

Figure 1: Life cycle diagram and census points for pre-reproduction census of gizzard shad used in a density-dependent integral projection model.

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- 4 U.S. Geological Survey, Upper Mississippi Environmental Sciences Center
- Graphical Abstract
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- $_{11}$ density-dependent age-0 survival
- J. Peirce^{1,3}, G. Sandland^{2,3}, B. Bennie¹, R.A. Erickson⁴

14 Highlights

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- 16 density-dependent age-0 survival
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- We introduce an integral projection model that incorporates data from studies on 1) egg
 production in different size classes of adult gizzard shad, 2) density-dependent survival
 of age-0 shad, and 3) length distributions of gizzard shad in the Upper Mississippi
 River system that span nearly thirty years.
- We establish survival parameters using gizzard shad data collected from the 5 sites along the Mississippi River and validate our model using empirical information collected from the La Grange Reach of the Illinois River.

An integral projection model for gizzard shad (*Dorosoma cepedianum*) utilizing density-dependent age-0 survival

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1 Abstract

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Gizzard shad (Dorosoma cepedianum) is a common freshwater fish species found throughout the central and eastern portions of North America. Within these regions, gizzard shad play a number of critical roles in the freshwater community such as serving as prey for other fish species and translocating nutrients from substrates into the water column. Because of this, it is important that we understand how gizzard shad populations respond to environmental changes and what these changes may mean for aquatic communities in general and fish assemblages in particular. Here, we introduce an integral projection model (IPM) for gizzard shad that incorporates empirical information from a number of sources including Long Term Resource Monitoring (LTRM) data. IPMs are a generalization of stage-based, matrix population models that have been used to describe a wide range of organisms and, as such are a natural choice for gizzard shad since many aspects of their life cycle have been studied. We tested model outputs against empirical patterns reported for gizzard shad from a different location along the Illinois River (La Grange Reach). Results of our work suggest that our model could serve as an important tool for predicting gizzard shad population responses to changing environmental conditions, including those mediated through species invasions.

- 32 Keywords: population dynamics, fisheries, Mississippi River basin, population ecology,
- 33 invasive species impact

1. Introduction

- Gizzard shad (*Dorosoma cepedianum*) is a laterally compressed, deep-bodied fish species
- that occupies numerous aquatic systems throughout central, southern and eastern regions of

the United States (Pierce et al., 1981; Vanni et al., 2005). They have the potential to reach high abundances in more eutrophic habitats, such as reservoirs, leading to its dominance within fish assemblages. Because of this, gizzard shad have the potential to influence freshwater systems in a number of ways. First, young shad often serve as a critical food source for many fish species, including those of commercial and recreational importance (such as walleye and largemouth bass) (Jester et al., 1972). Thus, gizzard shad can serve as an important trophic link within aquatic food webs. Second, because detritus can serve as a primary food source throughout much of gizzard shad development (i.e., from the age-0 stage onward), these fish can transport nutrients from benthic regions into pelagic habitats (Mather et al., 1995; Schaus and Vanni, 2000; Vanni et al., 2005). This process can result in an increase in the nutrients available to organisms within the water column leading to increases in phytoplankton biomass, algal blooms, and, due to these conditions, shifts in freshwater community structure (Aday et al., 2003; Schaus and Vanni, 2000). Finally, the fact that detritus can comprise a substantial portion of gizzard shad diet also makes this species an important connection between terrestrial inputs and aquatic processes (Schaus and Vanni, 2000). Given its potentially important role in aquatic ecosystems, interest has intensified in understanding how gizzard shad populations respond to environmental changes (both nat-53 ural and anthropogenic) and what these changes may mean for freshwater communities in general and fish assemblages in particular. 55

Interactions within and between species throughout interconnected environments can have important consequences for fish populations across space and time (Thorp et al., 2006). For gizzard shad, previous work has suggested that fish densities can play an important role in both the growth and survival patterns observed in populations of these fish. For example, (Buynak et al., 1992) reported an inverse relationship between densities and the lengths of age-0 gizzard shad. Similarly (Welker et al., 1994) found that high densities of age-0 shad were negatively associated with both fish length and survival under both field and seminatural conditions. Finally, (Michaletz, 2010) reported that the densities of age-0 gizzard shad were negatively correlated with survival in two Missouri lakes. These patterns were attributed to intraspecific competition among young shad for prey (zooplankton) resources. Although intraspecific competition may be influencing life-history traits in subsequent stages

of gizzard shad development, little work has actually been conducted to address this (Di-Cenzo et al., 1996). There is also evidence that the densities of other co-occurring fish species (such as invasive carps) may negatively influence aspects of gizzard shad biology, such as body condition (Irons et al., 2007; Love et al., 2018).

Although substantial empirical work on gizzard shad biology has accumulated over the decades, few if any studies have attempted to use these data to model population dynamics in this species. Work by Catalano and Allen (2010, 2011) used empirically-based simulations of gizzard shad to assess population-level responses in this species. As part of this process, the authors investigated population-structure using fish lengths but did not consider the impacts of gizzard-shad densities on population patterns. Here, we introduce an integral projection model (IPM) for gizzard shad based on empirical data with density-dependent survival in age-0 fish. We then compare model outcomes to the dynamics reported for this fish species in a well-studied pool of the Illinois River (specifically, the La Grange Reach). Results from this work suggest that our model could be an important tool for predicting gizzard shad population responses to changing environmental conditions, including those mediated through species invasions (i.e., silver and bighead carp).

83 2. Model development

84 2.1. Gizzard shad life history

Mature gizzard shad tend to mate between May and June, although this can vary based on water temperatures. Males and females aggregate and then broadcast gametes into the surrounding water; fertilized eggs then settle and adhere to the bottom substrates. After a period of days, eggs hatch and fish develop from the larval stage to juveniles and eventually to adults. In many habitats, individuals can reach sexual maturity within a year. As gizzard shad mature, their diet preferences typically shift from phytoplankton and zooplankton early in development to detritus and zooplankton as adults. Given the large number of eggs produced by shad females (> 300,000/year), there is evidence that intraspecific competition can be intense during early developmental stages in this species leading to density-dependent mortality. The strength of competition may then subside as fish transition to feeding on different food types during later stages of development.

Table 1: A summary of parameters, their biological meaning, and source for mean values.

Parameter	Meaning (units)	Mean	Source
Logistic survival probability func-			
tion, $s(z)$			
s_{\min}	minimum survival	0.002	(Bodola, 1955)
$s_{ m max}$	maximum survival	$1 - 8.871K^{0.73}L_{\infty}^{-0.33}$	(Then et al., 2015)
α_s	inflection point	80.01	Estimated from LTRM dataset
β_s	slope	-139.93	Estimated from LTRM dataset
Growth function, $G(z, z')$			
L_{∞}	maximum length (in mm)	394.30	(Catalano and Allen, 2010)
K_g	individual growth rate	0.26	(Michaletz, 2017)
σ_g	growth standard deviation	25	(Michaletz, 2017)
Normal distribution of length of age-			
$1, C_1(z')$			
μ_r	mean length of recruitment (in	105	(Michaletz, 2017)
	mm)		
σ_r	standard deviation of length	25	(Michaletz, 2017)
Eggs produced, $\operatorname{egg}(\mathbf{z})$			
$\mathrm{egg}_{\mathrm{max}}$	maximum number of eggs pro-	737512.1	Estimated from (Jons and Mi-
	duced		randa, 1997)
$lpha_e$	inflection point	314.44	Estimated from (Jons and Mi-
			randa, 1997)
eta_e	slope	-7.18	Estimated from (Jons and Mi-
			randa, 1997)
Survival of age-0, $s_0(d(t))$			
a_0	intercept	0.27	Estimated from (Michaletz, 2010)
b_0	decay rate	0.003	Estimated from (Michaletz, 2010)
Spawning			
ν	probability that egg becomes vi-	0.002	(Bodola, 1955)
	able		
p_b	probability that female spawns	0.90	

6 2.2. Equations

We used an IPM to describe the life history of gizzard shad in the Upper Mississippi 97 River system (UMRS). IPMs were first introduced by Easterling (Easterling et al., 2000), as 98 a generalization of stage-based, matrix population models. Since that time, IPMs have been 99 used to describe a wide range of organisms (Ellner et al., 2016; Merow et al., 2014; Rees et al., 100 2014), but have only recently be used to model fish populations, such as (Erickson et al., 101 2017; Liao et al., 2019; White et al., 2016; Pollesch et al., 2022). Fish are an ideal group 102 to investigate using this approach as many species have been well studied and are therefore 103 associated with robust information about life-history traits (i.e., growth and reproduction). 104 Gizzard shad are one such species and have been the subject of numerous laboratory and field studies over the decades making it a logical choice for IPMs. In our model specifically, 106 we incorporate data from studies on 1) egg production in different size classes of adults, 2) 107 density-dependent survival of age-0 shad, and 3) length distributions of gizzard shad in the 108 UMRS that span nearly thirty years. 109 We assumed that variations among individual gizzard shad can be summarized by its 110 length z (in mm) ranging from the minimum possible length L to the maximum value U. 111 The state of the population at time t (in years) is described by the length distribution 112

We assumed that variations among individual gizzard shad can be summarized by its length z (in mm) ranging from the minimum possible length L to the maximum value U. The state of the population at time t (in years) is described by the length distribution n(z,t). Specifically, for each time t, n(z,t) is a smooth function of z such that the number of individuals of length z in the interval [a,b] at time t is $\int_a^b n(z,t) \, dz$. Between times t and t+1, individual gizzard shad may grow, die, and produce offspring that vary in length depending on the individual's current length (Figure 2).

2.2.1. Growth and survival

For an individual of length z at time t, $P(z',z)\Delta z$ is the probability that the individual is alive at time t+1, and its size is in the interval $[z',z'+\Delta z]$ (as with n(z,t) this is an approximation that is valid for small Δz , and the exact probability is given by an integral like the one above). We define P(z',z)=s(z)G(z',z) where s(z) is the annual adult survival probability and the growth G(z',z) describes the annual length transitions. The survival function is a logistic function,

$$s(z) = s_{\min} + \frac{s_{\max} - s_{\min}}{1 + e^{\beta_s(\ln(z) - \ln(\alpha_s))}},\tag{1}$$

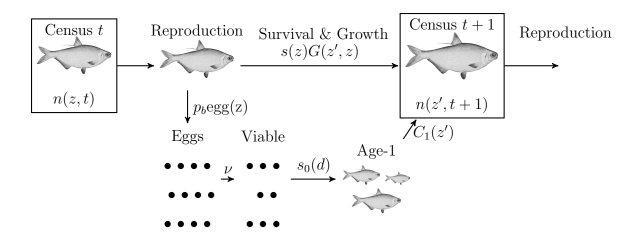


Figure 2: Life cycle diagram and census points for pre-reproduction census of gizzard shad.

with four parameters: the minimum survival rate s_{\min} ; a maximum survival rate, s_{\max} ; an intercept parameter, α_s ; and a slope parameter, β_s (Bolker, 2008). The growth function is a two-variable normal distribution centered around a modified von Bertalanffy function of the length at time t (Erickson et al., 2017). Consequently if L_{∞} is maximum asymptotic length and K_g is the individual growth rate of gizzard shad, then the growth kernel

$$G(z', z) = \text{Prob}(z' \mid z, L_{\infty}, K_g) = \text{NormalPDF}(\mu_g, \sigma_g)$$

where $\mu_g = L_{\infty} \left(1 - e^{-K_g} \right) + z(t)e^{-K_g}$ and σ_g is the standard deviation.

125 2.3. Fecundity

 $F(z',z)\Delta z$ is the number of new offspring in the length interval $[z',z'+\Delta z]$ present at time t+1, per length-z individual at time t. The fecundity kernel is

$$F(z', z) = p_b \operatorname{egg}(z) \nu s_0(d(t)) C_1(z')$$
(2)

where p_b is the probability of reproducing, egg(z) is the mean number of eggs produced, ν is the probability that an egg is viable, $s_0(d(t))$ is the density-dependent probability of surviving to age-1, and $C_1(z')$ is the length distribution of new recruits at age-1 (when they are first censused in the model).

The mean number of eggs produced by females of a certain length is a three-parameter logistic function,

$$\operatorname{egg}(z) = \frac{\operatorname{egg}_{\max}}{1 + e^{\beta_e(\ln(z) - \ln(\alpha_e))}}.$$
(3)

The probability of gizzard shad survival during their first year can depend on many factors (Michaletz, 2010) including predation, temperature, the mean total length of fish, and the density of age-0 fish. In this study, we focused only on the density factor and assumed the probability of survival of age-0 fish is the exponential function,

$$s_0(d(t)) = a_0 e^{-b_0 d(t)}, (4)$$

where a_0 is the intercept, b_0 the decay rate, and d(t) is the density at time t of age-0 gizzard shad per 1000 m³,

$$d(t) = 10^{-3} \int_{L}^{U} p_b \operatorname{egg}(z) \nu n(z, t) dz.$$

Finally, the total number of eggs that survive to be an age-1 fish is multiplied with a normal distribution of length, $C_1(z') = \text{NormalPDF}(\mu_r, \sigma_r)$ where μ_r is the mean length of age-1 gizzard shad and σ_r is the standard deviation.

143 2.4. Dynamical model

The population at time t+1 is the sum of the contributions from each individual alive at time t,

$$n(z', t+1) = \int_{L}^{U} K(z', z) n(z, t) dz,$$
 (5)

where K(z',z) = s(z)G(z',z) + F(z',z) and [L,U] is the range of all possible lengths.

147 3. Methods

148 3.1. LTRM data and model parameterization

The LTRM element of the Upper Mississippi River Restoration monitors the UMRS to 149 provide an understanding of the system's ecology, resource changes, and inform management 150 (Bouska et al., 2018; Maher et al., 2015). In order to achieve this goal, numerous features 151 of the UMRS, such as aquatic vegetation, bathymetry, fish, land use/land cover, and water 152 quality are continually surveyed from Navigation Pool 1 (at Minneapolis, Minnesota) south 153 to the confluence of the Mississippi and Ohio Rivers at Cairo, Illinois. LTRM fish surveys are 154 conducted at five locations along the main channel of the Upper Mississippi River (Pools 4, 155 8, 13, 26, and the Open River Reach) and at one location along the Illinois River (La Grange 156



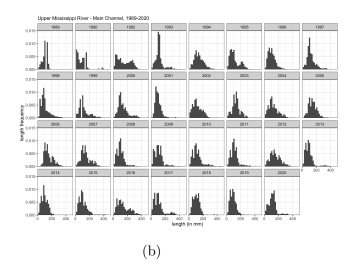


Figure 3: (a) The LTRM fish survey sites in the Upper Mississippi River system. (b) Length frequency of sampled gizzard shad in the main channel of the Upper Mississippi River system.

Reach) (Figure 3a). Fish are captured using a multiple gear approach (which includes netting and electrofishing) in order to monitor the responses and health of fish communities along 158 these two very important waterways over time (Gutreuter et al., 1995). Specific capture 159 methodologies, protocols and modifications to the LTRM can be found in Gutreuter et al. 160 (1995), and Ickes and Burkhardt (2002). In terms of gizzard shad, fish traits (such as total 161 length) have been recorded since 1989 with approximately 3000 collections occurring per year along the Mississippi River (Figure 3b). For the La Grange Reach of the Illinois River, 163 gizzard shad have been sampled and measured since 1990 with approximately 500 collections 164 occurring per year. 165

To parameterize our model, we used gizzard shad data collected from the 5 sites along the Mississippi River. We then validated our model using empirical information collected from the La Grange Reach of the Illinois River. We undertook this approach as the La Grange Reach is upstream of the Mississippi River making it a more distinct location compared to the other sites.

3.1.1. Fecundity and recruitment

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Female gizzard shad begin reproducing at approximately 140 mm in length and egg numbers tend to increase with fish size (Jons and Miranda, 1997). The logistic parameters for the mean number of eggs produced by females of a certain length were obtained by

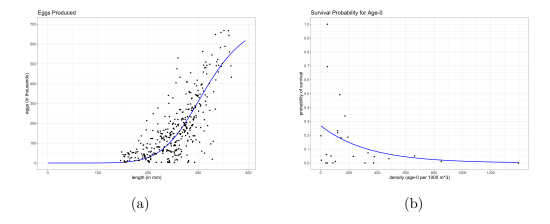


Figure 4: (a) Mean number of eggs produced by female gizzard shad egg(z). Data from (Jons and Miranda, 1997). (b) Age-0 density-dependent survival function $s_0(d)$. Data from (Michaletz, 2010).

fitting the three-parameter logistic function (Equation 3) to the data for batch fecundity 175 versus length (Jons and Miranda, 1997) (Figure 4a). Fish survival to age-1 was assumed to be dependent on the density of age-0 gizzard shad (Figure 4b). Parameters for the exponentially 177 decaying age-0 survival function (Equation 4) were determined by fitting equation 4 to the 178 survival means for 2003-2007 cohorts of gizzard shard in five Missouri reservoirs (Michaletz, 179 2010). To complete the recruitment process we assign a length to the recruited individuals 180 by simulating a Gaussian random variable with mean μ_c and standard deviation σ_c . The 181 parameters for the size distribution of age-1 fish were gleaned from a study of gizzard shad 182 located in large impoundments (Michaletz, 2017) and the historic 1990-2020 LTRM dataset 183 from the main channel of the UMRS (discussed in Section 3.1). 184

3.1.2. Growth and survival of adults

The parameters for the growth function were chosen as the mean values published on a 186 study of gizzard shad located in large impoundments in Missouri U.S.A. (Michaletz, 2017). 187 The association between adult lengths and survival have not been well-resolved in gizzard 188 shad leading us to make a number of assumptions. First, we assume that the probability 189 of adult survival is related to the length by a four-parameter logistic function (Equation 190 1). An investigation of gizzard shad in Lake Eerie (Bodola, 1955) provided the minimum 191 and maximum survival rate of adults. Based on the observed length distributions of gizzard 192 shad sampled from the main channel of the Mississippi River (Figure 3b), we assumed 193

that solutions of our model will exhibit periodic behavior every 8-9 years. We used a least squares method to estimate the α_s and β_s parameters that minimized the total square-195 distance between the (observed) pre-carp LTRM length distribution in the main channel 196 of the UMRS and (predicted) model equilibrium, n(z,t) during a 8-year period occurring 197 100 years after initialization. The slope parameter β_s was found to be large in magnitude resulting in a primarily two-valued survival probability. Gizzard shad less than α_s mm 199 in length have a very low survival rate (s_{\min}) while lengths larger than α_s mm approach 200 the maximum survival rate s_{max} . This survival pattern has been reported for a number of 201 fish species and can arise due to a number of biotic (i.e. predation) and/or abiotic (i.e. 202 temperature) factors (Pepin et al., 1992; Nowlin et al., 2006).

204 4. Analysis and results

We numerically solved the integral model using the Midpoint Rule with large approximating matrices (Burden and Faires, 2005). The Midpoint Rule has been commonly used for integral projection models because of its simplicity and effectiveness (Ellner and Rees, 2006; Ramula et al., 2009; Merow et al., 2014). During the course of model development, we explored different step sizes for the Midpoint Rule and found that about 50 points provided numerically stable results. We integrated over lengths from 0 mm to 500 mm. The upper limit was chosen based upon numerical stability and consistency of the system (e.g., avoiding eviction or the loss of individuals due to numerical errors (Williams et al., 2012)).

4.1. Initial conditions

We assumed that the initial density of gizzard shad was $d_0 = 964.7$, the annual average density of gizzard shad observed in La Grange Reach from 1993-2020. The probability of an individual being length z at time t = 0 was assumed to be normally distributed with mean $0.5L_{\infty}$ and standard deviation $\sigma_0 = 30$. As a result, we initialized our model with length distribution

$$n(z,0) = d_0 \text{Norm}(0.5L_\infty, \sigma_0) = 964.7 \text{Norm}(197.15, 30).$$
 (6)

The model was coded in R (R Core Team, 2017) and the scripts are published on JP's GitHub page https://github.com/jppeirce.

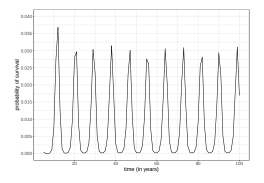


Figure 5

Figure 6: Survival probability of age-0 gizzard shad.

4.1.1. Effect of density-dependence on the survival probability of the age-0 cohort

In the simulated solutions to the IPM, the density of age-0 Fish strongly influenced the 222 density of gizzard shad at subsequent developmental stages. When adult densities are large, 223 there may be more fish of longer lengths that can produce a greater number of eggs. More 224 eggs leads to a higher density of age-0 fish and reflectively a reduction in the survival to 225 age-1. If reduced survival continues over subsequent years, the overall density of fish within 226 the population may decline and result in a smaller number of longer length fish that are 227 reproducing. Fewer fish spawning could result in a smaller age-0 class which, in turn, could 228 enhance survival probability in this cohort (through reduced competition). We would then expect the overall density of fish to increase over the following years, until large numbers 230 of eggs are again produced by larger, adult fish. This oscillatory pattern is reflected in our 231 model by the time-dependent survival probability of age-0 recruits (Figure 5). 232

4.1.2. Periodic orbit and validation with external dataset

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The total number of gizzard shad in our simulation reached a stable periodic orbit (Figure 7a) within 50 years. After simulating an additional 50 years, we fit a periodic function to the annual density of gizzard shad and determined the period of approximately 8.74 years. Figure 7b illustrates the periodic length dynamics within the gizzard shad population during a 9-year window of the periodic orbit. As a validation of the model, the simulated length distributions during a periodic orbit (Figure 7b) have similarities to fish lengths observed from the La Grange Reach (Figure 8a). In addition, the ebb and flow of the frequency of the

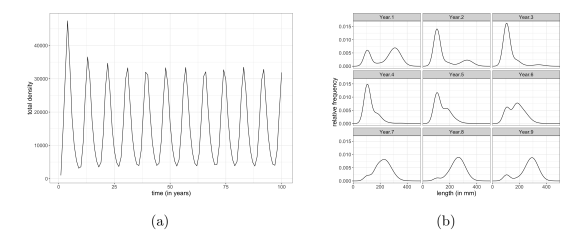


Figure 7: (a) The total density of gizzard shad in La Grange Reach predicted by the IPM in the first 100 years. (b) Simulated length distributions during a 9 year interval of time (approximately 1 period of the total density function).

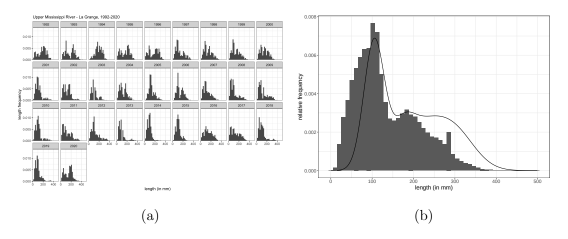


Figure 8: (a) Length frequencies of sampled gizzard shad in La Grange for each year, 1992-2020. (b) Overall length frequencies of gizzard shad sampled in La Grange Reach from 1992-2020 compared with the average (over 1 period) simulated length frequency.

²⁴¹ age-1 cohort can be observed and is associated with the density-dependent survival function ²⁴² for age-0 fish (explained in Section 4.1.1).

5. Discussion

Gizzard shad are common in freshwater systems of North America where they contribute to the integrity of aquatic communities. Given this, it is important to understand how populations of this species vary with changing environmental circumstances, such as the occurrence of invasive species. Herein, we use both LTRM data and parameters gleaned from other empirical studies to develop an integral projection model for gizzard shad and test model outputs against patterns reported from a well-studied population of this species from the La Grange Reach of the Illinois River.

After fitting two adult survival parameters using LTRM data from the main channel 251 of the Mississippi River, we compared a simulated length distribution with empirical data 252 collected from the La Grange Reach of the Illinois River from 1992 to 2020. The resulting 253 simulated length distributions (Figure 7b) reflected a number of the patterns observed in gizzard shad captured from the La Grange Reach over the same timeframe (Figure 8a). 255 For example, the predicted transition from bimodal length distributions to a single peak 256 in smaller gizzard shad in our simulations was reflected in the empirical data from 1996 to 257 2000. This trend was also observed from 2004-2008. It should be noted, that there were 258 also discrepancies between model simulations and the empirical data. In particular, we did not notice a distinct, single peak in intermediate sized fish from field collections even 260 though this was predicted by our model. Temporal and spatial variations in the La Grange 261 Reach environment and their subsequent impacts on growth within the shad population may 262 help explain why certain trends in our IPM outputs were not well represented in the field collections. 264

The average (over 1 period) simulated length frequency compared well with the overall length frequencies of gizzard shad sampled in La Grange Reach from 1992-2020 (Figure 8b). Our simulated size frequencies of gizzard shad captured the general trends seen in fish collected from the study site; however the model did predict a slightly higher density of adult lengths and fewer juvenile (smaller) lengths than in the empirical data. This may

be explained by gear type and capture method which vary from site to site potentially introducing bias in the observed length distributions year to year. In addition, studies also suggest that environmental stochasticity and food variability may alter recruitment densities which are difficult to measure accurately (Rose, 2000; Okamoto et al., 2016).

While our model uses the density of age-0 gizzard shad to affect the survival probability in their first year, we assumed constant viability at subsequent developmental stages, which may also be sensitive to density-related factors. That being said, there is little information available on the role that density plays in the life-history responses of adult gizzard shad. The location of the maximum length and the variation in the effect the survival probability in the LTRM data suggests that there may be smaller age-0 fish in La Grange Reach than in the study location (Michaletz, 2017) used to parameterize the model.

Gaining an understanding of how length distributions of gizzard shad emerge under density-dependent survival in the age-0 class will serve as a foundation for investigating density effects at subsequent stages in the life cycle. In addition, this single-species model could also be expanded to incorporate interspecific interactions between gizzard shad and species such as invasive carp, which appear to negatively impact gizzard shad life-histories through competition for food resources.

287 6. Acknowledgments

These data are a product of the U.S. Army Corps of Engineer's Upper Mississippi River 288 Restoration Program (UMRR) Long Term Resource Monitoring (LTRM) element imple-280 mented by the U.S. Geological Survey in collaboration with the five Upper Mississippi River 290 System (UMRS) states of Illinois, Iowa, Minnesota, Missouri, and Wisconsin. The U.S. 291 Army Corps of Engineers (Corps) provides guidance and has overall program responsibility. 292 We thank the U.S. Geological Survey Biological Threats and Invasive Species Program 293 and Great Lakes Restoration Initiative for funding. In addition, research was supported 294 by NSF-DMS Grant #1852224, "REU Site: Ecological Modeling of the Mississippi River 295 Basin". Any use of trade, firm, or product names is for descriptive purposes only and does 296 not imply endorsement by the U.S. Government. 297

298 References

- R. J. Pierce, T. E. Wissing, B. A. Megrey, Aspects of the feeding ecology of gizzard shad in Acton Lake, Ohio, Transactions of the American Fisheries Society 110 (3) (1981) 391–395.
- M. J. Vanni, K. K. Arend, M. T. Bremigan, D. B. Bunnell, J. E. Garvey, M. J. Gonzalez,
- W. H. Renwick, P. A. Soranno, R. A. Stein, Linking landscapes and food webs: effects of
- omnivorous fish and watersheds on reservoir ecosystems, BioScience 55 (2) (2005) 155–167.
- D. B. Jester, B. L. Jensen, et al., Life history and ecology of the gizzard shad, *Dorosoma* cepedianum (Le Sueur) with reference to Elephant Butte Lake.
- M. E. Mather, M. J. Vanni, T. E. Wissing, S. A. Davis, M. H. Schaus, Regeneration of nitrogen and phosphorus by bluegill and gizzard shad: effect of feeding history, Canadian Journal of Fisheries and Aquatic Sciences 52 (11) (1995) 2327–2338.
- M. H. Schaus, M. J. Vanni, Effects of gizzard shad on phytoplankton and nutrient dynamics: role of sediment feeding and fish size, Ecology 81 (6) (2000) 1701–1719.
- D. D. Aday, R. J. H. Hoxmeier, D. H. Wahl, Direct and indirect effects of gizzard shad on bluegill growth and population size structure, Transactions of the American Fisheries Society 132 (1) (2003) 47–56.
- J. H. Thorp, M. C. Thoms, M. D. Delong, The riverine ecosystem synthesis: biocomplexity in river networks across space and time, River Research and Applications 22 (2) (2006) 123–147.
- G. L. Buynak, R. S. Hale, B. Mitchell, Differential growth of young-of-year gizzard shad in several Kentucky reservoirs, North American Journal of Fisheries Management 12 (3) (1992) 656–662.
- M. T. Welker, C. L. Pierce, D. H. Wahl, Growth and survival of larval fishes: roles of competition and zooplankton abundance, Transactions of the American Fisheries Society 123 (5) (1994) 703–717.

- P. H. Michaletz, Overwinter survival of age-0 gizzard shad in Missouri reservoirs spanning a productivity gradient: roles of body size and winter severity, Transactions of the American Fisheries Society 139 (1) (2010) 241–256.
- V. J. DiCenzo, M. J. Maceina, M. R. Stimpert, Relations between reservoir trophic state and gizzard shad population characteristics in Alabama reservoirs, North American Journal of Fisheries Management 16 (4) (1996) 888–895.
- K. S. Irons, G. Sass, M. McClelland, J. Stafford, Reduced condition factor of two native fish species coincident with invasion of non-native Asian carps in the Illinois River, USA Is this evidence for competition and reduced fitness?, Journal of Fish Biology 71 (2007) 258–273.
- S. A. Love, N. J. Lederman, R. L. Anderson, J. A. DeBoer, A. F. Casper, Does aquatic invasive species removal benefit native fish? The response of gizzard shad (Dorosoma cepedianum) to commercial harvest of bighead carp (Hypophthalmichthys nobilis) and silver carp (H. molitrix), Hydrobiologia 817 (1) (2018) 403–412.
- M. J. Catalano, M. S. Allen, A size-and age-structured model to estimate fish recruitment, growth, mortality, and gear selectivity, Fisheries Research 105 (1) (2010) 38–45.
- M. J. Catalano, M. S. Allen, A whole-lake density reduction to assess compensatory responses
 of gizzard shad *Dorosoma cepedianum*, Canadian Journal of Fisheries and Aquatic Sciences
 68 (6) (2011) 955–968.
- A. Bodola, The life history of the gizzard shad, *Dorosoma cepedianum*, in western Lake Erie,
 Ph.D. thesis, Doctoral dissertation. Ohio State University, Columbus, 1955.
- A. Y. Then, J. M. Hoenig, N. G. Hall, D. A. Hewitt, H. editor: Ernesto Jardim, Evaluating the predictive performance of empirical estimators of natural mortality rate using information on over 200 fish species, ICES Journal of Marine Science 72 (1) (2015) 82–92.
- P. Michaletz, Variation in characteristics among gizzard shad populations: The role of impoundment size and productivity, Fisheries Management and Ecology 24 (5) (2017) 361–371.

- G. Jons, L. Miranda, Ovarian weight as an index of fecundity, maturity, and spawning periodicity, Journal of Fish Biology 50 (1) (1997) 150–156.
- M. R. Easterling, S. P. Ellner, P. M. Dixon, Size-specific sensitivity: applying a new structured population model, Ecology 81 (3) (2000) 694–708.
- S. P. Ellner, D. Z. Childs, M. Rees, et al., Data-driven modelling of structured populations:
- A practical guide to the Integral Projection Model., Springer International Publishing,
- Switzerland, 2016.
- ³⁵⁷ C. Merow, J. P. Dahlgren, C. J. E. Metcalf, D. Z. Childs, M. E. Evans, E. Jongejans,
- S. Record, M. Rees, R. Salguero-Gómez, S. M. McMahon, Advancing population ecology
- with integral projection models: a practical guide, Methods in Ecology and Evolution
- ³⁶⁰ 5 (2) (2014) 99–110.
- M. Rees, D. Z. Childs, S. P. Ellner, Building integral projection models: a user's guide,

 Journal of Animal Ecology 83 (3) (2014) 528–545.
- R. A. Erickson, E. A. Eager, M. K. Brey, M. J. Hansen, P. M. Kocovsky, An integral projection model with YY-males and application to evaluating grass carp control, Ecological
- 365 Modelling 361 (2017) 14–25.
- B. Liao, X. Shan, C. Zhou, Y. Han, Y. Chen, Q. Liu, A dynamic energy budget—integral
- projection model (DEB-IPM) to predict population-level dynamics based on individual
- data: a case study using the small and rapidly reproducing species *Engraulis japonicus*,
- Marine and Freshwater Research 71 (4) (2019) 461–468.
- J. W. White, K. J. Nickols, D. Malone, M. H. Carr, R. M. Starr, F. Cordoleani, M. L.
- Baskett, A. Hastings, L. W. Botsford, Fitting state-space integral projection models to
- size-structured time series data to estimate unknown parameters, Ecological Applications
- 26 (8) (2016) 2677–2694.
- N. Pollesch, K. Flynn, S. Kadlec, J. Swintek, S. Raimondo, M. Etterson, Developing integral projection models for ecotoxicology, Ecological Modelling 464 (2022) 109813.

- B. M. Bolker, Ecological models and data in R, Princeton University Press, 2008.
- K. L. Bouska, J. N. Houser, N. R. De Jager, J. Hendrickson, Developing a shared understanding of the Upper Mississippi River, Ecology and Society 23 (2).
- R. Maher, M. Griffin, K. Stauffer, J. Sternburg, J. Fischer, K. Hagerty, M. Hubbell, B. John-
- son, K. McCain, K. Mitvalsky, T. Novak, K. Westlake, B. Clevenstine, J. Duyvejonck,
- J. Hauser, M. Jawson, B. Johnson, J. Stoner, G. Benjamin, D. Buntin, K. Mickelsen, En-
- hancing Restoration and Advancing Knowledge of the Upper Mississippi River, A Strate-
- gic Plan for the Upper Mississippi River Restoration Program 2015-2025, Tech. Rep., U.S.
- Army Corps of Engineers, 2015.
- S. Gutreuter, R. Burkhardt, K. S. Lubinski, Long Term Resource Monitoring Program procedures [microform] .
- B. S. Ickes, R. W. Burkhardt, Evaluation and proposed refinement of the sampling design for the Long Term Resource Monitoring Program's fish component, Tech. Rep., Geological Survey La Crosse WI Upper Midwest Environmental Sciences Center, 2002.
- R. Burden, J. D. Faires, Numerical analysis 8th ed, Thomson Brooks/Cole.
- S. P. Ellner, M. Rees, Integral projection models for species with complex demography, The
 American Naturalist 167 (3) (2006) 410–428.
- S. Ramula, M. Rees, Y. M. Buckley, Integral projection models perform better for small demographic data sets than matrix population models: a case study of two perennial herbs, Journal of Applied Ecology 46 (5) (2009) 1048–1053.
- J. L. Williams, T. E. Miller, S. P. Ellner, Avoiding unintentional eviction from integral projection models, Ecology 93 (9) (2012) 2008–2014.
- R Core Team, R: A Language and Environment for Statistical Computing, R Foundation for Statistical Computing, Vienna, Austria, URL https://www.R-project.org/, 2017.
- K. A. Rose, Why are quantitative relationships between environmental quality and fish populations so elusive?, Ecological Applications 10 (2) (2000) 367–385.

D. K. Okamoto, R. J. Schmitt, S. J. Holbrook, Stochastic density effects on adult fish survival and implications for population fluctuations, Ecology letters 19 (2) (2016) 153–162.