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GENIC STUDIES OF LASIURUS (CHIROPTERA: VESPERTILIONIDAE)

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Bats of the genus Lasiurus present a number of interesting systematic problems that are difficult to resolve by traditional techniques. Members of the genus share a suite of derived morphological (Hall and Jones, 1961; Handley, 1960) and karyotypic (Bickham 1979, 1988) characteristics. However, until 1960 (Handley, 1960), members were placed in two genera—Lasiurus and Dasypterus—based primarily upon the presence or absence of the small, first upper premolar. Handley (1960) analyzed the differences and similarities among these two genera and concluded they were not distinct even at a subgeneric level. One goal of this study was to provide an estimate of genetic differentiation among the more divergent taxa in Lasiurus.

Additionally, a number of species-level taxonomic problems exist within the genus. Lasiurus borealis and L. seminolus are broadly sympatric in the eastern United States. They are morphologically similar, both externally and cranially, to the extent that they properly may be described as sibling species. Some workers, in fact, have suggested that these two taxa may represent only color phases of a single species.

The zoogeographic affinities of bats of the Antillean Islands were reviewed by Baker and Genoways (1978) and several problem species groups were noted. One of the taxa that needed more study included the several populations recognized by Varona (1974) as Lasiurus borealis. Representatives of this group of bats

are found on all Greater Antillean Islands and populations from each island have, at some time in the past, been accorded specific distinction. Varona (1974), without providing any supporting data, reduced all red bats from the Antillean Islands to subspecies of *L. borealis*.

A chromosomal difference exists between two currently recognized subspecies of Lasiurus ega that may signal these two taxa as specifically distinct (Baker and Patton, 1967; Baker et al., 1971). The X-chromosome of L. e. xanthinus from western México is submetacentric and resembles that of most vespertilionid bats, whereas in L. e. panamensis from southern Texas and eastern and southern México the X is acrocentric or subtelocentric, having undergone a pericentric inversion (Bickham, 1979, 1988).

This study examines the genic relationships of Lasiurus borealis (including specimens from Jamaica, Venezuela, Baja California, and the eastern United States), L. seminolus, L. cinereus, L. ega (including specimens from Suriname, Venezuela, Central America, and México), and L. intermedius. The choice of taxa was designed to give the kind of data necessary to examine the problems outlined above. Also, representatives of other vespertilionid genera were examined to provide outgroups for cladistic analysis (Hennig, 1966) in an attempt to better document the evolutionary relationships of the taxa of Lasiurus studied.

METHODS AND MATERIALS

Methods for tissue preparations, starch gel electrophoresis, and enzyme designations were similar to those of Selander et al. (1971) except for creatine kinase (CK) and peptidase (PEPT), which were described by Avise et al. (1980). PEPT-1 represents the most cathodally-migrating peptidase using the substrate L-leucyl-Lalanine; PEPT-2 and -3 represent the two most anodal zones of activity using the substrate leucyl-glycyl-glycine. Twenty-two presumptive loci, consisting of enzymatic and nonenzymatic proteins, were assayed (Table 1) as follows: CK-1, CK-2, CK-3, alpha-glycerophosphate dehydrogenase (α -GPD), glucose-6-phosphate isomerase (GPI), amino asparate transaminase-1, 2 (AAT-1, AAT-2) superoxide dismutase-1, 2 (SOD-1, SOD-2), isocitrate dehydrogenase-1, 2 (ICD-1, ICD-2), lactate dehydrogenase-1, 2 (LDH-1, LDH-2), malate dehydrogenase-1, 2 (MDH-1, MDH-2), mannosephosphate isomerase (MPI), PEPT-1, PEPT-2, PEPT-3, phosphoglucomutase-1, 2 (PGM-1, PGM-2), 6-phosphogluconate dehydrogenase (6-PGD).

TABLE 1.—Relative mobility of alleles for loci determined polymorphic within the genus Lasiurus. Where samples were polymorphic frequency of each allele is given in parenthesis. Monomorphic loci for Lasiurus were LDH-1,2; AAT-1,2; MDH-1,2 CK-2,3; SOD-1; PEPT-3.

Locus	(1) blossevillii	(2) blossevillii	(3) borealis	(4) degelidus	(5) seminolus	(6) cinereus	(7) xanthinus	(8) ega (Mx)	(9) ega (SA)	(10) intermedius	(11) P. subflavus
ICD-1	105	105	100	120	120	Null	145(.25) 140(.68) 120(.07)	120	130(.08) 120(.92)	130	150
ICD-2	-100	-100	-100	-90	-90	-90	-90	-90	-90	-90	-60
6PGD	100	100(.64) 90(.36)	100(.79) 90(.21)	50(.50) 30(.50)	80	100	110(.06) 90(.94)	110(.75) 90(.25)	110(.16) 90(.84)	110(.33) 90(.67)	140
α-GPD	40	100(.67) 80(.33)	100	100	100	105	103(.10) 60(.90)	60	60	103	120
GPI	50	50	100	100	100	48	45	125	125	125	130
MPI	110	110	100	110	100	100	140	100	100	100	80
PGM-1	100	100	100	90	90	95(.75) 90(.25)	92(.15) 90(.85)	105(.16) 92(.68) 80(.16)	92	85	140
PGM-2	100	100	100	100	100	80	75	100	100	100	125
CK-1	100	_	100	100	100	_	_	_	130	130	_
SOD-2	100	100	100(.86) -300(.14)	200	200	Null	250	200	200	200	150
PEPT-1	100	100	100	100	100	80	80	80	80	80	210
PEPT-2	100	100	100	100	105(.25) 100(.75)	100	95	100(.33) 95(.67)	100(.25) 95(.75)	100(.17) 95(.83)	100

Electromorph (allele) frequencies of 21 loci (CK-1 was excluded) were calculated from banding patterns. Nei's Identity (I) and Distance (D) matrices (Nei, 1972) were generated using modifications suggested by Hillis (1984). Cladistic analysis (Buth, 1984: Derr et al., 1987; Patton et al., 1981) was performed by hand using discrete character-state coding in which the locus was considered the character and the allelic composition of the locus was the character state. Additionally, side-by-side comparisons of alleles in Lasiurus were run with samples from Myotis velifer, M. thysanodes, M. yumanensis, M. nigricans, M. dominicensis, Pipistrellus subflavus, Nycticeius humeralis, and Eptesicus fuscus. Except as related to genic evolution in the genus Lasiurus (identification of unique alleles and the primitive and derived conditions for outgroup comparison), the details of the electrophoretic data from the other genera of vespertilionids are beyond the scope of this report.

RESULTS

Twenty-two electrophoretic loci were assayed. Loci found to be monomorphic for all Lasiurus examined, were as follows: LDH-1, -2; AAT-1, -2; MDH-1, -2; CK-2, -3; PEPT-3; SOD-1. Of these 10, three (AAT-1, CK-2, and SOD-1) distinguish Lasiurus from samples of the other four genera of Vespertilioninae examined. Electrophoretic data for the 12 polymorphic loci from the 10 samples are summarized in Table 1. None of the loci that was found to be polymorphic in Lasiurus shared an allele with other species of Vespertilioninae except PEPT-2 of Pipistrellus. Pairwise comparisons for Nei's Identity (I) and Distance (D) for the 10 samples are given in Table 2. The electrophoretic data are summarized phenetically (Fig. 1) by use of the unweighted pairgroup method of analysis (UPGMA—Sneath and Sokal, 1973) and cladistical analysis (Fig. 2) by the methods of Hennig (1966), Patton et al. (1981), and Buth (1984).

Discussion

Two aspects of our biochemical data support Handley's (1960) conclusion that yellow bats and red bats are congeneric. First, representatives from the two formerly recognized genera, Dasypterus and Lasiurus, are not more divergent from each other than L. borealis is from L. cinereus (species that were considered congeneric in the older classification). Second, the magnitude of biochemical divergence that distinguishes the three lineages in

TABLE 2.—Genetic distances (upper right) computed using the modification of Hillis (1984) of the formulae. Genetic identities (Nei, 1982) lower left, for data given in Table 1 and text.

		bl o (1)	blo (2)	bor (3)	deg (4)	sem (5)	cin (6)	xan (7)	ega (8)	ega (9)	int (10)	sub (11)
ı.	L. blossevillii (Ven)		.055	.222	.405	.484	.560	.742	.607	.618	.629	1.10
2.	L. blossevillii (NA)	.946		.168	.343	.417	.540	.694	.694	.577	.590	1.10
3.	L. borealis	.801	.845		.337	.275	.482	.738	.518	.515	.526	1.10
4.	L. degelidus	.667	.709	.714		.103	.530	.550	.347	.383	.464	1.10
5.	L. seminolus	.617	.695	.760	.902		.456	.550	.330	.321	.398	1.11
6.	L. cinereus	.571	.565	.617	.589	.634		.533	.446	.756	.469	1.10
7.	L. xanthinus	.476	.500	.478	.577	.577	.587		.366	.323	.405	1.25
8.	L. ega (Mex)	.545	.551	.596	.688	.736	.640	.694		.028	.123	1.18
9.	L. ega (SA)	.539	.562	.597	.682	.725	.634	.724	.973		.152	1.20
10.	L. intermedius	.533	.554	.591	.629	.672	.629	.667	.841	.859		1.22
11.	P. subflavus	.333	.333	.333	.333	.331	.333	.286	.307	.301	.295	

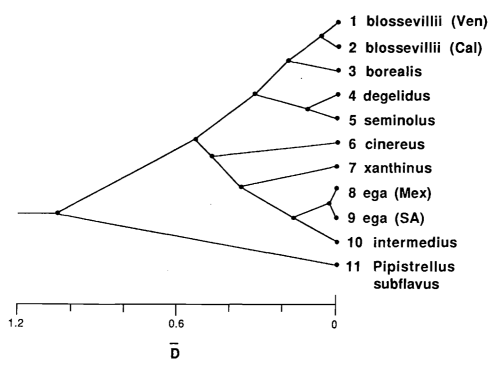


Fig. 1.—Phenogram generated from the electrophoretic data using average Nei's distance values (\bar{D}) and a UPGMA clustering analysis.

the genus Lasiurus is well within the range of divergence that characterizes comparisons of congeneric species of bats as well as other mammals (Arnold et al., 1982, 1983; Avise, 1974; Baker et al., 1981, 1985; Honeycutt et al., 1981; Koop and Baker, 1983; Straney et al., 1979). If only biochemical data were used as a basis for a systematic arrangement, the best alternative (because of the low level of genic differences that distinguish the three groups) would be to recognize a single genus with no subgenera (Fig. 1) and the second best arrangement would be to recognize three subgenera—1) Lasiurus, containing the red bats (distinguished by three shared fixed differences), 2) Dasypterus, including the yellow bats (distinguished by six shared fixed differences), and 3) a third subgenus containing the hoary bats (distinguished by six shared fixed differences). Essentially, our biochemical data are in agreement with Hall and Jones (1961), who proposed the early phylogeny of Lasiurus as consisting of three primary lineages.

Species-level Problems

Red bats.—As only PEPT-2¹⁰⁰ was shared among Lasiurus and other vespertilionine genera examined, it was rarely possible to determine which of the electromorphs was primitive or derived in

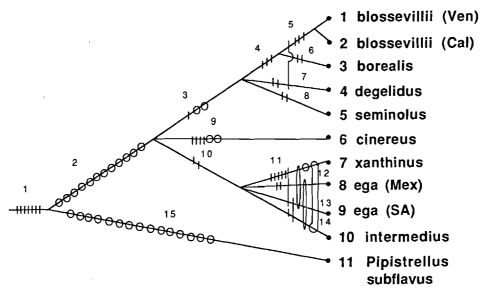


Fig. 2.—Phylogenetic tree generated by qualitative analysis of characters given in Table 3 and text using the method of Hennig (1966) and Patton *et al.* (1980). See Table 3 for definition of character states.

Lasiurus. Therefore, the functional outgroups for the red bats (L. borealis, L. seminolus, and the Jamaican Lasiurus) were restricted to the yellow bats (L. ega and L. intermedius) and L. cinereus, and for the yellow bats, the red bats and hoary bat served as the outgroups. Nonetheless, our data reveal patterns that have systematic implications.

Within *L. borealis* (as currently recognized), there is a significant genic demarcation between our samples from the eastern United States (Texas, South Carolina, and Georgia) and those from New Mexico, México, and South America. Eastern United States samples are separated from the New Mexican, Mexican, and Venezuelan samples by identity values at the 0.80 and 0.85 levels. Although the New Mexican, Mexican, and Venezuelan samples are separated by much greater geographic distance, their similarity values are much higher (0.95). Differences in ICD-1, GPI, and MPI are fixed between the populations in our samples. Cryptic species of other mammalian groups have similarity values in the range found in our comparison of South American-Mexican-New Mexican samples with those from the eastern United States (Avise, 1974).

Schmidly and Hendricks (1984) have demonstrated morphometric differences between eastern and western populations of *L. borealis*. Their samples from eastern Texas, representing *L. b. borealis*, were significantly larger in five of six cranial measure-

Table 3.—Electromorphs defining root and branches of phylogenetic tree represented by Fig. 2 as defined by Hennig (1966) and modified by Patton, et al. (1981). Characters representing assumed apomorphs (phenetically placed characters) are enclosed in brackets. Characters that must be strictly interpreted (in a cladistic sense) as ambiguous are italicized.

- 1. LDH-1¹⁰⁰, LDH-2⁻¹⁰⁰, MDH-1¹⁰⁰, MDH-2⁻¹⁰⁰, AAT-2⁻¹⁰⁰, CK-3¹⁰⁰, PEPT-2¹⁰⁰
- 2. [ICD- 1^{120} , ICD- 2^{-90} , 6PGD 100 , AAT- 1^{100} , CK- 2^{100} , MPI 100 , PGM- 1^{90} , PGM- 2^{100} , SOD- 1^{100} , SOD- 2^{200} , PEPT- 2^{100} , PEPT- 1^{80}]
- 3. αGPD^{100} , [GPI¹⁰⁰, PEPT-1¹⁰⁰]
- 4. ICD-2⁻¹⁰⁰, PGM-1¹⁰⁰, SOD-2¹⁰⁰
- 5. ICD-1¹⁰⁵, αGPD⁸⁰, GPI⁵⁰, MPI¹¹⁰
- 6. ICD-1¹⁰⁰, SOD-2⁻³⁰⁰
- 7. 6PGD⁵⁰, 6PGD³⁰, MPI¹¹⁰
- 8. 6PGD⁸⁰, PEPT-2¹⁰⁵
- 9. ICD-1^{null}, PGM-1⁹⁵, PGM-2⁸⁰, SOD-2^{null}, [αGPD¹⁰⁵, GPI⁴⁸]
- 10. 6PGD¹¹⁰, PEPT-2⁹⁵
- 11. ICD-1¹⁴⁵, ICD-1¹⁴⁰, MPI¹⁴⁰, PGM-2⁷⁵, SOD-2²⁵⁰, PGM-I⁹², [GPI⁴⁵, α GPD¹⁰³, α GPD⁶⁰]
- 12. PGM-1¹⁰⁵, PGM-1⁸⁰, PGM-1⁹², [GPI¹²⁵, α GPD⁶⁰]
- 13. $ICD-1^{130}$, $PGM-1^{92}$, $[GPI^{125}$, $\alpha GPD^{60}]$
- 14. PGM-1⁸⁵, $ICD-1^{130}$, $[GPI^{125}, \alpha GPD^{103}]$
- 15. [ICD-1¹⁵⁰, ICD-2⁻⁶⁰, 6PGD¹⁵⁰, α GPD¹²⁰, AAT-1¹²⁰, GPI¹³⁰, MPI⁸⁰, PGM-1¹⁴⁰, PGM-2¹⁷⁵, SOD-1¹⁷⁵, SOD-2¹⁵⁰, PEPT-2¹⁰⁵, PEPT-1²¹⁰]

ments of males and all six measurements of females than three samples of *L. b. teliotis*, including two from Tamaulipas in northeastern México. The western populations also differ from those to the east in pelage characteristics, including rusty-red rather than brownish dorsal coloration, noticeably fewer frosted dorsal hairs, and the posterior margin of the uropatagium is bare or only sparsely haired rather than well furred to the posterior margin (Bogan and Williams, 1970).

Based on these significant morphological and genic differences, we believe that the western and eastern populations of *L. borealis* are best considered distinct species. The specific name *L. borealis* is here restricted to eastern populations designated *L. b. borealis* by Hall (1981), but regarded by us as a monotypic species. The senior synonym for the western populations is *Vespertilio blossevillii* Lesson and Garnot, 1826 (type locality Montevideo, Uruguay). The appropriate trinomials for populations examined

in our study would be Lasiurus blossevillii teliotis and Lasiurus blossevillii frantzii. Researchers should be alert for sympatric populations or indication of hybridization between these two species in southwestern New Mexico, western Texas, and northeastern México.

Lasiurus borealis has a similarity level with L. seminolus of 0.76 (including five fixed differences—ICD-1, -2; 6 PGD; PGM-1; SOD-2), which is compatible with the conclusion that the seminolus and borealis represent distinct species, not sympatric color phases of a single species. Specimens of Lasiurus from Jamaica have a similarity with mainland populations of L. borealis of 0.71 and with L. blossevillii of 0.67 (Table 2), which implies that L. degelidus is best recognized as a species distinct from both borealis and blossevillii. However, Lasiurus from Jamaica have a much higher similarity level (0.90) with L. seminolus; therefore, another possibility would be to recognize L. degelidus as a race of L. seminolus. Cladistic analysis of the alleles (ICD-1¹²⁰, ICD-2⁹⁰, and SOD-2²⁰⁰) shared by seminolus and degelidus, but which are distinct from those of L. borealis, failed to provide any data that document these shared alleles as derived (synapomorphies). Additionally, a cladistical analysis of the one character (MPI¹¹⁰) shared by borealis and degelidus, but not present in seminolus, indicates that MPI¹⁰⁰ of seminolus is primitive. This means that, although there is a higher similarity value for degelidus and seminolus, cladistic characters (synapomorphies) ally degelidus more closely with borealis than with seminolus. However, due to the possibility of an ancestral MPI¹⁰⁰, ¹¹⁰ polymorphism, it still is possible that degelidus arose from a seminolus stock rather than a borealis stock. Specimens of borealis and seminolus differ morphologically in that borealis possesses a protuberance along the anterior border of the lachrymal ridge (Hall, 1981: fig. 178). Examination of a specimen from St. Ann Parish, Jamaica (TTU 22080), and one from Department du Sud, Haiti (TTU 22804), revealed that the condition of lachrymal ridge in these specimens most closely resembles that of L. seminolus.

Specimens of borealis and seminolus traditionally have been distinguished on the basis of pelage color, but this character is not definitive in that the specimen from Jamaica most closely resembles seminolus and the one from Haiti most closely resembles borealis. We conclude that, in light of the above data, the best course is to recognize L. degelidus as a distinct species,

but future data should be evaluated in light of the possibility that degelidus, as well as other Antillean populations, may be subspecies of L. seminolus. Of course, data from Cuban, Hispaniolan, Puerto Rican, and Bahamian red bats are needed before final decisions can be made.

Yellow bats.—Electrophoretic data for yellow bats suggest a dichotomy within Lasiurus ega that, in our opinion, signals specific differences. Although specimens of L. ega from Venezuela and Suriname are geographically widely separated from those from Chiapas and Guerrero, similarity values are at the level (0.97) expected for conspecific populations and no fixed differences were found between the two groups. On the other hand, specimens of L. e. xanthinus from Baja California and Neuvo León, are fixed for four different alleles from other samples currently recognized as L. ega (GPI⁴⁵, SOD-2²⁵⁰, MPI¹⁴⁰, PGM-2⁷⁵) and have a low (0.69 to 0.72) similarity to the other Mexican and South American samples.

Also of interest is the high level of similarity 0.84 and 0.86 between L. intermedius and the South American and southern Mexican specimens of L. ega. There is no doubt that ega and intermedius are recognizable, widely sympatric species. However, if electrophoretic data were used to indicate systematic position, we would conclude that L. ega (which has an acrocentric X cytotype) is more closely related to L. intermedius than to what currently is known as L. e. xanthinus (which has a biarmed X cytotype) (Fig. 1). Lasiurus intermedius possesses an acrocentric X chromosome that apparently has evolved by a pericentric inversion. Within vespertilionids, a submetacentric X chromosome is considered the primitive condition with the acrocentric condition having evolved independently in several genera (Baker, 1970; Bickham, 1979, 1988; McBee et al., 1986).

The most parsimonious explanation of the evolution of the inverted X in two species of Lasiurus is to postulate a common origin for those taxa (L. intermedius and L. e. panamensis) as indicated also by electrophoretic data. However, it is also obvious that an acrocentric X has evolved at least twice (McBee et al., 1986) in vespertilionids (to explain its presence in some species of Plecotus and in some species of Lasiurus), and the possibility of convergent evolution in Lasiurus cannot be ruled out. That congruence occurs within the electrophoretic and chromosomal data sets for the yellow bats suggests the possibility of common ancestry for the taxa of Lasiurus with an inverted acrocentric X

(L. ega and L. intermedius shared a common ancestry after separating from L. xanthinus) should remain a viable systematic hypothesis.

We believe that the appropriate interpretation of these data is to recognize L. xanthinus (type locality Sierra Laguna, Baja California) as a species distinct from L. ega. It is distinguished from ega by a submetacentric X-chromosome and genically by four fixed electromorphs (Table 1). Morphologically the two species are distinguished by pelage coloration, which is a brighter yellow, especially on the anterior third of the uropatagium, in most specimens of L. xanthinus. Comparing measurements of the two taxa from the published literature, it appears that the only measurement that may distinguish them is length of the maxillary toothrow, means for females (with extremes in parentheses) are as follows: L. xanthinus from Baja California, 5.7 (5.4 to 5.9) (Jones et al., 1965) and Arizona, 5.9 (5.8-6.0) (Hoffmeister, 1986) as compared to L. ega from Texas, 5.4 (5.1 to 5.6) (Baker et al., 1971) and Tamaulipas, 5.4 (5.4 to 5.5) (Schmidly and Hendricks, 1984, who originally assigned this population to L. e. xanthinus but we believe it is best considered as L. e. panamensis). Although the level of morphological distinctiveness for xanthinus and ega is not as great as is usually characteristic of currently recognized mammalian species, the degree of genic differences, which are fixed in our samples, is similar to that found in sympatric species of another vespertilionid bat, Rhogeessa, for which no morphological differences have been found (Baker, 1984).

Ecologically, L. xanthinus seems to be associated with the dry thorny vegetation of the Mexican Plateau, coastal western México including parts of Baja California, and the deserts of the southwestern United States. In the data available to us, the easternmost record of this species is from 20 mi. N Santa Anna, Nuevo León (this paper), and the southernmost record is from Oaxtepec, Morelos (Baker and Patton, 1967). We would expect potential sympatry or hybridization between L. xanthinus and L. ega along the eastern and southern edges of the Mexican Plateau. We believe that L. e. panamensis occupies the Gulf versant as far north as 5 mi. SE Brownsville, Texas; in southern México this taxon occupies both versants as well as most if not all of the intervening highlands.

Hoary bats.—Although our sample of L. cinereus included specimens from three states within the United States and two

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