1	
2	An Early Nimravid from California and the Rise of Hypercarnivorous Mammals after the Middle
3	Eocene Climatic Optimum
4	
5	
6	ASHLEY W. POUST ^{1,2} , PAUL Z. BARRETT ³ , and SUSUMU TOMIYA ^{2,4,5,6}
7	¹ Department of Paleontology, San Diego Natural History Museum, San Diego, California,
8	92101, USA
9	² University of California Museum of Paleontology, Berkeley, California, 94720, USA
10	³ Department of Earth Sciences, University of Oregon, Eugene, OR, 97403, USA
11	⁴ Center for International Collaboration and Advanced Studies in Primatology, Kvoto University
12	Primate Research Institute, Inuvama, Aichi, 484-8506, Japan
13	⁵ Center for the Evolutionary Origins of Human Behavior, Kvoto University, Inuvama, Aichi,
14	484-8506, Japan
15	⁶ Negaunee Integrative Research Center, Field Museum of Natural History, Chicago, Illinois,
16	60605, USA
17	
18	OrcIDs: AWP 0000-0001-7955-613X; PZB 0000-0001-5717-9193; ST 0000-0001-6038-899X
19	
20	
21	Subject Areas: palaeontology, evolution, taxonomy and systematics
22	Keywords: sabretooth, carnassial, Hoplophoneus, Duchesnean, evolution, Eocene-Oligocene
23	biotic transition
24	
25	RH: POUST et al.—EARLY NIMRAVID
26	
27	Carnivoraforms (crown carnivorans and their closest relatives) first occupied hypercarnivorous
28	niches near the dawn of the late Eocene, 40-37 million years ago. This followed the decline or
29	extinction of earlier carnivorous groups, Mesonychia and Oxyaenodonta, leaving carnivoraforms
30	and hyaenodontan meat-eaters as high trophic level consumers. The pattern of this change and
31	the relative contributions of the taxonomic groups has hitherto been unclear. We report a new
32	genus and species of the sabretoothed mammalian carnivore family Nimravidae, Pangurban
33	egiae, from the Eocene Pomerado Conglomerate of southern California, with strongly derived
34	hypercarnivorous features. While geochronologically the oldest named nimravid in North
35	America, Pangurban egiae is recovered as phylogenetically derived, with affinities to
36	Hoplophoneus. This provides unequivocal evidence for rapid radiation and spread of nimravid
37	carnivores across Asia and North America and constrains the timing of early divergences within
38	the family. Pangurban egiae narrows the gap between convergent iterations of sabretoothed
39	mammalian carnivores and demonstrates swift diversification of the hypercarnivorous nimravids
40	during a period of global climatic instability. Furthermore, it highlights the top-to-bottom
41	restructuring North American ecosystems underwent during the Eocene-Oligocene transition,
42	resulting in carnivoraforms taking positions as trophic specialists for the first time, a niche they
43	still occupy today.

45 Electronic supplementary material is available at http://dx.doi.org/XXXXX or via

46 <u>http://rsbl.royalsocietypublishing.org</u>.

47

48 1. Introduction

49 The climatically dynamic period from the late middle Eocene to the early Oligocene (ca. 41–30 50 Ma) saw not only the reorganisation of biotas worldwide that is known as the Eocene–Oligocene biotic transition [1] but also the initial rise of members of crown Carnivora and their closest 51 52 relatives as hypercarnivores [2,3]. The early nimravids joined the Hyaenodonta in this specialised 53 niche, which carnivorans would later occupy alone until the present day. Due to the eventual 54 extinction of these groups and the resulting lack of genetic data, fossils provide the only window 55 into an important period of faunal change presaging the organisation of modern trophic structure. 56 However, the lack of early data from this critical period has presented a challenge to

interpreting this transition. In particular, a general paucity of mammalian fossils from the
Duchesnean North American Land Mammal 'Age' (NALMA; 41–37 Ma) has long obscured an

early phase of the Eocene–Oligocene faunal transition in North America, particularly with
 respect to mammalian carnivores [4]. Recent taxonomic and biostratigraphic studies, however,

have unveiled the late-middle-Eocene diversity of carnivores in southwestern North America
 sufficiently to permit its reassessment [5–9]. The turnover of mammalian carnivores during this
 period is noteworthy for the replacement of early meat-eating lineages by crown-clade

64 carnivorans and their close carnivoraform relatives—an episode that broadly coincided with the 65 global climatic fluctuations surrounding the Middle Eocene Climatic Optimum (MECO; [9]).

We report a new genus and species of nimravid, *Pangurban egiae*, on the basis of a
partial maxilla with the upper 3rd and 4th premolars (P3–4) (Figure 1) from the likely upper
middle Eocene portion of the Pomerado Conglomerate of San Diego County, California. *Pangurban* is one of the earliest diagnosable nimravids in the world, and preserves the earliest
nimravid cheek teeth from North America, allowing its inclusion in phylogenetic analysis.

71 Bearing similarities to the derived later nimravid Hoplophoneus, its early occurrence and

72 hypercarnivorous adaptations demonstrate the rapid diversification of the group.

73 2. Methods

74 The new fossil is housed at the San Diego Natural History Museum (San Diego Society 75 of Natural History collections, SDSNH). We scored the holotype specimen (SDSNH 60554) into 76 the Bayesian tip-dating analysis of Barrett (2021) [10], with a uniform tip-date of 40.0–37.0 Ma 77 based on the likely middle- to late-Duchesnean age of the specimen. The age of the holotype 78 (and the only known specimen) of Maofelis cantonensis was updated to the late Eocene, 79 following [11]. The Bayesian analysis utilized a fossilized birth-death process [12–14] with 80 complex models of morphological evolution (e.g. ordered, multipath, Dollo) best supported for 81 the nimravid character matrix of [10]. Additional details can be found in the electronic 82 supplementary material. Markov Chain Monte Carlo runs were sampled every 1,000 generations 83 in Beast 2 v. 2.6.3 ([15]), while trees were logged every 10,000 generations until Effective 84 Sample Sizes were > 200, as determined in Tracer version 1.7.1 [16]. From the output of Tracer, 85 sampling and timing (generation number) of the stability of the run was assessed. Thus, the first 86 10% of the generations were discarded (burn-in), and topology and posterior probabilities were

- 87 estimated from the remaining generations. The maximum clade credibility (MCC) tree of this
- analysis is presented in Figure 2a (barbourofelins and most outgroups not shown; see Fig. S2 for
- 89 complete tree) and the electronic supplementary material, along with the XML run file. We
- 90 reconstructed synapomorphies in TNT version 1.5, which only reports synapomorphies
- supported by both accelerated and delayed transformations. Body mass of the new taxon was
- 92 estimated through linear regression analysis of P4 lengths and corresponding nimravid body
- masses (based originally on lower first molar [m1] lengths) in [10]. We estimated ancestral body
- 94 masses along the branches of the MCC tree using the 'fastANC' function found in the 'phytools'
- 95 [17] package for the R programming environment, v. 4.1.2 (R Development Core Team, 2021).
- 96

97 3. Results

- 98 3.1. Systematic Palaeontology
- 99
- 100 Class MAMMALIA sensu Rowe [18]
- 101 unranked clade CARNIVORAMORPHA sensu Bryant [19] (amended from Wyss and Flynn
- 102 [20])
- 103 unranked clade CARNIVORAFORMES Flynn, Finarelli, and Spaulding [21]
- 104 Family NIMRAVIDAE sensu Bryant [22]
- 105 *Pangurban egiae* gen. et sp. nov.
- 106 Fig. 1
- 108
- 109 **Holotype.** SDSNH 60554, right maxillary fragment with P3–P4; collected in 1997 by Stephen L.
- Walsh and Robert Gutzler, who also prepared the specimen, and housed at the San Diego NaturalHistory Museum (San Diego, U.S.A.).
- 112 **Type locality and horizon.** SDSNH Locality 4042 ("Spring Canyon Site 2"). Upper member of
- 113 Pomerado Conglomerate, San Diego, California, U.S.A., here considered middle to late
- 114 Duchesnean in age (ca. 40–37 Ma; see SI Text).
- 115 **Diagnosis**. Anterior cutting edge of P4 expressed as sharp and distinct preparacristal blade, not
- 116 reaching apex of paracone; anterior face of P4 flattened and nearly vertical giving the paracone a
- 117 steep angle. Differs from: *Maofelis* and MA-PHQ 348 (undet. nimravid specimen housed at
- 118 Museum d'Histoire Naturelle Victor Brun, Montauban, France [23]) in infraorbital foramen
- posterior to P3, double rooted P3, P3 width to length ratio 0.4–0.65; *Maofelis*, MA-PHQ 348,
- and "nimravines" of [10] which includes barbourofelins, in reduced or absent P4 protocone, P4
- 121 metastylar blade length to paracone length ratio of 1.0–1.3, P3 crown height 48–70% that of P4;
- hoplophoneins (*Hoplophoneus*, *Eusmilus*, and *Nanosmilus*) in absence of parastyle on P4.
- 123 **Etymology.** The generic name is from the Old Irish, in reference to the cat in the 9th century CE
- 124 poem of the same name, whose hunting is likened to academic pursuit of truth. The species name
- 125 honours the contribution of palaeontologist Naoko Egi to the study of terrestrial carnivore
- 126 evolution on both sides of the northern Pacific Ocean, mirroring the nimravid distribution.
- 127 **Remarks.** SDSNH 60554 was previously identified as "cf. *Hyaenodon* sp." in the collection, and
- 128 was apparently the basis for the previously reported occurrence of that hyaenodontid genus in the 129 upper Pomerado Conglomerate [24]. We re-identify this specimen as a nimravid based on the

130 combination of: low height of P3; presence of serration on P3; elongate posterior basal cusp of

- P3; absence of cuspidate protocone on P4; absence of parastyle from the anterolabial base of P4
- (cf. [25,26]); presence of prominent P4 preparacrista; and absence of deep incision associated
 with carnassial notch on P4 (cf. [25]).
- 134

135 3.2. Morphology and Phylogeny

136

137 The double-rooted P3 measures 11.2 mm in length and 4.6 mm in width. It is labiolingually compressed, and its concave lingual wall descends steeply to its base without 138 139 forming an appreciable lingual bulge (Fig. 1; SI Fig. 2). The anterior slope of the paracone forms 140 a shallow angle with the palate near its base, but steepens toward the tip of the cusp. The anterior 141 ridge is finely serrated (~3 slightly dorsally-inclined incisions per mm on preserved portion) except for ~1 mm at the base of the crown. No anterior cuspule is present. The presence of a 142 143 prominent and likely trenchant posterior basal cusp, separated from the main cusp by a notch, is 144 indicated by a groove on the labial wall and the outlines of the breakages.

145 The largely intact P4 measures 16.8 mm in length and 12.6 mm in width, which yielded 146 an estimated body mass of 28 kg. Its anterolingual projection lacks a distinct protocone. The 147 parastyle, cingular or otherwise in origin (cf. [27]), is absent. The preparacrista is notably 148 trenchant. The carnassial notch lacks a deep incision. The metastylar blade is posterolabially 149 deflected. There is no appreciable development of a cingulum. A very light facet of attritional 150 wear is visible in inclined light on the lingual side of the P4. It surrounds the carnassial notch and 151 rises onto the postparacrista, generally resembling wear found in *Dinictis* in both extent and 152 severity.

153 The posteroventral curvature of the infraorbital foramen is preserved above the anterior 154 edge of the P4. This differs from most "nimravines" of [10] and *Eusmilus*, in which it is located 155 more anteriorly. A deep embrasure pit for the occlusion of the m1 protoconid is partially 156 preserved lingual to the metastylar blade of P4.

157 In the MCC tree (Fig. 2*a*), we recovered *Pangurban egiae* as the sister taxon to the node-158 based tribe Hoplophoneini sensu Barrett [28], with which it was united by three

159 synapomorphies: reduced or absent P4 protocone; P4 metastylar blade 1.0–1.3 times the length

160 of paracone; and P3 crown height 48–70% of that of P4. All of these features would appear again

161 in Miocene barbourofelins, and are thus not unambiguous synapomorphies, but they do indicate

162 comparably derived hypercarnivory within this clade in the late middle Eocene.

- 163 The remaining topology of the MCC tree differed minimally from [10]. The
- 164 hoplophoneins were recovered in largely the same arrangement, with successive divergences of
- 165 *Hoplophoneus* (here recovered as a paraphyletic taxon), *Nanosmilus*, and *Eusmilus*.

166 4. Discussion

167

Published occurrences of sabretooth material have generally suggested a middle Eocene
origin of nimravids within East Asia [11,29]. Subsequent dispersal(s) to North America later in
the Eocene generated a cradle of diversification in disparate ecologies, from which additional

immigrations into Asia and Europe seem to have occurred [10,11]. The holotype of *Pangurban*

172 egiae, SDSNH 60554, is significant as one of the oldest securely-identified nimravid fossils

- 173 globally (Fig. 2*b*; see SI Text for further discussion). Although a few likely-older (ca. 40 Ma)
- specimens are known from the Pondaung Formation of Myanmar [30,31] and the Clarno
- Formation of Oregon, USA [32,33], it is the oldest nimravid fossil that preserves enough of the
- 176 upper dentition to be meaningfully incorporated into a phylogenetic analysis. While not
- 177 falsifying the hypothesis of East Asian origin for nimravids, the presence of a derived member in
- 178 North America at such an early time demonstrates not only swift circum-Pacific dispersal for the 179 group (in whatever direction), but also rapid evolution of the earliest nimravids. As such, it
- 180 provides key information on the early evolution and distribution of the family.
- 181 On the MCC tree, *Pangurban egiae* displays a zero-length branch, indicating that it is a 182 sampled ancestor. This status relates to *P. egiae* preserving derived character states that support 183 its affinity with Hoplophoneini, while not preserving distinct apomorphies in this matrix. 184 However, ~42% of sampled posterior trees have a tip-length greater than zero for *P. egiae*, 185 indicating that its status as a true sampled ancestor should be treated with some hesitancy—a 186 status that will require additional specimens to properly assess. In view of the unique 187 combination of traits in SDSNH 60554, we prefer to assign it to a new genus.
- 188 The discovery of *Pangurban egiae* accentuates the Eocene–Oligocene taxic turnover 189 across all trophic levels and highlights the late-middle-Eocene rise of hypercarnivorous 190 mammals in southwestern North America. Taking pronounced reduction of the carnassial 191 metaconid(s)—which is associated with a predominantly anteroposterior orientation of the 192 prevallid—and its correlates as taxon-independent proxies for hypercarnivory [4,34,35], 193 hypercarnivores are greatly outnumbered (both in species richness and abundance) by hypo- and 194 mesocarnivores in the late Uintan of southern California and western Texas combined (age 195 groups 1 & 2 in Fig. 2c; Table S1). The beginning of the Duchesnean NALMA is marked by the 196 immigration from Asia of the hypercarnivorous/durophagous Hyaenodon ([36–38]; age group 3 197 in Fig. 2c); interestingly, the earliest Duchesnean deposits of southern California have yielded 198 abundant fossils of Hyaenodon, whereas remains of carnivoraforms (none of whom are 199 hypercarnivores) are rare [6,36,39]. Middle to late Duchesnean assemblages from southwestern 200 North America contain more hypercarnivorous taxa (including *P. egiae*) than hypo- to 201 mesocarnivorous taxa (age group 4 in Fig. 2c).
- 202 The shifting balance within the carnivore guild cannot be readily explained as a 203 taphonomic artefact, and it seems likely to reflect ecological restructuring of the carnivore guild 204 across the climatically unstable earlier Duchesnean. Notably, the same time period saw a marked 205 decline in primate diversity in the region ([40]; Fig. 2c) and, in southern California, a $\sim 40\%$ loss 206 of primarily woody angiosperm taxa [41]. Substantial disassembly and reassembly of a regional 207 carnivore community concomitant with a major loss of primate diversity also characterises the 208 late Bridgerian-early Uintan (ca. 48-44 Ma) faunal succession in the central Rocky Mountain 209 region [42]. These events may reveal common faunal dynamics over evolutionary time in 210 response to reduced forest canopies and spatial redistribution of biomass, within the context of 211 the middle-to-late Eocene trend of declining precipitation in North America [43]. Perhaps the rise of carnivoraform hypercarnivores relates to the development of seasonally open canopies or 212 213 mosaic forested landscapes. Following the loss of sabretooth machaeroidines prior to the Middle 214 Eocene Climatic Optimum [44], a contingent opportunity may have occurred for circum-Pacific 215 immigration of nimravids into North America. Concerted declines in relatively small, arboreal 216 vertebrate prey such as primates (resulting in an upward shift in the body size distribution of 217 vertebrate prey) and other food resources that are abundant in tropical/subtropical forests (e.g.,

- 218 fruits and insects) may have prompted evolution of, or invasion by, larger carnivoraforms with
- 219 more specialised adaptations for carnivory, as dictated by feeding energetics (cf.[45,46]). In view
- of the relative stability of the morphological composition of terrestrial mammals [47] and
- 221 carnivores in particular [48] across the Eocene–Oligocene boundary in North America,
- 222 *Pangurban egiae* points to the late-middle to early-late Eocene as a dynamic period of key
- 223 importance in the Eocene–Oligocene biotic transition and the origin of phylogenetically-modern
- 224 terrestrial carnivore guilds.
- 225

226 ACKNOWLEDGMENTS

- 227 We thank Kesler A. Randall and Thomas A. Deméré for SDSNH collections access; Gabriel
- 228 Vogeli for photogrammetry; Shawn P. Zack (University of Arizona), E. Bruce Lander (Paleo
- 229 Environmental Associates, Inc.), Anthony R. Friscia (UCLA), Patricia A. Holroyd (University of
- 230 California Museum of Paleontology,), and Naoko Egi (National Museum of Nature and Science,
- 231 Japan) for enlightening discussions; Kazue Takai (Kyoto University) for reference support;
- 232 reviewers Fabien L. Condamine and Steven Zhang and editors for helpful comments on the
- 233 manuscript; Joseph D. McDowell (Augustana College) for the introduction to Irish and
- 234 mediaeval literature: go raibh maith agat as do meantóireacht!
- 235

236 FIGURE CAPTIONS

237

Fig 1. Holotype SDSNH 60554 of *Pangurban egiae* gen. et sp. nov.

239 Right maxillary fragment with P3–P4 in lateral (a) and ventral (b) views. Approximate position

- 240 on cranium marked on (c).
- 241

Fig. 2. Phylogenetic, geographic, and faunal context of *Pangurban egiae*.

243 (a) Tip-dated Bayesian MCC tree of nimravids. Node values indicate posterior probabilities of

clades. Nimravid basal node marked by open circle (median age) and bar (95-percent credible

- interval). (*b*) Geographic distribution of Eocene nimravids (in part after [11,29]). Coloured (by land mammal 'ages' as in (*a*)) portions of icons indicate known dental elements. Taxonomically-
- 246 and mammal ages as in (*a*)) portions of icons indicate known dental elements. Taxonom 247 questionable occurrences marked as "?"; nimravid material from Gongkang Formation
- 248 apparently has never been described (cf. [49]). (c) Late-middle-Eocene carnivore (tooth icons)
- and primate (silhouettes) diversity in southern California and western Texas, divided into four
- age groups (data in Table S1; tooth icons modelled after *Lycophocyon hutchisoni* and
- 251 *Diegoaelurus vanvalkenburghae*) and based primarily on [5,6,8,36,39,40,44,50–53]). Time
- scales for magnetochrons and NALMAs follow [33,54], and that for Asian Land Mammal
- 253 'Ages' (ALMAs) after [55]. NALMA/ALMA abbreviations: Ch, Chadronian; Du, Duchesnean;
- Er, Ergilian; Ir, Irdinmanhan; Sh, Sharamurunian; Ui, Uintan; Ul, Ulangochuian. Timings of
- middle Eocene global climatic anomalies and angiosperm diversity loss in southern Californiafrom [56] and [41].
- 256 257
- Prothero DR, Berggren WA, editors. 1992 Eocene-Oligocene climatic and biotic
 evolution. Princeton University Press.
- Van Valkenburgh B. 1999 Major patterns in the history of carnivorous mammals. *Annu. Rev. Earth Planet. Sci.* 27, 463–493. (doi:10.1146/annurev.earth.27.1.463)

262 3. Sole F, Fischer V, Le Verger K, Mennecart B, Speijer RP, Peigne S, Smith T. 2022 263 Evolution of European carnivorous mammal assemblages through the Palaeogene. Biol. J. 264 *Linn. Soc.* **135**, 734–753. (doi:10.1093/biolinnean/blac002) 265 4. Wesley-Hunt GD. 2005 The morphological diversification of carnivores in North America. Paleobiology 31, 35–55. (doi:10.1666/0094-266 8373(2005)031<0035:tmdoci>2.0.co;2) 267 268 5. Tomiya S. 2011 A new basal caniform (Mammalia: Carnivora) from the middle Eocene of 269 North America and remarks on the phylogeny of early carnivorans. PLoS One 6. (doi:10.1371/journal.pone.0024146) 270 Tomiya S. 2013 New carnivoraforms (Mammalia) from the middle Eocene of California, 271 6. 272 USA, and comments on the taxonomic status of 'Miacis' gracilis. Palaeontol. Electron. 273 16. (doi:10.26879/364) 274 7. Tomiya S, Tseng ZJ. 2016 Whence the beardogs? Reappraisal of the middle to late eocene 275 'miacis' from Texas, USA, and the origin of amphicyonidae (mammalia, carnivora). R. 276 Soc. Open Sci. 3. (doi:10.1098/rsos.160518) 277 8. Lander EB. 2019 Early late Duchesnean (late middle Eocene) Titus Canyon fauna, Titus 278 Canyon Formation, Death Valley National Park, Inyo County, southeastern California. In 279 Exploring Ends of Eras in the Eastern Mojave Desert, Desert Symposium Field Guide and 280 Proceedings: Desert Hot Springs, California, Desert Symposium Inc, pp. 141–153. 281 Tomiya S, Morris ZS. 2020 Reidentification of Late Middle Eocene "Uintacyon" From 9. 282 the Galisteo Formation (New Mexico, U.S.a.) As an Early Beardog (Mammalia, 283 Carnivora, Amphicyonidae). Breviora 567, 1. (doi:10.3099/0006-9698-567.1.1) Barrett PZ. 2021 The largest hoplophonine and a complex new hypothesis of nimravid 284 10. 285 evolution. Sci. Rep. 11, 1–9. (doi:10.1038/s41598-021-00521-1) 286 Averianov A, Obraztsova E, Danilov I, Skutschas P, Jin J. 2016 First nimravid skull from 11. Asia. Sci. Rep. 6. (doi:10.1038/srep25812) 287 288 Stadler T. 2010 Sampling-through-time in birth-death trees. J. Theor. Biol. 267, 396–404. 12. 289 (doi:10.1016/j.jtbi.2010.09.010) 290 13. Didier G, Royer-Carenzi M, Laurin M. 2012 The reconstructed evolutionary process with 291 the fossil record. J. Theor. Biol. 315, 26–37. (doi:10.1016/j.jtbi.2012.08.046) 292 Stadler T, Kühnert D, Bonhoeffer S, Drummond AJ. 2013 Birth-death skyline plot reveals 14. 293 temporal changes of epidemic spread in HIV and hepatitis C virus (HCV). Proc. Natl. 294 Acad. Sci. U. S. A. 110, 228–233. (doi:10.1073/pnas.1207965110) 295 15. Bouckaert R, Heled J, Kühnert D, Vaughan T, Wu CH, Xie D, Suchard MA, Rambaut A, 296 Drummond AJ. 2014 BEAST 2: A Software Platform for Bayesian Evolutionary Analysis. 297 *PLoS Comput. Biol.* **10**, 1–6. (doi:10.1371/journal.pcbi.1003537) 298 16. Rambaut A, Drummond AJ, Xie D, Baele G, Suchard MA. 2018 Posterior summarization 299 in Bayesian phylogenetics using Tracer 1.7. Syst. Biol. 67, 901–904. 300 (doi:10.1093/sysbio/syy032) 301 17. Revell LJ. 2012 phytools: An R package for phylogenetic comparative biology (and other 302 things). Methods Ecol. Evol. 3, 217–223. (doi:10.1111/j.2041-210X.2011.00169.x) 303 Rowe T. 1988 Definition, diagnosis, and origin of mammalia. J. Vertebr. Paleontol. 8, 18. 304 241–264. (doi:10.1080/02724634.1988.10011708) 305 19. Bryant HN. 1996 Explicitness, stability, and universality in the phylogenetic definition and usage of taxon names: A case study of the phylogenetic taxonomy of the Carnivora 306 307 (mammalia). Syst. Biol. 45, 174–189. (doi:10.1093/sysbio/45.2.174)

- Wyss AR, Flynn JJ. 1993 A Phylogenetic Analysis and Definition of the Carnivora.
 Mammal Phylogeny, 32–52. (doi:10.1007/978-1-4613-9246-0_4)
- Flynn J, Finarelli J, Spaulding M. 2010 Phylogeny of the Carnivoramorpha, and the use of
 the fossil record to enhance understanding of evolutionary transformations. In *Carnivoran Evolution: New Views on Phylogeny, Form and Function* (eds A Goswami, A. Friscia),
 pp. 25–63. Cambridge Unversity Press.
- Bryant HN. 1996 Nimravidae. In *The Terrestrial Eocene-Oligocene Transition in North America* (eds DR Prothero, JR Emry), pp. 453–475. Cambridge University Press.
- Peigné S. 2001 A primitive nimravine skull from the quercy fissures, France: Implications for the origin and evolution of nimravidae (Carnivora). *Zool. J. Linn. Soc.* 132, 401–410.
 (doi:10.1111/j.1096-3642.2001.tb02467.x)
- Walsh SL, Gutzler RQ. 1999 Late Duchesnean-early Chadronian mammals from the
 upper member of the Pomerado Conglomerate, San Diego, California. J. Vertebr.
 Paleontol. 19(3), 82A.
- Flynn JJ, Galiano H. 1982 Phylogeny of Early Tertiary Carnivora, With Description of a
 New Species of Protictis From the Middle Eocene of Northwestern Wyoming. *Am. Museum Novit.*, 1–64.
- 325 26. Bryant HN. 1991 Phylogenetic Relationships and Systematics of the Nimravidae
 326 (Carnivora). J. Mammal. 72, 56–78.
- Peigné S. 2003 Systematic review of European Nimravinae (Mammalia, Carnivora,
 Nimravidae) and the phylogenetic relationships of Palaeogene Nimravidae. *Zool. Scr.* 32,
 199–229. (doi:10.1046/j.1463-6409.2003.00116.x)
- Barrett PZ. 2016 Taxonomic and systematic revisions to the North American Nimravidae
 (Mammalia, Carnivora). *PeerJ* 2016. (doi:10.7717/peerj.1658)
- Egi N, Tsubamoto T, Saneyoshi M, Tsogtbaatar K, Watabe M, Mainbayar B, Chinzorig T,
 Khatanbaatar P. 2016 Taxonomic revisions on nimravids and small feliforms (Mammalia,
 Carnivora) from the Upper Eocene of Mongolia. *Hist. Biol.* 28, 105–119.
 (doi:10.1080/08912963.2015.1012508)
- 336 30. Tsubamoto T *et al.* 2006 A summary of the Pondaung fossil expeditions. *Asian Paleoprimatology* 4, 1–66.
- 338 31. Zaw K *et al.* 2014 The oldest anthropoid primates in SE Asia: Evidence from LA-ICP-MS
 339 U-Pb zircon age in the Late Middle Eocene Pondaung Formation, Myanmar. *Gondwana*340 *Res.* 26, 122–131. (doi:10.1016/j.gr.2013.04.007)
- 341 32. Hanson CB. 2010 Stratigraphy and vertebrate faunas of the Bridgerian-Duchesnean
 342 Clarno Formation, north-central Oregon. In *The Terrestrial Eocene-Oligocene Transition*343 *in North America* (eds D Prothero, J Emry), pp. 206–239. Cambridge University Press.
 344 (doi:10.1017/cbo9780511665431.012)
- 345 33. Kelly TS, Murphey PC, Walsh SL. 2012 New Records of Small Mammals From the
 346 Middle Eocene Duchesne River. *Paludicola* 8, 208–251.
- 347 34. Butler BPM. 1946 The evolution of carnassial dentitions in the Mammalia. *Proc. Zool.*348 Soc. London 116, 198–220.
- 349 35. Van Valen L. 1969 Evolution of Dental Growth and Adaptation in Mammalian
 350 Carnivores. *Evolution (N. Y).* 23, 96–117.
- 351 36. Mellett JS. 1977 Paleobiology of North American Hyaenodon. Karger Publishers.
- 352 37. Bastl K, Morlo M, Nagel D, Heizmann E. 2011 Differences in the tooth eruption sequence 353 in Hyaenodon ('Creodonta': Mammalia) and implications for the systematics of the genus.

- 354 *J. Vertebr. Paleontol.* **31**, 181–192. (doi:10.1080/02724634.2011.540052)
- 355 38. Bastl K, Semprebon G, Nagel D. 2012 Low-magnification microwear in Carnivora and
 dietary diversity in Hyaenodon (Mammalia: Hyaenodontidae) with additional information
 on its enamel microstructure. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 348–349, 13–20.
 (doi:10.1016/j.palaeo.2012.05.026)
- 359 39. Tomiya S. 2012 Ecological Aspects of the Diversity Dynamics of North American Fossil
 360 Mammals. University of California, Berkeley.
- 40. Williams BA, Kirk EC. 2008 New Uintan primates from Texas and their implications for
 North American patterns of species richness during the Eocene. J. Hum. Evol. 55, 927–
 941. (doi:10.1016/j.jhevol.2008.07.007)
- Frederiksen NO. 1991 Pulses of middle Eocene to earliest Oligocene climatic
 deterioration in southern California and the Gulf Coast. *Palaios* 6, 564–571.
 (doi:10.2307/3514919)
- Tomiya S, Zack SP, Spaulding M, Flynn JJ. 2021 Carnivorous mammals from the middle
 Eocene Washakie Formation, Wyoming, USA, and their diversity trajectory in a postwarming world. J. Paleontol. 95, 1–115. (doi:10.1017/jpa.2020.74)
- 43. Eronen JT, Janis CM, Chamberlain CP, Mulch A. 2015 Mountain uplift explains
 differences in palaeogene patterns of mammalian evolution and extinction between north
 America and Europe. *Proc. R. Soc. B Biol. Sci.* 282. (doi:10.1098/rspb.2015.0136)
- 373 44. Zack SP, Poust AW, Wagner H. 2022 Diegoaelurus, a new machaeroidine (Oxyaenidae)
 374 from the Santiago Formation (late Uintan) of southern California and the relationships of
 375 Machaeroidinae, the oldest group of sabertooth mammals . *PeerJ* 10, e13032.
 376 (doi:10.7717/peerj.13032)
- 45. Carbone C, Mace GM, Roberts CR, Macdonald DW. 1999 Energetic constraints on the
 diet of terrestrial carnivores. *Nature* 402, 286–288.
- 46. Van Valkenburgh B, Wang X, Damuth J. 2004 Cope's rule, hypercarnivory, and
 extinction in North American canids. *Science (80-.).* 306, 101–104.
 (doi:10.1126/science.1102417)
- 47. Prothero DR, Heaton TH. 1996 Faunal stability during the early oligocene climatic crash. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 127, 257–283. (doi:10.1016/S0031-0182(96)00099-5)
- 48. Valkenburgh B Van. 1994 Extinction and Replacement Among Predatory Mammals in the
 North American Late Eocene and Oligocene: Tracking a Paleoguild Over Twelve Million
 Years. *Hist. Biol.* 8, 129–150. (doi:10.1080/10292389409380474)
- Tang Y, Qui Z. 1979 Vertebrate faunas of Baise, Guangxi. In *Mesozoic and Cenozoic Red Beds of South China* (ed Academia Sinica Institute of Vertebrate Paleontology and
 Paleoanthropology and Nanking Institute of Geology and Paleontology), pp. 407–415.
 Science Press.
- 50. López-Torres S, Silcox MT, Holroyd PA. 2018 New omomyoids (Euprimates, Mammalia)
 from the late Uintan of southern California, USA, and the question of the extinction of the
 paromomyidae (plesiadapiformes, primates). *Palaeontol. Electron.* 21. (doi:10.26879/756)
- 395 51. Gustafson EP. 1986 Carnivorous mammals of the late Eocene and early Oligocene of
 396 Trans-Pecos Texas. *Texas Meml. Museum Bull.* 33, 1–66.
- 397 52. Wesley GD, Flynn JJ. 2003 a Revision of Tapocyon (Carnivoramorpha), Including
- 398 Analysis of the First Cranial Specimens and Identification of a New Species. J. Paleontol.
- 399 **77**, 769–783. (doi:10.1666/0022-3360(2003)077<0769:arotci>2.0.co;2)

- Kirk EC, Williams BA. 2011 New adaption primate of Old World affinities from the
 Devil's Graveyard Formation of Texas. *J. Hum. Evol.* 61, 156–168.
 (doi:10.1016/j.jhevol.2011.02.014)
- 403 54. Speijer RP, Pälike H, Hollis CJ, Hooker JJ, Ogg JG. 2020 *The Paleogene Period*. 404 (doi:10.1016/b978-0-12-824360-2.00028-0)
- 405 55. Bai B, Wang YQ, Mao FY, Meng J. 2017 New Material of Eocene Helaletidae
 406 (Perissodactyla, Tapiroidea) from the Irdin Manha Formation of the Erlian Basin, Inner
 407 Mongolia, China and Comments on Related Localities of the Huheboerhe Area. Am.
 408 Museum Novit., 1–44. (doi:10.1206/3878.1)
- 409 56. Doria G, Royer DL, Wolfe AP, Fox A, Westgate JA, Beerling DJ. 2011 Declining
 410 atmospheric CO2 during the late Middle Eocene climate transition. *Am. J. Sci.* 311, 63–
 411 75. (doi:10.2475/01.2011.03)
- 412 57. Averianov A, Obraztsova E, Danilov I, Jin J. 2019 Anthracotheriid artiodactyl
 413 Anthracokeryx and an upper Eocene age for the Youganwo Formation of southern China.
 414 *Hist. Biol.* 31, 1115–1122. (doi:10.1080/08912963.2017.1421639)
- 415 58. Barrett PZ, Hopkins SSB, Price SA. 2021 How many sabertooths? Reevaluating the
 416 number of carnivoran sabretooth lineages with total-evidence Bayesian techniques and a
 417 novel origin of the Miocene Nimravidae. J. Vertebr. Paleontol. 41, 1–15.
 418 (doi:10.1080/02724634.2021.1923523)
- 419 59. Chow M. 1958 A record of the earliest sabre-tooth cats from the Eocene of Lushih,
 420 Honan. *Sci. Rec.* N.S. 2, 347–349.
- 42160.Ding S, Zhen J, Zhang Y, Tong Y. 1977 The age and characteristic of the Liuniu and the422Dongjun faunas, Bose Basin of Guangxi. Vertebr. Palasiat. XV, 35–45.
- 423 61. R Core Team. 2021 R: A language and environment for statistical computing.
- 424 62. Zack SP. 2019 The first North American propterodon (Hyaenodonta: Hyaenodontidae), a
 425 new species from the late Uintan of Utah. *PeerJ* 2019. (doi:10.7717/peerj.8136)
- 426 63. Zheng J, Tang Y, Zhai R, Ding S, Huang X. 1978 Early Tertiary strata of Lunan Basin,
 427 Yunnan. Prof. Pap. Stratigr. Paleontol. Beijing, 22–29.
- 428 429



