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# A faster escape does not enhance survival in zebrafish larvae

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## Abstract

An escape response is a rapid maneuver used by prey to evade predators. Performing this maneuver at greater speed, in a favorable direction, or from a longer distance have been hypothesized to enhance the survival of prey, but these ideas are difficult to test experimentally. We examined how prey survival depends escape kinematics through a novel combination of experimentation and mathematical modeling. This approach focused on zebrafish (*Danio rerio*) larvae under predation by adults and juveniles of the same species. High-speed 3D kinematics were used to track the body position of prey and predator and to determine the probability of behavioral actions by both fish. These measurements provided the basis for an agent-based probabilistic model that simulated the trajectories of the animals. Predictions of survivorship by this model were found by Monte-Carlo simulations to agree with our observations and we examined how these predictions varied by changing individual model parameters. Contrary to expectation, we found that survival may not be improved by increasing the speed or altering the direction of the escape. Rather, zebrafish larvae operate with sufficiently high locomotor performance due to the relatively slow approach and limited range of suction feeding by the fish predators. We did find that survival was enhanced when prey responded from a greater distance. This is an ability that depends on the capacity of the visual and lateral line systems to detect a looming threat. Therefore, performance in sensing, and not locomotion, is decisive for improving the survival of larval fish prey. These results offer a framework for understanding the evolution of predator-prey strategy that may inform prey survival in a broad diversity of animals.

## 1. Introduction

An escape response allows prey to evade predators with rapid locomotion [1]. Because of its potential to directly affect survivorship, natural selection may favor animals that can execute an escape response with high locomotor performance. Indeed, the physiology and mechanics of locomotion features many traits that appear to be adaptations for a rapid response and fast motion. Escape responses are controlled by large-diameter command neurons (e.g. the giant axon of squid [2]), which often recruit specialized muscles (e.g. the axial musculature of fish [3]), which may animate an appendage that is dedicated to escape behavior (e.g. the uropods of crayfish [4]). Prey may direct this escape in an optimal direction [5], or may alternatively benefit from moving randomly so as not to be predictable [6,7]. However, it does not necessarily follow that any enhancement in speed or variation in heading will have a positive effect on a prey's survival. Fish

33 predators commonly approach their prey at a relatively slow speed [8,9] and this could permit  
34 escape by a prey operating below its maximal capacity. The aim of the present study was to test  
35 whether improvements in kinematics related to locomotor performance and sensing affect prey  
36 survival by examining predator-prey interactions in zebrafish (*Danio rerio*).

37 We addressed this aim with a novel approach that combines experimentation with  
38 mathematical modeling. Our methodology was developed to meet the challenges to  
39 understanding the coupled dynamics of predators and prey. Due to the sensing of both animals,  
40 the actions of the prey may (or may not) be a response to the predator, which may (or may  
41 not) be a response to prior motion by the prey. Regression analysis is generally insensitive to  
42 such interdependency, yet may succeed in resolving dominant features of successful prey [10]  
43 or predators [11]. It has additionally been helpful to study behavioral responses to an artificial  
44 predator or prey that is experimentally controlled and therefore not coupled to the actions of the  
45 animal [11–15]. An alternative approach has attempted to formulate a behavioral algorithm of one  
46 animal by considering their responses to the measured kinematics of the other [16]. Our recent  
47 approach similarly included measurements of predator-prey kinematics, but these were used as  
48 a basis for an agent-based probabilistic model that calculated the trajectories of both animals  
49 from a series of behavioral actions. A series of simulations by this model allowed for a predictive  
50 consideration of the effects of differences in behavior on prey survival.

51 Zebrafish provide an excellent experimental subject for studying the mechanisms of predator-  
52 prey interactions. The larval stage of this species serves as a model for the neurophysiological [17–  
53 19] and biomechanical [20,21] basis of behavior. Predator-prey interactions may be experimentally  
54 replicated in the lab, where adults and juvenile zebrafish strike at larval zebrafish with suction  
55 feeding and the larvae respond with a fast-start escape response [22]. Therefore, the interactions  
56 between zebrafish of different life-history stages replicate the principle predator and prey  
57 behaviors that characterize a broad diversity of piscivorous interactions [5,10]. When approaching  
58 an evasive prey, adult zebrafish move much slower than their maximum speed [22], which is  
59 common among suction-feeding fishes [8,9]. A slow approach presumably allows greater control  
60 over the direction and timing of the suction feeding, which is limited to a brief duration over  
61 a small region in front of the mouth [23,24]. The prey, by contrast, respond with an explosive  
62 escape response with speed that generally exceeds that of the predator. As suggested by prior  
63 experiments [25] and theory [5], the relative speed and size of predator and prey greatly influences  
64 strategy. Therefore, we performed experiments with juvenile and adult predators with a nearly  
65 two-fold difference in body length to examine the effects of scale.

## 2. Material and methods

### (a) Animal husbandry

All experiments were conducted on zebrafish (*Danio rerio*, Hamilton 1922) with larvae (5–7 days post fertilization, dpf) that were preyed upon by older fish of the same species. We performed one set of experiments using adults ( $\geq 9$  months old, Mean  $\pm 1$  SD =  $3.4 \pm 0.5$  cm,  $N = 19$ ) and another using juvenile predators (3–4 months old,  $2.0 \pm 0.4$  cm,  $N = 19$ ). All fish were bred from wild-type (AB line) colonies housed in a flow-through tank system (Aquatic Habitats, Apopka, FL, USA) that was maintained at  $28.5^\circ\text{C}$  on a 14:10 h light:dark cycle. To produce larvae, the fertilized eggs from randomized mating were cultured according to standard techniques [26]. Predators were motivated to feed by fasting for a period of 7–14 days prior to an experiment.

### (b) Kinematics

We arranged the lights and cameras for high-speed recordings of both fish with high-contrast images. A hemispherical aquarium ( $\varnothing = 8.5$  cm) was composed of white acrylic, which served as a translucent diffuser of the IR illumination (940 nm) provided by three lamps (CM-IR200-940, CMVision, Houston, TX, USA), positioned below (Fig. 1a). These lamps provided high-intensity illumination that was invisible to the fish [27], while visible illumination at low intensity was provided by overhead fluorescent lights. Each camera (FASTCAM Mini UX50, Precision Photron Inc., San Diego, CA, USA) was fitted with a 55 mm lens (f/2.8 Micro Nikkon AIS, Nikon Inc., Melville, NY, USA) and positioned at a distance that permitted a view of the entire aquarium. The cameras were angled above the aquarium to allow both fish to be viewed by at least two cameras when the fish were close together. The cameras were synchronized to record at a 1000 fps (at  $1024 \times 1024$  pixels) with a common TTL trigger and controlled with the manufacturer's software (PhotronFASTCAM Viewer).

Predation experiments were performed by recording the swimming of one predator and one prey fish in the aquarium (Fig. 1A). This began by placing the fish on opposite sides of a partition. Following a 15 min acclimation period, we lifted the partition and observed the fish until the predator successfully ingested the prey. Using a end-trigger to the high-speed cameras, we saved recordings from  $\sim 0.5$  s before the first predatory strike and until  $\sim 0.5$  s after the prey was captured.

Our video recordings were used to perform measurements of 3D kinematics. We calibrated the cameras by recording a static body, with 48 landmarks of known relative position, in the center of the aquarium. A direct-linear transform (DLT) was calculated using ‘Digitizing Tools’ software in MATLAB (2015a, MathWorks, Natick, MA, USA) [28] from manually-selected coordinates of these landmarks from the perspective of the three cameras. Using a custom script in MATLAB, we found the body positions of predator and prey fish by selecting landmarks from two camera views and using the DLT to determine the coordinates in 3D space. We used the position of the predator’s two eyes to calculate a mean position that approximated the buccal cavity (Fig. 1a). The posterior margin of the swim bladder was found on the prey’s body, which approximates the center of mass [29]. The initial heading of the prey was approximated by matching an ellipsoid (using the ‘regionprops’ function in MATLAB) to the body of the prey and measuring the angle of the major axis of the ellipsoid. All subsequent heading measurements of the prey was defined as the average angular displacement during an escape relative to the prior heading. We acquired the landmark positions at five key events in each interaction between predator and prey: (1) the initiation of a predator’s approach toward the prey, the (2) opening and (3) closing of the predator’s mouth during a strike, and the (4) initiation and (5) completion of the prey’s escape response.

### (c) Descriptive statistics

Descriptive statistics were used to characterize the probability of actions by the predator and prey during predation experiments. We recorded the predator-specific parameters of the strike distance ( $s$ ), the distance from the prey at which a strike (i.e. a suction feeding event) was initiated, and the strike duration ( $\tau$ ), which was defined as the period between the opening and closing of the mouth during suction feeding. For the prey, we found the reaction distance ( $l$ ), the distance from the predator at which the escape response was initiated. The prey’s kinematics were additionally characterized by the escape angle ( $\theta$ ), the angular change in heading from the resting orientation to the escape path. The escape duration ( $\eta$ ) included the period for all stages of the C-start and subsequent undulatory swimming, until the larva ceased moving. The frequency distribution for each of these parameters was found to be well-approximated by the following log-normal probability density function:

$$f(x) = \frac{1}{x\sigma\sqrt{2\pi}} \exp \left[ -\frac{(\ln(x) - \mu)^2}{2\sigma^2} \right], \quad (2.1)$$

where  $x$  is a particular behavioral parameter ( $s, \tau, l, \theta$ , or  $\eta$ ),  $\mu$  is the log mean, and  $\sigma$  is the log standard deviation. We determined best-fit values for  $\mu$  and  $\sigma$  for each behavioral parameter by maximum-likelihood (the 'fitdist' function in MATLAB) to binned measured values. The bin size was determined by the Freedman-Diaconis rule (implemented by the 'histcount' function in MATLAB), which yielded a number of samples per bin of around 17 measurements. Parameters for the prey were not included in the dataset in instances where the prey was captured.

The probability that the strike of a zebrafish predator is successful depends critically on the distance between the mouth of the predator and the prey [22]. We therefore binned measurements of capture probability ( $C$ ) and approximated its dependency on distance with the following sigmoidal function:

$$C(d) = \left[ 1 + e^{-r(d-d_0)} \right]^{-1}, \quad (2.2)$$

where  $d$  is the strike distance,  $d_0$  is the decay distance, and  $r$  is the decay rate. This function additionally approximates the spatial variation in fluid forces that act on prey when subject to suction feeding [30]. The best-fit values for  $d_0$  and  $r$  were determined by least-squares (using the 'sqcurvefit' function in MATLAB).

All parameters for the prey and predators were compared between experiments with adult predators and juvenile predators. Because these measurements failed to conform to normal distributions, we performed statistical comparisons using non-parametric statistics. In particular, we used the two-sample Kolmogorov-Smirnov test (i.e. KS-test) [31], which does not assume any particular distribution.

#### (d) Mathematical model

An agent-based probabilistic model was developed to simulate the conditions of our experiments. This model predicted the 2D motion of a predator and prey [32] according to algorithms that were specific to the behavioral state of each of these agents (Fig. 1b). The predator's states were tracking (T) and striking (S) and the prey's were resting (R) and escaping (E). The duration of states, probability of transitioning between states, and probability of prey capture were determined by random-number generation that conformed to the probability distributions and range of values that we measured. Therefore, this model represents a Markov chain that treated the predator and prey's actions as probabilistic, but each outcome of an interaction also depended on the determinism of the kinematics of the two agents. Simulations were scripted in MATLAB to calculate the motion of both agents and their behavioral states, which consequently determined the number of evasions before prey capture.

Each simulation began with the predator in the tracking state, where it moved at a fixed approach speed with a direction that was always headed toward the prey, with perfect information about the prey's position (Fig. 1b). If the prey was motionless, then the solver would advance in time to the strike or escape initiation, whichever was found to occur first. Otherwise, a moving prey caused the solver to resolve both predator and prey kinematics at a fixed time step (5 ms). In this regime, the predator adjusted its heading to track the prey with a time delay ( $\lambda$ ). The predator's transition into the striking state occurred when the prey was within a particular value for the strike distance. This value was determined *a-priori* by the generation of a random value (using the 'random' function in MATLAB) according to the log-normal probability density function (Eqn. 2.1) for measured values of strike distance. The capture probability for a particular strike depended on the distance between the agents in the middle of a strike, according to our measured parameter values for this relationship (Eqn. 2.2). The simulation was terminated if a strike was successful, otherwise the predator reverted to the tracking state after completion of the strike duration (Fig. 1b). The value of strike duration was determined by the generation of a random value from the log-normal probability density function from measured values. Single values for the predator speed and delay were used for all simulations (Table 1) and were determined by trial-and-error to replicate the distribution of the measured number of unsuccessful strikes before prey capture. These values were found to approximate measurements reported in prior studies [22,33].

The model simultaneously determined the actions of prey (Fig. 1b). Prey behavior was modeled with resting and escaping states because larval zebrafish generally remain still between periods of rapid swimming initiated by an escape response [13,22]. The prey began each simulation in the resting state, where it was motionless and positioned at a random distance from the predator within one aquarium diameter ( $\varnothing = 8.5$  cm). The prey transitioned into the escaping state when the predator moved within the reaction distance, after a latency [34]. During an escape, the speed of prey varied as a single saw-toothed pulse, with the maximum value (the peak of the sawtooth) attained at  $0.2\eta$ , where  $\eta$  is the escape duration. We found that this function well-characterized prey speed using a frame-by-frame kinematic analysis of escape swimming for 12 larvae. The amplitude of the of the saw-toothed pulse represented the maximum escape speed,  $u$ , observed in our 12 recordings of time-resolved escape kinematics. During the escape, the prey was assumed to follow a straight path in a direction determined by the escape angle and escape direction (i.e. left or right sides of the body). The reaction distance, escape angle, and escape duration were determined by random numbers with probability density functions

188 matching experimental measurements. The escape angle ( $\theta$ ) was defined with respect to the prey's  
 189 frame of reference before the escape, with a zero angle corresponding to forward motion. This  
 190 angle was directed with respect to the right or left side of the body by the escape direction, with  
 191 a probability of being contralateral to the predator (Table 1), as previously measured [13].

192 This model simplified many aspects of the complexity of predator-prey interactions. It  
 193 assumed that the kinematics of the two fish may be approximated with two-dimensional motion  
 194 that is not bounded by an aquarium. Simulations were halted if prey successfully escaped on  
 195 20 occasions, which reflected the observed maximum and guarded against an errant simulation  
 196 of infinite duration. The model's use of random number generation considered the probabilistic  
 197 effects of biomechanics and neurophysiology without explicitly articulating those elements. For  
 198 example, capture success was treated as a distance-specific probability (Eqn. 2.2) that modeled  
 199 neither the effects of a predator's suction-feeding hydrodynamics, nor the propulsive forces  
 200 generated by an escaping prey. We tested the model's predictions by comparing its predictions  
 201 of survivorship for 1000 simulations against our measurements. Survivorship was defined as  
 202 the number of individuals surviving a particular number of strikes, divided by the initial  
 203 population. The comparison between measurements and simulations was executed by a two-  
 204 sample Kolmogorov-Smirnov test, which was chosen over a Kruskal-Wallis test because of its  
 205 emphasis on the shape of the distribution.

206 We performed an analysis of the model to evaluate the parameters that had the greatest effect  
 207 on prey survival. This was achieved by running a Monte-Carlo series of 1000 simulations where  
 208 individual parameters were varied in increments of 10% between -90% and 100% of their original  
 209 mean value. For parameters described by a probability distribution, the log-mean parameter,  $\mu$ ,  
 210 was adjusted to create the desired percent-change in the mean of the distribution, while  $\sigma$  was  
 211 held constant. The range of possible random values for each distribution was also adjusted to  
 212 retain the the same cumulative probability range in the probability distribution. The effect of  
 213 these manipulations were assessed by comparing survivorship against the model's prediction  
 214 without any parameter variation using a Kruskal-Wallis test. The results for these comparisons are  
 215 presented in the manuscript figures as the mean probability of surviving a strike, though results  
 216 are also included in supplemental materials (Fig. S2) with respect to the number of escapes prior  
 217 to capture.

### 218 3. Results



## (a) Kinematics

The behavior of both predator and prey were similar whether the predators were juvenile or adult zebrafish. Prey responded in both cases with indistinguishable differences in escape angle (KS-test:  $P = 0.86$ ,  $N = 164$ ) and significant differences in reaction distance (KS-test:  $P < 0.001$ ,  $N = 164$ ) and escape duration (KS-test:  $P = 0.04$ ,  $N = 153$ ) (Fig. 2*b–c*). For example, prey reacted at a mean distance to juvenile predators ( $\bar{l} = 0.84$  cm,  $N = 91$ ), that was about two-thirds the reaction distance to adults ( $\bar{l} = 1.26$  cm,  $N = 73$ ). Escape swimming lasted for about one-third of a second, with the response to juveniles ( $\bar{\eta} = 0.35$  s,  $N = 91$ ), which is only 50 ms longer than in response to adults ( $\bar{\eta} = 0.30$  s,  $N = 62$ ). Prey escaped earlier to adult predators (KS-test:  $P = 0.02$ ,  $N = 89$ ) by 41 ms, on average, relative to the timing of mid-gape (i.e. mean of times when the predator opened and closed their mouth) of suction feeding. Juvenile and adult predators were not significantly different in either their strike distance (KS-test:  $P = 0.08$ ,  $\bar{s} = 7.6$  mm,  $N = 154$ ), or strike duration (KS-test:  $P = 0.87$ ,  $\bar{\tau} = 44$  ms,  $N = 107$ ) (Fig. 2*d–e*). Therefore, much of the behavior of predator and prey were similar, despite the fact that the adults were nearly twice the body length of the juveniles.

Although they exhibited similar behavior, adult predators differed were more effective than the juveniles. Juveniles did not succeed in capturing prey beyond a distance of 3.2 mm ( $N = 91$ ), whereas adults were successful at a maximum distance that was about 3-times greater (10.4 mm,  $N = 77$ ). In the relationship between capture probability and distance (Eqn. 2.2), the decay distance indicates the spatial range of high capture probability. By this metric, the strike of adult predators also exhibited a range that was slightly greater than 3-times the distance of juveniles (Table 1, Fig. 2*f*). We tested whether this result was due to juveniles approaching the prey with inferior accuracy by measuring the bearing of the predators. The bearing corresponds to the radial position of the prey relative to the predator's heading, with perfect accuracy represented by a zero bearing. The bearing when prey initiated an escape was not significantly different (KS-test:  $P = 0.15$ ) between juveniles ( $N = 91$ ) or adults ( $N = 77$ ). At mid-gape, however prey succeeded in evading juvenile predators to the extent that the median bearing ( $13.1^\circ$ ), was less than half that of adult predators ( $30.0^\circ$ ), which was a significant difference (KS-test:  $P < 0.01$ ). Therefore, adult predators were more capable of adjusting their strike toward escaping prey than the juvenile predators. This is not an ability explicitly considered by our model, which assumed a fixed heading during a strike, but was encapsulated in the measured probability distributions of capture success (Fig. 2*f*).

## (b) Mathematical model

Predator and prey fish followed trajectories that were qualitatively similar to that predicted by our model. For most of the duration of our experiments, the predators swam toward the prey (Fig. 3a). In contrast, the prey were generally motionless, except when executing escape swimming. The predators and prey followed a more circuitous path in the predation experiments than the motion prescribed by our model. Nevertheless, the temporal sequence of events in the model offered a reasonable approximation of the kinematics of live predator-prey interactions (Fig. 3b).

The model accurately predicted our measurements of prey survivorship. Measured survivorship decreased monotonically from the first strike and extended to as many as 20 strikes (Fig. 3c). The ability of larval prey to more effectively evade juveniles than adults (Fig. 2f) was reflected in a slower decline in survivorship. The model was successful in replicating these trends, which were found to be statistically indistinguishable for both adult (KS-test:  $P = 0.93$ ,  $N = 73$ ) and juvenile (KS-test:  $P = 0.86$ ,  $N = 91$ ) predators. Furthermore, all trends from the parameter analysis of the pursuit-evasion model were similar between the adults (Fig. 4) and juveniles (Fig. S1).

Our parameter analysis revealed that escape speed and reaction distance were the only parameters for prey that showed any noteworthy effect on survival. This analysis adjusted the probability distribution of each parameter to generate a percent change in its mean value among 1000 simulations. Generating these changes in escape duration, escape direction, and escape angle yielded statistically insignificant or otherwise small changes in escape probability (Fig. 4a). An increase in escape speed similarly had a negligible effect on survival, but survival probability did decline when speed was reduced by 50% or more. Survival was most sensitive to reaction distance. An increase in reaction distance caused escape probability to elevate by as much as 16% and a decrease of at least 30% had a dramatically adverse effect on survival (Fig. 4a).

We tested for interactive effects between reaction distance and escape speed by conducting a two-dimensional parameter analysis (Fig. 4b). The relationship between reaction distance and survival was effectively unaltered by differences in escape speeds, unless this speed was reduced below 50% of its measured value (Fig. 4c). Irrespective of reaction distance, a reduction of escape speed by at least 50% was required to create a significant change in survival (Fig. 4d). This suggests little evidence for an interactive effect on survival between escape speed and response distance in the range of parameter values presently considered.

## 4. Discussion

We found that the survival of larval fish does not increase by escaping at a faster speed or by varying direction, but only by responding from a greater distance. These results were derived from an agent-based probabilistic model (Fig. 1*b*) that calculated the trajectories of predator and prey and the outcome of predatory strikes (Fig. 3*a,b*). This successfully replicated broad patterns of survivorship (Fig. 3*c*) by simulating behavioral actions that matched our measurements (Fig. 2). Our analysis of its predictions suggests that prey survival in fishes may only be enhanced by increasing the sensitivity of predator detection.

### (a) Locomotor performance and prey survival

The survival of prey depends largely on the actions of the predator. In contrast to the explosive speed of an escape response [20], adult zebrafish tend to approach their prey substantially slower than their capacity, often by braking [33]. We found that the approach speed amounts to less than one-third the maximum speed of escaping larvae (Table 1), which is consistent with previous measurements [22]. This approach relates strategically to the mechanics of feeding. The suction feeding of fishes succeeds in capturing prey in only a small region around the mouth over a duration of merely tens of milliseconds [35–37]. A slow approach is common among suction-feeding fishes and is likely a means of enhancing strike accuracy [8,9]. This style of predation is seen over many species of fish [38]. Furthermore, our data suggest that adult and juvenile zebrafish are more likely to capture when approaching larval zebrafish with a slower approach speed (Fig. S3). Therefore, the limited range of suction feeding may constrain some predators to a slow approach while offering prey an opportunity to escape [24]. Despite this strategic advantage for prey, adult zebrafish captured prey on the first strike more than one-quarter of the time and rarely needed more than three strikes to be successful (Fig. 3*c*).

The effectiveness of an escape has previously been considered by classic pursuit-evasion models of fish predation. This theory resolves how the direction of an escape affects the distance between predator and prey [5,32], generally by modeling a single encounter with the assumption that both animals move with a fixed heading and speed over time. A recently-developed version of this model suggests that animals like zebrafish operate in a ‘slow-predator’ strategic domain [39], where the predator moves more slowly than the prey. In this domain, no optimal escape angle exists and prey may evade predators with a broad range of effective escape directions. A faster escape speed serves only to modestly expand the range of effective escape angles.

Consistent with these predictions, our model found a monotonic decrease in survival as we reduced escape speed below half of the observed value (Fig. 4a). We additionally found only modest differences in survival between experiments using adult and juvenile predators (Figs. 2,3c), despite a nearly two-fold difference in body size and speed.

These predictions would not hold for cases where the predator is faster than the prey. In such a strategic domain, an optimal escape angle arises for the prey and failure to move in that direction is predicted to adversely affect survival [5,39]. Ram-feeding fishes strike at prey while swimming at a relatively high speed and may thereby place prey at a strategic disadvantage. Success in ram feeding may, in-turn, require superior coordination in directing and timing a strike [11]. Ram feeding in this way shows greater similarity in strategy to flying predators such as birds [15], bats [16], and insects [40]. Prey may benefit by escaping in a direction that conform to an optimal value [5], by being unpredictable [6], or escaping along a trajectory with a small radius of curvature [7,41].

## (b) Prey survival depends on reaction distance

The reaction distance has broad strategic significance. Classic pursuit-evasion models support the simple notion that prey are more evasive if they start from further away [5,32,39]. This principle is consistent with evolutionary models that contrast the fitness benefit of responding from a distance against its potential costs [42,43]. For example, escape responses that are frequently initiated unnecessarily may be energetically expensive, prohibit foraging, or succeed in revealing cryptic prey [44]. Responding from a great distance may even be inferior on purely strategic grounds. A prey that is slower than a predator, but capable of executing a tight turn, may be more evasive when initiating this maneuver at the final moments of a predatory strike, rather than providing the opportunity for the predator to adjust course [7]. Therefore, a greater reaction distance offers a clear strategic benefit in zebrafish (Fig. 4), but may not be universally advantageous.

The primacy of reaction distance underscores the strategic importance of predator detection. Responding to a predator from afar depends on the sensitivity of receptor organs and the capacity of the nervous system to rapidly recognize a threatening cue and trigger an escape response. As in invertebrate zooplankton [14], and insect prey of spiders [45,46], zebrafish larvae use flow sensing to detect the bow-wave of flow generated by an approaching predator [22]. Flow-sensing may be augmented by olfactory cues [47], though zebrafish do not acquire a sensitivity to the alarm pheromone Schreckstoff until a later stage of growth (>48 dpf) [47]. Using the lateral line system, zebrafish larvae may respond to this stimulus up to a distance of 1.3 cm [13] ahead of

a gliding zebrafish adult. This range encompasses many of the responses that we recorded (Fig. 2c), which supports a role of the lateral line in our experiments. This would agree with previous experiments that have shown flow sensing to be necessary for survival in zebrafish larvae [22]. Flow sensing offers the capacity to trigger an escape with a very brief ( $< 10$  ms) latency [48]. In contrast, a looming visual stimulus succeeds in stimulating an escape responses in zebrafish larvae [49], but the demands for visual processing necessitate a latency that is at least 20-fold longer than for flow sensing [50]. Nonetheless, the greatest reaction distances that we observed were outside the range of the lateral line and therefore likely generated by the visual appearance of the predator. The visual system consequently offers prey fish the means to enhance survival by responding to a predatory threat from a distance (Fig. 4).

## Summary

We found that zebrafish larvae operate in a slow-predator strategic domain when preyed upon by adults and juveniles of the same species. As a consequence, increasing the speed or varying the direction of an escape response shows a negligible effect on survival (Fig. 4). Survival may instead be enhanced by initiating the escape from greater distance by rapidly identifying the predator as a threatening visual stimulus. These findings offer valuable insight into the key strategic factors that govern predator-prey interactions in a diversity of similar fishes and other animals that operate with a similar strategy.

## Authors' contributions

The study was designed in collaboration between AN and MJM. AN and CN performed all experiments and kinematic analysis. The agent-based probabilistic model was created by AN, with guidance from MJM. The manuscript was written collaboratively by AN and MJM.

## Competing interests

We declare we have no competing interests.

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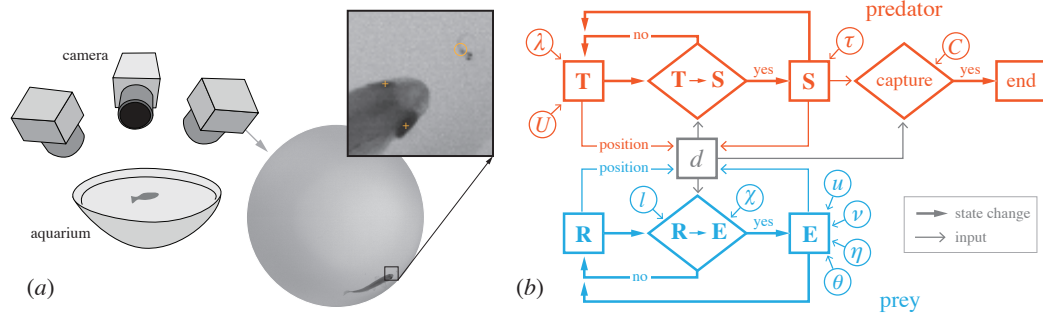
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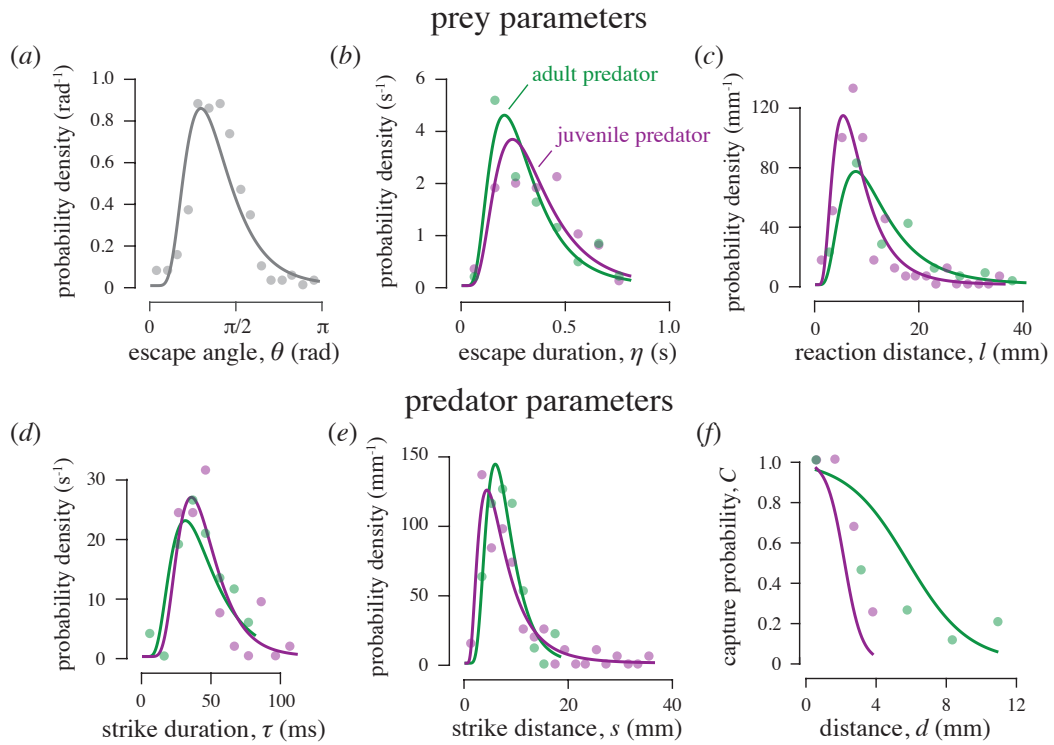
**Table 1.** Behavioral parameters and probability distributions

Variable	State	Adult predator	Juvenile predator
<i>Predator</i>			
Approach speed, $U$ ( $\text{m s}^{-1}$ )	T	$U = 0.13$	$U = 0.02$
Predator delay, $\lambda$ (ms)	T	$\lambda = 10$	$\lambda = 100$
Strike distance, $s$ (m)	T $\rightarrow$ S	$\mu_d = -4.980$ , $\sigma_d = 0.448$ ( $N = 51$ )	$\mu_d = -5.100$ , $\sigma_d = 0.648$ ( $N = 103$ )
Strike duration, $\tau$ (s)	S	$\mu_\tau = -3.166$ , $\sigma_\tau = 0.331$ ( $N = 53$ )	$\mu_\tau = -3.208$ , $\sigma_\tau = 0.399$ ( $N = 54$ )
Capture probability, $C$	S	$r = -0.573$ , $d_0 = 5.20$ ( $N = 77$ )	$r = 1.99$ , $d_0 = 1.60$ ( $N = 91$ )
<i>Prey</i>			
Reaction distance, $l$ (m)	R $\rightarrow$ E	$\mu_l = -4.546$ , $\sigma_l = 0.587$ ( $N = 73$ )	$\mu_l = -4.941$ , $\sigma_l = 0.582$ ( $N = 91$ )
Escape angle, $\theta$ (rad)	E	$\mu_\theta = 0.144$ , $\sigma_\theta = 0.449$ ( $N = 206$ )	$\mu_\theta = 0.144$ , $\sigma_\theta = 0.449$ ( $N = 206$ )
Escape duration, $\eta$ (s)	E	$\mu_\eta = -1.369$ , $\sigma_\eta = 0.552$ ( $N = 62$ )	$\mu_\eta = -1.167$ , $\sigma_\eta = 0.5234$ ( $N = 91$ )
Escape direction, $v$	E	$v = 0.696$ ( $N = 206$ )	$v = 0.696$ ( $N = 206$ )
Escape latency, $\chi$ (ms)	E	$\chi = 8$ ( $N = 15$ )	$\chi = 8$ ( $N = 15$ )
Escape speed, $u$ ( $\text{m s}^{-1}$ )	E	$u = 0.4$ ( $N = 12$ )	$u = 0.4$ ( $N = 12$ )

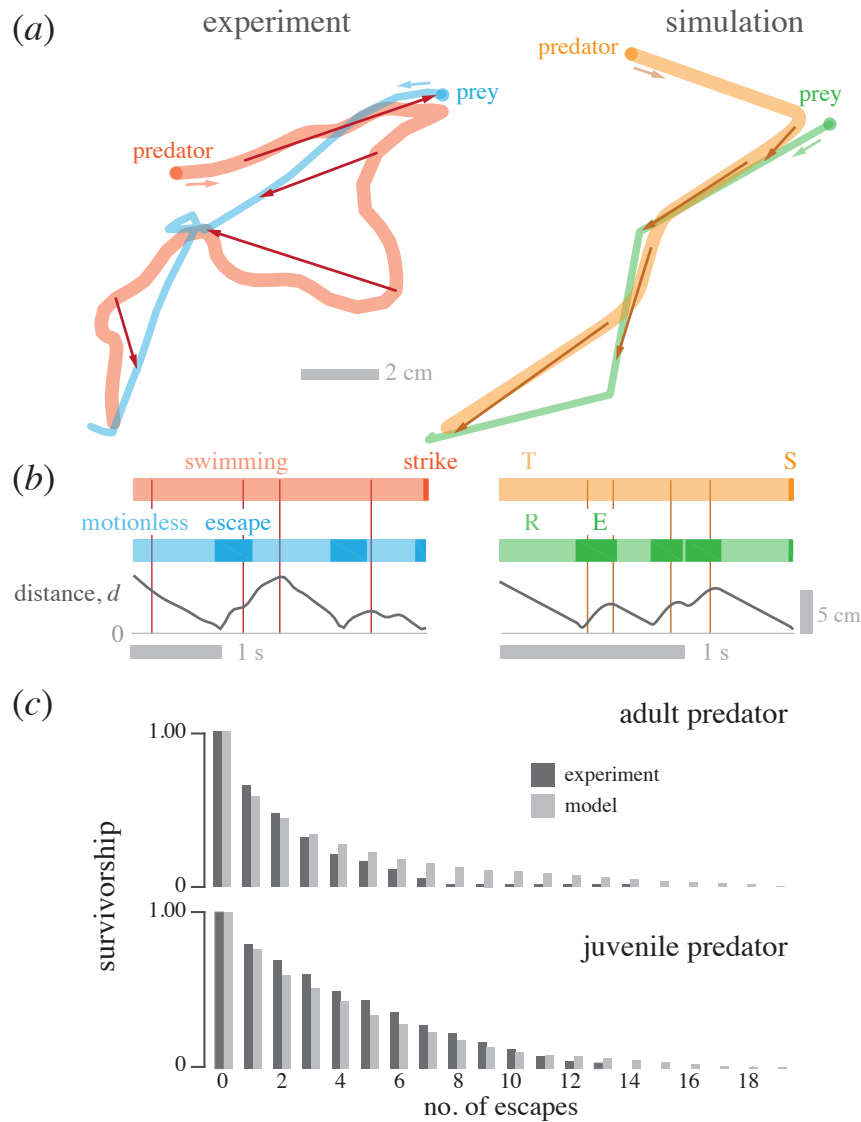
T, tracking; S, striking; R, resting; E, escaping;  $\mu$ , log mean;  $\sigma$ , log standard deviation;  $r$ , decay rate ( $\text{mm}^{-1}$ );  $d_0$ , decay distance (mm).



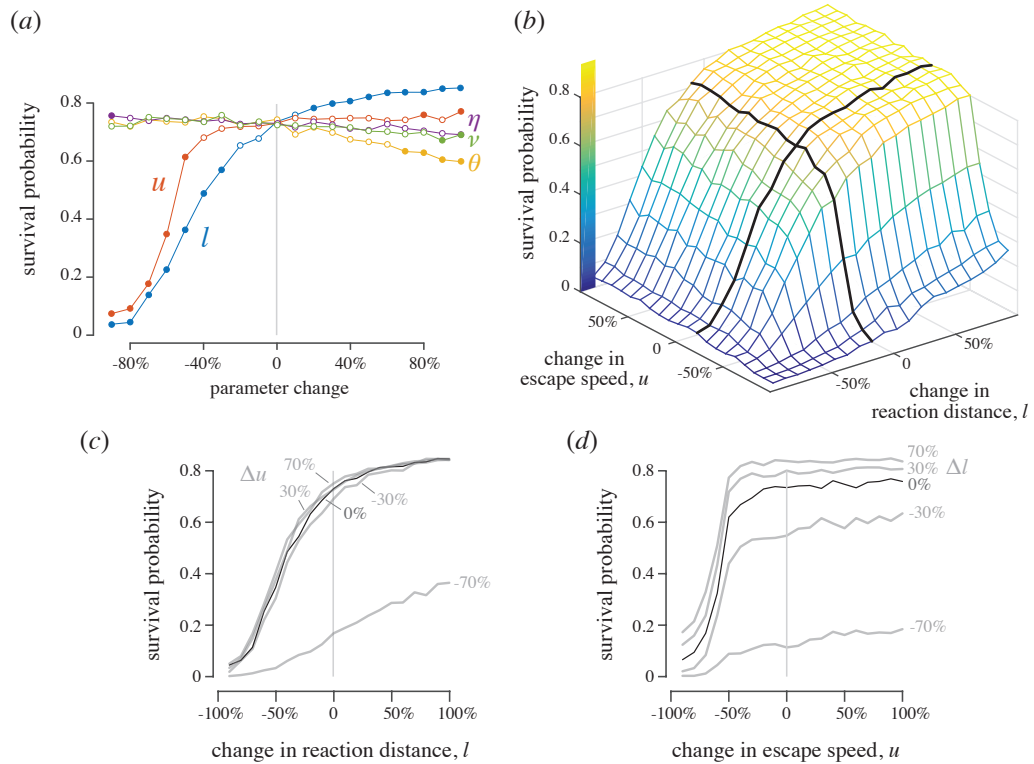
**Figure 1.** Kinematic measurements and mathematical modeling for studying predator-prey interactions in zebrafish. (a) Three high-speed video cameras recorded one larval prey and one predator fish (adult or juvenile) that were placed in a hemispherical aquarium. A representative video frame (cropped to the margin of the aquarium) shows an adult in close proximity to the prey. In the inset, orange markers denote the locations of morphological landmarks used to describe the position of the two fish. This consisted of the position of the two eyes for the predator ("+" ) and the posterior margin of the swim bladder in the prey (open circle). (b) A state diagram illustrates the major components of the agent-based probabilistic model used to simulate the interactions between predators and prey (see Table 1 for symbol definitions and parameter values). Each fish behaves according to an algorithm specific to a particular behavioral state and the probability of transitioning between states is determined by random-number generators with probability distributions matching our measurements (Fig 2). Predators (in red) operate between tracking (T) and striking (S) states and prey are either resting (R) or escaping (E). The outcome of a strike is determined by the capture probability ( $C$ , Eqn. 2.2). Simulations of this model were performed with a Monte-Carlo method to generate probability distributions of prey survival. See Material and methods for details.



**Figure 2.** Descriptive statistics of swimming kinematics. (a–e) Measurements of the probability density (circles) for the kinematics of the prey (a–c) and predator (d–e) for experiments that included either a juvenile (purple) or adult (green) predator. Points on the graphs denote measured binned values for probability density with a sample size determined using the Freedman-Diaconis rule, which yielded  $N \sim 17$  measurements per point (see Table 1 for total sample sizes). Measurements of escape angle (a) were not significantly different between the two types of experiments and these measurements were therefore combined (gray circles). For each parameter, we performed a non-linear least-squares fit to the measurements for a log-normal probability density function (Eqn. 2.1). The log-mean and log-standard deviation values from these fits (Table 1) were consequently used to describe the probability of events in our mathematical model (Fig. 1b). (f) The capture probability was measured as a function of distance between the predator and prey and we similarly performed a curve fit to approximate this relationship (Eqn. 2.2) for our model.



**Figure 3.** Comparison between experimental measurements and modeling. (a) Trajectories of predator and prey from a representative experiment (left) and simulation (right). The position of predator and prey that correspond to particular time points are shown with connecting arrows. (b) Ethograms for these trajectories illustrate the temporal changes in the predator's swimming and strike (left), which are respectively modeled by the tracking (T) and striking (S) (Fig. 1b) states (right). The prey's behavior while motionless and during escape (left) were respectively modeled as resting (R) and escape (E) modes (right). For both ethograms, the distance ( $d$ ) between predator and prey are shown. Particular moments in the trajectories are highlighted with vertical lines that correspond with the same-colored arrows in (a). (c) The probability that a prey survives over a particular number of strikes is shown for adult (above) and juvenile (below) predators for experiments (dark gray) and simulations (light gray).



**Figure 4.** Parameter analysis of the mathematical model to examine the effects on escape probability. (a) We individually varied the mean of a parameter among simulations by manipulating the distribution (Fig. 2) of our measurements (see Table 1 for parameter definitions and values). Each point represents the survival probability of prey among 1000 simulations and filled circles denote a significant difference (KS-test:  $P < 0.05$ ) from the observed probability. Simulations that varied in escape angle ( $\theta$ ) differed by a interval of 0.127 rad. (b) Variation in escape probability was examined with respect to both escape speed and reaction distance. The same simulation results are shown with respect to changes in escape speed (c) and reaction distance (d). All simulations shown here used an adult predator, although similar results were obtained with a juvenile predator (Fig. S1).