

INTRA-ANNUAL VARIATION IN FISH COMMUNITIES AND HABITAT ASSOCIATIONS IN A CHIHUAHUA DESERT REACH OF THE RIO GRANDE/RIO BRAVO DEL NORTE

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ABSTRACT.—Anthropogenic alterations to lotic systems are often implicated in global declines among stream-dwelling fishes. A primary step toward mediating fish declines is improving our understanding of species–environment relationships; unfortunately, such information is limited. The goal of this study was to assess relationships between fish communities and environmental variables in a relatively intact portion of the Rio Grande/Rio Bravo del Norte in the southwestern United States and northern Mexico. We quantified the relationship between riverine habitats and local fish communities at 7 sites in the Big Bend reach of the Rio Grande/Rio Bravo del Norte during 2006. Monthly collections yielded 10,565 individuals representing 20 species, including 6 species listed as threatened or otherwise at risk. Species richness varied among sites (range 13–19) as did diversity (1 – D: 0.49–0.72) and density (16–45 fish · 100 m⁻²). Fish community composition indicated spatial and temporal variations, but habitat characteristics indicated more variation among sampling sites than among months. Spatial variation in community structure correlated with site-specific habitat characteristics, and most threatened or at risk species were associated with run or riffle geomorphic units containing higher current velocities and gravel to cobble substrates. Temporal variation in community structure correlated with fall monsoonal flooding, during which composition of dominant species shifted from the habitat generalist red shiner (*Cyprinella lutrensis*) to the imperiled, regionally endemic Tamaulipas shiner (*Notropis braytoni*). Results from this study suggest that large flood pulses and maintenance of habitat heterogeneity are necessary for the persistence of both declining and intact local fish communities in the Rio Grande/Rio Bravo del Norte.

RESUMEN.—Las alteraciones antropogénicas a los sistemas lóticos están con frecuencia implicadas en las declinaciones globales entre peces de arroyos. Un paso primordial para mediar la disminución de los peces, es mejorar nuestro entendimiento de las relaciones entre las especies y su ambiente; desafortunadamente, dicha información es limitada. El objetivo de este estudio fue evaluar las relaciones entre las comunidades de peces y las variables ambientales en una parte relativamente intacta del Río Grande/Río Bravo del Norte al suroeste de los Estados Unidos y al norte de México. Cuantificamos la relación entre los hábitats ribereños y las comunidades locales de peces en 7 sitios del tramo Big Bend del Río Grande/Río Bravo del Norte durante el 2006. Las colecciones mensuales aportaron 10,565 individuos representando a 20 especies, incluyendo a 6 especies listadas como amenazadas en riesgo. La riqueza de especies varió entre sitios (con un rango de 13–19) al igual que la diversidad (1 – D: 0.49–0.72) y la densidad (16–45 peces · 100 m⁻²). La composición de la comunidad de peces indicó variaciones espaciales y temporales, pero las características del hábitat indicaron más variaciones entre los sitios de muestreo que entre los meses. La variación espacial en la estructura de la comunidad estuvo correlacionada con las características del hábitat específicas al sitio, y la mayoría de las especies amenazadas o en riesgo estuvieron asociadas con unidades geomórficas de correderas o rápidos con corrientes de velocidades mayores y sustratos de grava o guijón. La variación temporal en la estructura de la comunidad estuvo correlacionada con las inundaciones otoñales del monzón, durante las cuales la especie dominante cambió de la generalista carpita rojiazul (*Cyprinella lutrensis*), a la especie en peligro y regionalmente endémica sardinita pechona de Tamaulipas (*Notropis braytoni*). Los resultados de este estudio sugieren que los pulsos de inundación grandes y el mantenimiento de la heterogeneidad del hábitat son necesarios para la sobrevivencia de las comunidades locales de peces, tanto en declinación como las intactas, en el río Bravo del Norte.

Deterministic fitness, stochasticity, speciation, and dispersal regulate biotic communities within streams at local, regional, and continental scales (Vellend 2010). Anthropogenic modifications to streams often disrupt these regulatory processes and contribute to the current state of crisis facing freshwater biodiversity

(Dudgeon et al. 2006). Reductions in the abundance and distribution of freshwater fishes are particularly notable (Helfman 2007). In North America, nearly 40% of freshwater and diadromous fishes are imperiled (i.e., at risk of population decline or extinction; Jelks et al. 2008); in the southwestern USA, >50% of stream

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fishes are considered extirpated (Hughes et al. 2005); and in the Rio Grande/Rio Bravo del Norte Basin (hereafter, Rio Grande), 43% of native species are missing from all or portions of their historical range (Calamusso et al. 2005). Declines in abundance and distribution of native fishes are often associated with human-mediated changes (e.g., dam construction, water diversion) to naturally occurring fluvial conditions (Holden 1979, Winston et al. 1991, Hoagstrom et al. 2008), but specific mechanisms and causal factors of species declines remain largely unknown. Conservation approaches in which entire communities rather than specific species are considered hold the greatest potential for identifying and subsequently mediating factors contributing to the decline and extirpation of multiple organisms (Scott and Helfman 2001, Bunn and Arthington 2002).

Within the Rio Grande drainage of the southwestern United States and northern Mexico, numerous historically widespread and abundant native fishes are now less common and occupy a fraction of their historical ranges (Hubbs et al. 1977, Bestgen and Platania 1991, Edwards and Contreras-Balderas 1991, Edwards et al. 2002, Calamusso et al. 2005, Hoagstrom et al. 2008). For example, Rio Grande silvery minnow (*Hybognathus amarus*), speckled chub (*Macrhybopsis aestivalis*), Rio Grande shiner (*Notropis jemezianus*), and blue sucker (*Cyctepus elongatus*) are extirpated from numerous reaches (Anderson et al. 1995, Platania and Altenbach 1998); and phantom shiner (*Notropis orca*) and Rio Grande bluntnose shiner (*Notropis simus simus*) are extinct (Bestgen and Platania 1990, Hubbs et al. 2008). Declines in abundance and distribution of some Rio Grande fishes coincided with increases in habitat generalist and lentic-adapted species such as red shiner (*Cyprinella lutrensis*), fathead minnow (*Pimephales promelas*), gizzard shad (*Dorosoma cepedianum*), and western mosquito fish (*Gambusia affinis*) after stream alteration throughout the Rio Grande watershed (Calamusso et al. 2005). However, despite well-documented changes in species abundances and distributions, mechanisms contributing to the decrease of some native fishes and increase of other native or nonnative fishes remain unclear. Additional research is needed to establish the relationship between the ecology of local fish communities and the

anthropogenic alterations occurring throughout the basin.

Ichthyofaunal changes in the Rio Grande are associated with anthropogenic activities that modify instream habitat and cause concern for the long-term persistence of endemic taxa. Construction of reservoirs, channel straightening, water diversions for agricultural purposes, reduced stream-flow quantity, degraded water quality, and introduction of non-native species are common in the Rio Grande (Edwards et al. 2002, Calamusso et al. 2005). In some reaches, the mainstem river is completely dewatered because of agricultural and municipal withdrawals (Calamusso et al. 2005); and in other reaches, poor water quality and chemical pollution prompt advisories against water contact and fish consumption (Owens and Niemeyer 2006). Stream fragmentation is thought to be related to the extirpation of fishes belonging to the pelagic-spawning reproductive guild, and only a few remaining populations persist in relatively long fragments (>100 km in length; Dudley and Platania 2007). Such extensive alterations call into question the feasibility of conserving or restoring intact fish communities within highly modified mainstem reaches (Calamusso et al. 2005). Less impacted river reaches, including the Big Bend National Park reach of the Rio Grande, provide the opportunity to assess the ecology of relatively intact local fish communities (Moring 2002) or, more specifically, to assess habitat associations and suitability for declining or imperiled species (Helfman 2007). Knowledge of habitat associations and suitability is a primary step toward conserving imperiled species (Bond and Lake 2003). This knowledge provides targets for habitat conservation and restoration (Lake et al. 2007) and is necessary for ecoregions such as the Rio Grande where empirical data for instream flow requirements are lacking (Sansom 2008).

The goal of this study was to quantify intra-annual variation in the habitat ecology of local fish communities in the Big Bend reach of the Rio Grande. We specifically sought to assess occurrence and habitat associations of small-bodied species that are extirpated from Rio Grande reaches separated from the Big Bend region by impoundments and that occur in portions of New Mexico and the Rio Grande terminus near the Gulf of Mexico (e.g., Gido et al. 1997). Historical and contemporary literature

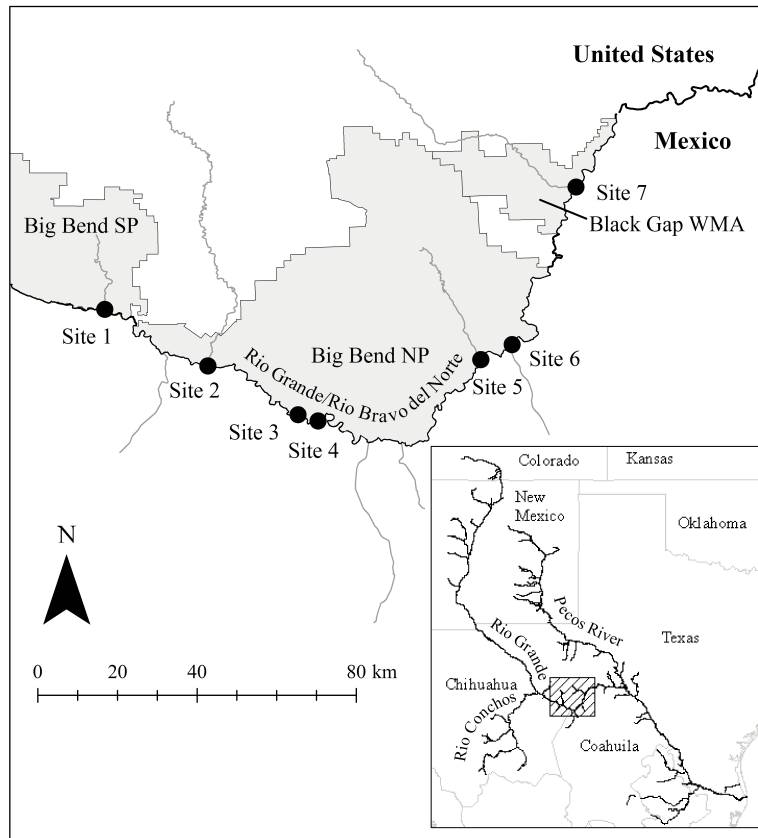


Fig. 1. Mainstem Rio Grande sites sampled monthly, January–December 2006. Site 1 was accessed through Big Bend State Park (SP), sites 2–6 through Big Bend National Park (NP), and Site 7 through Black Gap Wildlife Management Area (WMA).

suggests that many imperiled, small-bodied species inhabit shallow areas throughout a suite of geomorphic units in the Big Bend region (Hubbs et al. 1977, Moring 2002), so we targeted wadeable habitats accessible from within or near Big Bend National Park. Our objectives were to (1) quantify occurrence, abundance, and habitat associations among local communities, (2) assess local community and habitat change with respect to intra-annual variation in streamflow, and (3) describe relationships among abundant small-bodied taxa in terms of current velocity suitability and associated implications for community dominance.

METHODS

Study Area

The Big Bend reach of the Rio Grande generally refers to a section of the international

boundary from the confluence of the Rio Conchos (Chihuahua, Mexico) to the confluence of the Pecos River (Texas, USA). Aquatic and terrestrial environments within the reach are largely under the jurisdiction of various state and federal agencies, including Big Bend Ranch State Park (SP), Cañón de Santa Elena Protected Natural Area, Big Bend National Park (NP), Maderas del Carmen Protected Natural Area, Rio Grande Wild and Scenic River, and Black Gap Wildlife Management Area (WMA). Despite large-scale habitat degradation and homogenization throughout the basin, a diversity of habitats persist in the Big Bend reach, including multiple geomorphic units (e.g., riffles, runs, and pools), limited areas of braided channels, and substrates ranging from silt to boulders (Hubbs et al. 1977, Moring 2002). For this study, 7 sampling sites in a 200-km reach were selected because

of available habitat heterogeneity and accessibility (Fig. 1). Mainstem sites were located near the confluence of Contrabando Creek (Site 1; 29°16'44"N, 103°50'35"W), downstream from Santa Elena Canyon (Site 2; 29°09'52"N, 103°36'34"W), near the USGS gaging station (Site 3; 29°01'57"N, 103°23'15"W), near Johnson Ranch campground of Big Bend NP (Site 4; 29°01'17"N, 103°22'11"W), near the confluence of Tornillo Creek (Site 5; 29°10'41"N, 102°59'46"W), upstream from Boquillas Canyon (Site 6; 29°11'56"N, 102°55'02"W), and near the confluence of Maravillas Creek (Site 7; 29°33'45"N, 102°46'35"W). Study sites contained a mixture of wadeable geomorphic units characterized by a diversity of substrate types, depths up to 1.2 m, and current velocities up to $2 \text{ m} \cdot \text{s}^{-1}$. At all sites, riparian vegetation consisted of dense stands of exotic giant reed (*Arundo donax*) and saltcedar (*Tamarix* spp.; Moring 2002).

The Rio Grande flow regime in the Big Bend region is influenced by contributions from upstream portions of the Rio Grande proper and the largest tributary in the basin, the Rio Conchos. The Rio Conchos enters the Rio Grande near the city of Presidio, Texas, just upstream of the study area and now contributes >80% of water present in the Rio Grande downstream of the confluence (Kim et al. 2002). Patterns in flow variability from the Rio Conchos are driven by monsoonal weather patterns, with high flow events during fall months (August–November) and low base flows during remaining months of the year (Kim et al. 2002). Variability of flow contributions from the upper Rio Grande was historically driven by seasonal snowmelt patterns and was characterized by large flood pulses during spring months before regulation by reservoirs and subsequent storage and withdrawal of water upstream (Dean and Schmidt 2011). Consequently, characteristics of the contemporary flow regime of the Big Bend reach of the Rio Grande are dependent on contributions from the Rio Conchos (Dean and Schmidt 2011).

Sampling Methods

We sampled fishes and quantified wadeable habitats monthly from January through December 2006 at all sites, with the exception of sites 3 and 4, which were not accessible in July 2006. At each site, we established 4–20 transects. Transects were placed perpendicular

to the stream bank and spaced approximately one mean stream width apart (Simonson et al. 1994). Along each transect, we seined (seine dimensions: 3 m × 1.8 m; mesh size = 3.1 mm) discrete geomorphic units (run, riffle, pool, side channel, backwater, or eddy) with 5-m-long seine hauls in a downstream direction. In cases of fast runs and shallow riffles, 5-m kicks into blocking seines were used to sample fish communities. Along each transect, seine hauls were spaced approximately 10 m apart to avoid disturbing adjacent habitats, and a minimum of 20 seine hauls were conducted on each date (Matthews 1986a). Captured fish were identified to species when practical and released, except for those retained as voucher specimens. Within the area of each seine haul, current velocity ($\text{cm} \cdot \text{s}^{-1}$) and depth (cm) were measured from 2 random points, and substrate type at 10 random points was determined based on a modified Wentworth scale (Cummins 1962, McMahon et al. 1996).

Statistical Methods

Habitat characteristics among sites were described using principal component analysis (PCA; CANOCO version 4.5 © Micro Computer Power) to test the hypothesis that multiple habitat parameters differed through space and time (Ahmadi-Nedushan et al. 2006). For PCA, quantitative data (mean current velocity, mean depth, and percent substrate) were Z score transformed and qualitative data (geomorphic unit) were represented by dummy variables (Krebs 1999). Habitat similarities among sites and months were assessed by testing for differences among mean sampling scores along the principal components that described the most variation in the data (axes I and II) using single factor analyses of variance (ANOVA) and Fisher's test for least significant differences (LSD; $\alpha = 0.05$). Species–habitat associations were described using canonical correspondence analysis (ter Braak 1986). Relationships between environmental factors and species abundance and distribution were verified using an ordinal cross-validation technique following the methods of Hallgren et al. (1999). Relative importance of site, season, and environmental factors in determining species abundance and distribution were tested using variance-partitioning techniques in canonical correspondence analysis (Bean et al. 2007). For canonical correspondence analysis variance

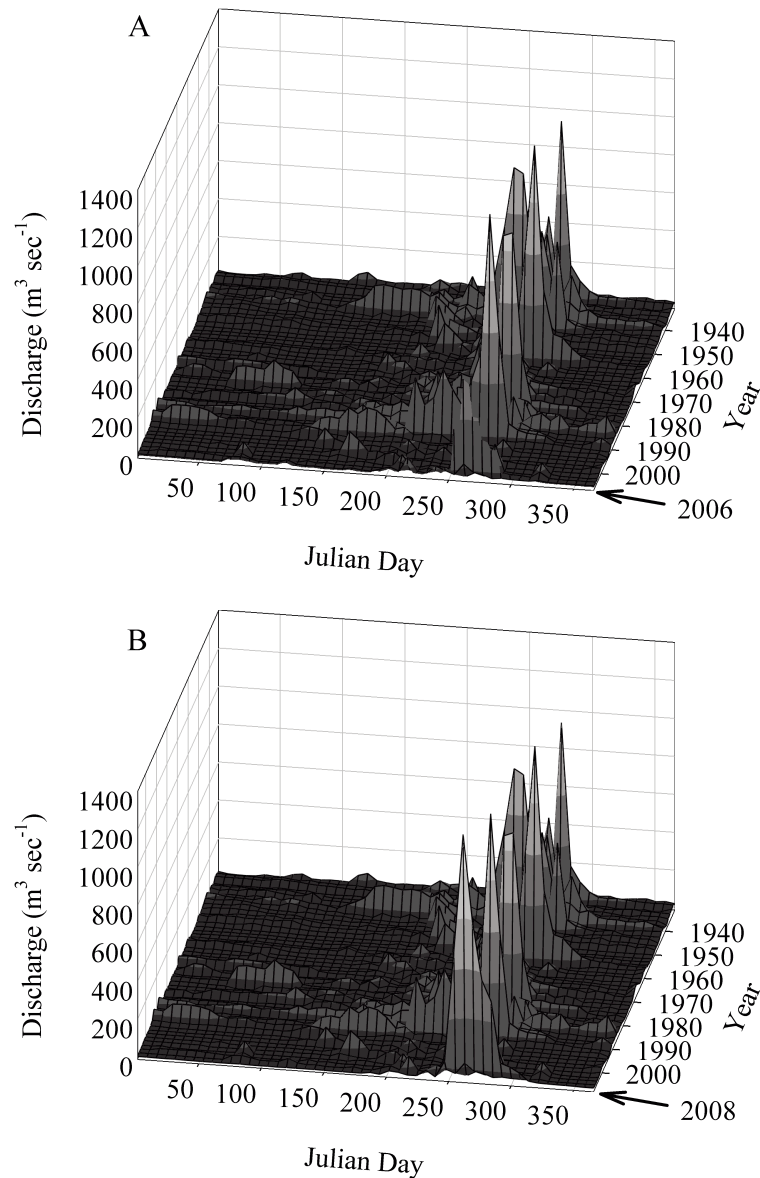


Fig. 2. Flow history for the Big Bend reach of the Rio Grande based on daily streamflows ($\text{m}^3 \cdot \text{s}^{-1}$) recorded at the International Boundary and Water Commission gage near Presidio, Texas, USA, and Ojinaga, Chihuahua, Mexico (gage #08-3742.00). The long-term pattern of monsoonal flooding is illustrated, including year of data collection (2006; A) and flooding that has occurred since (2008; B).

partitioning, we defined seasons according to monsoonal precipitation cycles in the arid southwestern USA and north central Mexico as pre-flood (January–July), flood (August–October), and post-flood (November–December; Kim et al. 2002; Fig. 2). Because some fishes are thought to be displaced or killed by high current velocities during flooding (Collins

et al. 1981, Harvey 1987, Minckley and Meffe 1987), we tested suitability of current velocities for 3 abundant small-bodied species (*C. lutrensis*, Tamaulipas shiner [*Notropis braytoni*], and *M. aestivalis*) using Kolmogorov–Smirnov (KS) tests to describe differences in expected and observed abundances among current velocities. Expected values for KS

TABLE 1. Species, canonical correspondence analysis (CCA) code, and total number and overall density (fish · 100 m⁻²) of fishes captured from the Rio Grande/Rio Bravo del Norte at 7 sites in the Big Bend National Park region. Species marked with asterisks are listed as imperiled or threatened by Hubbs et al. (2008). Two *Lepomis* species (*L. megalotis* and *L. macrochirus*) were collected (total $S = 20$), but individuals <12 mm were not identified to species.

Species	CCA code	Total	Site 1	Site 2	Site 3	Site 4	Site 5	Site 6	Site 7
<i>Lepisosteus osseus</i>	<i>Lep oss</i>	15		0.01	0.05	0.06	0.02	0.04	0.14
<i>Cyprinella lutrensis</i>	<i>Cyp lut</i>	4895	16.06	12.42	8.18	17.10	13.71	19.36	7.40
<i>Cyprinus carpio</i>	<i>Cyp car</i>	69	0.08	0.53	0.16	0.08	0.15	0.11	0.09
<i>Macrhybopsis aestivalis</i> *	<i>Mac aes</i>	333	0.44	1.71	1.17	1.28	0.67	0.84	0.07
<i>Notropis braytoni</i> *	<i>Not bra</i>	3695	4.76	7.18	3.05	5.66	21.78	20.73	7.47
<i>Notropis chíhuahua</i> *	<i>Not chi</i>	13		0.17		0.02			
<i>Notropis jemezianus</i> *	<i>Not jem</i>	5		0.04		0.02	0.02		
<i>Rhinichthys cataractae</i> *	<i>Rhi cat</i>	38		0.54					
<i>Carpionodes carpio</i>	<i>Car car</i>	433	0.42	2.73	0.23	1.81	1.63	0.33	0.21
<i>Ictiobus bubalus</i>	<i>Ict bub</i>	2				0.02		0.02	
<i>Cycleptus elongatus</i> *	<i>Cyc elo</i>	22	0.02	0.03			0.02	0.31	0.09
<i>Astyanax mexicanus</i>	<i>Ast mex</i>	141	0.13	0.04	0.18	0.02	0.91	0.53	1.05
<i>Ictalurus furcatus</i>	<i>Ict fur</i>	171	0.55	0.16	0.18	0.47	0.63	0.64	0.78
<i>Ictalurus punctatus</i>	<i>Ict pun</i>	249	0.40	0.43	0.47	0.34	1.87	0.77	0.44
<i>Pylodictis olivaris</i>	<i>Pyl oli</i>	31	0.02	0.14	0.21		0.02	0.13	0.11
<i>Fundulus zebrinus</i>	<i>Fun zeb</i>	54		0.54	0.08	0.04	0.17		0.02
<i>Gambusia affinis</i>	<i>Gam aff</i>	315	0.50	0.40	1.80	1.36	1.22	0.79	0.37
<i>Menidia beryllina</i>	<i>Men ber</i>	64		0.54	0.03	0.02	0.21	0.13	0.14
<i>Lepomis</i> spp.	<i>Lep spp</i>	20	0.02	0.04	0.13	0.12	0.02		0.09
Total number individuals		10,565	1225	1938	611	1441	2511	2033	806
Species richness (S)		20	12	18	14	16	16	14	15
Diversity (1-D)		0.66	0.49	0.72	0.68	0.6	0.64	0.6	0.67
Number of seine hauls		2393	349	467	256	338	389	303	291

tests were calculated as the total number of individuals collected among habitats if distribution was uniform, and observed values were calculated as the total number of individuals captured within current velocity frequency bins calculated using the Sturges (1926) equation (Williams and Bonner 2006). Although river carpsucker (*Carpionodes carpio*) was the third most abundant species, we excluded it from further testing because 99% of those captured were juveniles (<200 mm TL; Jester 1972, Ross 2001), opting instead to provide more information for a Rio Grande endemic (*M. aestivalis*).

Differences in fish community composition among sites and months were tested using analysis of similarity (ANOSIM). Bray-Curtis similarity matrices (Bray and Curtis 1957) created in Primer 6.1.6 were tested with ANOSIM ($\alpha = 0.05$; 9999 permutations) using permutations to assess average rank dissimilarity in fish density between sites and months (Clarke 1993, Bean et al. 2007). Abundances of individual species among sites and months were quantified by calculating density (fish · 100 m⁻²) for species collected within each seine haul. Before analysis, density data were fourth-root transformed to standardize contributions of high-

and low-abundance species (Clarke and Green 1988). Spatiotemporal variability in fish communities was quantified by plotting mean similarities among all pairwise site comparisons for each month of sampling (Gido et al. 1997), and differences were assessed using ANOVA and Fisher's LSD. Similarly, changes in proportional density (density of individual species / density of all species) summed across sites were plotted by month for the 3 most dominant small-bodied species and overlaid with discharge values to illustrate changes in dominance that corresponded with changes in discharge.

RESULTS

We collected a total of 10,565 fishes representing 20 species from 7 sites along the Big Bend reach of the Rio Grande during monthly sampling in 2006 (Table 1). Two *Lepomis* species (*Lepomis megalotis* and *Lepomis macrochirus*) were collected (total $N = 20$) but individuals <12 mm were not identified to species. Therefore we grouped larvae and adult individuals into *Lepomis* spp. Site 1 had the lowest species richness ($S = 12$) and diversity ($1 - D = 0.49$), whereas Site 2 had the highest species richness ($S = 18$) and diversity ($1 - D = 0.72$). We

TABLE 2. Habitat variable loadings, percent explained variance, and distribution of mean site and monthly scores along principal component (PC) I and II for habitat parameters measured in the Rio Grande/Rio Bravo del Norte. Standard deviations are in parentheses. Bolded values indicate parameters shown in Figure 3 and letters indicate statistical differences ($\alpha = 0.05$) among PC scores.

Parameter	PC I	PC II
Current velocity ($m \cdot s^{-1}$)	-1.163	-1.288
Depth (m)	-1.432	0.619
Run	-2.248	0.428
Riffle	1.396	-1.537
Side channel	0.929	0.561
Backwater	0.859	0.816
Pool	0.740	0.595
Eddy	0.499	0.212
Boulder (%)	0.278	-0.274
Cobble (%)	-0.021	-2.107
Gravel (%)	-0.889	0.730
Sand (%)	-0.458	0.747
Silt (%)	0.952	1.841
Variance explained (%)	15.5	13.8
Site 1	1.107 (1.51) a	-0.146 (1.60) c
Site 2	0.017 (1.53) b	0.101 (1.37) b
Site 3	-0.334 (1.20) d	-0.791 (0.87) d
Site 4	-0.423 (1.52) d	1.421 (0.93) a
Site 5	-0.009 (1.37) bc	-0.615 (1.20) d
Site 6	-0.345 (1.43) d	-0.054 (1.43) bc
Site 7	-0.199 (1.45) cd	-0.062 (1.32) bc
Jan	0.116 (0.40) a	-0.013 (0.43) ab
Feb	0.097 (0.37) a	-0.035 (0.36) ab
Mar	0.028 (0.32) ab	-0.084 (0.34) b
Apr	-0.005 (0.47) ab	0.032 (0.34) ab
May	0.066 (0.40) ab	0.030 (0.37) ab
Jun	0.035 (0.37) ab	-0.077 (0.41) ab
Jul	-0.069 (0.33) ab	0.004 (0.32) ab
Aug	-0.115 (0.34) ab	0.032 (0.29) ab
Sep	-0.166 (0.33) b	0.055 (0.36) a
Oct	-0.005 (0.46) ab	0.055 (0.40) a
Nov	-0.053 (0.39) ab	0.020 (0.37) ab
Dec	-0.060 (0.38) ab	0.003 (0.36) ab

attributed high richness and diversity at Site 2 to the occurrence of uncommon species such as Chihuahua shiner (*Notropis chihuahua*), *N. jemezianus*, longnose dace (*Rhinichthys cataractae*), and *C. elongatus*. Habitat generalist *C. lutrensis* was the most abundant fish (46% relative abundance) across sites and months and most abundant at sites 1–4 (45%–69% relative abundance). Endemic *N. braytoni* was second most abundant (35%) across sites and months and most abundant at sites 5–7 (40%–51% relative abundance). Excluding *N. braytoni*, imperiled fishes constituted 4% of the total fish community. Nonnative common carp (*Cyprinus carpio*), inland silverside (*Menidia beryllina*), and plains killifish (*Fundulus zebrinus*)

represented <1% of the total assemblage, and although exotic blue tilapia (*Oreochromis aureus*) were not collected, we observed them during a fish kill in December 2006 near Site 2.

Wadeable habitats within the study area consisted of backwater, eddy, pool, riffle, run, and side-channel geomorphic units. Within geomorphic units, substrate particle sizes ranged from silt (0.0038 mm) to boulder (>256 mm), depths ranged between 5 and 110 cm, and current velocities were 0–1.8 $m \cdot s^{-1}$. The first 2 principal components (PCs) of PCA explained 29.3% of variation in habitat parameters (Table 2). The remaining PCs explained little additional variation (PC III 9%; PC IV 8%) and were disregarded. The first principal component explained 15.5% of variation and represented a geomorphic unit, depth, and velocity gradient. Habitat parameters with large negative loadings along PC I were characterized by run geomorphic units (loading = -2.24), greater depths (-1.43), and greater current velocities (-1.16); whereas habitat parameters with large positive loadings were characterized by riffle geomorphic units (1.39), silt substrate (0.95), and side-channel geomorphic units (0.93). The second PC explained 13.8% of variation and represented a substrate, geomorphic unit, and velocity gradient. Habitat parameters with large negative loadings along PC II were characterized by cobble substrate (-2.11), riffle geomorphic units (-1.54), and greater current velocities (-1.29); whereas habitat parameters with large positive loadings were characterized by silt substrate (1.84) and backwater geomorphic units (0.82). Sites differed among habitat parameters along PC I ($F_{6,2386} = 45.0$, $P < 0.01$) and PC II ($F_{6,2386} = 101.7$, $P < 0.01$) and by month along PC I ($F_{11,2386} = 9.25$, $P < 0.01$) and PC II ($F_{11,2386} = 3.14$, $P < 0.01$). Site 1 consisted of riffle and side-channel geomorphic units; site 4 consisted primarily of backwater geomorphic units with silt substrate; sites 3 and 5 consisted primarily of riffle geomorphic units, cobble substrate, and higher current velocities; and sites 2, 6, and 7 had greatest habitat heterogeneity and consisted of all available geomorphic units and substrate sizes (Fig. 3A). Monthly variability in available habitat characteristics was less notable than variation among sites (Fig. 3B). However, March and June were characterized by riffle geomorphic units with cobble substrate and higher current velocities, whereas October and September

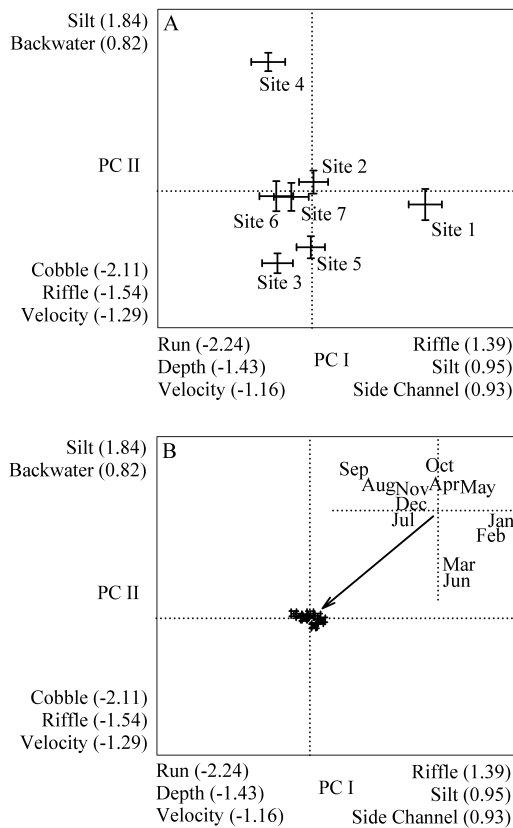


Fig. 3. Principal components analysis biplots for Rio Grande/Rio Bravo del Norte sites sampled monthly, January–December 2006: A, spatial variation among 7 sites; B, temporal variation among 12 months for habitat gradients along principal component (PC) I and II. Dotted lines represent the origin (panels are scaled similarly) and points represent mean (\pm SE) PC scores.

were predominantly characterized by run geomorphic units with greater depths and higher current velocities (see Table 2 for all pairwise comparisons).

We characterized species–habitat associations using canonical correspondence analysis. Physical habitat, season, and site collectively explained 6.2% of the total taxonomic variability within the sampled fish assemblage, partitioned among physical habitat (4%; $P < 0.01$), season (1.6%, $P < 0.01$), site (0.4%, $P < 0.01$), and 2- and 3-way interactions (0.2%). Ordinal cross-validation identified habitat parameters that explained significant ($P < 0.05$) portions of species distributions, including current velocity (14% of explained variation), flood season (8%), backwater geomorphic units (7%), depth (5%),

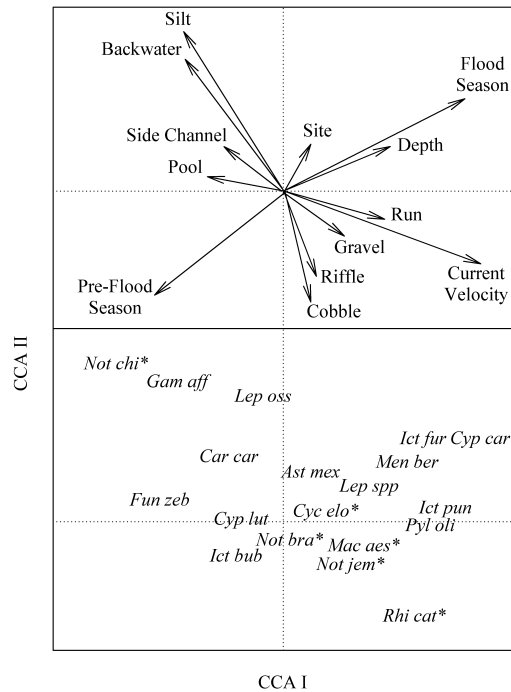


Fig. 4. Canonical correspondence analysis biplots illustrating species–habitat associations along canonical correspondence axis (CCA) I and II for fish and habitat data collected from 7 Rio Grande/Rio Bravo del Norte sites, January–December 2006. Asterisks denote imperiled species listed by Hubbs et al. (2008).

site (4%), and silt substrate (3%). Among these parameters, those with large negative loadings along canonical correspondence axis (CCA) I were pre-flood season (-0.45), silt substrate (-0.34), backwater geomorphic units (-0.32), and pool geomorphic units (-0.25). Parameters with large positive loadings were current velocity (0.69), depth (0.37), run geomorphic units (0.35), and flood season (Fig. 4). Habitat parameters with large negative loadings along CCA II were cobble (-0.49), pre-flood season (-0.47), riffle geomorphic units (-0.37), and current velocity (-0.32). Parameters with large positive loadings were silt substrate (0.69), backwater geomorphic units (0.59), and flood season (0.40). Species with strong associations with habitat parameters included *M. aestivalis*, *N. jemezianus*, and *R. cataractae*, which associated with riffle to run geomorphic units characterized by gravel substrate and greater current velocities. Blue catfish (*Ictalurus furcatus*), *Cyprinus carpio*, and *M. beryllina* indicated strong associations with flood season, and *G. affinis* indicated

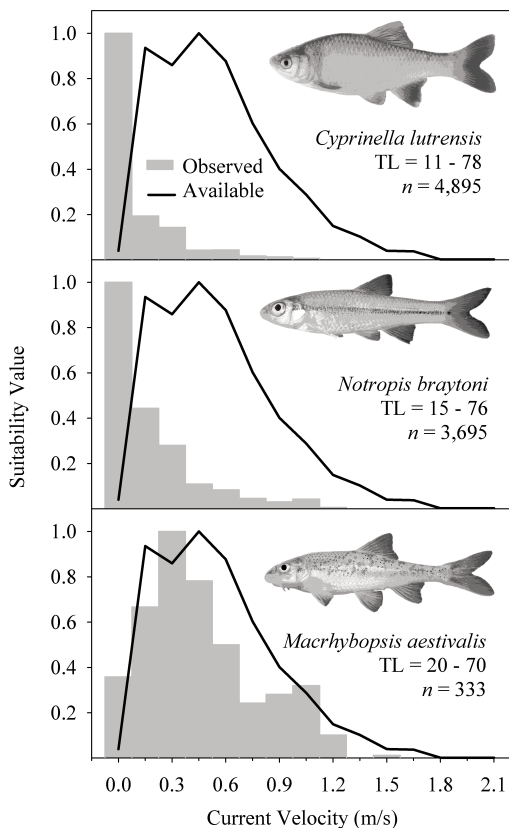


Fig. 5. Current velocity associations and suitability values for 3 dominant fish species collected from the Rio Grande/Rio Bravo del Norte, January–December 2006. Black lines represent distribution of available current velocities and gray bars represent distribution of observed associations by species. Ranges for total lengths (TL) of captured individuals are shown in mm; n is the frequency of occurrence.

strong associations with backwater geomorphic units containing silt substrate. Habitat associations were most general for *C. lutrensis* and *N. braytoni*, both of which occurred near the origin of CCA I and CCA II. The 3 most abundant small-bodied species collected during the study—*C. lutrensis*, *N. braytoni*, and *M. aestivalis*—partitioned along a current velocity gradient. *Cyprinella lutrensis* frequency distribution among current velocities differed from expected ($D = 0.76$, $P < 0.001$) and indicated association with slow current velocities; *N. braytoni* distribution differed from expected ($D = 0.62$, $P < 0.001$) and indicated association with increasingly intermediate velocities; and *M. aestivalis* distribution differed from expected ($D = 0.27$, $P < 0.001$) and indicated

association with relatively faster current velocities (Fig. 5).

Community composition differed spatially (among sites) and temporally (among months) over the course of the study. Analysis of similarity (ANOSIM) detected differences in community composition among sites (global $R = 0.01$, $P < 0.01$), and the magnitude of these differences varied temporally. Mean percent similarity among all possible pairwise site comparisons differed through time ($F_{11,229} = 40.37$, $P < 0.01$). During January and February, mean similarity among sites was 4.6%–4.8% but increased to 7.9% in March and increased again during April through July when similarity values ranged from 14.7% to 23.9% (Fig. 6A). Following large flood pulses in August, which marked the onset of the monsoonal flooding season, mean similarity among sites significantly decreased to 7.3%. Mean similarity values continued to decrease during September and October (4.4% and 4.0%, respectively) and November and December (2% and 1.4%, respectively). Higher percent similarity among sites during the months between April and July corresponded with increases in proportional density of *C. lutrensis* and decline of *N. braytoni*, while mean densities of *M. aestivalis* remained relatively stable (Fig. 6B). Monthly density among all species correlated most with changes in mean density of *C. lutrensis* ($r = 0.97$, $P < 0.01$) and to a lesser extent *N. braytoni* ($r = 0.84$, $P < 0.01$), but not with *M. aestivalis* ($r = 0.46$, $P = 0.13$), indicating dominance by *C. lutrensis* and *N. braytoni*. During all months from January through July, proportional density of *C. lutrensis* outranked *N. braytoni*; however, proportional density of *C. lutrensis* declined notably between July and August, after the onset of monsoonal flooding. Following the largest flood pulse observed during 2006 (September, $186 \text{ m}^3 \cdot \text{s}^{-1}$), proportional density of *N. braytoni* outranked *C. lutrensis* for the remainder of the study.

DISCUSSION

Fish communities at 7 sites in the Big Bend reach of the Rio Grande exhibited spatial and temporal variation in 2006. During this time, streamflows followed a consistent pattern of increased discharges in fall associated with monsoonal weather patterns in the region. Similar habitat types persisted through time despite

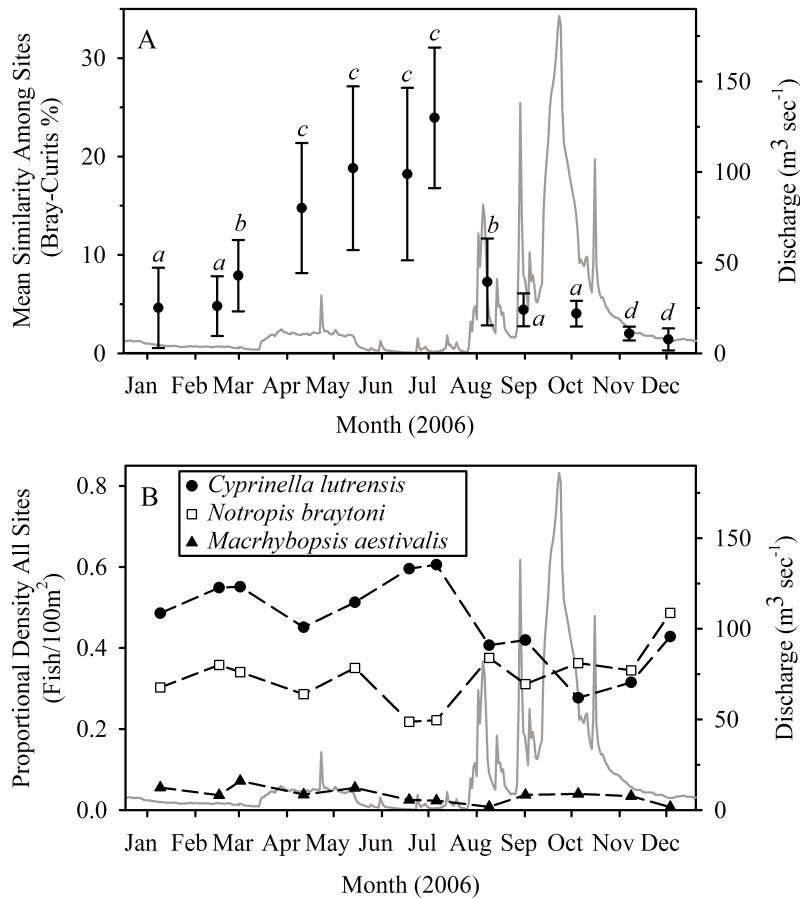


Fig. 6. Spatiotemporal variability in fish communities at 7 Rio Grande/Rio Bravo del Norte sites, January–December 2006: A, monthly mean (\pm SD) and Bray–Curtis percent similarity for all pairwise comparisons among sites (solid circles) and daily discharge ($\text{m}^3 \cdot \text{s}^{-1}$; gray line). Letters indicate significant ($\alpha = 0.05$) differences among means; B, monthly proportional density across all sites (fish \cdot 100 m^{-2}) for 3 dominant small-bodied species and daily discharge.

large changes in flow regime, as evidenced by greater spatial variation relative to temporal variation detected during PCA procedures. Consequently, temporal differences in community composition were likely influenced by flow regime, often regarded as the master variable in stream ecosystems (Poff and Ward 1989), whereas spatial differences were likely related to available habitats among sampling sites and species-specific associations with habitat components. These conclusions are supported by species-specific associations we detected in canonical correspondence analysis that illustrated seasonal and spatial associations among fishes. For example, imperiled species indicated associations with higher current velocities in run to riffle geomorphic units dominated by gravel and cobble substrates with little silt; imperiled

species tended to occur at higher densities among sites exhibiting a mix of these habitats (i.e., sites 2, 6, and 7).

Species-specific habitat associations were dependent upon habitat heterogeneity within the Big Bend reach of the Rio Grande, especially for imperiled species. Many of the imperiled Rio Grande fishes listed by Hubbs et al. (2008) and collected during this study were associated with gravel substrates in run to riffle geomorphic units with greater current velocities. These species were *C. elongatus*, *M. aestivalis*, *N. braytoni*, *N. jemezianus*, and *R. cataractae*. Although canonical correspondence analysis explained only 6.2% of variation among 2393 seine hauls, the resulting gradients are ecologically relevant based on Økland's (1999) conclusion that even low levels of explained percent variation can

establish ecologically meaningful gradients. Cross-validation techniques indicated that habitat variables explained a significant portion of taxonomic variation. Some potential causes for the observed low levels of explained variation include processes operating at larger spatial scales (e.g., a riverscape perspective) or environmental parameters not measured during our study. For example, the only imperiled species not listed above but collected during this study was *N. chihuahuana*, a species known to inhabit tributary streams and disperse infrequently into mainstem habitats (Edwards et al. 2002). Canonical correspondence analysis classified this species as associated with backwaters characterized by silt substrate, but other factors such as distance to nearest tributary might have explained the occurrence of this species more appropriately. *Cyprinella lutrensis* dominance at upstream sites and *N. braytoni* dominance at downstream sites is another example of a pattern that our environmental data were likely not capable of explaining. Based on broad physiological tolerances of *C. lutrensis* (Matthews 1986b) and occurrence of poor water quality and documented fish kills in upstream reaches of the Big Bend reach of the Rio Grande (USGS 2004), reduced abundance of *C. lutrensis* at downstream sites might be an artifact of longitudinal improvement of water quality and the return of a more natural fish assemblage (Kinsolving and Bain 1993). Regardless, our findings support the assertion that habitat heterogeneity is correlated with increased species diversity (Guégan et al. 1998), and preservation of heterogeneity is likely necessary for imperiled species conservation (Lake et al. 2007), which might explain why the Big Bend reach maintains a relatively intact fish assemblage.

Fish community composition in the Big Bend reach was correlated with high seasonal streamflows associated with monsoonal precipitation. During the pre-flood season, streamflows ranged between 0.22 and 31.90 m³ · sec⁻¹, and assemblage composition was dominated by *C. lutrensis* and *N. braytoni*. During the flood season, streamflows ranged from 8.7 to 186.0 m³ · s⁻¹, and density decreased by 48.0% for *C. lutrensis* and 24.2% for *N. braytoni*. Conversely, during the flood season, density was 4 times greater for *Cyprinus carpio*, 3 times greater for *I. furcatus*, 8 times greater for *I. punctatus*, and 1.5 times greater for *M. beryllina*. Based on the susceptibility of *C. lutrensis*, *Cyprinus*

carpio, *I. furcatus*, *I. punctatus*, and similar species to displacement or mortality during high streamflows (Minckley and Meffe 1987), we suspect that observed density decreases for *C. lutrensis*, *N. braytoni*, and river carpsucker (*Carpiodes carpio*) were related to downstream displacement (Collins et al. 1981) and that observed density increases for *Cyprinus carpio*, *I. furcatus*, *I. punctatus*, and *M. beryllina* were likely related to transport from upstream reservoirs during water releases (Ward and Stanford 1979, Schultz et al. 2003). Over 80% of water in the Big Bend region of the Rio Grande enters from the Rio Conchos Basin of Mexico, where monsoonal precipitation events during the flood season cause inundation of reservoirs (Edwards et al. 2002, Kim et al. 2002), and large-scale water releases from reservoirs likely transported many individuals from upstream portions of the Rio Conchos (e.g., Minckley and Meffe 1987).

Large flood pulses during the monsoon season correlated with dynamic changes in dominant species within communities, despite consistency of sampled habitats. *Notropis braytoni* proportional density indicated little change during the onset of monsoonal flooding during August–November, whereas *C. lutrensis* density decreased until the postflood season (November–December). Monsoonal flooding in the Big Bend region typically begins during late July and continues through late October. The timing of these flood pulses combined with life history attributes and habitat associations of *C. lutrensis* and *N. braytoni* might explain the observed changes in dominance during the post-flood season. Age-0 cyprinids are susceptible to downstream transport during flooding because of small body size and poor swimming ability, but susceptibility to displacement decreases with ontological growth (Harvey 1987). Spawning seasons differ for *C. lutrensis* (April–September; Farringer et al. 1979) and *N. braytoni* (February–August; T. Heard, unpublished data), and these differences might allow for an additional month of growth for *N. braytoni* before the onset of monsoonal flooding. Earlier spawning is advantageous in that individuals hatched early in a reproductive season exhibit greater growth rates compared to individuals hatched during the later part of a reproductive season (Durham and Wilde 2005), and early spawning may offer additional benefits by allowing individuals to achieve a larger body size before arrival of flood pulses during the flood season.

From a habitat suitability standpoint, *N. braytoni* was more tolerant of higher current velocities than was *C. lutrensis*, as evidenced by the separation of the 2 species along CCA I and by frequency distributions of observed current velocity associations. These observed habitat associations, combined with the coincidental onset of the flood season and declines in *C. lutrensis* proportional density, support published accounts of natural (and dynamic) flow regimes benefiting native and endemic species more than habitat generalist or introduced species (Propst and Gido 2004). Moreover, persistent low-flow conditions typically favor nonnative or native-invasive species by removing displacing floods and effectively limiting habitat heterogeneity that might otherwise be necessary for the persistence of native or imperiled species (Scott and Helfman 2001, Propst and Gido 2004, Perkin and Bonner 2011).

The fish assemblage in the Big Bend reach of the Rio Grande maintains high integrity compared to assemblages in other reaches in the basin. Exotic fishes comprise >50% of species richness in the upper reaches of the Rio Grande from its headwaters downstream to the Big Bend reach (Calamusso et al. 2005), 37% of species richness and 63% of relative abundance from Amistad Reservoir to Falcon Reservoir downstream of the Big Bend reach, and 33% of species richness and 71% of relative abundance downstream from Falcon Reservoir to Hidalgo, Texas (M. Bean, Texas Parks and Wildlife Department, and T. Bonner, unpublished data). Imperiled taxa downstream of Big Bend reach have a relative abundance of <5%. In contrast, >30% (6 of 19 species) of the Big Bend reach fish assemblage sampled during this study consisted of imperiled species, which collectively constituted a relative abundance of 39%. Exotic species constituted <2% of the sampled assemblage.

This pattern of imperiled species persistence is likely related to the relatively intact nature of the river in the Big Bend region. For example, stream fragmentation is implicated in declines and extirpations of species in many Rio Grande Basin fragments <100 km in length (Dudley and Platania 2007), but the distance between Caballo Dam (upstream of the study reach) and Amistad Dam (downstream) is >1000 km. Complete dessication of the Rio Grande near the city of El Paso, Texas, is associated with water withdrawal and retention

(Calamusso et al. 2005), but stable base flows of at least $12 \text{ m}^3 \cdot \text{sec}^{-1}$ persist at the International Boundary and Water Commission (IDWC) streamflow gage near Presidio, Texas, USA, and Ojinaga, Chihuahua, Mexico (gage #08-3742.00). Collectively, these factors contribute to instream habitat, stream connectivity, and streamflow regime attributes that allow the persistence of imperiled species.

Reductions in abundance and distribution are widespread for imperiled freshwater fishes (Jelks et al. 2008) and coincide with hydrologic alteration in the Rio Grande (Small et al. 2009). Despite persistence of monsoonal flood seasons, anthropogenic alteration of flow regime has caused substantial changes in stream morphology and instream habitat availability in the Big Bend reach (Dean and Schmidt 2011). Channel width has been reduced by 36%–52% since as early as the 1940s, while vertical accretion within floodplains and propagation of exotic riparian vegetation have caused increased bank sedimentation and narrower channels with steeper banks (Schmidt et al. 2003, Dean and Schmidt 2011). Reduction in channel width and increased bank slope effectively limit abundance and availability of shallow runs and riffles characterized by greater current velocity and gravel to cobble substrates (Poff et al. 1997), which we found were associated with numerous imperiled fishes. Although costly, mechanical removal of exotic riparian vegetation along reaches might improve habitat heterogeneity by increasing sediment transport, providing potential for greater stream widths and reducing evapotranspiration by riparian vegetation by 35% (Zavaleta 2000). Artificially mimicking natural timing without magnifying or amplifying flood pulses might enhance native fish recruitment via displacement of introduced and habitat generalist species (Gido et al. 1997, Propst and Gido 2004); however, experimental flooding of the highly regulated Grand Canyon reach of the Colorado River did not displace habitat generalist species effectively at low magnitudes, and recurring floods were likely necessary to effectively reduce competition of habitat generalists with native obligate riverine species (Valdez et al. 2001). Even so, each of these management options might ultimately improve the conservation of Big Bend reach fish communities that still maintain imperiled species populations.

Since completion of this study, a large magnitude flood ($1450 \text{ m}^3 \cdot \text{s}^{-1}$) during the monsoonal flood season of 2008 (Fig. 2B) drastically influenced instream habitat and removed exotic riparian vegetation in much of the Big Bend reach. Due to the relative quality of instream habitat and riverscape connectivity in the Big Bend reach, an experimental population of threatened *H. amarus* was reintroduced in 2009 (Shiley and Lamberti 2010). In the years since, *H. amarus* eggs have been collected in drift samples (Roberson 2010) and adult individuals have been collected on at least 4 occasions (M. Bean, Texas Parks and Wildlife Department, personal communication). These anecdotal observations support our conclusions that maintaining a diversity of instream habitats coupled with dynamic flows likely allow for species-specific associations and persistence of imperiled fishes within the Big Bend reach of the Rio Grande.

ACKNOWLEDGMENTS

We thank C.S. Williams, D.T. Runyan, B.M. Fordham, and M.G. Bean for assistance in the field. Sampling permits and river access were granted by Texas State University–San Marcos (IACUC 06-0620_802), the Texas Parks & Wildlife Department (Permit SPR-0601-159; issued to THB, 1 Jun 2001–1 Jun 2007), Big Bend National Park (Permit BIBE-2005-SCI-0071; issued to THB, 1 May 2005–30 Dec 2007), and Black Gap Wildlife Management Area (BG 06-01; issued to TCH, 1 Jan 2006–31 Dec 2006). We extend a special thanks to the late C. Hubbs for his assistance during early portions of this study and for his lifelong dedication to Texas fishes. We thank C.S. Williams and C.W. Hoagstrom for comments on earlier versions of this manuscript. This project was funded by the Texas Rivers Systems Institute at Texas State University–San Marcos and, in part, by Fred and Yetta Richan Aquatic Biology Scholarships awarded to TCH.

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Received 3 August 2011
Accepted 10 November 2011