Bandicoot fossils and DNA elucidate lineage antiquity amongst xeric-adapted Australasian marsupials

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Bandicoots (Peramelemorphia) are a unique order of Australasian marsupials whose sparse fossil record has been used as prima facie evidence for climate change coincident faunal turnover. In particular, the hypothosed replacement of ancient rainforest-dwelling extinct lineages by antecedents of xeric-tolerant extant taxa during the late Miocene (~10 Ma) has been advocated as a broader pattern evident amongst other marsupial clades. Problematically, however, this is in persistent conflict with DNA phylogenies. We therefore determine the pattern and timing of bandicoot evolution using the first combined morphological + DNA sequence dataset of Peramelemorphia. In addition, we document a remarkably archaic new fossil peramelemorphian taxon that inhabited a latest Quaternary mosaic savannah-riparian forest ecosystem on the Aru Islands of Eastern Indonesia. Our phylogenetic analyses reveal that unsuspected dental homoplasy and the detrimental effects of missing data collectively obscure stem bandicoot relationships. Nevertheless, recalibrated molecular clocks and multiple ancestral area optimizations unanimously infer an early diversification of modern xeric-adapted forms. These probably originated during the late Palaeogene (30–40 Ma) alongside progenitors of other desert marsupials, and thus occupied seasonally dry heterogenous habitats long before the onset of late Neogene aridity.

Bandicoots (Peramelemorphia) are a speciose order of Australasian marsupials that appeared early in the evolutionary history of Australidelphia1. Most are small to medium sized (up to 5 kg) terrestrial omnivores occupying a spectrum of rainforest to desert habitats2,3. Molecular studies1,4 have defined three taxonomic subdivisions within the crown superfamily Perameloidea (Fig. 1): Chaeropodidae (pig-footed bandicoots), comprising the Central Australian dry-grassland, and possibly herbivorous Chaeropus ecaudatus, which became extinct as recently as the 1950’s; Thylacomyidae (bilbies), a monogenic classification for the genetically divergent6 Macrotis, which occurs in the arid and semi-arid zones of Australia7; and Peramelidae (typical bandicoots), an ecologically diverse radiation incorporating the primarily Australian Peramelinae and New Guinean (including surrounding islands and tropical far northern Australia) Peroryctinae and Echymiperinae8. DNA-based cladogenic scenarios for these groups envisage a latest Oligocene–early Miocene split (~20–30 Ma) between the xeric-adapted chaeropodids and thylacomyids versus predominantly mesic peramelids9,10. This contrasts with published morphological data, which posits both a late Miocene–Pliocene diversification of Australian Peramelinae concurrent with increasing aridity8,10, and late Miocene vicariant origins for Peroryctinae and Echymiperinae (historically united as Peroryctidae)11 accompanying New Guinean tectonic uplift. Such arguments are inferred from the fossil record, which has yielded no generically referable crown taxa older than the early Pliocene11,12. Fossil evidence of extant higher-level clades is also extremely sparse, consisting of isolated dental remnants from the middle Miocene putative basal peramelid Crash12 and thylacomyid Liyamayi12, as well as cranial-mandibular elements of the early–middle Miocene stem perameloids Madju13 and Kutjamarcoot14. Conversely, all other pre-Pliocene peramelomorphians are usually placed outside of Perameloidea, including the early Miocene Bulungu15,16.
Galadi\textsuperscript{17,18}, and a markedly plesiomorphic family-level grouping Yaralidae, which is composed of two species, \textit{Yarala burchfieldi} from the early–middle Miocene\textsuperscript{19,20}, and the late Oligocene \textit{Y. kida}\textsuperscript{21}. Yaralids are traditionally considered the basal sister radiation (= Yaraloidea)\textsuperscript{20} to all other bandicoots, and are thus important for calibrating molecular clocks within Peramelemorphia\textsuperscript{1,4} and Marsupialia as a whole\textsuperscript{22–24}. However, yaralids are united by a single unique symplesiomorphy – retention of a ‘complete’ centrocrista formed by the postparacrista and premetacrista on the upper molars\textsuperscript{21}. This has since been identified in a range of Oligocene – Miocene bandicoots\textsuperscript{16–18}, and is ancestral amongst marsupials\textsuperscript{25}, but otherwise lost in crown perameloids (defined by separated postparacristae–premetacristae) prompting assertions of a Miocene ‘bottleneck’ within Peramelemorphia, as well as other Australasian marsupials\textsuperscript{26}, whereby the radiation of crown taxa occurred only after the decline of rainforest-restricted stem forms, perhaps in response to climate change, ecological competition and/or vicariance events\textsuperscript{3,9,10,15,19–24,26,27}.

Here we test the long-standing hypotheses of Miocene faunal turnover and the recent origin of modern Australasian marsupials, by using the first total evidence morphological + multi-gene sequence dataset of Peramelemorphia incorporating examplars of all extinct and living genera named in the published literature. We also report on a remarkably primitive new fossil bandicoot from cave deposits on Pulau Kobroor in the Aru Islands, which has been radiocarbon and U-series dated to the latest Pleistocene–Holocene (28–9 Ka)\textsuperscript{28–31}. This significant discovery represents the geologically youngest stem-grade peramelomorphian, and implies not only extreme longevity but also undetected complexity affecting state acquisitions within this pivotal australidelphian order. Furthermore, our improved molecular clock constraints and coupled ancestral area analyses enable more precise timing for the nascent dispersal of crown bandicoots into open mosaic habitats, and thus refines burgeoning DNA-based evidence for Australia’s modern arid-zone marsupials as members of enduring, adaptable clades\textsuperscript{32}.

**Results**

Systematic Palaeontology
Marsupialia (Illiger, 1811) Cuvier, 1817
Australidelphia Szalay, 1982

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**Figure 1.** Geochronologically calibrated consensus hypothesis of peramelemorphian interrelationships. Colour gradient demarcates \textit{Lemdubioctyes aruensis} and other stem-grade bandicoots (pink) from xeric-adapted extant lineages (ochre), and the crown peramelid radiation (green). The peramelomorphian root node (●) and named subclades (■) are also correlated against a schematic of changing habitats through time. Dashed lines indicate phylogenetic uncertainty. †Fossil; *recently extinct arid-zone taxon. Graphics produced with Adobe CS5 by B. P. K.
Peramelemorphia (Kirsch, 1968) Aplin & Archer, 1987

**Lemdubuoryctes aruensis** gen. et sp. nov.

**Diagnosis.** *Lemdubuoryctes* is distinguished from all currently extant bandicoot genera (plus *Chaeropus*), the early Pliocene *cf. Peroryctes tedfordi*33, early–middle Miocene *Kutjamarcot*14, *Madju*15, *Liyamayi*15, *Crash*12, *Galadi amplus*15, and late Oligocene *Bulungu campbelli*16 by its retention of a ‘complete’ centrocrista with continuous postparacristae-premetacristae on all upper molars. The centrocrista is incomplete on M3 of the early Miocene *Galadi speciosus*17, and is formed by residual buccal crests on M3 of the early Miocene *Bulungu palara*15. *Galadi grandis*15 also from the early Miocene, and *Bulungu muirheadae*12 the oldest known late Oligocene bandicoot, possess ‘complete’ centrocristae along their upper molar rows but differ from *Lemdubuoryctes* in the presence of two mental foramina on the dentary, and the absence of anterior cingulae on M2–4 respectively. The lack of an elevated talonid separates *Lemdubuoryctes* from the latest Miocene–early Pliocene *Ischnodon*. Finally, oblique orientation of the posthypocristid relative to the lower molar row contrasts with late Oligocene–early Miocene *Yarala*19,21.

**Etymology.** ‘Lemdubu’ from the type locality, and ‘oryctes’ (ορυκτης, masculine) for ‘digger’; species name refers to its endemic occurrence on the Aru Islands.

**Holotype.** WAM 14.9.6, left maxilla containing P3 and M1–2.

**Additional material.** WAM 14.9.1–WAM 14.9.5 and WAM 14.9.7–WAM 14.9.20 (dentigerous elements, petrosals and calcanea).

**Type locality and horizon.** Liang Lemdubu cave (‘Layer 4’, Spit 24) on Pulau Kobroor, Aru Islands group, Eastern Indonesia29. Additional material was recovered from Spit 6, 14 and 18–25 (‘Layers 3–5’) at Liang Lemdubu, and Spits 33–40 (‘Layer 4’) in a second cave on Pulau Kobroor – Liang Nabulei Lisa29. A combination of radiocarbon and U-series age determinations place these deposits within the latest Pleistocene to Holocene, between ca 28,000–9,000 BP for Liang Lemdubu, and ca 16,200–12,000 BP for Liang Nabulei Lisa30,31.

**Description of the new fossil taxon.** Both the holotype (Western Australian Museum [WAM] 14.9.6) and referred (WAM 14.9.9) maxillae of *Lemdubuoryctes* (Fig. 2) display tooth eruption and molar wear indicative of adult animals (this is most extreme in WAM 14.9.9). The remnant palatal shelf on WAM 14.9.9 preserves a vacuity in the molar region. The base of the zygomatic arch is level with the alveolar margin. The antorbital fossa extends posteriorly from above the M3 to behind the M4; this differs from many extant peramelemorphians, as well as *Bulungu palara*15 and *Galadi speciosus*17, but can be intraspecifically variable13. Posterior expansion of the antorbital fossa

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**Figure 2.** *Lemdubuoryctes aruensis* referred (WAM 14.9.9) and holotype (WAM 14.9.6) maxillae. (a) SEM image of P3–M4 from WAM 14.9.9 shown in oblique occlusal view. (b) SEM image of the hypertrophied P3 and M1–M2 from WAM 14.9.6 in oblique occlusal view. Scale bars: (a) 3 mm, (b) 4 mm. Anatomical abbreviations: acd, anterior cingulid; ecf, ectoflexus; mcl, metaconule; mtc, metacone; pac, paracone; paf, area of the palatal fenestra; pmtc, premetacrista; ppac, postparacrista; pprc, postprotocrista; prc, protocone; rcc, residual centrocrista; stB, stylar cusp B; stylar cusp D. Imaging by K.P.A.
otherwise occurs only in living *Peroryctes* (Supplementary Fig. S1), some *Echymipera rufescens* (e.g. Australian Museum [AM] S1866)\(^{22}\), *Yarala burchfieldi*\(^{20}\), and osteologically mature *Madjyu*\(^{21}\) specimens. The antorbital fossa of *Macrotris* is uniquely elevated above the tooth row\(^{21}\). The anterior opening of the infraorbital canal in WAM 14.9.6 extends to the P3 alveolus (or the posterior margin of the M1 in WAM 14.9.9). This is similar to most bandicoots (including *B. palara*)\(^{15}\) in which the infraorbital canal usually has an expansive exit over the M1–P3. The infraorbital canal opens immediately above the P3 in *Y. burchfieldi*\(^{22}\) and species of *Galadji*\(^{27,28}\).

The upper premolars of *Lemdubuoryctes* are double-rooted with distemata between P1 and P2 (WAM 14.9.11), suggesting an elongate maxillary rostrum. The P3 exhibits marked size dimorphism consistent with sexual variation observed in some extant peramelids, most notably species of *Peroryctes* and *Echymipera* (Supplementary Fig. S1). We therefore interpret WAM 14.9.6 as a probable male because the P3 exceeds the M1 in occlusal area (Fig. 2B). The P3 is smaller than the M1 in WAM 14.9.9 and thus represents a potential female (Supplementary Table S1). Both P3 morphotypes are otherwise identical in their triangular basal outline with conical central cusp, weak posteroangular cingulum and incipient anterobasal cuspule.

At up to 14.89 mm in length (Supplementary Table S1), the complete M1–M4 row of *Lemdubuoryctes* (WAM 14.9.9) is equal to the largest living bandicoot *Peroryctes broadbenti*\(^{34}\). The M1 (Fig. 2; Supplementary Fig. S2) is triangular in occlusal outline unlike the more quadrangular molars of peramelines, *Chaeropus* and *Macrotris*; the latter further distinguished by extreme lingual displacement of the metacone. In *Lemdubuoryctes*, the metacone is positioned at the posteroangular margin of the trigon basin, which is bounded buccally by the paracone. A prominent protocone is situated lingually. There is no protoconule. The metaconule (metacanula) is laterally weak only a small spur that connects the postprotocrista to the base of the metacone. This is compatible with extreme metacanical reduction seen in the stem peramelomorphs *Yarala*\(^{9,21}\), *Bulungu*\(^{15,16}\), and *Galadji*\(^{27,18}\), together with the early Pliocene cf. *Peroryctes tedfordi*\(^{33}\). Alternatively, living bandicoots\(^{22}\) as well as species of *Crash*\(^{13}\), *Madjyu*\(^{13}\) and *Kutjammarcool*\(^{11}\) elaborate the metaconule into an enamel flange that is demarcated from the protocone via a vertical trough (Supplementary Fig. S1). Dasyuromorphian marsupial carnivores have a more prominent cusp-like metacone. Amongst bandicoots only *Macrotris* completely lacks a metacanical structure, but a small metaconule is present in the putative thylacomyid *Liyanangi*\(^{12}\).

The anterior cingulum on the M1 of *Lemdubuoryctes* is formed by the preprotocrista, which connects to the parastyle base. There is no posterior cingulum. The paracone lies directly behind the parastyle and the preparacrista runs posterobuccally towards the parastyle tip (a common trait amongst *Peramelemorphia*: see the parastylar base. There is no posterior cingulum. The paracone lies directly behind the parastyle and the preparacrista runs posterobuccally towards the parastyle tip (a common trait amongst *Peramelidae*: see Supplementary Data, character 14). The postparacrista merges with the premetacrista to create a ‘complete’ centrocrista. The opposing buccal ectoflexus is shallowly incised between the remnants of stylar cusps B and D (there is no discernible stylar cusp E). Remarkably, there are broad ectolophs evident on the M2 and M3 that closely resemble those of the most ancient peramelomorphs *Yarala*\(^{9,21}\), *Bulungu*\(^{15,16}\), and *Galadji*\(^{27,18}\). In other fossils, the postparacrista and premetacrista gradually retract resulting in a ridge-like centrocrista on the M3 of *B. palara*\(^{15}\), and total division of the ectoloph in *Bulungu campbelli*\(^{16}\), *Galadji*\(^{18}\), *Madjyu*\(^{13}\) and *Kutjammarcool*\(^{11}\). Remnants of the centrocrista also occur on the M1–3 of extant *P. broadbenti* (Supplementary Fig. S1), less prominently on the M1–2 of *Peroryctes raffrayana*, and occasionally in *E. rufescens*\(^{16}\). Crests appear elsewhere on the M1–2 of *Crash*\(^{12}\) and cf. *P. tedfordi*\(^{33}\), which also has a small ‘stylar’ cusp C, perhaps constituting another terminal component.

The M2 and M3 of *Lemdubuoryctes* (Fig. 2; Supplementary Fig. S2) differ from the M1 in their lingually positioned paracone, less distinct ectoflexus, and transversely oriented preparacrista that trends towards stylar cusps B and D, but retains contact with the parastyle tip via a subsidiary crest. This forms a truncated anterior cingulum comparable to that on the M2–3 of *P. raffrayana*\(^{34}\).

The M4 of *Lemdubuoryctes* (Fig. 2; Supplementary Fig. S2) is reduced relative to the anterior molars and bears both a paracone and diminutive protocone. An anterobuccal cingulum is not visible but could be covered by matrix in WAM 14.9.9. The postprotocrista forms the posterior margin of the trigon basin and meets the postparacrista at stylar cusps B.

Mandibular elements were referred to *Lemdubuoryctes* based on obvious morphological distinction from the sympatric bandicoots* Isoodon macrourus, E. rufescens* and *E. kalulu*. Relative hypertrophy of the p3 differentiates presumed male (WAM 14.9.1) and female (WAM 14.9.3) specimens (Fig. 3; Supplementary Fig. S3). The mandibular rami of *Lemdubuoryctes* (Supplementary Fig. S4) are ventrally convex and up to 9.7 mm deep below the m3 (WAM 14.9.3). The single mental foramen is level with the midline of p1, and the mandibular symphysis extends to the middle of p2. The ascending ramus in WAM 14.9.1 is angled at ~45°; the mandibular foramen opens low on the medial surface and the masseteric fossa is well defined. The i3 root on WAM 14.9.3 is separated from the canine alveolus (3.9/1.8 mm in maximum length/width) by a 2 mm diastema. Another diastema (3.5 mm) separates between c1 and p1 with a narrower gap between p1 and p2. The length and height of p1–3 decrease anteriorly (Supplementary Table S2) and are coupled with progressive migration of the blade-like central cusp forward over the anterior root. There are no accessory cusps.

The complete m1–4 row of *Lemdubuoryctes* (Fig. 3; Supplementary Fig. S3) was up to 17.43 mm long (WAM 14.9.3: Supplementary Table S2), with marked constriction evident at the enamel crown–root interface (also visible on p1–3); this is typical of peramelomorphs except for *Isoodon* and *Macrotris*\(^{27}\). The m1 is laterally compressed with a bulbous trigonid incorporating a prominent paracanid, which is absent in *Ishmodon*\(^{37}\), living peramelines and *Echymipera*\(^{27}\). There are no median buccal cusps between the trigonids and talonids as reported in *Y. burchfieldi*\(^{19}\). The anterior cingulid is reduced on m1–4 and the labial cingulids are weakly developed, similar to *Peroryctes* (Supplementary Fig. S1). The cristid obliqua terminates buccally against the posterior wall of the protoconid on m1–2 (rather than the metaconid as in many peramelines\(^{27,28}\) but is more lingually positioned on the m3, and immediately adjacent to the metacristid notch on m4. The hypoconulid is situated distally anterior to the entoconid, and sunken well below the talonid basin on all lower molars (synapomorphies for *Peramelemorphia*\(^{27}\)). The posthypocristid is oblique to the molar row like that of *P. raffrayana*\(^{34}\).
Figure 3. *Lemdubuoryctes aruensis* referred dentaries. (a) SEM image of p1–p3, and m2–m4 from WAM 14.9.1 shown in oblique occlusal view. (b) p3 and m1–m4 from WAM 14.9.3 in oblique occlusal view. Scale bars: 4 mm. Anatomical abbreviations: cob, cristid obliqua; dia, diastema; ecd, entoconid; hcd, hypoconid; hcdl, hypoconulid; mcd, metaconid; pcd, paraconid; phcd, posthypocristid; prcd, protoconid, tal, talonid; tgd, trigonid. Imaging by K.P.A.
and the respective classifications of *Kutjamarcoot* and *Crash* as either a stem perameloid or stem peramelids\(^\text{1,3}\), were likewise uncertain (Supplementary Figs S11 and S12). Similarly, the earliest dated peroryctine\(^\text{1,4}\), *cf. Peroryctes tedfordi*, was only intermittently nested within *Peroryctes* (Supplementary Fig. S10), a result that compromises existing molecular clock calibrations\(^\text{1,4}\). The affinity of *Perameles bowensis*\(^\text{3,15}\), which has also previously been used for dating constraints\(^\text{1,4}\), was unresolved\(^\text{33}\) (Supplementary Figs S8 and S9); however, this taxon manifests a posthypocristid-entoconid contact on its lower molars, which is distinctive for both *Perameles* and the Pleistocene species *P. sobbei*\(^\text{16}\) (topographically grouped with *Perameles nasuta* and *P. gunnii*: Supplementary Figs S8 and S9), as well as the enigmatic Miocene–Pliocene genus *Ischnodon*. Notably, neither *Ischnodon* nor *Liyamayi* were recovered as ancestral bilbies\(^\text{1,12,16}\), and constraint tests on these placements were inconclusive (Supplementary Table S3). Such results corroborate the original taxonomic assessment of *Ischnodon*, which reported thylocomyid dental similarities but refrained from definitive classification\(^\text{12}\).

Our assessments of outant bandicoot morphology were consistent with DNA\(^\text{1,4}\) in returning the xeric-adapted *Macrotris lagotis* as the most divergent living peramelemorphian (Supplementary Fig. S13). On the other hand, alternate grouping of the extinct arid-zone chaeropodid *Chaeropus* within Peramelinae (Supplementary Fig. S14) suggests that either extensive dental/osteological convergence\(^\text{3}\), or incomplete characterization of its scant mitochondrial sequence data\(^\text{30}\) confound its relationships. Interestingly, inclusion of the extinct desert-adapted thylocomyid, *Macrotris leucura*, promoted extensive topological degradation (Supplementary Fig. S15). This might be due to its curious ‘peramelid-like’ dental attributes (see matrix scores in Supplementary Data), which could again denote either homoplasy, or the retention of ancestral perameloid states.

Trees generated by the concatenated dataset of morphology + DNA were identical to those produced by DNA alone\(^\text{1}\), but with amplified support values for weak nodes demonstrating overall signal congruence. This was most notable at the nodes excluding *Macrotris* from *Bulungu* (Supplementary Table S4). Successful deletion of molecular information for major clades\(^\text{44}\) pinpointed residual morphological conflict over a paraphyletic *Peroryctinae* + *Echymiperinae*, and repositioning of *Chaeropodidae* within *Peramelinae* (Supplementary Table S4). This concurs with previous studies\(^\text{14-17,45}\), which have placed *Macrotris* and *Chaeropus* outside of *Peramelinae* using a molecular backbone, but not with morphology on its own. As expected, the introduction of fossils completely degraded node support (Supplementary Fig. S17), and revealed long-branch effects in the clumped redistribution of taxa (Supplementary Fig. S18). Sequential deletion of highly homoplastic dental–dependent terminals did improve these results (Supplementary Figs S19 and S20), but still failed to yield stable positioning of fossils, perhaps because they integrate insufficient cranial–postcranial skeletal data to accurately discriminate relationships.

**Divergence times and ancestral areas.** We utilized a DNA dataset with expanded outgroup sampling of diprotodontian, notoryctemorphian and dasyuromorphian taxa to correlate the timeframes and settings for peramelemorphian intra-clade divergences. Alternative fossil constraints (Supplementary Table S5) and Bayesian random local clocks\(^\text{46}\) (Supplementary Fig. S21) were also implemented to assess possible sources of overestimation\(^\text{16}\). Despite these tests, our analyses demonstrated an unequivocal origination of the crown bandicoot total-group during the mid-Paleocene around 60 Ma (Table 1; Supplementary Tables S6–S8; Supplementary Figs S22–S30). This corroborates the discovery of possible stem peramelemorphian fossils from the early Eocene\(^\text{1,2}\), but massively predates previous molecular estimates\(^\text{1,4}\) by up to 40 Ma. In accordance, diversifications amongst chaeropodid, thylocomyid and peramelid family-level clades seem to have commenced in the middle Eocene (~40–30 Ma). These epochs coincide with the tectonic isolation of Australia and instigation of the circum-Antarctic current, which propagated seasonally cool–dry climates and the spread of sclerophyllous vegetation\(^\text{42}\). Compellingly, our S-DIVA/Bayesian Binary MCMC ancestral area optimisations onto morphological (Supplementary Figs S31 and S32), total evidence (Supplementary Fig. S33), and DNA trees (Supplementary Fig. S34) decisively correlated the basal peramelemorphian split with dispersal into open habitats and an early occupation of xeric ecosystems (Supplementary Tables S9–S12). Unanimous inference of a post early–middle Miocene (after ~20 Ma) rainforest–woodland radiation amongst peroryctines and echymiperines likewise coincides with uplifting of the New Guinean landmass and onset “greenhouse” climates\(^\text{48}\), which propagated higher rainfall and coastal/riparian vegetation\(^\text{50}\). In contrast, our analyses failed to pinpoint an emergent habitat for peramelines. We attribute this to their rapid expansion into openly vegetated environments\(^\text{1,3,51}\), compounded by methodological dependence of our probability matrices upon predefined species distribution codes. These are particularly sensitive to highly dispersive organisms, as well as significant area changes through time\(^\text{52}\). In our case this included the pronounced middle–late Miocene (after ~16 Ma) resurgence of cool-dry climates, and Pliocene predominance of mosaic vegetation, especially incorporating intra–continental grasslands which proliferated across Australia during this interval\(^\text{16}\).

**Discussion**

The ‘complete’ centrocristae delimiting broad ectolophs, and extreme metaconular reduction on the M1–3 of *Lembuduoryctes* are virtually identical to the conditions found in the most ancient fossil bandicoots *Yarala kida*\(^\text{21}\) and *Bulungu muirheadae*\(^\text{16}\). As shown here, these unexpected state expressions have radical implications for bandicoot phylogeny in placing *Lembuduoryctes* as an exceptionally late-surviving stem-grade peramelemorphian. Moreover, the presence of both residual centrocrista and metaconules on the upper molars of the early–middle Miocene *Bulungu*\(^\text{15,16}\) and *Galadi*\(^\text{17,18}\), as well as the early Pliocene *cf. Peroryctes tedfordi*\(^\text{33}\), and extant species of *Peroryctes and Echymipera*\(^\text{24}\) shows that these symplesiomorphies were persistent throughout bandicoot evolution, and could represent examples of repeated convergent atavism. Although postulated\(^\text{15,16,18,27}\), such rampant homoplasy has never previously been demonstrated within the fundamental discriminative features of the peramelomorphian dentition. Equally as significant is our topological nesting of *Bulungu campbelli* amongst living perameloids\(^\text{33}\),
which implies a corresponding appearance of advanced dental traits within the stratigraphically earliest bandicoot lineages. *Bulungu campbelli* is a late Oligocene species (*Etadunna Formation Zone C*: 24.6–24.1 Ma) that approximates the oldest known fossil peramelemorphian taxon *B. mutirheadae* (*Etadunna Formation Zone B*: 24.9–24.6 Ma). *Bulungu campbelli* is also important because it predates what is usually regarded as the most plesiomorphic bandicoot *Yarala kita* (*Wipajiri Formation equivalent: 24.1 Ma*). However, the stratigraphical horizons containing *Bulungu campbelli* fossils date from prior to inundation of the Torresian Plain after the Last Glacial Maximum (26–31). At this time, the Aru Islands were a limestone plateau surrounded by open savannah plains with dense riparian forest restricted to topographic lows along fault-controlled ‘sungai’ channels. The fossil bandicoot species from these settings are dominated by both *Lemdubuoryctes* and *Isodon macrourus*, the latter being an extant grassland–open woodland inhabitant. On the other hand, *Echymipera rufescens* which presently occupies lowland rainforests on the Aru Islands is comparatively rare, and *E. kalulu* which typifies rainforests and anthropogenic grasslands in high rainfall areas, has been tentatively identified from a few teeth but these post-date the late Pleistocene Aru Islands marine transgression. The numerical abundance of *Lemdubuoryctes* at Liang Lemdubu and Liang Nabulei Lisa, coupled with palynomorph evidence, and its associated open savannah–moist forest vertebrate assemblage, could therefore suggest a preference for heterogeneous habitats. This pointedly compliments zoogeographic correlations of the late Pleistocene Aru Islands with mosaic ecotones in northern Australia and the Trans–Fly region of southern New Guinea, as well as the reconstructed palaeoenvironments ascribed to other plesiomorphic *Pliocene–Pleistocene bandicoots*.

## Table 1. Molecular divergence date estimates for peramelemorphian clades (million years BP).

| Node                                      | Divergence estimates                                      | No constraints | Ingroup constraints |
|-------------------------------------------|-----------------------------------------------------------|----------------|---------------------|
| Dasyuromorpha + Peramelemorphia          | 60.68 (54.61–65.7)                                        | 60.34 (54.74–65.79) |
| Chaeropus v Macrotis + Peramelidae        | 37.87 (29.78–46.05)                                       | 38.06 (30.14–46.32) |
| *Peroryctinae v Peramelidae*              | 31.45 (25.38–38.03)                                       | 31.55 (25.76–38.24) |
| *Peroryctinae v Echymiperinae*            | 20.24 (16.41–24.23)                                       | 20.49 (16.77–24.43) |
| *Peroryctinae v Echymiperinae*            | 17.37 (13.96–21.15)                                       | 17.55 (14.08–21.08) |
| *Peroryctes broadbeniti v P. raffrayana*   | 8.01 (5.12–11.4)                                          | 8.08 (5.08–11.62) |
| *Blychomelus + Echymipera v Microperoryctes* | 12.08 (9.53–14.97)                                       | 12.21 (9.62–15.14) |
| *Blychomelus v Echymipera*                | 10.6 (7.81–13.62)                                         | 10.7 (7.92–13.8)  |
| *E. clara v E. kalulu v E. rufescens*     | 9.76 (7.36–12.51)                                         | 9.85 (7.36–12.59) |
| *E. kalulu v E. rufescens*                | 6.34 (4.22–8.73)                                          | 6.4 (4.18–8.85)   |
| *Microperoryctes sp. (Tembagapura) v M. ornata v M. longicauda (Sol) + M. papuensis* | 8.11 (5.83–10.58)                                       | 8.18 (5.84–10.74) |
| *M. ornata v M. longicauda (Sol) v M. papuensis* | 4.95 (3.29–6.9)                                         | 5.01 (3.35–6.95)  |
| *M. ornata v M. longicauda (Sol) + *I. macrourus* | 2.16 (1.16–3.4)                                         | 2.17 (1.15–3.42)  |
| *Isodon v Perameles*                      | 13.2 (10.34–16.5)                                        | 13.73 (10.84–16.91) |
| *I. auritus v I. macrourus + I. obesus*   | 5.18 (3.42–7.2)                                          | 5.25 (3.46–7.3)   |
| *I. macrourus v I. obesus*                | 5.27 (4.94–9.93)                                         | 5.31 (4.94–9.4)   |
| *Peramelinae v Peroryctinae*              | 11.38 (8.62–14.44)                                        | 11.83 (9.14–14.8) |
| *T. bougainvillei v P. eremiana v P. gunnii + P. nasuta* | 8.5 (5.07–12.15)                                       | 8.81 (5.27–12.51) |
| *P. gunnii v P. nasuta*                   | 4.71 (2.81–7.04)                                         | 4.77 (2.82–7.12)  |

The survival of *Lemdubuoryctes* on what is today a rainforest prevalent island refuge, seemingly accords with the most ancient peramelemorphian habitats. This key premise underlies the Miocene ‘bottleneck’ hypothesis, under which environmentally constrained stem taxa were replaced by crown perameloid lineages, such as those belonging to the terminal Pleistocene–Holocene. In addition, our demonstration of profound antiquity for modern desert-living bandicoot lineages (a result unaffected by alternative constraint parameters...
or variation in substitution rate\(^4\); Supplementary Tables S6–S8; Supplementary Figs S21–S30), indicates that increasing aridity during the late Neogene likely did not initiate the genesis of crown Peramelidea, although it probably assisted in peramelina intra-clade habitat expansion and localized speciation events. The undeniable rarity of definitive crown perameloid fossils in pre-Pliocene sediments might therefore be explained by sampling biases and/or incompletely documented collections\(^3,27,58,59\), as well as ecological underrepresentation\(^60\). Indeed, our tree-based ancestral area optimisations (Supplementary Tables S9–S12; Supplementary Figs S31–S34) infer that the seminal radiation of modern bandicoots accompanied widespread australidelphian niche dispersals into drier mosaic settings, perhaps such as mallee (Eucalyptus) woodlands that spread through Central Australia from the late Oligocene\(^60,61\). The scarcity of crown bandicoot antecedents in intensively studied fluvial or karstic contexts\(^54,57,59\) with higher preservation potential thus becomes understandable, as does the ecologically disjunct DNA-based phylogeny of living peramelomorphs. This now clearly captures one of the most deeply divergent radiations of xeric-adapted marsupials\(^1,3,4,22–24,32\), and reinforces a biota-wide exaptive response to late Neogene aridity\(^62\), including diversification amongst clades that had already maintained substantial habitat disparity for many millions of years.

Methods

Dataset construction. Peramelemorphian morphological phylogenies have suffered from persistently inadequate resolution\(^12–18\) prompting weighting of dental data via homologous sets\(^4\) and incremental qualitative sub-divisions\(^43\). We therefore compiled a de novo matrix that emphasized partition sampling\(^5\) across 93 cranial-dental and postcranial characters assembled for demonstrable extant outgroup dasyurid/didelphid marsupials\(^66\) and 36 ingroup bandicoot species including 13 fossil taxa: Bulungu campbelli and Yarala kida (late Oligocene), Bulungu palara, Kutjamarcoot brevirostrum, Madju variae (late Oligocene—early Miocene), Crash bandicoot, Galadji amplus, Galadji speciosus, Kutjamarcoot brevirostrum, Liamaya dayi and Yarala burchfieldii (middle Miocene), Ichnodon australis, cf. Peroryctes tedfordi and Perameles bowensis (early Pliocene), P. sobbei (late Pleistocene), and Ledmuboryctes aruensis (late Pleistocene—Holocene). Scores were derived from original specimens (all crown taxa, G. speciosus, Y. burchfieldii, I. australis, P. sobbei, L. aruensis), high-definition casts (B. campbelli), and/or reference to relevant published works (Y. kida, B. palara, K. brevirostrum, M. variae, C. bandicoot, G. amplus, L. dayi; the type references of cf. P. tedfordi and P. bowensis could not be located by their housing institutions). State definitions were either redrafted from earlier studies (1–63)\(^11,18,25,27,34,37,67\), or determined exclusively for this phylogeny (64–93). A complete character list with annotations describing modified state definitions, redundancy and/or resoring, the molecular data matrix, and a catalogue of examined specimens are provided in the Supplementary Data.

The concatenated series of 9977 DNA sequence nucleotides representing five nuclear (ApoB, BRCAl, IRBP, RAG1, vWF) and three mitochondrial genes (12S rRNA, cytochrome b and the 3’ portion of 16S rRNA) was used to: (1) produce a backbone tree (Supplementary Fig. S7) that determined the best-supported position of fossils relative to the living species topology; and (2) compute a combined (non-weighted) total evidence analysis that examined effects of morphological data on molecular nodes. Laboratory procedures, DNA sequence derivation/alignment, and model testing of separate gene/codon regions were described in Westerman et al.\(^11,18\).

Phylogenetic analysis. We implemented a six-stage strategy to manage the detrimental effects of incomplete fossils and characters\(^19\). (1) Initial selection of operational taxonomic units [OTUs] specifically targeted extant taxa with overlapping coverage of morphological and DNA sampling\(^9\). Fossil OTUs included only the most complete genus-level exemplars for branching lineages (Bulungu\(^14\), Galadji\(^10\), Madju\(^11\)) as well as those species critical for molecular clock calibrations (Yarala kida, cf. Peroryctes tedfordi, Perameles bowensis)\(^3,4\) or uncontested crown clade referral (Perameles sobbei)\(^3,8,45\). (2) Excessively incomplete DNA characters were removed in morphology-only analyses to examine the effects of extinct taxa with numerous missing entries\(^8\). A manual screen for redundant taxa (safe taxonomic reduction\(^40\)) identified all OTUs that degraded strict consensus resolution and pinpointed instability caused by missing data versus character conflict\(^75\). (3) The ‘amb’ option was implemented during all PAUP\(^*\) v4.0b10\(^73\) parsimony searches to eliminate ambiguous zero length branches\(^74\). (4) A posteriori screening of wildcard taxa produced a strict reduced consensus profile based on the semi-strict Adams consensus (where wildcards do not obscure adequately supported nodes\(^40,41\)) and assessments of relative character support at affected nodes\(^74\). (5) Bootstrapping and branch (Bremer) decay indices were alternately employed with and without wildcard exclusion to test the impact of mobile OTUs upon support measures. (6) Sequential exclusion of fossils incorporating numerous missing entries for cranial and postcranial characters was used to assess the effects of sub-sampling and long-branch attraction\(^76\) within the total evidence framework\(^70\).

 Parsimony trees and bootstrap frequencies (1000 repetitions) were computed using heuristic searches with TBR (tree-bisection-reconnection) branch swapping and 100 random-addition replicates. The molecular scaffold enforced monophyly for clades receiving ≥70% partitioned maximum likelihood bootstrap, and ≥0.95 Bayesian posterior probability support. Bremer values were calculated for unconstrained morphological data with TNT v1.1\(^75\), which also cross-referenced MP topologies via a 'New Technology Search' with sectorial searches, drift, and tree fusing enabled. Results were then processed using a 'Traditional Search' option with TBR. DELTRAN character state optimisation was preferentially employed for tree construction, but unequivocal synapomorphies were shared by both DELTRAN and ACCTRAN outputs (see Supplementary Data).

Bayesian analyses used MrBayes v3.2\(^4\) with the obligate standard discrete model (Mrv\(\gamma\))\(^77\) for morphology. Because this assumes equal change between all character states, we set the rates variation parameter to ‘gamma’ (with coding to ‘variable’), thereby introducing heterogeneity, and also tested data partitioning via cranial/postcranial (1–6, 32–55, 59, 65), dental (7–31, 56–58, 60–64), and postcranial (66–93) character sets\(^65\). Gene partitions
followed Westerman et al.\textsuperscript{1,48}. Two simultaneous runs and four Markov Chains (one cold and three heated using default heating values) were applied for $5 \times 10^8$ generations with default settings and the burn-in fraction set at 0.25.

**Molecular clocks.** Time-trees were generated in BEAST\textsuperscript{29} with uncorrelated relaxed lognormal clocks. A Bayesian random local clock model was also used to test for overestimation imposed by punctuated shifts in substitution rate\textsuperscript{46}. We used traditional node dating to assess contested\textsuperscript{16} ingroup constraints for Peramelemorpha, Chaeropodidae, Thylacomyidae, Peroryctinae, Echymiperinae, Peramelinae and Isoodon + Peramelaceae. Minimum and maximum fossil calibrations (Supplementary Table S5) were compiled according to best practice protocols\textsuperscript{79}. Analyses incorporated a birth-death model and normal priors imposed on soft bounds with 95% distribution between the minimum and maximum. MCMC analyses were run for $5 \times 10^8$ generations with a 25% burn-in for tree summaries. Runs were terminated when ESS values reached $>200$ for all estimated parameters.

**Ancestral areas.** Because the precise topological placement and habitat preferences of fossil peramelemorphians are uncertain, we used a non-ultrametric Bayesian DNA tree sub-sample of extant taxa, and alternative S-DIVA (tree dataset) versus Bayesian Binary MCMC (condensed tree accommodating topological uncertainty) approaches implemented in RASP ver. 3.2\textsuperscript{100} to infer ancestral habitat dispersal patterns. Four Markov Chains (default heating values) were run twice over $5 \times 10^8$ generations with sampling frequency and burn-in fixed at 500. Among-site rate variation was set to ‘gamma’ and state frequencies utilized a ‘fixed (JC)’ model. Habitat codings were generalized to accommodate for non-exclusivity, but broadly adhered to defined vegetation units\textsuperscript{50}: A = rainforest; B = woodland (referring primarily to sclerophyll forests); C = shrubland (including ‘open’ and xeromorphic vegetation); and D = arid/semi-arid vegetation and desert.

**Nomenclatural acts.** ZooBank (http://zoobank.org/) registration of nomenclatural acts in this published work can be accessed using the Life Science Identifier: urn:lsid:zoobank.org:pub:7AC0A046-A507-4AA2-933D-F1A36132F2FD.

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**Author Contributions**

B.P.K., K.P.A. and M.W. wrote the manuscript. B.P.K. prepared the figures. K.P.A. prepared the fossil material, compiled the descriptions, photography and SEM imaging. B.P.K. and M.W. performed the phylogenetic, molecular clock, and ancestral areas analyses.

**Additional Information**

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