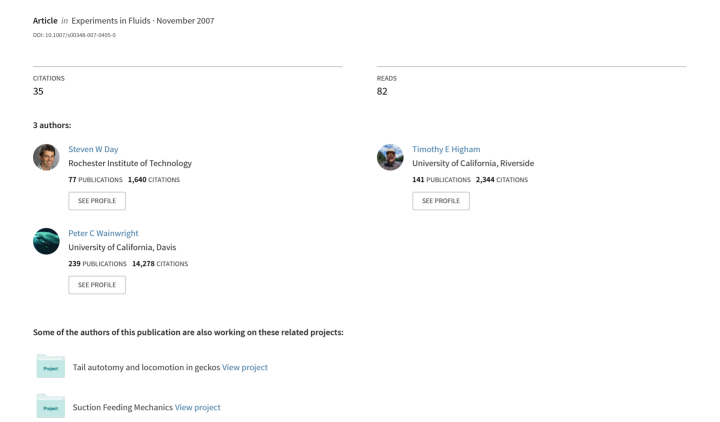
Time resolved measurements of the flow generated by suction feeding fish



RESEARCH ARTICLE

Time resolved measurements of the flow generated by suction feeding fish

Steven W. Day · Timothy E. Higham · Peter C. Wainwright

Received: 20 February 2007/Revised: 25 August 2007/Accepted: 15 September 2007/Published online: 12 October 2007 © Springer-Verlag 2007

Abstract The majority of aquatic vertebrates are suction feeders: by rapidly expanding the mouth cavity they generate a fluid flow outside of their head in order to draw prey into their mouth. In addition to the biological relevance, the generated flow field is interesting fluid mechanically as it incorporates high velocities, is localized in front of the mouth, and is unsteady, typically lasting between 10 and 50 ms. Using manometry and high-speed particle image velocimetry, this is the first study to quantify pressure within and outside the mouth of a feeding fish while simultaneously measuring the velocity field outside the mouth. Measurements with a high temporal (2 ms) and spatial (<1 mm) resolution were made for several feeding events of a single largemouth bass (Micropterus salmoides). General properties of the flow were evaluated, including the transient velocity field, its relationship to pressure within the mouth and pressure at the prey. We find that throughout the feeding event a relationship exists for the magnitude of fluid speed as a function of distance from the predator mouth that is based on scaling the velocity field according to the size of the mouth opening and the

magnitude of fluid speed at the mouth. The velocity field is concentrated within an area extending approximately one mouth diameter from the fish and the generated pressure field is even more local to the mouth aperture. Although peak suction pressures measured inside the mouth were slightly larger than those that were predicted using the equations of motion, we find that these equations give a very accurate prediction of the timing of peak pressure, so long as the unsteady nature of the flow is included.

1 Introduction

Most aquatic vertebrates, including fish, feed by generating a flow of water outside the head that acts to draw the prey into the mouth, a behavior known as suction feeding. The flow is interesting from a fluid mechanical perspective as it includes high spatial gradients of velocity and pressure and dramatic temporal accelerations with the entire prey capture lasting ~ 10 ms for many species. As is the case with locomotion through a fluid medium, feeding involves some direct fluid-structure interactions. Forces are a function of the biomechanical forces within the fish and the fluid mechanical pressure within mouth. The speed of cranial expansion is a function of the biomechanical kinematics within the animal and also the volumetric flow rate of the fluid. Force and speed of fish movement are related through muscle physiology and gearing within the animal, and pressure and fluid speed are related through the governing equations of fluid mechanics. As an example, an attempt by a fish to increase the speed of mouth opening increases the required force through interaction of the fluid field; increased fluid speeds lead to increased magnitude of generated pressures.

S. W. Day (🖂)
Department of Mechanical Engineering,
Rochester Institute of Technology,
76 Lomb Memorial Dr, Rochester, NY 14623, USA

e-mail: steven.day@rit.edu

T. E. Higham

Department of Organismic and Evolutionary Biology, Concord Field Station, Harvard University, 100 Old Causeway Road, Bedford, MA 01730, USA

P. C. Wainwright Section of Evolution and Ecology, University of California, One Shields Avenue, Davis, CA 95616, USA



Species of fish exhibit variations in suction feeding behavior, including some with relatively large mouths that swim rapidly towards the prey when feeding and some that have small mouths and hold their body still while drawing the prey into the mouth, as have been investigated by Norton and Brainerd (1993), Norton (1991), and Higham et al. (2007). Individuals within a given species can modulate aspects of this feeding behavior with musculoskeletal control, including the speed with which they open the mouth and the maximum size of the mouth opening. For example, Nemeth (1997) demonstrated that more aggressive feedings (i.e. faster mouth opening) are employed when feeding on more evasive prey. Extensive research by Aerts et al. (1987), Aerts (1990), Lauder (1980), Liem (1973), Carroll et al. (2004), and Svanback et al. (2002) has investigated the musculoskeletal basis of suction feeding and Muller et al. (1982), Ferry-Graham et al. (2003), and van Wassenbergh et al. (2006) have devoted some attention to the study of the generated fluid flow.

Initial investigations into the fluid mechanics of suction feeding based on mathematical models using potential flow theory include those by Drost et al. (1988), Muller et al. (1982), Weihs (1980) and van Leeuwen and Muller (1984b). Earlier empirical studies included qualitative flow visualization by Muller and Osse (1984), van Leeuwen and Muller (1984a) and quantitative measurements of speed by particle streaking at a few locations in front of the fish Lauder and Clark (1984). Recent studies by our group, including Day et al. (2005) and Higham et al. (2005, 2006a), used particle image velocimetry (PIV) to resolve the flow field in front of two species of fish with sufficient spatial and temporal resolution to quantify the distribution of fluid speed as a function of distance in front of the fish and investigated the effect of swimming speed (or "ram speed"), mouth size, and time of mouth expansion on this flow. Nauwelaerts et al. (2007) used PIV to investigate the effect of substrate on water flow patterns generated by sharks.

This fluid flow field as described by distributions of velocity and pressure is central to the suction feeding event because it imparts all of the forces that act to draw the prey into the predator's mouth. The traditional view is that pressure within the mouth drives the velocity field and the forces imparted onto the prey are caused by drag and the acceleration reaction force, both of which are a consequence of the generated velocity field. Recently, Wainwright and Day (2007) proposed that there are three forces that the flow can exert onto a prey item: drag, acceleration reaction, and the pressure gradient force. Drag and the acceleration reaction are caused by the relative motion of the fluid and prey, both of which are a result of the velocity field. The pressure gradient force (Batchelor (1967) is, however, a direct result of gradients of pressure

within the flow field and would be present even in the absence of the prey. While the characteristics of the velocity field in front of a fish's mouth have been investigated previously, and it is assumed that there is a pressure gradient within this flow field, no study has empirically determined the difference in pressure between the prey and predator's mouth. It is supposed that a strong gradient of pressure exists as the result of fluid velocity varying in both space, being higher at the mouth aperture and decreasing with distance from the mouth, and in time.

The generated flow is very local to the fish's mouth and ephemeral, typically lasting 10–50 ms. The predator must manipulate the timing and location of the feeding to locate a small region of influence near the prey, some of which are very evasive. Suction feeding success depends heavily on the details of these water flow patterns and how the fish uses them to capture prey as has been described in Nyberg (1971), van Leeuwen and Muller (1984b) and Weihs (1980). Thus, a clear understanding of the time-course of the suction flow and the spatial region over which it operates is required before it will be possible to fully interpret the extensive morphological and behavioral diversity that exists among suction feeding species.

Manometry has frequently been used to measure the magnitude of negative pressure generated within the mouth cavity of feeding fish, for examples see Lauder (1980), van Leeuwen and Muller (1983), Sanford and Wainwright (2002) and Higham et al. (2006b). Peak sub-ambient pressure within the mouth is still the most common metric of suction feeding performance, but is only related to the prey through the flow field within, see van Wassenbergh et al. (2006), and outside the head. One way of conceptualizing this flow field is as follows: the feeding fish uses its musculoskeletal apparatus to pull against the volume of water within its mouth, thereby generating a sub-ambient "suction" pressure. The flow both within and outside the mouth is then driven by the gradient of pressure between the ambient surroundings and the sub-ambient pressure within the mouth. Because the pressure changes suddenly and resulting fluid accelerations are high, the fluid momentum is substantial and velocity as a function of time is temporally delayed as compared to the pressure. Experimentally, the measurement of pressure involves considerably less apparatus than measurements of velocity, so that many authors, such as Wainwright et al. (2006), Norton and Brainerd (1993), van Leeuwen and Muller (1983) and Carroll et al. (2004), have presented measurements of the timing and magnitude of maximum subambient intra-oral pressure (aka suction pressure) within a variety of species of suction feeding fish.

In Day et al. (2005) and Higham et al. (2005), we used particle image velocimetry to investigate the flow field generated by bluegill and largemouth bass. In the current



study we couple manometry with the PIV method used in prior studies in order to empirically determine the temporal relationship between the velocity field in front of suction feeding largemouth bass and pressure at two critical locations, within the mouth and at the prey. We demonstrate that, while substantial fluid velocities are confined to a region near the mouth of the fish, the sub-ambient pressures extend an even smaller distance into the flow. We describe the temporal patterns of velocity and pressure and demonstrate directly that maximum sub-ambient pressures occur before peak fluid speeds. We show that, although the velocity field varies with time, the distribution of fluid velocity in front of the fish is still a function of only the instantaneous gape and fluid speed at the mouth. We are able to demonstrate the effectiveness of applying the momentum equation to predict the relationship between the generated velocity field outside the mouth and pressure within the mouth and to demonstrate the contribution of fluid momentum to pressure within the mouth.

2 Materials and methods

2.1 Experimental animals

One largemouth bass, Micropterus salmoides Rafinesque, with a standard length of 18.5 cm was used in this study. The fish was caught in Yolo County, near Davis, CA and housed in a 100 l aquarium. The fish was fed a variety of squid (Loligo) and live ghost shrimp (Palaemonetes) daily. All fish maintenance and experimental procedures used in this research followed a protocol approved by the University of California, Davis Animal Care and Use Committee. Six feeding sequences were analyzed for this individual. The low inter-individual variance with respect to fluid speed-pressure relationships found in a previous study using this species (Higham et al. 2006b) suggests that using a single individual is a valid approach for measuring the relationship between the fluid velocity field in front of the feeding fish and pressures within the mouth. All values are reported as the mean \pm standard deviation for these six feedings.

2.2 Experimental setup

Experiments were conducted in a 2001 experimental aquarium that was integrated with a PIV system. Digital particle image velocimetry (DPIV) is a well-established technique (Adrian 1991) that measures the nearly instantaneous velocity field within an illuminated plane of the fluid field using light scattered from particles seeded into the fluid. This method is very versatile and ideally suited

for non-uniform transient flows. Similarly, miniature pressure transducers developed for blood vessel catheterization are commercially available. Details of both the PIV and the manometry system used in these experiments follow.

2.2.1 Particle image velocimetry

The illumination used in this experiment consists of an Innova I-90 continuous argon-ion laser with an output power of ~3 W (Coherent, Inc., Santa Clara, CA, USA). A matched set of a single cylindrical and spherical lens created a collimated light sheet about 10 cm wide and was focused to a waist thickness of approximately 1 mm within the measurement region. The laser sheet was directed upwards by a mirror located below the tank and passed into the aquarium through the tank bottom (Fig. 1). The resulting sheet was parallel to the sagittal plane (plane of symmetry) of an approaching fish. After traversing the depth of the tank, a mirror near the surface of the tank reflected the laser sheet back down within the same plane as the upward directed sheet, but angled towards the posterior of the fish. In this experiment, the downward directed sheet illuminated the fluid field above the upper lip of the fish that was in the shadow of the upward directed beam, but the methods may be useful for illuminating the flow in the shadow of any opaque object. Additionally, most of the flow field is illuminated by both the "upward" and the "reflected" sheets, thereby nearly doubling the amount of scattered light.

The seed particles used for light scattering were nominally $14 \mu m$ silver-coated glass beads manufactured for light scattering in reflective paints (Potter Industries, Inc., Carlstadt, NJ). The particles are hollow and have an

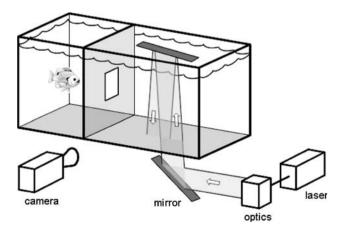


Fig. 1 Schematic of the experimental setup, showing experimental tank, position of laser sheet, optics, mirrors, camera, fish prior to a feeding, tank divider and door



average specific gravity of approximately 1.05 which, when used in fresh water, results in a calculated settling velocity of 5e-4 mm/s (Durrani and Greated 1977). In practice, it was apparent that a fraction of the particles settled out of the flow within 10-15 min, so we suspect that the density of the particles is variable.

The scattered light was imaged and collected by a NAC Memrecam ci digital high-speed video camera (Tokyo, Japan) through an f/1.2 c-mount video lens. The Memrecam is a high speed CMOS based camera with full resolution (512×462 pixels) at 500 frames per second. As the laser was not shuttered or pulsed, a mechanical shutter that is built into the camera was used to give an exposure time of 1/3,000 s. This was sufficiently short to prevent blurring and long enough to provide adequate signal (~ 1 mJ/image).

2.2.2 Manometry

Two pressure transducers were used in these experiments. The first, hereafter referred to a $p_{\rm mouth}$, was surgically implanted into the fish skull. The second, referred to as $p_{\rm prey}$, was attached to the prey mount and so moved with the prey up until the point of ingestion. Typically, as the prey moved into the mouth aperture, the prey mount wire contacted the upper lip of the fish so that the prey came off the mount and was carried into the mouth as the mount and transducer remained in the mouth aperture (see Fig. 2c, d).

In order to implant the mouth transducer, the fish was anesthetized by submerging it in a 0.3 g l⁻¹ solution of buffered MS-222 (Carroll et al. 2004). Once anesthetized, as determined by a cessation of gill ventilation and lack of a response to tactile stimulation, the fish was positioned in

clean water in a dissection tray and a 15 gage (1.8 mm) biopsy needle was forced through the neurocranium of the fish caudal to the ascending process of the pre-maxilla but rostral to the braincase. The needle emerged within the buccal cavity just lateral to the midline. A plastic cannula was constructed from PE-90 tubing and threaded into the needle (0.034" ID, 0.050" OD). The end of the cannula that was inside the buccal cavity had been flared prior to the procedure, enabling the cannula to be pulled up against the dorsal surface of the cavity with its opening positioned about 1-2 mm away from the buccal wall. A small sleeve of Tygon tubing (Cole-Parmer, Vernon Hills, IL, USA) was friction fit to the cannula where it protruded from the head of the fish to prevent it from sliding into to the skull. A Millar SPR-407 micro catheter-tipped pressure transducer (Millar Instruments, Inc., Houston, TX, USA) was threaded into the cannula and held in place by inserting the tip of the cannula into a piece of silicon sealant that had previously been allowed to set around the pressure transducer cable. The tip of the pressure transducer was positioned such that it was flush with the buccal cavity or slightly dorsal to the opening of the cannula. Surgery took no more than 15 min. The experiment began within 2-4 h after surgery.

The Millar SPR-407 is a solid state strain gauge type pressure transducer designed for intravenous measurements in small laboratory animals. The tip of the probe is cylindrical with a diameter of 0.67 mm and length ~ 5 mm. The sensor is located on the side of this cylinder and measures about 0.5×2 mm. The flange of the cannula that housed the transducer was designed to lay flush with the interior of the mouth. The transducer was recessed slightly in the cannula. The sensing element was exposed by a short fluid path to pressure in the buccal cavity so that it measures the static pressure outside the cannula and has a frequency

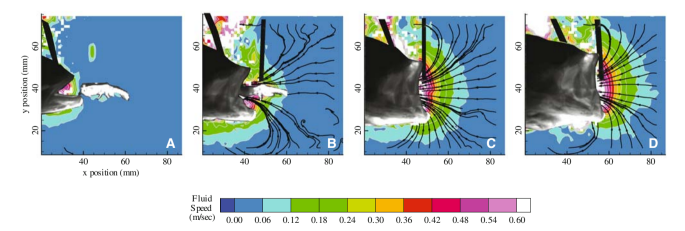


Fig. 2 PIV measurements in front of a bass feeding on a live, but mounted ghost shrimp at four instants during the feeding. Times are measured from the time that mouth opening began, the time of each is **a** 6 ms, **b** 18 ms, **c** 28 ms, **d** 50 ms. The position of the predator and prey are both shown as overlays onto the calculated velocity field.

The contour levels represent the magnitude of fluid speed and direction is parallel to the streamlines. The cannula that holds the mouth pressure transducer is clear as a black object protruding from the fish head. A second cannula that houses the pressure transducer located at the prey is also seen in the image



response of 10 kHz. Pressure transducers were calibrated prior to the surgery by placing them into a sealed flask. The pressure within the flask was varied over a range of -60-0 kPa using a vacuum pump and was measured with a commercial (World Precision Instruments, Sarasota, FL, USA) pressure transducer that came with an NIST (National Institute of Standards and Testing) certificate of calibration. The voltage output of the transducer is a linear function of pressure ($r^2 = 0.99$).

The transducer at the prey was housed in and protected by a short length of stainless steel tubing in order to both maintain the position of the transducer and to protect it when contacted by the feeding fish. The tubing was attached with glue and thread to the prey mount, so that the open and exposed end of the housing was just above the prey. Thin plastic tubing (PE-90) connected the top of the stainless tubing to a syringe located outside the tank. This was used to purge the cannula before each feeding.

The output of the mouth and prey pressure transducers were sent to linear op-amp based amplifiers with gains of 10 and 25 respectively. The amplified voltages were on the order of 1 V and were recorded at a sampling rate of 5 kHz using a National Instruments analog to digital converter (ADC) and LabView software (DAQPad-6070E National Instruments, Austin, Texas). Since the baseline pressure varied depending on the depth of the fish, we refer to pressure in this paper as the difference between the instantaneous pressure and the baseline value prior to the feeding event. Both the camera and ADC have a circular buffer, so that when armed they are continually acquiring data until triggered. In these experiments the pressure and video recordings were synchronized by using an external manual button to trigger both devices.

2.3 Experimental protocol

The fish was easily trained over the course of several days to reliably feed within the laser sheet. Prior to beginning the experiment, the fish was confined to one end of the tank by a physical barrier. The prey was suspended and positioned in the laser sheet and the camera field of view by a flexible (0.15 in. diameter) wire inserted under the exoskeleton, so that the shrimp generally remained alive and initiated an escape in response to the approaching predator. To keep the predator motivated, unrestrained prey were occasionally introduced during feeding sessions. The fish was confined to one end of the tank until a trap door was opened at the beginning of each experiment (Fig. 1). The fish swam through the door and towards the prey, suspended in the measurement region. The fish's movement is not confined in any direction, but due to the relative location of the trap door and prey, generally swam down the midline of the tank and located the center of its mouth at the laser sheet. A standard 30 Hz camcorder (Sony, Inc., Tokyo, Japan) recorded an anterior view of the fish in order to determine the relative position of the laser sheet and the mouth of the feeding fish. Only those trials for which the laser sheet was located less than 10% of the mouth diameter from the center line of the fish were analyzed.

2.3.1 PIV processing

Images were transferred from NAC proprietary format into a series of TIFF images. The entire sequence from the beginning of mouth opening until mouth closure lasted from 20 to 150 images (40-300 ms) depending on the speed of the event. An adaptive mesh cross-correlation algorithm created by Scarano and Riethmuller (1999) was used to calculate fluid velocities from image pairs, each pair consisting of two successive images from the highspeed video sequence. This is an iterative algorithm that includes resolution refinement, discreet window offsets, and deformation. All of the measurements for this work used an initial interrogation window size of 64×64 pixels and two refinement steps for a final resolution of 16×16 pixels. Two additional refinement steps are applied with no further refinement, but iterative offset and final image deformation. The interrogation used 16×16 pixel interrogation regions with 50% overlap, so that the measurement grid spacing was eight pixels, corresponding to approximately 1.3 mm for the camera field of view $(7 \times 5 \text{ cm})$. Each image pair lead to a simultaneous measurement of two components (u and v) of velocity at every location on a regularly spaced i, j ordered measurement grid with overall dimensions of i = 53 and j = 70. In addition to two components of velocity, the algorithm returned the signal-to-noise ratio (SNR) for each correlation. During all refinement steps, vectors falling below a fixed SNR or neighboring mean criterion are removed and replaced by interpolation. In the final interrogation step, removed vectors are not replaced. The majority of vectors lying within interrogation regions located on the fishes body are invalidated, but >95% of vectors lying in front of the fish pass the validation criteria.

Determining the uncertainty of correlation-based PIV methods complicated, but well studied. Generally, uncertainty increases as a result of poor seeding in that region of the image, high velocity gradients, solid boundaries that scatter light, and particle displacements that are large relative to the size of the interrogation region, as is described in Adrian (1997). The following two-step validation scheme was implemented.

First, vectors with a signal-to-noise ratio (SNR) of less than 2.0 were removed, without replacement, and no



smoothing was applied to the final velocity field. Some spurious measurements that are not representative of fluid particle displacement still passed the SNR validation criterion. The second step of the validation scheme deals with these spurious vectors. For velocities extracted from a transect extending away from the mouth along the centerline of the fish, measurements both directly on the transect (i, j) and at 2 grid points above (i, j + 2) and 2 grid points below (i, j-2) were considered at each horizontal position. Measurements located 2 grid points away from the primary measurement location are used because these do not overlap the primary measurement region. The neighboring points in the i direction were not used for validation because of the very steep gradients of fluid velocity in the horizontal direction. If at least two of the three measurements considered had not been removed based on the SNR criterion (step one of the validation scheme), then the mean of the remaining measurements was used as the value of speed for that given position along the transect.

This validation scheme resulted in the removal of some measurements near the mouth for all sequences. Measurements near the mouth were the most likely to fail the validation scheme because particle displacements (high fluid speed), velocity gradients, and occasional glare from the predator were all highest in this region. The camera frame rate and spatial resolution used for the measurements placed an effective upper bound of measured fluid speed at approximately 1.5 m/s, corresponding to 16 pixel displacement between images. For the majority of feedings, all measurements greater than 2–3 mm from the mouth passed the validation procedure and measurements further than 5 mm from the mouth were validated for all feedings.

2.3.2 Data analysis

The position of the eye and upper and lower jaws were manually identified and measured for each frame of the acquired video sequence using Image J (NIH, Washington, DC, USA). The x, y position of each was used to calculate several kinematic variables using and Excel spreadsheet. The position of the center of the mouth was defined as the midpoint between the upper and lower jaw tips. Gape was the distance from tip of the upper jaw to tip of the lower jaw and Peak Gape (PG) was the maximum value of gape during the feeding. Ram speed was calculated as the horizontal component of the temporal derivative of eye position. The measurements of horizontal position were smoothed with a three-point moving average before calculating derivative quantities. The onset of mouth opening and the onset of peak gape are defined as the time at which the mouth has opened to 20 and 95% of its maximum value, respectively. The time between these is referred to as time to peak gape, TTPG.

This extraction of speed along the centerline of the fish from the vector field output by the PIV algorithm was automated with a custom program written in Visual Basic because the absolute position of the fish, and therefore positions of the transects varied during the feeding. The program probed the PIV velocity data, accounting for both the angle of the fish and the position of mouth as determined from manual digitization.

In Day et al. (2005), we showed that the profiles of speed in front of the fish scale with the magnitude of fluid speed at the mouth and the size of the mouth at the time of peak fluid speed. In this study, the profile of speed along the centerline throughout the feeding event was scaled by dividing spatial dimensions (x) by the magnitude of instantaneous gape (G) and speeds by the magnitude of fluid speed at a distance of 1/2 gape in front of the fish. The speed at this relative position is used throughout as a reference because at this location fluid speed is substantial and the PIV measurements meet the validation criteria described above, and is referred to as $FS_{1/2}$ gape from here on. Profiles of scaled speed were compared to one another to see if the function of fluid speed holds true throughout the duration of the feeding.

The pressure transducer located at the prey mount moves towards the mouth during the feeding, resulting in pressure measurements as a function of time and position in front of the fish. For each acquired data point, the magnitude and distance between the mouth aperture and transducer were recorded. In a manner similar to our scaling of the velocity field, the acquired pressures at the prey are scaled. The magnitude of each measurement of pressure at the prey, p_{prey} , was divided by the instantaneous pressure in the fish mouth, p_{mouth} , and the spatial position of each was divided by the instantaneous gape. This results in a distribution of scaled pressure as a function of nondimensional position $(x^* = x/G)$ in front of the mouth. Is should be remembered that the profile of pressure versus distance was not acquired at a single instantaneous time for each feeding, but is the compilation of single point measurements taken over the period of the feeding. Nonetheless, each is scaled by the instantaneous pressure in the fish mouth and instantaneous gape.

Finally, the measured distribution of the velocity in front of the fish was used to calculate a predicted pressure within the buccal cavity according to the momentum equation along the centerline of the fish. We integrated the differential form of one-component of the momentum equation (Eq. 1) along a path extending from a point far away from the fish and ending at the mouth aperture in order to find the relationship between p_{mouth} and p_{ambient} (Eq. 2).



$$\frac{dp}{dx} = -\rho \left(\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} \right) \tag{1}$$

$$p_{\text{mouth}} - p_{\text{ambient}} = \int_{x_{\text{ambient}}}^{x_{\text{mouth}}} \frac{dp}{dx} dx = \int_{x_{\text{ambient}}}^{x_{\text{mouth}}} \rho \left(\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} \right) dx$$
(2)

Implementation of the momentum equation directly on the raw PIV data leads to high uncertainty because of the derivative terms, so we instead first fit a function to the empirical data and then applied the equation to this function. The spatial distribution of fluid speed is specified by a polynomial fit to the empirical data. The curve-fit for the time course of fluid speed is based on the form of equation 11 from Muller et al. (1982) with parameters manually adjusted to agree with the empirical data for that particular feeding. This implementation resulted in a function of predicted pressure at the mouth aperture that was arrived at independently of the measured $p_{\rm mouth}$. This predicted pressure was compared to the measured $p_{\rm mouth}$ in order to validate the utility of this relationship.

3 Results

The suction feeder creates a velocity and pressure field that both have high spatial and temporal gradients. Speeds are highest near the mouth $(1.5 \pm 0.36 \text{ m/s})$ and decrease as a function of distance away from the mouth. There is a general shape of contour lines of constant velocity throughout the feeding that we describe as the top of a mushroom (Fig. 2). This pattern is symmetric about the long-axis of the fish. The size of the pattern varies in proportion to the diameter of the mouth and the magnitude of velocity at the mouth varies throughout the feeding. Both the size of the affected region and magnitude of fluid speed are highest at a time shortly after reaching peak gape. The fish continues to ingest water throughout the duration of the feeding, even as the mouth is closing, as shown in Figs. 2 and 3.

3.1 Temporal pattern

There is a general temporal pattern of generated fluid speed, gape, and the recorded pressures (see Fig. 3 for a representative sequence). All times are given as both the absolute time (relative to the beginning of mouth opening) and as a fraction of TTPG, the time between beginning of mouth opening and peak gape, as the relative timing of events scales in proportion of the total duration of the feeding. Maximum sub-ambient pressure in the mouth $(-5.7 \pm 1.9 \text{ kPa})$ occurs

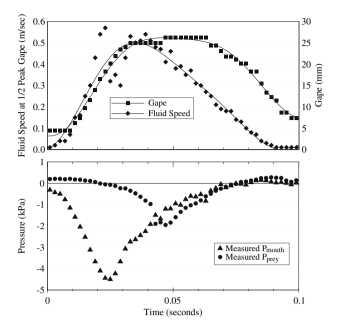


Fig. 3 Relative time course of fluid speed compared to gape distance and pressures within and outside the mouth for the representative sequences. Fluid speed is measured at a constant distance in front of the fish equal to 1/2PG. This is a moderate speed feeding (time to open mouth ~ 25 ms). The relative timing of events is similar for all recorded feedings. Peak fluid speed occurs slightly before the onset of peak gape. The pressure within the mouth, p_{mouth} reaches is maximum sub-ambient pressure during mouth opening and has already decayed significantly by the time of mouth opening. The pressure at the prey, p_{mouth} , is near ambient, but gradually decreases as both fluid speed increase and the prev moves closer to the predator's mouth. Near 48 ms, the prey enters the mouth. The transducer and mouth are in contact with the upper lip of the fish, so the transducer remains in the aperture until the end of the feeding. During this time, the pressure measured near the mouth aperture and within the buccal cavity are nearly equal

at a time $(24 \pm 18 \text{ ms}, 58 \pm 18\% TTPG)$ when the mouth is still opening and has typically decayed to approximately 1/2 the peak amplitude at the onset of peak gape (38 \pm 21 ms, $100 \pm 0\% TTPG$), consistent with Sanford and Wainwright (2002). Peak fluid speed (1.50 \pm 0.36 m/s) reached a maximum just after (38.3 \pm 20.3 ms, 113 \pm 41% TTPG) the onset of peak gape, consistent with Day et al. (2005) and Higham et al. (2006a). Peak gape is maintained for some time $(25 \pm 9.7 \text{ ms})$ before the mouth begins to close. At the end of peak gape $(0.63 \pm 0.20 \text{ m/s})$, as indicated by the mouth diameter having reached it's maximum and then decreased to 95% of its maximum size, fluid speed at the mouth has decreased to a fraction (32 \pm 28%) of the maximum speed for that feeding. The pressure at the prey is initially very small and decreases exponentially as a result of both the flow speed increasing and the prey moving towards the mouth. At the time when the prey is at the mouth opening (48 ms in this feeding), the pressure transducer is located in the mouth aperture and remains there for the duration of the feeding



after the prey is drawn off the wire and ingested. The transducer located at the prey, $p_{\rm prey}$, records a nearly identical pressure as $p_{\rm mouth}$. It should be emphasized that for all feedings that we recorded, the second transducer, $p_{\rm prey}$, is located at the aperture only after peak gape and not during mouth expansion.

3.2 Spatial pattern

After creating a non-dimensional distance (x^*) and speed (SS, scaled speed) by dividing all measured distances (x)by the instantaneous mouth diameter and dividing all fluid speeds (FS) by the fluid speed at a reference location located at a distance 1/2 of gape ($x^* = 1/2$) in front of the fish ($FS_{1/2 \text{ gape}}$), the velocity profile along the centerline for all feedings collapse to one empirical relationship (Fig. 4; Eq. 3). At all times during the feeding, fluid speed decays with distance in front of the feeding fish, being equal to approximately 25% the speed at the mouth at a distance of 1/2 the mouth diameter and 5% at one mouth diameter. Because the spatial distribution of velocity scales in proportion to the mouth diameter, the spatial distribution of absolute velocity is dependent on gape diameter and the magnitude of fluid speed generated at the mouth (Day et al. 2005). The feeding *Micropterus* can modulate both of these, so that there is significant feeding-to-feeding variation. A fourth order polynomial fit to the data assembled for all feedings is given as

$$FS(x^*) = FS_{\text{mouth}} (0.215x^{*4} - 1.26x^{*3} + 2.73x^{*2} - 2.63x^* + 1)$$
(3)

The magnitude of fluid speed and gape vary throughout the feeding. After normalization based on FS at the mouth and the mouth diameter, both measured as a function of time, transects of scaled speed in front of the fish have a very similar shape at all times during the feeding (Fig. 5), suggesting that Eq. 3 is an appropriate description of the spatial pattern of the velocity throughout the feeding.

3.3 Pressure outside the mouth

For most feedings, the onset of mouth opening occurs when the prey is approximately one mouth diameter in front of the fish and the prey enters the mouth of the fish slightly before the onset of peak gape (40 ± 16 ms, $58 \pm 18\%$ TTPG). This allows for measurements of the pressure in front of the fish during this period as the transducer attached to the prey moved towards the mouth. The distribution of pressure decays even more dramatically than

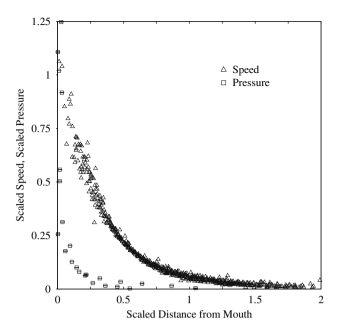


Fig. 4 Profiles of scaled speed and pressure along the centerline in front of the fish. Additionally, measurements from the pressure transducer located at the prey are presented. As the prey moved towards the predator during each feeding, the distance from mouth to prey was calculated and scaled by the instantaneous gape. The magnitude of pressure was scaled by the magnitude of measured pressure within the mouth cavity at that instant in time. Note that the decay of pressure in front of the fish is even more dramatic than that of fluid velocity. There is a very steep pressure gradient very near the mouth and substantially less away from the mouth

velocity in front of the fish (Fig. 4). The pressure is approximately 10% the pressure within the mouth at a distance of 1/4 gape, and 5% at 1/2 gape, thereby only influencing a region that is extremely local to the fish mouth. The spatial gradient of pressure is steepest at the mouth aperture and very small at a distance more than 1/4 gape from the fish.

3.4 Model results

The existence of a general form of the spatial pattern of fluid speed (Eq. 3) based on fluid speed at the mouth (FS_{mouth}) and the mouth aperture (G), allows a mathematical function for the spatio-temporal pattern of fluid speed in front of the fish so long as FS_{mouth} and G are specified as a function of time. The resulting spatio-temporal pattern of fluid speed along a line extending away from the fish's mouth is shown in Fig. 6 for a representative feeding.

Predictions of the pressure within the mouth based on the solution of Eq. 2 both neglecting (steady form) and including (unsteady form) the temporal derivative of velocity are compared in Fig. 7 for this same representative



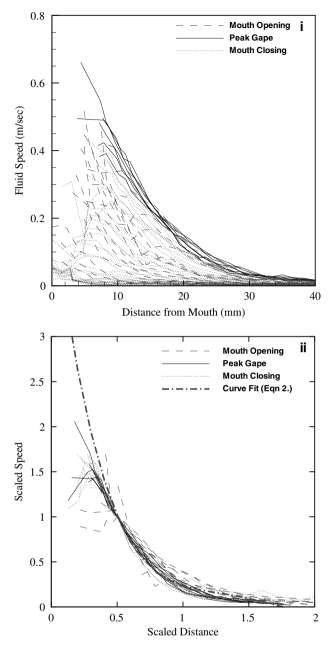


Fig. 5 Profiles of speed (a) and scaled speed (b) measured along the centerline at all times during the feeding, including mouth opening, the duration of peak gape and mouth closure. a Both the magnitude and size of the velocity profile vary during the feeding. b The same data as in a, but after scaling. The distance in front of the fish was scaled by the mouth diameter at the time of velocity measurement. Magnitudes of velocities were scaled by the FS measured at a distance of 1/2 gape at that time (FS1/2 gape). Variations in mouth size and the speed of flow at the mouth have a very significant effect on the absolute speeds in front of the fish, but the scaled profiles are similar at all times during the feeding

feeding. For all feedings, the time of peak sub-ambient pressure predicted based on the unsteady form of the model $(25 \pm 18 \text{ ms}, 61 \pm 19\%TTPG)$ agreed well with the empirical data $(24 \pm 18 \text{ ms}, 58 \pm 18\%TTPG)$, as shown in

Fig. 8. The steady form $(30 \pm 19 \text{ ms}, 77 \pm 21\%TTPG)$ predicted peak pressure to occur later than the unsteady form, and always after it was measured (Figs. 7, 8). The unsteady form predicts greater magnitude of sub-ambient pressures than the steady form $(-3.9 \pm 1.1 \text{ kPa} \text{ vs}. -3.2 \pm 1.0 \text{ kPa})$. For some feedings there is very good agreement between the magnitude of predicted and measured $(-5.7 \pm 1.9 \text{ kPa})$ mouth pressures but the predicted pressure is smaller magnitude than the observed for most feedings (as is the case for the feeding shown in Fig. 7).

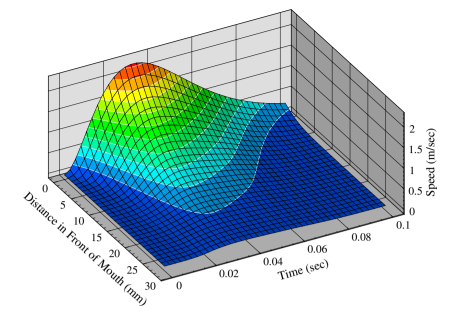
4 Discussion

The simultaneous measurement of fluid speed and pressure presented here is one of the first empirical attempts to bridge the relationship between pressure and fluid speed in the complex, unsteady flow field that characterizes suction feeding. During suction feeding, the affected flow is confined to a region close to the mouth of the fish and the size of this region varies in direct proportion to mouth diameter throughout the feeding. The area of substantial fluid speed forms a three-dimensional shaped region similar to the top of a mushroom. This result is consistent with models by Muller and Osse (1984) and the empirical findings of Ferry-Graham et al. (2003) and Day et al. (2005) for other species. This study also shows that the area of affected pressure is confined to an even smaller region than the area affected by the velocity field. The pressure gradient force, as opposed to velocity-based forces such as drag, has recently been proposed to be the dominant forces in many suction feeding scenarios by Wainwright and Day (2007). This study underscores the conclusion of previous studies that the predator must locate the prey in very close proximity to its mouth in order to have any effect on the prey.

Day et al. (2005) showed that a single polynomial fit to empirical data describes the distribution of fluid speed in front of the fish and showed that this relationship holds true at the time of peak fluid speed across a wide range of mouth size and fluid speed in a bluegill. Here we show that a similar polynomial function (Eq. 3) can be used to describe the spatial distribution of fluid speed at all times during the feeding for the largemouth bass. Velocity profiles scaled by the instantaneous gape and instantaneous fluid speed all fit one generalized function well. This result is consistent with previous findings that peak fluid speeds measured at three locations along the centerline transect all occurred nearly simultaneously with one another and with the onset of peak gape (95% opening) Higham et al. (2006b). Further, this allows a description of the velocity field using only two parameters, the mouth diameter and fluid speed, which are specified as functions of time.



Fig. 6 Curve fit to the spatiotemporal pattern of fluid speed along the centerline transect in front of the feeding fish. Fluid speed in front of the fish is always given (Eq. 3) as $FS(x^*) = FS_{mouth}$ (0.215 x^{*4} $-1.26x^{*3} + 2.73x^{*2}$ $-2.63x^* + 1$). FS_{mouth} is the fluid speed at the mouth (distance equals 0) and is specified as a function of time and is based on the form of equation 11 from Muller et al. (1982)



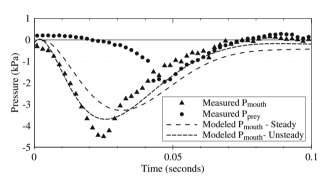


Fig. 7 The *symbols* show measured pressures within the mouth and in front of the fish. The *dashed lines* show predictions of pressure at the mouth aperture calculated by applying Eq. 3 to the time sequence of measured speeds along the centerline

Fluid speed was not constant during the course of the feeding event, increasing from zero to peak fluid speed in 40 ± 18 ms. The time of peak fluid speed occurred near the onset of peak gape (38 \pm 21 ms). The synchronization of fluid speed and gape is potentially a very effective feeding strategy because the fish simultaneously maximizes flowinduced forces acting on the prey and the space over which the flow occurs. One mechanism that can facilitate this that continued expansion of the posterior portion of the buccal and opercular cavities after peak expansion of the anterior portion prolongs the period of volumetric expansion of the total mouth past peak gape. A distinct delay in the posterior expansion (as measured by suspensory and opercular abduction) relative to gape was shown consistently throughout feedings of three species of centrarchid fishes by Lauder (1980). The anterior-to-posterior delay of expansion of major functional components of the head has been shown to hold true across a wide range of taxa by

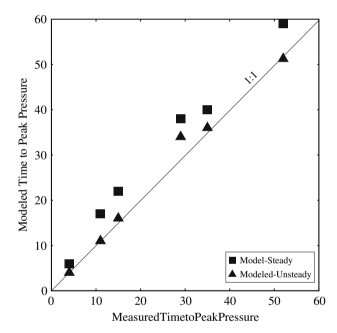


Fig. 8 Comparison of the time to peak pressure, as predicted by both the steady and unsteady models, plotted versus the measured peak pressure

Lauder (1982). The second is that the opening of the opercular slits allows fluid to continue to flow into and through the mouth, driven only by fluid momentum, after volumetric expansion of the buccal cavity has ceased. At the beginning of the feeding, when the opercular slits are closed, the volumetric flow rate into the mouth aperture is exactly equal to the instantaneous volumetric expansion of the combined buccal and opercular cavities. After the opercular slits are open, the flow into the mouth is equal to any remaining expansion of the mouth cavity in addition to



the volumetric flow rate out of the opercular slits. After the mouth has reached full expansion, the velocity of the fluid cannot continue to increase. However, the flow will continue on its own momentum, slowing due to losses, while the mouth is held open. It is likely that both the anterior-to-posterior expansion and timing of opercular opening contribute to the observed temporal relationship between fluid speed and gape.

Although a relationship between the velocity and pressure field through the equations of motion necessarily exists, the relationship in this particular flow is complicated. According to van Leeuwen and Muller (1983) "Flow patterns cannot be derived from the pressure regime as the process is essentially unsteady (Muller, 1982)." As a result of statements to this effect and support from numerical models that show that the magnitude of generated buccal pressure is very sensitive to the shape of the mouth cavity van Wassenbergh et al. (2006), measurements of pressure within the mouth are only related to the velocity field in front of the fish with great caution. This is, however, problematic when trying to study and evaluate feeding performance because the purpose of the predator is to exert forces on the prey and these forces are a result of the flow outside the mouth and the mechanism for doing this is to directly affect the fluid within the mouth. We have shown in Higham et al. (2006b) that the magnitude of peak fluid speed is correlated to the magnitude of peak suction pressure.

This study quantifies the degree to which the time course of buccal pressure is affected by the unsteady nature of the flow. When the unsteady terms are included in the relationship between the velocity field and pressure, the prediction of pressure at the mouth leads to excellent temporal agreement with the measured buccal pressure. Exclusion of these terms (a quasi-steady assumption) leads to slightly lower magnitude and temporal latency of the prediction.

In this study we have demonstrated that in at least one species of fish, there is a clear relationship between pressure within the mouth and fluid speed outside the head. We propose that because of properties and characterization of this flow field, measured pressures within the mouth could, in fact, be used to derive the flow outside the fish. This is primarily due to the fact that there is a general function for the velocity distribution in front of the fish that is valid for all times during the feeding and over a wide range of behavioral variations, as was shown in Day et al. (2005) and Higham et al. (2006a). By accounting for only two variables, i.e. mouth size and fluid speed at the mouth, we can characterize velocity in front of the fish. Assuming that mouth size is known as a function of time, the pressure at the mouth aperture can be used to predict the velocity at the mouth aperture or vice versa. In this study, we have demonstrated the latter. Our measurements demonstrate that the pressure at the mouth aperture is very nearly the same as at the posterior location of $p_{\rm mouth}$ for all times after the onset of peak gape, which is perhaps one reason why our prediction of the pressure at the mouth aperture agrees well with the measured pressure.

Although the predictions of the timing of peak pressure agree well with observations, the magnitude of estimated pressure was always lower than observed. This could be due to radial movement of the mouth walls resulting in streamline curvature, losses at the mouth opening, or viscous losses both within and outside the mouth, none of which are accounted for in the integration of one-dimensional form of the momentum equation given in Eq. 2.

The flow field is characterized by high spatial and temporal gradients of velocity. The local acceleration near the mouth aperture is on the order of 50 gs and even higher in some species, such as the bluegill, as was shown in Day et al. (2005). Both the spatial gradients of velocity and accelerations contribute to pressure gradient within the flow field, which Wainwright and Day (2007) have shown exerts substantial forces on the prey. This study represents a significant advance in understanding the relationship between pressure and velocity in this dynamic fluid flow because it measures both simultaneously and demonstrates good temporal agreement between the fluid velocity external to the fish and the pressure measured within the fish mouth. In addition to the intra-oral pressure, this inclusion of pressure experienced by the prey is a novel perspective on this flow. The study confirmed the localized and short lasting generated velocity field described previously and demonstrated that substantial gradients of pressure are only present in an even smaller region in front of the fish. Despite the spatial and transient nature of the flow, a fairly simple pattern of fluid speed exists throughout the feeding and the fluid velocity field outside the fish can be successfully related to pressure within the mouth.

In conclusion, we find that the equations of motion can, in fact, be used to relate pressure measured inside the mouth cavity, to the flow field outside the mouth, at least up until the time when the opercula are opened at the back of the head. This is the first such demonstration for suction feeding fishes and establishes an important physical linkage between the pressures that are generated during expansion of the head and the flow that results and is used to capture prey. Our understanding of the fluid mechanics of suction feeding has advanced considerably in recent years, but as yet there has not been a detailed treatment of the mechanics of internal expansion of the oral cavity and how this pattern relates to flow entering the mouth. These results are an important step in this direction that will lead to a greater understanding of how internal head movements can be regulated to control flow in front of the fish.



Acknowledgments This research was supported by NSF grants IOB-0444554 and 0610310.

References

- Adrian RJ (1991) Particle imaging techniques for fluid mechanics. Annu Rev Fluid Mech 23:261–304
- Adrian RJ (1997) Dynamic ranges and spatial resolution of particle image velocimetry. Meas Sci Technol 8:1393–1398
- Aerts P (1990) Variability of the fast suction feeding process in Astatotilapia elegans (Teleostei, Cichlidae)—a hypothesis of peripheral feedback control. J Zool 220:653–678
- Aerts P, Osse JWM, Verraes W (1987) Model of jaw depression during feeding in *Astatotilapia elegans* (Teleostei, Cichlidae)—mechanisms for energy-storage and triggering. J Morphol 194:85–109
- Batchelor GK (1967) An introduction to fluid mechanics. Cambridge Carroll AM, Wainwright PC, Huskey SH, Collar DC, Turnigan RG (2004) Morphology predicts suction feeding performance in centrachid fishes. J Exp Biol 207:3873–3881
- Day SW, Higham TE, Cheer AY, Wainwright PC (2005) Spatial and temporal patterns of water flow generated by suction-feeding bluegill sunfish *Lepomis macrochirus* resolved by particle image velocimetry. J Exp Biol 208:2661–2671
- Drost MR, Muller M, Osse JWM (1988) A quantitative hydrodynamical model of suction feeding in larval fishes: the role of frictional forces. Proc R Soc Lond Ser B 234:263–281
- Durrani TS, Greated CA (1977) Laser systems in flow measurements. New York
- Ferry-Graham LA, Wainwright PC, Lauder GV (2003) Quantification of flow during suction feeding in bluegill sunfish. Zoology 106:159–168
- Higham TE, Day SW, Wainwright PC (2005) Sucking while swimming: evaluating the effects of ram speed on suction generation in bluegill sunfish *Lepomis macrochirus* using digital particle image velocimetry. J Exp Biol 208:2653–2660
- Higham TE, Day SW, Wainwright PC (2006a) Multidimensional analysis of suction feeding performance in fishes: fluid speed, acceleration, feeding accuracy and the ingested volume of water. J Exp Biol 209:2713–2725
- Higham TE, Day SW, Wainwright PC (2006b) The pressures of suction feeding: the relation between buccal pressure and induced fluid speed in centrarchid fishes. J Exp Biol 209:3281–3287
- Higham TE, Hulsey CD, Rican O, Carroll AM (2007) Feeding with speed: prey capture evolution in cichlids. J Evol Biol 20:70–78
- Lauder GV (1980) The suction feeding mechanism in sunfishes (*Lepomis*): an experimental analysis. J Exp Biol 88:49–72
- Lauder GV (1982) Patterns of evolution in the feeding mechanism of actinopterygian fishes. Am Zool 22:275–285

- Lauder GV, Clark BD (1984) Water flow patterns during prey capture by teleost fishes. J Exp Biol 113:143–150
- Liem KF (1973) Modulatory multiplicity in the feeding mechanism in cichlid fishes, as exemplified by the invertebrate pickers of Lake Tanganyika. J Zool Soc Lond, pp 93–125
- Muller M, Osse JWM (1984) Hydrodynamics of suction feeding fish. Trans Zool Soc Lond 37:51–135
- Muller M, Osse JWM, Verhagen JHG (1982) A quantitative hydrodynamical model of suction feeding in fish. J Theor Biol 95:49–79
- Nauwelaerts S, Wilga C, Sanford C, Lauder G (2007) Hydrodynamics of prey capture in sharks: effects of substrate. J R Soc Interface 4(13):341–345
- Nemeth DH (1997) Modulation of buccal pressure during prey capture in *Hexagrammos decagrammus* (Teleostei: Hexagrammidae). J Exp Biol 200:2145–2154
- Norton SF (1991) Capture success and diet of cottid fishes—the role of predator morphology and attack kinematics. Ecology 72:1807–1819
- Norton SF, Brainerd EL (1993) Convergence in the feeding mechanics of ecomorphologically similar species in the Centrarchidae and Cichlidae. J Exp Biol 176:11–29
- Nyberg DW (1971) Prey capture in largemouth bass. Am Midl Nat 86:128–144
- Sanford CPJ, Wainwright PC (2002) Use of sonomicrometry demonstrates the link between prey capture kinematics and suction pressure in largemouth bass. J Exp Biol 205:3445–3457
- Scarano F, Reithmuller ML (1999) Iterative multigrid approach in PIV image processing with discrete window offset. Exp Fluids 26:513-523
- Svanback R, Wainwright PC, Ferry-Graham LA (2002) Linking cranial kinematics, buccal pressure, and suction feeding performance in largemouth bass. Physiol Biochem Zool 75:532–543
- van Leeuwen JL, Muller M (1983) The recording and interpretation of pressures in prey-sucking fish. Neth J Zool 33:425–475
- van Leeuwen JL, Muller M (1984a) A quantitative study of flow in prey capture by rainbow trout, *Salmo gairdneri*, with general consideration of the actinopterygian feeding mechanism. Trans Zool Soc Lond 37:171–227
- van Leeuwen JL, Muller M (1984b) Optimum sucking techniques for predatory fish. Trans Zool Soc Lond 37:137–169
- van Wassenbergh S, Aerts P, Herrel A (2006) Hydrodynamic modelling of aquatic suction performance and intra-oral pressures: limitations for comparative studies. J R Soc Interface 3:507–514
- Wainwright PC, Day SW (2007) The forces exerted by aquatic suction feeders on their prey. J R Soc Interface 4:553-560
- Wainwright PC, Huskey SH, Turingan RG, Carroll AM (2006) Ontogeny of suction feeding capacity in snook, *Centropomus undecimalis*. J Exp Zool A 305A:246–252
- Weihs D (1980) Hydrodynamics of suction feeding fish in motion. J Fish Biol 16:425-433

