Generalized locomotion of brittle stars with an arbitrary number of arms

Daiki Wakita¹, Katsushi Kagaya^{2,3}, Hitoshi Aonuma^{1,4}*

¹Graduate School of Life Science, Hokkaido University, Sapporo 060-0810, Japan

²The Hakubi Center for Advanced Research, Kyoto University, Yoshida-Konoe 606-8501, Japan

³Seto Marine Biological Laboratory, Field Science, Education and Research Center, Kyoto University, Shirahama 649-2211, Japan

⁴Research Institute for Electronic Science, Hokkaido University, Sapporo 060-0812, Japan

*Correspondence to Hitoshi Aonuma

Research Center of Mathematics for Social Creativity, Research Institute for Electronic Science, Hokkaido University, Sapporo 060-0812, Japan.

Tel/Fax: +81-11-706-3832, Email: aon@es.hokudai.ac.jp

bioRxiv preprint doi: https://doi.org/10.1101/616383; this version posted April 23, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

Abstract

Typical brittle stars have five radially symmetrical arms, which coordinate to move their body in a certain direction. However, some species of them show individual difference in the number of arms: commonly five or six, rarely four or seven. We find this trait unique since intact legged animals each own a fixed number of limbs in general. How does a single species manage such different numbers of motile organs to realize adaptive locomotion? Our study aims to describe four- to seven-armed locomotion to explore a common rule across different arm numbers in brittle stars. Gathering several quantitative indices obtained from *Ophiactis brachyaspis*, we figured out an average locomotion where a front position emerges at one of the second neighboring arms to a mechanically stimulated arm, while side arms adjacent to the front synchronously work as left and right rowers, regardless of the total number of arms. The idea would provide a general scheme of how 'left and right' emerges in a radially symmetrical body.

Main text

Legged animals utilize appendages to move around on the ground. In most cases, intact adults of each species have a constant number of motile organs, such as four in turtles, six in beetles, and eight in spiders. Supposedly, each species adopts a number-dependent mode of locomotion which is optimal for a situation. In this context, some species of brittle stars (Echinodermata: Ophiuroidea) exhibit an appealing individual difference, where some intact individuals have five appendages or less, while others have six or more (Fig. 1). This difference in number is usually found in fissiparous species, which undergo asexual reproduction by fission and regeneration (Mladenov et al., 1983; Mladenov and Emson, 1984).

As typical echinoderms show pentaradial symmetry, most ophiuroid species standardly have five multi-jointed appendages called "arms," which extend from the "disk" at the center. Previous studies have described arm movements in the locomotion of five-armed species in qualitative terms (Romanes and Ewart, 1881; Preyer, 1887; von Uexküll, 1904; Glaser, 1907; Arshavskii et al., 1976a,b; Clark et al., 2019) as well as in quantitative terms (Astley, 2012; Kano et al., 2017). Several locomotor modes have been known even in a single species. An often reported mode, referred to as "breast stroke" by Arshavskii et al. (1976a,b) or "rowing" by Astley (2012), is characterized by a leading arm facing forward, two side arms working as left and right rowers, and two back arms dragged passively (Romanes and Ewart, 1881; Preyer, 1887; Glaser, 1907; Arshavskii et al., 1976a,b; Astley, 2012; Kano et al., 2017). Some studies have observed another locomotor mode, called as "paddling" by Arshavskii et al. (1976a) or "reverse rowing" by Astley (2012), where a backmost arm is dragged while the other four actively row (Prever, 1887; von Uexküll, 1904; Glaser, 1907; Arshavskii et al., 1976a; Astley, 2012). The ophiuroid body creeps in a certain direction with such bilaterally coordinated manners (Astley, 2012). Since the 'role' of each arm switches when the body changes moving direction (Astley, 2012), brittle stars do not have consistent antero-posterior and left-right axes in behavior.

Although the five-armed locomotion in common brittle stars and the individual difference in specific species have been viewed in different contexts, none has combined them to spotlight ophiuroid locomotion under the different numbers of arms.

Referring to the human body, whether the body comprises five, six, or other numbers of modules seems to bring a huge issue in individual function. How do these animals manage the difference in the number of motile organs to realize adaptive locomotion within a species? The aim of our study is to quantitatively describe the four-, five- six-, and seven-armed locomotion in the intact individual of an ophiuroid species, Ophiactis brachyaspis Clark, 1911 (Fig. 1), to explore a common rule across different arm numbers. Figuring out a control law which is flexible with the number of limbs, we would understand what sorts of body structure and interaction are of great or little importance to adaptive movements, and further apply such capacity in nature to a more flexible design in robotics. We will also provide a general scheme of how 'left and right'—'front and back' in the same time—emerges in a multidirectional body with an arbitrary number of rays to make a unidirectional behavior. One conclusion through our study is that a mechanical stimulus to an arm averagely makes one of its second neighboring arms be a leading arm, with the leader's side arms working as left and right synchronous rowers. Thus regardless of the total number of arms, ophiuroid locomotion shows a common anterior pattern, which could be positioned by counting how many arms some signal passes along a circular pathway.

Results

Moving direction (Θ)

The measured data of the post-stimulus moving direction Θ (Figs 2, 3; Eqn 1 in Methods) are shown in Fig. 4 by dot plots. For all the four-, five-, six-, and seven-armed cases, the model assuming a mixture of two von Mises distributions in Θ yielded smaller WAICs—better predictabilities—than that assuming a single distribution (Table 1). Compared with the small Δ (difference from the best one; Eqn 12 in Methods) of the one-distribution model in four- and five-armed animals, the six- and seven-armed Δ values were larger enough to interpret that bimodality was more obvious in more arms. Following the better model in terms of WAIC, we hereafter show the results on the assumption of two distributions for all the cases.

The posterior medians of two distributions' means, which were calculated

separately for the negative and positive ranges, were ± 17 , ± 29 , ± 46 , and ± 70 in four- to seven-armed animals, respectively. These estimated values signify that the more arms a brittle star had, the further two distributions of Θ were apart from each other (Fig. 4). In other words, the average moving direction of individuals with more arms was more angled from the opposite direction of the stimulated arm. Predictive distribution of Θ indeed depicted this trend (Fig. 4).

Left or right rower (B_{α})

The measured data of B_{α} —the α -th arm's degree of being a left or right rower (Figs 2, 3; Eqn 3 in Methods)—are schematized trial-by-trial in Figs S1–S4. As for the five- and six-armed populations, no-individuality models consistently took smaller WAICs than their counterparts where individuality was assigned to the mean of B_{α} (Table 1). We thus avoid mentioning individual difference within the same arm number.

Among L_a , S, Θ , Θ_{sign} , and F_a , the five-armed B_a was better explained by the continuous moving direction Θ , whereas the six- and seven-armed cases rather emphasized its sign Θ_{sign} (Eqn 2 in Methods) in discrete terms (Table 1). In four arms, the arm length L_a was chosen for a best explanatory variable although Θ showed a close performance. Given the dominance of the moving direction indicators as well as Θ 's bimodality (Fig. 4), we present the data of B_a separately by Θ_{sign} —in which side moving direction angled from the midline of the stimulated arm. Two groups were here defined by whether it angled clockwise ($\Theta_{sign} = 0$) or anticlockwise ($\Theta_{sign} = 1$).

The Θ_{sign} -based grouping exhibited a common locomotor mode among four-, five-, six-, and seven-armed animals in regards to B_a 's posterior means. The directional property of each arm could be explained by how many arms we count from the stimulated arm. Primarily, one of the *first* neighboring arms to the stimulated arm consistently took the largest or second largest $|B_a|$ —absolute values of posterior means (Figs 5–8A,C). This *first* arm corresponds to the anticlockwise neighbor of the arm 1 when $\Theta_{\text{sign}} = 0$ (Figs 5–8A) and the clockwise one when $\Theta_{\text{sign}} = 1$ (Figs 5–8C). In the next place, the *second* neighbor from the stimulus—next to the *first* in the same detour—took the smallest or second smallest $|B_a|$. Then, the *third* neighbor of the stimulus—next to the *second*—took the largest or second largest $|B_a|$ which was opposite in sign to that of the *first*. One exception was the seven-armed case when Θ_{sign} = 0 (Fig. 8A); the *second* (arm 3) and the *third* (arm 4) respectively had the fourth smallest and the third largest, probably due to the outlying trial shown at the row 1 of column 4 in Fig. S4. Replacing the ordinary cases' values with actual movements, the *first* actively pushed in the direction of the stimulated arm, while the *third* actively pushed oppositely to the *first*. These movements could make the *second* face forward, which indeed corresponded to the ranges of Θ in all the cases (Figs 5–8A,C).

Synchronization between two arms $(E_{\alpha\beta})$

The higher explanatory power of Θ_{sign} could also apply to the instance of the degree of synchronization between the α - and β -th arms, $E_{\alpha\beta}$ (Eqn 4 in Methods), because five-, six-, and seven-armed animals each brought the smallest WAIC in the model assuming Θ_{sign} 's effect (Table 1). In the four-armed case, the model without an explanatory variable best performed while the presence of Θ or Θ_{sign} resulted in similar predictabilities. Accenting the significance of Θ_{sign} as with B_{α} 's situation, we here show the resultant values of $E_{\alpha\beta}$ discretely by the sign of Θ .

A side-by-side comparison with the Θ_{sign} -based results of B_{α} shows us that the pair of the *first* and *third* rowers counting from the stimulus had the negatively largest medians of $E_{\alpha\beta}$'s posterior means in most cases (Figs 5–8B,D). Although one exception was found in the seven-armed with $\Theta_{sign} = 0$, the pair's value E_{24} leaned negatively as well (Fig. 8B). These values gave a quantitative indication that these two arms tended to simultaneously push in the opposite direction, regardless of the number of arms.

Discussion

Our study newly described the locomotion of brittle stars in a comparative context of four-, five-, six-, and seven-armed intact individuals in a single species. For this purpose, not stereotyping a discrete role of each arm, we introduced a quantitative index which can visualize each arm's degree of being a left or right rower, namely B_{α} . Coupled with other supportive values, this assessment would bring a unique idea of how 'left and right' emerges in the locomotion of a radially symmetrical animal (Fig. 9).

Locomotor modes

In past quantitative studies using five-armed brittle stars, antiphase synchronization of two distant arms has been supported by assessing the stop and start timing of arm movements (Astley, 2012) and by evaluating $E_{\alpha\beta}$ as in our study (Kano et al., 2017). This locomotor mode, which is referred to as "breast stroke" or "rowing," is characterized by a leading arm and its side rowing arms (Romanes and Ewart, 1881; Preyer, 1887; Glaser, 1907; Arshavskii et al., 1976a,b; Astley, 2012; Kano et al., 2017; Clark et al., 2019). Our study figured out that the triplet of left-front-right could appear even in four-, six-, and seven-armed individuals, suggesting that this locomotor mode is determined anteriorly, not laterally or posteriorly. In addition, the two back arms in the five-armed locomotor mode have been often interpreted as passively dragged ones (Romanes and Ewart, 1881; Preyer, 1887; Arshavskii et al., 1976a,b; Watanabe et al., 2012); nevertheless, our study showed that these arms rather worked as weaker rowers since their values of B_{α} ranged either negatively or positively (Fig. 5A,C). In six- and seven-armed ophiuroids, back arms following the two strong rowers similarly exhibited a trend of rowers, whereas the backmost ones were usually neutral as to the leftward or rightward bias just like the leading arm (Figs 6A,C, 8A,C). Thus more arms could take charge of 'rowers' especially when a brittle star has more arms.

Although "breast stroke" or "rowing" is a frequently reported locomotor mode in five-armed brittle stars, some studies have also described patterns where there is no leading arm. One is called as "paddling" or "reverse rowing," where a backmost arm is dragged while the other four actively row (Preyer, 1887; von Uexküll, 1904; Glaser, 1907; Arshavskii et al., 1976a; Astley, 2012). Such patterns without leading arms have been observed in free movement without experimental stimuli (Arshavskii et al., 1976a; Astley, 2012) as well as in escape behavior for a short time (Yee et al., 1987). In our study using *Ophiactis brachyaspis*, each trial seldom showed such a non-leading pattern (Figs S1–S4). Assuming this brittle star actually switches different locomotor modes, non-leading patterns might be employed only for several seconds after stimuli. In this case, our study might overlook or underestimate this urgent phase since we evenly analyzed one-minute duration after the beginning of the disk's movement. In either case, as far as post-stimulus locomotion was quantified for the fixed period, it seems the locomotor mode with a leading arm is more usual in the intact individuals of the Ophiactis species regardless of how many arms they have.

Decision of moving direction

Since brittle stars show no consistent front in behavioral terms as in most echinoderms, every arm can be a leading arm. Astley (2012) described their turning behavior in a short-term series, which was made by changing the roles of arms, not by rotating their body axis. As to an escape situation, several studies have observed that brittle stars avoid open or bright spaces (Cowles, 1910; Matsuzaka et al., 2017), a predator extract (Yee et al., 1987), and a KCl solution (Clark et al., 2019). However, few have probed into how such repellents make a certain reaction per arm to decide the moving direction of a whole individual. Since light and liquid diffuse in water, it is difficult to stimulate only a single target arm. Especially for small brittle stars such as *Ophiactis* species, mechanical stimuli would perform better with the aim to understand how signals from a stimulated arm affect the movements of the other arms.

In our study, two quantitative indices calculated from the filtered angular velocity of arms— B_a and $E_{a\beta}$ —and one obtained from the original coordinate data— Θ —could together visualize the ophiuroid locomotion without contradiction (Figs 5–8). Postulating each average of the two Θ_{sign} -based patterns as a representative, our numerical results suggest the most frequent locomotion pattern after a mechanical stimulus, in which a leading arm emerges at the second neighbor of a stimulated arm while side arms adjacent to the leader synchronously push backward. To realize this bilateral distribution with a high probability, it can be assumed that an afferent signal from an arm makes one of the *first* neighboring arms be an active rower which pushes in the direction of the signaling arm, the *second* neighboring arm be an inactive one which has a less directional preference, and the *third* neighboring arm be another active one which pushes synchronously but oppositely to the *first*'s pushing (Fig. 9). Accordingly, the second faces forward while the first, third, and some rear arms work on its both sides. In this model, whether the clockwise or anticlockwise second arm becomes a leading arm depends on in which detour the signal dominantly transfers from the stimulated arm, which is determined by some perturbation.

Under our model shown in Fig. 9, brittle stars with more arms would have a more risk of 'escape to stimulus.' If the front is placed ideally around the *second*

neighboring arm from the stimulus, four-, five-, six-, and seven-armed animals will respectively show 0, 36, 60, and 77 deg in average $|\Theta|$. In fact, the estimation from measured data copied it reasonably—17, 29, 46, and 70 deg, respectively—, and trials where moving direction rather inclined toward the stimulated arm (90 < $|\Theta| \le 180$) were more frequent as a body had more arms: 0/15, 1/30, 3/30, and 5/15, respectively (Fig. 4). Although the 'escape to stimulus' behavior is considered less adaptive, an evolutionary background would explain it. It has been proposed that primitive ophiuroids showed pentaradial symmetry (Paul and Smith, 1984; Sumrall and Wray, 2007), implying that brittle stars had developed a locomotion mechanism which worked optimally for the five-armed body. Some exceptional individuals in arm number, at least the four-, six-, and seven-armed bodies, probably have kept following this initial plan without vital issues, although escaping direction could be more or less bent as a side effect.

Our study has significance to understand how behavioral direction is expressed in a body without antero-posterior and left-right axes. Even when the individual body is round, some direction-making signal could transfer linearly at a local view (Fig. 9), just like a wave on a string or neural transmission in the spinal cord. Suppose brittle stars use this strategy, it seems not important how many segments with identical function are counted in the pathway. Otherwise, animal species would never allow individual difference in the number of motile organs.

Inter-arm interaction

Even if arms' function determines as represented in Fig. 9, our study remains questions of what kind of interactions mediates such coordination. In neurological aspects, the ophiuroid nervous system principally comprises a circumoral nerve ring in the disk and radial nerve cords extending into each arm (Cobb and Stubbs, 1981, 1982; Ghyoot et al., 1994; Bremaeker et al., 1997; Zuava et al., 2018). Some behavioral studies have supported the essential roles of the circumoral nerve ring in locomotion; menthol-anesthetic experiments indicated its function in initiating locomotion (Matsuoka et al., 2017); nerve cut experiments have demonstrated its necessity for coordinating arms (Mangold, 1909; Diebschlag, 1938; Arshavskii et al., 1976b; Clark et al., 2019). For such cases, the inter-arm connection depicted in Fig. 9 is recognizable as the circumoral nerve ring. We can assume that the movement of each arm directly reflects neural

activity in each radially symmetrical sector, which could be explained even by a couple of neurons. For instance, ophiuroid locomotion would be a useful material for testing "neuron ring" models (Suzuki et al., 1971; Matsuoka, 1985) to know how circularly arranged neurons work in the real world. Taking advantage of the unique individual difference in fissiparous brittle stars, we are able to demonstrate them with different neuron numbers as in computer, which would build a new bridge between theoretical biology and experimental biology.

Besides the crucial role of neural interactions, Kano et al. (2017) found the ophiuroid's ability to immediately change their locomotion patterns after the loss of certain numbers of arms, and then built an ophiuroid-like robot which imitated the adaptive locomotion via a local feedback without any preprogrammed control. Other robotics studies have also suggested the importance of physical interactions in movement coordination which is independent to electrical circuits (Owaki et al., 2013; Owaki and Ishiguro, 2017). Taking account of these researches as well, it is not likely that four- to seven-armed individuals each employ a different central control system while counting the total number of arms. Each arm would just refer to the states of its neighboring arms to realize a coordinated pattern at an individual level, no matter how many arms they own. A trial-by-trial variability in moving direction and other indices (Figs S1–S4) might reflect the influence of physical properties such as arms' posture at each moment, although a circular neural network might dominantly design the average orientation, where the stimulated arm's *second* neighbor faces forward (Fig. 9).

Such decentralized autonomous systems must contribute to the ophiuroid evolution that is flexible with the appendage number. It may be a reason why some species such as *Ophiactis brachyaspis* have acquired fissiparity, being capable of drastic morphological changes in a life cycle while retaining its locomotive ability.

Methods

Animals

We used the fissiparous brittle star *Ophiactis brachyaspis* (Fig. 1). In nature, this species densely inhabits the upper and side surfaces of rough rocks or other adherent

organisms such as sponges. Some of its arms lie in interstices while some rise from the substrate; suspension feeding ophiuroids show such a posture to capture particles (Warner, 1971). Animals collected in Shirahama Aquarium, Kyoto University, were reared in a laboratory aquarium ($600 \times 600 \times 600$ mm) filled with artificial seawater at 25–28°C (TetraMarin Salt Pro, Tetra Japan Co, Tokyo, Japan; salinity, 32–35‰). Body size ranged 1.5–3.0 mm in disk diameter and 5–15 mm in arm length.

Behavioral experiments

To investigate locomotor behavior, we chose 10 five-armed individuals and 10 sixarmed individuals, in each of which the lengths of arms did not differ by more than twice (c.f. Fig. 1). Four- and seven-armed individuals were also targeted; we obtained one for each. Each individual was put in a horizontal flat acrylic case $(105 \times 75 \times 22$ mm) filled with 100 mL of artificial seawater from the laboratory aquarium. There were no strong light gradient and no strong wind. Locomotion was recorded in aboral view using a digital camera (EOS8000D, Canon, Tokyo, Japan) with videos saved in MP4 format. We applied mechanical stimuli to arm tips using a toothpick. Stimulating an arm with subsequent observation was defined as one trial. The next trial came at the anticlockwise neighboring arm with an interval of more than two minutes. With repeating this rotation in order, every arm was stimulated at least three times for each individual.

Measurements

Per five- or six-armed individual, we analyzed three trials which showed the longest moving distances of the disk. In the four- and seven-armed cases, we picked out 15 trials with the longest moving distances. Analyzed duration for each trial was one minute after beginning to move the disk following each stimulus. The stimulated arm in each trial was numbered 1, which was followed anticlockwise by the other arms; α is the index of arms ($\alpha = 1, 2, 3, 4, 5$ in the five-armed instance). We tracked two coordinate points of the α -th arm using a tracking software Kinovea ver. 0.8.27 (http://www.kinovea.org/, accessed 4 December 2018) at 10 f.p.s.: $P_{\alpha}(t) = (x_{\alpha}(t), y_{\alpha}(t))$ —the attachment point between the α -th arm and the disk viewed aborally—and $P'_{\alpha}(t) = (x'_{\alpha}(t), y'_{\alpha}(t))$ —the point at half the length of the α -th arm, in terms of the range from the center of disk to the arm tip—at the *t*-th frame (Fig. 2). For the latter, we did not choose each arm's tip because it often rose and showed casual movements seeming irrelevant to locomotion as Matsuzaka et al. (2017) indicated. $P_{cent}(t)$ was defined as the center of gravity of all arms' $P_{\alpha}(t)$ (Fig. 2). The α -th arm's length (L_{α}) was defined as the maximum length of the segment $P_{\alpha}(t)P'_{\alpha}(t)$ in the analyzed duration. Note that L_{α} is a variable sampled in each trial, not accounting for the constant length of each arm. Moving distance (*S*) was measured as the length of $P_{cent}(1)P_{cent}(T)$, where *T* is the total number of frames, i.e. 600 (Fig. 2). We assessed moving direction (Θ) as follows:

$$\Theta = \frac{1}{T} \sum_{t=1}^{T} \left(\theta(t) \right) \quad (1),$$

where $\theta(t)$ is the angle made by the two segments $P_c(1)P_c(T)$ and $P_1(t)P_{cent}(t)$ (Figs 2, 3). Θ , which takes the range from -180 to 180 deg, is 0 deg when the disk moves in the opposite direction of the stimulated arm. A negative or positive value of Θ represents that the disk movement is angled clockwise or anticlockwise, respectively, from the opposite direction of the stimulated arm. For later uses in statistics, the dummy variable Θ_{sign} is defined as

$$\label{eq:sign} \boldsymbol{\theta}_{\mathrm{sign}} = \begin{cases} 0 \ (-180 \leq \boldsymbol{\Theta} < 0) \\ 1 \ (0 \leq \boldsymbol{\Theta} < 180) \end{cases} \ (2).$$

The segment $P_{\text{cent}}(t)P'_{a}(t)$ during locomotion swung around $P_{\text{cent}}(t)P_{a}(t)$, so the α -th arm's angle at the *t*-th frame ($\varphi_{a}(t)$) was defined as the angle made by these two segments (Fig. 2). $\varphi_{a}(t)$ is negative or positive when $P_{\text{cent}}(t)P'_{a}(t)$ is angled clockwise or anticlockwise, respectively, from $P_{\text{cent}}(t)P_{a}(t)$. $\varphi_{a}(t)$'s angular velocity ($\omega_{a}(t)$) was calculated with a five-point moving average method, and then smoothened with a low-pass filter with the cutoff frequency of 1.0 Hz (Fig. 3). To quantify to what extent each arm functions as left or right rower, we focused on that returning was faster than pushing in rowing arms. The filtered $\omega_{a}(t)$ was thus analyzed to evaluate the degree of a leftward or rightward bias in movement, which is represented by B_{a} (named after "bias"; Fig. 3):

$$B_{\alpha} = \frac{1}{T} \sum_{t=1}^{T} \left(\omega_{\alpha}(t)^{2} \operatorname{sign}(\omega_{\alpha}(t)) \right) \quad (3).$$

Assuming that a directional bias results from a speed difference between pushing and returning in each arm, we can rephrase B_{α} as the α -th arm's degree of being a left or right rower. A largely negative value of B_{α} represents that the α -th arm moves clockwise

faster than anticlockwise, indicating that it slowly pushes leftward and fast returns rightward viewed proximally from the disk. On the contrary, B_{α} is largely positive when the α -th arm pushes rightward (clockwise). Its value is close to zero when the α -th arm pushes leftward and rightward equally or is dragged without actively returning. We also extracted frequency components in the non-filtered $\omega_{\alpha}(t)$ of each arm using Fourier transforms. F_{α} was defined as the frequency at the peak amplitude in the α -th arm.

Besides for B_a , we used the filtered $\omega_a(t)$ to calculate Kano et al.'s (2017) " E_{ij} ," namely, the degree of synchronization between two arms:

$$E_{\alpha\beta} = \frac{1}{T} \sum_{t=1}^{T} \omega_{\alpha}(t) \omega_{\beta}(t) \quad (4)$$

A negative or positive value of $E_{\alpha\beta}$ represents that the movements of the α - and β -th arms synchronize in the opposite or same direction, respectively. A value around zero represents that the two arms move without strong correlation or are static.

Statistical modeling

We built statistical models for later comparative assessments with the following procedure. Firstly, to examine the structure of a possible bimodality in moving direction, we assume that Θ is subjected to a single von Mises distribution (f_{vM} , 'circular normal distribution'),

$$\Theta[n] \sim f_{\rm vM}(\mu_{\Theta}, \kappa_{\Theta}), -\pi \le \mu_{\Theta} \le \pi, \kappa_{\Theta} \ge 0 \quad (5)$$

or a mixture of two von Mises distributions,

$$\Theta[n] \sim \frac{1}{2} f_{\rm vM}(-\mu_{\Theta}, \kappa_{\Theta}) + \frac{1}{2} f_{\rm vM}(\mu_{\Theta}, \kappa_{\Theta}), -\pi \le \mu_{\Theta} \le \pi, \kappa_{\Theta} \ge 0 \quad (6).$$

Hereafter, *n* takes one to the total number of trials, so that $\Theta[n]$ denotes the *n*-th element of Θ . The parameters as random variables μ_{Θ} —converted to radians for modeling—and κ_{Θ} are analogous to the mean and the reciprocal of variance, respectively, in normal distribution. For the mixed case, we assume that the two distributions are symmetrical to each other with respect to the position of 0 deg.

Secondly, to understand what brings a trial-by-trial variability of B_{α} , we parametrize L_{α} , S, Θ , Θ_{sign} , and F_{α} each as an explanatory variable for B_{α} . We assume the normal distribution $f_{norm}(\mu, \sigma)$, where μ and σ respectively represent the mean and standard deviation (s.d.), as follows:

$$B_{\alpha}[n,\alpha] \sim f_{\text{norm}}(\mu_{\text{Bi}}[\alpha] + \mu_{\text{Bs}}[\alpha]X, \sigma_{\text{Bi}}[\alpha]), \sigma_{\text{Bi}} \ge 0 \quad (7).$$

Here, μ_{Bi} , μ_{Bs} , and σ_{Bi} are arm-by-arm parameters and *X* is an explanatory variable to which $L_{\alpha}[n, \alpha]$, S[n], $\Theta[n]$, $\Theta_{sign}[n]$, or $F_{\alpha}[n, \alpha]$ is assigned. *S*, Θ , and Θ_{sign} are common values for all the arms in the same trial. The categorical index Θ_{sign} is to know whether B_{α} varies continuously by Θ or switches discretely by the sign of Θ . In this instance, μ_{Bs} represents the mean's difference between the negative and positive cases since this variable disappears when Θ_{sign} is zero ($-180 \le \Theta < 0$) and appears when Θ_{sign} is one ($0 \le \Theta < 180$). The model without the member $\mu_{Bs}[\alpha]X$, i.e. without an explanatory variable, is for comparison. In parallel, let us consider whether B_{α} is better explained by individuality, namely, the quality made by some individual difference other than arm number as to five- and six-armed animals. Consideration of individuality is given to the mean's intercept μ_{Bi} :

$$\mu_{\mathrm{Bi}}[i,\alpha] \sim f_{\mathrm{norm}}(\mu_{\mathrm{B0}}[\alpha],\sigma_{\mathrm{B0}}), \sigma_{\mathrm{B0}} \ge 0 \quad (8),$$
$$B_{\alpha}[n,\alpha] \sim f_{\mathrm{norm}}(\mu_{\mathrm{Bi}}[i,\alpha] + \mu_{\mathrm{Bs}}[\alpha]X,\sigma_{\mathrm{Bi}}[\alpha]), \sigma_{\mathrm{Bi}} \ge 0 \quad (9),$$

where *i* takes one to the total number of individuals (i.e. 10) and the hyperparameters μ_{B0} and σ_{B0} are random variables. Let σ_{B0} , which is common in all arms, have a weakly informative prior as

$$\sigma_{\rm B0} \sim f_{\rm t}^+(3,0,20)$$
 (10),

where f_t^+ denotes the half *t* distribution and the parenthetical parameters represent the degree of freedom (*v*), location (mean when *v* > 1), and scale (s.d. divided by $\sqrt{3}$ when *v* = 3), respectively.

The final modeling is to examine which of Θ and Θ_{sign} is a better explanatory variable for $E_{\alpha\beta}$ in four- to seven-armed animals:

 $E_{\alpha\beta}[n,p] \sim f_{\text{norm}}(\mu_{\text{Ei}}[p] + \mu_{\text{Es}}[p]X, \sigma_{\text{Ei}}[p]), \sigma_{\text{Ei}} \ge 0 \quad (11),$

where μ_{Ei} , μ_{Es} , and σ_{Ei} are pair-by-pair parameters and the explanatory variable *X* takes *S*[*n*], Θ [*n*] or $\Theta_{\text{sign}}[n]$. Also considered is the model without the explanatory member $\mu_{\text{Es}}[p]X$.

Employing the Bayesian approach, posterior distribution of each parameter was estimated by the no-U-turn sampler (NUTS; Hoffman and Gelman, 2014)—a variant of Hamiltonian Monte Carlo (HMC) algorithm. In each sampling, we totally obtained 10,000 NUTS samples from four Markov chains, in each of which every 40th generation was sampled in 100,000 iterations after a warmup of 5,000, with the target

acceptance rate of 0.8. Convergence of each parameter was checked by trace plots, the potential scale reduction factor $\hat{R} \leq 1.1$, and the effective sample size $\hat{n}_{eff} \geq 40$, i.e. at least 10 per chain (Gelman et al., 2013). We assessed the predictability of the models based on WAIC (Watanabe, 2009, 2010), as this criterion is applicable to our models containing mixed distributions (Eqn 6) or hierarchical parameters (Eqns 8, 9). We developed the resultant statements according to better predicting models, which yielded smaller WAICs than the others considered. For comparison between the models, we referred to the difference as

$$\Delta = 2N(W - W_{\min}) \quad (12).$$

N is the total number of measured samples; multiplication by 2*N* is for the AIC scaling (Gelman et al., 2013). *W* is a given model's WAIC while W_{\min} is the smallest WAIC among those of the proposed models, so that Δ is zero in the best performed models. In presenting figures, the posterior predictive distributions of Θ are shown based on the parameters' posterior distributions in a better performed model. To visualize B_{α} and $E_{\alpha\beta}$ dependent on a better explanatory variable, we obtained the median of each posterior distribution under a model including the explanatory variable not only in the mean but also in the s.d.; Eqn 7 or Eqn 9 was modified to

 $B_{\alpha}[n, \alpha] \sim f_{\text{norm}}(\mu_{\text{Bi}}[\alpha] + \mu_{\text{Bs}}[\alpha]X, \exp(\sigma'_{\text{Bi}}[\alpha] + \sigma'_{\text{Bs}}[\alpha]X)) \quad (13),$ while Eqn 11 was replaced by

 $E_{\alpha\beta}[n,p] \sim f_{\text{norm}}(\mu_{\text{Ei}}[p] + \mu_{\text{Es}}[p]X, \exp(\sigma'_{\text{Ei}}[p] + \sigma'_{\text{Es}}[p]X)) \quad (14).$

Exponentiation in scale is to make the s.d. positive while σ'_{Bi} , σ'_{Bs} , σ'_{Ei} , and σ'_{Es} are random variables without constraints. We did not consider scale's explanatory variables in WAIC comparing terms because the Markov chain simulation failed to converge in many cases. All statistical computation was performed in the software environment R ver. 3.5.1 (R Core Team, 2018), where Stan codes were compiled and executed by the R package "rstan" (Stan Development Team, 2018). All source codes and data are available from the Figshare repository (Wakita et al., 2019).

References

1. Arshavskii YI, Kashin SM, Litvinova NM, Orlovskii GN, Fel'dman AG (1976a)

Types of locomotion in ophiurans. Neurophysiology 8: 398-404.

- Arshavskii YI, Kashin SM, Litvinova NM, Orlovskii GN, Fel'dman AG (1976b) Coordination of arm movement during locomotion in ophiurans. *Neurophysiology* 8: 404–410.
- 3. Astley HC (2012) Getting around when you're round: quantitative analysis of the locomotion of the blunt-spined brittle star, *Ophiocoma echinata*. *J Exp Biol* 215: 1923–1929.
- Bremaeker ND, Deheyn D, Thorndyke MC, Baguet F, Mallefet J (1997) Localization of S1- and S2-like immunoreactivity in the nervous system of the brittle star *Amphipholis squamata* (Delle Chiaje 1828). *P Roy Soc Lond B Bio* 264: 667–674.
- Carpenter B, Gelman A, Hoffman MD, Lee D, Goodrich B, Betancourt M, Brubaker MA, Guo J, Li P, Riddell A (2017) Stan: a probabilistic programming language. *J Stat Softw* 76: 1–32.
- Clark EG, Kanauchi D, Kano T, Aonuma H, Briggs DE, Ishiguro A (2019) The function of the ophiuroid nerve ring: how a decentralized nervous system controls coordinated locomotion. *J Exp Biol* 222: jeb192104.
- Cobb JL, Stubbs TR (1981) The giant neurone system in ophiuroids I: the general morphology of the radial nerve cords and circumoral nerve ring. *Cell Tissue Res* 219: 197–207.
- 8. Cobb JL, Stubbs TR (1982) The giant neurone system in ophiuroids III: the detailed connections of the circumoral nerve ring. *Cell Tissue Res* 226: 675–687.
- 9. Cowles RP (1910) Stimuli produced by light and by contact with solid walls as factors in the behavior of ophiuroids. *J Exp Zool* 9: 387–416.
- Diebschlag E (1938) Ganzheitliches verhalten und lernen bei echinodermen. Z Vergl Physiol 25: 612–654.
- 11. Gelman A, Stern HS, Carlin JB, Dunson DB, Vehtari A, Rubin DB (2013) *Bayesian Data Analysis*. 3rd edition. Florida, USA: CPC Press.
- Ghyoot M, Cobb JLS, Thorndyke MC (1994) Localization of neuropeptides in the nervous system of the brittle star *Ophiura ophiura*. *Phylos T Roy Soc B* 346: 433– 444.
- 13. Glaser OC (1907) Movement and problem solving in Ophiura brevispina. J Exp

Zool 4: 203-220.

- 14. Hoffman MD, Gelman A (2014) The No-U-Turn sampler: adaptively setting path lengths in Hamiltonian Monte Carlo. *J Mach Learn Res* 15: 1593–1623.
- 15. Kano T, Sato E, Ono T, Aonuma H, Matsuzaka Y, Ishiguro A (2017) A brittle starlike robot capable of immediately adapting to unexpected physical damage. *Roy Soc Open Sci* 4: 171200.
- 16. Mangold E (1909) Studien zur physiologie des nervensystems der echinodermen. *Pflug Arch Eur J Phy* 126: 371–406.
- 17. Matsuoka K (1985) Sustained oscillations generated by mutually inhibiting neurons with adaptation. *Biol Cybern* 52: 367–376.
- Matsuzaka Y, Sato E, Kano T, Aonuma H, Ishiguro A (2017) Non-centralized and functionally localized nervous system of ophiuroids: evidence from topical anesthetic experiments. *Biol Open* 6: 425–438.
- Mladenov PV, Emson RH (1984) Divide and broadcast: sexual reproduction in the West Indian brittle star *Ophiocomella ophiactoides* and its relationship to fissiparity. *Mar Biol* 81: 273–282.
- Mladenov PV, Emson RH, Colpit LV, Wilkie IC (1983) Asexual reproduction in the West Indian brittle star *Ophiocomella ophiactoides* (H.L. Clark) (Echinodermata: Ophiuroidea). *J Exp Mar Biol Ecol* 72: 1–23.
- 21. Owaki D, Ishiguro A (2017) A quadruped robot exhibiting spontaneous gait transitions from walking to trotting to galloping. *Sci Rep* 7: 277.
- 22. Owaki D, Kano T, Nagasawa K, Tero A, Ishiguro A (2013) Simple robot suggests physical interlimb communication is essential for quadruped walking. *J R Soc Interface* 10: 20120669.
- 23. Paul CRC, Smith AB (1984) The early radiation and phylogeny of echinoderms. *Biol Rev* 59: 443–481.
- 24. Preyer WT (1887) Die Bewegungen der Seesterne. Berlin, Germany: Friedländer.
- 25. R Core Team (2018) R: a language and environment for statistical computing. https://www.R-project.org (accessed 15 November 2018).
- 26. Romanes GJ, Ewart JC (1881) Observations on the locomotor system of Echinodermata. *Philos T R Soc Lond* 172: 829–885.
- 27. Stan Development Team (2018) Stan modeling language users guide and reference

manual, version 2.18.0. http://mc-stan.org (accessed 15 November 2018).

- Sumrall CD, Wray GA (2007) Ontogeny in the fossil record: diversification of body plans and the evolution of "aberrant" symmetry in Paleozoic echinoderms. *Paleobiology* 33: 149–163.
- 29. Suzuki R, Katsuno I, Matano K (1971) Dynamics of "neuron ring". *Kybernetik* 8: 39–45.
- 30. von Uexküll J (1905) Studien über den tonus II: die bewegungen der schlangensterne. Z Biol 46: 1–37.
- Wakita D, Kagaya K, Aonuma H (2019) Data from "Data and codes of "Generalized locomotion of brittle stars with an arbitrary number of arms"" Figshare. https://doi.org/10.6084/m9.figshare.8019827.v2 (deposited 23 April 2019).
- 32. Warner GF (1971) On the ecology of a dense bed of the brittle-star *Ophiothrix fragilis*. *J Mar Biol Assoc UK* 51: 267–282.
- 33. Watanabe S (2009) *Algebraic Geometry and Statistical Learning Theory*. Cambridge, UK: Cambridge University Press.
- 34. Watanabe S (2010). Asymptotic equivalence of Bayes cross validation and widely applicable information criterion in singular learning theory. *J Mach Learn Res* 11: 3571–3594.
- 35. Watanabe W, Kano T, Suzuki S, Ishiguro A (2011) A decentralized control scheme for orchestrating versatile arm movements in ophiuroid omnidirectional locomotion. *J R Soc Interface* 9: 102–109.
- 36. Yee A, Burkhardt J, Gilly WF (1987) Mobilization of a coordinated escape response by giant axons in the ophiuroid, *Ophiopteris papillosa*. J Exp Biol 128: 287–305.
- 37. Zueva O, Khoury M. Heinzeller T, Mashanova D, Mashanov V (2018) The complex simplicity of the brittle star nervous system. *Front Zool* 15: 1.

Acknowledgments

We thank Mr. Keita Harada in Shirahama Aquarium, Kyoto University, for sending us the fresh individuals of brittle stars from the aquarium. We are grateful to the member of Stan Community (https://mc-stan.org/community/) for suggesting better methods of bioRxiv preprint doi: https://doi.org/10.1101/616383; this version posted April 23, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

statistical modeling. This work was partly supported by JSPS KAKENHI (Grant Number 16KT0099) and by JST CREST (Grant Number JPMJCR14D5), Japan.

Author contributions

D.W. and H.A. designed the study and conducted behavioral experiments and measurements. D.W. and K.K. performed statistical modeling. All authors wrote the manuscript.

Competing interests

The authors declare no competing interests.

Data availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Material and correspondence

Hitoshi Aonuma

Tables

Madel	Specification		Four-armed			Five-armed				Six-armed			Seven-armed		
Model			Rank	WAIC	Δ	Rank	WAIC	Δ	Rank	WAIC	Δ	Rank	WAIC	Δ	
Θ	Distribution number														
1	one		2	0.876	0.917	2	1.192	0.764	2	1.518	5.55	2	1.860	10.7	
2	two		1	0.845	0*	1	1.179	0*	1	1.425	0*	1	1.502	0*	
B_{a}	Explanatory variable†	Individ- uality†													
1	no	no	3	4.217	4.94	8	4.671	60.4	5	5.338	58.9	4	4.940	25.7	
2	no	yes				9	4.676	61.9	10	5.352	64.0				
3	L_{α}	no	1	4.176	0*	10	4.679	62.9	9	5.350	63.2	5	4.947	27.1	
4	L_{a}	yes	—			12	4.688	65.4	12	5.382	74.9			_	
5	S	no	5	4.267	10.9	7	4.670	60.3	7	5.338	59.2	3	4.938	25.2	
6	S	yes				11	4.680	63.1	11	5.360	67.0				
7	Θ	no	2	4.187	1.29	1	4.469	0*	2	5.191	6.25	2	4.827	2.03	
8	Θ	yes				2	4.477	2.16	4	5.208	12.2				
9	$\Theta_{\rm sign}$	no	4	4.230	6.41	3	4.501	9.43	1	5.174	0*	1	4.818	0*	
10	$\Theta_{ m sign}$	yes				4	4.505	10.8	3	5.193	6.89				
11	F_a	no	6	4.271	11.3	5	4.640	51.0	6	5.338	59.1	6	4.955	28.8	
12	F_{a}	yes				6	4.644	52.3	8	5.347	62.4				
$E_{\alpha\beta}$	Explanatory	variable†													
1	no		1	3.951	0*	4	4.327	25.6	4	4.440	42.0	3	4.294	9.30	
2	S		4	3.974	4.14	3	4.321	22.0	2	4.413	17.5	4	4.320	25.7	
3	Θ		2	3.953	0.365	2	4.292	4.90	3	4.416	20.0	2	4.282	1.58	
4	$\Theta_{ m sign}$		3	3.959	1.45	1	4.284	0*	1	4.394	0*	1	4.279	0*	

Table 1. WAICs of statistical models for Θ , B_{α} , and $E_{\alpha\beta}$.

* $\Delta = 0$, bolded, indicating a best supportive model. †Not considered if "no"; otherwise, considered in the mean of normal distribution.

bioRxiv preprint doi: https://doi.org/10.1101/616383; this version posted April 23, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

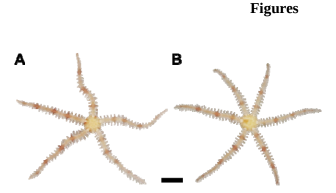
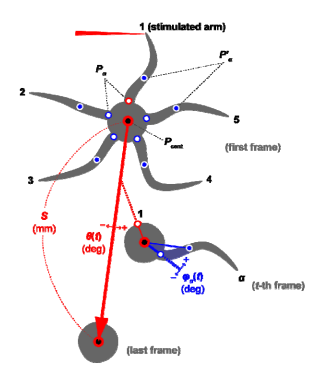
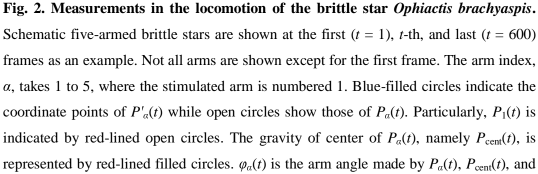


Fig. 1. The fissiparous brittle star *Ophiactis brachyaspis*. A: a five-armed individual. B: a six-armed individual. The scale bar represents 2 mm.





 $P'_{\alpha}(t)$. $\theta(t)$ is the angle made by the segment $P_{\text{cent}}(1)P_{\text{cent}}(600)$ and the segment $P_1(t)P_{\text{cent}}(t)$, representing the direction of the stimulated arm compared to moving direction. The moving distance *S* corresponds to the length of the segment $P_{\text{cent}}(1)P_{\text{cent}}(600)$.

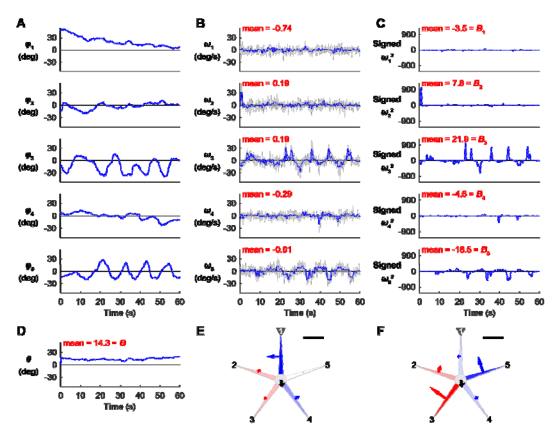


Fig. 3. Calculation and visualization in a five-armed example of the locomotion of the brittle star *Ophiactis brachyaspis*. A: temporal change of $\varphi_a(t)$ deg (c.f. Fig. 2). B: temporal change of $\omega_a(t)$ deg/s—angular velocity of $\varphi_a(t)$. Background gray plots represent the original data while thicker blue plots show low-pass filtered data. Each plot's "mean" shows the mean value of the filtered $\omega_a(t)$ for t = 1, ..., 600. C: temporal change of signed $\omega_a(t)^2$. Each plots' "mean" shows its mean value for t = 1, ..., 600, corresponding to B_a —the degree of being a left or right rower in the α -th arm. D: temporal change of $\theta(t)$ deg (c.f. Fig. 2). The "mean" shows its mean value for t = 1, ..., 600, corresponding to Θ (deg)—moving direction. E: schematized brittle star reflecting the mean $\omega_a(t)$ calculated in B and Θ in D. F: schematized brittle star reflecting B_a in C

and Θ in D. In E and F, each gray arrowhead indicates the stimulated arm numbered 1, with the number followed anticlockwise in order. The angles of black arrows at the disks represent Θ . An arm with a negative/positive mean value extends a blueleftward/red-rightward arrow, respectively, with its length corresponding to the absolute value of its mean. Compared to the mean values of the original $\omega_{\alpha}(t)$ in E, B_{α} in F well explains actual locomotion (c.f. Movie S1). Note that B_{α} originally reflects a returning direction by its sign (positive B_{α} denotes anticlockwise returning), but its schematized arrow here indicates a 'pushing direction' for simply imagining force to the ground (positive B_{α} denotes clockwise pushing, so apparently opposing the sign in Fig. 2). Scale bars represent 1.0 for the mean $\omega_{\alpha}(t)$ in E and 20 for B_{α} in F.

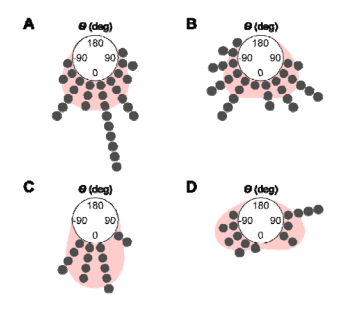
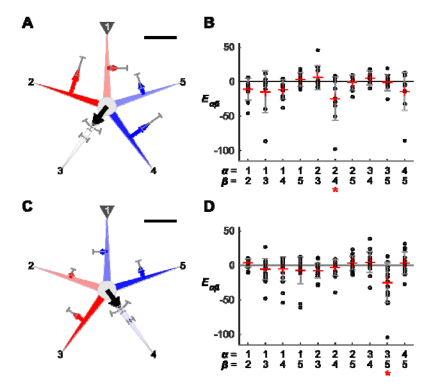


Fig. 4. Circular plots of moving direction after mechanical stimuli in the brittle star *Ophiactis brachyaspis*. A: five-armed case (10 individuals, 30 trials). B: six-armed case (10 individuals, 30 trials). C: four-armed case (one individual, 15 trials). D: seven-armed case (one individual, 15 trials). The moving direction Θ is the measured angle based on the position of a mechanically stimulated arm (c.f. Figs 2, 3). Θ is 0 deg when the disk moves in the opposite direction of the stimulated arm, and is negative/positive when the disk movement is angled clockwise/anticlockwise, respectively, from the 0 deg. Each point represents Θ in each trial, which is grouped in a bin divided per 22.5 deg. Density plots on the background represent predictive distributions on the



assumption of two symmetrical von Mises distributions.

Fig. 5. Five-armed locomotion grouped by moving direction in the brittle star *Ophiactis brachyaspis*. A,B: case where moving direction (Θ ; c.f. Figs 2, 3) is angled clockwise from the opposite direction of the stimulated arm, i.e. Θ is negative and $\Theta_{sign} = 0$ (eight individuals, 11 trials). C,D: case where Θ is positive (angled clockwise), i.e. $\Theta_{sign} = 1$ (10 individuals, 19 trials); an exemplary locomotion in this case is shown in Movie S1. A,C: schematized brittle stars reflecting the resultant quantitative values. Black arrows at the disks represent the measured means of moving distance (*S*; c.f. Fig. 2) by length and the measured means of Θ by angle. Error bars parallel to the disks' arrows show *S*'s standard deviation (s.d.) and arc-shaped error bars represent Θ 's s.d. in data. The blue or red arrow at each arm represents the degree of being a left or right rower (B_{α} ; c.f. Figs 2, 3), reflecting the absolute median of each posterior mean was negative/positive, its blue-leftward/red-rightward arrow extends from its arm, indicating that the arm pushed leftward/rightward (anticlockwise/clockwise), respectively. In each panel, the arm with the maximum absolute value in posterior mean

is colored with the most vivid blue/red, while the other arms show lighter blue/red corresponding to the relative values to the maximum. Scale bars represent 40 mm for *S* and 50 for B_{α} . B,D: degree of synchronization between two arms ($E_{\alpha\beta}$ for the α - and β -th arms). Small circles represent measured values. Pair-by-pair red pluses indicate the medians of posterior means while error bars show the medians of posterior s.d. parameters. Negative/positive values represent that the paired movement of the α - and β -th arms synchronized in the opposite/same direction, respectively. Each asterisk indicates the pair with the negatively largest estimated mean, showing remarkable antiphase synchronization. All posterior distributions for both B_{α} and $E_{\alpha\beta}$ were estimated under a better performed model in terms of WAIC, where Θ_{sign} is an explanatory variable for the mean and s.d.

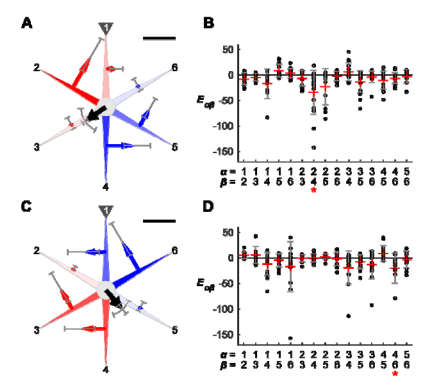
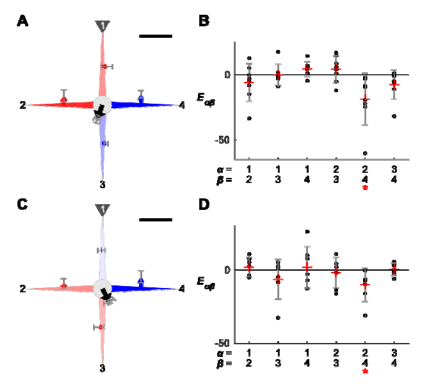


Fig. 6. Six-armed locomotion grouped by moving direction in the brittle star *Ophiactis brachyaspis*. A,B: case where $\Theta_{sign} = 0$ (eight individuals, 16 trials). C,D: case where $\Theta_{sign} = 1$ (eight individuals, 14 trials); an exemplary locomotion in this case is shown in Movie S2. A,C: schematized brittle stars reflecting the resultant quantitative values, as explained in Fig. 5. B,D: degree of synchronization between two arms ($E_{\alpha\beta}$



for the α - and β -th arms), as explained in Fig. 5.

Fig. 7. Four-armed locomotion grouped by moving direction in the brittle star *Ophiactis brachyaspis*. A,B: case where $\Theta_{sign} = 0$ (one individuals, eight trials). C,D: case where $\Theta_{sign} = 1$ (one individuals, seven trials). A,C: schematized brittle stars reflecting the resultant quantitative values, as explained in Fig. 5. B,D: degree of synchronization between two arms ($E_{\alpha\beta}$ for the α - and β -th arms), as explained in Fig. 5.

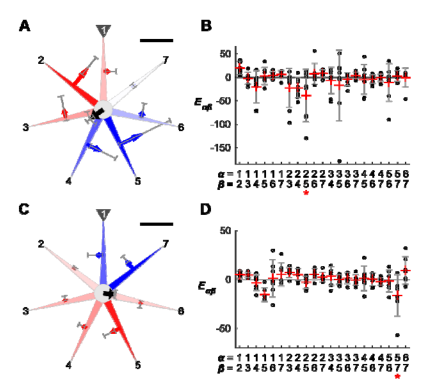


Fig. 8. Seven-armed locomotion grouped by moving direction in the brittle star *Ophiactis brachyaspis*. A,B: case where $\Theta_{sign} = 0$ (one individuals, eight trials). C,D: case where $\Theta_{sign} = 1$ (one individuals, seven trials). A,C: schematized brittle stars reflecting the resultant quantitative values, as explained in Fig. 5. B,D: degree of synchronization between two arms ($E_{\alpha\beta}$ for the α - and β -th arms), as explained in Fig. 5.

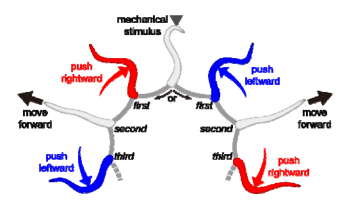
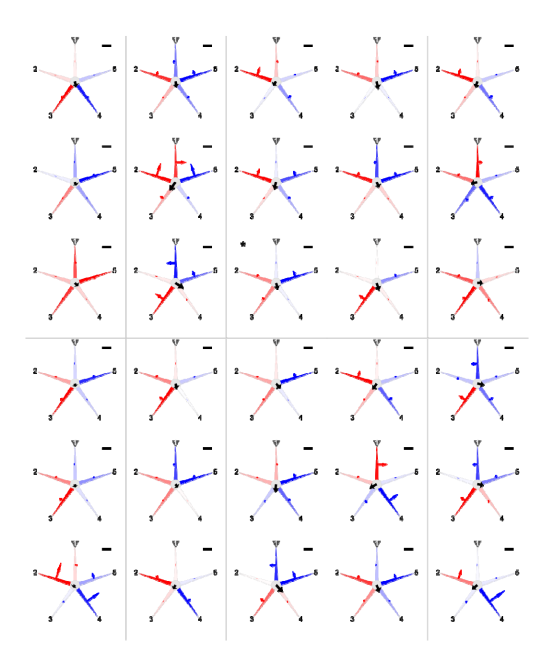


Fig. 9. Model of how a mechanical stimulus makes arm-by-arm locomotive movements in brittle stars with an arbitrary number of arms. An afferent signal from the stimulated arm transfers clockwise or anticlockwise to other arms through inter-arm connections made by neurons and/or other physical properties. Subsequently,

one of the *first* neighboring arms to the stimulated arm pushes actively in the stimulus direction, while the *third* neighbor in the same detour pushes oppositely to the *first*. As a result, the *second* arm between the *first* and *third* faces forward in behavioral terms.

bioRxiv preprint doi: https://doi.org/10.1101/616383; this version posted April 23, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



Supplementary information

Fig. S1. Five-armed trial-by-trial locomotion in the brittle star *Ophiactis brachyaspis*. Three trials were obtained from each of 10 individuals, which are partitioned by gray lines. Black arrows at the disks represent moving distance (*S*; c.f. Fig. 2) by length and moving direction (Θ ; c.f. Figs 2, 3) by angle. An arm with a negative/positive value for the degree of being a left or right rower (B_{α} ; c.f. Figs 2, 3)

extends a blue-leftward/red-rightward arrow, respectively, with its length corresponding to $|B_{\alpha}|$. In each panel, the arm with the maximum $|B_{\alpha}|$ is colored with the most vivid blue/red, while the other arms show lighter blue/red corresponding to the relative values to the maximum. Scale bars represent 20 mm for *S* and 50 for B_{α} . The asterisked trial (row 3, column 3) is shown in Movie S1.

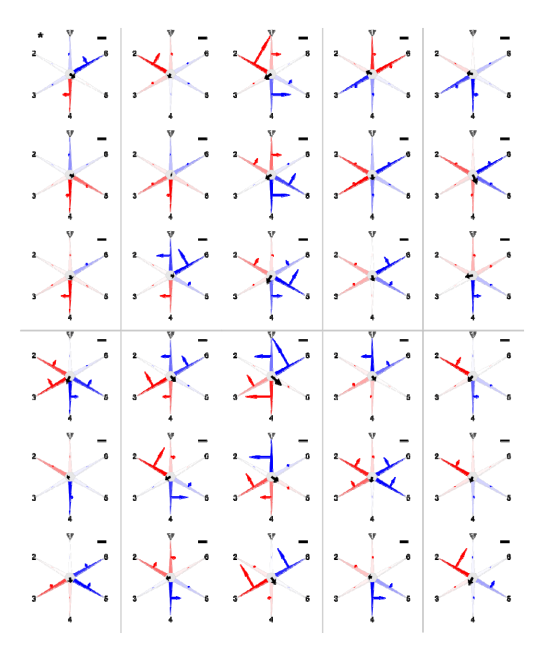


Fig. S2. Six-armed trial-by-trial locomotion in the brittle star Ophiactis brachyaspis.

Three trials were obtained from each of 10 individuals, which are partitioned by gray lines. Results are shown as in Fig. S1. The asterisked trial (row 1, column 1) is shown in Movie S2.

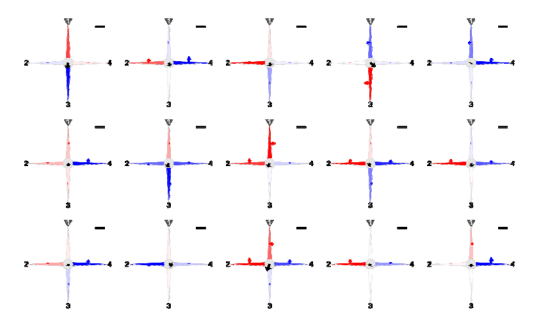


Fig. S3. Four-armed trial-by-trial locomotion in the brittle star *Ophiactis brachyaspis*. Fifteen trials were obtained from one individual. Results are shown as in Fig. S1.

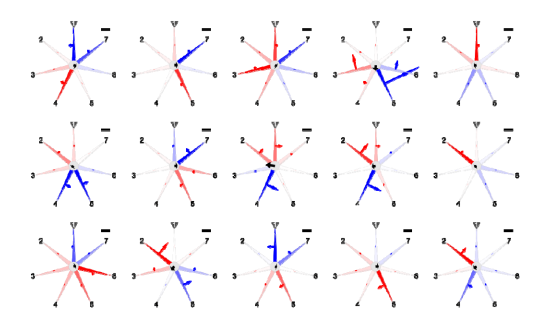


Fig. S4. Seven-armed trial-by-trial locomotion in the brittle star *Ophiactis brachyaspis*. Fifteen trials were obtained from one individual. Results are shown as in Fig. S1.

Movie S1. Locomotion of a five-armed individual of the brittle star *Ophiactis brachyaspis*. Quantitative analysis of this trial is presented in Fig. 3. Resultant values are schematized at the asterisked panel in Fig. S1.

Movie S2. Locomotion of a six-armed individual of the brittle star *Ophiactis brachyaspis*. Resultant values are schematized at the asterisked panel in Fig. S2.