

RESEARCH ARTICLE

Body ram, not suction, is the primary axis of suction-feeding diversity in spiny-rayed fishes

Sarah J. Longo^{1,*}, Matthew D. McGee¹, Christopher E. Oufiero², Thomas B. Waltzek³ and Peter C. Wainwright¹

ABSTRACT

Suction-feeding fishes exhibit diverse prey-capture strategies that vary in their relative use of suction and predator approach (ram), which is often referred to as the ram–suction continuum. Previous research has found that ram varies more than suction distance among species, such that ram accounts for most differences in prey-capture behaviors. To determine whether these findings hold at broad evolutionary scales, we collected high-speed videos of 40 species of spiny-rayed fishes (Acanthomorpha) feeding on live prey. For each strike, we calculated the contributions of suction, body ram (swimming) and jaw ram (mouth movement relative to the body) to closing the distance between predator and prey. We confirm that the contribution of suction distance is limited even in this phylogenetically and ecologically broad sample of species, with the extreme suction area of prey-capture space conspicuously unoccupied. Instead of a continuum from suction to ram, we find that variation in body ram is the major factor underlying the diversity of prey-capture strategies among suction-feeding fishes. Independent measurement of the contribution of jaw ram revealed that it is an important component of diversity among spiny-rayed fishes, with a number of ecomorphologies relying heavily on jaw ram, including pivot feeding in syngnathiforms, extreme jaw protruders and benthic sit-and-wait ambush predators. A combination of morphological and behavioral innovations has allowed fish to invade the extreme jaw ram area of prey-capture space. We caution that while two-species comparisons may support a ram–suction trade-off, these patterns do not speak to broader patterns across spiny-rayed fishes.

KEY WORDS: Ram–suction, Continuum, Prey capture, Jaw ram, Constraint, Acanthomorpha

INTRODUCTION

The ability to produce suction is an important adaptation for capturing prey in aquatic environments. Suction-feeding organisms take up food by generating a flow of water into the mouth through rapid expansion of the oral cavity. Such mechanisms have evolved multiple times in aquatic groups of vertebrates and are found today in sharks and rays, fishes, turtles, amphibians, birds and mammals (Wainwright et al., 2015). By using suction to draw prey (and water) towards their mouth, predators use the viscosity of water to their advantage, as these flows and the forces they exert on prey are difficult to overcome during prey escape attempts (Van Leeuwen

and Muller, 1984; Holzman and Wainwright, 2009). However, suction flows are only significant roughly a single mouth diameter in front of the predator's mouth (Muller et al., 1982; Muller and Osse, 1984; Van Leeuwen, 1984; Ferry-Graham et al., 2003; Day et al., 2005), making the approach and positioning of the mouth near prey key to a successful prey-capture strategy (de Jong et al., 1987; Holzman et al., 2012).

Suction-feeding predators use a variety of mechanisms to quickly move their mouth in close proximity to the prey, including overtaking them with a burst of swimming, ambushing them from a concealed location in close quarters, or protruding their jaws toward the prey while the body remains motionless. A central challenge in understanding the diversity of feeding behaviors in aquatic feeding vertebrates has been to place this behavioral diversity into a mechanistic framework that captures the major axes of diversity. Movements of an aquatic predator towards prey are often summarized as the ram components of a feeding strike (Liem, 1980a), which are then split into two sources of movement: body ram, or movements of the predator's body towards the prey by swimming or coasting, and jaw ram, or movements of the predator's mouth towards the prey relative to the rest of the predator's body (Liem, 1980b; Norton and Brainerd, 1993; Nyberg, 1971; Osse, 1985). Body ram and jaw ram can be used in combination with suction to decrease the distance between predator and prey. The amount of body ram also influences initial predator–prey distance, strike speed and strike duration, as well as the shape and volume of ingested water during the strike (Weihs, 1980; Harper et al., 1991; Higham et al., 2005; Tran et al., 2010; Oufiero et al., 2012). Jaw protrusion is the most common mechanism of jaw ram and has been shown to decrease the hydrodynamic disturbance detectable by prey while significantly increasing the suction forces on prey (Holzman et al., 2008; Holzman and Wainwright, 2009; Staab et al., 2012).

In this study, we explored the diversity of suction-feeding behaviors shown by spiny-rayed fishes (Acanthomorpha). To characterize this diversity, we separated the contributions of suction, body ram and jaw ram to closing the distance between predator and prey. There is a long tradition of quantifying the relative contributions of ram and suction to prey capture (Norton, 1991, 1995; Norton and Brainerd, 1993; Cook, 1996; Gibb, 1997; Nemeth, 1997; Ferry-Graham et al., 2001a; Kerfoot and Turingan, 2011; Wainwright et al., 2001; Wintzer and Motta, 2005; Wilga et al., 2007; Tran et al., 2010; Ferry et al., 2012; Staab et al., 2012). It is well recognized that fishes vary in the relative amount of ram and suction employed during feeding, and hence the 'ram–suction continuum' is a pervasive framework used to characterize diversity (Norton and Brainerd, 1993). This framework has helped clarify ecomorphological traits that are often associated with the extremes of this continuum. For instance, two members of Centrarchidae, bluegill (*Lepomis macrochirus*) and largemouth bass (*Micropterus salmoides*), have often been held up as examples of species that feed with suction- or ram-dominated strikes, respectively (Norton and

¹Department of Evolution and Ecology, University of California, Davis, One Shields Avenue, Davis, CA 95616, USA. ²Department of Biological Sciences, Towson University, Towson, MD 21252, USA. ³Department of Infectious Diseases and Pathology, College of Veterinary Medicine, University of Florida, PO Box 110880, Gainesville, FL 32611, USA.

*Author for correspondence (sjlongo@ucdavis.edu)

Brainerd, 1993; Higham et al., 2006a,b; Higham, 2007; Wainwright et al., 2007).

Recent studies have shown that the continuum between ram and suction may be more complex than was originally thought. The limited reach of suction has been found to lead to very little diversity in suction distance (Wainwright et al., 2001), while there is much more variation possible in ram distance (Ferry et al., 2001b; Wainwright et al., 2001; Tran et al., 2010). Instead of a straightforward continuum between ram speed and suction capacity, the diversity of suction capacity was found to be dependent on attack speed among 30 serranid species (Oufiero et al., 2012). Others have recently focused on the duration of suction and predatory movements during feeding strikes in ray-finned fishes and found that the majority of variation among strikes is driven by the duration of ram movement (Ferry et al., 2015). Collectively, these studies suggest that because of the hydrodynamic constraints of suction, variation in prey-capture diversity is driven almost entirely by the amount of ram employed.

Although jaw ram is part of the suction-feeding paradigm in acanthomorph fishes (Motta, 1984), most comparative analyses focus on jaw ram's relationship to suction or how jaw ram contributes to overall ram (Norton and Brainerd, 1993; Wainwright et al., 2001; Oufiero et al., 2012). However, the kinematics of jaw ram are morphologically independent from those of body ram (fish can protrude their jaw without swimming), and therefore jaw ram provides an independent axis on which fish can diversify their prey-capture strategies. Multiple independent origins of jaw protrusion among actinopterygian fishes (Wainwright et al., 2015) and novel morphological adaptations for extreme jaw protrusion in acanthomorphs (Ferry-Graham et al., 2001a; Waltzek and Wainwright, 2003; Westneat and Wainwright, 1989) attest to the adaptive significance of jaw ram in prey capture. However, reliance on jaw ram alone or in large proportion may be constrained in a similar fashion to suction; there should be inherent structural limits to how far and fast a fish can protrude or rotate the mouth or head relative to the rest of the body (Westneat, 1991). Adding a jaw ram axis to our analyses allowed us to look for patterns of variation in jaw ram that may not be otherwise apparent if jaw ram is simply lumped into total ram.

In this study, we asked whether recent comparative findings hold for acanthomorphs in general and whether the isolation of jaw ram contributes to clarifying the diversity of attack strategies used by suction-feeding fishes. Across this group, do we see specialists that make almost exclusive use of suction, body ram or jaw ram during prey capture? Is diversity governed by a continuum from suction to ram, or is it more accurately defined by the relative amount of ram? How constraining is the spatial limitation of suction feeding on prey-capture behaviors? Evolution has a knack of finding ways around biomechanical constraints thought to be insurmountable (e.g. elastic-recoil mechanisms circumvent the limitations of the force–velocity trade-off of muscles; Roberts and Azizi, 2011). Indeed, new evidence suggests that simple trade-offs do not limit the evolutionary diversification of complex mechanisms as expected (Holzman et al., 2011; Oufiero et al., 2012). Therefore, we might expect to find some acanthomorphs that have evolved morphological or behavior traits that allow them to be suction specialists.

MATERIALS AND METHODS

Collection and husbandry

The majority of specimens included in this study were commercially obtained from the aquarium industry. *Lepomis macrochirus* and *Micropterus salmoides* were collected locally in Yolo County, CA, USA.

With the exception of *Macroramphosus scolopax*, fish were housed in the laboratory between 22 and 23°C; *M. scolopax* required colder temperatures and was housed at 17°C. Fish were kept in 18–110 l aquaria, depending on their size. We filmed 40 species in 33 families of acanthomorph fishes (Table 1) covering a wide range of ecologies. More than one representative was included for some families to capture some diversity at this scale, including cichlids (Liem, 1973), haemulids (Tavera et al., 2012) and serranids (Oufiero et al., 2012). Among the prey-capture strategies we targeted were sit-and-wait ambush predators (*Antennarius hispidus*, *Inimicus didactylus*), pivot-feeding syngnathiforms (*Aulostomus maculatus*, *Aeoliscus strigatus*), high-ram suction feeders (*Ephinephelus ongus*, *Caranx sexfasciatus*), ram-biters (*Sphyrna barracuda*), benthic invertebrate pickers or foragers from both freshwater and saltwater environments (*L. macrochirus*, *Dactylopus dactylopus*), water-column zooplanktivores (*M. scolopax*, *Emmelichthys atlanticus*), and some species with morphological adaptations for extreme jaw protrusion (*Epibulus insidiator*, *Caquetaia kraussi*). All fish were filmed feeding on live prey. Because of their extremely small gapes, pipefish (*Doryrhamphus exicus*) and shrimpfish (*A. strigatus*) were fed freshly hatched brine shrimp (*Artemia*). All other species were fed live cyprinid or poeciliid fish (mostly *Danio rerio* or *Gambusia affinis*).

The feeding kinematics of many species in this dataset have not been previously published and deserve detailed attention, but that is beyond the scope of this study. Because the purpose of this dataset was to sketch the diversity of acanthomorph feeding in prey-capture space and determine the axes of variation underlying their distribution, a single sequence representing a typical feeding event was chosen for each species. In practice, typical strikes approximated species means calculated from multiple individuals. We made one exception and included a somewhat atypical strike by *Serranocirrhitis latus* to visually demonstrate the highest suction proportion we found, although including this strike or a typical one did not change the patterns we describe for acanthomorphs. We also investigated the sensitivity of our approach to variation in the choice of a typical strike using a resampling method (see 'Data visualization and statistical analysis', below).

Isolated comparisons focused on the prey-capture strategy of three pairs of species previously identified as exemplars of closely related species at extremes of the ram–suction continuum. *Lepomis macrochirus* and *M. salmoides* (Centrarchidae) were filmed feeding on tethered ghost shrimp (Higham et al., 2006a,b; Norton and Brainerd, 1993; Wainwright et al., 2007). The cichlids *Heros severus* (previously *Cichlasoma severum*) and *Cichla ocellaris* were filmed feeding on live *Daphnia* (Norton and Brainerd, 1993; Wainwright et al., 2001). *Serranocirrhitis latus* and *E. ongus* were recently identified as occupying opposing ends of the ram–suction continuum and sequences from that study were included of these species feeding on live zebra fish (*D. rerio*) (Oufiero et al., 2012). For each species in a pair, we analyzed multiple individuals (2–3 per species) and multiple strikes per individual (7–11 feeding strikes), totaling 169 videos. For each individual, the five fastest strikes (shortest time between strike onset and prey capture) were kept for downstream analyses. Because principal component analyses (PCA) can be skewed by unequal representation between groups, equal numbers of sequences were retained for each species in a pair (e.g. five videos each from two *C. ocellaris* and five videos each from two *H. severus*); in cases where there were extra individuals in one pair, the individual with the slowest mean strike was discarded until representation was equal. Seventy videos were included in the final analyses for ram–suction pairs. Means and standard deviations were first calculated for each individual, which were then used to calculate species means and standard deviations.

High-speed video

All fish were filmed using either a NAC Memrecam ci digital system (Tokyo, Japan) high-speed camera at 500 frames s^{−1} or a Fastec HiSpec 1 system (San Diego, CA, USA) at 1000–2000 frames s^{−1}. Two 120 W halogen lights were used to light the field of view during filming. Fish were not fed for at least 24 h prior to filming and were filmed in their housing tank. Videos were selected for analysis based on these criteria: the predator's entire head and a portion of the body were in view, the predator appeared to

Table 1. Mean (\pm s.d.) time to prey capture and relative contributions of suction, body ram and jaw ram to prey capture

Species	Family	ΔT_{pc} (s)	Body ram	Jaw ram	Suction
<i>Ctenopoma kingsleyae</i>	Anabantidae	0.041	0.527	0.138	0.335
<i>Antennarius hispidus</i>	Antennariidae	0.011	0.122	0.635	0.243
<i>Aulostomus maculatus</i>	Aulostomidae	0.013	0.021	0.938	0.041
<i>Opsanus beta</i>	Batrachoididae	0.017	0.104	0.545	0.351
<i>Pterocaesio pisang</i>	Caesionidae	0.014	0.334	0.502	0.164
<i>Dactylopus dactylopus</i>	Callionymidae	0.022	0.175	0.611	0.214
<i>Caranx sexfasciatus</i>	Carangidae	0.013	0.810	0.157	0.033
<i>Lepomis macrochirus</i>	Centrarchidae	0.018	0.355	0.442	0.204
<i>Aeoliscus strigatus</i>	Centriscidae	0.007	0.082	0.907	0.011
<i>Macroramphosus scolopax</i>	Centriscidae	0.010	0.269	0.677	0.054
<i>Centrogenys vaigiensis</i>	Centrogenyidae	0.008	0.065	0.657	0.278
<i>Lates niloticus</i>	Centropomidae	0.018	0.275	0.502	0.223
<i>Boulengerochromis microlepis</i>	Cichlidae	0.009	0.484	0.240	0.276
<i>Caquetaia kraussi</i>	Cichlidae	0.106	0.241	0.602	0.157
<i>Pterophyllum scalare</i>	Cichlidae	0.015	0.083	0.532	0.385
<i>Oxyrrhites typus</i>	Cirrhitidae	0.010	0.275	0.452	0.273
<i>Datnioides microlepis</i>	Datnioididae	0.038	0.178	0.548	0.273
<i>Butis butis</i>	Eleotridae	0.022	0.731	0.099	0.169
<i>Stigmatogobius pleurostigma</i>	Gobiidae	0.027	0.619	0.193	0.187
<i>Emmelichthys atlanticus</i>	Haemulidae	0.008	0.181	0.677	0.141
<i>Haemulon aurolineatum</i>	Haemulidae	0.009	0.377	0.267	0.356
<i>Haemulon vittatum</i>	Haemulidae	0.008	0.161	0.545	0.294
<i>Pristilepis oligolepis</i>	Holocentridae	0.015	0.367	0.460	0.173
<i>Epibulus insidiator</i>	Labridae	0.030	0.129	0.760	0.111
<i>Ocyurus chrysurus</i>	Lutjanidae	0.025	0.341	0.268	0.391
<i>Malacanthus purpureus</i>	Malacanthidae	0.010	0.376	0.432	0.192
<i>Ptereleotris heteroptera</i>	Microdesmidae	0.016	0.411	0.444	0.145
<i>Nandus nandus</i>	Nandidae	0.094	0.409	0.407	0.184
<i>Oplegnathus fasciatus</i>	Oplegnathidae	0.040	0.798	0.199	0.003
<i>Betta pugnax</i>	Osphronemidae	0.016	0.182	0.393	0.425
<i>Plesiops caeruleatus</i>	Plesiopidae	0.021	0.242	0.532	0.226
<i>Monocirrhus polyacanthus</i>	Polycentridae	0.034	0.182	0.596	0.222
<i>Inimicus didactylus</i>	Scorpaenidae	0.019	0.059	0.683	0.258
<i>Serranocirrhitis latus</i>	Serranidae	0.005	0.101	0.392	0.507
<i>Hypoplectrus puella</i>	Serranidae	0.040	0.529	0.274	0.197
<i>Epinephelus ongus</i>	Serranidae	0.018	0.733	0.148	0.119
<i>Sphyrna barracuda</i>	Sphyrnidae	0.034	0.883	0.047	0.070
<i>Synanceia</i> sp.	Synanceiidae	0.024	0.084	0.615	0.301
<i>Doryrhamphus excisus</i>	Syngnathidae	0.002	0.044	0.784	0.172
<i>Paracentropogon rubripinnis</i>	Tetrarogidae	0.012	0.300	0.527	0.173

ΔT_{pc} , time to prey capture. Contributions of body ram, jaw ram and suction were calculated as proportions.

be oriented nearly perpendicular to the plane of the camera and was in-focus, and prey movement due to swimming or escape maneuvers was minimal. The last criterion is important as these movements can result in apparent negative suction distances in downstream analyses.

Kinematic analysis

For each video, the x,y coordinates of five landmarks were digitized on two frames corresponding to the onset of the strike (T_0) and the time at prey capture (T_{pc}) (Fig. 1). T_0 was defined as one frame before the start of craniofacial movement relative to the body. In most strikes, the first craniofacial movement was the onset of lower jaw depression, but in some fishes the first craniofacial movement was jaw protrusion or hyoid rotation. T_{pc} was defined as the first frame in which the center of mass of the prey passed into the mouth. The difference in T_{pc} and T_0 was used to determine the time to prey capture in seconds (ΔT_{pc}). Points were digitized using the DLTdv3 package in Matlab (Hedrick, 2008), and all downstream analyses on their coordinates were carried out in the R statistical computing platform. Point 1 represented the anterior tip of the upper jaw, point 2 the anterior tip of the lower jaw, point 3 a stationary point on the body such as a scale or spot, point 4 the approximate center of mass of the prey, and point 5 a stationary background point that was used to control for camera movement during the course of the video sequence.

Using the coordinates from these five landmarks at T_0 and T_{pc} , we calculated the contributions of suction, body ram and jaw ram to prey

capture. The contribution of suction to prey capture is the decrease in distance between the predator's mouth and the prey between time T_0 and T_{pc} . We first calculated the distance between point 4 and the midpoint of points 1 and 2 at T_0 , then found the distance between point 4 at T_{pc} and the midpoint of points 1 and 2 at T_0 . The difference between these two distances is the suction contribution to prey capture. Performing the calculation this way focuses on movements of the prey that decrease predator-prey distance in the earthbound frame of reference at T_0 and ensures that escape movements of the prey away from the predator result in negative calculations, which would indicate that the video sequence was not suitable for analysis. We calculated the body ram contribution to prey capture, or how much closer the predator's body moved towards the prey as a result of swimming and coasting between T_0 and T_{pc} . This calculation used the stationary point on the body (point 3) at T_0 and T_{pc} . To control for movement of the prey due to suction, point 4 was only used from time T_0 in this calculation. We calculated the contribution of jaw ram to prey capture independent of body ram and suction. Because any change in the location of the mouth at T_{pc} can be due to body ram and jaw ram, the movement of the body (point 3) was first subtracted from points 1 and 2 at T_{pc} . Then we calculated the change in the distance between the midpoint of the mouth and prey at T_0 from the distance between the translated gape midpoint at T_{pc} and the original location of the prey at T_0 . Finally, the suction, body ram and jaw ram components were totaled, and we calculated their contributions as proportions of the total prey-capture

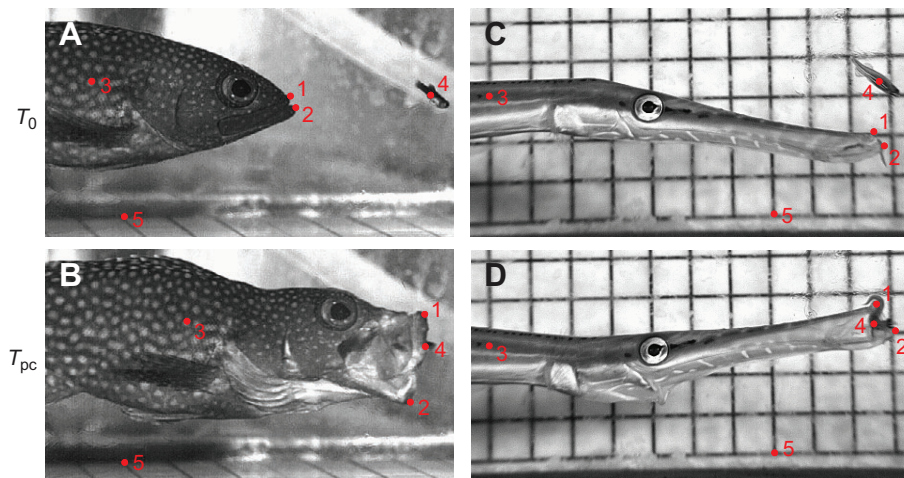


Fig. 1. Kinematic analysis. Five landmarks were digitized at two time points in each feeding sequence, as demonstrated by a ram-ambush predator, *Epinephelus ongus* (A,B), and a pivot-feeder, *Aulostomus maculatus* (C,D). Onset of the strike (T_0 ; A,C) was one frame before the first observed craniofacial movement, and time at prey capture (T_{pc} ; C,D) was the first frame in which the prey passed into the mouth. See Materials and methods for details regarding point placement.

distance. Note that because all contributions were calculated as proportions, our analyses are scale independent, which is a useful characteristic for large comparative studies.

Data visualization and statistical analysis

We used ternary plots to visualize the contribution of suction, body ram and jaw ram to prey capture. Ternary plots are a convenient two-dimensional representation when three variables add to a constant value. In our case, three prey-capture contributions are represented as proportions and therefore all add to one. A ternary plot has three axes oriented at 120 deg with respect to one another. Here, the axes are the suction, body ram and jaw ram independent proportional contributions to prey capture. All ternary plots were created in a custom-modified version of the R package *robCompositions* (Templ et al., 2011) and are shown in the same orientation: the suction axis runs from the midpoint of the bottom edge of the triangle to the top vertex, the body ram axis runs from the midpoint of the left edge of the triangle to the lower right vertex, and the jaw ram axis runs from the midpoint of right edge of the triangle to the lower left vertex.

Robust PCA for compositional data were calculated using a centered log-ratio transformation as implemented in the *pcaCoDa* function in the *robCompositions* R package (Filzmoser et al., 2009; Templ et al., 2011). Note that as body ram, jaw ram and suction contributions to prey capture all add to a constant value, our dataset is two-dimensional and so only two principal component axes were obtained. Principal components plotted into the ternary diagrams appear as curves as a result of the log-ratio transformation used in their computation. For the large acanthomorph dataset, we calculated Pearson's correlation coefficients using the *cor.test* function in R, with a two-tailed P set to 0.05 to see how strike duration (natural logarithm of ΔT_{pc}) was associated with prey-capture strategy variables. For the ram-suction species-pairs dataset, we were primarily interested in determining how different components of prey-capture strategy varied between the species in each pair. We used a nested mixed model ANOVA with species as the fixed independent effect and individual as the random independent factor, as has been done in previous studies (Norton and Brainerd, 1993). Nested mixed models were carried out using the *lmer* function in the *lme4* package in R, and numerator degrees of freedom, denominator degrees of freedom, F -statistics and P -values were calculated with the *anova* function in the *lmerTest* package. We were unable to incorporate phylogenetic information in this analysis because of the lack of published trees including all the species in our dataset. However, we do not find reason to believe that evolutionary history has strongly biased our analysis as closely related species are not necessarily near one another in the ternary plots.

We investigated the sensitivity of our PCA on representative strikes to variation in the choice of a typical strike using a resampling method and the species-pairs dataset for which we had multiple strikes per species (10–15 sequences per species, 70 sequences total). A single video was randomly selected from each of the six species to simulate choosing a representative

strike for a species. This was repeated 10,000 times and we performed PCA as described above to determine the loadings of the jaw ram, body ram and suction proportions on PC1 for each replicate. The resulting distributions of loadings were then compared with PC1 loadings based on species means to judge whether our results are rigorous to variation in representative strike choice

RESULTS

The kinematics of feeding events from 40 species of spiny-rayed fishes are summarized in Table 1. ΔT_{pc} , or the time between the onset of craniofacial movement and prey capture, varied from 0.002 to 0.094 s. Body ram proportion ranged from 0.021 to 0.883 (mean \pm s.d. 0.316 ± 0.24), jaw ram from 0.047 to 0.938 (0.471 ± 0.22) and suction from 0.003 to 0.507 (0.213 ± 0.11). Thus, when considered as a proportion of the total strike distance, body ram and jaw ram had similar, large ranges (0.862 and 0.891, respectively), while the range of suction proportion was smaller (0.503), and the maximum contribution of suction was markedly lower than that of either of the ram components. The 40 species measured fill a large proportion of prey-capture space, as visualized using a ternary plot (Fig. 2), and reached very near the extremes of high jaw ram (lower left vertex) and high body ram (lower right vertex); however, there were no points at or near the extreme suction area of prey-capture space (top vertex). Strikes in which more than half of the strike distance was covered by suction were very rare.

For suction-feeding spiny-rayed fishes, principal component 1 (PC1) explained 86.2% of the variation in the data (Fig. 2), with body ram loading positively and heavily on this axis (0.82), and jaw ram and suction loading negatively with equal magnitude (-0.405 and -0.411 , respectively). PC1 therefore largely reflects the amount of body ram compared with the other strike components. Principal component 2 (PC2) represents the remaining 13.8% of variation in the data. On this axis, jaw ram and suction have strong but opposite loadings of -0.709 and 0.705 , respectively, while body ram is negligible (-0.003).

In the acanthomorph data, the natural logarithm of ΔT_{pc} was significantly correlated with PC1 ($r=0.369$, $P=0.020$; Fig. 3) and with body ram proportion ($r=0.351$, $P=0.026$), and negatively correlated with jaw ram proportion ($r=-0.313$, $P=0.049$). There was no relationship between strike duration and PC2 ($r=0.115$, $P=0.479$) or with suction proportion ($r=-0.120$, $P=0.461$).

The proportions of body ram, jaw ram and suction contributions to prey capture were calculated for six species belonging to three ram-suction species pairs (Table 2). There was variation within

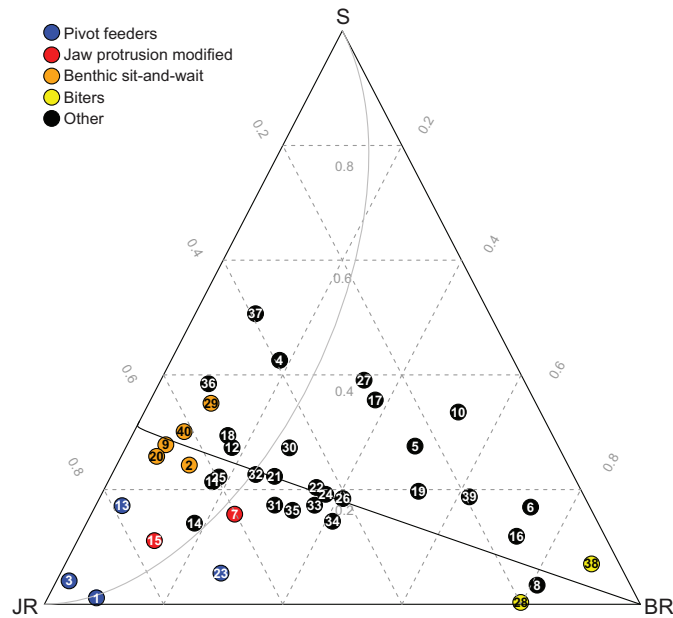


Fig. 2. Prey capture diversity in suction-feeding acanthomorphs. Ternary plot showing the diversity of strike behaviors as determined by the suction proportion (S), jaw ram proportion (JR) and body ram proportion (BR) that contributed to prey capture. Suction distances are constrained across acanthomorphs, and prey-capture diversity does not follow a strict ram–suction trade-off. The first principal component represents a strong trade-off between body ram and the combined contributions of jaw ram and suction and is shown by the black curve. The second principal component is largely a trade-off in jaw ram and suction and is shown by the gray curve. The distribution of acanthomorphs yields new insights into how behavioral and morphological convergence shapes prey-capture diversity, and we have highlighted some of the examples mentioned in the Results (colored circles). Species are numbered as follows: (1) *Aeolisus strigatus*, (2) *Antennarius hispidus*, (3) *Aulostomus maculatus*, (4) *Betta pugnax*, (5) *Boulengerochromis microlepis*, (6) *Butis butis*, (7) *Caquetaia kraussi*, (8) *Caranx sexfasciatus*, (9) *Centrogenys vaigiensis*, (10) *Ctenopoma kingsleyae*, (11) *Dactylopus dactylopus*, (12) *Datnioides microlepis*, (13) *Doryrhamphus excisus*, (14) *Emmelichthyops atlanticus*, (15) *Epibulus insidiator*, (16) *Epinephelus ongus*, (17) *Haemulon aurolineatum*, (18) *Haemulon vittatum*, (19) *Hypoplectrus puella*, (20) *Inimicus didactylus*, (21) *Lates niloticus*, (22) *Lepomis macrochirus*, (23) *Macroramphosus scolopax*, (24) *Malacanthus purpureus*, (25) *Monocirrhus polyacanthus*, (26) *Nandus nandus*, (27) *Ocyurus chrysurus*, (28) *Oplegognathus fasciatus*, (29) *Opsanus beta*, (30) *Oxyrrhites typus*, (31) *Paracentropogon rubripinnis*, (32) *Plesiops caeruleolineatus*, (33) *Pristilepis oligolepis*, (34) *Ptereleotris heteroptera*, (35) *Pterocaesio pisang*, (36) *Pterophyllum scalare*, (37) *Serranocirrhites latus*, (38) *Sphyrna barracuda*, (39) *Stigmatogobius pleurostigma* and (40) *Synanceia* sp.

individuals, between individuals, and between species. Individuals within pairs seemed to cluster as expected, such that ‘high-ram’ species (*M. salmoides*, *C. ocellaris*, *E. ongus*; squares in Fig. 4) had higher body ram, lower jaw ram and lower suction proportions than their ‘high-suction’ counterparts (*L. macrochirus*, *Heros severus*, *S. latus*; circles in Fig. 4).

Nested mixed models for the centrarchid pair found that only body ram proportion was significantly different between *L. macrochirus* and *M. salmoides* (body ram $F_{1,4}=15.4$, $P=0.017$). Jaw ram proportion and suction proportion did not differ between species (jaw ram $F_{1,4}=6.04$, $P=0.070$, suction $F_{1,4}=3.78$, $P=0.124$), which reflects the large amount of overlap along these axes in strikes from both species (Fig. 4A). PC1 for the centrarchid data explained

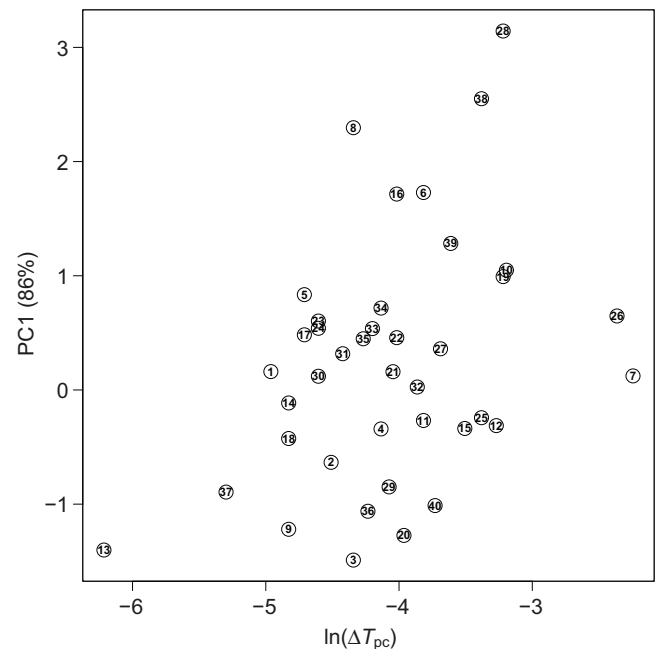


Fig. 3. Strike duration is correlated with PC1 ($r=0.369$, $P=0.020$). Species with smaller values on PC1 (lower body ram proportions and higher combined suction and jaw ram proportions) tend to have quicker strikes overall. Time to prey capture (ΔT_{pc}) was measured in seconds. Species are numbered as in Fig. 2.

86.3% of the variation among strikes; with body ram and suction both loading heavily on PC1, this axis appears to represent a body ram–suction trade-off (body ram 0.717, suction -0.696 , jaw ram -0.021). Nest mixed models showed that there was a significant difference between *L. macrochirus* and *M. salmoides* on PC1 ($F_{1,28}=10.75$, $P=0.003$).

The cichlids (Fig. 4B) were the only ram–suction pair for which all three proportions were significantly different between species (body ram $F_{1,18}=23.4$, $P=0.0001$; jaw ram $F_{1,18}=28.7$, $P=4.3e-05$; suction $F_{1,18}=6.1$, $P=0.024$). PC1 explained 90.8% of the variation in the data, and body ram and suction loaded heavily and oppositely on PC1 (body ram 0.712, suction -0.702 , jaw ram -0.010). *Cichla ocellaris* and *H. severus* were significantly different on PC1 ($F_{1,18}=11.7$, $P=0.003$), reflecting a body ram–suction trade-off underlying the majority of variation in this species pair.

In the serranid pair, only body ram proportion was significantly different between the high-ram (*E. ongus*) and high-suction (*S. latus*) species (body ram $F_{1,2}=25.8$, $P=0.037$; jaw ram $F_{1,2}=4.6$, $P=0.169$; suction $F_{1,2}=8.4$, $P=0.101$). PC1 explained the majority of variation in the data (97.8%), and body ram and suction proportions loaded heavily and oppositely (body ram 0.766, suction -0.627). Though to a small degree, jaw ram also loaded on PC1 in the same direction as suction (jaw ram -0.139). Scores along PC1 were only marginally different between *S. latus* and *E. ongus* ($F_{1,2}=13.9$, $P=0.065$), despite the separation of these species in prey-capture space (Fig. 4C).

A PCA on the means for the six species represented in the ram–suction pairs revealed a major axis of variation for the six species that closely mirrored the results for the 40 species dataset (Fig. 4D). PC1 explained almost all variation in the dataset (99.6%) with body ram loading heavily in one direction and jaw ram and suction loading in the opposite direction (body ram 0.813, jaw ram -0.341 , suction -0.472). We compared these loadings based on species means with the distribution of loadings on PC1 from 10,000

Table 2. Mean (±s.d.) time to prey capture and prey-capture contributions for individuals and species within three ram–suction pairs

	ID	ΔT_{pc} (ms)	Body ram	Jaw ram	Suction		ID	ΔT_{pc} (ms)	Body ram	Jaw ram	Suction
Centrarchidae											
<i>L. macrochirus</i>						<i>M. salmoides</i>					
	1	19.6±3.3	0.23±0.161	0.583±0.091	0.192±0.090		1	62.8±6.1	0.546±0.078	0.247±0.048	0.207±0.071
	2	12.8±2.3	0.122±0.034	0.563±0.074	0.315±0.107		2	30.4±3.8	0.389±0.074	0.422±0.032	0.189±0.062
	3	16.4±3.0	0.284±0.069	0.425±0.040	0.291±0.089		3	48.8±3.6	0.556±0.125	0.372±0.082	0.072±0.052
	sp	16.3±3.4	0.214±0.083	0.524±0.086	0.266±0.065		sp	47.2±16.1	0.497±0.094	0.347±0.090	0.156±0.073
Cichlidae											
<i>H. severus</i>						<i>C. ocellaris</i>					
	1	27.0±6.0	0.358±0.125	0.424±0.103	0.217±0.131		1	015.2±2.5	0.608±0.135	0.247±0.060	0.145±0.092
	2	25.6±11.7	0.290±0.213	0.476±0.121	0.234±0.102		2	18.6±6.7	0.652±0.081	0.252±0.032	0.095±0.056
	sp	26.3±1.0	0.324±0.063	0.450±0.013	0.226±0.020		sp	16.9±2.4	0.630±0.038	0.249±0.020	0.120±0.025
Serranidae											
<i>S. latus</i>						<i>E. ongus</i>					
	1	5.6±0.5	0.167±0.141	0.416±0.052	0.417±0.104		1	12.6±1.5	0.568±0.147	0.311±0.092	0.121±0.065
	2	5.4±0.5	0.090±0.061	0.652±0.083	0.258±0.079		2	14.0±2.0	0.766±0.110	0.184±0.059	0.051±0.052
	sp	5.5±0.1	0.129±0.054	0.534±0.167	0.337±0.112		sp	13.3±1.0	0.667±0.140	0.248±0.090	0.086±0.050

The centrarchid pair (*Lepomis macrochirus*, *Micropterus salmoides*) was filmed feeding on tethered ghost shrimp, the cichlids (*Heros severus*, *Cichla ocellaris*) on live *Daphnia*, and the serranids (*Serranocirrhitis latus*, *Epinephelus ongus*) on live fish prey. ΔT_{pc} , time to prey capture. sp, species mean and s.d. Contributions of body ram, jaw ram and suction were calculated as proportions.

resampled datasets consisting of a single representative strike for each of the six species. Although the resulting distributions (see Fig. S1) are skewed and therefore not amenable to parametric significance testing, the density curves show that the majority of replicates are quantitatively similar to the loadings calculated from species means. PC1 explained the vast majority of variation in replicates (mean 94.6%, median 97.4%) and the mean loadings were: body ram 0.743 (median 0.786), jaw ram −0.251 (median −0.252) and suction −0.491 (median −0.548). PC loadings are

subject to qualitative interpretation, and the distributions indicate that most analyses would support a trade-off between body ram versus the combination of jaw ram and suction. For instance, jaw ram and suction only load in opposite directions in 1790 out of 10,000 replicates (17.9%). There is some tendency for the representative datasets to underestimate the importance of jaw ram; the jaw ram peak is displaced towards zero relative to the species-mean loading (Fig. S1, green line), and there is a relatively high frequency with loadings near zero. Accordingly, the loading of

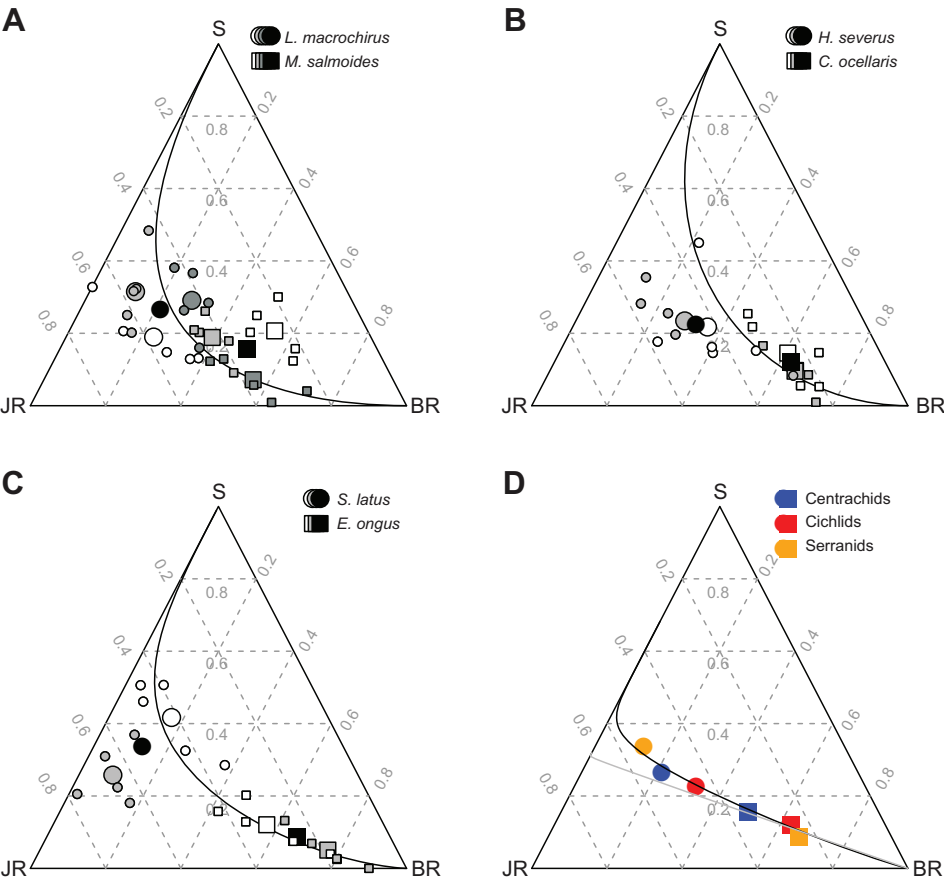


Fig. 4. Comparison of prey capture from species pairs used as exemplars of extremes of the ram–suction continuum in previous studies. (A) Centrarchids, *Lepomis macrochirus* and *Micropterus salmoides*; (B) cichlids, *Heros severus* and *Cichla ocellaris*; and (C) serranids, *Serranocirrhitis latus* and *Epinephelus ongus*. Within each pair, circles represent the suction-dominated species and squares represent the ram-dominated species. Strikes from different individuals are represented by small circles/squares of different shades of white and gray, and individual means are plotted as larger circles/squares of the same shade. Black coloration designates species means. Black curves represent the first principal component for each dataset. (D) Mean prey-capture proportions for each species, plotted for comparison. Note that PC1 (black curve, D) for the species means is similar to PC1 from the acanthomorph dataset (gray curve, D).

suction on PC1 tends to be overestimated in the resampled data compared with the value obtained using species means (Fig. S1, pink line). If these tendencies are true of our method in general, we would expect to be biased towards finding a strict body ram–suction trade-off when using representative strikes instead of species means. However, this is not what we found for our acanthomorph dataset. It should be noted that the variation in the distributions for jaw ram and suction is based on six species, but we would expect variation to be less if the same analysis was performed on the 40 species in our acanthomorph dataset, as a larger sample size will decrease the ability of one arbitrarily chosen strike to significantly influence the PC loadings. Not knowing of any bias in our selection of typical strikes and based on the findings of our resampling method, we conclude that our results regarding the major axis of diversity in acanthomorphs are robust to variations in representative strike choice.

DISCUSSION

The contribution of suction distance to prey capture is greatly limited compared with ram in acanthomorph fishes. The high-suction area of prey-capture space is unoccupied, and the highest contribution of suction to prey-capture distance exhibited by any fish in our dataset was only about half (Fig. 2). In contrast, there were strikes occupying the full range of jaw and body ram proportions. This is the largest published kinematic study to date for suction-feeding fishes in terms of family representation and number of species, and we purposefully included fishes from a range of trophic niches (e.g. benthic invertebrates, zooplankton, fish) and prey-capture behaviors (e.g. water-column zooplanktivores, benthic sit-and-wait predators, pelagic ram-biters). Amongst this diversity, we found scant evidence that evolutionary innovation has surmounted the hydrodynamic constraints imposed on suction distance to bring any species into the extreme suction area of prey-capture space.

Does the absence of suction-dominated strikes in our analysis reflect a constraint on suction feeding or did this feeding mode elude our investigation because it is relatively rare? We suggest that a combination of these factors is responsible. Suction feeders generate flows that are spatially limited, with flow velocity dropping by 95% at a distance of one mouth diameter from the predator (Day et al., 2005). Because suction distances are limited in this way, any forward movement of the mouth aperture during the strike by swimming or rotation of cranial linkages is likely to make a significant contribution to prey-capture distance. Suction-dominated strikes require that the body and mouth do not advance toward the prey during the strike. One might expect to see this feeding mode in sit-and-wait predators that strike from a position resting on the substratum. However, the representatives of this feeding mode that we studied all used considerable jaw protrusion to close in on their prey (see below).

Because our study was limited to acanthomorphs feeding on mobile prey, it is possible that there are other taxa or prey types that could exhibit feeding strikes with high suction proportions. Jaw protrusion, while a synapomorphy of spiny-rayed fishes, is not a universal trait of suction feeders (Wainwright et al., 2015). Perhaps there are non-acanthomorph fishes or other aquatic vertebrates lacking jaw protrusion that have evolved strategies to feed on evasive prey using high proportions of suction. However, even these taxa may generate ram by sucking themselves toward the prey (Summers et al., 1998). Furthermore, some of these lineages have independently evolved jaw protrusion (Wilga et al., 2007; Wainwright et al., 2015) and many use large amounts of body ram to lunge forward at the last moment, even if their prey-capture

strategies appear to be sit-and-wait. We also note that our study focused on strikes at mobile prey. When approaching prey that cannot escape, fishes can move to within less than a mouth diameter before initiating the strike, because there is no risk of disturbing the prey into an escape response. Such a strike could potentially reach the suction-dominated region of the continuum.

Our findings challenge the traditional view that a fundamental trade-off between ram and suction underlies the diversity of prey-capture behaviors in suction-feeding fishes. Across spiny-rayed fishes, PC1 represented a strong trade-off between body ram and the combined contribution of suction and jaw ram; suction and jaw ram load in the same direction and with near-equal magnitudes, such that the major axis of variation is not simply a ram–suction continuum. Instead, variation is better described by the relative amount of body ram involved in the strike, or a continuum between low and high body ram, as confirmed by the strong correlation between PC1 and body ram proportion. Combined with the apparent lack of suction-specialized strikes, our results corroborate previous studies that questioned the role of suction in generating diversity in prey-capture distance (Wainwright et al., 2001; Ferry et al., 2015).

Simultaneously comparing the contributions of jaw ram, body ram and suction clarifies the importance of jaw ram in the diversification of prey-capture strategies among acanthomorph fishes. Body ram and jaw ram are often combined into a measurement of total ram, which implies that they function similarly and trade-off with suction in a comparable manner. However, PC1 reveals that jaw ram and suction combined trade-off with the relative amount of body ram in a strike. This sets jaw ram apart from body ram and illustrates that jaw ram provides a separate axis along which to generate variation in prey capture. The interaction between suction and jaw ram appears to be particularly important in strikes with low contributions of body ram: close-range strikes where both jaw ram and suction have the opportunity to make large proportional contributions to prey capture. We expect the synergistic effect of jaw protrusion on suction forces (Holzman et al., 2012) to be most important in close-range strikes where a predator has less distance to accelerate its mouth opening but must still approach the prey fast enough to capture it despite its attempts at escape. Indeed, many high jaw ram strikes in our dataset are from zooplanktivores and sit-and-wait predators that are known to only strike at close range. The close relationship between PC1 and time to prey capture also suggests that strikes dominated by short-range suction and jaw ram components are quicker than strikes relying more on body ram (Fig. 3).

We found that some species with reputations as suction specialists due to their rapid and powerful strikes are actually relying on jaw ram more than suction (or body ram) to decrease the distance between their mouth and prey. Pivot-feeding seahorses and pipefish were identified as high jaw ram feeders previously (Flammang et al., 2009), but we found that syngnathiforms as a whole are specialized jaw ram feeders and include the most extreme jaw ram strikes in this study (Fig. 2, blue). In fact, trumpetfish (*A. maculatus*) and shrimpfish (*A. strigatus*) exhibited higher proportions of jaw ram than the slingjaw wrasse (*E. insidiator*), which holds the record for the highest jaw protrusion relative to head length among fishes (Westneat and Wainwright, 1989). Frogfish (*A. hispidus*) are another group of fish with very rapid feeding (Grobeck and Pietsch, 1979) that we found have greater jaw ram than suction distances during prey capture. In fact, all the benthic sit-and-wait predators included in our study grouped closely together in prey-capture space (Fig. 2, orange). Although these fish are generally cryptic and may have large upturned mouths, they are morphologically and taxonomically diverse, in our dataset

representing five families (Antennariidae, Batrachoididae, Centrogeniidae, Scorpaenidae, Synanceiidae) that have converged in kinematics. Predators like frogfish and pipefish may seem like evolutionary oddballs, but this study suggests that these are the type of fish that we should be studying to learn more about the interaction between jaw ram and the ability to produce fast, powerful suction (Van Wassenbergh et al., 2013).

In contrast to suction, a combination of morphological and behavioral adaptations have allowed fish to invade the extreme jaw ram area of morphospace. These include the slingjaw wrasse, *E. insidiator* (Fig. 2, number 15), which has evolved a novel linkage allowing unusually high jaw protrusion (Westneat and Wainwright, 1989; Westneat, 1991; Ferry-Graham et al., 2001a,b; Waltzek and Wainwright, 2003). Syngnathiforms (Fig. 2, blue) use a novel jaw ram mechanism referred to as pivot feeding, which relies on rapid rotation of their head and long snout (Bergert and Wainwright, 1997; de Lussanet and Muller, 2007; Van Wassenbergh et al., 2008; Flammang et al., 2009; Roos et al., 2009). At least some syngnathiforms power-amplify this pivoting motion using tendon elastic recoil (Van Wassenbergh et al., 2008, 2009), which is the only known elastic recoil feeding mechanism in fishes.

However, novelty in one trait does not guarantee that a species will have extreme strikes as morphology and behavior interact to produce kinematics. This is demonstrated by the cichlid *C. kraussi*, which is from a lineage known to have a modified suspensorial linkage allowing high jaw protrusion relative to other cichlids (Waltzek and Wainwright, 2003). *Caquetaia kraussi* (Fig. 2, number 7) may be extreme when compared with cichlids, but this individual's strike was not particularly unusual when compared across acanthomorphs and had a jaw ram proportion similar to or less than 12 other species. Therefore, adaptations for high jaw protrusion alone do not make a strike extreme. In contrast, some benthic sit-and-wait predators and water-column zooplanktivores, such as *I. didactylus*, and *E. atlanticus* (Fig. 2, numbers 20 and 14) had relatively extreme kinematics and high values of jaw protrusion (as much as 48% of head length in *E. atlanticus*) without any obvious morphological innovations. Therefore, while a combination of behavior and unusual morphology are necessary for acanthomorphs to become extreme jaw ram specialists, many acanthomorph clades have achieved relatively high jaw ram prey-capture behaviors despite potential biomechanical and kinematic constraints on jaw function and strike distance.

Specific pairs of closely related species were used to illustrate a trade-off in ram and suction contributions to prey capture. Even with jaw ram as a separate source of variation, we found strong evidence that species within the centrarchid and cichlid pairs fell out along a ram–suction continuum. This is at odds with our findings from the larger sample of acanthomorph diversity where we did not recover a simple continuum between body ram and suction. We caution that focusing only on ram–suction pairs gives a skewed interpretation of the diversity of prey-capture strategies in acanthomorph fishes. By looking at a larger taxonomic and ecomorphological sample of acanthomorph suction feeders and incorporating another source of variation in prey-capture behavior, we show that the apparent ram–suction trade off in closely related pairs of fishes does not govern feeding diversity at broader evolutionary scales. In agreement with this conclusion, PC1 for the six species averages converges on the same axis found in the large-scale acanthomorph study (Fig. 4D). This suggests that as you add diversity, a strict body-ram versus suction continuum breaks down, and jaw ram and suction contributions combined trade-off with changes in body ram.

Our findings highlight the importance of considering ram, especially body ram, when studying how suction-feeding fishes diversify across feeding niches on evolutionary time scales. It is worth pointing out that laboratory studies that record feeding events may greatly underestimate the maximum body ram that some species can exhibit, because laboratory feeding arenas are quite cramped compared with most natural settings. Also, methods that select for highly motivated strikes based on time to prey capture may also underestimate ram, because strikes with greater ram distances tend to increase prey-capture times despite high ram speeds (Fig. 3; Tran et al., 2010). Further studies on locomotion during predator–prey interactions, and how different ram strategies affect feeding accuracy and suction performance, will be important in developing a better understanding of the diversity of prey-capture strategies (Kane and Higham, 2014, 2015; Rice et al., 2008). Additionally, predator–prey interactions generally involve unsteady swimming modes, such as fast starts and quick turns (Harper et al., 1991; Domenici, 2001), which may not be best characterized by body ram speed as reported in most fish feeding studies. A better understanding of locomotor performance of predators in the context of prey capture will be important in understanding what behavioral options are available to suction-feeding fishes and how locomotion is modified during evolution to enhance prey-capture performance.

While relative measures of suction, body ram and jaw ram reveal new insights into how acanthomorph fish diversify their prey-capture strategy at broad evolutionary scales, the use of proportions can be misleading and provides an incomplete view of attack strategies. For instance, *Monocirrhus polyacanthus* (Fig. 2, number 25) is a freshwater fish that blends in among leaves and branches in the water column before striking with rapid forward jaw protrusion and is located in prey-capture space almost on top of *D. dactylopus* (Fig. 2, number 11), a forager that hovers along the benthos before striking with rapid ventral jaw protrusion. Our study found that both fish rely largely on jaw ram (approximately 60%), but obscures other differences in prey-capture strategy and kinematics that would become apparent with other metrics. Using proportions also tends to downplay strikes that employ a combination of approaches and can lead to the inference that two strikes are similar even though the absolute distances covered may be very different. Depending on the question being asked, absolute instead of relative measures may better characterize the difference in strikes between species (Wainwright et al., 2001).

The distance from which fishes draw prey into their mouth during feeding is significant to predator–prey encounters, but should not be interpreted as a measure of suction performance (Wainwright et al., 2007). The high accelerations reached by suction flows may be spatially restricted and temporally ephemeral, but suction forces over these small distances are crucial to successfully capture evasive aquatic prey (Holzman and Wainwright, 2009; Yen et al., 2015). Suction also serves an important role in prey transport within the oral cavity that was not captured by this study. For instance, we found that trumpetfish (Fig. 2, number 3) are extreme jaw ram feeders that use head rotation to place their jaws very close to prey. However, once prey pass into the mouth, suction continues to transport the prey down what is essentially a long sealed tube. In the sequence included in this study, the distance that prey traveled down the snout after prey capture was more than 52 times greater than the distance suction moved prey outside the mouth. Intraoral transport can also be important in high-ram and ram-biting fish, which often show large excursions of the hyoid and prolonged hyoid depression and can use suction to position prey during swallowing after capture (Liem, 1990; Porter and Motta, 2004; Gibb and Ferry-Graham,

2005; Tran et al., 2010). Suction distances outside the mouth may be constrained, so the diversity of suction among suction feeders may lie along axes that are rarely explored, including the ability to generate strong suction pressure gradients, high fluid accelerations, and the volume of water ingested during the strike (Nemeth, 1997; Higham et al., 2006a,b; Van Wassenbergh et al., 2006; Wainwright et al., 2007; Motta et al., 2008; Kane and Higham, 2015).

Acknowledgements

We thank Will Seah (Pan Ocean, CA, USA) and Forest Young (Dynasty Marine, FL, USA) for help obtaining many species used in this study. To the postdocs, graduate students and undergraduates who over many years helped collect the videos used in this project, we thank you for your time, effort and perseverance. We are grateful to Bruce Draper (UC Davis) for providing live evasive prey.

Competing interests

The authors declare no competing or financial interests.

Author contributions

S.J.L. and P.C.W. designed the research approach. S.J.L., M.D.G., C.E.O. and T.B.W. collected data (high-speed video). S.J.L. analyzed the data. C.E.O. assisted with statistical analyses. S.J.L. and P.C.W. wrote the paper with input from M.D.G., C.E.O. and T.B.W.

Funding

This research was supported by National Science Foundation (NSF) Grant IOS-0924489, the NSF Graduate Research Fellowship Program under Grant No. 1148897, and the Center for Population Biology at the University of California, Davis.

Supplementary information

Supplementary information available online at <http://jeb.biologists.org/lookup/suppl/doi:10.1242/jeb.129015/-DC1>

References

- Bergert, B. A. and Wainwright, P. C. (1997). Morphology and kinematics of prey capture in the syngnathid fishes *Hippocampus erectus* and *Syngnathus floridae*. *Mar. Biol.* **127**, 563–570.
- Cook, A. (1996). Ontogeny of feeding morphology and kinematics in juvenile fishes: a case study of the cottid fish *Clinocottus analis*. *J. Exp. Biol.* **199**, 1961–1971.
- Day, S. W., Higham, T. E., Cheer, A. Y. and Wainwright, P. C. (2005). Spatial and temporal patterns of water flow generated by suction-feeding bluegill sunfish *Lepomis macrochirus* resolved by Particle Image Velocimetry. *J. Exp. Biol.* **208**, 2661–2671.
- de Jong, M. C., Sparenberg, J. A. and de Vries, J. (1987). Some aspects of the hydrodynamics of suction feeding in fish. *Fluid Dyn. Res.* **2**, 87–112.
- de Lussanet, M. H. E. and Muller, M. (2007). The smaller your mouth, the longer your snout: predicting the snout length of *Syngnathus acus*, *Centriscus scutatus* and other pipette feeders. *J. R. Soc. Interface* **4**, 561–573.
- Domenici, P. (2001). The scaling of locomotor performance in predator–prey encounters: from fish to killer whales. *Comp. Biochem. Physiol. A Mol. Integr. Physiol.* **131**, 169–182.
- Ferry, L. A., Konow, N. and Gibb, A. C. (2012). Are kissing gourami specialized for substrate-feeding? Prey capture kinematics of *Helostoma temminckii* and other Anabantoid fishes. *J. Exp. Zool. A Ecol. Genet. Physiol.* **317**, 571–579.
- Ferry, L. A., Paig-Tran, E. M. and Gibb, A. C. (2015). Suction, ram, and biting: deviations and limitations to the capture of aquatic prey. *Integr. Comp. Biol.* **55**, 97–109.
- Ferry-Graham, L. A., Wainwright, P. C., Hulsey, C. D. and Bellwood, D. R. (2001a). Evolution and mechanics of long jaws in butterflyfishes (Family Chaetodontidae). *J. Morphol.* **248**, 120–143.
- Ferry-Graham, L. A., Wainwright, P. C. and Bellwood, D. R. (2001b). Prey capture in long-jawed butterflyfishes (Chaetodontidae): the functional basis of novel feeding habits. *J. Exp. Mar. Biol. Ecol.* **256**, 167–184.
- Ferry-Graham, L. A., Wainwright, P. C. and Lauder, G. V. (2003). Quantification of flow during suction feeding in bluegill sunfish. *Zoology* **106**, 159–168.
- Filzmoser, P., Hron, K. and Reimann, C. (2009). Principal component analysis for compositional data with outliers. *Environmetrics* **20**, 621–632.
- Flammang, B. E., Ferry-Graham, L. A., Rinewalt, C., Ardizzone, D., Davis, C. and Trejo, T. (2009). Prey capture kinematics and four-bar linkages in the bay pipefish, *Syngnathus leptorhynchus*. *Zoology* **112**, 86–96.
- Gibb, A. (1997). Do flatfish feed like other fishes? A comparative study of percomorph prey-capture kinematics. *J. Exp. Biol.* **200**, 2841–2859.
- Gibb, A. C. and Ferry-Graham, L. (2005). Cranial movements during suction feeding in teleost fishes: Are they modified to enhance suction production? *Zoology* **108**, 141–153.
- Grobecker, D. B. and Pietsch, T. W. (1979). High-speed cinematographic evidence for ultrafast feeding in antennariid anglerfishes. *Science* **205**, 1161–1162.
- Harper, D. G., Blake, R. W. and Harper, B. Y. D. G. (1991). Prey capture and the fast-start performance of northern pike *Esox Lucius*. *J. Exp. Biol.* **155**, 175–192.
- Hedrick, T. L. (2008). Software techniques for two- and three-dimensional kinematic measurements of biological and biomimetic systems. *Bioinspir. Biomim.* **3**, 034001.
- Higham, T. E. (2007). Feeding, fins and braking maneuvers: locomotion during prey capture in centrarchid fishes. *J. Exp. Biol.* **210**, 107–117.
- Higham, T. E., Day, S. W. and Wainwright, P. C. (2005). Sucking while swimming: evaluating the effects of ram speed on suction generation in bluegill sunfish *Lepomis macrochirus* using digital particle image velocimetry. *J. Exp. Biol.* **208**, 2653–2660.
- Higham, T. E., Day, S. W. and Wainwright, P. C. (2006a). The pressures of suction feeding: the relation between buccal pressure and induced fluid speed in centrarchid fishes. *J. Exp. Biol.* **209**, 3281–3287.
- Higham, T. E., Day, S. W. and Wainwright, P. C. (2006b). Multidimensional analysis of suction feeding performance in fishes: fluid speed, acceleration, strike accuracy and the ingested volume of water. *J. Exp. Biol.* **209**, 2713–2725.
- Holzman, R. and Wainwright, P. C. (2009). How to surprise a copepod: strike kinematics reduce hydrodynamic disturbance and increase stealth of suction-feeding fish. *Limnol. Oceanogr.* **54**, 2201–2212.
- Holzman, R., Day, S. W., Mehta, R. S. and Wainwright, P. C. (2008). Jaw protrusion enhances forces exerted on prey by suction feeding fishes. *J. R. Soc. Interface* **5**, 1445–1457.
- Holzman, R., Collar, D. C., Mehta, R. S. and Wainwright, P. C. (2011). Functional complexity can mitigate performance trade-offs. *Am. Nat.* **177**, E69–E83.
- Holzman, R., Collar, D. C., Mehta, R. S. and Wainwright, P. C. (2012). An integrative modeling approach to elucidate suction-feeding performance. *J. Exp. Biol.* **215**, 1–13.
- Kane, E. A. and Higham, T. E. (2014). Modelled three-dimensional suction accuracy predicts prey capture success in three species of centrarchid fishes. *J. R. Soc. Interface* **11**, 20140223.
- Kane, E. A. and Higham, T. E. (2015). Complex systems are more than the sum of their parts: using integration to understand performance, biomechanics, and diversity. *Integr. Comp. Biol.* **55**, 146–165.
- Kerfoot, J. R. and Turingan, R. G. (2011). Similarity and disparity in prey-capture kinematics between the invasive pike killifish (*Belonesox belizanus*) and the native Florida largemouth bass (*Micropterus floridanus*). *Fla. Sci.* **74**, 137–150.
- Liem, K. F. (1973). Evolutionary strategies and morphological innovations: Cichlid Pharyngeal jaws. *Syst. Zool.* **22**, 425–441.
- Liem, K. F. (1980a). Adaptive significance of intra- and interspecific differences in the feeding repertoires of cichlid fishes. *Am. Zool.* **20**, 295–314.
- Liem, K. F. (1980b). Acquisition of energy by teleosts: adaptive mechanisms and evolutionary patterns. In *Environmental Physiology of Fishes* (ed. M. A. Ali), pp. 299–334. New York: Plenum Publishing Cooperation.
- Liem, K. F. (1990). Aquatic versus terrestrial feeding modes: possible impacts on the trophic ecology of vertebrates. *Integr. Comp. Biol.* **30**, 209–221.
- Motta, P. J. (1984). Mechanics and functions of jaw protrusion in teleost fishes: a review. *Copeia* **1984**, 1–18.
- Motta, P. J., Hueter, R. E., Tricas, T. C., Summers, A. P., Huber, D. R., Lowry, D., Mara, K. R., Matott, M. P., Whitenack, L. B. and Wintzer, A. P. (2008). Functional morphology of the feeding apparatus, feeding constraints, and suction performance in the nurse shark *Ginglymostoma cirratum*. *J. Morphol.* **269**, 1041–1055.
- Muller, M. and Osse, J. W. M. (1984). Hydrodynamics of suction feeding in fish. *Trans. Zool. Soc. Lond.* **37**, 51–135.
- Muller, M., Osse, J. W. M. and Verhagen, J. H. G. (1982). A quantitative hydrodynamical model of suction feeding in fish. *J. Theor. Biol.* **95**, 49–79.
- Nemeth, D. H. (1997). Modulation of attack behavior and its effect on feeding performance in a trophic generalist fish, *Hexagrammos decagrammus*. *J. Exp. Biol.* **200**, 2155–2164.
- Norton, S. F. (1991). Capture success and diet of cottid fishes: the role of predator morphology and attack kinematics. *Ecology* **72**, 1807–1819.
- Norton, S. F. (1995). A functional approach to ecomorphological patterns in cottid fishes. *Environ. Biol. Fishes* **72**, 1807–1819.
- Norton, S. F. and Brainerd, E. L. (1993). Convergence in the feeding mechanics of ecomorphologically similar species in the Centrarchidae and Cichlidae. *J. Exp. Biol.* **176**, 11–29.
- Nyberg, D. W. (1971). Prey capture in the Largemouth Bass. *Am. Midl. Nat.* **86**, 128–144.
- Osse, J. W. M. (1985). Jaw protrusion, an optimization of the feeding apparatus of teleosts? *Acta Biotheor.* **34**, 219–232.
- Oufiero, C. E., Holzman, R. A., Young, F. A. and Wainwright, P. C. (2012). New insights from serranid fishes on the role of trade-offs in suction-feeding diversification. *J. Exp. Biol.* **215**, 3845–3855.
- Porter, H. T. and Motta, P. J. (2004). A comparison of strike and prey capture kinematics of three species of piscivorous fishes: Florida gar (*Lepisosteus platyrhincus*), redbfin needlefish (*Strongylura notata*), and great barracuda (*Sphyrna barracuda*). *Mar. Biol.* **145**, 989–1000.

- Rice, A. N., Cooper, W. J. and Westneat, M. W. (2008). Diversification of coordination patterns during feeding behaviour in cheilne wrasses. *Biol. J. Linn. Soc.* **93**, 289–308.
- Roberts, T. J. and Azizi, E. (2011). Flexible mechanisms: the diverse roles of biological springs in vertebrate movement. *J. Exp. Biol.* **214**, 353–361.
- Roos, G., Van Wassenbergh, S., Herrel, A. and Aerts, P. (2009). Kinematics of suction feeding in the seahorse *Hippocampus reidi*. *J. Exp. Biol.* **212**, 3490–3498.
- Staab, K. L., Ferry, L. A. and Hernandez, L. P. (2012). Comparative kinematics of cypriniform premaxillary protrusion. *Zoology* **115**, 65–77.
- Summers, A. P., Darouian, K. F., Richmond, A. M. and Brainerd, E. L. (1998). Kinematics of aquatic and terrestrial prey capture in *Terrapene carolina*, with implications for the evolution of feeding in cryptodire turtles. *J. Exp. Biol.* **4**, 280–287.
- Tavera, J., Acero, P. A., Balart, E. F. and Bernardi, G. (2012). Molecular phylogeny of grunts (Teleostei, Haemulidae), with an emphasis on the ecology, evolution, and speciation history of New World species. *BMC Evol. Biol.* **12**, 57.
- Templ, M., Hron, K. and Filzmoser, P. (2011). robCompositions: an R-package for robust statistical analysis of compositional data. In *Compositional Data Analysis. Theory and Applications* (ed. E. Pawlowsky-Glahn and A. Buccianti), pp. 341–355. Chichester, UK: John Wiley & Sons.
- Tran, H. Q., Mehta, R. S. and Wainwright, P. C. (2010). Effects of ram speed on prey capture kinematics of juvenile Indo-Pacific tarpon, *Megalops cyprinoides*. *Zoology* **113**, 75–84.
- Van Leeuwen, J. L. (1984). A quantitative study of flow in prey capture by Rainbow trout, *Salmo gairdneri* with general consideration of the actinopterygian. *Trans. Zool. Soc. Lond.* **37**, 171–227.
- Van Leeuwen, J. L. and Muller, M. (1984). Optimum sucking techniques for predatory fish. *Trans. Zool. Soc. Lond.* **37**, 137–169.
- Van Wassenbergh, S., Aerts, P. and Herrel, A. (2006). Hydrodynamic modelling of aquatic suction performance and intra-oral pressures: limitations for comparative studies. *J. R. Soc. Interface* **3**, 507–514.
- Van Wassenbergh, S., Strother, J. A., Flammang, B. E., Ferry-Graham, L. A. and Aerts, P. (2008). Extremely fast prey capture in pipefish is powered by elastic recoil. *J. R. Soc. Interface* **5**, 285–296.
- Van Wassenbergh, S., Roos, G., Genbrugge, A., Leysen, H., Aerts, P., Adriaens, D. and Herrel, A. (2009). Suction is kid's play: extremely fast suction in newborn seahorses. *Biol. Lett.* **5**, 200–203.
- Van Wassenbergh, S., Leysen, H., Adriaens, D. and Aerts, P. (2013). Mechanics of snout expansion in suction-feeding seahorses: musculoskeletal force transmission. *J. Exp. Biol.* **216**, 407–417.
- Wainwright, P. C., Ferry-Graham, L. A., Waltzek, T. B., Carroll, A. M., Hulsey, C. D. and Grubich, J. R. (2001). Evaluating the use of ram and suction during prey capture by cichlid fishes. *J. Exp. Biol.* **204**, 3039–3051.
- Wainwright, P., Carroll, A. M., Collar, D. C., Day, S. W., Higham, T. E. and Holzman, R. A. (2007). Suction feeding mechanics, performance, and diversity in fishes. *Integr. Comp. Biol.* **47**, 96–106.
- Wainwright, P. C., McGee, M. D., Longo, S. J. and Hernandez, L. P. (2015). Origins, innovations, and diversification of suction feeding in vertebrates. *Integr. Comp. Biol.* **55**, 134–145.
- Waltzek, T. B. and Wainwright, P. C. (2003). Functional morphology of extreme jaw protrusion in neotropical cichlids. *J. Morphol.* **257**, 96–106.
- Weih, D. (1980). Hydrodynamics of suction feeding of fish in motion. *J. Fish Biol.* **16**, 425–433.
- Westneat, M. W. (1991). Linkage Biomechanics and evolution of the unique feeding mechanism of *Epibulus insidiator* (Labridae: Teleostei). *J. Exp. Biol.* **159**, 165–184.
- Westneat, M. W. and Wainwright, P. C. (1989). Feeding mechanism of *Epibulus insidiator* (Labridae: Teleostei): evolution of a novel functional system. *J. Morphol.* **202**, 129–150.
- Wilga, C. D., Motta, P. J. and Sanford, C. P. (2007). Evolution and ecology of feeding in elasmobranchs. *Integr. Comp. Biol.* **47**, 55–69.
- Wintzer, A. P. and Motta, P. J. (2005). A comparison of prey capture kinematics in hatchery and wild *Micropterus salmoides floridanus*: effects of ontogeny and experience. *J. Fish Biol.* **67**, 409–427.
- Yen, J., Murphy, D. W., Fan, L. and Webster, D. R. (2015). Sensory-Motor Systems of copepods involved in their escape from suction feeding. *Integr. Comp. Biol.* **55**, 121–133.

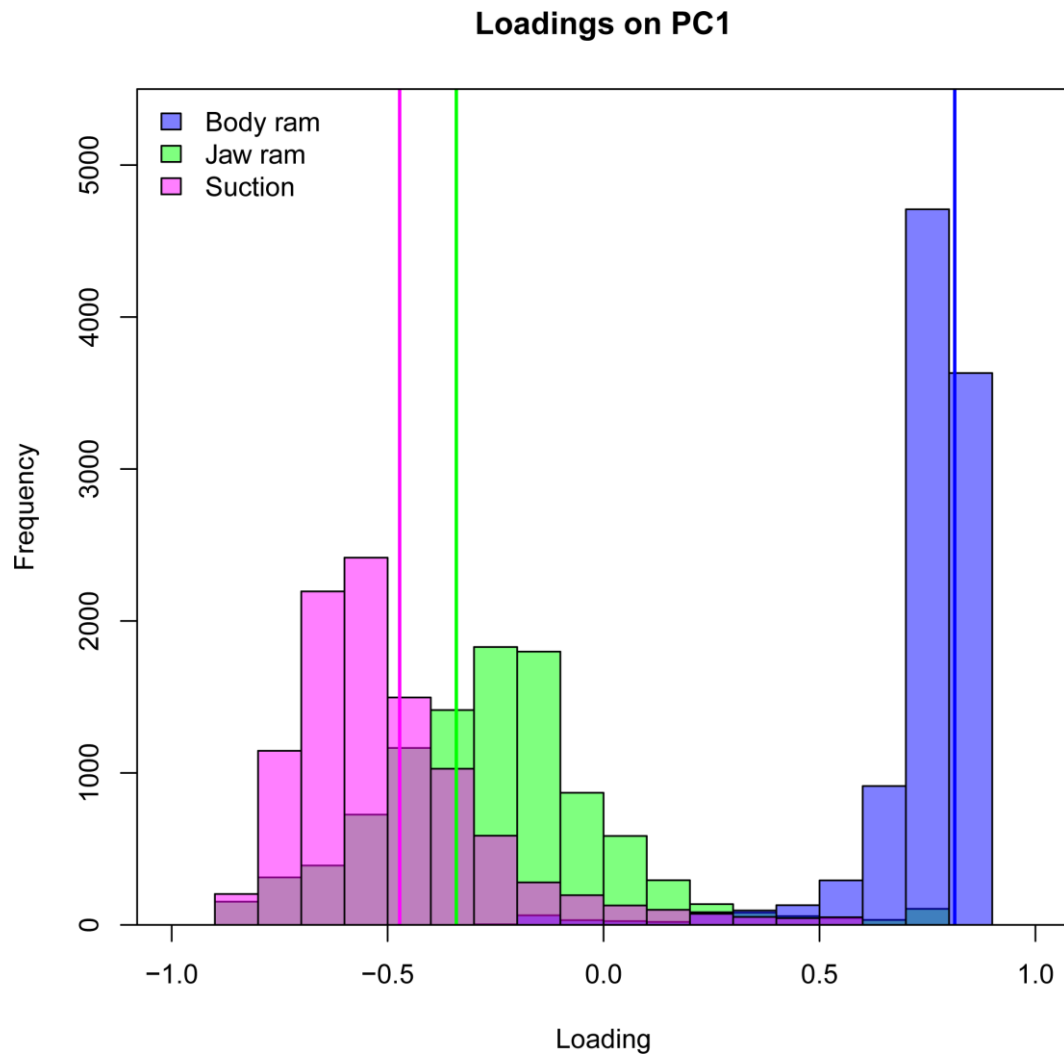


Fig. S1. Histograms of the loadings of prey capture contributions on PC1 from 10,000 replicate datasets with a representative strike per species compared to PC1 loadings obtained from species means (lines). Our resampling analysis shows that, in general, datasets composed of representative strikes for each of the 6 species agree with the analysis performed on species means calculated from multiple individuals per species and multiple strikes per individual (Fig 4D). We would expect there to be less variation in loadings, especially for jaw ram (green histogram) and suction (pink histogram), in our acanthomorph dataset composed of forty species instead of six. The body ram loadings are shown in blue.