could reach heights up to 4 m [7,12,13,33,87,88] is supported by our results. The largest hadrosaurids from the DPF, Gryposaurus and Prosaurolophus, probably could reach heights approaching 5 m in a bipedal posture (Figure 3). Despite these maxima, some authors [87,89] have proposed that hadrosaurid feeding was probably concentrated below 2 m, which would accord with the quadrupedal feeding postures calculated here.

Both vertebrate microfossil and skeletal remains suggest that hadrosaurids formed approximately 40% of the herbivorous dinosaur assemblage in the DPF ecosystem [7,61], which likely translates to a greater proportion of the herbivore biomass because hadrosaurids are the largest members of the fauna. The remainder of herbivores, notably the large ceratopsids and ankylosaurs which combined form approximately an equivalent proportion in terms of relative abundance, fed at or below the 1 m mark. Although feeding heights could not be discriminated below 1 m, the fact that hadrosaurids could reach up to 5 m and were therefore segregated from all other herbivores is likely fundamental in partitioning the resource base. Importantly, hadrosaurids were capable of reaching shrubs and low-growing trees that were beyond the reach of ceratopsids, ankylosaurs, and other small herbivores, effectively dividing the herbivores in terms of relative abundance. This may also have allowed hadrosaurids to escape resource stresses imparted by low browsers, and may have facilitated the co-existence of large herds of ceratopsids and highly abundant hadrosaurids [20,90] in DPF palaeoecosystems.

Dinosaur browsing and vegetation structure

The prevailing climate of the DPF has been described as warm temperate, as revealed by tree-, tooth-, and bone-growth ring data [91,92], sedimentological data [93], and biogeographic data [94,95]. Regional leaf physiognomic data have also reinforced this interpretation [96]. This climate is, in part, thought to have given rise to both open and closed habitats [96-99] akin to those of modern ecosystems [36]. Braman and Koppelhus ([100]:124) describe the landscape of the DPF as having been "wet everywhere, at least for portions of the year", with dense vegetation lining the rivers, and more open habitats occurring further distally.

The open habitats of the DPF and surrounding regions were likely dominated by ferns and low-growing angiosperms. Coe et al. ([13]: 235) even proposed the existence of extensive "fern prairies", analogous to modern grasslands, but Tiffney [101] stressed that evidence for such fern-dominated communities is lacking. Nonetheless, Wing et al. [102] subsequently reported one exceptional fossil flora from the mid-Maastrichtian in which

Table 5 Medium-scale Mann–Whitney pairwise comparisons of the MAZ-1 analysis

|     | An  | No  | Ce  | Ha (Q) | Ha (B) | La (Q) | La (B) |
|-----|-----|-----|-----|--------|--------|--------|--------|
| An  | 0.82| 0.11| 0.15| 0.15   | 0.04   | 0.04   |
| No  | 1   | 0.67| 0.05| 0.05   | 0.01   | 0.01   |
| Ce  | 1   | 1   | 0.05| 0.05   | 0.01   | 0.01   |
| Ha (Q)| 1 | 1 | 1 | 0.08 | 0.55 | 0.01 |
| Ha (B)| 1 | 1 | 1 | 1 | 0.01 | 0.80 |
| La (Q)| 0.87 | 0.12 | 0.12 | 0.30 | 1 | 0.00 |
| La (B)| 0.87 | 0.12 | 0.12 | 0.30 | 1 | 0.00 |

N = 36, H = 31.25, p = 2.269 × 10⁻⁵. Bonferroni corrected p-values shown in lower left triangle; uncorrected pairwise comparison p-values shown in upper right triangle. Significant results reported in bold. Abbreviations as in Tables 1 and 2.

Table 6 Finer-scale Mann–Whitney pairwise comparisons of the MAZ-1 analysis

|     | Pan | Cen | Gry (Q) | Gry (B) | Cor (Q) | Cor (B) | Lam (Q) | Lam (B) | Par (Q) | Par (B) |
|-----|-----|-----|---------|---------|---------|---------|---------|---------|---------|---------|
| Pan | 0.67| 0.05| 0.05    | 0.01    | 0.01    | 0.11    | 0.11    | 0.11    | 0.11    |
| Cen | 1 | 0.05 | 0.05 | 0.01 | 0.01 | 0.11 | 0.11 | 0.11 | 0.11 |
| Gry (Q)| 1 | 1 | 0.08 | 0.90 | 0.03 | 0.39 | 0.15 | 0.77 | 0.15 |
| Gry (B)| 1 | 1 | 1 | 0.03 | 0.90 | 0.15 | 0.77 | 0.15 | 0.77 |
| Cor (Q)| 0.64 | 0.64 | 1 | 1 | 0.01 | 0.01 | 0.11 | 0.11 | 0.11 |
| Cor (B)| 0.64 | 0.64 | 1 | 1 | 0.03 | 0.87 | 0.07 | 0.07 | 0.40 |
| Lam (Q)| 1 | 1 | 1 | 1 | 1 | 0.23 | 0.07 | 0.25 | 0.25 |
| Lam (B)| 1 | 1 | 1 | 1 | 1 | 1 | 0.25 | 0.70 | 0.25 |
| Par (Q)| 1 | 1 | 1 | 1 | 1 | 1 | 0.25 | 0.25 | 0.25 |
| Par (B)| 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

N = 34, H = 29.26, p = 5.86 × 10⁻⁴. Bonferroni corrected p-values shown in lower left triangle; uncorrected pairwise comparison p-values shown in upper right triangle. Significant results reported in bold. Abbreviations as in Tables 1 and 3.

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ferns and other “pteridophytes” account for nearly 50% of the total ground cover. By comparison, these same plants account for approximately 40% of the total palynomorph abundance in the DPF [103].

The Late Cretaceous saw the radiation of the angiosperms, which typically took the form of “weedy” herbs and shrubs growing in open or marginal habitats [12,33,97,98,101,104,105]. Angiosperm trees, although inferred to have existed elsewhere [96], probably did not occur in the DPF, as evidenced by the lack of diagnostic fossil wood [100]. It is commonly argued ([98]:125 [104,106]) that angiosperms occurred most regularly in coastal and fluvial depositional settings, occupying “stream-side and aquatic habitats, the forest understory and early successional thickets”; however, Wheeler and Lehman [107] noted the existence of angiosperm-dominated communities in southern upland environments as well, where conifers were otherwise thought to have dominated [98,104]. By virtue of their r-selected life history strategies, it is likely that angiosperms were capable of growing in a wide variety of habitats [101].

A common theme in the literature is the persistence of open-habitat cycadophytes (bennettitaleans and cycads) as forage for Late Cretaceous herbivores [34,97,108-113]. However, bennettitaleans went extinct by the Santonian [114], and cycads were probably absent from the DPF ([65,115,116], D. R. Braman, pers. comm., 2012), having been replaced by angiosperms [98,117]. It is by no means clear that cycadophytes were common enough elsewhere during the Late Cretaceous to support dense megarhivore populations, but their seed coats may have served to supplement dinosaur diets [97,109-111].

Unlike tropical forests, temperate forests typically exhibit limited stratification [118], and there is little reason to suspect that the temperate forests of the DPF were any different. Wolfe and Upchurch [96] proposed that such forests were, in fact, relatively sparse, with sunlight often penetrating fully through to the ground. Palynofloral and macroplant evidence from the DPF suggests that the forest canopy was formed primarily by taxodiaceous, cupressaceous, and podocarpaceous conifers [100,119,120], a composition typical of most Late Cretaceous warm temperate forests [117]. Angiosperm shrubs may [98] or may not [121] have formed an understory, alongside tree ferns and gymnosperm saplings [100]. The herb layer would have included ferns, lycopods, angiosperm herbs, and gymnosperm saplings, and ground cover comprised mosses, lichens, fungi, hornworts, and decaying vegetable matter [100].

Opinions vary about the degree to which habitat structure influenced the regional distribution of the Late Cretaceous herbivorous dinosaurs. Some [97,101,112] argued that the megaherbivorous forms were likely restricted to feeding in open habitats, partly as a result of their large sizes. However, it appears that the forests of the Late Cretaceous were not particularly dense [96], and probably did not inhibit the movement of even the more massive herbivores [13,96]. Alternatively, Baszio [122] suggested that, within the DPF, ankylosaurs and ceratopsids occupied open habitats, whereas hadrosaurids lived in forested environments. His reasoning was that ankylosaurs and ceratopsids, being limited in their range of vertical movements, could not have taken full advantage of stratified forest vegetation in the same way that...
hadrosaurids presumably could. However, it is unlikely that hadrosaurids could have accessed the entire forest structure; the canopy was almost certainly out of reach, particularly if Late Cretaceous taxodiaceous and cupressaceous conifers grew as tall as their modern descendants (> 90 m). Hadrosaurids likely could forage among the shrubs of the forest understory, but shrubs were abundant in more open habitats as well [12,33,97,98,101,105]. In that case, there is little reason to suspect that hadrosaurids could not have occupied both open and closed habitats [36], alongside ankylosaurs and ceratopsids.

This does not contradict the idea that certain groups may have preferred certain environments over others. Various lines of sedimentological evidence have been brought to bear on the matter [7,12,60,61,91,123,124]. There simply does not appear to have been any major structural obstacles to impede the movement of these animals. Consider that elephants, which are comparable in size to the megaherbivorous dinosaurs considered here, regularly occupy even dense forests and thickets in search of food [125]. In fact, their movements and feeding habits typically result in the creation of new, more navigable habitats [126,127].

Regardless of where hadrosaurids spent most of their time, it is likely that they usually foraged quadrupedally on abundant, low-lying herbage [87,89,128], occasionally rearing up onto their hindlimbs to feed among the angiosperm shrubs. Additional evidence for bipedal feeding in these animals comes from the Campanian aged Blackhawk Formation of Utah, where hundreds of dinosaur footprints are preserved in association with taxodiaceous conifer and palm roots and fallen logs [129]. In many places, pes prints attributed to hadrosaurids are found straddling the roots. The fact that manus prints are not also found in these areas suggests that these animals were rearing up to feed on the high foliage. This bipedal feeding behaviour would have been particularly beneficial in instances where large herds of low-browsing ceratopsids were passing through the same area [17,20-22]. Dietary niche partitioning could have been achieved among hadrosaurids if they utilized different levels within the shrub layer, as do living ruminants [82]. This may also have served to limit niche overlap between different ontogenetic stages of the same species [130]. The larger feeding heights of the hadrosaurids suggest that these animals were able to reach a wider variety of plant types than other sympatric herbivores. Circumstantial evidence for diet in these animals comes from multiple examples of fossil gut contents [131-133], which preserve conifer and angiosperm browse, including twigs and stems, bark, seeds, leaves, and fruit. Probable hadrosaurid coprolites [134,135] also contain abundant fungally-degraded conifer wood, which would presumably indicate that hadrosaurids fed at ground level at least occasionally. However, in light of the problems associated with the attribution of some of these fossils [32,132,136,137], their interpretation as dietary residues must be regarded with due caution.

Ankylosaurs, ceratopsids, and small ornithischians may have partitioned the herb layer by feeding height, as do the ungulates of the Serengeti today [78]. Ceratopsids, being slightly taller, may have even facilitated the existence of the other forms by cropping the herb layer to expose new growth. Of course, this is a highly speculative scenario requiring further investigation. Unfortunately, no ceratopsid gut contents are known by which to gauge these ideas, but an ankylosaurid cololite from Australia is reported to contain fibrous tissue (probably leaves), angiosperm fruits or endocarps, small seeds, and possible fern sporangia [138]. Ankylosaurs, it would seem, consumed less woody browse than hadrosaurids, which is in line with the interpretation given here.

One final aspect of herbivorous dinosaur ecology bears consideration. Elephants are known to regularly fell trees up to 10 m tall to feed on the otherwise unreachable browse, effectively increasing the feeding envelope of these animals up to three times [13,125]. It is possible that the megaherbivores of the DPF were capable of the same behaviour [13,65]. If so, tree felling may have served to increase dietary overlap between these animals, with the squat ankylosaurs and ceratopsids consuming foliage otherwise in reach of hadrosaurids alone. Unfortunately, while tree felling behaviour among dinosaurs is plausible, there is not yet any evidence supporting this speculation. Similarly, scenarios involving hypsilophodontids and pachycephalosaurids climbing trees to increase their feeding heights [122], while not impossible, are implausible owing to a lack of appropriate skeletal adaptation [139,140]. For these reasons, such highly speculative behaviours are not considered further.

**Evolutionary palaeoecological implications**

There is no convincing evidence that feeding height stratification, as revealed by reconstructed MFH, played as significant a role in facilitating herbivorous dinosaur niche partitioning in the DPF as previously assumed [12,13]. Despite the 18 genera considered here—six or more of which typically coexisted at a time [51]—only four statistically distinct MFHs are detected. If niche partitioning did allow herbivorous dinosaurs from the DPF to coexist, it may have been achieved by other means in addition to feeding height stratification. Although this hypothesis has yet to be subjected to rigorous testing, multiple morphological features have been proposed to have fostered the coexistence of these herbivores. For example, Carpenter [38-41] suggested that differences between the tooth and beak shapes of ankylosaurs and nodosaurids may have allowed these taxa to specialize on different plant types. Similarly,
differences between centrosaurines and chaostosaurines in cranial [37], mandibular [45], and beak [8,43] morphology have been cited as evidence for dietary niche partitioning. Finally, variations in beak shape [7,34,36,44], tooth morphology [36,46], and skeletal proportions [36] are thought to have enabled hadrosaurines and lambeosaurines to forage differentially. Many of these assumptions have not been tested and require further examination, particularly in light of questions regarding the significance of intraspecific variation and the influence of time-averaging.

The disappearance of ankylosaurs from the upper intervals of MAZ-2 of the DPF [51] suggests the possibility that some change in their habitat structure caused their displacement. Although it is by no means obvious whether such a change did occur, the gradual transgression of the Western Interior Seaway over the approximately 1.5 Ma span of the DPF undoubtedly would have had some influence on the palaeoflora. It may be that some of the herbaceous plants preferred by the ankylosaurs disappeared, but this scenario is difficult to test at present.

Overall, the distribution of herbivore MFHs changed minimally over the course of the DPF. Rather, MFHs were quite stable in spite of rapid and continual species turnover, and roughly the same ratio of low to high browsers was upheld (Figure 3B,C). This, in turn, suggests that time-averaging does not completely obscure palaeoecological signals within the DPF, other than to artificially inflate estimates of standing crop biodiversity. It also suggests that the MFHs maintained by their respective species were evolutionarily stable strategies, and may reflect correlated stability in the growth habits of the surrounding plants. Major changes in habitat structure do not appear to have occurred until the beginning of the Paleocene [101,105], underscoring the importance of low-growing herbage in sustaining Late Cretaceous herbivore faunas.

Finally, it must be noted that, while differences in estimated MFHs are consistent with the hypothesis of feeding height stratification, they are not sufficient for this hypothesis to be true. It is possible that, despite these differences, all herbivore taxa from the DPF spent most of their time feeding at ground level [33,34]. In this sense, they may be compared to grazing ungulates, which spend most of their time feeding on low grasses, despite being physically capable of reaching higher browse. Therefore, although it seems likely from an ecological perspective that the herbivorous dinosaur fauna of the DPF exhibited some form of feeding height stratification, competing hypotheses about the role of this mechanism in the facilitation of niche partitioning are underdetermined [141] by the available evidence. To reject the null hypothesis of no feeding height stratification, it would be necessary to show that the herbivorous dinosaurs browsed to their full potential, utilizing their entire reconstructed MFHs, and did not simply spend all their time feeding at ground level. Unfortunately, this type of behaviour simply does not fossilize. Nevertheless, it may be possible to approximate the amount of time spent feeding at different heights by observing other aspects of morphology. For example, it has been shown that low-level grazers often possess a suite of cranial characteristics that allow them to efficiently crop short grass, including wide, ventrally deflected muzzles, elongate faces, transversely wide paroccipital processes, deep mandibles, and tall withers [142-150]. Similarly, primates feeding close to the ground generally possess narrower dental microwear scratches than those feeding higher up in the forest canopy, a function of mean particle size and the ratio of soil particles to phytoliths [151-153]. These same features might be sought among the herbivorous dinosaurs to more accurately determine their browsing habits.

Conclusions

Niche partitioning between herbivorous dinosaurs from the DPF could have been facilitated in part by feeding height stratification, as is commonly observed among the ungulates of the Serengeti [78,82]. Small ornithischians (cf. Orodromeus, Unescoceratops, pachycephalosaurids), ankylosaurs, and ceratopsids were generally restricted to feeding below 1 m, and although there is limited data available for the first group, ankylosaurs and ceratopsids may have partitioned the herb layer. Hadrosaurids may also have fed on herbs, but could have avoided competing with the other forms by feeding on taller angiosperm shrubs in times of duress. Sympatric hadrosaurid species, or ontogenetic stages, also could have avoided competing with one another in this way. There is, as yet, no evidence to suggest that any of these taxa were restricted to feeding in either open or closed habitats.

Despite the evidence for feeding height stratification among ankylosaurs, ceratopsids, and hadrosaurids, there is currently no evidence for feeding height stratification within these clades (with the possible exception of quadrupedal versus bipedal hadrosaurids described above). We note, however, that the problem of underdetermination does not allow us to reject the possibility that some form of feeding height stratification occurred within the herb layer. It is also possible that the coexistence of the herbivorous dinosaurs from the DPF was facilitated by other means in addition to feeding height stratification. The prolonged stability of the palaeoecological relationships identified here supports the contention that the fossil assemblage from the Dinosaur Park Formation constitutes a chronofauna.

Availability of supporting data

The data sets supporting the results of this article are included within the article (and its additional files).
Additional files

Additional file 1: Data used in this study.

Additional file 2: Reduced major axis regression results used to estimate missing elements in this study.

Competing interests
The authors declare that they have no competing interests.

Authors' contributions
JCM planned the study, gathered the data, ran the statistics, and wrote the manuscript. DCE contributed to planning the study, collecting the data, and writing the manuscript. MJR and JSA contributed to planning the study and writing the manuscript. MJR and JSA contributed to planning the study and writing the manuscript. All authors reviewed the final draft of the manuscript. All authors read and approved the final manuscript.

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References
1. Archibald JD. Dinosaur Extinction and the End of an Era: What the Fossils Say. New York: Columbia University Press; 1996.
2. Weishampel DB, Young L: Dinosaurs of the East Coast. Baltimore: Johns Hopkins University Press; 1996.
3. Zanno LE, Makovicky PJ: Herbivorous ecomorphology and specialization patterns in theropod dinosaur evolution. Proc Natl Acad Sci USA 2011, 108:232–237.
4. Lehman TM: Late Maastrichtian paleoenvironments and dinosaur biogeography in the Western Interior of North America. Palaeogeogr Palaeoclimatol Palaeoecol 1987, 60:189–217.
5. Lehman TM: Late Campanian dinosaur biogeography in the Western Interior of North America. In Dinofest International: Proceedings of a Symposium Sponsored by Arizona State University. Edited by Wolberg DL, Stump E, Stenger GD. Philadelphia: Academy of Natural Sciences; 1997:223–240.
6. Lehman TM. Late Cretaceous dinosaur provinciality. In Mesozoic Vertebrate Life. Edited by Tanke DH, Carpenter K. Bloomington: Indiana University Press; 2001:310–328.
7. Dodson P: A faunal review of the Judith River (Oldman) Formation, Dinosaur Provincial Park, Alberta. Mosasaur 1983, 189–118.
8. Sampson SD, Loewen MA: Unraveling a radiation: a review of the diversity, stratigraphic distribution, biogeography, and evolution of horned dinosaurs (Ornithischia: Ceratopsidae). In New Perspectives on Horned Dinosaurs: The Royal Tyrell Museum Ceratopsian Symposium. Edited by Ryan MJ, Chinney-Allgeier BJ, Eberth DA. Bloomington: Indiana University Press; 2010:405–427.
9. Gates TA, Sampson SD, Zanno LE, Roberts EM, Eaton JG, Nydam RL, Hutchison JH, Smith JA, Loewen MA, Getty MA: Biogeography of terrestrial and freshwater vertebrates from the Late Cretaceous (Campanian) Western Interior of North America. Palaeogeogr Palaeoclimatol Palaeoecol 2010, 291:371–387.
10. du Toit JT, Cumming DNH: Functional significance of ungulate diversity in African savannas and the ecological implications of the spread of pastoralism. Biodiversity Conserv 1999, 8:1643–1661.
11. Farlow JO: A consideration of the trophic dynamics of a Late Cretaceous large-dinosaur community (Oldman Formation). Ecology 1976, 57:841–857.
12. Béland P, Russell DA: Paleoeocology of Dinosaur Provincial Park (Cretaceous), Alberta, interpreted from the distribution of articulated vertebrate remains. Can J Earth Sci 1978, 15:1012–1024.
13. Coe MJ, Dilcher DL, Farlow JO, Jarzen DM, Russell DA: Dinosaurs and land plants. In The Origins of Angiosperms and their Biological Consequences. Edited by Frisri EM, Chaloner WG, Crane PR. Cambridge: Cambridge University Press; 1987:225–258.
14. Farlow JO, Dodson P, Chinnery-Allgeier A: Dinosaur biology. Annu Rev Ecol Syst 1995, 26:445–471.
15. Horner JR, Makela R: Nest of juveniles provides evidence of family structure among dinosaurs. Nature 1979, 282:296–298.
16. Lockley MG, Young BH, Carpenter K: Hadrosaur locomotion and herding behavior: evidence from footprints in the Mesaverde Formation, Grand Mesa Coal Field, Colorado, Mountain Geol 1983, 20:5–14.
17. Currie PJ, Dodson P: Mass death of a heard of ceratopsian dinosaurs. In Third Symposium of Mesozoic Terrestrial Ecosystems. Edited by Reif WE, Westphal F. Tübingen: Attempto Verlag; 1984:52–60.
18. Varriochio DJ, Horner JR: Hadrosaurid and lambeosaurid bone beds from the Upper Cretaceous Two Medicine Formation of Montana: taphonomic and biologic implications. Can J Earth Sci 1993, 30:997–1006.
19. Lockley MG, Hunt AP: Dinosaur Tracks and Other Fossil Footprints of the Western United States. New York: Columbia University Press; 1995.
20. Ryan MJ, Russell AP, Eberth DA, Currie PJ: The taphonomy of a Centrosaurus (Ornithischia: Ceratopsidae) bone bed from the Dinosaur Park Formation (Upper Campanian), Alberta, Canada, with comments on cranial ontogeny. PALAIOS 2001, 16:482–506.
21. Eberth DA, Brinkman DB, Bakas V: A centrosaurine mega-bonebed from the Upper Cretaceous of southern Alberta: implications for behavior and death events. In Perspectives on Horned Dinosaurs: The Royal Tyrell Museum Ceratopsian Symposium. Edited by Ryan MJ, Chinney-Allgeier BJ, Eberth DA. Bloomington: Indiana University Press; 2010:495–508.
22. Hunt RK, Farke AA: Behavioral interpretations from ceratopsid bonebeds. In Perspectives on Horned Dinosaurs: The Royal Tyrell Museum Ceratopsian Symposium. Edited by Ryan MJ, Chinney-Allgeier BJ, Eberth DA. Bloomington: Indiana University Press; 2010:447–455.
23. Sampson SD: Dinosaur Odyssey: Fossil Threads in the Web of Life. New York: Columbia University Press; 1990.
24. Ostrom JH: A reconsideration of the paleoecology of hadrosaurian dinosaurs. Am J Sci 1964, 262:975–997.
25. Janis CM, Damuth J, Theodor M: Miocene ungulates and terrestrial primary productivity: where have all the browsers gone? Proc Natl Acad Sci USA 2000, 97:7899–7904.
26. Janis CM, Damuth J, Theodor M: The species richness of Miocene browsers, and implications for habitat type and primary productivity in the North American grassland biome. Palaeogeogr Palaeoclimatol Palaeoecol 2004, 203:371–398.
27. Miller C, LaPasha C: Flora of the Early Cretaceous Kootenai Formation in Montana, confiners. Palaeonto Abt B 1984, 1931:17.
28. Beervel DJ: Modelling palaeophylosynthesis: Late Cretaceous to present. Phil Trans R Soc B 1994, 346:421–432.
29. Beervel DJ: Global terrestrial productivity in the Mesozoic era. Geol Soc London, Spec Pub 2000, 181:17–32.
30. Dechert S: Primary productivity and forage quality of Ginkgo biloba in response to elevated carbon dioxide and oxygen – an experimental approach to mid-Mesozoic paleoecology. PhD thesis. North Carolina State University, Department of Marine, Earth, and Atmospheric Sciences; 2006.
31. Sinclair ARE: Does interspecific competition or predation shape the African ungulate community? J Anim Ecol 1985, 54:899–918.
32. Sander PM, Gee CT, Hummel J, Claus LA: Mesozoic plants and dinosaur herbivory. In Plants in Mesozoic Time: Morphological Innovations, Phylogeny,
Ecosystems. Edited by Geij CT. Bloomington: Indiana University Press; 2011:331–359.

33. Bakker RT. Dinosaur feeding behaviour and the origin of flowering plants. Nature 1978, 274:661–663.

34. Bakker RT. The Dinosaur Heresies: New Theories Unlocking the Mystery of the Dinosaurs and their Extinction. New York: Zebra Books; 1986.

35. Dodson P. Taxonomic implications of relative growth in lambeosaurine hadrosaurs. Syst Biol 1975, 24:37–54.

36. Carrano MT, Janis CM, Sepkoski JJ Jr. Hadrosaurs as unigulate parallels: lost lifestyles and deficient data. Acta Palaeontol Pol 1999, 44:237–261.

37. Henderson DM. Skull shapes as indicators of niche partitioning by sympatric hadrosaurine and centrosaurine dinosaurs. In Perspectives on horns in Dinosaurs: The Royal Tyrrell Museum Ceratopsian Symposium. Edited by Ryan MJ, Chinnery Allgeier BJ, Bether DA. Bloomington: Indiana University Press; 2010:299–307.

38. Carpenter K. Skeletal and dermal armor reconstructions of Europlophus tutus (Ornithischia: Ankylosauridae) from the Late Cretaceous Oldman Formation of Alberta. Can J Earth Sci 1982, 19:689–697.

39. Carpenter K. Ankylosaurus. In Encyclopedia of Dinosaurs. Edited by Currie PJ, Padlan K. San Diego: Academic Press; 1997:16–20.

40. Carpenter K. Ankylosaur. In The Complete Dinosaur. Edited by Farlow JO, Brett-Surman MK. Bloomington: Indiana University Press; 1997:307–316.

41. Carpenter K. Redescription of Ankylosaurus magniventris Brown 1908 (Ornithischia) from the Upper Cretaceous of the Western Interior of North America. Can J Earth Sci 2004, 41:961–980.

42. Campione NE, Evans DC. Cranial growth and variation in Edmontosaurus (Dinosauria: Hadrosauridae): implications for latest Cretaceous megarhivory in North America. PLoS One 2011, 6(9)e25186. doi:10.1371/journal.pone.0025186.

43. Lull RS: A revision of the Ceratopsia or horned dinosaurs. Mem Peabody Mus Nat Hist 1933, 1:1–175.

44. Whitlock JA. Inferences of diplodocoid (Sauropoda: Dinosauria) feeding behavior from snout shape and microwear analyses. PLoS One 2011, 6(4)e18304. doi:10.1371/journal.pone.0018304.

45. Ostrow JH. Functional morphology and the evolution of the ceratopsian dinosaurs. Evolution 1966, 20:290–308.

46. Erickson GM, Kirk BA, Hamilton M, Bourne GR, Norell MA, Lilleodden E, Sawyer WG. Complex dental structure and wear biomechanics in hadrosaurid dinosaurs. Science 2012, 337:98–101.

47. Ryan MJ, Evans DC. Ornithischian dinosaurs. In Dinosaur Provincial Park: a Spectacular Ancient Ecosystem Revealed. Edited by Currie PJ, Koppelhus EP. Bloomington: Indiana University Press; 2005:312–348.

48. Brown CM, Evans DC. Megaherbivorous dinosaur turnover in the Dinosaur Park Formation (upper Campanian) of Alberta, Canada. Palaeogeogr Palaeocl 2012, 350–352:124–138.

49. Calvo JO. Jaw mechanisms in sauropod dinosaurs. Gaia 1994, 10:183–194.

50. Fiorelli AR. Dental microwear on the teeth of Camarasaurus and Diplodocus: implications for sauropod paleoecology. In Fifth Symposium on Mesozoic Terrestrial Ecosystems and Biota. Edited by Kielan-Jaworowska Z, Heintz N, Nakrem HA. Oslo: Paleontologisk Museum, University of Oslo; 1991:23–24.

51. Fiorelli AR. Dental microwear patterns of the sauropod dinosaurs. In The Complete Dinosaur. Edited by Farlow JO, Brett-Surman MK. Bloomington: Indiana University Press; 2010:537–569.

52. Calvo JO: Jaw mechanisms in sauropod dinosaurs. Gaia 1994, 10:183–194.

53. Fiorelli AR. Dental microwear patterns of the sauropod dinosaurs. Acta Palaeontol Pol 2009, 54:213–230.

54. Fiorelli AR. Lack of variability in feeding patterns of the sauropod dinosaurs Diplodocus and Camarasaurus (Late Jurassic, western USA) with respect to climate as indicated by tooth wear features. In Vertebrate Microfossil Assemblages: their Role in Paleontology and Paleoecobiography. Edited by Sankey JB, Basilo S. Bloomington: Indiana University Press; 2008:104–116.

55. Stevens KA, Parrish JM. Neck posture and feeding habits of two Jurassic sauropod dinosaurs. Science 1999, 284:798–800.

56. Stevens KA, Parrish JM. Digital reconstructions of sauropod dinosaurs and implications for feeding. In The Sauropods: Evolution and Paleobiology. Edited by Currie Rogers KA, Wilson JA. Berkeley: University of California Press; 2005:178–200.

57. Upchurch P, Barrett PM. The evolution of sauropod feeding mechanisms. In Evolution of Herbivory in Terrestrial Vertebrates: Perspectives from the Fossil Record. Edited by Sues H-D. Cambridge: Cambridge University Press; 2000:79–122.

58. Taylor MP, Wedel MJ, Nash D. Head and neck posture in sauropod dinosaurs inferred from extant animals. Acta Palaeontol Pol 2009, 54:213–230.

59. Brinkman DB. Paleoecology of the Judith River Formation (Campanian) of Dinosaur Provincial Park, Alberta, Canada: evidence from vertebrate microfossil localities. Palaeogeogr Palaeoclimatol Palaeoecol 1990, 78:37–54.

60. Brinkman DB, Ryan MJ, Eberth DA. The paleogeographic and stratigraphic distribution of ceratopsids (Ornithischia) in the upper Judith River Group of western Canada. PALAIOS 1998, 13:160–169.

61. Brown CM, Russell AP. Homology and architecture of the caudal basket of Pachycephalosaurus (Dinosauria: Ornithischia): the first occurrence of myorrhabilids in Tetrapoda. PLoS One 2012, 7(1):e30212. doi:10.1371/journal.pone.0030212.
Sedimentology and taphonomy of the Oldman Formation revisited. In Evolution of Herbivory in Terrestrial Vertebrates: Perspectives from the Fossil Record. Edited by Sues H-D. Cambridge: Cambridge University Press; 2000:123–143.

Horner JR, Weishampel DB, Forster CA: Weishampel DB, Jianu C-M: Cameron EZ, du Toit JT: Barrett PM, Upchurch P: Sauropod feeding mechanisms: their bearing on palaeoecology. In Sixth Symposium on Mesozoic Terrestrial Ecosystems and Biota. Short Papers. Edited by Sun A, Wang Y. Beijing: China Ocean Press; 1995:107–110.

Weishampel DB, Norman DB: Vertebrate herbivory in the Mesozoic: jaws, plants, and evolutionary metrics. Geol Soc Am Spec Pap 1989, 238:87–100.

Weishampel DB, Jianu C-M: Plant-eaters and ghost lineages: dinosaurian herbivory revisited. In Evolution of Herbivory in Terrestrial Vertebrates: Perspectives from the Fossil Record. Edited by Sues H-D. Cambridge: Cambridge University Press; 2000:123–143.

Hommel JR, Weishampel DB, Forster CA: Weishampel DB, Jianu C-M: Cameron EZ, du Toit JT: Barrett PM, Upchurch P: Sauropod feeding mechanisms: their bearing on palaeoecology. In Sixth Symposium on Mesozoic Terrestrial Ecosystems and Biota. Short Papers. Edited by Sun A, Wang Y. Beijing: China Ocean Press; 1995:107–110.

83. Arsenukt R, Owen-Smith N: Resource partitioning by grass height among grazing ungulates does not follow body size relation. Okos 2008, 117:1711–1717.
84. Cameron EZ, du Toit JT: Winning by a neck: tall giraffes avoid competing with shorter browsers. Amer Nat 2007, 169:130–135.
85. Fiorillo AR: Time resolution at Carnegie Quarry (Morrison Formation: Dinosaur National Monument, Utah). Univ Wyo Contib Geol 1994, 30:149–156.
86. Barreto PM, Upchurch P: Sauropod feeding mechanisms: their bearing on palaeoecology. In Sixth Symposium on Mesozoic Terrestrial Ecosystems and Biota. Short Papers. Edited by Sun A, Wang Y. Beijing: China Ocean Press; 1995:107–110.
87. Weishampel DB, Norman DB: Vertebrate herbivory in the Mesozoic: jaws, plants, and evolutionary metrics. Geol Soc Am Spec Pap 1989, 238:87–100.
88. Weishampel DB, Jianu C-M: Plant-eaters and ghost lineages: dinosaurian herbivory revisited. In Evolution of Herbivory in Terrestrial Vertebrates: Perspectives from the Fossil Record. Edited by Sues H-D. Cambridge: Cambridge University Press; 2000:123–143.
89. Hommel JR, Weishampel DB, Forster CA: Weishampel DB, Jianu C-M: Cameron EZ, du Toit JT: Barrett PM, Upchurch P: Sauropod feeding mechanisms: their bearing on palaeoecology. In Sixth Symposium on Mesozoic Terrestrial Ecosystems and Biota. Short Papers. Edited by Sun A, Wang Y. Beijing: China Ocean Press; 1995:107–110.
134. Chin K, Gill BD: Dinosaurs, dung beetles, and conifers: participants in a Cretaceous food web. PALAIOS 1996, 11:280–285.

135. Chin K: The paleobiological implications of herbivorous dinosaur coprolites from the Upper Cretaceous Two Medicine Formation of Montana: why eat wood? PALAIOS 2007, 22:554–566.

136. Abel O: Diskussion zu den Vorträgen R. Krausel und F. Versluys. Paläont Z 1922, 4:87.

137. Chin K: Coprolites. In Encyclopedia of Dinosaurs. Edited by Currie PJ, Padian K. San Diego: Academic Press; 1997:147–150.

138. Molnar RE, Clifford HT: An ankylosaurian cololite from the Lower Cretaceous of Queensland, Australia. In The Armored Dinosaurs. Edited by Carpenter K. Bloomington: Indiana University Press; 2001:399–412.

139. Galton PM: Hypsilophodon, the cursorial nonarboreal dinosaur. Nature 1971, 231:159–161.

140. Galton PM: The mode of life of Hypsilophodon, the supposedly arboreal ornithopod dinosaur. Lethaia 1971, 4:453–465.

141. Turner D: Paleontology: a Philosophical Introduction. Cambridge: Cambridge University Press; 2011.

142. Janis CM, Ehrhardt D: Correlation of relative muzzle width and relative incisor width with dietary preference in ungulates. Zool J Linn Soc 1988, 92:267–284.

143. Solounias N, Dawson-Saunders B: Dietary adaptations and paleoecology of the Late Miocene ruminants from Pikermi and Samos in Greece. Palaeogeogr Palaeoclimatol Palaeoecol 1988, 65:149–172.

144. Janis CM: Correlation of cranial and dental variables with dietary preferences in mammals: a comparison of macropodoids and ungulates. Mem Queensl Mus 1990, 28:349–366.

145. Janis CM: Correlations between craniodental morphology and feeding behavior in ungulates: reciprocal illumination between living and fossil taxa. In Functional Morphology in Vertebrate Paleontology. Edited by Thomason JJ. Cambridge: Cambridge University Press; 1995:76–98.

146. Solounias N, Moelleken SMC: Dietary adaptation of some extinct ruminants determined by premaxillary shape. J Mammal 1993, 74:1059–1071.

147. Spencer LM: Morphological correlates of dietary resource partitioning in the African Bovidae. J Mammal 1995, 76:448–471.

148. Mendoza M, Janis CM, Palmqvist P: Characterizing complex craniodental patterns related to feeding behaviour in ungulates: a multivariate approach. J Zool 2002, 258:223–246.

149. Fraser D, Theodor JM: Anterior dentary shape as an indicator of diet in ruminant artiodactyls. J Vertébr Paleontol 2011, 31:1366–1375.

150. Fraser D, Theodor JM: Comparing ungulate dietary proxies using discriminant function analysis. J Morphol 2011, 272:1513–1526.

151. Walker A: Wear striations on the incisors of ceropithecid monkeys as an index of diet and habitat preference. Am J Phys Anthropol 1976, 45:299–308.

152. Ungar PS: Incisor microwear of Sumatran anthropoid primates. Am J Phys Anthropol 1994, 94:339–363.

153. Ungar PS: Dental microwear of European Miocene catarrhines: evidence for diets and tooth use. J Hum Evol 1996, 31:335–366.