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Isolating the hydrodynamic triggers of the dive response in eastern oyster larvae

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Abstract

Understanding the behavior of larval invertebrates during planktonic and settlement phases remains an open and intriguing problem in larval ecology. Larvae modify their vertical swimming behavior in response to water column cues to feed, avoid predators, and search for settlement sites. The larval eastern oyster (Crassostrea virginica) can descend in the water column via active downward swimming, sinking, or "diving," which is a flick and retraction of the ciliated velum to propel a transient downward acceleration. Diving may play an important role in active settlement, as diving larvae move rapidly downward in the water column and may regulate their proximity to suitable settlement sites. Alternatively, it may function as a predator-avoidance escape mechanism. We examined potential hydrodynamic triggers to this behavior by observing larval oysters in a grid-stirred turbulence tank. Larval swimming was recorded for two turbulence intensities and flow properties around each larva were measured using particle image velocimetry. The statistics of flow properties likely to be sensed by larvae (fluid acceleration, deformation, vorticity, and angular acceleration) were compared between diving and non-diving larvae. Our analyses showed that diving larvae experienced high average flow accelerations in short time intervals (approximately 1-2 s) prior to dive onset, while accelerations experienced by nondiving larvae were significantly lower. Further, the probability that larvae dove increased with the fluid acceleration they experienced. These results indicate that oyster larvae actively respond to hydrodynamic signals in the local flow field, which has ecological implications for settlement and predator avoidance.

Many marine invertebrates have a planktonic larval dispersal period before settling to the seafloor as adults. Our understanding of how larval behavior may influence dispersal and transport across a range of spatial scales is limited (Metaxas and Saunders 2009), and larval responses to a variety of physical, chemical, and biological cues remain ongoing areas of research. Larval swimming can be impacted by turbulent flow fields, especially in the turbulent bottom boundary layer as larvae move toward the substratum (e.g., Butman 1987; Butman et al. 1988). However, the impact of turbulent flow on the behavior of individual larvae is not well characterized due to technical challenges in simultaneously quantifying larval swimming and the motion of the surrounding flow field. Recent advances (Fuchs et al. 2013; Wheeler et al. 2013) are now making such studies feasible.

Small swimming organisms in a turbulent ocean experience a complex fluid environment, and may potentially respond to different components of ambient flow conditions, such as temporal velocity gradients (acceleration), spatial velocity gradients governing fluid deformation and rotation (strain rate and vorticity, respectively), and temporal vorticity gradients (angular acceleration). Rapid behavioral responses to local flow conditions are better studied for zooplankton than for larvae: threshold flow deformation has been observed to trigger escape responses in copepods (Kiørboe et al. 1999) as well as multiple protists (Jakobsen 2001). Acceleration, meanwhile, has not been observed to produce a similar response, although both acceleration and deformation are strong components of the suction flow fields produced by feeding predators (Kiørboe et al. 1999; Jakobsen 2001; Holzman and Wainwright 2009). In vortical flows, small organisms (ranging from bacteria to larvae) tilt and reorient, a response that has been attributed to a physical mechanism involving the balance of viscous and

Additional Supporting Information may be found in the online version of this article.

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gravitational torques acting on the organism (*see* e.g., Jonsson et al. 1991; Pedley and Kessler 1992; Chan 2012). In this study, we focus on the larvae of the eastern oyster, *Crassostrea virginica*, to increase our understanding of rapid behavioral responses of marine invertebrate larvae, and bivalves particularly, to flow conditions that they might experience in the field.

We chose oyster larvae for this study because they exhibit intriguing swimming behaviors in turbulent flows characteristic of coastal benthic habitats. They swim using a ciliated velum and so control their own swimming direction in still water, likely sensing their orientation and swimming direction with respect to gravity using a statocyst structure (Galtsoff 1964). A specific behavior of interest in oyster larvae is a response known as "dive-bombing" or "diving" (Finelli and Wethey 2003; Wheeler et al. 2013). Herein, we consider diving as a transient response occurring over timescales of approximately 1 s, where larvae abruptly accelerate downward, achieving speeds up to 1 cm s^{-1} , or approximately 50 body lengths s⁻¹, which is distinct from the sustained slower downward swimming behavior defined as diving in Fuchs et al. (2013). Diving, as we have defined it, has been observed in a moderately turbulent channel flow (Finelli and Wethey 2003), and in low turbulence induced by a grid-stirred tank (Wheeler et al. 2013). The cue or cues triggering the onset of the dive response are not well understood: some population-level estimates of larval swimming velocity in flow suggest that downward swimming increases in high turbulence (Fuchs et al. 2013), while others suggest that larvae persist in upward swimming in high turbulence, and further, that the dive response disappears in highly turbulent flow (Wheeler et al. 2013). As larval swimming responses in turbulence appear to be highly variable at the population level, we seek to identify specific triggers experienced consistently by larvae immediately prior to dive onset. It is important to identify these cues because through diving, a larva can rapidly displace itself downward through the water column. This behavior may therefore impact larval supply to the benthos, as diving may help larvae avoid predators and/or identify and approach suitable settlement sites.

Larvae settling into oyster reefs and other complex benthic structures experience a complex fluid environment which may impact settlement patterns (e.g., Nowell and Jumars 1984; Butman 1987; Koehl 2007). Current field research on oyster reefs suggests a link between oyster larval settlement patterns and turbulent flow over regions of settlement. Whitman and Reidenbach (2012) observed that turbulent drag and shear fields were considerably higher over live oyster reefs than mud flats and restoration reefs made of broken oyster or whelk shells. Larvae were observed to settle preferentially on oyster reefs, followed by whelk shell restoration sites, then oyster shell restoration sites, and not at all on mud flats. Settlement patterns suggest that flow fields generated by rough relief and low levels of turbulence in interstitial spaces may abet larval recruitment. Because oyster larvae display a dive response in turbulent conditions, we want to determine whether or not larvae dive in response to local hydromechanical cues in the turbulent flow field, such as flow acceleration, deformation, vorticity, or angular acceleration.

When transitioning out of the water column to the benthos, oyster larvae experience turbulent flow fields that may induce rapid downward diving responses. In this study, we actively quantify the diving response observed in two turbulence regimes, and determine which (if any) local hydromechanical signals induce the response, as well as the response timescales. Further, we use a Bayesian approach to calculate probabilities of larval diving conditioned on specified local hydromechanical conditions (e.g., the probability of a larva diving, supposing it has experienced a specified flow acceleration for a specified length of time). This relationship may be useful for understanding the ecological implications of larval responses in specific field conditions, and for the integration of behavior into larval models. We determine these diving triggers by identifying diving larvae and their local flow conditions in experimentally generated grid-stirred turbulence, then comparing the conditions experienced by diving and non-diving larvae as they move through the turbulent fluid environment.

Methods

Experimental organism and larval culturing

C. virginica, the eastern oyster, is a mollusc species native to the North Atlantic. Adults inhabit coastal shallow waters and broadcast spawn into the plankton, where larvae reside as free-swimming planktotrophs for 2–3 weeks (Kennedy 1996). Larvae entering the final planktonic stage, referred to as pediveligers, develop a foot and commonly a pronounced eyespot which are used in aquacultural practice to denote competency to settle (Thompson et al. 1996).

We obtained such competent larvae from the Aquaculture Research Corporation in Dennis, Massachusetts, U.S.A., in three separate spawns in the summers of 2011, 2012, and 2013. All spawns were retained prior to experiments in identical culture conditions: 3 µm-filtered, aerated seawater at ambient field temperature (20-22°C) and salinity (33 psu), in covered 16 L plastic buckets. Larvae were kept at low densities to minimize interactions (~ 3000 larvae L⁻¹) and fed a suspension of haptophyte Isochrysis sp. once per day (375 mL filtered seawater with $\sim 9 \times 10^5$ cells mL⁻¹.) Larvae were given a minimum period of 8h to acclimate post-transport from the aquaculture facility, and used for experiments within 2d of competency onset. A representative sample of larvae from the 2013 spawn were measured and examined for evespots prior to their use in experiments: average larval width (perpendicular to hinge) was $\sim 277 \ \mu m$, average height (parallel to hinge) was $\sim 264 \ \mu m$, and percentage of larvae with eyespots was > 80%.

Experimental setup

The turbulence tank used in the experiments (*see* Wheeler et al. 2013 for schematic) consists of a \sim 180L plexiglass tank (44.5 × 44.5 × 90 cm) with two horizontal grid structures set equidistant from the centre of the tank, connected by vertical rods in each corner. The grid structures are made from 1 cm × 1cm plexiglass bars spaced 5 cm apart. Both grids are connected to a motor above the tank by a vertical rod, which drives a simultaneous vertical oscillation in the grids. The oscillation amplitude is 5 cm and the oscillation frequency is specified by the user to induce flow fields of different turbulence intensity.

In the analysis described in this study, the larvae were subjected to two turbulence levels, hereafter referred to as "unforced" and "forced" regimes: the first regime has no flow induced in the tank (i.e., the grid frequency is 0 Hz) and the second regime has low forcing conditions with a grid frequency of 0.25 Hz. The forced regime has an estimated energy dissipation rate of 2×10^{-3} cm² s⁻³, and has Kolmogorov and integral length scales of 0.14 cm and 3.02 cm, respectively, roughly comparable to calm field conditions in tidal channels and estuarine flows (Gross and Nowell 1985). Note that although the grid was not operating in the unforced case, there was weak turbulent flow in the tank due to residual motions and possibly convection. The original experiments additionally subjected larvae to more highly turbulent flow conditions with dissipation rates ranging from $0.017 \text{ cm}^2 \text{ s}^{-3}$ in a moderate turbulence regime to $0.667 \text{ cm}^2 \text{ s}^{-3}$ in the most highly turbulent regime, and associated Kolmogorov and integral length scales ranging from 0.08 cm to 0.03 cm and 3.64 cm to 3.59 cm, respectively. These regimes were not examined in our present study because the larval diving behavior disappears in more highly turbulent flow (see Wheeler at el. 2013).

A vertical cross-section in the centre of the tank was illuminated by a pulsed near-infrared laser (Oxford Lasers, Firefly 300 W, 1000 Hz, 808 nm) in a plane approximately 1 mm thick. A high-speed monochrome camera (Photron Fastcam SA3, 1024×1024 pixel resolution) was trained perpendicularly to the laser sheet, recording a $\sim 3 \times 3$ cm twodimensional (2D) field of view (FOV).

The tank was maintained in an environmental chamber of fixed temperature (20°C) and filled with surface seawater filtered to particle size < 1 μ m. Larvae were gently introduced into the tank using a beaker to densities of 0.5–0.62 larvae mL⁻¹. The tank was subsequently seeded with a 2.5 mL suspension of neutrally buoyant polystyrene passive particles (3.0–3.4 μ m diameter, 1.05 g cm⁻³density, 5% weight by volume, Spherotech, Lake Forest, Illinois, U.S.A.) to a density of ~ 4.2 × 10⁴ particles mL⁻¹ for flow quantification by particle image velocimetry (PIV). Preliminary experiments showed no effects of these artificial particles on larval swimming in still water, when compared to both swimming in control filtered seawater and seawater seeded with natural *Isochrysis* algae (of roughly comparable size and concentration), leading us to conclude that artificial particles could be used in turbulence experiments without affecting behavior.

Larval behavior was recorded for 5-6 separate 45s intervals at 60 fps (with the number of intervals depending on the spawn and the turbulence level). These intervals were separated in time by approximately 5 min each to transfer images from the camera to the computer as TIFF files (e.g., Fig. 1A). Experiments were conducted under identical conditions over three separate 2 day periods in the summers of 2011, 2012, and 2013, corresponding to three separate spawns. Larvae were subjected to multiple randomly ordered turbulence levels, although only the two lowest turbulence regimes were examined in this study. Turbulence treatment order has no observed effect on larval swimming velocity (Wheeler et al. 2013), so eliminating measurements from these higher turbulence levels should not affect our results. Separate batches of larvae were also pooled for this analysis. Analyses of mean vertical swimming velocities in higher turbulence regimes, and separated by larval batches, are presented for the 2011 and 2012 data in Wheeler et al. (2013).

Larval tracking and local flow subtraction

The following methodology for isolating larval swimming velocity from advection in the local flow field was presented in Wheeler et al. (2013) and is summarized here, with the added refinement of interpolating local flow velocities to larval positions. First, larvae were identified by the following method: all TIFF files were imported into LabVIEW 2010 (National Instruments) and average background intensity was subtracted. Larval centroid positions (*x* and *z* coordinates) were identified using a fixed threshold particle size and intensity and recorded along with larval size, in the frame which they appeared.

Second, observed larval trajectories were computed using an in-house MATLAB script which tracked identified larvae from frame to frame according to a subsequent-frame tolerance distance radius set by the user. Larval trajectories were truncated by five frames at both the beginning and end of the trajectories due to uncertainties in centroid estimates in cases where larvae passed laterally into and out of the focal plane, which caused larvae to appear diffuse and out of focus. Instantaneous observed larval velocities, denoted u_{obs} = $[u_{obs}, w_{obs}]$ for each larva, were computed using a central difference scheme of larval centroid position in time, so that the velocity is defined centered in time between the two images.

Third, fluid velocity fields in the FOV were quantified using PIV imaging software LaVision DaVis (v.7.2). All TIFF files were imported into the software and velocity fields were computed using correlations (default FFT with Whittaker reconstruction) of most likely passive particle positions from



Fig. 1. (A) Sample image from FOV in the turbulence tank: larvae are bright white spots and polystyrene passive particles are small dim white specks. (B) Close up of individual larva (white spot) overlaid with annulus of local flow velocity field (white arrows) estimated using PIV. (C–E) Sample time series of diving (black curve) vs. non-diving larva (grey curve), where the vertical black dashed line denotes dive onset time: vertical displacement due to larval swimming (C), vertical swimming velocity w_s (D), and flow acceleration magnitude |a| experienced by each larva (E).

frame to frame, using 16×16 pixel interrogation windows (with 7–8 particles per window, not distinguishable by eye in Fig. 1A,B). This process yielded two 64×64 spatial grids of horizontal and vertical flow velocity for each time step, corresponding to a grid spacing of 0.039–0.046 cm (varied slightly by spawn).

Fourth, fluid velocities local to larvae were subtracted from observed larval velocities to obtain larval swimming velocities by the following method. The velocity fields estimated by PIV were imported and converted to MATLAB data files and velocity vectors in an annulus around each larva were used to estimate the fluid velocity at the larval position at each time step. The radius of the annulus changed dynamically for each larva: the inner radius was the sum of the maximum individual larval radius and the grid spacing of the PIV data (16 pixels), and the outer radius was four times greater than the inner radius (Fig. 1B). The inner radius of the annulus masked the larval presence in the PIV data, which might otherwise contaminate the PIV analysis for fluid velocity. The velocity data in the annulus were fit to a 2D, second-order Taylor series function by least-squares. The flow velocity $\boldsymbol{u}=[\boldsymbol{u},\boldsymbol{w}]$ local to a larva was then obtained by evaluating the function at the larval centroid position. This interpolated fluid velocity was subtracted from the observed larval velocity at that time step to obtain the larval swimming velocity $\boldsymbol{u}_s=[\boldsymbol{u}_s, \boldsymbol{w}_s]$. For each larva,

$$\boldsymbol{u}_{s} = \boldsymbol{u}_{obs} - \boldsymbol{u}_{s}$$

Identification of dive response

The dive response was initially observed by eye in experimental footage and in individual larval vertical swimming velocity time series, where it was characterized by a rapid drop to high downward swimming velocities, followed by a slow deceleration over the span of several seconds to nearzero vertical swimming velocity. We described a larva as diving if it performed downward accelerations of at least 3.0 cm s^{-2} (approximately 150 body lengths s^{-2}) for minimally two time steps (1/30 s) and achieved negative vertical swimming velocities of at least -0.4 cm s^{-1} . These thresholds in vertical swimming acceleration and velocity were used to separate diving larvae from non-diving larvae in the subsequent analysis (example difference between diving and non-diving larvae velocity time series, Fig. 1D).

Hydromechanical parameters detectable by larvae

In this section, we propose a suite of hydromechanical cues in the turbulent flow that are likely to be detectable by larvae. Because larvae can be divided into divers and nondivers, relevant potential cues experienced by these two groups (Fig. 1E) can then be compared for statistical differences. Following Kiørboe and Visser (1999), one may isolate the various aspects of a turbulent flow to which a larva might respond. Potentially relevant hydromechanical triggers are fluid acceleration, deformation (strain rate), rotation (vorticity), and angular acceleration. Given a flow velocity \boldsymbol{u} local to a larva having swimming velocity \boldsymbol{u}_s , on any given time step, we can calculate the following acceleration, strain rate, vorticity, and angular acceleration fields.

Acceleration measures the rate of change in fluid velocity and could potentially be perceived by a larva through its statocyst structure: a calcareous statolith would be displaced into the wall of the statocyst cavity due to inertia in an accelerating flow (Chia et al. 1981; Fuchs et al. 2013). To characterize the temporal changes in flow velocity near an individual larva, we use the magnitude of the 2D acceleration of the fluid flow following the larval position (Maxey and Riely 1983) (*see* the Supporting Information for a derivation):

$$|\mathbf{a}| = \left| \frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} + \mathbf{u}_s) \cdot \nabla \mathbf{u} \right|$$

We use acceleration magnitude, with magnitude denoted by | · |, as a hydromechanical metric to incorporate both dimensions of the acceleration vector. This acceleration metric excludes the acceleration that a larva experiences due to its own swimming motion, accounting only for the acceleration the larva experiences due to the local flow field. Larval swimming velocity \boldsymbol{u}_s is present in $|\boldsymbol{a}|$ because both larval swimming and flow velocity contribute to larval position, hence the inclusion of both in the advection term. If larvae perceive acceleration using a statocyst, they would feel the total acceleration from both the flow and their own swimming (see Supporting Information). However, we focus on the externally imposed fluid acceleration because it is independent of all larval behavior: this simplifies the interpretation of our results, as we do not conflate the larval responses to internally imposed and externally imposed motion.

In practice, the flow acceleration above is calculated by interpolating flow velocity to the larval position at each time step, then using a central difference scheme to compute the temporal derivative along the larval path. While the acceleration magnitude used in this analysis uses only the two known dimensions (x, z) available from our PIV setup, the unknown *y*-acceleration component will be similar to that of *x*, due to tank and forcing symmetries. We estimated a three-dimensional (3D) acceleration magnitude by doubling the *x*-acceleration component and found that the 2D and 3D fluid acceleration estimates yield similar statistical results, so we report only the 2D results in the subsequent sections.

The velocity gradients in a fluid flow lead to shear stresses on the surface of any object or fluid parcel in that flow. The net effect of these shear stresses can be to strain (i.e., deform) and rotate the object or fluid parcel. The strain rate (quantified using the rate of strain tensor) determines how a fluid parcel is stretched or sheared in different spatial dimensions, and could potentially be detected by a larva at sufficiently high signal strength by a deformation of cilia along the velum. The rotation rate (quantified using the vorticity) is likely detectable through a larva's statocyst structure (Chia et al. 1981), as the statolith is displaced and rolls steadily along the statocyst cavity wall, imposing a centrifugal force.

Strain rate is quantified in a 3D flow by the symmetric strain rate tensor e_{ij} , elements of which describe the deformation of the flow along two axes. Because we have only two dimensions of velocity data, the full strain rate tensor cannot be computed, and we are restricted to the examination of three of the elements of the tensor: the shear strain rate e_{xz} and the normal strain rates e_{xx} and e_{zz} . We use the 2D shear strain rate magnitude at the larval position:

$$|e_{xz}| = \left|\frac{1}{2}\left(\frac{\partial u}{\partial z} + \frac{\partial w}{\partial x}\right)\right|.$$

This metric represents the shearing, or deformation, of a fluid parcel in the focal plane, and is calculated using flow velocities local to the larval position. We use the magnitude of the shear strain rate because the sign of this term simply governs the direction in which the shear deformation occurs, and we do not expect larvae to recognize or respond to this directionality.

Normal strain rates are

$$e_{xx} = \frac{\partial u}{\partial x}$$

and

$$e_{zz} = \frac{\partial w}{\partial z}$$
,

where these quantities measure how fluid is stretched in the x and z dimensions, respectively, calculated local to the

larval position at each time step. Unlike the shear strain rate, the signs of the normal strain rates are retained; positive normal strain rates indicate divergence in the specified spatial dimension, while negative normal strain rates indicate convergence in the specified spatial dimension, and these are physically distinct phenomena. For all strain rates, the spatial derivatives are calculated at the fluid velocity points in the annulus around each larva and then interpolated to the larval position using the method described for the velocity field in the local flow subtraction section.

Vorticity measures the rotation of a fluid parcel, and is likely detectable through a larva's statocyst structure, as described above. Vorticity is a 3D vector for a 3D flow, with each element describing the rotation of the fluid normal to a plane described by the other two dimensions. Because we have only two dimensions of velocity data, we are restricted to using the vorticity element normal to the focal plane as our vorticity metric:

$$|\omega_{\gamma}| = \left| \frac{\partial w}{\partial x} - \frac{\partial u}{\partial z} \right|$$

The vorticity is calculated local to larval position at each time step, with spatial derivatives calculated as described above for the strain rate metrics. Similarly to shear strain rate, we define our vorticity metric by the magnitude of the vorticity element: the sign of vorticity denotes the direction of rotation of the local fluid (clockwise vs. anticlockwise), which we do not expect the larvae to distinguish. In a simple parallel shear flow, vorticity is equal to the velocity gradient in a single direction, and we use vorticity in this study because it generalizes the shear metric commonly reported in simpler flows (Kiørboe and Visser 1999). Similar to the acceleration term defined above, this vorticity term accounts only for the fluid rotation around the larva and not the larva's own rotation term. The larval rotation term is not considered in this analysis; as above, the rationale is to separate external forcing imposed by the fluid from the internal forcing of the larva's own swimming motion.

Angular acceleration measures the rate of rotation of a fluid parcel, and may be detectable in the larval statocyst structure through the onset of statolith motion along the statocyst wall. To characterize the temporal changes in flow vorticity near an individual larva, we compute the magnitude of the angular acceleration of the fluid flow following the larval position:

$$|\alpha| = \left| \frac{\partial \omega_{\gamma}}{\partial t} + (\boldsymbol{u} + \boldsymbol{u}_{s}) \cdot \nabla \omega_{\gamma} \right|.$$

In practice, the angular acceleration is calculated by interpolating flow vorticity to the larval position at each time step, then using a central difference scheme to compute the temporal derivative along the larval path. To avoid confusion, in the following analysis and discussion, *acceleration* always refers to **a**, the rate of change of fluid velocity following larval paths, while *angular acceleration* specifically will be used to refer to α , the rate of change of fluid vorticity following larval paths.

Statistical analysis

In this study, one of our objectives was to determine differences in hydromechanical parameters (flow acceleration, normal and shear strain rates, vorticity, and angular acceleration) experienced by diving larvae and non-diving larvae. To determine this, we calculated mean hydromechanical parameters experienced by all diving larvae in a set temporal interval immediately prior to dive onset, and mean hydromechanical parameters in the same temporal interval (randomly selected in the individual larval trajectory) for non-diving larvae. We used means instead of maxima, as using mean values in short time intervals allowed us to capture peak hydromechanical parameter values while filtering out PIV noise that distorts the maxima. A randomly subsampled group of non-diving larvae were then selected to compare to the diving larvae, so that the sample size in both groups would be identical. Two conditional probability distributions were then constructed for comparative purposes: P(T| larva dives) and P(T| larva does not dive) for each mean hydromechanical parameter *T*.

The distributions of mean hydromechanical parameters experienced by diving larvae and non-diving larvae were then compared statistically using the following methods. If T was strictly non-negative (i.e., all magnitude terms) we used a nonparametric two-sided Wilcoxon rank sum test to compare the medians of the diving vs. non-diving distributions. If the distributions were drawn from both positive and negative values, we used a modified two-tailed *t*-test (Welch's approximate *t*-test statistic and Satterthwaite's approximation for the degrees of freedom) to compare the means of the distributions instead.

If a parameter was found to differ significantly between diving and non-diving larvae, both distributions were compared to the background distribution of the hydromechanical parameter, P(T), which was determined by computing Tthrough four fixed spatial points in the FOV over the three experiments (over comparable spatial and temporal scales to which T was computed for the larvae). The comparisons of diving, non-diving, and background T distributions were carried out using a nonparametric Kruskal–Wallis test. A multiple comparison test was subsequently carried out to identify whether hydromechanical parameters experienced by diving and/or non-diving larvae differed significantly from the average parameter values in the background flow. All statistical tests were carried out using MATLAB.

For any hydromechanical parameter which differed significantly between diving and non-diving larvae, the conditional probability of diving given a specified mean parameter value was calculated using Bayes theorem:



Fig. 2. Diving larval vertical swimming velocity time series in the unforced regime, aligned by dive onset time. Larvae display strong uniformity in time spent accelerating downward, maximum downward velocity, and time spent decelerating out of the dive. Larvae exhibit a range of vertical swimming velocities prior to dive onset.

$$P(\text{larva dives} \mid T) = \frac{P(\text{larva dives}) \cdot P(T| \text{ larva dives})}{P(T)}.$$

The probability of larval diving, P(larva dives), is the number of diving larval trajectories divided by the total number of trajectories observed, while P(T| larva dives) and P(T) are described above. The conditional probability of larval diving given a mean hydromechanical parameter value, P(larva dives | T), is an ecologically relevant function as it predicts larval behavior in response to specific environmental conditions.

A 95% confidence interval for this conditional probability was computed by summing in quadrature the independent confidence intervals from each term in the equation. Confidence intervals for P(T| larva dives) and P(T) were estimated by bootstrapping the distributions and directly computing the confidence interval for each value of T. The confidence interval for the scalar P(larva dives) was computed using the Clopper–Pearson method for binomial confidence intervals as the diving probability is a probability of success in a binomial trial (i.e., diving vs. non-diving).

Results

Identification of dive response

Using our quantitative definition of diving, we found that 82 larvae (of 874 total larvae) dove at least once during their observed trajectory in the unforced regime, and 57 larvae (of 1019 total larvae) dove at least once in the forced regime. We overlaid the diving trajectories aligned by dive onset time in the unforced regime (Fig. 2) to identify similarities in diving trajectories, and found similar timescales in the downward acceleration for all larvae, on the order of 0.1 s. Larvae reached peak downward velocities ranging from -0.5 to -0.7 cm s⁻¹ and decelerated to zero velocity in approximately 1 s. Prior to dive onset, larvae engaged in a range of vertical velocities, centered near zero, but both upward and downward swimming were observed, suggesting that larvae

had no fixed pre-dive behavior. As larvae decelerated from the dive and resumed a more constant vertical velocity, they exhibited a similar range of vertical velocities, indicating that larvae also had no fixed post-dive behavior. Vertical displacement from a single dive was of order 10^{-1} cm, or approximately four body lengths, and comparable to the Kolmogorov scale, the length scale of the smallest eddies in the forced regime.

Hydromechanical parameters triggering the dive response

A range of temporal intervals prior to the dive onset was investigated, from 0.33 s to 3 s, in intervals of 0.33 s (*see* Supporting Information) to identify potential reaction time-scales for diving larvae. A hydromechanical parameter was considered to be a consistent trigger to the dive response only if (1) it differed significantly between diving and non-diving larvae in the specified temporal interval, and (2) this significant response held in both flow regimes, unforced and forced, for identical temporal intervals.

Diving larvae consistently experienced significantly higher mean fluid acceleration than non-diving larvae. In the unforced regime, mean accelerations were significantly higher for diving larvae in the 1s, 1.33s, 1.66s, and 2s time intervals and intermittently significant for longer time intervals (Table 1; Fig. 3A). In the forced regime, mean accelerations were significantly higher for diving larvae in all time intervals from 1.33s to 3s prior to the dive onset (Table 1; Fig. 3A). The intersection of these temporal intervals is 1.33-2.33 s, representing the consistent response range in which diving larvae experienced significantly higher acceleration than non-diving larvae. For subsequent analyses presented in the main text, we used a central point of this interval, 1.66s prior to dive onset, as the averaging window and denote the mean acceleration experienced by a larva in this interval as **a**|1.66.

No other hydromechanical parameter differed significantly prior to dive onset between diving and non-diving larvae (Fig. 3B–F, Supporting Information Tables A1, A2) in Wheeler et al.

Table 1. Wilcoxon rank sum test comparing medians of mean acceleration distributions experienced by diving vs. non-diving larvae, where means are computed in the stated window prior to dive onset. The null hypothesis states that medians $M_d = M_{nd}$ while the alternate hypothesis states that they differ. Significance level is $\alpha = 0.05$, with bold results indicating significant *p*-values. The medians of mean acceleration distributions are significantly higher for diving larvae than non-diving larvae in both flow regimes, given at least a 1.33 s window over which local acceleration is averaged.

Time interval prior to dive	Turbulence	Rank		
onset (s)	regime	sum	Z	<i>p</i> -value
0.33	Unforced regime	6314	1.84	0.06
0.66	$\epsilon \to 0 cm^2 \; s^{-3}$	6270	1.67	0.09
1.00		6490	2.48	0.01
1.33		6735	3.39	< 0.001
1.66		6697	3.25	0.001
2.00		6406	2.17	0.02
2.33		6148	1.22	0.21
2.66		6626	2.99	0.002
3.00		6318	1.88	0.06
0.33	Forced regime	3095	1.63	0.10
0.66	$\epsilon = 10^{-3} \text{ cm}^2 \text{ s}^{-3}$	3076	1.51	0.13
1.00		3015	1.13	0.25
1.33		3166	2.08	0.03
1.66		3164	2.07	0.03
2.00		3325	3.08	0.002
2.33		3232	2.50	0.01
2.66		3144	1.94	0.05
3.00		3244	2.57	0.009

contrast to acceleration (Fig. 3A; Table 1). That is, none of shear deformation, normal deformation (horizontal or vertical), vorticity, or angular acceleration induced a diving response in larvae in any temporal window examined.

Flow accelerations experienced by diving and non-diving larvae in the 1.66s interval were then compared to background acceleration fields (Fig. 4). These three distributions of flow acceleration, $P(|\boldsymbol{a}|_{1.66}|$ larva dives), $P(|\boldsymbol{a}|_{1.66}|$ larva does not dive), and $P(|\boldsymbol{a}|_{1.66})$, were significantly different in the unforced regime (Table 2). A post hoc multiple comparison test of these distributions demonstrated that diving larvae experienced significantly higher average flow accelerations than both non-diving larvae and the average background acceleration. Non-diving larvae experienced flow accelerations that were indistinguishable from the background acceleration. In the forced regime, a similar pattern was observed: diving larvae experienced higher accelerations than did non-diving larvae, as well as higher accelerations than those occurring in the background flow. However, the result in this regime was non-significant (Table 2), likely

due to the smaller sample size of dives and lower power of the multiway comparison.

These distributions were then used to compute $P(\text{larva dives} | |\boldsymbol{a}|_{1.66})$, the conditional probability that larvae dove for a given acceleration averaged over the 1.66 s pre-dive window in the unforced regime (Fig. 5). The positive relationship between this probability and the acceleration demonstrates that diving became a more probable response as mean fluid acceleration experienced by larvae increased. The bounds on the 95% confidence intervals increased for high acceleration values due to the rarity of high acceleration events, which likely also accounted for overestimates of the conditional diving probability (i.e., greater than 1) for high accelerations. The computation is omitted for the forced regime as the large decrease in number of dives observed renders estimates much more uncertain.

Discussion

Comparisons of flow fields experienced by diving and non-diving larvae strongly support a conclusion that flow acceleration triggers the dive response in oyster larvae. Diving larvae experienced significantly higher mean fluid accelerations than did non-diving larvae during a short period leading up to the dive onset in both turbulence regimes. The other candidate hydromechanical parameters did not differ significantly between diving and non-diving larvae: none of mean normal strain rates, shear strain rate, vorticity, or angular acceleration triggered the dive response. An examination of diving in the central 1.66s response window demonstrated that not only did diving larvae experience higher accelerations than non-diving larvae, but that these accelerations were anomalously high compared to the background (significantly so in the unforced regime). The correspondence between probability of diving and increasing fluid acceleration further reinforces the interpretation that diving is triggered by acceleration. Further, the time interval over which the threshold mean acceleration was experienced was important for triggering the dive response. When acceleration was averaged over temporal windows shorter than 1.33 s, higher acceleration did not appear to induce diving preferentially. This analysis suggests that the reaction timescale of the larvae to the fluid acceleration field they experience was at least 1.33 s. A lack of pattern in timescales longer than 2s suggests that the larvae are responding to an acceleration event, roughly 1.5s before the dive, rather than to mean acceleration over a longer interval.

The observation that a mean acceleration of 0.035 cm s^{-2} triggered a dive in the unforced case, but not in forced case, indicates that the required threshold acceleration changes with the turbulence level. In the low-forcing regime, an average acceleration of 0.06 cm s^{-1} triggered a dive, while non-diving larvae experienced mean accelerations of 0.04 cm s^{-2} .

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Fig. 3. Values of hydromechanical parameters (mean and 95% confidence intervals) for diving larvae (black) and non-diving larvae (grey) in unforced and forced regimes. Values are calculated in a 1.66s time interval prior to the dive onset in diving larvae, and a randomly selected 1.66s time interval in the trajectories of non-diving larvae. Sample sizes are n = 82 for both groups in the unforced regime, and n = 57 in the forced regime. (A) Mean acceleration magnitude |a| is significantly different between diving and non diving larvae for both turbulence regimes (*see* Table 1). (B) Mean shear strain rate magnitude $|e_{xz}|$ experienced by diving and non-diving larvae is not significantly different in either turbulence regime (Supporting Information Table A2). (C–D) Mean horizontal and vertical normal strain rates e_{xx} and e_{zz} experienced by diving and non-diving larvae is not significantly different in either turbulence regime (Supporting Information Table A1). (E) Mean vorticity magnitude $|\omega_y|$ experienced by diving and non-diving larvae is not significantly different in either turbulence regime (Supporting Information Table A2). (F) Mean angular acceleration magnitude $|\alpha|$ experienced by diving larvae is not significantly different in either turbulence regime (Supporting Information Table A2). (F) Mean angular acceleration magnitude $|\alpha|$ experienced by diving larvae is not significantly different in either turbulence regime (Supporting Information Table A2). (F) Mean angular acceleration magnitude $|\alpha|$ experienced by diving and non-diving larvae is not significantly different in either turbulence regime is not significantly different in either turbulence regime (Supporting Information Table A2). (F) Mean angular acceleration magnitude $|\alpha|$ experienced by diving and non-diving larvae is not significantly different in either turbulence regime (Supporting Information Table A2).

This result suggests that larvae become conditioned to the flow regime in which they find themselves, and the dive response is triggered by anomalously high accelerations compared to the background acceleration. This interpretation is supported by the finding that the accelerations experienced by diving larvae were significantly higher than both nondiving larvae and the background field. In a previous study (Wheeler et al. 2013), the dive response was found to disappear entirely in highly turbulent flow conditions (having energy dissipation rates greater than 10^{-1} cm² s⁻³). While our experimental results do not provide a complete explanation for this disappearance, we offer several possibilities.



Fig. 4. Probability distributions of mean flow acceleration magnitude experienced by larvae in a 1.66s time interval (prior to dives for diving larvae, randomly selected for non-diving larvae), in unforced (A) and forced (B) regimes, respectively. The black bar distributions are those of diving larvae, $P(|\boldsymbol{a}|_{1.66}|$ larva dives), the grey bar distributions are those of non-diving larvae, $P(|\boldsymbol{a}|_{1.66}|$ larva does not dive), and the black dashed curves are background mean acceleration magnitudes $P(|\boldsymbol{a}|_{1.66}|$. Note the different acceleration scales in unforced and forced regimes.

Table 2. Kruskal–Wallis test comparing median average accelerations experienced by the following three groups: diving larvae in a 1.66 s window prior to dive onset, non-diving larvae in a random 1.66 s window, and four fixed spatial points over all three experiments in a random 1.66 s window. The null hypothesis states that medians of all three mean acceleration distributions are equal, and the alternate hypothesis states that the mean accelerations experienced by these groups are different. Significance level is $\alpha = 0.05$, with bold results indicating significant p-values.

Source	SS	df	MS	χ ²	р
Unforced r	regime				
Group	5.54×10^4	2	2.77×10^4	12.40	0.002
Error	9.71×10^5	228	4.26×10^4		
Total	1.02×10^{6}	230			
Forced reg	ime				
Group	9.76×10^{3}	2	4.88×10^3	4.38	0.11
Error	3.51×10^5	160	2.19×10^3		
Total	3.60×10^5	162			

First, larvae may simply stop reacting to an acceleration trigger above a certain threshold which occurs in the higher turbulence regimes. Second, recall that larvae respond to anomalously high accelerations within a turbulence level, and this threshold increases with turbulence intensity, at least in unforced and low forcing conditions. The frequency



Fig. 5. Probability of larval dive conditioned on $|\boldsymbol{a}|_{1.66}$, the local mean acceleration field (averaged over 1.66 s window), i.e., $P(\text{larva dives}||\boldsymbol{a}|_{1.66})$, for the unforced regime. Larvae were more likely to dive when they encountered higher local flow acceleration. Shaded grey region represents the 95% confidence interval for all mean accelerations.

with which larvae encounter sufficiently high acceleration anomalies in more turbulent regimes may be lower, which would explain the lack of diving in these regimes. However, we cannot quantify the diving threshold accelerations for these higher flow regimes (beyond supposing the thresholds are greater than that observed in our low flow forced regime), and as such, this explanation for the lack of diving in high turbulence remains speculative. Alternatively, it is possible that the experimental setup precluded detection of dives because larvae are advected quickly in more highly turbulent flow. It is possible that it becomes more difficult to observe the diving response because larvae remain in the FOV for shorter time periods (although more larvae are observed in higher turbulence regimes).

The dive response for all observed larvae was highly uniform in terms of acceleration and deceleration timescales (Fig. 2), and the response is predictable based on fluid acceleration through the probability $P(\text{larva dives} | |\mathbf{a}|)$. These characteristics make the dive response well suited for inclusion into individual based models of larval behavior in complex flow fields (see for instance Koehl et al. 2007). Such models would be very useful for testing whether diving affects settlement success in simulated turbulent flow fields over rough bottom topography. The strong uniformity of the dive further suggests that the response, once instigated, is regulated by biomechanical constraints, as all larvae emerge from the dive and resume swimming on comparable timescales. In this way, the diving response triggered by acceleration may differ from the sinking response to waterborne chemical cues observed in the larval sea slug Phestilla sibogae (Hadfield and Koehl 2004). These larvae retract velar lobes instantly in response to coral-conditioned seawater, and continue to sink unless the cue is absent on timescales of 1 s or longer. Our larvae, conversely, cease to dive after approximately 1 s regardless of local flow conditions. While the larvae are capable of diving multiple times in succession, their behavior appears distinct from the sustained sinking observed in P. sibogae larvae.

The effects of local environmental conditions on the behavior of mollusc larvae have been previously studied in a few species with varying results. Two bivalve larvae (Crassostrea gigas and Mytilus edulis) exposed to horizontal suction flow demonstrated no discernible swimming response as they approached a suction tube (Troost et al. 2008), a flow that would have a strong acceleration signal. However, the flow fields experienced by these larvae were quantified in a separate experiment from the larval observations. This technique can make it difficult to isolate larval behavior (Wheeler et al. 2013), as small scale temporal and spatial variations in the flow field that larvae might experience are not captured. P. sibogae retract their velar lobes in response to mechanical stimulus (Hadfield and Koehl 2004), and potentially to local hydrodynamic conditions (M. Koehl pers. comm.), as well as the potentially distinct response to chemical cues, as discussed above. The similarity of the response (retraction of ciliated swimming organ into a shell) in different mollusc groups suggests that larval diving in response to acceleration may be common to multiple species.

A dive response when larvae are experiencing anomalously high accelerations could potentially be a beneficial strategy if they need to settle onto rough bottom surfaces, or to avoid predator feeding currents. We consider both possibilities, beginning with the ecological implications of diving as a settlement response. PIV measurements over rough topography in an oscillating flow tank have demonstrated that the highest accelerations occur up to 5 cm from the bottom, and decay rapidly farther above (R. Pepper, J. Jaffe, E. Variano, and M. Koehl pers. comm.). Further, simulated larvae in the PIV-measured flow experience peak accelerations of short duration that are much higher in magnitude than the mean values, much like the anomalously high accelerations experienced by the diving larvae in our study. The threshold accelerations experienced by larvae in our unforced and forced regimes are small compared to the fluid accelerations near the bottom reported by Pepper et al., but may help larvae navigate downward through the water column at heights above 5 cm from the bottom. The dive response disappears in more highly turbulent flow regimes that more closely mimic the energetics of flow immediately above preferred settlement sites (e.g., Whitman and Reidenbach 2012), which offers further evidence the dive response is likely to be employed by larvae higher in the water column.

Larvae could alternatively experience flow acceleration due to suction feeding flows from predators in the plankton (Kiørboe et al. 1999; Jakobsen 2001; Holzman and Wainwright 2009) or even from the feeding currents of adult oysters on reefs (Troost et al. 2008). In this way, the dive could act as an escape response analogous to the jumping behavior of copepods (e.g., Waggett and Buskey 2007; Lee et al. 2010) or the rapid downward swimming of insect larvae and pupae (e.g., Awasthi et al. 2012) observed in the presence of predators. Larval dive responses to flow acceleration in the water column could thus increase larval supply to the seafloor, by either increasing the rate of downward flux, or decreasing the proportion lost to predators.

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