Synchrony and symmetry-breaking in active flagellar coordination

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Living creatures exhibit a remarkable diversity of locomotion mechanisms, evolving structures specialized for interacting with their environment. In the vast majority of cases, locomotor behaviours such as flying, crawling and running are orchestrated by nervous systems. Surprisingly, microorganisms can enact analogous movement gaits for swimming using multiple, fast-moving cellular protrusions called cilia and flagella. Here, I demonstrate intermittency, reversible rhythmogenesis and gait mechanosensitivity in algal flagella, to reveal the active nature of locomotor patterning. In addition to maintaining free-swimming gaits, I show that the algal flagellar apparatus functions as a central pattern generator that encodes the beating of each flagellum in a network in a distinguishable manner. The latter provides a novel symmetry-breaking mechanism for cell reorientation. These findings imply that the capacity to generate and coordinate complex locomotor patterns does not require neural circuitry but rather the minimal ingredients are present in simple unicellular organisms.

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1. The locomotor gaits of organisms

Since the invention of photography, the natural habits of organisms have come under increasing scrutiny. Modern optical technologies have enabled resolution of ever-finer detail, so that we can visualize and track behaviour across scales [1,2]. Consider for example the gallop of a horse: how are its four feet coordinated? This fast action cannot be resolved by the human eye, so the question remained unanswered until 1878, when (reputedly to settle a bet) the photographer Eadweard Muybridge successfully imaged the gait sequences of a horse [3]. Control of rhythmic limb movements in both vertebrates and invertebrates is determined by neuronal networks termed central pattern generators (CPG), where motor feedback is not required [4]. Indeed, CPGs in in vitro preparations can produce activity patterns that are identifiable with the in vivo behaviour. For instance, in the pteropod Clione limacine (sea butterfly), which swims with two dorsal wings, the neuronal pattern for basic synchronous movement (fictive swimming) can be reproduced in isolated pedal ganglia [5]. A striking example of the genetic basis of gait-control and CPGs pertains to the Icelandic horse, where the ability to pace (legs on the same side of the body moving synchronously) was shown to be associated with a point mutation [6].

At the microscale, organisms interact with their world following very different physical principles. Here, inertia is negligible and many species developed motile appendages that are slender and suited to drag-based propulsion through fluids. From these simplest designs, myriad locomotion strategies have evolved. The model microswimmer Escherichia coli is peritrichously flagellated—bundles of rigid (prokaryotic) flagella rotate synchronously in the same sense to elicit forward swimming (a ‘run’), but lose synchrony and unbundle (a ‘tumble’) when one or more of the motors reverse direction (see [7] and references therein).

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The latter results in rapid changes in reorientation. In this way, a bacterium controls its propensity for reorientation to bias its swimming towards or away from chemo-attractants or repellents [8]. It has been shown that bundling is largely owing to hydrodynamic interactions between the rotating filaments [9], but unbundling is stochastic. By contrast, eukaryotes exhibit more deterministic responses to vectorial cues [10]. Phototactic reorientation by the biflagellate alga *Chlamydomonas reinhardtii* requires bilateral symmetry-breaking in a pair of apparently identical flagella [11], though differences in signal transduction must be present [12–14]. Eukaryotic flagella and cilia possess a flexible and distributed molecular architecture allowing for many more degrees of freedom than prokaryotic flagella. This versatility, particularly when multiple appendages are attached to the same cell, allows algal flagellates to orchestrate diverse swimming gaits such as the breaststroke, trot or gallop [15].

Extensive research has already shown that distinct waveforms and beat frequencies can be produced by the same ciliary structure [16–18], depending on intrinsic and extrinsic forcing [19–21]. This ability to modulate dynein motor cooperativity to produce distinct beating modes on a single cilium does not however explain the higher-level coordination over a network of such oscillating structures. To achieve the latter, intracellular control mechanisms are implicated (reviewed in [22]). In this article, I present new findings on gait patterning in microalgae, which reveal that single-celled multiflagellates can actively dictate the dynamics and activity of each individual flagellum. I further propose that symmetry-breaking processes in the flagellar apparatus are causal to this distinguishability between flagella. The attainment of control specificity of locomotor appendages may be a key innovation in the evolution toward increasingly deterministic movement in eukaryotes.

2. Gait species-speciality

We focus on unicellular algae bearing 2, 4, 8 or 16 flagella and study the patterns of flagellar actuation and movement. Representative gaits are compared to animals with analogous limb positioning (figure 1), for which the neuronal basis of movement control has been clearly demonstrated (except in the sea star, which is yet to be subject to systematic investigation). In all cases, the movement operates in a very different physical and size regime [23,24] from that encountered by the microeukaryotes from this study. Species-speciality of locomotor behaviours also applies to flagellates: organisms with an identical number and arrangement of flagella can still assume different locomotor gaits [15]. Concurrently, a single species can be capable of multiple gaits associated with significant changes in the flagellar beat pattern. For instance, sudden environmental perturbations elicit so-called shock responses in multiple species, in which flagella reverse their beating direction (but not the direction of wave propagation) to produce transient backward movement or even cessation of swimming [25,26]. Some species have a stop state [27], in which all flagella become reversibly quiescent. We organize this section by characterizing first the gaits used for forward swimming, and then separately, introducing the phenomenon of selective beat activation in free-swimming cells.

(a) Run gaits for forward swimming

The biflagellate *C. reinhardtii* is favoured for biophysical models of swimmers [28]. Cells execute a forward breaststroke gait in which the two flagella are synchronized, interrupted by phase slips. Stochastic switching between this characteristic in-phase breaststroke and a perfectly antiphase (or freestyle) gait was also discovered in a phototaxis mutant [29]. Experimental and theoretical evidence overwhelmingly implicates intracellular coupling as the dominant mechanism for flagellar coupling [15,30–32]. However, phase synchrony is not a given: heterotrophic biflagellates such as *Polytoma uvella* have two flagella of slightly unequal lengths and different beat frequencies, which remain largely asynchronous. With more than two flagella, new gaits become accessible. Depending on flagella...
configuration and ultrastructure, different interflagellar phase patterns are sustained during steady, forward swimming. Comparison with quadrupedal locomotion is instructive here [33]. Among species with four flagella, a robust trotting gait comprising two alternating pairs of synchronous breaststrokes is assumed by two marine Pyramimonas species: P. parkeae and P. tychotreta, whereas galloping gaits are consistently displayed by Tetraselmis sp. and Carteria crucifera (see electronic supplementary material, videos S1–S5). Only three extant species are known to have eight flagella [34]. Of these, Pyramimonas octopus coordinates its eight flagella into a unique forward propulsion gait that we term the rotary breaststroke (see §4). Identifying equivalence between this and octopod animal movement is more challenging. The spider possesses obvious bilateral symmetry not present in the algae, though alternating patterns of leg movements are observed [35]. Jellyfish ephyrae use cycles of contraction and relaxation involving the simultaneous movement of their eight soft appendages, but move in an inertial regime [36]. Finally, the Arctic species Pyramimonas cyrtoptera has 16 flagella [37] that can be actuated metachronously rather than in discrete patterns. This may be the critical number at which the flagella begin to interact hydrodynamically [15]. In this limiting scenario, the wave-like coordination of many closely separated appendages may help to avoid collisions and mirrors the locomotion mechanisms adopted by arthropods [38,39].

(b) Gait-switching and partial activation

In animals, the capacity to produce multiple locomotor modes is critical for survival. Gait often depends on the speed of the desired movement. For example, the ghost crab (Ocypode ceratophthalmus) becomes bipedal at the highest speeds [40]. Likewise, we find that unicellular flagellates dynamically reconfigure their motor apparatus to produce different gaits that are coupled to photo- or mecano-sensory pathways. Next, we compare the motion repertoire of different multiflagellate species. Freely swimming cells were imaged at kilohertz and sub-micrometre resolution, so that flagella motion could be discerned at the same time as the cell body movement. Strong behavioural stereotypy was observed within each species, but heterogeneity across species (figure 2). Consistently, tracks presenting high curvature loops were found to result from a previously uncharacterized motility mechanism—namely, the partial activation of flagella. For the same cell, beating can be restricted to a subset of flagella with the remaining flagella quiescent. Asymmetries arising from the actively beating flagellar subgroup produce large-angle turns in the free-swimming trajectory. A previously quiescent flagellum can subsequently become active again, and vice versa (figure 2b and electronic supplementary material, video S1). Four different quadriflagellate species showed selective activation of flagella (P. parkeae, Tetraselmis suecica, T. subcordiformis and P. tychotreta). In cells exhibiting motion intermittency such as P. parkeae (figure 2d), flagella alternate between periods of bursting activity and...
total quiescence (figure 2g; electronic supplementary material), analogous to the physiological behaviour of motor neurons.

Next, we resolve gait transients and interflagellar phase dynamics while keeping track of flagellum identity. Automated methods could not follow the flagellum movement reliably. Owing to the three-dimensional movement and continuous cell axial rotation, features of interest do not always remain in focus. Instead, flagella tip positions were manually annotated in TrackMate [41]. The distance between flagellum tip and cell centroid undergoes oscillations for an actively beating flagellum but remains constant for a quiescent flagellum. We use this distance as proxy for flagellar phase (figure 3). The quadriflagellate C. crucifera displays stochastic gait-switching between forward swimming and transient shocks, as does P. octopus [27]. Shocks involve simultaneous conversion of all four flagella to a hyperactivated undulatory state (figure 3a). After approximately 0.3 s, the canonical ciliary beat pattern is recovered. A rhythmic galloping gait resumes after only a few asynchronous beat cycles. The rapidity of phase-resetting responses in these organisms further implies that gait-control is active [42]. In figure 3b, we demonstrate partial activation in P. tychotreta [43]. Here, full beat cycles are sustained in flagella f1,3, while beating in the diametrically opposite pair f2,4 is suppressed throughout. Small fluctuations detected in the inactive flagella are owing to passive movement induced by the beating of nearby flagella. Further examples of quadriflagellates in which beating is dynamically constrained to one, two or three out of a total possible four flagella are provided in the electronic supplementary material.

3. Intermittency and temporal ordering in a flagellar network

Free-swimming individuals are not amenable to long-time imaging. In order to investigate the long-time statistics of locomotor patterning, experiments were also conducted on micropipette-held (body-fixed) organisms. Single or double micropipette configurations were used to position, manipulate single cells, and to apply localized mechanical perturbations to single flagella. Not all species were suitable for the micropipette technique; however in some cases, captured cells prematurely shed their flagella. This is likely related to a stress-induced, calcium-dependent deflagellation response [44,45]. We identified two quadriflagellate species that were minimally affected by pipette-capture. These are P. parkeae and T. suecica, which are representative of the two known quadriflagellar arrangements: type I and II in [15] (see also figure 4). In P. parkeae, the flagella form a cruciate arrangement, while in T. suecica they align into two anti-parallel pairs. Both flagellates show partial activation and beat intermittency.

For pipette-fixed individuals, flagellar phases could be extracted automatically from imaging data to characterize the emergence and decay of coordination in the flagellar network. Phase patterns are reminiscent of classical studies of footfall patterns and limb locomotion in animals [46,47]. In long-time, high-speed recordings, reversible transitions in locomotor patterning were found in both species. These occurred spontaneously, but could also be induced by mechanical forcing. This touch-dependent gait-sensitivity is a novel manifestation of a mechanoresponse in motile cilia and flagella. At the moment of capture, contact with the pipette can induce rhythmogenesis in a cell with initially quiescent flagella (figure 4a; electronic supplementary material, video S6). Similarly, changes in locomotor patterning were induced when fluid was injected manually (approx. 1 s pulses) via a second pipette in the vicinity of a quiescent cell (figure 4b). The first pulse administered at time t1 induced beating in f1,3 only. This biflagellate state continued until the second pulse (time t2), which then resulted in beating in all four flagella (electronic supplementary material, video S7). In both cases, even after cessation of external perturbations, the cells

Figure 3. Quadriflagellate gaits resolved at the single-flagellum level in free-swimming cells. Displacement from tip to centre of the cell body was used as proxy for flagellar phase for each flagellum, labelled (f1–4). Frames corresponding to specific timepoints are highlighted. Scalebars for cell size reference. (a) Carteria crucifera has a galloping gait during forward runs, but exhibits noise-induced shocks involving reorientation of the flagella (time 3). Recovery of the original rhythm (phase-resetting), completed within four beat cycles. (b) Pyraminonas tychotreta has a canonical trotting gait during forward runs but exhibits partial activation of flagella. Beating is restricted to the pair f1,3, but suppressed in f2,4. Synchrony of f1,3 drifts from antiphase (times 3,4) to in-phase (time 5) over the course of this recording (electronic supplementary material, videos S4 and S5).
Figure 4. Reversible rhythmogenesis and gait mechanosensitivity in quadriflagellates. (a) The four flagella of *P. parvus* have a cruciate arrangement (type I, top view). The captured cell has two flagella in the focal plane (f1,3) and two others moving transversely—these are labelled as f2 and are tracked as one (see electronic supplementary material, video S6). (b) The four flagella of *T. suecica* have an anti-parallel arrangement (type II, top view). Mechanical perturbations are introduced at times t1,2, and spontaneous shocks are detected at times indicated by red pointers. (c–f) are zoomed-in plots of (a,b), at time points of interest. At the moment of capture, the flagella begin to beat, but not simultaneously; a regular trot gait emerges within three beat cycles (c). Spontaneous termination of beating does not occur at the same time in all flagella (d). Induced changes in locomotor patterning (e) are compared with the regular phase patterning (f) observed during the trotting gait. (See electronic supplementary material, video S7.) Inset: flagellar phase is defined according to progression through the beat cycle.

4. Symmetry-breaking in the flagellar apparatus

In this section, we elaborate on the non-identity of flagella in multiflagellate algae. Adapting terminology from gait analyses of limbed animal locomotion, we distinguish between symmetric gaits (e.g. the trot, prong) and asymmetric gaits (e.g. the gallop, bound). A quadrupedal gait is referred to as being symmetric if footfall patterns of both pairs of feet are evenly spaced in time and as asymmetric if the activity of at least one pair is unevenly spaced in time. Different gaits can therefore arise from the same underlying locomotor circuit. The presence of more than two flagella also allows for diverse possibilities for symmetry-breaking even during regular gait patterning (not transitional gaits).

We focus now on the rotary breaststroke of *P. octopus*, which has not been described in any other organism. Gait ordering in multiflagellates is particularly difficult to visualize, owing to the highly three-dimensional swimming. Additionally, *P. octopus* rotates slowly clockwise about its own axis when viewed from the anterior end (electronic supplementary material, video S8). All eight flagella are distinguishable simultaneously only when the cell is swimming into or out of the focal plane (figure 5; electronic supplementary material, video S9). Recordings of such transients were made, from which the extremal positions of each flagellum could be followed in time and tracked manually in ImageJ [41]. As before, a normalized flagellar phase was computed for each flagellum. We find that phase ordering propagates directionally, in the same sense as the about-axis rotation (figure 5c). (Gaits are similarly directional in the related *Pyramimonas cryoptera*, which swims with 16 flagella.) A pairwise Pearson-correlation matrix was computed for each tracked cell to quantify the degree of correspondence between the eight flagella (figure 5d). A strong positive correlation was measured between diametrically opposite flagella (dashed white lines are guides), which tend to move synchronously to produce a noisy sequence of intercalated breaststrokes (first 1 : 5, then 2 : 6 and so on).

Given the directional nature of this swimming gait, asymmetries in developmental patterning must be present. This is corroborated by the only ultrastructural study available on this species [49]. The flagella emerge from separate basal bodies (centrioles during cell division), which are localized to the anterior of the organism (figure 5a). Electron microscopy...
sections reveal a complex basal architecture lacking in rotational symmetry. Individual basal bodies are positioned uniquely according to generational age [49]. A contractile network of thick and thin fibres (some are striated) connect specific numbered microtubule triplets to provide intracellular coupling between flagella [15]. No obvious structural or morphological differences have been reported in the flagellar axonemes. Consequently, it is not possible from light microscopy data to prescribe the identity of the individual flagella in accordance with the numbering system devised from TEM images. Flagella numbering is thus distinguishable only up to circular permutation. Nonetheless, robust in-phase synchrony during free-swimming pertains to diametrically opposite flagella (positive phase correlation, figure 5d). This is in good agreement with the presence of direct physical coupling between distinguished pairs of flagella. Particularly, the principal pair b1 and b5 is connected by a thick striated structure known as the synistosome, which is functionally related to the distal fibre in C. reinhardtii [50]. The lack of correlation between certain basal body pairs (figure 4d) will be investigated further in future studies to determine if this is related to the detailed topology of the fibre network (e.g. b2 and 8 appear unconnected).

Finally, we find that in approximately 5% of cases, free-swimming cells exhibit a distinguished flagellum that is held extended in front of the cell body (electronic supplementary material, video S10). Figure 5 compares this ‘search gait’ with the canonical forward swimming gait of this organism. The beat pattern in this single flagellum is undulatory (sperm-like), in contrast to the lateral (ciliary) stroke used by the other seven flagella. These observations further highlight the capacity for producing heterodynamic behaviour in a network of outwardly identical flagella.

5. Discussion and outlook

The present article is devoted to the gait dynamics of free-living unicellular algal flagellates, advancing the state-of-the-art from single-cell to single-flagellum resolution. Performing a comparative study, I showed that gait coordination in algal flagellates is not only intracellular but also active. The same network of flagella can undergo dynamic reconfiguration over fast timescales to generate different locomotor patterns. Multiple species share an ability to reversibly activate beating in a subset of flagella, while suppressing the activity of the remainder. This is an extreme instantiation of symmetry-breaking in the algal flagella apparatus. Individually, a eukaryotic flagellum or cilium can be in an oscillatory or non-oscillatory state, while beating can be modulated by extrinsic factors including mechanosensitivity [27,51,52], but how can drastically distinct beat patterns be produced in different flagella of the same cell? We suggest two sources of this apparent control specificity in coupled flagella: (i) physical asymmetries in the basal apparatus and (ii) differences in signal transduction. The two contributions are not mutually exclusive.

In some species, physical asymmetries are obvious. Heterokont biflagellates can have one long flagellum and one short, or one smooth and one hairy (bearing mastigonemes). In our case, control specificity is all the more surprising as we only considered species whose flagella exhibit no exterior morphological differences. During flagellar assembly and lengthening, tubulin subunits are shuttled from an intracellular pool by molecular motors and added to the tips of growing flagella by a process known as intraflagellar transport [53]. This process is unlikely to result in structural heterogeneities between the individual flagella axonemes. Instead, asymmetries arising from physiological differences in basal body age [54] and associated contractile structures may be more significant. In flagellate green algae, basal body duplication is semi-conservative. The two flagella of C. reinhardtii are termed cis or trans according to their proximity to a single eyespot: the trans basal body is inherited from the parental cell but the cis is formed anew. This distinction likely underlies their differential calcium response, important during phototaxis [11]. The octoflagellate exhibits an extreme form of this asymmetric ultrastructural patterning, where the eight flagella are again distinguishable by the age of the attached basal body and organization of accessory structures (figure 5a).
Asymmetries in the control architecture itself are sufficient but not necessary to template asymmetric dynamics. A radially symmetric nerve ring innervates the 5-legged starfish [55], yet the organism routinely performs a breaststroke gait differentiating between two pairs of side arms and a single leading arm. It would therefore be interesting to explore whether the ‘special’ flagellum in *P. octopus* is always the same flagellum (as uniquely identified by basal body numbering), or whether different flagella can take turns to assume the ‘exploratory’ position (electronic supplementary material, video S10). Given the dual sensory and motor capabilities of cilia and flagella, this could signify an early evolution of division of labour. We suggest that symmetry-breaking in excitable or inhibitory signalling in the algal flagella apparatus specifies the local state of contractility of intracellular fibres to control the activation state of individual flagella. This may be likened to the case of vertebrate limbs, where antagonistic control of flexor and extensor muscles is provided by motor neurons or to marine invertebrate larvae, where specialized ciliomotor neurons induce coordinated ciliary arrest [56]. We showed that in multiflagellates the ability to activate subsets of flagella provides a novel mechanism for trajectory reorientations that is distinct from steering in sperm [57] and in other uniflagellates that rely upon a change in the beat pattern of a single flagellum. A quadriflagellate can turn left, right, up or down, depending upon which of its four flagella is active. However, the extent to which organisms make use of these capabilities warrants further study.

The diverse gait of multiflagellates may have arisen from different evolutionary pressures associated with a need to occupy different ecological niches. Even small genetic changes could cause sufficient genetic rewiring of the basal apparatus and differences in locomotor pattern. With increasing flagella number, we suggest that new gait arises from superpositions of gait from flagellates with fewer flagella. In this perspective of gait modularity, the octoflagellate rotary breaststroke may have originated from intercalation of two quadriflagellate trotting gait, and the trot or gallop from intercalation of two pairs of biflagellate gait. Indeed, it is thought that *P. octopus* branched from a related quadriflagellate *Pyramimonas* species via incomplete cell division [49].

In conclusion, the capacity to generate and coordinate complex behaviour is by no means exclusive to organisms possessing neural circuitry. In diverse unicellular flagellates, phase-resetting processes and transitions in locomotor pattern occurring much faster than is possible by any passive means, implicating excitable signalling. In future work, we will seek to identify structures capable of functioning as pacemakers for locomotor pattern in flagellates. By analogy with CPGs, we suggest that parts of the algal cytoskeleton may be independently capable of generating coupled oscillations. This has enabled these organisms to dynamically reconfigure their gait in response to environmental changes and uncertainty, without the need for a central controller. The latter feat harkens to the embodiment perspective applicable to designing bioinspired robots, in which ‘control of the whole’ is ‘outsourced to the parts’ [58]. As modern technology strives towards greater automation, engineering adaptability into artificial systems has remained a formidable challenge. In this sense, cell motility is in fact a form of physical embodiment, wherein the compliant cytoskeleton and appendages enact morphological computation. In the light of these findings, we may wish to extend the scope of gait research beyond model vertebrates and invertebrates [24,59] to include aneural organisms. Much can be gained from exploring how simple unicellular organisms sculpt motor output and achieve sensor–motor integration—for herein lie the evolutionary origins of decentralized motion control.

Data accessibility. Please refer to the electronic supplementary material for videos S1–10, and supplementary text file for more information. Additional datasets are available at https://doi.org/10.5281/zenodo.3551942.

Competing interests. We declare we have no competing interests.

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References

1. Taute KM, Gude S, Tans SJ, Shimizu TS. 2015 High-throughput 3D tracking of bacteria on a standard phase contrast microscope. Nat. Commun. 6, 8776. (doi:10.1038/ncomms9776)
2. Catavitello G, Ivanenko Y, Lacquaniti F, Calabrese RL, Chowdhury R. 2018 A kinematic synergy for terrestial locomotion shared by mammals and birds. *eLife* 7, e38190. (doi:10.7554/eLife.38190)
3. Maybridge E. 1979 *Maybridge’s complete human and animal locomotion*, vol. 1. New York, NY: Dover Publications.
4. Ijspeert AJ. 2008 Central pattern generators for locomotion control in animals and robots: a review. *Neural Netw.* 21, 642–653. (doi:10.1016/j.neunet.2008.03.014)
5. Archavsky YI, Delligaia TG, Ofrovsky GN, Panchin YV, Pospelov LB, Sadreyev RI. 1998 Analysis of the central pattern generator for swimming in the mollusk *Clione*. *Ann. NY Acad. Sci.* 860, 51–69. (doi:10.1111/j.1749-6632.1998.tb09038.x)
6. Anderson LS et al. 2012 Mutations in DMRT3 affect locomotion in horses and spinal circuit function in mice. *Nature* 488, 642–646. (doi:10.1038/nature11399)
7. Lauga E. 2019 Bacterial hydrodynamics. *Annu. Rev. Fluid Mech.* 48, 105–130. (doi:10.1146/annurev-fluid-122414-034606)
8. Blair DF. 1995 Hoxa and cilia in thorax and swim. *Annu. Rev. Microbiol.* 49, 489–522. (doi:10.1146/annurev.micro.49.100195.002421)
9. Man Y, Page W, Poole RJ, Lauga E. 2017 Bundling of elastic filaments induced by hydrodynamic interactions. *Phys. Rev. Fluids* 2, 123101. (doi:10.1103/PhysRevFluids.2.123101)
10. Jikeli JF et al. 2015 Sperm navigation along helical paths in 3D chemosensory landscapes. *Nat. Commun.* 6, 7985. (doi:10.1038/ncomms9895)
11. Ruffer U, Nultsch W. 1987 Comparison of the beating of cis- and trans-flagella of *Chlamydomonas* cells held on micropipettes. *Cell Motil. Cytokeleton.* 7, 87–93. (doi:10.1002/cm.970070111)
12. Hegemann P. 1997 Vision in microalgae. *Planta*. 203, 265–274. (doi:10.1007/s004250050191)
13. Ruffer U, Nultsch W. 1991 Flagellar responses of *Chlamydomonas* cells held on micropipettes II. Change in flagellar beat pattern. *Cell Motil. Cytokeleton.* 18, 269–278. (doi:10.1002/cm.970180404)
14. Wan KY, Leptos KC, Goldstein RE. 2014 Lag, lock, sync, slip: the many ‘phases’ of coupled flagella. *J. R. Soc. Interface*. 11, 20131160. (doi:10.1098/rsif.2013.1160)
15. Wan KY, Goldstein RE. 2016 Coordinated beating of algal flagella is mediated by basal coupling. *Proc. Natl Acad. Sci. USA* 113, E2784–E2793. (doi:10.1073/pnas.1518527113)
16. Woolley DM, Vernon GG. 2001 A study of helical and planar waves on sea urchin sperm flagella, with a theory of how they are generated. J. Exp. Biol. 204, 1333–1345.

17. Sartori P, Geyer VF, Scholich A, Julicher F, Howard J. 2016 Dynamic curvature regulation accounts for the symmetric and asymmetric beats of Chlamydomonas flagella. Elife 5, e13258. (doi:10.7554/eLife.13258)

18. Man Y, Ling F, Kano E. 2019 Cilia oscillations. Phil. Trans. R. Soc. B 375, 20190157. (doi:10.1098/rstb.2019.0157)

19. Tsang AKH, Lam AT, Riedel-Kruse IH. 2018 Polygonal motion and adaptable phototaxis via flagellar beat switching in the microswimmer Euglena gracilis. Nat. Phys. 14, 1216–1222. (doi:10.1038/s41567-018-0277-7)

20. Qin B, Gopinath A, Yang J, Gollub JP, Aratia PE. 2015 Flagellar kinematics and swimming of algal cells in viscoelastic fluids. Sci. Rep. 5, 9190. (doi:10.1038/srep09190)

21. Sun SY et al. 2018 Flagellum couples cell shape to motility in Trypanosoma brucei. Proc. Natl Acad. Sci. USA. 115, E5916–E5925. (doi:10.1073/pnas.1722618115)

22. Wan KY. 2018 Coordination of eukaryotic cilia and flagella. Essays Biochem. 62, 829–838. (doi:10.1042/BEC20180029)

23. Nawroth JC, Feit KE, Colin SP, Costello JH, Dabiri JO. 2010 Phenotypic plasticity in juvenile jellyfish medusae facilitates effective animal–fluid interaction. Bioll. Lett. 6, 389–393. (doi:10.1098/rsbl.2010.0068)

24. Katz PS. 2015 Evolution of central pattern generators and rhythmic behaviours. Phil. Trans. R. Soc. B. 371, 20150057. (doi:10.1098/rstb.2015.0057)

25. Schmidt JA, Eckert R. 1976 Calcium couples flagellar reversal to photostimulation in Chlamydomonas reinhardtii. Nature 262, 713–715. (doi:10.1038/262713a0)

26. Kunita I, Kunoda S, Ohki K, Nakagaki T. 2014 Attempts to retreat from a dead-ended long capillary by backward swimming in Paramecium. Front. Microbiol. 5, 270. (doi:10.3389/fmicb.2014.00270)

27. Wan KY, Goldstein RE. 2018 Time irreversibility and criticality in the motility of a flagellate microorganism. Phys. Rev. Lett. 121, 058103. (doi:10.1103/PhysRevLett.121.058103)

28. Jeanneret R, Contino M, Polin M. 2016 A brief introduction to the model microswimmer Chlamydomonas reinhardtii. Eur. Phys. J. Spec. Top. 225, 2141–2156. (doi:10.1140/epjst/e2016-60065-3)

29. Leptos KC, Wan KY, Polin M, Tuval I, Pesci AI, Goldstein RE. 2013 Antiphase synchronization in a flagellar-dominance mutant of Chlamydomonas. Phys. Rev. Lett. 111, 158101. (doi:10.1103/PhysRevLett.111.158101)

30. Kildidt GS, Rufoluf C, Wagner C, Friedrich BM. 2017 In-phase and anti-phase flagellar synchronization by wavefront compliance and basal coupling. New J. Phys. 19, 113052. (doi:10.1088/1367-2630/aae031)

31. Quaranta G, Aubin-Tam ME, Tam D. 2015 Hydrodynamics versus intracellular coupling in the synchronization of eukaryotic flagella. Phys. Rev. Lett. 115, 238101. (doi:10.1103/PhysRevLett.115.238101)

32. Liu YJ, Claydon R, Polin M, Brumley DR. 2018 Transitions in synchronization states of model cilia through basal-connection coupling. J. R. Soc. Interface. 15, 20180450. (doi:10.1098/rsif.2018.0450)

33. Golubkis M, Stewart I, Buono L, Collins JJ. 1998 A modular network for legged locomotion. Physica D 115, 56–72. (doi:10.1016/S0167-2789(97)00222-4)

34. Moestrup O, Hill DRA. 1991 Studies on the Genus Pyramimonas (Prasinophyceae) from Australian and European waters: P. propulsa sp. nov and P. mitra sp. nov. Phycologia 30, 534–546. (doi:10.2216/i0031-8848-30-6-534.1)

35. Wilson D. 1967 Stepping patterns in tarantula spiders. J. Exp. Biol. 47, 133–151.

36. Ren Z, Hu W, Dong X, Sitti M. 2019 Multi-functional soft-bodied jellyfish-like swimming. Nat. Commun. 10, 2703. (doi:10.1038/s41467-019-10549-7)

37. Daugbjerg N, Moestrup O. 1992 Ultrastructure of Pyramimonas cyrtoperota sp nov (prasinophyceae), a species with 16 flagella from Northern Fose Basin, Arctic Canada, including observations on growth-rates. Can. J. Bot. 70, 1259–1273. (doi:10.1139/b92-159)

38. Anderson B, Shultz J, Jayne B. 1995 Axial kinematics and muscle activity during terrestrial locomotion of the centipede Scolopendra heros. J. Exp. Biol. 198, 1185.

39. Kano T, Sakai K, Yasa K, Okawa D, Ishiguro A. 2017 Decentralized control mechanism underlying interlimb coordination of millipedes. Bioinspir. Biomim. 12, 036007. (doi:10.1049/bi.201746-3190/ a6546)

40. Burrows M, Hoyle G. 1973 The mechanism of rapid running in the ghost crab, Ocypode ceratophthalma. J. Exp. Biol. 58, 327–349.

41. Tinevez JY, Perry N, Schindelin J, Hoopes GM, Williams EA, Shahidi R, Jekely G. 2017 Ciliomotor decentralization, signal-transduction pathways can trigger transitions in the synchronization modes of elastic filaments through basal-connection coupling. Phil. Trans. R. Soc. Lond. B. 375, 20190376. (doi:10.1098/rstb.2019.0376)

42. Robinson JM, Witman GB. 2002 Intrflagellar transport. Nat. Rev. Mol. Cell Biol. 3, 813–825. (doi:10.1038/nrm9552)

43. Bornens M. 2018 Cell polarity: having and making sense of direction—on the evolutionary significance of the primary cilium/centrosome organ in Metazoa. Open Biol. 8, 180052. (doi:10.1098/rsob.180052)

44. Watanabe W, Kano T, Suzuki S, Ishiguro A. 2011 A decentralized control scheme for orchestrating versatile arm movements in opiniroid omnidirectional locomotion. J. R. Soc. Interface. 9, 102–109. (doi:10.1098/rsif.2011.0317)

45. Veraszto C, Ueda N, Bezares-Calderon LA, Panzera A, Daun-Gruhn S, 2019 The steering gaits of model cilia-based mechanosensory systems and their functions in marine animal behaviour. Phil. Trans. R. Soc. B 375, 20190376. (doi:10.1098/rstb.2019.0376)

46. Tinevez JY, Perry N, Schindelin J, Hoopes GM, Williams EA, Shahidi R, Jekely G. 2017 Ciliomotor decentralization, signal-transduction pathways can trigger transitions in the synchronization modes of elastic filaments through basal-connection coupling. Phil. Trans. R. Soc. Lond. B. 375, 20190376. (doi:10.1098/rstb.2019.0376)

47. Williams EA, Shahidi R, Jekely G. 2017 Ciliomotor decentralization, signal-transduction pathways can trigger transitions in the synchronization modes of elastic filaments through basal-connection coupling. Phil. Trans. R. Soc. Lond. B. 375, 20190376. (doi:10.1098/rstb.2019.0376)