

ECOLOGICAL CORRELATES OF MORPHOLOGICAL EVOLUTION IN A DARWIN'S FINCH, *GEOSPIZA DIFFICILIS*

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Morphological variation among different island populations is a familiar attribute of many species of Darwin's finches. Lack (1947) provided the first detailed arguments for the adaptive significance of morphological variation in this group. He stressed two aspects: first, that several morphological features are related to feeding, and second, that morphological differences between populations may be associated with inter-island differences in the type and number of potentially competing species.

A specific example is the sharp-beaked ground finch, *Geospiza difficilis*. This species is widely distributed in the Galápagos archipelago and occurs with a variety of combinations of congeneric species (Fig. 1). Lack argued that the population on Genovesa resembles the absent species *G. fuliginosa*. Two other populations on the low islands of Darwin and Wolf resemble the cactus finch, *G. scandens*, where both this species and *G. fuliginosa* are absent (Lack, 1947). *Geospiza difficilis* populations on the more central highland islands are intermediate between these two extremes. From these data Lack inferred that on Genovesa *G. difficilis* occupies the feeding niche exploited by *G. fuliginosa* on other islands, and that on Darwin and Wolf it combines the niches of *G. fuliginosa* and *G. scandens* (Fig. 1). Lack considered this as evidence in support of his general thesis that competition had influenced the adaptive radiation of the finches.

Lack based these conclusions largely

upon morphological study of museum specimens. Hence a central assumption of his arguments is that morphological differences between *G. difficilis*, *G. fuliginosa* and *G. scandens* adequately reflect differences in diet. In this report we use morphological and ecological data from *G. difficilis* populations to investigate this central assumption. *Geospiza difficilis* is particularly useful for this type of study because it is morphologically one of the most differentiated species of Darwin's finches (Lack, 1947; Grant, 1981; Grant and Schluter, 1984). Our purposes are threefold:

1. To characterize, more quantitatively than Lack did, interpopulation variation in the morphology of *G. difficilis* and the morphology of this species in relation to the two most similar congeners, *G. fuliginosa* and *G. scandens*.
2. To compare diets of different *G. difficilis* populations with their morphologies, and with the diets of the two morphologically similar species, *G. fuliginosa* and *G. scandens*.
3. To combine information on feeding and morphology with data on food supply in order to interpret trends of morphological evolution in *G. difficilis*.

Lack's method of inferring feeding niches from morphological data is commonly used, especially to examine patterns in the assembly and evolution of ecological communities (e.g., Keast, 1972; Gatz, 1979; Ricklefs and Travis, 1980; Ricklefs et al., 1981; Simberloff and Boecklen, 1981 and references therein; Grant and Schluter, 1984). Since the validity of such inference has rarely been

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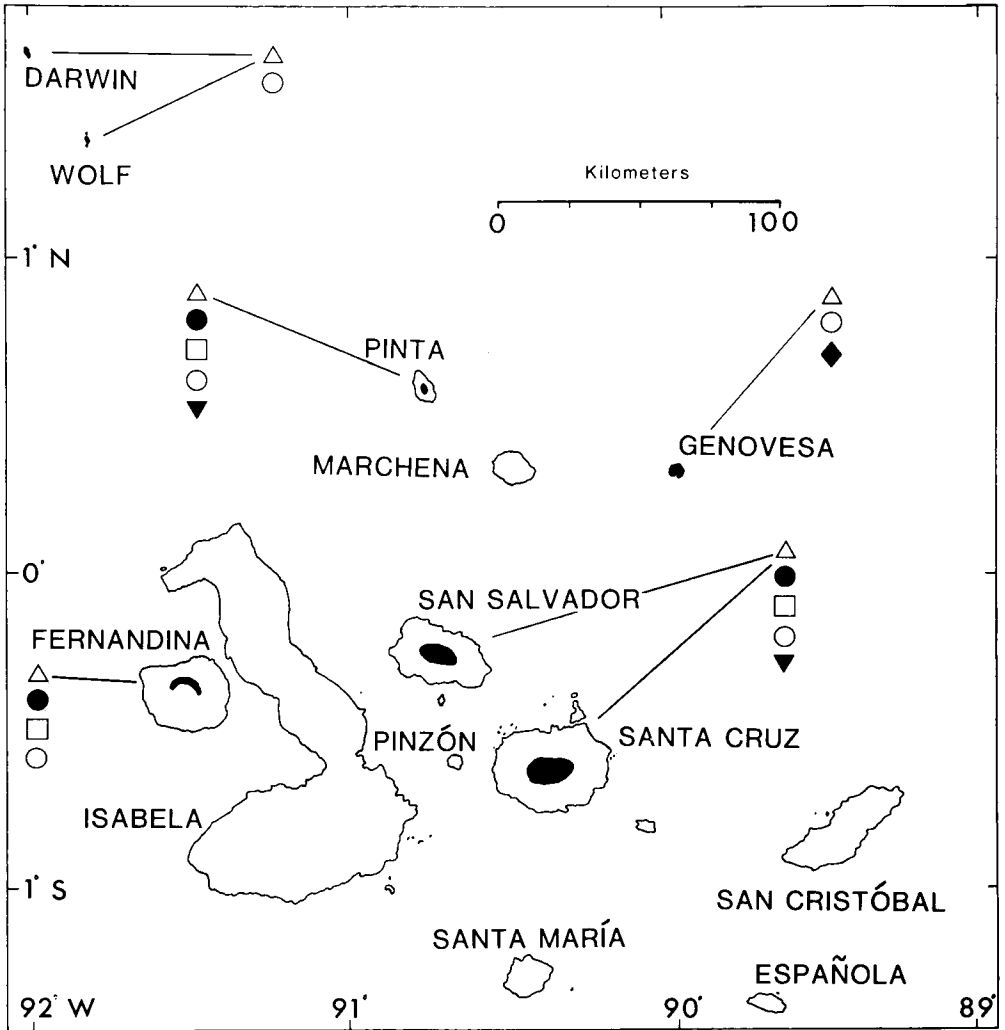


FIG. 1. The distribution of *G. difficilis* (Δ ; darkened regions) in relation to congeneric species (after Lack, 1947). Lack (and Harris, 1973) were uncertain as to the presence of *G. difficilis* on Fernandina: it has a large population there (this study). Lack also mentioned the possibility of a *G. difficilis* population on southern Isabela. Habitat destruction may have caused its extinction there, as apparently happened on Santa María (Lack, 1947). The species is now extinct on Santa Cruz. Congeners are the small (*G. fuliginosa*; ●), medium (*G. fortis*; □) and large (*G. magnirostris*; ○) ground finches, and the cactus finches, *G. scandens* (▼) and *G. conirostris* (◆). As Lack noted, *G. difficilis* is found mainly in the highlands except on the low islands of Darwin, Wolf and Genovesa, where *G. fuliginosa* is absent.

examined (James, 1982), the present analysis may be of general significance.

METHODS

Nine islands were visited during the study (Table 1), and measurements were

taken on morphology, density, diets and food supplies of finches.

Study Sites

The islands visited include all six islands on which *G. difficilis* is presently

TABLE 1. Dates of visits to islands, and sample sizes for finch measurements. Values in parentheses refer to birds in partly or fully black plumage (adult males). Measurements, made by both authors and D. J. Anderson (DA), were not all made on the dates shown. Maximum elevations are from Wiggins and Porter (1971).

Island	Maximum elevation	Dry-season dates	No. specimens measured			Person measuring
			<i>difficilis</i>	<i>fuliginosa</i>	<i>scandens</i>	
High						
Pinta	777 m	28 Oct–3 Dec 1979	35 (22)	149 (72)	29 (19)	DS
San Salvador	905 m	13 Sept–8 Oct 1981	25 (13)	56 (28)	—	DS
Fernandina	1,494 m	9 Oct–5 Nov 1981	16 (11)	26 (15)	—	DS
Low						
Darwin	168 m	12 Sept 1981	22 (4)	—	—	DS
Wolf	253 m	10–11 Sept 1981	45 (21)	—	—	PG
Genovesa	76 m	17 Sept–13 Oct 1979	92 (49)	—	—	PG
Marchena	343 m	4–27 Oct 1979	—	18 (11)	4 (2)	DS
Española	198 m	11–26 Nov 1981	—	23 (12)	—	DA
Tortuga	186 m	7–10 Nov 1981	—	25 (8)	—	DA

found (Fig. 1). Three additional islands, Marchena, Española and Tortuga, were added to the list in order to increase the sample size of *G. fuliginosa* measurements and observations. At least one, and up to six, 1-ha study sites were established on each island depending upon time constraints and the diversity of habitats. On high elevation islands (Table 1), sites were altitudinally spaced so as to represent each of the three major vegetation zones: arid, transition, and humid forest (Wiggins and Porter, 1971). The approximate locations of sites are: Pinta, 6 sites spaced ca. .5 km apart along S trail to summit, from 110 m elevation to 510 m; San Salvador, sites at Caleta Bucanero (coast), at 670 m elevation near trail on N slope, and at 830 m on S slope immediately below main summit; Fernandina, 3 adjacent sites along NW crater edge at summit above Cabo Douglas (1,300 m), and one site along coast 3 km NE C. Douglas; Darwin, 1 site on landslide area above landing on NW side; Wolf, 1 site at the eastern half of NW plateau; Genovesa, 3 sites at Bahía Darwin, in the same location as the one used by Abbott et al. (1977); Marchena, 1 site behind E point of Playa Grande, and a second on hilltop above lava plains 3 km

uphill and NE (170 m); Española, 1 site behind Bahía Gardner in the same location as the one used by Abbott et al. (1977), and 2 others 1 km and 2 km directly inland (30 and 85 m); Tortuga, 1 site N slope immediately below ridge (130 m). Site descriptions for Pinta, Marchena, and Genovesa are given in Schluter (1982a) and Schluter and Grant (1982).

Climate in the Galápagos is highly seasonal, with a hot/wet season (ca. January–May) and a cool/dry season (June–December) (Grant and Boag, 1980). All islands were visited in a dry season (Table 1), though several were visited also during a wet season. Effort was concentrated in the dry season because at that time standing crop and production of foods reach their lowest values. There is evidence that finch populations become food-limited then, and at this time diet should reflect morphology most closely (Smith et al., 1978; Boag and Grant, 1981; Schluter, 1982a). Finches usually breed only in the wet season.

Goats have been introduced onto the islands of Pinta, San Salvador and Española. Extermination programs have eliminated their populations on Pinta (almost) and Española (completely). Vege-

TABLE 2. Untransformed measurement means and standard deviations (in parentheses) for adult male *G. difficilis*. Measurements for male *G. fuliginosa* and *G. scandens*, averaged over populations, are provided for comparison. Mass is in g, the remainder are in mm. Sample sizes are in Table 1.

	Body mass	Wing length	Tarsus length	Beak length	Beak depth	Beak width	Beak length at 4 mm
<i>difficilis</i>							
Pinta	19.0 (1.1)	63.1 (1.4)	19.4 (.57)	9.8 (.25)	7.2 (.19)	7.1 (.16)	4.6 (.14)
San Salvador	26.8 (2.3)	71.2 (1.9)	21.6 (.68)	10.2 (.49)	7.6 (.21)	8.0 (.17)	4.1 (.23)
Fernandina	19.6 (1.1)	65.6 (1.7)	20.7 (.56)	9.3 (.40)	6.9 (.22)	7.0 (.23)	4.5 (.30)
Darwin	25.5 (2.5)	74.0 (1.4)	22.7 (.62)	11.5 (.45)	7.9 (.24)	7.8 (.43)	4.4 (.21)
Wolf	20.5 (.79)	74.6 (2.3)	20.8 (.40)	11.1 (.35)	7.1 (.23)	7.2 (.28)	4.9 (.26)
Genovesa	11.6 (.66)	64.0 (1.8)	18.1 (.47)	9.6 (.44)	6.4 (.27)	6.5 (.28)	5.1 (.28)
<i>fuliginosa</i>	13.3	62.1	17.9	8.6	6.9	6.8	3.8
<i>scandens</i>	21.8	73.1	21.1	14.5	9.1	8.7	4.8

tation and soil have recovered significantly on Pinta, but not on Española where they are still visibly affected, and not on San Salvador where goats have been present since 1813 (Hamann 1979, 1981; pers. observ.). Pigs, donkeys and rats have also been introduced to San Salvador. Undoubtedly the disturbance has affected finch feeding patterns to some extent, especially on the latter islands.

Finch Diets and Food Supply

Study procedures are described in Schluter (1982a and references therein). Briefly, an observer walked along a pre-determined route and recorded the amount of time that individual finches spent feeding on specific food types. These finches are tame and may be approached closely, and food items could usually be identified directly in the beak, or indirectly from the parent plant. A maximum of 300 observations per bird was set so that information could be obtained from as many birds as possible. All data were collected by two observers, with consistent results.

Food supply was assessed on all islands in the dry season, excepting Darwin where time did not permit this. Twenty-five (occasionally 50) random 1 m² quadrats were located in each site. In each quadrat, the percent cover of plant species, the abundance of seeds and fruits on the vegetation, and the number of flowers

were determined. In addition seeds on rock surfaces in two subquadrats of .125 m² were counted. All litter and surface soil (to a depth of 2.5 cm) within these subquadrats was collected in bags and subsequently examined for seeds and invertebrates. Dry mass of both these food types was later assessed in the laboratory.

Finch Morphology

Seven characters were measured on live finches captured in nets. Characters represent different aspects of body or beak size, and hence they are among the traits most likely to be related to feeding. Body Mass was determined to the nearest .1 g with a spring balance. The remaining characters were measured to the nearest .1 mm: Wing Length, from the metacarpal bend to the tip of the longest primary, with the wing held flat against a rule; Tarsus Length, with dividers, from the tibiotarsal joint to the distal end of the first scute surrounding the tarsus that was wider than its length; Beak Length, measured with dividers, from the anterior end of the nares to the tip of the upper mandible. The remaining three traits were measured using vernier calipers: Beak Depth, measured at the anterior end of the nares perpendicular to the commissure; Beak Width, the width of the lower mandible at the widest part, near the base; and Beak Length at 4 mm, the distance from the point along the commissure at

TABLE 3. Factor loadings for the first three principal components derived from mean morphology in populations of *G. difficilis* (adult males). Values in parentheses give the fraction of the variation accounted for in each trait (Pimentel, 1979). Also given is the cumulative total variance accounted for by the addition of successive components. $N = 6$ populations.

Variables	Scores and variance fractions		
	PC1	PC2	PC3
Wing length	.37 (.67)	.42 (.24)	-.56 (.07)
Tarsus length	.47 (.92)	-.04 (.00)	-.44 (.04)
Beak length	.38 (.57)	.60 (.40)	.41 (.03)
Beak depth	.41 (.91)	-.11 (.02)	.55 (.07)
Beak width	.45 (.92)	-.21 (.06)	.12 (.00)
Beak length at 4 mm	-.36 (.53)	.63 (.47)	.08 (.00)
% variance	74.7	95.3	98.8

which the depth of the beak is 4 mm, to the tip of the upper mandible. Unlike previous 'size' variables, Beak Length at 4 mm is essentially a 'shape' variable, measuring beak pointedness at the tip. Its magnitude within species is dependent upon Beak Length and Beak Depth, but differences between species, especially *G. fuliginosa* and *G. difficilis*, are due mainly to differences in beak curvature independent of differences in Beak Length and Depth.

Sample sizes are given in Table 1, and descriptive statistics are provided in Table 2. Measurements from different populations were made by three persons and at different periods over 1978 to 1981. Identical methods were used throughout. Boag (1983) has shown high repeatabilities (Falconer, 1981) for these traits in *G. fortis* and *G. scandens* individuals measured at different times and by different persons.

Measurements of adult male *G. scandens* from Santa Fe ($N = 5$), Champion ($N = 6$) and Daphne ($N = 105$) by P.R.G. and P. T. Boag were included in the comparative analyses to improve the sample size for this species.

RESULTS

We begin with a brief description of the general morphology of *G. difficilis*, and compare it to the morphologies of *G. fuliginosa* and *G. scandens*. Diets are then described in relation to food avail-

ability, and diet differences between populations are compared with differences in morphology.

Multivariate Patterns

Principal components analysis on log(10)-transformed measurements (PCA; Pimentel, 1979) was used to represent multivariate differences among populations of *G. difficilis*, and similarities with other species. Two separate PCA's were performed. In the first, components were extracted from a six-variable covariance matrix computed from all *G. difficilis* individuals combined. The second PCA used the covariances computed from the population means for adult males only. Means for males differ significantly among populations (ANOVA, $P < .001$), and are larger than female means (Price, 1984). Body Mass was not included in either analysis because it is given in different units from the other variables. The two analyses gave almost identical results, and we use only the second.

The first component (PC1) explains a large amount of the variance in all of the original 'size' variables, and accounts for a large amount of the total variation among *G. difficilis* populations (Table 3). Body Mass, highly variable among these populations (Table 2) is also highly correlated with PC1 ($r = .95$). PC1 may thus be considered a general body size variable (Pimentel, 1979).

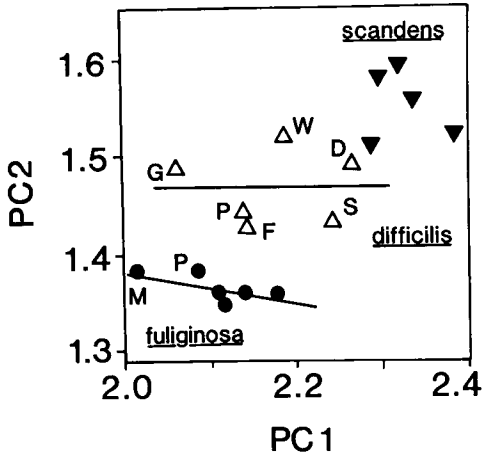


FIG. 2. Population means for adult male *G. difficilis*, *G. fuliginosa*, and *G. scandens* along the first two morphological principal components. Loadings for original variables are given in Table 3. Components are derived from population means for adult male *G. difficilis* only. Solid line through *G. fuliginosa* points is the first principal axis derived from population means ($N = 6$) projected onto the PC1-PC2 plane through the overall *G. fuliginosa* mean. The axis accounts for 89% of the variance among *G. fuliginosa* means. PC1 was redrawn through the overall *G. difficilis* mean for comparison. Island symbols for *G. difficilis* are Pinta (P), San Salvador (S), Fernandina (F), Darwin (D), Wolf (W), and Genovesa (G). *Geospiza fuliginosa* forms are unlabelled except for those on Pinta (P) and Marchena (M).

PC2 is essentially a function of only three traits, Wing Length, Beak Length, and Beak Length at 4 mm, and hence may be said to incorporate population differences in shape. Together PC1 and PC2 account for over 95% of total variation among populations; PC3 and subsequent components contribute substantially less ($\leq 3.5\%$).

Figure 2 shows the positions of means for individual populations along PC1 and PC2. In size (PC1) the Genovesa population is the smallest form, and San Salvador and Darwin are the largest. Populations on Darwin and Wolf possess the relatively longest wings, and the longest, most pointed beaks (PC2). In the three highland forms the wing is relatively short, and the beak relatively short and blunt.

TABLE 4. Morphological distance (M) and diet difference (D) between each *G. difficilis* population and the unweighted mean of *G. fuliginosa* and *G. scandens* populations. M is the Euclidean distance based on the six morphological variables from which principal components were derived (Table 3). D is $\frac{1}{2}$ the Manhattan distance, and varies from 0 (no difference) to 1 (maximum difference). $1 - D$ is the familiar Renkonen index of niche overlap (Hurlbert, 1978). Data are from Table 5. Underlined values indicate islands where *G. fuliginosa* and *G. scandens* are absent, but their major foods are present. *Geospiza scandens* is absent from Genovesa but is replaced there by *G. conirostris* (Fig. 1). Fernandina also lacks *G. scandens*, but cactus is uncommon there (pers. observ.).

<i>difficilis</i>	<i>fuliginosa</i>		<i>scandens</i>	
	M	D	M	D
Pinta	.11	.77	.23	.97
Fernandina	.10	.74	.25	.95
Genovesa	.14	.33	.28	.91
San Salvador	.15	.69	.19	.95
Wolf	.19	.56	.18	.88
Darwin	<u>.21</u>	<u>.52</u>	<u>.14</u>	<u>.93</u>

Mean scores for adult males were also determined on the same axes for populations of *G. fuliginosa* and *G. scandens* (Fig. 2) to allow a direct comparison with *G. difficilis*. These two axes account for 80.4% of the variance among population means for all three species combined. The relationships among populations and species shown are the same when the PC's are instead determined from the combined species pool. *Geospiza scandens* is larger than *G. difficilis* and has a relatively long and pointed beak, and relatively long wings. *Geospiza fuliginosa* is slightly smaller, with a relatively short, blunt beak.

The data in Figure 2 are largely consistent with Lack's (1947) original comparison. The Genovesa form of *G. difficilis* is similar to *G. fuliginosa* in size, notably Tarsus Length and also Body Mass (Table 2). The Darwin and Wolf populations are similar to *G. scandens* in both size and shape. However, the morphological relations among populations and species are more complex than Lack realized. For example, populations of *G.*

TABLE 5. Dry-season proportions of time spent feeding on foods of different types and in different feeding positions. Dates are given in Table 1. Values are averages over all sites, weighted by the density of birds (census walk estimates). Daphne data are from Sept. 1979, provided by S. J. Millington. *Geospiza scandens* diets are not available for other sites. See text for description of diet categories. Values $\geq 10\%$ of diet are given in boldface.

	Ground			Vegetation		Pollen nectar	Opuntia	Scavenging	Blood	Other	No. sec. obs.
	Invertebrates	Concealed seeds	Exposed seeds	Invertebrates	Seeds						
<i>difficilis</i>											
Pinta	.78	.17	.01	.04	.01						38,915
San Salvador	.59	.30	.04	.07							14,150
Fernandina	.02	.93	.05								9,705
Darwin	.29	.28	.34		.06			.02	.01		5,045
Wolf		.60	.17	.04			.05	.04	.07	.02	17,535
Genovesa	.04	.29	.36		.02	.23	.02	.05			41,990
<i>fuliginosa</i>											
Pinta		.05	.16	.05	.63	.10	.01			.02	103,990
San Salvador		.24	.54	.01	.19					.02	26,405
Fernandina		.07	.61	.01	.29	.01					25,870
Marchena	.01		.49		.21	.28					51,960
Española		.56	.08	.01						.34	25,365
Tortuga			.59	.30				.12			10,868
<i>scandens</i>											
Pinta							1.00				2,900
Daphne			.11	.03	.03		.84				†

† Based on 37 point-observations.

difficilis on Fernandina (not examined by Lack) and Pinta are also similar to *G. fuliginosa* in size and shape (Fig. 2). This is confirmed in Table 4, which summarizes morphological distances between species. The Fernandina, Pinta, and Genovesa forms of *G. difficilis* are all quite close to the overall mean for *G. fuliginosa*. The Darwin form of *G. difficilis* is most similar to *G. scandens*, with Wolf and San Salvador farther off and essentially equidistant. These distance relationships are the same if Body Mass (cube root) is included with the other six variables, or if the analysis is restricted to the three dimensions that Lack measured: Wing Length, Beak Length, and Beak Depth.

Diets and Food Availability

Geospiza populations are most likely to be food-limited in the dry season, and at this time diets should most closely reflect morphological attributes (Smith et al., 1978; Grant and Grant, 1980a). Dry

season diets are shown in Table 5. The food spectrum is divided on the basis of both food type and foraging position; previous study has shown that both aspects of the feeding niche distinguish *G. difficilis* from *G. fuliginosa* and *G. scandens* (Schluter, 1982a). This classification obscures some differences between species within categories (e.g., food size differences) but these are small compared to differences among categories. In general *G. fuliginosa* and *G. difficilis* do not differ in the ranges of seed size consumed. However, *G. scandens* usually takes larger seeds than both these species (Abbott et al., 1977; pers. observ.).

Concealed seeds (Table 5) refer to those obtained by digging in the litter or by shoving aside small stones. Most ground invertebrates are obtained in the same way. Exposed seeds are removed directly from the soil, rock, or litter surface. Scavenged items include decaying fish, seabird eggs, and undigested fecal remains from seabirds. Blood is a special cate-

TABLE 6. Factor loadings for the first four principal diet components (DC1-4) in populations of *G. fuliginosa*, *G. difficilis*, and *G. scandens* (Table 5). The cumulative variance accounted for by successive components is given below. *N* = 14 populations.

Variables	Scores			
	DC1	DC2	DC3	DC4
Ground invertebrates	-.16	.35	-.82	.07
Ground concealed seeds	-.43	.55	.54	-.01
Ground exposed seeds	-.15	-.62	.08	.50
Vegetation invertebrates	-.01	-.07	-.05	.19
Vegetation seeds	-.03	-.34	-.02	-.82
Pollen and nectar	-.02	-.13	-.01	-.09
Opuntia	.87	.22	.13	.07
Scavenging	-.01	-.03	-.01	.11
Blood	-.01	.00	-.02	.01
% Variance	38.8	68.0	87.6	95.6

gory: *G. difficilis* on Wolf is known to draw and consume blood from the base of the retrices in seabirds, especially boobies (*Sula* spp.; Bowman and Billeb, 1965). Our observations are the first of this behavior on Darwin. *Opuntia* includes food resources obtained from this plant which are most important in the diet of *G. scandens*: seeds, fruits, insect larvae and pupae, nectar, and pollen (Grant and Grant, 1980b, 1981). The remaining vegetation categories, seeds, invertebrates, and pollen/nectar include foods produced on plants other than *Opuntia*.

Table 6 summarizes major differences among populations in the use of foods and feeding positions. Principal components were derived from covariances computed from untransformed diet proportions for populations of all three species combined. Only the first nine diet categories were used (Table 5). Values were rescaled to sum to 1 before computation. Population positions along the first four diet components (DC's) are shown in Figure 3.

DC1 strongly represents variation among populations in the use of *Opuntia* (Table 6). *Geospiza scandens* is essentially an *Opuntia* specialist (Table 5; see also Grant and Grant, 1981) and it is widely separated along DC1 from all populations of *G. fuliginosa* and *G. dif-*

ficilis (Fig. 3a). DC2 summarizes major differences between the latter two species. Overall, *G. difficilis* ate concealed invertebrates and seeds more often than did *G. fuliginosa*, which consumed more exposed seeds on the ground and vegetation (Tables 5, 6). *Geospiza difficilis* populations on the high islands of Pinta, San Salvador and Fernandina were most distinct from *G. fuliginosa* in this regard.

DC3 summarizes additional differences among populations of *G. difficilis* (Fig. 3b): forms on San Salvador and Pinta take invertebrates (particularly snails) more often, and concealed seeds less often, than populations on other islands (Tables 5, 6). DC4 represents variation among *G. fuliginosa* populations in the relative degree to which seeds are taken from the surface of the ground or vegetation. Subsequent components account for only minor amounts of the total variation in feeding ($\leq 3\%$).

Food availability differed among islands. This explains some interpopulation variation in feeding. Table 7 provides ranges of food density estimates, and demonstrates some important qualitative differences among islands. For example, ground invertebrates were relatively rare on Genovesa and Wolf, and this may explain their infrequent occurrence in *G. difficilis* diets there (Table 5). Pollen and nectar were likewise rare on

some islands, and hence were rare or absent from *G. fuliginosa* diets there. Food availability does not simply explain all inter-island variation in diets. For example exposed seeds on rock and soil surfaces are present on all islands where *G. difficilis* occurs, yet these are heavily exploited only on the three low islands.

Associations Between Morphology and Diet

Lack (1947) discussed morphological variation among populations of *G. difficilis* primarily in terms of their morphological similarities with *G. fuliginosa* and *G. scandens*. He suggested that *G. difficilis* populations similar to either *G. fuliginosa* or *G. scandens* in morphology would resemble them in diet. This can be examined with data in Table 4 which compares morphological (*M*) and diet (*D*) differences between individual *G. difficilis* populations and the mean for *G. fuliginosa* and *G. scandens*.

As Lack predicted, *G. difficilis* on Genovesa is the population most similar in diet to *G. fuliginosa* (see also Fig. 3). The data also provide partial support for the idea that *G. difficilis* on Darwin and Wolf combine the feeding niches of the absent *G. fuliginosa* and *G. scandens*. *Geospiza difficilis* on both Darwin and Wolf are moderately similar to *G. fuliginosa* in diet, more so than any other *G. difficilis* population except Genovesa. However, while the Darwin and Wolf populations are more similar to *G. scandens* than are most other *G. difficilis* populations, the resemblance is unexpectedly low. Also, the small amount of diet overlap which exists between these *G. difficilis* forms and the *Opuntia* specialist, *G. scandens*, is due more to their similar use of seeds and invertebrates (DC2 and DC3) than to their similar use of *Opuntia* (DC1; Fig. 3).

The feeding characteristics of *G. difficilis* on Darwin and Wolf contribute to a weak correlation between overall morphology and diet (Table 4). *M* and *D* are only weakly positively correlated between individual *G. difficilis* populations

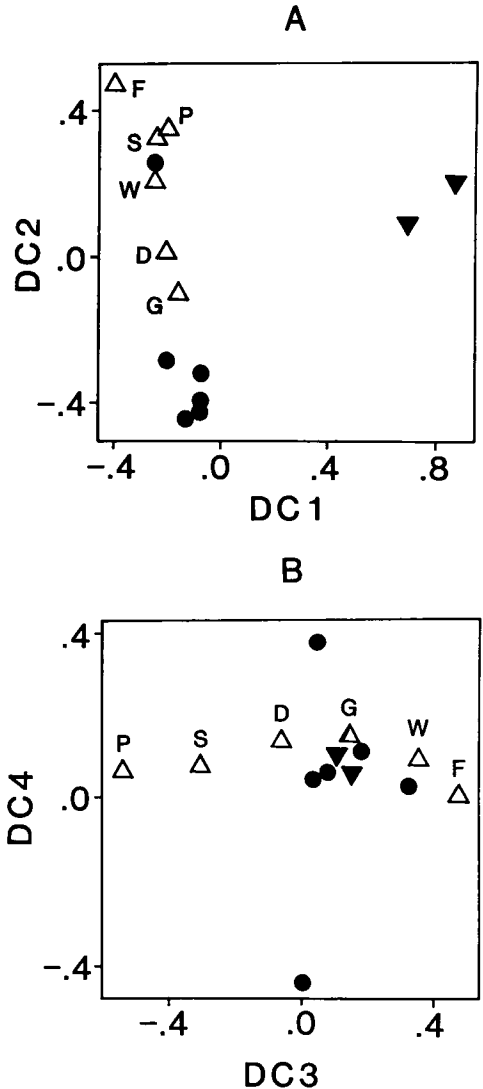


FIG. 3. Positions of *G. difficilis*, *G. fuliginosa*, and *G. scandens* populations along the first four principal diet components (DC). Loadings for original variables are given in Table 6. Symbols as in Figure 2.

($N = 6$) and *G. scandens* ($r = .15$). *M* and *D* are negatively correlated between individual *G. difficilis* populations and *G. fuliginosa* ($r = -.49$). This unexpected trend results from the Darwin and Wolf forms of *G. difficilis* being morphologically the most distant from *G. fuliginosa*, while they are among the forms most

TABLE 7. Food characteristics in the dry season. Figures indicate the range of variation in mean values for different sites on the study islands. No quantitative data are available for Darwin. Small seeds include all seeds commonly eaten by *G. fuliginosa* or *G. difficilis* when available. Certain abundant seeds eaten infrequently are excluded (see Schluter, 1982a).

	Volume litter (l/m ²)	Density ground inverts. (mg/m ²)	Density small seeds (mg/m ²)	Density nectar- producing flowers (no./m ²)	Percent cover <i>Opuntia</i>
Pinta	2.5-9.0	2.4-162	23.9-302	.0-1.3	.0-3.4
San Salvador	1.0-14.1	10.7-470	1.5-156	0	0
Fernandina	.5-9.3	1.1-55.4	11.5-506	0	0
Darwin	?	?	?	0	0
Wolf	.3	39.9	66.2	0	23.0
Genovesa	1.9-2.5	7.2-15.5	1.1-24.1	.0-.7	6.4-12.8
Marchena	.6-9	13.6	5.0-6.2	.0-2.7	.0-6.4
Española	.0-.8	2.1-5.1	12.5-28.4	0	.0-6.4
Tortuga	.2	8.0	12.7	0	0

similar to *G. fuliginosa* in diet. Also, while *G. difficilis* on Pinta and Fernandina are as close to *G. fuliginosa* in morphology as is the population on Genovesa, they are the most distant from *G. fuliginosa* in diet.

DISCUSSION

Some general features of Lack's hypothesis concerning the evolution of *G. difficilis* populations in relation to the distribution of competitors are supported by our field studies. As Lack (1947) predicted from specific morphological considerations, lowland populations of *G. difficilis* are very like *G. fuliginosa* in diet. *Geospiza difficilis* occurs on low islands only where *G. fuliginosa* is not present, and this is best attributed to competitive exclusion on low islands (Schluter and Grant, 1982). On high islands the diet of *G. difficilis* is quite different from the diet of the lowland form, and this is associated with food differences between the habitat types. The implication is that the diversity of habitats occupied by *G. difficilis* has been influenced by competitors, and this in turn has directly influenced the degree of morphological variation exhibited.

Attempts to understand more precisely how food conditions have influenced morphological variation in *G. difficilis* are complicated by a weak overall correlation between morphology and diet (Table

4). For example, it is not clear from the diet data why the Darwin and Wolf forms of *G. difficilis* are close to *G. scandens* in morphology. Nor is it clear why, relative to its highland populations, *G. difficilis* on Genovesa has converged to *G. fuliginosa* in some morphological traits (e.g., Body Mass, Tarsus Length) but not in others (Beak Length, Beak Length at 4 mm; Table 2, Fig. 2). Below we discuss possible factors which may influence diet-morphology correlations, and we attempt to account for these factors in an explanation for morphological patterns in *G. difficilis*.

Environmental Variation

Morphological traits may not always reflect diet closely because of opportunistic feeding under variable food supply conditions. A correlation is expected only under food limitation, when natural selection may most strongly influence morphology (Wiens, 1977; Smith et al., 1978; Grant and Grant, 1980a; Wiens and Rotenberry, 1980, 1981; Boag and Grant, 1981).

If selection in *G. difficilis* is related to low food supplies, then the most critical period in any year is likely to be near the end of the dry season, after most foods have been depleted, but before new flowers and fruits appear. Our studies were conducted in the dry season, but possibly some visits to islands preceded the period

of greatest resource scarcity. Darwin and Wolf were visited rather early in the dry season compared to other islands (Table 1) and seed density was still quite high on Wolf compared to other low islands (Table 7).

Possibly, *G. difficilis* on Darwin and Wolf exploit *Opuntia* more heavily later in the dry season (Köster and Köster, 1983), and this would explain why they have a long beak, similar to that in *G. scandens*. Or, the long beak in these two forms may be an adaptation for probing *Opuntia* flowers which bloom in the wet season (and possibly late in the dry season). Beck (Rothschild and Hartert, 1902) reported *G. difficilis* on Wolf feeding on flowers in the breeding season, and we observed the same behavior there in early April, 1978.

Alternatively, a long beak in *G. difficilis* on these two islands might be an adaptation to blood feeding. Visitors to Wolf in the late dry season of different years note the high frequency of blood-feeding by *G. difficilis* (Köster and Köster, 1983; S. Harcourt, pers. comm.). In September 1981 this behavior constituted 7% of the diet on Wolf (Table 5), but by December of the same year the proportion had increased substantially (S. Harcourt, pers. comm.). A long beak would probably facilitate probing through feathers and skin of seabirds. It may also be advantageous for a related reason: *G. difficilis* on both Darwin and Wolf is known to use its beak to pierce the eggs of seabirds which nest there in large numbers (Köster and Köster, 1983; R. I. Bowman, pers. comm.).

However, data suggest that the environmental variation hypothesis is not a sufficient explanation for the weak diet-morphology correlation recorded. First, most observations for this study were collected in the late dry season, by which time finch numbers had declined in correspondence with dwindling food supplies (Schluter, 1982a; Schluter and Grant, 1982, 1984). Food supply was therefore probably limiting. Second, repeat observations on dry season diets re-

corded at different times in the same year and in different years give similar results to those in Table 5. *Geospiza difficilis* on Genovesa is one of the best-studied populations in the Galápagos, and it has been found to be invariably highly similar to *G. fuliginosa* in diet (Smith et al., 1978; Grant and Grant, 1980a; Schluter and Grant, 1982; pers. observ.). *Geospiza difficilis* on Santa Fe and other highland islands is generally quite different from *G. fuliginosa* in diet (Schluter, 1982a; pers. observ.). Finally, during the course of this study *G. scandens* and *G. fuliginosa* were generally observed to be feeding on those foods to which they are thought to be morphologically adapted (Table 5; Abbott et al., 1977; Smith et al., 1978; Grant and Schluter, 1984). It is therefore unlikely that this would not also be true for most *G. difficilis* populations observed at the same time.

These observations suggest that some intrinsic factors may influence diet-morphology correlations, a possibility discussed next.

Morphological Interdependence

Previous comparisons of *Geospiza* morphology and diet have concentrated on populations differing mainly in general beak and body size, and this variable is usually correlated with size and hardness of seeds consumed (Abbott et al., 1977; Schluter, 1982b; Schluter and Grant, 1984). However, populations of *G. difficilis* differ from one another and from related species in both size and shape (Fig. 2), and they use different foraging positions and food types. It is possible that the poor association between morphology and diet in this study is partly a consequence of evolutionary interdependence among those traits responsible for size-shape differences.

There are at least two ways in which traits may be evolutionarily interdependent. First, traits may be genetically correlated (Boag, 1983; Grant, 1983), and some traits will evolve as a correlated response to natural selection on other

traits (Lande, 1979). Second, there may be alternative morphological solutions to the problem of exploiting particular food types. A simple consequence of such interactions is that final morphology resulting from response to a particular selection pressure may vary depending on the initial morphological and genetical characteristics of that population. Two species initially different morphologically may thus evolve to two different morphological forms in the same environment.

These considerations may help explain why populations of *G. fuliginosa* and *G. difficilis* differ in some morphological traits, even though their diets may be similar. In each species traits are correlated among populations, and lines of interpopulation allometry are largely parallel (Fig. 2). Thus the Genovesa form of *G. difficilis* has converged in general size to *G. fuliginosa*, but it retains the relatively long and pointed beak that is characteristic of the species (Fig. 2). Interestingly, in general size *G. difficilis* on Genovesa is most like the Pinta and Marchena forms of *G. fuliginosa*, which are themselves the smallest forms of this species. This is associated with a nectar-feeding habit in all three populations, in addition to the consumption of exposed seeds (Table 5). Since nectar is frequently removed from small flowers on thin branches, metabolic efficiency as well as perching ability are probably major selective factors in the evolution of small size in these three populations.

The different beaks in these three populations are used in a different manner, but to the same end. One of the most important sources of nectar on Pinta, Marchena and Genovesa is the small flower of *Waltheria ovata*. While the sharp-beaked *G. difficilis* on Genovesa probes the flowers with the entire beak to obtain nectar, the more blunt-beaked *G. fuliginosa* on the other two islands uses only the lower mandible (pers. observ.). We do not know whether the two species remove nectar with equal efficiency, although they appear to do so;

each probe at a maximal rate of 30–40 flowers/min (pers. observ.).

Other Factors

While diet per se is probably the major determinant of morphological variation in *G. difficilis*, other environmental factors may also be important. A possible example involves Wing Length, which varies among populations partly independently of general size (Table 3). Relative Wing Length is greatest in the lowland populations of *G. difficilis* (Fig. 2). This is associated with an arid-zone distribution (Fig. 1), and may allow for more efficient heat-loss in such habitats (Hamilton, 1961; James, 1970). Flight energetics could also influence variation in relative Wing Length (e.g., Feinsinger et al., 1979). Lowland forms of *G. difficilis* seem to fly more frequently, and they are less restricted in their movements in space than are highland forms (Schluter, 1984). This may be a consequence of differences among habitats in the dispersion and renewal patterns of resources exploited.

SUMMARY

The morphology of *G. difficilis* varies on different islands in the Galápagos (Fig. 2). Lack (1947) argued that beak and body size in this species are related to feeding, and noted that variation in morphology is associated with inter-island differences in the presence or absence of potential competitor species. We investigated the relationship between inter-island morphological variation in *G. difficilis* and in two potential competitor species, *G. fuliginosa* and *G. scandens*, and their dry season diets.

Our observations support some of Lack's general conclusions concerning the significance of morphological variation in *G. difficilis*. For example, populations of *G. difficilis* are similar in diet to *G. fuliginosa* on islands where this species is absent. However, overall morphological distances between populations of *G. difficilis* and the other two species did not successfully predict diet differences. There are at least two reasons for discrepancies.

First, environments vary seasonally and annually, and populations may not all have been studied during the most critical periods of food limitation. Second, traits have not evolved independently in *G. difficilis*, and this complicates diet-morphology comparisons, for example when there are alternative morphological solutions to a single foraging problem. Consideration of these factors helps clarify the significance of variation among *G. difficilis* populations in certain morphological traits. Beak length and body size appear to be particularly important in relation to food. On the low island of Genovesa, *G. difficilis* has become smaller, and this is correlated with a diet of small seeds and flower nectar. On the low islands of Darwin and Wolf *G. difficilis* has a relatively longer beak than other populations have. This may be an adaptation to cactus-feeding, but some data indicate that it may have evolved to facilitate blood- and egg-feeding.

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LITERATURE CITED

- ABBOTT, I., L. K. ABBOTT, AND P. R. GRANT. 1977. Comparative ecology of Galápagos ground

- finches (*Geospiza* Gould): evaluation of the importance of floristic diversity and interspecific competition. *Ecol. Monogr.* 47:151-184.
- BOAG, P. T. 1983. The heritability of external morphology in the Darwin's finches (*Geospizinae*) of Daphne Major Island, Galápagos. *Evolution* 37:877-894.
- BOAG, P. T., AND P. R. GRANT. 1981. Intense natural selection in a population of Darwin's finches (*Geospizinae*) in the Galápagos. *Science* 214:82-85.
- BOWMAN, R. I., AND S. I. BILLEB. 1968. Blood-eating in a Galápagos finch. *Living Bird* 4:29-44.
- FALCONER, D. S. 1981. Introduction to Quantitative Genetics, 2nd ed. Longman, London.
- FEINSINGER, P., R. K. COLWELL, J. TERBORGH, AND S. B. CHAPLIN. 1979. Elevation and the morphology, flight energetics, and foraging ecology of tropical hummingbirds. *Amer. Natur.* 113:481-497.
- GATZ, A. J. 1979. Community organization in fishes as indicated by morphological features. *Ecology* 60:711-718.
- GRANT, B. R., AND P. R. GRANT. 1981. Exploitation of *Opuntia* by birds on the Galápagos. *Oecologia* 49:179-187.
- GRANT, P. R. 1981. Speciation and the adaptive radiation of Darwin's finches. *Amer. Sci.* 69:653-663.
- . 1983. Inheritance of size and shape in a population of Darwin's finches, *Geospiza conirostris*. *Proc. Roy. Soc. London. B* 220:219-236.
- GRANT, P. R., AND P. T. BOAG. 1980. Rainfall on the Galápagos and the demography of Darwin's finches. *Auk* 97:227-244.
- GRANT, P. R., AND B. R. GRANT. 1980a. The breeding and feeding characteristics of Darwin's finches on Isla Genovesa, Galápagos. *Ecol. Monogr.* 50:381-410.
- . 1980b. Annual variation in finch numbers, foraging and food supply on Isla Daphne Major, Galápagos. *Oecologia* 46:55-62.
- GRANT, P. R., AND D. SCHLUTER. 1984. Interspecific competition inferred from patterns of guild structure. In D. R. Strong, D. S. Simberloff, L. G. Abele, and A. B. Thistle (eds.), *Ecological Communities: Conceptual Issues and the Evidence*. Princeton Univ. Press, Princeton.
- HAMANN, O. 1979. Regeneration of vegetation on Santa Fe and Pinta islands, Galápagos, after the eradication of goats. *Biol. Conserv.* 15:215-236.
- . 1981. Plant communities of the Galápagos islands. *Dansk Botanisk Arkiv* 34:1-163.
- HAMILTON, T. H. 1961. The adaptive significance of intraspecific trends of variation in wing length and body size among bird species. *Evolution* 15:180-195.
- HARRIS, M. P. 1973. The Galápagos avifauna. *Condor* 75:265-278.
- HURLBERT, S. H. 1978. The measurement of niche overlap and some relatives. *Ecology* 59:67-77.
- JAMES, F. C. 1970. Geographic size variation in

- birds and its relationship to climate. *Ecology* 51: 365-390.
- . 1982. The ecological morphology of birds: a review. *Ann. Zool. Fennici* 19:265-275.
- KEAST, A. 1972. Ecological opportunities and dominant families, as illustrated by the neotropical Tyrannidae (Aves). *Evol. Biol.* 5:229-277.
- KÖSTER, F., AND H. KÖSTER. 1983. Twelve days among the "vampire finches" of Wolf island. *Noticias de Galápagos* 38:4-10.
- LACK, D. 1947. *Darwin's Finches*. Cambridge Univ. Press, Cambridge.
- LANDE, R. 1979. Quantitative genetic analysis of multivariate evolution, applied to brain: body size allometry. *Evolution* 33:402-416.
- PIMENTEL, R. A. 1979. *Morphometrics, the Multivariate Analysis of Biological Data*. Kendall/Hunt, Dubuque, Iowa.
- PRICE, T. D. 1984. The evolution of sexual size dimorphism in Darwin's finches. *Amer. Natur.* 123:500-518.
- RICKLEFS, R. E., D. COCHRAN, AND E. R. PIANKA. 1981. A morphological analysis of the structure of communities of lizards in desert habitats. *Ecology* 62:1474-1483.
- RICKLEFS, R. E., AND J. TRAVIS. 1980. A morphological approach to the study of avian community organization. *Auk* 97:321-338.
- ROTHSCHILD, W., AND E. HARTERT. 1902. Further notes on the fauna of the Galápagos Islands. *Notes on the birds*. *Novit. Zool.* 9:381-418.
- SCHLUTER, D. 1982a. Distributions of Galápagos ground finches along an altitudinal gradient: the importance of food supply. *Ecology* 63:1504-1517.
- . 1982b. Seed and patch selection by Galápagos ground finches: relation to foraging efficiency and food supply. *Ecology* 63:1106-1120.
- . 1984. Feeding correlates of breeding and social organization in two Galápagos finches. *Auk* 101:59-68.
- SCHLUTER, D., AND P. R. GRANT. 1982. The distribution of *Geospiza difficilis* in relation to *G. fuliginosa* in the Galápagos islands: tests of three hypotheses. *Evolution* 36:1213-1226.
- . 1984. Determinants of morphological patterns in communities of Darwin's finches. *Amer. Natur.* 123:175-196.
- SIMBERLOFF, D., AND W. BOECKLEN. 1981. Santa Rosalia reconsidered: size ratios and competition. *Evolution* 35:1206-1228.
- SMITH, J. N. M., P. R. GRANT, B. R. GRANT, I. J. ABBOTT, AND L. K. ABBOTT. 1978. Seasonal variation in the feeding habits of Darwin's ground finches. *Ecology* 59:1137-1150.
- WIENS, J. A. 1977. On competition and variable environments. *Amer. Sci.* 65:590-597.
- WIENS, J. A., AND J. T. ROTENBERRY. 1980. Patterns of morphology and ecology in grassland and shrubsteppe bird populations. *Ecol. Monogr.* 50:287-308.
- . 1981. Morphological size ratios and competition in ecological communities. *Amer. Natur.* 117:592-599.
- WIGGINS, I. L., AND D. M. PORTER. 1971. *Flora of the Galápagos Islands*. Stanford Univ. Press, Stanford.

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