# Evolutionary trends and the origin of the mammalian lower jaw

Christian A. Sidor

Abstract.—The single bony element forming the lower jaw of living mammals, the dentary, has been interpreted as representing the culmination of a long and gradual evolutionary trend. Numerous fossil nonmammalian synapsids ("mammal-like reptiles") show varying degrees of enlargement of the dentary and concomitant reduction in the postdentary bones. To quantitatively reexamine patterns of morphological change in the evolution of the mammalian lower jaw, measurement and discrete character data were collected from 322 fossil synapsid mandibles spanning Late Carboniferous through Jurassic time. Measurements confirm that the relative contribution of the dentary increased in theriodont (advanced therapsid) evolution with regard to both stratigraphic and phylogenetic position. However, dentary enlargement and postdentary reduction failed to typify all therapsid subclades. Qualitative characters of the mandible were used to quantify morphological similarity with regard to the early mammal Morganucodon. Analyses contrasting stratigraphic and phylogenetic position with mammalian similarity indicate that mandibular evolution was primarily conservative, with only anomodont therapsids evolving substantial morphological novelty. Scaling analyses comparing the area of the dentary and postdentary regions to jaw length uniformly show isometry or slight positive allometry, although cynodont therapsids have a smaller postdentary region than any other therapsid subgroup. These results suggest that body size decreases cannot fully explain the reduction of the postdentary bones. Finally, step size bias was tested as a mechanism for explaining long-term trends. Qualitative data reveal no significant difference in the magnitude of character changes occurring in mammalian and nonmammalian directions.

Christian A. Sidor. Department of Anatomy, New York College of Osteopathic Medicine, Old Westbury, New York 11568-8000. E-mail: casidor@iris.nyit.edu

Accepted: 31 January 2003

# Introduction

Mammals are unique among extant vertebrates in possessing a lower jaw (mandible) formed by a single bony element, the dentary. By contrast, the lower jaws of other vertebrates retain a host of postdentary bones (e.g., four to six in most lizards, five in crocodiles and many birds, and typically even greater numbers in fishes). Recorded from rocks dating from over 300 Ma, the mandibles of the earliest nonmammalian synapsids possessed up to seven postdentary bones (Fig. 1), whereas stratigraphically more recent taxa show various stages in the reduction and eventual loss of these bones (Fig. 2) (Romer and Price 1940; Crompton 1963; Allin 1975). The evolutionary fate of the mammalian postdentary bones has been well established; Reichert (1837) used embryological evidence to homologize the incus and malleus of the mammalian middle ear with the quadrate and articular, respectively, of nonmammalian vertebrates. The transformation of several postdentary jaw bones into sound-conducting middle ear bones within

synapsids is one of the best-documented examples of a major evolutionary transformation in the vertebrate fossil record (Hopson 1966; Allin 1975; Allin and Hopson 1992; Luo and Crompton 1994). Indeed, synapsid mandibular evolution has come to be regarded as recording a gradual trend whereby enlargement of the dentary occurs at the expense of the postdentary bones (Crompton and Jenkins 1973; Kemp 1982; Hopson 1987). In this study, I use measurement and discrete character data to: (1) quantify the morphological changes that occurred in the evolution of the lower jaw between pelycosaur-grade synapsids and their mammalian descendants, and (2) address several previously proposed hypotheses concerning the nature and magnitude of morphological trends during the first ~100 Myr of synapsid history.

Background to Study Taxa.—Theories of synapsid evolution have traditionally been couched in terms of several adaptive radiations or grades of organization representing successive steps in the mammalian direction. However, a recent proliferation of numerical

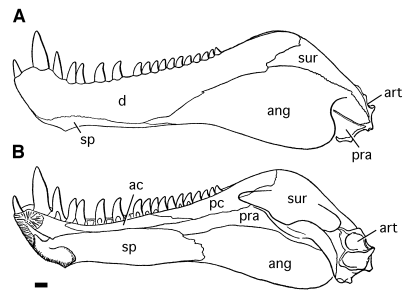


FIGURE 1. The lower jaw of the Late Carboniferous and Early Permian pelycosaur-grade synapsid *Dimetrodon* in lateral (A) and medial (B) views. Scale bar, 1 cm. Anatomical abbreviations: ac = anterior coronoid, ang = angular, art = articular, d = dentary, pc = posterior coronoid, pra = prearticular, sp = splenial, sur = surangular. Illustration modified from Romer and Price 1940.

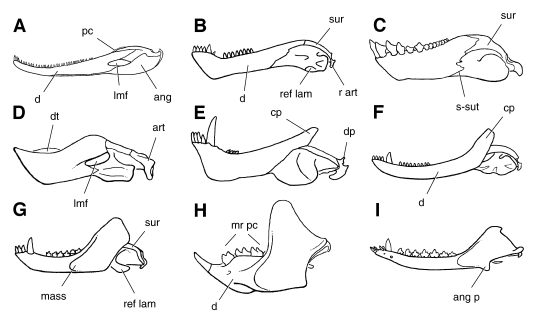


FIGURE 2. Representative synapsid mandibles in lateral view (not to same scale). A, The Late Pennsylvanian and Early Permian "pelycosaur" *Ophiacodon*. B, The Late Permian biarmosuchian *Biarmosuchus*. C, The Late Permian tapinocephalid dinocephalian *Ulemosaurus*. D, The Late Permian advanced dicynodont *Diictodon*. E, The Late Permian gorgonopsid *Arctognathus*. F, The Late Permian therocephalian *Ictidosuchoides*. G, The Early Triassic primitive cynodont *Thrinaxodon*. H, The late Early Jurassic or early Middle Jurassic tritylodontid *Bocatherium*. I, The Early Jurassic primitive mammal *Morganucodon*. Anatomical abbreviations: ang = angular, ang p = angular process, art = articular, cp = freestanding coronoid process, d = dentary, dp, dorsal process of the articular; dt = dentary tables, lmf = lateral mandibular fenestra, mass = masseteric fossa, mr pc = multirooted postcanines, pc = posterior coronoid, r art = retroarticular process, ref lam = reflected lamina, s-sut = s-shaped dentary/angular suture, sur = surangular. Illustration sources: Romer and Price 1940 (A), Efremov 1940 (C), and Hopson 1994 (remaining figure parts).

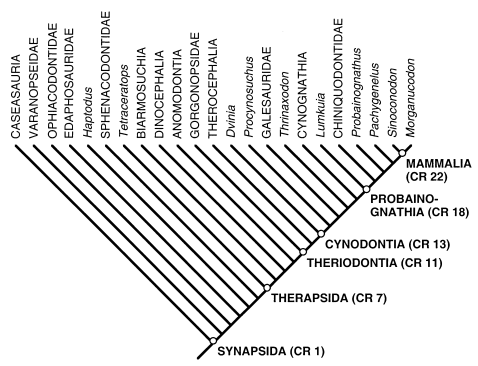


FIGURE 3. Cladogram of higher-level synapsid relationships used herein. This topology is based primarily on those proposed by Reisz (1986) and Sidor and Hopson (1998). See Appendix 1 for cladogram details and Figure 4 for lower-level relationships. This cladogram includes 22 nodes along its spine that define the 22 clade ranks (CR) used in later analyses. For example, taxa included within the Varanopseidae and Galesauridae have CRs of 2 and 15, respectively.

cladistic analyses has contributed greatly to our understanding of synapsid phylogeny (Fig. 3), and regions of broad consensus are gradually emerging (Rubidge and Sidor 2001).

The earliest occurring and phylogenetically most primitive synapsids are the "pelycosaurs" of traditional terminology. These taxa form a paraphyletic series and are primarily known from Upper Carboniferous to Lower Permian rocks in Europe and North America (Reisz 1986) although several taxa persisted into the Middle Permian in Russia and South Africa (Reisz et al. 1998; Modesto et al. 2001). Sphenacodontids, such as the familiar sailback *Dimetrodon*, are among the most advanced pelycosaur subgroups (Reisz et al. 1992). All more derived synapsids form the clade Therapsida.

All of the major therapsid clades first appear in the fossil record during the Middle and Late Permian (e.g., Biarmosuchia, Dinocephalia, Anomodontia, Gorgonopsia, Thero-

cephalia, and Cynodontia) and—except for dicynodont anomodonts, some advanced therocephalians, and cynodonts—went extinct in this time interval as well. Therapsids taxonomically and ecologically dominated the end-Paleozoic Pangaean landscape and established the first herbivore-based food chains among vertebrates in the terrestrial realm (Olson 1962; King et al. 1989; Reisz and Sues 2000). The presence of several derived features recently led Laurin and Reisz (1990, 1996) to suggest that *Tetraceratops insignis*, from the Early Permian of Texas, is phylogenetically the most primitive therapsid (but see Conrad and Sidor 2001).

Cynodonts are first recorded from Upper Permian strata in southern Africa and Russia and represent the therapsid subclade that includes mammals. Numerous derived features associated with obtaining food and its mastication characterize Cynodontia, including the presence of a fossa for the neomorphic masseter muscle on the lateral surface of the dentary, postcanine teeth with accessory cusps and lingual cingula, and a complete sagittal crest for the origin of temporalis musculature. According to the phylogenetic hypothesis proposed by Hopson (1991b, 1994; Hopson and Kitching 2001), a key dichotomy in cynodont phylogeny occurred with the Triassic divergence of the cynognathian and probainognathian lineages. Terminal cynognathians (tritylodontids) range stratigraphically upwards into the Lower Cretaceous (Tatarinov and Matchenko 1999), whereas terminal probainognathians (mammals) first appear in Upper Triassic or Lower Jurassic rocks and survive until the Recent (Lucas and Luo 1993; Luo 1994).

Vertebrate paleontologists have traditionally defined mammals as possessing a wellformed dentary-squamosal jaw joint (Simpson 1960). Taxa included under this (apomorphybased) definition include Morganucodon and Kuehneotherium, although these and other early Mesozoic forms (e.g., Sinoconodon) probably possessed a functional quadrate-articular jaw joint as well (Hopson 1991b; Luo and Crompton 1994). More recently, Rowe (1988) and Rowe and Gauthier (1992) have advocated using a crown-group definition for Mammalia, and they have termed the larger clade-including traditional mammals that lie phylogenetically outside the clade bounded by extant forms-Mammaliaformes. My use of Mammalia and of the terms "mammal" and "mammalian" correspond to traditional usage (see also Luo et al. 2002).

# **Data Collection**

Taxon Sampling.—Fossil synapsids included in this study range from the earliest-appearing (Late Carboniferous) pelycosaur-grade taxa through some of the most primitive mammals, such as the Early Jurassic genera Morganucodon and Sinoconodon. In total, 19 "pelycosaurs," six basal therapsids, 13 dinocephalians (including five anteosaurians and eight tapinocephalians), 25 anomodonts, ten gorgonopsians, ten therocephalians, and 25 cynodonts were included. The cynodonts include six non-eucynodonts, 11 cynognathians (including six tritylodontids), and eight probainognathians (including two Mesozoic

mammals). All taxa were at the genus level or, in several instances, below.

The stratigraphic range of each taxon was collected from original museum locality information or the literature (e.g., Kitching 1977; Rubidge 1995; Ivachnenko et al. 1997) and then binned into one or more of 18 age ranks (AR) for the purpose of analysis. ARs are nonoverlapping stratigraphic bins in an ordered sequence (Gauthier et al. 1988). Importantly, ARs are not necessarily of equal duration; some ARs are equivalent to a single geological formation, whereas others encompass several formations or groups. The goal of this type of binning is a single, resolved sequence of the synapsid fossil record despite its derivation from a variety of widely separated continental deposits (see also Sidor 2001). One major drawback to the AR approach is that gaps in the synapsid record are effectively ignored; time periods lacking synapsid fossils are not represented in the analysis. For example, a major hiatus in the synapsid record occurs between the youngest continental deposits in North America (e.g., the San Angelo and Flowerpot Formations) and the oldest in Russia and South Africa (e.g., Mezen and the Eodicynodon Assemblage Zone, respectively) (Lucas and Heckert 2001). This approximately 2-Myr hiatus encompasses much of Roadian time, but is not evident between ARs 6 and 7. Appendix 4 reports the geological formations and vertebrate biozones making up each AR.

From a recent compilation of synapsid cladistic analyses, I also collected phylogenetic inference data, which consisted of each taxon's clade rank (CR) (Gauthier et al. 1988) and the number of branch points from the root of the cladogram (i.e., patristic distance, PD) (Figs. 3, 4). A rationale for this specific arrangement of synapsid relationships is provided in Appendix 1. CR equals the number of branching points a taxon is positioned up the phylogenetic trajectory from Synapsida to Mammalia (Fig. 3). Branching within a terminal taxon on this pectinate tree is not considered. For example, every species within Gorgonopsia has a CR of 11. In contrast, PD measures the total number of nodes passed from the root of the cladogram to the taxon in question because branching within side-branches is taken into

account. Only when a singleton attaches directly to the primary spine of the cladogram (e.g., *Tetraceratops* or *Dvinia*) are CR and PD equal.

Data were collected from study of fossil specimens at the following institutions: Albany Museum, Grahamstown, South Africa; American Museum of Natural History, New York; Field Museum of Natural History, Chicago; Museum of Comparative Zoology, Harvard University, Cambridge, Massachusetts; National Museum of Natural History, Washington, D.C.; University of California Museum of Paleontology, Berkeley; Bernard Price Institute for Palaeontological Research, University of the Witwatersrand, Johannesburg; National Museum, Bloemfontein; South African Museum, Cape Town; Transvaal Museum, Pretoria; The Natural History Museum, London; Museum of Zoology, University of Cambridge, Cambridge, United Kingdom; Oxford University Museum, Oxford; Paleontological Institute, Moscow; Bayerische Staatssammlung für Paläontologie und Historische Geologie, Munich; Humboldt Museum für Naturkunde, Berlin; and Institut und Museum für Geologie und Paläontologie, Tübingen. Only four taxa were coded exclusively from the literature: Bienotheroides wanhsienensis, Ecteninion lunensis, Probelesodon sanjuanensis, and Sinoconodon sp. (Sun 1984; Crompton and Luo 1993; Martinez and Forster 1996; Martinez et al. 1996).

Qualitative Data.—I used 82 qualitative characters to describe morphological variation within the synapsid mandible. These characters included 22 pertaining to the dentary and 41 to the postdentary bones, six general shape features, and 13 dentition-related features. The last set specifically did not include characters describing the morphology of the dentition itself (such as cusp patterns). Rather, these characters focused on dental features manifesting themselves on the form of the lower jaw itself (e.g., whether or not the teeth were set in sockets). Many of the characters and character states were taken from previous cladistic analyses of synapsid relationships. Characters, character state descriptions, and literature references are presented in Appendix 2. The corresponding taxon/character data matrix is in Appendix 3.

The data matrix in Appendix 3 was converted to a taxon/taxon similarity matrix using the Simple Matching Coefficient ( $S_{\rm SM}$ ) (Sokal and Sneath 1963), which is equal to the number of characters for which two taxa match divided by the number of characters for which they could possibly match (in order to accommodate missing or inapplicable characters). Appendix 4 contains the final line in the similarity matrix, which compares the early mammal *Morganucodon* with every other taxon.

Quantitative Data.—Two areas and four linear measurements constitute the quantitative data set (Fig. 5). The area of the dentary and postdentary bones was calculated by digitizing their respective outlines in NIH Image. Synapsid mandibles were oriented norma lateralis, and then the following measurements were taken parallel to the long axis of the jaw: (1) jaw length, measured from the anteriormost point on the dentary to the midpoint of the craniomandibular joint; (2) dorsal length of the dentary, measured from the anterior tip of the dentary to the sutural contact between the dentary and surangular along the dorsal margin of the mandible; (3) ventral length of the dentary, measured from the anterior tip of the dentary to the sutural contact between the dentary and angular along the ventral margin of the mandible; and (4) perpendicular to the previous measurements, height of the coronoid region, equal to the distance from the mandibular joint to the dorsalmost point on the lower jaw. Measurements under 200 mm were taken with digital calipers and recorded to the nearest one-tenth millimeter. Measurements over 200 mm were taken with a measuring tape and recorded to the nearest halfmillimeter. Raw data are presented in Appendix 5.

From the six original variables, four ratios were calculated: (A) the relative position of the dentary/surangular suture, calculated as the ratio of measurement (2) to measurement (1), (B) the relative position of the dentary/angular suture, calculated as the ratio of measurement (3) to measurement (1); (C) the relative height of the coronoid process, calculated as the ratio of measurement (4) to measurement (1); and (D) the relative area of the

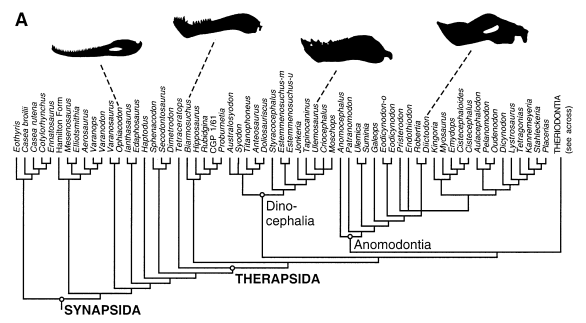


FIGURE 4. Cladogram of lower-level relationships among synapsids used herein. Appendix 1 includes a rationale for this specific topology. A taxon's patristic distance (PD) is calculated as the number of nodes passed from the base of this cladogram. For example, the ophiacodontids *Varanosaurus* and *Ophiacodon* each have a PD of 4. The most primitive cynodont, *Dvinia*, has a PD of 13 and illustrates the fact that singleton taxa attaching directly to the spine of the cladogram have PD and clade ranks (CR) of equal value. PDs for each terminal taxon are given in Appendix 4. CGP 1/61 refers to a new burnetiamorph housed in the collections of the Council for GeoSciences, Pretoria (Sidor 2000). SAM-PK-K9954 refers to a new galesaurid housed in the South African Museum, Cape Town (Sidor and Smith in press). Hamilton Form refers to a new, primitive varanopseid (Reisz and Dilkes 2003). "Estemmenosuchus-m" refers to E. mirabilis, "Estemmenosuchus-u" to E. uralensis, "Probelesodon-lew." to P. lewisi, and "Probelesodon-san." to P. sanjuanensis.

dentary, calculated as the area of the dentary divided by the total area of the lower jaw. Measurements were originally collected from 764 mandibles (Sidor 2000: App. 6.1). Of these, 322 lower jaws preserved at least two measurements and were used to calculate mean values for each taxon for each of the four dentary ratios (Appendix 5). For the purpose of summarizing changes in all four ratios, each taxon's dentary index (DI) was calculated as the average of the four ratios when each was standardized to have a mean of zero and unit variance. Appendix 4 contains the four original ratios and the summary DI, in addition to each taxon's first and last appearances (in ARs), CR, and PD. Because of fossil incompleteness, not every taxon has a complete set of measurements and so could not be included in all analyses.

# Measurement-based Analyses

Stratigraphic Results.—Figure 6A summarizes the results of the measurement-based anal-

yses. The dentary index (DI) is plotted on the abscissa, with increasingly positive values indicating an overall larger contribution of the dentary to the composition of the lower jaw. Although only the DI is depicted, the four individual dentary ratios show similar patterns (Table 1). The ordinate represents the stratigraphic distribution of each taxon in ARs. Significant, positive correlation between the two axes corresponds to a preferential positioning of dentary relative sizes through time.

The pattern depicted in Figure 6A confirms that the earliest-occurring, pelycosaur-grade taxa had the relatively smallest dentaries and largest complement of postdentary bones, and that the latest-occurring synapsids (e.g., tritylodontids and mammals) had mandibles that were almost exclusively formed by the dentary. Importantly, both the maximum and average dentary size increased over time in this study sample. However, it is interesting to note that several late-occurring synapsids retained relatively small dentaries.

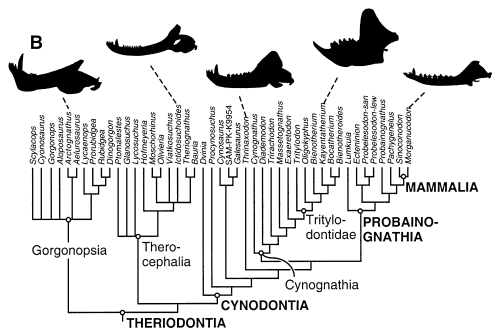


FIGURE 4. Continued.

Table 1 displays the results of a series of analyses that examine evolutionary patterns within several synapsid subclades. These analyses show that a significant positive correlation between DI and AR is nearly uniformly present in those clades encompassing mam-

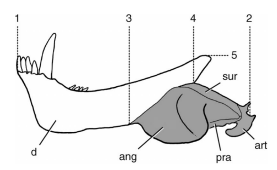


FIGURE 5. Lower jaw of the gorgonopsid *Arctognathus* in lateral view illustrating the five landmarks and two areas measured for this study. Landmarks included (1) the anteriormost tip of the dentary, (2) the midpoint of the jaw joint, (3) the anteriormost contact between the angular and dentary along the ventral margin of the lower jaw, (4) the anteriormost contact between the surangular and dentary along the dorsal surface of the lower jaw, and (5) the dorsalmost point on the coronoid eminence or process. The measurements taken between these landmarks are described in the text. The dentary is unshaded, whereas the area of the postdentary bones is shaded. Anatomical abbreviations as in Figure 1. Figure modified from Kemp 1982.

mals (e.g., Synapsida, Therapsida, Theriodontia, Cynodontia). Probainognathia is the exception to this pattern, but this may be due to the relatively few intervals that this clade spans. In contrast, clades not encompassing mammals (i.e., side branches such as anomodonts) generally have nonsignificant correlations. This crucial disagreement suggests that only the ancestral lineage leading to mammals (i.e., along the backbone of the cladogram) shows a consistent dentary enlargement (see below), and that clades budded off from this line retained their ancestral proportions but did not systematically continue the trend. It is worth noting that pelycosaur-grade synapsids show little indication of directionality, even though they span seven long intervals (ARs 1-7; Late Carboniferous to early Middle Permian, or approximately 35 Myr) and represent the primitive morphotype from which all subsequent change was derived.

Phylogenetic Results.—Figure 6B plots the relationship between DI and each taxon's cladogram position, as measured by CR (see Table 2 for complete results). It is clear from this graph that the degree to which a synapsid clade shares ancestry with mammals has a strong, positive relationship with that clade's

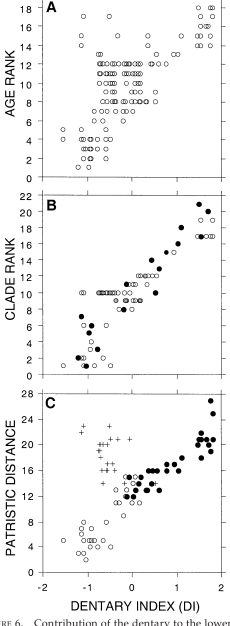


FIGURE 6. Contribution of the dentary to the lower jaw in synapsid evolution. (A) compares the stratigraphic range (measured in Age Ranks) to each fossil synapsid's dentary index (DI). The DI is computed as the average score of the four original dentary ratios when each is scaled to a mean of zero with unit variance. Low DIs correspond to mandibles with relatively small dentaries and low coronoid regions, whereas high DIs correspond to the opposite. In (B), filled circles represent the inferred primitive condition at each clade rank. Grayed circles indicate the two most primitive members for clades with no single most primitive taxon. The following taxa were used in these cases: Ianthasaurus for edaphosaurids (Modesto 1995), Syodon and Styracocephalus for dinocephalians (Rubidge 1994; Rubidge and van den Heever 1997), Patranomodon (in A) or Ulemica for anom-

average dentary size. However, when the inferred primitive condition for each consecutive clade is highlighted (filled circles), this point does not consistently reside in the left tail of that clade's range of DI values. This position suggests that diversification within each synapsid subclade expanded the range of DI values but did not uniformly increase the relative contribution of the dentary.

Directionality within subclades is more fully considered in Figure 6C (and Table 3), which contrasts the number of branch points separating each taxon from the root of the cladogram (patristic distance; PD) with its DI. Because stratigraphic and phylogenetic position show a strong relationship in synapsids (Sidor and Hopson 1998), Figure 6C is very similar to 6A. Taxa diverging relatively early (i.e., with low PDs) tend to have small dentaries, whereas phylogenetically more derived taxa show a wider range of values. The expanding range of values observed at high PDs can be attributed to the persistence of smalldentaried anomodonts (plus signs) with theriodonts (filled circles) that consistently enlarge the dentary. However, just as with the stratigraphic analyses, the individual theriodont subclades that lack mammals as a subgroup lack a corresponding trend (Table 3). Again, this suggests that increasing the dentary size was not a universal feature of synapsid evolution.

 $\overline{\underline{\phantom{a}}}$ 

odonts (Rubidge and Hopson 1996), Cyonosaurus for gorgonopsids (Sigogneau 1970), Ptomalestes and Glanosuchus for therocephalians (Hopson and Barghusen 1986), and Sinoconodon for mammals (Luo 1994). A strong trend for increasing the dentary's overall contribution to the composition of the mandible is present among the primitive members of each consecutively more advanced clade. Interestingly, however, the most primitive member of each subgroup does not tend to be positioned at the low end of its group's distribution, suggesting that a within-subclade evolution does not display the same pattern. In (C), open circles are stem taxa, plus signs are anomodonts, and filled circles are theriodonts. Theriodonts show the strongest relationship between PD and each of the four dentary size measurements whereas anomodonts consistently display none. Note that the variance in DI observed in anomodonts in (C) is collapsed in to a single horizontal line in (B). Statistics for (A), (B), and (C) are in Tables 1, 2, and 3, respectively.

included here to represent the early portion of synapsid evolution. All other subgroups represent clades. Only first appearances were used in this analysis (mean taxon Results of Spearman rank correlation tests for lower jaw measurements versus stratigraphic range (in age ranks). "Pelycosaurs" are not a clade, but they are duration = 1.6 age ranks). The test statistic (Rho) and p-value are corrected for ties in this and subsequent Tables.  $n^* p = 0.0570$ .

Rho p  0.77 <0.0001 0.29 ns 0.61 <0.0001 0.41 ns -0.02 ns 0.86 <0.0001 -0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns		Oorsal	Ver	ıtral	Corc	noid	Ι	Σ
0.77 <0.0001 0.29 ns 0.61 <0.0001 0.41 ns -0.02 ns 0.86 <0.0001 0.25 ns 0.25 ns 0.25 ns 0.25 ns 0.64 ns	р Rho	d	Rho	d	Rho	d	Rho	d
0.29 ns 0.61 <0.0001 0.41 ns -0.02 ns 0.86 <0.0001 -0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns	0.0001 0.42	<0.0001	0.61	<0.0001	0.55	<0.0001	0.64	<0.0001
0.61 <0.0001 0.41 ns -0.02 ns 0.86 <0.0001 -0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns	ns 0.24	ns	0.26	ns	-0.08	us	0.14	su
0.41 ns -0.02 ns 0.86 <0.0001 -0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns		0.0005	0.52	< 0.0001	0.35	0.0040	0.46	0.0003
-0.02 ns 0.86 <0.0001 -0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns		ns	0.49	ns	-0.06	us	0.39	su
0.86 <0.0001 -0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns		ns	-0.30	ns	-0.19	us	-0.33	su
-0.46 ns 0.25 ns 0.83 <0.0001 0.64 ns		<0.0001	0.82	< 0.0001	0.47	0.0055	0.89	< 0.0001
0.25 ns 0.83 <0.0001 0.64 ns		ns	-0.66	0.0475	0.78	ns*	-0.53	ns
0.83 <0.0001 0.64 ns		ns	0.01	ns	0.11	ns	0.64	ns
0.64 ns		0.0012	0.65	0.0027	0.29	ns	0.81	0.0008
		ns	0.27	ns	0.32	ns	0.52	ns
Tritylodontidae $-1.00$ ns $0$ .	ns 0.36	0.0008	0.50	ns	-0.40	ns	-1.00	ns
Probainognathia 0.99 0.0275 0.		ns	0.36	ns	-0.55	ns	0.36	su

# **Character-based Analyses**

The six measurements used above can provide only a limited view of morphological changes occurring within synapsid mandibular evolution. Potentially more informative is quantifying morphological similarity with reference to an exemplar primitive mammal (Morganucodon, in this case) using discrete characters (Appendix 4). Phenetic similarity is an appropriate metric to use in this case because the convergent acquisition of a certain phenotype pertains to net, rather than total, morphological change (Foote 1996). As with the measurement-based analyses, significantly positive correlations between the degree of similarity to mammals and stratigraphic or phylogenetic position would support the hypothesis of a morphological trend toward gaining mammal-like features, whereas nonsignificant correlations would refute it. Furthermore, negative correlations correspond to increasing dissimilarity; i.e., the morphological modifications experienced by a clade's lower jaw consistently distance it from the mammalian position in morphospace.

Stratigraphic and Phylogenetic Results.—Figure 7 and Table 4 contain the principal results of the discrete character-based analyses, which are remarkably similar to those based on measurements (compare with Fig. 6). This similarity implies that both data sets are capturing a common signal from synapsid evolution. When compared with stratigraphic position (Fig. 7A), Late Carboniferous and Early Permian pelycosaur-grade synapsids begin with approximately 60% of their (comparable) lower-jaw characters matching the condition in Morganucodon (AR 1-6). By the middle of Late Permian times (AR 9), however, therapsid diversification expanded this range of values, with anomodonts becoming increasingly dissimilar to mammals, and theriodonts becoming increasingly similar (presumably through synapomorphy). The "increase in variance" pattern continued until the demise of anomodonts in the Late Triassic (AR 17), whereby only the advanced cynodonts (i.e., the right tail of the distribution) remained. As with the measurement-based results, subclades encompassing mammals typically show signifi-

TABLE 2. Results of Spearman rank correlation tests for lower jaw measurements versus inferred phylogenetic position in clade ranks (CR). All taxa except "pelycosaurs" and "non-theriodonts" represent clades.  $ns^* p = 0.0601$ 

		Area	DC	Oorsal	Ver	/entral	Core	Coronoid		DI
Subgroup	Rho	d	Rho	d	Rho	d	Rho	d	Rho	d
Synapsida	0.89	<0.0001	0.54	<0.0001	0.78	<0.0001	0.64	<0.0001	0.82	<0.0001
''Pelycosaurs''	0.05	ns	-0.34	us	-0.03		0.22		0.17	us
Therapsida	0.83	<0.0001	0.57	< 0.0001	0.78		0.49		0.74	< 0.0001
"Non-theriodonts"	0.68	<0.0001	-0.38	0.0060	0.13		0.44		0.37	0.0099
Theriodontia	0.88	<0.0001	0.87	< 0.0001	0.90		0.25		0.91	< 0.0001
Cynodontia	0.56	<0.0001	0.81	0.0004	0.48	0.0237	-0.09	ns	0.67	0.0055
Probainognathia	0.75	ns	0.84	ns*	0.00		-0.81		0.21	su

cant correlations between AR and the degree of similarity to mammals, whereas sidebranches often do not (Table 4).

Figure 7B compares the degree of mammal mandibular similarity against each taxon's CR, with the inferred primitive condition at each point highlighted. An increasingly mammal-like lower jaw is expected to correlate with higher CRs, given that some of the features used in this analysis have been proposed as synapomorphies diagnosing higher-level synapsid clades. An interesting result is the relatively low degree of divergence (i.e., range of values) from the presumed ancestral condition at each CR (filled circles). Only anomodonts, and in particular their derived dicynodont subclade (e.g., Fig. 2D), show substantial morphological divergence. Although I attempted to be as exhaustive as possible in my character selection, doubtless additional characters could be discovered and affect this low degree of subclade morphological divergence.

Presuming that the characters used herein are an unbiased sample from the total pool of possible lower-jaw characters, an interesting pattern emerges: except for caseasaurs (CR 1), the inferred primitive condition at each CR lies at the mammal-like (right-hand) tail for non-theriodonts (CRs 2-10), shifts to an intermediate value within gorgonopsians and therocephalians (CRs 11 and 12), and then lies on the non-mammal-like (left-hand) end for cynodonts onward (CRs 13-22). This implies that morphological change within each subclade went from being primarily divergent, within non-cynodonts, to convergent, within cynodonts (but see below). It is also worth noting that discontinuities between the inferred ancestral condition between adjacent CRs could indicate gaps in the fossil record, if synapsid evolution was predominantly monotonic (Sidor and Hopson 1998), or variation in the rate of character acquisition, if taxon sampling probabilities were relatively constant (Sereno et al. 1999).

Figure 7C plots the number of inferred branch points from the root of the cladogram to each terminal taxon (PD) against the degree to which each taxon's lower jaw is similar to that of *Morganucodon*. Taxa positioned near the base of the tree (with low PDs) hover around

Results of Spearman rank correlation tests for lower jaw measurements versus inferred degree of phylogenetic advancement based on the number of branch points passed from the root of the cladogram (PD). All taxa except "pelycosaurs" represent clades. Only first appearances were used in this analysis (mean taxon = 0.0668,  $ns^{**} p = 0.0845$ ,  $ns^{***} p = 0.0738$ . duration = 1.6 intervals). ns\* pTABLE 3.

	A	Area	Dc	Oorsal	Ve	Ventral	Cor	Coronoid		DI
Subgroup	Rho	d								
Synapsida	0.65	<0.0001	0.15	ns	0.38	0.0002	0.48	<0.0001	0.44	<0.0001
"Pelycosaurs"	0.01	us	-0.13	su	-0.17	su	0.44	su	0.07	ns
Therapsida	0.39	0.0016	0.04	ns	0.17	ns	0.23	0.0512	0.07	ns
Dinocephalia	-0.55	ns	-0.21	ns	-0.13	su	0.77	0.0210	0.34	ns
Anomodontia	-0.13	ns	-0.06	su	-0.37	ns	-0.13	su	-0.25	ns
Theriodontia	0.87	< 0.0001	0.82	< 0.0001	0.84	< 0.0001	0.50	0.0031	0.00	< 0.0001
Gorgonopsidae	0.48	ns	0.29	ns	-0.31	su	0.08	ns	0.26	ns
Therocephalia	0.19	ns	0.38	su	-0.10	ns	0.17	su	0.73	ns***
Cynodontia	0.85	0.0003	69.0	0.0027	0.63	0.0029	0.36	ns	0.27	0.0371
Cynognathia	0.57	us	0.17	ns	0.21	ns	0.38	su	0.46	ns
Tritylodontidae	-1.00	ns	1.00	ns	0.50	su	-0.11	su	-1.00	ns
Probainognathia	0.82	ns*	0.77	ns**	0.45	ns	-0.62	su	0.89	ns

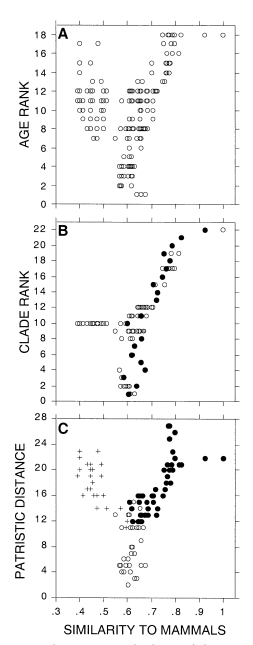


FIGURE 7. Plots comparing the degree of phenetic similarity with the early mammal *Morganucodon* for each taxon against its stratigraphic position (A) and phylogenetic position (B, C). Points highlighted in (B) correspond to the same taxa as in Figure 6B, except that *Anomocephalus* (Modesto et al. 1999) was used in place of *Patranomodon*. See Table 4 for details of correlation statistics.

a mammal mandibular similarity of 60%, whereas anomodonts and nonmammalian theriodonts expand this range by roughly 20% in negative and positive directions, respec-

TABLE 4. Results of Spearman rank correlation tests comparing the overall similarity of each taxon's lower jaw
with that of the early mammal Morganucodon, to stratigraphic position (AR) and two measures of phylogenetic
position (CR and PDR). All taxa except "pelycosaurs" represent clades. The clade rank comparison can only be
made for those taxa encompassing mammals. $ns^* p = 0.0953$ .

	Age	rank	Cla	de rank	Patristi	c distance
Subgroup	Rho	р	Rho	р	Rho	р
Synapsida	0.43	< 0.0001	0.64	< 0.0001	0.16	ns
"Pelycosaurs"	-0.218	ns	0.20	ns	0.04	ns
Therapsida	0.49	< 0.0001	0.77	< 0.0001	0.13	ns
Dinocephalia	-0.19	ns	_	_	-0.55	0.0087
Anomodontia	-0.33	ns	_	_	-0.37	ns
Theriodontia	0.80	< 0.0001	0.93	< 0.0001	0.83	< 0.0001
Gorgonopsidae	-0.33	ns	_	_	-0.09	ns
Therocephalia	-0.27	ns	_	_	-0.33	ns
Cynodontia	0.63	0.0026	0.81	0.0001	0.69	0.0009
Cynognathia	0.18	ns	_	_	0.46	ns
Tritylodontidae	-0.10	ns	_	_	-0.41	ns
Probainognathia	0.63	ns*	0.87	0.0211	0.87	0.0221

tively. Thus, prior to the early mammal *Sinoconodon*, the lower jaws of advanced nonmammalian cynodonts such as *Probainognathus* maintained substantial differences from those of early mammals. This plot most clearly depicts the Y-shaped pattern hinted at in several other graphs (compare Figs. 6A,C and 7A,C), where anomodonts and theriodonts morphologically diverge from one another after an early phase of nondirectionality within more basal synapsids. The gap between the branches of the Y is due to the relatively low diversity and short temporal duration of the clades with intermediate similarity values (viz. gorgonopsians and therocephalians).

# **Scaling Patterns**

Synapsids diversified into an impressive array of body sizes and presumed ecologies during the Permian and Triassic. However, the possibility that changes in body size were important factors in the reduction of the postdentary bones has received scant attention in the literature. Instead, most analyses have focused on the detailed morphology of several exemplar taxa assumed to be phylogenetically close to the line leading toward mammals (Allin 1975).

Figure 8A compares dentary area and total jaw length for the 160 synapsids with both measurements (Appendix 5). A line with a slope of two indicates isometry in this case because an area is being plotted against a linear measurement. A reduced major axis regres-

sion (RMA) indicates that synapsids as a whole conform to this expectation (slope =  $2.031 \pm 0.101$ ). Furthermore, various synapsid subgroups show either near isometry or slight positive allometry (Table 5).

Figure 8B plots postdentary area against total jaw length for 154 fossil synapsids. In contrast to the dentary area results, the RMA regression for Synapsida is significantly greater than isometry (slope =  $2.617 \pm 0.180$ ). However, this apparent allometry is due to the mixing of two regressions. When synapsids are subdivided into cynodonts and non-cynodont components, both of these groups show a relationship between postdentary area and jaw length that is indistinguishable from isometry (slopes of 2.174  $\pm$  0.184 and 2.108  $\pm$ 0.180, respectively). However, the cynodont regression is offset below that of all other synapsids, indicating that this group had a proportionally more gracile postdentary region. See Table 6 for complete results.

## **Step-Size Analysis**

The analyses presented above show a complicated pattern of results but do not address the underlying mechanisms by which trends could develop. One such mechanism could be a bias in step size (Fisher 1986; McShea 1994; Wagner 2000b). For example, even if dentary increases and decreases were equally likely to occur during the course of synapsid evolution, if increases were twice the magnitude of decreases, then the dentary would be predicted

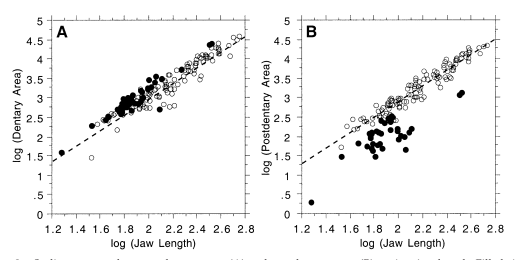


FIGURE 8. Scaling patterns between dentary area (A) and postdentary area (B) against jaw length. Filled circles denote cynodonts, whereas open circles denote non-cynodont synapsids. Dashed line indicates isometric scaling (slope = 2). Regressions for most synapsid subgroups either are indistinguishable from isometry or show slight positive allometry. See Tables 5 and 6 for regression results.

to enlarge over time. The possibility that unequal degrees of mammalian versus nonmammalian morphological change brought about a trend for an increasingly mammalian lower jaw is examined below.

Methods.—To address the hypothesis of step-size bias with the discrete character data, I used MacClade (Maddison and Maddison 1992) to contrast the number of internodal character state changes leading to each pair of sister taxa at every CR along the spine of the cladogram in Figure 3. For example, at CR 7 (Therapsida) between three and ten character-state changes (depending on optimization) oc-

Table 5. Log-log regressions of dentary area and jaw length in synapsids. Regressions for all of these synapsid subgroups are either isometric (slope = 2) or slightly positively allometric. The slope and intercept are based on reduced major-axis regression. The 95% confidence limit (CL) and correlation coefficient ( $r^2$ ) are estimates based on the results of simple linear regressions. All slopes are significantly different from zero for at least p < 0.05 level.

Subgroup	Slope ± CL	Intercept	$r^2$	n
Synapsida	$2.031 \pm 0.101$	-0.991	0.900	160
"Noncynodont"	$2.116 \pm 0.100$	-1.221	0.909	120
"Pelycosaurs"	$2.313 \pm 0.105$	-1.918	0.903	23
Biarmosuchia	$2.203 \pm 0.120$	-1.523	0.928	7
Dinocephalia	$1.915 \pm 0.110$	-0.695	0.917	13
Anomodontia	$2.040 \pm 0.102$	-0.940	0.981	52
Gorgonopsidae	$2.193 \pm 0.113$	-1.386	0.979	10
Therocephalia	$2.304 \pm 0.108$	-1.594	0.936	15
Cynodontia	$2.249 \pm 0.103$	-1.266	0.901	40

cur along the branch to *Tetraceratops*, whereas six to nine occur along the branch to CR 8. If evolution along the mammalian line typically produced larger than average changes, then we might expect the number of character-state changes between CRs to be consistently larger than between CR nodes and side branches.

These comparisons are based on the premise that morphological changes occurring between consecutive nodes on this cladogram produce increased similarity to mammals (because they are synapomorphic), whereas changes accumulated on the side-branches (i.e., toward the individual terminal taxa)

Table 6. Log-log regressions of postdentary area and jaw length in synapsids. Regressions for all of these synapsid subgroups are either isometric (slope = 2) or slightly positively allometric. The slope and intercept are based on reduced major axis regression. The 95% confidence limit (CL) and correlation coefficient ( $r^2$ ) are estimates based on the results of simple linear regressions. All slopes are significantly different from zero for at least p < 0.05 level.

Subgroup	Slope ± CL	Intercept	$r^2$	п
Synapsida	$2.617 \pm 0.180$	-2.540	0.818	154
"Noncynodont"	$2.108 \pm 0.180$	-1.325	0.926	118
"Pelycosaurs"	$2.198 \pm 0.188$	-1.521	0.875	23
Biarmosuchia	$2.472 \pm 0.215$	-2.107	0.810	7
Dinocephalia	$1.900 \pm 0.198$	-0.806	0.854	12
Anomodontia	$2.046 \pm 0.182$	-1.127	0.969	52
Gorgonopsidae	$2.232 \pm 0.202$	-1.701	0.941	10
Therocephalia	$2.068 \pm 0.195$	-1.430	0.878	14
Cynodontia	$2.177\pm0.184$	-2.086	0.643	36

TABLE 7. Mammalian versus nonmammalian step-size contrasts. The 21 clade ranks (CR) are derived from the cladograms in Figures 3 and 4. "Mammal" refers to the number of character state changes occurring between consecutive CRs (e.g., between CR 1 and CR 2). "Nonmammal" refers to the number of character state changes occurring between a particular node and the terminal taxon attaching to it (e.g., between the node at CR 7 and Tetraceratops). Minimum and maximum numbers of character state changes were computed in MacClade (Maddison and Maddison, 1993). In order to calculate the maximum character state changes, polychotomies in Figure 5 were arbitrarily resolved. Results of a Wilcoxon signed rank test indicate no significant difference in the sign or magnitude of mammalian versus nonmammalian changes for either the minimum or maximum of discrete changes (p = 0.7405).

CR	Mammal Char <sub>min/max</sub>	Non-mammal Char <sub>min/max</sub>
1	0/2	0/2
2	1/1	1/3
3	2/6	3/6
4	3/6	0/4
5	1/5	0/2
6	1/6	1/6
7	3/10	6/9
8	0/1	1/3
9	1/5	2/4
10	1/1	4/7
11	2/4	6/8
12	3/6	1/3
13	0/0	1/2
14	1/5	3/6
15	1/2	0/0
16	4/5	2/2
17	2/4	0/1
18	2/6	1/2
19	1/4	0/3
20	4/10	2/6
21	5/10	2/7

should produce increased dissimilarity. One shortcoming of this type of analysis is that it only uses the first possible comparison at each CR (i.e., changes occurring on the first internode in either direction) and thereby disregards subsequent (i.e., more deeply nested) changes within each sister clade.

Results.—Wilcoxon sign-rank tests found no significant difference between the number of character changes in mammalian and non-mammalian directions (Table 7, Fig. 9). This result was the same regardless of whether minimum or maximum numbers of character-state changes were used.

## Discussion

The hypothesis that disparate groups of synapsids independently acquired mammallike characteristics has a long pedigree (Olson 1944, 1959, 1962; Romer 1965; Simpson 1959). However, some examples of "convergence" probably arose from the taxonomic framework accepted at that time—one that recognized paraphyletic and polyphyletic grades of organization (Hopson 1994). The application of cladistic methods to synapsid systematics has dispelled some cases of morphologic homoplasy as unnecessary when viewed from the standpoint of total character congruence (Hopson and Barghusen 1986; Rowe 1986; Kemp 1988b). Here, I have readdressed the oft-noted observation that the size of the dentary increased during the course of synapsid evolution. Both quantitative and discrete data indicate that a lower jaw of increasingly mammalian cast was a prevalent feature of premammalian synapsid evolution (Fig. 10), but finer scales of phylogenetic resolution yield more complex patterns.

In Theriodontia and its subordinate clades that encompass mammals, the pattern of both measurements and similarity values is suggestive of a driven trend (in that both the minimum and maximum values steadily increase). Thus, the measurement results accord well with Allin's (1975) hypothesis that reduction of the postdentary bones improved highfrequency hearing in these taxa and was therefore selectively advantageous. However, corresponding directionality is not apparent within the "side-branch" clades (Tables 1-4), which suggests that a common driving force is doubtful. In the most extreme case, anomodonts show the exact opposite trend: decreasing dentary size and increasing their lower jaw's distinctiveness from that of mammals. This suggests either that high-frequency hearing was not important to anomodonts or that selection for this feature was not exclusively molding mandibular evolution in this group.

The specialized structure of the anomodont mandible is an interesting exception to another result of these analyses—the relative scarcity of divergent lower-jaw morphologies among synapsid side-branches. Although there are certainly some features that are autapomorphic for the clades that do not encompass mammals (e.g., the extremely slender dentary of varanopseids, the near-vertical

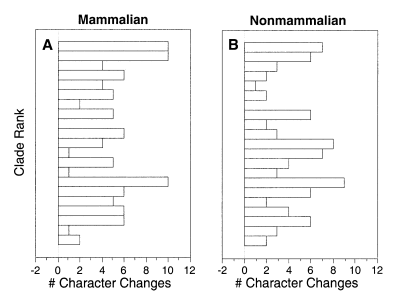


FIGURE 9. Step-size comparisons contrasting the maximum and minimum number of character-state changes along the phylogenetic trajectory toward mammals (mammalian) versus those made toward side-branches (nonmammalian). The vertical axis represents clade rank (CR). The final 21 comparisons were made for discrete characters. A and B display the maximum number of discrete character changes in mammalian and nonmammalian directions, respectively. See text and Table 7 for details.

ridge on the reflected lamina of gorgonopsians, or the elongate angular process of the dentary in some advanced cynognathians), no one synapsid subgroup amasses more than a few such specializations, except for the anomodonts. Importantly, this lack of mandibular autapomorphy indicates that the acquisition of only a few mammalian characters would be sufficient to drive an apparent trend toward a mammal-like jaw.

Disruptive Patterns.—The Y-shaped pattern of dentary size and mammalian similarity (Figs. 6, 7) that emerged from several analyses is strikingly similar to that of disruptive selection within modern populations (i.e., when selection acts against intermediates and favors morphological extremes). Foote (1993) showed that blastoids exhibit a similar disruptive pattern, but he suggested that if a bias against intermediates were present, then its explanation would require investigation at finer scales. In the case of synapsids, the lack of intermediates is due to the early extinction of gorgonopsians and therocephalians, compared with the relatively long-lived anomodont and cynodont clades.

Combining Methods.—Both stratigraphyand phylogeny-based methods have been

used to examine patterns of morphological change in fossil lineages (Gingerich 1976; Benton 1990; McShea 1994; Wagner 1996). Importantly, the potential weaknesses of either approach might be overcome by using both methods in a study. For example, if cladistic estimates of synapsid phylogeny have been led astray by rampant homoplasy, then the stratigraphic distribution of the taxa may yield a more informative measure of relatively primitive and derived taxa. Conversely, if the fossil record does not accurately portray the first appearances of synapsids because preservation rates vary widely, then phylogenetic measures might yield a more reliable sequence of branching events. The concordant results found in this study suggest that the synapsid fossil record is relatively well sampled and that the cladistic hypothesis of synapsid relationships presented here is in line with the distribution of fossil finds (Sidor and Hopson 1998).

## Conclusions

The prevalence of homoplasy in synapsid evolution has been a hotly contested topic (Kemp 1988a; Rowe 1988; Hopson 1991a). Hopson (1994: p. 212) suggested that although

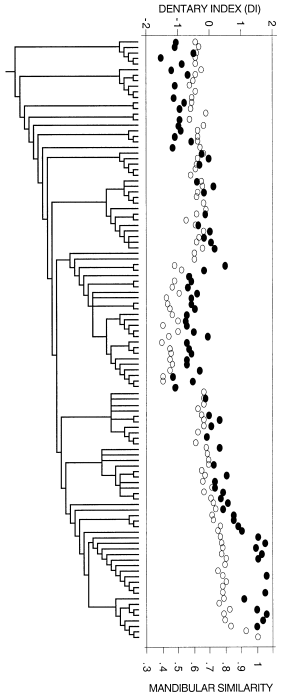


FIGURE 10. Changes in dentary size and mammal mandibular similarity plotted against a cladogram of synapsid relationships. DI values are denoted by filled ellipses (left axis) and similarity values by hollow ellipses (right axis).

"[t]he polyphyletic origin of mammals is no longer a tenable hypothesis... this is not to say that parallelism and convergence have not been significant aspects of pre-mammalian synapsid evolution."

The present study supports the following main conclusions:

- 1. The lack of a well-supported phylogeny has exaggerated previous estimates of morphological convergence or parallelism in the synapsid fossil record. The hypothesis of multiple therapsid groups arising independently from pelycosaur-grade ancestors (e.g., Olson 1962; Boonstra 1972) necessitated rampant homoplasy and are now considered untenable (Rubidge and Sidor 2001). Certain lower jaw characteristics and proportions are better viewed as broadly distributed synapomorphies indicative of common ancestry.
- 2. Despite the striking differences between the lower jaws of basal synapsids (i.e., "pelycosaur") and mammals, mandibular evolution within synapsids was predominantly conservative. Except for dicynodont anomodonts, most therapsid subclades do not acquire substantial morphological novelty in their lower jaw structure.
- 3. The area of the dentary and postdentary regions scales either isometrically or with slight positive allometry when compared with jaw length. This suggests that bodysize trends are not sufficient to drive the reduction of the postdentary bones in synapsid evolution. Importantly, when compared with other synapsid subgroups, cynodonts are characterized by smaller-thanpredicted postdentary areas.
- 4. Selection acting to decrease the size of the postdentary bones, and thereby improving high-frequency hearing, is still the most tenable mechanism for the evolution of the mammalian lower jaw (Allin 1975; Allin and Hopson 1992). However, this mechanism by itself has difficulty explaining the converse pattern in anomodont therapsids (i.e., decreasing the size of the dentary and increasing the size of the postdentary bones).

These conclusions, in combination with

those of recent studies on long-term patterns of epipodial (Hopson 1995) and cranial (Sidor 2001) evolution, suggest that morphological trends within synapsids should be reinvestigated within a quantitative and phylogenetic framework.

# Acknowledgments

This project was part of my dissertation research at the University of Chicago. I thank my committee, M. Foote, R. Reisz, P. Sereno, P. Wagner, and especially my advisor, J. Hopson, for comments on previous versions of the manuscript. M. Carrano's review of the first draft helped considerably. I also acknowledge the support and assistance provided by my fellow graduate students, including H. Larsson, J. Wilson, R. Blob, J. Socha, R. O'Keefe, D. Croft, P. Magwene, F. Lando, A. Beck, J. Conrad, E. Love, and J. Tsao. Data for this project were gathered on trips to several domestic and foreign research collections. For their help in this critical aspect of my dissertation research, I sincerely thank the following curators and museum personnel: N. Hotton, M. Brett-Surman, G. Wilson, K. Padian, M. Norell, G. Gaffney, F. Jenkins, A. Crompton, C. Schaff, M.-A. Turner, H.-D. Sues, R. Reisz, S. Modesto, D. Scott, J. Bolt, O. Rieppel, B. Rubidge, M. Raath, C. Gow, F. Thackeray, H. Fourie, J. Neveling, J. Welman, R. Smith, S. Kaal, A. Milner, S. Chapman, A. Friday, R. Symonds, T. Kemp, M. Wills, D. Sigogneau-Russell, D. Dutheil, M. Maisch, P. Wellnhofer, D. Unwin, M. Ivachnenko, N. Kalandadze, A. Kurkin, V. Bulanov, and V. Golubev. I also acknowledge the following granting organizations and institutions for supporting my research: National Science Foundation Doctoral Dissertation Improvement Grant (NSF DEB-9801342), Hinds Fund (University of Chicago), the American Museum of Natural History, and the Richard Estes Memorial Award from the Society of Vertebrate Paleontology. This manuscript profited from reviews by T. Kemp, S. Wing, J. Lillegraven, and D. McShea.

# Literature Cited

- Allin, E. F. 1975. Evolution of the mammalian middle ear. Journal of Morphology 147:403–438.
- Allin, E. F., and J. A. Hopson. 1992. Evolution of the auditory system in Synapsida ("mammal-like reptiles" and primitive

- mammals) as seen in the fossil record. Pp. 587–614 *in* D. B. Webster, R. R. Fay, and A. N. Popper, eds. The evolutionary biology of hearing. Springer, New York.
- Angielczyk, K. D. 2001. Preliminary phylogenetic analysis and stratigraphic congruence of the dicynodont anomodonts (Synapsida: Therapsida). Palaeontologia Africana 37:53–79.
- Battail, B. 1991. Les Cynodontes (Reptilia, Therapsida): une phylogénie. Bulletin du Muséum National d'Histoire Naturelle 13:17–105.
- Benton, M. J. 1990. Reptiles. Pp. 279–300 in K. J. McNamara, ed. Evolutionary trends. Belhaven, London.
- Berman, D. S., R. R. Reisz, J. R. Bolt, and D. Scott. 1995. The cranial anatomy and relationships of the synapsid *Varanosaurus* (Eupelycosauria: Ophiacodontidae) from the Early Permian of Texas and Oklahoma. Annals of the Carnegie Museum 64: 99–133.
- Bonaparte, J. F. 1962. Descripción del cráneo y mandibula de *Exaeretodon frenguellii*, Cabrera, y su comparación con Diademodontidae, Tritylodontidae y los cinodontes sudamericanos. Publicaciones del Museo Municipal de Ciencias Naturales de Mar del Plata "Lorenzo Scaglia" 1:135–202.
- Boonstra, L. D. 1972. Discard the names Theriodontia and Anomodontia: a new classification of the Therapsida. Annals of the South African Museum 59:315–338.
- Brinkman, D. 1981. The structure and relationships of the dromasaurs (Reptilia: Therapsida). Breviora 465:1–34.
- Clark, J. M., and J. A. Hopson. 1985. Distinctive mammal-like reptile from Mexico and its bearing on the phylogeny of the Tritylodontidae. Nature 315:398–400.
- Cluver, M. A., and G. M. King. 1983. A reassessment of the relationships of the Permian Dicynodontia (Reptilia, Therapsida) and a new classification of dicynodonts. Annals of the South African Museum 91:195–273.
- Conrad, J., and C. Sidor. 2001. Re-evaluation of *Tetraceratops insignis* (Synapsida: Sphenacodontia). Journal of Vertebrate Paleontology 21:42A.
- Crompton, A. W. 1963. The evolution of the mammalian jaw. Evolution 17:431–439.
- ——. 1972. Postcanine occlusion in cynodonts and tritylodontids. Bulletin of the British Museum (Natural History) Geology 21:30–71.
- Crompton, A. W., and F. Ellenberger. 1957. On a new cynodont from the Molteno Beds and the origin of the tritylodontids. Annals of the South African Museum 44:1–14.
- Crompton, A. W., and N. Hotton. 1967. Functional morphology of the masticatory apparatus of two dicynodonts (Reptilia, Therapsida). Postilla 109:1–51.
- Crompton, A. W., and F. A. Jenkins Jr. 1973. Mammals from reptiles: a review of mammalian origins. Annual Review of Earth and Planetary Sciences 1:131–155.
- Crompton, A. W., and Z. Luo. 1993. Relationship of the Liassic mammals Sinoconodon, Morganucodon oehleri, and Dinnetherium. Pp. 30–44 in F. S. Szalay, M. J. Novacek, and M. C. McKenna, eds. Mammal phylogeny: Mesozoic differentiation, multituberculates, monotremes, early therians, and marsupials. Springer, New York.
- Efremov, I. A. 1940. *Ulemosaurus svijagensis* Riab.—ein Dincephale aus den Ablagerurgen des Perm der UdSSR. Nova Acta Leopoldina 9:155–205.
- Fisher, D. C. 1986. Progress in organismal design. Pp. 99–117 inD. M. Raup and D. Jablonski, eds. Patterns and processes in the history of life. Springer, Berlin.
- Foote, M. 1993. Contributions of individual taxa to overall morphological variation. Paleobiology 19:403–419.
- ——. 1996. Evolutionary patterns in the fossil record. Evolution 50:1–11.
- Gauthier, J., A. Kluge, and T. Rowe. 1988. Amniote phylogeny and the importance of fossils. Cladistics 4:105–209.

- Gingerich, P. D. 1976. Paleontology and phylogeny: patterns of evolution at the species level in Early Tertiary mammals. American Journal of Science 276:1–28.
- Grine, F. E. 1997. Dinocephalians are not anomodonts. Journal of Vertebrate Paleontology 17:177–183.
- Hopson, J. A. 1965. Tritylodontid therapsids from Yunnan and the cranial morphology of *Bienotherium*. Ph.D. dissertation. University of Chicago, Chicago.
- ——. 1966. The origin of the mammalian middle ear. American Zoologist 6:437–450.
- ——. 1987. The mammal-like reptiles: a study of transitional fossils. American Biology Teacher 49:16–26.
- ——. 1991a. Convergence in mammals, tritheledonts, and tritylodonts. Journal of Vertebrate Paleontology 11:36A.
- ——. 1991b. Systematics of the nonmammalian Synapsida and implications for patterns of evolution in synapsids. Pp. 635– 693 in H.-P. Schultze and L. Trueb, eds. Origins of the higher groups of tetrapods: controversy and consensus. Comstock, Ithaca.
- ——. 1994. Synapsid evolution and the radiation of non-eutherian mammals. Pp. 190–219 in D. B. Prothero and R. M. Schoch, eds. Major features of vertebrate evolution. Paleontological Society, Knoxville, Tenn.
- ——. 1995. Patterns of evolution in the manus and pes of nonmammalian therapsids. Journal of Vertebrate Paleontology 15:615–639.
- ——. 1999. Therapsids. Pp. 1256–1266 *in* R. Singer, ed. Encyclopedia of paleontology. Fitzroy Dearborn, Chicago.
- Hopson, J. A., and H. Barghusen. 1986. An analysis of therapsid relationships. Pp. 83–106 in N. Hotton, P. D. MacLean, J. J. Roth, and E. C. Roth, eds. The ecology and biology of the mammal-like reptiles. Smithsonian Institution Press, Washington, D.C.
- Hopson, J. A., and J. W. Kitching. 1972. A revised classification of cynodonts (Reptilia; Therapsida). Palaeontologia Africana 14:71–85.
- ——. 2001. A probainognathian cynodont from South Africa and the phylogeny of nonmammalian cynodonts. Bulletin of the Museum of Comparative Zoology 156:3–35.
- Huelsenbeck, J. P. 1991. When are fossils better than extant taxa in phylogenetic analysis? Systematic Zoology 40:458–469.
- Ivachnenko, M. F., V. K. Golubev, Y. M. Gubin, N. N. Kalandadze, I. V. Novikov, A. G. Sennikov, and A. S. Rautian. 1997. Permskie i Triasovye tetrapody Vostochnoi Evropy (Permian and Triassic tetrapods of Eastern Europe). Geos, Moscow.
- Kemp, T. S. 1982. Mammal-like reptiles and the origin of mammals. Academic Press, New York.
- ——. 1983. The relationships of mammals. Zoological Journal of the Linnean Society 77:353–384.
- ——. 1988a. Haemothermia or Archosauria? The interrelationships of mammals, birds and crocodiles. Zoological Journal of the Linnean Society 92:67–104.
- 1988b. Interrelationships of the Synapsida. *In* M. J. Benton, ed. The phylogeny and classification of the tetrapods, Vol.
  2. Mammals. Systematics Association Special Volume 35B:1–22. Clarendon, Oxford.
- Kermack, D. M., F. Mussett, and H. W. Rigney. 1973. The lower jaw of Morganucodon. Zoological Journal of the Linnean Society 53:87–175.
- King, G. M. 1988. Anomodontia. Pp. 1–174 in P. Wellnhofer, ed. Encyclopedia of paleoherpetology, Part 17C. Gustav Fischer, Stuttgart.
- King, G. M., B. W. Oelofsen, and B. S. Rubidge. 1989. The evolution of the dicynodont feeding system. Zoological Journal of the Linnean Society 96:185–211.
- Kitching, J. W. 1977. The distribution of the Karroo vertebrate fauna, Memoir 1. Bernard Price Institute for Palaeontological Research, University of the Witwatersrand, Johannesburg.

- Laurin, M. 1993. Anatomy and relationships of *Haptodus garnettensis*, a Pennsylvanian synapsid from Kansas. Journal of Vertebrate Paleontology 13:200–229.
- Laurin, M., and R. R. Reisz. 1990. Tetraceratops is the oldest known therapsid. Nature 345:249–250.
- ——. 1996. The osteology and relationships of *Tetraceratops insignis*, the oldest known therapsid. Journal of Vertebrate Paleontology 16:95–102.
- Lucas, S. G., and A. B. Heckert. 2001. Olson's gap: a global hiatus in the record of Middle Permian tetrapods. Journal of Vertebrate Paleontology 21:75A.
- Lucas, S. G., and Z. Luo. 1993. Adelobasileus from the Upper Triassic of West Texas: the oldest mammal. Journal of Vertebrate Paleontology 13:309–334.
- Luo, Z. 1994. Sister-group relationships of mammals and transformations of diagnostic mammalian characters. Pp. 98–128 in
   N. C. Fraser and H.-D. Sues, eds. In the shadow of the dinosaurs: early Mesozoic tetrapods. Cambridge University Press, New York.
- Luo, Z., and A. W. Crompton. 1994. Transformation of the quadrate (incus) through the transition from non-mammalian cynodonts to mammals. Journal of Vertebrate Paleontology 14: 341–374.
- Luo, Z., and X.-C. Wu. 1995. Correlations of vertebrate assemblage of the Lower Lufeng Formation, Yunnan, China. Pp. 83–88 *in* A. Sun and Y. Wang, eds. Sixth symposium of Mesozoic terrestrial ecosystems and biota, short papers. China Ocean Press, Beijing.
- Luo, Z., Z. Kielan-Jaworoska, and R. L. Cifelli. 2002. In quest for a phylogeny of Mesozoic mammals. Acta Palaeontologica Polonica 47:1–78.
- Maddison, W. P., and D. R. Maddison. 1992. MacClade 3.0.8a: interactive analysis of phylogeny and character evolution. Sinauer, Sunderland, Mass.
- Martinez, R. N., and C. A. Forster. 1996. The skull of *Probelesodon sanjuanensis*, Sp. Nov., from the Late Triassic Ischigualasto Formation of Argentina. Journal of Vertebrate Paleontology 16:285–291.
- Martinez, R. N., C. L. May, and C. A. Forster. 1996. A new carnivorous cynodont from the Ischigualasto Formation (Late Triassic, Argentina), with comments on eucynodont phylogeny. Journal of Vertebrate Paleontology 16:271–284.
- McShea, D. W. 1994. Mechanisms of large-scale evolutionary trends. Evolution 48:1747–1763.
- Modesto, S. P. 1995. The skull of the herbivorous synapsid *Edaphosaurus boanerges* from the Lower Permian of Texas. Palaeontology 38:213–239.
- Modesto, S. P., B. S. Rubidge, and J. Welman. 1999. The most basal anomodont therapsid and the primacy of Gondwana in the evolution of the anomodonts. Proceedings of the Royal Society of London B 266:331–337.
- Modesto, S., C. A. Sidor, B. S. Rubidge, and J. Welman. 2001. A second varanopseid skull from the Upper Permian of South Africa: implications for Late Permian 'pelycosaur' evolution. Lethaia 34:249–259.
- Olson, E. C. 1944. The origin of mammals based on the cranial morphology of the therapsid suborders. Geological Society of America Special Paper 55:1–136.
- ——. 1959. The evolution of mammalian characters. Evolution 13:344–353.
- ——. 1962. Late Permian terrestrial vertebrates, U.S.A. and U.S.S.R. Transactions of the American Philosophical Society 52:1–224.
- Parrington, F. R. 1955. On the cranial anatomy of some gorgonopsids and the synapsid middle ear. Proceedings of the Zoological Society of London 125:1–40.
- Reichert, C. 1837. Über die Visceralbögen der Wirbeltiere im Allgemeinen und deren Metamorphosen bei den Vögeln und

- Säugetieren. Müllers Archiv für Anatomie, Physiologie und wissenschaftliche Medizin 1837:120–222.
- Reisz, R. R. 1986. Pelycosauria. Pp. 1–102 in P. Wellnhofer, ed. Encyclopedia of paleoherpetology, Part 17A. Gustav Fischer, Stuttgart.
- ——. 1988. Two small reptiles from a Late Pennsylvanian quarry near Hamilton, Kansas. Kansas Geological Survey Guidebook 6:189–194.
- Reisz, R. R., and D. W. Dilkes. 1995. A small varanopseid (Synapsida: Eupelycosauria) from the uppermost Carboniferous (Upper Pennsylvanian) of Kansas. Journal of Vertebrate Paleontology 15:49A.
- ——. 2003. Archaeovenator hamiltoni, a new varanopid (Synapsida: Eupelycosauria) from the Upper Carboniferous of Kansas. Canadian Journal of Earth Sciences 40:667–668.
- Reisz, R. R., and H.-D. Sues. 2000. Herbivory in Late Paleozoic and Triassic terrestrial vertebrates. Pp. 9–41 in H.-D. Sues, ed. Evolution of herbivory in terrestrial vertebrates. Cambridge University Press, New York.
- Reisz, R. R., D. S. Berman, and D. Scott. 1992. The cranial anatomy and relationships of Secodontosaurus, an unusual mammal-like reptile (Synapsida: Sphenacodontidae) from the Early Permian of Texas. Zoological Journal of the Linnean Society 104:127–184.
- Reisz, R. R., D. W. Dilkes, and D. S. Berman. 1998. Anatomy and relationships of *Elliotsmithia longiceps* Broom, a small synapsid (Eupelycosauria: Varanopseidae) from the Late Permian of South Africa. Journal of Vertebrate Paleontology 18:602–611.
- Renaut, A. J. 2000. A re-evaluation of the cranial morphology and taxonomy of the Triassic dicynodont genus *Kannemeyeria*. Ph.D. dissertation. University of the Witwatersrand, Johannesburg.
- Romer, A. S. 1965. Possible polyphylety of the vertebrate classes. Zoologische Jahrbuch 92:143–156.
- ——. 1969. The Chañares (Argentina) Triassic reptile fauna V. A new chiniquodontid cynodont, *Probelesodon lewisi*: cynodont ancestry. Breviora 333:1–24.
- ——. 1970. The Chañares (Argentina) Triassic reptile fauna VI. A chiniquodontid cynodont with an incipient squamosaldentary jaw articulation. Breviora 344:1–18.
- Romer, A. S., and L. I. Price. 1940. Review of the Pelycosauria. Geological Society of America Special Paper 28:1–538.
- Rowe, T. 1986. Osteological diagnosis of Mammalia, L. 1758, and its relationship to extinct Synapsida. Ph.D. dissertation. University of California, Berkeley.
- ———. 1988. Definition, diagnosis, and the origin of Mammalia. Journal of Vertebrate Paleontology 8:241–264.
- ——. 1993. Phylogenetic systematics and the early history of mammals. Pp. 129–145 in F. S. Szalay, M. J. Novacek, and M. C. McKenna, eds. Mammal phylogeny: Mesozoic differentiation, multituberculates, monotremes, early therians, and marsupials. Springer, New York.
- Rowe, T., and J. Gauthier. 1992. Ancestry, paleontology, and the definition of the name Mammalia. Systematic Biology 41:372– 378.
- Rubidge, B. S. 1991. A new primitive dinocephalian mammallike reptile from the Permian of southern Africa. Palaeontology 34:547–559.
- ——. 1994. Australosyodon, the first primitive anteosaurid dinocephalian from the Upper Permian of Gondwana. Palaeontology 37:579–594.

- ——. 1995. Biostratigraphy of the Beaufort Group (Karoo Series), South Africa. Government Printer, Pretoria.
- Rubidge, B. S., and J. A. Hopson. 1996. A primitive anomodont therapsid from the base of the Beaufort Group (Upper Permian) of South Africa. Zoological Journal of the Linnean Society 117:117–139.
- Rubidge, B. S., and C. A. Sidor. 2001. Evolutionary patterns among Permo-Triassic therapsids. Annual Review of Ecology and Systematics 32:449–480.
- Rubidge, B. S., and J. A. van den Heever. 1997. Morphology and systematic position of the dinocephalian *Styracocephalus pla-tyrhinus*. Lethaia 30:157–168.
- Rybczynski, N. 2000. Cranial anatomy and phylogenetic position of *Suminia getmanovi*, a basal anomodont (Amniota: Therapsida) from the Late Permian of Eastern Europe. Zoological Journal of the Linnean Society 130:329–373.
- Sereno, P. C., A. L. Beck, D. B. Dutheil, H. C. E. Larsson, G. H. Lyon, B. Moussa, R. W. Sadleir, C. A. Sidor, D. J. Varricchio, G. P. Wilson, and J. A. Wilson. 1999. Cretaceous sauropods from the Sahara and the uneven rate of skeletal evolution among dinosaurs. Science 286:1342–1347.
- Sidor, C. A. 2000. Evolutionary trends and relationships within the Synapsida. Ph.D. dissertation. University of Chicago, Chicago.
- ——. 2001. Simplification as a trend in synapsid cranial evolution. Evolution 55:1419–1442.
- Sidor, C. A., and J. A. Hopson. 1998. Ghost lineages and "mammalness": assessing the temporal pattern of character acquisition in the Synapsida. Paleobiology 24:254–273.
- Sidor, C. A., and R. M. H. Smith. In press. A new galesaurid (Therapsida: Cynodontia) from the Early Triassic of South Africa. Palaeontology.
- Sigogneau, D. 1970. Révision systématique des gorgonopsiens Sud-Africains. Cahiers de Paléontologie, Paris.
- Simpson, G. G. 1959. Mesozoic mammals and the polyphyletic origin of mammals. Evolution 13:405–414.
- ——. 1960. Diagnosis of the classes Reptilia and Mammalia. Evolution 14:388–392.
- Sokal, R. R., and P. H. A. Sneath. 1963. Principles of numerical taxonomy. W. H. Freeman, San Francisco.
- Sues, H.-D. 1985. The relationships of the Tritylodontidae (Synapsida). Zoological Journal of the Linnean Society 85:205–217.
- Sun, A. 1984. Skull morphology of the tritylodont genus *Bienotheroides* of Sichuan. Scientia Sinica 27:970–984.
- Tatarinov, L. P., and E. N. Matchenko. 1999. A find of an aberrant tritylodont (Reptilia, Cynodontia) in the Lower Cretaceous of the Kemerovo region. Paleontological Journal 33:422–428.
- van den Heever, J. A. 1994. The cranial anatomy of the early Therocephalia (Amniota: Therapsida). Universiteit van Stellenbosch Annale 1994/1:1–59.
- Wagner, P. J. 1996. Contrasting the underlying patterns of active trends in morphologic evolution. Evolution 50:990–1007.
- ——. 2000a. The quality of the fossil record and the accuracy of phylogenetic inferences about sampling and diversity. Systematic Biology 49:65–86.
- 2000b. Rate heterogeneity in shell character evolution among lophospiroid gastropods. Paleobiology 27:290–310.
- Watson, D. M. S., and A. S. Romer. 1956. A classification of therapsid reptiles. Bulletin of the Museum of Comparative Zoology 114:37–89.

## Higher-Level Relationships

Sidor and Hopson (1998) presented the most recent analysis of higher-level synapsid relationships, and their analysis was chosen as the baseline cladogram for this compilation. Not surprisingly, Sidor and Hopson's results largely conform to the topology previously put forward by Hopson and Barghusen (1986) and Hopson (1991b, 1994). Where they overlap, the Sidor and Hopson analysis corroborates the results of Reisz (1986) and Laurin (1993), with regard to the basal pelycosaur-grade taxa. The remainder of higher-level pelycosaur relationships was adopted from Reisz (1986). Higher cynodont relationships are discussed below.

Rowe (1986, 1988) and Gauthier et al. (1988) proposed a phylogenetic arrangement for synapsids that differs from that adopted here in terms of (1) the position of varanopseids relative to caseasaurs and ophiacodontids, (2) the position of anomodonts (dicynodonts in their terminology) relative to gorgonopsians, and (3) higher cynodont relationships. The position of varanopseids has since been resolved by Reisz et al. (1998), although Modesto et al. (2001) dispute the position of Elliotsmithia (a viewpoint upheld here). The position of Anomodontia within the therapsid tree has been surprisingly labile (Gauthier et al. 1988; King 1988; Rubidge and Sidor 2001). Indeed, the grouping that combines anomodonts and theriodonts (Neotherapsida sensu Hopson 1999) was supported by relatively few characters in Sidor and Hopson's (1998) analysis. Grine (1997) has demonstrated that at least one of the proposed anomodont sister-groupings, that with dinocephalians (Watson and Romer 1956; King 1988), is unsupported. I have chosen to maintain anomodonts in the position advocated by Sidor and Hopson (1998), but it is important to note that advancing them one clade rank (as proposed by Gauthier et al. 1988) has a negligible effect on the overall results presented in the text.

Nonmammalian cynodont systematics still lack consensus, but most phylogenetic hypotheses fall into one of three camps. The first supports the traditional view that tritylodontids are derived from a lineage of gomphodont cynodonts with transversely expanded cheek teeth, whereas mammals arose from a lineage with a persistently sectorial dentition (Crompton and Ellenberger 1957; Crompton 1972; Hopson and Kitching 1972, 2001; Sues 1985; Hopson 1991b). The second contends that most gomphodonts (i.e., diademodontids, trirachodontids, and traversodonts) form a clade, but that tritylodontids are distinct and more closely related to mammals (and possibly tritheledonts) (Kemp 1982, 1983). The final permutation completely dissolves the tooth-type dichotomy and intersperses gomphodont with non-gomphodont taxa as successive mammal outgroups (Rowe 1993). Battail (1991) groups gomphodonts and tritylodontids but suggests that mammals evolved from a Thrinaxodon-grade ancestor, a hypothesis unlike that any of the previous workers.

Hopson and Kitching (2001) have provided the most recent investigation of the higher cynodont problem. Their results support the traditional hypothesis and also have the benefit of the most extensive taxon sampling—an important factor in accurately reconstructing phylogenies (Huelsenbeck 1991; Wagner 2000a). Thus, it stands as the most current understanding of cynodont phylogeny, and I have used Hopson and Kitching's (2001) cladogram here. Placing tritylodontids as the sister taxon to mammals has little effect on the overall pattern of results.

### Lower-Level Relationships

Less effort has been devoted to understanding lower-(i.e., genus-) level relationships within nonmammalian synapsids. Among pelycosaur-grade synapsids, I have followed the lower-level relationships put forward by Modesto et al. (2001), Berman et al. (1995), Modesto (1995), Laurin (1993), and Reisz et al. (1992) for varanopseids, ophiacodontids, edaphosaurids, *Haptodus*, and sphenacodontoids, respectively. "Hamilton Form" refers to KUVP 12483, a specimen described by Reisz (1988) as a small reptile, and then later by Reisz and Dilkes (1995, 2003) as the most primitive varanopseid. *Tetraceratops* was positioned between sphenacodontids and traditional therapsids (Laurin and Reisz 1996), although the poor preservation of the sole, holotypic specimen makes a confident acceptance of this placement problematic (Conrad and Sidor 2001).

Hopson and Barghusen (1986) were the first to propose that taxa such as *Biarmosuchus*, *Hipposaurus*, and *Ictidorhinus* were among the phylogenetically most primitive therapsids. They did not, however, attempt to resolve the relationships among these taxa, and no work has been published since. The cladistic topology used here is based on the results of Rubidge and Sidor (unpublished data).

I based dinocephalian interrelationships on the published analyses of Hopson and Barghusen (1986), Rubidge (1991, 1994), and Rubidge and van den Heever (1997). Although *Ulemosaurus, Criocephalus*, and *Moschops* have been proposed to form a clade more derived than *Tapinocaninus* (Rubidge 1991), their precise relationships have not yet been proposed in print. The topology for these tapinocephalids, depicted in Figure 4, is based on my unpublished cladistic analyses.

The discovery in recent years of new, primitive anomodonts has prompted several investigations into the early evolution of this clade (Rubidge and Hopson 1996; Modesto et al. 1999; Rybczynski 2000). Within anomodonts, dicynodont interrelationships have been examined by Cluver and King (1983), King (1988), and Angielczyk (2001). For the purpose of this analysis, I have used the basal anomodont topology of Modesto et al. (1999), which recognizes a clade of Russian venyukovioids (e.g., Ulemica + Suminia), but is otherwise similar to that of Rubidge and Hopson (1996) and, earlier, Hopson and Barghusen (1986). Among dicynodonts, most analyses have yielded fairly congruent results; conflicting opinions as to the position of Diictodon/Robertia, Pristerodon, and Endothiodon have been represented by an unresolved basal trichotomy in Figure 4.

Gorgonopsian interrelationships have yet to be examined within the cladistic paradigm. The topology used here is based on the evolutionary scheme put forth by Sigogneau (1970). Similarly, except for van den Heever's (1994) work on relatively primitive forms, knowledge of therocephalian systematics has been at a standstill since Hopson and Barghusen's (1986) initial treatment. The relationships used here therefore come directly from these two sources.

As discussed above, the cynodont relationships used here are based on the results of Hopson and Kitching's (2001) most recent analysis. However, several taxa included here were not included in Hopson's analysis. *Cynosaurus* and a new genus housed in the South African Museum (SAM-PK-K9954) are grouped with *Galesaurus* on the basis on their possession of galesaurid synapomorphies (Sidor and Smith in press). Tritylodontid interrelationships are based on the findings of Clark and Hopson (1985), with further resolution provided by Luo and Wu (1995).

The following is a list of the characters and character states used in this analysis. Following the last character state for each character is a citation for previous uses of the character in the literature. An asterisk following a citation means that the character definition has been modified or that an additional character state(s) has been added. Citations are in the form: (author: character number), except for those of Hopson and Barghusen (1986), which are (author:table.clade.character number), and King (1988), which are (author:suite.character number). Abbreviations for authors are as follows: B, Berman et al. (1995); GKR, Gauthier et al. (1988); HB, Hopson and Barghusen (1986); K, King (1988); M, Modesto (1995); MRW, Modesto et al. (1999); R, Rowe (1988); SH, Sidor and Hopson (1998); S, Sidor (2001).

- 1. Dentary symphysis: unfused (0), fused (1) (HB:1.6.2; HB: 4.42.2; SH:81; MRW:33; S:25).
- 2. Ventral protuberance on the anterior portion of the dentary: absent (0), present, obscuring symphysis in lateral view (1).
- Depth of anterior ramus of dentary: moderate to deep (0), extremely slender (1).
- Shape of anterior portion of dentary: tapering continuation of posterior regions (0), dorsoventrally deepened compared with posterior portions (1), or beak shaped (2). (HB:2.21.1; SH:79\*)
- 5. Inclination of the anterior portion of the dentary: nearly horizontal (0), tipped anterodorsally producing procumbency (1), or strongly recurved (2).
- 6. Boss on lateral surface of dentary (dentary shelf of King 1988): absent (0), present (1). (K:E.12; MRW:34)
- Large boss positioned halfway along ventral margin of dentary: absent (0), present (1).
- 8. Posterodorsal edge of dentary thickened into laterally overhanging shelf: absent (0), present (1).
- 9. Pit along tooth row formed to receive upper canine: absent (0), present (1).
- Occlusal surface has parallel ridges bounding median groove (longitudinal dentary groove or sulcus of Crompton and Hotton 1967): absent (0), present (1). (K:J.1\*; K:C'.2\*)
- 11. Angle between coronoid eminence/process to long axis of jaw: less than 70 degrees (0), greater than 80 degrees (1).
- 12. Dentary coronoid region: basically flat (0), convex eminence (1), forms distinct freestanding coronoid process (2). (HB: 1.7.1\*; GKR:92\*; SH:80\*; B:61\*)
- Coronoid region dorsal extent: below middle of orbit (0), in upper half of orbit (1), extends above orbit (2). (GKR:97\*; SH:86\*)
- 14. Coronoid process extends posteriorly beyond level of jaw articulation: absent (0), present (1).
- Dentary masseteric fossa: absent (0), present (1). (HB:1.6.7\*; HB:1.11.1\*; GKR:88; SH:82)
- Dentary masseteric fossa extent: high on coronoid region (0), extends to lower border of dentary (1). (HB:4.38.1; SH: 83)
- Dentary possesses a freestanding, posteriorly directed (angular) process along its posterior margin: absent (0), present but small (1), present and elongate (2). (GKR:89\*)
- Dentary articular process: absent (0), present as posterior eminence (1), present as a distinct process (2). (GKR:96\*; SH:87\*)
- 19. Dentary boomerang or banana shaped: absent (0), present (1). (HB:3.25.1)
- Dentary—angular suture runs smoothly anteroventrally (0), or S-shaped (1).
- 21. Dentary tables: absent (0), present (1). (K:E.3\*; K:L.1\*; K: D'.3\*)
- 22. Direction of angular process: posterior (0), ventral (1).
- 23. Splenial: present (0), absent (1). (S:26\*)

- 24. Splenial symphysis: unfused (0), fused (1). (S:26\*)
- 25. Mandibular symphysis: dentary and splenial (0), dentary only (1), (B:58)
- 26. Splenial appearance: visible near symphysis in lateral view (0), visible between dentary and angular in lateral view (1), or not visible in lateral view (2). (B:59\*; SH:90\*)
- Splenial with triangular dorsal process in symphysial region: absent (0), present (1).
- 28. Splenial pinches out dentary anteriorly at symphysis: absent (0), present (1). (M:22\*; B:59\*)
- 29. Angular distinct (0), or at least partially fused to adjacent postdentary bones (1). (S:27\*)
- 30. Ventral margin of angular: rounded (0), keeled (1). (GKR: 98: B:60)
- 31. Angular reflected lamina: absent (0), present (1). (HB:1.1.1; GKR:102; SH:95)
- 32. Angular reflected lamina shape: flat and platelike (0), or ringlike (1). (GKR:103\*; SH:99\*)
- 33. Angular reflected lamina posterior emargination: short (0), long with free dorsal margin (1), long but lacking free dorsal margin (2). (HB:1.2.5\*; HB:1.8.9\*; SH:96\*)
- 34. Angular reflected lamina with pattern of radiating ridges and grooves: absent (0), present (1). (SH:98\*)
- 35. Angular reflected lamina with near vertical ridge: absent (0), present (1). (HB:1.8.10)
- 36. Lateral surface of angular with a thickened region (boss) adjacent to dentary: absent (0), present (1). (HB:2.13.1)
- Ventral margins of angular and dentary confluent (0) or angular offset dorsally from that of dentary (1). (HB:1.9.9\*; GKR:93\*; SH:84)
- 38. Reflected lamina of angular lies far anterior to jaw articulation: absent (0), present (1). (HB:1.8.8; SH:97\*)
- 39. Posteroventral margin of lateral surface of angular supports large boss: absent (0), present (1).
- Angular anterior ramus extends anteriorly to level of jaw symphysis: absent (0), present (1).
- 41. Surangular: present (0), absent (1). (S:28\*)
- 42. Surangular distinct (0), or at least partially fused to adjacent postdentary bones (1). (S:28\*)
- 43. Surangular abuts (0) or dorsally overrides (1) the dentary along dorsal margin of lower jaw.
- 44. Lateral surface of surangular with fossa for lateral slip of adductor mandibulae externus: absent (0), present (1).
- 45. Surangular vertical lamina lateral to articular: absent (0), or present (1). (MRW:37)
- Surangular participation in craniomandibular joint: absent (0), present as articular recess or fossa (1), or as condylar process (2). (HB:4.42.1\*)
- 47. Surangular position: exposed laterally (0), confined medially (1).
- 48. Articular distinct (0), or at least partially fused to prearticular (1). (S:30 $^{*}$ )
- 49. Articular-prearticular complex at least partially fused to surangular: absent (0), present (1). (S:29\*; S:30\*)
- 50. Dorsolateral surface of articular forms lateral shelf: absent (0), present (1).
- 51. Articular bone with prominent posterolateral process (dorsal process of Parrington 1955), which contacts the posterior surface of the quadrate above the lateral condyle: absent (0), present (1). (HB:1.8.6)
- 52. Level of jaw articulation: set below dentary tooth row (0), roughly at level of dentary tooth row (1), or above dentary tooth row (2). (M:15\*)
- 53. Shape of articular glenoid: elongate oblique troughs (0), screw-shaped hinge (1), elongate flat plate (2), elongate convex curve (3), non-screw-shaped hinge (4), longitudinal troughs (5). (HB:2.21.8\*; SH:101\*)

- 54. Articular surface of lower jaw slopes steeply posteroventrally: absent (0), present (1). (HB:1.6.3)
- Articular glenoid expanded anteroposteriorly: absent (0), present (1). (HB:2.21.8)
- 56. Prearticular with (0), or without (1) lateral exposure posteriorly. (MRW:39)
- 57. Prearticular teeth: absent (0), present (1).
- 58. Anterior coronoid: present (0), absent (1). (HB:1.6.10\*; K: A.4; GKR:100; SH:89; MRW:48; S:31\*)
- 59. Anterior coronoid teeth: absent (0), present (1).
- 60. Posterior coronoid: present (0), absent (1). (S:32\*)
- 61. Posterior coronoid exposed on lateral surface of lower jaw: absent (0), present (1).
- 62. Posterior coronoid dentition: absent (0), present (1).
- Posterior coronoid mediolaterally thickened: absent (0), present (1). (SH:92)
- 64. Lateral mandibular fenestra: absent (0), present (1), present as a small foramen (2). (HB:1.6.4\*; K:A.6\*; GKR:87\*, SH:64\*; SH:94\*; B:57\*; MRW:36\*)
- Lateral mandibular fenestra bordered by dentary, angular, and surangular (0), or dentary and angular (1). (HB:1.6.4\*; HB:3.24.2\*; SH:64\*)
- 66. Dentary-squamosal: not in contact (0), articulating (1) (HB: 4.51.1; GKR:91)
- 67. Position of postdentary bones: broadly exposed laterally (0), narrow and in medial groove (1). (SH:88\*)
- 68. As indicated by wear facets, mandibular movement: primarily orthal (0), with medial component (1), with strong longitudinal component (2). (R:79\*; K:E.6; MRW:40\*)
- 69. Foramen present between prearticular, angular, and splen-

- ial on medial surface of lower jaw: absent (0), present (1), or present between angular and prearticular (2).
- 70. Number of dentary teeth: zero (0), one to ten (1), 11 to 20 (2), 21 to 30 (3), 31 or greater (4). (SH:112\*)
- 71. Dentary tooth row: absent (0), single (1), double or multiple (2).
- 72. Dentary tooth row set at lateral margin of dentary (0), or more medially (1).
- Terminal lower tooth: absent (0), present and subequal in size to remaining teeth/precanines (1), present and enlarged compared with remaining teeth/precanines (2). (HB:2.18.1\*; HB:4.50.2\*; SH:107\*)
- 74. Position of anteriormost dentary tooth: terminal (0), non-terminal (1), absent (2).
- 75. Number of lower incisors: zero (0), one or two (1), three (2), four (3), five or greater (4), undefined owing to lack of lower caniniform (5). (SH:103\*)
- Lower canine: absent (0), incisiform/postcaniniform (1), present (2). (HB:1.6.6\*; HB:2.17.2\*)
- Posteriormost dentary tooth: visible in lateral view (0), or obscured by dentary coronoid process (1).
- 78. Medial surface of lower jaw with large crushing plates: absent (0), present (1). (M:19\*)
- 79. Tooth roots: undivided (0), divided (1). (HB:4.51.3; GKR: 118\*; SH:117\*)
- 80. Postcanine tooth implantation: subthecodont (0), thecodont (1). (SH:121)
- 81. Lower canine passes external to lateral border of maxilla: absent (0), present (1). (HB:2.15.2)
- 82. Lower postcanine tooth row (and/or incisors) passes medial to lower canine: absent (0), present (1). (HB:2.15.4)



The following is the data matrix for the qualitative characters of lower jaw shape. Information regarding characters and character states is provided in Appendix 2. Taxa arranged as in Figure 4. "?" denotes missing data and "n" denotes that a character is inapplicable for that taxon. Electronic copies of this matrix are available upon request.

\_\_\_\_\_\_ 1111111112 222222223 1234567890 Taxon 1234567890 1234567890 Eothyris parkeyi 000000000 00000n0000 0n00000000 Casea broilii 0000000000 00000n0000 0n000000000 000000000 00000n0000 0n00000000 Casea rutena Cotylorhynchus romeri 0000000000 00000n0000 0n00000100 Ennatosaurus tecton 0000000000 00000n0000 0n00000100 00000n0000 Hamilton Form 0010000000 0n00120002 Mesenosaurus romeri 0010000000 00000n0000 0n00020000 Elliotsmithia longiceps ?010000000 00000n000? 0n??????00 Aerosaurus wellesi 0010000000 00000n0000 0n00120000 Varanops brevirostris 0010000000 00000n0000 Varanodon agilis 0010000000 00000n0000 0n00000000 Varanosaurus acutirostris 0010000000 00000n0000 0n00120001 Ophiacodon spp. 000000000 00000n0000 0n00020001 Ianthasaurus hardestii 0????00000 01000n0000 0n??????01 0000000000 01000n0000 0n00000101 Edaphosaurus spp. 000000000 01000n0000 0n00000001 Haptodus garnettensis Sphenacodon spp. 0001000000 01000n0000 0n0000001 0000000000 01000n0000 0n00100001 Secodontosaurus obtusidens 0001000000 01000n0000 0n00000001 Dimetrodon spp. Tetraceratops insignis 0000000000 01000n0000 0n0000000? Biarmosuchus tener 0001000000 01000n0000 0n00020001 01000n0100 0n00020001 0001000100 Hipposaurus spp. 0n00020001 Rubidgina augusticeps 0001000100 01000n0100 0001000100 01000n0001 0n00020001 CGP 1/61 Proburnetia viatkensis 0000001100 01000n0001 0n000200?1 01000n0000 Anteosaurus magnificus 0001000200 0n00220001 Titanophoneus potens 0001000000 01000n0000 0n00020001 Doliosauriscus yanshinovi 0001000000 01000n?001 022222201 Australosyodon nyaphuli 0001000?00 01000n0?0? 0n00020001 Syodon efremovi 0001000000 01000n0000 0n00020001 01000n0001 Styracocephalus platyrhynchus 0001000?00 0n0011000? Estemmenosuchus uralensis 0001000000 01000n0000 0n00000001 Estemmenosuchus mirabilis 0001011000 01000n0001 0n000??001 Jonkeria ingens 0001000000 01000n0001 0n00020001 Tapinocaninus pamelae 000000000 01000n0001 0n00020001 Criocephalus vanderbyli 000000000 01000n0001 0n00020001 Ulemosaurus svijagensis 000000000 01000n0001 0n00020001 Moschops spp. 000000000 01000n0001 0n00020001 Anomocephalus africanus 3300000000 0100100?00 0n?????00? Patronomodon nyaphulii 0n0?00??01 3303300000 00000n0000 Ulemica spp. 0100011010 0100100001 0n000n1001 1 0000010000 0n0002?001 Suminia getmanovi 0100100001 Galeops whaitsi 000000000 01000n0000 0n0002??01 Eodicynodon oosthuizeni 0002210001 01000n0001 1n01001101 Eodicynodon oelofseni 1002210000 01000n0000 0n010011?1 Pristerodon sp. 1002210001 01000n0001 0n01001101 Endothiodon uniseries 1002211001 0000100001 0n01001101 1002210000 01000n0001 1n01001001 Diictodon spp. Robertia broomiana 1002210000 01000n0000 1n010????1 Myosaurus gracilis 1002210000 00000n0001 0n02022001 Emydops sp. 1002210001 00000n000? 0n010????1 Cistecephalus microrhinus 1002210001 01000n0000 0n01001101

Appendix 3. Extended.

					====
3333333334	444444445	555555556	6666666667	777777778	88
1234567890	1234567890	1234567890	1234567890	1234567890	12
0nnnn00n00	00000?00n0	0150010??0	1000n000?3	1010510000	nı
0nnnn00n00	00000000n0	005001???0	?000n000?2	1010510000	nı
0nnnn00n00	00000?00n0	00500100?0	??00n000??	101??10000	?1
0nnnn00n00	00100000n0	00500101n0	1000n00022	1010510000	nı
0nnnn00n00	0000000n0	0050011000	0000n00022	1010510000	nı
0nnnn00n00	0000000330	01?0010??0	1000n000?4	1010510000	nı
0nnnn00n00	00100000110	01500101.0	0000n00003	1010510000	nı
0nnnn00n00	00100000110	?1???1????	???0n000?2	1010510000	nı
0nnnn00n00		0150010000	1?00n00023	1010510000	
	00100000n0				nı
0nnnn00n00	00?0?00??0	0150010???	???0n000??	???????00?	?:
0nnnn00n00	00100000n0	0150010??0	1000n00003	1010510000	n
0nnnn00n00	00100000n0	0150010??0	1001000024	1010510000	n
0nnnn $0$ 0n $0$ 0	00100000n0	0150010000	1001000024	1010510000	n
0nnnn00n00	00000000n0	0150010???	???0n000??	10????0000	?:
0nnnn00n00	00000000n0	0050111010	0100n00213	1010510100	0:
0nnnn00n00	00000000n0	0210010000	0100n00003	1010120000	0:
1000000000	00000000n0	0110000000	1000n00003	1010120000	0
1000000000	00100000n0	0210000000	1000n00004	1010510000	n
1000000000	00000000n0	0010000000	1000n00003	1010120000	0
?????00?10	00100000n0	020001????	???0n000?2	1010510000	n:
1011000000	0000300330	10100????0	0000n000?2	1010320000	0
1011000000	0000100??0	1010000???	???0n000?2	1010320000	0
1011000000	00001000n0	10100001n0	0000n000?2	1010220000	0
1011?00000	00?0?00??0	10100001n0	?000n000?2	3 1010?20000	0
		?0100001110		1010:20000	0
10?1000000	00001?00n?		???0n000??		
101?010000	0330330333	?0???????	???0n000??	1020320000	0
1010010000	00101000n0	00100001n0	0??0n00012	1020320000	0
1????10??0	0010??0???	333333330	0??0n000?2	1020320000	0
1010000000	00?0?000n0	00100?0??0	?000n000?2	1020220000	0
1010000000	00101000n0	00100101n0	0??0n00012	1020320000	0
3333300300	0000??0???	3033330330	0000n00??2	10?0320000	?
1010000000	0000100???	0030033330	0??0n00023	1120320000	1
1010000000	0000030333	?0100??1n?	???0n000?3	1120320000	1
1010000000	00001000n0	00100101n0	0000n00023	1120320001	1
1010000000	00001000n?	00?00?01n1	nnn0n000?2	1020320001	1
1010000000	00001000n	00100001n1	nnn0n00022	1020320001	n
1010000000	00101000n0	00100101n1	nnn0n00012	1020510001	n
1010000000	00001000n0	00100101n0	0000n000?2	1020510001	n
3333333303	0001?000??	3033333333	???1?000?1	1010510?01	n
101100010?	0??01000n0	00100001n0	?001100001	10???10001	n
1011000000	00011000n1	00210101n0	0001100002	2120511000	n
1011000000	00011000n0	00311001n0	0001100212	1120510000	n
					n
101??00100	00001000n0	00311001n1	nnn1000201	10n1510001	n
1011000101	00001?0100	00311101n1	nnn10002?1	11n1510000	n
101?000100	0??01?0??1	00311?0???	???1?002?1	11n1510000	n
1011000100	0000000101	00311101n1	nnn1000201	11n1510000	n
10110?0101	0010000101	??311001n1	nnn10002?1	11n1511000	n
1011000101	0100000101	00211101 1	2	2	
1011000101	01000001?1	00311101n1	nnn11002?0	0n0200n0nn	n
101100010?	0?00??0??1	00311?????	???00002?1	11n1511000	n
101?00010?	00000001?1	01311?01n1	1 nnn1?00200	0n0200n0nn	n
		00311?????	???11002?1		
101100010?	0??0??01?1			11n1511000	n
1011000101	0000000101	00311101n1	nnn1000200	0n0200n0nn	n
			1		

Appendix 3. Continued.

=======================================		=========	======
	1	1111111112	222222223
Taxon	1234567890	1234567890	1234567890
Cistecephaloides boonstrai	1002210001	01000n0000	0n0?0???01
Kingoria nowacki	1002210001	01000n0000	1n01001101
Oudenodon spp.	1002210001	01000n0001	1n00001101
Pelanomodon sp.	1002210001	01000n0001	1n01001101
Aulacephalodon spp.	1002210001	00000n000?	1n000?1101
Dicynodon spp.	1002210001	00000n0001	1n01001101
Lystrosaurus spp.	1002210001	00000n0000	1n01001101
Tetragonias njalilus	1002210001	00000n0001	1n01010001
Kannemeyeria spp.	1002210001	00000n0001	1n01001101
Placerias gigas	100221000?	00000n0000	1n01001101
Stahleckeria potens	100221000?	00000n0001	1n0100??01
Aelurosaurus felinus	0001000000	02000n0000	0n01000001
Arctognathus spp.	0001000000	02000n0000	0n00000001
Cyonosaurus spp.	0001000000	02000n0001	0n000?0001
Scylacops spp.	0001000?00	02000n000?	0n00000001 0n01000001
Lycaenops spp.	000100000? 0001000000	02000n0000 02000n0000	0n00000001
Prorubidgea spp.	000100000	02000n0000	0n00000001
Dinogorgon spp. Rubidgea spp.	0001100001	02000n0000	0n00000011
Aloposaurus spp.	0001000000	02000n0000	0n01000001
Gorgonops spp.	000100000?	02000n0000	0n00000001
corgenops spp.	000200000	02000110000	2
Hofmeyria atavus	000000000	02100n0010	0n00020001
Moschorhinus kitchingi	0001000000	02000n0000	0n00020001
Olivieria parringtoni	0001000000	02000n0000	0n00020001
Theriognathus spp.	000000000	02000n0010	0n00020001
Ictidosuchoides spp.	000000000	02100n0000	0n00020001
Viatkosuchus sumini	000000000	02000n0000	0n000?0001
Bauria cynops	000000000	02100n0000	0n00020001
Lycosuchus vandereti	0001000000	02000n0000	0n00000001
Glanosuchus macrops	0001000000	02000n0000	0n00000001
Ptomalestes avidus	0001000000	02000n0000	0n00020001
Dvinia prima	0000000000	0200100100	0n00020001
Procynosuchus delaharpeae	0000100000	0200100100	0n00000001
Cynosaurus suppostus	0001000000	02??110100	0n00020001
SAM-PK-K9954	000000000	0200110100	0n00020001
Galesaurus planiceps	000000000	0210110100	0n00020001
Thrinaxodon liorhinus	0001000000	0210110000	0n00020001
Cynognathus spp.	1000000000	0210110100	0n01020001
Diademodon spp.	1000000000	0210110100	0n0102000?
Trirachodon spp.	1000000000	0220111100	0001020001
Massetognathus spp.	1000000000	0221110200	0n01020001
Exaeretodon frenquelli	1000000000	0220112200	00011200?1
Tritylodon longaevus	0000100000	12?0110200	0n00020011
Oligokyphus spp.	0000100000	1220110200	0n000200??
Bienotherium elegans	0000100000	1220111200	0000020011
Kayentatherium wellesi	0000100000	1220111200	000?020001
Bocatherium mexicanum	0000110000	1220111200	0055550055
Bienotheroides wanhsienensis	0000100000	1220112200	000002000?
Lumkuia fuzzi	1000000000	0210110100	0n01020001
Ecteninion lunesis	100??0000?	??2111??00	0?01??00??
Probelesodon lewisi Probelesodon sanjuanensis	1000000000	0220111200	0001020001
Probelesodon sanjuanensis Probainognathus jenseni	100??00000 1000100000	0220112?00 0220110200	00???????0? 0n01020001
Pachygenelus monus	0000000000	0210110200	0n00020001
Sinoconodon spp.	0000000000	0210110200	011n1nnn??
Morganucodon spp.	0000100000	0210111200	011n1nnn01
- <del></del>			

Appendix 3. Extended.

333333334	444444445	555555556	6666666667	777777778	8
1234567890	1234567890	1234567890	1234567890	1234567890	
101100010?	0100??0111	?0311101n1	nnn11002?0	0n0200n0nn	
1011000100	0110000111	00311?01n1	nnn0n00200	0n0200n0nn	
1011000100	0000010101	00311101n1	nnn1000200	0n0200n0nn	
1017000101	0000070101	00311701m1 00311?01m1	nnn10002?0	0n0200n0nn	
101?000100	3333333333	00311?01n1 00311?01n1	nnn1?002?0	0n0200n0nn	
				0n0200n0nn	
1011000101	00000001?1	00311101n1	nnn11002?0		
1011000101	0000010101	00311101n1	nnn11002?0	0n0200n0nn	
1010000101	00000?0111	00311?01n1	nnn1000200	0n0200n0nn	
1011000101	0100010111	00311001n1	nnn1000200	0n0200n0nn	
1010000001	0000001?1	00311101n1	nnn1000200	0n0200n0nn	
101?000001	01000?0111	00311?01n1	nnn1?002?0	0n0200n0nn	
1021100100	0000330003	?0100?01n0	0000n00001	1010320000	
1021100100	00000?0100	10100001n0	0000n00011	1010320000	
1021100100	0000000330	?0100?01n?	???0n000??	1010320000	
1021100100	00000003555	3333330333	???0n000??	101032?000	
1021100100	0000030000	?0???1????	0??0n000??	1010320000	
1021100100	0000000330	1010000???	???0n000??	???????0??	
1021100100	??00??010?	?????0????	???0n000??	???????0??	
1021100100	000000100	?0?00101n0	0000n00011	1010320000	
1021100100	0000?00???	??????0???	???0n000??	?????????	
1021100100	0000001??	1??00?0??0	0000n00011	1010?20000	
1011001000	00001?0000	01100101n?	0??10000??	1010?20000	
		00100101n? 00100101n0		1070?20000	
1011001000	0000100001	?0100101n0	00011000??	1010320000	
1011001000	00001?0000		0??10000?1		
1011001000	0000100000	00100001n0	0001100001	101022n00n	
1011001000	0000100???	02100001n?	0??10000?2	101032?000	
1011001000	00001?0??0	0010010???	???10000??	?????2?000	
1011001000	0.503.30333	01?00?0???	0??10000?2	1010220000 3	
1011001???	0000??000?	3333330333	0??0n000?2	1010220000	
1011001000	0000110000	01100001n0	0000n00002	1010220000	
1011001000	0000100000	01100001n0	0000n0002	1010220000	
1????01100	0000100000	01100001n0	0002100002	1010420000	
1?11?01100	0000100000	00100001n0	0002100002	1010320000	
1011001100	0000170770	?110??0???	0??0n000??	1010220000	
1011001?00	00001?0000	01?00?0???	???0n000?2	1010220000	
1011001100	0000100000	01100101n0	0000n?0002	1010220000	
1111001100	0000100000	01100101n0	0000n00002	1010220000	
1111001100	0000020000	01100101n0	0000n01002	1010220000	
1111001100	00000201?0	01100101n0	0000n01202	1010221001	
1111001100	0000020100	02100101n0	0000n01002 2	1010221001	
111??01100	0000021100	01200101n0	0000n01202	1110221001	
111?001100	0000021100	02200101n0	00?0n01202	1120221001	
1110001100	0100001110	02400101n0	0010n012?1	1120111011	
?????01?00	0100001110	02400?01n?	???0n012?1	1120211011	
111??01100	0100001110	02400?01n0	0010n01201	1120111011	
1110001100	0100001110	02400101n0	0010n01201	1120111011	
?????01?00	??????1???	?2????01n0	0010n31201	1120111011	
111??01100	0100001110	0240?101n0	0010n01201	1120111011	
111?001100	000000001	02?0010???	???0n01001	1010220000	
????0????0	000002010?	00?00?????	???0n010??	???????0??	
1110001100	0000021101	02400101n0	0000n01002	1010220000	
????001?00	000002011?	02?001????	???0n010??	3333333030	
1110001100	0100021111	01400101n0	0000n01002	1010220000	
1110001100	01000n1110	02400101n0	0010n11001	1110121000	
??????1??0	?????????	?2400??1n0	0010n111?1	1010320001	
1110001100	0000001100	02400101n0	0010n11102	1010320011	

Appendix 4

graphic and phylogenetic data used elsewhere. Taxa are arranged as in Figure 4. Average ratios are provided where a taxon is represented by multiple specimens preserving two or more measurements. FA = First Appearance (in age ranks); LA = Last Appearance (in age ranks); CR = Clade Rank; PD = Patristic Distance from the root of the cladogram shown in Figure 4; D/J = ratio of distance to surangular suture to jaw length; V/J = ratio of distance to angular suture to jaw length; C/R = ratio of coronoid height to jaw length; Area = area of dentary relative to entire lower jaw area; DI = cumulative Dentary Index (average of the previous four ratios when each is z-transformed);  $S_{SM} = Simple \ matching \ coefficient for every taxon when the data in Appendix 3 are compared with the early mammal Morganucodon. The$ following geologic formations or biozones correspond to age rank values: 1 = Late Pennsylvanian (Stanton Formation); 2 = Moran, Cutler Abo, and Putnam Formations; Chañares Formation; 17 = Ischigualasto, Santa Maria, Los Rastros, and Chinle Formations; 18 = Lower Lufeng (dull purplish beds and dark red beds), Kayenta, La The following table provides a summary of the results for the measurement-based analyses, the similarity to Morganucodon metric used in Figure 7, and the strati-3 = Admiral and Belle Plains Formations; 4 = Clyde, Arroyo, Vale, Hennesey, and Choza Formations; 5 = San Angelo, Flower Pot, and Chickasha Formations; 6 = Mezen nathus Assemblage Zone; 10 = Tropidostonna Assemblage Zone; 11 = Cistecephalus Assemblage Zone and Kotelnich Assemblage; 12 = Dicynodon Assemblage Zone and Assemblage; 7 = Eodicynodon Assemblage Zone, Middle Dinocephalian Complex; 8 = Tapinocephalus Assemblage Zone, Upper Dinocephalian Complex; 9 = Pristerog-Sokolov Assemblage; 13 = Lystrosaurus Assemblage Zone; 14 = Cynognathus Subzone A; 15 = Cynognathus Subzone B + C, Manda and N'twere formations; 16 Boca, Upper Elliot, and Clarens Formations, and the Shazimiao Series. Electronic copies of this matrix are available upon request.

Taxon	FA	LA	CR	PD	D/J	٢/٦	C/J	Area	DI	$_{ m SSM}$
Eothyris parkeyi	ĸ	m	П	2	.73	.41	0.	.38	.03	.60
Casea broilii	4	4	Н	М	0.618	0.365	0.106	0.544	-1.085	0.629
Casea rutena	4	4	_	4	.75	.38	۲.	.50	.48	9
Cotylorhynchus romeri	4	5	_	2	.50	.35	.08	.47	1.53	.60
Ennatosaurus tecton	9	7	7	2	.74	.31	.18	.39	.83	.60
Hamilton Form	Н	Т	7	m	.71	.42	.05	.35	.19	.63
Mesenosaurus romeri	9	9	7	5	. 74	.53	.06	. 52	9.	9.
Elliotsmithia longiceps	<b>∞</b>	œ	7	5			-			. 59
Aerosaurus wellesi	7	7	7	Ŋ	.72	.46	0.097	. 28	-1.086	. 56
Varanops brevirostris	4	4	7	9	. 79	.39		.44		.60
Varanodon agilis	2	2	2	9	. 73	.41	.05	.38	۲.	. 58
Varanosaurus acutirostris	m	4	m	4	9.	9.	.01	0.578	-0.771	. 58
Ophiacodon spp.	7	4	m	4	.72	.45	.04	.48	.93	.57
Ianthasaurus hardestii	Н	1	4	2	.64	.30	.07			.67
Edaphosaurus spp.	7	4	4	2	. 69	.36	. 18	.37	σ.	. 56
Haptodus garnettensis	Н	7	Ŋ	വ	. 78	.30	.12	.39	σ.	.65
Sphenacodon spp.	7	7	9	7	.71	ς.	.18	.40	σ.	.61
Secodontosaurus obtusidens	М	4	9	∞	. 63	.44	.08	.46	0.	9.
Dimetrodon spp.	7	Ŋ	9	∞	0.701	0.441	0.213	0.446	-0.571	.61
Tetraceratops insignis	4	4	7	7	.60	.41	.12	.44	۲.	.63
Biarmosuchus tener	9	ω	ω	თ	.81	. 53	.13	.57	.20	.65
Hipposaurus sp.	∞	0	∞		.84	. 52	.21	.49	00.	. 62
Rubidgina augusticeps	12	12	∞		.76	. 56	.14	. 52	٣.	.61
CGP 1/61	σ	0	∞		-			-		.60
Proburnetia viatkensis	11	11	∞	12	.77	.55	0.206			.5
Anteosaurus magnificus	∞	œ	σ		0.794	0.478				.60
Titanophoneus potens	∞	∞	0		.73	.51	0.193	0.626	-0.153	9.
Doliosauriscus yanshinovi	∞	œ	6							.65
Australosyodon nyaphuli	7	7	0				0.198		ω.	9.
Syodon efremovi	∞	œ	0		.85	.47	0.2	. 68	0.164	.64

Appendix 4. Continued.

										FIG C
Styracocephalus platyrhynchus Estemmenosuchus uralensis	8 7	8 7	9	11	0.809	0.653	12.2	0.572	-0.117	0.667
	7	7	n 01		.	:	0.199	. 60	!	. 54
Jonkeria ingens	∞ ι	∞ ι	o (		. 75	.45	.16	.60	.35	.60
Tapinocaninus pamelae Orioganhaliis mandarhiili	- α	- α	ത ര		) ) (	. 5 L		2 . 2 .		გა
ciiocephaius vahueibyii Ulemosaurus svijadensis	0 00	0 00	n 01		0.749	0.552	. 25	. 50	0	. 60
Moschops capensis	, ∞	- ∞	, O		. 79	. 52	30	. 53	. 2	. 64
Anomocephalus africanus	œ	∞	10		-		-	1		. 59
Patronomodon nyaphulii	<b>-</b> 0	7	10		0	1	1	. 53	1	. 6
Ulemica spp.	ω -	ے ہے ر	7 -			0.633	0.285	し. 6g 7/		٠
	-l 00		) C		. 6		1 6	5 C	0.10	
Sarcers miarcs Eodicynodon oosthuizeni	) C	) C	10		. 59	. 3	30	49	0.55	.47
	7		10		.54	.48	.21	. 55	0.68	4.
Pristerodon sp.	ω	12	10		.70	.44	.18	.62	0.38	٠.
Endothiodon uniseries	o 1		10		.71	.36	. 11	. 70	0.56	.41
Diictodon spp.	∞ (		10		99.	4.2	.20	. 63	0.43	.44
Kobertia broomiana Myosaurus gradilis			) C		٥.	 	. L 4	5 C	9 4	•
Myosautus graciiis Fmydons sn			) C		# <del>-</del>	T C T				1, 4
us microrhinu	10	11	10		.56	. 4	30	. 50	0.48	. 49
Cistecephaloides boonstrai			10		.72	.47	.25	.62	0.04	43
Kingoria nowacki			10		.65	.39	.15	. 59	0.70	.45
Oudenodon spp.			10		9.	.41	.20	.60	0.55	.44
			0 7		9	ص د	. 7.	. 54	0.63	444
Aulacephalodon spp. Dicynodon spp.			1 C		. 63 66	4 . 2 0	1.4	٠ د ر	0./1	•
Liczniczni zpp. Lystrosaurus spp.			10		.67	. 3	. 19	.57	0.70	.45
Tetragonias njalilus			10		.68	.51	.21	. 56	0.31	.44
Kannemeyeria spp.			10		9.	.34	0.	. 56	1.14	4.
Placerias gigas			10		.64	08.	.12	. 5.	1.08	
stanieckeria potens belinosauris felinus			) T		σ σ	4. 0.	. L &	٠ ر	0.50	4. 7. 7.
			-, -		7.5	, 7.	2.5	. 70	0.1	6.4
Cyonosaurus spp.	10		111		.73	.58	. 22	.51	.12	. 65
Scylacops spp.			11		.73	. 59	.23	1	1	.65
Lycaenops spp.			11		.77	. 54	.20	0.646	0.068	. 65
Prorubidgea spp.			11		.80	. 51	.33	1		.61
Dinogorgon spp.			7		.67	.45		0.631	1	.60
Rubidgea spp.			11		.76	.43	0.231	. 63	-0.078	. 68
$\widetilde{A}loposaurus$ spp.			, ,		.73	.42	-	-		. 62
Gorgonops spp.			T 7		9 1	75.	'	1	,	.65
Hormeyria atavus Mogaboahimma hitahimmi			17		20.5	. 6	0.196	. 6	0.138	ο,
MOSCHOININGS KICCHINGI Olivieria parringtoni	7 1	7 F	7 1		0.00	6.0	7.	0.639	7.7	•
Theriognathus spp.			17		7.00	. 64	. 23	.70	44.	. 65
Ictidosuchoides spp.			12		.78	.64	0.202	.60	0.183	.70
Viatkosnehus sumini	,				1	(	Ċ	,	7	6.4

Appendix 4. Continued.

Taxon	FA	LA	CR	PD	ה/ם	٥/ر	C/J	Area	DI	SSM
Bauria cynops	14	15	12	16	.70	.69	0.22	0.752	0.355	
Lycosuchus vandereti	ω	∞	12	13		•				. 68
Glanosuchus macrops	∞	∞	12	13	.77	. 66		٠.		.67
Ptomalestes avidus	∞	∞	12	13	.71	•	.26	9.	0.32	.67
Dvinia prima			13	13	.83	.67	.20	.77		.71
Procynosuchus delaharpeae		12	14	14	0.8	0.672	0.222	0.685	0.438	۲.
Cynosaurus suppostus	12		15	16		.64	.23	.83	7	.70
SAM-PK-K9954	12		15	17	۲.	. 68	.30	.75	۲.	.71
Galesaurus planiceps	13		15	17	∞.	. 73	. 28	. 78	6	.75
Thrinaxodon liorhinus	13		16	16	∞.	. 73		. 82	.04	. 74
Cynognathus spp.	14		17	18	6.	.80	.27	6.	1.539	۲.
Diademodon spp.	14		17	19	6.	. 79	.32	.95	9	. 76
Trirachodon spp.		15	17	20	٥.	. 78	. 29	.89	.46	.77
Massetognathus spp.	16		17	21	٥.	•	.32	.94	1.673	.76
Exaeretodon frenguelli	17		17	22	٥.	•	ς.	. 92	1.549	. 79
Tritylodon longaevus			17	23	∞.	. 78	.34			. 78
Oligokyphus spp.			17	24	-	-	-	-	1	. 74
Bienotherium elegans	18	18	17	25	0.958	0.819	0.317	0.988	1.806	0.775
Kayentatherium wellesi			17	26		-	1		1	. 79
Bocatherium mexicanum			17	27	-			-		.77
Bienotheroides wanhsienensis			17	27		.81	.30	. 97	1.759	.77
Lumkuia fuzzi			18	18	ο.	. 69		0.872	1.106	.77
Ecteninion lunesis			19	20	.94	.84				. 75
Probelesodon lewisi			19	21	0.962	٠.	.27	. 93	3	.81
Probelesodon sanjuanensis			19	21	.95	. 79	.35	.94	П	. 78
Probainognathus jenseni			20	20	96.	. 74	.34	0.938	1.702	. 78
Pachygenelus monus			21	21	.98	.75	. 25	. 95	0	. 82
Sinoconodon spp.			22	22	-	0.814	0.195	-		6.
Morganucodon spp.			22	22		.76	.17	0.952	1	00.

This table provides the six raw measurements used for this study. Entries are arranged alphabetically by taxon name. The four linear measurements (1-4) and two areas (A<sub>d</sub> and A<sub>pd</sub>) are described in the text. All measurements are in mm and areas in mm<sup>2</sup>. A literature reference under the specimen heading means that the measurements were taken from a reconstruction or specimen drawing in that cited work. The following institutional abbreviations are used: AM, Albany Museum, Grahamstown, South Africa; AMNH, American Museum of Natural History, New York; BMNH, The Natural History Museum, London; BP, Bernard Price Institute for Palaeontological Research, University of the Witwatersrand, Johannesburg; BSP, Bayerische Staatssammlung für Paläontologie und Historische Geologie, Munich; CGP, Council for Geosciences, Pretoria (formerly Geological Survey of South Africa); FMNH, Field Museum of Natural History, Chicago; GPIT, Institut und Museum für Geologie und Paläontologie, Tübingen; IGM, Instituto de Geología, Universidad Nacional Autónoma de México, Mexico City; IVPP, Institute for Vertebrate Palaeontology and Palaeoanthropology, Beijing; MB, Humboldt Museum für Naturkunde, Berlin; MCZ, Museum of Comparative Zoology, Harvard University, Cambridge, Massachusetts; MNHN, Muséum National d'Histoire Naturelle, Paris; NM and NMQR National Museum, Bloemfontein; NMNH, National Museum of Natural History, Washington, D.C.; PIN, Paleontological Institute, Moscow; PVSJ, Museo de Ciencias Naturales, Universidad Nacional de San Juan, San Juan, Argentina; RC, Rubidge Collection, Graaff-Reinet, South Africa; ROZ, Collection of Roy Oosthuizen, now housed at the South African Museum; SAM, South African Museum, Cape Town; SGU, Saratov Geological Institute, Saratov, Russia; TM, Transvaal Museum, Pretoria; TSK, Oxford University Museum, Oxford; UCMP, University of California Museum of Paleontology, Berkeley; and UCMZ, Museum of Zoology, University of Cambridge, Cambridge, United Kingdom.

Taxon	Specimen	1	2	3	4	$A_{d}$	$A_{ad}$
Aelurosaurus felinus	BMNH R339	79.4	65.8	47.3	17.8	710	387
Aerosaurus wellesi	UCMP 40097	149	106.1	58.1	14.5	602	1521
Aerosaurus wellesi	UCMP 40096	84.6	62.3	45			
Aloposaurus ?tenuis	BP/1/709	107.3	78.3	52.7			
Aloposaurus tenuis	BP/1/789	111.2		40			
Anomocephalus africanus	BP/1/5582		10.6			4205	
Anteosaurus magnificus	Unpublished data	560		315			
Anteosaurus magnificus	SAM-PK-9329	561.1	445.4	221.1		38,164	
Arctognathus cf. curvimola	MCZ 4357	162.5		86.2	44.9		
Arctognathus sp.	UCMZ FRP91	282	213	138.4	63.4	8392	5713
Aulacephalodon moschops	UCMP 42699	377.5	238.6	165.8	69.1	19,536	14,022
Aulacephalodon sp.	USNM 24621	391		212	43.7	16,805	16,702
Australosyodon nyaphuli	NMQR 3152	261.5			58		
Australosyodon nyaphuli	Rubidge 1994	258.8	192.5	115	45		
Bauria cynops	AMNH 5622	102.8		79	20.3	1527	580
Bauria cynops	BP/1/1180	95.9		65.5	19.5	886	250
Bauria cynops	BP/1/3770	100.4		70.6	20.4		
Bauria cynops	USNM 23331	81.4	57.7	50.6	22.5		
Biarmosuchus tener	PIN 1758/2	143.9	109	77.4	22.2	1386	1014
Biarmosuchus tener	PIN 1758/7	176	138.4				
Biarmosuchus tener	PIN 1758/8	177.7	158		19.5		
Bienotherium yunnanense	Hopson 1965	114.7	109.1	93.6	31.8	3649	45
Bienotherium yunnanense	Unpublished data	112	108	92	40		
Bienotheroides wanhsienensis	Sun 1984	100.7	97.9	81.7	30.6	2832	84
Bocatherium mexicanum	IGM 3492	38			15.4		
Casea rutena	MNHN MCL2	81.9	62	31.5	16.3	688	663
Cistecephaloides boonstrai	SAM-PK-6243	43.3	31.5	20.5	11.2	252	152
Cistecephalus microrhinus	SAM-PK-K7852	28.7	16.3	14.3			
Cistecephalus microrhinus	SAM-PK-K6814	37.6	21.4	12.1	11.4	215	186
Cistecephalus sp.	USNM 22942	46		21.5			
Cotylorhynchus romeri	Unpublished data	151	76.5	54.2	12.5	1626	1827
Criocephalus sp.	SAM-PK-K319	330	265	130	82	12,440	10,256
Cynognathus merenskyi	BP/1/1181	284.2	278	227	76.1	12,110	10,200
Cynognathus platyceps	BSP 1934 VIII 1	237	219	198.4	62.6		
Cynognathus platyceps	BSP 1934 VIII 2	207	217	170.1	02.0	12,661	505
Cynognathus sp.	UCMP 42749	318.6	289.5	261	95.6	23,495	1161
Cynognathus sp.	Kermack et al. 1973	330.4	304.6	254.1	87.6	24,312	1395
Cynosaurus suppostus	BP/1/4469	46.7	38.7	29.9	07.0	309	62
Cynosaurus kitchingi	TM 279	46	40.2	30.2	10.7	307	02
Cyonosaurus longiceps	FMNH WM1515	134	10.2	85	10.7		
Cyonosaurus longiceps	BP/1/137	146.2	104.6	78.7			
Cyonosaurus longiceps	BP/1/2598	117.6	87.4	73.9		1067	758
Cyonosaurus rubidgei	BP/1/2867	139	102.6	72.5	30.9	1845	2268
Diademodon polyphagus	BMNH R2578	188.2	104.0	153.5	30.9	5356	2200
Diademodon rhodesiensis	BP/1/3639	188.8	185	162.8	62.3	5550	
Dimenional modestensis	DI / I/ 3037	100.0	100	102.0	02.3		

Appendix 5. Continued.

Taxon	Specimen	1	2	3	4	$A_d$	$A_{ad}$
Diademodon sp.	MB R1004		160			5791	270
Diademodon sp.	MCZ 7843	162.7	151.9	121.7	48.8		
Diademodon tetragonus	RC 112	112	112.4	90.2	41.6		
Diademodon tetragonus	USNM 22937	255.8	237	194.8	78.9		
Dicynodon lacerticeps	USNM 25183	78.9	53.6	31.1	14.2		
Dicynodon leoniceps	BP/1/349	363.4	226.8	161	43.9	19,154	13,019
Dicynodon leoniceps	MB R992	325	230	110	65		
Dicynodon leontops	AMNH 5582	232	152.4	77.4	40.4	1.400	0.47
Dicynodon sp. Dicynodon trigonocephalus	Cluver and King 1983	104.6 147.9	63.4 101.4	50.8	9 21.4	1400 3828	946 2621
Dictodon feliceps	TSK 14 RC 100	80	49	58.3 34.4	6	3626	2021
Diictodon feliceps	CGP RS97	168	104.4	73	20		
Diictodon grimbeeki	AMNH 1991	70.9	104.4	29.4	16.6		
Diictodon grimbeeki	USNM 25157	90.3	54.1	38.6	18.2	1096	640
Diictodon grimbeeki	USNM 452057	86.6	67.4	34.9	25.4		
Diictodon grimbeeki	UCMP 42396	70.4	42.4	22.4	10.2	675	312
Diictodon sollasi	USNM 25217	72.3	45.3	34.2	14.3	432	310
Diictodon sp.	BMNH R11184	74.2	53.3	39.3	18.7		
Diictodon sp.	MB R1000	73	49.6	20.2	18	809	485
Diictodon sp.	UCMP 42837	68	39.8	29.1	10.2		
Diictodon sp.	UCMP 41757	100.6	62.8	35.4			
Diictodon sp.	UCMP 42049	80.8	50	35.9	11.1		
Diictodon sp.	USNM 22915	44.4	30.9	24.1	10.4		
Diictodon sp.	USNM 22939	75.2	50.9	31.9	16.5	876	545
Diictodon sp.	USNM 22949	79.7	56.2	38.1	17.6	836	382
Diictodon sp.	USNM 452060	84.8	65.4	41.2	21.5	1675	897
Diictodon sp.	SAM-PK-K6873	83.1	56.7	38.3	24.1		
Diictodon sp. Diictodon sp.	SAM-PK-K6929 SAM-PK-K7084	96.3 60.3	64.5 37.9	41 25.9	25 8.4		
Diictodon testudirostris	USNM 22982	62	42.6	14.4	10.7	505	302
Diictodon testudirostris	SAM-PK-10086	68.2	48.5	33.8	15.8	643	457
Dimetrodon limbatus	MCZ 2779	336.1	269	174	80.9	8887	13,082
Dimetrodon limbatus (m)	AMNH 4081	383.5	230.9	139.5	71.3	13,775	14,448
Dinogorgon quinquemolaris	RC 103	292		144		,	,
Dinogorgon quinquemolaris	GPIT K16	377	255.1	157.3		17,736	10,367
Dvinia prima	Unpublished data	86.3	70.9	57.8	17	823	240
Dvinia prima	PIÑ 2005/2469	83.3	70.8	56.4	17.5		
Ecteninion lunensis	Martinez et al. 1996	91.3	86.3	77.5			
Edaphosaurus boanerges	Modesto 1995	123	88	47	22	1087	2078
Edaphosaurus pogonias	AMNH 4009	160.6	110	45.8	20.6		
Edaphosaurus pogonias	Romer and Price 1940	164	105.9	56.9	23.4	1699	2955
Edaphosaurus sp.	AMNH 21326	154			20.3		
Edaphosaurus sp.	MCZ 3417	138.7	106.9	64.5	39.8	2233	3085
Edaphosaurus sp.	USNM 299844	140.7	88.9	51.2	34.8	1593	2738
Emydops arctatus	BMNH R1690	40.5	21.0	21.3	6 E		
Emydops sp.	AMNH 8209	35.9	21.9	11.6 93.3	6.5 28.2	11 660	4833
Endothiodon uniseries Endothiodon uniseries	AMNH 5334 BMNH R4044	256.2 206	194.2 141.7	93.3	20.2	11,668 7072	3050
Endothiodon uniseries	AMNH 5612	278	200.7	91.9	34.7	12,250	4979
Endothiodon whaitsi	AMNH 5565	393.8	276.5	162.2	47.3	12,230	17/7
Ennatosaurus tecton	PIN 1580/16	146.1	103.7	51.5	35.4		
Ennatosaurus tecton	PIN 1580/14	163.8	135.8	51.3	21	2871	4685
Ennatosaurus tecton	PIN 1580/24	95.2	65.9	38.2		549	781
Ennatosaurus tecton	PIN 1580/17	165.7		34.4			
Eodicynodon oelofseni	NMQR 2913	51.6	28.3	24.8	11.2	373	296
Eodicynodon oostuizeni	NMQR 2991	82	49.5	27.2	20.2	682	717
Eodicynodon oostuizeni	NMQR 2911	64.9	37.4	29	24	631	633
Eothyris parkeyi	MCZ 1161	55.3	40.7	23.1	4.4	145	235
Estemmenosuchus mirabilis	PIN 1758/6	281.6			56	7984	5230
Estemmenosuchus uralensis	PIN 1758/4	460	385	190	115	21,148	18,436
Estemmenosuchus uralensis	PIN 1758/327	415	325	206	85		
Exaeretodon freguelli	Bonaparte 1962	284	240	192	40.0		
Exacretodon freguelli	MCZ 4493	197.4	188	132	49.8		
Exaeretodon freguelli	MCZ 4469	184	163.6	133.1			

Appendix 5. Continued.

Taxon	Specimen	1	2	3	4	$A_d$	A <sub>ad</sub>
Galeops whaitsi	Brinkman 1981	42.3	26.5	22.5	7.1	283	274
Galesaurus planiceps	BP/1/4714	65.2		46.4	17.4		
Galesaurus planiceps	BP/1/5064	89.8	70.3	65.5	21.2	1015	313
Galesaurus planiceps	CGP 1/74	88.6			29		
Galesaurus planiceps	NMQR 860	99.1	81.6	68.8	24.9		
Galesaurus planiceps	NMQR 1451	74	72.1	56.2	21.2	1.400	222
Galesaurus planiceps Galesaurus planiceps	NMQR 3340 SAM-PK-K9956	86.8 60.2	73.1 49.4	63.2 42.6	31.3 16.2	1490 435	333 127
Galesaurus planiceps	TM 83	86.7	47.4	70.6	10.2	433	127
Glanosuchus macrops	van den Heever 1994	221.1	171.6	146.3		4867	1849
Gorgonops torvus	BP/1/4089	155.8	107.9	85.3		100,	1019
Gorgonops whaitsi	BP/1/1426	285.7		173.9			
Gorgonopsid indet sp.	SAM-PK-6417	113.1	93.9	75.7	18	1691	541
Hamilton form	KUVP 12483	31.5	22.5	15	1.6		
Hamilton form	KUVP 12483	33.4	23.8	12.7	1.9	28	51
Haptodus garnettensis	ROM 43602	94.6	71.1	26.4	10.2		
Haptodus garnettensis	ROM 30099	100.9	81	31.4			
Haptodus garnettensis	Laurin 1993	102.1	81.6	34.4	13.7	566	865
Hipposaurus boonstrai	SAM-PK-8950	137	119.1	73.4	38.5	1974	1623
Hipposaurus boonstrai	SAM-PK-9081	193.7	168.2	87.8	39.6	3602	3743
Hipposaurus sp. Hofmeyeria avatus	CGP 1/66 TM 254	173.3 79.3	140.3	99.6 47.4	29 17.2	2215	2867
Hofmeyeria avatus	BP/1/4404	52.8	61.4 41.6	47.4 34.9	9.2	387	240
Ianthasaurus hardestii	Unpublished data	61.1	39.6	18.5	4.5	307	240
Ictidosuchoides intermedius	BP/1/218	89.8	75	60.6	21.3		
Ictidosuchoides longiceps	USNM 336444	79.9	65	60	16.9		
Ictidosuchoides longiceps	RC 646	70.5	54.5	42	10		
Ictidosuchoides sp.	GPIT K70	115.7	92	74.5	27.3		
Ictidosuchoides sp.	SAM-PK-K6731	84.5	57	45.5		546	524
Ictidosuchoides sp.	SAM-PK-K8659	92.5	77.8	60.1	15.3	608	262
Ictidosuchoides sp.	SAM-PK-K6886	130.1	102.5	84.7	28.5		
Jonkeria augusticeps	AMNH 5633	545.3	396.3	265.7	57.4	30,810	21,186
Jonkeria sp.	SAM-PK-12030	508	395	210	112.6	34,377	20,792
Kannemeyeria simocephalus	UCMP 38371	301.7	181.4	112.1	23.3	9912	9204
Kannemeyeria simocephalus	Renaut 2000	374.7	220.5	130.5	35.6	18,328	14,393
Kannemeyeria sp.	BMNH R3602 UCMP 42916	303.9 299.4	193.1 181	95.6 106	43.6 17.6	10,912	8609 7106
Kannemeyeria vanhoepeni Kayentatherium wellesi	MCZ 8812	299.4	208	168.9	82.1	12,029 13,067	367
Kingoria nowacki	UCMZ T747	99.4	63.2	40.6	18	1237	943
Kingoria nowacki	Cluver and King 1983	102.6	67.5	43.6	11.4	1154	751
Kingoria nowacki	UCMZ T748	95.4	68.8	37.2	17.3		
Kingoria nowacki	UCMZ T746	100.6	61.6	37.5	15.6	1272	836
Lumkuia fuzzi	BP/1/2669	33.9	30.5	23.7	8.7	197	29
Lycaenops angusticeps	RC 60	247.2		126.9			
Lycaenops angusticeps	AMNH 5537	250.8	193.1	143.2	49.6	7590	3010
Lycaenops minor	BP/1/209	135.3	113.7	74.9			
Lycaenops ornatus	BP/1/881	201	158.3	107.1	44.2	4984	3327
Lycaenops ornatus	BP/1/2470	151	107.8	81.9		2235	1356
Lycosuchid sp.	van den Heever 1994 AMNH 5600	185.6	155.6	119.6	24.7	5004	2000
Lystrosaurus latirostris Lystrosaurus declivis	NM C403	127.8 87.9	90.3 52.5	63.5 43.7	$\frac{24.7}{14.2}$	2941 1113	2000 969
Lystrosaurus murrayi	MB R2880	81.7	55.5	30.1	14.2	1113	707
Lystrosaurus murrayi	BP/1/269	83.9	56	15.2	16.7		
Lystrosaurus murrayi	BP/1/3908	73	51.1	22.3	12		
Lystrosaurus murrayi	BP/1/4798	75.4	53.8	24.7	10.3		
Lystrosaurus murrayi	MB R2881	80.7	51.9	17	20.7		
Lystrosaurus murrayi	NM C150	82.4	52.1	19.8	16.9		
Lystrosaurus murrayi	NM C6457	89.8	63.4	25.3	22.1	1251	821
Lystrosaurus murrayi	NMQR 3300	65.8	52.6	25.4	14.8		
Lystrosaurus murrayi	NMQR 3239	97	62.9	27.8	21.5	1059	695
Lystrosaurus sp.	UCMP 31359	118.3	82.3	48.2	21.7		
Lystrosaurus sp.	UCMP 42870	87	52.3	19.9	8.8	948	808
Massetognathus sp.	BP/1/4245	77.2	76.9	61.4	25.6		

Appendix 5. Continued.

Taxon	Specimen	1	2	3	4	$A_d$	$A_{ad}$
Massetognathus pascuali	BMNH R8430	121.8	119.8	94.9	42.3		
Massetognathus pascuali	MCZ 3800	111.6	102.2	85.9	43.7	2566	100
Massetognathus pascuali	MCZ 4258	70.5	68	55	16.5	792	62
Massetognathus pascuali	MCZ 4214	65.5	60.6	53.8	18.6	837	
Massetognathus pascuali	MCZ 3999	116.1	110.1	83.7	36.3		
Massetognathus teruggii	MCZ 3807	103.9	98.6	79.2	30.1	1893	101
Massetognathus teruggii	MCZ 3812	129.8	122.5	97.7	47.9		
Massetognathus teruggii	MCZ 4047	117.6		88	42.8		
Massetognathus sp.	PVL 3671		116.3	101.7	47.9	3262	
Mesenosaurus romeri	PIN 158/1		37.6	27.8		139	101
Mesenosaurus romeri	PIN 4541/22	51.4		27.5			
Mesenosaurus romeri	SGU 104v/1558		47.9	28		154	171
Morganucodon sp.	Kermack 1973	19		14.6	3.4	40	2
Moschops capensis	AMNH 5550	243.1	189.1	113.1	59.3	7567	8400
Moschops capensis	AMNH 5553	250.7	192.1	133.8	94	9528	9466
Moschops whaitsi	AMNH 5602	268	217.4	146	81.6	11,234	6958
Moschorhinus sp.	NMQR 3351	215.9	182.9	128		7685	3283
New galesaurid sp.	SAM-PK-K9954	76.9	59.9	52.8	23.4	724	230
Olivieria parringtoni	NMQR 62	78.4	70.6	48.2	20.3	613	331
Olivieria sp.	BP/1/3849	85.5	72.7	54	28.8	631	359
Olivieria sp.	Unpublished data	84.2	67.9	59	14	515	303
Ophiacodon retroversus	MCZ 1203	475.9	328.2	153.3	30	11,300	13,694
Ophiacodon sp.	USNM 487096					3394	3778
Ophiacodon sp.	USNM 487098	282.1	229.9	217.2	10		
Ophiacodon uniformis	Romer and Price 1940	272	185.6	72.1	13.3	3709	3190
Oudenodon halli	BMNH R4067	166.1	109.1	60	48.5		
Oudenodon sp.	SAM-PK-3414	98.9	69.5	58.8	31.2		
Oudenodon sp.	SAM 6045	86.2	51.4	29.7	11.9	765	648
Oudenodon sp.	TSK 67	102.8	66.7	38.7	19.2	1420	892
Oudenodon sp.	TSK 104	106.8	65.7	40.2	22.7		
Oudenodon sp.	USNM 22814	250.6	143.7	96.8	42.9		
Oudenodon sp.	USNM 24626	213	130.6	89.1	36.5	6072	3584
Oudenodon sp.	USNM 24922	271	173	99.3	62.9	10,408	6119
Oudenodon sp.	USNM 335338	82	52.6	31.9	13.4	1062	645
Oudenodon sp.	USNM 452032	110.6	69.9	62.1	19.8		
Pachygenelus monus	Unpublished data	63.9	63.1	47.5	11.3	605	30
Pachygenelus sp.	BP/1/4761	36.2		27.4	11.9		
Patranomodon nyaphulii	NMQR 3000				9.6	217	190
Pelanomodon sp.	UCMZ T981	118.4	80.8	43.9	31.3	1918	1639
Pelanomodon sp.	GPIT K114	191	113.9	81.7	31.6		
Pelanomodon sp.	GPIT K14					4545	3797
Placerias gigas	UCMP 32405	302	193.5	91.8	37.6	11,822	10,242
Pristerodon raniceps	BMNH R1650	69.9	50.7	31.1	9.2	614	347
Pristerodon sp.	MB R985	61.4	46.2	24	10.8	361	205
Pristerodon sp.	SAM-PK-10153	49.7	33.2	20.3	8	402	271
Pristerodon sp.	SAM-PK-10161	64.3	42.6	31.6	13.5		
Pristerodon sp.	SAM-PK-10322	41.2	31	15.2			
Pristerodon sp.	SAM-PK-K1658	48.3	34.3	28.6	10.2		
Pristerodon sp.	USNM 23580	39	25.2	15.6	8.9		
Probainognathus jenseni	MCZ 4069	62.6	60.2	47.5			
Probainognathus jenseni	MCZ 4096					747	71
Probainognathus jenseni	MCZ 4274	61.5	58	45.4	25.3	938	65
Probainognathus jenseni	MCZ 4276	61	60	45.1	19.6	708	42
Probainognathus jenseni	MCZ 4293	60.5	57.3	43.3	17.9	746	50
Probainognathus jenseni	Romer 1970	72.7	70.4	56.5	25.5	1148	48
Probelesodon lewisi	BMNH R8429	126.8	120.3	103.9	37.6	3125	157
Probelesodon lewisi	Romer 1969	98	95.5	71.4	24.6	1694	137
Probelesodon sanjuanensis	PVSJ 411	67	63.8	53.4	23.9	1014	60
Proburnetia viatkensis	PIN 2416/1	150.9	117.6	84.2	31.1	-011	
Procynosuchus cf. delaharpeae	MCZ 8967	76.1	59.6	50.4			
Procynosuchus delaharpeae	BP/1/226	76.2	59.9	52.5	20.5		
			59.4	48.1	15.8	485	255
Procunosuchus delaharneae	BP / 1 / 59 I						
Procynosuchus delaharpeae Procynosuchus delaharpeae	BP/1/591 BP/1/2600	72.2 76.7	39.4	53.9	15.3	103	200

Appendix 5. Continued.

Procymosuchus delaharpeae	Taxon	Specimen	1	2	3	4	$A_d$	A <sub>ad</sub>
Procymosuchus delaharpeae         SSAM-PK-K338         72.2         58.4         15.4         73.4         293           Procymosuchus sp.         MCZ 8968         76.7         63.8         54.7         15.3         74.1         293           Prorubidgea phe         BMNH 89750         260         155         15.7         15.3         15.2	Procynosuchus delaharpeae	RC 87	56	43.5	34	16		
Procymosuchus delahrpae         TSK 34         82.4         69.1         58.4         15.4         73.4         293           Prorubidgea sp.         MCZ 8968         76.7         63.8         54.7         15.5           Prorubidgea alticeps         BP/11/1566         225         181         109         74.9         74.9           Prorubidgea alticeps         BP/11/1566         225         181         109         74.9         74.9           Prorubidgea arbousta         BR C 34         262.2         111.1         185.8         76.7         6608         3255           Robertia broomiana         SAM-PK-11885         41.3         25.7         23.6         5.4         80.8         18.1         18.1         80.8         18.1         18.1         80.9         25.7         18.1         80.9         25.7         80.0         32.5         80.0         80.7         80.0         32.5         80.0         80.7         80.0         80.0         80.7         80.0         80.7         80.0         80.7         80.0         80.7         80.0         80.0         80.7         80.0         80.0         80.0         80.0         80.0         80.0         80.0         80.0         80.0         80.0 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>								
Procymsuchus sp.         MCZ 8968         76.7         63.8         54.7         15.3           Prornubidgea pl.         BMNN H8750         260         195           Prorubidgea alticeps         BP/1/813         193.6         93.4         PVA           Prorubidgea robusta         BP/1/2190         139.4         111.4         78.5           Prorubidgea robusta         BP/1/2190         139.4         111.4         78.5           Prorubidgea robusta         SAM-PK-11942         240.6         171.1         158.9         63.7         6608         3255           Robertia fromorima         SAM-PK-11942         240.6         171.1         158.9         63.7         6608         3255           Robertia fromorima         SAM-PK-11883         41.3         25.7         23.6         5.4         5.5           Robitia sp.         CISS         41.3         24.2         58.6         5.0         5.0         5.0         7.0         1.0         2.1 </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>=</td> <td></td>							=	
Prorubidge alticeps							734	293
Prorubidge alticeps				63.8		15.3		
Prorubidgea alticeps								
Prorubidge anacabei				181		74 9		
Promublisgea maccabei						7 1.7		
Pomalesies avidus								
Robertia sp.   USNM 410241   52.4   35.1   8.1   Rubidgea attrox   RC 13   39.2   181.6   Rubidgea platyrhina   BP/1/803   360   277   148   83   19.31   11.021   Rubidgina sp.   BP/1/8924   102.9   83.2   83.6   81.8   1035   570   Rubidgina sp.   CGP 1/67   121   87.8   68.8   18.1   1035   1216   Scylacops capensis   UCMZ 1356   118.5   86.9   70.1   27.6   12.5   25.0   20.0   27.6   27.5   27.0   27.5   2		SAM-PK-11942	240.6	171.1	158.9	63.7	6608	3255
Rubidgea alrax   RC 13   392.5   181.6   Start Rubidgea platyrhina   BP 1/803   360   277   148   83   19,310   11,021   Rubidgina sp.   BP 1/803   360   277   148   83   19,310   11,021   Rubidgina sp.   CGP 1/67   121   87.8   68.8   18.1   10.5   1216   Scylacops capensis   UCMZ T356   118.5   86.9   70.1   27.6   Scylacops capensis   UCMZ T356   118.5   86.9   70.1   27.6   Scoodnotosaurus obtusidens   MCZ 1124   268.2   170.4   120   22.8   2934   3404   Sinoconodon sp.   IVPP 4727   44   34.4   8.1   268.   Sinoconodon sp.   IVPP 8688   46   38.9   9.5   341   59.5   Sphenacodon frox   UCMP 83459   238.4   69.3   39.5   341   59.5   Sphenacodon frox   UCMP 34219   238.3   171   85.9   5136   6715   Sphenacodon frox   UCMP 34219   238.3   171   85.9   5136   6715   Sphenacodon frox   UCMP 34226   200.7   142.7   86.2   42.1   27.11   4738   Stableckeria potens   GPIT 1   380   245   180   65   24,638   18,852   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.3   88.8   40.3   6250   4567   Stableckeria potens   GPIT 2   206.8   148.2   30.8   48.8   40.3   6250   4567   Stableckeria potens   GPIT 2   207.4   27.4   27.1   27.1   47.5   Stableckeria potens   GPIT 2   207.4   27.4   27.1   27.1   47.5   Stableckeria potens   GPIT 2   207.4   27.4   27.1   27.1   27.5   Stableckeria potens   GPIT 2   207.4   27.4   27.1   27.1   27.5   Stableckeria potens   GPIT 2   207.4   27.4   27.4   27.4   27.4   27.4   27.4   27.4   2	Robertia broomiana	SAM-PK-11885	41.3	25.7	23.6	5.4		
Rubis   Rubi	1			35.1		8.1		
Rublidgina sp.         GP/1/3924         102.9         83.2         58.6         13.9         828         570           Rublidgina sp.         CGP 1/67         121         87.8         68.8         18.1         1035         1216           Scylacops capensis         UCMZ T356         118.5         86.9         70.1         27.6         34.4           Sinoconodon sp.         IVPP 4277         44         34.4         8.1         26.8           Sinoconodon sp.         IVPP 4884         46         38.9         9.5         341           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34226         200.7         142.7         86.2         42.1         2711         478           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         18.1         88.8         40.3         24.63         18.1         84.6         50.2         42.1         111         83.6	O .							
Rubidgina sp.         CGP 1/67         121         87.8         68.8         18.1         1035         1216           Scyclacops capensis         UCMZ T356         118.5         86.9         70.1         27.6         Secodontosaurus obtusidens         MCZ 1124         268.2         170.4         120         22.8         2934         3404           Sinoconodon sp.         IVPP 8688         46         38.9         9.5         3411         5925           Sphenacodon ferox         UCMP 34219         23.8.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         23.8.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715         7514         4738           Sphenacodon ferox         UCMP 34219         23.8.3         171         85.9         5136         6715         4714         4738           Sphenacodon ferox         UCME 34219         23.8.3         171         85.9         5136         6715         5468         481.8         42.1         42.1         42.1         42.1         42.1         42.1         42.1         42.1         4								
Scylacops capensis         UCMZ 1356         118.5         86.9         70.1         27.6           Scocodontosaurus obtusidens         MCZ 1124         268.2         170.4         120         22.8         2934         3404           Sinoconodon sp.         IVPP 4727         44         34.4         8.1         268           Sinoconodon sp.         IVPP 8688         46         38.9         9.5         341           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34226         200.7         142.7         56.2         42.1         2711         4738           Stableckeria potens         GPIT 1         380         245         180         65         24.638         18.852           Stableckeria potens         GPIT 2         206.8         148.3         8.8         40.3         625.0         4567           Stypacocephalus platyrhynchus         SAM-PK-8936         262.3         212.1         171.2         525         153           Suminia gelmanovi         Rybezynski 2000         46         30         26.6         10.5         71.3           Sydao effemovi         PIN 157/2         213								
Secondinosaurus obtusidens							1035	1216
Simoconodon sp.         IVPP 8688         46         38.9         5.341         268           Sphenacodon ferox         UCMP 83459         238.4         69.3         39.5         4119         5925           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         421         2711         4738           Stallackeria potens         GPIT 1         380         245         180         65         24,638         18,882           Stallackeria potens         GPIT 2         206.8         148.3         88.8         40.3         6250         4567           Stallackeria potens         GPIT 2         206.8         148.3         88.8         40.3         6250         4567           Stamina germanoor         Rybezynski 2000         46         30         26.6         10.5         255         153           Sydon efremooi							2034	3404
Sinconodon sp.         IVPP 8688         46         38.9         9.5         341           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34226         200.7         142.7         56.2         42.1         2711         4738           Stablackeria potens         GPIT 1         380         245         180         65         24,638         18,852           Stablackeria potens         GPIT 2         206.8         148.3         88.8         40.3         6250         4567           Styracocephalus platyrhynchus         SAM-PK-8936         262.3         212.1         171.2         82.2           Suminia getmanovi         PIN 2212/10         40.2         27.4         22.1         82.2           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Tapinocaninus pamelae         ROZ K95         353.4         299.9         185.6         71.4         75.6           Tapinocaninus pamelae         NMQR 2986         353.1         278.1         184.4         75.6         71.4           Tetraceratops insiginis         AMNH 4526         <				170.4				3404
Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34226         200.7         142.7         56.2         42.1         2711         4738           Stallackeria potens         GPIT 1         380         245         180         65         24,638         18,852           Stallackeria potens         GPIT 2         206.8         148.3         88.8         40.3         6250         4567           Styracocephalus platyrhynchus         SAM-PK-8936         262.3         212.1         171.2         82           Suminia getmanovi         PIN 157.7         213         182.7         100.8         42.6         5004         2325           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Tapinocaninus pamelae         NMQR 2986         353.1         279.9         185.6         71.4         75.6         <	1							
Sphenacodon ferox         UCMP 34219         238.3         171         85.9         5136         6715           Sphenacodon ferox         UCMP 34226         200.7         142.7         56.2         42.1         2711         4738           Stahleckeria potens         GPTT 1         380         245         180         65         24,638         18,852           Stahleckeria potens         GPTT 2         206.8         148.3         88.8         40.3         6250         4567           Styracocephalus platyrhynchus         SAM-PK-8936         262.3         212.1         171.2         2           Suminia getmanovi         PIN 2212/10         40.2         27.4         22.1         8.2         590         153           Suminia getmanovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Sydon of fermovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Tapinocanimus pamelae         NMQR 2986         353.1         228.1         44.4         75.6         71.4         71.4         71.4         71.4         71.4         71.4         71.4         71.4         71.4         71.4	1							5925
Sphenacodom ferox         UCMP 34226         200.7         142.7         56.2         42.1         2711         4738           Stahleckeria potens         GPIT 1         380         245         180         65         24,638         18,852           Stahleckeria potens         GPIT 2         206.8         148.3         88.8         40.3         6250         4567           Styracocephalus platyrhynchus         SAM-PK-8936         262.3         212.1         171.2         7           Suminia getmanovi         PIN 2212/10         40.2         27.4         22.1         8.2         5           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         500.4         2325           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         500.4         2325           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         500.4         2325           Syodon efremovi         PIN 157/2         213         182.7         101.0         86.7         14.4         36.4         491.2         36.4         491.2         32.1         47.2         45.2         46.2         42.2	,			171				
Stahleckeria potens         GPT 1         380         245         180         65         24,638         18,852           Stahleckeria potens         GPT 2         206.8         148.3         88.8         40.3         6250         4567           Stypracocephalus platyrhynchus         SAM-PK-8936         262.3         212.1         171.2         171.2           Suminia getmanovi         PIN 221/10         40.2         27.4         22.1         8.2           Suminia getmanovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Sydon efremovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Tapinocanimus pamelae         NMQR 2986         353.1         228.1         144.4         75.6         71.4         71.4         71.4         71.4         75.6         71.4         71.4         75.6         70.2         7	,					42.1		
Styracocephalus platyrhynchus   SAM-PK-8936   262.3   212.1   171.2		GPIT 1	380	245	180	65	24,638	18,852
Suminia getmanovi         Rybczynski 2000         46         30         26.6         10.5         255         153           Suminia getmanovi         PIN 2212/10         40.2         27.4         22.1         8.2         150           Syodon efremovi         PIN 157/2         213         182.7         100.8         42.6         5004         2325           Tapinocanimus pamelae         ROZ K95         353.4         29.9         185.6         71.4         71.6           Tapinocanimus pamelae         NMQR 2987         346.8         270         170.1         86.7           Tetraconian signis         AMNH 4526         90.1         54.3         37         11.2         656         822           Tetragonias njalilus         GPIT 292         136.4         91.1         64.9         31         2462         1887           Theriognathus sp.         AMNH 8226         140.8         110.8         90.5         23.2         1887           Theriognathus sp.         BP/1/512         111.9         83         79.1         32.7           Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6           Theriognathus sp.         BP/1/184         135.4         110.	Stahleckeria potens	GPIT 2	206.8	148.3	88.8	40.3	6250	4567
Suminia getmanovi		SAM-PK-8936	262.3	212.1				
Syodon efremovi							255	153
Tapinocaninus pamelae		'						
Tapinocaninus pamelae   NMQR 2986   353.1   278.1   184.4   75.6   Tapinocaninus pamelae   NMQR 2987   346.8   270   170.1   86.7   Tapinocaninus pinalitus   UCMZ T753   169.6   118.2   93.5   34.6   4912   3915   3915   Tapinocaninus njalilus   UCMZ T753   169.6   118.2   93.5   34.6   4912   3915   Tapinocaninus njalilus   GPIT 292   136.4   91.1   64.9   31   2462   1887   Theriognathus sp.   AMNH 8226   140.8   110.8   90.5   23.2   Theriognathus sp.   BP/1/512   111.9   83   79.1   32.7   Theriognathus sp.   BP/1/747   116.8   95.7   70.2   Theriognathus sp.   BP/1/844   135.4   110.3   89.2   36.8   2810   1026   Theriognathus sp.   BP/1/844   135.4   110.3   89.2   36.8   2810   1026   Theriognathus sp.   BP/1/82   55.5   44.9   33   12.6   Theriognathus sp.   BP/1/82   55.5   44.9   33   12.6   Theriognathus sp.   BMNH R511a   67.9   61.2   52.2   26.3   1542   762   Thrinaxodon liorhinus   BMNH R511a   67.9   61.2   52.2   Thrinaxodon liorhinus   BMNH R511a   67.9   61.2   52.2   Thrinaxodon liorhinus   BMNH R5480   65.5   53.6   46.7   16.9   575   134   Thrinaxodon liorhinus   BP/1/4263   62.2   55.9   45.9   16.5   Thrinaxodon liorhinus   BP/1/4280   53.8   44.4   36.5   14.3   Thrinaxodon liorhinus   BP/1/4280   53.8   44.4   36.5   14.3   Thrinaxodon liorhinus   MCZ 4282   60.8   53.6   43.6   16.3   Thrinaxodon liorhinus   MCZ 4282   60.8   53.6   43.6   16.3   Thrinaxodon liorhinus   MCZ 4282   60.8   53.6   43.6   16.3   Thrinaxodon liorhinus   MCZ 2179   51.1   35.5   Thrinaxodon liorhinus   MCZ 2184   66.1   52.9   Thrinaxodon liorhinus   MCZ 2184   66.1   52.9   Thrinaxodon liorhinus   MCZ 2266   61.5   51.8   43.3   55.5   51.8   45.3   51.5   52.3   52.5   53.3   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53.5   53							5004	2325
Tapinocaninus pamelae								
Tetraceratops insignis         AMNH 4526         90.1         54.3         37         11.2         656         822           Tetragonias njalilus         UCMZ T753         169.6         118.2         93.5         34.6         4912         3915           Tetragonias njalilus         GPIT 292         136.4         91.1         64.9         31         2462         1887           Theriognathus sp.         AMNH 8226         140.8         110.8         90.5         23.2           Theriognathus sp.         BP/1/512         111.9         83         79.1         32.7           Theriognathus sp.         BP/1/844         135.4         110.3         89.2         36.8         2810         1026           Theriognathus sp.         BP/1/82         55.5         44.9         33         12.6         120           Theriognathus sp.         BP/1/82         55.5         44.9         33         12.6         762           Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6         762           Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6         762           Theriognathus sp.         BP/1/184         135.4								
Tetragonia's njalilius         UCMZ T753         169.6         118.2         93.5         34.6         4912         3915           Tetragonias njalilius         GPIT 292         136.4         91.1         64.9         31         2462         1887           Theriognathus sp.         AMNH 8226         140.8         110.8         90.5         23.2         23.2           Theriognathus sp.         BP/1/512         111.9         83         79.1         32.7         32.7           Theriognathus sp.         BP/1/747         116.8         95.7         70.2							656	822
Tetragonias njalilus         GPIT 292         136.4         91.1         64.9         31         2462         1887           Theriognathus sp.         AMNH 8226         140.8         110.8         90.5         23.2           Theriognathus sp.         BP/1/512         111.9         83         79.1         32.7           Theriognathus sp.         BP/1/747         116.8         95.7         70.2         70.2           Theriognathus sp.         BP/1/844         135.4         110.3         89.2         36.8         2810         1026           Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6         1026           Theriognathus sp.         NMQR 3375         112.3         87.8         75.2         26.3         1542         762           Thrinaxodon liorhinus         BMNH R511         68.7         59.6         48.6         19.6         726         129           Thrinaxodon liorhinus         BMNH R3731         59.5         51.7         45.9         11.9         147           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5         134           Thrinaxodon liorhinus         BP/1/4280         53.8	, 0							
Theriognathus sp.         AMNH 8226         140.8         110.8         90.5         23.2           Theriognathus sp.         BP/1/512         111.9         83         79.1         32.7           Theriognathus sp.         BP/1/747         116.8         95.7         70.2         70.2           Theriognathus sp.         BP/1/844         135.4         110.3         89.2         36.8         2810         1026           Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6         762           Theriognathus sp.         NMQR 3375         112.3         87.8         75.2         26.3         1542         762           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         762         119         762         119         762         119         762         129         762         119         762         129         762         129         762         119         762         129         762         119         762         129         762         129         762         129         762         129         762         129         762         129         762         129         762         129         129	,							
Theriognathus sp.         BP/1/512         111.9         83         79.1         32.7           Theriognathus sp.         BP/1/747         116.8         95.7         70.2           Theriognathus sp.         BP/1/182         35.5         70.2           Theriognathus sp.         BP/1/182         35.5         44.9         33         12.6           Theriognathus sp.         NMQR 3375         112.3         87.8         75.2         26.3         1542         762           Thrinaxodon liorhinus         BMNH R511a         68.7         59.6         48.6         19.6         726         129           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         75.1         45.9         11.9         75.5         134           Thrinaxodon liorhinus         BMNH R53731         59.5         51.7         45.9         11.9         57.5         134           Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         57.5         134           Thrinaxodon liorhinus         BP/1/4280         53.8         44.4         36.5         14.3           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6								
Theriognathus sp.         BP/1/844         135.4         110.3         89.2         36.8         2810         1026           Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6           Theriognathus sp.         NMQR 3375         112.3         87.8         75.2         26.3         1542         762           Thrinaxodon liorhinus         BMNH R511         68.7         59.6         48.6         19.6         726         129           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         726         129           Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         575         134           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5         14.3           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5         14.3           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5           Thrinaxodon liorhinus         MCZ 4282         60.8								
Theriognathus sp.         BP/1/182         55.5         44.9         33         12.6           Theriognathus sp.         NMQR 3375         112.3         87.8         75.2         26.3         1542         762           Thrinaxodon liorhinus         BMNH R511         68.7         59.6         48.6         19.6         726         129           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         726         129           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         726         129           Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         575         134           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5         14.3	Theriognathus sp.	BP/1/747	116.8	95.7	70.2			
Theriognathus sp.         NMQR 3375         112.3         87.8         75.2         26.3         1542         762           Thrinaxodon liorhinus         BMNH R511         68.7         59.6         48.6         19.6         726         129           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         52.2         52.2         52.2         53.6         46.7         16.9         575         134         57.5         15.7         45.9         11.9         575         134         57.5         15.7         45.9         11.9         575         134         57.7         45.9         11.9         575         134         57.7         45.9         11.9         575         134         57.7         45.9         11.9         575         134         57.7         45.9         16.5         57.5         134         57.7         45.9         16.5         57.5         134         57.7         45.9         16.5         57.5         134         57.7         45.9         16.5         57.5         134         57.7         16.5         57.5         45.9         16.5         57.5         14.3         57.7         45.9         16.5         57.5         45.9         16.3		BP/1/844	135.4	110.3	89.2		2810	1026
Thrinaxodon liorhinus         BMNH R511         68.7         59.6         48.6         19.6         726         129           Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2         11.9           Thrinaxodon liorhinus         BMNH R3731         59.5         51.7         45.9         11.9           Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         575         134           Thrinaxodon liorhinus         BP /1/4263         62.2         55.9         45.9         16.5         755         134           Thrinaxodon liorhinus         BP /1/4280         53.8         44.4         36.5         14.3         14.3           Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3         16.3           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3         16.3           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         49.5         21.9           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9         46.3         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75								
Thrinaxodon liorhinus         BMNH R511a         67.9         61.2         52.2           Thrinaxodon liorhinus         BMNH R3731         59.5         51.7         45.9         11.9           Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         575         134           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5           Thrinaxodon liorhinus         BP/1/4280         53.8         44.4         36.5         14.3           Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3           Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ 8377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9      <	0 1							
Thrinaxodon liorhinus         BMNH R3731         59.5         51.7         45.9         11.9           Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         575         134           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5         16.3           Thrinaxodon liorhinus         BP/1/4280         53.8         44.4         36.5         14.3           Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3           Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ 8377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         46.3           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         TM 80a         60.5         50.5						19.6	726	129
Thrinaxodon liorhinus         BMNH R5480         65.5         53.6         46.7         16.9         575         134           Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5           Thrinaxodon liorhinus         BP/1/4280         53.8         44.4         36.5         14.3           Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3           Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ 8377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523						11.0		
Thrinaxodon liorhinus         BP/1/4263         62.2         55.9         45.9         16.5           Thrinaxodon liorhinus         BP/1/4280         53.8         44.4         36.5         14.3           Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3           Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ 8377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         35.5         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9         52.9         52.9           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.2         18.9           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5							575	124
Thrinaxodon liorhinus         BP/1/4280         53.8         44.4         36.5         14.3           Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3           Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ K377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129							373	134
Thrinaxodon liorhinus         BSP 1934 VIII 506         51.8         43.3         38.7         16.3           Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ K377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25								
Thrinaxodon liorhinus         FMNH UR156         56.3         51.5         45           Thrinaxodon liorhinus         MCZ 4282         60.8         53.6         43.6         16.3           Thrinaxodon liorhinus         MCZ K377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMZ 7815         58.1         48.2         43.4<								
Thrinaxodon liorhinus         MCZ K377         67.5         58.9         49.5         21.9           Thrinaxodon liorhinus         MCZ 2179         51.1         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3         116           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98								
Thrinaxodon liorhinus         MCZ 2179         51.1         35.5           Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3         116           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98	Thrinaxodon liorhinus	MCZ 4282				16.3		
Thrinaxodon liorhinus         MCZ 2184         66.1         52.9           Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3         18.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98	Thrinaxodon liorhinus	MCZ K377	67.5	58.9	49.5	21.9		
Thrinaxodon liorhinus         MCZ 2226         61.5         51.8         46.3           Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3         18.4           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98		MCZ 2179	51.1		35.5			
Thrinaxodon liorhinus         MCZ uncat.         75.1         63.2         52.7         18.9           Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98								
Thrinaxodon liorhinus         SAM-PK-K1461         70.9         63.8         55.2         24.2         904         118           Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98						46-		
Thrinaxodon liorhinus         TM 80a         60.5         50.5         42.9         15         523         116           Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98							004	110
Thrinaxodon liorhinus         TM 80b         45         39.5         32.5         13.3           Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98								
Thrinaxodon liorhinus         UCMP 40466         123.3         108.4         91.2         25         499         129           Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98							523	116
Thrinaxodon liorhinus         UCMP 42866         57.8         48.3         39.9         14.1         441         123           Thrinaxodon liorhinus         UCMZ T815         58.1         48.2         43.4         16.2         401         98							100	120
<i>Thrinaxodon liorhinus</i> UCMZ T815 58.1 48.2 43.4 16.2 401 98								
	Titanophoneus potens	PIN 157/1	374.9	276.6	193	72.5	13,398	8016

Appendix 5. Continued.

Taxon	Specimen	1	2	3	4	$A_d$	$A_{ad}$
Trirachodon berryi	BP/1/4658	85	82.1	66	24.6	1519	152
Trirachodon berryi	MCZ 8896	80.7	74.3	61.8	27.2		
Trirachodon sp.	unpublished data	54.5	49.9	42.2	15.9	513	57
Trirachodon sp.	NMQR 3256	71.2	65	59.4	24.9		
Trirachodon sp.	AM 461	76.7	74.9	60.2	21		
Trirachodon sp.	BP/1/4535	40.9	36.6	30	7.3		
Trirachodon sp.	BP/1/5362	97.4		79.4	35.8	1845	155
Trirachodon sp.	BP uncat.	69.8		53.9	19		
Trirachodon sp.	SAM-PK-11481	59.2	53.9	46.9	16.2	575	92
Tritylodon longaevus	BP/1/4778	110	98	86	38		
Tritylodon longaevus	BP/1/5288		116.6	94.4	37	3650	
Tritylodon longaevus	SAM-PK-K1411		84.3	70.5	28	2067	
Ulemica invisa	PIN 157/5	127.5	97.1	74.9	41	2992	1187
Ulemica prima	PIN 157/1112	145.3	115.2	98.7	36.2	3949	1883
Ulemosaurus svijagensis	Efremov 1940	298.8	202.5	164.8	58.1	12,757	9003
Ulemosaurus svijagensis	PIN 2207/2	335	275		105	18,749	12,111
Varanodon agilis	FMNH UR986	163.7	120.9	67.6	8.5	618	1000
Varanops brevirostris	Romer and Price 1940	134.2	106.8	52.8		675	847
Varanosaurus acutirostris	Berman et al. 1995	156.2	110.3	94.9	6.4	1241	905
Varanosaurus acutirostris	BSP 1901 XV 20	150	103.8				
Varanosaurus acutirostris	FMNH PR1760	123	78	81	2.2		
Viatkosuchus sumini	PIN 2213/13	140.5	109.9	85.4	29.6	1912	1042