New Notharctine (Primates, Adapiformes) Skull From the Uintan (Middle Eocene) of San Diego County, California

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ABSTRACT  A new genus and species of notharctine primate, Hesperolemur actius, is described from Uintan (middle Eocene) aged rocks of San Diego County, California. Hesperolemur differs from all previously described adapiforms in having the anterior third of the ectotympanic anulus fused to the internal lateral wall of the auditory bulla. In this feature Hesperolemur superficially resembles extant cheirogaleids. Hesperolemur also differs from previously known adapiforms in lacking bony canals that transmit the internal carotid artery through the tympanic cavity. Hesperolemur, like the later occurring North American cercamoniine Mahgarita stevensi, appears to have lacked a stapedial artery. Evidence from newly discovered skulls of Notharctus and Smilodectes, along with Hesperolemur, Mahgarita, and Adapis, indicates that the tympanic arterial circulatory pattern of these adapiforms is characterized by stapedial arteries that are smaller than promontory arteries, a feature shared with extant tarsiers and anthropoids and one of the characteristics often used to support the existence of a haplorhine-strepsirhine dichotomy among extant primates. The existence of such a dichotomy among Eocene primates is not supported by any compelling evidence. Hesperolemur is the latest occurring notharctine primate known from North America and is the only notharctine represented among a relatively diverse primate fauna from southern California. The coastal lowlands of southern California presumably served as a refuge area for primates during the middle and later Eocene as climates deteriorated in the continental interior. Hesperolemur probably was an immigrant taxon that entered California from either the northern (Wyoming/Utah) or southern (New Mexico) western interior during the middle Eocene  © 1995 Wiley-Liss, Inc.
fieldwork, a primate skull was discovered from rocks of early Uintan age. The purpose of this paper is to describe the new primate, to compare it with other notharctid and adapid primates from North America and Europe, and to provide a summary of primate faunas from San Diego and Ventura counties.

Institutional abbreviations and designations used in the text are as follows: AMNH, American Museum of Natural History, New York, NY; BMNH, Natural History Museum, London, United Kingdom; CM, Carnegie Museum of Natural History, Pittsburgh, PA; Halle, Geiseltal Museum, Halle, Germany; Louis, Private collection of P. Louis, Cormicy, France; MNHN, Muséum National d'Histoire Naturelle, Paris, France; MPM, Milwaukee Public Museum, Milwaukee, WI; NHB, Basel, Naturhistorisches Museum, Basel, Switzerland; PLV, Laboratorium voor Actuopalaeontologie, Katholieke Universiteit, Louvain, Belgium; SDSNH, San Diego County Geologic Survey, San Diego, CA; TMM, Texas Memorial Museum, Austin, TX; UALP, University of Arizona, Laboratory of Paleontology, Tucson, AZ; UCMP, University of California, Museum of Paleontology, Berkeley, CA; UL, University of Lyon, Lyon, France; UM, University of Michigan, Museum of Paleontology, Ann Arbor, MI; USGS, United States Geological Survey, Denver, CO; USNM, United States National Museum, Washington, DC; YPM, Yale Peabody Museum, New Haven, CT.

MATERIALS AND METHODS

Measurements and nomenclature

All tooth measurements were taken with dial calipers under a binocular microscope and recorded to the nearest tenth millimeter. Length (L) measurements record maximum mesiodistal tooth dimensions, and width (W) measurements record maximum buccolingual breadth. Upper teeth are designated by tooth position with superscripts and lower teeth by subscripts with I = incisor, C = canine, P = premolar, and M = molar. All measurements of arterial canal widths were made using a binocular microscope with an optical micrometer at ×10. Arterial pathways referred to as canals are those surrounded by bone and manifest as a hollow tube, while those that are not enclosed in bone are referred to as grooves.

Comparative samples

Comparisons of the new California primate were made with a wide range of adapiforms, either with original specimens or high quality casts. Included in these comparisons were the following: European adapiforms: Adapis parisiensis (UM 63301, 63302, Cambridge M.538, Montauban-4); Agerinia roSELLI (Unnumbered holotype and Cecile 4241); Anchomomys gaillardi (UL L-46bis); Anchomomys stehlini (Basel En-1); Caenopithecus lemuroides (NHB Eh 597-728); Cantius eppsi (BMNH 13773, 15145, 15147, 29639); Cantius savagei (Louis Collection Mu 155-158, 160, MNHN Av 4846, 5907, 7702, Gr 98); Cercamonius brachyrhynchus (Basel Qv 619); Donrussellia gallica (MNHN Av many unnumbered teeth); Donrussellia lousi (MNHN Av 4731, 4845, 5664 and unnumbered teeth from Avenay and Grauves); Europolemur klatti (Halle—unnumbered dentary from Geiseltal and Halle 4238, 4292, 4304, 7325, 7396); Leptadapis magnus (Basel Qv 545, 920, MNHN Qv 10943); Periconodon (= Anchomomys?) pygmaeus (Basel En 367, Halle 7418, MNHN Bchs 494); Pronycticebus gaudryi (MNHN unnumbered holotype skull, Qv 11057); Protoadapis curvicuspides (AL 5182, 5719); Protoadapis filholi (PLV 35); Protoadapis russelii (MNHN Av 4644, 5759); Protoadapis sp. (MNHN Gr 150 and unnumbered teeth from Bouxwiller); North American adapiforms: Cantius abditus (many specimens in UM collections, AMNH 4734); Cantius angulatus (AMNH 55505, 55510, 55515); Cantius frugivorus (AMNH 16210, 55501, 86296, CM 37448); Cantius mckennai (many specimens in UM collections); Cantius ralstoni (many specimens in UM collections); Cantius torresi (many specimens in UM collections); Copelemur australotutus (USNM 22261, 411833); Copelemur praetutus (USNM 411-882, 411886, YPM 14698); Copelemur tutus (AMNH 16205, 55462, UALP 11377-8); Mahgarita stevensi (TMM 41578-9); Notharctus robinsoni (many specimens in UM collections); Notharctus tenebrosus (many specimens in UM collections); Pelycodus jarrovi (CM 37453, USGS 6549); Smilodectes graci-
**RESULTS**

Systematic paleontology

Order Primates Linnaeus, 1758  
Suborder Prosimii Illiger, 1811  
Infraorder Adapiformes Szalay and Delson, 1979  
Family Notharctidae Trouessart, 1879  
Subfamily Notharctinae Trouessart, 1879

**Hesperolemur, gen. nov.**

*Notharctus* (Lillegraven, 1980).  
**Type species.** *Hesperolemur actius*., sp. nov.

**Diagnosis.** *Hesperolemur* differs from all other known adapiforms in lacking canals enclosing the internal carotid arterial system within the tympanic cavity and in having the anterior third of the ectotympanic and the anterior crus fused to the internal surface of the lateral wall of the auditory bulla; it differs from adapids (*Adapis* and *Leptadapis*) in having a smaller sagittal crest, in having less massive zygomatics, in having a premolariform *P₄*, and in having upper molars with a protocone fold (= "pseudohyopocone") and a well-developed metaconule; it differs from cercamoniines (*Cercamonia* and *Northarctus*) in having upper molars with protocone folds, metaconules, and mesostyles on upper molars. The lone North American exception is *Mahgarita*, which is placed within cercamoniines based on the presence of a cingular ("true") hypocone and an absence of metaconules and mesostyles on upper molars.

While *Hesperolemur* is generically distinct from other North American notharctines, it clearly shares common ancestry with the taxa in this subfamily (*Cantius, Pelycodus, Copelemur, Notharctus*, and *Smilodectes*). Like these taxa, *Hesperolemur* has upper molars with protocone folds and strong metaconules, both characteristics that stand in contrast to cercamoniines and adapids. Cranially, *Hesperolemur* is similar to *Notharctus* in most features, differing substantially only in the disposition of the tympanic cavity.

**Hesperolemur actius, sp. nov.**

*Notharctus* sp. near *N. robustior* (Lillegraven, 1980).

**Holotype.** SDSNH 35233, compressed skull with left and right *P₄*-*M₃*, collected by R.A. Cerutti, December 7, 1987, from SDSNH locality 3413, Azuaga II, site 5.

**Referred specimens.** SDSNH 42415, right *M₃*, from SDSNH locality 3380, Carmel Mountain Ranch, Unit 16, site 3; UCMP 113256, broken left *M₂* from UCMP locality V-72157.

**Horizon.** The type and referred specimens were collected from early Uintan (middle Eocene) strata originally mapped as Mission Valley Formation by Kennedy and Moore (1975) and regarded as a new litho-
stratigraphic unit by Walsh (1991). These strata are now tentatively correlated by Walsh (personal communication) with the type Friars Formation.

**Diagnosis.** As for genus.

**Etymology.** Latin, *aktios*, coastal, referring to the coastal setting of the San Diego localities.

**Description.** SDSNH 35233 is a dorsoventrally compressed and somewhat plastically deformed skull of a relatively large notharctine primate (Fig. 1). The anterior portion of the skull is missing in front of P4. The skull now measures 70.2 mm in length and 48.2 mm in maximum width but obviously would have been both longer and wider had it not been crushed and broken. No sutures are evident anywhere on the skull because of crushing and step-fracturing of cranial elements.

**Lateral view (Fig. 1A).** Only a few features of *Hesperolemur* can be discerned in lateral view due to crushing and breakage. The maxillary, from the base of the orbit to the gingival margin, is relatively narrow dorsoventrally, more so than in other North American notharctines. It is much narrower than in *Adapis*. The infraorbital foramina are relatively larger than in *Notharctus*, *Smilodectes*, *Cantius* (UM 93938), and *Adapis* and are positioned above P4 as in those taxa. The root of the jugal originates dorsal to the posterior root of M1, making it more anteriorly placed than in *Notharctus*, *Smilodectes*, or *Adapis* but about as in *Cantius*.

The orbits are crushed, but, judging from relatively undistorted portions of the orbital margin, they would have been about the relative size of those in North American notharctines, relatively larger than those exhibited in *Adapis*. The lacrimal canal appears to have been positioned just inside the orbit as in *Notharctus*, *Smilodectes*, and *Adapis*, where it interrupts the continuity of the anterior orbital margin. This condition differs from *Cantius* (UM 93938 (Fig. 2), where the lacrimal canal is positioned anterior to the orbital margin such that the anterior orbital rim forms a continuous, unbroken border. The morphology of the orbital fossa in *Hesperolemur* is completely obscured by broken and overlapping bone fragments.

**Dorsal view (Fig. 1B).** The skull exhibits its relatively strong postorbital constriction dividing it into distinct splanchnocranium and neurocranium. The nasals are not preserved. The frontals are relatively broad as in *Smilodectes* (Gazin, 1958) and *Notharctus* (Gregory, 1920) but unlike *Adapis*, in which the frontals are constricted and concave.

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**Fig. 1.** Skull of *Hesperolemur actius* (SDSNH 35233, holotype). A: Lateral view. Small black triangle to left indicates position of infraorbital foramen, white arrowhead indicates position of confluence of frontal lines, and black arrow indicates position of parietosquamosal foramen. B: Dorsal view. Small black arrow to left indicates position of confluence of frontal lines; larger arrow to right indicates position of parietosquamosal foramen. C: Ventral view. Small black triangle indicates position of basisphenoid-basioccipital suture. Anterior to left in all views. Scale in 1 cm increments.
Fig. 2. Rostrum of Cantius abditus (UM 93938) in lateral view, anterior to left. Small black arrow above indicates position of the lacrimal canal; larger arrow below indicates position of infraorbital foramen. Scale in 1 cm increments.

(Stehlin, 1912; Gingerich, 1981a; Gingerich and Martin, 1981). There is no evidence to suggest that the frontals were inflated as in Smilodectes (Gazin, 1958), but instead they were probably flattened as in Notharctus. The frontals taper laterally to form relatively broad superior rami of the postorbital bars, very similar to the condition exhibited in Smilodectes. The frontal lines converge at the midline just posterior to the postorbital constriction much as in Adapis and unlike Smilodectes or Notharctus, where the frontal lines join farther posteriorly.

Hesperolemur has a well-developed sagittal crest extending from the confluence of the frontal lines posteriorly to the nuchal crest. The sagittal crest is less developed than is typical of Adapis (Gingerich, 1981a) but is better developed than it is in most specimens of Smilodectes (which almost never has a sagittal crest) or Notharctus (where a weak to moderate sagittal crest often is present). The braincase is pear-shaped, being relatively broad and rounded posteriorly, tapering anteriorly to the postorbital constriction. The anterior biparietal width (20.9 mm) is comparable to that of Smilodectes gracilis (21.0 mm) but much broader than it is in Adapis parisiensis (11.3 mm). The braincase is not inflated anteriorly as it is in Smilodectes but tapers more gradually to the postorbital constriction.

The anterior, posterior, and ventral extent of the parietals is obscured by crushing. Hesperolemur appears to have distinct parietosquamosal foramina as in Smilodectes, Notharctus, and Adapis, but they are positioned more anteriorly than in those taxa. If interpreted correctly, these foramina are located more dorsally than in Smilodectes or Notharctus, which suggests that the dorsolateral portion of the squamosal was relatively more extensive than in either of these taxa, more like the condition exhibited in Adapis. The root of the zygomatic arch in Hesperolemur is relatively broad anteroposteriorly as in Notharctus and Smilodectes. Although broken, it does not appear to have extended posteriorly to form deep temporal gutters as in Adapis (Stehlin, 1912).

Ventral view (Fig. 1C). The skull of Hesperolemur preserves the left and right maxillary tooth rows (P4 to M3), small portions of the basisphenoid, crushed and plastically deformed petrosals, and a somewhat distorted basioccipital. The anterior portions of the basisphenoid are missing. Posteriorly, the basisphenoid contacts the basioccipital at about the anteriormost point of the auditory bullae as in Notharctus and Smilodectes, relatively more posteriorly than in Adapis. The basisphenoid contacts the anteromedial portion of the bullae anterior to its contact with the basioccipital as in Notharctus. The basisphenoid does not appear to overlap the bullae as in Smilodectes. The central ridge of the basisphenoid is broad and relatively flat as in Notharctus, not narrower and more elevated as in Adapis or laterally compressed and sharply defined as in Smilodectes. The mesopterygoid fossa appears to have been only moderately excavated laterally as in Adapis and Notharctus, not deep pocketed as in Smilodectes (Gazin, 1958).

The basioccipital of Hesperolemur is narrow anteriorly and widens posteriorly, forming a relatively broad surface approaching the condition in Notharctus but not as broad as in Adapis and not as laterally impinged upon by the bullae as in Smilodectes. The
basioccipital does not overlap the medial wall of the auditory bullae as it does in Notharctus and especially Smilodectes but does contact it more broadly than in Adapis. The basioccipital has a relatively weak median ridge extending from its anterior contact with the basisphenoid to the margin of the foramen magnum much as in Notharctus but unlike Adapis, where this ridge appears to be absent, or Smilodectes, where a median ridge is well developed.

The auditory bullae are relatively large, continuous with the petrosal, and rounded anteromedially. These taper slightly posterolaterally but do not form an ossified external auditory meatus. The anteromedial extent of the petrosal reaches beyond the contact of the basisphenoid and basioccipital as in Notharctus, but this is not as anteriorly extended as in Adapis. Anteriorly, the bullae overlap the posterior margins of the basisphenoids, unlike Smilodectes, where the basisphenoids overlap the bullae.

**Tympanic cavity.** Morphological features of the tympanic cavity can be discerned from both the left and right side of the skull of Hesperolemur. The right tympanic cavity has been crushed anteriorly, tilting all of the preserved features anterodorsally to posterovertrally (Fig. 3A). The left tympanic cavity has been crushed dorsoventrally and anteriorly, pushing most of the middle ear dorsally (Figs. 3B, 4). Even with such distortion, many of the relevant features of middle ear morphology are preserved on one side or the other.

The tympanic cavity of Hesperolemur is separable into four distinct regions. There is a relatively large anteromedial cavity (Simons and Rasmussen, 1989; Rasmussen, 1990) roofed by the epitympanic wing of the petrosal. This cavity is divided into anterior and posterior areas by a medial secondary septum (MacPhee, 1981). A relatively large anterolateral cavity is present, roofed by the "tegmen tympani." Posterolaterally, the tympanic cavity is mediodorsally limited by the facial canal and laterally by a deep epitympanic recess. The postero medial portion of the tympanic cavity is occupied by the promontorium and associated structures.

Although crushed and broken, many of the features of the tympanic roof of Hesperolemur can be discerned by examining both left and right ear regions (Fig. 3C). The promontorium (best seen in the right ear region [Fig. 3A]) is a rounded eminence. At the dorsal base of the posterior portion of the promontorium is the cochlear fenestra, somewhat hidden by a posterior extension of the promontorium. Just lateral to the cochlear fenestra is a small fossa that may represent the point of origin for the stapedius muscle. The vestibular fenestra (best seen in the left ear region [Figs. 3B, 4]) is located along the lateral margin of the promontorium. The epitympanic recess is lateral to the vestibular fenestra. The tensor tympani fossa is located just anterior to the epitympanic recess.

Relatively large stylomastoid foramina are preserved on both left and right bullae located posterolaterally. It is possible to follow the course of the facial canal through the stylomastoid foramen and into the tympanic cavity. However, the course of the facial canal cannot be traced farther due to breakage. It presumably crossed the tympanic cavity at the medial margin of the epitympanic recess as in other notharcines, but this is impossible to determine with certainty.

The circulatory pattern of the internal carotid arterial system in Hesperolemur appears quite different from that documented for all previously known notharcines (Gregory, 1920; Szalay, 1975; Szalay and Delson, 1979). Like other notharcines, the internal carotid artery presumably entered the bulla posteriorly, ventral and medial to the stylomastoid foramen (neither ear region preserves the posterior carotid foramen). Unlike other known adapiforms (and other Eocene primates), there is no evidence of any bony canals enclosing arteries within the tympanic cavity (Figs. 3A,B, 4). It is difficult to determine the exact path of the internal carotid artery until it reaches the promontorium. There is a large, deep groove (not a canal) along the lateral aspect of the promontorium that represents the promontorial pathway of the promontory artery, but no tube or canal is present that is continuous with this groove either anteriorly or posteriorly. Some specimens of Smilodectes and Notharctus have canals that are open across the lateral part of the promontorium (manifest as a groove), but well-developed, closed
Fig. 3. Stereophotographs of tympanic regions of *Hesperolemur actius* (SDSNH 35233, holotype). A: Ventral view of right tympanic cavity with anterior to top and lateral to left showing promontorium (p), stylomastoid foramen (white arrow), groove across lateral surface of promontorium for promontory artery (white arrowhead), and medial secondary septum (black arrowhead). B: Ventral view of left tympanic cavity with anterior to top and lateral to right showing stylomastoid foramen (white arrowhead at bottom right), groove across lateral surface of the promontorium for the promontory artery (white arrowhead to left), anterior portion of the eustympanic anulus (white arrow at top), and the position of the auditory meatus (large white arrow to right). C: Photograph of entire basicranial region of *H. actius*, anterior to top. Scale = 1 cm.
canals are present throughout the rest of the internal carotid circulatory pathway through the tympanic cavity. It is possible that these canals are simply missing in *Hesperolemur* because of breakage, but the areas where such canals would have connected are smooth and show no signs of any bony portions having been broken off. In addition, there were no broken bone fragments within the tympanic cavities that could be
attributed to these canals, which seems unlikely given that other delicate structures (ear ossicles, portions of the ectotympanic anulus) were found within the tympanic cavities.

*Hesperolemur* also differs from known notharctids in apparently lacking a canal or groove for the stapedial artery. It is difficult to be certain of this because of the condition of the ear regions, but there is no evidence apparent for the existence of a stapedial artery (Figs. 3A,B, 4). The lack of a stapedial artery in *Hesperolemur* seems incongruous with previous notions of the disposition of arterial circulation through the middle ear in “extinct strepsirhines” (Szalay, 1975; Szalay and Delson, 1979). It has been suggested that the strepsirhine condition is one in which the stapedial artery is larger than the promontory artery, while the opposite holds in haplorhines (Gregory, 1920; Szalay, 1975; Szalay and Delson, 1979; but also see MacPhee and Cartmill, 1986; Ross, 1994). There has been some evidence to suggest that the supposed strepsirhine condition is not representative of notharctines (Gingerich, 1973; MacPhee and Cartmill, 1986), *Adapis* (Gingerich and Martin, 1981), or *Mahgarita* (Rasmussen, 1990), but this evidence has been disputed (Wilson and Szalay, 1976; Szalay and Delson, 1979).

Gingerich (1973) provided measurements of diameters of bony canals in one specimen of *Notharctus* (YPM 11466) that indicated that the promontory artery was, in fact, larger than the stapedial artery (see Table 1). Gingerich and Martin (1981) provided measurements of the Cambridge skull of *Adapis* where the same condition holds. MacPhee and Cartmill (1986) note that two skulls of *Smilodectes* (MPM 2612 and 5409) both have promontory canals much (their italics) larger than stapedial canals. Fieldwork in the Bridger Formation of southwestern Wyoming has produced six additional notharctine skulls (UM numbers 98828 (Notharctus tenebrosus), 100000, 100044, 100521, 101212 (Smilodectes gracilis), and 100603 (Smilodectes mcgrewi) (Figs. 5, 6) that provide additional support for the morphology cited by Gingerich (1973) and MacPhee and Cartmill (1986). In all cases, by either external or internal diameters, the promontory canal is larger than the stapedial canal in these notharctine specimens. While this does not necessarily prove that the arteries in these canals reflected the same relationship, it does call into question the usefulness of such a character for recognition of a haplorhine-strepsirhine dichotomy among middle Eocene primates. If relative size of promontory and stapedial arteries is a valid character differentiating strepsirhines from haplorhines (this is in dispute, see Beard and MacPhee, 1994), then adapiiforms are haplorhines, a conclusion few proponents of Strepsirhini-Haplorhini would support (although see Cartmill and Kay, 1978). Other lines of evidence do not support the existence of a strepsirhine-haplorhine dichotomy among Eocene primates. The reality is that such a dichotomy simply is not usefully applied to the early primate radiation. In any event, the relatively large size of the promontory arterial groove and the apparent lack of a stapedial artery in *Hesperolemur* is not out

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* From Gingerich (1973).
* Measured at bifurcation of internal carotid artery.
of step with findings based on other known adapiforms. Broken pieces of the left ectotympanic anulus, incus, and malleus and a nearly complete right incus were found within the tympanic cavities of *Hesperolemur*. The ectotympanic fragment consists of approximately the anterior third of the anulus (Figs. 3B, 4). This ectotympanic portion appears to have been solidly fused to the internal bullar wall just medial to the postglenoid foramen along the anterodorsal margin of the auditory meatus. It is attached to the bullar wall by a solid, bony anular bridge (MacPhee and Cartmill, 1986; MacPhee, 1987). There is a small suture or crack running along the lateral margin of the ectotympanic suggesting the presence of a recessus dehiscence (MacPhee, 1987) and indicating that the anular bridge is petrosal in origin. A grooved surface that served as the attachment area for the tympanic membrane is formed by a
Unlike *Smilodectes* (MacPhee and Cart-
Fig. 7. Stereophotographs of right ear region of *Smilodectes gracilis* showing disposition of ectotympanic. A: UM 100521 in lateral view with anterior to right and dorsal to top. Small white arrowheads indicate posterior crus (left) and broken “free” segment (right) of ectotympanic; white arrow at lower left indicates position of the internal carotid foramen. B: UM 100521 in ventrolateral view with anterior to top and lateral to left. Small white arrowhead indicates posterior crus of ectotympanic fused to anular bridge, larger white arrowhead indicates broken “free” segment of ectotympanic, and black arrow indicates anterior crus located along dorsal surface of the auditory meatus. C: UM 101212 in ventral view with anterior to top and lateral to left. Black arrow indicates position of internal carotid foramen; white arrowhead indicates broken “free” portion of ectotympanic lying anteromedial to promontorium. Scale = 1 cm.

disposition of the ectotympanic in *Smilodectes gracilis*. The posterior crus of *Smilodectes* is connected to the lateral bullar wall via an anular bridge. After the ectotympanic passes the level of the internal carotid canal, it is no longer attached to the bulla, exhibiting the “free” ring condition of extant *Lemur* (anular bridge incomplete sensu MacPhee and Cartmill, 1986). The ectotympanic anulus may contact a small portion
of the lateral bullar wall along with its anterior curvature as in *Lemur* but is not fused to the lateral wall anteriorly as in *Hesperolemur*. The anterior crus of *Smilodectes* rests in the auditory meatal fossa (the crus is clearly visible in UM 100521 [Fig. 7B]). The anterior crus of the ectotympanic of *Hesperolemur* does not appear to lie in a meatal fossa but is instead attached to the internal surface of the lateral bullar wall at the auditory meatus similar to extant *Otolemur*.

The presence or absence of a continuous bony connection between the ectotympanic and the lateral bullar wall (a complete anular bridge [see MacPhee and Cartmill, 1986; MacPhee, 1987; Beard and MacPhee, 1994]) cannot be determined. If *Hesperolemur* was like other known notharctids it would have lacked a complete anular bridge (MacPhee and Cartmill, 1986; MacPhee, 1987; Franzen, 1987). However, the fact that the anterior third of the ectotympanic is solidly attached to the lateral bullar wall via an anular bridge suggests that the entire ectotympanic may have been similarly connected to the bulla. There is a small piece of bullar wall from the ventrolateral portion of the right bulla of *Hesperolemur* that has a smooth, shallow groove running anteroposteriorly across its dorsal surface. This may represent a band or groove for the ectotympanic and if so suggests that the ectotympanic was in close proximity to the dorsal bullar surface, but there is no evidence to indicate that the ectotympanic was directly connected to the bulla in this area. If *Hesperolemur* had an ectotympanic anulus attached to the dorsal surface of the bulla throughout its course by a complete or nearly complete anular bridge, it would have resembled the condition exhibited by some extant cheirogaleids (Cartmill, 1975; MacPhee and Cartmill, 1986).

The malleus fragment of *Hesperolemur* is represented only by the incudal articular surface and provides little information about the complete disposition of this ossicle. Compared to *Smilodectes gracilis* (UM 101212), the malleus of *H. actius* is somewhat smaller, and the incudal articular surface is flatter and less saddle-shaped.

The right incus of *Hesperolemur* is nearly complete, missing only the long crus, while the left incus is missing both the long and short crura. Comparisons with a left incus of *Smilodectes gracilis* (UM 100000) indicate that *Hesperolemur* and *Smilodectes* shared most features of incudal morphology (Fig. 8A, B). The body of the incus in both taxa has distinct medial and lateral mallear facets that are oriented at nearly 90 degrees to one another. The medial mallear facet is more concave than the lateral facet, especially in *Smilodectes*, although the lateral facet is somewhat saddle-shaped in both taxa. The short crus is relatively shorter and less robust in *Hesperolemur*, and, although the long crus is missing, judging by the angle of the broken surface, it must have been more inferiorly oriented than in *Smilodectes*, resulting in the short and long crura being relatively farther apart in *Hesperolemur*.

Other characteristics of the basicranium are also evident on the skull of *Hesperolemur*. The mastoid region, while not relatively large, did enclose at least two substantial air cells. The mastoids are not as inflated as in *Adapis* but approach the condition exhibited in *Smilodectes* and *Notharctus*. The glabel fossa is relatively wider mediolaterally than in *Smilodectes* but is approached by *Adapis* in this characteristic. The postglenoid process is robust and angled slightly posteriorly as in *Smilodectes*. It may have contacted the bulla, unlike *Smilodectes*, but does not appear to have been fused to the bulla anterior to the external auditory meatus as in *Adapis*. There is a distinct postglenoid foramen located on the posteromedial aspect of the postglenoid process abutting against the anterior portion of the auditory meatus.

**Dentition.** The skull of *Hesperolemur* preserves P4–M3 on both sides (Figs. 1C, 9B). P4 is a mesiodistally compressed tooth with a well-developed protocone that is positioned near the mesiolingual base of the paracone. The paracone is nearly twice the height of the protocone. The paracone is centrally placed along the buccal margin producing a preparacrista and postparacrista of nearly equal length. There is no metacone. P4 has a small parastyle and no metastyle. There is a weak cingulum that encircles the tooth.

The molars are quite worn across the lingual half of their occlusal surfaces. M2 is
Fig. 8. Line drawings of (A) right incus of *Hesperolemur actius* (SDSNH 35233, holotype) and (B) left incus of *Smilodectes gracilis* (UM 100000). In A, lateral view is bottom left, medial view is bottom right, and anterior view is top. In B, medial view is bottom left, lateral view is bottom right, and anterior view is top. Small arrows point dorsally in each view. Scale = 2 mm.

Fig. 9. Stereophotographs of left upper dentitions of (A) *Notharctus tenebrosus* (UM 32335, P\textsuperscript{3}–M\textsuperscript{3}) and (B) *Hesperolemur actius* (SDSNH 35233, holotype, P\textsuperscript{3}–M\textsuperscript{3}) in occlusal view. Scale = 2 mm.

the largest molar, with M\textsuperscript{1} being noticeably smaller than M\textsuperscript{2} and M\textsuperscript{3} slightly smaller than M\textsuperscript{1}. It is difficult to determine the position and relationships of upper molar protocones and hypocones with precision, but extensive comparisons with other adapiforms indicate that *Hesperolemur* almost certainly had a protocone fold and lacked a cingulum hypocone. M\textsuperscript{1–2} have well-developed paracones and metacones that are separated by a rather wide, shallow valley formed by gently sloping postparacristae and premetacristae. M\textsuperscript{2} has a very weak, low metacone. Relatively robust metaconules are present on M\textsuperscript{1–2}, while M\textsuperscript{3} has a relatively smaller metaconule. The presence and disposition of molar paraconules cannot be determined due to obscuring wear. M\textsuperscript{1} has a weakly formed mesostyle that is somewhat better developed on M\textsuperscript{2} and absent on M\textsuperscript{3}. Relatively heavy
mesial and distal cingula are present on all three molars as well as relatively weaker buccal and lingual cingula. Stylar shelves are essentially absent. M3 is longest buccally and tapers somewhat lingually but retains a generally square outline. M2 is wider and more rectangular in outline. M1 is generally squared but the mesiobuccal corner of the tooth is slightly distended. and the hypocone lobe is reduced compared to the other molars.

Comparing the upper dentition with other North American notharctines indicates that Hesperolemur is generally similar to most taxa but differs in important ways from each (Figs. 9, 10). Hesperolemur differs from Notharctus in having P4 with a more mesially and buccally placed protocone and a centrally placed paracone. Upper molars of Hesperolemur differ from those of Notharctus in having less well-developed mesostyles with little or no stylar shelf development and in lacking distinct hypocone lobes that are separated from the protocone by a distinct lingual groove (especially true of more derived Notharctus species). Hesperolemur differs from Smilodectes in having weaker mesostyles, in having very shallow trigon basins, and in having relatively larger hypocone lobes.

Hesperolemur differs from derived Cantius species in much the same manner as it does from Notharctus, but more primitive species of Cantius have weaker mesostyles and more mesial P4 protocones like those of Hesperolemur. Hesperolemur also has a more mesiodistally compressed P4 than most species of Cantius. Hesperolemur differs from Copelemur (C. tutus only) in lacking a well-developed mesostylar cusp (Hesperolemur has a crest-like mesostyle instead of an isolated cuspule), in lacking a lingually expanded protocone lobe, in having a weaker stylar shelf, in having a shallower trigon basin, and in having more bulbous and robust metaconules and stronger upper molar cingula. Hesperolemur differs from Pelycodus jarrovi in having upper molar mesostyles and in having the M3 paracone less buccally distended.

SDSNH 42415 is a right M3 referred to Hesperolemur actius (Fig. 10F). The protoconid and metaconid are widely spaced, with the metaconid taller, more basally inflated, and much more massive. The protoconid and metaconid are connected buccally by a well-developed crest. There is no paraconid, but a short, arcuate paracristid connects the mesial flanks of the protoconid and metaconid forming the mesial wall of a distinct trigonid fovea. There is a distinct, robust hypoconid that is connected to the postprotocristid by a sigmoid cristid obliqua. A small ectocingulid is present beneath the hypoflexid. A small entoconid and an elongate, faintly bilobate hypoconulid are present.

The M3 of Hesperolemur differs from that of Notharctus and Cantius (Fig. 10D,E) in having a smaller trigonid fovea with a shorter, more robust paracristid and no paraconid, in having a very massive metaconid and a more inflated protoconid, in having a weaker buccal cingulid, in lacking multiple entoconid cuspules, and in lacking the distinctive lingual extension of the postprotocristid at its juncture with the cristid obliqua. Hesperolemur differs from Copelemur in lacking a distinct paraconid (C. tutus has a paraconid: C. praetutus does not), in lacking an entoconid notch, in lacking the lingually positioned entoconid typical of Copelemur, and in having more massive and inflated protoconid and metaconid. C. praetutus shares a sigmoidal cristid obliqua-postprotocristid flexure with Hesperolemur, while C. tutus has this complex less well developed.

The M3 of Hesperolemur differs from that of Pelycodus jarrovi in lacking a mesiodistally compressed trigonid, in lacking a paraconid, in having a sigmoid cristid obliqua-postprotocristid flexure, in having a massive metaconid, and in having a better developed trigonid fovea. Hesperolemur differs from Smilodectes (Fig. 10C) in having the trigonid of M3 closed lingually by the paracristid, in having a much more massive metaconid and an inflated protoconid, in having a sigmoidal cristid obliqua-postprotocristid flexure, in having a shallower talonid basin, and in having a relatively smaller entoconid.

UCMP 113256, a broken left M2, was described and figured by Lillegraven (1980) as Notharctus sp. near N. robustior. Although the talonid is damaged, the trigonid is nearly complete. The trigonid consists of a rela-
tively robust protoconid widely separated from a massive, basally inflated metaconid and a short, relatively straight paracristid that connects the mesial flanks of the protoconid and metaconid. There is no paraconid developed. All of these features are shared with SDSNH 42415, and therefore this specimen is referred to *H. actius*.

Measurements of the teeth of *Hesperolemur actius* are presented in Table 2.

**RELATIONSHIPS OF HESPEROLEMUR**

*Hesperolemur* is the latest surviving notharctine primate known. The only later adapiform known from North America is the cercamoline notharctid *Mahgarita stevensi* from the Skyline Channels, Devil’s Graveyard Formation, Duchesnean Land-Mammal Age (late Eocene) of southwestern Texas (Wilson and Szalay, 1976; Rasmussen, 1990). Comparison of *Hesperolemur* and *Mahgarita* does not appear to support a close relationship. *Mahgarita* and *Hesperolemur* do share an apparent lack of a stapedial artery along with a large promontory artery and possibly the presence of a “fused” ectotympanic anulus (a complete anular bridge [see Rasmussen, 1990]). These character states are shared in common with primitive anthropoids such as *Aegyptopithecus* (Rasmussen, 1990; Simons and Rasmussen, 1989; but also see Ross, 1994). However, as pointed out above, relatively small stapedial arteries also appear to be the rule in most known adapiforms, so this character state could be viewed as primitive (symplesiomorphic) for the infraorder. The apparent presence of an anteriorly fused ectotympanic and the lack of a stapedial artery may both represent synapomorphies for *Mahgarita* and *Hesperolemur*, but the evidence is such that definitive interpretation remains difficult.

In nearly all other dental and cranial features, *Hesperolemur* and *Mahgarita* appear quite different (Wilson and Szalay, 1976; Rasmussen, 1990). *Mahgarita*, like all other known Eocene primates, had a bony tube that carried the promontory artery through the tympanic cavity. *Mahgarita* has true (circular) hypocones on upper molars in contrast with all other North American notharctids and lacks upper molar mesostyles (unlike *Hesperolemur*). *Mahgarita* also differs from *Hesperolemur* in having the following: more cuspat e teeth that lack robust, bulbous cusps; *P* to a centrally placed protocone and more steeply sloping pre- and postpara-crista; lower molars with metaconid positioned posterior to the protoconid and lacking basal inflation; lower molar trigonids that are open lingually with a short, sloping paracristid and a very shallow, small trigonid fovea; and lower molars with a straight cristid obliqua and a well-developed entoconid positioned posterior to the hypoconid.

*Hesperolemur* shares some character states with adapids. Like *Adapis parisiensis*

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**TABLE 2. Dental measurements Hesperolemur actius compared with selected North American notharctines and the cercamoline Mahgarita stevensi**

<table>
<thead>
<tr>
<th>Genus/species</th>
<th>P* L x W</th>
<th>M1 L x W</th>
<th>M2 L x W</th>
<th>M3 L x W</th>
<th>M4 L x W</th>
<th>M5 L x W</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Hesperolemur actius</em></td>
<td>4.4 x 6.0 (1)</td>
<td>5.2 x 6.2 (1)</td>
<td>5.3 x 7.2 (1)</td>
<td>5.0 x 6.2 (1)</td>
<td>6.9 x 4.5 (1)</td>
<td></td>
</tr>
<tr>
<td><em>Notharctus tenebrosus</em></td>
<td>4.8 x 6.7 (12)</td>
<td>6.4 x 7.8 (16)</td>
<td>6.4 x 8.2 (14)</td>
<td>5.2 x 6.9 (10)</td>
<td>8.5 x 4.4 (11)</td>
<td></td>
</tr>
<tr>
<td><em>Notharctus robinsoni</em></td>
<td>4.5 x 6.6 (3)</td>
<td>6.0 x 7.6 (4)</td>
<td>6.1 x 8.0 (3)</td>
<td>—</td>
<td>6.8 x 4.2 (10)</td>
<td></td>
</tr>
<tr>
<td><em>Smilodectes gracilis</em></td>
<td>3.4 x 4.2 (5)</td>
<td>4.4 x 5.5 (7)</td>
<td>4.6 x 6.0 (9)</td>
<td>4.0 x 5.3 (10)</td>
<td>5.7 x 3.5 (10)</td>
<td></td>
</tr>
<tr>
<td><em>Smilodectes megrewi</em></td>
<td>—</td>
<td>4.4 x 5.6 (2)</td>
<td>4.8 x 6.1 (3)</td>
<td>4.2 x 5.4 (2)</td>
<td>5.1 x 3.6 (2)</td>
<td></td>
</tr>
<tr>
<td><em>Cantius abditus</em></td>
<td>3.9 x 5.3 (5)</td>
<td>4.9 x 6.4 (3)</td>
<td>5.1 x 7.6 (4)</td>
<td>4.2 x 5.9 (3)</td>
<td>6.6 x 4.3 (7)</td>
<td></td>
</tr>
<tr>
<td><em>Copelemur tutus</em></td>
<td>—</td>
<td>—</td>
<td>5.6 x 8.1 (1)</td>
<td>—</td>
<td>—</td>
<td>5.3 x 3.1 (1)</td>
</tr>
<tr>
<td><em>Copelemur praetutus</em></td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><em>Pelycodus jarrosoi</em></td>
<td>—</td>
<td>5.5 x 6.5 (4)</td>
<td>5.4 x 7.2 (2)</td>
<td>4.8 x 6.1 (3)</td>
<td>6.6 x 4.4 (2)</td>
<td></td>
</tr>
<tr>
<td><em>Mahgarita stevensi</em></td>
<td>2.9 x 3.8 (1)</td>
<td>4.6 x 4.5 (1)</td>
<td>4.0 x 4.7 (2)</td>
<td>3.2 x 4.4 (1)</td>
<td>4.2 x 2.5 (1)</td>
<td></td>
</tr>
</tbody>
</table>

* The number in parentheses = N; measurement mean is given for samples larger than 1.

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and *Leptadapis magnus*, the skull of *H. ac-tius* has frontal lines that converge anteriorly and are confluent with a well-developed sagittal crest. *Hesperolemur* also shares a relatively large dorsolateral expansion of the squamosal with *Adapis*. *Hesperolemur* differs dentally from *Adapis* and *Leptadapis* in having upper molars with postprotocingula and well-developed metaconules, a premo-
lariform P4, closed lower molar trigonids, and basally inflated protoconids and meta-
conids that are opposite one another, not off-
set with the metaconid posterior to the pro-
toconid.

*Hesperolemur* is also similar to European cercamoniines and European *Cantius eppsi* and *Cantius savagei* in some ways. Like European *Cantius* species, *Hesperolemur* has upper molars with well-developed metacoon-
ules and lower third molars with closed tri-
gonids, small entoconids, flexed cristid obli-
que, and relatively inflated trigonid cusps. As in all cercamoniines except *Donrussellia*, *Hesperolemur* has lower molars that lack a paraconid, but it differs from most cercamoniines (except *Periconodon* and *Agerinia*) by having lower molar trigonids closed lingually instead of being broadly open as in the former. *Caenopithecus lemuroides* shares upper molar mesostyles and lower molar hypoconulids with *Hesperolemur* but differs in most other dental details. Like *Pro-
toadapis*, *Hesperolemur* has relatively bul-
bous cusps, but *Protoadapis* lacks the basal in-
fation of lower molar cusps exhibited by *Hesperolemur*. *Hesperolemur* differs from all cercamoniines in having a protocone fold and a well-developed metaconule on upper molars.

Among North American notharctines, *Hesperolemur* seems to be most similar to *Pelycodus*, *Notharctus*, and *Cantius*. Like *Notharctus*, *H. actius* has relatively broad frontals and shares similar basisphenoid and basioccipital morphologies. Dentally, *H. actius* shares a flexed cristid obliqua on M3 with *Notharctus*, *Cantius*, *Pelycodus*, and some species of *Copelemur*. *Hesperolemur* shares relatively heavy upper molar cingula with *Pelycodus* as well as relatively robust and basally inflated lower molar protoconids and metaconids and a small, single-cusped entoconid on M3. *Hesperolemur* shares in-
flated lower molar protoconids and metacon-
ids and relatively weak upper molar meso-
styles with some derived species of *Cantius* (from both North America and Europe). *Hesperolemur* seems more distantly related to *Smilodectes* and *Copelemur* among North American notharctines.

An analysis of the relationships among the seven North American adapiforms was car-
rried out using the branch-and-bound option of PAUP (Phylogenetic Analysis Using Par-
simony, version 3.1.1 [Swofford, 1993]). This analysis is based on 21 unordered cranial and dental characters (Tables 3, 4) using the primitive European cercamoniine *Donrus-
sellia* as the outgroup to root trees. No a pri-
ori or a posteriori weighting of characters was attempted. Three most parsimonious trees were derived from the character ma-
trix, each consisting of 29 steps with a consis-
tency index of 0.724 and a retention index of 0.600. Figure 11 presents one of the three hypothesized branching sequences for North American notharctines and also represents the 50% Majority Rule Consensus Tree. This cladogram differs from the strict consensus tree by resolving a polytomy consisting of all North American notharctines except *Can-
tius*. In all trees *Mahgarita* and *Adapis* form a clade that is the sister group of North American notharctines, and *Cantius* is the sister.taxon to all other North American no-
tharctines. *Smilodectes* and *Copelemur* are more closely related to each other than to any other notharctine in all trees. In two of the three trees, *Hesperolemur* is the sister taxon of *Pelycodus*, while in the third *Hesperolemur* is the sister to a clade con-
sisting of *Notharctus*, *Smilodectes*, and *Co-
pelemur*.

**SOUTHERN CALIFORNIAN PRIMATE FAUNAS**

Middle to late Eocene primates are well rep-
resented in southern California from both San Diego and Ventura counties (Stock, 1933a, 1934a; Gazin, 1958; Szalay, 1976; Lil-
legraven, 1980; Kelly, 1990; Mason, 1990; Honey, 1990; Gunnell, 1995). Two separate regions in San Diego County have middle to late Eocene rocks that have produced pri-
mate specimens. In the northwestern part
of San Diego County the Lower and Upper units of the Santiago Formation contain five local faunas ranging in age from early Uintan to early Duchesnean (Walsh, 1991). In the Poway/Greater San Diego area a series of at least five rock units spans the early to late Uintan (Kennedy and Moore, 1971; Peterson and Kennedy, 1974; Kennedy and Peterson, 1975; Walsh, 1991, personal communication). In Ventura County, Kelly (1990) and Kelly et al. (1991) have recently revised the biostratigraphy of the Eocene portion of the Sespe Formation. These authors recognize five local faunas spanning the late Uintan and Duchesnean, each of which contains primate taxa.

Figure 12 summarizes the distribution of primate taxa in San Diego and Ventura counties. By far the greatest number of primate taxa known from southern California come from the Poway fauna of early Uintan age in San Diego County. At least nine different primates are found in these rocks. All but two taxa are referable to omomyid primates, and most are represented by genera also known from earlier (Bridgerian) rocks in the interior of western North America (the exceptions being Stockia powayensis, whose ancestry can be linked to taxa in Wyoming and Utah [Honey, 1990; Gunnell, 1995], and Ourayia, a taxon also known from the Uintan of the western interior).

One adapiform primate was described from the early Uintan Poway faunas by Lillegraven (1980). It is represented by a single maxilla (UCMP 113210) containing M1-2 and
Fig. 11. Cladogram depicting one possible branching sequence for North American notharctine primates using the primitive cercamomine Donrussellia as outgroup. This also represents the 50% Majority Rule Consensus Tree. The analysis is based on 21 unordered cranial and dental characters (see Table 3) and represents one of three most parsimonious trees obtained by PAUP 3.1.1 using the branch-and-bound option on the resulting data matrix (see Table 4). Tree length is 29 steps; consistency index = 0.724; retention index = 0.600.

a broken M3. Lillegraven (1980) compared this specimen carefully with Hemiacodon and “Pelycodus” (now Cantius) and concluded that UCMP 113210 showed greater overall similarity to “Pelycodus” (= Cantius). Upon further consideration, I believe a third possibility exists. Comparisons of stereophotographs of UCMP 113210 with other omomyid taxa leads to the conclusion that this specimen represents a species of Macrotarsius, close to the recently described M. roederi (Kelly, 1990) from the Sespe Formation.

UCMP 113210 differs from Macrotarsius most notably in lacking well-developed mesostyles on upper molars (Robinson, 1968; Kristhalka, 1978). In other features noted by Lillegraven (1980)—the broadly rounded anterior and posterior buccal corners, the lower, more rounded cusps and crests, the cingular development of the hypocone, and the presence of small pericones on M1–2—UCMP 113210 is very similar to Macrotarsius. In addition, both UCMP 113210 and Macrotarsius share a broad, well-developed stylar shelf.

The absence of molar mesostyles does not rule out inclusion of UCMP 113210 in Macrotarsius. M. roederi from the Brea Canyon Local Fauna, while only known from a single lower jaw, has molar morphology suggesting that upper molars may have lacked or had only small mesostyles. During mastication the postcristid and hypoconulid regions of the lower molars shear along surfaces formed by the premetacrista and mesostyle of the upper molars. In Macrotarsius siegerti (CM 15056, 18646), where a well-developed mesostyle exists, the postcristid and hypoconulid of the lower molars are well developed, closing off the posterior end of the talonid (Kristhalka, 1978). Kelly (1990) describes M. roederi lower molars as lacking hypoconulids (except on M3) and instead as having a notch in the postcristid where the hypoconulid would normally be found. This suggests that M. roederi upper molars may not have had mesostyles. If interpreted correctly, this suggests that UCMP 113210 may well represent a specimen of M. roederi or a closely related taxon. In any event, UCMP 113210 is better interpreted as an omomyid than an adapiform, leaving Hesperolemur as the sole adapiform primate known from southern California.

BIOGEOGRAPHY AND PALEOCLIMATES

Lillegraven (1979) has hypothesized that mammalian migration in the middle Eocene between southern California and the western interior was accomplished via a lowland route, the Sevier orogenic belt, that extended from southern California through northern Arizona and southern Nevada into Utah and Wyoming during the middle Eocene. Early Uintan mammalian faunas are similar in taxonomic makeup from both the western
interior and southern California. By the later Uintan, notable faunal differences occur between these areas, suggesting the closure of the Sevier orogenic belt leading to the development of more endemic faunas in southern California and the western interior (Lillegraven, 1979). Emry (1990) has recently described a middle Bridgerian fauna from Nevada that shares taxa in common with both California and the western interior, suggesting that some mammalian migration was possible between these areas, at least in the Bridgerian.

The climate along the southern California coastal lowlands during the middle Eocene ranged from tropical in the early Uintan to somewhat more arid and subtropical by the latter part of the Uintan (Lillegraven, 1979).
The western interior paleoclimate had begun to deteriorate by the early Uintan from a tropical, moist environment in the Bridgerian to a drier, more open environment in the Uintan, followed by a collapse of paleotemperatures in the latest Eocene/earliest Oligocene (Berggren and Prothero, 1992; Leopold et al., 1992; Wolfe, 1992). The warmer coastal lowlands may have served as a refuge area for tropical dwelling primates during the early Uintan. As paleoclimates continued to deteriorate towards the end of the Uintan in California, primates became less and less important members of paleocommunities. By the latest Eocene primates were essentially gone from North America with only the rare Mahgarita and Rooneyia, along with isolated occurrences of Omomys, Macrotauris, and Ourayia (West, 1982; Westgate, 1988, 1990) being represented from the southern part of Texas. The only possible later occurrence of a North American primate is the enigmatic Arikareean genus Ekgmowechashala (Macdonald, 1963, 1970; Rose and Rensberger, 1983) from South Dakota and Oregon (recently placed in the order Dermoptera by McKenna, 1990).

Hesperolemur, as with the many omomyid taxa known from southern California, seems to have found refuge in this warm, coastal region during the later portion of the middle Eocene. Hesperolemur may have migrated from the western interior, either from the Wyoming/Utah/Colorado area or perhaps from New Mexico along the Sevier orogenic belt during the late Bridgerian, reaching California by the early Uintan. Alternatively, Hesperolemur may have evolved from an unknown adapiform stock previously present in southern California.

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