# SYSTEMATICS OF THE FUSCUS GROUP OF THE FROG GENUS LEPTODACTYLUS (AMPHIBIA, LEPTODACTYLIDAE) 

By W. RONALD HEYER



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# SYSTEMATICS OF THE FUSCUS GROUP OF THE FROG GENUS LEPTODACTYLUS (AMPHIBIA, LEPTODACTYLIDAE) ${ }^{1}$ 

By W. Ronald Heyer ${ }^{2}$


#### Abstract

Thirteen characters of external morphology are analyzed in detail for the species comprising the fuscus group (genus Leptodactylus). The major method of data analysis is application of the multivariate stepwise discriminant function analysis. Results of the morphological analysis are compared with known information on mating calls, larvae, and karyotypes. Based on all available data, taxonomic conclusions are drawn. The nomenclature of the group is described in detail, associating proposed names with the species units recognized in this study. Wherever possible, the original type material was re-examined for this study. Of the 19 species recognized in the fuscus group, 4 are described as new. For each species, the following information is provided: a synonymy of primary names, a diagnosis for adults, adult and larval morphological characteristic summaries, diagnostic description of the mating call, diagnostic description of the karyotype, and distribution including localities and associated specimen museum numbers for the specimens examined. A key is provided at the end of the species accounts.

The composite range of the group is extensive, ranging from Texas to Argentina, on both sides of the Andes, and certain islands of the West Indies. Several characters used in the analysis are sexually dimorphic. It is postulated that sexual dimorphism in hind limb proportions is due to differential selection, the shorter male limb the result of selection for the burrowing activity of incubating chamber formation, the longer female limb the result of selection for avoiding above ground vertebrate predators. Sexual dimorphism occurring in the lip and thigh stripes of some species is explained by the hypothesis that males are using the information to discriminate among females in mate recognition. The ancestral stock of the fuscus group is presumed to have been fossorially adapted to an area with a vegetation type similar to that now found in the Gran Chaco. Evolutionary events within the species group correlate with adaptations to more mesic environments.


## INTRODUCTION

This study is the third in a series (Heyer 1970a, 1973) treating the systematics of the species groups of the Leptodactylus complex.
The aim of this study is to set a new baseline for the systematic understanding of the fuscus group based on museum specimens and field observations. The study is based on all available specimens, exclusive of five new species in the group that are being described by South American workers.

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The paper has benefitted from the constructive comments of the assigned reviewers. They are, as usual, not responsible for any flights into fantasy on my part.

The Smithsonian Research Foundation supported the research.

Museum abbreviations as used in the text are:

| AMNH | ican Museum of Natural History, New York |
| :---: | :---: |
|  | A. Schwartz private collection, Miami |
| BMNH | British Museum (Natural History), Lon |
| CAS-SU | California Academy of Sciences, Stanford University Collection |
| CHINM | Collección Herpetólogica del Instituto Nacional de Microbiología, Buenos Aires |
|  | Carnegie Museum, Pittsburgh |
| CRE | University of Southern Californi |
| MN | Field Museum of Natural Hist |
| IML | Fundación |
| KU | University of Kansas Museum of Natural History, Lawrence |
| LACM | Natural History Museum of Los Angeles County, Los Angeles |
| LES | J. Lescure private collection, Paris |
| MACN | Museo Argentino de Ciencias Naturales, Buenos Aires |
| MCZ | Museum of Comparative Zoology, Harvard University, Cambridge |
| MNRi | Museu Nacional, Rio de Janeiro |
| MZUSP | Museu de Zoologia, Universidade de São Paulo, São Paulo |
| RMN | Rijksmuseum van Natuurlijke Historie, Leiden |
| CWC | Texas Cooperative Wildlife Collection, Texas A\&M University, College Station |
| UMMZ | University of Michigan Museum of Zoology, Ann Arbor |
| UPR | University of Puerto Rico, Mayaguez |
| USNM | National Museum of Natural History, Washington, D. C. |
| UT | University of Texas at Arlington, Arlington |
| CAB | W. C. A. Bokermann private collection, São P |

## METHODS AND MATERIALS

The study represents several stages of analysis. Briefly, as many museum specimens as could be reasonably borrowed were initially analyzed with respect to extemal morphology. Other known biological information was added to the results of the morphological analyses. In some cases, information at that point was adequate to
draw systematic conclusions. In other cases, the data were inconclusive and additional field work and/or morphological data were gathered. After the first draft of this paper was completed, Izecksohn's description of a new species of Leptodactylus was published. As he had allowed me to examine the specimens, the data are included in the species accounts, but are not included in the population analysis section.
The following characters were recorded for every adult specimen examined.

1) Dorsal pattern. Standards were prepared for dorsal patterns and the specimens were placed in the category they most closely resembled (fig. 1).
2) Lip stripe. The lip was coded as either having a distinct light stripe or not. In some species, information was also recorded on the distinctiveness of a dark subocular bar.
3) Thigh stripe. The posterior face of the thigh was coded as having a distinct, indistinct, or no light stripe.
4) Dorsolateral folds. The total number of dorsolateral folds was recorded for each specimen.
5) Sex.

6-8) Tibia, tarsal, and foot texture. The relative presence or absence of white tubercles was recorded separately for the tibia, tarsus, and foot elements.
9) Snout-vent length (SVL). The SVL is the distance from the tip of the snout to behind the vent.

10-14) Head length, head width, femur length, tibia length, foot length ratios. Measurements were taken for each variable and divided by the SVL of the same animal. Head length was measured from behind the angle of the jaw to the tip of the snout. Head width was measured at the angle of the jaws. The leg measurements were taken with the leg positioned in a Z pattern with the femur element at right angles to the vertebral column. The foot was measured from behind the inner metatarsal tubercle to the tip of the third digit.

In addition, the tibia pattern was recorded for members of the L. gracilis complex (fig. 2).

All measurements were taken with vernier calipers. A series of 10 L. albilabris of diverse conditions of preservation were measured on two occasions to determine the repeatability of measurements. The average differences of measurements ranged from .2 to .4 mm ; measurements are repeatable within a tolerance of .5 mm . The actual error in measurement may be greater, particularly in SVL, femur, tibia, and foot length where the position of the animal in preservative may not allow the accurate measurement of the variable.

The above data were analyzed by the Stepwise Discriminant Analysis, BMDO7M, in the Biomed package produced by the University of California. Justification for using this multivariate approach to aid in distinguishing species in leptodactylid frogs, using the type of data analyzed herein, has been presented elsewhere (Heyer 1977). The number of dorsolateral folds was not used in the computer analysis because the condition could not
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Figure 1. Dorsal pattern standards utilized for the Leptodactylus fuscus species group.


Figure 2. Tibia pattern standards utilized for the Leptodactylus gracilis-complex. Left, barred condition; right, striped condition.
be determined in a number of poorly preserved individuals. Tibial texture was also omitted from all analyses except for $L$. labialis because of slight interspecific variation. The number of variables used differs slightly from group to group. The information on group size and number of variables analyzed is presented case by case in the next section. Some members of the study group are sexually dimorphic; the male and female data were run separately. For the female L. albilabris-complex data, standardized and non-standardized data were analyzed. The non-standardized data were simply the raw values punched on the computer cards. The data were standardized so that the total range of variation of each character fell between 0 and 1 . The discriminant function analysis results were exactly the same using the standardized and non-standardized data; the remaining analyses were run using non-standardized data.

Atchley, Gaskins, and Anderson (1976) presented theoretical ärguments against the use of ratios as variables in discriminant function analysis. In terms of the ratios used here, their argument is that dividing through by SVL does not entirely eliminate size as a factor in the variable involved. Atchley et. al. (1976) compared the results of analysis of original untransformed hypothetical data with the analysis of ratios and found striking differences. As the paper by Atchley et. al. appeared after my computer runs had been made, I tested their conclusions by reanalyzing data for four members of the mystaceus-complex, using the measurements as originally recorded.

Overall, the results of the two runs are very similar.

The posterior classifications are identical for the female data and differ by one specimen for the male data. The plots of the first two discriminant axes are essentially the same. The cumulative proportions of total dispersion accounted for by successive discriminant axes are nearly identical in both runs, in marked contrast to the runs of Atchley et. al. For example, for the female data using ratios, the cumulative proportion of dispersion of the first discriminant axis is .807 (.817 for data using measurements), .977 for the first and second axes (.978) and 1.00 for the first, second and third (1.00).

The only noticeable differences are in the entering order of the variables (Table 1). The F levels of significance cannot be interpreted literally because not all of the variables are normally distributed (see Heyer 1977, for discussion). However, the critical F-level (5\%) can be used at least to screen out variables that are not adding information to the analysis. Variables having a low F value are labelled as not important (NI) in the analysis section, indicating that they are probably not statistically significant contributors to inter-group discrimination in a particular run. However, rigorous statistical interpretation is not possible. The most striking difference in variable entering order is with SVL, but overall, the orders are similar.

Corruccini (1977), in response to Atchley et. al. (1976), found analysis of ratios to be meaningful for real data sets. As Atchley et. al.'s arguments are not substantiated by real data sets, ratios are used in the discriminant function analyses of this paper.

A discriminant function analysis requires pre-formed

Table 1
Entering order of variables for members of the $L$. mystaceus-complex.
Line indicates F significance at the $5 \%$ level (see text).

|  | Head and limb variables entered as ratios | Head and limb variables entered as measurements |
| :---: | :---: | :---: |
| Female data | .tarsal texture | tarsal texture |
|  | foot texture | head width |
|  | foot/SVL | foot length |
|  | SVL | foot texture |
|  | head length/SVL | head length |
|  | femur/SVL | femur length |
|  | head width/SVL | dorsal pattern |
|  | dorsal pattern | lip stripe |
|  | lip stripe | tibia length |
|  | tibia/SVL | SVL |
|  | thigh stripe | thigh stripe |
| Male data | .tarsal texture | tarsal texture |
|  | foot texture | foot texture |
|  | foot/SVL | foot length |
|  | dorsal pattern | SVL |
|  | tibia/S VL | dorsal pattern |
|  | lip stripe | tibia length |
|  | femur/SVL | lip stripe |
|  | head width/SVL | femur length |
|  | SVL | tibia length |
|  | head length/SVL | head width |
|  | thigh stripe | thigh stripe |

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entering of signifnot all of rer 1977, (5\%) can not addng a low ; analysis atistically nation in interpresrence in zrall, the
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groups for analysis. The groups used are what I believed to be species units based on my observations during the data taking phase. The discriminant function analysis is used to determine whether there are demonstrable morphological differences among the units analyzed. In ten years of experience working with frogs of the genus Leptodactylus, I have found that consistent morphological differences among populations is indicative of species level differentiation. For purposes of this paper, if the discriminant function analysis demonstrates that the species units are morphologically distinct, no further explanation is required. If the discriminant function analysis only partly separates the groups being analyzed, then other data where available are added to see if the additional data support the species groupings as originally determined.

The use of discrete variables in the discriminant function analysis places two restrictions on the results. First, the discriminatory power of the analysis is reduced. A two state character can only discriminate two groups, a continuous character can discriminate many groups. Second, the posterior classification of individuals involves confidence limits around the centroid values for the groups as analyzed. Discrete variables do not lend themselves to meaningful confidence limits. The results of the posterior classifications are thus not robust and should not be overinterpreted. The net result of the use of discrete variables is that the discriminant function analysis results are conservative. Any differences observed are real, but there may be more differences among groups than the results indicate.

The single most useful output of the discriminant function analysis as used herein is the plot of the first two discriminant axes. This gives a visual presentation of the distinctiveness of the groups being analyzed. It is this feature that is used to demonstrate the relative morphological distinctiveness of the groups being analyzed. The results are not used to test whether or not my original sorting into species was correct. The results are used to demonstrate the relative morphological distinctiveness of the groups. For the species represented by adequate geographic samples, discriminant function analyses are performed using locality samples as groups to determine whether any of the geographic samples are morphologically distinctive. These results are interpreted very conservatively. That is, a geographic sample would have to be clearly distinctive to warrant further analysis.

The criteria used to determine the species limits for members of the fuscus group in the order in which I have confidence in them follow.

1. Mating calls.-The mating calls of members of this group are species specific and the kinds of differences coding species specificity have been commented on (Straughan and Heyer 1976). Where mating call information is known, those data are considered of prime importance and take precedence over the other data uti-
lized in this study. Because mating calls are known for relatively few populations, the mating call data are used operationally in conjunction with the data of the second criterion.
2. External adult morphology.-Consistent, discrete - morphological differences among populations of members of the fuscus group usually correlate with the mating call data. In this study, the discriminant function analysis was applied in two different ways for which I have two levels of confidence.
A. Use of the multivariate analysis with the populations I consider to represent distinct species. This analysis is utilized to show the kinds of morphological differences among the species recognized herein. Morphological overlap can be extensive for species which are clearly distinct (figs. 25 and 26 for two species which have very distinctive mating calls and karyotypes). In some cases, data not coded further separate the species groupings, particularly information on dorsolateral folds. Because all the coded data are used in these analyses, the results are interpreted liberally. That is, species groupings are considered to be morphologically distinctive and distinguishable even with a moderate amount of overlap on the discriminant axis plots.
B. Use of the multivariate analysis with geographic samples of what I consider to be the same species. In all cases, some of the variables are uniform for the analyses; thus, the analyses are based upon smaller data sets. In addition, there are no other morphological data that were not coded that will allow further discrimination. For these reasons, the results of these analyses are interpreted very conservatively. Wherever the results of this analysis show a distinctive population unit that conflicts with the mating call information, the mating call information is given priority. Where mating calls are not available, the distinctive morphological units are pointed out, but not accorded specific level recognition. I do not have enough confidence in this level of analysis to recognize species levels based on the results. The value of the technique is to point out distinctive populations that should then be sampled for mating calls before a final taxonomic decision is made. If there are taxonomic errors in this paper, they involve recognition of too few, not too many species, in my opinion.
3. Larval morphology and karyotypes.-Information from these systems is not useful in determining species limits for members of the fuscus species group. Too few larval samples are available to determine whether apparent differences in denticle number has systematic value. The general shapes and color patterns of all known larvae are similar. The known karyotypes for members of this group are very similar, with but a single exception. The exception is the karyotype of L. latinasus which is interpreted as indicating a species level difference. All other kinds of karyotypic differences reported
are as likely due to differences of preparation or interpretation as to differences of systematic value (Heyer and Diment 1974).

Within the fuscus group, a number of species complexes are apparent. The following complexes are recognized for purposes of discriminant function anaylses: albilabris, labialis, fuscus, bufonius, latinasus.

## POPULATION ANALYSES

The coding of characters for computer analysis results in a loss of information in some cases. For character 1, dorsal pattern, two different codes were used. For L. labialis, the presence of a double dorsal chevron (fig. 1, A) was coded as a 2 , any other pattern was coded as a 1 . For the other species, the presence of a light middorsal stripe was coded as a 2 , absence was coded as a 1 . For the only analysis in which $L$. labialis is analyzed with another species group (latinasus), the dorsal pattern is omitted from analysis. Character 2, lip stripe, was uniformly coded as 1 for an indistinct light lip stripe, 2 for a distinct lip stripe. Character 3, thigh stripe, was uniformly coded as 1 for a distinct light stripe, 2 for an indistinct, but still discernable stripe, 3 for no stripes. Characters 6 to 8 , textures of the tibia, tarsus, and sole of foot were uniformly coded as 1 for presence of any white tubercles, 2 for no white tubercles. The actual numbers for the SVL, head, and hind limb measurements were punched on cards; the head and hind limb measurements were each divided through by SVL and a new card deck punched by computer.

## L. ALBILABRIS—COMPLEX

Morphology. - Members of the L. albilabris complex are distributed on the West Indian islands. Morphologically the group is distinct from all mainland species populations. Most taxonomic questions concerning the L. albilabris complex center on the question whether the different island bank systems have different species. The following variables were used in the stepwise discriminant function analysis: 1-3, 9-14. Characters 7-8 are uniform in L. albilabris.

Female data.-Seventy-two individuals were analyzed from five localities in Puerto Rico, two localities from the Dominican Republic and one locality each from St. Croix, St. Thomas, and Tortola. The smallest sample used consisted of three individuals from a single locality; the largest contained 16 individuals. The results (fig. 3) indicate that the Dominican Republic samples are the most distinctive, but that there is overlap with the other samples. Overlap, as used throughout, means overlap of the polygons on the plot figures of the first two discriminant axes. The first two axes account for $68 \%$ of the total variation. The variables were entered in the program in the following order (i.e. in order of descending contribution to the intergroup variation): dorsal pat-
tern, SVL, head width ratio, tibia ratio, thigh stripe, head length ratio, foot ratio (NI), femur ratio (ND), and lip stripe (NI).

Male data.—One hundred thirty five individuals were analyzed from 7 localities in Puerto Rico and one locality each from the Dominican Republic, St. Croix, St. John's, St. Thomas, and Tortola. Four individuals from a single locality was the smallest group used, the largest was comprised of 24 individuals. The results (fig. 4) indicate that as with the females, the Dominican Republic samples are the most distinctive, but there is morphological overlap with the other samples. The first two axes account for $73 \%$ of the total variation. The variables entered in the program in the following order: dorsal pattern, tibia ratio, SVL, head width ratio, head length ratio, femur ratio, thigh stripe (NI), lip stripe (NI), foot ratio (NI).

The results of the male and female analyses both indicate that the Dominican Republic samples are the most distinctive. There is sexual dimorphism in patterns of geographic variation, as some of the variables entered the program in different orders. Part of this may be due to the fact that different numbers of localities were used for the two sexes, and only 4 localities were represented in common in the two samples.

Larvae.-Tadpole samples were examined from Puerto Rico (ASFS 7901, UMMZ 125168, 125174), St. Thomas (USNM 119038) and the Dominican Republic (USNM field 41052). All larvae examined are indistinguishable.

Mating calls.-Two calls were available for analysis: Puerto Rico: El Yunque (AMNH tape) and Dominican Republic: El Seibo Prov; 3.2 km E Sabana de la Mar (USNM tape). The calls sound similar to the human ear, but representative calls analyzed in detail show some differences. Sonagrams (fig. 5) indicate the calls have the same frequency and basic structure. The pattern of frequency modulation differs between the two calls (fig. 5). The strip chart records of individual calls (fig. 6) indicate that the initial part of the calls differ, as well as the shape of the initial part of the second portion of the call. These differences are of the kind that code spe-cies-specific information in Leptodactylus (Straughan and Heyer 1976), but the magnitudes of the differences (figs. 5 and 6) are not great.

No information is available on call variation within island populations or among individuals in a given population. While the calls available for analysis differ, the evidence for specific differentiation is not decisive.

Taxonomic conclusion.-The adult morphology and calls (sample size of only 2 ) are different for the populations from the Dominican Republic with respect to all other populations. The evidence indicates that all West Indian populations had a common ancestor: the question revolves about the degree of differentiation. I interpret the available evidence to indicate the degree of differentiation has not reached the species level.

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Figure 3. Discriminant axis plot for geographic samples of females of Leptodactylus albilabris. A-E = Puerto Rico, $3=$ St. Croix, $5=$ St. Thomas, $6=$ Tortola, $7-8=$ Dominican Republic. Letters and numbers placed at group means. Envelopes contain all group members by islands.


Figure 4. Discriminant axis plot for geographic samples of males of Leptodactylus albilabris. $\mathrm{A}-\mathrm{G}=$ Puerto Rico, $1=$ St. Croix, $2=$ St. Johns, $3=$ St. Thomas, $4=$ Tortola, $5=$ Dominican Republic. Letters and numbers placed at group means. Envelopes contain all group members by islands.



Figure 6. Strip chart records of the mating call of Leptodactylus albilabris. Line equals 0.01 s . Upper figure is note of specimen from Puerto Rico, El Yunque, lower is note of specimen from Dominican Republic, Sabana de la Mar. See legend of Figure 5 for further specimen data.

## LEPTODACTYLUS LABIALIS

Morphology.-Groupings used in the computer analysis consist of specimens from single localities unless otherwise indicated. The following variables were used: $1-3,6,9-14$. Variables 7 and 8 are uniform for $L$. labialis.

Female data.-Specimens from localities in the following political units were analyzed as follows (number of specimens in parentheses): Mexico, Campeche (47), Mexico, Michoacán (4), Mexico, Oaxaca (10), Mexico, San Luis Potosí (7), Mexico, Tamaulipas (6), Mexico, Veracruz (5), Mexico, Yucatán (4), Guatemala (3), Belize (36), Honduras, Francisco Morazán (10), Honduras (8), Costa Rica (5), Panama (4), Colombia (4), Venezuela, Apure (21), Venezuela (5). The plot of the first two discriminant axes (fig. 7) shows a complex pattem, mostly of overlapping groups. The first two axes account for $61 \%$ of the variation. The variables entered in the following order: SVL, tibia ratio, tibia texture, head
width ratio, thigh stripe, foot ratio, femur ratio, lip stripe, head length ratio, dorsal pattern (NI). The northernmost Michoacán sample is the only group showing no overlap with other groups. The Costa Rican sample is also relatively distinctive. All other samples show broad overlap; generally, samples from adjacent localities are close to each other in the discriminant axis plot (fig. 7).

Male data.-Specimens from localities in the following political units were analyzed as follows (number of specimens in parentheses): Texas ( 3 from 2 localities), Mexico, Campeche (11), Mexico, Colima (7), Mexico, Guerrero (7), Mexico, Michoacán (6), Mexico, Morelos (3), Mexico, Tamaulipas (5), Mexico, Tamaulipas (6), Mexico, Yucatán (8), Guatemala (15), Belize (5), Honduras (7), Costa Rica, Guanacaste (6), Costa Rica, Puntarenas (5), Panama, Canal Zone (11), Panama, Coclé (7), Panama, Veraguas (8), Colombia, Antioquia (5), Colombia, Santander (5), Venezuela (9). The plot of the


Figure 7. Discriminant axis plot for geographic samples of females of Leptodactylus labialis. $1-7=$ Mexico, $\mathrm{G}=$ Guatemala, $\mathrm{B}=$ Belize, $\mathrm{H}-\mathrm{I}=$ Honduras, $\mathrm{R}=$ Costa Rica, $\mathrm{P}=$ Panama, $\mathrm{C}=$ Colombia, $\mathrm{V}-\mathrm{W}=$ Venezuela. Numbers and letters are placed at group means. Envelopes contain all group members. intergroup variation, and (2) the northernmost populations from west coastal Mexico are the most distinctive based on external morphology.

Larvae.-Larvae have previously been described for L. labialis (e.g. Heyer 1970b). During that previous study, I found no differences between larval samples from Mexico and Middle America. To my knowledge,
no larval samples are available from any South American localities.

Mating call.-Straughan and Heyer (1976) summarized the call information for labialis, indicating a clinal trend in call characteristics from Mexico to Panama. The differences are not of the magnitude demonstrated by different species of Leptodactylus. No calls were available for any South American populations.

Taxonomic conclusion.-The discriminant function analysis indicates that the northwest coast Mexico population is morphologically distinguishable from all other groups. The mating call information indicates that the call of the northwest coast Mexican population is not specifically distinct from the Panamanian population call. In this case, I place more confidence in the mating call data and conclude that differentiation has not reached the species level.

## Leptodactylus fuscus - COMPlex

Computer analysis of the morphological data was done in two stages. The first analysis is based on data from museum specimens assembled in the laboratory.

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Figure 8. Discriminant axis plot for geographic samples of males of Leptodactylus labialis. $1=$ Texas, $2-9=$ Mexico, $\mathrm{G}=$ Guatemala, $\mathrm{B}=$ Belize, $\mathrm{H}=$ Honduras, $\mathrm{R}-\mathrm{S}=$ Costa Rica, $\mathrm{N}-\mathrm{P}=$ Panama, $\mathrm{C}-\mathrm{D}=$ Colombia, $\mathrm{V}=$ Venezuela. Numbers and letters are placed at group means. Envelopes contain all group members.

## 1 Ameri-

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function ico popall other that the n is not pulation : mating reached oratory.

In some cases, sample sizes were small and attempts were made to gather more data on specimens located in South American museums.
Morphology.-As discussed earlier, specimens were sorted into what appeared to be different species. The first analytic procedure was to enter each of these species units as predefined groups to determine the relative morphological distinctiveness of each of the groups. The following variables were used: 1-3, 7-14.
Female data.-the following groups were analyzed (number of specimens in parentheses): fuscus (178), barred gracilis (referring to tibial pattern) (10), striped gracilis (6), longirostris (following Rivero's (1971) identification) (15), northern mystaceus (76), southem mystaceus (12), coastal Brasil mystaceus (3), poecilochilus (83). The results (fig. 9), indicate good separation of some groups, but considerable overlap in others. Posterior classification of cases into group results are discussed below with the male data. The first two axes account for $82 \%$ of the variation. The variables entered in the following order: foot texture, tibia ratio, foot ratio, tarsal texture, head width ratio, SVL, dorsal pattern, lip
stripe, head length ratio, thigh stripe, and femur ratio (NI).

Male data.-The groups analyzed were (number of specimens in parentheses): fuscus (214), barred gracilis (21), striped gracilis (18), longirostris (34), northern mystaceus (75), southern mystaceus (15), coastal Brasil mystaceus (3), poecilochilus (50). The results (fig. 10) are comparable to the female results. Seventy seven percent of the variation is accounted for in the first two axes. The variables entered in the following order: foot texture, tarsal texture, foot ratio, tibia ratio, SVL, dorsal pattern, head length ratio, lip stripe, head width ratio, thigh stripe, and femur ratio (NI).

The results of the a posteriori classification routine which assigns cases to their "most probable"' groups are similar for males and females (Table 2). As indicated previously, because discrete variables were used, the results of the posterior classification should not be interpreted too finely. The results indicate that separation of the groups is good. As more specimens of fuscus were placed in other groups than any other species unit, the fuscus unit is discussed as an example to show that other


Figure 9. Discriminant axis plot of females of the fuscus complex. $\mathrm{F}=$ fuscus, $\mathrm{A}=$ striped gracilis, $\mathrm{B}=$ barred gracilis, $\mathrm{L}=$ longirostris, $1=$ northern mystaceus, $2=$ southern mystaceus, $3=$ south coast mystaceus, $\mathrm{P}=$ poecilochlus. Letters and numbers placed at group means. Envelopes contain all group members.
evidence can be used to further separate the analytic units. There are two reasons why several fuscus specimens were assigned to other groups: (1) the variables analyzed are not sufficient in themselves to completely separate the fuscus specimens from specimens of the other groups, and (2) the foot texture coding is very dependent on state of preservation in this group. As noted above, foot texture was the most important distinguishing factor in the analysis for both males and females. In most of the other species, white tubercles are prominent and obviously present or conspicuously absent. In fuscus, however, the tubercles are at best small, are often the same color as the rest of the foot, and therefore not conspicuous. All fuscus probably have a tubercular foot texture, but the texture is often lost in preservation. All fuscus specimens classified as northern and southern mystaceus were coded as having foot tubercles present. Only 4 additional specimens that were coded as having foot tubercles were computer assigned to fuscus. Because of geographic ranges, some of the computer assignments are improbable, for example, some
fuscus specimens from Argentina were assigned to poecilochilus (found in Middle America and northern South America). Improbable assignments account for $59 \%$ of the wrong assignments. As stated earlier, the information on dorsolateral folds was not included in the computer analysis because the information was missing from several specimens due to preservation. Leptodactylus fuscus specimens always have 6 dorsolateral folds, mystaceus specimens always have 4, and only longirostris and poecilochilus specimens with a light mid-dorsal stripe have 6 dorsolateral folds. When the original data were checked on the fuscus specimens assigned to other groups by the computer, the dorsolateral fold information resolved $77 \%$ of the cases where the computer assignments were geographically possible. Thus, out of the 129 cases in which the computer assigned fuscus specimens to other groups, the additional information concerning geographic improbability and state of dorsolateral folds resolved all but 14 cases.

Additional data were gathered for the mystaceus and gracilis complexes from South American museums.

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reus and ums.

Figure 10. Discriminant axis plot for males of the fuscus complex. $\mathrm{F}=$ fuscus, $\mathrm{A}=$ striped gracilis, $\mathrm{B}=$ barred gracilis, $\mathrm{L}=$ longirostris, $1=$ northern mystaceus, $2=$ southern mystaceus, $3=$ south coast mystaceus, $\mathrm{P}=$ poecilochilus. Letters and numbers placed at group means. Envelopes contain all group members.

Table 2
Posterior classification of members of the fuscus complex.

| MALES |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of cases classified into group |  |  |  |  |  |  |  |  |
| Group |  |  |  |  |  |  |  |  |
|  | A | B | C | D | E | F | G | H |
| A-fuscus | 153 | 0 | 1 | 17 | 9 | 6 | 5 | 23 |
| B-striped gracilis | 0 | 20 | 1 | 0 | 0 | 0 | 0 | 0 |
| C-barred gracilis | 0 | 2 | 16 | 0 | 0 | 0 | 0 | 0 |
| D-longirostris | 0 | 0 | 0 | 33 | 0 | 0 | 1 | 0 |
| E- northern mystaceus | 0 | 0 | 0 | 0 | 75 | 0 | 0 | 0 |
| F - southern mystaceus | 0 | 0 | 0 | 0 | 1 | 14 | 0 | 0 |
| G-coastal mystaceus | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 |
| H-poecilochilus | 6 | 0 | 0 | 0 | 4 | 1 | 0 | 39 |
| FEMALES |  |  |  |  |  |  |  |  |
| Number of cases classified into group |  |  |  |  |  |  |  |  |
| Group |  |  |  |  |  |  |  |  |
|  | A | B | C | D | E | F | G | H |
| A-fuscus | 110 | 1 | 0 | 31 | 9 | 4 | 3 | 20 |
| B-striped gracilis | 0 | 8 | 2 | 0 | 0 | 0 | 0 | 0 |
| C-barred gracilis | 0 | 0 | 6 | 0 | 0 | 0 | 0 | 0 |
| D-longirostris | 3 | 1 | 0 | 11 | 0 | 0 | 0 | 0 |
| E-northern mystaceus | 0 | 0 | 0 | 0 | 75 | 1 | 0 | 0 |
| F-southern mystaceus | 0 | 0 | 0 | 0 | 4 | 8 | 0 | 0 |
| G-coastal mystaceus | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 |
| H -poecilochilus | 2 | 0 | 0 | 2 | 4 | 0 | 0 | 75 |

As more specimens were examined from coastal Brasil, it became evident that two taxa were present. The discriminant function analyses were performed to determine the morphological distinctiveness of these two species from the previously determined species, northern and southern mystaceus.

Female mystaceus-complex data.-The following groups were analyzed (number of specimens in parentheses): south coast mystaceus (9), east coast mystaceus (14), southern mystaceus (11), northern mystaceus (76). The results (fig. 11) show good separation of the groups. The first two axes account for $98 \%$ of the total dispersion. The variables entered in the following order: tarsal texture, foot texture, foot ratio, SVL, head length ratio, femur ratio, head width ratio (NI), dorsal pattern (NI), lip stripe (NI), tibia ratio (NI), thigh stripe (NI). All south coast mystaceus were classified posteriorly as south coast mystaceus, 1 east coast mystaceus was assigned to southern mystaceus, 1 southern mystaceus was
assigned to east coast mystaceus and 1 southern mystaceus was assigned to northern mystaceus, 3 northem mystaceus were assigned to south coast mystaceus and 1 northern mystaceus was assigned to east coast mystaceus.

Male mystaceus-complex data.-The following groups were analyzed (number of specimens in parentheses): south coast mystaceus (9), east coast mystaceus (24), southern mystaceus (32), northern mystaceus (72). The results (fig. 12) show reasonably good separation of groups. The first two axes account for $98 \%$ of the total dispersion. The variables entered in the following order: tarsal texture, foot ratio, foot texture, head length ratio, tibia ratio, dorsal pattern, SVL (NI), femur ratio (NI), head width ratio (NI), thigh stripe (NI), lip stripe (NI). Two of the nine south coast mystaceus were posterionly classified as northern mystaceus, 1 east coast mystaceus was assigned to south coast mystaceus and 3 east coast mystaceus were assigned to southern mystaceus, 5


Figure 11. Discriminant axis plot for females of the mystaceus complex. $\mathrm{E}=$ east coast mystaceus, $\mathrm{N}=$ northern mystaceus, S $=$ south coast mystaceus, $\mathrm{W}=$ southern mystaceus. Letters placed at group means. Envelopes contain all group members.

Figure
dthern mys3 northem staceus and east coast
wing groups arentheses): aceus (24), $s$ (72). The paration of of the total iwing order: ength ratio, : ratio (NI), stripe (NI). : posteriorly it mystaceus 3 east coast งstaceus, 5


FIGURE 12. Discriminant axis plot for males of the mystaceus complex. $\mathrm{E}=$ east coast mystaceus, $\mathrm{N}=$ northern mystaceus, $\mathrm{S}=$ south coast mystaceus, $\mathrm{W}=$ southern mystaceus. Letters placed at group means. Envelopes contain all group members.
southern mystaceus were assigned to east coast mystaceus and 1 southern mystaceus was assigned to northern mystaceus, all northern mystaceus were assigned to northem mystaceus.

The male and female mystaceus-complex results are comparable. The east coast form is nearly always morphologically distinguishable from the south coast form and the northem form is nearly always morphologically distinguishable from the southern form. In a few cases the forms are not morphologically distinguishable by the discriminant function analysis, but these mostly involve completely allopatric species pairs.

The discriminant function analysis run on the larger sample sizes of barred and striped gracilis is similar to the analysis run on a smaller data set. For females, 22 of 23 individuals of striped gracilis were posteriorly classified as striped gracilis, 10 of 11 barred gracilis were classified as such. For males, 34 of 37 striped gracilis were classified as striped gracilis, 23 of 24 barred gracilis were classified as barred gracilis. The entering order of variables differs between males and females. The first three variables entered for females are head width ratio, foot ratio, and thigh stripe; the first three variables entered for males are SVL, foot ratio, and head width ratio.

The overall analysis indicates that the predetermined species units are generally separable on the basis of the morphological characters used. In some cases, additional information such as dorsolateral folds is required to make the proper species assignment.

Larvae.-Tadpoles are available or have been described for the following taxa from this complex: L. fuscus (Lescure 1972), striped gracilis (Fernandez and Fernandez 1921), northern mystaceus (see species accounts) and poecilochilus (Heyer 1970b). All larvae have similar shapes and patterns. Based on limited material, the number of denticles in the split tooth row anterior to the beak appears diagnostic at the species level, but the available larval data are not adequate to add any information to a species level discrimination analysis.

Mating calls.-Calls are known for L. fuscus (discussed after the following intraspecific morphological variation section), striped gracilis, longirostris, northem and southern mystaceus, and poecilochilus.
Rivero (1971) demonstrated the distinctiveness of calls of L. fuscus, longirostris, and poecilochilus in Venezuela. Straughan and Heyer (1976) indicated that the differences between calls of specimens from northern mystaceus and southern mystaceus populations (as used here) are indicative of species differentiation.

Barrio (1973) described the calls of $L$. gracilis and geminus (also see species accounts). Both species are morphologically striped gracilis (see nomenclature section under $L$. geminus for further discussion). W. C. A. Bokermann kindly gave me a copy of a recording of barred gracilis. The sonagram of this recording is visually distinctive from the sonagrams of the striped gracilis calls that Barrio published, confirming the distinctiveness at the species level of barred and striped gracilis (Bokermann and Sazima are describing the call of barred gracilis).

Taxonomic conclusions.-Mating calls are known for all but two members of this complex: east coast mystaceus and south coast mystaceus. All of the known calls are distinct, supporting the species level of differentiation hypothesized for these units. The east coast mystaceus and south coast mystaceus units are as morphologically distinctive as the other species in this complex and are considered to be specifically distinct.

Enough morphological data are available to study intragroup variation in the following: L. fuscus, northem mystaceus, and poecilochilus.

## Leptodactylus fuscus

Morphology. -The groups used for analysis of variation consist of samples of individuals from single localities except for Panama. The variables used in analysis were: 1-3, 7-14.
Female data.-Groups composed of individuals from single localities were analyzed from the following political areas (numbers of specimens of each group in parentheses): Panama ( 3 individuals from 3 localities), Colombia (3), (10), (4), Guyana (6), (3), (6), (7), (3), (11), Surinam (3), (9), (5), (8), (9), French Guiana (5), Tobago (5), Trinidad (3), Bolivia (4), (4), (9), Brasil (3), (4), Argentina (5). The first two discriminant axes account for $69 \%$ of the variation (fig. 13). The variables entered in the following order: SVL, tibia ratio, head width ratio, foot ratio, lip stripe, head length ratio, tarsal texture (NI), dorsal pattern (NI), foot texture (NI), thigh stripe (NI), femur ratio (NI). The plot of the first two discriminant axes (fig. 13) demonstrates a complex pattern of variation, with pronounced overlap of groups. The most distinctive groups are mostly at the edges of the geographic range; Panama, Colombia, Tobago, and

FIGURE 13. Discriminant axis plot for geographic samples of females of Leptodactylus fuscus. $\mathrm{P}=$ Panama, $\mathrm{C}-\mathrm{E}=$ Colombia $1-6=$ Guyana, $\mathrm{H}-\mathrm{L}=$ Surinam, $\mathrm{F}=$ French Guiana, $\mathrm{T}=$ Tobago, $\mathrm{R}=$ Trinidad, $\mathrm{M}-\mathrm{O}=$ Bolivia, $8-9=$ Brasil, $\mathrm{A}=$ Argentina . Letters and numbers placed at group means. Envelopes contain all group members.
rsis of varia single losed in anal-
'iduals from llowing po:h group in localities), j), (7), (3), Guiana (5), (9), Brasil ninant axes re variables ratio, head ratio, tarsal (NI), thigh te first two mmplex patof groups. ae edges of 'obago, and
= Colombia, = Argentina.

Argentina. The single distinctive group from Surinam, based on three small females, is not at the edge of the geographic range.
Male data.-Groups composed of individuals from single localities were analyzed from the following political areas (numbers of specimens of each group in parentheses): Colombia (5), (4), (7), Venezuela (3), Guyana (3), (5), (3), (9), Surinam (11), (5), (3), French Guiana (13), Tobago (9), Trinidad (5), Bolivia (14), (5), (5), Brasil (3), (5), (3), (3), (4), Argentina (10), (8). The first two discriminant axes account for $72 \%$ of the variation (fig. 14). The variables entered in the following order: SVL, tibia ratio, dorsal pattern, head width ratio, head length ratio, foot texture, femur ratio, thigh stripe, lip stripe, foot ratio, tarsal texture (all variables important). One group is very distinct and 4 other groups are moderately distinctive (fig, 14). All other groups overlap in a complex manner. The single distinctive group is from a geographically extreme population in an interandean valley in Colombia and is composed of quite large individuals. The other two groups analyzed from Colombia are moderately distinctive (fig. 14), as is another geographically extreme population from Argentina. A moderately distinct group of three individuals from Brasil: Bahia is not geographically extreme.

The combined male and female results are similar in the following points: (1) The two factors which account for the most intergroup variation are SVL and tibia ratio; (2) The most distinctive populations are from the periphery of the geographic range, Panama and Colombia in the north, Argentina in the south. As these peripheral populations are the only ones that are distinctive in both male and female analyses, the populations that are distinctive in individual analyses may well be due to sampling error, as both cases involved but three specimens.

Larvae.-Geographic samples of larvae are not available. Comparisons of literature descriptions indicate no apparent differences (Kenny 1969, for Trinidad, Lescure 1972, for French Guiana),

Mating Calls.-Calls are available from a few localities throughout the geographic range. Comparison of the sonagrams (fig. 15) and strip chart records (fig. 16) with the published analyses of Lescure (1972) for French Guiana and Rivero (1971) for coastal Venezuela indicate that all calls are similar.

Taxonomic conclusion.-The morphological evidence indicates differentiation of the geographically peripheral northern and southern populations. The mating call of the southern population is not distinctive from calls throughout the range. On this basis, the conservative approach of recognizing but a single species is taken.


Figure 14. Discriminant axis plot for geographic samples of males of Leptodactylus fuscus. $\mathrm{C}-\mathrm{E}=\mathrm{Colombia}, \mathrm{V}=\mathrm{Venezuela}$, $1-4=$ Guyana, $J-L=$ Surinam, $F=$ French Guiana, $T=$ Tobago, $R=$ Trinidad, $M-O=$ Bolivia, $5-9=$ Brasil, $A-B=$ Argentina. Letters and numbers placed at group means. Envelopes contain all group members.
-f $A$

Figure 15. Sonagrams of representative mating calls of Leptodactylus fuscus, narrow band filter. Vertical scale marks at 1000 hz intervals. Horizontal scale mark at $1 \mathrm{~s} . \mathrm{A}=$ specimen from Colombia, nr. Villavicencio, air temperature $23.5^{\circ} \mathrm{C}$ (UTA tape); B $=$ specimen from Brasil, Manaus (USNM tape and specimen number 202506); C = Bolivia (AMNH tape recorded by William P McLean III); $\mathrm{D}=$ specimen from Argentina, Embarcación (LACM tape and specimen field number WRH 1399).


Figure 16. Strip chart records of the mating call of Leptodactylus fuscus. Line equals 0.01 s . Upper to lower figures are representative calls for specimens from Colombia, Brasil, Bolivia, and Argentina respectively. See legend of Figure 15 for further specimen data except specimen from Argentina is LACM field number WRH 1363 recorded at $21.3^{\circ} \mathrm{C}$ air temperature.

## Northern mystaceus

Morphology.-The groups used for analysis of variation consist of individuals from single localities. The variables used in the analysis were: $1-3,7-14$. Variables $1,7,8$ were constant for both the male and female data and do not appear in any of the analyses.

Female data.-Individuals (number in parentheses) from localities from the following areas were used as groups for analysis: Colombia (4), Guyana (5), (4), French Guiana (3), (6), Ecuador (19), Bolivia (9). The first two discriminant axes account for $81 \%$ of the variation (fig. 17). The variables entered in the following order: tibia ratio, SVL, femur ratio, head length ratio (NI), thigh stripe (NI), foot ratio (NI), lip stripe (NI), head width ratio (NI). The discriminant axis plot (fig. 17), demonstrates overlap of groups with no group distinct from any other group.

Male data.-Individuals (number in parentheses) from localities in the following areas were used as groups for analysis: Colombia (4), (3), Guyana (12), French Guiana (5), Ecuador (21). The variables entered in the stepwise discriminant function program in the following order: femur ratio, SVL, head length ratio, tibia ratio, head width ratio, lip stripe (NI), foot ratio (NI), thigh stripe (NI). The first two discriminant axes (fig. 18) account for $80 \%$ of the variation. The discriminant axis plot (fig. 18) shows extensive group overlap with the single exception of a group of three males from Vaupés, Colombia. The second Colombian sample, from Caquetá, which
borders Vaupés, is well within the variation of the other samples analyzed.

The male and female data differ in the importance of variables describing the patterns of variation. This may be due to the different number of groups analyzed in each data set. Both data sets agree in that most geographic samples overlap each other with respect to morphological variation. The single exception is the sample from the state of Vaupés, Colombia. No females were available from this locality for analysis. The distinctiveness of the Vaupés sample may be due to the small sample size.

Larvae.-Tadpoles are known only from Ecuador (see species account for description).
Mating call.-Mating calls from Colombia and Ecuador are similar (Straughan and Heyer 1976).
Taxonomic conclusion.-The available evidence indicates a single species is involved.

## Leptodactylus poecilochilus

Morphology.-The groups used for analysis of variation consisted of individuals from single localities. The variables used were $1-3,7-14$. Variables 2 and 7 were constant for both female and male data sets; variable 8 was constant for the male data set.
Female data.-Individuals (number in parentheses) from localities in the following areas were used for analysis: Costa Rica (9), (4), Panama (3), (4), (4), Colombia (4), (14), (20). Variables entered in the stepwise dis-


Figure 17. Discriminant axis plot for geographic samples of females of northern mystaceus. $\mathrm{C}=$ Colombia, $\mathrm{G}, \mathrm{Y}=$ Guyana, $\mathrm{F}, \mathrm{R}$ $=$ French Guiana, $\mathrm{E}=$ Eclador, $\mathrm{B}=$ Bolivia. Letters placed at group means. Envelopes contain all group members.


FIgure 18. Discriminant axis plot for geographic samples of males of northern mystaceus. $\mathrm{C}-\mathrm{D}=$ Colombia, $\mathrm{G}=$ Guyana, $\mathrm{F}=$ French Guiana, E = Ecuador. Letters placed at group means. Envelopes contain all group members.
criminant function analysis in the following order: SVL, dorsal pattern, head width ratio, tibia ratio, femur ratio, head length ratio, thigh stripe, foot texture (NI), foot ratio (NI). The first two discriminant axes account for $78 \%$ of the variation (fig. 19). The discriminant axis plot (fig. 19) shows overlap of all groups with the exception of the sample from Córdoba, Colombia.

Male data.-Individuals (number in parentheses) from localities in the following areas were used for analysis: Costa Rica (5), Colombia (5), (7), (4). Variables entered in the stepwise discriminant analysis in the following order: SVL, head length ratio, thigh stripe, head width ratio, tibia ratio, dorsal pattern, femur ratio (NI), foot ratio (NI). The first two discriminent axes account for $99 \%$ of the variation. The discriminant axis plot (fig. 20), shows overlap of the two samples from Antioquia, Colombia, and distinctive samples from Costa Rica and Córdoba, Colombia.

As there are few samples available for analysis, especially of males, the above results should be treated cautiously. Both data sets indicate the distinctiveness of
the population from Córdoba, Colombia, which is a geographically peripheral population in terms of the specimens available for the present analysis.
Larvae.-Tadpoles have been previously described (Heyer 1970b) based on Middle American samples. No samples are available from South America.

Mating call.—Fouquette (1960) described the call for specimens from Panama and Rivero (1971) described the call for specimens from Venezuela. The sonagrams figured by these two authors are very different and likely represent two distinct species. At present, no calls are available from Colombia to determine whether there is a cline in call characteristics. Neither author indicated whether voucher specimens were kept for the recordings; it is therefore possible the species identifications used by Fouquette (1960) and Rivero (1971) differ from mine. Until such time as the significance of the observed call differences is resolved, I assume the reported call for L. poecilochilus in Venezuela to refer to a different species than that indicated as poecilochilus in this study.
Taxonomic conclusion.-Clearly, more information


Figure 19. Discriminant axis plot for geographic samples of females of Leptodactylus poecilochilus. $\mathrm{R}-\mathrm{S}=$ Costa Rica, $\mathrm{N}-\mathrm{P}=$ Panama, A-C $=$ Colombia. Letters placed at group means. Envelopes contain all group members.
bufonius are analyzed. The variables entered in the following order: foot texture, SVL, femur (NI), tibia ratio is required to understand the apparent variation in morphology and mating call. For the present, the conservative approach of recognizing a single species is taken until further field work clarifies the situation.

## Leptodactylus bufonius-COMPLEX

Morphology.-The species recognized during the data gathering procedure were used as predetermined groups for the discriminant function analysis. Additional data were gathered for some group members from South American museums after the first discriminant function analysis was completed. The variables used in the computer analysis for the first set of available data were 1 -$3,7-14$. Variable 1 does not appear in the stepwise discriminant function results as it is uniform throughout the group. Likewise, variable 2 does not appear in the female results.

Female data.-The following groups were analyzed (number of specimens in parentheses): bufonius (34), läbrosus (23), mystacinus (13), ventrimaculatus (16). The discriminant axis plot results (fig. 21) indicate good
separation of the predetermined species groupings. The first two axes account for $97 \%$ of the variation. The variables entered in the following order: foot ratio, foot texture, tibia ratio, head length ratio, SVL, tarsal texture, femur ratio (NI), head width ratio (NI), and thigh pattern (NI).
Male data.--The following groups were analyzed: bufonius (53), northern bufonius (4), labrosus (9), mystacinus (44), ventrimaculatus (22). The discriminant axis plot results (fig. 22) also indicate good separation of the species groupings. The first two axes account for $92 \%$ of the variation. The variables entered in the following order: foot texture, lip stripe, foot ratio, head length ratio, head width ratio, tarsal texture, SVL (NI), femur ratio (NI), thigh stripe (NI), and tibia ratio (NI).
The results of the posterior classification into groups for female and male data (Table 3) also indicate that the species are morphologically distinguishable.
Data were taken on more northern and southern bufonius from specimens in South American museums to determine whether the initial separation based on very few northern bufonius specimens was substantiated.

For females, 54 southem bufonius and 15 northern


Figure 20. Discriminant axis plot for geographic samples of males of Leptodactylus poecilochilus. $\mathrm{R}=$ Costa Rica, $\mathrm{A}-\mathrm{C}=$ Colombia. Letters placed at group means. Envelopes contain all group members.


Figure 21. Discriminant axis plot of females of the bufonius complex. $\mathrm{B}=$ bufonius, $\mathrm{L}=$ labrosus, $\mathrm{M}=$ mystacinus, $\mathrm{V}=$ ventrimaculatus. Letters placed at group means. Envelopes contain all group members.


Figure 22. Discriminant axis plot of males of the bufonius complex. $\mathrm{B}=$ bufonius, $\mathrm{C}=$ northern bufonius, $\mathrm{L}=$ labrosus, $\mathrm{M}=$ mystacinus, $V=$ ventrimaculatus. Letters placed at group means. Envelopes contain all group members.

Table 3
Posterior classification of members of the bufonius complex.
MALES

| Group | Number of cases classified into group |  |  |  |  |
| :--- | ---: | ---: | ---: | ---: | ---: |
|  | A | B | C | D | E |
| A-bufonius | 52 | 1 | 0 | 0 | 0 |
| B-northern bufonius | 0 | 4 | 0 | 0 | 0 |
| C-labrosus | 0 | 0 | 8 | 0 | 1 |
| D-mystacinus | 1 | 5 | 0 | 37 | 1 |
| E-ventrimaculatus | 0 | 2 | 0 | 0 | 20 |

FEMALES

| Group | Number of cases classified into group |  |  |  |
| :--- | ---: | ---: | ---: | ---: |
|  | A | B | C | D |
| A-bufonius | 33 | 0 | 1 | 0 |
| B-labrosus | 0 | 21 | 0 | 2 |
| C-mystacinus | 2 | 0 | 11 | 0 |
| D-ventrimaculatus | 0 | 0 | 1 | 15 |

(NI), head length ratio (NI), head width ratio (NI). Separation of the groups is good, but not complete. In the posterior classification, 4 of 54 southern bufonius are assigned to northern bufonius, all northem bufonius are assigned to northern bufonius.

Eighty five southern bufonius and 27 northern bufonius males comprise the groups for analysis. The variables entered in the following order: foot texture, head length ratio, head width ratio (NI), foot ratio (NI), femur ratio (NI), tibia ratio (NI), SVL (NI). As for females, separation of the groups is good, but not complete. In the posterior classification, 4 of 85 southern bufonius are assigned to northern bufonius, all northern bufonius are assigned to northern bufonius.

For both the female and male data, virtually all of the separation of groups is accounted for by the first variable, foot texture. The F values, although not statistically interpretable, give the order of magnitude differences between the importance of the first and second variables in separating the groups. For females, the F value for the first variable is 182 , for the second, 10 ; for males, the F value for the first variable is 537 , for the second, 7.

In summary, the variables used in the computer analysis distinguish the predetermined species groupings quite well. Enough data are available to study geographic trends in L. mystacinus only.

Female mystacinus data.-The groups and sample sizes analyzed are: Brasil, Bahia (pooled localities), 3; Brasil, São Paulo (pooled localities), 7; Brasil, Rio Grande do Sul (pooled localities), 8; Uruguay (pooled localities), 4. The variables entered in the following order: foot texture, tibia ratio, head width ratio (NI), thigh stripe (NI), foot ratio (NI), SVL (NI), femur ratio (NI), tarsal texture (NI), head length ratio (NI). The first discriminant axis accounts for $71 \%$ of the total dispersion, the first two axes account for $90 \%$. The plot of the first against second discriminant axes indicates that the sample from the state of São Paulo is distinctive (fig. 23).

Male mystacinus data.-The groups and sample sizes analyzed are: Bolivia (pooled localities), 3, Brasil, Rio Grande do Sul (single locality), 19; Brasil, Rio Grande do Sul (remaining cases, pooled localities) 4; Brasil, São Paulo (single locality), 7; Brasil, São Paulo (remaining cases, pooled localities), 6; Argentina, Misiones (single locality), 5; Argentina (remaining cases, pooled localities), 9; Uruguay (pooled localities), 7. The variables entered in the following order: foot texture, SVL, tibia ratio, tarsal texture, thigh stripe, head width ratio, foot ratio (NI), lip stripe (NI), femur ratio (NI), head length ratio (NI). The first discriminant axis accounts for $50 \%$ of the total dispersion, the first two axes account for $80 \%$. The plot of the first against second discriminant axes (fig. 24) gives a pattern of separation best described together with the female data.

The pictorial results of group separation for female (fig. 23) and male (fig. 24) data show similar patterns in that the samples from the state of São Paulo are distinctive. Except for the Săo Paulo groups, the male data groups show a geographic trend of differentiation from Rio Grande do Sul - Uruguay - Argentina. The female data indicate that this trend is not complete, as the Bahia group is morphologically similar to the Rio Grande do Sul group. The São Paulo groups thus do not fit a clinal pattern of geographic differentiation.

Larvae.-The only species in this complex for which the larvae are adequately described is mystacinus (Sazima 1975).

Mating calls --Barrio (1965) figured and described the mating calls of $L$. bufonius and $L$. mystacinus. Werner C. A. Bokermann kindly gave me a copy of a recording of a northern bufonius. The call is very distinctive from southern bufonius (see species accounts for bufonius and troglodytes).

Taxonomic conclusions.-The five species recognized, bufonius, northem bufonius, labrosus, mystacinus, and ventrimaculatus, are morphologically distinguishable. The available mating call evidence supports


Figure 23. Discriminant axis plot for geographic samples of females of Leptodactylus mystacinus. $1=$ Brasil, Bahia, $2=$ Brasil, Sāo Paulo, $3=$ Brasil, Rio Grande do Sul, U = Uruguay. Numbers and letters placed at group means. Envelopes contain all group members.


Figure 24. Discriminant axis plot for geographic samples of males of Leptodactylus mystacinus. $1-2=$ Brasil, Rio Grande do Sul, 3-4 = Brasil, São Paulo, $Y-Z=$ Argentina, $U=$ Uruguay, $B=$ Bolivia. Numbers and letters placed at group means. Envelopes contain all group members.
recognition of these units. Until more call information becomes available for mystacinus, I prefer to treat it as a single species.

## LEPTODACTYLUS LATINASUS -LABIALIS

Specimens of Leptodactylus latinasus bear a striking resemblance to specimens of $L$. labialis. Both species usually lack well defined dorsolateral folds, are small, have prominent white tubercles on the tibia, tarsus, and foot, and have a distinct light thigh stripe. The two species are allopatric, one with a primarily Middle American distribution, the other with a primarily Chacoan distribution. Although there has never been much question regarding the specific distinctness of labialis and latinasus, I was curious to see how the stepwise discriminant function analysis would treat the morphological data. The following variables were used: 2-3, 9-14.

Female data.-The variables entered in the stepwise discriminant function analysis in the following order: lip stripe, head length ratio, head width ratio, tibia ratio, foot ratio, thigh stripe (NI), SVL (NI), femur ratio (NI). The first two discriminant axes account for all the variation. There is considerable overlap of the two groups (fig. 25). The percentage of specimens posteriorly clas-
sified in the other group amounts to $9 \%$ labialis assigned to latinasus and $24 \%$ latinasus assigned to labialis.
Male data.-The variables entered the stepwise discriminant analysis in the following order: SVL, head length ratio, tibia ratio, foot ratio, lip stripe, thigh stripe ( NI ), head width ratio ( NI ), femur ratio ( NI ). The first two discriminant axes account for $100 \%$ of the variation. There is considerable overlap of the two groups (fig. 26). The percentage of specimens assigned to the other group amounts to $14 \%$ for labialis and $9 \%$ for latinasus.

In contrast to the other species complexes analyzed by the stepwise discriminant function analysis (figs. 912, 21-22), there are no additional morphological features that were omitted from the analysis which will serve to further differentiate the two groups. The two species are very difficult to distinguish only on the basis of external morphology. The karyotypes and mating calls are distinctive however (see species accounts for fragilis and latinasus), amply verifying the specific level of distinction between the two.

## LEPTODACTYLUS LATINASUS

Morphology.-The data are analyzed geographically, using the following variables: $1-3,7-14$. Three of these are constant and do not appear in the analyses: $1,7,8$.


Figure 25. Discriminant axis plot for females of Leptodactylus labialis and latinasus. $\mathrm{F}=$ labialis, $\mathrm{L}=$ latinasus. Letters placed at group means. Envelopes contain all group members.
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Figure 26. Discriminant axis plot for males of Leptodactylus labialis and latinasus. $\mathbf{F}=$ labialis, $\mathrm{L}=$ latinasus. Letters placed at group means. Envelopes contain all group members.

Female data.-The following groups and numbers of specimens comprise the samples available for geographic analysis: Argentina, Buenos Aires (pooled localities), 4; Argentina, Catamarca (pooled localities), 4; Argentina, Formosa (single locality), 7; Argentina, Salta (single locality), 7; Argentina, Salta (single locality), 9; Argentina, Tucumán (single locality), 7; Brasil, Rio Grande do Sul (pooled localities), 8; Uruguay (pooled localities), 8. The variables entered in the following order: tibia ratio, SVL, head width ratio, head length ratio, thigh stripe, femur ratio (NI), foot ratio (NI), lip stripe (NI). The first two discriminant axes account for $72 \%$ of the total dispersion. The plot of the first discriminant axis against the second (fig. 27) indicates moderate separation of the groups, generally with geographically close samples being morphologically closest also. There is no overall trend of geographic variation.

Male data.-The following groups and numbers of specimens comprise the samples, available for geographic analysis: Argentina, Buenos Aires (pooled localities), 6; Argentina, Catamarca (single locality), 10;

Argentina, Chaco (single locality), 7; Argentina, Corrientes (single locality), 4; Argentina, Formosa (single locality), 4; Argentina, Jujuy (single locality), 5; Argentina, Jujuy (single locality), 10; Argentina, Salta (single locality), 22; Argentina, Salta (single locality), 6; Argentina, Salta (single locality), 6; Argentina, Tucumán (single locality), 8; Argentina, Tucumán (single locality), 29; Brasil, Bahia and Espirito Santo (pooled localities), 5; Brasil, Rio Grande do Sul (pooled localities), 10; Uruguay (pooled localities), 14. The variables entered in the following order: tibia ratio, SVL, head length ratio, femur ratio, head width ratio, foot ratio, thigh stripe (NI), lip stripe (NI). The first two discriminant axes account for $69 \%$ of the total dispersion. The plot of the first discriminant axis against the second (fig. 28) shows a complex pattern in which no geographic samples are distinctive nor are any geographic trends clearly discernable.

Larvae .-Fernandez and Fernandez (1921) described the larvae of $L$. latinasus (as prognathus) from Argentina. Mating calls.-Barrio (1965) described the call from


Figure 27. Discriminant axis plot for geographic samples of females of Leptodactylus latinasus. B = Brasil, Rio Grande do Sul, $\mathrm{U}=$ Uruguay, $\mathrm{Z}=$ Argentina, Buenos Aires, $1=$ Argentina, Catamarca, $2=$ Argentina, Tucuman, 3-4 = Argentina, Salta, 5
$=$ Argentina, Formosa. Letters and numbers placed at group means. Envelopes contain all group members.


Figure 28. Discriminant axis plot for geographic samples of males of Leptodactylus latinasus. $\mathrm{B}=$ Brasil, Bahia and Espirito Santo, $\mathrm{R}=$ Brasil, Rio Grande do Sul, $\mathrm{U}=$ Uruguay, $\mathrm{Z}=$ Argentina, Buenos Aires, $\mathrm{M}=$ Argentina, Corrientes, $\mathrm{I}=$ Argentina, Catamarca, 2-3 = Argentina, Jujuy, 4-5 = Argentina, Tucuḿan, 6-8 = Argentina, Salta, $9=$ Argentina, Formosa, $0=\mathrm{Ar}$ gentina, Chaco. Letters and numbers placed at group means. Envelopes contain all group members.
populations in two physiographically distinct areas in Argentina and concluded the calls represented the same species.

Taxonomic conclusion.-A single species is recognized.

## SUMMARY OF TAXONOMIC CONCLUSIONS

Based on the available data, 17 species are recognized in the fuscus species group (names as used in the analysis section):
albilabris
northern bufonius
southern bufonius
fuscus
barred gracilis
striped gracilis
labialis
labrosus
latinasus
longirostris
northern mystaceus
southern mystaceus
east coast mystaceus
south coast mystaceus
mystacinus
poecilochilus
ventrimaculatus

## NOMENCLATURE

Each name proposed for a member of the fuscus species group is discussed in chronological order.

Rana fusca Schneider 1799.-The confusion regarding this name has been commented on previously (Heyer 1968a). The neotype, Paris Museum 680, has been compared with recent material from French Guiana by Lescure (1972). He finds the specimens conspecific. This name applies to the species referred to as $L$. fuscus in the previous section.
Rana typhonia Daudin 1803.-Heyer (1968a) designated the male cotype of Rana typhonia Daudin as the neotype of Rana fusca Schneider. Lescure (1972) compared the type with recent specimens from French Guiana;
all apply to the species referred to as $L$. fuscus in the analysis section.

Rana mystacea Spix 1824.-Spix's description is based on a specimen from Bahia; he compares the described specimen with a second from Solimoens (female?). Peters (1873), the last person to examine the Spix types before they were lost, concluded that the specimens were synonyms of Rana typhonia Daudin. Peters (1873) clarified the Spix figure legends. Figure 3, plate 3, is an adult male from Bahia. Figure 1, plate 3, is an adult female from Solimoens. Bokermann (1966) gives the type locality of Rana mystacea as Salvador, Bahia. Spix's plate figures clearly pertain to members of the mystaceus complex as analyzed previously herein. The diagnostic characters for the mystaceus complex are the white tubercular conditions of the tibia, tarsus, and foot. Neither Spix nor Peters mentions these characters for any of the type specimens. In this case, geographic location of the two Spix types is sufficient for proper allocation. The taxon referred to as "east coast mystaceus" in the preceding analysis is the only member of the complex found in coastal Bahia; the taxon I termed "northern mystaceus"' is the only species found along the Rio Solimōes. Thus it appears that the two Spix type specimens represent different species. As all members of this complex have traditionally been called mystaceus, nomenclatural stability will not be improved by choosing one or the other of the two specimens as the form to which the name applies. As Spix gave the detailed description of the specimen from Bahia, I choose the specimen figured in figure 3 , plate 3 as the name bearer of mystacea. Thus mystacea applies to the species called east coast mystaceus in the analysis section. There is enough confusion in this case that a designation of a neotype would appear to be in order. Unfortunately, no museum specimens are available from near the type locality of Salvador. As the species of the mystaceus complex are for the most part allopatrically distributed, the result of the action taken here should be clear to any subsequent worker in spite of not designating a neotype.

Rana sibilatrix Wied-Neuwied 1824.-Wied describes sibilatrix in several publications; the figure published in 1824 (Wied 1824) is usually cited for the original description of the name. The type specimen is apparently no longer extant. The figure, together with the restricted type locality of Marobá (= Vila Viçosa), Rio Peruipé (Müller 1927 as clarified by Bokermann, 1966) clearly allocates the name to the species identified as $L$. fuscus in the analysis. The figure shows a spotted dorsum with several dorsolateral folds. The species identified as L. fuscus is the only species along coastal Bahia to which the name can apply.

I have examined AMNH 485, a specimen from the Wied-Neuwied collection originally identied as sibilatrix. The specimen is a male with obvious, dark, vocal sacs; no vocal sacs are indicated in the figure of the type specimen. There is no convincing evidence that associates or disassociates AMNH 485 with Wied-Neu-
wied's figure. The locality given for the specimen is simply Brasil.
I can find no mention of Wied clearly associating Vila Viçosa as the collecting locality for Rana sibilatrix (Wied-Neuwied 1820). I have not examined museum specimens from this locality.

As this study shows no marked differences between specimens of L. fuscus from coastal Brasil and French Guiana, there is no need to make a final decision on whether AMNH 485 is actually the type of Rana sibilatrix or whether Vila Viçosa should be accepted as the type locality for the taxon.

Cystignathus gracilis Duméril and Bibron 1841.The holotype, Paris Museum 4490, still contains the salient features to allocate the name properly. The holotype is a member of the gracilis complex as used by previous authors. The question is whether it is a barred or striped gracilis, as those terms are used in this analysis. The tibias, although soft and partly faded, clearly show the light longitudinal stripes; the name applies to the population identified as striped gracilis in the analysis section.

Cystignathus typhonius Duméril and Bibron 1841.As pointed out previously (Heyer 1968a), although Duméril and Bibron indicated that the description they provided was of a new species, the name dates back to Daudin. The same specimens are involved; the lectotype of typhonia was designated as the neotype of Rana fusca. The name applies to the species identified as $L$. fuscus in the previous analysis.

Cystignathus schomburgkii Troschel 1848.—Attempts to locate the type material of this taxon have been unsuccessful. The types are not at any of the major German museums at present. The most likely depository was the collection in Leipzig. All of the herpetological material in this collection was transferred to the Staatliches Museum für Tierkunde in Dresden in 1972. F. J. Obst, the curator at the Dresden Museum, kindly informs me that Troschel's type material of $C$. schomburgkii is not in the Leipzig collection now housed at Dresden. Further, he has no knowledge of where Troschel's material might be. Troschel described two other new species in the same paper where he described $C$. schomburgkii: Podocnemis unifilis and Hyla calcarata. Duellman (1973, p. 522) was not able to locate the type of $H$. calcarata. In a brief literature search on Podocnemis unifilis, I find no one who refers to the type specimens. In all probability, Troschel's type specimens are lost.

Troschel's description of C. schomburgkii is brief and inconclusive. Three statements in the description give possible clues to the identity of $C$. schomburgkii: (1) the species is closest to C. gracilis; (2) color above uniform brown; (3) commonly found in dense, damp woods and in woodland swamps. The following fuscus group species are known from Guyana: L. fuscus, L. longirostris, northern mystaceus. Troschel's statement of close relationship with gracilis suggestsfuscus. The uniform brown
dorsal color could only be longirostris of the Guyana members of the fuscus group, assuming that Troschel excluded the dorsal chevron as a uniform pattern. There is no reason to assume that Troschel and I mean the same thing by a uniform pattern, however. To my knowledge, no member of the fuscus group is commonly found in forests. Leptodactylus longirostris likely comes closest, being found in open situations in conjunction with forests. Thus C. schomburgkii most likely refers either to L. fuscus or longirostris. If the name applies to fuscus, it is a junior synonym; if it applies to longirostris, it is the oldest available name for that species. Historically, the name has been treated as a synonym of sibilatrix ( $=$ fuscus). As there is no proof that the types of schomburgkii have been destroyed, there is the remote possibility that the types still exist. As long as this possibility exists, I think the proper position to take at this point is to consider schomburgkii as a synonym of $L$. fuscus until such time as the status of the types can be resolved.

Cystignathus albilabris Günther 1859.-I have examined two syntypes from the series of specimens from St. Thomas, West Indies. Both specimens clearly represent the same taxon as the species referred to as albilabris in the analysis section. Günther's description reads as if he were looking at all the specimens, rather than describing one individual from the series. I hereby designate BMNH 1947.2.1760, an adult 35.3 mm male as the lectotype of Cystignathus albilabris.

Cystignathus mystacinus Burmeister 1861.-The holotype is in the collections of the Martin-LutherUniversität, Halle (Saale). Apparently no precise type locality was ever associated with the specimen other than Argentina. The specimen, although faded, is in a good state of preservation. It is a male, 51.8 mm SVL with the following diagnostic characteristics still visible: a pair of dorsolateral folds, white tubercles dorsally present in the sacral region, as well as on the dorsal surface of the tibia and lower surfaces of the tarsus and foot. The specimen is too faded to state for certain whether it has a light lip or thigh stripe, but there is no doubt that it belongs to the species identified in the analysis section as mystacinus.

Cystignathus poecilochilus Cope 1862.—The type specimen from Colombia is soft and faded and generally in poor condition. The light stripe on the posterior face of the thigh is still evident and the foot and tarsal surfaces are smooth. The name applies to the species identified as L. poecilochilus in the previous analysis.

Leptodactylus labrosus Jiménez de la Espada 1875.Heyer and Peters (1971) discussed the type specimen. The name applies to the species identified as $L$. labrosus in the analysis.
Leptodactylus latinasus Jiménez de la Espada 1875.— Heyer (1969) discussed the type specimen. The name applies to the species identified as $L$. latinasus in the analysis.

Cystignathus fragilis Brocchi 1877.-The holotype, Paris Museum 6316, from Tehuantepec, Mexico, is
clearly the same species analyzed as L. labialis. The specimen has tubercles on the tarsus and sole of foot, distinct light stripe on the posterior face of the thigh, indistinct light lip stripe, and somewhat distinct dorsolateral folds. The holotype compares well with other specimens collected from the Tehuantepec region.

Leptodactylus fragilis (Brocchi) is the oldest name for the species in question. Previously (Heyer 1971), I incorrectly cited the date of publication of Cystignathus labialis Cope as 1877. (Brocchi 1881, also thought that Cope's name predated fragilis.) The year 1877 is when the paper was read at the meeting, but the description was published in 1878. As discussed next, C. labialis Cope applies to a South American species, not to any species found in Mexico. The priority of the name, together with the misapplication of $C$. labialis for a Mexican species, leads to the conclusion that the best course of action is to use the name L. fragilis Brocchi for the species in question. See C. labialis (next) for further discussion.

Cystignathus labialis Cope 1878.-The juvenile holotype and 5 paratypes are so faded that no patterns are visible. The series demonstrates the following diagnostic character states: distinct dorsolateral folds from eye to groin; tubercles on tibia, tarsus, and sole of foot; posterior face of thigh lacking a light stripe; skin warty along sides and posterior dorsum. These states best match the species identified as L. mystacinus in the analysis section. Occasional individuals of specimens identified as $L$. labialis in the analysis section have uniform posterior faces of the thighs, but a series of 3 or 4 spec imens always shows the light stripe. The same is true for $L$. albilabris and members of the mystaceus complex. The types were directly compared to faded juvenile specimens of $L$. albilabris and $L$. mystacinus (both as used in analysis section). Even in faded juvenile albilabris, the light thigh stripe is evident. The types match juvenile specimens of L. mystacinus. A faded L. mystacinus even has a white pin stripe along the dorsolateral folds, as do most of the labialis types. The type specimens thus do not pertain to the Middle American species as has always been assumed, but to the species usually called mystacinus.

In the original description, Cope gives the following diagnostic character states: one dermal fold on each side; skin rough; color chocolate brown; a brilliant white band extends from the anterior part of the upper lip, and describing a curve upwards, bounds the orbit below and descends to the canthus oris, from which point it continues in a straight line to the humerus, and ceases. All these states fit the species identified as $L$. mystacinus in the analysis section. The statement "color chocolate brown'' better fits mystacinus than labialis (as used in analysis), as the former often has a uniform dorsum and the latter has some sort of spotting or mottling. All mystacinus have distinct light lip stripes; few labialis (as used in analysis) have brilliant white lip stripes. Cope gives the following measurements (my measurements in
meters of holotype in parentheses): length of head and body, .020 (.018); of head, .007 (. 007 from tip of snout to posterior tympanum); of hind limb, 028 (.027), of hind foot, .013 (.018). The description and specimen match except for the hind foot length, which might have been a typographical error in the description.

Cope makes the following statement concerning locality, 'The precise habitat of this species is at present uncertain. It is probably a part of Sumichrast's Mexican collection." The introductory paragraph of the paper states, "The greater number of the species described in this paper were sent to the Smithsonian Institution by its correspondents, and submitted to my examination by its Secretary, Professor Henry.' Included in the paper are Mexican specimens collected by Sumichrast, Xantus, and others, Costa Rican material collected by Gabb and Franzius, Panamanian material collected by Selfridge, and a collection of 9 species, two described as new, from the following locality, 'Habitat unknown, but supposed to be the Argentine Confederation." The species from this collection are indeed known from Argentina. Thus, it is as reasonable to assume that the specimens Cope described as $C$. labialis were part of the Argentina collection as the Sumichrast Mexican collection.

There has been confusion regarding some of the Cope types in the Smithsonian Collection. The specimens were returned to the Smithsonian after Cope died, but were not indicated especially as types. The specimens were originally recorded in the catalogue as Cystignathus labialis and no locality information was originally entered. Under remarks, the statement 'Ret. from Cope's Estate" is recorded. Later, the following data were entered in pencil, 'Probably Tehuantepec (?), [Collector] (?) Francis Sumichrast." Presumably this information was entered based on the information Cope gave in the description. Other material returned from Cope's Estate, entered in the catalog at the same time, includes some, but not all of the types Cope described in the same paper as labialis. Even though the specimens are now labelled as holotype and paratypes, this action seems to have been taken by a cataloguer, not a revisor in print. No further action need be taken, for if the series were still syntypes, the specimen now labelled as the holotype would clearly be the best choice to designate as lectotype.

As I have previously stated in print (Heyer 1971) that I examined the holotype and applied it to the Middle American species, comment on my previous decision is required. The 1971 statement is based upon a 1967 examination of the type specimen, when I first started systematic work on the genus. My original notes are, '"Type examined 3 September 1967. The specimen is so faded that it is virtually impossible to see any pattern. Using all my imagination, I could perhaps make out a posterior thigh light stripe. The toes are not fringed, and the tarsus and sole of foot are covered with white (they could be nothing else due to the fading of the specimen)
tubercles." At that time I did not look at the other type specimens. It is these specimens which clearly show the dorsolateral folds (the holotype is wrinkled along the sides due to tying the tag tightly around the waist and the dorsolateral folds do not distinctly stand out from the wrinkles). In 1967 I assumed that Mexico was the correct locality and knew that there were only two species of Leptodactylus in Mexico and that the holotype was certainly not $L$. melanonotus. Now that I have studied all members of the fuscus group, it is clear that the type specimens of labialis are certainly not from Mexico and that they are the same as species found in Argentina.

The evidence is reasonably conclusive. The specimens and description separately match the species identified as $L$. mystacinus in the analysis section. The holotype matches the description with the exception of the foot measurement. None of the Cope specimens can conclusively be demonstrated to be the holotype, but the evidence is most consistent with acceptance of the specimens as the types. Thus, the name Cystignathus labialis applies to the species identified as L. mystacinus in the analysis section and not to L. labialis as used in the analysis section.

The proper allocation of the name $C$. labialis will cause some confusion, as Leptodactylus labialis as currently understood is a well known species concerning which there is a sizeable body of literature. The Middle American species is mostly known to professional herpetologists, however, not to physiologists or general anatomists. The impact of the proper allocation of the name will not be felt outside of the herpetological community. The herpetological community has already endured one name change, as Kellogg's (1932) influential work considered the species in question a junior synonym of L. albilabris. In the long run, nomenclatural stability will best be served by the proper allocation of the type of Cystignathus labialis Cope.

Leptodactylus longirostris Boulenger 1882.—Boulenger based the new species upon two specimens, BMNH 76.5.26.4 and 76.5.26.5. Allocation of these specimens with the species recognized in the analysis section is not straightforward and requires discussion of the two specimens.

BMNH 76.5.26.4 is the better preserved of the two specimens. It is a 48.5 mm female, with smooth feet from Santarem, Brasil. A member of the L. mystaceus complex occurs in the Santarem region, but other specimens referred to as "longirostris' in the analysis section have not been taken that far south, the distribution of 'longirostris'" being broadly associated with the Guiana shield. The size of the specimen matches members of the Amazonian mystaceus complex, but is about 3 mm larger than any 'longirostris', examined. The smooth foot matches 'longirostris' and differs from the Amazonian mystaceus complex species. BMNH 76.5.26.4 has a narrow head ( $31 \%$ SVL) and long femur ( $51 \%$ SVL), tibia ( $59 \%$ SVL), and foot ( $60 \%$ SVL).

The head width is narrower than the average head widths of both 'longirostris'" and the Amazonian mystaceus complex species. The head width is matched by 6 individual "longirostris" and 2 individual Amazonian mystaceus complex species. The hind limb is longer than the average hind limb of both species, but individuals of 'longirostris'' match the limb proportions of the type for the femur, tibia, and foot, whereas no individuals of the Amazonian mystaceus complex species have the same length of either the femur or tibia. Two details of color pattern of the type specimen are matched by "longirostris' specimens and not by the Amazonian mystaceus complex species: (1) a postorbital dark triangle with the apex pointing toward the angle of the jaw, and (2) posterior continuations of the dark mid-dorsal scapular chevron.

The second specimen, BMNH 76.5.26.5 has an anomalous right leg, but is otherwise in good condition. The right femur appears shortened, the tibia is only a few millimeters long terminating in 3 misshapen toes. The female specimen resembles the other type in the following features: head length, hind limb proportions, foot texture, postorbital dark triangle. The specimen differs from the other type specimen in SVL ( 43.4 mm ) and head width ( $33 \% \mathrm{SVL}$ ): both of these measurements are the same as found in specimens analyzed as 'longirostris" in the previous section.

Both types of L. longirostris clearly represent the same taxon. The combined information on both specimens is most consistent with the specimens analyzed as longirostris. Two aspects do not allow a certain allocation of the types with the Guiana shield species analyzed as longirostris at this time: the large size of one of the female types and locality. Two possible conclusions may be drawn. 1) The types of $L$. longirostris represent a distinct species from the Guiana shield species analyzed as longirostris. This conclusion would be supported by the locality data and size differences and would require the recognition of the two as sibling species essentially indistinguishable morphologically. 2) The types of $L$. longirostris represent the same species as the Guiana shield species identified as $L$. longirostris in the analysis section. This conclusion would be consistent with the morphological data except for female size, which would have to be explained as due to small sample size of longirostris museum specimens or geographic variation, etc. This conclusion would also suggest that Santarem was the shipping port and the specimens were actually collected from the upper Mapuera or Trombetas rivers, for example.

I know of only two other museum specimens that resemble the types of $L$. longirostris in form and geographic provenance (MZUSP 24880, Ponta Negra, Rio Negro, Amazonas, MZUSP 37518, Tapera, Rio Negro, Amazonas) and these specimens are close geographically to the Guiana shield region. I therefore hesitate to recognize two distinct species in this assemblage, rec-
ognizing that additional data may require a re-evaluation of this position. Thus, for present purposes, I consider the types of $L$. longirostris to represent the same species as the species identified as $L$. Iongirostris in the analysis section. As specimen BMNH 76.5.26.4 is clearly the specimen described and figured by Boulenger, I hereby designate it as the lectotype.
Leptodactylus prognathus Boulenger 1888.-The holotype is clearly the same species identified as latinasus in the analysis section. Boulenger's description is misleading in one respect. He states that the 33 mm specimen is a half-grown male specimen. The specimen has vocal slits and external lateral vocal sac folds: it is a fully adult male.
Leptodactylus andicola Boettger 1891.-This name has been associated with members of the fuscus group (Heyer, 1974). Dr. John Lynch concurs with my current opinion that this name applies to the genus Eleutherodactylus. The type has been destroyed.
Leptodactylus quadrivittatus Cope 1893.-The holotype is apparently lost. The description matches recent specimens from Costa Rica. The name can only pertain to the species identified as $L$. poecilochilus in the analysis, as it is the only species in Costa Rica to have the mid-dorsal stripe color pattern phase. There is no taxonomic confusion surrounding this name and the holotype may yet be identified as such, thus there is no reason to designate a neotype for quadrivittatus.
Leptodactylus bufonius Boulenger 1894.-I have examined two of the four syntypes; the types are the same species called bufonius or southern bufonius in the analysis section. Boulenger gives the snout to vent measurement as 48 mm . I measure 46.4 mm on specimen BMNH 1947.2.17.72, the first specimen in the series. As this specimen is likely the one Boulenger's description is based upon, and is still well preserved, I hereby designate this female specimen as the lectotype of Leptodactylus bufonius Boulenger.
Leptodactylus maculilabris Boulenger 1896.-I have examined the type specimen and concur with previous workers that it represents the same species identified as L. poecilochilus in the analysis section.

Leptodactylus raniformis Werner 1899.-The holotype, an adult male, is clearly a member of the Amazonian Colombian population of fuscus as analyzed previously. The dorsolateral folds are indistinct, but the spotting pattern on the dorsum is characteristic of fuscus as is the tarsal and foot surfaces (smooth with light pigment spots).
Leptodactylus ventrimaculatus Boulenger 1902.—The type series was previously commented on (Heyer and Peters, 1971). The name applies to the species identified as $L$. ventrimaculatus in the analysis section.
Leptodactylus diptychus Boulenger 1918.-The holotype has the following diagnostic character states: light stripe on posterior face of thigh, smooth tarsus and foot; 2 distinct dorsolateral folds; lips with dark brown spots;
45.4 mm female. There is only one species in Venezuela that this combination of character states can apply to: the species identified as $L$. poecilochilus in the analysis section. It is the only species with brown lip spots with the other character states mentioned. The type locality, Andes of Venezuela, is probably in slight error, as $L$. poecilochilus is known only from the coastal plain region of Venezuela. As discussed in the analysis section, the presumed mating calls of Middle American and Venezuelan poecilochilus are distinctive. If two species are actually involved, diptychus would apply to the Venezuelan form.
Leptodactylus curtus Barbour and Noble 1920. Heyer and Peters (1971) discussed the allocation of this name. The name applies to the species identified as $L$. labrosus in the analysis section.
Leptodactylus dominicensis Cochran 1923.-The type is clearly a member of the L. albilabris complex. As but one species is recognized in this complex, the name is a synonym of albilabris. As noted before, the Dominican Republic population is distinctive. If further work demonstrates that the Dominican Republic population is distinct at the species level, then dominicensis would apply to this form.
Leptodactylus troglodytes Lutz 1926.-The specimen labelled as the type in the Adolfo Lutz collection is completely faded in any exposed areas. It is impossible to tell whether the foot and tarsus are tuberculate or not. It is clear, however, that the name applies to the form identified as northern bufonius in the analysis section as the type locality is Pernambuco. As northern bufonius is recognized as a distinct species herein, troglodytes applies to this species.
Leptodactylus plaumanni Ahl 1936.-I have been unable to examine the holotype. Mertens (1967) considered plaumanni a synonym of sibilator ( $=$ fuscus as used here). The original description appears to better match what is here recognized as striped gracilis, particularly in details of dorsal and tibia color pattern. In order to point out that plaumanni may refer to another species than fuscus, I prefer to place L. plaumanni in the synonym of $L$. gracilis until such time that I am able to examine the holotype. Also see $L$. geminus.
Leptodactylus anceps Gallardo 1964.—Gallardo described anceps, indicating that it had a Chacoan distribution, whereas prognathus (= latinasus) had a coastal distribution. I have examined paratypes of $L$. anceps and specimens identified by Gallardo as anceps. The name certainly applies to the group of frogs analyzed herein as latinasus. In order to examine the morphological distinctiveness of anceps, a discriminant function analysis was run, using specimens from geographic areas clearly within the range of anceps or latinasus as defined by Gallardo (1964).

The sample sizes for females are 20 latinasus and 34 anceps. The variables entered in the following order: tibia ratio, SVL, femur ratio, head length ratio (NI), foot
ratio (NI), thigh stripe (NI), head width ratio (NI). Two of 20 latinasus were posteriorly classified as anceps, 1 of 34 anceps classified as latinasus. This classification separation is also shown in the plot of the first two discriminant axes, where $100 \%$ of the dispersion is accounted for by the first axis (fig. 29).

Male sample sizes are 35 latinasus and 107 anceps. The variables entered as follows: tibia ratio, SVL, head width ratio, head length ratio (NI), lip strip (NI), femur ratio (NI), thigh stripe (NI). Three latinasus were posteriorly classified as anceps, 8 anceps were classified as latinasus. All of the dispersion is accounted for by the first discriminant axis (fig. 30).

The female and male results complement one another. The species Gallardo described as anceps is morphologically distinctive from latinasus, but there is some morphological overlap. The complexity of the overlap is better seen in figs. 27 and 28 where the pattern of geographic variation is complex and not easily interpretable. Interestingly, Gallardo distinguished anceps in large part by differences in snout shape. The head ratios used here do not reflect those differences. Although the Chacoan populations are morphologically differentiated from the coastal populations, there is some morphological overlap (as specimens from questionable, intermediate localities were omitted from the analysis, the degree of overlap may be even greater than evidenced in the analysis). This morphological overlap, together with similarity of mating call (Barrio 1965) is interpreted to mean that a single species is involved and that anceps is a synonym of latinasus.

Leptodactylus gualambensis Gallardo 1964. Gallardo (1964) described gualambensis as a species of the fuscus group with a Chacoan distribution. I have examined specimens Gallardo identified as gualambensis and find them morphologically identical to fuscus. The geographic analysis of fuscus (figs. 13 and 14) did not demonstrate a morphological distinctiveness of the Chaco populations of fuscus. In order to specifically test for the morphological distinctiveness of gualambensis, a discriminant function analysis was run on individuals from the Chaco, including specimens Gallardo identified as gualambensis and non-Chaco individuals. Any specimens from possible intermediate localities were not used in this analysis. There are enough data for analysis of males only.

Sample sizes are 30 gualambensis and 163 fuscus. The variables entered in the following order: tibia ratio, SVL, dorsal pattern, thigh stripe, foot texture, femur ratio (NI), head width ratio (NI), head length ratio (NI), tarsal texture (NI), lip stripe (NI). Four gualambensis were posteriorly classified as fuscus, 22 fuscus were classified as gualambensis. The first discriminant axis accounts for $100 \%$ of the total dispersion (fig. 31). The results show that the Chaco populations are moderately distinctive, but fall within the range of morphological variability of the other populations of fuscus. Barrio


Figure 29. Discriminant axis plot for females of Leptodactylus anceps and latinasus. $\mathrm{A}=$ anceps, $\mathrm{L}=$ latinasus. Letters placed at group means. Envelopes contain all group members.


Figure 30. Discriminant axis plot for males of Leptodactylus anceps and latinasus. $\mathrm{A}=$ anceps, $\mathrm{L}=$ latinasus. Letters placed at group means. Envelopes contain all group members.


Figure 31. Discriminant axis plot for males of Leptodactylus fuscus and gualambensis, $\mathrm{F}=$ fuscus, $\mathrm{G}=$ gualambensis. Letters placed at group means. Envelopes contain all group members.
(1965) indicated that the call of gualambensis is not distinctive from the call of southeastern Brasilfuscus. This, together with the fact that there is morphological overlap of gualambensis and fuscus leads to the conclusion that but one species is involved.
Leptodactylus gracilis delattini Müller 1968.-Data were taken on the holotype and two paratypes (MZUSP). The data on the three specimens are compared with data on male barred and striped gracilis in a discriminant function analysis. The results of the plot of the first two discriminant axes are adequate for discussion (fig. 32). The subspecies is clearly morphologically closest to striped gracilis. The dorsal surface of the tibia has the folds characteristic of striped gracilis, but lacks the stripes themselves. The tibia pattern of the types are thus barred gracilis. Most morphological evidence is consistent with striped gracilis: the subspecies is considered here as belonging to striped gracilis, but the island population is distinctive.

Leptodactylus geminus Barrio 1973.-Barrio differentiated geminus from gracilis based solely upon distinctive features of the mating call, pointing out that the external morphologies and karyotypes of the two taxa are practically identical. I have not been able to examine the type specimens of geminus in detail. Barrio's figures
show light longitudinal stripes on the tibia on both geminus and the gracilis he compared geminus with. Barrio's call data clearly indicate that two species are included in what I have recognized as "striped gracilis" in the analysis section. Pending further morphological analysis, $L$. geminus is recognized on the basis of its distinctive call, but subsequent discussions of specimens will not differentiate between geminus and gracilis. A further nomenclatural complication is that $L$. plaumanni may refer to either striped gracilis or geminus. The type description suggests that plaumanni would pertain to striped gracilis.
Leptodactylus marambaiae Izecksohn 1976.—Izecksohn distinguished the new species from L. gracilis on the basis of a shorter leg, smaller size, and color pattern. The species is very similar to striped gracilis as analyzed herein. Werner C. A. Bokermann kindly gave me a copy of a recording of the mating call (recorded by Izecksohn). The call is distinctive from striped gracilis and geminus (see species accounts).

All of the proposed names apply to 13 of the species recognized in the analysis section plus two species not recognized in that section. Thus, four species remain unnamed. Descriptions for these new species are included in the accounts below.


Figure 32. Discriminant axis plot for males of Leptodactylus gracilis delattini, striped gracilis, and barred gracilis. B = barred gracilis, $\mathrm{D}=$ gracilis delattini, $\mathrm{S}=$ striped gracilis. Letters placed at group means. Envelopes contain all group members.

## SPECIES ACCOUNTS

Members of the fuscus group of Leptodactylus are small to moderate sized frogs; the toes lack fringe or web, the head is of normal width proportions, and the males lack thumb spines. All members of the melanonotus and ocellatus groups have fringe on the toes. All members of the pentadactylus group are moderate to large frogs having broad heads and in most species the males have thumb spines.

In the descriptions, only the holotypes of new species are described in detail. The characteristics used to describe the adults and larvae are those which differentiate the member species (which are the same characters used in the analysis section for the most part).

In the adult characteristics sections, N refers to the number of adult individuals used for statistical analyses. Numerical summaries are means plus or minus one standard deviation.

All known tadpoles are quite similar morphologically; all have a denticle formula of $\frac{1-1}{3}$ or $\frac{1}{\frac{1-1}{1-1}}$, an entire oral disk with an anterior papillary gap, a median anus, sinistral spiracle, a pond type larval morphology, and an indistinct, mottled body and tail pattern. These features are not repeated in the species accounts.

Locality data are recorded as nearly as possible to the original catalog data and are not standardized in terms of distances or altitudes. Numbers in parentheses after museum numbers indicate the number of specimens with the same museum number.

The maps are computer generated and are based on localities for which longitudes and latitudes could be found.

LEPTODACTYLUS ALBILABRIS (GÜNTHER) 1859
Cystignathus albilabris Günther 1859:217. (Type locality, West Indies, St. Thomas. Lectotype BMNH 1947.2.1760, adult male.)
Leptodactylus dominicensis Cochran 1923:184-185. (Type locality, Dominican Republic; El Seibo Province, Las Cañitas. Holotype USNM 65670, adult male.)

Diagnosis-The other species which have a light stripe on the posterior face of the thigh and obvious white tubercles on the tarsus and foot are elenae, fragilis, latinasus, mystaceus, and mystacinus. The dorsal surface of the tibia is covered with obvious white tubercles in albilabris, white tubercles are lacking or indistinct on the dorsal surface of the tibia in elenae. Leptodactylus albilabris has a pair of distinct dorsolateral folds, in fragilis and latinasus the folds, if present, are indistinct. Leptodactylus albilabris is also larger (males $30.3-43.1 \mathrm{~mm}$, females $34.9-45.4 \mathrm{~mm}$ ) than either fra-
gilis (males 25.9-43.0, females 28.7-43.6 mm) or latinasus (males 27.9-37.9 mm, females 29.0-36.3 mm), but smaller than $L$. mystacinus (males $43.6-58.8 \mathrm{~mm}$, females 53.8-64.3 mm). Leptodactylus albilabris has a shorter tibia (male mean $43 \%$ SVL, female $44 \%$ ) and foot (male mean 49\% SVL, female $50 \%$ ) than $L$. mystaceus (tibia-male mean $51 \%$ SVL, female $52 \%$; footmale mean 55\% SVL, female 55\%). Leptodactylus albilabris is the only group member found in the West Indies.

Adult Characteristics $(\mathrm{N}=268)$-Dorsum with dorsal chevrons, blotches, or confluent chevrons and blotches (fig. 1, A, B); mid-dorsal light stripe present in $17 \%$ of individuals, presence not sexually dimorphic ( $X^{2}=1.05$, $P>.05$ ); light lip stripe almost always distinct (94\%), rarely indistinct ( $6 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=0.58, P>.05$ ); dark suborbital bar absent; light stripe on posterior face of thigh present, distinct $(56 \%)$ or indistinct ( $44 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=0, P>.05$ ); tibia barred; 2 distinct dorsolateral folds; dorsal surface of tibia usually covered with many white tubercles, sometimes scattered with white tubercles; posterior surface of tarsus with many white tubercles; sole of foot with many white tubercles; male SVL $35.2 \pm 2.7 \mathrm{~mm}$, female $40.7 \pm 3.1$ mm , females larger than males ( $t=15.19, P<.01$ ); male head length/SVL ratio $.380 \pm .013$, female .372 $\pm .015$, male head longer ( $t=4.64, P<.01$ ); male head width/SVL ratio $.354 \pm .014$, female $.354 \pm .015$, not sexually dimorphic ( $t=.354, P>.05$ ); male femur/ SVL ratio $.400 \pm .024$, female $.414 \pm .024$, female femur longer $(t=4.68, P<.01)$; male tibia/SVL ratio $.431 \pm .022$, female $.442 \pm .025$, female tibia longer ( $t=3.80, P<.01$ ); male foot/SVL ratio $.487 \pm .025$, female $.495 \pm .028$, female foot longer $(t=2.45, P$ $<.02$ ).
Larval Characteristics-Eye diameter 9-11\% headbody length; oral disk width $20-25 \%$ head-body length; oral papilla gap $48-56 \%$ oral disk width; $42-67$ denticles in anterior split tooth row on one side; head-body length 33-43\% total length; total length, stage 36, 42.6 mm (fig. 33).

Mating Call (figs. 5 and 6)-Dominant ( $=$ fundamental) frequency modulated between $2000-2800 \mathrm{hz}$; no harmonic structure in call; each note of two pulses, a lower frequency and intensity pulse of .004 to .013 $s$ duration followed without pause by a pulse of higher frequency and intensity of .038 to .040 s duration.

Karyotype - Bogart (1974) described the karyotype as diploid number 22; 7 pair median, 3 pair submedian, 1 pair subterminal; secondary constriction in chromosome pair 8.

Distribution - Known from the Virgin Islands, Puerto Rico and eastern Dominican Republic (fig. 34).

ANEGADA. MCZ 4198-4202, 4208-224.
DOMINICAN REPUBLIC. No specific locality, AMNH 20925, 20937, 20943, 20951-52, 20958, 20961-64.

EL SEIBO: Sabana de La Mar, AMNH 34402-411; 3.2 km E Sabana de La Mar, USNM field 41045-051.

PUERTO RICO. Adjuntas, USNM 25607; Aguada, CM 36058 (10); Aguadilla Rincon, CM 46462; Aguas Buenas, USNM 25628-631; Aibonito, AMNH 10030-39, USNM 25759, Añasco, USNM 25726-27; Arecibo, 23 km W, USNM 86560; Arroyo, USNM 25728-731; Bayamón, MCZ $4104-$ 110, 21865 (5), USNM 25772-75; Cabo Rojo, 3 km E, MCZ 30790; Canóvanas, $10 \mathrm{~km} \mathrm{~S}, \mathrm{MCZ}$ 19023-050; Caguas, USNM 25740; Cartagena Lagoon, MCZ 19014-19; Catalina Plantation, USNM 26894; Cayey, MCZ 18976, 18978-985 (5); Cayo Santiago, MCZ 31576-78; Coamo Springs, MCZ 19020-22; Culebra Island, MCZ 18963-65; Desengano, FMNH 1238486; Húcares, USNM 26091-92; Humaco, USNM 27775, 8660209; Lares, USNM 62933; Luquillo, USNM 27053; Mameyes, USNM 26820-23, 26825-833, 26835, 26981-82; Mayagüez, CM 46418-425, 46457, 46500, FMNH 12379, 12413-15, MCZ 30791, 34052-57, USNM 27749-758, 29357-362, 29390-91, 100901; Ponce, MCZ 2756 (3), USNM 27313; Pueblo Viejo, USNM 26817-19, 86559; Río Piedras, FMNH 38582, MCZ 19001-013, 21893; Santa Barbara, USNM 31089; San Germán, USNM 86561-64; San Juan, MCZ 2187 (52); Utuado, MCZ 9352, USNM 27227-236; Vieques Island, USNM 27084-099, 27103-138; El Yunque, KU 79231; Zugillo Mtns, MCZ 18986-19000 (3).

ST. CROIX. No specific locality, CM 18821 (11), KU 94395-96, MCZ 3706-09, USNM 115898-5906; Bethlehem, USNM 162238-243; Caledonia, MCZ 24146.

ST. JOHN ISLAND. No specific locality, MCZ 18949958 (27); Annaberg, KU 45629.

ST. THOMAS. No specific locality, AMNH 52653-57 (11), FMNH 11290 (3), 42076-080, MCZ 18959-962, USNM 15403-08, 52499-2503, 52512-524, 52527, 119036-37, 161011; near Magens Bay, USNM 52504-06; near Smith Bay, USNM 103163 (tadpoles).

TORTOLA. No specific locality, AMNH 77503-05, FMNH 11284 (6), MCZ 4225-235, 189666-69, 189671, 189673-75 (15).

## Leptodactylus amazonicus new species

## Figure 35

Holotype: LACM 92111, an adult male from Ecuador; Napo Province, Limoncocha, $0^{\circ} 24^{\prime} \mathrm{S}, 76^{\circ} 37^{\prime} \mathrm{W}$, elevation 260 m . Collected by Keith A. Berven and W. Ronald Heyer on 15 July 1971.

Paratopotypes: LACM 92067-070, 92072-75, 92077-085, 92087, 92090-92, 92094-95, 92098, 92102-05, 92108, 92112-$15,92117-20,92122-25, \mathrm{MCZ} 56309,56312$, a series of adult specimens collected by various collectors on different dates from the type locality. LACM 92067, 92072, 92090 , 92105 were karyotyped.

Diagnosis.-The only species in which some or all individuals share the combination of a distinct light posterior thigh stripe, posterior surface of the tarsus smooth and sole of foot with prominent white tubercles are amazonicus, mystaceus, and notoaktites. Most individuals of mystaceus also have white tubercles on the posterior surface of the tarsus. Some individuals of mystaceus have a mid-dorsal light stripe, no amazonicus have a mid-dorsal light stripe. Leptodactylus amazonicus are found throughout the greater Amazon Basin, mystaceus occur along the east coast of Brasil from Bahia to Rio de Janeiro. Some individuals of notoaktites


Figure 33. Lateral view and mouthparts of tadpole of Leptodactylus albilabris. Semidiagrammatic figure based on specimens from
Puerto Rico.


Figure 34. Distribution map of Leptodactylus albilabris (squares), amazonicus (triangles) and bufonius (Xs).
lack white tubercles on the sole of the foot; some notoaktites have light mid-dorsal stripes; the distribution of notoaktites is southern coastal Brasil from São Paulo to Santa Catarina.

Description of Holotype.-Snout subelliptical from above, acute in profile; canthus rostralis indistinct; loreal slightly concave; tympanum distinct, greatest diameter about $1 / 2$ eye diameter; vomerine teeth in arched series posterior to choanae; vocal slits present; pair of external lateral vocal folds; finger lengths in order of decreasing size $\mathrm{I} \simeq \mathrm{III}>\mathrm{II} \simeq \mathrm{IV}$, first finger much longer than second; inner metacarpal tubercle ovoid, flat, smaller than large, flat, rounded outer metacarpal tubercle; no nuptial asperities; dorsum smooth; one pair of dorsolateral folds
from back of eye to groin; supratympanic fold from eye to humerus; ventral surfaces smooth; belly disk fold well developed; toe tips not expanded; toes free, lacking fringe or web; subarticular tubercles moderately well developed; outer metatarsal tubercle small, round, about $1 / 4$ large, ovoid inner metatarsal tubercle; tarsal fold indistinct; no metatarsal fold; posterior surface of tarsus smooth; sole of foot with several large, white tubercles.

SVL 50.3 mm , head length 19.7 mm , head width 18.0 mm , interorbital distance 3.1 mm , eye-nostril distance 4.6 mm , femur 24.5 mm , tibia 26.5 mm , foot 26.8 mm .

Dorsum brown with darker brown markings including interorbital bar and 2 mid-dorsal chevrons; dorsolateral


Figure 35. Dorsal view of the holotype of Leptodactylus amazonicus.
folds outlined with dark and light pin stripes; light upper lip stripe indistinct; limbs barred; underside of chin dark edged; belly light; posterior surface of thigh blotched above, dark with distinct light longitudinal stripe below.

Etymology-Named in reference to the distribution pattern characteristic of the species.

Remark.-This is the species referred to as "northern mystaceus'" in the morphological analysis.

Adult Characteristics ( $N=148$ ).-Dorsum spotted, spots very rarely fused (fig. 1, A, B, C); no light middorsal stripe, light upper lip stripe usually distinct ( $56 \%$ ) (fig. 57), often indistinct (44\%), more females with distinct lip stripes than males ( $X^{2}=8.66, P=.003$ ); no dark suborbital bar; light stripe on posterior face of thigh almost always distinct (93\%), rarely indistinct (7\%), distinctiveness not sexually dimorphic ( $X^{2}=1.81, P=$ .18); tibia barred; usually 2 or 4 well defined dorsolateral folds; dorsal surface of tibia lacking white tubercles; posterior surface of tarsus almost always lacking white tubercles ( $99 \%$ ), very rarely present ( $1 \%$ ), presence not sexually dimorphic ( $X^{2}=.001, P=.98$ ); sole of foot with many or scattered white tubercles ( $100 \%$ ); male SVL $47.4 \pm 2.4 \mathrm{~mm}$, female $50.2 \pm 2.6 \mathrm{~mm}$, females larger than males ( $\mathrm{F}_{1}, 146=47.8, \mathrm{P}<.001$ ); male head length/SVL ratio $.386 \pm .011$, female $.380 \pm .013$, male head longer than female ( $\mathrm{F}_{1,146}=7.38, .005<$ $P<.01$ ); male head width/SVL ratio $.352 \pm .013$, female $.347 \pm .013$, male head broader than female ( $\mathrm{F}_{1,146}=4.34, .025<P<.05$ ); male femur/SVL ratio $.433 \pm .027$, female $.447 \pm .033$, not sexually dimorphic ( $\mathrm{F}_{1,146}=.43, P>.05$ ); male tibia/SVL ratio
$.515 \pm .020$, female $.526 \pm .026$, female tibia longer than male ( $\mathrm{F}_{1,146}=7.82, .005<P<.01$ ); male foot/ SVL ratio $.532 \pm .021$, female $.539 \pm .026$, not sexually dimorphic ( $\mathrm{F}_{1,146}=2.96, P>.05$ ).
Larval Characteristics.-Eye diameter $9-14 \%$ headbody length; oral disk width $22-26 \%$ head-body length; anterior oral papilla gap $50-70 \%$ oral disk width; $50-$ 70 denticles on one side of split tooth row anterior to beak; head-body length $33-40 \%$ total length; total length, stage $40,36.2 \mathrm{~mm}$ (fig. 36).
Mating Call.-Dominant frequency modulated from $700-1400 \mathrm{hz}$ (fig. 37); call without harmonic structure; call pulsatile, about 15 pulses per note (fig. 38); note duration about 0.2 s ; note repetition rate 1.78 per second.

Karyotype.-Diploid number 22, 5 pair median, 3 pair submedian, 3 pair subterminal (Bogart 1974) or 4 pair median, 4 pair submedian, 3 pair subterminal (Heyer and Diment 1974); secondary constriction in chromosome pair 8.
Distribution.-Throughout the greater Amazon Basin, Guianas, northem Atlantic forest, and cerrados bordering the Amazon Basin (fig. 34).

BOLIVIA. BENI: Boca del Baures, AMNH 79096; Reyes, UMMZ 64107; Rurrenabaque, UMMZ 64109.

SANTA CRUZ: Buenavista, CM 3885, 3967, MCZ 12896, UMMZ 63832 (4), 64025 (3), 66478, 66481, 66492.
BRASIL. ALAGOAS: Usina Sinimbu, S. Miguel, WCAB 2775.

AMAPÁ: Serra do Navio, LACM 44711-12, WCAB 2308, 35229-231.

AMAZONAS: Rio Canabari, Rio Tucano, WCAB 34225; Ducke Reserve, KU 129942; Prainha, Aripuanã R., MZUSP 36886.

GOIÁS: Flôres, MZUSP 25348, USNM 121271; Mun. de Aliança, Jatobasinho, MNRio 2699 (5); mouth São Domingos River, MZUSP 25347.

MARANHÃO: Aldeia Araçu, igarapé Gurupi-Una, MZUSP 24954, 24958; Aldeia Javariuhu, igarapé Gurupi-Una, MZUSP 25014; Carolina, WCAB 6692-93.

MATO GROSSO: Chapada dos Guimarães, WCAB 15382; Pimentel River, Serra do Roncador, MZUSP 1358; mouth Tapirapés River, MZUSP 25276.

MINAS GERAIS: Uberlândia, MZUSP 12136.
PARÂ: As Pedras, Cuminá-Miri River, MZUSP 28400; Belém, MNRio 1470, MZUSP 11478; Belém-Brasilia road, km 43, MZUSP 24946; Benevides, KU 127397; Cachimbo, MZUSP 21835, 21876; IPEAN, KU 127395-96; Jacareacanga, WCAB 6645-46.

PERNAMBUCO: Agua Azal, Vicência, MZUSP 36837; Bonito, UMMZ 132459-460; Iguarassú, MNRio 2363; Recife, MZUSP 25029.

RONDÔNIA: Lg. Marmelo (near Abunã), WCAB 9840.
RORAIMA: Serra de Parima, MZUSP 24937-941.
COLOMBIA. CAQUETÁ: Florencia, USNM 147039-047.
META: Acacías, USNM 17048-050; Caño Losada, upper Río Guayabero, USNM 146346, 150488-89; Villavicencio, USNM 146433-35, 147396.

PUTUMAYO: about 7 km SE Mocoa, near Río Pepino, AMNH 84862-64; Santa Rosa de los Kofanes, about 30 minutes walking below San Antonio del Guamés, along middle course of Río Guamés, tributary of Upper Putumayo, CM 50647-650.

VAUPÉS: Río Ariari and Río Guaviare, UTA 2777, 2780, 3717, 3938-941, 3954.


FIGURE 36. Lateral view and mouthparts of tadpole of Leptodactylus amazonicus. Semidiagrammatic figure based on specimens from Santa Cecilia, Ecuador.


Figure 37. Sonagram of the mating call of Leptodactylus amazonicus, narrow band filter. Vertical scale marks at 1000 hz intervals.


FIGURE 38. Strip chart record of the mating call of the holotype of Leptodactylus amazonicus. Line equals 0.01 s .

VICHADA: Anaben, UPR 91.
ECUADOR. NAPO: Limoncocha, LACM 92067-095, 92098-2125, MCZ 56308-317; Santa Cecilia, MCZ 5632021, UMMZ 129282.

FRENCH GUIANA. Antécume-Pata (Haut Maroni), LES 1210-11; Cacao (Riv. Comté), LES 216-18; Embouchure, Haut Oyapock, Riv. Yaroupi, LES 1501; Saut Verdun et Grigel (Ouaque, Haut Maroni), LES 1447-49; Trois-Sauts (Haut Oyapock), LES 177; Village Pina (Haut Oyapock), LES 115054, 1161; Village Zidok (Haut Oyapock), LES 1285-89, 1291, 11204-05, 11207.

GUYANA. Issano, UMMZ 83584 (2); Kalacoon, Mazaruni River, AMNH 3988; Kartabo, AMNH 39593-94, 39651-56, 39658-59, 39661-62, 39665-68, 39672, CM 4065, UMMZ 83583 (3), USNM 118058; Kurupung, Upper Mazaruni Dist., UMMZ 83585; Rupununi, $N$ of Acarahy Mts., W of New River, KU 69675-680, 69701-07; Shudi-kar-wau, AMNH 49250, 53488 (7); Yacarascine, LES 1494.

PERU. LORETO: Valley of Río Huallaga, AMNH 43198; Tapiche-Río Utoquinia, AMNH 43225, 43381.

SAN MARTÍN: Tocache Nuevo, Río Huallaga, USNM 195998-99.

SURINAM. Brokolonko Loksihattie, Saramacca, FMNH 134736-38, 134740; Kaiserberg Airstrip, Zuid River, FMNH 128826, 128841-42, 128924; Mataway, CM 44267-270; Paramaribo, USNM 158961-62.

VENEZUELA. AMAZONAS: Capibara, 106 km SW Esmeralda, Brazo Casiquiare, 130 m , USNM field 19585, 19588; La Culebra, UPR 3048; Monte Marahuaca, UPR 98-99.

ARAGUA: Maracay, near Rancho Grande, AMNH 7066566.

## Leptodactilus bufonius Boulenger 1894

Leptodactylus bufonius Boulenger, 1894: 348. (Type locality, Paraguay, Asunción. Lectotype BMNH 1947.2.17.72, female.)

Diagnosis.-The species with a combination of no distinct light stripe on the posterior face of the thigh and the posterior surface of the tarsus covered with obvious white tubercles are: bufonius, labrosus, mystacinus, troglodytes, and ventrimaculatus. A pair of distinct dorsolateral folds (indicated at least in color pattern in poorly preserved specimens) characteristic of labrosus, mystacinus, and ventrimaculatus, distinguish them from bufonius, which lacks well defined dorsolateral folds. The sole of the foot is covered with white tubercles in troglodytes, the sole of the foot is almost always smooth in bufonius. Leptodactylus bufonius has a Chacoan distribution, troglodytes occurs in NE Brasil.

Adult Characteristics $(N=139)$.-Dorsum spotted or blotched (fig. 1, C, E, F, L, M); mid-dorsal light stripe absent $(100 \%)$; light lip stripe absent $(100 \%)$; dark suborbital bar present; light stripe on posterior face of thigh absent ( $100 \%$ ); tibia barred; a pair of indistinct dorsolateral folds present or (usually) absent; dorsal sur-
face of tibia with white tubercles; posterior surface of tarsus with many or scattered white tubercles ( $100 \%$ ); sole of foot rarely with white tubercles ( $6 \%$ ), usually absent ( $94 \%$ ), presence not sexually dimorphic ( $X^{2}=$ $0.08, P=.77$ ); male SVL $51.6 \pm 2.0 \mathrm{~mm}$, female 53.6 $\pm 2.3 \mathrm{~mm}$, females larger ( $\mathrm{F}_{1,137}=29.86, P<.001$ ); male head length/SVL ratio $.366 \pm .011$, female .361 $\pm .011$, male head longer ( $\mathrm{F}_{1,137}=6.58, .01<P<$ .025 ); male head width/SVL ratio $.346 \pm .011$, female $.341 \pm .012$, not sexually dimorphic ( $\mathrm{F}_{1.137}=3.89$, $.10>P>.05$ ); male femur/SVL ratio $.374 \pm .024$, female $.377 \pm .017$, not sexually dimorphic ( $\mathrm{F}_{1,137}=$ $0.82, P>.05$ ); male tibia/SVL ratio $.400 \pm .018$, female $.398 \pm .019$, not sexually dimorphic ( $\mathrm{F}_{1,137}=$ $0.41, P>.05$ ); male foot/SVL ratio $.381 \pm .019$, female $.382 \pm .020$, not sexually dimorphic ( $\mathrm{F}_{1,137}=$ $0.08, P>.05$ ).

Larval Characteristics.-Available materials are insufficient for an adequate description.
Mating Call.-Dominant frequency modulated from 1000 to 2000 hz (fig. 39); note lacking harmonic structure (fig. 40); note either non-pulsed (fig. 40) or partially pulsed (Straughan and Heyer 1976, fig. 1); note duration $0.2 \mathrm{~s} ; 1.25$ notes/second.

Karyotype.—Diploid number 22, 7 pair median, 2 pair submedian, and 2 pair subterminal (Bogart, 1974) or 6 pair median, 2 pair submedian, and 3 pair subterminal (Heyer and Diment, 1974); secondary constriction in chromosome pair 8.
Distribution.-Found throughout the Gran Chaco ąd surrounding areas (fig. 34).

ARGENTINA. CHACO: Laguna Limpia, IML 562; Río Teuco, Estancia La Fidelidad, IML 122 (18); Roque Sáenz Peña, IML 588 (5).

FORMOSA: Ingeniero Juárez, IML 980 (55), LACM 91935, 91945-47, 91959-962, MCZ 35584; Bañados del Río Teuco, Depto. Bermejo, IML 1050 (29); La Florencia, Teuquito, IML 968 (9); Palma Sola, IML 1057.

JUJUY: Valle Grande, IML 1790.
LA RIOJA: Between Olta and Chamical, MCZ 33970-79.
SALTA: Abra Grande-Orán, IML 1696 (5); Aguaray, IML 560 (2); Embarcación, LACM 91925-27, 91929-932, 91934, 91937-940, 91942-44, 91948-950, 91953-58, 91963-65; Hickmann, IML 442 (39), 841 (128), 844 (64), 981 (27), KU 128857-58, MCZ 35336-345, USNM 159753; La Unión, IML 1755; Pocitos, MACN 4495; Saucelito, IML 1517.

SANTIAGO DEL ESTERO: Huyapampa, IML 819 (7), MCZ 32766-68; 46 km S Loreto, MCZ 33710-13; Ojo de Agua, IML 1139; Simbol Bajo, MACN 4999.

TUCUMAN: Los Gómez, IML 634 (3); Río Urueña (nr. Salta), IML 1761 (3); Tucumán, MZUSP 13783-84.

BOLIVIA. CHUQUISACA: 30 km SE Carandaiti, LACM 37705-06.


Figure 39. Sonagram of the mating call of Leptodactylus bufonius, narrow band filter. Vertical scale marks at 1000 hz intervals. Horizontal scale mark at 1 s . Specimen from Argentina, Embarcación, air temperature $24.8^{\circ} \mathrm{C}$ (LACM tape and specimen field number WRH 1411).


Figure 40. Strip chart record of the mating call of Leptodactylus bufonius. Line equals 0.01 s . See legend of Figure 39 for specimen data.

SANTA CRUZ: El Carmen, CM 36159-160, 36187, MCZ 29962-66; San José de Chiquitos, CM 36229, MCZ $29967-$ 973, MZUSP 21340-41; Parapetí, KU 92902-04.

BRASIL. MATO GROSSO: Carandázal, MZUSP 127; Fazenda Cruzeiro, Aquidauana, MZUSP 16201.

PARAGUAY. Colonia Nueva Italia, MCZ 25806; Río Pilcomayo, MCZ 25819-821.

## Leptodactylus elenae new species

Figure 41
Holotype: LACM 92096, an adult female from Argentina; Salta, Embarcación. Collected by Keith A. Berven, Laura M. Heyer, Miriam H. Heyer, and W. Ronald Heyer on 4 January 1972.

Diagnosis.-The species sharing the combination of a distinct light stripe on the posterior surface of the thigh and obvious white tubercles on the sole of the foot in some or all individuals are albilabris, amazonicus, elenae, fragilis, fuscus, latinasus, mystaceus, notoaktites. Leptodactylus elenae has no white tubercles on the dorsal surface of the tibia, differing from albilabris, fragilis, latinasus, and mystaceus. Leptodactylus elenae has 2 or 4 distinct (at least indicated in color pattern) dorsolateral folds, fuscus has 6. Leptodactylus elenae usually has white tubercles on the posterior tarsus, the tarsus is smooth in amazonicus and notoaktites. Lepto-


Figure 41. Dorsal view of the holotype of Leptodactylus elenae.
dactylus elenae has a Chacoan distribution, L. notoaktites a SE Brasilian distribution.

Description of Holotype.-Snout subelliptical from above, rounded-acute in profile; canthus rostralis slightly
obtuse; loreal slightly convex; tympanum distinct, greatest diameter just more than $1 / 2$ eye diameter; vomerine teeth in slightly arched series posterior to choanae; finger lengths in order of decreasing length $\mathrm{I} \simeq \mathrm{III}>\mathrm{II} \simeq$ IV, I >> II; inner metacarpal tubercle flat, oval, smaller than flat, heart shaped outer metacarpal tubercle; dorsal surfaces smooth; 2 pair of dorsolateral folds (indicated by color pattern); ventral texture smooth; belly disk fold distinct; toe tips just wider than adjacent portion of toes; toes free, lacking fringe or web; subarticular tubercles moderately distinct; outer metatarsal tubercle small, round, about $1 / 3$ oval inner metatarsal tubercle; tarsal fold extends about $3 / 4$ length of tarsus; no metatarsal fold; posterior surface of tarsus with scattered, barely visible light tubercles; sole of foot with many distinct white tubercles.

SVL 43.5 mm , head length 15.6 mm , head width 14.3 mm , interorbital distance 2.6 mm , eye-nostril distance 4.2 mm , femur 18.4 mm , tibia 20.6 mm , foot 22.1 mm .

Dorsum tan with darker tan markings consisting of an irregular interorbital bar and 2 dorsal blotches; inner broken light pin stripe bordered by outer irregular dark brown stripe along dorso-lateral fold from back of eye to groin; broken light pin stripe along lateral fold; dark brown canthal stripe from tip of snout across upper tympanum to humeral region; distinct light upper lip stripe; limbs faintly barred; venter immaculate; tarsal fold highlighted by a white line; posterior surface of thigh blotched with distinct light longitudinal stripe.
Etymology.-Named for my daughter, Elena, who shares my enthusiasm for encountering frogs in nature.
Remarks.-This is the species referred to as "southern mystaceus" in the morphological analysis.

Adult Characteristics ( $N=43$ ).-Dorsum spotted, spots rarely fused (fig. 1, A, B, C); no light mid-dorsal stripe; light lip stripe usually distinct ( $77 \%$ ), sometimes indistinct ( $23 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=.77, P=.38$ ); dark suborbital bar absent; light stripe on posterior face of thigh distinct ( $100 \%$ ); tibia barred; usually 4 well defined dorsolateral folds; no white tubercles on dorsal surface of tibia; many or scattered white tubercles usually present on posterior surface of tarsus ( $91 \%$ ), sometimes lacking ( $9 \%$ ), presence not sexually dimorphic ( $X^{2}=3.16, P=.08$ ); sole of foot with many or scattered white tubercles ( $100 \%$ ), male SVL $42.7 \pm 2.5 \mathrm{~mm}$, female $42.8 \pm 3.1 \mathrm{~mm}$, not sexually dimorphic ( $\mathrm{F}_{1,41}=.02, P>.05$ ); male head length/SVL ratio $.375 \pm .011$, female $.374 \pm .009$, not sexually dimorphic ( $\mathrm{F}_{1,41}=.15, P>.05$ ); male head width/SVL ratio $.338 \pm .013$, female $.336 \pm .019$, not sexually dimorphic ( $\mathrm{F}_{1,41}=.20, P>.05$ ); male femur/ SVL ratio $.406 \pm .021$, female $.404 \pm .034$, not sexually dimorphic ( $\mathrm{F}_{\mathrm{L}, 41}=.08, P>.05$ ); male tibia/SVL ratio $.468 \pm .020$, female $.470 \pm .030$, not sexually dimorphic ( $\mathrm{F}_{1,41}=.06, P>.05$ ); male foot/SVL ratio $.501 \pm .027$, female $.491 \pm .030$, not sexually dimorphic ( $\mathrm{F}_{1,41}=.95, P>.05$ ).

Larval Characteristics.-Larvae unknown.
Mating Call.-Barrio (1965) described and figured the call (as L. mystaceus from Villa Angela, Chaco). The fundamental frequency modulates from 700-1500 hz , note duration 0.3 s , call repetition rate 2 notes per second.

Karyotype.—Diploid number 22, no terminal pairs fno further interpretations can be made from the karyotype prepared from LACM 92097, from the type locality).
Distribution.-Found in the Gran Chaco and adjacent areas to central Brasil and Río Huallaga, Peru (fig. 42).

ARGENTINA. JUJUY: Ruta Yuto-Ledesma, near Arroyo Quemado, IML 1275, Ruta Yuto-Ledesma, 7 km from bifurcation, IML 1274.

SALTA: Campo Aguaray, IML 1472 (2); El Saucelito, 50 km S Orán, IML 1624 (5) near Embarcación, LACM 9202627, 92127; Río Pescado, IML 1401 (9).

BOLIVIA. BENI: Lake Rogoagua, UMMZ 64108; Río Mamoré, about 10 km W San Pedro, AMNH 79095. LA PAZ: Ixiamas, UMMZ 64106 (2).
SANTA CRUZ: Buenavista, CM 3889, 4345, 4352, 4432, UMMZ 63832 (4), 66491,66543 (2); El Carmen, MCZ 29985; San José de Chiquitos, CM 36118, MCZ 29987-88.

BRASIL. MATO GROSSO: Carandázal, MZUSP 139; Corumbá, CM 36162; Rosario Oeste, WCAB 15628-632; Salobra, USNM 133011-12; Santo Antônio do Leverger, WCAB 1510205; Parque Indigena do Xingu, Posto Diauarum, MZUSP 49543, WCAB 37186.
PERU. SAN MARTÍN: Tocache Nuevo, Río Huallaga, USNM 195998-99.

## LEPTODACTYLUS FRAGILIS (Brocchi) 1877

Cystignathus fragilis Brocchi 1877:182-184. (Type locality, Mexico, Tehuantepec. Holotype Paris Museum 6316, female.)

Remark.-This is the species referred to as labialis in the analysis section and in the herpetological literature for the past 30 years.

Diagnosis.-The species demonstrating a combination of a distinct light stripe on the posterior surface of the thigh, and obvious white tubercles on the posterior surface of the tarsus and sole of foot in some or all individuals are albilabris, elenae, fragilis, latinasus, and mystaceus. Leptodactylus elenae has a smooth dorsal tibial surface, the dorsal surface of the tibia is covered with white tubercles in fragilis. Leptodactylus albilabris and mystaceus have distinct dorsolateral folds (indicated by color pattern in poorly preserved specimens), fragilis has indistinct dorsolateral folds or lacks them. Leptodactylus fragilis and latinasus have considerable morphological and color pattern overlap (fig. 43), fragilis being a slightly larger species (maximum male SVL 43 mm , female 43.6 mm ) than latinasus (maximum male SVL 37.9 mm , female 36.3 mm ). Leptodactylus fragilis has a Middle American and north coast South American distribution, L. latinasus has a southern South American distribution.

Adult Characteristics ( $N=591$ ).-Dorsum spotted or blotched, blotches rarely confluent (fig. 1, A, B, C,


FIGURE 42. Distribution map of Leptodactylus elenae (squares) and fragilis (triangles).

D, E); mid-dorsal light stripe absent; light lip stripe usually indistinct ( $97 \%$ ), rarely distinct ( $3 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=.01, P=.94$ ); dark suborbital bar absent; light stripe on posterior fade of thigh usually very distinct ( $66 \%$ ), often moderately distinct ( $33 \%$ ), rarely absent ( $1 \%$ ), expression not sexually dimorphic ( $X^{2}=2.70, P=.26$ ); tibia barred; dorsolateral folds usually indistinct, 2 or 4 present when visible; dorsal surface of tibia usually covered with many white tubercles, sometimes scattered with white tubercles; posterior surface of tarsus with many white tubercles $(89 \%)$, rarely absent ( $11 \%$ ), presence of tubercles not sexually dimorphic ( $X^{2}=2.50, P=.11$ ); sole of foot always with many white tubercles ( $100 \%$ ); male SVL $34.7 \pm 2.9 \mathrm{~mm}$, female $34.2 \pm 2.6 \mathrm{~mm}$, not
sexually dimorphic ( $\mathrm{F}_{1,589}=3.23, P>.05$ ); male head length/SVL ratio $.379 \pm .017$, female $.376 \pm .013$, male head longer ( $\mathrm{F}_{1,589}=9.22, .001<P<.005$ ); male head width/SVL ratio $.336 \pm .019$, female .333 $\pm .016$, male head wider ( $\mathrm{F}_{1,589}=4.64, .025<P<$ .05 ); male femur/SVL ratio $.389 \pm .028$, female .399 $\pm .025$, female femur longer ( $\mathrm{F}_{1}, 589=22.19, P<$ .001 ); male tibia/SVL ratio $.451 \pm .026$, female .456 $\pm .026$, female tibia longer ( $\mathrm{F}_{1,589}=44.43, P<.001$ ); male foot $/ \mathrm{SVL}$ ratio $.494 \pm .033$, female $.502 \pm .030$, female foot longer ( $\mathrm{F}_{1,589}=10.03, .001<P<.005$ ).
Larval Characteristics.-Eye diameter $12-16 \%$ headbody length; oral disk width $17-22 \%$ head-body length; oral papilla gap $53-67 \%$ oral disk width $46-101$ denticles in one side of split tooth row anterior to beak;


Figure 43. Dorsal views of Leptodactylus fragilis (left, KU 116833) and latinasus (right, LACM 92039).
head-body length $31-40 \%$ total length; total length, stage $41,41 \mathrm{~mm}$ (Heyer 1970b, figs. 7, 12, 17).

Mating Call.-Dominant frequency modulates from $600-1200 \mathrm{hz}$ (Texas) to $1000-2200 \mathrm{hz}$ (Panama); call lacking harmonic structure; call with a pulsatile and nonpulsatile portion; pulsatile portion .170 s duration immediately followed by non-pulsatile portion of .023 s duration; note repetition rate 1.5 per second (Straughan and Heyer 1976, fig. 3).

Karyotype .-Bogart (1974) described the karyotype as diploid number 22; 7 pair median, 1 pair submedian, 3 pair subterminal; secondary constriction in chromosome pair 8.

Distribution.-From southernmost Texas throughout lowland Middle America along the north coast of South America as far as Venezuela, including the Magdalena Valley of Colombia (fig. 42).

BELIZE. Belize, FMNH 4392, 4398, 4732, UMMZ 124744 (3); Corozal, (A. Ross collection numbers) 2975-981, 2984, 2986-88, 2991-92, 2998-3036, 3046-058; Gallon Jug, MCZ

37863, 37873-74; Kates Lagoon, FMNH 49060; Manatee, FMNH 4263; Monkey River, Swazey Branch, MCZ.37867872; Otro Benque, USNM 194891-99, 194931; Tower Hill, USNM 167739, 194081-83.

COLOMBIA. ANTIOQUIA: Casabe, USNM 147079; Chigorodó, near Turbo, USNM 153915-17; Golfo de Urabá, N Turbo, LACM 50198, USNM 150491-0515; Nechí, FMNH 54572, 54575-76.

BOLIVAR: Cartagena, Bocagrande, CM 50603; Isla Fuerte, FMNH 74937, USNM 150516-19.

CHOCÓ: Atrato, Sautatá, FMNH 74920 (2).
CUNDINAMARCA: Beltrán, USNM 145743.
GUAJRA: near Pájaro, USNM 151306.
MAGDALENA: Fundación, UMMZ 48489-492, 4849599, 48503-04, 48509, 48511, USNM 102409.

NORTE DE SANTANDER: Catatumbo, USNM 145088092; Río Zulia, USNM 147071, 147074-75.

SANTANDER: El Centro, FMNH 81760, USNM 144839_ 842, 147091-92; Río Zulia, USNM 147051-52.

TOLIMA: Espinal, MCZ 15065-66, 15069-070, 15072-75 (2); Mariquita, FMNH 81835, 81838, USNM 150516-19.

COSTA RICA. ALAJUELA: Los Chiles, CRE 7215, 7217 (2).

GUANACASTE: Arenal, CRE 6251 (2); 2.4 mi N. Bagaces, CRE 8193; near Cañas, CRE 2902 (2), 8009, 8181; 50

km S Cañas, CRE 249; Finca Jiménez, CRE 3088 (2), 3091, 3094 (2), 3095 (8), 3097, 3099 (2); $7.6 \mathrm{mi} \mathrm{S} \mathrm{La} \mathrm{Cruz}$, 8091; near Liberia, CRE 107, 2888, 8015 (3), 8140, 8153, 8162 (2), 8163-64; 21.3 mi SW Liberia, CRE $8215 ; 35.5 \mathrm{mi}$ N Liberia, CRE 8196; between Liberia and Cañas, USNM 192558; near Nicoya, CRE 8229-230; near Playa del Coco, CRE 6504, 6512-14, 6516 (2), 8012 (4); near Santa Cruz, CRE 8218; Hacienda Taboga, CRE 6297, 6439.

LIMÓN: Los Diamantes, FMNH 176916.
PUNTARENAS: near Barranca River, CRE 254, 739 (4); base of Peninsula, CRE 253 (4); Coto, km 47 on rail from Golfito, CRE 176-78, 180 (5); Esterillos Oeste, 15 km SE Jacó, CRE 2873 (3); Golfito, CRE 7231; Río Grande de Tárcoles, 6.1 km NE mouth, SW Orotina, CRE 817; Villa Neily, 75 m , CRE 8039 (6); 13.6 mi NW Villa Neily, CRE 8005.

EL SALVADOR. LA PAZ: Los Blancos, FMNH 6512122.

MORAZÁN Divisadero, USNM 73285-86.
GUATEMALA. EL PETEN: near La Libertad, CM 13020, MCZ 21455; Pacomon, USNM 71333; near Poptún, UMMZ 117989 (7), 124377 (8), 124378 (2), 124379 (4), 124380 (2); Tikal UMMZ 117988.

RETALHULEU: Hacienda Casa Blanca, UMMZ 107886, 107887 (7).

HONDURAS. CHOLUTECA: 2 mi NE Choluteca, LACM 60525-28.

COMAYAGUA: $2-3 \mathrm{mi} \mathrm{S}$ Comayagua, LACM 47522, 47532, 3112 mi WSW Siguatepeque, LACM 47531.

COPÁN: Copán, FMNH 40864; 6 mi SW La Florida, LACM 47523.

CORTÉS: Lake Ticamaya, E San Pedro, FMNH 4656-660; Lake Yojoa, MCZ 26407-09 (8); 2112-41/2 mi ENE Villanueva, LACM 47524-530.

EL PARAISO: 1 km N Santa Maria, LACM 45084-85.
FRANCISCO MORAZÁN: 8.6 mi NW Comayaguela, LACM 60529-532; El Hatillo, 1400 m, LACM 72074; El Picacho, Tegucigalpa, MCZ 28887-898; El Zamorano, 2700 ft , LACM 39757; near Río Yeguare, MCZ 25964-69 (8), 26466 (30).

GRACIAS A DIOS: Ahuás, LACM 45245; Tansín, 15 km NW Puerto Lempira, LACM 47511-16.

OLANCHO: 8.6 mi E Catacamas, LACM 45101-03; 7.6 mi SW Juticalpa, LACM 45238-240; $1 / 2 \mathrm{mi}$ SE San José de Río Tinto, LACM 45178, 45194-99.

SANTA BÁRBARA: Quimistán, USNM 128058-59.
VALLE: near Río Guascorán, LACM 45064, 47520-21; 5.5 mi E San Lorenzo, LACM 48374.

YORO: 1.5 km W Olanchito, LACM 47517-19; Subirana Valley, FMNH 21821-24, MCZ 21260-69 (4).

MEXICO. CAMPECHE: Balchacah, FMNH 108273, 108276, 108279-280, 108282-87, 108289-291, 108298-8301, 108306, 108310-11, 108315-16, 108319, 108322-27, 108329-330, 108333-35, 108337, 108340-41, 108344, 108355, 108357, 108362, 108364-371, 108376, 108380-81 108385, 108388, 108390-92, 108395-98, 108401, 108408, 108410-11, 10841315, 108422-24, 108426; Champotón, MCZ 21452; Escárcega, 5 mi W 'EI Tormento,' CM 40106-07; Matamoros, FMNH 38588; Pital, FMNH 108271, 108312, 108348, 108352, 108361, 108373, 108383, 108409, 108421; Tres Brazos, FMNH 108288, 108320.

CHIAPAS: near Asunción, FMNH 108443; El Censo, MCZ 28255; El Real, $600 \mathrm{~m}, \mathrm{MCZ} 28271$ (4); near San Ricardo, FMNH 108436, 108444-45, 108461, 108467; near Tapachula, FMNH 108435; near Tonda, FMNH 108460; near Tuxtla Gutiérrez, FMNH 108449.

COLIMA: 7.5 mi SW Colima, LACM 37265-66, 37430_ 34.

GUERRERO: 2 mi S Garropata, 44 mi S Chilpancingo,

FMNH 108454; near Palo Blanco, S of Chilpancingo, FMNH 108427, 108430, 108432, 108446, 108452, 108459, 108462, 108464-65.

MICHOACÁN: Apatzingán, FMNH 38806-817; 11.7 mi S Cuatro Caminos, LACM 37049, 37429; Hacienda El Sabino, FMNH 108438, 108448, 108458, 108466.

MORELOS: near Antiguo, FMNH 108431, 108434, 108450, 108457, 108483.

OAXACA: Barrio, USNM 30241-42; Matías Romero, AMNH 52139-140 (3), 69508; Mixtequilla, AMNH 13922; Niltepec, CM 52739-742; 10 mi W Río Ostuta, FMNH 7242728; Tehuantepec, AMNH 65633-35, MCZ 15767, USNM 10018-19, 27765, 114229-231; Tolosa, AMNH 53608-09; Tuxtepec Soyaltepec, LACM 74753.

QUINTANA ROO: Isla Cozumel, 12 km SW San Miguel, CM 41315; Laguna Chacanacab, 86 km W Chetumal, CM 45231-32.

SAN LUIS POTOSÍ: El Salto Falls, 12 mi W Nuevo Morelos, UMMZ 99518 (7).

TABASCO: 2.5 mi NE Comalcalco, AMNH 60317; Encarnación, FMNH 106351, 108272, 108274-75, 108278, 108281, 108292-97, 108302-05, 108307, 108309, 108313-14, 10831718, 108321, 108328, 108331-32, 108336, 108338-39, 10834243, 108345-47, 108349-351, 108353, 108358-360, 108363, $108372,108374-75,108377-79,108382,108384,108386-$ 87, 108389, 108393-94, 108399-8400, 108402-07, 108412, 108416-18, 108420, 108425, 108456; Tenosique, USNM 114217-18; 43 mi N Villa Hermosa, USNM 192539.

TAMAULIPAS: Arroyo Los Almos, 3 mi SE Rio Grande City, FMNH 108429; Ciudad Victoria, N on Highway 101, LACM 64151; La Laguna Doña Ana, MCZ 24982-86 (2); between Monterrey and Ciudad Victoria, FMNH 108481; Ocampo, AMNH 62065-66; Pano Ayuctle, 5 mi NE Gómez Farías, UMMZ 98948, 102913, 110714 (4); Rancho Sta. Ana, MCZ 24973-77 (2); Río Corona, MCZ 24966-68; 3 mi W San Gerardo, UMMZ 110712 (2), 110713 (3); 10 mi E jct highways 80 and 85 to Tampico, LACM 65752; 10 mi N Victoria, FMNH 105279, 108442, 108451, 108453, 108472, 108477, 108479-480, 108482, 108487.

VERACRUZ: Hacienda La Oaxaqueña, 30 km S Jesús Carranza, on Coatzacoalcos River, AMNH 43926-29; Orizaba, USNM 16547; Potrero near Córdoba, USNM 32410-12; Potrero Viejo, FMNH 108277, 108468-69, 108471, USNM 114210-16; Río Chiquito at San Lorenzo, USNM 123528-29; near San Andrés Tuxtla, AMNH 69505, FMNH 108437; near San Gerónimo, FMNH 108433, 108440, 108455; Tierra Colorado, FMNH 108428; Veracruz, MCZ 4651-52.

YUCUTÁN: Chichén Itzá, FMNH 26962-64, 36567 (7); Cozumel Id., UMMZ 78548 (14); Dzibichaltún, Cenote Xcalah, CM 45230.

NICARAGUA. MATAGALPA: near Sebaco, LACM 943034.

PANAMA. CANAL ZONE: Alhajuela, CM 7391, 7407; Balboa, AMNH 41759, MCZ 17378, Fort Kobbe, USNM 193338 (3); Fort Sherman, MCZ 16017; Gatun, MCZ 35645; Laguna to Mendoza, Madden Dam road, AMNH 55398; 2 mi W Locona, KU 67949-951; Majanál, MCZ 10729; Rosseau, KU 67948; Summit, FMNH 22976-77, KU 115292-301.

CHIRIQUÍ: 3.3 mi E Concepción, AMNH 69723.
COCLE: 3.2 km W Aguadulce, 15 m , KU 115291; 1 km NE El Caño, 40 m , KU 115290 ; El Valle de Antón, AMNH 59585-87, 59589, 69721-22, FMNH 22985, KU 76519; near Penonomé, $30-70 \mathrm{~m}, \mathrm{KU} 115281-89$, 116833, 116835-37.

COLÓN: Achiote, 40 m , KU 76517-18; 3.5 km SE Puerto Pilón, 260 m , KU 116834.

LOS SANTOS: Los Santos, CM 43570; Tonosí, 40 m, KU 108617-624.

PANAMÁ: 14.4 km SSW Bejuco, 40 m , KU 115302; Capitán, near Chepo, USNM 192622; near La Chorrera, CM

23483; Nueva Gorgona, AMNH 69720; Panama City, MCZ 17580; about 8 mi W Playa Coronado, road to Laguna, FMNH 67893-95; near Puerto la Chorrera, UMMZ 95477; Tapia, AMNH 18932, 22838.

VERAGUAS: Cerro Lute, near Santa Fe, FMNH 678967908; Mojara, USNM 129843-44; Río Corobora, USNM 140877.

UNITED STATES. TEXAS: Cameron County; Brownsville, FMNH 27150 (12); Hidalgo County; 10 mi NW Edinburg, USNM 101143-44; 15 mi W Mission, UMMZ 98905; Starr County; 13 mi SE Rio Grande City, Arroyo El Salado, AMNH 46014, FMNH 107556-57.

VENEZUELA. APURE: Hato La Cuanota, 4 km W San Fernando de Apure, TCWC 45229-260, 45262-67, 45269278; Río Apure at San Fernando de Apure, UMMZ 85112.

FALCÓN: Boca de Yaracuy, 28 km WNW Puerto Cabello, USNM field 1543-48; 19 km NW Urama, km 40, USNM field 5176-182.

MONAGAS: 42 km SE Maturín, LACM 31380-81.
TRUJILLO: Sabana de Mendoza, UMMZ 57478-482.
YARACUY: San Felipe, BMNH 1973.2274-75.

## Leptodactylus fuscus Schneider 1799

Rana fusca Schneider 1799:130-131. (Type locality not specified. Neotype Paris Museum 680, male [lectotype of Rana typhonia Daudin 1803 and Cystignathus typhonius Duméril and Bibron 1841]).
Rana typhonia Daudin 1803:55-56, plate 17, fig. 3. (Type locality, Surinam. Lectotype Paris Museum 680, male.)
Rana sibilatrix Wied-Neuwied 1824: fig. 2. (Type locality originally undesignated, designated by Müller (1927) as Vila Viçosa, nio Peruipé. Type material apparently lost.)
Cystignathus typhonius Duméril and Bibron 1841:402-404. (Type locality, French Guiana and Surinam. Lectotype Paris Museum 680, male.)
Cystignathus schomburgkii Troschel 1848:659. (Type locality, British Guiana. Type material apparently lost.)
Leptodactylus raniformis Werner 1899:479-480. (Type locality, Colombia; Llanos, Río Meta. Holotype II Zoologisches Institut und Museum der Universität, Göttingen, no number, male.)
Leptodactylus gualambensis Gallardo 1964:46-50, plate 2, fig. 1. (Type locality, Argentina; Salta, Urundel, 43 km W Orán, Río Santa María. Holotype MACN 9752, male.)

Diagnosis. -The species having a combination of a light stripe on the posterior surface of the thigh and 6 distinct dorsolateral folds (almost always recognizable in fuscus) in some or all individuals arefuscus, geminus, gracilis, laurae, longirostris, marambaiae, mystaceus, notoaktites, and poecilochilus. Of these, only fuscus has individuals that have 6 dorsolateral folds without a light mid-dorsal stripe; in all of the other species, the individuals with 6 dorsolateral folds also have a light middorsal stripe (this feature allows positive identification when series of specimens are available). Individuals of L. fuscus rarely have distinct white tubercles on the sole of the foot and posterior surface of the tarsus, but small light spots are present on these surfaces indicating the presence of weakly developed tubercles. The posterior surface of the tarsus and sole of foot are smooth and uniform in coloration in geminus, gracilis, laurae, longirostris, marambaiae, and poecilochilus. Leptodactylus mystaceus usually has well developed, distinct white
tubercles on the posterior surface of the tarsus and sole of foot. Leptodactylus notoaktites has a smooth posterior surface of the tarsus.

Adult Characteristics ( $N=392$ ). -Dorsum spotted or blotched (fig. 1, C, D, E, F, G, H, I); mid-dorsal light stripe sometimes present ( $20 \%$ of individuals), presence not sexually dimorphic ( $X^{2}=1.98, P=.15$ ); light lip stripe usually indistinct $(81 \%)$, sometimes distinct ( $19 \%$ ), more females with distinct light lip stripes than males $\left(X^{2}=19.18, \ddot{P}<.001\right)$; dark suborbital bar absent; light stripe on posterior face of thigh usually very distinct ( $77 \%$ ), often moderately distinct ( $23 \%$ ), distinctiveness not sexually dimorphic $\left(X^{2}=3.67, P=\right.$ .06); tibia barred; usually 6 distinct dorsolateral folds; dorsal surface of tibia usually lacking white tubercles, few rarely present; posterior surface of tarsus rarely with distinct white tubercles ( $4 \%$ ), but scattered light spots associated with tubercles almost always present, presence of distinct tubercles not sexually dimorphic ( $X^{2}=$ $.39, P=.53$ ); sole of foot rarely with distinct white tubercles ( $8 \%$ ), but many to scattered light spots associated with tubercles almost always present, presence of distinct tubercles not sexually dimorphic $\left(X^{2}=.83, P\right.$ $=.36$ ); male SVL $42.8 \pm 4.0 \mathrm{~mm}$, female $43.6 \pm 4.4$ mm , not sexually dimorphic $\left(\mathrm{F}_{1,390}=3.47, P>.05\right)$; male head length/SVL ratio $.374 \pm .015$, female .376 $\pm .021$, not sexually dimorphic $\left(F_{1,390}=1.12, P>\right.$ .05 ); male head width/SVL ratio $.336 \pm .015$, female $.332 \pm .017$, male head broader than female ( $\mathrm{F}_{1,390}=$ 4.55, . $025<P<.05$ ); male femur/SVL ratio $.426 \pm$ .024 , female $.436 \pm .028$, female femur longer than male $\left(\mathrm{F}_{1,390}=14.61, P<.001\right)$; male tibia/SVL ratio $.510 \pm .031$, female $.521 \pm .030$, female tibia longer than male ( $\mathrm{F}_{1,390}=12.46, P<.001$ ); male foot/SVL ratio $.509 \pm .028$, female $.514 \pm .032$, not sexually dimorphic $\left(\mathrm{F}_{1,390}=3.75, P>.05\right)$.

Larval Characteristics.-Lescure (1972) described and figured the larvae.

Mating Call.-Dominant frequency modulates between $1000-2800 \mathrm{hz}$ (fig. 15); no harmonic structure in call; call pulsed or partially pulsed (fig. 16); note duration $.16-.17 \mathrm{~s}$, note repetition rate 1 per second.

Karyotype. -Diploid number 22; 7 pair median, 3 pair submedian, 1 pair subterminal (Bogart, 1974) or 5 pair median, 3 pair submedian, 3 pair subterminal (Heyer and Diment 1974); secondary construction on chromosome pair 8.
Distribution.-Known from a broad geographic range from Panama, throughout lowland South America east of the Andes (fig. 44).

ARGENTINA. CORRIENTES: Ituzaingó, Isla Apipé, IML 711, 768, 914; Manantiales, IML 778, MACN 13422-23, MCZ 35586.

FORMOSA: Esteros Lacuna Oca, IML 2195; Ingeniero Juárez, IML 700, 1102, 2194.

JUJUY: Ruta Yuto-Ledesma, near Arroyo Quemado, IML 1277-78, 1280.

MISIONES: Caraguatay, FMNH 9304; El Bonito, IML

1 sole terior rotted lorsal uals), .15); $s$ distripes al bar ' very , dis$P=$ folds; rcles, $r$ with spots pres$X^{2}=$ white assoice of 83, $P$ $\pm 4.4$ .05); .376 $P>$ male


Figure 44. Distribution map of Leptodactylus fuscus.

2045; Mártires, MACN 13644-46; Oberá, IML 769; Río Paranay, FMNH 9395-98, 9400, 9454, 9456-57; San Ignacio, IML 707.

SALTA: Agua Blanca, IML 1686; Campo Aguaray, IML 1470; near Embarcación, LACM 92010-030; near Hickmann, IML 448, 454, 660, 699; Los Toldas, Santa Victoria, IML 2252; Orán, Abra Grande, IML 1584, 1691, 2212; Río Pescado, IML 1403 (3); Tobantirenda, N Aguaray, IML 555, 1481.

BOLIVIA. BENI: Beni, FMNH 140212; Rurrenabaque, MCZ 10092-93, UMMZ 58831 (18).

COCHABAMBA: Villa Tumarí Road, km 58, Chapare Prov., USNM 146508-512.

SANTA CRUZ: Buenavista, CM 3883-84, 3954-55, 3957, 3976, 4241, 4392, 4431, 4433-34, 4436, 4439-440, UMMZ 60633 (5), 60634 (4), 60637; El Carmen, CM 36247; San José de Chiquitos, CM 36230 (8), MCZ 30032-38.

BRASIL. AMAZONAS: Coarí, MZUSP 28129, 39845861, 40805-0950; Igarapé Belém, no Solimões, MZUSP 245994600; Manacapuru, MZUSP 15951; Manaus, FMNH 64220, MCZ 295; Tefé, MZUSP 39919.

BAHIA: near Barreiras, UMMZ 109999, 110000 (2), 110001-03; Bom Jesus da Lapa, UMMZ 109996-98 (3), 110004.

CEARÁ: Crato, MNRio 409.
DISTRITO FEDERAL: Brasilia, MNRio 2716 (3).
ESPÍRITO SANTO: Itá, MZUSP 24601-610, 24669, USNM 121267-69; Linhares, MZUSP 25096.

GOIÅS: Aruanā, MZUSP 4993, 7549-551; Barra R. S. Domingos, MZUSP 24622-25, USNM 121299; Cana Brava, MZUSP 24628-646, USNM 121289-291, 121297-98; Fazenda Transvaal, Rio Verde, MZUSP 12511; Flôres, MZUSP 24626; Jaraguá, MZUSP 1419; Rio Verde, MZUSP 2534042; Sta. Isabel, Ilha do Bananal, MZUSP 24627.

MARANHAO: Perimirim, WCAB 8816
MATO GROSSO: Burití, Chapada dos Guimarães, MZUSP 37459-460; Corumbá, CM 36231-32, MCZ 30030-31, MZUSP 25496; Dumbá, MZUSP 1433; Fazenda Cruzeiro, Aquidauana, MZUSP 16206-07; Local do Massacre, MZUSP 4279-280, 14747-48; Mato Verde, rio Araguaia, MZUSP 24611-621; Santa Luzia (ex. Juti), MZUSP 28552; São Domingos, rio das Mortes, MZUSP 1080, 1093, 1394, 1397; São Luiz de Cáceres, MNRio 3073 (6), MZUSP 3638, 22160-67; mouth, Tapirapés River, MZUSP 25277-281; Três Lagoas, MZUSP 25221, 25227; Urucum, MZUSP 21382-85; Utiariti, MZUSP 25207-211; Parque Indigena do Xingu, Posto Diauarum, MZUSP 49491-9500.

MINAS GERAIS: Arinos, MZUSP 25050-51; Belo Horizonte, MNRio 1060-62, UMMZ 109994 (3); Januária, rio Pandeiros, MZUSP 24647-668, USNM 121287; Lagoa de Curralinho, Lassance, USNM 97015, 98210; Lagoa Santa, MZUSP 25076, 25080-81; Morro da Garça, MZUSP 25087; Ouro Preto, USNM 98028-044; Passa Quatro, MNRio 3898 (4); Pirapora, USNM 98268-271; Piraporinha, UMMZ 109995, USNM 98548; Rib. Confins, Buritís, MZUSP 25070; Rio Pandeiros, MZUSP 24152, USNM 121288, 121294-96; Sete Lagoas, MZUSP 25085; Uberlândia, MZUSP 12090-2122.

PARÁ: Barreira, no Tapajós, MZUSP 35805; Cachimbo, MNRio 2860 (3), MZUSP 21596-97, 21836-37, 21849-853, 21874; Cachoeira do Ararí, Iha de Marajó, MZUSP 2497374; Igarapé Taperebá, Ilha de Marajó, MZUSP 24961, 2496364, 24966-67, 24969-971; João Cativo, km 149 da Rede Cearense, 12 km a oeste de Itapipoca, MNR'o 3901; Rio Trombetas, headwaters, 15 km from Surinam, KU 128029-033.

RIO DE JANEIRO: Caxias, MNRio 2240; Campo Belo, USNM 96944-45; Itaguaí, MCZ 32699-2700; Itatiaia, Vale Pará́ba, MNRio 3555, 3561; Niteroi, Saco de Săo Francisco, USNM 99119; São João da Barra, MNRio 2539 (6); Teresópolis, USNM 97678-79.

RIO GRANDE DO SUL: Rio Pardo, MZUSP 21682-83; 39 km W Rio Pardo, FMNH 80332-33; Santa Maria, UMMZ 83133; Santo Augusto, Baixada da Olaria, MNRio 3844 (6).
RONDÔNIA: Pôrto Velho, MZUSP 16917-17365; Forte Príncipe da Beira, MZUSP 25160-61.

SÃO PAULO: Botucatú, MZUSP 4156; Emas, USNM 129176-77; Eugênio Lefévre, MZUSP 14906; Ituai, FMNH 83274; Jurumirim, MZUSP 24672; Piracicaba, MZUSP 1302; Piraçununga, Cachoeira de Emas, CM 33436, MNRio 2114; MZUSP 2294, 2440, 2442, 2860, 2862-65, 2867-68, 4606612, 4614-15, 4617-626, 9035-37, 11105-155, 11224, 2467071; Pôrto Martins, MZUSP 116-17, 281, 1963, 1966; Rio Pardo, Botucatú, MZUSP 3868; Rio Preto, MZUSP 24674; São Paulo, MZUSP 24677-686.

COLOMBIA. ANTIOQUIA: Nechí, FMNH 54569-570, 54573-74, 54577-78; Villa Arteaga, FMNH 78141.

BOLÍVAR: Río Viejo, USNM 145777-79; Tierra Alta, FMNH 61806; Tolu Viejo, MZUSP 5438, 5442.

BOYACÁ: Miraflores, USNM 153920-21.
CUNDINAMARCA: Cambao, USNM 147080-82; Villeta, USNM 151878.

MAGDALENA: Fundación, MCZ 8968-69; Ciénaga, USNM 144159-160.

META: 11.6 mi E Candilejas, UTA 3951; Granada, on Río Ariari, S of Villavicencio, USNM 151495-97; Loma Linda, UTA 3716; Macarena, upper Río Guejar and El Meco, USNM 144894; Mapiripán, UTA 3943, 3945; Menegua, E Puerto López, USNM 147275; near Puerto López, USNM 14619799, UTA 3713, 3942; San Juan de Arama, Los Micos, FMNH 81330-31; Villavicencio, FMNH 30574, 30813, 174078 , 174081, 174085-86, MCZ 64699, UMMZ 71223, USNM 144895, 147083-87, 147397-98, UTA 3748, 3944, 3946-47, 3949-950, 3952-53.
NORTE DE SANTANDER: Astillero, USNM 147088-89.
SANTANDER: Puerto Wilches, USNM 142805.

TOLIMA: Mariquita, FMNH 81836-37, USNM 1448964900, 147093-94.

VAUPÉS: Cerro Yapoboda, Río Cuduyarí, USNM 146432.
VICHADA: Puerto Carreño, CM 55655 (5).
FRENCH GUIANA. Kourou, LES 50-61, 285-290; Montsinéry, LES 664-669, 752-53; Rochambeau, LES 1018; Stoupan, LES 62.

GUYANA. Atkinson, USNM 162880-88, 162890-93; Demerara, FMNH 3299; Essequibo River, UMMZ 79476 (7); Georgetown, FMNH 174462-471, UMMZ 43968, 80416; Lethem, MCZ 50710-13; Manari, near Lethem, FMNH 17460204; upper Rupununi River, AMNH 46495 (4); Wismar, UMMZ 76680 (19), 104473.

PANAMA. HERRERA: Parita, USNM 127261.
PANAMÁ: Capitán, near Chepo, USNM 192621; Nueva Gorgona, AMNH 69735; Río Tocumen, MCZ 10036.

SURINAM. Berlijn, RMNH 15054; Blakawatra, RMNH 17522; Christian Kondre, MZUSP 24759-760, 24762-63; 24766; Coronie Road, RMNH 17568; Enmore Estate, USNM 16294349, 162951-965; Langaman Kondre, Marowijne, MZUSP 24583598; Lawa River, MZUSP 24773, 24777; Lelydorp, RMNH 17551 (3), 17561 (3); Moengo Tapoe, RMNH 17536; Paramaribo, CM 49483, RMNH 15132 (2), 15140, 15158, 17534, 17548, 17550 (2), 17552, USNM 158953-960; Powakka, CM 44266, 4949 [-92; Sipaliwini, RMNH 15171, 15189, 17523, 17541-46; Tibiti, RMNH 17553, 17554 (2), 17556-57, 17559560, 17562, 17564, 17567; 43 km S Paramaribo on Zanderij Highway, CM 49487; Zanderij, CM 50484-85, 50559.

TOBAGO. Bloody Bay, Charlotteville Road, USNM 167493, 167507-08, 167513, 192748 (5), 194989, 195005, 195017-19.

TRINIDAD. Aripo Savanna, MCZ 3299-3302; Brickfield, FMNH 49666; La Veroraca, USNM 141545; Piarro, USNM 166617-621; Port of Spain, USNM 102392-99; Quare River, CM 4535 .

VENEZUELA. AMAZONAS: Misión Coromoto-Atures, USNM 137193; Puerto Ayacucho, FMNH 175466-67, 190627 (3).

APURE: Hato Cariben, 46 km NE Puerto Páez, Río Cinaruco, USNM field 5482, 5668.

ARAGUA: Pie del Cerro (La Victoria), USNM 121148.
BOLÍVAR: Los Patos, 25 km SE El Manteco, USNM field 7587, 7589, 7590, 7592.

GUÁRICO: Calabozo, MCZ 50709; Estación Biológica de los Llanos, 9 km SE Calabozo, 100 m , USNM field 24619; Hato La Palmita, USNM 162700-01; Laguna de los Patos, UMMZ 131704-05; 10 km SE Valle de la Pascua, USNM 128839.

MONAGAS: 42 km SE Maturín, LACM 31375-79.
NUEVA ESPARTA: Salamanca, Margarita Island, USNM 137346.

SUCRE: Cumanacoa, CM 9064; Guaraúnos, KU 150799.
TÁCHRA: Maracoi, FMNH 125406, 176331.
YARACUY: San Felipe, BMNH 2272-73.

## Leptodactylus geminus Barrio 1973

Leptodactylus geminus Barrio 1973:199-206, figs. 2, 4, 6, 8. (Type locality, Argentina; Misiones, Bernardo de Irigoyen. Holotype CHINM 5860, male.)

Diagnosis. - Apparently morphologically identical to and indistinguishable from L. gracilis (see diagnosis for gracilis). At present geminus and gracilis can be differentiated only on the basis of call; the note repetition rate for geminus is faster ( 22 per second) than for gracilis (4 per second).

Adult Characteristics. - Specimens not examined by author.

Larval Characteristics.--Unknown.
Mating Call.-Dominant frequency modulated from ' $2700-3100 \mathrm{hz}$ (fig. 45); 16-31 notes per call group; call without harmonic structure; call pulsatile, 2-4 pulses per note (fig. 46); note duration from about 0.02 s (beginning notes) to 0.03 s (mid-call); note repetition rate 22 per second.
Karyotype.—Diploid number 22; 5 pair median, 4 pair submedian, 2 pair subterminal; secondary constriction in chromosome pair 8 (Barrio 1973).
Distribution. -Known from the northeastern part of the province of Misiones, Argentina (fig. 47).

Leptodactylus gracilis Duméril and Bibron 1841
Cystignathus gracilis Duméril and Bibron 1841:406-407. (Type locality, Uruguay, Montevideo. Holotype Paris Museum 4490, male.)

Leptodactylus plaumanni Ahl 1936:389-390. (Type locality, Brasil; Santa Catarina, Nova Teutônia. Holotype Senckenberg Museum 22469, male.)
Leptodactylus gracilis delattini Müller 1968:48-52, figs. 2, 3. (Type locality, Brasil; Ilha Campeche. Holotype originally Saarbrucken 4080 now in MZUSP.)

Diagnosis. -The species with light longitudinal stripes on skin folds on the dorsal surface of the tibia (fig. 48) (if light stripes indistinct, folds are present where stripes occur in other individuals) are geminus, gracilis, and marambaiae. Leptodactylus gracilis has a longer leg (e.g. tibia average $58 \% \mathrm{SVL}$ in males, $57 \%$ in females) than marambaiae (e.g. tibia $50 \%$ SVL). At present, geminus and gracilis can be differentiated only on the basis of call. The note repetition rate for gracilis is slower (4 per second) than for geminus (22 per second).

Figure 45. Sonagram of the mating call of Leptodactylus geminus, narrow band filter. Vertical scale marks at 1000 hz intervals; Horizontal scale mark at 1 s. Specimen from Argentina (Barrio tape copy).


Figure 46. Strip chart records of the mating call of Leptodactylus geminus. Upper figure of initial two notes in call sequence, lower figure of two notes in middle of call sequence. Line equals 0.01 s . See legend of Figure 45 for specimen data.


Figure 47. Distribution map of Leptodactylus geminus (circle) and gracilis (triangles).

Adult Characteristics ( $N=60$ ). -Dorsum spotted or striped (fig. 1, F, G, H, striped pattern not figured); mid-dorsal light stripe always present ( $100 \%$ ), light upper lip stripe almost always distinct ( $95 \%$ ), rarely indistinct ( $5 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}$ $=.18, P=.67$ ); no dark suborbital bar; light stripe on posterior face of thigh usually distinct ( $72 \%$ ), sometimes indistinct ( $27 \%$ ), rarely absent ( $2 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=1.19, P=.55$ ); tibia partially barred with light longitudinal pin stripes present; 6 well defined dorsolateral folds, sometimes an additional 2 or 4 ill defined folds (total 8 or 10 ); upper surface of tibia lacking white tubercles; posterior surface of tarsus lacking white tubercles $(100 \%)$; sole of foot lacking white tubercles ( $100 \%$ ); male SVL $43.0 \pm 4.8 \mathrm{~mm}$, female
$43.0 \pm 3.7 \mathrm{~mm}$, not sexually dimorphic $\left(\mathrm{F}_{1,58}=.002\right.$, $P>.05$ ); male head length/SVL ratio $.373 \pm .013$, female $.369 \pm .014$, not sexually dimorphic ( $\mathrm{F}_{1,58}=$ 1.37, $P>.05$ ); male head width/SVL ratio $.324 \pm$ .013 , female $.317 \pm .010$, male head broader than female ( $\mathrm{F}_{1,58}=5.79, .01<P<.025$ ); male femur/SVL ratio $.476 \pm .031$, female $.480 \pm .028$, not sexually dimorphic ( $\mathrm{F}_{1,58}=.31, P>.05$ ); male tibia/SVL ratio $.579 \pm .038$, female $.573 \pm .032$, not sexually dimorphic ( $\mathrm{F}_{1,58}=.38, P>.05$ ); male foot/SVL ratio $.593 \pm .029$, female $.598 \pm .022$, not sexually dimorphic ( $\mathrm{F}_{1,58}=.52, P>.05$ ).

Larval Characteristics. -Fernandez and Fernandez (1921) described and figured the larvae.

Mating Call. -Dominant frequency modulates be-

tween $500-2400 \mathrm{hz}$ (fig. 49); call without harmonic structure; call partially pulsed (fig. 50); note duration 0.04 to 0.05 s ; note repetition rate about 4 per second.

Karyotype.—Diploid number 22; 5 pair median, 4 pair submedian, 2 pair subterminal (Barrio 1973) or 5 pair median, 5 pair submedian, 1 pair subterminal (Bogart 1974); secondary constriction on chromosome pair 8.

Distribution.-The following records and distribution are based on museum specimens which may contain both $L$. geminus and L. gracilis. The (combined) distribution is Argentina through southeast Brasil (fig. 47).

ARGENTINA. BUENOS AIRES: Emestina, Ptdo. 25 de Mayo, MACN 20970-74; Lincoln, Estancia Triunfo, MACN 4688; Tigre, AMNH 11959, MACN 3692.

CATAMARCA: near Balcosna, IML 2263.
CHACO: Laguna Limpia, IML 406.
CÓRDOBA: Achiras, AMNH 51906; btwn La Falda and Río Ceballos, IML 1340; Puesto EI Cura, IML 25.

CORRIENTES: Colonia Carlos Pellegrini, MACN 4760; Ituzaingó, IML 916, MACN 4352.

MISIONES: Yacú-poí, 30 km E Pto. Bemberg, on Río Uruguaí, MACN 12341.

SANTA FÉ: Roldán, MACN 4911; Tostado, MACN 1831. TUCUMÁN: Tacanas, IML 532.
BRASIL. PARANÁ: Bituruna, MNRio 3712 (5).
RIO GRANDE DO SUL: Corrientes, MCZ 32701; Gramado, Taquara, MZUSP 16038; Ipanema, MZUSP 16051; Itaqui, MZUSP 348; Osório, CAS 85689-690, 94560, CM 39033 , MNRio 2723, 3781, MZUSP 21684-85; near Pôrto Alegre,

KU 154549, MZUSP 16059; Restinga Sêca, MZUSP 24692; Santa Maria, MCZ 22954, 22958, 22959, MZUSP 24691, USNM 121265-66; São Lourenço, MZUSP 90, 96; Tramandaí, MZUSP 26801.

SANTA CATARINA: Bôca da Serra, mun. Bom Jardim da Serra, 1200 m , MZUSP 35572-580; Nova Teutônia, MZUSP 8711-12; Novo Horizonte, 400-800 m, MZUSP 35307-330; São Bento, USNM 97174-75.

SÃO PAULO: França, MZUSP 610; Ipiranga, CM 33791; Perus, MZUSP 604; Ribeirão Pires, MZUSP 584; São Paulo, MZUSP 33, 453, 4532, 14897; Serra da Cantareira, MZUSP 24676.

URUGUAY. CERRO LARGO: 6 km SE Melo, AMNH 71176.

COLONIA: Santa Ana-Artilleros, MZUSP 22926.
DURAZNO: 18 km NE Paloma, Arroyo del Estado, CM 57038-39.

## Leptodactylus labrosus Espada 1875

Leptodactylus labrosus Jimenez de la Espada 1875:36. (Type locality, Ecuador; Los Ríos, Pimocha, shores of Río Daule. Lectotype Museo Nacional de Ciencias Naturales, Madrid, no number, female.)
Leptodactylus curtus Barbour and Noble 1920:405-406. (Type locality, Peru; Cajamarca, Bellavista. Holotype MCZ 5281.)

Diagnosis. -The species lacking a distinct light stripe on the posterior surface of the thigh in some or all individuals are albilabris, bufonius, labrosus, mystacinus, and troglodytes. The sole of the foot is usually smooth in labrosus (fig. 69), the sole of the foot has distinct


Figure 49. Sonagram of the mating call of Leptodactylus gracilis, narrow band filter. Vertical scale marks at 1000 hz intervals. Horizontal scale mark at 1 s . Specimen from Argentina, Buenos Aires (Barrio tape copy).


Figure 50. Strip chart record of the mating call of Leptodactylus gracilis. Line equals 0.01 s . See legend of Figure 49 for specimen data.
white tubercles in albilabris, troglodytes, ventrimaculatus and some mystacinus individuals. Leptodactylus labrosus is found along dry coastal South America from mid-Ecuador to Peru; albilabris is in the West Indies, troglodytes in NE Brasil, ventrimaculatus along wet east coast South America from Colombia to mid-Ecuador, mystacinus occurs in southern South America east of the Andes. Leptodactylus labrosus has a pair or two of distinct dorsolateral folds (indicated at least by color pattern in poorly preserved specimens), bufonius lacks distinct dorsolateral folds.

Adult Characteristics ( $N=32$ ). -Dorsum spotted or rarely uniform, spots rarely fused (fig. $1, \mathrm{~A}, \mathrm{~B}, \mathrm{C}, \mathrm{J}$ ); no light mid-dorsal stripe; no distinct light upper lip stripe; dark suborbital bar present or absent; light stripe on posterior face of thigh almost always absent (94\%), rarely indistinct ( $6 \%$ ), presence not sexually dimorphic ( $X^{2}=.01, P=.92$ ); tibia barred; dorsolateral folds often absent or 4 indistinct folds present; dorsal surface of tibia usually with many or scattered white tubercles, sometimes absent; posterior surface of tarsus usually with scattered white tubercles $(78 \%)$, sometimes absent $(22 \%)$, presence not sexually dimorphic ( $X^{2}=.26, P$ $=.61$ ); sole of foot usually lacking white tubercles $(91 \%)$, rarely present $(9 \%)$, not sexually dimorphic ( $X^{2}$ $=.22, P=.64$ ); male SVL $54.6 \pm 5.3 \mathrm{~mm}$, female $53.3 \pm 6.3 \mathrm{~mm}$, not sexually dimorphic $\left(F_{1,30}=.30\right.$, $P>.05$ ); male head length/S VL ratio $.356 \pm .013, \mathrm{fe}-$ male $.361 \pm .016$, not sexually dimorphic $\left(F_{1,30}=.60\right.$, $P>.05$ ); male head width/SVL ratio $.344 \pm .008$, female $.347 \pm .013$, not sexually dimorphic $\left(F_{1,30}=.68\right.$, $P^{\cdot}>.05$ ); male femur/SVL ratio $.400 \pm .019$, female $.417 \pm .022$, female femur longer than male $\left(\mathrm{F}_{1,30}=\right.$ 4.38, . $025<P<.05$ ); male tibia/SVL ratio $.430 \pm$ .007 , female $.442 \pm .025$, not sexually dimorphic ( $F_{1,30}$ $=1.84, P>.05$ ); male foot $/ \mathrm{SVL}$ ratio $.459 \pm .014$, female $.481 \pm .032$, not sexually dimorphic $\left(\mathrm{F}_{1,30}=\right.$ 3.81, $P>.05$ ).

Larval Characteristics. -Unknown.
Mating Call. -Unknown.
Karyotype. -Unknown.
Distribution.-Mostly associated with dry west coast South America from mid-Ecuador to northern Peru, including the dry interandean valley of northern Peru (fig. 51).

ECUADOR. EL ORO: 7 km SE Buena Vista, USNM 196728; near Machala, USNM 196729-730 (3); Río Jubones, AMNH 16241.

GUAYAS: near Guayaquil, KU 120296, USNM 66876880, 164327-333; Río Puyango, AMNH 16206, 16228.

LOJA: Casanga Valley, USNM 196731; La Toma, USNM 196732.

LOS RÍOS: Quevedo, WCAB 40161; 56 km N Quevedo, KU 146181-85, 147569, 152576-77.

PERU. ANCASH: 4.5 km SSE Río Casma, LACM 49161.
CUZCO: Río Cosñipata, 4 km SW Santa Isabel, 1700 m , KU 46443, 46603.

LIBERTAD: Río Jequetepeque, 2 km N Cruce de San José, LACM 49148-154, 49280.

PIURA: 1.5 km S Las Lomas, Río Chipillico, LACM 49155160, 49281, 77005; near Sullana, USNM 153799.

## LEPTODACTYLUS LATINASUS ESPADA 1875

Leptodactylus latinasus Jiménez de la Espada 1875:40. (Type locality, Uruguay; Montevideo. Holotype Museo Nacional de Ciencias Naturales, Madrid, jar number 335, female.)
Leptodactylus prognathus Boulenger 1888:187. (Type locality, Brasil; Rio Grande do Sul. Holotype BMNH 1947.2.17.52, male.)
Leptodactylus anceps Gallardo 1964:100-105, plate 1, fig. 2, plate 2, fig. 2. (Type locality, Argentina; Tucumán, Tucumán. Holotype MACN 531, male.)

Diagnosis. -The species having a combination of a distinct light stripe on the posterior face of the thigh and obvious white tubercles on the posterior surface of the tarsus and sole of foot in some or all individuals are albilabris, elenae, fragilis, latinasus, mystaceus. Leptodactylus latinasus lacks distinct dorsolateral folds, distinct dorsolateral folds (indicated by color pattern in poorly preserved individuals) are found in albilabris, elenae, and mystaceus. Leptodactylus latinasus and fragilis have considerable morphological and color pattern overlap (fig. 43), fragilis being a slightly larger species (maximum male SVL 43.0 mm , female 43.6 mm ) than latinasus (maximum male SVL 37.9 mm , female 36.3 mm ). Leptodactylus latinasus has a southern South American distribution, fragilis has a Middle American and north coast South American distribution.

Adult Characteristics $(N=233)$.-Dorsum spotted or blotched (fig. 1, A, B, C, E, N); mid-dorsal light stripe absent; light lip stripe usually indistinct ( $66 \%$ ), more females with distinct lip stripes than males $\left(X^{2}=\right.$ $11.67, P<.001$ ); dark suborbital bar absent; light stripe on posterior face of thigh usually very distinct ( $90 \%$ ), sometimes indistinct ( $10 \%$ ), distinctiveness not sexually dimorphic $\left(X^{2}=.11, P=.74\right)$; tibia barred; dorsolateral folds indistinct, when present usually 2 , sometimes 4 ; dorsal surface of tibia usually with many, sometimes scattered, white tubercles, very rarely absent; posterior surface of tarsus with many distinct white tubercles $(100 \%)$; sole of foot with many distinct white tubercles ( $100 \%$ ); male SVL $31.2 \pm 1.7 \mathrm{~mm}$, female $33.0 \pm 1.9$ mm , females larger than males $\left(\mathrm{F}_{1,231}=51.75, P<\right.$ .001 ); male head length/SVL ratio $.372 \pm .012$, female $.368 \pm .013$, male head longer than female $\left(F_{1,231}=\right.$ $3.85, P=.05$ ); male head width/SVL ratio $.343 \pm$ .013 , female $.343 \pm .012$, not sexually dimorphic $\left(\mathrm{F}_{1,231}=.06, P>.05\right) ;$ male femur/SVL ratio $.388 \pm$ .028 , female $.394 \pm .023$, not sexually dimorphic $\left(\mathrm{F}_{1,231}=2.08, P>.05\right)$; male tibia/SVL ratio $.451 \pm$ .023 , female $.455 \pm .029$, not sexually dimorphic $\left(\mathrm{F}_{1,231}\right.$ $=1.36, P>.05$ ); male foot $/ \mathrm{SVL}$ ratio $.476 \pm .024$, female $.475 \pm .027$, not sexually dimorphic $\left(F_{1,231}=\right.$ $.02, P>.05$ ).

Larval Characteristics.-Fernandez and Fernandez (1921) described and figured the larvae as L. prognathus.

Mating Call.-Dominant frequency modulates from


Figure 51. Distribution map of Leptodactylus labrosus (squares) and latinasus (triangles).
$3100-4000 \mathrm{hz}$ (fig. 52); call lacking harmonic structure; note non-pulsed (fig. 53); note duration 0.06 s ; note repetition rate 2.3 per second.

Karyotype. -Bogart (1974) described the karyotype as diploid number 22; 5 pair median, 3 pair submedian, 2 pair subterminal, 1 pair terminal; secondary constriction on chromosome pair 8.

Distribution.--Found throughout the Gran Chaco and littoral zone of Argentina, central and coastal Brasil (fig. 51).

ARGENTINA. BUENOS AIRES: Bancalari, UMMZ 98837 (3); Ensenada, UMMZ 98838; José C. Paz, IML 854, 4475,

CATAMARCA: El Alto, IML 1442; near La Merced, IML 2261 (13).

CHACO: Barranqueras, MZUSP 26325-333.
CORRIENTES: Corrientes, MACN 4605 (4).
ENTRE RÍOS: Concepción del Uruguay, MACN 4530 (3).
FORMOSA: Ingeniero Juárez, LACM 92044; Río Bermejo, La Florencia, IML 651 (11).

JUJUY: Arroyo Los Naranjos, 8.3 km SSW Perico del Carmen, KU 43866-870; Ruta Yuto-Ledesma, near Arroyo Quemado, IML 1276 (11).

SALTA: Campo Aguaray, IML 1479 (3); El Saucelito, 50 km S Orán, IML 1627 (2); near Embarcación, LACM 9203132, 92034-043, 92045-46, 92048-054, 92057-065; near Hickmann, IML 307 (3), 311 (4), 652 (8); Parque El Rey, Pozo Los Lobitos, IML 2393 (2); Tobantirenda, $N$ of Aguaray, IML 1482 (7); Urundel, IML 9.

SANTA FÉ: Bañados del Rincón, CM 38728.
SANTIAGO DEL ESTERO: Bañado de Figueroa, $\pm 6 \mathrm{~km}$ N Caspi Corral, KU 43961-62; S. Loreto, MCZ 33714-16. TUCUMÁN: El Cadillal, IML 2297, 2410 (15), KU $44045-$ 062, 44065-66; El Durazno, IML 1902; Hualinchai, 8 km W S. P. de Colalao, IML 1783; Río Urueña, near Salta, IML 1429 (2); Saladillo, IML 467 (3); San Javier, IML 1599 (2); Soledad de Maria-Lamadrid, IML 37; Tacanas, IML 534; Tucumán FMNH 69077, IML 1427, UMMZ 109751 (8).

BRASIL. BAHIA: Bom Jesus da Lapa, UMMZ 109991 (2); Itiúba, MZUSP 38556-564.

ESPÍRITO SANTO: São Mateus, MCZ 1298.
MINAS GERAIS: Rio Grande at São José, UMMZ 109992.
RIO GRANDE DO SUL: Pôrto Alegre, FMNH 80347-351, 80354-59, 83289; 39 km N Rio Pardo, MZUSP 21699-1701; 39 km W Rio Pardo, FMNH 80352-53; São Lourenço, MZUSP 93; Vila Nova, São Sepé, MZUSP 27303-06.

URUGUAY. 30 Y 3: 8 mi E 30 y 3, FMNH 10435, 10463 ; Boca del Rio Tacuarí, AMNH 71189; Quebrada de los Cuervos, 45 km N 30 y 3, FMNH 10489-491.

ARTIGAS: 6 km NNW Belén, AMNH 71181.
CANELONES: Montevideo, USNM 196655.
COLONIA: Nueva Palmira, Arroyo del Sauce, CM 5704647.

LAVALLEJA: Río de Averías, Depto. Minas, FMNH 10251. MALDONADO: Sierra de Animas, MZUSP 24567-570. RÍO NEGRO: Arroyo Neapo, 15 km S Paysandú, AMNH 71178-180.

ROCHA: Arroyo Garzón, 10 km NW Garzón, FMNH 10251. SAN JOSE: Arazatí, S of Cocilda, FMNH 10623, 10628.
TACUAREMBÓ: 40 km NW Tacuarembó, AMNH 71182 88; 3 km NE Tambores, Pozo Hondo, CM 55394-98.

## Leptodactylú laurae new species

Figure 54
Holotype: MZUSP 130, an adult male from Brasil: Minas Gerais; Agua Limpa, Juiz de Fora. Collected by Joaquim Venancio on 27 November 1947.

Diagnosis. -The species with a combination of a distinct light stripe on the posterior surface of the thigh and smooth surfaces on the posterior tarsus and sole of the foot in some or all individuals are fuscus, geminus, gracilis, laurae, longirostris, marambaiae, notoaktites, and poecilochilus. Leptodactylus laurae lacks light stripes on the dorsal surface of the tibia, such stripes are found in geminus, gracilis, and marambaiae (fig. 48). All individuals of $L$. laurae have a light mid-dorsal stripe and at least 6 dorsolateral folds; only individuals with a light mid-dorsal stripe have 6 dorsolateral folds in longiros-


Figure 54. Dorsal view of the holotype of Leptodactylus laurae.
tris, notoaktites, and poecilochilus, most fuscus individuals lack a light mid-dorsal stripe. The leg of laurae is longer (e.g. male foot/SVL ratio $.649 \pm .039$, female $.628 \pm .028$ ) than fuscus (male foot/SVL ratio $.509 \pm$ .028 , female $.514 \pm .032$ ), longirostris (male foot/SVL ratio $.545 \pm .026$, female $.553 \pm .031$ ), notoaktites male foot/SVL ratio $.587 \pm .033$, female; $.583 \pm .036$ ), and poecilochilus (male foot/SVL ratio $.514 \pm .029$, female $.508 \pm .029$ ). Leptodactylus laurae has a mideast and southern South American distribution, L. poecilochilus and L. longirostris occur in northem South America. Many individuals of notoaktites have distinct white tubercles on the sole of the foot.
Description of Holotype. -Snout pointed from above, rounded-acute in profile; canthus rostralis indistinct; loreal slightly concave; tympanum distinct, greatest diameter just greater than $1 / 2$ eye diameter; vomerine teeth in slightly arched series posterior to choanae; vocal slits present; internal vocal sac; finger lengths in order of decreasing size $\mathrm{I} \simeq \mathrm{III}>\mathrm{II} \simeq \mathrm{IV}, \mathrm{I} \gg \mathrm{II}$; inner metacarpal tubercle flat, oval, smaller than large, flat, outer metacarpal tubercle; no nuptial asperities; dorsal sur-
faces smooth; 3 pair of dorsolateral folds; ventral texture smooth; belly disk fold distinct; toe tips not expanded; toes free, lacking fringe or web; subarticular tubercles moderately developed; outer metatarsal tubercle small, indistinct, smaller than indistinct, oval, inner metatarsal tubercle; tarsal fold indistinct; no metatarsal fold; posterior surface of tarsus smooth; sole of foot smooth.

SVL 40.4 mm , head length 14.5 mm , head width 12.1 mm , interorbital distance 1.5 mm , eye-nostril distance 3.4 mm , femur 19.5 mm , tibia 24.6 mm , foot 27.4 mm .

Dorsum tan with lighter and darker brown markings including a light mid-dorsal stripe bordered by an irregular dark stripe on either side; series of dark spots parallel other dorsolateral folds; upper lip dark edged bordered above by distinct light stripe from tip of snout passing under eye to tympanum; limbs barred; venter immaculate; posterior surface of thigh blotched, light stripe more distinct on left than right side; light tarsal fold stripe; posterior tarsus and sole of foot mottled.
Etymology. - Named for my daughter Laura, a friend to all animals, including frogs.
Remarks. -This is the species referred to as "barred gracilis" in the morphological analysis.

Adult Characteristics ( $N=35$ ).—Dorsum spotted or striped (fig. 1, F, H, striped pattern not figured); middorsal light stripe always present ( $100 \%$ ); light upper lip stripe usually distinct ( $71 \%$ ), often indistinct ( $29 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=.27, P=$ .60); no dark suborbital bar; light stripe on posterior face of thigh distinct ( $51 \%$ ) or indistinct ( $49 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=.71, P=.40$ ); tibia barred; 6 well defined dorsolateral folds; dorsal surface of tibia without white tubercles; posterior surface of tarsus without white tubercles ( $100 \%$ ), sole of foot without white tubercles ( $100 \%$ ); male SVL $36.1 \pm 2.3 \mathrm{~mm}$, female $40.7 \pm 3.3 \mathrm{~mm}$, females larger than males ( $\mathrm{F}_{1,33}=22.5, P<.001$ ); male head length/SVL ratio $.379 \pm .014$, female $.365 \pm .011$, male head longer than female ( $\mathrm{F}_{1,33}=7.53, .01<P<.025$ ); male head width/SVL ratio $.313 \pm .013$, female $.304 \pm .007$, male head broader than female ( $\mathrm{F}_{1}, 33=4.34, .025<P<$ .05 ); male femur/SVL ratio $.485 \pm .030$, female .489 $\pm .023$, not sexually dimorphic ( $\mathrm{F}_{1,33}=.12, P>.05$ ); male tibia/SVL ratio $.591 \pm .038$, female $.594 \pm .032$, not sexually dimorphic ( $\mathrm{F}_{1,33}=.03, P>.05$ ); male foot/SVL ratio $.649 \pm .039$, female $.628 \pm .028$, not sexually dimorphic ( $\mathrm{F}_{1}, 33=2.55, P>.05$ ).

Larval Characteristics.-W. C. A. Bokermann and Ivan Sazima are in the process of describing the larvae of $L$. laurae.

Mating Call.-W. C. A. Bokermann and Ivan Sazima are describing the call of this species (pers. comm.).

Karyotype. -Unknown.
Distribution.-Southeast and central Brasil (fig. 55).
BRASIL. DISTRITO FEDERAL: Brasilia, MZUSP 25349.
MATO GROSSO: Serrinha, 120 km W Mortes River, MZUSP 4275.
ral texture expanded; - tubercles cle small, metatarsal fold; posmooth. ead width ye-nostril mm , foot markings $y$ an irregoots paraldged bor, of snout ed; venter hed, light ight tarsal 1ottled.
a, a friend
is "barred
spotted or red); midt upper lip ct (29\%), .27, $P=$ terior face listinctive.40); tibia sal surface ace of tarot without 3 mm , fetan males SVL ratio sad longer male head
.007, male
$15<P<$
male .489
$P>.05$ );
$4 \pm .032$, .05); male .028, not
mann and the larvae

1 Ivan Sa 3. comm.). 1 (fig. 55 ). USP 25349. urtes River,


Figure 55. Distribution map of Leptodactylus laurae (squares), longirostris (triangles), änd marambaiae (circle).

MINAS GERAIS: Agua Limpa, MZUSP 130; Serra do Caraça, MZUSP 13516.
PARANÁ: Curitiba (Xaxim), USNM 125499.
RIO GRANDE DO SUL: Santa Maria, MCZ 22951-53, 22955-57, 22959 (2), MZUSP 24688-690.
SÃO PAULO: Botucatú, MZUSP 14482; Campo Grande, Santo André, CAS 93822-23, KU 9209-212, 74215-16, MZUSP 516; Emas, MZUSP 9034; Eugênio Lefévre, MZUSP 11328; Itanhaem, MZUSP 625; Paranapiacaba, MZUSP 846; Rio Grande, MZUSP 1967; São Paulo, MZUSP 906; Serra da Bocaina, MCZ 15849, MZUSP 24136-39, 25467, USNM 81133, 96614-16; Alto da Serra de Cubatäo, USNM 96813, 124588-89.

## Leptodactylus longirostris Boulenger 1882

Leptodactylus longirostris Boulenger 1882:240, plate 16, fig. 3. (Type locality, Brazil; Santarem. Lectotype BMNH 76.5.26.4, female.)

Diagnosis. -The species having a combination of a distinct light stripe on the posterior surface of the thigh and smooth surfaces on the posterior tarsus and sole of foot in some or all individuals are fuscus, geminus, gracilis, laurae, longirostris, marambaiae, mystaceus. notoaktites, and poecilochilus. Leptodactylus longirostris has a barred tibia, the dorsal surface of the tibia has light longitudinal stripes in geminus, gracilis, and marambaiae. Only individuals of $L$. longirostris with light middorsal stripes have 6 dorsolateral folds (fig. 56), all fuscus have 6 dorsolateral folds (individual $L$. longirostris with the light mid-dorsal stripe are morphologically difficult to distinguish from fuscus). All individuals of laurae have a light mid-dorsal stripe and 6 dorsolateral folds; the leg of longirostris is shorter (e.g. male foot/


FIGURE 56. Dorsal views of striped (left, FMNH 128831) and unstriped (right, UPR 2641) Leptodactylus longirostris.


Figure 57. Lateral view of the heads of Leptodactylus amazonicus (left, lip stripe very distinct), longirostris (center, lip stripe moderately distinct), poecilochilus (right, no lip stripe, lip bar present).

SVL ratio $.545 \pm .026$, female $.553 \pm .031$ ) than laurae (male foot/SVL ratio $649 \pm .039$, female .628 $\pm .028$ ); longirostris occurs in northern South America, laurae in mid-east and southern South America. Most individuals of mystaceus have distinct white tubercles on the sole of the foot; mystaceus occurs along coastal Brasil. Many individuals of notoaktites have white tubercles on the sole of the foot; notoaktites occurs in SE Brasil. Leptodactylus longirostris often has a distinct light lip stripe and lacks a dark suborbital bar, poecilochilus lacks a distinct light lip stripe and often has a dark suborbital bar (fig. 57).

Adult Characteristics ( $N=70$ ).-Dorsum uniform or spotted, spots sometimes elongate, fused (fig. 1, A, B, C, E, J); light mid-dorsal stripe present in $17 \%$ of individuals, presence not sexually dimorphic $\left(X^{2}=.09\right.$, $P=.76$ ); light lip stripe usually indistinct ( $60 \%$ ), often distinct ( $40 \%$ ), distinctiveness not sexually dimorphic ( $X^{2}=2.94, P=.09$ ); dark suborbital bar absent; light stripe on posterior face of thigh usually distinct ( $80 \%$ ), sometimes indistinct ( $20 \%$ ), more females ( $100 \%$ ) have distinct light stripes than males ( $X^{2}=6.80, P=.009$ ); tibia barred; usually 4 well defined dorsal folds, 6 dorsolateral folds present when light mid-dorsal stripe present; dorsal surface of tibia lacking white tubercles; pos-
terior surface of tarsus almost always (99\%) lacking white tubercles, presence not sexually dimorphic ( $X^{2}=$ $.14, P=.71$ ); sole of foot lacking white tubercles $(100 \%)$; male SVL $38.2 \pm 1.8 \mathrm{~mm}$, female $41.8 \pm 2.4$ mm , females larger than males $\left(\mathrm{F}_{1,68}=49.7, P<\right.$ .001); male head length/SVL ratio $.394 \pm .012$, female $.387 \pm .014$, male head longer than female ( $\mathrm{F}_{1,68}=$ $5.28, P=.025$ ); male head width/SVL ratio $.338 \pm$ .015 , female $.334 \pm: 013$, not sexually dimorphic ( $\mathrm{F}_{1,68}=1.12, P>.05$ ); male femur/SVL ratio .446 $\pm .041$, female $.457 \pm .033$, not sexually dimorphic $\left(\mathrm{F}_{\mathrm{t}, 68}=1.31, P>.05\right)$; male tibia/SVL ratio $.512 \pm$ .024 , female $.527 \pm .031$, female tibia longer than male $\left(\mathrm{F}_{1,68}=5.28, P=.025\right)$; male foot/SVL ratio $.545 \pm$ .026 , female $.553 \pm .031$, not sexually dimorphic ( $\mathrm{F}_{1,68}=1.21, P>.05$ ).

Larval Characteristics.-Unknown.
Mating Call.-Dominant frequency modulated between $1500-3600 \mathrm{hz}$; note duration about 0.8 s ; note repetition rate 1.4 per second (from Rivero, 1971, fig. 58 reproduced here from same sonagram described by Rivero).
Karyotype.—Unknown.
Distribution. - Centered upon the Guiana Shield (fig. 55).

Figure 58. Sonagram of the mating call of Leptodactylus longirostris. Vertical scale marks at 1000 hz intervals. Horizontal scale mark at 1 s . Specimen from Venezuela, La Escalera (sonagram courtesy of Juan A. Rivero).

BRASIL. AMAZONAS: Ponta Negra, Negro River, MZUSP 24880; Tapera, Rio Negro, MZUSP 37518.

PARÁ: Igarapé Jaramacaru, Campos do Ariramba, MZUSP 28401-04; Rio Mapuera, at equator, AMNH 46189-190 (3); Rio Mapuera, R. Trombetas, AMNH 46187-88.

GUYANA. Kartabo, USNM 118065-66; Kuyuwini Landing, AMNH 49349-351, 49353-54 (4); upper Rupununi River, AMNH 81355-56.

SURINAM. Brownsberg Nature Park, Brokopondo Dist., MCZ 89648; Brownsweg, RMNH 17531, 17535; Christian Kondre, MZUSP 24758, 24761, 24765, 24767-772; Kaiserberg Airstrip, Zuid River, FMNH 128827-832, 128913-18, 128920-23, RMNH 17527 (4), 17530, 17549 (5); Krakka, RMNH 17540 (2); road between Krakka and Phedra, RMNH 17537, 17539 (2); Powakka, CM 49482, 49484, 44265, 44272, 44274; Matta, RMNH 17558; Sabakoe Creek, between Berlijn and Zanderij, RMNH 15106; Sipaliwini, RMNH 15176, 15178 (2), 17524-26, 17528-29, 17532-33, 17547, 17569; Tibiti, RMNH 17555, 17563 ; Troeli Cr., 6 km S Matta, RMNH 15115 (2), 15133 (4); Zanderij, MCZ 35642, MZUSP 15869-870, USNM 159066-67.

VENEZUELA. BOLÍVAR: km 104-151 on El DoradoSanta Elena de Uairén Road, KU-WED 40072, 40078, 40080, 40085, 40151, 40181-82, 40208-09, 40263, 40281-87, 40381; La Escalera, Serrania de Lema, MCZ 79907, UPR 2641, 264345, 2647.

## LEPTODACTYLUS MARAMBALAE IZECKSOHN 1976

Leptodactylus marambaiae Izecksohn 1976:527-530, fig. 1. (Type locality, Brasil: Rio de Janeiro; Restinga da Marambaia. Holotype personal collection of Izecksohn 4123, adult male.)

Diagnosis. -The species with light longitudinal stripes on skin-folds on the dorsal surface of the tibia (fig. 48) (if light stripes indistinct, folds are present where stripes occur in other individuals) are geminus, gracilis, and marambaiae. Leptodactylus marambaiae has a shorter leg (e.g. tibia $50 \%$ SVL) than gracilis (e.g. tibia average $58 \%$ SVL in males, $57 \%$ SVL in females). At present, marambaiae cannot be morphologically distinguished from geminus. The note repetition rate of the mating call is slower for marambaiae ( 6 per second) than for geminus (22 per second).

Adult Characteristics. -Dorsum striped; mid-dorsal light stripe always present; light upper lip stripe distinct; no dark suborbital bar; light stripe on posterior face of thigh usually distinct, sometimes indistinct; tibia partially barred with light longitudinal pin stripes present; 6 well defined dorsolateral folds; upper surface of tibia lacking white tubercles; posterior surface of tarsus lacking white tubercles; sole of foot lacking white tubercles; male SVL 36.8 mm , female 40.2 mm ; male head length/ SVL ratio .40, female . 36 ; male head width/SVL ratio . 34 , female .34 ; male femur/SVL ratio .44 , female .44 ; male tibia/SVL ratio . 50 , female .50 ; male foot/SVL ratio .56, female . 54.

## Larval Characteristics. - Unknown.

Mating Call.-Dominant frequency modulates between $3000-3700 \mathrm{hz}$ (fig. 59); call without harmonic structure; call not pulsed (fig. 60); note duration about 0.02 s ; note repetition rate about 6 per second.

Karyotype. -Unknown.
Distribution. -Known only from the type locality (fig. 55).

BRASIL. RIO DE JANEIRO: Restinga da Marambaia.

## LEPTODACTYLUS MYSTACEUS (SPIX) 1824

Rana mystacea Spix 1824:27, plate 3, fig. 3. (Type locality, Brasil: Bahia [Salvador as designated by Bokermann 1966]. Types lost.)

Diagnosis. -Most individual mystaceus have a combination of a distinct light stripe on the posterior surface of the thigh and distinct white tubercles on the surfaces of the posterior tarsus and sole of foot; these states are shared with albilabris, elenae, fragilis, and latinasus. Leptodactylus mystaceus have distinct dorsolateral folds (at least indicated by color pattern), fragilis and latinasus lack distinct dorsolateral folds. Leptodactylus mystaceus has white tubercles on the dorsal surface of the tibia, the tibia is smooth in elenae. Leptodactylus mystaceus is found in east coastal Brasil, albilabris occurs in the West Indies.

Some individuals of mystaceus lack the white tubercles on the tarsus and sole of foot (light thigh stripe present), these states are shared with at least some individuals of fuscus, geminus, gracilis, laurae, longirostris, notoaktites, and poecilochilus. The tubercles on the dorsal surface of the tibia distinguishes mystaceus from all these species.

Adult Characteristics ( $N=38$ ). -Dorsum spotted or rarely uniform (fig. $1, \mathrm{~A}, \mathrm{C}, 0$ ); light mid-dorsal stripe usually absent ( $97 \%$ ), presence not sexually dimorphic ( $X^{2}=.08, P=.78$ ); light lip stripe usually distinct (79\%), distinctiveness not sexually dimorphic $\left(X^{2}=\right.$ $.14, P=.71$ ); dark suborbital bar absent; light stripe on posterior face of thigh distinct ( $100 \%$ ); tibia barred; usually 4 or 2 well defined dorsolateral folds, 6 dorsolateral folds present when light mid-dorsal stripe present; dorsal surface of tibia usually with many distinct white tubercles; posterior surface of tarsus usually with many distinct white tubercles ( $87 \%$ ), tubercles sometimes lacking ( $13 \%$ ), presence not sexually dimorphic ( $X^{2}=.43, P=.51$ ); sole of foot usually with many distinct tubercles ( $87 \%$ ), tubercles sometimes lacking ( $13 \%$ ), presence not sexually dimorphic $\left(X^{2}=.43, P\right.$ $=.51$ ); male SVL $42.7 \pm 2.3 \mathrm{~mm}$, female $43.6 \pm 3.0$ mm , not sexually dimorphic $\left(\mathrm{F}_{1,36}=1.18, P>.05\right)$; male head length $/ \mathrm{SVL}$ ratio $.379 \pm .015$, female .375 $\pm .022$, not sexually dimorphic $\left(\mathrm{F}_{1,36}=.40, P>.05\right)$; male head width/SVL ratio $.344 \pm .018$, female .342 $\pm .029$, not sexually dimorphic $\left(\mathrm{F}_{1,36}=.15, P>.05\right)$; male femur/SVL ratio $.434 \pm .032$, female $.461 \pm$ .037, female femur longer than male ( $\mathrm{F}_{1,36}=5.61, .025$ $<P<.01$ ); male tibia/SVL ratio $.509 \pm .015$, female $.517 \pm .029$, not sexually dimorphic $\left(\mathrm{F}_{1,36}=1.42, P\right.$ $>.05$ ); male foot/SVL ratio $.554 \pm .022$, female .548 $\pm .034$, not sexually dimorphic $\left(\mathrm{F}_{1,36}=.42, P>.05\right)$. mmann 1966].
have a comsrior surface the surfaces se states are d latinasus. lateral folds lis and latieptodactylus ll surface of optodactylus lbilabris oc-
white tuber1 stripe pres)me individlongirostris, $s$ on the doreus from all
m spotted or dorsal stripe ly dimorphic tally distinct phic $\left(X^{2}=\right.$ ; light stripe tibia barred; Ids, 6 dorso1 stripe presnany distinct usually with ercles somely dimorphic $y$ with many imes lacking $X^{2}=.43, P$ 2 $43.6 \pm 3: 0$ $8, P>.05$ ); female .375 10, $P>.05$ ); female .342 $15, P>.05$ ); nale $.461 \pm$ $=5.61, .025$ .015, female ${ }_{36}=1.42, P$ , female . 548 42, $P>.05$ ).


Figure 59. Sonagram of the mating call of Leptodactylus marambaiae. Vertical scale marks at 1000 hz intervals. Horizontal scale mark at 1 s . Specimen from Brasil, Restinga da Marambaia (tape courtesy of W. C. A. Bokermann).


Figure 60. Strip chart record of the mating call of Leptodactylus marambaiae. Line equals 0.01 s . See legend of Figure 59 for specimen data.

Larval Characteristics.—Unknown.
Mating Call.-Unknown.
Karyotype. -Unknown.
Distribution.-East coast of Brasil (fig. 61).
BRASIL. BAHIA: Copec. Ilhéus, MNRio 1724 (4), WCAB 45899-5919, 46570-6601, 47066-69; Itapetinga, WCAB 44885.

ESPÍRITO SANTO: Santa Teresa, CAS-SU 11787-88; São Mateus, MCZ 1298 (5).

RIO DE JANEIRO: Caxias, MNRio 1809 (5), 2374, 2861; Cidade dos Meninos, MNRio 1656 (3); Meriti, USNM 96222; Niteroi, Saco de São Francisco, USNM 96407-411, 99120 ; road to Sáo Paulo, km 40, D. F., 97572; Serra de Friburgo, USNM 96467; Teresópolis, KU 92927-931, MNRio 397 (4), WCAB 12252.

## Leptodactylus mystacinus Burmeister 1861

Cystignathus mystacinus Burmeister 1861:532. (Type locality, Argentina. Holotype Martin-Luther-Universität, Halle (Saale), no number, male.)
Cystignathus labialis Cope 1878:90. (Type locality unknown. Presumed holotype USNM 31302, juvenile.)

Diagnosis.-The species having a combination of no light stripe on the posterior surface of the thigh and distinct white tubercles on the posterior surface of the tarsus are bufonius, labrosus, mystacinus, troglodytes, and
ventrimaculatus. Leptodactylus mystacinus has distinct dorsolateral folds (at least indicated by color pattern), dorsolateral folds are indistinct or lacking in bufonius and troglodytes. Leptodactylus mystacinus occurs east of the Andes, labrosus and ventrimaculatus occur west of the Andes.

Adult Characteristics ( $N=87$ ). -Dorsum uniform, striped, or slightly spotted (fig. 1, A, C, J, K); no light mid-dorsal stripe; light lip stripe usually distinct $(86 \%)$, sometimes indistinct ( $14 \%$ ), more females ( $100 \%$ ) with distinct lip stripes than males $\left(X^{2}=4.10, P=.04\right)$; dark suborbital bar absent; light stripe on posterior face of thigh usually absent ( $94 \%$ ), rarely indistinct ( $6 \%$ ), presence not sexually dimorphic ( $X^{2}=1.17, P=.28$ ); tibia barred; usually 2 or 4 well defined dorsolateral folds; dorsal surface of tibia with many or scattered distinct white tubercles; posterior surface of tarsus almost always ( $94 \%$ ) with many or scattered distinct white tubercles, absence not sexually dimorphic ( $X^{2}=.004, P=.95$ ); sole of foot usually with distinct scattered or many white tubercles ( $75 \%$ ), sometimes absent ( $25 \%$ ), presence not sexually dimorphic ( $X^{2}=.41, P=.52$ ); male SVL 53.0 $\pm 4.6 \mathrm{~mm}$, female $56.5 \pm 2.7 \mathrm{~mm}$, females larger than males ( $\mathrm{F}_{1,85}=12.59, P<.001$ ); male head length/ SVL ratio $.371 \pm .013$, female $.358 \pm .013$, male head


Figure 61. Distribution map of Leptodactylus mystaceus (squares) and notoaktites (triangles).
longer than female ( $\mathrm{F}_{1,85}=18.17, P<.001$ ); male head width/SVL ratio $.351 \pm .015$, female $.348 \pm .013$, not sexually dimorphic ( $\mathrm{F}_{1}, 85=.88, P=.94$ ); male femur/SVL ratio $.388 \pm .023$, female $.389 \pm .024$, not sexually dimorphic ( $\mathrm{F}_{1,85}=1.52, P>.05$ ); male tibia/ SVL ratio $.421 \pm .013$, female $.416 \pm .018$, not sexually dimorphic ( $\mathrm{F}_{1,85}=2.34, P>.05$ ); male foot/ SVL ratio $.428 \pm .021$, female $.423 \pm .022$, not sexually dimorphic ( $\mathrm{F}_{1,85}=1.06, P>.05$ ).

Larval Characteristics. -Sazima (1975) described and figured the larvae.

Mating Call.-Dominant frequency modulates between $2200-2500 \mathrm{hz}$; note duration 0.1 s ; note repetition rate $5-6.5$ per second (Barrio 1965).

Karyotype.-Diploid number 22; 7 pair median, 3 pair submedian, 1 pair subterminal; secondary constriction on chromosome pair 11 (Bogart 1974).

Distribution.-Interior Brasil to and including the Gran Chaco, coastal southeast Brasil and Argentina (fig. 62).

ARGENTINA. BUENOS AIRES: Buenos Aires, MACN 4150.

CHACO: Ciervo Petizo, IML 243.
ENTRE RÍOS: Concepción del Uruguay, MACN 4530.
JUJUY: Sobre ruta entre Rio San Francisco y La Realidad ( 5 km from Yuto), IML 1272; Ruta Yuto-Ledesma, IML 1273.

LA PAMPA: Conelo, MACN 1166; General Pico, MACN 4479, 4505, 4513.
MISIONES: Dos de Mayo, IML 2356; Puerto Piray, km 18,

median, 3 y constricluding the :ntina (fig.
es, MACN

N 4530 .
-a Realidad IML 1273. ico, MACN ', km 18,


Figure 62. Distribution map of Leptodactylus mystacinus (triangles) and poecilochilus (squares).

MACN 2956; Río Paranay, FMNH 9462-66; 10870; San Javier, Puerto Londero, MACN 2072; Santa Ana, MACN 5548.

SALTA: Campo Aguaray, IML 1473; near Hickmann, IML 148, 433.

SANTIAGO DEL ESTERO: Caspi Corral, 96 km , IML 2188; Pajares, Simbol, Chichi Huarcunay y Guanaco, Depto. Atamisqui, IML 2230.

TUCUMÁN: Río Urueña, near border of Salta, IML 1428.
BOLIVIA. SANTA CRUZ: Buenavista, MCZ 12897, UMMZ 66479 (2), 66480, 66488; El Carmen, CM 36097, MCZ 29986; Río Surutú, CM 3811.

BRASIL. BAHIA: Maracás, WCAB 31825-28.
DISTRITO FEDERAL: Brasilia, USNM 121292.
GOIÁS: Anapolis, AMNH 43847; Flôres, USNM 121270. MATO GROSSO: Aquidauana, MZUSP 15800.
MINAS GERAIS: Lapa Vermelha, Lagoa Santa, MZUSP 15877; Urucuia Riv., first waterfall, Buritís, MZUSP 25069.

PARANÁ:St. Antonio da Platina, MZUSP 24155
RIO DE JANEIRO: Niteroi, Saco de São Francisco, AMNH 20308 USNM 99121.

RIO GRANDE DO SUL: Albardão, WCAB 16843; Bagé, WCAB 3878; 18 km S Farroupilha, FMNH 80374; Montenegro, MZUSP 16050; Pôrto Alegre, FMNH 80360-371, KU 92921-23, MZUSP 16048-49, 21688-89, WCAB 3876; 39 km N Rio Pardo, FMNH 80372-73; Sta. Maria, MZUSP 24153-54, USNM 121272, WCAB 5259; São Leopoldo, MZUSP 25478; São Lourenço, MZUSP 91, 1970; Viamão, MCZ 32695-96, WCAB 7137-178; Vila Nova, São Sepé, MZUSP 23707-08.

SANTA CATARINA: Nova Teutônia, MZUSP 8694-98.
SÃO PAULO: Botacatú, WCAB 4351; Ermelindo Matarazzo, MZUSP 8106; Faveiro, MZUSP 25423-26; Guapiara, WCAB 6119; Itu, FMNH 83235, KU 92923-24, WCAB 4306, 4311, 4314, 6223, 8230; Nova Itaperuna, WCAB 13660; Pe-
rus, MZUSP 49; Rio Pardo, Botucatú, MZUSP 7132; Santa Branca, Rio Paraíba, MZUSP 25456; Santo Antonio do Pinhal, MZUSP 14907; São Paulo, USNM 121293.

URUGUAY. CANELONES: Carrasco, MZUSP 22640-41.
DURAZNO: 18 km NE Paloma, Arroyo del Estado, CM 57041-42.

LAVALLEJA: Río de Averías, Depto. Minas, FMNH 10400 01.

MALDONADO: Maldonado, FMNH 10155; Sierra de Animas, WCAB 7273.

ROCHA: 22 km SE Lascano, AMNH 71177.
TACUAREMBO: 3 km NE Tambores, Pozo Hondo, CM 55392-93.

30 Y 3: 8 mi E 30 y 3, FMNH 10465, 10470-72; Quebrada de los Cuervos, 45 km N 30 y 3, FMNH 10500.

## Leptodactylus notoaktites new species

Figure 63
Holotype: MZUSP 25428, a female from Brasil; São Paulo, Iporanga. Collected by Nelson Papavero on 2 November 1963.

Diagnosis. -The species having a combination of a distinct light stripe on the posterior face of the thigh and a smooth posterior surface of the tarsus in some or all individuals are amazonicus, fuscus, geminus, gracilis,


Figure 63. Dorsal view of the holotype of Leptodactylus notoaktites.
laurae, longirostris, marambaiae, mystaceus, notoaktites, and poecilochilus. Leptodactylus notoaktites has a barred tibial pattern, the dorsal surface of the tibia has light stripes in geminus, gracilis, and marambaiae. Only individual notoaktites with a mid-dorsal light stripe have 6 dorsolateral folds; all fuscus and laurae individuals have 6 dorsolateral folds. Leptodactylus notoaktites has a shorter leg (e.g. male foot/SVL ratio $.587 \pm .033$, female $.583 \pm .036$ ) than laurae (male foot/SVL ratio $.649 \pm .039$, female $.628 \pm .028$ ). Leptodactylus notoaktites does not have the dorsal blotching of $L . f$ fuscus. Most mystaceus have white tubercles on the posterior surface of the tarsus. Some individual notoaktites have white tubercles on the sole of the foot, the sole of the foot is smooth in longirostris and poecilochilus. Leptodactylus notoaktites occurs in southeast Brasil, longirostris and poecilochilus are found in northern South America. Some notoaktites have a smooth sole of the foot and/or a light mid-dorsal stripe, all amazonicus have white tubercles on the sole of the foot and lack light mid-dorsal stripes; amazonicus occurs throughout the Amazon basin.

Description of Holotype. -Snout rounded-subelliptical from above, rounded in profile; canthus rostralis indistinct; loreal slightly concave; tympanum distinct, greatest diameter about $1 / 2$ eye diameter; vomerine teeth in slightly arched series posterior to choanae; finger lengths in order of decreasing size $\mathrm{I} \simeq \mathrm{III}>\mathrm{II} \simeq \mathrm{IV}$, I $\gg \mathrm{II}$; inner metacarpal tubercle oval, smaller than rounded outer metacarpal tubercle; dorsum smooth above anteriorly, warty on sides and posteriorly; 1 pair of distinct dorsolateral folds from eye to groin, 1 pair of indistinct lateral folds; ventral texture smooth; belly disk fold distinct; toe tips not expanded; toes free, lacking fringe or web; subarticular tubercles moderately developed; outer metatarsal tubercle small, round, about $1 / 4$ oval inner metatarsal tubercle; tarsal fold indistinct; no metatarsal fold; posterior surface of tarsus smooth; sole of foot with 1 or 2 indistinct white tubercles.

SVL 56.1 mm , head length 20.6 mm , head width 18.4 mm , interorbital distance 3.7 mm , eye-nostril distance 5.0 mm , femur 27.0 mm , tibia 31.5 mm , foot 31.8 mm .

Dorsum brown with faint darker markings including an interorbital blotch and dorsal chevron; dorsolateral folds light outlined posteriorly; upper lip edge dark, bordered above by distinct light stripe from tip of snout passing under eye to angle of. jaw; dark canthal stripe above light lip stripe from tip of snout to eye; venter immaculate; posterior surface of thigh mottled above, dark below with distinct light longitudinal stripe.

Etymology. -From the Greek notos, south, and aktites, coast dweller, in reference to the geographic distribution of the species in Brasil.

Remark. -This species was analyzed as south coastal mystaceus.

Adult Characteristics ( $N=18$ ).-Dorsum spotted,
s, notoakiktites has e tibia has ziae. Only itripe have ndividuals iktites has $7 \pm$.033, SVL ratio ctylus noL. fuscus. posterior tites have ole of the ilus. Leprasil, lonlem South ole of the nazonicus and lack hroughout
subellipti, stralis in1 distinct, arine teeth ae; finger $I I \simeq I V$, laller than roth above air of dis,air of inbelly disk 2, lacking ely devel, about $1 / 4$ istinct; no ooth; sole
ead width ye-nostril mm , foot
including rsolateral dark, bor-- of snout thal stripe ye; venter ed above; ipe. 1 , and $a k$ aphic disth coastal a spotted,
blotched, or striped (fig. 1, A, C, O, striped pattern not figured); light mid-dorsal stripes present in $11 \%$ of individuals, presence not sexually dimorphic (Fisher's exact test $P=1.0$ ); light upper lip stripe usually distinct ( $78 \%$ ), sometimes indistinct ( $22 \%$ ), distinctiveness not sexually dimorphic (Fisher's exact test $P=.29$ ); no dark suborbital bar; distinct light stripe on posterior face of thigh present ( $100 \%$ ); tibia barred; upper surface of tibia lacking white tubercles; posterior surface of tarsus lacking white tubercles $(100 \%)$; sole of foot usually with scattered or very few white tubercles ( $78 \%$ ), sometimes absent ( $22 \%$ ), presence not sexually dimorphic (Fisher's exact test $P=1.0$ ); male SVL $47.4 \pm 3.4 \mathrm{~mm}$, female $49.1 \pm 3.0 \mathrm{~mm}$, not sexually dimorphic ( $\mathbf{F}_{1,16}=1.21$, $P>.05$ ); male head length/SVL ratio $.368 \pm .014$, female $.375 \pm .011$, not sexually dimorphic ( $\mathrm{F}_{1,16}=$ $1.60, P>.05$ ); male head width/SVL ratio $.336 \pm$ .013 , female $.334 \pm .014$, not sexually dimorphic ( $F_{1,16}$ $=.10, P>.05$ ); male femur/SVL ratio $.470 \pm .037$, female $.450 \pm .032$, not sexually dimorphic ( $\mathrm{F}_{1,16}=$ $1.57, P>.05)$; male tibia/S VL ratio $.533 \pm .024$, female $.549 \pm .021$, not sexually dimorphic ( $\mathrm{F}_{1,16}=$ $2.11, P>.05$ ); male foot/SVL ratio $.587 \pm .033$, female $.583 \pm .036$, not sexually dimorphic ( $\mathrm{F}_{1,16}=.04$, $P>.05$ ).

Larval Characteristics.-Unknown.
Mating Call. - Unknown.
Karyotype. -Unknown.
Distribution.-Southeast Brasil (fig. 61).
BRASIL. PARANÁ: Paranaguá, WCAB 35170.
SANTA CATARINA: Colonia Hansa, Joinville, MZUSP 459, 1295; Humboldt ( $=$ Corupá), AMNH 15555; Rio Vermelho, WCAB 6717-723, 7929; Santa Luzia, prope Serra do Mar, MNRio 2148; São Bento, USNM 97176-78.

SÃO PAULO: Engenheiro Ferraz, MZUSP 25420; Iporanga, MZUSP 24149-150, 25428; Piraçununga, Cachoeira de Emas, MNRio 2107.

## Leptodactylus poecilochilus (Cope) 1862

Cystignathus poecilochilus Cope 1862:156-157. (Type locality, Colombia; Antioquia, Turbo. Holotype USNM 4347, male.)
Leptodactylus quadrivittatus Cope 1893:339-340. (Type locality, Costa Rica; Puntarenas, Buenos Aires. Holotype apparently lost.)
Leptodactylus maculilabris Boulenger 1896:404-405. (Type locality, Costa Rica; Guanacaste, Bebedero. Holotype BMNH 94.11.15.27.)
Leptodactylus diptychus Boulenger 1918:431. (Type locality, Andes of Venezuela. Holotype BMNH 94.8.31.11, female.)

Diagnosis. - The species having a combination of a distinct light stripe on the posterior surface of the thigh and smooth surfaces on the posterior tarsus and sole of the foot in some or all individuals are fuscus, geminus, gracilis, laurae, longirostris, marambaiae, mystaceus, notoaktites, and poecilochilus. The dorsal surface of the tibia lacks light longitudinal stripes in poecilochilus, such stripes are present in geminus, gracilis, and mar-
ambaiae. Only individuals of poecilochilus with light mid-dorsal stripes have 6 dorsolateral folds (fig. 64), all individuals of fuscus and laurae have 6 dorsolateral folds. The leg of poecilochilus is shorter (e.g. male foot/ SVL ratio $.514 \pm .029$, female $.508 \pm .029$ ) than laurae (male foot/SVL ratio $.649 \pm .039$, female .628 $\pm .028$ ). Leptodactylus poecilochilus lacks the scattered dorsal blotches characteristic of fuscus, does not have a light lip stripe, and often has a dark suborbital bar (fig. 57). No longirostris, mystaceus, or notoaktites have a dark suborbital bar and individuals often have distinct light lip stripes.
Adult Characteristics ( $N=133$ ). -Dorsum spotted, spots sometimes elongate, rarely fused (fig. 1, A, B, C, D, E) light mid-dorsal stripe present in $13 \%$ of individuals, presence not sexually dimorphic ( $X^{2}=.35, P=$ .55); lip stripe indistinct; dark suborbital bar usually present ( $67 \%$ ) or often absent ( $33 \%$ ); light stripe on posterior face of thigh usually distinct ( $77 \%$ ), sometimes indistinct ( $21 \%$ ), rarely absent ( $2 \%$ ), expression not sexually dimorphic ( $X^{2}=1.31, P=.52$ ); tibia barred; usually 2 or 4 well defined dorsolateral folds present, 6 dorsolateral folds present when light mid-dorsal stripe present; dorsal surface of tibia lacking white tubercles; posterior surface of tarsus almost always lacking white tubercles ( $99 \%$ ), presence not sexually dimorphic $\left(X^{2}=\right.$ $.07, P=.80$ ); sole of foot almost always lacking white tubercles (93\%), presence not sexually dimorphic ( $X^{2}=$ $.63, P=.43$ ); male SVL $44.8 \pm 2.2 \mathrm{~mm}$, female 45.9 $\pm 3.4 \mathrm{~mm}$, not sexually dimorphic ( $\mathrm{F}_{1,131}=3.75, P$ $>.05$ ); male head length/SVL ratio $.380 \pm .010$, female $.376 \pm .011$, not sexually dimorphic ( $\mathrm{F}_{1}, 131=$ $3.48, P>.05$ ); male head width/SVL ratio $.340 \pm$ .013 , female $.340 \pm .011$, not sexually dimorphic ( $\mathrm{F}_{1,131}$ $=.03, P>.05$ ); male femur/SVL ratio $.424 \pm .024$, female $.427 \pm .025$, not sexually dimorphic, ( $\mathrm{F}_{1,131}=$ $.32, P>.05$ ); male tibia/SVL ratio $.489 \pm .024$, female $.488 \pm .024$, not sexually dimorphic ( $\mathrm{F}_{1,131}=.06, P$ $>.05$; male foot/SVL ratio $.514 \pm .029$, female .508 $\pm .029$, not sexually dimorphic ( $\mathrm{F}_{1,131}=1.47, P>$ .05).

Larval Characteristics. - Eye diameter 9-14\% headbody length; oral disk width $15-27 \%$ head-body width; oral papilla gap $45-65 \%$ oral disk width; 64-142 denticles on one side of split tooth row anterior to beak; head-body length $35-45 \%$ total length; total length, stage 41, 37 mm (Heyer 1970b, figs. 10, 15, 20).
Mating Call. -Dominant frequency môdulates from $350-550 \mathrm{hz}$; call lacks harmonic structure; note non-pulsatile; note duration 0.055 to 0.080 s ; note repetition rate. 1.7 per second (Straughan and Heyer 1976).

Karyotype.-Unknown.
Distribution. -Lowlands of Costa Rica to north coastal South America as far as Venezuela (fig. 62).

COLOMBIA. ANTIOQUIA: Belén, $23 / 4 \mathrm{~h}$ upstream Pto. Palacios, Río Arquia, LACM 51090-1110, 51138-148; Finca Chibiguí, Río Arquia, LACM 51112-137; Finca Los Llanos,

176 (6),
Grande
CRE 71
8279; n
(2), 65

Villa N
SSE Sa del Ge PAF
tun, U:
(3), 12

Río Si
Río M
9161; :

Río Arquia, LACM 51111; Pto. Palacios, Río Arquia, LACM 51089; Villa Arteaga, USNM 146437-38. CHOCO: Golfo de Urabá, Unguía, FMNH 63846. CÓRDOBA: Río Manso, trib. Río Sinú, USNM 151034058.

GUAJRA: Río Barbacoa, UMMZ 54599, 54602-03.
MAGDALENA: Fundación, UMMZ 48505-06, 48508, 51106, USNM 102408, 102410; Río Frio, MCZ 16069; Valencia, UMMZ 54604-08.

NORTE DE SANTANDER: Río Zulia, USNM 147070, 147072-73.

COSTA RICA. ALAJUELA: 3 km W La Fortuna, CRE 8078.

GUANACASTE: Arenal, CRE 6254; Finca Comelco, 30 km NNW Cañas, UMMZ 131908; near Liberia, CRE 8207; near Playa del Coco, CRE 8143, UMMZ 129248 (2); Río Sandillal, UMMZ 131909; 2 mi W Santa Cruz, CRE 8233; Hacienda Taboga, CRE 3086.

HEREDIA: Cariblanco, FMNH 175200.
PUNTARENAS: Coto, km 47 on rail from Golfito, CRE 176 (6), 178 (6), 180 (11); Finca Helechales, 15 km NE Potrero Grande, CRE 3126 (2), 8267-68; 6 km ESE Golfito, 10 m , CRE 7105; 8 km NE Potrero Grande, Finca del Sr. Treño, CRE 8279; near Rincón de Osa, CRE 705 (4), 750 (2), 3108, 6391 (2), 6545, 7228, 7236, LACM 53998-99, UMMZ 129258 (2); Villa Neily, 75 m, CRE 179, 8031, 8039.

SAN JOSÉ: Pozo Azul de Pirrís, MCZ 7997-8001; 3 mi SSE San Isidro del General, CRE 8001; 13 mi WSW San Isidro del General, on Dominical road, 710 m , CRE 687.

PANAMA. CANAL ZONE: Cocoli, USNM 193340; Gatun, USNM 54177; near Madden Dam, FMNH 174061; near Paca, Military Road, FMNH 43577; Rosseau, KU 67960; Summit, MCZ 21834.

CHIRIQUÍ: Progresso, UMMZ 58267-272, 58275-283, USNM 118673.

COCLÉ: 1 km NE El Caño, 40 m , FMNH 22986.
DARIEN: Camp Creek, Camp Townsend, AMNH 41022; Ortiga, FMNH 170465, 170467; Río Canglón, UMMZ 125021 (3), 125022-29; Río Lara, FMNH 170304, 170392, 170436; Río Silugandí, UMMZ 113120-22 (3), 113123; Río Tuira at Río Mono, KU 116829-831; Sambu Valley, Río Esaupe, MCZ 9161; Santa Fe Camp, FMNH 170269, 170308; S 6 VIII Camp, FMNH 170343.

PANAMÁ: Cermeño, MCZ 24880; Cerro Campana, FMNH 60500, MCZ 82072, USNM 13970I; Río Itarare, FMNH 28856; Tapia, AMNH 18931.

SAN BLAS: SG VIII site, FMNH 170374.
VERAGUAS: Mojara, USNM 129841-42.
VENEZUELA. ARAGUA: near Maracay, Rancho Grande, AMNH 70688; near Ocumare, UMMZ 122374.

FALCÓN: 5 km S Palma Sola, UMMZ 55554; Soute Parriji, MCZ 25989; 19 km NW Urama, km 40, USNM field 1808, 5217, 5243, 5246.

GUÁRICO: Hato La Palmita, USNM 162702.
TRUJLLLO: Sabana de Mendoza, UMMZ 57483.

## Leptodactylus troglodytes A. Lutz 1926

Leptodactylus troglodytes A. Lutz 1926:149-150, plate 32, fig.
12. (Type locality, Brasil; Pernambuco, Procedencia.

Holotype Adolfo Lutz collection, no number, female.)

Diagnosis.-The species lacking a distinct thigh stripe and having distinct white tubercles on the posterior surface of the tarsus in some or all individuals are albilabris, bufonius, labrosus, mystacinus, troglodytes, and ventrimaculatus. Leptodactylus albilabris usually has at
least an indication of a light stripe on the posterior surface of the thigh. Leptodactylus troglodytes lacks distinct dorsolateral folds; distinct dorsolateral folds (indicated at least by color pattern) occur in albilabris, labrosus, mystacinus, and ventrimaculatus. Leptodactylus troglodytes and bufonius are morphologically similar and have similar dorsal patterns (fig. 65). All individuals of troglodytes have distinct white tubercles on the sole of the foot, almost all bufonius have smooth surfaces on the sole of the foot. Leptodactylus troglodytes occurs in northeast Brasil, bufonius has a distribution centered upon the Gran Chaco.
Remark.-This is the species referred to as "northern bufonius" in the morphological analysis.
Adult Characteristics ( $N=42$ ). -Dorsum with chevrons, spots, or blotches (fig. 1, A, B, C, G, L, N); no light mid-dorsal stripe; no light upper lip stripe; dark suborbital bar always present; light stripe on posterior face of thigh absent ( $100 \%$ ); tibia barred; dorsolateral folds usually absent, 2 weak indistinct folds rarely present; dorsal surface of tibia with many distinct white tubercles; posterior surface of tarsus with distinct white tubercles ( $100 \%$ ); sole of foot with white tubercles ( $100 \%$ ); male SVL $48.8 \pm 2.2 \mathrm{~mm}$, female $49.9 \pm 1.8$ mm , not sexually dimorphic ( $\mathrm{F}_{1,40}=2.67, P>.05$ ); male head length/SVL ratio $.385 \pm .008$, female .374 $\pm .010$, male head longer ( $\mathrm{F}_{1,40}=16.17, P<.001$ ); male head width/SVL ratio $.344 \pm .011$, female .339 $\pm .011$, not sexually dimorphic ( $\mathrm{F}_{1,40}=1.92, P>$ .05); male femur/SVL ratio $.400 \pm .020$, female .393 $\pm .015$, not sexually dimorphic ( $\mathrm{F}_{1,40}=1.38, P>$ .05); male tibia/SVL ratio $.406 \pm .012$, female $.397 \pm$ .014, male tibia longer ( $\mathrm{F}_{1,40}=5.51, .01<P<.025$ ); male foot/SVL ratio $.395 \pm .011$, female $.386 \pm .016$, male foot longer ( $\mathrm{F}_{1,40}=5.38, .025<P<.05$ ).

Larval Characteristics. -Unknown.
Mating Call. -Dominant frequency modulates from $2600-3200 \mathrm{hz}$ (fig. 66); call without harmonic structure (fig. 67); call not pulsed; note duration .042 s ; note repetition rate 1 per second.

Karyotype. -Unknown.
Distribution.-Northeast Brasil (fig. 68).
BRASIL. BAHIA: Andaraí, WCAB 43766-67; Barreiras, UMMZ 109980-81 (2); Carnaíba, WCAB 43867; Cocorobó, MZUSP 38278-79; Feira de Santana, WCAB 44085; Jeremoabo, MZUSP 38167; Maracás, WCAB 31813-824; Salvador, MZUSP 10715.

CEARÁ: Açude Amanarí, Maranguapé, MZUSP 13589; Fortaleza, WCAB 19149.

GOIÁS: Cana Brava, MZUSP 20441-42.
MINAS GERAIS: Rio Pandeiros, MZUSP 24695, USNM 121300.

PARAÍBA: Piancó, WCAB 3626, 4976.
PERNAMBUCO: Bonito, UMMZ 132461; Exú, WCAB 39218.

PIAUÍ: 35 km N Valença, MZUSP field 750647-652.
RIO GRANDE DO NORTE: Natal, Areia Preta, USNM 97048-49; Ponta Negra, MZUSP 25017.

SERGIPE: Areia Branca, MZUSP 37825-837.



FIGURE 66. Sonagram of the mating call of Leptodactylus troglodytes, narrow band filter. Vertical scale marks at 1000 hz intervals. Horizontal scale mark at 1 s . Specimen from Brasil, Andaraí, air temperature $24^{\circ} \mathrm{C}$ (WCAB tape).


Figure 67. Strip chart record of the mating call of Leptodactylus troglodytes. Line equals 0.01 s . See legend of Figure 66 for specimen data.

## Leptodactylus ventrimaculatus Boulenger 1902

Leptodactylus ventrimaculatus Boulenger 1902:53. (Type locality, Ecuador, Bulun, $160^{\prime}$. Lectotype BMNH 1947.2.17.78, female.)

Diagnosis. - The species having a combination of no light stripe on the posterior surface of the thigh and distinct white tubercles on the posterior surface of the tarsus and sole of foot (fig. 69) in some or all individuals are bufonius, labrosus, mystacinus, troglodytes, and ventrimaculatus. Leptodactylus ventrimaculatus has distinct dorsolateral folds (indicated at least by color pattern), dorsolateral folds are absent or indistinct in bufonius and troglodytes. Some individuals of mystacinus lack white tubercles on the sole of the foot; $L$. mystacinus occurs east of the Andes, $L$. ventrimaculatus occurs west of the Andes along the wet coastal regions of Colombia to midEcuador. Most $L$. labrosus have a smooth sole of the foot (fig. 69); labrosus occurs along the dry west coasts of South America from mid-Ecuador to Peru, including the northem interandean valley of northerm Peru.

Adult Characteristics ( $N=38$ ). -Dorsum spotted, striped, or rarely uniform (fig. 1, A, B, C, J, K, striped
pattern not figured); no light mid-dorsal stripe; no light upper lip stripe; dark suborbital bar almost always present; light stripe on posterior face of thigh almost always absent $(97 \%)$, rarely indistinct ( $3 \%$ ), presence not sexually dimorphic ( $X^{2}=.03, P=.87$ ); tibia barred; usually 2 dorsolateral folds present; dorsal surface of tibia with many white tubercles; posterior surface of tarsus with many white tubercles ( $100 \%$ ); sole of foot with scattered or very few white tubercles (at least some tubercles present in $100 \%$ of study sample); male SVL $50.4 \pm 3.5 \mathrm{~mm}$, female $51.9 \pm 4.8 \mathrm{~mm}$, not sexually dimorphic ( $\mathrm{F}_{1,36}=1.23, P>.05$ ); male head length/ SVL ratio $.363 \pm .013$, female $.360 \pm .015$, not sexually dimorphic ( $\mathrm{F}_{1,36}=.38, P>.05$ ); male head width/SVL ratio $.343 \pm .010$, female $.341 \pm .010$, not sexually dimorphic ( $\mathrm{F}_{1,36}=.11, P>.05$ ); male femur/ SVL ratio $.389 \pm .023$, female $.384 \pm .024$, not sexually dimorphic ( $\mathrm{F}_{\mathrm{L}, 36}=.35, P>.05$ ); male tibia/SVL ratio $.420 \pm .019$, female $.409 \pm .020$, not sexually dimorphic ( $\mathrm{F}_{1,36}=2.50, P>.05$ ); male foot/S VL ratio $.457 \pm .022$, female $.447 \pm .021$, not sexually dimorphic ( $\mathrm{F}_{1,36}=2.0, P>.05$ ).

Larval Characteristics. -Unknown.


Figure 68. Distribution map of Leptodactylus troglodytes (squares) and ventrimaculatus (triangles).

## Mating Call.-Unknown.

Karyotype. -Unknown.
Distribution.-Western South America, primarily west of the Andes, from mid-Ecuador to northem Peru (fig. 68).

COLOMBIA. CAUCA: Quebrada Guanguí, $1 / 2 \mathrm{~km}$ above Río Patía (upper Saija drainage), 100-200 m, AMNH 88529. CHOCÓ: 2 km above Playa de Oro, upper Río San Juan, AMNH 87124-132; Quebrada Bochoramá, $180-190 \mathrm{~m}$, LACM 44383; upper Río Buey, $110-160 \mathrm{~m}$, LACM 44381.

NARINO: Imbilí, Río Mira, USNM 147457-480; near La Guayacana, LACM 50173-74; Río Satinga, USNM 14748385; N Tumaco, Río Rosario, USNM 147488-89.

VALLE: Buenaventura (islet in Pacific), USNM 14707778.

ECUADOR. ESMERALDAS: 1 km N Cachaví, USNM 196757 (7), 196758 (2); Hacienda Equinox, 30 km NNW Santo Domingo de los Colorados, 1000 ft , USNM $196755 ; 1 \mathrm{~km}$ NW Lita Station, USNM 196756; Río Pilatón, WCAB 276.

IMBABURA: Cachaco, USNM 196769; Lita, $520 \mathrm{~m}, \mathrm{KU}$ 132805-06.

PICHINCHA: Hacienda Espinosa, 9 km W Santo Domingo de los Colorados, road to Chone, CAS-SU 10455-466; 5 km E La Palma, KU-WED 48227-233; 1 km E Mindo, farm of Julio Goetschel, 1400 m, USNM 196764-66 (2); Río Blanco, near mouth of Rio Yambi, 700 m , USNM 196767; Río Toachi, USNM 196768; near Santo Domingo de los Colorados, KU 117794, 146186-87, USNM 196759-60 (6), 196761 (2), $196762(2), 196763$.


Figure 69. Tarsal and foot textures. Left, tarsus smooth, foot scattered with white tubercles (based on L. fuscus, LACM 92015, extreme development of foot tubercles for this species). Center, tarsus with many white tubercles, foot smooth (based on $L$. labrosus, LACM 49161). Right, tarsus and foot with many white tubercles (based on L. ventrimaculatus, AMNH 88529).

## AN ARTIFICIAL KEY TO THE ADULT MEMBERS OF THE LEPTODACTYLUS FUSCUS GROUP

This key is designed to be used in conjunction with the diagnoses. For those species demonstrating variation in key characters, the most frequent condition is presented in the key; the diagnoses incorporate the range of variation of key characters.

1 A. Dorsal surface of tibia with distinct light longitudinal stripes (figs. 2, 48) ......................... 17
1 B. Dorsal surface of tibia barred, lacking light longitudinal stripes (figs. 2, 48) ........................ 2
2 A. Posterior surface of thigh lacking distinct light longitudinal stripe .......................... 3
2 B. Posterior surface of thigh with distinct light longitudinal stripe ............................. 7
3 A. Dorsolateral folds indistinct or absent ..................................................................... . . 4
3 B. Dorsolateral folds distinct (indicated at least by color pattern) ....................................... 5
4 A. Sole of foot smooth, Chacoan distribution ............................................. . . . bufonius
4 B. Sole of foot with white tubercles, northeast Brasil ................................. troglodytes
5 A. Light upper lip stripe usually distinct, never a dark suborbital bar (fig. 57) . . . . . . . . . . . . . mystacinus
5 B. No light upper lip stripe, usually a dark suborbital bar (fig. 57)
6
6 A. Sole of foot smooth, dry west coastal South America and interandean valley in northern Peru labrosus
6 B. Sole of foot with white tubercles, wet west coastal Colombia to mid-Ecuador ventrimaculatus

7 A. Posterior surface of tarsus smooth (fig. 69) . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 8
7 B. Posterior surface of tarsus with distinct white tubercles (fig. 69) ................................. . . 13
8 A. Sole of foot with white tubercles (fig. 69) ...................................................... 9
8 B. Sole of foot smooth (fig. 69) ..................................................................... 10
9 A. No individuals with a mid-dorsal light stripe, Amazonian distribution .................. amazonicus
9 B . Some individuals with a mid-dorsal light stripe, southeast Brasil ........................ notoaktites
10 A. Leg noticeably long, all individuals with a mid-dorsal light stripe, southeast Brasil .... laurae
10 B . Leg of normal proportions, most individuals without a mid-dorsal light stripe ............. 11
11 A. All individuals with 6 dorsolateral folds, dorsum usually with many irregular spots or blotches, widespread
 Most individuals with 2 or 4 dorsolateral folds, only individuals with a mid-dorsal light stripe having 6 dorsolateral folds, dorsum usually with a few more or less regular spots or blotches, Middle America and northern South America 12

12 B. Light lip stripe indistinct, dark suborbital bar often present, Costa Rica through coastal Venezuela poecilochilus
13 A. Dorsolateral folds indistinct or absent ..... 14
13 B. Dorsolateral folds distinct (at least indicated by color pattern) ..... 15
14 A. Texas throughout Middle America along coastal northern South America to Venezuela fragilis14 B. Gran Chaco and coastal Brasillatinasus
15 A. Dorsal surface of tibia smooth, Chacoan distribution ..... elenae
15 B. Dorsal surface of tibia with white tubercles ..... 16
16 A. Tibia (male mean $43 \%$ SVL, female $44 \%$ ) and foot (male mean $49 \%$ SVL, female $50 \%$ ) shorter, West Indies ..... albilabris
16 B. Tibia (male mean $51 \%$ SVL, female $52 \%$ ) and foot (male and female mean $55 \%$ SVL) longer,east coastal Brasil mystaceus
17 A. Leg shorter, tibia $50 \%$ SVL marambaiae
17 B. Leg longer, tibia $52-62 \%$ SVL geminus \& gracilis

## THE SIGNIFICANCE OF SEXUAL DIMORPHISM IN MEMBERS OF THE LEPTODACTYLUS FUSCUS GROUP

A surprising amount of sexual dimorphism was encountered in the morphological analysis. Many frogs demonstrate sexual dimorphism in size and various other secondary sexual characteristics, but to my knowledge, no one previously has demonstrated sexual dimorphism in limb proportions in frogs. The characters involved in secondary sexual dimorphism correlate with aspects of ecology and breeding biology in certain cases. Against the patchy background of available ecological information, tentative predictions can be made for some species for which ecological data are as yet unavailable.

In the species accounts section, sexual dimorphism was established at a significance level of $5 \%$. Due to the degree of measurement error combined with the small sample sizes available, the $10 \%$ level of significance is used here to establish presence of sexual dimorphism in measurement and ratio characters (Table 4). Leptodactylus geminus is not included in this discussion as no validated specimens were available for analysis. The available series of $L$. marambaiae is too small to analyze.

Size. -In many species of frogs the female is larger than the male. The usual explanation for this phenomenon is that the larger female size allows for a greater clutch size and hence an increase in reproductive effort. In Leptodactylus elenae, fragilis, fuscus, gracilis, la-
brosus, mystaceus, notoaktites, poecilochilus, troglodytes, and ventrimaculatus, the sexes are not dimorphic with respect to size. Both sexes of all these species must be under some environmental or developmental constraint selecting for the same size. What the constraint(s) is, is not known at present.

Head Length.-Members of the fuscus group deposit their eggs in underground chambers. In at least some species, these incubating chambers are constructed by the male. Males of the following species have either been observed constructing an incubating chamber, or have been observed calling in association with an incubating chamber: amazonicus (pers. obs.), bufonius (Philibosian, et. al. 1974), fragilis (Dixon and Heyer 1968), mystacinus (Sazima 1975). In all of these species the male's head is longer than the female's, reflecting the development of a rigid chisel-like snout that is used in the construction of the incubating chamber. The chamber is apparently excavated only in damp ground by males of these species. On the basis of snout morphologies, albilabris, latinasus, laurae, longirostris, poecilochilus, and troglodytes males are also predicted to excavate the nest chamber in damp ground. For those species in which the head lengths of the sexes are the same, one would predict either that both sexes were involved in formation of the incubating chamber (or burrow construction) or that naturally occurring depressions are used for deposition of the foam nest with little modification on the part of the males. Based on head mor-

Table 4
Occurrence of sexual dimorphism in members of the fuscus group. $-=$ no sexual dimorphism, $F=$ sexual dimorphism present, female elements longer or more distinct, $M=$ sexual dimorphism present, male elements longer.

|  | Variables |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | SVL | Head Length | Head Width | Femur | Tibia | Foot | Mid-dorsal Stripe | $\underset{\text { Stripe }}{\text { Lip }}$ | Thigh Stripe | Tarsal Texture | Foot Texture |
| albilabris | F | M | - | F | F | F | - | - | - | - | - |
| amazonicus | F | M | M | - | F | F | - | F | - | - | - |
| bufonius | F | M | M | - | - | - | - | - | - | - | - |
| elenae | - | - | - | - | - | - | - | - | - | - | - |
| fragilis | - | M | M | F | F | F | - | - | - | - | - |
| fuscus | - | - | M | F | F | - | - | F | - | - | - |
| gracilis | - | - | M | - | - | - | - | - | - | - | - |
| labrosus | - | - | - | F | - | - | - | - | - | - | - |
| latinasus | F | M | - | - | - | - | -- | F | - | - | - |
| laurae | F | M | M | - | - | - | - | - | - | - | - |
| longirostris | F | M | - | - | F | - | - | - | F | - | - |
| mystaceus | - | - | - | F | - | - | - | - | - | - | - |
| mystacinus | F | M | - | - | - | - | - | F | - | - | - |
| notoaktites | - | - | - | - | - | - | - | - | - | - | - |
| poecilochilus | - | M | - | - | - | - | - | - | - |  | - |
| troglodytes | - | M | - | - | M | M | - | - | - | - | - |
| ventrimaculatus | - | - | - | - | - |  | - |  |  |  |  |

phology, labrosus is predicted to be a species in which both sexes are involved in the formation of incubating chambers or burrows. On the other hand, elenae, fuscus, gracilis, mystaceus, notoaktites, and ventrimaculatus are good candidates for foam nest deposition in natural depressions with little subsequent modification.

Head Width.-Not all species which show dimorphism of head length are also dimorphic with respect to head width, and vice versa. This suggests that head width is not associated with incubating chamber formation, but instead may be important with respect to such aspects as food niche separation or mating call broadcasting. Data are not available to test these hypotheses.

Limb Proportions. -The longer any of the hind limb elements are, the better a frog is at jumping (Zug 1972). Extreme jumping ability is usually associated with avoidance of vertebrate predators. The sexes of species demonstrating sexual dimorphism in leg length might have different abilities to escape predation, suggesting different selective forces operating on the two sexes. The most parsimonious explanation for the development of sexual dimorphism in hind limb length for members of the fuscus group takes into account: (1) relative fossoriality as relates to the general niche adaptations of the species; (2) fossoriality only in terms of incubating chamber formation by the males; and (3) exposure to above ground vertebrate predation. Members of the fuscus group appear to segregate into six groupings based on these three variables. The following groupings are not presented as groupings of fact, but rather as hypotheses which can account for limb length dimorphism. Hopefully, the hypotheses will focus attention on gathering data on differential predator success or exploration
of alternate hypotheses accounting for dimorphic limb lengths such as weight differences.

1. Fossorial niche, male under more vertebrate predatory pressure than female: L. troglodytes. This pattern suggests that most of the life activities of the species takes place under ground and the nest construction and/ or calling activity of the male are the longest above ground activities in the adult life history. The longer limb of the male results from selective pressures exerted by vertebrate predation.
2. Fossorial niche, both sexes responding in the same way to vertebrate predatory pressure: L. bufonius, mystacinus, ventrimaculatus. This pattern indicates that most life activities take place fossorially but that both species spend about the same amount of time in above ground activities.
3. Fossorial niche, shorter leg of male the result of selection for fossorial activity of incubating chamber construction, longer leg of female the result of selection from vertebrate predators when above ground: L. labrosus. This pattern suggests that while many of the activities of the species are fossorial, relatively more time is spent above ground than for the species in the preceding pattern.
4. Above ground niche, longer head and shorter leg of male the result of selection for fossorial activity of incubating chamber construction, longer leg of female the result of selection pressure from vertebrate predators: L. albilabris, amazonicus, fragilis, fuscus, longirostris, mystaceus. Rather than being primarily fossorial as in the previous patterns, members showing this pattern are active above ground and incubating chamber construction is an important male activity.
5. Above ground niche, longer head of male the re-
sult of selection for fossorial activity of incubating chamber construction, both sexes responding in same way to selection from vertebrate predators: L. latinasus, laurae, poecilochilus. This pattern implies that selection from vertebrate predators is not important (latinasus, short legs), very important (laurae, long legs in both sexes), or that a different selective force is operating on limb length (all three species).
6. Above ground niche, male spends little or no energy in incubating chamber construction, both sexes responding equally to selection from vertebrate predators: L. elenae, gracilis, notoaktites. This pattern implies that neither sex is under selection that would result in shorter limb elements; males probably locate available depressions, holes, or burrows and make few if any modifications of them preparatory for use as incubating chambers.

Mid-dorsal Stripe. -This characteristic is not sexually dimorphic, suggesting that the character state is not involved in mate recognition.

Distinct Lip and Thigh Stripes.-A few species are dimorphic for these characteristics. In all cases, the stripes are more distinct in the females than in the males. Straughan (1966) has demonstrated that thigh pattern is important in mate recognition, acting as a species isolating mechanism. The Leptodactylus lip and thigh stripe data indicate that the males utilize this information in mate discrimination in several members of the fuscus group.

Texture of Tarsus and Foot.-The presence or absence of white tubercles is not sexually dimorphic in any species, indicating that these structures are not used in mate recognition but are probably important in how the frog physically interacts with the environment.

## RELATIONSHIPS

The purpose of this section is to determine whether a pattern of phyletic relationship can be inferred among the species. Detailed relationships cannot be analyzed at this time for two reasons. The first is that there are several as yet undescribed species in this group, and the morphological information on L. geminus and marambaiae needs to be clarified. The second reason that detailed relationships cannot yet be determined is methodological. I prefer to deduce relationships on the basis of shared derived character state patterns (see Heyer, 1975, for fuller explanation). This method requires more derived character states than taxa for any detailed analysis. With the small number of characters presently available for analysis, only general hypotheses regarding relationships are possible.

The outgroup used for comparative purposes in determining the primitive states consists of members of the other species groups of Leptodactylus and members of the genera Adenomera, Lithodytes, and Vanzolinius. States common to the outgroup but variable within
members of the fuscus group are considered primitive. These four genera likely had a common ancestor. Only those characters for which the states are known for all species (geminus and marambaiae excluded) are analyzed in detail.

Character Analysis-Mid-dorsal stripe. Character 1. -State $0=$ light mid-dorsal stripe absent in all individuals; State $1=$ light mid-dorsal stripe present in some individuals; State $2=$ light mid-dorsal stripe present in all individuals. Of the outgroup, only some members of the genus Adenomera have a light mid-dorsal stripe. The Adenomera light mid-dorsal stripe differs among Adenomera species and differs from the mid-dorsal stripe of members of the fuscus group, however. The most parsimonious explanation is that the common ancestor to all of the taxa considered had the genetic potential for a light mid-dorsal stripe. The direction of change of states is:

$$
0 \rightarrow 1 \rightarrow 2
$$

Lip stripe. Character 2.—State $0=$ light lip stripe indistinct; State $1=$ light lip stripe distinct in at least some individuals. Members of the genera Adenomera, Lithodytes, Vanzolinius and the melanonotus species group lack light lip stripes. One species of the ocellatus group and two species of the pentadactylus group have light lip stripes. The pentadactylus group members have a stripe differing in detail from the fuscus group light stripe. The situation is analogous to the mid-dorsal stripe character. State 0 is considered the primitive state.

Thigh stripe. Character 3.-State $0=$ no distinct light longitudinal stripe on the posterior surface of the thigh; State $1=$ distinct posterior thigh stripe. Only some individuals of the other species groups of Leptodactylus approach state 1 in the outgroup. State 0 is considered the primitive state.

Dorsolateral folds.-The presence of at least a pair of distinct dorsolateral folds is common throughout the outgroup; the primitive state can not be determined from the outgroup. The condition of 6 dorsolateral folds is probably derived, but this character state is associated with the light mid-dorsal stripe in all fuscus group members except for fuscus itself. As the mid-dorsal stripe information is being analyzed, the dorsolateral fold information is not analyzed or used further.
Tarsal and foot texture.-Several members of the outgroup demonstrate all states regarding tarsal and foot texture. The outgroup provides no information on which state is primitive.

Size.-Members of the outgroup are both larger and smaller than members of the fuscus group; the primitive state can not be determined. In all likelihood, the moderate size of most of the fuscus group members is the primitive state.
Sexual dimorphism in size, head and limb propor-tions.-As discussed in the previous section, any sexual dimorphism of head and leg proportions is uncommon
in frogs and is here considered the derived state. The exception is SVL, in which sexual dimorphism in size is considered the primitive state as discussed previously. For all characters listed below, state 0 is the primitive state.
SVL. Character 4.—State $0=$ sexually dimorphic; State $1=$ not sexually dimorphic.

Head length. Character 5.-State $0=$ not sexually dimorphic; State $1=$ sexually dimorphic.
Head width. Character 6.-State $0=$ not sexually dimorphic; State $1=$ sexually dimorphic.
Femur/SVL ratio. Character 7.-State $0=$ not sexually dimorphic; State $1=$ sexually dimorphic.
Tibia/SVL ratio. Character 8.-State $0=$ not sexually dimorphic; State $1=$ sexually dimorphic. As troglodytes is unique in that it is the male with the longer tibia, it is coded as 0 for analytic purposes.

Foot/SVL ratio. Character $9 .-$ State $0=$ not sexually dimorphic; State $1=$ sexually dimorphic. See above for troglodytes.

The distribution of states among the species is presented in Table 5 . With only 9 characters, detailed relationships can not be drawn, but the distribution of states and clustering pattems allow certain generalizations to be made. 1) There is a cluster of taxa characterized by having very few derived states, which do not demonstrate any meaningful patterns of relationships among themselves. These species, L.• bufonius, labrosus, troglodytes, and ventrimaculatus are likely similar to the ancestor of the fuscus group and demonstrate the basic adaptive features of the ancestral stock of the entire fuscus group. Assuming this to be true, the ancestral stock of the fuscus group had a basic semi-fossorial adaptive set. As all members of this assemblage have white tubercles either on the tibia, tarsus, or foot, the fuscus group ancestor likely had tubercles also. 2) A
second assemblage of species is characterized by sharing the derived states of lip and thigh stripes: L. albilabris, amazonicus, elenae, fragilis, fuscus, gracilis, latinasus, laurae, longirostris, mystaceus, and notoaktites. Within this assemblage, albilabris, amazonicus, and fragilis together share the most derived states (5) within the data set. 3) Leptodactylus mystacinus and poecilochilus are intermediate between these two assemblages. There is no parsimonious way to include both of these species in the same evolutionary sequence leading to the second assemblage. As L. mystacinus bears more morphological similarity to members of the first, supposed primitive, assemblage of species, it does provide at least an example of how the transition between the first two assemblages could have occurred.

## ZOOGEOGRAPHY

When the distributions of species (excluding those known only from single localities) are outlined and overlayed, two results are apparent. First, most species of the fuscus group occur south and east of the Amazon basin. Second, the areas of greatest present species densities do not appear to coincide with local areas of speciation. There are four areas where the ranges of five species overlap. Two of these are in the dry interior portions of Argentina. The species coexisting in these two areas are: (A) bufonius, elenae, fuscus, latinasus, mystacinus, and (B) elenae, fuscus, gracilis, latinasus, mystacinus. A third area is in southeast Brazil in the São Paulo region. The species that occur in sympatry there are fuscus, gracilis, laurae, notoaktites, mystacinus. The fourth area is the border region between southeast Brasil and Uruguay. The species are fuscus, gracilis, latinasus, laurae, mystacinus. Clearly, the high numbers of species that coexist in these regions reflect overlap in the ranges of widespread species. There are no

Table 5
Distribution of character states among members of the fuscus group. Character numbers and states as used in text.

|  | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| albilabris | 1 | 1 | 1 | 0 | 1 | 0 | 1 | 1 | 1 |
| amazonicus | 0 | 1 | 1 | 0 | 1 | 1 | 0 | 1 | 1 |
| bufonius | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 |
| elenae | 0 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 |
| fragilis | 0 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| fuscus | 1 | 1 | 1 | 1 | 0 | 1 | 1 | 1 | 0 |
| gracilis | 2 | 1 | 1 | 1 | 0 | 1 | 0 | 0 | 0 |
| labrosus | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 |
| latinasus | 0 | 1 | 1 | 0 | 1 | 0 | 0 | 0 | 0 |
| laurae | 2 | 1 | 1 | 0 | 1 | 1 | 0 | 0 | 0 |
| longirostris | 1 | 1 | 1 | 0 | 1 | 0 | 0 | 1 | 0 |
| mystaceus | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 0 | 0 |
| mystacinus | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 |
| notoaktites | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 |
| poecilochilus | 1 | 0 | 1 | 1 | 1 | 0 | 0 | 0 | 0 |
| troglodytes | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 |
| ventrimaculatus | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 |

small circumscribed geographic areas characterized by having a number of endemic species.

When the distributions of fuscus group species are compared with the distributions of broad vegetation types, a general correlation is evident. There are certain distributions that do not correlate, however.

Two peripheral populations of $L$. amazonicus do not fit the general distribution pattern of the remaining populations: the population in north coastal Venezuela and the population in northeast Brasil (fig. 34). Both of these populations occur in mesic forest regions, but are separated from the mesic forest associated Amazon populations by dry forests. No information is available on the mating calls of individuals from these populations to determine whether they are sibling species or disjunct populations of $L$. amazonicus. Vanzolini (1974) suggested that there was broad continuity between the Atlantic and Amazonian forests until relatively recently. If the population in northeast Brasil is a disjunct population of $L$. amazonicus, its presence there can be accounted for if the Amazonian and Atlantic forests were in recent contact. I have no explanation for the Venezuelan population.

The distribution of L. elenae is Chacoan with a single exception of a sample of two frogs from Tocache Nuevo, Río Huallaga, Peru. Further sampling and knowledge of the mating call of frogs from this area are needed.

Two disjunct populations of $L$. latinasus are apparent (fig. 51), a southern series and a northeast Brasil series. No mating calls are available from the northeast Brasil specimens. They may represent a sibling species of the southern latinasus.

The individual distribution patterns (figs. 34, 42, 44,
$47,51,55,61,62,68)$ were compared with the vegetation map of South America by Hueck and Seibert (1972). Because most Leptodactylus locality records do not include specific data on associated vegetation, only broad associations can be made. The occurrence by species within Hueck and Seibert's (1972) broad categories are shown in Table 6. Most species occur in more than one broad vegetation type. The three species which occur within only one vegetation category are restricted to tropical and subtropical rainforest. Vanzolini (1970) grouped the individual vegetation units of Hueck (1966) into broad units which differ in part from the broad categories later recognized by Hueck and Seibert (1972). For those species occurring east of the Andes and in the greater Amazon basin southeastward, excluding those species known from but a single locality, distributions by broad vegetation types are shown in Table 7. According to the Vanzolini modification, several species are associated with a single broad vegetation category; thus the Vanzolini modification (1970) describes the distributions by vegetation types of Leptodactylus species better than the Hueck and Seibert (1972) classification.

Three conclusions may be drawn from the data in Tables 6 and 7: (1) Some species are restricted to wet forest or open habitat vegetation formations, (2) more species are associated with mesic forest vegetation types than xeric vegetations, (3) several species show distribution patterns associated with more than one major vegetation type. In the discussion that follows, open formations as used here contrast with closed canopy forests. Open formations include the open vegetation formations such as cerrado and caatinga, natural and man-

Table 6
Species occurrence within general vegetation types of Hueck and Seibert (1972). Occurrence of L. elenae in tropical rainforest is at single Peruvian locality, see text.

|  | Tropical and Subtropical Rainforest | Deciduous, <br> Mesophytic, <br> Tropical and Subtropical Forests | Dry Forests | Savannas, Palm <br> Savannas, and Palm Forests | Bush and Grass Steppe; Half and Full Deserts |
| :---: | :---: | :---: | :---: | :---: | :---: |
| albilabris | X | X | X |  |  |
| amazonicus | X |  |  |  |  |
| bufonius |  | X | X |  | X |
| fragilis | X | X | X |  |  |
| elenae | (X) |  | X | X |  |
| fuscus | X | X | X | X | X |
| gracilis | X | X | X |  | X |
| labrosus | X |  | X |  | X |
| latinasus |  | X | X |  | X |
| longirostris | X | X |  | X |  |
| laurae | X | X |  |  | X |
| mystaceus | X |  |  |  |  |
| notoaktites | X | X |  |  |  |
| mystacinus | X | X | X | X | X |
| poecilochilus | X |  | X |  |  |
| troglodytes |  |  | X |  | X |
| ventrimaculatus | X |  |  |  |  |

Table 7
Species occurrence within general vegetation types using Vanzolini's (1970) modification of Hueck's (1966) scheme. Species with West Indian, distributions west of the Andes, and primarily Middle American distribution patterns omitted.

|  | Hylea | Atlantic Forest <br> and Araucaria | Cerrado | Caatinga |
| :--- | :---: | :---: | :---: | :---: |
| amazonicus | X | $(\mathrm{X})$ |  |  |
| bufonius | $(\mathrm{X})$ |  | X |  |
| elenae | X | X | X |  |
| fuscus |  | X | X |  |
| gracilis | latinasus | X | X | X |
| longirostris |  | X | X |  |
| laurae | mystaceus |  | X |  |
| notoaktites |  | X |  |  |
| mystacinus | troglodytes |  |  | X |

made openings in closed canopy forests, and river flood plains. If the only data one had were museum specimens, localities, and a vegetation map, one would logically look in the forest for such species as $L$. amazonicus or longirostris on a field trip. But there is a paradox: Although several species in the fuscus group are associated geographically with wet forest formations, not one (to my knowledge) is found within the forests themselves. These species occur in the open formations within the forest systems, such as along river banks. The paradox can be resolved by recognizing a scheme of two basic zoogeographic patterns.

The first zoogeographic pattern involves species occurring only within dry forest vegetations (e.g. cerrado and caatinga). The distributions of $L$. bufonius and troglodytes are typical of species associated with the diagonal band of open formations (Vanzolini 1974, which see for comparable distribution patterns in lizards). Characteristically, two closely related populations are found in the diagonal; one in the Chaco, one in northeast Brasil. Speciation of these xeric adapted forms undoubtedly followed the classical allopatric model. However, the stability of the open formations throughout the probable period of speciation of members of the fuscus group (Solbrig 1976) has provided limited opportunities for speciation. Looking at it another way, if one removed the members of the fuscus group that are associated with mesic forest, the group would consist of only four or five species. If bufonius, labrosus, troglodytes, and ventrimaculatus reflect the original distribution of members of the group as postulated in the relationships section, then the ancestral group must have had a broader distribution historically than is reflected by the four remaining species populations. The intervening areas (Amazonia) must have had dry corridors; extensive dry corridors may have existed up until the Miocene, when there was still a lowland area in the forming Andean chain, about where Ecuador is today (e.g. Solbrig 1976, fig. 2.2). After the uplift of the Andes, any mesic period
(such as now) would eliminate the extensive dry corridors in Amazonia, resulting in elimination of the ancestral fuscus group stock from the Amazon basin.

A second zoogeographic pattern involves the species associated with wet forests. The open formations found within wet forest systems are distinctive from the open formations of dry forest systems. Further, the open formations of different wet forest units must differ in soil characteristics, standing water, etc., so that given time, the adaptations to open formations within different wet forest units will differ. The key to the relatively large number of species in the fuscus group is that evolutionary histories of many of the species have been associated with the histories of the Neotropical mesic forests. The mesic forest units have been a dynamic system, providing greater opportunity for speciation than the dry forests for the fuscus group. The critical point is that the open formation Leptodactylus species of the mesic forests have the same evolutionary histories as the fauna of the forests. In other words, the evolutionary-environmentalgeographic unit consists of the wet forests and their associated open habitats, not just the forests themselves. The distributions of Leptodactylus species associated exclusively with wet forests can be correlated with the location of supposed Pleistocene forest refugia (e.g. compare fig. 4, Vuilleumier 1971, with the distributions of longirostris, notoaktites, and ventrimaculatus). The dynamic expansion, contraction and fragmentation of the mesic forests and their associated open formations has provided the opportunity for speciation in many members of the fuscus group.

In regions characterized by both wet and dry forests, it is likely that the differentiation of species has been associated with presence of the open formations within a given wet forest system. The species subsequently spread to adjacent open habitats in drier forests. The distribution in drier forest open habitats could occur in association with the gallery forests along the rivers. For example, if amazonicus only occur in association with
gallery forests along the rivers in cerrado, the result would be a case of symmetry to the network of open formations in closed forests. In such instances, the microhabitats utilized by the species within the open formations of mesic closed forest and galleries of dry forest would be similar.

The zoogeographic hypotheses invoke mesic and xeric associations. For didactic purposes, the zoogeographic patterns have been explained separately. This does not infer that the zoogeographic patterns are the result of two separate processes. The single process of historic climatic fluctuations has produced all of the zoogeographic patterns.

In a previous section (see Relationships), L. mystacinus was cited as an example of how evolution could have proceeded from the more primitive member species to the more derived. The distribution of $L$. mystacinus is also exemplary in this regard. The species occurs in open formations in dry and wet forests. The pattern demonstrates that an ancestral member of the fuscus group, which was adapted to dry forest open habitats could have invaded the openings within wet forests. Once such open habitats were occupied, the species range could expand during periods when the forests were extensive. During drier times, some of the populations were likely isolated in open habitats within forest islands.
One difficulty in understanding the zoogeography of this group (or any other large species group) is what might be termed the palimpsest factor (term and following discussion suggested by P. E. Vanzolini). There are three possible historical times to date certain zoogeographic distributions for the fuscus group: (1) A possible Miocene distribution of the ancestral fuscus stock, (2) A Pliocene distribution event for fragilis and poecilochilus, and (3) A very recent (hundreds of years) wet climax providing continuity of the hylaean and Atlantic forests accounting for the present distribution of amazonicus. So much of what happened between these end points has been erased and written over, there is no hope of unravelling the history.

## EVOLUTIONARY HYPOTHESES

The fuscus group ancestral stock was semi-fossorial, adapted to the kind of open, xeric vegetation formation that now occurs in the Gran Chaco (this does not infer that the origin of the group was necessarily in the Chaco). The extant species of the fuscus group that are the most primitive in morphology and habits are still primarily associated with this ancestral vegetation formation. The burrowing adaptations of the semifossorial ancestral stock served as a preadaptation for placement of the foam nest in an underground chamber. It is possible that the ancestral stock formed their own underground burrows for retreats or aestivating sites and the males simply made use of these chambers as calling sites with consequent deposition of the foam nest. The place-
ment of the foam nest in an underground chamber was a preadaptation for the expansion of the group into adjacent, less harsh, vegetation formations. In more mesic habitats, more activities were carried out above ground, and the principal function of burrowing became the formation of the chamber in which the foam nest is placed. This nesting activity had by now become solely a male activity. Distinctive lip and thigh stripes presumably became important at this evolutionary stage because the male is expending considerable energy into reproductive activities, the success of which depend on selection of a proper mate. In the previous evolutionary stage, there was much less energy specifically channeled into reproductive activities by males, for the latter utilized chambers formed for another purpose. Presumably females also made the same kind of burrows or chambers, for the snout shapes of $L$. labrosus suggest that individuals of both sexes engage in burrow or chamber formation in this species. Once the formation of an incubating chamber becomes a strictly male activity, selection should reinforce any mechanism that assures that the male makes the correct species choice in mate selection. The female chooses the male on the basis of call; but the male must make a choice based upon the females that he encounters. Observations on L. mystacinus (Sazima 1975) corroborate this series of events. The male calls to attract a female. The male does not initiate chamber formation until a female approaches. Once the male starts chamber formation, he stops frequently and makes contact with the female. Apparently this frequent interruption of chamber formation is for reinforcement from the female, either tactile or visual. Males of some species of the fuscus group form the incubating chamber before females are called in. In these cases, proper species mate recognition would be at a premium; it would appear that the thigh and lip stripes function in this role.

## PRELIMINARY COMMENT ON SIBLING SPECIES

There are two sibling species complexes in the fuscus group as now constituted (the new species being described by South American workers may provide additional cases). A sibling species complex is operationally defined herein as a group of biological species which are indistinguishable morphologically, with or without the aid of sophisticated statistical techniques. The two cases of sibling species pair complexes are L. fragilis - latinasus and L. geminus - gracilis - marambaiae. Although more data are needed concerning the geminus - gracilis marambaiae problem, it appears that the fragilis - latinasus pair and geminus - gracilis - marambaiae group had very different evolutionary origins. All that is essentially required as a mechanism for sibling species formation in frogs is the evolution of distinctive mating calls. If polyploidy accompanied mating call differentiation, such as in the sibling species pair Hyla chrysoscelis - versicolor, reproductive isolation would be
immediate. The geminus - gracilis - marambaiae sibling triad suggests that call differentiation, unaccompanied by polyploidy, has led to reproductive isolation. In contrast, the fragilis - latinasus morphologies apparently are due to convergence. As indicated in the relationships section (and data in Table 5), the two species are not particularly closely related to each other. In fact fragilis has several closer related species than latinasus. This is substantiated by the karyotype data, in that latinasus is unique in the fuscus group in having a pair of terminal chromosomes. The similarities in size and morphology of these two species apparently are due to parallel selective pressures in similar habitat types. The fragilis latinasus example points out the need for caution in assuming that because two species of frogs are morphologically most similar to each other, they are necessarily most closely related to each other.

## RESUMEN

Se analizan detalladamente trece caracteres de la morfologia externa para las especies comprendiendo el grupo fuscus (género Leptodactylus). El método principal del análisis de los datos es el aplicación del análisis multivariante de función selectiva en serie (multivariate stepwise discriminant function analysis). Se comparan los resultados del análisis morfologico con la información conocida respeto a los cantos nupciales, las larvas, y los cariotipos. Basándose sobre todos los datos obtenibles, se extraen conclusiones taxonómicos.

La nomenclatura del grupo se describe detalladamente, asociando los nombres propuestos con las unidades de especies reconocidas en este estudio. Cada y cuando que fuese posible fué re-examinado el material de los tipos originales para este estudio. De los diez y
nueve especies reconocidos en el grupo fuscus, cuarto se describen como especies nuevas.

Para cada especie, se provee la siguente información: una sinonimía de los nombres primarios, un diagnóstico para los adultos, sumarios de las caracteristicas morfológicas de los adultos y las larvas, descripciones diagnósticas de los cantos nupciales, descripción diagnóstico del cariotipo, y distribución incluyendo localidades y los respectivos numeros de clasificación de los ejemplares de museos para las especies examinadas. Se provee una clave al final de las descripciones de las especies.

El orden compuesta del grupo es enorme, con distribución de Texas hasta Argentina en ambos lados de la Cordillera de los Andes y ciertas islas de las Antillas.

Varios caracteres utilizados en el análisis son sexualmente dimorfos. Queda postulado que el dimorfismo sexual en las proporciones de los miembros traseros se debe a la selección diferencial, el miembro más corto del macho es el resultado de la selección para la actividad de hacer madrigueras relacionada a la formación de camaras de incubación, el miembro más largo de la hembra es el resultado de la selección para evitar vivientes de repaña. El dimorfismo sexual que occurre en las rayas del labio y del muslo de varias especies es explicado por el hipótesis que los machos estan usando la información a diferenciar entre las hembras en el reconociemiento aparear.

El linaje hereditario del grupo fuscus es presumibido haber sido cavadoramente adaptado a una área con un tipo vegetivo parecido a este ahora encuentro en el Gran Chaco. Los hechos evolutivos dentro el grupo de las especies tienen correlación con las adaptaciones a las ambientes más húmedas.

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