

Evolutionary systematic of three species of troglobitic beetles: electrophoretic and morphological evidence

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Summary – Genic and morphological variations were compared for 3 allopatric and endemic troglobitic beetles of the genus *Speonomus*, by use of an allozyme data set (18 putative loci) and 1 based on morphometric characters (16 morphological variables). Allozymic and morphometric relationships were also compared with some aspects of the mating behaviour, and considered in relation to the ecology and biogeography of these species. The extent of agreement between the assessments of evolutionary divergence at the genic and morphological levels is discussed. Enzyme analysis revealed pronounced differences in degree of genetic differentiation between the 3 species: *S colluvii* is the most divergent species with large genetic distances ($D \simeq 1$). Morphometric differentiation between the 3 species, assessed on 16 characters, is important between the 3 species, principally between *S zophosinus* and the 2 others. The level of congruence is strongly data – set dependent and these results reveal independent trends for the 2 sets of characters. While *S zophosinus* has diverged little from *S hydrophilus* on the molecular level, the morphometric differentiation between them is high. This pattern may result from different sensitivities of each character set to different components of the environment.

troglobitic beetle / systematic / electrophoretic and morphometric variations

Résumé – Systématique évolutive de trois espèces de Coléoptères troglobies: analyses électrophorétiques et morphologiques. Nous avons comparé les variations génétiques et morphologiques de trois espèces de Coléoptères troglobies allopatriques et endémiques du genre *Speonomus* sur la base des données allozymiques (18 locus) et des caractères morphométriques (16 variables). Les distances basées sur les données génétiques et les divergences morphologiques et biométriques sont également comparées à certains aspects du comportement lors de l'accouplement et discutées en fonction du contexte écologique et des caractéristiques biogéographiques de ces espèces. L'analyse des locus, correspondant à 19 protéines enzymatiques, met en évidence des différenciations importantes entre ces espèces: *S colluvii* est l'espèce la plus divergente avec une distance génétique voisine de 1. L'étude morphométrique estimée sur 16 caractères sépare les populations des trois espèces selon la largeur du pronotum et la longueur des élytres (*S zophosinus* les plus petits, *S hydrophilus* les plus grands), la largeur des articles antennaires et la forme du 8^e article antennaire. Il n'y a pas concordance entre les données moléculaires (gènes structuraux) et morphométriques: *S zophosinus* sur le plan moléculaire a peu divergé de *S hydrophilus*

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alors que la différenciation morphométrique est importante entre ces espèces. L'action différentielle des pressions de sélection aux différents niveaux, protéique et morphologique, peut rendre compte de ces discordances.

coléoptère troglobie / systématique / variations électrophorétiques et morphométriques

INTRODUCTION

Closely related species provide opportunities to study how the evolutionary process operates in natural populations. Nevertheless, it is often difficult to assign taxonomic ranks or to reconstruct phylogenetic relationships. A high degree of independence is often observed between molecular and morphological evolution (Gorman and Kim, 1976; Sene and Carson, 1977; Schnell *et al*, 1978; Turner *et al*, 1979; Lessios, 1981; Berlocher and Bush, 1982; Allegrucci *et al*, 1987); this may be due to varying selection pressures on different traits (Turner *et al*, 1979) or merely related to the number of loci segregating for the different characters (Lewontin, 1984). Electrophoretic separation of enzymes has become a powerful tool for investigating systematic and evolutionary problems (Avisé, 1974). Morphological studies alone are usually inadequate to determine evolutionary relationships in closely related species. Integrative studies utilizing allozyme variability and data from behavioural breeding systems in addition to morphometric variability are most powerful in such complex groups; then we can estimate the importance of gene flow, genetic drift and selection in shaping population structures.

The 3 species of Bathysciinae (Coleoptera) of this study belong to a monophyletic group of the genus *Speonomus* which contains many cave-dwelling species from the more primitive to the more specialized (Jeannel, 1924). Studies on these species have been the subject of repeated analyses by various approaches, including the study of morphology (Jeannel, 1924; Juberthie *et al*, 1980a; Juberthie *et al*, 1981; Delay *et al*, 1983), karyology (Durand and Juberthie-Jupeau, 1980) and allozymic variation (Crouau-Roy, 1989a, b). The relationship of these beetles to one another and to other species groups of *Speonomus* has not been adequately resolved. The 3 allopatric species *S hydrophilus*, *S zophosinus* and *S colluvii*, narrowly endemic to central Pyrenees (fig 1), have undergone some ecological divergence and present a high degree of specialization to underground life (*eg* life cycle of 1 larval stage with suppression of larval feeding, Deleurance-Glaçon, 1963). These blind and wingless troglobitic species occur in caves and deep soil (Juberthie *et al*, 1980a, b) in the massif Arize (*S hydrophilus* at about 430 - 1440 m), in the valleys of Salat and Arac rivers (*S zophosinus* at about 480 - 620 m) and on the north face of the massif des Trois Seigneurs (*S colluvii* at about 700 m - 1350 m) at the foothill of the French central Pyrenees. The southern extent of the massif Arize (between the area of *S hydrophilus* and *S zophosinus*) is composed of colmated metamorphic rocks which seem to be geological barriers to underground colonization by cave fauna. Their distribution strongly suggests that speciation occurred after gene flow was interrupted by physical barriers: there were probably geomorphological and pedological barriers after climatic shifts during the ice ages and interglacials of the quaternary.

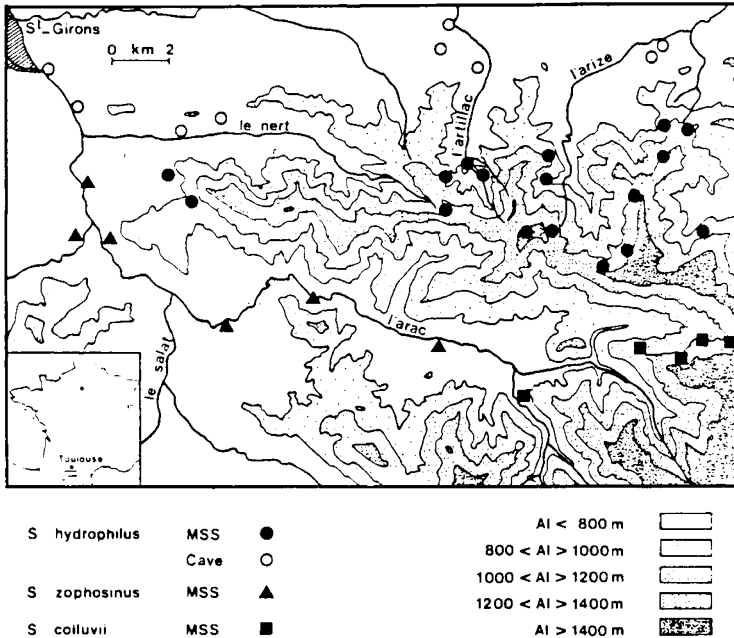


Fig 1. Geographical location of sampled populations for the electrophoretic and the morphometric study (Map of the distribution of the 3 species of *Speonomus* with the sample sites).

The phylogeny of these 3 closely related troglobitic (*ie*, obligate cave dwelling) beetles was investigated using isozyme electrophoresis and biometric analysis. Patterns of allozyme and biometric variation were compared with some aspects of the mating behaviour and aspects of the ecology of these organisms to address the question of intrageneric relationships of species within this group of beetles.

The present study provides answers to the following questions: 1) how much genetic variation is contained in these populations? 2) how is the genetic variation partitioned within and among the isolated populations? 3) what phylogenetic relationships can be deduced from the isozyme data and how congruent are they with biometrical based relationships at the species level? 4) how do the different approaches contribute to our knowledge of evolution in this group?

MATERIALS AND METHODS

Twenty-three populations of *S. hydrophilus*, 6 of *S. zophosinus* and 5 of *S. colluvii* were studied for electrophoretic, morphometric and behavioural studies.

Electrophoretic analyses

Electrophoresis was performed in slabs of 12% hydrolized starch for the following enzymes: esterases (Est-1, Est-6, EC 3.1.1.2), alkaline phosphatase (Aph, EC 3.1.3.1),

leucine aminopeptidase (Lap-1, EC 3.4.11.1), malic enzyme (Me, EC 1.1.1.40) hexokinases (Hk-1, Hk-3, Hk-4, EC 2.7.1.1) phosphohexose isomerase (Phi-1, Phi-2, EC 5.3.1.9) and α glycerophosphate dehydrogenase (α -Gpdh, EC 1.1.1.8); in slabs of acrylamide for fumarase (Fum, EC 4.2.1.12), hydroxybutyrate dehydrogenase (Hbdh-1, Hbdh-2, EC 1.1.1.30), acid phosphatase (Pac-1, EC 3.1.3.2) malate dehydrogenase (Mdh, EC 1.1.1.37), and lactate dehydrogenase (Ldh, EC 1.1.1.28), aldehyde oxidase (Ao, EC 1.2.3.1). Additional loci that could not be scored unequivocally were deleted from further analysis. Loci and alleles were numbered, respectively, in order of decreasing anodal migration. Electrophoresis buffers and stains are described in Crouau-Roy (1986). Individuals from different species were run together on the same gel to determine whether corresponding electromorphs had similar mobilities. Allozyme frequencies for each sample were derived from the electrophoretic results. These data were employed to compute genetic distance estimates (Nei, 1972, 1978) which were used for a phenetic averaging (UPGMA; Sneath and Sokal, 1973). The apportionment of genetic diversity was determined using genetic diversity analysis (Wright, 1965; Nei, 1977, 1986). Total gene diversity ($H_T = 1 - \sum x_i^2$: weighted average allele frequencies over all populations) is subdivided into gene diversity within populations (H_S : weighted average over all populations of the values $1 - \sum x_i^2$ for each population) and gene diversity among populations. Differentiation among populations is calculated as $F_{ST} = H_T - H_S / H_T$.

Morphometric analysis

Sample sizes for all morphological analyses were 30 adults per population. For *S. hydrophilus* only 18 populations out of the 23 were examined. The following 16 morphological variables were measured from each specimen: length (L) and width (W) of 7 antennal segments (5th, 6th, 7th, 8th, 9th, 10th and length only of the 11th), length of the tibia of the 3rd pair of legs (LT), length of the elytra (LE) and width of the pronotum (WP). Measurements were recorded in micrometres. Because of statistically significant correlations between values for the sexes, the measurements were only made in males. Euclidean distances were calculated on the basis of the 16 morphometric variables; a dendrogram was drawn up using these distances, according to the UPGMA method of cluster analysis (Sneath and Sokal, 1973). Data were also analyzed using a principal components analysis (PCA) on the standardized variables.

RESULTS

Electrophoretic differentiation

Genetic structure of populations at enzyme loci

The allele frequencies for each putative locus (18 loci) are given in table I. Each locus containing more than 1 variant was considered polymorphic. Fourteen of these were polymorphic (Est-6, Lap-1, Phi-1, Phi-2, Pac-1, Hk-3, Hk-4, Hbdh-1, Me, Mdh, α -Gpdh, Ldh-2, Fum, Ao) and the remaining 4 were monomorphic with the same electromorph fixed in all populations of the 3 species (Hbdh-3, Aph,

Table I. Gene frequencies for loci coding for monomorphic and/or polymorphic proteins within each species of *Speonomus*. *N* is the number of beetles sampled. Anodal mobilities relative to the common allele (termed 100) are used for allozyme nomenclature.

* Monomorphic loci among the 3 species: *Aph*, *Hbdh-3*, *Est-1* and *Hk-1*.

<i>Loci</i> *	<i>S hydrophilus</i> (23 localities)	<i>S zophosinus</i> (6 localities)	<i>S colluvii</i> (5 localities)
	84	—	0.354
	86	—	0.126
	88	—	0.046
	90	—	0.217
	92	—	0.231
	94	0.004	0.029
Est-6	96	0.011	—
	98	0.085	0.463
	100	0.473	0.514
	102	0.196	0.022
	104	0.192	—
	106	0.038	—
	N	2053	844
	98	0.020	—
	100	0.732	0.381
Lap-1	102	0.088	0.619
	104	0.160	—
	N	1447	648
	96	—	0.857
	98	0.004	0.143
Phi-1	100	0.450	0.741
	102	0.463	0.259
	104	0.083	—
	N	2074	511
	98	0.002	—
	100	0.986	1
Hk-3	102	0.013	0.112
	104	—	0.495
	N	1327	560
	100	0.994	1
Hk-4	102	0.006	0.090
	104	—	0.796
	N	1315	560
	96	—	1
	98	0.004	—
ME	100	0.979	0.946
	102	0.014	—
	104	0.004	—
	M	1467	672
	96	—	212
	98	—	0.638
α -Gpdh	100	1	—
	N	1301	422
			206

Table I. continued

<i>Loci</i> *		<i>S hydrophilus</i> (23 localities)	<i>S zophosinus</i> (6 localities)	<i>S colluvii</i> (5 localities)
MDH	100	1	—	—
	102	—	1	—
	104	—	—	0.574
	106	—	—	0.426
	N	1187	306	263
Pac-1	98	0.010	0.989	—
	100	0.990	0.011	—
	102	—	—	1
	N	1352	394	250
Hbdh-1	100	1	1	—
	102	—	—	1
	N	921	240	200
Fum	100	1	1	—
	102	—	—	1
	N	988	238	203
Ao	100	1	—	—
	102	—	1	—
	104	—	—	1
	N	920	240	203
Ldh-2	100	1	—	—
	102	—	1	—
	104	—	—	1
	N	920	244	200
Phi-2	98	—	—	1
	100	1	—	—
	102	—	1	—
	N	2050	500	211

Est-1, Hk-1). Six loci (Mdh, Hbdh-1, Fum, Ao, Ldh-2 and Phi-2) are diagnostic for at least 1 species, and 2 additional loci (Est-6, Pac-1) are diagnostic with a 1% probability of error. Estimates of the proportion of polymorphic loci per population (P) and the average frequency of heterozygous loci per individual (H) indicate some differences in genetic variability between the 3 species. *Speonomus colluvii* displays the highest overall percentage of polymorphic loci (41.2%). For the other 2, percent polymorphic loci are 23.0% in *S hydrophilus* and 25.5% in *S zophosinus*. Expected average heterozygosities, H_{exp} , are listed in table II. Genotypic frequencies are

compared to expected Hardy-Weinberg proportions by χ^2 or G -test (Sokal and Rohlf, 1969); χ^2 or G values show significant differences between observed genotypic frequencies and those expected under Hardy-Weinberg equilibrium. The genotypic fixation index (Wright, 1965), which measures the relative difference between expected (H_{exp}) and observed (H_{obs}) heterozygosity, shows a significant deficiency of heterozygotes ($F_{IS} > 0$) (table II) (Crouau-Roy, 1988).

Table II. F -statistics and heterogeneity χ^2 values for the 3 species of *Speonomus*. ns: non significant; *: $P < 0.01$. *S. hydrophilus*: $P = 0.230$ (0.015); $H_{obs} = 0.062$ (0.014); $H_{exp} = 0.107$ (0.029).; *S. zophosinus*: $P = 0.255$ (0.012); $H_{obs} = 0.054$ (0.006); $H_{exp} = 0.088$ (0.008).; *S. colluvii* $P = 0.412$ (0.000); $H_{obs} = 0.097$ (0.002); $H_{exp} = 0.184$ (0.004).

	<i>Est-6</i>	<i>Lap-1</i>	<i>Phi-1</i>	<i>Me</i>	<i>Pac-1</i>	<i>Hk-3</i>	<i>Hk-4</i>	α - <i>Gpdh</i>	<i>Mdh</i>
<i>S. hydrophilus</i>									
F_{IS}	0.409	0.365	0.405	0.520	0.447	0.456	0.545	-	-
F_{IT}	0.556	0.497	0.452	0.542	0.475	0.492	0.582	-	-
F_{ST}	0.249	0.208	0.080	0.047	0.052	0.068	0.083	-	-
χ^2	1022.4*	601.9*	347.8*	137.9*	140.3*	192.7*	218.3*	-	-
<i>S. zophosinus</i>									
F_{IS}	0.351	0.481	0.428	0.403	0.497	-	-	-	-
F_{IT}	0.581	0.483	0.430	0.403	0.518	-	-	-	-
F_{ST}	0.354	0.005	0.004	0	0.043	-	-	-	-
χ^2	597.5*	6.5 ns	4.1 ns	0 ns	30.4*	-	-	-	-
<i>S. colluvii</i>									
F_{IS}	0.367	0.486	0.442	-	-	0.443	0.385	0.507	0.468
F_{IT}	0.379	0.546	0.448	-	-	0.449	0.485	0.539	0.477
F_{ST}	0.019	0.118	0.012	-	-	0.012	0.164	0.006	0.018
χ^2	4.7 ns	38.5*	4.5 ns	-	-	4.9 ns	61.6*	2.2 ns	6.8 ns

Genetic differentiation between populations

The genetic differentiation within the *Speonomus* species complex was determined using genetic diversity analysis (F -statistics: Wright, 1965; Nei, 1977, 1986) and a χ^2 contingency analysis of heterogeneity (Workman and Niswander, 1970). Nire loci are variable in some or all populations of each species (table II). Significant heterogeneity in gene frequencies χ^2 was observed among *S. hydrophilus* populations at all variable loci. Significant heterogeneity in allele frequencies was observed among populations of *S. zophosinus* (only at the Est-6 locus). For *S. colluvii*, 2 variable loci (*Lap-1* and *Hk-4*) show significant heterogeneity. There is significant genetic variation between populations in the complex of 3 species (table III). Genetic diversity for 5 loci is due totally to the between-species component ($F_{ST} = 1$; populations fixed for different alleles: *Hbdh-1*, *Fum*, *Ao*, *LDh-2* and *Phi-2*). For 3 additional loci (*Mdh*, *Pc-1*, *Me*) almost all the diversity is due to between-species variation rather than within-species variation ($F_{ST} = 0.978$, 0.979 and 0.902 , respectively). The mean of 0.790 (for polymorphic loci only) for differentiation

Table III. Genetic diversity among populations in the species complex of *speonomus* F_T = total heterozygosity, H_S : average heterozygosity per population, F_{ST} : fixation index. *Values are given only for polymorphic loci.

<i>Differentiation among populations</i>			
<i>Locus</i>	H_T	H_S	$F_{ST} = \left(\frac{H_T - H_S}{H_T} \right)$
Est-6	0.857	0.506	0.409
Lap-1	0.502	0.366	0.266
Phi-1	0.751	0.389	0.482
Hk-3	0.465	0.198	0.574
Hk-4	0.445	0.103	0.769
Me	0.478	0.047	0.902
Gpdh	0.497	0.150	0.698
Mdh	0.722	0.160	0.978
Pac-1	0.667	0.014	0.979
Fum	0.446	0	1.000
Ao	0.667	0	1.000
Hbdh-1	0.446	0	1.000
Ldh-2	0.667	0	1.000
Phi-2	0.667	0	1.000
Mean	0.591	0.138	0.790

among species indicates that a large portion of the genetic variability in *Speonomus* is not present among the individuals of a single species.

Extensive isozyme divergence between species is further indicated by their low Nei genetic identity values.

Nei's standard genetic distances corrected for sample size between the populations of each species are listed in table IV. Variances of Nei's distances (Nei and Roychoudhury, 1974) were all on the order of 1.7%-2.5% of the distance values. Beetles from the 3 species are well differentiated: coefficients of genetic distance range between 1.021 and 0.400. The genetic differentiation observed between the species *S. hydrophilus* and *S. zophosinus* is significant ($D = 0.431$) compared to that observed within each of them (intraspecific values of Nei's index: 0.036 and 0.050 respectively). *S. zophosinus* differed completely from *S. hydrophilus* at 5 isozyme loci (Mdh, Ao, Ldh-2, Phi-2, Est-6) and differed substantially at 2 others (Phi-1, pac-1). The distribution of the loci with respect to genetic identity exhibits the U-shaped pattern characteristic of comparisons between good species. Of 18 loci, most are either identical in their allelic composition (55%) or completely different (32% of its loci are diagnostic for the other). For *S. colluvii*, the genetic differentiation from the other 2 species is greater and affects an important part of the genome (50% of its loci are diagnostic: Ao, Fum, Pac-1, Me, Mdh, Hbdh-1, Ldh-2, α -Gpdh, Phi-2). The genetic distances between pairs of populations are summarized in figure 2 in the form of a dendrogram by using the UPGMA clustering method: this analysis of the data indicates the pattern of phenetic clustering of the species into 3 major groups.

Table IV. Nei's standard genetic distance (below the diagonal) based on allozymic variables and Euclidean distances (above the diagonal) based on morphometric variables. Values in parentheses are standard errors.

	<i>S hydrophilus</i> (23 populations)	<i>S zophosinus</i> (6 populations)	<i>S colluvii</i> (2 populations)
<i>S hydrophilus</i>	—	263.09 (5.68)	116.70 (5.82)
<i>S zophosinus</i>	0.430	—	187.55 (16.47)
<i>S colluvii</i>	1.002	1.020	—
Mean Nei's distance	0.036	0.050	0.014
Mean Euclidean distance	54.33 (1.71)	76.70 (10.09)	71.99 (0.00)

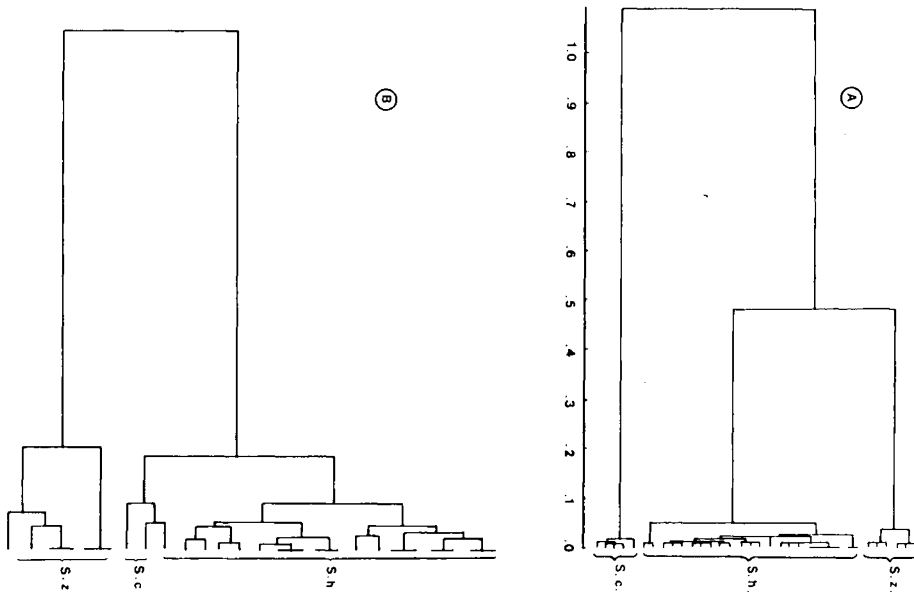


Fig 2. Comparison of the results of the allozymic (A: Nei's genetic distances) and morphometric analysis (B: Euclidean distances) of the 3 species of *Speonomus*, using the UPGMA clustering method. *Sz*, *S zophosinus*; *Sh*, *S hydrophilus*; *Sc* *S colluvii*.

Morphometric differentiation

The average values for the 16 morphological characters are given in table V and relationships between populations of the 3 species have been tested by a Student test. The *S hydrophilus* and *S zophosinus* populations display significant differences for all measured morphological characters; between *S hydrophilus* and *S colluvii*, 4 comparisons are significant (L8, L7, L6, L5) and only 2 are significant (L11, W9) between *S zophosinus* and *S colluvii*. In particular, the shape of the 8th antennal segment (L8 and W8) differentiates the 3 species from each other (fig 3).

Table V. Average values of 16 biometrical traits for the troglotic beetles and comparison by a *t*-test of populations coming from different species. Units for all measurements are micrometres. *** $p \leq 0$. ** $p \leq 0.0001$. ()^a: degrees of freedom.

Character	S Hydrophilus		S zophosinus (Standard error)		S colluvii		Hyd zop (22) ^a	t-test	
								Hyd coll (18) ^a	Zop coll (6) ^a
WP	1258	(5.2)	1152	(12.6)	1188	(20.9)	8.24**	3.76	1.44
LT	896	(3.6)	789	(9.7)	870	(2.0)	11.97***	2.13	4.48
L11	183.61	(0.6)	160	(1.5)	185	(1.3)	20.64**	0.67	12.73**
L10	124	(0.6)	105	(1.4)	117	(2.5)	14.11***	3.35	4.34
W10	78	(0.6)	72	(1.1)	80	(2.8)	4.64**	0.38	2.50
L9	143	(1.2)	117	(2.2)	133	(1.9)	9.45**	1.92	4.26
W9	78	(3.4)	66	(0.7)	77	(1.4)	7.53**	3.25	9.53**
L8	129	(0.6)	105	(1.6)	107	(0.4)	14.18**	8.49**	0.71
W8	55	(0.4)	51	(0.9)	58	(0.5)	4.31*	1.20	3.67
L7	178	(1.0)	144	(2.6)	155	(0.4)	13.37***	6.84**	1.92
W7	70	(0.4)	62	(0.7)	74	(0.3)	6.74**	1.39	4.74
L6	176	(0.7)	139	(1.0)	143	(0.5)	27.10***	16.10***	0.87
W6	53	(1.1)	45	(0.5)	51	(0.3)	9.08**	1.09	3.46
L5	202	(1.2)	167	(2.9)	189	(0.5)	12.66***	4.94**	2.65
W5	51	(0.4)	45	(0.6)	51	(0.3)	7.34**	0.62	3.46
LE	1826	(6.0)	1641	(18.1)	1796	(28.8)	12.60***	1.31	4.36

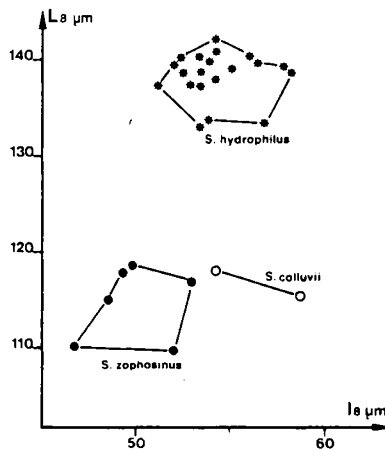


Fig 3. Separation of the populations according to the form of the 8th antennal segment (L_8 and W_8).

Table IV reports, with Nei's distances, matrices of intra- and interspecific Euclidean distances on morphometric measurements. These data indicate that variations in morphology are more important between species than they are within each species. Figure 2 compares the results of the clustering based on allozyme

(A, Nei's distances) and morphometric data (B, Euclidean distances). On the 2 bases, populations are clustered into 3 groups corresponding to the 3 species except for 1 population of *S hydrophilus* which clusters with the group of *S colluvii* populations. Nevertheless, the morphometric differences between the 3 allopatric species of *Speonomus* are discordant with the pattern displayed by electrophoretic dissimilarities. While on the molecular level, *S zophosinus* has diverged less from *S hydrophilus* than has *S colluvii*, the morphological distance between these 2 species is greater (Euclidean distance: 263.09 ± 5.68). The marked dissimilarity of *S zophosinus* is distinctly illustrated in a plot of the 3 species (conspecific populations pooled) on the 1st 2 axes generated by the principal components' analysis (fig 4). The raw data, and the principal-component scores, are dominated by size: the 1st principal component, explaining 82% of the variance, is associated with general size, with near-equal contributions from each character. The 3 species differ according to the general size of the individuals (*S zophosinus* individuals are the smallest and *S hydrophilus* individuals the largest) and also according to the width of the 8 and 10 antennal segments (explained by the 2nd principal component).

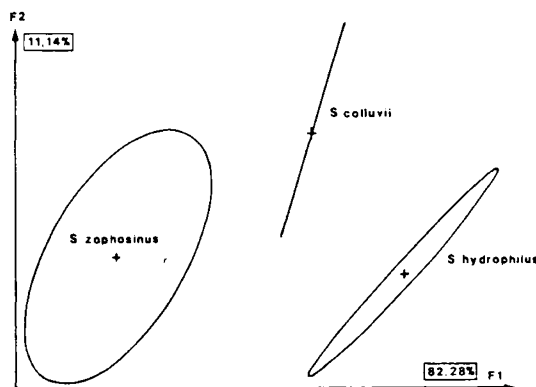


Fig 4. Principal components analysis of *Speonomus* populations based on morphometric data. Ellipses of equiprobability (95%) are calculated according to Healy (1972).

DISCUSSION

In systematic studies, the choice of parameters may substantially influence the patterns observed. Since selection intensity is difficult to assess, consideration of more than a single class of traits can allow a more reasonable evaluation of the accuracy of systematic schemes. Several investigations have demonstrated that speciation can occur with little or no divergence at structural gene loci coding for allozymes: classes of genes other than those studied by electrophoresis (eg regulatory genes) might be involved in this process to a greater extent Ayala,

1975; Wilson, 1976; Nevo, 1978). These species of *Speonomus*, show high degrees of genetic divergence; this genetic differentiation does not correspond to the differences in allele frequencies at the polymorphic loci but to the fixation of specific alleles. The greater part of the isozyme loci examined contain a large portion of their genetic diversity among species rather than within species (table III) despite the high level of within – population variation in some local populations of the 3 species (table IV). Electrophoresis usually underestimates the amount of variability in a species or population. Nevertheless, electromorphs can provide valuable taxonomic information and we found 32 and 50% of the loci studied to be diagnostic for any of the 3 sibling species of *Speonomus*. Moreover, an obvious disparity in size and behaviour during mating between the 3 species is visible. Ethological criteria have a great importance in speciation events. These *Speonomus* show within a single species a very low experimental frequency of matings (Crouau-Roy, 1986) in contrast to the 2 larval stages of *S delarouzei* (Juberthie-Jupeau and Cazals, 1984). Thus, the occurrence of reproductive isolation between the 3 species cannot be shown. Nevertheless, during mating, constant and sometimes important differences appear between the species; they concern the length of mating, the different phases of the mating and the amplitude and frequency of abdominal movements of males during mating (Crouau-Roy, 1986). These different types of mating behaviour may enable one to discriminate the 3 species.

The consistent differences in electrophoretic differentiation, morphometric and behavioural differences, in conjunction with biogeographical distributions and ecological separation, offer convincing evidence of the separate species status of *S hydrophilus*, *S zophosinus* and *S colluvii*. The results of this study applied to the taxonomic question are in complete agreement with the previously determined specific status of these beetles based on male genitalia (Jeannel, 1924), and on other morphological characters (Juberthie *et al*, 1980a, 1981; Delay *et al*, 1983).

What is the congruence between biochemical and morphological data? There is no congruence between the 2 sets of characters at the species level. Relative degrees of molecular divergence between the 3 species do not concur with degrees of morphological differentiation between the same species. The highest genetic identity values are found among the species *S hydrophilus* and *S zophosinus*. This is in contrast to morphological and biometrical analysis in which these 2 species exhibit the greatest difference (table IV; figs 2 and 3). *S colluvii* and *S hydrophilus*, morphologically more similar, show greater biochemical separation (fig 2). What factors would account for the observed discrepancy in degrees of differentiation? The genetic differences between these species, at structural genes measured by electrophoresis of the gene products, are the results of mutations accumulated since the time of divergence. This does not imply that they are critical in evolutionary terms. Electrophoresis can measure reduced gene flow or absence of gene flow, but cannot say anything about changes at the regulatory level, which may be very important in some speciation events (Wilson *et al*, 1974). Application of electrophoretic methods to taxonomic studies requires caution in interpretation of experimental data. It is not easy to assign limits to the electrophoretic variability disclosed experimentally in relation to biological criteria used in differentiation between species. The processes of evolution of different aspects, represented by different data sets (molecular, morphological, morphometrical and behavioural),

may have been substantially uncoupled and proceeded independently or at different rates. For example, *S colluvii* appears to have evolved biochemically much more than it has changed morphotypically, in particular from *S hydrophilus*. The incongruence between extent of divergence in the 2 sets of characters could be the result of different sensitivities of each to different components of the environment. The 3 species studied here differ in their genetic structure (Crouau-Roy, 1989a, b) and in their ecological characteristics (Juberthie *et al* 1981; Crouau-Roy, 1986). The external morphometrical characteristics of *Speonomus*, traditionally used in systematic studies, could be under extensive selective pressures imposed by the local environment. Without heritability data, 1 suggestion is that the morphological variation has been more immediately shaped by ecological differences between the species. Further, an unknown proportion of the variation exhibited by many of these traits could be environmental rather than genetic in origin. The morphometrical data may reflect historical processes but are much more under the influence of differential selective pressures (micro - and macro-environmental influences) than the biochemical data.

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