

Running head: *Body size and trophic position*

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Body Size and Trophic Position in a Diverse Tropical Food Web

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*Abstract.* We use stomach contents and stable isotope ratios of predatory fishes, collected over a ten year time span from a species-rich river in Venezuela, to examine potential body size-trophic position relationships. Mean body size of predator taxa and their prey (determined by stomach content analyses) were significantly correlated, but trophic position of predators (estimated by stable isotope ratios) was not correlated with body size. This reflects no apparent relationship between body size and trophic position among prey taxa. Primary consumer taxa (algivore/detritivores) in this system are characterized by diverse size and morphology, and thus predatory fish of all body sizes and feeding strategies are able to exploit taxa feeding low in the food web. Regardless of relative body size, predators exploit short, productive food chains. For any given food chain within a complex web where predators are larger than their prey, trophic position and body size are necessarily correlated. But in diverse food webs characterized by a broad range of primary consumer body size, apparently there is no relationship between trophic position and body size across all taxa in the web.

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*Keywords:* diet analyses, detritivore, diversity, food-chain length, fish, morphology, Neotropics, predation, prey vulnerability, river, stable isotope ratios, trophic level.

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## INTRODUCTION

It long has been acknowledged that body size is central to the structure and function of food webs (Elton 1927), as body size is a fundamental determinant of energy flow, species diversity, and population densities (Pimm 1982, Peters 1983, Hairston and  
5 Hairston, Elser et al. 1996). Size-based descriptions of trophic structure have been used increasingly over the last twenty years to provide generalizations regarding food web properties (Warren and Lawton 1987, Cohen et al. 1993, Memmott et al. 2000, Williams and Martinez 2000, Kerr and Dickie 2001, Leaper and Huxham 2002, Cohen et al. 2003). Body size analyses may be especially useful in species-rich, reticulate food webs in  
10 which there may be thousands of individual feeding links and complex trophic pathways (Winemiller 1990, Polis and Strong 1996). Some of the most threatened ecosystems on the planet (e.g., rain forests, coral reefs, freshwater lotic systems) are characterized by extremely complex food webs, and body size analyses may provide insights necessary to assess community- and ecosystem-level responses to human-driven perturbations.

15 Predators are typically larger than their prey (but for exceptions see Leaper and Huxham 2002), and thus trophic position (i.e., the average level, relative to primary producers, at which an organism feeds) often increases with body-size within a given food web (Cohen et al. 1993, France et al. 1998, Jennings et al. 2001, Woodward and Hildrew 2002, Jennings and Mackinson 2003). The size-structured nature of food webs  
20 may allow specific predictions regarding ecological responses to perturbations. For example, in marine systems, commercial fisheries drive an overall reduction in food chain length because the large individuals (i.e., those targeted in most fisheries) are positioned high in the food web (Rice and Gislason 1996, Pauly et al. 1998). Examining

the body size-trophic position relationship in other systems should reveal whether a positive correlation between the variables is an intrinsic property of food webs that transcends different underlying web structures (e.g., ecosystems dominated by simple linear food chains vs. those with reticulate food webs).

5            In the present study, we examined the relationship between trophic position and body size for fish in the species-rich Cinaruco River (Venezuela). Specifically we asked: (1) Are predator and prey body size related?, and (2) Does trophic position increase with increasing predator body size? Results are evaluated with respect to specific characteristics of the complex food web, and extended to assess how the food  
10 web may be affected by commercial net fishing in the river.

## METHODS

The Cinaruco River is a tributary of the Orinoco River that drains the llanos (savanna) of southern Venezuela (6° 32' N, 67° 24' W). The Cinaruco is an oligotrophic,  
15 moderate-blackwater floodplain river with high levels of dissolved organic carbon and low pH. Hydrology is strongly seasonal, with water levels fluctuating more than 5 m annually (Arrington and Winemiller 2003). Maximum width of the main channel during the dry season is 50-200 m. The river supports a diverse fish community (>280 species), with taxa representing a wide range of ecological attributes and life history strategies  
20 (Jepsen 1997, Winemiller et al. 1997, Arrington and Winemiller 2003, Hoeninghaus et al. 2003, Layman and Winemiller 2004, Layman et al. 2005).

From 1993-2003, predatory fishes were collected with gill nets, cast nets, and by hook and line using artificial lures, and data from all years are pooled for these analyses.

Pooling data across years enabled us to examine general trends that transcend potential annual variations in predator-prey dynamics, thereby providing a general model of food web structure in the system. Importantly, the Cinaruco River has a highly predictable annual hydrology, with distinct wet and dry seasons, in which ecological interactions are set (Arrington and Winemiller 2003, Layman and Winemiller 2004). General trends in predator-prey dynamics are consistent across years, despite inter-annual variation in the food web (due to extent of flooding during the wet season, varying levels of fauna recruitment, etc.). For example, Jepsen et al. (1997) reported that the detritivore *Semaprochilodis kneri* made up almost 50% of the diet of the abundant piscivore *Cichla temensis* during the falling water season, but was absent from diets in the peak dry season (April). Stomach content analyses in subsequent falling water periods (1999, 2001, 2003, 2004) have revealed strikingly consistent results with *S. kneri* making up ~50% of *Cichla* diets; additional stomach content analyses in the peak dry season (April of 2001, 2002, 2003, and 2004) have not revealed a single *S. kneri* in piscivore stomachs (Layman et al., unpublished manuscript). Data reported here are based on large sample sizes, taken throughout the annual hydrological cycle, from all river habitats; no taxa analyzed for stable isotope ratios or stomach contents were collected only in a given season or habitat. Thus, emergent food web patterns should be robust in relation to overall temporal and spatial variation within the web.

20           Contents from *Cichla* spp. and *Pseudoplatystoma fasciatum* stomachs were removed by pressing down the posterior region of the tongue and applying pressure on the fish's stomach while holding the fish in a head-down position (Layman and Winemiller 2004). All other species were euthanized, and stomachs removed for

examination. Stomach contents were quantified volumetrically in the field using graduated cylinders. Most of the identifiable prey items were fishes (Table 1), so we restrict size-based analysis of prey to fish taxa. Often prey items were partially digested and could only be identified to genus, particularly among closely related and morphologically similar taxa. Thus, fishes identified in stomach content analyses were lumped at the generic level, and only those identified to genus were included in prey trophic position estimates. Body mass and length produce similar predator-prey size-based patterns (Jennings et al. 2001, Cohen et al. 2003), and we present size data based on the latter. All predator and prey individuals were measured (standard length, SL,  $\pm 1.0$  mm) in the field.

Collection, preservation, and preparation of tissues for stable isotope analyses follow Arrington and Winemiller (2002), and estimation of trophic position follows Layman et al. (2005). Primary consumers provide the best baseline for estimation of trophic position using  $\delta^{15}\text{N}$  values, because they integrate temporal and spatial variation in isotopic signatures of basal resources (Post 2002). We identified two fish species, *Semaprochilodus kneri* (Characiformes: Prochilodontidae) and *Metynnis hypsauchen* (Characiformes: Characidae) that integrate the dominant source pools of primary production in the system: (1) *S. kneri* for autochthonous algal/detrital resources, and (2) *M. hypsauchen* for allochthonous  $\text{C}_3$  plant material. We used a standard 2-source mixing model (Post 2002) to estimate trophic position of predators using these baseline taxa. Following the meta-analysis of Vanderklift and Ponsard (2003) we employ a mean  $\delta^{15}\text{N}$  enrichment of 2.54 ‰ to estimate trophic position. This is a lower enrichment value than suggested by Post (2002), and thus our estimates of predator trophic position should

represent upper estimates. Samples were not available for some rare taxa, and these taxa are not included in trophic position analyses (see Table 1).

## RESULTS

5 We analyzed 5,439 stomachs of 31 predator species representing 8 fish families (Table 1). Mean predator size ranged from  $69.2 \pm 5.9$  to  $458.2 \pm 89.7$  mm (mean  $\pm$  1 SD) for individual species. Identifiable prey in stomachs included individuals from 35 genera of fish. Measurable prey individuals varied from 5 to 225 mm. There was no correlation between mean predator size and the total volumetric proportion of stomach contents  
10 (summed across all individuals of that species) identifiable as fish remains (Table 1,  $F_{1,30} = 0.13$ ,  $P = 0.72$ ,  $R^2 = 0.005$ ). The number of predator species in which each prey genus was identified (an estimate of prey “vulnerability”, see Schoener 1989) decreased with increasing mean prey size (Fig. 1,  $F_{1,33} = 10.5$ ,  $P = 0.003$ ,  $R^2 = 0.24$ ), based on measurable prey recovered in stomachs. Eighteen predator species had multiple  
15 identifiable and measurable fish in their stomachs and were included in size-based analyses. The mean size of prey recovered from each predatory species was significantly correlated to the mean size of the predator (Fig. 2A,  $F_{1,17} = 19.4$ ,  $P < 0.001$ ,  $R^2 = 0.55$ ).

Predator trophic position, estimated using stable isotopes, was independent of predator size (Fig. 2B,  $F_{1,20} = 0.19$ ,  $P = 0.68$ ,  $R^2 = 0.001$ , total isotope samples analyzed =  
20 403). To examine this pattern further, we used stable isotope ratios to estimate mean trophic position for each of the prey genera that were identified in piscivore stomachs. Taxa analyzed for isotopes were collected independently (i.e., individuals analyzed were not from predator stomachs), and typically were in the same size range (mean  $\pm$  1 SD) as

individuals recovered from stomachs. Mean body size and estimated trophic position were not correlated for prey genera (Fig. 2C,  $F_{1,30} = 0.006$ ,  $P = 0.94$ ,  $R^2 = 0.0002$ ; total isotope samples analyzed = 617). For example, a genus of detritivorous fishes (*Curimata* spp., estimated trophic position =  $2.08 \pm 0.38$ ) had the third largest mean body size in piscivore stomachs (150 mm), whereas the second smallest prey genera (*Apistogramma* sp., 15 mm) had the second highest estimated trophic position (3.5).

## DISCUSSION

Although individual food chains within the Cinaruco food web were size-structured, i.e., prey selection was influenced by size-based morphological constraints (Figures 1, 2A), there was no relationship between body size and trophic position across the range of predator body sizes examined. In tropical rivers, primary consumers (i.e., herbivores and algivore/detritivores) typically are diverse in terms of size, morphology, and habitat affinity. In the Cinaruco River, primary consumer fishes exhibit a range of body sizes from some of the smallest (e.g., loricariid catfishes, adults of various species <1g) to among the largest (*Semaprochilodus laticeps*, adults >5kg) fish species in the river, a range of five orders of magnitude. This diversity is a function of far greater niche diversification in tropical than temperate freshwater fishes (Goulding 1980, Lowe-McConnell 1987, Winemiller 1990, Winemiller 1991, Flecker 1992). For example, many niches exploited by insects in temperate lentic waters are filled by fishes in the tropics. Larger scope for growth in fishes (relative to insects) allows for utilization of a broad range of primary resources (e.g. nuts/seeds and different components of benthic flora),

and the greater motility of fishes allows for exploitation of resources over large spatial scales (Winemiller 1990, Flecker 1992).

Piscivore species exploit prey individuals throughout the range in primary consumer body size in the Cinaruco River (Jepsen et al. 1997, Winemiller et al. 1997, 5 Layman et al. 2005). Piscivores of all body sizes are thus able to maximize profitability (sensu Scharf et al. 2002) both by: (1) exploiting short, productive, food chains and (2) feeding on optimally-sized prey. This appears to be a principal reason why tropical floodplain rivers support such high levels of secondary production (Lewis et al. 2001, Layman et al. 2005). Variation in primary consumer body size, morphology, habitat 10 affinity, and behavior affords the opportunity for the suite of predators, characterized by diverse feeding strategies, to occupy low trophic positions (i.e., at or just above trophic position 3, when 1 represents primary producers). And since primary consumers range from among the smallest to largest individuals in the community, all predators are able to exploit prey at energetically-optimal sizes.

15 Our analyses provide an example of how complex food webs may obscure patterns apparent in simple food chains. In linear food chains with a single basal resource and a single predator for each prey, predators are necessarily larger than their prey (excluding parasites and other exceptional cases such as pack hunters, see Leaper and Huxham 2002). In more complex webs, each individual food chain from primary 20 producers to secondary consumers is also size-structured. But if there is a substantial range in prey body size, especially among primary consumer taxa, there may not be a positive relationship between body size and trophic position across all taxa in the web, as large predators may target relatively large primary consumers. The exact nature of the

relationship between consumer body size and trophic position also will vary as a function of predator-prey body size ratios and minimum body size (D. M. Post, *unpublished manuscript*), but, in general, webs with a wide range of primary consumer body sizes, and piscivores that exploit a wide range of prey body sizes, will result in the lack of a strong relationship between body size and trophic position.

Food web structure is a primary determinant of ecosystem responses to perturbations. For example, fishery exploitation has been shown to reduce food chain length in marine fish communities, i.e., “fishing down the food web” (Pauly et al. 1998), because of the size-structured food web in which the largest fishes are positioned at the highest trophic levels. Commercial netters in the Cinaruco target large fishes, including both piscivores and algivore-detritivores (Layman and Winemiller 2004, Layman et al. 2004). Removal of large piscivorous fishes will not decrease mean food chain length of the system, because smaller predators (that are not removed by netters) feed at a similar trophic position as the large species. Removal of abundant, large, algivore/detritivores (e.g., *S. kneri*), however, may result in a net increase in food chain length, as generalist predators shift their diets and consume taxa that occupy higher trophic positions (e.g., the small-bodied predators). Food web structure thus may be affected directly by removal of species, as well as indirectly by an overall decrease in secondary productivity as predators shift to feeding at trophic levels further removed from primary producers (Elton 1927). The relationship among body size and trophic position in complex food webs warrants further attention, especially in species-rich systems where human alterations to food web structure and ecosystem function are great and on-going.

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**Table 1.** Species for which stomach content analyses were conducted. Data in the “SL” column are mean standard lengths of individuals for which stomach contents were analyzed, “N” represents the total number of stomachs examined, and “% Fish” the volumetric proportion of stomach contents identifiable as fish remains. An X in the “Measurable Prey” column indicates species for which >1 measurable prey individuals were recovered in stomach content analyses (depicted in Fig. 2A), and an X in the “Isotope Analyses” column indicates piscivore species for which stable isotope analyses were conducted (depicted in Fig. 2B).

Species	SL (mm, Mean $\pm$ 1 SD)	N	% Fish	Measurable Prey	Isotope Analyses
<i>Pseudoplatystoma fasciatum</i>	458.2 ( $\pm$ 89.7)	77	98.1	X	X
<i>Platynemichthys notatus</i>	428.2 ( $\pm$ 74.5)	30	100	X	
<i>Raphiodon vulpinus</i>	397.5 ( $\pm$ 21.2)	11	40		X
<i>Pseudoplatystoma tigrinum</i>	382.7 ( $\pm$ 100.0)	3	100		
<i>Boulengerella cuvieri</i>	361.5 ( $\pm$ 59.1)	292	100	X	X
<i>Pellona castelnaeana</i>	356.6 ( $\pm$ 86.5)	52	61.9	X	X
<i>Cichla temensis</i>	356.3 ( $\pm$ 87.1)	1365	100	X	X
<i>Boulengerella lucius</i>	343.1 ( $\pm$ 50.5)	411	100	X	X
<i>Ageniosis brevifilis</i>	340.5 ( $\pm$ 97.9)	4	33.3		X
<i>Cichla intermedia</i>	333.6 ( $\pm$ 50.9)	307	97.6	X	X
<i>Leiarius marmoratus</i>	312.2 ( $\pm$ 46.8)	5	100		
<i>Plagioscion squamosissimus</i>	310.2 ( $\pm$ 53.1)	115	94.7	X	X
<i>Hydrolycus armatus</i>	304.9 ( $\pm$ 64.3)	510	100	X	X
<i>Hoplias malabaricus</i>	302.0 ( $\pm$ 56.9)	39	100	X	X
<i>Cichla orinocensis</i>	301.5 ( $\pm$ 49.7)	755	99.8	X	X
<i>Brycon</i> sp. A	297.5 ( $\pm$ 116.7)	4	16.7	X	X
<i>Boulengerella maculata</i>	283.7 ( $\pm$ 36.0)	22	100	X	X
<i>Serrasalmus manuelei</i>	236.8 ( $\pm$ 65.2)	597	93.1		X
<i>Crenicichla</i> af. <i>lugubris</i>	228.6 ( $\pm$ 31.8)	24	90.5		X
<i>Serrasalmus rhombeus</i>	217.5 ( $\pm$ 37.6)	67	91.8		X
<i>Brycon falcatus</i>	210 ( $\pm$ 34.5)	11	80		X
<i>Pygocentris caribe</i>	193.1 ( $\pm$ 36.4)	17	91.6		
<i>Cynodon gibbus</i>	191.2 ( $\pm$ 34.7)	44	50		
<i>Acestrorhynchus falcirostris</i>	189.2 ( $\pm$ 73.2)	12	100	X	
<i>Agoniatas anchovia</i>	170.0 ( $\pm$ 17.7)	14	100	X	X
<i>Serrasalmus medinai</i>	161.7 ( $\pm$ 33.6)	16	38.3		
<i>Serrasalmus altuvei</i>	161.0 ( $\pm$ 35.4)	29	84.6		
<i>Pristobrycon striolatus</i>	145.9 ( $\pm$ 26.6)	58	61.6		X
<i>Acestrorhynchus microlepis</i>	107.9 ( $\pm$ 19)	464	96.9	X	X
<i>Acestrorhynchus grandoculis</i>	90.0 ( $\pm$ 8.9)	19	100	X	X
<i>Acestrorhynchus minimus</i>	69.2 ( $\pm$ 5.9)	65	100	X	

**FIGURE LEGENDS**

**Figure 1** Relationship between mean prey taxa size (SL of prey recovered from predator stomachs) and the number of predator species in which each prey taxa (genus) was identified.

**Figure 2** Relationship between (A) mean predator and prey body size (SL), (B) estimated mean trophic position of predators (based on stable isotope analysis) and body size, and (C) estimated trophic position of prey and their body size. In (A) and (B) each data point represents one predator species, and in (C) each point represents one prey genus identified in stomach content analyses. In (A), mean prey and predator sizes are plotted (instead of every individual value) to provide for the most direct comparison with the mean data presented in (B) and (C). Error bars represent  $\pm 1$  SD.

Figure 1

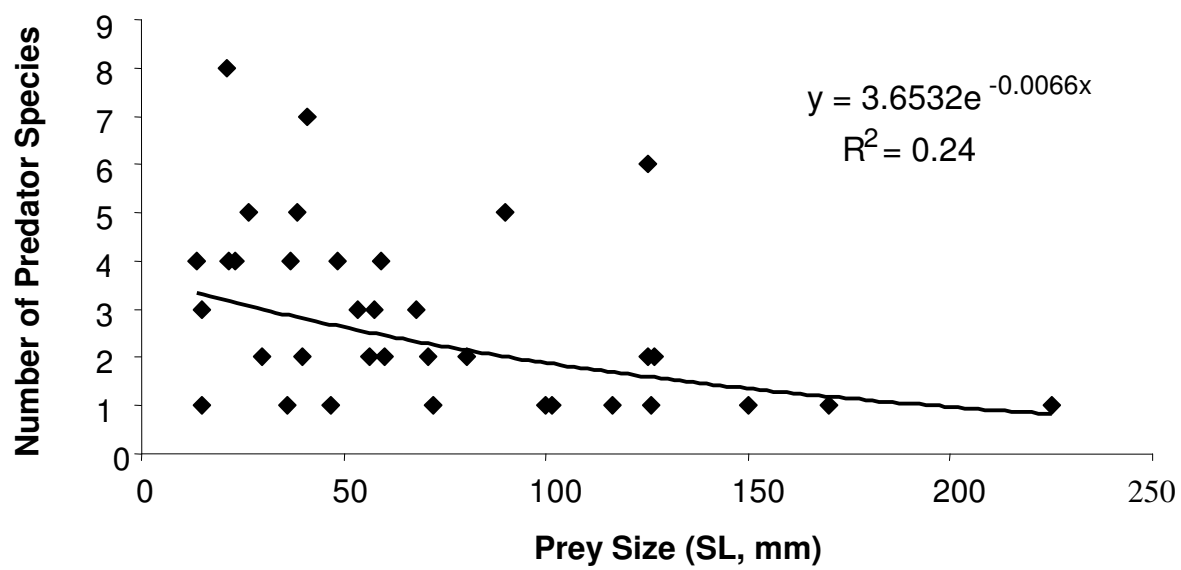


Figure 2

