

# STRATIGRAPHY AND PHYLOGENY OF THE GIANT EXTINCT RAT KANGAROOS (PROPLEOPINAE, HYPSPRYMNODONTIDAE, MARSUPIALIA)

S. WROE

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The Giant Rat-kangaroos were placed in the Propleopinae by Archer & Flannery (1985) and in the Hypsiprymnodontidae by Ride (1993). Cladistic analysis of *Ekaltadeta* material from Riversleigh, northwestern Queensland (Wroe, 1996) suggested that a middle to late Miocene dichotomy in *Ekaltadeta* may have produced two lineages of Plio-Pleistocene *Propleopus*, indicating polyphyly for *Propleopus* and paraphyly for *Ekaltadeta*. Metrical data for propleopines and stratigraphic information support Wroe's (1996) cladistic analysis of propleopines. □ *Propleopinae, Hypsiprymnodontidae, Riversleigh, Ekaltadeta, cladistics*.

S. Wroe, School of Biological Science, University of New South Wales, N.S.W. 2052 Australia; received 4 November 1996.

Giant Rat-kangaroos (Hypsiprymnodontidae: Propleopinae) may be the plesiomorphic sister group of potoroids (Flannery, 1987). Archer & Flannery (1985) considered *Ekaltadeta ima* (Fig. 1) the sister group to *Propleopus* De Vis, 1888 with *P. oscillans* De Vis, 1888 (Fig. 2) the more plesiomorphic and *P. chillagoensis* Archer et al. (1978) (Fig. 2) the more apomorphic within *Propleopus* (Fig. 3). Propleopine species described since 1985 are *Ekaltadeta jamiemulvaneyi* Wroe, 1996, (Fig. 4) and *Jackmahoneya toxoniensis* Ride, 1993. Wroe (1996) suggested another possible phylogeny for the Propleopinae with *E. ima* and *P. chillagoensis* forming the sister group to another clade containing a new species, *E. jamiemulvaneyi*, as the sister taxon to *P. wellingtonensis* and *P. oscillans* (Fig. 5). As an adjunct to the cladistic analysis (Wroe, 1996), metric and stratigraphic data for propleopines are used to clarify intrasubfamilial relationships.

Dental homology for premolars follows Flower (1867) and Luckett (1993) for molars. Higher level systematics of kangaroos follows Flannery (1987) and Ride (1993). Specimens are housed in the Queensland Museum (QMF). Other prefixes include; UCM (University of California Museum), NMV (Museum of Victoria).

## METHODS

Specimens of *Ekaltadeta* from Riversleigh represent 30 individuals from several stratigraphic levels. The relative paucity of specimens and chronological data precludes a strictly stratophenetic approach (*sensu* Gingerich, 1976, 1979; Bown & Rose, 1987) to propleopine phylogeny.

However, a more general consideration of stratigraphy in phylogenetic analysis may be appropriate in association with cladistic treatment where specimens are stratigraphically disjunct or sparsely distributed (Gingerich, 1990).

Sites with *Ekaltadeta* are late Oligocene to early late Miocene (Archer et al., 1989, 1994, 1995). A number of characters were analysed to assess the development of time-dependent changes. Specimens were ranked to indicate relative age (Appendix 1). Stratigraphic levels are from Archer et al. (1989, 1995): level 1=late Oligocene early Miocene; level 2=early Miocene; level 3=late early Miocene; level 4=mid Miocene; level 5=late mid Miocene; level 6=early late Miocene; level 7=Pliocene; level 8=Pleistocene.

I included all propleopines possible, although Pliocene and Pleistocene *Jackmahoneya* and *Propleopus* are known from material often limited to portions of upper and/or lower dentitions. Most *Propleopus* are from the Pleistocene, although material has been recorded from early Pliocene local faunas (Archer & Flannery, 1985). *Propleopus chillagoensis* was described as Pleistocene (Archer et al., 1985), but could be older, possibly late Miocene or early Pliocene (Archer pers. comm.). *Jackmahoneya toxoniensis* is Pliocene (Ride, 1993).

Differences in molar gradient were used by Archer & Flannery (1985) and Wroe (1996) to distinguish propleopine species. Molar gradient reflects both the surface area and length of the molar tooth row. In propleopines a high molar gradient correlates with a reduction in both molar surface area and the length of the tooth row.

Reducing the distance between condyle and sectorial tooth maximizes leverage applicable to the tooth (Young et al., 1989). Through shortening the molar row, leverage on the large shearing  $P^3/3$  of propleopines is increased. This effect is achieved at the cost of molar length.

Relative  $P_3$  size and molar gradient for upper and lower dentitions has been quantified. Distinct reduction in tooth size posteriorly occurs in upper and lower dentitions of *E. ima*. In the upper dentition this steep gradient begins with a reduced posterior width (pw) relative to the anterior width (aw) of  $M^2$  which then ramifies through  $M^{3-4}$ . In *E. ima*  $M^4$  pw is  $<1/2$   $M^2$  aw. The upper dentition of *P. chillagoensis* is similar to that of *E. ima*. Lower dentition is not known for *P. chillagoensis*. For *P. oscillans*  $M^{3-4}$  are missing but  $M^2$  pw is only slightly less than  $M^2$  aw suggesting a less extreme gradient. This supposition is strongly supported by the lower dentition in which molar gradient contrasts strongly with *E. ima*.  $M_{1-4}$  tooth widths decrease steadily anteroposteriorly in *E. ima* but are reversed in *P. oscillans* where tooth width increases posteriorly for  $M_{1-3}$ , with only a slight decrease in  $M_4$ .

Several methods to quantify molar gradient have been considered. Accurate determination of individual molar surface areas and comparisons between teeth would be useful but would require 2 or more teeth/ specimen, greatly limiting data sets, particularly for upper dentitions. Molar gradient might also be estimated geometrically by determining the angle at which a line drawn buccally or lingually through the faces of the crown intersects the mid-line of the dentary or skull.

In this study the clear initiation of a marked molar gradient at  $M^2$  in the upper dentitions of *E. ima* and *P. chillagoensis* permitted estimation of the gradient from a single molar by comparing aw to pw. In lower dentitions the gradient is less

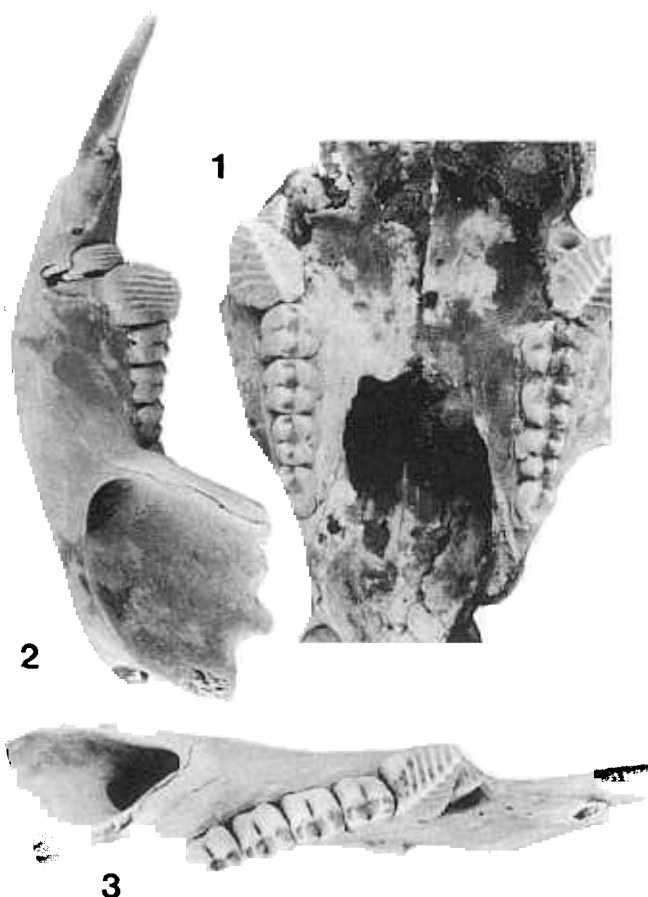


FIG. 1. *Ekaltadeta ima*, x 2. A, occlusal view of QMF12436 (uppers). B, buccal view of QMF12435, left dentary containing  $I_1$ , alveolus for  $I_2$ ,  $P_{2-3}$ ,  $M_{1-4}$ . C, occlusal view of QMF12435.

distinct and 2 molars were required to demonstrate a gradient. Measurements were made using a Wild MMS 235 Digital Length-Measuring Set attached to a Wild M5A Stereomicroscope. Abbreviations are: l=length, w=width, aw=anterior width, pw=posterior width, dd=depth of dentary, G-value=ratio of anterior to posterior tooth width.

## RESULTS

$M^2$  aw /  $M^2$  pw VS STRATIGRAPHIC LEVEL. (Fig. 6). For upper dentitions the ratio  $M^2$  aw:  $M^2$  pw (G-value) was used as an arbitrary measure of molar gradient, with  $M^2$  being common to the largest number of specimens.

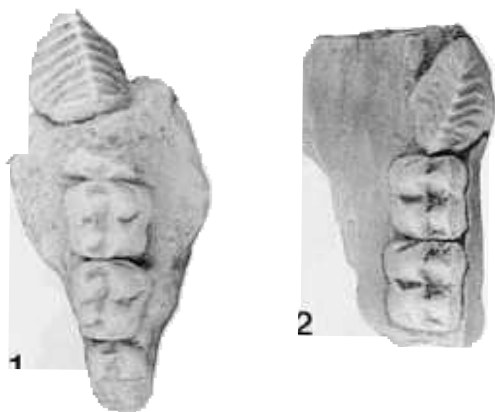


FIG. 2. *Propleopus chillagoensis*, x 2. A, occlusal view of NMV P15917, right maxillary fragment (juvenile), containing unerupted P<sup>3</sup>, partial M<sup>1</sup>, M<sup>2-3</sup>, partial M<sup>4</sup> (cast of holotype). B, *Propleopus oscillans*, x 2, occlusal view of QMF6675, left maxillary fragment, containing P<sup>3</sup>, M<sup>1-2</sup>.

A trend is apparent in this scatter graph of G-value against stratigraphic level. *P. chillagoensis* and *P. oscillans* represent 2 extremes with G-values of 1.23 and 1.06 respectively, with the lower number indicating a lesser molar gradient. *Ekaltadeta ima* from levels 3 and 4 has a limited range of G-values (1.09-1.15).

The 2 *Ekaltadeta* from level 6 both fell outside the range of *E. ima* from older strata. *E. jamiemulvaneyi* (QMF24212; Cleft of Ages 4 Site) had a low G-value of 1.05, slightly less than that of *P. oscillans*. *E. ima* (QMF24211; Henk's Hollow Site) had a relatively high G-value of 1.19 approaching that of *P. chillagoensis*. These results indicate a divergence in the *Ekaltadeta* lineage with one population leading to *P. oscillans* and another leading to *P. chillagoensis*.

**M<sub>1</sub> pw / M<sub>2</sub> pw VS STRATIGRAPHIC LEVEL.** (Fig. 7). The molar gradient of the dentary was estimated by dividing M<sub>1</sub> pw by M<sub>2</sub> pw (G-value). *P. oscillans* had the lowest G-value at 0.93. The G-values for *P. wellingtonensis* and *J. toxoniensis* were slightly higher at 0.96. At levels 3 and 4 the G-values for *Ekaltadeta* were 1.01-1.08. The G-value for *E. jamiemulvaneyi*, from level 6 (QMF24200, Encore Site) was 0.97. This placed *E. jamiemulvaneyi* about halfway between the lowest G-value from levels 3 and 4 and *P. oscillans*. Again the highest degree of divergence among *Ekaltadeta* was for the *E. jamiemulvaneyi* from level 6, possibly indicating a trend toward

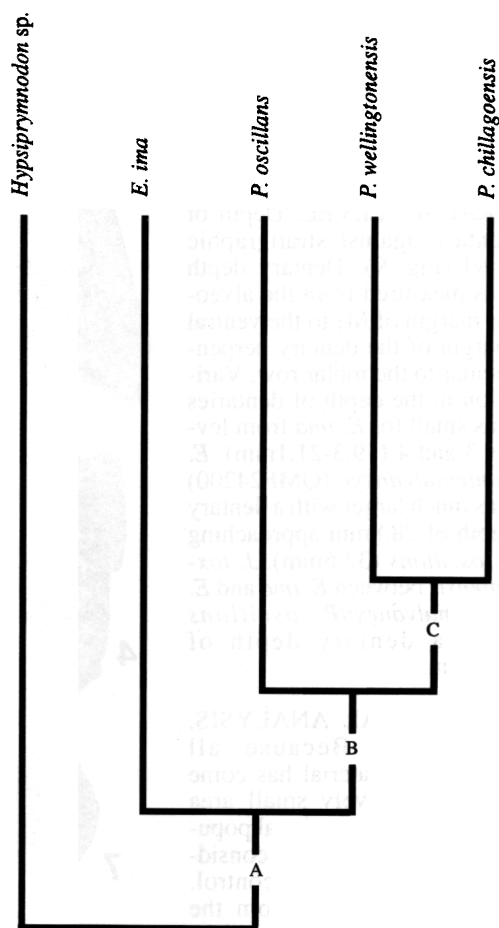


FIG. 3. Cladogram for the propleopines from Archer & Flannery (1985). Character states at nodes: A=gain of an anterior cristid emanating from the metaconid of M<sub>1</sub>, gain of derived I<sub>1</sub> morphology; B=incorporation of the protolophid into the anterior lophid of M<sub>1</sub> loss of P<sub>2</sub> with eruption of P<sub>3</sub>, dentary deeper anteriorly than posteriorly; C=reduction of metacone/entoconid, P<sub>3</sub> hypertrophy.

the species with low molar gradients (*J. toxoniensis*, *P. oscillans* and *P. wellingtonensis*).

**P<sub>3</sub> w/M<sub>1</sub>pw VS STRATIGRAPHIC LEVEL.** (Fig. 8). In *P. oscillans* P<sub>3</sub> width was small compared to M<sub>1</sub> posterior width (1.09). For *J. toxoniensis* relative P<sub>3</sub> width was greater (1.27). *E. ima* from levels 3 and 4 had ratios of P<sub>3</sub> w / M<sub>1</sub> pw of 1.35-1.52. *E. jamiemulvaneyi* from Encore

site (QMF24200) again positioned between *E. ima* from lower strata and *J. toxoniensis* / *P. oscillans*, with a ratio of 1.28.

**DEPTH OF DENTARY AGAINST STRATIGRAPHIC LEVEL.** Depth of dentary against stratigraphic level (Fig. 9). Dentary depth was measured from the alveolar margin of  $M_1$  to the ventral margin of the dentary perpendicular to the molar row. Variation in the depth of dentaries was small for *E. ima* from levels 3 and 4 (19.3–21.1mm). *E. jamiemulvaneyi* (QMF24200) was much larger with a dentary depth of 28.8mm approaching *P. oscillans* (32.6mm). *J. toxoniensis* between *E. ima* and *E. jamiemulvaneyi* / *P. oscillans* with a dentary depth of 23.3mm.

**STATISTICAL ANALYSIS.** (Table 1). Because all *Ekaltadeta* material has come from a relatively small area (Riversleigh), a regional population of potoroids was considered an appropriate control. Sixteen specimens from the Australian museum of *Potorous tridactylus* collected around Hobart were used, this being the largest potoroid specimen sample available. Variation in the G-values of *Ekaltadeta* from levels 3 and 4 approached that of *P. tridactylus*. When G-values from the 2 *Ekaltadeta* from level 6 were included the variation fell well outside that of the local *P. tridactylus* population.

## DISCUSSION

Increases in premolar and molar shear within the Propleopinae appear to be mutually exclusive and their relative importance probably reflects dietary preference. A requirement for high premolar shear might be associated with carnivory

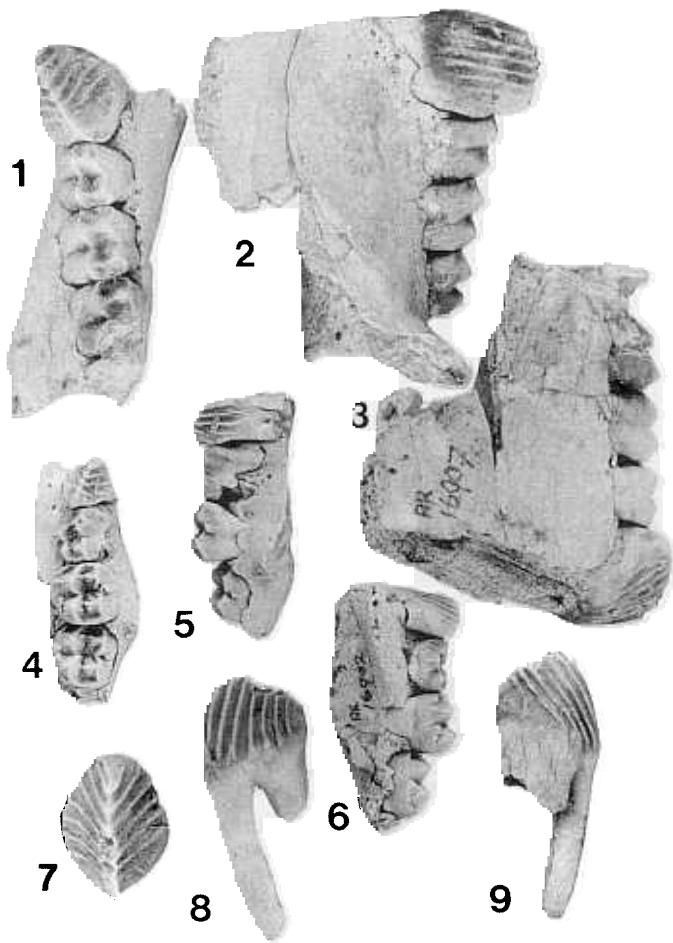


FIG. 4. *Ekaltadeta jamiemulvaneyi* x 2. A, occlusal view of QMF24200, left dentary containing  $P_3$ ,  $M_{1-3}$ , holotype. B, buccal view of QMF24200. C, lingual view of QMF24200. D, occlusal view of QMF24212, left maxillary fragment, containing  $P^2$ ,  $dP^3$ ,  $M^{1-2}$ , referred specimen. E, buccal view of QMF24212. F, lingual view of QMF24212. G, occlusal view of QMF20842, left  $P^3$ , referred specimen. H, buccal view of QMF20842. I, lingual view of QMF20849.

(Abbie, 1939), while a more extensive molar array may indicate a more herbivorous diet (Wells et al., 1982).

Species with a large molar surface area and low molar gradient (*P. oscillans*, *P. wellingtonensis*, *J. toxoniensis*) have relatively small premolars. Species with high molar gradients and reduced molar shear (*E. ima*, *P. chillagoensis*) are characterised by  $P_3$  hypertrophy. The extraordinary change in function for  $P_2$  shown by individ-

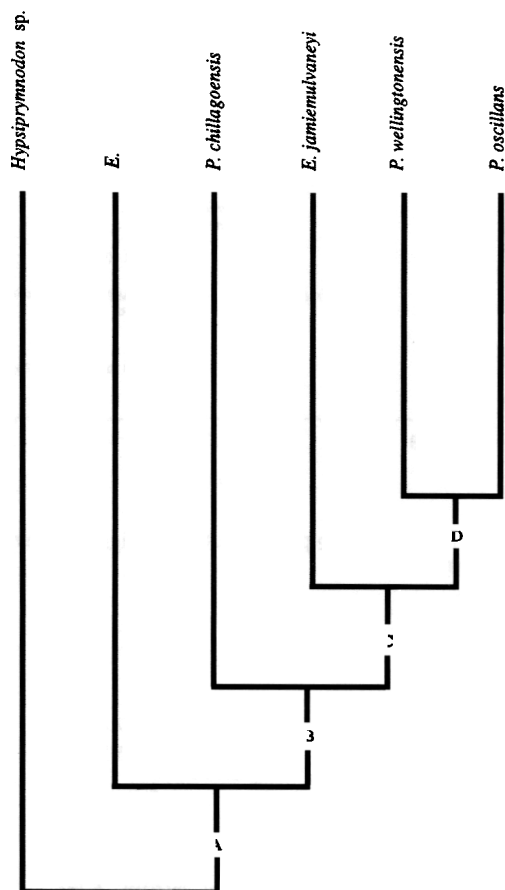


FIG. 5. Minimal tree produced by Wagner analysis for the Propleopinae (from Wroe, 1996). Character states at nodes: A = gain of an anterior cristid emanating from the metaconid of  $M_1$ ; basally broad conical upper molars; B = presence of lingual cingula on the upper molars; C = reduced molar gradient, reduced  $P_3$ ; D = incorporation of the protolophid into the anterior lophid of  $M_1$ , a dentary deeper posteriorly than anteriorly.

ual *E. ima* (Wroe & Archer, 1995) probably constitutes a response to the increased loading placed on  $P_3$ . Regarding molar gradient and relative size of the  $P_3$ , *E. jamiemulvaneyi* is intermediate, falling between *E. ima* specimens from lower levels and *P. oscillans*/*P. wellingtonensis*/*J. toxoniensis*. Using the same criteria *J. toxoniensis* lies between *E. jamiemulvaneyi* and *P. oscillans*/*P. wellingtonensis*. In terms of variation in  $P_3$  size and molar gradient *P. chillagoensis* and *P. oscillans* represent opposite extremes in propleopine evolution and it is suggested that *P. oscillans*

TABLE 1. Statistical summaries for  $M^2$  aw /  $M^2$  pw (G-value) for propleopines and a local *P. tridactylus* population.

	N	SD	CV	SE
Propleopines to level 8	13	0.05	4.53	0.01
<i>Ekaltadeta</i> to level 6	11	0.38	3.35	0.01
<i>Ekaltadeta</i> to level 4	9	0.02	2.07	7.71 E-3
<i>P. tridactylus</i>	16	0.02	1.70	4.61 E-3

was largely if not wholly herbivorous. Other derived features interpreted as adaptations to herbivory for *P. oscillans* include a large diastema between  $P_3$  and  $I_1$ , and large spatulate lower incisors (Wroe, 1996). Regarding dentary depth *E. ima* is the smallest propleopine with a general increase in depth for taxa at higher stratigraphic levels probably reflecting a general increase in body size.

Stratigraphic and metric analysis support the proposal of a late Miocene dichotomy in *Ekaltadeta* producing 2 lineages of *Propleopus*, and a reversal of previous assumptions on relative apomorphy within *Propleopus*, with *P. oscillans* considered the most derived and *P. chillagoensis* the most plesiomorphic (Wroe, 1996). However, broad trends suggested in this study are not interpreted here chronoclines in the stratophenetic sense (*sensu* Bown & Rose, 1987). The scarcity of material and uncertain chronology of both the Oligo-Miocene Riversleigh deposits and the Plio-Pleistocene local faunas from which most propleopine specimens are known necessitates caution in the interpretation of results. A considerable temporal gap exists between estimated ages of the most recent *Ekaltadeta* specimens and all other propleopines. As noted by Ride (1993), the period separating the latest known incidence of *Ekaltadeta* from Plio-Pleistocene *Propleopus* and *Jackmahoneya* may be sufficient to have permitted a secondary reversal of character states within *Propleopus* to produce *P. chillagoensis*.

Many questions remain concerning the age, stratigraphy and method of deposition of Riversleigh's Oligocene-Miocene limestone deposits (Archer, 1994, 1995; Megirian, 1992, 1994). If the phylogeny for propleopines suggested by Wroe (1996) reflects evolutionary events, then it provides tacit support for Archer et al.'s (1989) proposed stratigraphy, with an agreement of hypothesised superpositional and phylogenetic patterns.

The capacity of stratigraphic occurrence to explicitly mirror phylogenies is questionable (Engelmann & Wiley, 1977). Although strong

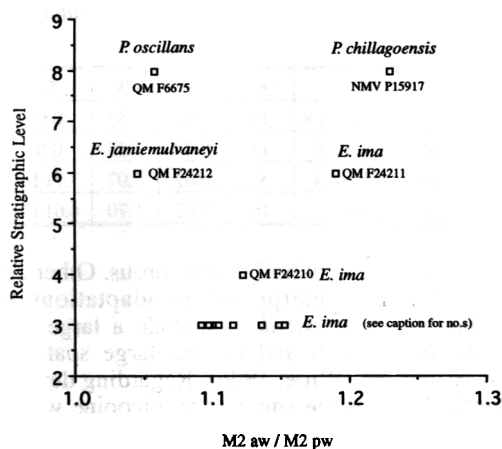


FIG. 6.  $M^2$  aw /  $M^2$  pw vs relative stratigraphic level for propleopines. *Ekaltadeta ima* from level 3, left to right, QMF24207, 24204, 24205, 12436, 24203, 24208, 24209, and 24206.

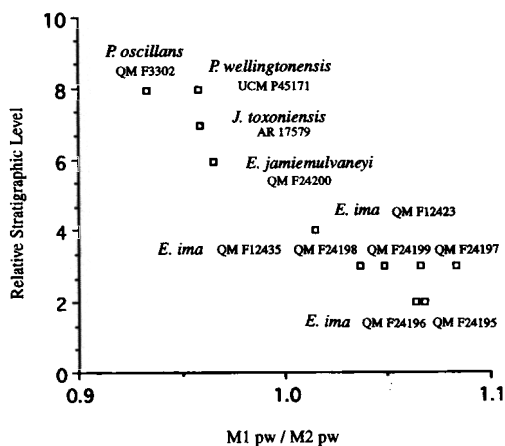


FIG. 7.  $M_1$  pw /  $M_2$  pw vs relative stratigraphic level for propleopines.

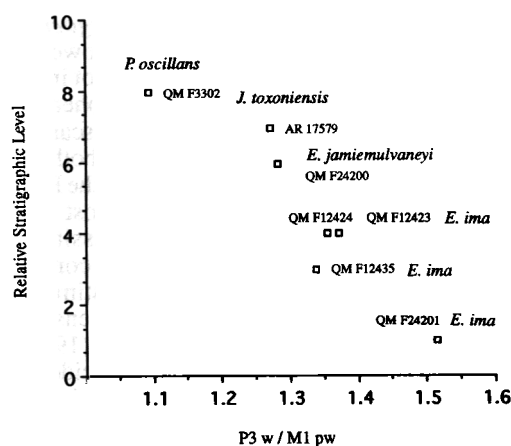


FIG. 8.  $P_3$  w /  $M_1$  pw vs relative stratigraphic level for propleopines.

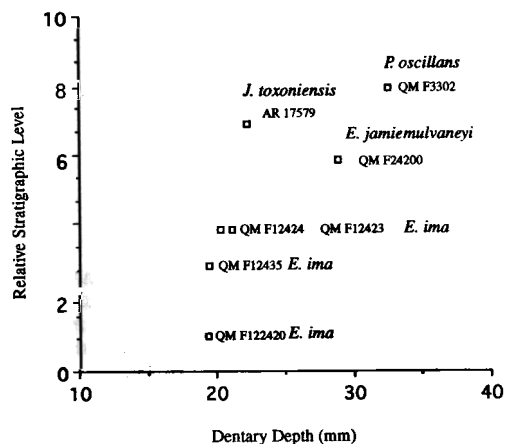


FIG. 9. Depth of dentary vs relative stratigraphic level for propleopines.

congruence between cladistic and stratigraphic arrangements has been demonstrated for many vertebrate taxa by Norell & Novacek (1992a,b), the same authors advised that correlation between the 2 diminishes rapidly where cladistic or stratigraphic data is poorly resolved. Debate over conformity of age and cladistic information commonly centres around the value of superpositional data as an adjunct to phylogenetic reconstruction. Where cladistic analysis is sound it

may be useful as a test of stratigraphic interpretations.

The propleopine phylogeny of Wroe (1996) is based on analysis of an incomplete data matrix, with important characters unknown for several species. Consequently the cladistic data presented cannot be viewed as a robust basis for testing superpositional pattern. However, the productivity of the Oligocene-Miocene deposits of

Riversleigh engenders reasonable expectation for the reliable resolution of phylogenies.

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### LITERATURE CITED

- ABBIE, A. 1939. A masticatory adaptation peculiar to some diprotodont marsupials. *Proceedings of the Zoological Society* 109: 261-279.
- ARCHER, M. 1994. Introduction: background and controversy about the Riversleigh geological and palaeontological resource. *Riversleigh Symposium 1994*, Abstracts: p 5-6.
- ARCHER, M., BARTHOLOMAI, A. & MARSHALL, L.G. 1978. *Propleopus chillaensis*, a new north Queensland species of extinct giant Rat-kangaroo (Macropodidae: Potoroinae). *Memoirs of the National Museum of Victoria* 39: 55-60.
- ARCHER, M. & FLANNERY, T.F. 1985. Revision of the extinct gigantic Rat-kangaroo (Potoroidae: Marsupialia), with description of a new Miocene genus and species, and a new Pleistocene species of *Propleopus*. *Journal of Paleontology* 59: 1331-1349.
- ARCHER, M., GODTHELP, H., HAND, S.J. & MERGIRIAN, D. 1989. Fossil mammals of Riversleigh, northwestern Queensland: preliminary overview of biostratigraphy, correlation, and environmental change. *Australian Zoologist* 25: 29-65.
- ARCHER, M., GODTHELP, H. & HAND, S.J. 1994. 'Riversleigh. The story of animals in ancient rain forests of inland Australia, Second Edition'. (Reed Books Pty Ltd. Sydney).
1995. Tertiary environmental and biotic change in Australia. Pp. 77-90. In Vrba, E.S., Denton, G.H., Partridge, T.C. & Burkle, L.H. (eds) 'Paleoclimate and evolution, with emphasis on human origins'. (Yale University Press: New Haven).
- BOWN, T.M. & ROSE, K.D. 1987. Patterns of evolution in early Eocene anaptomorphine primates (Omomyidae) from the Bighorn basin, Wyoming. *Journal of Paleontology* 23: 1-128.
- DE VIS, C.W. 1888. On an extinct genus of the marsupials allied to *Hypsiprymnodon*. *Proceedings of the Linnean Society of N.S.W.* 3: 5-8.
- ENGELMANN, G.F. & WILEY, E.O. 1977. The place of ancestor-descendant relationships in phylogeny reconstruction. *Systematic Zoology* 26: 1-11.
- FLANNERY, T. 1987. The relationships of the macropodoids (Marsupialia) and the polarity of some morphological features within the Phalangeriformes. Pp. 741-747. In Archer, M. (ed.) 'Possums and opossums: studies in evolution'. (Surrey Beatty & Sons and Royal Zoological Society of New South Wales: Sydney).
- FLOWER, W.H. 1867. On the development and succession of teeth in the Marsupialia. *Philosophical Transactions of the Royal Society of London* 157: 631-641.
- GINGERICH, P. D. 1976. Cranial anatomy and evolution of North American Plesiadapidae (Mammalia, Primates). *University of Michigan Papers on Paleontology* 15: 1-140.
1979. The stratophenetic approach to phylogeny reconstruction in vertebrate paleontology. (Columbia University Press: New York).
1990. Stratophenetics. Pp. 437-442. In Briggs, D.E.G. & Crowther, P.R. (eds), *Palaeobiology, a synthesis*. (Blackwell Scientific: Oxford).
- LUCKETT, P. 1993. An ontogenetic assessment in dental homologies in the therian mammals. Pp. 182-204. In Szalay, F.S., Novacek, M.J. & McKenna, m.c. (eds) 'Mammal Phylogeny: Mesozoic Differentiation, Multituberculates, Monotremes, Early Therians, and Marsupials'. (Springer-Verlag: New York).
- MEGIRIAN, D. 1992. Interpretation of the Carl Creek Limestone, northwestern Queensland. *The Beagle* 9: 219-248.
1994. Approaches to marsupial biochronology in Australia and New Guinea. *Alcheringa* 18: 259-274.
- MYERS, T. & ARCHER, M. 1997. *Kuterintja ngama* (Marsupialia, Ilariidae): a revised and extended systematic analysis based on fossil material from the late Oligocene of Riversleigh, northwestern Queensland, Australia. *Memoirs of the Queensland Museum* 41: 379-392.
- NORELL, M.A. & NOVACEK, M.J. 1992a. Congruence between superpositional and phylogenetic patterns: comparing cladistic patterns with fossil records. *Cladistics* 8: 319-337.
- 1992b. The fossil record and evolution: comparing cladistic and paleontologic evidence for vertebrate history. *Science* 255: 1690-1693.

- RIDE, W.D.L. 1993. *Jackmahoneya* gen. nov., and the genesis of the macropodiform molar. *Memoirs of the Association of Australasian Palaeontologists* 15: 441-459.
- WELLS, R.T. HORTON, D.R., & ROGERS, P. 1982. *Thylacoleo carnifex* Owen (Thylacoleonidae): Marsupial carnivore? Pp. 573-586. In Archer, M. (ed.) 'Carnivorous Marsupials'. (Surrey Beatty and Sons Pty Ltd and the Royal Zoological Society of New South Wales: Sydney).
- WROE, S. 1996. An investigation of phylogeny in the Giant Rat-kangaroo *Ekaltadeta* (Propleopinae, Hypsiprymnodontidae, Marsupialia). *Journal of Paleontology* 70: 681-690.
- WROE, S. & ARCHER, M. 1995. Extraordinary diphyodonty-related change in dental function for a tooth of the extinct marsupial *Ekaltadeta ima* (Propleopinae, Hypsiprymnodontidae). *Archives of Oral Biology* 40: 597-603.
- YOUNG, W.G., JUPP, R. & KRUGER, B.J. 1989. Evolution of the skull, jaws, and teeth in vertebrates. (University of Queensland Press: Brisbane).

## APPENDIX TO FIGURES 6-9

Data for Fig. 6.  $M^2$  aw divided by  $M^2$  pw (G-value) vs. relative stratigraphic level for propleopines. Measurements in mm. \* = skull, (R) = right tooth row, (L) =

Species	Cat. no.	$M^2$ aw	$M^2$ pw	G-value	Level
<i>E. ima</i>	QMF24203	6.80	6.10	1.12	3
<i>E. ima</i>	QMF24204	6.80	6.20	1.10	3
<i>E. ima</i> *	(R)QMF12436	6.40	5.80	1.10	3
<i>E. ima</i> *	(L)QMF12436	+	5.90		3
<i>E. ima</i>	QMF24205	6.50	5.90	1.10	3
<i>E. ima</i>	QMF24206	6.80	5.90	1.15	3
<i>E. ima</i>	QMF24207	7.20	6.60	1.09	3
<i>E. ima</i>	QMF24208	6.70	5.90	1.14	3
<i>E. ima</i>	QMF24209	6.20	5.40	1.15	3
<i>E. ima</i>	QMF24210	7.40	6.60	1.12	4
<i>E. ima</i>	QMF24211	6.90	5.80	1.19	6
<i>E. jamie mulvaneyi</i>	QMF24212	6.90	6.60	1.05	6
<i>P. oscillans</i>	QMF6675	9.20	8.70	1.06	8
<i>P. chillagoensis</i>	NMVP15917	10.7	8.70	1.23	8

Data for Fig. 7.  $M_1$  pw divided by  $M_2$  pw (G-value) vs. stratigraphic level. Measurements in mm.

Species	Cat. no.	$M_1$ pw	$M_2$ pw	G-value	Level
<i>E. ima</i>	QMF24195	6.70	6.30	1.06	2
<i>E. ima</i>	QMF24196	6.30	5.90	1.07	2
<i>E. ima</i>	QMF24197	6.50	6.00	1.08	3
<i>E. ima</i>	QMF24198	5.70	5.50	1.04	3
<i>E. ima</i>	QMF12435	6.50	6.20	1.05	3
<i>E. ima</i>	QMF24199	6.50	6.10	1.07	3
<i>E. ima</i>	QMF12423	7.00	6.90	1.01	4
<i>E. jamie mulvaneyi</i>	QMF24200	8.20	8.50	0.97	6
<i>P. wellingtonensis</i>	UCMP45171	9.20	9.60	0.96	8
<i>P. oscillans</i>	QMF3302	9.70	10.4	0.93	8
<i>J. toxoniensis</i>	AR17579	7.00	7.40	0.96	7

Data for Fig. 8.  $P_3$  w divided by  $M_1$  pw (G-value) vs. stratigraphic level for propleopines. Measurements in mm.

Species	Cat. no.	$P_3$ w	$M_1$ pw	G-value	Level
<i>E. ima</i>	QMF24201	10.3	6.80	1.56	1
<i>E. ima</i>	QMF12435	8.70	6.50	1.34	3
<i>E. ima</i>	QMF12424	8.80	6.50	1.35	4
<i>E. ima</i>	QMF12423	9.60	7.00	1.37	4
<i>E. jamie mulvaneyi</i>	QMF24200	10.5	8.20	1.28	6
<i>P. oscillans</i>	QMF3302	10.6	9.70	1.09	8
<i>J. toxoniensis</i>	AR17579	8.9	7.00	1.27	7

Data for Fig. 9. Depth of dentary vs. stratigraphic level for propleopines. Measurements in mm.

Species	Cat. no.	Dentary depth	Level
<i>E. ima</i>	QMF24201	19.3	1
<i>E. ima</i>	QMF12435	19.4	3
<i>E. ima</i>	QMF12424	21.1	4
<i>E. ima</i>	QMF12423	20.3	4
<i>E. jamie mulvaneyi</i>	QMF24200	28.9	6
<i>P. oscillans</i>	QMF3302	32.6	8
<i>J. toxoniensis</i>	AR17579	23.3	7