

Active tail flexion in concert with passive hydrodynamic forces improves swimming speed and efficiency

Haotian Hang¹, Sina Heydari¹, John H. Costello^{2,3} and Eva Kanso^{1,†}

¹Department of Aerospace and Mechanical Engineering, University of Southern California, 854 Downey Way, Los Angeles, CA 90089, USA

²Biology, Providence College, Providence, RI 02918, USA

³Whitman Center, Marine Biological Laboratory, Woods Hole, MA 02543, USA

(Received 19 April 2021; revised 19 September 2021; accepted 20 October 2021)

Fish typically swim by periodic bending of their bodies. Bending seems to follow a universal rule; it occurs at about one-third from the posterior end of the fish body with a maximum bending angle of about 30°. However, the hydrodynamic mechanisms that shaped this convergent design and its potential benefit to fish in terms of swimming speed and efficiency are not well understood. It is also unclear to what extent this bending is active or follows passively from the interaction of a flexible posterior with the fluid environment. Here, we use a self-propelled two-link model, with fluid–structure interactions described in the context of the vortex sheet method, to analyse the effects of both active and passive body bending on the swimming performance. We find that passive bending is more efficient but could reduce swimming speed compared with rigid flapping, but the addition of active bending could enhance both speed and efficiency. Importantly, we find that the phase difference between the posterior and anterior sections of the body is an important kinematic factor that influences performance, and that active antiphase flexion, consistent with the passive flexion phase, can simultaneously enhance speed and efficiency in a region of the design space that overlaps with biological observations. Our results are consistent with the hypothesis that fish that actively bend their bodies in a fashion that exploits passive hydrodynamics can at once improve speed and efficiency.

Key words: flow–structure interactions, swimming/flying, wakes

1. Introduction

Millions of years of natural selection have endowed fish with remarkable abilities to swim efficiently compared with underwater man-made propulsors (Sfakiotakis, Lane &

† Email address for correspondence: kanso@usc.edu

© The Author(s), 2021. Published by Cambridge University Press. This is an Open Access article, distributed under the terms of the Creative Commons Attribution licence (<https://creativecommons.org/licenses/by/4.0/>), which permits unrestricted re-use, distribution, and reproduction in any medium, provided the original work is properly cited.

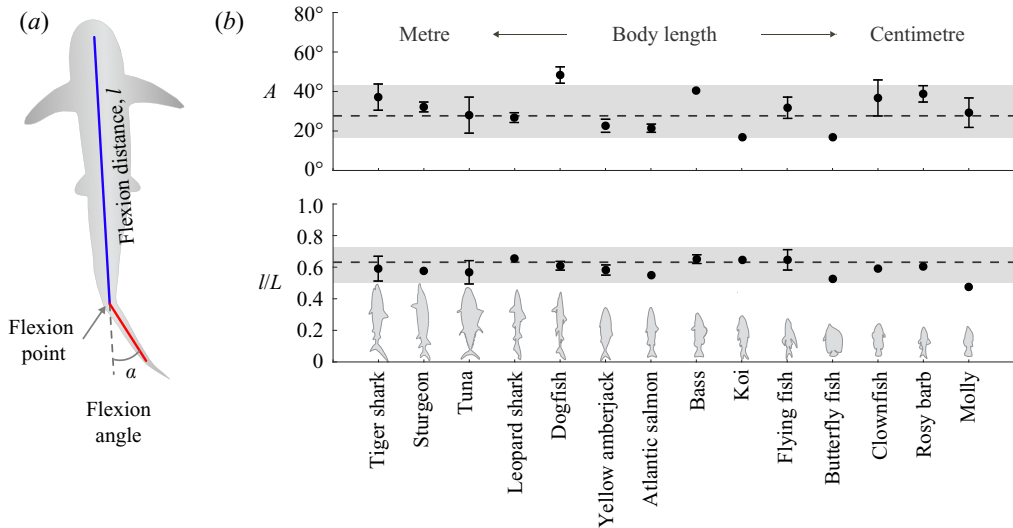


Figure 1. Bending rules of swimming fish. (a) Schematic of flexion parameters in fish with flexion distance l , ratio l/L and maximum angle defined as in Lucas *et al.* (2014). (b) Flexion ratio l/L and maximum flexion angle A observed in different fish species; data taken from Lucas *et al.* (2014). The observed flexion ratios and angles are fairly consistent among different fish species, despite large variations in length scales.

Davies 1999; Lauder 2015). Several fish use body and caudal fin (BCF) deformations for propulsion. Details of BCF deformations have been used to classify fish swimming modes (Sfakiotakis *et al.* 1999; Shadwick & Gemballa 2005; Low & Chong 2010; Smits 2019). Invariably, in most BCF swimming modes, anterior to posterior bending of the fish body seems to play an important role in swimming efficiency, and it is often linked to increased flexibility towards the fish tail or caudal peduncle (Combes & Daniel 2003; Tytell & Lauder 2004; Tytell *et al.* 2010; Gemmell *et al.* 2016; Tytell *et al.* 2016). However, the underlying bending mechanisms and optimal bending parameters remain opaque.

In an effort to document the bending rules of fluid-based propulsors, both aerial and aquatic, Lucas *et al.* (2014) collected morphometric data of the flexion parameters across length scales and animal taxa. They identified two parameters to characterize the bending behaviour: flexion ratio l/L between the flexion distance l and total length L of the propulsor and maximum flexion angle A . They found that flexion ratio and maximum flexion angle of all surveyed animals, including fish, clustered in a limited design space: bending occurs at about 70% of body length at maximum flexion angle of about 30° , as shown in figure 1 for swimming fish based on the data collected in Lucas *et al.* (2014). These fish vary in length, swimming speed and tailbeat frequency, and thus span a wide range of Reynolds numbers ($Re \sim 10^3\text{--}10^7$), but exhibit similar bending parameters. It is unclear the extent to which this anterior-to-posterior bending is active or whether it follows passively due to the interaction of a flexible posterior with the fluid motion. Either way, these findings raise the question of whether hydrodynamics could have provided a selective force for driving this convergent bending design.

To address these questions, we analyse the influence of bending on the swimming speed and efficiency of a simplified fish model that consists of anterior and posterior sections connected via a rotational joint at the flexion point (see figure 1a). The fish anterior undergoes periodic planar pitching while the posterior either (i) moves in synchrony with the anterior as if the two parts were a single rigid body, (ii) bends actively at distinct

amplitude and phase relative to the fish anterior or (iii) bends passively due to interactions with the flow generated by the fish anterior. We find that swimming with passive bending could be more efficient than rigid flapping but at the cost of diminished swimming speed. Active bending provides more possibilities to alter the swimming performance through not only the flexion ratio and maximum flexion angle reported in Lucas *et al.* (2014) but also the phase difference ϕ between the flapping motions of the anterior and posterior parts. Importantly, we find that antiphase anterior-to-posterior flexion can simultaneously enhance swimming speed and efficiency in a region of the design parameter space ($l/L, A$). We analyse in depth the hydrodynamic mechanisms underlying these improvements in swimming performance. Despite the major simplifications in our hydrodynamic model detailed later, we find that flexion ratios and angles that lead to significant improvements in speed and efficiency in our model overlap with the observations of real fish reported in Lucas *et al.* (2014).

Details of the flow field around swimming fish have received a great deal of attention. Several studies used particle image velocimetry to measure the flow field around live fish and analyse the interplay between body deformations and thrust production (see e.g. Müller *et al.* 1997; Müller, Stamuis & Videler 2002; Liao *et al.* 2003; Tytell & Lauder 2004; Gemmell *et al.* 2016). *In silico* models of various degrees of fidelity to fish morphology and kinematics have also been used to examine the offsets of body deformations on swimming speed and efficiency (Eldredge 2006; Kern & Koumoutsakos 2006; Tytell *et al.* 2010; Eloy 2013). Importantly, several experimental and numerical studies have shown that plates and foils undergoing pitching or heaving motions provide good approximations of the fluid–structure interactions in swimming fish (Blondeaux *et al.* 2005; Lauder *et al.* 2011; Wen & Lauder 2013; Menon & Mittal 2019), including the reverse von Kármán vortex wake left behind swimming fish and flapping foils (Taneda 1965; Triantafyllou & Triantafyllou 1993).

A variety of fluid–structure interaction models have been proposed to analyse the effect of body flexibility on bending in flows (see e.g. Shelley & Zhang (2011) and references therein). Here, we present a focused literature review. Heathcote & Gursul (2007) conducted experiments on a flapper with a rigid leading edge and flexible tail fixed in a water channel and found that flexibility can enhance both efficiency and thrust production. Eldredge (2008) and Eldredge, Toomey & Medina (2010) numerically simulated the flapping motion of articulated rigid links and found that joint flexibility can reduce the power required for flapping. Alben (2008) used a filament of uniform flexibility to model the tail of swimming fish in the context of the vortex sheet method and predicted enhancement in efficiency rather than thrust when choosing parameters (dimensionless rigidity and reduced pitching frequency) that are consistent with biological data. Quinn, Lauder & Smits (2015) conducted a large set of experiments on two-dimensional pitching and heaving flexible plates at various stiffness values, kinematic parameters and incoming flow speeds. By combining grid search and gradient-based optimization methods, they found that optimizing the pitching angle with heaving can almost double the propulsor efficiency compared with heave-only motions. Hoover *et al.* (2018) conducted simulations combining three-dimensional Navier–Stokes equation with one-dimensional Euler–Bernoulli beam theory to analyse the motion of heaving flexible plates, and identified local peaks in swimming speed over a parameter space consisting of the beam material property and heaving frequency. Also using three-dimensional simulations of pitching plates of uniform flexibility, Dai *et al.* (2012) found that the phase delay ϕ between the leading and trailing edge of the plate decreases with increasing stiffness κ . For large stiffness, the plate moves in no-neck mode (in phase in our notation), in

which thrust production is close to that of a rigid pitching plate with similar trailing-edge displacement.

The effects of uniform flexibility on swimming speed, thrust generation, swimming energetics and stability have been analysed in numerous other experimental and computational studies (see e.g. Shoele & Mittal 2016; Feilich & Lauder 2015; Michelin & Llewellyn Smith 2009; Combes & Daniel 2001; Hua, Zhu & Lu 2013; Wang 2020; Ryu *et al.* 2019). Specifically, Liu & Bose (1997), Heathcote, Wang & Gursul (2008) and Tangorra *et al.* (2010) have indicated that flexibility could lessen or prevent thrust production. Flexible propulsors have also been used in man-made biomimetic underwater autonomous vehicles (see e.g. Fujiwara & Yamaguchi 2017; Katzschmann *et al.* 2018; Gibouin *et al.* 2018; Zhu *et al.* 2019; White, Lauder & Bart-Smith 2021). Most notable is the Tunabot design of Zhu *et al.* (2019) and White *et al.* (2021) which mimics the shape and bending kinematics of yellowfin tuna (*Thunnus albacares*) and Atlantic mackerel (*Scomber scombrus*).

While most studies have focused on uniformly flexible bodies, Combes & Daniel (2003) and Lucas *et al.* (2014) noted that the stiffness along the fish body is not uniform, but decreases towards the tail, and that the propulsor becomes highly flexible at the flexion point of the body (see figure 1*a*). To explore the effects of non-uniform flexibility on efficiency and thrust production, Lucas *et al.* (2015) considered flexible plates of inhomogeneous stiffness undergoing heaving and pitching motions in a water tunnel and found that non-uniform stiffness can improve thrust production, and that in order to achieve optimal propulsion, the morphologic factor (flexion ratio) and kinematic factor (motion type and motion parameters) should be considered simultaneously. Vincent, Liu & Kanso (2020*a*) and Vincent *et al.* (2020*b*) analysed the effect of non-uniform flexibility on flight performance in the context of a tumbling wing model, and found that wing tip flexibility that follows the empirical rules reported in Lucas *et al.* (2014) leads to improved flight performance.

In this paper, we use a simplified two-link fish model to analyse the influence of active and passive bending on the swimming speed and efficiency. Two-link models are commonly used to study the effect of flexibility on the performance of flapping bodies (Eldredge *et al.* 2010; Wan, Dong & Huang 2012; Li, Dong & Liu 2015). We solve for fluid–structure interactions in the context of the vortex sheet method as described in § 2. This method is best suited for capturing hydrodynamic effects at intermediate Reynolds numbers ($Re \sim 10^3$ – 10^5), and has been used extensively to analyse problems of fluid–structure interactions in this Re number regime, including ring formation at the edge of a circular tube (Nitsche & Krasny 1994) and wakes of oscillating plates (Jones 2003; Sheng *et al.* 2012), falling cards (Jones & Shelley 2005), flapping flexible flags (Alben 2008, 2009), swimming plates (Wu 1971), hovering flyers (Huang, Nitsche & Kanso 2016; Huang *et al.* 2018) and schooling of swimming plates (Heydari & Kanso 2021) Here, we use the implementation of Nitsche & Krasny (1994). In § 3, we report the effects of both active and passive bending on the swimming performance of sinusoidally pitching swimmers compared with rigid flapping. In § 4, we discuss these findings in the context of existing work and highlight the implications of our results for the design of underwater autonomous vehicles.

2. Problem formulation

We model the flexible swimmer as a planar two-link body of total length L , negligible thickness $e \ll L$ and total mass per unit depth $m = \rho eL$, where ρ is both the swimmer

Antiphase tail flexion enhances swimming performance

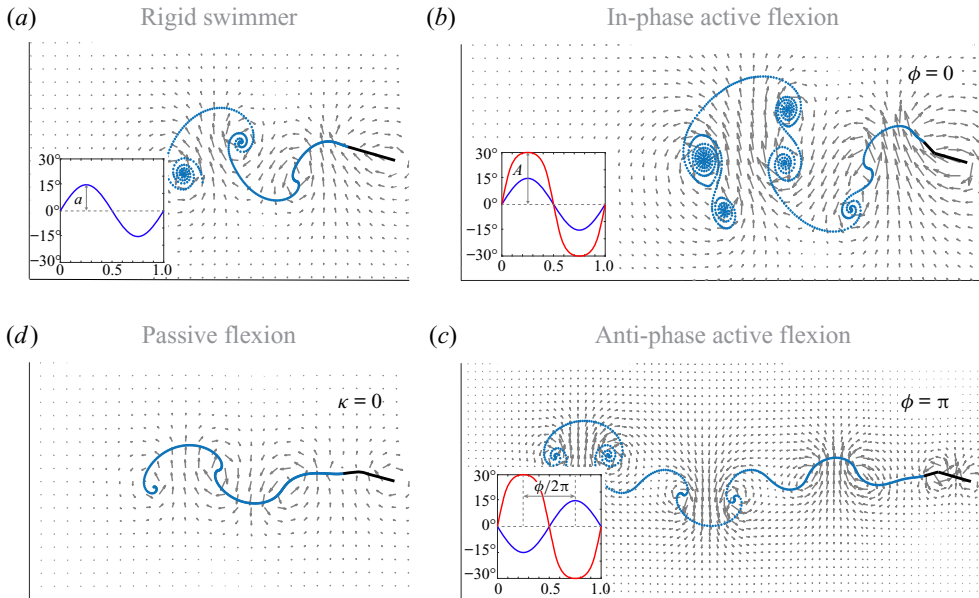


Figure 2. Wake and flow velocity of free swimmers. (a) Rigid swimmer undergoing periodic pitching (inset) of period $T = 1$ and amplitude $a = 15^\circ$. (b,c) Active bending with both anterior and posterior sections undergoing periodic pitching albeit at different amplitude and phase. The anterior follows the same pitching motion (blue line in inset) as the rigid swimmer while the relative rotation of the posterior follows a prescribed Jacobi elliptic sine function (red line in inset) with flexion amplitude $A = 30^\circ$, flexion ratio $l/L = 0.7$, elliptic modulus $M = 0.9$ and phase (b) $\phi = 0$ (in phase) and (c) $\phi = \pi$ (antiphase). (d) Passive bending of posterior while anterior follows the same prescribed pitching as the rigid swimmer. Joint parameters are set to $\kappa = 0$ and $c = 1$. The dissipation time is set to be $T_{diss} = 1.625T$ in (a–c) $\sqrt{2.09}T$ in (d).

and fluid density, assuming a neutrally buoyant fish. The flexion point indicates where the anterior link (of length l) is joined to the posterior link (see figure 1a). The anterior link undergoes sinusoidal pitching motion $\theta_a(t) = a \sin(2\pi ft)$, where θ_a is the angle relative to the swimming direction, taken to be parallel to the x axis. Here, a is the flapping amplitude, $f = 2\pi/T$ the flapping frequency and T the flapping period. When the posterior is connected rigidly to the anterior link at zero flexion, the posterior motion $\theta_p(t)$ is equal to $\theta_a(t)$ and the flexion angle $\alpha = \theta_p - \theta_a$ is identically zero for all time. The two links form a single rigid plate (figure 2a) whose swimming motion due to sinusoidal pitching has been extensively analysed (Jones 2003; Sheng *et al.* 2012; Moored & Quinn 2019; Labasse, Ehrenstein & Meliga 2020; Heydari & Kanso 2021). To explore the effects of body bending on swimming, we consider two cases: (i) active bending where the flexion angle $\alpha(t)$ is controlled by the swimmer and (ii) passive bending where the flexion angle $\alpha(t)$ is dictated by the physics of fluid–structure interactions.

When the two-link swimmer bends actively, we allow the anterior link to have a phase advantage of magnitude ϕ relative to the flapping motion of the posterior link. At $\phi = 0$, both anterior and posterior links flap in phase and the swimmer bends in the direction of flapping (figure 2b); for $\phi = \pi$, they flap antiphase resulting in bending in the opposite direction to the anterior pitching motion (figure 2c). The flexion angle $\alpha(t)$ follows a Jacobi elliptic sine function $\alpha(t) = A \operatorname{sn}(4Kft, M)$, where A is the maximum flexion angle and M is the elliptic modulus that controls the shape of the elliptic sine function. As $M \rightarrow 0$, the elliptic sine function tends to a sinusoidal function and as $M \rightarrow 1$, it approaches a square wave shape. The parameter K is introduced to ensure that the flapping frequency

of the posterior link is the same as that of the anterior link; namely, $K = F(\pi/2, M)$, where $F(\pi/2, M)$ is the incomplete elliptic integral of the first kind. In this paper, without other specification, we fix $M = 0.9$ and explore the effects of the anterior-to-posterior phase difference ϕ , flexion ratio l/L and maximum flexion angle A on the swimming performance.

We write the equations governing the self-propelled motion of the two-link swimmer in non-dimensional form. To this end, we scale all parameter values using $L/2$ as the characteristic length scale, $1/f$ as the characteristic time scale and $\rho(L/2)^2$ as the characteristic mass per unit depth. Accordingly, velocities are scaled by $Lf/2$, forces by $\rho f^2(L/2)^3$, moments by $\rho f^2(L/2)^4$ and power by $\rho f^3(L/2)^4$. The equation of motion governing the free swimming $x(t)$ is given by Newton's second law:

$$m\ddot{x} = F_x - D_x. \tag{2.1}$$

Here, F_x and D_x denote the x components, in the swimming direction, of the hydrodynamic pressure force normal to the swimmer and the drag force due to skin friction tangential to the swimmer. By definition, F_x and D_x can be either positive or negative, and thus can propel the swimmer forward or resist its motion. However, for notational convenience, we refer to F_x as thrust and D_x as drag, though technically they can be either; we refer to negative values of F_x as negative thrust.

We calculate the hydrodynamic pressure force in the context of the inviscid vortex sheet model (Nitsche & Krasny 1994; Huang *et al.* 2016, 2018; Heydari & Kanso 2021), and the drag force based on a skin friction model that emulates the effect of fluid viscosity (Fang 2016; Ramanarivo *et al.* 2016). A brief overview of the vortex sheet method and its numerical implementation is given in [Appendices A and C](#). Detailed expressions of the fluid forces acting on the swimmer are given in [Appendix B](#).

When the swimmer bends passively, the relative rotation $\alpha(t)$ of the posterior end is not prescribed *a priori* and follows from the physics of fluid–structure interactions. Considering that the rotational joint at the flexion point is equipped with a torsional spring of stiffness κ and damping coefficient c , we write the equation governing the rotational motion of the posterior link as

$$I_p(\ddot{\theta}_a + \ddot{\alpha}) + c\dot{\alpha} + \kappa\alpha = M_p + M_{inertia}, \tag{2.2}$$

where I_p and M_p are the moment of inertia and hydrodynamic moment acting on the posterior link about the flexion point and $M_{inertia}$ is an inertial moment that arises because the flexion point about which the moments are balanced is moving; see details in [Appendix B](#).

To assess the swimming performance of the two-link swimmer, we introduce four metrics: the period-average swimming speed $U = \int_t^{t+T} \dot{x} dt$ at steady state, the thrust force F_x , the period-average input power $P = \int_t^{t+T} P(t) dt$ required to maintain the prescribed flapping motions and the propulsion efficiency $mU^2/2PT$ defined as the kinetic energy of the swimmer divided by the input work over one flapping period; see [Appendix B](#) for more details.

3. Results

We compare the free swimming that results from flapping while undergoing active and passive bending with that of rigidly flapping. All swimmers have the same total length L and undergo the same sinusoidal pitching motion about their leading edge $\theta_a = a \sin(2\pi ft)$

with $a = 15^\circ$ and $f = 1$, albeit exhibiting distinct bending patterns. Figure 2 shows snapshots of the wake represented by the free vortex sheet and velocity field generated by a swimmer undergoing rigid flapping (figure 2a), in-phase active bending with flexion amplitude $A = 30^\circ$ and flexion ratio $l/L = 0.7$ (figure 2b), antiphase active bending at the same flexion amplitude and ratio (figure 2c) and passive bending (figure 2d). All snapshots are taken at the same instant in the flapping cycle (at $0.25T$ after steady state has been reached). Compared with the rigid swimmer, in-phase flexion produces wider wakes and larger leading-edge circulation and instantaneous flow speeds, while antiphase flexion is characterized by a leaner, longer wake with weaker leading-edge circulation and lower flow speeds. The main features of the instantaneous flow during antiphase flexion, namely the leaner wake and weaker leading-edge circulation, are also observed in passive bending of the swimmer body. These flow features are important for the hydrodynamic forces exerted on the swimmer as discussed later.

We quantitatively evaluate the steady-state motion of the rigid and actively bending swimmers in figure 2(a–c). In figure 3, we report the instantaneous (solid lines) and period-average (dashed lines) values of the swimming speed, thrust force, input power and circulation. On average, rigid flapping produces the lowest swimming speed while antiphase flexion the highest. Fluctuations around the average swimming speed are smallest for the swimmer undergoing antiphase flexion. The discrepancy in average swimming speeds between the three flapping modes is surprising at first sight given that the average values of the thrust force are comparable. Note that in all cases, average thrust F_x is positive. However, a closer look at the instantaneous thrust shows that the swimmer undergoing antiphase flexion hardly experiences negative thrust over its flapping cycle. In-phase flexion leads to negative thrust of high magnitudes over larger subintervals of the flapping cycle, as highlighted further in figure 4. Consequently, the required input power for in-phase flexion is largest compared to both rigid flapping and antiphase flexion. This is also true of the overall wake circulation. It is worth noting that, by Kelvin’s circulation theorem, circulation around the leading edge must be equal to the overall circulation in the wake. Therefore, compared to rigid flapping, in-phase flexion increases the circulation around the leading edge of the swimmer while antiphase flexion decreases leading-edge circulation as noted qualitatively in figure 2.

The results in figure 3 indicate that the swimmer undergoing antiphase flexion achieves higher swimming speed at lower power requirement and energetic cost. To elucidate the hydrodynamic forces at play, we report in figure 4 the force hodograph defined as a plot of the lateral pressure force F_y versus thrust F_x acting on each swimmer. Snapshots of the distribution of hydrodynamic pressure forces along the swimmers are depicted in bottom row of figure 4, and indicate that the x component of the forces on the anterior and posterior sections during antiphase flexion act opposite to each other as in a tug-of-war, leading to overall reduction in thrust values. Importantly, antiphase flexion also reduces the lateral force and negative thrust, with negative thrust experienced only over a small subinterval of the flapping period, as noted earlier. In contrast, in-phase flexion significantly increases the lateral force and negative thrust.

To explain the effect of flexion on the lateral force experienced by the swimmer, it is instructive to re-examine the flow field around the rigid and actively bending swimmers in figure 2(a–c). A large leading-edge vortex is known to generate large lift in flapping flight (see e.g. Ellington 1984; Dickinson, Lehmann & Sane 1999). In swimming, larger leading-edge circulation creates larger lateral force, which explains why, compared with rigid flapping, in-phase flexion increases the lateral force acting on the swimmer while antiphase flexion decreases it. Lift is beneficial for flight but large lateral forces are

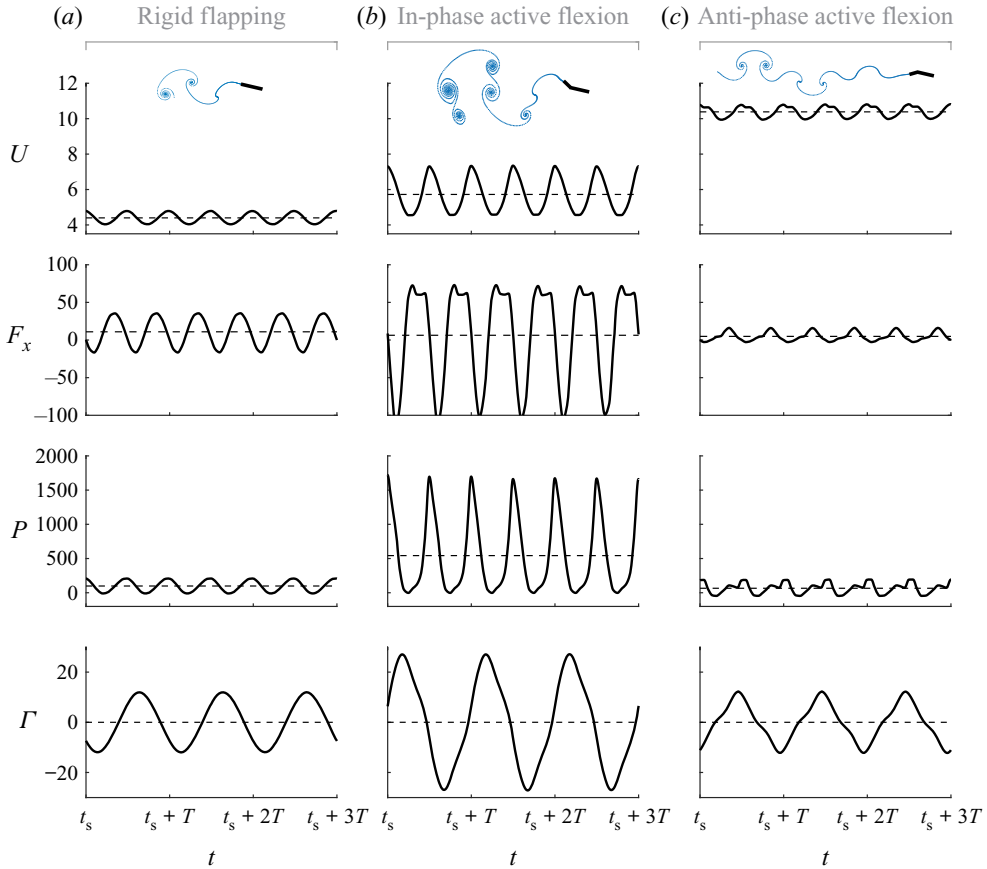


Figure 3. Active anterior-to-posterior bending of free swimmers. Time-dependent speed U , thrust F_x , input power P and circulation Γ for (a) rigid swimmer undergoing pitching at $a = 15^\circ$, $T = 1$, and active flexion (b) at phase difference $\phi = 0$ and (c) at phase difference $\phi = \pi$. In (b,c), flexion ratio $l/L = 0.7$ and flexion angle $A = 30^\circ$. Solid lines represent the instantaneous values and dashed lines represent time-period averages. Average thrust F_x is positive in all cases. The results are shown after the swimmers have reached steady state, $t_s = 15T$. The dissipation time is set to be $T_{diss} = 1.625T$.

detrimental to swimming speed, as noted in Drucker & Lauder (2000) for fish and recapitulated here in the context of our swimmer model.

To further analyse the difference in the swimming performance between rigid flapping and flapping with in-phase and antiphase active bending, we fix the swimmer in an oncoming uniform flow of speed U and we compute the hydrodynamic drag forces in each case. Unlike in the case of the free swimmer where the period-average of the total thrust and drag forces must be zero, here the swimmer may experience a net period-average force. In figure 5, we report the drag force, input power and circulation in the wake of the fixed swimmer. Period-average values are shown in dashed lines. Compared to rigid flapping, antiphase flexion reduces instantaneous drag, power and circulation, while in-phase flexion increases all three quantities. Reduced drag implies lower thrust requirement for steady-state swimming, which provides another perspective for understanding the improved performance of antiphase flexion.

Antiphase tail flexion enhances swimming performance

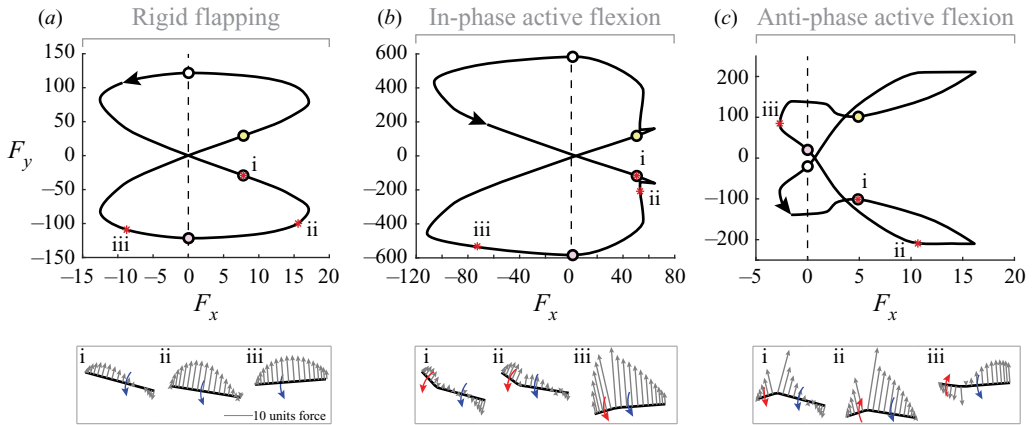


Figure 4. Active anterior-to-posterior bending can minimize lateral forces and negative thrust. Force hodograph of (a) rigid flapping, (b) in-phase flexion and (c) antiphase flexion for the same cases shown in figure 3. The arrow indicates the direction of time. The white, blue, red and yellow points are: $t/T = 0, 0.25, 0.5$ and 0.75 . Bottom row shows force distribution at the instants indicated by red stars. Blue arrows represent pitching direction of the anterior link motion and red arrows that of the posterior link.

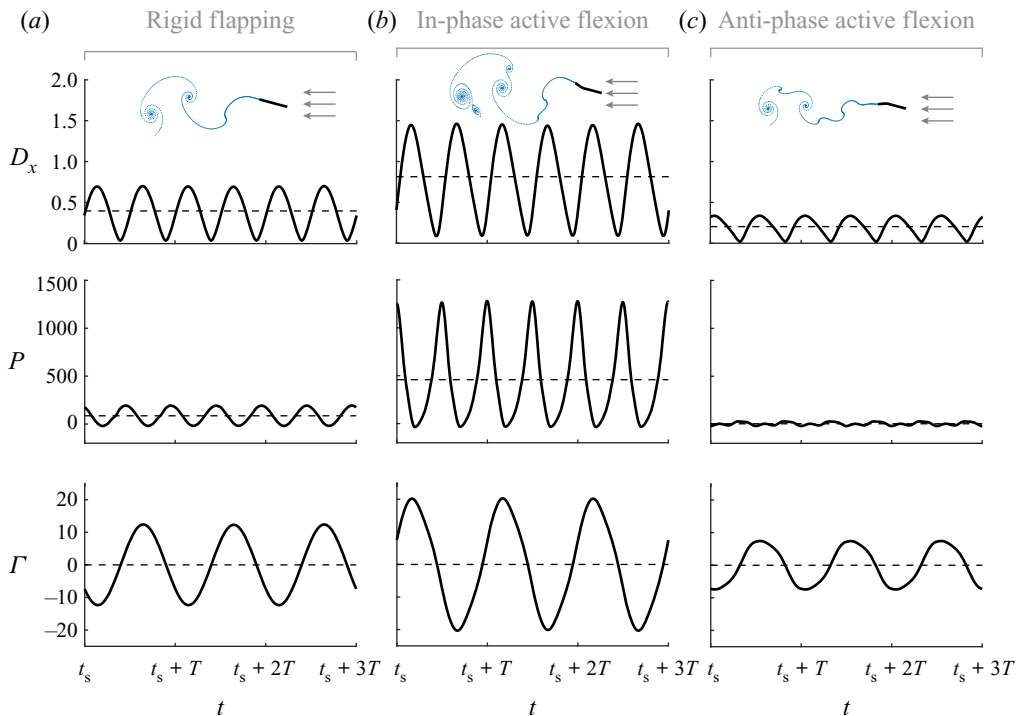


Figure 5. Active anterior-to-posterior bending of swimmers fixed in oncoming flow. Time-dependent drag force D_x , input power P and wake circulation Γ for (a) rigid swimmer, (b) in-phase active flexion ($\phi = 0$) and (c) antiphase active flexion ($\phi = \pi$). In all cases, the swimmer is fixed in a uniform oncoming flow at $U = 9$. Results are shown after the swimmers have reached steady state, $t_s = 11T$. Solid lines represent the instantaneous values and dashed lines represent time-period averages. The dissipation time is set to be $T_{diss} = 1.625T$.

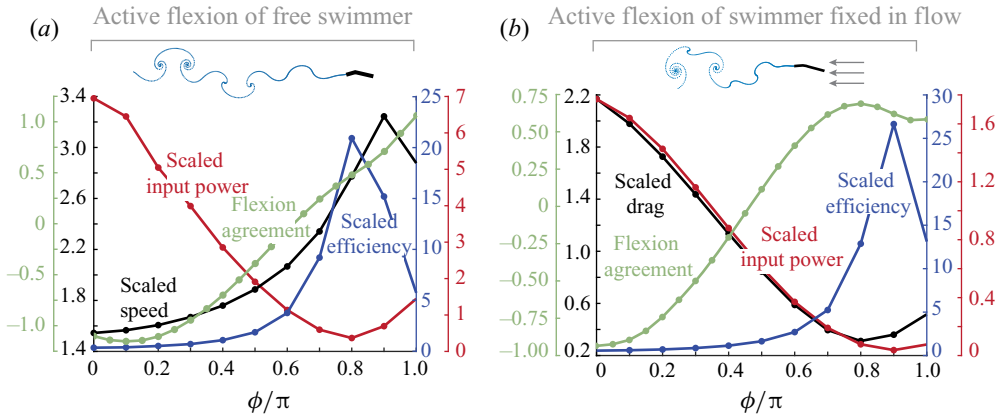


Figure 6. Performance of active anterior-to-posterior bending as a function of phase. (a) Free swimmer: swimming speed (black), power requirement (red) and efficiency (blue) are scaled by the corresponding values of a rigid swimmer. (b) Swimmer fixed in oncoming flow of uniform speed U : drag force (black), power requirement (red) and efficiency (blue) are scaled by the corresponding values of a pitching rigid plate. Parameter values are set to $\alpha = 15^\circ$, $l/L = 0.7$, $A = 20^\circ$, $M = 0.9$, $U = 9$. Flexion agreement parameter Z between the relative velocity of an actively bending posterior and the fluid velocity generated by a passively bending swimmer at zero stiffness $\kappa = 0$ (green) as a function of phase ϕ during (a) free swimming and (b) holding station in oncoming flow $U = 5$.

We next examine the period-average performance of actively bending swimmers as a function of anterior-to-posterior phase difference ϕ . In figure 6(a), we consider the case of free swimming, we fix the flexion ratio $l/L = 0.7$ and flexion amplitude $A = 20^\circ$ and we plot the swimming speed U , input power P and efficiency η versus ϕ , all scaled by the corresponding values of a rigidly flapping swimmer U_{rigid} , P_{rigid} and η_{rigid} , respectively. We find that active bending is always beneficial in terms of enhanced speed relative to rigid flapping, albeit at an increased power requirement. Importantly, as the anterior-to-posterior bending changes from in-phase flexion to flexion at a phase lag, the scaled speed increases and the scaled power requirement decreases. Optimal performance occurs at $\phi = 0.9$ and $\phi = 0.8$ in terms of maximum swimming speed and minimum input power and maximum efficiency, respectively. In figure 6(b), we fix the swimmer in oncoming flow of uniform speed U and compute the scaled drag force D_x , input power P and efficiency η as a function of ϕ scaled by the corresponding values of a fixed rigid flapper. We find that, as the anterior-to-posterior bending changes from in-phase flexion to flexion at a phase lag, the scaled drag decreases and so does the scaled power requirement. Specifically, analogous to the free swimmer, drag and input power are minimal at $\phi = 0.8$. Taken together, these results imply that antiphase active flexion is near optimal for enhancing speed and efficiency and reducing drag force and power requirement.

To complete this analysis, we also explored the effect of the flapping parameter M on the swimming performance. We found that for a fixed phase, the swimming speed and efficiency change monotonically with M with maximum speed and minimum efficiency as $M \rightarrow 1$ (see figure 12 in Appendix B). That is, reversing the bending direction with a quick flicker improves speed at the cost of decreasing efficiency.

Is active bending necessary for obtaining this enhancement in swimming speed and efficiency over rigid flapping? To address this question, we examine the free swimming of a passively bending swimmer, where the posterior end flaps passively under the effect of hydrodynamic forces and moments generated by the pitching motion of the anterior section. Elastic forces due to a spring of stiffness κ located at the flexion point are also

Antiphase tail flexion enhances swimming performance

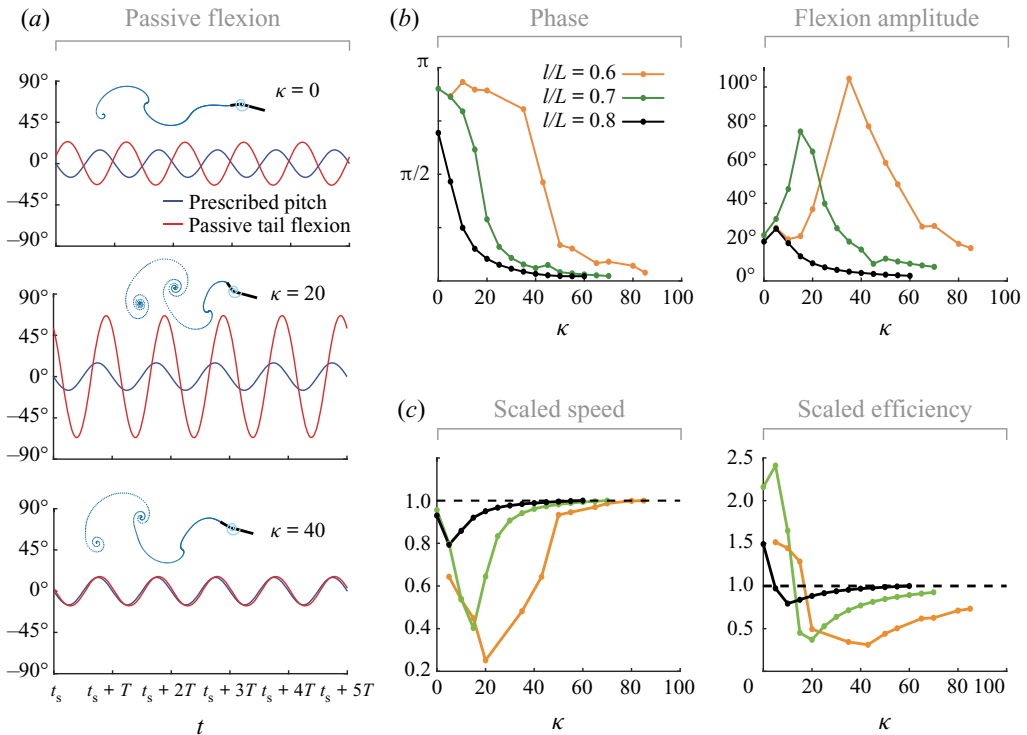


Figure 7. Passive anterior-to-posterior bending of free swimmers. (a) Top to bottom: passive flexion angle α of posterior end at three stiffness values: $\kappa = 0$, $\kappa = 20$ and $\kappa = 40$. The anterior link is pitching at $a = 15^\circ$. The damping ratio is set to $c = 1$, and flexion ratio to $l/L = 0.7$. (b) The bending parameters (phase ϕ (left) and maximum flexion angle A (right)) and (c) swimming speed U (left) and propulsion efficiency η (right) as a function of stiffness κ for three flexion ratios $l/L = 0.6, 0.7$ and 0.8 reported in orange, green and black, respectively. The results are shown after the swimmers have reached steady state, $t_s = 25T$. The dissipation time is set to be $T_{diss} = \sqrt{2.09}T$.

at play. In [figure 7\(a\)](#), we keep all parameter values the same as those used for the actively bending swimmer, and, from top to bottom, we report the flapping motions of the anterior and posterior ends for stiffness values $\kappa = 0, 20$ and 40 . At zero stiffness, flexion introduces no restoring forces and moments. The posterior part rotates antiphase relative to the flapping motion of the anterior part, at an amplitude comparable to the anterior pitching amplitude. The associated wake, shown in [figure 2\(d\)](#) and represented by the free vortex sheet in the inset of [figure 7\(a\)](#), shares similar features to the wake obtained during antiphase active flexion. At spring stiffness $\kappa = 20$, the flexion amplitude increases ($\alpha_{max} \approx 80^\circ$), and the wake also exhibits larger lateral dispersion. At large stiffness $\kappa = 40$, the posterior part rotates in phase with the anterior part at the same flapping amplitude in a way reminiscent of rigid flapping, as reported in [Dai et al. \(2012\)](#) for flexible pitching plates.

The relative motion of the posterior part is close to a sinusoidal function for all stiffness values κ . Therefore, for each κ value, we fit $\alpha(t)$ by a sine function $\alpha(t) = A \sin(\omega t - \phi)$ using a standard algorithm ([Moré & Sorensen 1983](#)). For all fitting, we have at least 95 % confidence and frequency $\omega \approx 2\pi$, thus ensuring convergence of the fitting. In [figure 7\(b\)](#), we report the fitted flexion amplitude A and phase difference ϕ as a function of spring stiffness κ for three distinct flexion ratios $l/L = 0.6, 0.7$ and 0.8 . We find that for all

l/L , as stiffness κ increases, the phase difference ϕ decreases monotonically from π to 0 implying that the posterior flapping motion changes from antiphase to in-phase motion. The maximum flexion angle A first increases with increasing κ , then decreases to nearly zero at large stiffness implying rigid flapping of both anterior and posterior ends.

We compute the associated swimming speed and efficiency for each stiffness value κ and we scale the results by those of a rigid swimmer (see figure 7c). Clearly, the swimmer with passive flexion never surpasses the swimming speed of a rigid swimmer. At very low stiffness ($\kappa \approx 0$), passive flexion results in a swimming speed close to that of rigid flapping while doubling the swimming efficiency. The increase in swimming efficiency at small stiffness comes purely from a decrease in power requirement compared to rigid flapping. This is in contrast to active flexion where the enhancement in swimming speed and efficiency noted in figure 6 comes at an increase in power requirement relative to rigid flapping. As κ increases, the scaled swimming speed and propulsion efficiency decrease, indicating that restoring elastic forces are detrimental to both speed and efficiency. For large κ , the speed and efficiency converge to the same speed and efficiency as the rigid swimmer, consistent with the results of Dai *et al.* (2012). We repeat this analysis for three flexion ratios $l/L = 0.6, 0.7, 0.8$. The scaled speed seems to increase monotonically with increasing l/L , but the scaled efficiency seems to peak at $l/L = 0.7$ but only for a range of small κ values. These findings imply that, unlike flapping insect wings (Ellington 1985; Huang, Nitsche & Kanso 2015), restoring elastic forces seem to be detrimental to swimming performance. Swimming efficiency peaks at low stiffness values when the restoring spring forces are weak and the posterior end is driven passively by the fluid forces.

Could the swimmer learn from passive flexion to improve its performance by bending actively in a way that exploits the hydrodynamic forces generated naturally during passive flexion? The results in figures 2–6 for actively bending swimmers suggest that maximum benefit occurs for near antiphase flexion, whereas the results in figure 7 show that maximum efficiency for passively bending swimmers occurs for zero stiffness ($\kappa = 0$) for which the posterior bends antiphase. Importantly, the main features of the instantaneous flow during antiphase active flexion (figure 2c) are also observed in passive bending at zero stiffness (figure 2d). We thus posit that active bending is most beneficial when the swimmer actively beats its tail in a direction that takes advantage of the natural flows that arise during passive bending. To test this hypothesis, we define a flexion agreement parameter Z that aims to relate passive and active bending. Starting from a swimmer bending passively at zero spring stiffness $\kappa = 0$ (figures 2d and 7a), we assume a hypothetical posterior that is actively flapping about the flexion point of the swimmer at a relative angle $\alpha = A \sin(2\pi t - \phi)$. We compute the fluid velocity $\mathbf{u}(s, t)$ induced by the passively bending swimmer along the hypothetical posterior that is bending actively. The flexion agreement parameter Z is given by

$$Z = \frac{1}{T} \frac{1}{L-l} \int_0^T \int_0^{L-l} \mathbf{u}(s, t) \cdot \mathbf{v}(s, t) \, ds \, dt, \quad (3.1)$$

where $\mathbf{v}(s, t) = s \dot{\alpha} \mathbf{n}$ is the relative velocity of the hypothetical actively flapping posterior and s is a dummy variable denoting the rectilinear distance from the flexion point. Positive values of the flexion agreement parameter imply a beneficial interaction between the flow generated during passive flexion and the velocity of the hypothesized posterior during active flexion, whereas negative values indicate a detrimental one.

In figure 6(a), we set A to be equal to the maximum flexion angle of the passively bending swimmer, and we vary ϕ from 0 to π . We find that the agreement parameter Z ,

Antiphase tail flexion enhances swimming performance

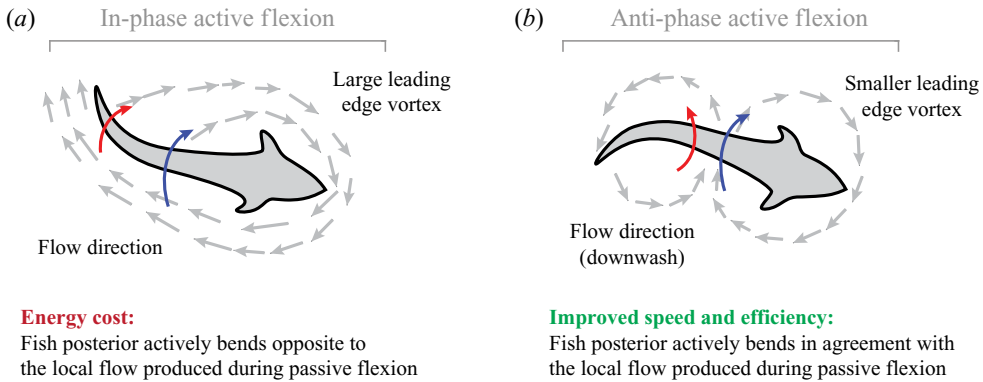


Figure 8. Active bending in agreement with passive hydrodynamics improves swimming performance. Illustration of the interaction with the flow field for (a) in-phase and (b) antiphase active flexion. Maximum improvement in swimming performance occurs when the tail benefits from flows created by the anterior portion of the fish body. Light grey arrows represent the flow direction (in accordance with figure 2). Red arrows and blue arrows represent the flapping direction of the posterior and anterior sections of the fish body, respectively.

normalized by its maximum value, is largest for antiphase active flexion and smallest near in-phase active flexion. This result indicates that antiphase active flexion matches best the local flow created during passive flexion, whereas in-phase active flexion acts opposite to these flows. That is, the swimmer during antiphase flexion can utilize better the flow field generated by the pitching motion of its anterior section, and thus it can achieve higher swimming speed and efficiency compared with in-phase flexion.

To emphasize the effect of the interaction between the flow field and kinematics of active flexion on swimming performance, we schematically summarize the two cases of in-phase and antiphase flexion in figure 8. During in-phase active flexion, the flow field is characterized by a strong leading-edge vortex around the anterior section of the fish and large lateral forces. Further, the posterior part bends opposite to the local flow field of a passively bending swimmer. During antiphase active flexion, the leading-edge vortex is smaller and it is followed by a counter-rotating vortex around the fish mid-section such that the posterior part is moving in synchrony with the downwash flow induced by this counter-rotating vortex. The flow field is helping the motion of the posterior end. These results indicate that active body deformations that are in agreement with local flows produced during passive deformations are more advantageous for enhancing swimming speeds and efficiencies.

Lastly, we explore the effect of maximum flexion angle A and flexion ratio l/L on the period-average values of the swimming speed and efficiency for both in-phase and antiphase active flexion. Specifically, we examine the range $l/L \in [0.4, 1.0]$ and $A \in [0^\circ, 60^\circ]$ for $\phi = 0$ and $\phi = \pi$. In figure 9, we report the period-average values normalized by the corresponding values for a rigid swimmer with pitching amplitude equal to the anterior part amplitude. We highlight in light and dark grey respectively the regions in the parameter space where the flexible swimmer outperforms and underperforms the rigid swimmer. The swimmer with in-phase flexion swims slower than the rigid swimmer for small flexion ratios ($l/L < 0.7$) and high flexion amplitudes ($A > 40^\circ$), and swims faster than the rigid swimmer otherwise. This swimmer, however, is always less efficient than the rigid swimmer for reasons explained previously. In figure 9(b), for most parameter values, the antiphase swimmer outperforms the rigid swimmer in terms of swimming speed and efficiency. Note that the region with the highest swimming speed advantage lies

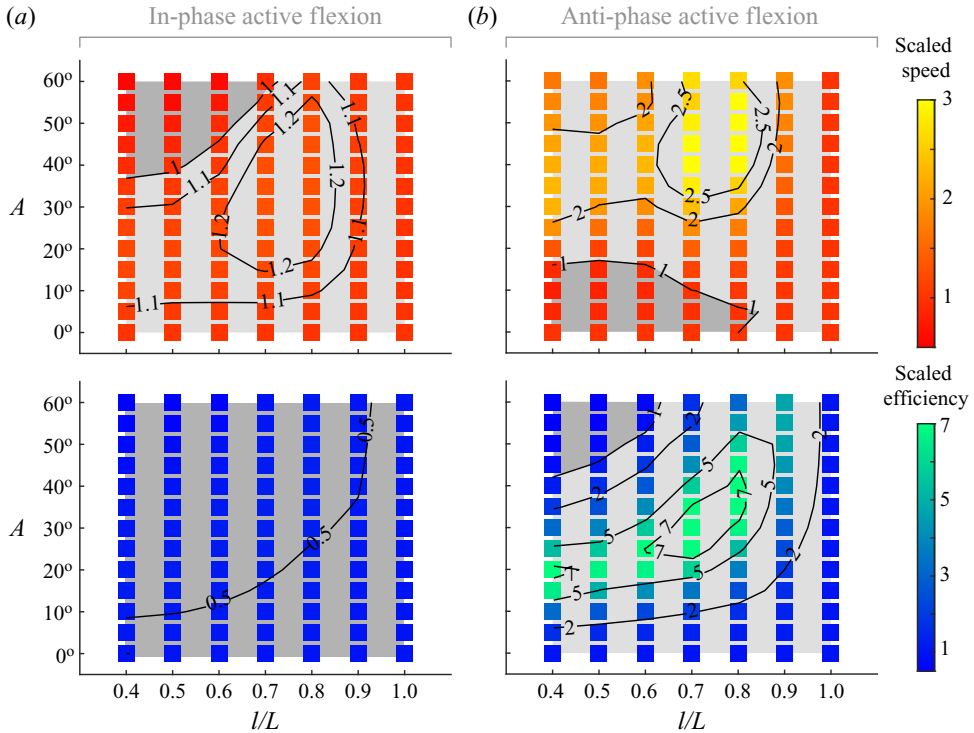


Figure 9. Swimming performance of actively flexing swimmer scaled by performance of rigid swimmer. Average speed and efficiency versus flexion ratio l/L and flexion angle A for (a) in-phase flexion ($\phi = 0$) and (b) antiphase flexion ($\phi = \pi$). The amplitude for the proximal part and the elliptic modulus are $a = 15^\circ$ and $M = 0.9$. Dark grey areas indicate regions of diminished performance while light grey areas indicate improved performance over a rigid swimmer. The dissipation time is set to be $T_{diss} = 1.625T$.

in $l/L \in [0.6, 0.8]$ and $A \in [30^\circ, 60^\circ]$, and the region with the highest efficiency advantage lies in $l/L \in [0.6, 0.8]$ and $A \in [25^\circ, 50^\circ]$.

We compare the regions of highest swimming speed and efficiency obtained during antiphase flexion in figure 9(b) with the design parameters of biological fish reported in figure 1. In figure 10, we plot the flexion angle A as a function of the flexion ratio l/L for the fish data in figure 1 and we superimpose on this design space the regions of 200% enhancement in speed and 300% enhancement in efficiency from figure 9(b). As shown in figure 10, there is significant overlap between these regions of improved performance and the biological data. Indeed, all biological data lie within the region of improved efficiency.

Many of these fish are known to exhibit migratory behaviour that requires efficient swimming. Even baby clownfish are reported to migrate over long distances (Simpson *et al.* 2014). Tuna can cover 7600 km in one travelling phase (Itoh, Tsuji & Nitta 2003) and tiger shark are capable of travelling long distances in short times (Simpfendorfer 2009). Our results in figure 10 are consistent with these facts: tiger shark and clownfish lie in the intersection region of improved speed and efficiency, and tuna lies in the region with highest increase in efficiency. On the other hand, butterflyfish, which are only known to migrate over short distances during spawning (Yabuta 1997), and Koi, for which there is no evidence of migration, lie in the regions characterized by smaller increase in speed and efficiency.

Antiphase tail flexion enhances swimming performance

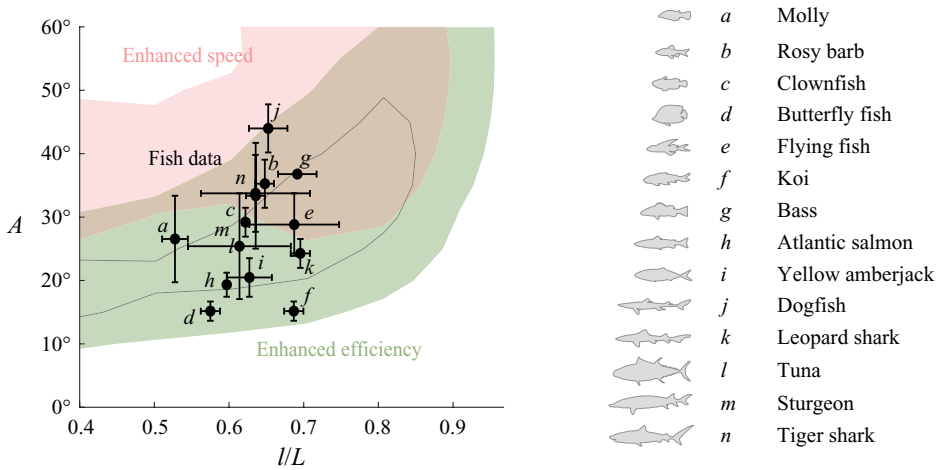


Figure 10. Relation to fish swimming behaviour. Comparison of fish flexion parameters (black dots, data from Lucas *et al.* (2014)) and regions of optimal performance predicted by our model (antiphase active flexion swimmer): pink region corresponds to 200 % enhancement in swimming speed and green region corresponds to 300 % enhancement in swimming efficiency, both compared with a swimmer of the same total length rigidly flapping with no flexion. Overlap of the two regions is indicated in beige. The contour grey line encloses a region of 600 % enhancement in efficiency.

To conclude this section, a few remarks on Reynolds numbers are in order. The fish listed in figure 10 vary in length, swimming speed, tailbeat frequency and cross-sectional geometry, whereas the model abstracts these details into a simple two-link fish and flows represented by the vortex sheet method. Specifically, the biological data span a wide range of Reynolds numbers ($Re \sim 10^3\text{--}10^7$). In the model, we used non-dimensional parameters with fluid density $\rho = 1$, fish length $L = 2$ and tailbeat frequency $f = 1$, and we obtained a range of dimensionless swimming speeds $U = 4.5\text{--}10$ by varying the bending kinematics. Here, $U = 4.5$ is the speed of the rigid flapper. Because skin friction is accounted for in the model, it is possible to calculate an effective $Re = \rho LU/\mu$ in the context of our dimensionless vortex sheet model, where μ is a dimensionless viscosity. Starting from the density (10^3 kg m^{-3}) and viscosity (10^{-3} Pa s) of water, and using the range of length scales and flapping frequencies from the biological fish data, we arrive at a range of non-dimensional viscosity $\mu \sim 10^{-8}\text{--}10^{-2}$. However, in our model, our choice of the drag coefficient $C_d = 0.664\sqrt{\rho\mu L} = 0.04$ (see Appendix B) fixes the value of the dimensionless viscosity to $\mu \approx 10^{-3}$. Thus, Re for the range of swimming speeds ($U = 4.5\text{--}10$) obtained in the model is of the order $Re \sim 10^4$.

4. Conclusion

We analysed the swimming performance of flapping swimmers undergoing active and passive deformations. Whereas fish exhibit a variety of swimming modes (Sfakiotakis *et al.* 1999), we simplified body deformations to account for only anterior-to-posterior bending, with one degree of freedom describing the relative rotation between the two sections. We explored the effects of morphological and kinematic parameters on the swimming speed and efficiency.

We found that passive body bending, at negligible body stiffness and minor elastic forces, caused anterior-to-posterior antiphase flexion. This antiphase flexion is dictated

by the flow physics and causes the swimmer's morphology to get more streamlined compared to rigid flapping with no flexion, thus creating leaner wakes that reduce drag and power requirement and increase efficiency. While drag reduction is desirable for improved efficiency, passive bending also reduced thrust production, thus diminishing swimming speed. Interestingly, restoring elastic forces seemed detrimental to both swimming speed and efficiency for a range of intermediate stiffness values. These findings are consistent with the hypothesis that for maximum efficiency, the fish tail and posterior body should flex like water, exhibiting little or no resistance to the flows generated by the flapping motion of the anterior portion of the body. This hypothesis could explain how anaesthetized fish, with no muscle activity, placed in periodic wakes generate oscillatory body deformations that allow the fish to swim upstream (Beal *et al.* 2006).

We also found that a swimmer that actively creates antiphase anterior-to-posterior bending enjoyed the same benefits of leaner wake and reduced drag as a passively bending swimmer while mitigating the reduction in thrust and swimming speed. To quantify the hydrodynamic mechanisms leading to this improved swimming performance, we introduced a flexion agreement parameter that compares the active flexion velocity of the swimmer's posterior to the local flow velocity during passive flexion. We found that during active in-phase flexion, the posterior beats opposite to the local flow that would naturally arise during passive flexion, leading to a negative flexion agreement parameter, and thus lower swimming speed and efficiency. During active antiphase flexion, the posterior flaps consistently with the local flow, leading to a positive flexion agreement parameter, and improved speed and efficiency.

These findings suggest tremendous versatility in swimming performance, even when accounting only for coarse anterior-to-posterior bending motions. They indicate that fish can readily and fluidly transition from efficient (passive bending) to fast (active bending) by actively beating their tail in agreement with the local flow generated during passive bending.

To explore the role of flow physics in the convergent bending rules of Lucas *et al.* (2014), we examined the effect of flexion ratio and maximum flexion angle on the performance of swimmers undergoing active antiphase anterior-to-posterior bending. We found an optimal region in this design space that simultaneously enhances swimming speed and efficiency. Importantly, we found that this region has a significant overlap with the fish bending parameters reported in Lucas *et al.* (2014); see figures 10 and 9(b). Fish are able to adjust their swimming speed by altering their tailbeat frequency (Hoover & Tytell 2020); thus, fish could in principle maintain kinematic flapping patterns that optimize efficiency, while increasing their tailbeat frequency to achieve higher swimming speeds.

Taken together, our results have two major implications for understanding the role of body bending in fish swimming. They are consistent with the hypothesis that fish that actively bend their bodies in a fashion that exploits the local hydrodynamics can at once improve speed and efficiency. They also support the hypothesis that flow physics could have provided a selective force for driving the evolution of fish bending patterns.

Beyond the trade-offs between speed and efficiency explored here, our model could be generalized in future studies to examine transient swimming manoeuvres such as turning. Pollard & Tallapragada (2019) proposed that a passive posterior not only improves efficiency, but also improves fish manoeuvrability compared to a rigid body. Drucker & Lauder (2000) pointed out that although large lateral forces are detrimental to fish swimming speed, they can improve fish manoeuvrability. Our model predicts large lateral forces during in-phase active flexion, suggesting that this bending pattern, while not optimal for forward swimming, could be beneficial for turning motions.

These considerations, as well as models of higher fidelity to the fish biomechanics and fluid environment, will be explored in future research.

Finally, our finding that active body bending and tailbeat patterns that match local flow velocities that would be produced naturally by anterior sections of the fish body could lead to improved performance and energy savings might have important implications for understanding the mechanisms driving BCF deformations in swimming (Bozkurttas *et al.* 2007; Ramakrishnan *et al.* 2011), schooling (Becker *et al.* 2015; Li *et al.* 2020; Heydari & Kanso 2021), and navigating ambient unsteady flows (Liao *et al.* 2003).

Supplementary movies. Supplementary movies are available at <https://doi.org/10.1017/jfm.2021.984>.

Acknowledgements. E.K. would like to thank M.J. McHenry for interesting conversations.

Funding. This work was partially supported by the National Science Foundation (grant no. NSF CBET-2100209), the Army Research Office (grant nos. N00014-14-1-0421, N00014-17-1-2287, N00014-17-1-2062) and the Office of Naval Research (grant no. W911NF-16-1-0074).

Declaration of interests. The authors report no conflict of interest.

Data availability statement. The data that support the findings of this study are included in the figures and are available at <https://github.com/haotianh9/Data-for-Paper--Antiphase-tail-flexion-enhances-swimming-speed-and-efficiency>.

Author ORCIDs.

 Haotian Hang <https://orcid.org/0000-0001-5217-8124>;

 Sina Heydari <https://orcid.org/0000-0001-8907-5751>;

 John H. Costello <https://orcid.org/0000-0002-6967-3145>;

 Eva Kanso <https://orcid.org/0000-0003-0336-585X>.

Author contributions. E.K. and J.H.C. designed the study, H.H. and S.H. performed the simulations. All authors contributed equally to analysing data and reaching conclusions, and in writing the paper.

Appendix A. Vortex sheet model

The coupled fluid–structure interaction between the two-link swimmer and the surrounding fluid is simulated using an inviscid vortex sheet model. In viscous fluids, boundary layer vorticity is formed along the sides of the swimmer, and it is swept away at the swimmer’s tail to form a shear layer that rolls up into vortices. In the vortex sheet model, the swimmer is approximated by a bound vortex sheet, denoted by l_b , whose strength ensures that no fluid flows through the rigid plate, and the separated shear layer is approximated by a free regularized vortex sheet l_w at the trailing edge of the swimmer. The total shed circulation Γ in the vortex sheet is determined so as to satisfy the Kutta condition at the trailing edge, which is given in terms of the tangential velocity components above and below the bound sheet and ensures that the pressure jump across the sheet vanishes at the trailing edge.

To express these concepts mathematically, it is convenient to use the complex notation $z = x + iy$, where $i = \sqrt{-1}$ and (x, y) denote the components of an arbitrary point in the plane. The bound vortex sheet l_b is described by its position $z_b(s, t)$ and strength $\gamma(s, t)$, where $s \in [0, L]$ is a curvilinear coordinate along the swimmer starting from its leading edge. The curvilinear coordinate s is introduced since the swimmer is bending, and for the sake of further development of the method on more complex geometries. The separated sheet l_w is described by its position $z_w(\Gamma, t)$, $\Gamma \in [0, \Gamma_w]$, where Γ is the Lagrangian circulation around the portion of the separated sheet between its free end in

the spiral centre and the point $z_w(\Gamma, t)$. The parameter Γ defines the vortex sheet strength $\gamma = d\Gamma/ds$.

By linearity of the problem, the complex velocity $w(z, t) = u(z, t) - iv(z, t)$ is a superposition of the contributions due to the bound and free vortex sheets:

$$w(z, t) = w_b(z, t) + w_w(z, t). \tag{A1}$$

In practice, the free sheet l_w is regularized using the vortex blob method to prevent the growth of the Kelvin–Helmholtz instability. The bound sheet l_b is not regularized in order to preserve the invertibility of the map between the sheet strength and the normal velocity along the sheet. The velocity components $w_b(z, t)$ and $w_w(z, t)$ induced by the bound and free vortex sheets, respectively, are given by

$$w_b(z, t) = \int_0^L K_\delta(z - z_b(s, t))\gamma(s, t) ds, \quad w_w(z, t) = \int_0^{\Gamma_w} K_\delta(z - z_w(\Gamma, t)) d\Gamma, \tag{A2a,b}$$

where K_δ is the vortex blob kernel, with regularization parameter δ :

$$K_\delta(z) = \frac{1}{2\pi i} \frac{\bar{z}}{|z|^2 + \delta^2}, \quad \bar{z} = x - iy. \tag{A3}$$

If z is a point on the bound sheet for which $\delta = 0$, w_b is to be computed in the principal value sense. The position of the bound vortex sheet z_b is determined from the plate’s flapping $(\theta_a(t), \theta_p(t))$ and swimming $x(t)$ motions. The corresponding sheet strength $\gamma(s, t)$ is determined by imposing the no penetration boundary condition on the plate, together with conservation of total circulation. The no penetration boundary condition is given by

$$\text{Re}[wn]_{z_b} = \text{Re}[w_{swimmer}n], \tag{A4}$$

where

$$n = \begin{cases} -\sin \theta_a + i \cos \theta_a, & s \in [0, l], \\ -\sin \theta_p + i \cos \theta_p, & s \in [l, L], \end{cases} \tag{A5}$$

and

$$w_{swimmer} = \begin{cases} \dot{x} - i\dot{y} - i\dot{\theta}_a [\bar{z}_b - (x - iy)], & s \in [0, l], \\ \dot{x} - i\dot{y} - i\dot{\theta}_a [\bar{z}_b - (x - iy)] - i\dot{\alpha} [\bar{z}_b - (x_A - iy_A)], & s \in [l, L]. \end{cases} \tag{A6}$$

Conservation of the fluid circulation implies that $\int_{l_b} \gamma(s, t) ds + \Gamma_w(t) = 0$.

The circulation parameter Γ along the free vortex sheet $z_w(\Gamma, t)$ is determined by the circulation shedding rates $\dot{\Gamma}_w$, according to the Kutta condition, which states that the fluid velocity at the trailing edge is finite and tangent to the flyer (figure 11). The Kutta condition can be obtained from the Euler equations by enforcing that, at the trailing edge, the difference in pressure across the swimmer is zero. To this end, we integrate the balance of momentum equation for inviscid planar flow along a closed contour containing the vortex sheet and trailing edge:

$$[p]_{\mp}(s) = p_{-}(s) - p_{+}(s) = -\frac{d\Gamma(s, t)}{dt} - \frac{1}{2}(u_{-}^2 - u_{+}^2), \tag{A7}$$

where $\Gamma(s, t) = \Gamma_w + \int_0^s \gamma(s', t) ds'$, $0 \leq s \leq L$, is the circulation within the contour and $p_{\mp}(s, t)$ and $u_{\mp}(s, t)$ denote the limiting pressure and tangential slip velocities on both

Antiphase tail flexion enhances swimming performance

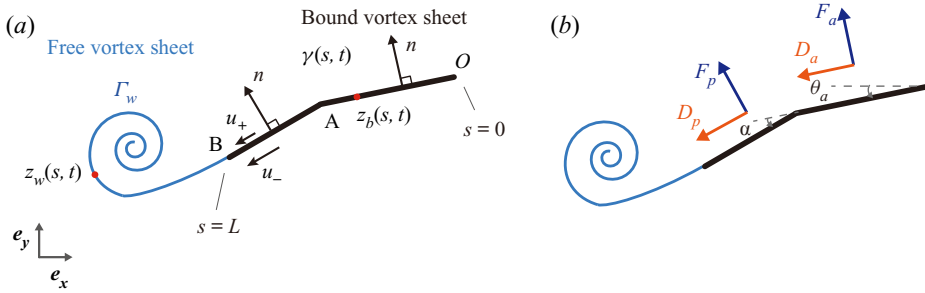


Figure 11. (a) Schematic of the vortex sheet model for a two-dimensional bending swimmer. (b) Depiction of the different hydrodynamic forces acting on the swimmer.

sides of the swimmer. Since the pressure difference across the free sheet is zero, it also vanishes at the trailing edge by continuity, which implies that

$$\dot{\Gamma}_w = -\frac{1}{2}(u_-^2 - u_+^2)|_{s=L}. \tag{A8}$$

The values of u_- and u_+ are obtained from the average tangential velocity component and from the velocity jump at the trailing edge, given by the sheet strength, evaluated at $s = L$:

$$\bar{u} = \frac{u_+ + u_-}{2} = \text{Im}[(w - w_{swimmer})n], \quad u_- - u_+ = \gamma. \tag{A9a,b}$$

Once shed, the vorticity in the free sheet moves with the flow. Thus the parameter Γ assigned to each particle $z_w(\Gamma, t)$ is the value of Γ_w at the instant it is shed from the trailing edge. The evolution of the free vortex sheet z_w is obtained by advecting it in time with the fluid velocity:

$$\dot{z}_w = w_w(z_w, t) + w_b(z_w, t). \tag{A10}$$

Appendix B. Forces and moments

As in figure 12, the hydrodynamic forces F_a and F_p acting on the anterior and posterior parts of the swimmer, respectively, are given by

$$F_a = F_{ax} + iF_{ay} = \int_0^l n[p]_{\mp} ds, \quad F_p = F_{px} + iF_{py} = \int_l^L n[p]_{\mp} ds. \tag{B1a,b}$$

The hydrodynamic moment M_a acting on the anterior part of the swimmer about its leading edge and the hydrodynamic moment M_p acting on the posterior part of the swimmer about the flexion point are given by

$$M_a = \int_0^l [p]_{\mp} s ds, \quad M_p = \int_l^L [p]_{\mp} (s - l) ds. \tag{B2a,b}$$

Note that the components F_{ax}, F_{ay} and F_{px}, F_{py} can be written explicitly as

$$\left. \begin{aligned} F_{ax} &= \int_0^l [p]_{\mp} (-\sin \theta_a) ds, & F_{ay} &= \int_0^l [p]_{\mp} \cos \theta_a ds, \\ F_{px} &= \int_l^L [p]_{\mp} (-\sin \theta_p) ds, & F_{py} &= \int_l^L [p]_{\mp} \cos \theta_p ds, \end{aligned} \right\} \tag{B3}$$

where $\theta_p = \theta_a + \alpha$ and α is the flexion angle.

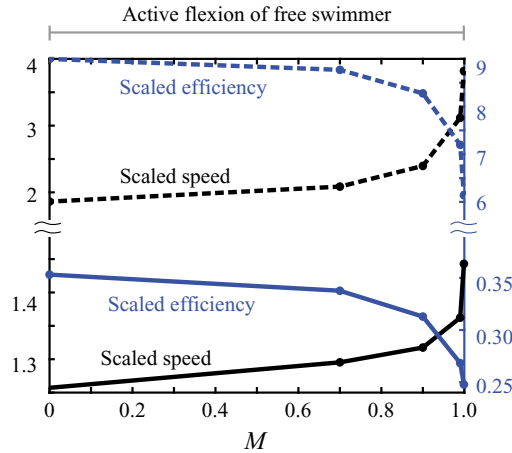


Figure 12. Swimming performance versus tailbeat profile parameter (elliptic modulus M). Swimming speed and efficiency of active flexion scaled by rigid flapping for $a = 15^\circ$, $l/L = 0.7$, $A = 30^\circ$. As M increases, the relative flapping of the posterior end goes from sinusoidal to square-like wave, with faster reversal of flapping direction during the flapping cycle. Dashed lines correspond to anti-phase active flexion and solid lines to in-phase flexion.

The total hydrodynamic force acting on the swimmer due to the pressure difference across the swimmer is given by

$$F = F_x + iF_y, \tag{B4}$$

where the components F_x and F_y are

$$\left. \begin{aligned} F_x = F_{ax} + F_{px} &= \int_0^l [p]_{\mp} (-\sin \theta_a) ds + \int_l^L [p]_{\mp} (-\sin \theta_p) ds, \\ F_y = F_{ay} + F_{py} &= \int_0^l [p]_{\mp} \cos \theta_a ds + \int_l^L [p]_{\mp} \cos \theta_p ds. \end{aligned} \right\} \tag{B5}$$

The total hydrodynamic moment acting on the swimmer about its leading edge is given by

$$M = \int_0^l [p]_{\mp} s ds + \int_l^L [p]_{\mp} (s - l + l \cos \alpha) ds. \tag{B6}$$

We introduce a drag force D that emulates the effect of skin friction due to fluid viscosity. This force is based on the Blasius laminar boundary layer theory as implemented by Fang (2016) in the context of the vortex sheet model. Blasius theory provides an empirical formula for skin friction on one side of a horizontal plate of length L placed in fluid of density ρ_f and uniform velocity U . In dimensional form, the Blasius formula is $D = \frac{1}{2} \rho_f C_f LU^2$, where the skin friction coefficient $C_f = 0.664/\sqrt{Re}$ is given in terms of the Reynolds number $Re = \rho_f UL/\mu$. Substituting back in the empirical formula leads to $D = C_d U^{3/2}$, where $C_d = 0.664\sqrt{\rho_f \mu L}$. Following Fang (2016), we write a modified expression of the drag force for a swimming plate:

$$D = C_d (\bar{U}_+^{3/2} + \bar{U}_-^{3/2}), \tag{B7}$$

where \bar{U}_\pm are the spatially averaged tangential fluid velocities on the upper and lower side of the plate, respectively, relative to the swimming velocity U :

$$\bar{U}_\pm(t) = \frac{1}{L} \int_0^L u_\pm(s, t) ds - U. \tag{B8}$$

Based on the experiments of Ramananarivo *et al.* (2016) with freely swimming flapping foils, they estimated $C_d = 0.04$ which corresponds to Reynolds number $Re \approx 5000$ for a rigid pitching swimmer.

The equation of motion governing the free swimming $x(t)$ is given by Newton’s second law:

$$m\ddot{x} = F_x - D_x, \tag{B9}$$

where D_x is the x component of the drag force D . When the swimmer bends passively, the relative rotation $\alpha(t) = \theta_p - \theta_a$ of the posterior end is not prescribed *a priori* and follows from the physics of fluid–structure interactions. Considering that the rotational joint at the flexion point is equipped with a torsional spring of stiffness κ and damping coefficient c , we write the equation governing the relative rotation of the posterior link as

$$I_p(\ddot{\theta}_a + \ddot{\alpha}) + c\dot{\alpha} + \kappa\alpha = M_p + M_{inertia}, \tag{B10}$$

where I_p is the moment of inertia of the posterior link about the flexion point and $M_{inertia}$ is an inertial moment acting on the posterior link due to the free motion of the flexion point. Namely,

$$M_{inertia} = m_p \text{Im}[-z_A \bar{a}_A], \tag{B11}$$

where $m_p = \rho e(L - l)$ is the mass of the posterior link, $z_A = ((L - l)/2)(\cos \theta_p + i \sin \theta_p)$ is the position of the flexion point relative to the mass centre of the posterior link and \bar{a}_A is the complex conjugate of the acceleration a_A at the flexion point. The latter is given by

$$a_A = (\ddot{x} - l\ddot{\theta}_a \sin \theta_a) - il\dot{\theta}_a^2 \sin \theta_a. \tag{B12}$$

For a swimmer undergoing active flexion, the flapping motion and body bending are produced by two active moments M_a and M_p acting by the swimmer on the fluid about the leading edge O and the hinge A, respectively. The power input by the swimmer to overcome the moment of all the hydrodynamic forces about the leading edge is given by

$$P(t) = \dot{\theta}_a M_a + \dot{\theta}_p (M_p + l|F_p| \cos \alpha). \tag{B13}$$

For a swimmer with passive flexion, the input power is given by

$$P(t) = \dot{\theta}_a (M_a + l|F_p| \cos \alpha - \kappa\alpha - c\dot{\alpha}). \tag{B14}$$

Note that the skin drag does not contribute to input power.

Appendix C. Numerical implementation

The bound vortex sheet is discretized by $2n + 1$ point vortices at $z_b(t)$ with strength $\Delta\Gamma = \gamma \Delta s$. These vortices are located at Chebyshev points that cluster at the two ends of the swimmer. Their strength is determined by enforcing no penetration at the midpoints between the vortices, together with conservation of circulation. The free vortex sheet is discretized by regularized point vortices at $z_w(t)$, which is released from the trailing edge at each time step with circulation given by (A8). The free point vortices move with the

discretized fluid velocity while the bound vortices move with the swimmer's velocity. For the actively bending swimmer, the discretization of (B9) and (A8), (A10) yields a coupled system of ordinary differential evolution equations for the swimmer's position, the shed circulation and the free vorticity, which is integrated in time using the fourth-order Runge–Kutta scheme. For the passively bending swimmer, the discretization of (B10) is added to the coupled system of equations to simultaneously solve for the rotational motion of the posterior link relative to the anterior link. The details of the shedding algorithm are given in Nitsche & Krasny (1994). The numerical values of the time step Δt , the number of bound vortices n and the regularization parameter δ are chosen so that the solution changes little under further refinement.

Finally, to emulate the effect of viscosity, we allow the shed vortex sheets to decay gradually by dissipating each incremental point vortex after a finite time T_{diss} ($T_{diss} = 1.625T$ for the swimmer with active flexion and $T_{diss} = \sqrt{2.09}T$ for passive flexion) from the time it is shed into the fluid. Larger T_{diss} implies that the vortices stay in the fluid for longer times, mimicking the effect of lower fluid viscosity. We refer the reader to Huang *et al.* (2018) for a detailed analysis of the effect of dissipation time on the hydrodynamic forces on a stationary and moving plate in the vortex sheet model. Details of the numerical validation in comparison with Jones (2003) and Jones & Shelley (2005) are provided in Huang *et al.* (2016).

REFERENCES

- ALBEN, S. 2008 Optimal flexibility of a flapping appendage in an inviscid fluid. *J. Fluid Mech.* **614**, 355–380.
- ALBEN, S. 2009 Wake-mediated synchronization and drafting in coupled flags. *J. Fluid Mech.* **641**, 489–496.
- BEAL, D.N., HOVER, F.S., TRIANTAFYLLOU, M.S., LIAO, J.C. & LAUDER, G.V. 2006 Passive propulsion in vortex wakes. *J. Fluid Mech.* **549**, 385–402.
- BECKER, A.D., MASOUD, H., NEWBOLT, J.W., SHELLEY, M. & RISTROPH, L. 2015 Hydrodynamic schooling of flapping swimmers. *Nat. Commun.* **6** (1), 8514.
- BLONDEAUX, P., FORNARELLI, F., GUGLIELMINI, L., TRIANTAFYLLOU, M.S. & VERZICCO, R. 2005 Numerical experiments on flapping foils mimicking fish-like locomotion. *Phys. Fluids* **17** (11), 113601.
- BOZKURTAS, M., DONG, H., MITTAL, R., TANGORRA, J., HUNTER, I., LAUDER, G. & MADDEN, P. 2007 CFD-based analysis and design of biomimetic flexible propulsor for autonomous underwater vehicles. In *37th AIAA Fluid Dynamics Conference and Exhibit, AIAA Paper 2007-4213*.
- COMBES, S.A. & DANIEL, T.L. 2001 Shape, flapping and flexion: wing and fin design for forward flight. *J. Exp. Biol.* **204** (12), 2073–2085.
- COMBES, S.A. & DANIEL, T.L. 2003 Flexural stiffness in insect wings. II. Spatial distribution and dynamic wing bending. *J. Exp. Biol.* **206** (17), 2989–2997.
- DAI, H., LUO, H., DE SOUSA, P.J.S.A.F. & DOYLE, J.F. 2012 Thrust performance of a flexible low-aspect-ratio pitching plate. *Phys. Fluids* **24** (10), 101903.
- DICKINSON, M.H., LEHMANN, F.-O. & SANE, S.P. 1999 Wing rotation and the aerodynamic basis of insect flight. *Science* **284** (5422), 1954–1960.
- DRUCKER, E.G. & LAUDER, G.V. 2000 A hydrodynamic analysis of fish swimming speed: wake structure and locomotor force in slow and fast labriform swimmers. *J. Exp. Biol.* **203** (16), 2379–2393.
- ELDREDGE, J.D. 2006 Numerical simulations of undulatory swimming at moderate Reynolds number. *Bioinspir. Biomim.* **1** (4), S19.
- ELDREDGE, J.D. 2008 Dynamically coupled fluid–body interactions in vorticity-based numerical simulations. *J. Comput. Phys.* **227** (21), 9170–9194.
- ELDREDGE, J.D., TOOMEY, J. & MEDINA, A. 2010 On the roles of chord-wise flexibility in a flapping wing with hovering kinematics. *J. Fluid Mech.* **659**, 94–115.
- ELLINGTON, C.P. 1984 The aerodynamics of hovering insect flight. VI. Lift and power requirements. *Phil. Trans. R. Soc. Lond. B* **305** (1122), 145–181.
- ELLINGTON, C.P. 1985 Power and efficiency of insect flight muscle. *J. Exp. Biol.* **115** (1), 293–304.
- ELOY, C. 2013 On the best design for undulatory swimming. *J. Fluid Mech.* **717** (25), 48–49.
- FANG, F. 2016 Hydrodynamic interactions between self-propelled flapping wings. PhD thesis, New York University.

Antiphase tail flexion enhances swimming performance

- FEILICH, K.L. & LAUDER, G.V. 2015 Passive mechanical models of fish caudal fins: effects of shape and stiffness on self-propulsion. *Bioinspir. Biomim.* **10** (3), 036002.
- FUJIWARA, S. & YAMAGUCHI, S. 2017 Development of fishlike robot that imitates carangiform and subcarangiform swimming motions. *J. Aero Aqua Bio-Mech.* **6** (1), 1–8.
- GEMMELL, B.J., FOGERSON, S.M., COSTELLO, J.H., MORGAN, J.R., DABIRI, J.O. & COLIN, S.P. 2016 How the bending kinematics of swimming lampreys build negative pressure fields for suction thrust. *J. Expl Biol.* **219** (24), 3884–3895.
- GIBOUIN, F., RAUFASTE, C., BOURET, Y. & ARGENTINA, M. 2018 Study of the thrust–drag balance with a swimming robotic fish. *Phys. Fluids* **30** (9), 091901.
- HEATHCOTE, S. & GURSUL, I. 2007 Flexible flapping airfoil propulsion at low Reynolds numbers. *AIAA J.* **45** (5), 1066–1079.
- HEATHCOTE, S., WANG, Z. & GURSUL, I. 2008 Effect of spanwise flexibility on flapping wing propulsion. *J. Fluids Struct.* **24** (2), 183–199.
- HEYDARI, S. & KANSO, E. 2021 School cohesion, speed and efficiency are modulated by the swimmers flapping motion. *J. Fluid Mech.* **922**, A27.
- HOOVER, A.P., CORTEZ, R., TYTELL, E.D. & FAUCI, L.J. 2018 Swimming performance, resonance and shape evolution in heaving flexible panels. *J. Fluid Mech.* **847**, 386–416.
- HOOVER, A.P. & TYTELL, E. 2020 Decoding the relationships between body shape, tail beat frequency, and stability for swimming fish. *Fluids* **5** (4), 215.
- HUA, R.-N., ZHU, L. & LU, X.-Y. 2013 Locomotion of a flapping flexible plate. *Phys. Fluids* **25** (12), 121901.
- HUANG, Y., NITSCHKE, M. & KANSO, E. 2015 Stability versus maneuverability in hovering flight. *Phys. Fluids* **27** (6), 061706.
- HUANG, Y., NITSCHKE, M. & KANSO, E. 2016 Hovering in oscillatory flows. *J. Fluid Mech.* **804**, 531–549.
- HUANG, Y., RISTROPH, L., LUHAR, M. & KANSO, E. 2018 Bistability in the rotational motion of rigid and flexible flyers. *J. Fluid Mech.* **849**, 1043–1067.
- ITOH, T., TSUJI, S. & NITTA, A. 2003 Migration patterns of young Pacific bluefin tuna (*Thunnus orientalis*) determined with archival tags. *Fish. Bull.* **101** (3), 514–534.
- JONES, M.A. 2003 The separated flow of an inviscid fluid around a moving flat plate. *J. Fluid Mech.* **496** (496), 405–441.
- JONES, M.A. & SHELLEY, M.J. 2005 Falling cards. *J. Fluid Mech.* **540**, 393–425.
- KATZSCHMANN, R.K., DELPRETO, J., MACCURDY, R. & RUS, D. 2018 Exploration of underwater life with an acoustically controlled soft robotic fish. *Sci. Robot.* **3** (16), eaar3449.
- KERN, S. & KOUMOUTSAKOS, P. 2006 Simulations of optimized anguilliform swimming. *J. Expl Biol.* **209** (24), 4841–4857.
- LABASSE, J., EHRENSTEIN, U. & MELIGA, P. 2020 Numerical exploration of the pitching plate parameter space with application to thrust scaling. *Appl. Ocean Res.* **101**, 102278.
- LAUDER, G.V. 2015 Fish locomotion: recent advances and new directions. *Annu. Rev. Mar. Sci.* **7** (1), 521–545.
- LAUDER, G.V., LIM, J., SHELTON, R., WITT, C., ANDERSON, E. & TANGORRA, J.L. 2011 Robotic models for studying undulatory locomotion in fishes. *Mar. Technol. Soc. J.* **45** (4), 41–55.
- LI, C., DONG, H. & LIU, G. 2015 Effects of a dynamic trailing-edge flap on the aerodynamic performance and flow structures in hovering flight. *J. Fluids Struct.* **58**, 49–65.
- LI, L., NAGY, M., GRAVING, J.M., BAK-COLEMAN, J., XIE, G. & COUZIN, I.D. 2020 Vortex phase matching as a strategy for schooling in robots and in fish. *Nat. Commun.* **11** (1), 5408.
- LIAO, J.C., BEAL, D.N., LAUDER, G.V. & TRIANTAFYLLOU, M.S. 2003 Fish exploiting vortices decrease muscle activity. *Science* **302** (5650), 1566–1569.
- LIU, P. & BOSE, N. 1997 Propulsive performance from oscillating propulsors with spanwise flexibility. *Proc. R. Soc. Lond. A* **453** (1963), 1763–1770.
- LOW, K.H. & CHONG, C.W. 2010 Parametric study of the swimming performance of a fish robot propelled by a flexible caudal fin. *Bioinspir. Biomim.* **5** (4), 046002.
- LUCAS, K.N., JOHNSON, N., BEAULIEU, W.T., CATHCART, E., TIRRELL, G., COLIN, S.P., GEMMELL, B.J., DABIRI, J.O. & COSTELLO, J.H. 2014 Bending rules for animal propulsion. *Nat. Commun.* **5** (1), 3293.
- LUCAS, K.N., THORNYCROFT, P.J.M., GEMMELL, B.J., COLIN, S.P., COSTELLO, J.H. & LAUDER, G.V. 2015 Effects of non-uniform stiffness on the swimming performance of a passively-flexing, fish-like foil model. *Bioinspir. Biomim.* **10** (5), 056019.
- MENON, K. & MITTAL, R. 2019 Flow physics and dynamics of flow-induced pitch oscillations of an airfoil. *J. Fluid Mech.* **877**, 582–613.

- MIAO, J.-M. & HO, M.-H. 2006 Effect of flexure on aerodynamic propulsive efficiency of flapping flexible airfoil. *J. Fluids Struct.* **22** (3), 401–419.
- MICHELIN, S. & LLEWELLYN SMITH, S.G. 2009 Resonance and propulsion performance of a heaving flexible wing. *Phys. Fluids* **21** (7), 071902.
- MOORED, K.W. & QUINN, D.B. 2019 Inviscid scaling laws of a self-propelled pitching airfoil. *AIAA J.* **57** (9), 3686–3700.
- MORÉ, J.J. & SORENSEN, D.C. 1983 Computing a trust region step. *SIAM J. Sci. Stat. Comput.* **4** (3), 553–572.
- MÜLLER, U.K., STAMUIS, E.J. & VIDELER, J.J. 2002 Riding the waves: the role of the body wave in undulatory fish swimming. *Integr. Compar. Biol.* **42**, 981–987.
- MÜLLER, U.K., VAN DEN HEUVEL, B.L.E., STAMHUIS, E.J. & VIDELER, J.J. 1997 Fish foot prints: morphology and energetics of the wake behind a continuously swimming mullet (*Chelon labrosus* Risso). *J. Expl Biol.* **200** (22), 2893–2906.
- NITSCHKE, M. & KRASNY, R. 1994 A numerical study of vortex ring formation at the edge of a circular tube. *J. Fluid Mech.* **276**, 139–161.
- POLLARD, B. & TALLAPRAGADA, P. 2019 Passive appendages improve the maneuverability of fishlike robots. *IEEE ASME Trans. Mechatron.* **24** (4), 1586–1596.
- QUINN, D.B., LAUDER, G.V. & SMITS, A.J. 2015 Maximizing the efficiency of a flexible propulsor using experimental optimization. *J. Fluid Mech.* **767**, 430–448.
- RAMAKRISHNAN, S., BOZKURTAS, M., MITTAL, R. & LAUDER, G.V. 2011 Thrust production in highly flexible pectoral fins: a computational dissection. *Mar. Tech. Soc. J.* **45** (4), 56–64.
- RAMANANARIVO, S., FANG, F., OZA, A., ZHANG, J. & RISTROPH, L. 2016 Flow interactions lead to orderly formations of flapping wings in forward flight. *Phys. Rev. Fluids* **1** (7), 071201.
- RYU, J., PARK, S.G., HUANG, W.-X. & SUNG, H.J. 2019 Hydrodynamics of a three-dimensional self-propelled flexible plate. *Phys. Fluids* **31** (2), 021902.
- SFAKIOTAKIS, M., LANE, D.M. & DAVIES, J.B.C. 1999 Review of fish swimming modes for aquatic locomotion. *IEEE J. Ocean. Engng* **24** (2), 237–252.
- SHADWICK, R.E. & GEMBALLA, S. 2005 Structure, kinematics, and muscle dynamics in undulatory swimming. *Fish Physiol.* **23**, 241–280.
- SHELLEY, M.J. & ZHANG, J. 2011 Flapping and bending bodies interacting with fluid flows. *Annu. Rev. Fluid Mech.* **43**, 449–465.
- SHENG, J.X., YSASI, A., KOLOMENSKIY, D., KANSO, E., NITSCHKE, M. & SCHNEIDER, K. 2012 Simulating vortex wakes of flapping plates, In *Natural Locomotion in Fluids and on Surfaces* (ed. S. Childress, A. Hosoi, W. Schultz & J. Wang), pp. 255–262. Springer.
- SHOELE, K. & MITTAL, R. 2016 Flutter instability of a thin flexible plate in a channel. *J. Fluid Mech.* **786**, 29–46.
- SIMPENDORFER, C. 2009 Galeocerdo cuvier. in iucn 2013. iucn red list of threatened species, version 2013.1. Website, <http://www.iucnredlist.org/>.
- SIMPSON, S.D., HARRISON, H.B., CLAEREBOUDT, M.R. & PLANES, S. 2014 Long-distance dispersal via ocean currents connects omani clownfish populations throughout entire species range. *PLoS One* **9** (9), e107610.
- SMITS, A.J. 2019 Undulatory and oscillatory swimming. *J. Fluid Mech.* **874**, P1.
- TANEDA, S. 1965 Experimental investigation of vortex streets. *J. Phys. Soc. Japan* **20** (9), 1714–1721.
- TANGORRA, J.L., LAUDER, G.V., HUNTER, I.W., MITTAL, R., MADDEN, P.G.A. & BOZKURTAS, M. 2010 The effect of fin ray flexural rigidity on the propulsive forces generated by a biorobotic fish pectoral fin. *J. Expl Biol.* **213** (23), 4043–4054.
- TRIANTAFYLLOU, G.S. & TRIANTAFYLLOU, M.S. 1993 Reverse kármán streets in the wake of flapping foils for application in fish propulsion. In *Bluff-Body Wakes, Dynamics and Instabilities* (ed. H. Eckelmann, J.M.R. Graham, P. Huerre & P.A. Monkewitz), pp. 11–14. IUTAM, Springer.
- TYTELL, E.D., HSU, C.-Y., WILLIAMS, T.L., COHEN, A.H. & FAUCI, L.J. 2010 Interactions between internal forces, body stiffness, and fluid environment in a neuromechanical model of lamprey swimming. *Proc. Natl Acad. Sci. USA* **107** (46), 19832–19837.
- TYTELL, E.D. & LAUDER, G.V. 2004 The hydrodynamics of eel swimming: I. Wake structure. *J. Expl Biol.* **207** (11), 1825–1841.
- TYTELL, E.D., LEFTWICH, M.C., HSU, C.-Y., GRIFFITH, B.E., COHEN, A.H., SMITS, A.J., HAMLET, C. & FAUCI, L.J. 2016 Role of body stiffness in undulatory swimming: insights from robotic and computational models. *Phys. Rev. Fluids* **1** (7), 073202.
- VINCENT, L., LIU, Y. & KANSO, E. 2020a Shape optimization of tumbling wings. *J. Fluid Mech.* **889**, A9.

Antiphase tail flexion enhances swimming performance

- VINCENT, L., ZHENG, M., COSTELLO, J.H. & KANSO, E. 2020*b* Enhanced flight performance in non-uniformly flexible wings. *J. R. Soc. Interface* **17** (168), 20200352.
- WAN, H., DONG, H. & HUANG, G.P. 2012 Hovering hinge-connected flapping plate with passive deflection. *AIAA J.* **50** (9), 2020–2027.
- WANG, L. 2020 Locomotion of a self-propulsive pitching plate in a quiescent viscous fluid. *Proc. Inst. Mech. Engrs C* 0954406220903338.
- WEN, L. & LAUDER, G. 2013 Understanding undulatory locomotion in fishes using an inertia-compensated flapping foil robotic device. *Bioinspir. Biomim.* **8** (4), 046013.
- WHITE, C.H., LAUDER, G.V. & BART-SMITH, H. 2021 Tunabot flex: a tuna-inspired robot with body flexibility improves high-performance swimming. *Bioinspir. Biomim.* **16** (2), 026019.
- WU, T.Y.-T. 1971 Hydromechanics of swimming propulsion. Part 1. Swimming of a two-dimensional flexible plate at variable forward speeds in an inviscid fluid. *J. Fluid Mech.* **46** (2), 337–355.
- YABUTA, S. 1997 Spawning migrations in the monogamous butterflyfish, *Chaetodon trifasciatus*. *Ichthyol. Res.* **44** (2–3), 177–182.
- ZHU, J., WHITE, C., WAINWRIGHT, D.K., DI SANTO, V., LAUDER, G.V. & BART-SMITH, H. 2019 Tuna robotics: a high-frequency experimental platform exploring the performance space of swimming fishes. *Sci. Robot.* **4** (34), eaax4615.