# Convergent evolution of weakly electric fishes from floodplain habitats in Africa and South America

Kirk O. Winemiller & Alphonse Adite Department of Wildlife and Fisheries Sciences, Texas A&M University, College Station, TX 77843, U.S.A.

Received 19.7.1995 Accepted 27.5.1996

Key words: diet, electrogenesis, electroreception, foraging, morphology, niche, Venezuela, Zambia

# **Synopsis**

An assemblage of seven gymnotiform fishes in Venezuela was compared with an assemblage of six mormyriform fishes in Zambia to test the assumption of convergent evolution in the two groups of very distantly related, weakly electric, noctournal fishes. Both assemblages occur in strongly seasonal floodplain habitats, but the upper Zambezi floodplain in Zambia covers a much larger area. The two assemblages had broad diet overlap but relatively narrow overlap of morphological attributes associated with feeding. The gymnotiform assemblage had greater morphological variation, but mormyriforms had more dietary variation. There was ample evidence of evolutionary convergence based on both morphology and diet, and this was despite the fact that species pairwise morphological similarity and dietary similarity were uncorrelated in this dataset. For the most part, the two groups have diversified in a convergent fashion within the confines of their broader niche as nocturnal invertebrate feeders. Both assemblages contain midwater planktivores, microphagous vegetation-dwellers, macrophagous benthic foragers, and long-snouted benthic probers. The gymnotiform assemblage has one piscivore, a niche not represented in the upper Zambezi mormyriform assemblage, but present in the form of *Mormyrops deliciousus* in the lower Zambezi and many other regions of Africa.

#### Introduction

It is widely recognized that electrogeneration and electroreception evolved independently in African fishes of the order Mormyriformes (elephant fishes, bottlenoses, stonebashers) and Neotropical fishes of the order Gymnotiformes (cuchillos, or knifefishes) (Roberts 1972, Lowe-McConnell 1975, Kirschbaum 1984, Marrero & Winemiller 1993). Both groups of tropical freshwater fishes generate weak electric fields that are used for navigation, prey detection, and species and sex identification (Lissman 1963, Hopkins & Bass 1981, Heiligenburg & Bastian 1984). The two phylogentic lineages are very distantly related, with mormyriforms members of an old lineage allied with the bony tongues

(infradivision Osteoglossimorpha) and gymnotiforms part of the more recent superorder Ostariophysi (infradivision Euteleostei) that also contains catfishes (Siluriformes) and tetras (Characiformes) (Fink & Fink 1981, Lundberg 1993). This convergent capacity to generate and perceive weak electromagnetic fields should result in convergent ecological niches, and, indeed, both groups are principally nocturnal, benthic, and predaceous on aquatic invertebrates. Both groups tend to be associated with lotic habitats and lentic habitats of river floodplains, and their species richness generally diminishes when rivers are impounded (Corbet 1961, Petr 1968, Balon 1974, Blake 1977, Winemiller 1989).

Recently, Marrero & Winemiller (1993) examined the marked morphological and apparent ecolog-

ical convergence of tube snouted mormyriform and gymnotiform fishes. Two species have independently evolved a highly specialized head morphology that enables the fish to extract benthic insect larvae from tiny holes and crevices in clay nodules and woody debris. We now ask the related question, to what degree do faunas of weakly electric fishes with similar numbers of species from similar habitats show morphological and ecological convergence? Because the two groups occupy the same nocturnal, invertebrate-feeder niche in speciose tropical faunas, one intriguing possibility is that they have evolved similar solutions for foraging and competing. Alternatively, patterns of morphological divergence and resource utilization might be entirely different in the two groups, even when the same number of species occur in very similar habitats in Africa and South America. The latter result is more probable, given the potential influence of differing geological and climatic histories, divergent continental fish faunas, specific differences in resources, and stochastic ecological influences. Here we compare morphological and dietary patterns of mormyriform and gymnotiform faunas from seasonal floodplain habitats. Given that complete (one-forone) evolutionary convergence at the level of local species assemblages is extremely unlikely, the degree of convergence detected is impressive. We briefly discuss some factors that influence the detection and interpretation of ecological convergence.

#### **Methods**

#### Electric fish assemblages

We define the spatial boundaries of the two local fish assemblages in the following manner: Caño Maraca, a low gradient creek and its associated floodplain in the savanna (llanos) of Venezuela's Portuguesa state (Winemiller 1990); and the Upper Zambezi river and associated floodplain wetlands in the savanna of Zambia's Western Province (Winemiller 1991a). The two sites were chosen for comparison because they have similar aquatic habitats, seasonal hydrology, and numbers of electric

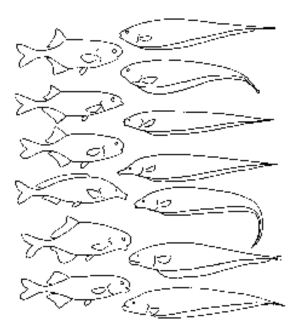


Figure 1. Mormyriform (left column) and gymnotiform (right column) electric fishes from the Zambian and Venezuelan study sites. Left column from top to bottom (all members of the Mormyridae): Petrocephalus catastoma, a small midwater swimmer; Marcusenius macrolepidotus, a medium size benthic and vegetation forager; Pollimyrus castelnaui, a small vegetation-dweller; Mormyrus lacerda, a large benthic and vegetation forager; Hippopotamyrus dischorhynchus, a medium-size benthic forager; and Hippopotamyrus ansorgii, a medium-size benthic forager. Right column from top to bottom: Eigenmannia virescens (Sternopygidae), a medium size midwater swimmer; Adontosternarchus devananzii (Apteronotidae), a medium-size vegetation dweller; Brachyhypopomus occidentalis (Hypopomidae), a medium-size vegetation dweller; Rhamphichthys marmoratus (Rhamphichthyidae), a large benthic and vegetation forager; Sternopygus macrurus (Sternopygidae), a large benthic forager; Apteronotus albifrons (Apteronotidae), a medium-size benthic forager; and Gymnotus carapo (Gymnotidae), a large midwater and vegetation forager.

fishes; 7 gymnotiforms from Caño Maraca and 6 mormyriforms from the Upper Zambezi (Figure 1). These similarities notwithstanding, the area of the Upper Zambezi floodplain is vastly greater than that of Caño Maraca. Most sites were sampled multiple times during long-term field studies (Winemiller 1996), and habitat data were recorded (channel width, depth, pH, temperature, etc.). Fishes were collected by a variety of methods, including seines, castnets, dipnets, traps, and gillnets, with the objective of capturing all species present at a site in

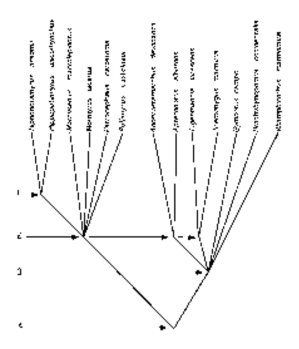


Figure 2. Branching diagram illustrating the presumed phylogentic relationships of species included in the study and the integers used as estimates of phylogenetic distance for the convergence index. This phylogeny is based on recent taxonomy (e.g. Daget et al. 1984, Mago-Leccia 1994) and is intended to have low resolution since our primary interest is interfaunal convergence and because new phylogenetic research is likely to alter relationships within orders. For example, a recent analysis of genetic, morphological, and electrophysiological data by Alves-Gomes et al. (1995) removed Eigenmannia from the family Sternopygidae and placed it in a clade (suprafamilial) containing Apteronotus and Adontosternarchus. A similar analysis by Van Der Bank & Kramer (1996) revealed greater phylogenetic divergence between Hippopotamyrus ansorgii and H. discorhynchus and suggested reinstatement of the latter within the genus Cyphomyrus.

numbers that reflected their relative abundances. With the exception of a presumptive second species of *Brachyhypopomus* that is virtually identical to *B. occidentalis* in gross morphology [possibly *H. diazi* listed as *Hypopomus* sp. 2 in Winemiller (1990)], we include all electric species collected at each site, however one of the included species was rare in the Upper Zambezi (*Hippopotamyrus ansorgii*) and two were rare at Caño Maraca (*Apteronotus albifrons, Sternopygus macrurus*).

#### Phylogenetic relationships

We coded the relative phylogenetic distances between each species pairing in the manner indicated by the nodes of the phylogeny in Figure 2. Species currently assigned to the same genus were given a distance of 1, species in the same family were given a distance of 2, and species in the same order were given a distance of 3. In this dataset, the most distantly related species were any African mormyrid versus any Neotropical gymnotiform; these pairings were assigned a phylogenetic distance of 4.

# Morphological data

Morphological measurements were made on fish specimens catalogued in the Texas Natural History Collection of the Texas Memorial Museum, Austin, Texas. Five specimens of each species were measured, except for Apternotus (N = 4), Rhamphichthys (4), Sternopygus (3), and H. ansorgii (1). The body plans of mormyriforms and gymnotiforms are radically different (e.g., gymnotiforms are very elongate, have elongate anal fins, usually lack dorsal fins, and caudal fins are absent or vestigial). Assuming these attributes are directly involved in feeding, we restricted our morphological analysis to anterior morphology and the gut. Eleven head attributes, pectoral fin length, and gut length were examined. Head attributes were maximum head depth, maximum head width, eye position, eye diameter, mouth orientation, mouth aperture orientation, mouth width, mouth height, snout length, gill raker length, and number of gill rakers. Most measurements are described fully in Winemiller (1991b). Mouth orientation is the angle (from horizontal) formed by an imaginary line passing through the center of the eye's pupil and the corner of the mouth where the upper and lower jaws meet. Aperture orientation is the angle (from horizontal) formed by an imaginary line tangent to the anterior tips of the upper and lower jaws when the mouth is closed. All distance measurements were taken as the straight line distance between points using vernier calipers (to nearest 0.1 mm).

Linear distance measures were standardized for size. We standardized linear measurements to conform to the interfaunal grand mean HL ( $\bar{X}_{HL}$  = 23.3 mm) using x' = cx, where c = 23.3/HL. The use of HL standardized linear measurements allows the first orthogonal axis from principal components analysis (PCA) to be interpreted as a shape deter-

minant rather than a body size axis. Since species scores on the PC axes provide the basis for ecomorphological ordination and the calculation of distance measures, the use of HL standardized measurements reduces the likelihood that the analysis is dominated by body size. Based on five adult specimens per species, species mean values were computed for each of the standardized attributes.

## Analysis of morphological data

Principal components analysis (PCA) was used as a basis for inter-assemblage comparisons of species distributions in morphological space, and to identify patterns of covariation among morphological characters related to feeding. PCA produces independent orthogonal axes (uncorrelated axes) from the original matrix of data points, such that the first several components model a major portion of the variation among the original variables. PCA was performed on the combined 13 species dataset based on the correlation matrix of morphological attributes ( $\log_{10}$ -transformed species mean values) and diets ( $\log_{10}$  + 1-transformed proportions).

Species scores on the PC axes with eigenvalues > 1.0 were used to calculate pairwise Euclidean distances among all species in each dataset. Euclidean distances between species pairs were calculated according to the formula:

$$D_{jk} = [\sum_{i=1}^{n} (x_{ij} - x_{ik})^{2}]^{1/2},$$

where n is the number of PC axes with eigenvalues greater than 1.0, and  $x_{ij}$  and  $x_{ik}$  are the character loadings on the same PC axis for the species pair j and k. From each matrix of Euclidean distances, we ranked each species' nearest neighbors. Mean nearest neighbor distance is an index of assemblage dispersion in morphological space, and the standard deviation of nearest neighbor distance serves as an index of the evenness of species distribution in morphological space (Findley 1973, Ricklefs et al. 1981).

## Ecomorphological convergence

We used the following modified version of Winemiller's (1991b, Winemiller et al. 1995) index of ecomorphological convergence:

$$C_{ij} = R_{ij}/M_{ij},$$

where  $C_{ij}$  is the convergence index for species (i) and nearest neighbor (j), Rii is the number of more closely related species that are more dissimilar to species (i) than ecomorphological nearest neighbor (j),  $M_{ij}$  is the maximum possible number of species that could be more closely related to species (i) relative to the jth nearest neighbor pairing. The index was calculated from the combined 13 species dataset using nearest neighbor distance (Euclidean distance) as the measure of ecomorphological similarity and Figure 2 as the hypothesis of phylogenetic relationships. The convergence index is equal to 1.0 whenever a phylogenetically distant species is ecomorphologically more similar to the target species than each of the species that are determined to to be more closely related. The convergence index equals 0 whenever a species' closest ecomorphological neighbor is actually the most closely related species in the dataset.

# Ecological data

Volumetric proportions of stomachs contents were estimated following the methods in Winemiller (1990). For interfaunal comparisons, diet items were reduced from 160 functional categories (Winemiller 1990) to 13 broad categories. This was done to reduce bias from having different specific resources in the two regions. Stomach contents data were summed across all available size classes within a species, and in most cases, across different seasons. Because larger individuals have greater stomach volumes, the aggregate diet data are very heavily influenced by adult diets. Species' sample sizes for stomach contents appear in the Table 1. Despite the fact that sample sizes for H. ansorgii, Apteronotus, and Sternopygus were insufficient to achieve highly resolved diet characterizations, we include those data to round out the inter-assemblage comparison of diet.

Analysis of morphological and ecological data We performed linear regressions on all combinations of morphological traits with diet items, as well as diet Euclidean distances with morphological Euclidean distances, both based on the combined species dataset. Due to the small number of species in this dataset (13) relative to the number of morphological and diet variables, we were unable to use canonical correspondence analysis (CCA) to examine the multivariate relationship between the dietary and morphological datasets.

### Results

Aquatic invertebrates were the principal food resource for both weakly electric fish assemblages (Table 1). Aquatic dipteran larvae, primarily Chironomidae, were the most important food for 4 of 6 mormyriforms and 3 of 7 gymnotiforms. All 13 spe-

cies consumed aquatic diptera and vegetation/detritus. None of these fishes have long guts or obvious morphological adaptations for herbivory/detritivory, therefore we conclude that most of the vegetation and detritus was probably debris consumed during suction feeding on benthic and macrophytedwelling invertebrates. In numerous instances, vegetative debris was identified as a component of the protective cases constructed by caddisfly larvae, and the volumetric abundances of the diet categories vegetation/detritus and Trichoptera were highly correlated (r = 0.89, p < 0.0001). Twelve species consumed mayfly nymphs (Ephemeroptera) and aquatic beetles (Coleoptera), and 11 species consumed zooplankton (primarily Cladocera, Copepo-

Table 1. Dietary composition [total volume (ml)] of mormyriform and gymnotiform fishes based on 13 food categories: Veget. = vegetation and detritus; Zoopl. = zooplankton and other microcrustacea; Dipt. = aquatic Diptera larvae; Ephem. = Epehemeroptera nymphs; Odona. = Odonata nymphs; Tricho. = Trichoptera larvae; Hemip. = aquatic Hemiptera; Coleop. = aquatic Coleoptera; Mollus. = molluscs (snails, mussels); Worms = Oligochaeta; Fishes = fishes; Shrimp = shrimp; Terr. Arthrop. = terrestrial arthropods.

Species	Sample size	Veget.	Zoopl.	Dipt.	Ephem.	Odona.	Tricho.	Hemip.	Coleop.	Mollus.	Worms	Fishes	Shrimp	Terr. arthrop.
Mormy	riformes:													_
Hippop	otamyrus	ansorgi	i											
	1	0.001	0.002	0.008	0.050	0	0	0	0	0	0	0	0.065	0
Hippop	otamyrus	dischor	hynchus											
	40	0.062	0.022	0.650	0.020	0.102	0.611	0.030	0.070	0.196	< 0.001	0.009	0	0
Marcus	enius mad	crolepido	otus											
	92	0.368	0.312	2.679	0.617	0.063	0.497	0	0.022	0.004	0.093	< 0.001	0.050	< 0.001
Mormy	rus lacero													
	120	4.197	0.011	4.666	2.346	1.745	2.825	0	0.030	0.022	0.049	0.046	0.012	0.075
Petroce	phalus ca													
	49	0.013	0.340	0.097	0.013	0.003	0.086	0	0.011	0	0.008	< 0.001	0	0
Pollimy	rus castel													
	61	0.389	0.073	0.741	0.079	0.005	0.065	0	0.001	0	0.003	0	0.025	0
U	tiformes:													
Adonto:	sternachu													
	58	0.081	0.945	1.906	0.143	0	0.023	0	0.029	0.003	0.008	0.010	0	0.093
Apteron	otus albi													
	2	0.001	0	0.252	0	0	0	0	0.010	0	0	0	0	0.700
Brachyl	hypopom													
	72	0.030	0.429	0.684	0.119	0	0.004	< 0.001	0.082	0	0	< 0.001	0	< 0.001
Eigenm	annia vir										_	_	_	
_	138	0.032	2.620	1.136	0.320	0.092	0.056	0.005	0.112	0.001	0	0	0	0.004
Gymno	tus carap													
	298	1.846	0.747	2.773	1.538	9.874	0.117	0.278	3.748	2.946	0.111	24.244	9.763	1.729
Rhampi	hichthys 1							_		_		_		_
α.	24	0.481	0.227	1.410	0.129	0.303	0.190	0	0.226	0	0.001	0	0	0
Sternop	ygus mac		_				_	_			_	_		_
	5	0.250	0	0.001	0.001	0.010	0	0	0.460	0.140	0	0	0.070	0

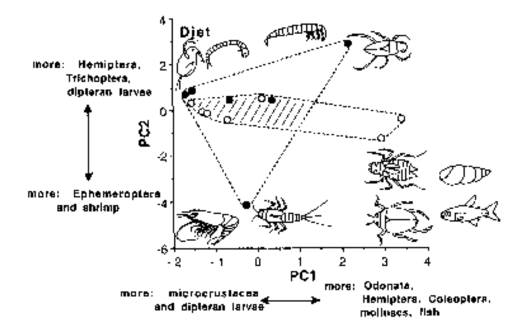


Figure 3. Plot of species scores on the first two PC axes from dietary data. Solid symbols are mormyriforms and open symbols are gymnotiforms.

da and Ostracoda). Because the three species that lacked records for mayflies, beetles, and zooplankton had very small sample sizes, these groups are probably consumed to varying degrees by all weakly electric fishes in these assemblages. Gymnotiforms consumed more terrestrial arthropods than mormyriforms, but terrestrial prey were not major diet components of either assemblage. Fishes were relatively unimportant prey for electric fishes, except for  $Gymnotus\ carapo\ which\ is\ strongly\ piscivorous\ at larger\ sizes\ (Winemiller\ 1989)$ . Correlations between individual diet items and morphological attributes tended to be moderate-low, and only the correlation between mouth height and fish prey was statistically significant (r = 0.64, p < 0.025).

Species ordination with dietary data (Figure 3) showed a larger area of inter-assemblage overlap on the first two PC axes compared with morphological data. Higher scores on the first PC axis were associated with more Odonata, Hemiptera, Coleoptera, Mollusca and fishes, and less zooplankton/microcrustacea and aquatic Diptera (Table 2). The second PC axis described a continuum of more Hemip-

Table 2. PCA statistics from analysis of mormyriform and gymnotiform diet data.

	PCA axis									
	1	2	3	4	5	6	7			
Eigenvalue	3.09	2.45	1.9	1.58	1.31	1.23	0.82			
Proportion of variation	0.24	0.19	0.15	0.12	0.10	0.09	0.06			
Cumulative variation	0.24	0.43	0.58	0.70	0.80	0.89	0.95			
Eigenvectos 1 & 2:										
Vegetation	0.19	0.11								
Zooplankton	-0.32	0.09	)							
Diptera	-0.35	0.34	ļ							
Ephemeroptera	-0.14	-0.44	ļ							
Odonata	0.32	0.06	i							
Trichoptera	0.08	0.45	i							
Hemiptera	0.28	0.31								
Coleoptera	0.37	7 -0.02	:							
Molluscs	0.49	0.14	ļ							
Worms	-0.24	0.15	,							
Fishes	0.29	-0.15	i							
Shrimp	0.12	2 - 0.55	i							
Terr. arthropods	-0.06	-0.06	i							

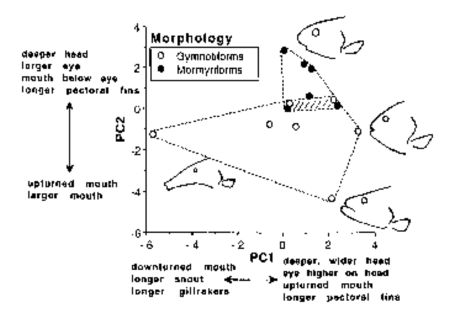


Figure 4. Plot of species scores on the first two PC axes from morphological data (solid symbols = mormyriforms, open symbols = gymnotiforms).

tera, Trichoptera and Diptera, and less Ephemeroptera and shrimp. Two mormyrids (Petrocephalus, Pollimyrus) and three gymnotiforms (Adonto-Eigenmannia) sternarchus. Brachyhypopomus, clustered in the region of low scores on axis 1 and intermediate scores on axis 2. Gymnotus and Sternopygus had highset scores on axis 1. The highest score on diet axis 2 was Hippopotamyrus discorhynchus, and H. ansorgii was lowest on axis 2. The degree of dietary variation was roughly comparable between the two assemblages, with gymnotiforms varying more along axis 1, and mormyriforms varying more along axis 2 (Figure 3). Based on the mean nearest neighbor Euclidean distances computed from species scores on the first six PC axes, the mormyriform assemblage had a more varied diet (mormyriform = 3.10, gymnotiform mean = 2.33). Based on the standard deviation of nearest neighbor Euclidean distances, the degree of dispersion within diet space was approximately the same (mormyriform = 1.71, gymnotiform = 1.61).

Species ordination by the first two principal components (PC1 and 2) from the morphological analysis showed a relatively small region of overlap between the two assemblages (Figure 4). The first PC axis described a gradient of increasing head depth

and width, eyes positioned higher on the head, more upturned mouth, longer pectoral fins, shorter snout, and shorter gill rakers (Table 3). High species

Table 3. PCA statistics from analysis of mormyriform and gymnotiform morphological data.

	PCA axis							
	1	2	3	4	5			
Eigenvalue	5.25	3.37	1.22	1.13	0.89			
Proportion of variation	0.40	0.26	0.10	0.09	0.07			
Cumulative variation	0.40	0.66	0.76	0.85	0.92			
Eigenvectors 1 & 2:								
head depth	0.36	0.26						
head width	0.40	-0.09						
eye position	0.31	0.17						
eye diameter	0.25	0.40						
mouth orientation	0.20	-0.39						
aperture orientation	-0.25	0.39						
mouth width	0.30	-0.30						
mouth height	0.33	-0.15						
snouth length	-0.30	-0.15						
pectoral fin length	0.25	0.37						
gill raker length	-0.26	0.01						
number of gill rakers	-0.17	0.19						
gut length	0.01	-0.35						

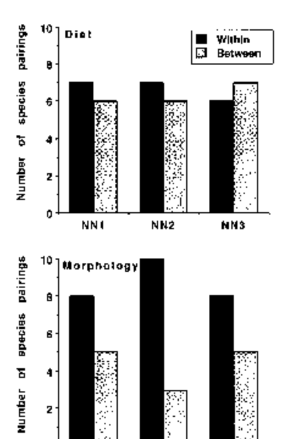


Figure 5. The distributions of within-fauna and between-fauna species pairings for the first three nearest neighbors based on Euclidean distances with dietary and morphological data.

NN2

MN3

NM1

scores on the second axis were associated with a deeper head, larger eyes, a mouth positioned lower on the head, a more downturned mouth aperture, and a smaller mouth. Four extreme morphotypes are illustrated in Figure 4. Adontosternarchus devananzii has a deep head, eyes positioned high on the head, an upturned mouth, relatively long pectoral fins, a short snout, and short gill rakers (highest score on PC1); the long snouted Rhamphichthys marmoratus is at the opposite end of this morphological gradient (lowest score on PC1). Petrocephalus catastoma has a deep head, large eyes, a small mouth low on the head, and relatively long pectoral fins (highest score on PC2); Gymnotus carapo has a large mouth, a dorsoventrally compressed head, and lies at the opposite end of this

gradient (lowest score on PC2). Based on the mean nearest neighbor Euclidean distances computed from species scores on the first four PC axes, the gymnotiform assemblage had greater morphological diversity than the mormyriform assemblage (gymnotiform = 3.37, mormyriform = 2.55). Based on the standard deviation of nearest neighbor Euclidean distances, gymnotiforms were more dispersed in morphological space (gymnotiform = 1.00, mormyriform = 0.60).

Despite the fact that virtually no correlation was observed between morphological Euclidean distance and dietary distance (r = 0.045, p < 0.001, based on 156 species pairings), evidence of inter-assemblage convergent evolution was uncovered. Forty six percent of the first nearest neighbors based on diet were inter-assemblage pairings (Figure 5). Inter-assemblage pairings comprised 46% of the second and 54% of the third dietary nearest neighbors. For morphology, 38%, 23% and 38% of the pairings were between assemblages for the first, second, and third nearest neighbors, respectively. Mean (with standard deviation) Euclidean distances for inter-assemblage morphological pairings were 2.67 (1.05), 2.74 (0.52), and 3.89 (1.06) for the first, second, and third nearest neighbors respectively. Mean (SD) distances for intra-assemblage morphological pairings for nearest neighbors one through three were 2.75 (0.67), 3.42 (0.95), and 3.44 (0.73). Mean (SD) Euclidean distances for inter-assemblage dietary pairings were 2.13 (1.47), 3.76 (1.66), and 2.56 (1.32) for the first, second, and third nearest neighbors, respectively. Mean (SD) distances for intra-assemblage dietary pairings for nearest neighbors one through three were 2.69 (1.91), 1.88 (0.80), and 3.89 (1.39).

The mean convergence index was approximately 0.5 for the first dietary neighbor and 0.4 for the nearest morphological neighbor (Figure 6). The mean convergence index ranged between 0.25 and 0.5 for the second and third dietary and morphological neighbors. The following inter-assemblage convergences were identified based on the criterion of having lowest Euclidean distance for the first nearest neighbor based on morphology: *Apternotus* with *H. ansorgii, Eigenmannia* with *Marcusenius, Rhamphichthys* with *Mormyrus*, and *Sternopygus* with *H.* 

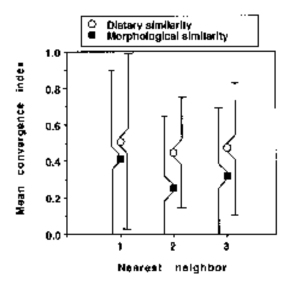


Figure 6. The mean convergence index (with SD) for the first three nearest neighbors based on dietary and morphological data.

ansorgii. Based on diets, the following inter-assemblage pairings had the lowest distances for the first nearest neighbor: *Brachyhypopomus* with *H. ansorgii, Adontosternarchus* with *Marcusenius, Rhamphichthys* with *Mormyrus, Eigenmannia* with *Petrocephalus,* and *Rhamphichthys* with *Pollimyrus.* 

## Discussion

The two groups of weakly electric fishes have similar arrays of ecological niches within the broader ecological classification of nocturnal invertebrate foragers. Macrophagous benthic foragers were represented by a single H. ansorgii from the Zambian samples and a few individuals of Apteronotus and Sternopygus from the Venezuelan samples. These fishes were highly convergent in morphology but weakly convergent based on diets. Diet information was based on very small sample sizes for these three species, and larger samples probably would reinforce the hypothesis of convergent evolution. Marrero and Taphorn (1991) listed Apteronotus albifrons as feeding on benthic insects, shrimp, and fish. Marrero (1987) found that Apteronotus bonapartii, an apternotid with morphology similar to A. albifrons, ate mostly large benthic mayfly nymphs. At other sites, *Sternopygus* feed on mayflies, odonata and other benthic insects, and very large individuals sometimes eat small fish (Winemiller personal observation; Marrero & Taphorn 1991). Skelton (1993) listed *H. ansorgii* as a benthic insect feeder.

Long-snouted electric fishes probe the substrate (e.g., crevices, leaf litter, vegetation mats) for aquatic invertebrates. One long-snouted fish occurred in each assemblage; *Mormyrus* (mormyriform) and *Rhamphichthys* (gymnotiform) were highly convergent in both morphology and diet. The two long-snouted fishes were also the largest weakly electric species captured from their respective assemblages. Species with more extreme long-snouted ecomorphologies exist in each continental fauna (Marrero & Winemiller 1993), but these species tend to inhabit deeper river channels of the Amazon/Orinoco Rivers and central/west Africa.

Several fishes in each assemblage fed heavily on microcrustaceans and micro-insects, chironomid larvae in particular. Among mormyriforms, Petrocephalus tends to be a midwater zooplanktivore (frequently captured by castnet in open water habitats at night), Pollimyrus is strictly a microphagus vegetation-dweller, and Marcusenius forages for aquatic invertebrates over soft bottom sediments and in dense vegetation. Among gymnotiforms, Eigenmannia is the midwater zooplanktivore, Brachyhypopomus is the microphagus vegetation-dweller, and Adontosternarchus devananzii captures invertebrates from the benthos and vegetation. The coloration of these fishes also shows convergence: Petrocephalus and Eigenmannia are light (transparent, white, or silvery) and the other four are mottled brown.

Hippopotamyrus discorhynchus was not strongly convergent with any gymnotiform based on morphology, but *Rhamphichthys* was its second dietary nearest neighbor based largely on the common heavy use of aquatic diptera larvae (chironomids). Whereas *Rhamphichthys* probes soft substrates and crevices with its long snout, the bulbous-snouted *H. discorhynchus* apparently forages for chironomid larvae buried in soft sediments (Skelton 1993). *Gymnotus*, the piscivore, was the other species in the dataset that did not show a strong pattern of in-

terfaunal morphological convergence. Gymnotus's second dietary nearest neighbor was Mormyrus, and this was largely a function of their large diet breadths. For example, Gymnotus fed on every food category, mostly in substantial volumes, and Mormyrus fed on every category except aquatic Hemiptera. Gymnotus has no piscivorous equivalent among upper Zambezi (i.e., above Victoria Falls) mormyriforms, however Mormyrops deliciousus (Mormyridae) is a piscivore that occurs in the middle and lower Zambezi as well as throughout central and western Africa. The head morphologies of Mormyrops and Gymnotus are very similar (e.g., wide head, large supraterminal mouth), and both species show shifts from juvenile diets that are invertebrate dominated to piscivory as adults (Winemiller 1989, Skelton 1993).

Convergent evolution of ecological attributes is not a dichotomous all-or-none phenomenon, rather it can be identified along a continuum by a variety of methods (Winemiller 1991b, Winemiller et al. 1995). The choice of attributes and quantitative analytical methods can influence results. For example, if measurements of detailed osteological attributes provide the basis for comparisons, then it would be very difficult to identify convergent patterns. In some cases, phylogenetically distant taxa might not even share certain osteological components. We chose to use relatively macro-scale morphological attributes that have inferred relationships to ecological performance, in this instance foraging. Given our methods, the identification of a certain amount of weak or intermediate convergence is to be expected. However, the extent of strong interfaunal convergences observed in this dataset is very significant, and indicates that the two groups of weakly electric fishes somehow have been selected to fill a similar array of ecological niches in their respective floodplain faunas.

Given the evidence for convergent ecomorphology between the two assemblages, the lack of correspondence between morphological distances and dietary distances appears problematic. The missing piece to this ecological puzzle is microhabitat use. Anterior morphology determines not only what can be efficiently eaten by a fish, but also how efficiently it forages in different kinds of microhabitats. We did

not have information on foraging frequency in different microhabitats, and this factor might have explained some of the scatter around the regression of morphological and dietary distances. We have already noted how some of the morphologically convergent species pairs forage in similar ways in the same habitats (e.g., substrate probing by long-snouted forms).

Despite the fact that mormyriforms are members of an older lineage, phylogenetic divergence is greater in gymnotiforms (6 families, sensu Mago 1994) than mormyriforms (2 families). The Venezuelan floodplain contained more families (5) and genera (7) of weakly electric fishes than the Zambian floodplain (1 family, 5 genera). Even so, the extent of ecomorphological divergence was similar within the two local fish assemblages, with gymnotiforms having slightly more morphological variation, and mormyriforms having slightly more diet variation. On a continental faunal basis, the two groups probably contain about the same number of species [ca. 200 mormyriforms (Daget et al. 1984); ca. 100 gymnotiforms are described (Mago-Leccia 1994), but new gymnotiforms are being discovered at a much faster rate than new mormyriforms] and show similar ranges of ecomorphological variation associated with a variety of aquatic habitats, including tube-snouted and other benthos-probing insectivores, planktivores, and piscivores. Notable differences do exist; mormyriforms are unique in having species with long lower lip protuberances used for benthic probing (e.g., Gnathonemus petersi), and a few large-mouthed, deep-water gymnotiforms seem to have no mormyriform equivalent (e.g. Apteronotus anas, A. bonapartii). The South American ichthyofauna contains more species than African fauna, but the resultant potential for more interspecific interactions in the Neotropics seems not to have greatly skewed ecomorphological radiation in the two groups of weakly electric fishes. Likewise, the much smaller Caño Maraca floodplain has more fish species (82) than the upper Zambezi floodplain (68 species, see also Winemiller 1996), yet weakly electric fish species richness and ecomorphological variation are similar. While provocative, the results of this study constitute a sample size of one. To test the generality of the findings and perhaps illuminate factors influencing ecological divergence, convergence, and the formation of local species assemblages, comparisons should be performed for weakly electric fish assemblages from other kinds of aquatic habitats and from floodplain habitats in other areas of Africa and South America.

# Acknowledgements

Field studies that contributed data for this comparative analysis were supported by grants to the first author from the National Geographic Society (research in Venezuela) and the Fulbright International Scholars program of the U.S. government (research in Zambia). We thank the numerous individuals who assisted in the field work, especially L. Kelso-Winemiller, D. Taphorn, L. Nico, A. Barbarino, J. Masinja, G. Milini, Mr. Sinda, and W. Ritter. Institutional support abroad was provided by D. Taphorn of the Universidad Experimental de los Llanos Occidentales in Venezuela and E. Muyanga and G. Milindi of the Department of Fisheries of Zambia. Collecting and fishing permits were obtained from the Dirección Administación y Desarollo Pesquero de la Republica de Venezuela and the Department of Fisheries and National Commission of Development Planning of the Republic of Zambia. We also thank D. Hendrickson of the Texas Natural History Collection for specimen loans.

#### References cited

- Alves-Gomes, J.A., G. Ortí, M. Haygood, W. Heiligenberg & A. Meyer. 1995. Phylogenetic analysis of the South American electric fishes (order Gymnotiformes) and the evolution of their electrogenic system: a synthesis based on morphology, electrophysiology, and mitochondrial sequence data. Mol. Biol. Evol. 12: 298–318.
- Balon, E.K. 1974. Fish production of the drainage area and the influence of ecosystem changes on fish distribution. pp. 459–497. *In:* E.K. Balon & A.G. Coche (ed.) Lake Kariba: A Man-Made Tropical Ecosystem in Central Africa, Monographiae Biologicae 24, Dr W. Junk Publishers, The Hague.
- Blake, B.F. 1977. Food and feeding of the mormyrid fishes of Lake Kainji, Nigeria, with special reference to seasonal variation and interspecific differences. J. Fish Biol. 11: 315–328.

- Corbet, P.S. 1961. The food of non-cichlid fishes in the Lake Victoria basin, with remarks on their evolution and adaptation to lacustrine conditions. Proc. Zool. Soc. Lond. 136: 1–101.
- Daget, J., J.-P. Gosse & D.F.E. Van Den Audenaerde (ed.). 1984.
  CLOFFA 1. Check-list of the freshwater fishes of Africa. Musee Royal de l'Afrique Centrale, Tervuren & ORSTOM, Paris. 410 pp.
- Fink, S.V. & W.L. Fink. 1981. Interrelationships of the ostariophysan fishes (Teleostei). Zool. J. Linn. Soc. 72: 297–353.
- Heiligenburg, W. & J. Bastian. 1984. The electric sense of weakly electric fish. Ann. Rev. Physiol. 46: 561–583.
- Hopkins, C.D. & A.H. Bass. 1981. Temporal coding of species recognition signals in an electric fish. Science 212: 85–87.
- Kirschbaum, F. 1984. Reproduction of weakly electric teleosts: just another example of convergent development? Env. Biol. Fish. 10: 3–14.
- Lissman, H.W. 1963. Electric location by fishes. Scientific American 208: 50–59.
- Lowe-McConnell, R.H. 1975. Fish communities in tropical freshwaters: their distribution, ecology, and evolution. Longman Press, London. 337 pp.
- Lundberg, J.G. 1993. African-American freshwater fish clades and continental drift: problems with a paradigm. pp. 157–199. *In:* P. Goldblatt (ed.) The Biotic Relationships Between Africa and South America, Yale University Press, New Haven.
- Mago-Leccia, F. 1994. Electric fishes of the continental waters of America. Fundacion para el Desarrollo de las Ciencias Fisicas, Matematicas y Naturales, Caracas. 225 pp.
- Marrero, C. 1987. Notas preliminares acerca de la historia natural de los peces del bajo llano. I. Comparación de los hábitos alimentarios de tres especies de Gymnotiformes del Río Apure (Edo. Apure) Venezuela. Rev. Hidrobiol. Trop. 20: 57–63.
- Marrero, C. & D.C. Taphorn. 1991. Notas sobre la historia natural y la distribución de los peces Gymnotiformes en la cuenca del Río Apure y otros rios de la Orinoquia. Biollania 8: 123–142.
- Marrero, C. & K.O. Winemiller. 1993. Tube-snouted gymnotiform and mormyriform fishes: convergence of a specialized foraging mode in teleosts. Env. Biol. Fish. 38: 299–309.
- Petr, T. 1968. Distribution, abundance and food of commercial fish in the Black Volta and the Volta man-made lake in Ghana during its first period of filling (1964–1966). I. Mormyridae. Hydrobiol. 32: 417–448.
- Roberts, T.R. 1972. Ecology of fishes in the Amazon and Congo basins. Bull. Mus. Comp. Zool. 143: 117–147.
- Van Der Bank, F.H. & B. Kramer. 1996. Phylogenetic relationships between eight African species of mormyriform fish (Teleostei, Osteichthyes): resolution of a cryptic species, and reinstatement of *Cyphomyrus* Myers, 1960. Biochem. Syst. Evol. (in press).
- Winemiller, K.O. 1989. Ontogenetic diet shifts and resource partitioning among piscivorous fishes in the Venezuelan llanos. Env. Biol. Fish. 26: 177–199.
- Winemiller, K.O. 1990. Spatial and temporal variation in tropical fish trophic networks. Ecol. Monogr. 60: 331–367.

- Winemiller, K.O. 1991a. Comparative ecology of *Serranochromis* species (Teleostei: Cichlidae) in the Upper Zambezi River. J. Fish Biol. 39: 617–639.
- Winemiller, K.O. 1991b. Ecomorphological diversification of freshwater fish assemblages from five biotic regions. Ecol. Monogr. 61: 343–365.
- Winemiller, K.O. 1996. Dynamic diversity in fish assemblages of
- tropical rivers. pp. 99–121. *In:* M.L. Cody & J.A. Smallwood (ed.) Long-Term Studies of Vertebrate Communities, Academic Press, Orlando.
- Winemiller, K.O., L.C. Kelso-Winemiller & A.L. Brenkert. 1995. Ecomorphological diversification and convergence in fluvial cichlid fishes. Env. Biol. Fish. 44: 235–261.