Challenging zebrafish escape responses by increasing water viscosity

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RESEARCH ARTICLE

Challenging zebrafish escape responses by increasing water viscosity

Nicole Danos* and George V. Lauder

Department of Organismic and Evolutionary Biology, Harvard University, 26 Oxford Street, Cambridge, MA 02138, USA

*Author for correspondence at present address: Ecology and Evolutionary Biology, 321 Steinhaus Hall, University of California Irvine, Irvine, CA 92697, USA (ndanos@post.harvard.edu)

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SUMMARY

Escape responses of fishes have long been studied as a model locomotor behavior in which hypothesized maximal or near-maximal muscle power output is used to generate rapid body bending. In this paper we present the results of experiments that challenged zebrafish (Danio rerio) to perform escape responses in water of altered viscosity, to better understand the effects that the fluid mechanical environment exerts on kinematics. We quantified escape kinematics using 1000 frames s\(^{-1}\) high-speed video, and compared escape response kinematics of fish in three media that differed in viscosity: 1 mPa s (normal water), 10 mPa s and 20 mPa s (20 times normal water viscosity). We hypothesized that because viscosity is increased but not density there will be a different effect on kinematic variables resulting from unsteady (acceleration-dependent) hydrodynamic forces and steady (velocity-dependent) ones. Similarly, we hypothesized that the kinematics of stage 1 will be less affected by viscosity than those of stage 2, as higher angular velocities are reached during stage 1 resulting in higher Reynolds numbers. Our results showed a significant overall effect of viscosity on escape response kinematics but the effect was not in accordance with our predictions. Statistical tests showed that increasing viscosity significantly decreased displacement of the center of mass during stage 1 and after 30 ms, and decreased maximum velocity of the center of mass, maximum angular velocity and acceleration during stage 1, but increased time to maximum angular acceleration and time to maximum linear velocity of the center of mass. Remarkably, increasing water viscosity 20 times did not significantly affect the duration of stage 1 or stage 2.

Key words: locomotion, fish, viscosity, zebrafish.

INTRODUCTION

The escape response of fishes is a widespread model of vertebrate locomotor behavior that has been used to clarify the design of neural circuits (e.g. Eaton et al., 1977; Foreman and Eaton, 1993; Liu and Fetcho, 1999), understand rapid activation of axial muscles (Jayne and Lauder, 1993; Rome et al., 1988; Tytell and Lauder, 2002; Westneat et al., 1998), clarify predator-avoidance dynamics (Walker et al., 2005) and investigate unsteady locomotor hydrodynamics (e.g. Borazjani et al., 2012; Frith and Blake, 1995; Tytell and Lauder, 2008). Escape behavior usually occurs in response to an impulsive hydrodynamic or visual stimulus and is usually (though not always) mediated by the Mauthner cells of the hindbrain (Zottoli, 1977). Escape responses by fishes are characterized by large body angular accelerations and displacements (Domenici and Blake, 1997) and are usually divided into two stages: an initial C-bend (stage 1) and a contralateral bend followed by one or more tailbeats (stage 2).

One assumption that has made escape responses a useful model for studying muscle physiology and mechanics is that this behavior appears to require maximum muscle power production. This is a reasonable assumption given the importance of this behavior in predator avoidance (Walker et al., 2005), although a difficult hypothesis to test directly given the assumptions needed to evaluate the in vivo power production of complexly arranged segmental body musculature with multiple fiber types. A number of studies have investigated the power output of fish muscle during escape behaviors and have suggested that the body musculature is activated in a near-maximal manner (e.g. Franklin and Johnston, 1997; Johnston et al., 1995; Wakeling and Johnston, 1998).

One approach that might reveal new features of fish escape response dynamics involves challenging fish to perform their escape behavior under increased loading, i.e. in a more viscous fluid environment. Specifically, we believe that fish induced to perform escape responses in water of substantially higher viscosity than normal should have to increase muscle power output to accomplish normal kinematics in a similar time frame. Alternatively, if muscle power output cannot be increased beyond that seen in normal escape behaviors, then substantial increases in water viscosity (of the order of 20 times normal) might result in longer times for the body to achieve the stage 1 and stage 2 kinematic bending patterns. Increased viscosity should also increase the thickness of the boundary layer around the bending fish and affect body acceleration patterns. Fish might perform the same sequence of kinematic events in water of high viscosity, but each stage could take longer to complete and accelerations might be lower. Altering increased viscosity could also have differential effects on stage 1 and stage 2 of the escape behavior. During stage 2 there are lower accelerations and angular displacements than in stage 1, and this stage is not controlled by the large Mauthner cells with their fast transmission times that activate myotomal muscle fibers (Eaton et al., 1977; Weiss et al., 2006).

Additionally, increased viscosity may also affect C-start kinematics by altering the hydrodynamics that govern fin function. Recent studies have demonstrated the importance of the dorsal and anal fins in producing forces during locomotion in general (Drucker and Lauder, 2005; Standen and Lauder, 2005; Standen and Lauder, 2007) and escape responses in particular (Tytell and Lauder, 2008).
The shedding of vortices from flapping foils is delayed when viscosity is increased (Faber, 1995) and in the case of an accelerating fish this could affect patterns of force production, resulting in different timings of events.

However, the high accelerations and displacements of this locomotor behavior indicate that unsteady forces govern its hydrodynamics, which are poorly understood. But, in addition to the steady-state forces such as skin and pressure drag, hydrodynamic analyses of escape responses need to consider the acceleration reaction force (Daniel, 1984). As the turning fish and its tail accelerate a mass of water, this reaction force acts opposite to the direction of body movement and resists the fish body acceleration. Only water density is expected to affect the magnitude of this force (Daniel, 1984), unless increased viscosity indirectly increases the added mass coefficient.

In this paper, we performed experiments that challenged zebrafish (Danio rerio) to perform escape responses in water altered to be 10 and 20 times more viscous than normal water. We found that while increasing water viscosity does have a significant overall effect on the escape response, many of the observed specific changes in the directions of body movement and resists the fish body acceleration.

The shedding of vortices from flapping foils is delayed when viscosity is increased (Faber, 1995) and in the case of an accelerating fish this could affect patterns of force production, resulting in different timings of events.

MATERIALS AND METHODS

Animals and viscosity treatments

Zebrafish, D. rerio (Hamilton 1822), were obtained from the Harvard University zebrafish facility and kept in an aquarium under 12h:12h light:dark conditions. Five fish were used for the experiments; the experimental design and statistical analyses are described below. Fish (mean total length 33.0±1.3 mm) were placed in a circular container (diameter 12 cm, filled to a depth of 10 cm) with an optically clear base and allowed to swim freely. Video images from below (ventral view) were captured at 1000 frames s⁻¹ using a Photron PCI 1024 camera (1024x1024 pixel resolution, Photron Inc., San Diego, CA, USA). A cylindrical object with an added flat base (approximate diameter 1.5 cm) to enhance the initial impact at the water surface was dropped into the tank holding an individual zebrafish to elicit escape responses from the fish. This follows the procedure that we have used in previous experiments on fish escape responses (Jayne and Lauder, 1993; Tytell and Lauder, 2002; Tytell and Lauder, 2008).

Each of the five fish was recorded performing escape responses in media of all three levels of viscosity: 1 mPa s (normal water), 10 mPa s or 20 mPa s. The sequence of viscosity treatments at which each fish was tested was randomized. To prepare high viscosity solutions, Dextran 500 (Pharmacosmos A/S, Holbaek, Denmark) was dissolved in aquarium water. Oxygen levels in the increased viscosity media were monitored using an SM 600 dissolved oxygen meter (Milwaukee Instruments, Rocky Mount, NC, USA) and never fell below 6.5 p.p.m. in any treatment, the equivalent of 84% saturation under our laboratory conditions. Viscosities were measured using shell cup viscometers (Norcross, Newton, MA, USA). Dextran solutions have Newtonian fluid dynamic properties even at the relatively high viscosities used here (Akashi et al., 2000). Newtonian fluids have a constant viscosity, independent of shear stress rate produced by the fish’s velocity during the escape response. Although the increased viscosity treatments differed from normal water by an order of magnitude, the differences in fluid density were negligible (Akashi et al., 2000). We caution that use of other compounds such as methyl cellulose to increase water viscosity may produce a non-Newtonian fluid in which the forces generated depend on shear rate, making interpretation of changes in kinematics due to changes in viscosity challenging.

Viscosity manipulations have been used in a number of previous studies to separate the effects of temperature and viscosity on locomotor performance and to examine the hydrodynamics of behaviors at low Reynolds numbers. Here, we report viscosity in Pa s, the SI unit for dynamic (or absolute) viscosity. The viscosity values of 10 and 20 mPa s were chosen because we expected that viscosity levels 10 and 20 times the normal viscosity of water would be outside the natural range any fish would experience and hence would represent a significant challenge to fish performing escape responses at already maximal power outputs.

All experimental protocols were approved by Harvard University’s institutional animal care and use committee.

Video analysis

At least three video sequences of escape responses were digitized in Matlab (The MathWorks Inc., Natick, MA, USA) for each individual at each viscosity using a custom-written program (Hedrick, 2008) for a total of 52 escape responses. Eleven points were digitized in each frame (Fig. 1): the tip of the snout, the base of each pectoral fin, the base of each pelvic fin, points on either side of the body 1/3 and 2/3 of the distance between the pelvic fin base and the caudal peduncle, the caudal peduncle and the tip of the dorsal caudal lobe. From these points a body midline was reconstructed in six segments and the angles between the segments were used to characterize body curvature.

The two stages of each escape response were identified as described by previous authors (e.g. Domenici and Blake, 1991; Tytell and Lauder, 2008). Stage 1 began in the frame of the first visible movement of the fish’s snout and ended when angular...
For all maxima calculated, we also recorded the time it took for the maximum linear velocity of the COM and of the tail (Table 1). The process was repeated for displacement at 30 ms, COM displacement in St2, COM angular velocity, and acceleration were calculated by dividing turn angle changes by the final turn angle, respectively. Instantaneous angular velocity and acceleration were calculated by dividing turn angle changes by the time between consecutive image frames (1 ms). From these values, we recorded maximum angular velocity and acceleration for the entire escape response and for each stage of the turn (Table 1). Instantaneous linear displacement of the center of mass (COM) in a stretched-straight fish (see Jayne and Lauder, 1993; Tytell and Lauder, 2002) was calculated as the linear distance traveled by the midpoint between the pectoral fins (end of body segment 1) relative to its initial position. Displacement at the end of each stage as well as after a predetermined amount of time was then recorded for each swimming sequence (Table 1; COM displacement in St1, COM displacement in St2, COM displacement at 30 ms). Linear velocity and acceleration were calculated by dividing displacement by the time between consecutive image frames. The process was repeated for displacement and velocity of the tail (Fig. 1). We then recorded the maximum linear velocity of the COM and of the tail (Table 1). For all maxima calculated, we also recorded the time it took for the maximum to be reached from the beginning of the escape response (e.g. time to maximum turn angle; Table 1).

### Statistics

To determine whether there was an overall viscosity effect, we performed three multivariate analyses of the dataset. First, the dataset was analyzed using a multivariate ANOVA with viscosity as a fixed effect and 21 kinematic response variables (Table 1). Maximum velocity and acceleration and the time to these values were not included in these analyses because they also appeared in the dataset as the maximum values in either stage 1 or stage 2. A significant multivariate ANOVA result for viscosity was then followed by post hoc univariate ANOVA on each of the kinematic variables with viscosity as a fixed effect and individual fish as a random factor (Zar, 1999). The results of the univariate ANOVA were deemed significant if supported by a P-value smaller than 0.01 (Table 1).

For each video frame the turn angle, $\theta$, was calculated. This was designated as the angle that the midline of the first body segment would make with the midline of this same segment in the first frame of the turn (Fig. 1). For each video sequence we then recorded the maximum turn angle, and the turn angle at the end of stage 1 and at the end of stage 2 (Table 1; St1 angle and final turn angle, respectively). Instantaneous angular velocity and acceleration were calculated by dividing turn angle changes by the time between consecutive image frames (1 ms). From these values, we recorded maximum angular velocity and acceleration for the entire escape response and for each stage of the turn (Table 1). Instantaneous linear displacement of the center of mass (COM) in a stretched-straight fish (see Jayne and Lauder, 1993; Tytell and Lauder, 2002) was calculated as the linear distance traveled by the midpoint between the pectoral fins (end of body segment 1) relative to its initial position. Displacement at the end of each stage as well as after a predetermined amount of time was then recorded for each swimming sequence (Table 1; COM displacement in St1, COM displacement in St2, COM displacement at 30 ms). Linear velocity and acceleration were calculated by dividing displacement by the time between consecutive image frames. The process was repeated for displacement and velocity of the tail (Fig. 1). We then recorded the maximum linear velocity of the COM and of the tail (Table 1). For all maxima calculated, we also recorded the time it took for

### Table 1. Univariate ANOVA results for 25 kinematic variables during zebrafish C-starts

<table>
<thead>
<tr>
<th>Variable</th>
<th>Viscosity</th>
<th>Individual</th>
<th>Viscosity × Individual</th>
<th>Steady or Unsteady</th>
<th>St1 or St2</th>
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<tbody>
<tr>
<td>Duration (ms): St1</td>
<td>0.59</td>
<td>0.64</td>
<td>0.18</td>
<td>Steady</td>
<td>St1</td>
</tr>
<tr>
<td>Duration (ms): St2</td>
<td>0.19</td>
<td>0.0003</td>
<td>0.0001</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Maximum turn angle (deg)</td>
<td>0.26</td>
<td>0.74</td>
<td>0.56</td>
<td>Steady</td>
<td>St1</td>
</tr>
<tr>
<td>Maximum angular velocity (deg s$^{-1}$)</td>
<td>&lt;0.0001</td>
<td>0.04</td>
<td>0.20</td>
<td>Steady</td>
<td>St1</td>
</tr>
<tr>
<td>Maximum angular acceleration (deg s$^{-2}$)</td>
<td>0.018</td>
<td>0.57</td>
<td>0.23</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
<tr>
<td>Final turn angle (deg)</td>
<td>0.11</td>
<td>0.79</td>
<td>0.51</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Maximum COM velocity (mm s$^{-1}$)</td>
<td>&lt;0.0001</td>
<td>0.04</td>
<td>0.37</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Maximum tail velocity (mm s$^{-1}$)</td>
<td>0.82</td>
<td>0.41</td>
<td>0.54</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>St1 angle (deg)</td>
<td>0.26</td>
<td>0.58</td>
<td>0.35</td>
<td>Steady</td>
<td>St1</td>
</tr>
<tr>
<td>Maximum angular velocity in St1 (deg s$^{-1}$)</td>
<td>&lt;0.0001</td>
<td>0.04</td>
<td>0.20</td>
<td>Steady</td>
<td>St1</td>
</tr>
<tr>
<td>Maximum angular acceleration in St1 (deg s$^{-2}$)</td>
<td>0.0057</td>
<td>0.32</td>
<td>0.04</td>
<td>Unsteady</td>
<td>St1</td>
</tr>
<tr>
<td>Maximum angular velocity in St2 (deg s$^{-1}$)</td>
<td>0.0028</td>
<td>0.27</td>
<td>0.25</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Maximum angular acceleration in St2 (deg s$^{-2}$)</td>
<td>0.0018</td>
<td>0.39</td>
<td>0.51</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
<tr>
<td>COM displacement at St1 (mm)</td>
<td>&lt;0.0003</td>
<td>0.46</td>
<td>0.47</td>
<td>Steady</td>
<td>St1</td>
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<tr>
<td>COM displacement at St2 (mm)</td>
<td>0.0020</td>
<td>0.30</td>
<td>0.12</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>COM displacement at 30 ms (mm)</td>
<td>0.0001</td>
<td>0.16</td>
<td>0.72</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum turn angle (ms)</td>
<td>0.27</td>
<td>0.67</td>
<td>0.05</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum angular velocity (ms)</td>
<td>0.92</td>
<td>0.81</td>
<td>0.62</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum angular acceleration (ms)</td>
<td>0.0061</td>
<td>0.89</td>
<td>0.52</td>
<td>Steady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum COM velocity (ms)</td>
<td>0.0007</td>
<td>0.71</td>
<td>0.14</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum tail velocity (ms)</td>
<td>0.25</td>
<td>0.10</td>
<td>0.10</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum angular velocity St1 (ms)</td>
<td>0.92</td>
<td>0.81</td>
<td>0.60</td>
<td>Unsteady</td>
<td>St1</td>
</tr>
<tr>
<td>Time to maximum angular acceleration St1 (ms)</td>
<td>0.026</td>
<td>0.90</td>
<td>0.96</td>
<td>Unsteady</td>
<td>St1</td>
</tr>
<tr>
<td>Time to maximum angular velocity St2 (ms)</td>
<td>0.61</td>
<td>0.15</td>
<td>0.15</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
<tr>
<td>Time to maximum angular acceleration St2 (ms)</td>
<td>0.08</td>
<td>0.04</td>
<td>0.11</td>
<td>Unsteady</td>
<td>St2</td>
</tr>
</tbody>
</table>

Data are P-values (see Results for discussion). St1 and St2 refer to stage 1 and stage 2 of the C-start escape response; ‘steady’ refers to velocity-dependent variables; ‘unsteady’ refers to acceleration-dependent variables.

*Significant at 0.01 level.
1 Mean value for water differed from both high viscosity treatments but the high viscosity means did not differ (Tukey HSD).
2 Mean value for water and 10 mPa s were indistinguishable but different from the mean at 20 mPa s (Tukey HSD).
3 Mean value for water was significantly different from the value at 10 mPa s but both were indistinguishable from the mean value at 20 mPa s.
RESULTS

Fifty-two escape responses were studied from 5 fish. Thirty-one of the turns were initiated after the dropped object impacted the surface of the water; these we called mechanically stimulated turns. Twenty-one of the turns began before the object hit the surface of the water; these we called visually stimulated turns. Mechanically stimulated turns constituted 71% of the escape turns in normal water (1 mPa s), 50% in 10 mPa s and 59% in 20 mPa s. The observed viscosity effects could be due to an alteration of the physics of mechanosensation (McHenry et al., 2009a). Therefore, for mechanically stimulated turns we also measured the time between the stimulus and the first frame of the turn (latency), the angle between the dorsal midline of the fish and a line connecting the center of mass of the fish to the stimulus location (angle to stimulus at turn start) as well as the linear distance between the fish center of mass at the beginning and end of the stimulus (distance to stimulus). Neither viscosity nor individual had any significant effect on the variables, although there was a tendency for latency to be shorter in higher viscosity media. We therefore grouped all C-starts together for subsequent analyses.

Reynolds number calculations using fish length, maximum linear velocity per sequence and the appropriate viscosity for each treatment show our manipulations altered the maximum Reynolds number of a sequence by an order of magnitude between consecutive viscosity levels. The mean maximum Reynolds number in 1 mPa s water was 27,800 (±2140 s.e.m.) compared with 2350 (±160 s.e.m.) and 820 (±60 s.e.m.) in 10 and 20 mPa s fluid, respectively. Fish escaping in normal water (1 mPa s) operated in the inertial hydrodynamic regime for the entire duration of the escape response (Fig. 2B). The effect of viscosity became apparent at 10 mPa s viscosity, where the initial angular acceleration took place under viscous hydrodynamic conditions, proceeded past the point of maximum angular velocity in an intermediate regime and ended with high linear velocity and constant angular acceleration in the inertial regime (Fig. 2B). Fish performing escape responses in the highest viscosity treatment, 20 mPa s, spent a very small proportion of the turn in the inertial regime, while most of the turn was under the complex hydrodynamics of the intermediate regime (Fig. 2B).

Escape responses at all viscosities followed the same general sequence of characteristic behavioral events typical of a fish C-start in normal water. Fig. 2 provides an example of a typical C-start from the same fish at each of the three viscosities tested. Each C-start shows the characteristic stage 1 and stage 2 components. The duration of each stage did not show a significant viscosity effect (Fig. 3; Table 1). Although the angular velocity profiles are similar
for all viscosities, linear velocity dropped sharply by the end of stage 2 in the highest viscosity treatment (Fig. 2B). The reduction in linear velocity during stage 2 was significantly higher at 20 mPa s viscosity compared to turns in water (Tukey’s HSD). In general, linear velocities at the end of stage 1 as well as the maximum velocity reached during that stage were significantly higher in water than in both increased viscosity treatments (Tukey’s HSD; Fig. 3).

As shown in Table 1, only 10 kinematic variables had a significant viscosity effect: maximum angular velocity, maximum COM velocity, maximum angular velocity and acceleration in both stages, COM displacement at the end of stage 1 and after 30 ms, time to maximum angular acceleration and time to maximum COM velocity. No variable showed an effect of individual, although the duration of stage 2 had a significant viscosity by individual effect (Table 1). Post hoc tests (Tukey HSD) indicated that the mean kinematic value never differed among all three viscosities (Table 1). Instead, two of the viscosities grouped together but differed from the third, or two viscosities were significantly different from each other but both were indistinguishable from the third (Table 1).

Maximum turn angles during the escape responses occurred in stage 2 while maximum angular velocity and acceleration both occurred during stage 1 (Fig. 3). Stage 1 angle correlated significantly with stage 2 angle in all viscosities ($R^2=0.80$, regression slope=1.2, $P<0.0001$ in 1 mPa s water; $R^2=0.52$, regression slope=1.4, $P=0.0007$ in 10 mPa s fluid; and $R^2=0.44$, regression slope=1.0, $P=0.0036$ in 20 mPa s fluid).
In all viscosities, maximum body bending proceeded anteroposteriorly without any significant differences among the viscosity treatments (Fig. 4A). The time to maximum angular acceleration and time to maximum COM velocity were significantly affected by the viscosity treatments (Fig. 4B; Table 1).

Univariate ANOVA tests were carried out for all 25 kinematic variables common to both visually and mechanically stimulated turns, as the overall MANOVA on the viscosity effect was significant (Wilks’s lambda, \( P=0.0006 \)). MANOVA of the first four principal components for the data set separated by each a priori kinematic category (steady, unsteady, stage 1 and stage 2) were all significant (Wilks’s lambda, \( P=0.0001, 0.0034, 0.0002 \) and 0.0048, respectively), indicating that changing viscosity had a significant overall effect on each stage as well as on variables separated by proposed locomotor pattern.

The discriminant function analysis was able to correctly categorize 94% of the escape responses based on the 21 kinematic variables used (Fig. 5). The first two canonical functions explained 100% of the variation (83.5% and 16.5%, respectively). The variables with the two highest positive and the two lowest negative coefficients for canonical 1 were the following: linear displacement of the COM after 30 ms (4.88), linear displacement of the COM at time to maximum angular velocity in stage 1 (–0.16). The variables with the two highest and two lowest coefficients for canonical 2 were: time to maximum turn angle (0.29), linear displacement of the COM at the end of stage 2 (0.11), linear displacement of the COM after 30 ms (–1.04) and time to maximum angular velocity in stage 1 (0.20).

**DISCUSSION**

In this paper, we explored the influence of viscosity on C-start escape behaviors by studying the response of a classic high-speed behavior to substantial changes in the external medium. Our multivariate statistical analyses show a statistically significant overall effect of viscosity, indicating that fish displayed altered kinematics as the viscosity of water was increased to 10 and 20 times that of normal water. And the discriminant function analysis showed that 94% of the C-starts could be correctly classified using the kinematic variables that we extracted from the escape responses. We expected a priori that duration and timing variables would show an increase in magnitude as viscosity increased and that there might be differences in how kinematic variables that reflect velocity responded to viscosity changes compared with those variables that reflect body accelerations.

Thus, we were surprised to find that variables such as the duration of stage 1 and stage 2 did not show a viscosity effect and that several other variables such as final turn angle and maximal turn angle, and the time to reach many of the maximal velocity and acceleration variables were also not affected by viscosity (Table 1). In addition, the finding that variables such as maximum angular acceleration and the timing of some acceleration maxima did not differ in the increased viscosity treatments was unexpected.

Nonetheless, increasing water viscosity did have several significant effects on escape kinematics and did not have the same effect on all aspects of the C-start. Whereas normal escape responses happened exclusively in the inertial regime based on Reynolds number calculations throughout the escape response (Fig. 2B), a significant proportion of escape responses in high viscosity occurred either in the viscous or the intermediate regimes. The Reynolds number (the ratio of inertial to viscous forces) cut-off points for these regimes in the zebrafish *D. rerio* were shown experimentally.
to be 300 for the upper end of the viscous regime and 1000 for the lower end of the inertial regime (McHenry and Lauder, 2005). In the viscous regime, motion should be dominated by water viscosity, body length and velocity. In contrast, motion in the inertial regime is affected by the water density, the wetted surface area of the fish body and the square of velocity (Vogel, 1994).

Escape responses in 10 times the viscosity of water only reached the inertial regime 75% into the duration of stage 1 (Fig. 2B), with approximately the first 10% of this stage under viscous conditions and the remaining 65% under intermediate hydrodynamic conditions. At 20 mPa s fluid, only a small fraction (~10%) of the turn occurred under dominantly inertial forces while the majority of the turn was performed in the intermediate hydrodynamic regime (Fig. 2B).

Evaluating hypotheses of viscosity effects
Escape responses are vital to a fish’s survival. For this reason, fish muscles are likely to operate near their limits for producing power during these brief events, and this view has been commonly argued in the literature (e.g. Franklin and Johnston, 1997; Johnston et al., 1995; Wakeling and Johnston, 1998). Consequently, we expected that an increase in fluid viscosity would result in slower escape responses. However, the duration of each stage and hence of the whole turn did not change in high viscosity (Table 1, Fig. 3G), hinting at similar timings of hydrodynamic force generation during stage 1 and 2 and for a limited effect on the hydrodynamics of fin function (Tytell and Lauder, 2008). This result also suggests that the duration of neuromuscular stimulation was unchanged despite the novel hydrodynamic environments, a finding in concert with the low amount of variation previously observed for this stage in normal water (Domenici and Blake, 1991; Tytell and Lauder, 2008). If higher viscosity environments require increased muscle power production for motions of the same duration, then myomeres may develop increased power production through an increase in stimulation frequency, or through increased muscle fiber recruitment. This may be true even if myotomal musculature is activated for similar durations in all viscosities. It is not currently known whether all myotomal muscle fibers are activated maximally during a zebrafish C-start escape behavior, but changes in activation of different regions of myotomes have been documented for other locomotor behaviors in other species (Jayne and Lauder, 1995). Models of myotomal muscle function commonly assume that all fibers within each myotomal segment are activated completely during behaviors that require white myotomal fiber recruitment, but increased power production could be achieved through recruitment of additional fibers not normally activated even during the C-start behavior, commonly believed to be a maximal behavior in normal viscosity water.

When Mauthner cells activating the escape response in fishes are directly stimulated, stage 1 duration and angle are constant (Wakeling, 2006) and both directly correlate with the duration of muscle activity in stage 1 (Eaton et al., 1988). Stage 2 angle (final angle in this study) has been shown to correlate with stage 1 angle, giving rise to the term ‘ballistic’ to describe the neural command of this behavior (Eaton and Emberley, 1991), as initiation of the turn by the Mauthner cells controls the amount of body bending and as a consequence the final angle of an escape turn. However, stage 2 angle responded differently from stage 1 angle to changing viscosity in our experiments, and also was not linearly related to viscosity: stage 2 angle decreased more in 10 mPa s fluid than it did in 20 mPa s fluid, while stage 1 angle decreased by the same amount in both high viscosity treatments (Fig. 3D). Stage 1 and 2 angles were also uncorrelated in a study of larval zebrafish raised in high viscosity environments (Danos, 2012).

Swimming kinematics are ultimately a product of the interaction between the fluid, the fish body and the contractile properties of the underlying musculature. An interesting simulation study recently demonstrated the effect of viscosity on lamprey swimming kinematics (Tytell et al., 2010). By keeping all parameters of the simulation constant, including neuromuscular input and body stiffness, but increasing viscosity to 10 times that of water (10 mPa s), the authors were able to observe the following kinematic changes in simulated swimming: a 50% reduction in swimming speed and initial acceleration, and an order of magnitude increase in the relative amount of muscle power required to swim at a steady speed (Tytell et al., 2010). Although the effect of viscosity on linear velocity is not nearly as pronounced in real fish performing escape turns, there is a significant effect of viscosity on maximum linear velocity and displacement of the center of mass (Table 1). This indicates that even under the unsteady and high Reynolds conditions of a fast-starting fish, viscous forces play a significant role.

Using viscosity as an environmental manipulation
The manipulation of water viscosity is a powerful tool to assist in the dissection of organismal biomechanics, and a number of studies have applied this approach on a wide range of organisms. In Table 2 we summarize a selection of recent studies manipulating viscosity and add notes on the specific viscous agent used and selected information on some of the conclusions drawn.

There are a variety of possible agents that could be added to water to increase viscosity, but we believe it is important to emphasize that for studies that involve rapid motion, a Newtonian agent such as Dextran should be used. The shear thinning that results from rapid motion in a non-Newtonian fluid could certainly affect results, and undermine attempts to maintain a constant treatment viscosity effect when motions differing in acceleration are studied.

A number of the studies mentioned in Table 2 have used viscosity manipulations to compensate for changes in water properties as temperature is altered. Temperature can have dramatic effects on organismal function and it is often useful to quantify these effects by calculating \( Q_{10} \) (e.g. Podolsky and Emlet, 1993). However, a 10°C increase in the temperature of water from 10°C to 20°C also causes a 23% decrease in both dynamic and kinematic viscosity. Kinematic viscosity decreases slightly faster than dynamic viscosity (e.g. 23.18% compared with 23.33% in the example above) and hence studies that use viscosity manipulations while testing for the effects of temperature tend to report kinematic viscosity (m² s⁻¹). Within the range of natural temperature and viscosity fluctuations, most studies found that the physiological effects of temperature far outweighed any physical effects of viscosity, except for organisms of very small sizes such as bacteria operating at extremely low Reynolds numbers (Beveridge et al., 2010).

In this study, we used a viscosity manipulation to show that fishes, even when challenged with a 20-fold increase in water viscosity, are able to perform escape behaviors of similar duration and stage 1 angle to those executed in normal water. We did observe several specific alterations in escape kinematics and multivariate analysis confirms a significant overall viscosity effect, although the effects were not along the predicted lines of steady versus unsteady kinematics. Our results therefore suggest that fishes may be able to generate greater muscular power than has been suspected when confronted with a more viscous medium, and that activations of myotomal musculature in normal water, even for the C-start, may not be maximal. Alternatively, the complex hydrodynamics of the
intermediate hydrodynamic regime in which a large proportion of the observed escape responses occurred may lead to an even more complex muscle–fluid interaction. Such an interaction may lead to changes in the timing of hydrodynamic force production interplay with the muscle’s intrinsic force–velocity properties.

**Sensory input and viscosity**

The role of sensory input during rapid fish behaviors such as the escape response is still a topic of active investigation, and there are several possible views of how sensory systems could modulate escape kinematics. On the one hand, there is a high correlation of stage 1 and stage 2 angles in escaping fish under normal viscosity conditions, suggesting a ballistic-type neural control of escape that is independent of sensory information gathered during the initial stages of the turn. On the other hand, there is a high degree of consistency in stage 1 angles of zebrafish larvae C-starts despite substantial differences in water viscosity (Danos, 2012), and the use of prey position information in predicting the kinematics of archer fish predatory attacks (Wohl and Schuster, 2007). Archer fish shoot insects out of trees with a bolus of water and then use escape response kinematics to move to the location where they predict the insect will fall on the water surface (Wohl and Schuster, 2006; Wohl and Schuster, 2007). The fact that archer fish can integrate sensory information with escape response kinematics suggests that not all neural control pathways of this behavior are deterministic.

Interesting recent work (McHenry et al., 2009a; Windsor and McHenry, 2009) has begun to examine the interaction between the mechanical properties of neuromasts and hydrodynamics. When quantifying the time between the impact of a dropped object onto the water surface and the first movement of the fish’s head (latency), we found a tendency (though non-significant) for latency to decrease as viscosity increases (Fig. 3A). Larval zebrafish had a latency of 13–15 ms when stimulated with a uniform flow field (McHenry et al., 2009b), while the mean latency of the adult zebrafish in this study was 8.9 ms in normal water, 8 ms in 10 mPa s fluid and 7.1 ms in 20 mPa s fluid. The tendency towards increased sensitivity at high viscosity likely represents the increased drag acting on superficial neuromasts resulting in increased wall shear stress around the superficial neuromasts, similar to the increase in sensitivity observed when the effective cross-sectional area of artificial neuromasts is increased (Peleshankov et al., 2007; Windsor and McHenry, 2009). Analysis of the role of fish sensory systems in modulating rapid behaviors such as C-starts is an interesting area for future investigation.

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**Table 2. Summary of selected studies that have used viscosity manipulations to study organismal movement**

<table>
<thead>
<tr>
<th>Study</th>
<th>Viscous agent</th>
<th>Kinematic viscosity (m² s⁻¹)</th>
<th>Dynamic viscosity (Pa s)</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beveridge et al., 2010</td>
<td>Ficoll</td>
<td>1.0 × 10⁻⁶ to 1.5 × 10⁻⁶</td>
<td></td>
<td>Temperature-dependent viscosity had a significant effect on the carrying capacity and growth rates of consumers (bacteria), as well as the average density of the top predator (ciliates).</td>
</tr>
<tr>
<td>Horner and Jayne, 2008</td>
<td>Poly-Bore</td>
<td>10, 100 and 1000</td>
<td></td>
<td>Shear-thinning, non-Newtonian thickening agent used to study lungfish locomotion in mud-like conditions. Lungfish did not switch to a terrestrial motor pattern at high viscosity.</td>
</tr>
<tr>
<td>Hubley et al., 2001</td>
<td>Ficoll</td>
<td>4% (w/v)</td>
<td></td>
<td>Distinguished the effects of temperature and viscosity on the locomotor capacity of juvenile annelid polychaetes. Found no effect of viscosity.</td>
</tr>
<tr>
<td>Hunt von Herbing and Keating, 2003</td>
<td>Methyl cellulose</td>
<td>1.32 × 10⁻⁸ to 2.2 × 10⁻⁸</td>
<td></td>
<td>Decreasing viscosity effects and increasing physiological effects of temperature on swimming performance with increasing size of larval Atlantic haddock.</td>
</tr>
<tr>
<td>Johnson et al., 1998</td>
<td>Dextran (M 242,000)</td>
<td>1, 1.6 and 3.4</td>
<td></td>
<td>Reduction in temperature from 20 to 5°C also causes a 1.5-fold increase in viscosity. However, there was little effect of viscosity on C-start kinematics even for a small fish like Poecilia.</td>
</tr>
<tr>
<td>Korta et al., 2007</td>
<td>Methyl cellulose</td>
<td>0.05–50</td>
<td></td>
<td>Swimming gait of C. elegans does not vary across the range of viscosities but the temporal frequency of the swimming gait decreases by ~20% with every 10-fold increase in viscosity.</td>
</tr>
<tr>
<td>Podolsky and Emlet, 1993</td>
<td>Polivinyl pyrrolidone</td>
<td>1.02 × 10⁻³ to 1.3 × 10⁻³</td>
<td>8 × 10⁻³</td>
<td>Calculated Qₙₜ for sand dollar larvae by factoring in viscosity effects.</td>
</tr>
<tr>
<td>Pate and Brokaw, 1980</td>
<td>Ficoll</td>
<td>1.1 × 10⁻² to 8 × 10⁻³</td>
<td></td>
<td>Compared locomotor effects of ficoll to methyl cellulose study in sea urchin spermatozoa. Found reduced beat frequency in higher viscosity.</td>
</tr>
<tr>
<td>McHenry and Lauder, 2005</td>
<td>Dextran</td>
<td>1.2 × 10⁻³ to 18 × 10⁻³</td>
<td></td>
<td>Coasting and drag over zebrafish ontogeny. Used viscosity to modify Reynolds number and its effect on drag coefficients.</td>
</tr>
<tr>
<td>Kanou et al., 2007</td>
<td>Methyl cellulose</td>
<td>32, 364 and 1344</td>
<td></td>
<td>The relative occurrence of walking in the insect Gryllus bimaculatus increased with viscosity suggesting that a reacting force from the substrate to the legs is one of the factors important in triggering walking.</td>
</tr>
<tr>
<td>Neugebauer et al., 1998; Jordan, 1998</td>
<td>Percoll, Poly-Ox</td>
<td>Up to 1.5 × 10⁻⁳ and 0.06</td>
<td></td>
<td>Central and peripheral graviperception in ciliates. Scale effects in the kinematics and dynamics of swimming leeches. Significant kinematic changes in high viscosity.</td>
</tr>
</tbody>
</table>
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