

Reviewed Preprint

v1 • April 21, 2026

Not revised

Reviewed Preprint

v2 • August 4, 2026

Revised by authors

✉ For correspondence:

yuuki-k@nagasaki-u.ac.jp

† equal contribution

Competing interests: No competing interests declared

Funding: See [page 11](#)

Reviewing editor: Yuuki Y Watanabe, Graduate University for Advanced Studies, SOKENDAI, Japan

© 2026, Taniguchi et al. This article is distributed under the terms of the [Creative Commons Attribution License](#), which permits unrestricted use and redistribution provided that the original author and source are credited.

Evolution of sideways locomotion in crabs

Junya Taniguchi^{1,2,†}, Tsubasa Inoue^{1,†}, Kano Kohara^{1,2,†}, Jung-Fu Huang², Atsushi Hirai³, Nobuaki Mizumoto⁴, Fumio Takeshita⁵, Yuuki Kawabata¹ ✉

¹Graduate School of Integrated Science and Technology, Nagasaki University, Nagasaki, Japan • ²National Kaohsiung University of Science and Technology, Kaohsiung, Taiwan • ³Susami Crustacean Aquarium, Wakayama, Japan •

⁴Department of Entomology & Plant Pathology, Auburn University, Auburn, United States • ⁵Kitakyushu Museum of Natural History & Human History, Kitakyushu, Japan

eLife Assessment

This **valuable** study presents a comparative dataset on crab locomotion to investigate the evolution of sideways walking. The evidence supporting the authors' claims is **convincing**. This work will be of broad interest to researchers in animal locomotion and evolutionary biology.

<https://doi.org/10.7554/eLife.110015.2.sa3>

Abstract

The evolutionary change in the mode of locomotion is often a major evolutionary event, triggering diversification. Sideways locomotion is a defining feature of true crabs (Brachyura) and may have contributed to their ecological success. Yet, the evolutionary origin of this unique behavior remains unknown. Here we show that the prevalence of sideways locomotion in true crabs reflects a single evolutionary origin from a forward-moving ancestor. Our behavioral analysis of 50 live crab species indicates that crab locomotion can be broadly separated into two predominant modes, sideways and forward locomotion. The phylogenetic comparative analysis revealed a single origin of sideways locomotion, with multiple independent reversions to forward locomotion in ecologically specialized groups. The species richness data show that the lineage in which sideways locomotion originated is far more diverse than its nearest outgroups. These results are consistent with the idea that sideways locomotion acted as a key innovation contributing to the evolutionary diversification of true crabs. Such a rare but innovative behavioral trait provides a framework for understanding how locomotor modes shape evolutionary diversification in animals.

Introduction

The evolution of novel functional traits can enable organisms to exploit previously inaccessible ecological niches (Stroud & Losos 2016 [↗](#); Miller *et al.* 2023 [↗](#)). Such key innovations have shaped biodiversity on Earth by facilitating adaptive radiation within lineages. Because locomotion is a fundamental behavior that is involved in most survival and reproductive processes (Alexander 2003 [↗](#); Domenici 2010 [↗](#)), innovations in locomotor mechanisms are often linked to adaptive radiations (Astudillo-Clavijo *et al.* 2015 [↗](#); Higham *et al.* 2015 [↗](#); Burress & Wainwright 2019 [↗](#); Hedrick *et al.* 2020 [↗](#); Feiner *et al.* 2021 [↗](#)). For example, the evolution of flight in insects opened aerial niches globally and contributed to their enormous radiation (Grimaldi & Engel 2005 [↗](#)), whereas modifications of the locomotor skeleton in *Anolis* lizards facilitated localized adaptive radiations within islands (Feiner *et al.* 2021 [↗](#)). Most comparative studies on such innovations in locomotor behavior have taken a functional morphological approach. However, despite the

recognized importance of behavioral aspects of key innovations (Miller *et al.* 2023 [DOI](#)), the actual locomotor behaviors have rarely been compared across species, largely due to the challenges of obtaining large, comparative datasets of animal behaviors.

True crabs (Infraorder Brachyura) are iconic for their sideways locomotion, enabling them to achieve fast bidirectional movements (Vidal-Gadea *et al.* 2008 [DOI](#)), which may be beneficial for escaping from predators (Wolfe *et al.* 2021 [DOI](#)). Sideways locomotion is associated with greater joint flexibility in the lateral direction and a thorax elongated along the preferred direction of locomotion (Vidal-Gadea *et al.* 2008 [DOI](#)). This unique locomotion could be a key innovation in decapod crustaceans, as it changes the behavioral axis through which animals interact with the environment (Miller 1949 [DOI](#)). Moreover, sideways locomotion could have contributed to the ecological success of true crabs. The number of species of true crabs (~7,904 species) far exceeds that of their sister group, Anomura (hermit crabs and others; ~3,437 species), or their closest relatives, Astacidea (clawed lobsters and crayfish; ~792 species) and Achelata (spiny and slipper lobsters; ~153 species), according to the World List of Decapoda, DecaNet (DecaNet eds. 2025 [DOI](#)). True crabs have also successfully colonized diverse habitats globally, including terrestrial, freshwater, and deep-sea environments (Wolfe *et al.* 2024 [DOI](#); DecaNet eds. 2025 [DOI](#)). In addition, the crab-like body plan has evolved repeatedly among decapod crustaceans, a phenomenon known as carcinization (Morrison *et al.* 2002 [DOI](#); Tsang *et al.* 2011 [DOI](#); Keiler *et al.* 2017 [DOI](#); Wolfe *et al.* 2021 [DOI](#)). Despite this rich diversity and extensive morphological information, however, data on actual locomotor behaviors of crabs are sparse, and no comparative studies based on large datasets have been conducted thus far, making it difficult to evaluate the role of this unique locomotor mode on crab evolution and diversity.

Although most true crabs use sideways locomotion, some groups—including raninids, majids, and mictyrids—move predominantly forward (Sleinis & Silvey 1980 [DOI](#); Faulkes 2006 [DOI](#); Vidal-Gadea *et al.* 2008 [DOI](#)). This raises key questions: when did sideways locomotion originate, how many times did it evolve, and how many times did it revert? With a recent comprehensive crab phylogeny based on genomic data (Wolfe *et al.* 2024 [DOI](#)), here, we conducted behavioral analyses of 50 live crab species. We aimed to (i) pinpoint the origin of the sideways locomotion within Brachyura, (ii) estimate the number of transitions and reversions between sideways and forward locomotion, and (iii) test whether the emergence of sideways locomotion is associated with species diversification. Our results highlight sideways locomotion as a rare but innovative behavioral trait, providing a framework to understand how locomotor modes shape evolutionary diversification in animals.

Methods

Experimental procedures

We obtained live crabs from multiple sources, including intertidal and subtidal field collections, public aquaria, and local fish markets. Animals were kept only as long as required to record locomotion and were returned to their habitat or handled according to institutional animal care guidelines. Animal care and experimental procedures were approved by the Animal Care and Use Committee of the Faculty of Fisheries, Nagasaki University (Permit No. NF-0060) in accordance with the Guidelines for Animal Experimentation of the Faculty of Fisheries and the Regulations of the Animal Care and Use Committee of Nagasaki University.

Locomotion was recorded in plastic circular arenas (diameter 80–140 cm) whose medium matched each species' native environment (dry, seawater, freshwater, or brackish, with or without bare sand) (Fig. 1a [DOI](#)). Individuals were acclimated for 5 min in a bucket and then for 1 min inside a transparent cylinder placed at the arena center to minimize startle responses. After removing the cylinder, each trial was filmed for 10 min using a standard video camera (DSC-RX0, Sony Corporation, Tokyo, Japan) at 30 frames s⁻¹. The locomotion data were obtained from one representative individual per species due to logistical constraints and the low expected within-species variation. Our preliminary observations on several species for which many individuals were available suggest that locomotor direction is a species-level trait that is typically conserved.

Nevertheless, this design does not capture possible ontogenetic, size-dependent, or allometric variation within species. Accordingly, our conclusions are intended to identify broad interspecific patterns of locomotor direction rather than the full range of within-species variation.

Video analysis

Obtained videos were converted and downsampled to 5 frames s^{-1} for analysis using XMedia Recode 3.5 (www.xmedia-recode.de). For each frame, two landmarks along the longitudinal body axis (anterior and posterior carapace margins) were digitized using Kinovea 0.8.27 (www.kinovea.org) to estimate the instantaneous body axis and the centroidal position of the animal (Fig. 1b).

To standardize the evaluation of movement directions, we used a reference circle centered on the animal's starting position (Fig. 2). Each displacement bout was defined as the movement from the starting point to the point where the trajectory of the body center crossed the circle, with the next bout beginning once the reference circle was crossed. For each displacement bout, we computed the angle between the pre-movement body axis and the line to the crossed point as the movement direction (Fig. 2). Movements occurring to the left of the body axis were mirrored to the right before classification. Bouts were classified as forward (0–60° relative to the body axis), sideways (60–120°), or backward (>120°). This classification reflects three equal directional sectors in the full 360° movement space (forward/sideways/backward; 120° each), which provides a consistent reference under a null expectation of uniform movement directions. Backward movements were rare and therefore excluded from the analyses (Fig. S1). We calculated behavior using the Forward–Sideways Index (FSI): $FSI = (F - S) / (F + S)$, where F denotes the number of forward bouts and S denotes the number of sideways bouts. FSI values range from –1 (completely sideways) to +1 (completely forward). Species with $FSI < 0$ were classified as sideways movers, and those with $FSI > 0$ as forward movers.

The radius of the reference circle was determined systematically for each species (Table S1). For each species, we tested circle radii from 2 mm to 200 mm in 2-mm increments and computed the FSI at each radius. When the circle is too small, digitizing noise and body wobble near the start point tend to drive FSI toward 0. When the circle is too large, animals may turn within the circle before crossing, making the estimate unstable and unreliable. To capture the biologically meaningful motions, we selected the smallest radius at which the FSI showed a clear local maximum or minimum away from zero and then remained approximately stable as the radius increased further.

Ancestral state reconstruction

We extracted and pruned a recently published crab phylogeny (Wolfe *et al.* 2024), which was based on sequences of 10 genes (2 mitochondrial ribosomal RNA coding genes, 2 nuclear rRNA genes, and 6 nuclear protein-coding genes) and included 344 species across most major brachyuran lineages. Because our behavioral dataset did not always perfectly match the species included in (Wolfe *et al.* 2024), we reduced this tree to 44 genera, five families, and one superfamily, allowing closely related taxa to represent the observed species when the same species were unavailable. This approach enabled us to retain the terminals present in our dataset while preserving the placements of major clades relevant to this study (e.g., Eubrachyura, Raninoidea).

All terminals (44 genera, five families, and one superfamily) were coded as discrete states of either forward or sideways, based on the observed FSI values (positive: forward, negative: sideways). The tree was rooted with a hermit crab terminal (Anomura), for which we assigned the observed locomotor mode of *Coenobita purpureus* from our behavioral data. The node representing the common ancestor of Brachyura and Anomura (Meiura, starting/root node) was fixed to forward as a root prior. This assumption is based on the fact that further outer groups (e.g., crayfish, Astacidea) exhibit forward locomotion (Vidal-Gadea *et al.* 2008). Ancestral states were estimated on the pruned tree using maximum likelihood under equal-rates (ER) and all-rates-different (ARD) models, with model fit compared by Akaike Information Criterion (AIC). To summarize node-wise uncertainty and transition counts, we performed stochastic character mapping (500 replicates)

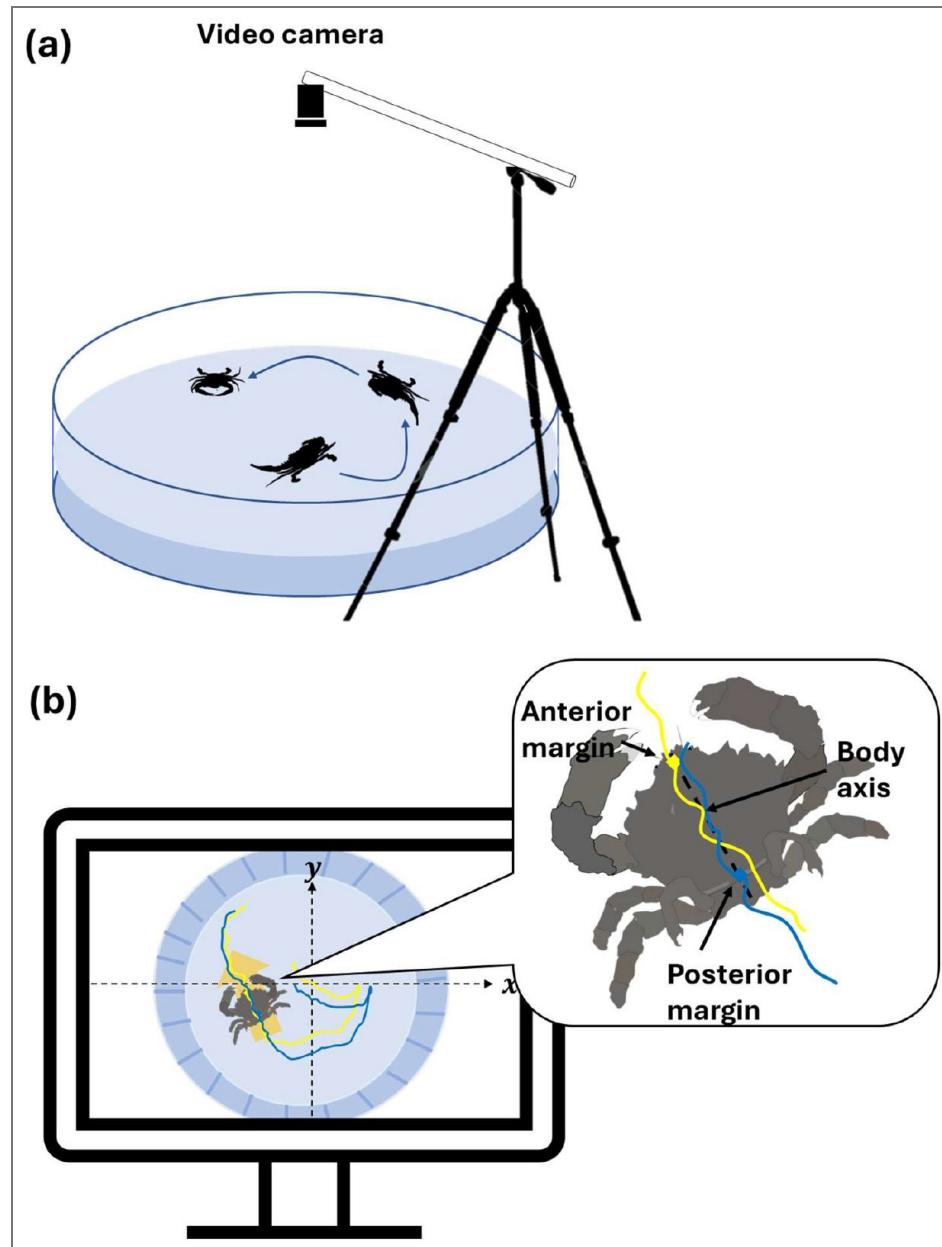


Figure 1. Video acquisition and analysis workflow.

(a) Experimental setup used to record each crab's behavior. (b) Extraction of two-dimensional position coordinates from video frames.

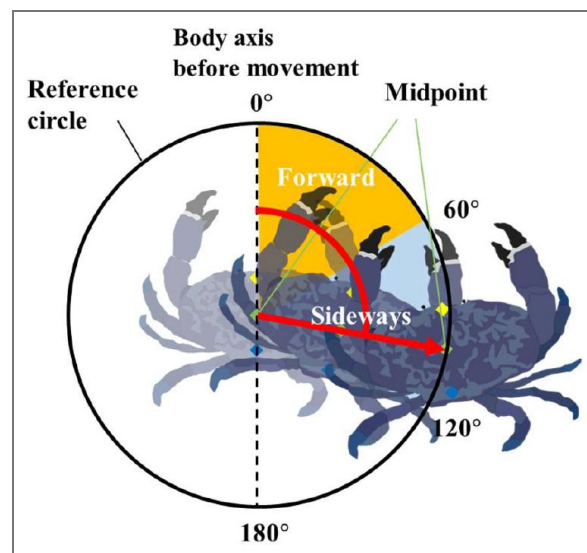


Figure 2. Method for determining movement directions of each crab.

For each movement bout, movement direction was defined as the angle between the previous body axis (from tail to head) and the displacement vector of the body's center (referred to as the midpoint). Displacement was measured when the midpoint reached the reference circle. These values across all movement bouts were then used to calculate the Forward-Sideways Index (FSI).

and reported posterior probabilities at key nodes and the posterior distributions of transitions and reversals. All analyses were conducted in R version 4.3.2 (R Core Team 2023) using the packages *ape*, *phytools*, and *geiger*.

Morphological analysis

To examine the relationship between body shape and locomotor mode, we used two simple carapace shape indices: relative carapace length (CL/CW, carapace length divided by carapace width) and relative carapace depth (CD/CS, carapace depth divided by carapace size), where carapace size (CS) was defined as the geometric mean of CL, CW, and CD, where CL is carapace length, CW is carapace width, and CD is carapace depth. We used phylogenetically informed ANOVA to test whether these indices differed between forward- and sideways-moving taxa.

Results

Of the 50 species, 35 were classified as sideways movers and 15 as forward movers based on FSI (Fig. 3; Table S1). For example, *Ranina ranina* showed an FSI of 0.89, indicating forward movement, whereas *Geothelphusa dehaani* showed an FSI of -0.70, indicating sideways movement (Fig. 4). FSI values showed a clear separation between these locomotion modes: forward movers had a median FSI of 0.82 (range: 0.24–0.94), while sideways movers had a median FSI of -0.80 (range: -1.00 to -0.39) (Fig. 3). This separation was supported by Hartigan's dip test ($D = 0.083$, $n = 50$, $p = 0.007$), indicating significant deviation from unimodality. All species-level circular histograms are provided in Supplementary Figure S1. To further examine the underlying angle distributions, we fitted one- and two-component Gaussian mixture models to the continuous bout-angle distributions of each taxon and examined peak locations and mixture weights (Table S2). Fourteen taxa were best described by a one-component model, whereas 36 taxa were best described by a two-component model. Among the 36 taxa best described by a two-component model, 33 had a dominant component explaining at least 70% of the distribution, whereas only three taxa showed relatively balanced two-component distributions. Thus, although some taxa showed mixed directional tendencies, most taxa were dominated by a primary directional component. As an additional, data-driven check of the original FSI-based classification, we estimated a boundary from the distribution of dominant peak locations (Fig. S2; Gaussian mixture cutoff $\approx 49.4^\circ$). This peak-based classification was identical to the original FSI-based forward/sideways classification (15/35), indicating that the main classification was not dependent solely on FSI.

In the morphological analysis (Fig. S3), relative carapace length (CL/CW) differed significantly between forward- and sideways-moving taxa (phylogenetically informed ANOVA: $F = 26.90$, $p < 0.001$), whereas relative carapace depth (CD/CS) did not differ significantly between the two groups ($F = 1.18$, $p = 0.403$). These results suggest that locomotor mode is associated with some aspects of carapace shape, but is not explained by simple carapace flattening alone.

Model comparison favored the ARD model over the ER model ($\Delta\text{AIC} = 6.23$; AIC weights = 0.957 vs. 0.043). Under the preferred ARD model, the transition rate from sideways to forward locomotion was estimated as 0.0029, slightly higher than the forward-to-sideways rate (0.0026). Ancestral state reconstruction placed the origin of sideways locomotion at the base of Eubrachyura (Fig. 5). Stochastic character mapping estimated the posterior probability of forward locomotion as 0.91 for the common ancestor of Brachyura, 0.79 for the common ancestor of Raninoida and Eubrachyura, and 0.24 for the eubrachyuran stem lineage. These results indicate that early-diverging lineages (i.e., Homoloida, Dromiacea, and Raninoida) retained the forward locomotion present in the common ancestor of Brachyura, and that sideways locomotion first appeared at the divergence between Raninoida and Eubrachyura. The ER model yielded a qualitatively similar placement, with only minor differences in the number of inferred transitions (Fig. S4).

To place these findings in a broader phylogenetic context, we examined locomotor states alongside species richness patterns in early-diverging brachyuran lineages. Cyclodorippoida is identified as the sister group of Eubrachyura, with Raninoida forming the next outgroup (Tsang *et al.* 2011);

Figure 3. Distribution of Forward-Sideways Index (FSI) values among crab species exhibiting forward and sideways locomotion.

Gold bars represent species classified as forward movers, and blue bars represent species classified as sideways movers.

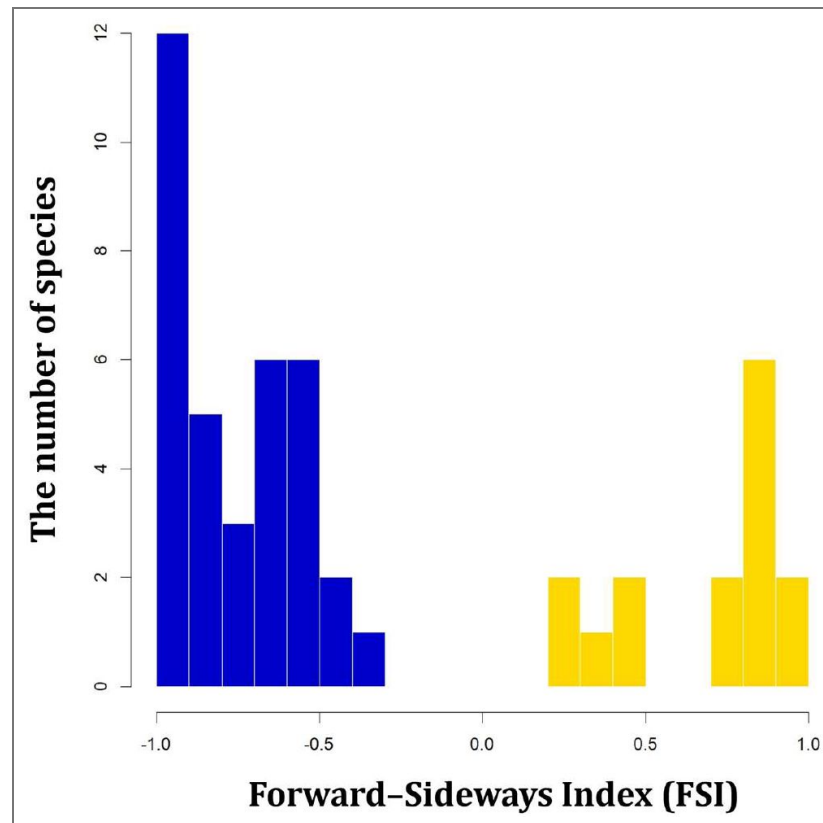
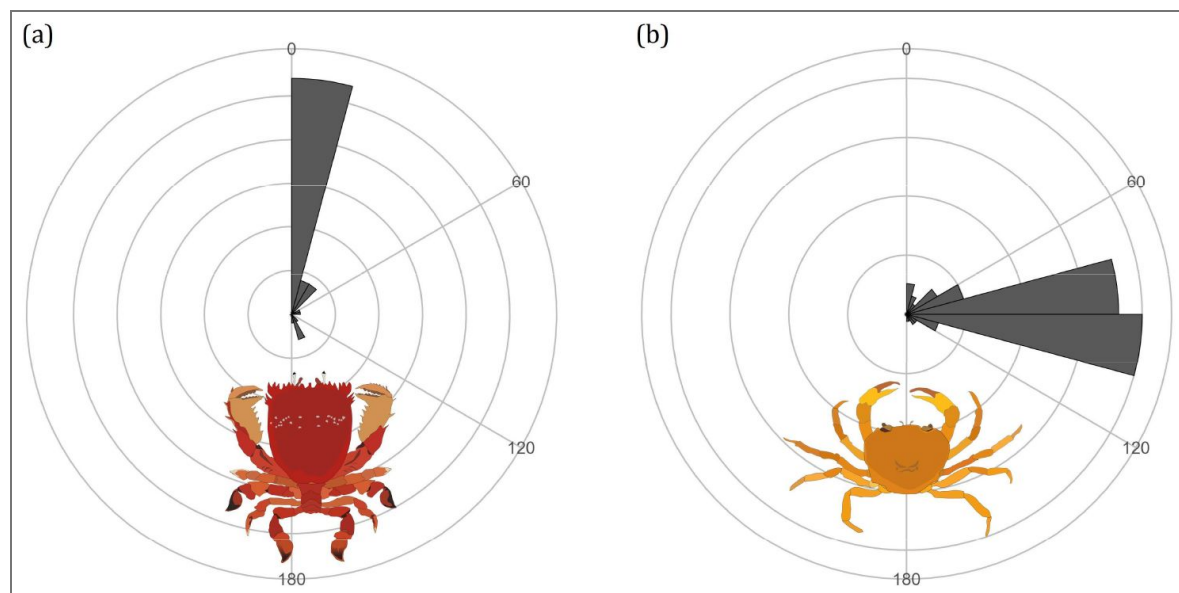


Figure 4. Representative circular histograms of movement directions in crabs.

(a) Forward movement in *Ranina ranina* (FSI = 0.89). (b) Sideways movement in *Geothelphusa dehaani* (FSI = -0.70). The 0°-180° axis denotes the crab's body axis before movement, with bars indicating the frequency of movement direction.



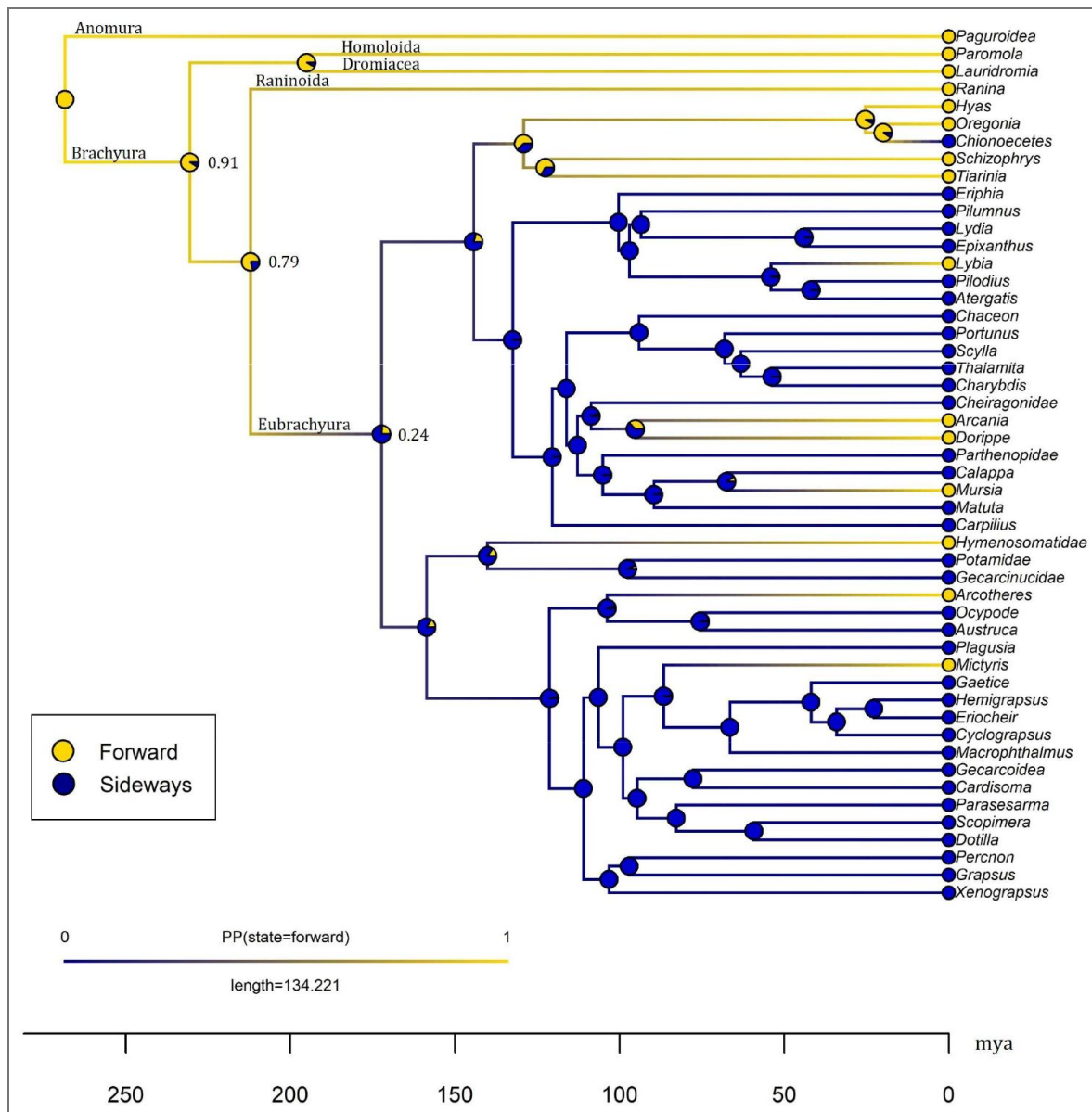


Figure 5. Ancestral state reconstruction of locomotion in crabs under the all-rates-different (ARD) model.

Gold circles at the tips indicate forward locomotion, whereas blue circles indicate sideways locomotion. Pie charts at internal nodes and along branches represent the posterior probabilities of each locomotor state, estimated from 500 stochastic character maps. The x-axis shows geological time, scaled in millions of years before present (mya).

Wolfe *et al.* 2024 [↗](#)). Behavioral data from Raninoida indicate that these crabs move forward (Fig. 3 [↗](#); Table S1). In contrast, the locomotor mode of Cyclodorippidae remains unknown because these crabs inhabit deep-sea environments that preclude direct behavioral observations. Based on DecaNet (World List of Decapoda) counts of accepted extant species (DecaNet eds. 2025 [↗](#)), Eubrachyura contains approximately 7,468 described species, whereas Cyclodorippoida includes about 110 species and Raninoida only ~46 species. This sharp disparity highlights that the clade in which sideways locomotion is fixed—at the base of Eubrachyura or potentially in the common ancestor of Cyclodorippoida and Eubrachyura—is associated with far greater taxonomic diversity than the lineages retaining forward locomotion.

Our results also indicate that the sideways mode was largely retained across major eubrachyuran lineages. However, despite this overall stability, we inferred multiple independent reversions to forward locomotion distributed across the tree, including lineages leading to Majidae (*Hyas*, *Oregonia*, *Chionoecetes*, *Schizophrys*, *Tiarinia*), *Lybia*, *Arcania*, *Dorippe*, *Mursia*, Hymenosomatidae, *Arcotheres*, and *Mictyris*. Interestingly, within Majidae, *Chionoecetes* likely underwent a secondary reversion back to sideways locomotion from a forward-moving ancestor within the group. Stochastic character mapping under the ARD model estimated sideways → forward reversions at a mean of 10.2 (interquartile range, IQR = 9–12) and forward → sideways gains at a mean of 4.1 (IQR = 3–5). Under the ER model, the corresponding estimates were 10.5 (9–12) and 4.6 (3–5), respectively. These results indicate that the evolution of sideways locomotion is rare but, once established, tends to be stably retained. Reversions to forward locomotion (and subsequent reversions back to sideways locomotion) occur under particular ecological specializations, producing a complex evolutionary pattern across true crabs.

Discussion

Our ever-biggest behavioral dataset on crab locomotion reveals that sideways locomotion originated only once from the forward locomotion ancestor, rather than through multiple independent origins (Fig. 5 [↗](#)). In other words, the widespread sideways locomotion across true crabs is highly conserved after being inherited from the common ancestor at the base of Eubrachyura. This suggests that modern sideways-moving crabs likely share the same anatomical, neurological, and developmental mechanisms, such as the reduction of motor neurons that control muscles of proximal legs (Vidal-Gadea & Belanger 2013 [↗](#)). The single transition event of sideways locomotion contrasts with carcinization, which has occurred repeatedly across decapods (Tsang *et al.* 2011 [↗](#); Keiler *et al.* 2017 [↗](#); Tan *et al.* 2018 [↗](#); Wolfe *et al.* 2021 [↗](#)). Carcinization has produced crab-like morphologies in several lineages of Anomura, including porcelain crabs (Porcellanidae), king crabs (Lithodidae), and the coconut crab *Birgus latro*. However, our behavioral observations indicate that porcelain crabs move predominantly backward rather than sideways, and king crabs and the coconut crab move predominantly forward (unpublished data). These examples demonstrate that even when crab-like body forms evolve, the characteristic locomotor mode (i.e., moving sideways) does not necessarily accompany them. This highlights a distinction between morphological convergence and behavioral innovation: while body forms may converge multiple times, fundamental behavioral transitions can be rare.

The species richness data show that Eubrachyura is far more diverse than its nearest outgroups, and the origin of sideways locomotion is inferred to lie around the base of the Eubrachyura. These results are consistent with the idea that this unique locomotor mode acted as a key innovation that contributed to the evolutionary diversification of Eubrachyura. Note, however, that a key innovation is not the only process driving adaptive radiation, and the innovation may not always result in radiation (Fürsich & Jablonski 1984 [↗](#); Miller *et al.* 2023 [↗](#)). External factors, such as ecological opportunity provided by mass extinction, are also critical for evolutionary diversification (Stroud & Losos 2016 [↗](#)). Based on the divergence times reported by (Wolfe *et al.* 2024 [↗](#)), the origin of sideways locomotion falls around ~200 Mya (earliest Jurassic, immediately post-Triassic–Jurassic extinction), a recovery interval marked by Pangaeon rifting, expansion of shallow-marine habitats, and the early Mesozoic Marine Revolution—conditions that typically increase ecological opportunity (Buatois *et al.* 2016 [↗](#); Schoepfer *et al.* 2022 [↗](#)). Disentangling the

relative roles of intrinsic innovation and extrinsic environmental change will require trait-dependent diversification analyses (Maddison *et al.* 2007), fossil-informed timelines, and performance tests that link sideways movement to adaptive advantages.

Sideways locomotion may confer several adaptive advantages. One likely advantage is the ability to move rapidly at similar speeds in both lateral directions (Vidal-Gadea *et al.* 2008; Wolfe *et al.* 2021), which is also supported by an experiment using crab-like robots (Chen *et al.* 2022). Having multiple locomotor directions is highly advantageous for escaping from predators, not only by making the escape direction unpredictable but also by providing multiple optimal escape routes (Domenici *et al.* 2011; Kawabata *et al.* 2023). Other possible advantages may include movement through confined spaces and visual-field sampling during locomotion, although these possibilities remain untested and will require future biomechanical and sensory studies. An alternative explanation is that sideways locomotion is a secondary phenomenon resulting from a flattened body structure that limits the distance between the legs and thereby makes forward movement difficult. However, our additional morphological analysis showed that relative carapace depth (CD/CS) did not differ significantly between forward- and sideways-moving taxa, whereas relative carapace length (CL/CW) differed significantly. Therefore, sideways locomotion cannot be explained solely by carapace flattening, and the morphological correlates of sideways locomotion are more complex.

Despite these potential adaptive advantages, sideways locomotion appears to have evolved rarely across the animal kingdom; it has occurred only in true crabs (Fig. 5), and potentially also in crab spiders (Wilcox 2017) and leafhopper nymphs (Chasen *et al.* 2014). One possible reason is that this locomotor mode fundamentally changes the behavioral axis, potentially affecting a wide range of behaviors such as predator avoidance, shelter use, burrowing, mating, and foraging (Atkinson & Eastman 2015; Crane 2015; Asakura 2016; Takeshita & Nishiumi 2022). In this sense, the origin of sideways locomotion may represent a rare and major behavioral transition rather than a simple change in movement direction alone.

Nevertheless, sideways locomotion does not appear to be universally favored across all ecological contexts. After the transition to sideways locomotion, crabs have experienced at least six independent reversions to forward locomotion (Fig. 5). These reversions are particularly associated with major changes in life history traits. For example, soldier crabs (*Mictyris*) predominantly use forward walking (Fig. S1; Table S1) in a way biomechanically similar to other forward-walking animals rather than sideways-walking crabs (Sleininis & Silvey 1980). Soldier crabs are unique for their gregarious nature and coordinated collective movements (Murakami *et al.* 2014), which may have brought them back to forward locomotion. Similarly, majoid crabs (e.g., *Oregonia*, *Tirarinia*) camouflage themselves with seaweed (Sato & Wada 2000; Hultgren & Stachowicz 2009), and pea crabs (e.g., *Arcotheres*) live hidden inside bivalves and other invertebrates, relying on their hosts for protection (de Gier & Becker 2020). Given that the major benefit of sideways locomotion is rapid escape from predators (Vidal-Gadea *et al.* 2008; Wolfe *et al.* 2021), these examples with alternative strategies of predator avoidance may no longer need sideways locomotion, resulting in secondary losses. These exceptional forward-moving species imply that it is costly to maintain sideways locomotion in crabs, and that they retain evolutionary flexibility to lose this locomotor mode under certain ecological pressures.

Modes of locomotion—such as walking, swimming, and flying—fundamentally shape how animals interact with the environment, affecting behaviors related to foraging, predator avoidance, and reproduction (Alexander 2003; Domenici 2010). Sideways locomotion in true crabs is also a critical change in the mode of locomotion, whose evolutionary history is characterized by rarity, stability, and occasional reversions. Our case study illustrates how major innovations can open new adaptive opportunities, yet remain constrained by both phylogenetic history and ecological context. By integrating direct behavioral observations with a robust phylogenetic framework, this study expands our understanding of how animal locomotor modes diversify and persist through evolutionary time.

Data availability

Supplementary figures and tables contain the data used to generate the figures.

Acknowledgements

We sincerely thank the staff and the students of the Kawabata Laboratory, Nagasaki University, and the Huang Laboratory, National Kaohsiung University of Science and Technology, for their assistance with crab sampling, experiments, and video analysis. We are deeply grateful to the staff of the Susami Crustacean Aquarium and the Wakayama Prefectural Museum of Natural History for generously providing experimental animals, assisting with field sampling, and allowing us to use their facilities for behavioral observations. We sincerely thank Takehiro Sato and the staff of the Kanagawa Prefectural Museum of Natural History for their generous assistance in measuring crab morphology. We are also grateful to Akinori Yamada, Nagasaki University, for his assistance with the preliminary analysis of crab phylogeny construction using genomic data.

Additional files

[Supplementary Table S1.](#)

[Supplementary Table S2.](#)

[Supplementary Figures.](#)

Additional information

Funding

Funder	Grant reference number	Author
Japan Science Society (JSS)	2022-4086	Junya Taniguchi
Japan Science Society (JSS)	2025-4060	Kano Kohara
MEXT Japan Society for the Promotion of Science (JSPS)	24H01444	Yuuki Kawabata
MEXT Japan Society for the Promotion of Science (JSPS)	19H04936	Yuuki Kawabata

Author ORCID iDs

Yuuki Kawabata:  <https://orcid.org/0000-0001-8267-5199>

References

- Alexander RM (2003) *Principles of animal locomotion* New Jersey, USA: Princeton University Press.
- Asakura A (2016) The evolution of mating systems in decapod crustaceans. In: Martin JW, Crandall KA, Felder DL (Eds). *Decapod crustacean phylogenetics* CRC Press. pp. 133-194
<https://doi.org/10.1201/9781420092592>
- Astudillo-Clavijo V, Arbour JH, López-Fernández H (2015) Selection towards different adaptive optima drove the early diversification of locomotor phenotypes in the radiation of Neotropical geophagine cichlids. *BMC Evol. Biol* **15**:77 <https://doi.org/10.1186/s12862-015-0348-7> | PubMed
- Atkinson RJA, Eastman LB (2015) Burrow dwelling in Crustacea. *The Natural History of the Crustacea* **2**:78-117
- Buatois LA, Carmona NB, Curran HA, Netto RG, Mángano MG, Wetzel A (2016) The Mesozoic Marine Revolution. In: Mángano MG, Buatois LA (Eds). *The Trace-Fossil Record of Major Evolutionary Events: Volume 2: Mesozoic and Cenozoic* Springer Netherlands Dordrecht. pp. 19-134
https://doi.org/10.1007/978-94-017-9597-5_2

- Burrer ED, Wainwright PC (2019) Adaptive radiation in labrid fishes: A central role for functional innovations during 65 My of relentless diversification. *Evolution* **73**:346-359 <https://doi.org/10.1111/evo.13670> | PubMed
- Chasen EM, Dietrich C, Backus EA, Cullen EM (2014) Potato leafhopper (Hemiptera: Cicadellidae) ecology and integrated pest management focused on Alfalfa. *Journal of Integrated Pest Management* **5**:A1-A8 <https://doi.org/10.1603/ipm13014>
- Chen Y, Grezmak JE, Graf NM, Daltorio KA (2022) Sideways crab-walking is faster and more efficient than forward walking for a hexapod robot. *Bioinspir Biomim* **17** <https://doi.org/10.1088/1748-3190/ac6847> | PubMed
- Crane J (2015) *Fiddler crabs of the world: Ocypodidae: genus Uca* Princeton University Press.
- de Gier W, Becker C (2020) A review of the ecomorphology of pinnotherine pea crabs (Brachyura: Pinnotheridae), with an updated list of symbiont-host associations. *Diversity* **12**:431 <https://doi.org/10.3390/d12110431>
- DecaNet eds (2025) DecaNet. Accessed 2025-09-16 <https://www.decanet.info>
- Domenici P (2010) *Fish locomotion: an eco-ethological perspective* CRC Press.
- Domenici P, Blagburn JM, Bacon JP (2011) Animal escapology I: Theoretical issues and emerging trends in escape trajectories. *J. Exp. Biol* **214**:2463-2473 <https://doi.org/10.1242/jeb.029652> | PubMed
- Fürsich FT, Jablonski D (1984) Late triassic naticid drillholes: Carnivorous gastropods gain a major adaptation but fail to radiate. *Science* **224**:78-80 <https://doi.org/10.1126/science.224.4644.78> | PubMed
- Faulkes Z (2006) The locomotor toolbox of the spanner crab, *Ranina ranina* (Brachyura, Raninidae). *Crustaceana* **79**:143-155 <https://doi.org/10.1163/156854006776952874>
- Feiner N, Jackson ISC, Stanley EL, Uller T (2021) Evolution of the locomotor skeleton in Anolis lizards reflects the interplay between ecological opportunity and phylogenetic inertia. *Nature Communications* **12**:1525 <https://doi.org/10.1038/s41467-021-21757-5> | PubMed
- Grimaldi D, Engel MS (2005) *Evolution of the Insects* Cambridge University Press.
- Hedrick BP, Dickson BV, Dumont ER, Pierce SE (2020) The evolutionary diversity of locomotor innovation in rodents is not linked to proximal limb morphology. *Sci. Rep* **10**:717 <https://doi.org/10.1038/s41598-019-57144-w> | PubMed
- Higham TE, Birn-Jeffery AV, Collins CE, Hulse CD, Russell AP (2015) Adaptive simplification and the evolution of gecko locomotion: Morphological and biomechanical consequences of losing adhesion. *Proceedings of the National Academy of Sciences* **112**:809-814 <https://doi.org/10.1073/pnas.1418979112> | PubMed
- Hultgren K, Stachowicz J (2009) Evolution of decoration in majoid crabs: a comparative phylogenetic analysis of the role of body size and alternative defensive strategies. *The American Naturalist* **173**:566-578 <https://doi.org/10.1086/597797> | PubMed
- Kawabata Y, Akada H, Shimatani KI, Nishihara GN, Kimura H, Nishiumi N, et al. (2023) Multiple preferred escape trajectories are explained by a geometric model incorporating prey's turn and predator attack endpoint. *eLife* **12**:e77699 <https://doi.org/10.7554/eLife.77699> | PubMed
- Keiler J, Wirkner CS, Richter S (2017) One hundred years of carcinization – the evolution of the crab-like habitus in Anomura (Arthropoda: Crustacea). *Biol. J. Linn. Soc* **121**:200-222 <https://doi.org/10.1093/biolinnean/blw031>
- Maddison WP, Midford PE, Otto SP (2007) Estimating a binary character's effect on speciation and extinction. *Syst. Biol* **56**:701-710 <https://doi.org/10.1080/10635150701607033>
- Miller AH (1949) Some ecologic and morphologic considerations in the evolution of higher taxonomic categories. *Ornithologie als biologische Wissenschaft* **84**

- Miller AH, Stroud JT, Losos JB (2023) The ecology and evolution of key innovations. *Trends Ecol. Evol* **38**:122-131 <https://doi.org/10.1016/j.tree.2022.09.005> | PubMed
- Morrison CL, Harvey AW, Lavery S, Tieu K, Huang Y, Cunningham CW (2002) Mitochondrial gene rearrangements confirm the parallel evolution of the crab-like form. *Proc. R. Soc. Lond. B* **269**:345-350 <https://doi.org/10.1098/rspb.2001.1886> | PubMed
- Murakami H, Tomaru T, Nishiyama Y, Moriyama T, Niizato T, Gunji Y.-P (2014) Emergent runaway into an avoidance area in a swarm of soldier crabs. *PLoS one* **9**:e97870 <https://doi.org/10.1371/journal.pone.0097870> | PubMed
- R Core Team (2023) R: The R project for statistical computing.
- Sato M, Wada K (2000) Resource utilization for decorating in three intertidal majid crabs (Brachyura: Majidae). *Mar. Biol* **137**:705-714 <https://doi.org/10.1007/s002270000389>
- Schoepfer SD, Algeo TJ, van de Schootbrugge B, Whiteside JH (2022) The Triassic– Jurassic transition – A review of environmental change at the dawn of modern life. *Earth-Sci. Rev* **232**:104099 <https://doi.org/10.1016/j.earscirev.2022.104099>
- Sleinis S, Silvey GE (1980) Locomotion in a forward walking crab. *Journal of comparative physiology* **136**:301-312 <https://doi.org/10.1007/bf00657350>
- Stroud JT, Losos JB (2016) Ecological opportunity and adaptive radiation. *Annual Review of Ecology, Evolution, and Systematics* **47**:507-532 <https://doi.org/10.1146/annurev-ecolsys-121415-032254>
- Takeshita F, Nishiumi N (2022) Social behaviors elevate predation risk in fiddler crabs: quantitative evidence from field observations. *Behav. Ecol. Sociobiol* **76**:162 <https://doi.org/10.1007/s00265-022-03268-5>
- Tan MH, Gan HM, Lee YP, Linton S, Grandjean F, Bartholomei-Santos ML, et al. (2018) ORDER within the chaos: Insights into phylogenetic relationships within the Anomura (Crustacea: Decapoda) from mitochondrial sequences and gene order rearrangements. *Mol. Phylogenet. Evol* **127**:320-331 <https://doi.org/10.1016/j.ympev.2018.05.015> | PubMed
- Tsang LM, Chan TY, Ah Yong ST, Chu KH (2011) Hermit to king, or hermit to all: multiple transitions to crab-like forms from hermit crab ancestors. *Syst. Biol* **60**:616-629 <https://doi.org/10.1093/sysbio/syr063> | PubMed
- Vidal-Gadea AG, Belanger JH (2013) The evolutionary transition to sideways-walking gaits in brachyurans was accompanied by a reduction in the number of motor neurons innervating proximal leg musculature. *Arthropod Struct. Dev* **42**:443-454 <https://doi.org/10.1016/j.asd.2013.07.003> | PubMed
- Vidal-Gadea AG, Rinehart MD, Belanger JH (2008) Skeletal adaptations for forwards and sideways walking in three species of decapod crustaceans. *Arthropod Struct. Dev* **37**:95-108 <https://doi.org/10.1016/j.asd.2007.06.002> | PubMed
- Wilcox JT (2017) Crab spider: grasslands predator hiding in plain sight. *Grasslands* 3-4
- Wolfe JM, Ballou L, Luque J, Watson-Zink VM, Ah Yong ST, Barido-Sottani J (2024) Convergent adaptation of true crabs (Decapoda: Brachyura) to a gradient of terrestrial environments. *et al Syst. Biol* **73**:247-262 <https://doi.org/10.1093/sysbio/syad066>
- Wolfe JM, Luque J, Bracken-Grissom HD (2021) How to become a crab: Phenotypic constraints on a recurring body plan. *BioEssays* e2100020 <https://doi.org/10.1002/bies.202100020> | PubMed
- Wolfe J, Ballou L, Luque J, Watson-Zink V, Ah Yong S, Barido-Sottani J, Chan T, Chu K, Crandall K, Daniels S, et al. (2023) Convergent adaptation of true crabs (Decapoda: Brachyura) to a gradient of terrestrial environments. Dryad Digital Repository. <https://doi.org/10.5061/dryad.tmpg4f52z>

Peer reviews

Reviewer #1 (Public review):

Summary:

This is an interesting and well-written manuscript in which the authors set out to answer a simple, longstanding question with a modern comparative approach. Namely where in crab evolution did sideways walking arise, how often has it been lost or regained, and is its evolution plausibly associated with the ecological and taxonomic success of true crabs. To address these questions the authors recorded locomotion from 50 live species, quantified the predominant direction of locomotion, and mapped these behavioral states onto a recent crab phylogeny to reconstruct the likely evolutionary history of sideways walking. The revised manuscript also includes analyses of the underlying movement-angle distributions and tests whether the evolutionary conclusions depend on the original behavioral classification scheme.

Strengths:

The strongest part of the study remains the dataset itself. Comparable behavioral measurements across dozens of crab species are rare, and obtaining and recording live representatives from this range of taxa required substantial field, aquarium, and husbandry effort. The overall pattern that emerges, in which most true crabs are strongly biased toward sideways locomotion while several specialized lineages move predominantly forward, is interesting and likely to be useful to researchers studying animal locomotion, functional morphology, and behavioral evolution.

The revised analyses substantially strengthened the manuscript. In the original version, I was concerned that the main behavioral classification depended too strongly on first assigning individual movements to forward or sideways bins using a fixed angular boundary. The authors have now analyzed the underlying continuous movement-angle distributions and have shown that, although mixed directional tendencies are present in some species, most taxa have a dominant directional preference. They also derived a separate, data-informed boundary from the distribution of dominant movement directions. I favor this alternative approach and it produces the same classification of species as the original index-based method, providing useful evidence that the main evolutionary reconstruction is not simply an artifact of the original 60{degree sign} cutoff.

The authors also responded appropriately to the limitation that locomotion was measured from one individual per species. This sampling design cannot establish the full extent of within-species, ontogenetic, or size-dependent variation, but the revised manuscript now states this limitation clearly and restricts its conclusions to broad interspecific patterns in predominant locomotor direction. This is a more appropriate interpretation of the available sampling.

The phylogenetic analysis provides a reasonable framework for addressing the main evolutionary question. Taken together, the behavioral and phylogenetic results support the conclusion that sideways locomotion likely arose once within the lineage leading to true crabs and was followed by multiple reversions toward predominantly forward locomotion in specialized groups. The manuscript therefore makes a convincing case that sideways walking is not simply an inevitable consequence of possessing a crab-like body plan.

Weaknesses:

My main remaining reservation concerns the interpretation of forward and sideways locomotion as two discrete biological modes. The revised analyses convincingly show that

species can be classified according to their predominant direction of locomotion and that this classification is robust to alternative analytical approaches. However, this does not necessarily demonstrate that forward and sideways locomotion represent two intrinsically discrete or mutually exclusive behavioral modes. Indeed, the new analyses show that many taxa are better described by two-component movement-angle distributions, even though most of these have one dominant component. This is consistent with strong directional preferences, but it also indicates that mixed movement strategies are common. The supplementary circular distributions similarly show considerable variation in the shape and breadth of directional preferences among taxa. Having said that, I do not think this substantially weakens the central evolutionary conclusion. The phylogenetic analysis requires a defensible classification of predominant locomotor direction, and the revised analyses now provide one. The evolutionary story remains interesting whether the underlying behavioral variation consists of two sharply discrete modes or a broader continuum of directional strategies with strong clustering toward forward and sideways movement.

A second limitation is that the proposed relationship between sideways locomotion and diversification remains necessarily correlational. The revised manuscript handles this more cautiously than the original version and now frames sideways locomotion as a plausible key innovation whose emergence is associated with the exceptional diversity of true crabs, rather than as a demonstrated causal driver of diversification. This distinction is important because differences in species richness among lineages can also reflect ecological opportunity, extinction history, and other lineage-specific factors. The revised framing is therefore better aligned with the strength of the evidence.

Final assessment: Overall, this is a valuable comparative study with an unusually broad behavioral dataset. The revisions have addressed the principal methodological concerns raised in the original review, particularly by analyzing continuous movement directions and demonstrating that the main phylogenetic classification is robust to an alternative, data-informed approach. The evidence now convincingly supports the central conclusion concerning the evolutionary origin and repeated reversal of predominant locomotor direction in crabs, although the stronger interpretation that forward and sideways locomotion represent two strictly discrete biological modes remains less certain.

<https://doi.org/10.7554/eLife.110015.2.sa2>

Reviewer #2 (Public review):

Summary:

The current work investigates the evolution of sideward locomotion in Brachyura in light of a single evolutionary origin. To this end, the authors first analysed the mode of locomotion in 50 crab species and observed mutually exclusive presence of sideways vs. forward movement. The phylogenetic analysis confirmed that there is indeed a single evolutionary origin for sideways movement, which was sometimes followed by several reversions to forward locomotion. This way, authors demonstrate how locomotor movement modes shape evolutionary diversification in animals by showing that species richness is much higher in side-ways-moving crabs than in the nearest groups. This is an interesting work that integrates behavioural analysis and phylogenetic relations, capitalising largely on crabs.

Original questions/suggestions:

Firstly, I think the paper spends too much time on a straightforward analysis of the mode of locomotion. I was also wondering whether the phylogenetic analysis could be simply achieved by maximising an objective function in which the modes of movement are inversely coded for two putative groups, with all values calculated at all possible nodes.

Unfortunately, I find that the authors did not sufficiently discuss differences in the ecological niches of species with forward vs. sideways locomotion modes (including challenges of locomotion and substrate).

Likewise, what are the anatomic correlates of forward vs. sideways locomotion? For instance, how are the advantages assumed for sideways movement associated with a flattened body? Is it possible that the mode of motion is secondary to flattened/narrow body structure, which basically limits the distance between legs and thus makes the forward movement difficult - under this logic, the mode of movement would be a secondary phenomenon to body shape traits. How can one differentiate between this alternative and the one that puts the mode of movement in the centre of the story? On a related note, how do different modes of movement relate to the ability to fit into tight spaces - how does it relate to differences in leg joints?

Is it possible that the sideways movement maximises the scanned visual field per unit time/displacement, which may be beneficial for mostly forward-moving predators?

Briefly, although I find the study interesting, the presented complexity may not be necessary given the endpoints; it can be achieved much more simply. Furthermore, the degree to which the conceptual analysis of different modes of locomotion was exercised was limited. The general approach may serve as a good model for the evolutionary analysis of other traits. The demonstration of traceability of the relations in question is a major contribution of the work.

Comment on revised version:

I am not fully convinced by the authors' handling of the complexity of the paper, but this seems like a moot point.

<https://doi.org/10.7554/eLife.110015.2.sa1>

Author response:

The following is the authors' response to the original reviews.

Public Reviews:

Reviewer #1 (Public review):

Summary:

This is an interesting and well-written manuscript in which the authors set out to answer a simple, old question with a modern toolkit: where in crab evolution did sideways walking arise, how often has it been lost or regained, and is it plausibly linked to the ecological and taxonomic success of true crabs. To do this, they record locomotion from 50 live species, convert each species' movements into a quantitative index that compares forward versus sideways bouts, and then map the resulting states onto a recent crab phylogeny to infer the most likely evolutionary history of locomotor direction.

We thank the reviewer for this positive summary of the study and for recognizing the value of our comparative behavioral dataset and phylogenetic approach.

Strengths:

The strongest part of the study is the dataset itself. Comparable behavioral measurements across dozens of crab species are rare. The authors have done the field and husbandry work needed to make this possible. The overall pattern they recover, that most true crabs are strongly biased toward sideways movement (while a smaller set of lineages move predominantly forward), is interesting and likely to be useful to others. The phylogenetic mapping is also a reasonable way to address the "how many times"

question (although this is peripheral to my expertise). The manuscript makes a convincing case that sideways locomotion is not simply a trivial byproduct of a crab-like body plan.

We appreciate the reviewer's recognition of the dataset and the overall value of the study. We have revised the manuscript to make the conclusions more robust and better aligned with the strength of the evidence.

(1) Where I am less convinced is in how strongly the authors describe the discreteness of the behavioral categories and the absence of intermediates. The manuscript states that the Forward-Sideways Index shows a clear separation between two locomotor types with little evidence for intermediates, and it cites a statistical test rejecting a single peak in the distribution. However, the histogram in Figure 3 appears structured within each labeled category, with subclusters inside both the forward and sideways groups rather than a single tight peak per group. This matters because the index is built by first placing each movement bout into "forward" versus "sideways" bins using a fixed angle boundary and then collapsing the result into a single ratio. That approach is simple and transparent enough, but it can also hide mixed strategies. For example, a species that produces substantial amounts of both forward and sideways walking can still end up with a strongly positive or negative index, and therefore be classified as a pure "type," even though the underlying behavior is mixed. In that context, rejecting a single peak in the across-species distribution does not, by itself, justify the stronger claim that intermediates are rare or absent.

Related to this, a key methodological choice is the use of 60 degrees as the cutoff between forward and sideways bouts. This boundary may be reasonable as a convention, but the paper does not explain why it is the right place to draw the line, and there is a plausible biological concern that a fixed angular cutoff does not mean the same thing across taxa.

Crabs vary in body shape and in how the legs are arranged around the body. In my own comparative work, for example, some species show an elliptical stance pattern elongated along the preferred direction of travel, while others show a more circular leg arrangement, and the latter can express more mixed forward and sideways behavior. When limb arrangement and body geometry differ across species, the same measured angle can correspond to different underlying mechanics and different functional "degree of sidewaysness." The practical implication is that the reported binary separation may partly reflect the imposed classification rule, rather than a sharp biological divide.

We thank the reviewer for this important point. We agree that the across-species distribution of FSI values alone does not justify a strong statement that intermediate or mixed locomotor tendencies are absent. We also agree that reducing continuous bout-angle distributions to a single index could potentially obscure mixed directional strategies. We have therefore revised the manuscript to avoid implying a strict absence of intermediates and have added an additional analysis of the underlying continuous angle distributions (Abstract, lines 27-29; Results, lines 191-208; Table S2).

Specifically, we fitted one- and two-component mixture models to the continuous bout-angle distributions of each taxon and examined the supported number of components, peak locations, and mixture weights (Results, lines 196-204; Table S2). This analysis showed that 14 taxa were best described by a one-component model, whereas 36 taxa were best described by a two-component model. Importantly, among the 36 taxa best described by a two-component model, 33 had a dominant component explaining at least 70% of the distribution, whereas only three taxa showed relatively balanced two-component distributions. Thus, although some taxa do show mixed directional tendencies, most taxa are dominated by a primary

directional component rather than showing an even mixture of forward and sideways locomotion.

We also clarified the rationale for the 60° threshold used in the FSI calculation (Methods, lines 125-130). This threshold was not intended to represent a taxon-specific biological boundary between forward and sideways locomotion. Rather, it was used to divide the 360° space into three equal directional sectors: forward, sideways, and backward. This equal partitioning provides a consistent reference under a null expectation of uniformly distributed movement directions.

To further assess whether our classification depended on the original FSI-based classification, we performed an additional data-driven check based on continuous bout-angle distributions (Results, lines 204-208; Fig. S2). We extracted the dominant peak location from each taxon's continuous bout-angle distribution (Table S2) and estimated a boundary from the distribution of these dominant peak locations using a Gaussian mixture model. This yielded a data-informed cutoff of approximately 49.4°. The resulting peak-based classification was identical to the original FSI-based classification, with 15 forward-moving and 35 sideways-moving taxa. Thus, no taxon changed category under this independent classification approach.

(2) Another limitation that affects interpretation is the decision to use one individual per species. I understand the logistics, and for some questions, a single representative individual can be a reasonable first pass. But it is not strong support for negative claims about intermediates, especially in a group where individuals can change substantially with growth and allometry. Crabs can grow dramatically, often with pronounced allometric shifts in limb proportions that can alter the center of mass location. Size alone can alter the kinematics and choice of locomotor behaviors in crustaceans. In species where appendage proportions change with size, or where certain legs become disproportionately large (or calcified), it is plausible that locomotor direction and the distribution of movement angles shift across ontogeny. That makes it hard to treat a single individual as a complete description of a species-level strategy, particularly for species that fall closer to the boundary between categories.

We thank the reviewer for raising this important limitation. We agree that using one representative individual per species cannot capture the full range of within-species variation, including ontogenetic, size-dependent, or allometric changes in locomotor behavior. We also agree that this limitation is particularly relevant to strong claims about the absence of intermediates.

As described in our response to Comment #1, we have therefore toned down statements implying a strict absence of intermediates and added analyses of the underlying continuous angle distributions (Abstract, lines 27-29; Results, lines 191-208; Table S2). These additional analyses showed that some taxa do exhibit mixed directional tendencies, although most taxa were dominated by a primary directional component.

We have also revised the manuscript to clarify the scope of our conclusions. Specifically, we now state that our single-individual sampling design does not capture possible ontogenetic, size-dependent, or allometric variation within species (Methods, lines 105-107). We also clarify that our conclusions are intended to identify broad interspecific patterns in the predominant direction of locomotion across major brachyuran lineages, rather than to describe the full range of locomotor variation within each species (Methods, lines 107-108). Thus, we no longer treat a single individual as providing a complete description of species-level behavioral variation, but instead use it as a standardized representative observation for broad comparative and phylogenetic analyses.

In sum, this is a valuable and useful behavioral comparative study with a dataset that many in the field will appreciate. The main conclusions about the likely evolutionary

placement of sideways walking are plausible, but several of the stronger claims about discrete locomotor types, the absence of intermediates, and the relationship to diversification would be more convincing if the analysis were less dependent on a fixed angular cutoff and on single individuals per species, or if the manuscript framed those points more cautiously so the conclusions track the strength of the evidence.

We thank the reviewer for this constructive summary and for recognizing the value of our behavioral comparative dataset. We have addressed these concerns in detail in our responses above and revised the manuscript to make the main claims better aligned with the strength of the evidence.

Reviewer #2 (Public review):

Summary:

The current work investigates the evolution of sideward locomotion in Brachyura in light of a single evolutionary origin. To this end, the authors first analysed the mode of locomotion in 50 crab species and observed mutually exclusive presence of sideways vs. forward movement. The phylogenetic analysis confirmed that there is indeed a single evolutionary origin for sideways movement, which was sometimes followed by several reversions to forward locomotion. This way, authors demonstrate how locomotor movement modes shape evolutionary diversification in animals by showing that species richness is much higher in side-ways-moving crabs than in the nearest groups. This is an interesting work that integrates behavioural analysis and phylogenetic relations, capitalising largely on crabs. I have a few suggestions and questions.

We thank the reviewer for the positive assessment of the study and for recognizing the value of integrating behavioral analysis with phylogenetic relationships. We address the specific suggestions and questions below.

(1) Firstly, I think the paper spends too much time on a straightforward analysis of the mode of locomotion.

We agree that the final classification of taxa into predominantly forward- and sideways-moving groups is conceptually simple. However, because our study compares locomotor behavior across a broad range of crab taxa and then uses these behavioral data for phylogenetic reconstruction, we considered it important to describe the behavioral quantification in a transparent and reproducible way. The purpose of this section is therefore not to make a simple endpoint unnecessarily complex, but to show how discrete locomotor states were derived from raw trajectory data using a standardized procedure. For this reason, we retained the current analytical description.

(2) I was also wondering whether the phylogenetic analysis could be simply achieved by maximising an objective function in which the modes of movement are inversely coded for two putative groups, with all values calculated at all possible nodes.

The proposed objective-function approach may be useful for identifying a node that best separates two predefined locomotor groups. However, in the present study, we aimed not only to locate a possible boundary between forward- and sideways-moving lineages, but also to reconstruct the evolutionary history of locomotor transitions under an explicit phylogenetic model.

For this reason, we used standard ancestral state reconstruction and stochastic character mapping rather than maximizing an ad hoc objective function across possible nodes. This approach allowed us to compare alternative transition-rate models (ER and ARD), estimate uncertainty in ancestral states at internal nodes, and quantify the posterior distribution of gains and reversals. We therefore retained the current phylogenetic framework, as it

provides a model-based and more informative reconstruction of locomotor evolution across true crabs.

(3) Unfortunately, I find that the authors did not sufficiently discuss differences in the ecological niches of species with forward vs. sideways locomotion modes (including challenges of locomotion and substrate).

Likewise, what are the anatomic correlates of forward vs. sideways locomotion? For instance, how are the advantages assumed for sideways movement associated with a flattened body? Is it possible that the mode of motion is secondary to flattened/narrow body structure, which basically limits the distance between legs and thus makes the forward movement difficult - under this logic, the mode of movement would be a secondary phenomenon to body shape traits. How can one differentiate between this alternative and the one that puts the mode of movement in the centre of the story? On a related note, how do different modes of movement relate to the ability to fit into tight spaces - how does it relate to differences in leg joints?

Is it possible that the sideways movement maximises the scanned visual field per unit time/displacement, which may be beneficial for mostly forward-moving predators?

We thank the reviewer for this helpful comment. We agree that the previous version did not sufficiently address the possible relationship between locomotor mode and body shape, especially the alternative explanation that sideways locomotion may be secondary to carapace flattening. In response, we added a new morphological analysis using two carapace shape indices: relative carapace length (CL/CW) and relative carapace depth (CD/CS) (Methods, lines 178–185). In the revised Results, we report that relative carapace length differed significantly between forward- and sideways-moving taxa (phylogenetically informed ANOVA: $F = 26.90$, $p < 0.001$), whereas relative carapace depth did not differ significantly between the two groups ($F = 1.18$, $p = 0.403$) (Results, lines 209–214; Fig. S3). We also added this interpretation to the Discussion, noting that locomotor mode is associated with some aspects of carapace shape but is not explained by simple carapace flattening alone (Discussion, lines 318–325).

We also revised the Discussion to address the reviewer's suggestions about possible functional advantages of sideways locomotion beyond rapid bidirectional escape. Specifically, we now mention that other possible advantages may include movement through confined spaces and visual-field sampling during locomotion (Discussion, lines 310–318).

Finally, we retained the existing discussion of ecological specializations in forward-moving lineages, including coordinated collective movement in soldier crabs, decoration and concealment in majoid crabs, and life inside confined host spaces in pea crabs. This discussion supports the broader point that the adaptive value of sideways locomotion may depend on ecological context.

(4) It is really difficult to decipher the information contained in the nodes (circles) in the printed black-and-white version of the manuscript.

We have changed the color scheme and strengthened the outlines of the node pie charts so that the ancestral-state probabilities can be more easily distinguished (Fig. 5). We also applied the same revised color scheme to Figure 3 and Figure S4 for consistency across the manuscript.

(5) Briefly, although I find the study interesting, the presented complexity may not be necessary given the endpoints; it can be achieved much more simply. Furthermore, the degree to which the conceptual analysis of different modes of locomotion was exercised was limited. The general approach may serve as a good model for the evolutionary

analysis of other traits. The demonstration of traceability of the relations in question is a major contribution of the work.

We thank the reviewer for this constructive summary and for recognizing the broader value of our approach. We have addressed the methodological and conceptual points raised here in our responses to the specific comments above.

Strengths:

The research question and the novel combination of different data types.

We thank the reviewer for highlighting the research question and the novel combination of different data types as strengths of the study. We have revised the manuscript to further strengthen this integrative framework.

Weaknesses:

The complexity of the methods used, along with a limited discussion of the potential dynamics that may underlie the evolution of the sideways movement mode.

We have addressed these concerns in our responses to the specific comments above, particularly by clarifying the rationale for the behavioral quantification and expanding the discussion of morphology, ecological context, and functional hypotheses.

Recommendations for the authors:

Reviewer #1 (Recommendations for the authors):

(1) Unbiased analysis of angle distribution. The authors already extract continuous bout angles prior to binning. I recommend using these distributions directly to assess modality at the species level (e.g., unimodal vs bimodal, peak locations, mixture weights) before collapsing behavior into the Forward-Sideways Index. Even a simple circular density estimate or mixture model would clarify whether species classified as "forward" or "sideways" are behaviorally pure or mixed, and would provide a quantitative basis for claims about intermediacy.

We added mixture-model analyses of the continuous bout-angle distributions, including modality, peak locations, and mixture weights (Results, lines 196-204; Table S2).

(2) Justify or stress-test the 60° cutoff. The manuscript should either provide a clear biological or data-driven justification for using 60° as the boundary between forward and sideways bouts, or demonstrate that the main conclusions are robust to reasonable alternative cutoffs (e.g., 45°, 75°). A brief sensitivity analysis in the supplement would be sufficient and would greatly strengthen confidence in the classification. Alternatively (my preference) would be to let the data inform the cutoff.

We clarified the rationale for the 60° sector definition used to calculate FSI (Methods, lines 125-130; Fig. 2) and added an independent data-driven boundary analysis based on dominant peak locations, which yielded the same forward/sideways classification (Results, lines 204-208; Fig. S2).

(3) Sampling justification. I recommend explicitly acknowledging that sampling a single individual per species limits the ability to detect ontogenetic, size-dependent, or allometric variation in locomotor strategy. If feasible, adding even limited replication across size classes or individuals for a small subset of taxa (particularly those near the classification boundary) would substantially strengthen the conclusions; otherwise, the manuscript should more clearly delimit which claims do and do not rely on the assumption of within-species invariance.

We clarified that our conclusions concern broad interspecific patterns of predominant locomotor direction, rather than the full range of within-species variation (Methods, lines 102-108).

(4) I would suggest toning down or reframing statements about "no intermediates". If additional analyses are not added, I recommend revising statements that imply a strict absence of intermediates to language that reflects what is directly shown (e.g., bimodality in an index derived from binned data). This would better align the claims with the current evidence.

We revised the manuscript to avoid implying a strict absence of intermediates and now acknowledge that some taxa show mixed directional tendencies (Abstract, lines 27-29; Results, lines 191-208).

(5) Framing and claims about diversification. The discussion of sideways locomotion as a key innovation would benefit from clearer separation between observed correlations and causal inference. If trait-dependent diversification analyses are not added, I suggest consistently framing this section as a hypothesis supported by comparative patterns rather than a demonstrated mechanism.

We revised the Discussion to more clearly frame sideways locomotion as a possible key innovation associated with diversification, rather than as a demonstrated causal mechanism (Abstract, lines 31-34; Discussion, lines 294-309).

<https://doi.org/10.7554/eLife.110015.2.sa0>