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Collective dynamics in entangled worm and robot blobs

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Living systems at all scales aggregate in large numbers for a variety of functions including mating, predation, and survival. The majority of such systems consist of unconnected individuals that collectively flock, school, or swarm. However, some aggregations involve physically entangled individuals, which can confer emergent mechanofunctional material properties to the collective. Here, we study in laboratory experiments and rationalize in theoretical and robophysical models the dynamics of physically entangled and motile self-assemblies of 1-cm-long California blackworms (Lumbriculus variegatus) and C. elegans. Thousands of individual worms form braids with their long, slender, and flexible bodies to make a three-dimensional, soft, and shape-shifting “blob.” The blob behaves as a living material capable of mitigating damage and assault from environmental stresses through dynamic shape transformations, including minimizing survival area for survival against desiccation and enabling transport (negative thermotaxis) from hazardous environments (like heat). We specifically focus on the locomotion of the blob to understand how an amorphous entangled ball of worms can break symmetry to move across a substrate. We hypothesize that the collective blob displays rudimentary differentiation of function across itself, which when combined with entanglement dynamics facilitates directed persistent blob locomotion. To test this, we develop a robophysical model of the worm blobs, which displays emergent locomotion in the collective without sophisticated control or programming of any individual robot. The emergent dynamics of the living functional blob and robophysical model can inform the design of additional classes of adaptive mecanofunctional living materials and emergent robotics.

Significance

Living organisms form collectives across all scales, enabling biological functions not accessible by individuals alone. In a few cases, the individuals are physically connected to each other, forming an additional class of entangled active matter systems with emergent mechanofunctionalities of the collective. Here, we describe the dynamics of macroscopic aquatic worms that braid their long, soft bodies to form large entangled worm blobs. We discover that the worm blob behaves as a living material to undergo dynamic shape transformations to reduce evaporation or break-symmetry and locomote to safety against thermal stresses. We validate our biological hypotheses in robophysical swarming blobs, which pave the way for additional classes of mecanofunctional active matter systems and collective emergent robotics.


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and how they spontaneously move as whole. By developing roboticphysical blobs (39, 40) consisting of three-link robots (smarticles) (38), we describe how variation of gaits and mobility of simple individuals in a physically entangled collective can lead to varying levels of locomotive performance, without the need for sophisticated central control.

Results and Discussion
Worm Blob as Complex Material. Under certain environmental conditions (evaporation, cold temperature, etc.) individual worms spontaneously aggregate to form three-dimensional blobs that range from small collectives \((N = 10 \text{ worms})\) to large macroscopic entangled networks \((N = 50,000)\), both in water and in air (Fig. 1 and Movie S1). The blobs display non-Newtonian material properties (41) and can flow at long timescales (SI Appendix, Fig. S1D), while retaining shape to short timescale disturbances (SI Appendix, Fig. S1 A and C). The underlying material timescale is set by the physical entanglement of the worm bodies through self-braiding (Fig. 1). The blob formation and disintegration are reversible through changes in ambient fluid temperature (SI Appendix, Fig. S1B). Increasing the fluid temperature “melts” the blob (fluid-like phase), while decreasing the temperature leads to a more entangled, solid-like phase (SI Appendix, Fig. S1B). We posit that the underlying principle for this emergent collective phase behavior lies in the changing activity of individual cold-blooded worms in response to external temperature and cooperative effects of active phase domains (41–46). At lower temperatures \((T < 25^\circ \text{C})\), worms are less motile and elongated, while at higher temperatures \((25 < T \leq 30^\circ \text{C})\), they are more active and coiled up, with a peak activity at \(T = 30 \pm 2^\circ \text{C}\) (SI Appendix, Fig. S5) (47). Thus, an entangled solid-like blob is observed at lower temperatures through slow-moving, elongated worms, while a disentangled fluid-like aggregate is observed at higher temperatures (SI Appendix, Fig. S2) due to highly active, coiled-up worms, similar to motility-induced phase separations in active matter (46, 48).

Worms in a Blob Survive Longer against Desiccation. Worms are cold-blooded animals and their activities are greatly influenced by variations in temperature and light intensity of their surroundings (47, 49). Since blackworms naturally inhabit shallow aquatic regions (42–44), we hypothesize that forming a blob provides survival benefit to individual worms by reducing desiccation in air. To test this hypothesis, we expose blobs of different numbers \((N = 1 \text{ to } 1,000 \text{ worms}, 10 \text{ replicates per condition})\) on a dry plate at controlled temperature and humidity \((24 \pm 5^\circ \text{C}, 48\%\) and track the projected blob area, \(A\), using time-lapse imaging over a few hours (SI Appendix, Fig. S4 and Movie S2). We observe that a single worm \((N = 1)\) perishes in less than 1 h, while worms in a blob \((N = 1,000)\) are alive even after 10 h (Movie S2). Additionally, the blobs are not static. Larger blobs \((N > 20)\) undergo exploratory searches for potentially favorable conditions, during which the projected blob area can almost double (40 min, \(N = 100)\) (Fig. 2A and B and Movie S2). This phase is followed by a shrinking mode during which the blobs become increasingly hemispherical to minimize surface area to volume ratio. In contrast, smaller blobs \((N < 20)\) monotonically shrink into hemispherical structures (Fig. 2B). We note that this dynamic shape transformation behavior of worm blobs is reversible: Once the blob has shrunk into a spherical shape, addition of water restarts the search mode (Movie S2).

To support the hypothesis that the worm blobs reduce evaporative losses through surface area minimization, we quantify the steady-state projected surface area \((A_{\text{ss}}\), defined as the projected surface area is invariant, i.e., \(\text{d}A/\text{d}t < 1\%\) as a function of blob size \(N\) as shown in Fig. 2 C and D. Theoretically, we estimate the volume of \(N\) worms as \(V_w = N \times (\pi r_w^2 h)\), where each worm is assumed to have a cylindrical body of radius \(r_w\) and length \(h\). If these \(N\) worms minimized surface area by bending and coiling to form a hemisphere, then the theoretical projected surface area of the hemisphere (i.e., area of the circle) is given by \(A_{\text{th}} = \pi r_b^2\), where \(r_b = \sqrt{\frac{3V_w}{4\pi}}\) is the equivalent blob radius. Note that, since the worms have a hydrostatic skeleton, the radius of worms \((r_w)\) decreases with evaporation from their nominal value 0.6 ± 0.1 to 0.3 ± 0.1 mm (SI Appendix, Fig. S7). Indeed, the experimental data and theoretical model are in good agreement over three orders of blob size \(N\) between the ranges of evaporated worm quantities.

Fig. 1. Worm blobs form via physical entanglements. (A) An entangled worm blob composed of \(~50,000 \text{ worms})\). (B) The worm blob behaves as a non-Newtonian fluid, which can flow at long timescales and maintain shape as a solid at short timescales (Movie S1). (C and D) Blob formation in water. The experiment starts in water at \(~30^\circ \text{C}\) in which the worms are mainly untangled with each other. As the water cools down to \(25^\circ \text{C}\), the worms aggregate initially into two smaller blobs \((t = 1 \text{ min})\), which ultimately merge to form one large blob \((t = 20 \text{ min}; \text{Movie S1})\). (E) Close view of braid formation within a blob.
radius, validating the hypothesis that worm blobs form hemispherical shapes to reduce evaporation losses (Fig. 2 C and D). Beyond *L. variegatus*, other annelids that we tested, such as *Lumbricus terrestris* and *Eisenia fetida* (*SI Appendix*, Fig. S6 and Movie S2), as well as past observations on nematodes (*Caenorhabditis elegans*) (50–54), suggest forming entangled collectives may be a general biological strategy to survive desiccation for extended periods by these organisms in fluctuating arid environments.

**Emergent Locomotion of Worm Blobs.** Since the worms are sensitive to temperature (as described earlier) as well as light (42, 45), we next investigate the emergent behavior of the worm blobs in response to a combination of light and temperature cues in a custom setup shown in Fig. 3 A (SI Appendix, Fig. S8). In laboratory cultures at constant temperature (~15 °C), we observe that the worms form tightly entangled blobs at high light intensities and form loose dispersions in the dark. A sudden increase in light intensity results in a rapid blob contraction (33 ± 6% reduction, after 15 h in dark) of a short duration (t < 5 s; SI Appendix, Fig. S9 and Movie S3), which is expected as individual worms are known to exhibit a rapid contractile escape response to shadows and photic stimulations arising from movements of overhead predators in the water (55). Thus, bright light serves as a cue for worms to aggregate and entangle tightly.

Under room light intensity (~400 lux) if we next expose the blob to an approximately linear temperature gradient (see the experimental setup in Fig. 3 A), the blob dissipates and worms individually crawl to the cold side (Fig. 3 B, Left column; SI Appendix, Fig. S10; and Movie S3). Without changing the temperature gradient when we increase the light brightness to 5,500 lux, we discover a surprising behavior: Worms collectively move as a blob toward the cold side (negative thermotaxis; Fig. 3 B, Right column; SI Appendix, Fig. S11; and Movie S3). In both cases, the majority of the worms (>70% for 400 lux and >90% for 5,500 lux) are successfully able to move toward the colder side, with a similar average speed (0.6 ± 0.1 cm/min, calculated when 70% of worms reach cold side) as shown in Fig. 3 C.

To examine whether moving together as a blob vs. moving individually conferred additional benefits, we utilize a quasi–two-dimensional (quasi-2D) experimental apparatus, which facilitates worm tracking (SI Appendix, Fig. S12 A and B). We find that worms crawling individually can move at faster speed to the cold side (N = 10, v<sub>i</sub> = 1.2 ± 0.3 cm/min) compared to worms in a blob (N = 80, v<sub>b</sub> = 0.4 ± 0.1 cm/min). However, moving as a blob despite being slow enabled all of the worms to be transported safely to the cold side. Not all worms moved safely when they crawled individually (SI Appendix, Fig. S12 C). Thus, for an individual worm, being in a blob confers multiple survival benefits: reduced evaporation when water is scarce and reliable transport to safety when the environmental temperature becomes fatal.

**Mechanism of Blob Locomotion.** How does an entangled worm blob spontaneously break symmetry and move? We hypothesize that the locomotion of the blob toward the cold side emerges as a consequence of the individual worm response to temperature. To test this hypothesis, we recorded close-up videos of small worm blobs (N = 20) while moving under a thermal gradient (Fig. 4 A). Using their circular and longitudinal muscles, the worms can apply contractile pulling forces along the length of

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**Fig. 2.** Water evaporation response of the worm blob. (A) When water is scarce, worms spontaneously form hemispherical blobs as a survival strategy to minimize evaporative losses. Shown is a time snapshot from the experiment (side view, N = 100 worms) for 450 min (Movie S2). The worms first undergo a stereotypical search mode for a water source and, after a certain time, spontaneously transition into a shrink mode to reduce surface area. (B) The shape changes of the worm blobs in the air as a function of blob size (N = 5, 10, 20, 100, 1,000). See Movie S2 for the example experiments with N = 1 and N = 1,000. (C) Projected surface area (A) as a function of cluster size (N = 1, 5, 10, 20 100, and 1,000 worms, 10 replicates per condition) under controlled laboratory conditions (24 °C, humidity 48%). The red star on the light blue curve indicates the time when the shrink mode starts. The worm blobs achieve a steady-state projected surface area (A<sub>s</sub>) (black) with theoretical estimation of surface area (red) across three orders of magnitude of blob size (N) reveals good agreement between model and experiments (SI Appendix, Fig. S7).
their body to crawl forward (56). As described earlier, the motility (and activity) of the worms increases at higher temperatures (SI Appendix, Fig. S5) (47). Thus, depending on the position in a blob (front, rear, or top), individual worms encounter different thermal stimuli. We observe that worms at the top of the blob act as entangling binders to keep the blob as a cohesive unit. Worms facing the cold side (leading edge) act as pullers and use their elongated bodies to apply slow, periodic pulling forces on the blob (Movie S4 and Fig. 4 B and C). The worms closer to the hot side (trailing edge) are more coiled up, lifting up the back of the blob potentially to reduce friction (wrigglers, Fig. 4 A–C).

To explore this hypothesis further, we first evaluate whether the pulling/wiggling behaviors are correlated with blob displacement. We plot the contraction in length of the puller worms $\Delta L = f(t_f) - f(t_i)$ with the forward displacement of the center of blob $\Delta x = x_c(t_f) - x_c(t_i)$, where $t_i$ and $t_f$ represent the initial and final times (Fig. 4D). Each symbol represents an event where a forward blob movement occurred ($x_c > 1 \text{ mm}$) for a total 17 events from $n = 3$ trials each of $N = 20$ worm blobs (Fig. 4D). We observe that in the majority of the cases (13/17 events), the forward blob displacement $\Delta x$ is correlated with both a pulling event (negative $\Delta L$ due to contraction of puller worms) and vertical lifting of the blob by the wiggler worms (positive $y_i$). Thus, our data suggest that functional differentiation of worms in a blob into puller worms at the leading edge and friction-reducing worms (wriggling) is a possible mechanism for blob motility. For example, the pull events are visually evident in the accompanying videos (Movie S4) as well as in jumps in blob displacement during locomotion of small blobs ($N = 20$) across a substrate (Fig. 4E).

However, since each worm can individually exhibit a wide array of dynamic locomotor behaviors (42, 45, 47), the collective blob locomotion may not be limited to just a binary gait differentiation; more complex mechanisms and emergent cooperative behaviors that build upon this motif are possible. For example, in Fig. 4D there are a few points (4/17) where although the leading puller worms did not contract (positive $\Delta L$), the blob still exhibits significant forward displacement. A closer look at one of these events (point 4 in Fig. 4D) reveals that a few puller worms intertwine to form a braided chain and collectively pull together (Movie S6). Moreover, as the blobs become larger ($N > 300$), we observe that the blob moves at a slower speed compared to smaller blobs ($N = 20$), but the movement becomes more consistent compared to the relatively jerky pull events observed in smaller blobs (Fig. 4E). Visualizing the exact behavior (puller vs. wiggler) in these larger blobs is challenging due to the opacity and three-dimensional entanglement in these blobs (Movie S4). Thus, although we propose and provide evidence below (using robotic blobs) for a simplistic mechanism for blob motility through gait differentiation by the puller and wiggler worms, we believe that more complex behaviors are possible and yet to be discovered that may further depend on the blob size, external stimulus, and substrate properties.

To support our proposed basic mechanism and test whether a single worm can indeed generate the necessary force to pull a blob, we tether individual worms to a calibrated cantilever beam and measure the pulling force exerted by a single worm on a rigid peg (Fig. 4F and SI Appendix, Fig. S13). At low temperatures ($20 ^\circ C$), the worms are more elongated and use their bodies to exert large pulling forces ($F_p = 178.2 \pm 52.5 \mu N$, $\sim 2.5$ times the weight of a single worm). While at high temperatures ($30 ^\circ C$) the forces were nominal $F_p = 28.0 \pm 10.9 \mu N$ (Fig. 4G and H).
Collective dynamics in entangled worm and robot blobs

Fig. 4. Mechanism of emergent locomotion of a worm blob through differentiation of activity. (A) Schematic of the experimental setup with enforced temperature gradient. The black dot shows the center of blob (CoB). (B) Snapshots from the quasi-2D experiment, where a small worm blob \((N = 20)\) exhibits negative thermotaxis, crawling toward the cold side (right) in response to a temperature gradient (Movie S4). (C) Proposed mechanism of how an entangled worm blob breaks symmetry to exhibit directed motion. The worm blob moves through differentiation of activity: Worms at the leading edge (blue) act as pullers, while worms at the trailing edge (red) curl up and lift the blob to potentially reduce friction. \(\Delta x\) is the contraction amount of leading worms and \(\Delta x_c\) is the forward displacement of a worm blob. (D) Correlation plot of changes in the length of the leading-edge puller worms, \(\Delta L = l(t_f) - l(t_i)\), and the blob’s forward position, \(\Delta x_c = x_c(t_f) - x_c(t_i)\), where \(t_i\) and \(t_f\) represents the initial and final times of the peak forward movement (total 17 events from \(N = 3\) trials). Color represents the maximum height of the blob (max \(y_c\)) during the forward displacement. (E) Horizontal displacement of the center of blob for \(N = 20\) (dark and light purple) and \(N > 300\) (dark and light yellow) in response to thermal gradients. Red dots on the \(N = 20\) case indicate where the pulling events begin (the events [1–7] and [8–12] are shown in D) (Movie S4). (F) Schematic of the experimental setup to measure pulling force of individual worms. The pegs are mounted on a plastic petri dish (100 \(\times\) 15 mm) and the tail of the worm is glued to a force-calibrated elastic beam (SI Appendix, Fig. S13). By measuring the deflection of the beam by the worms, pulling force is estimated. (G) Illustration describing the observed behavior of worms during measurements at 20 °C (blue) and 30 °C (red) as shown in Movie S4. (H) Force measurements for single worms in cold (20 °C, blue) and hot water (30 °C, red). The black dots on the blue curve indicate the start time of successive pulling events by worms as seen in Movie S4. Inset shows the mean and standard deviation of the maximum pulling forces in cold (five trials) and hot water (three trials).

At higher temperatures, we hypothesize that the worms cannot apply enough force to pull the pegs due to rapid body movements and gait reversal at elevated temperatures (47). Assuming a low coefficient of static friction for wet acrylic (\(\mu_s = 0.3\), substrate used in our experiments) and a blob mass of \(m = 0.14\) g \((N = 20\) worms), we estimate two to three individual worms could generate sufficient traction force \((F = \mu_s mg = 412\) \(\mu\)N) to move a small blob under certain conditions (low friction, existence of a rigid object to grab and pull the body, etc.). This estimate qualitatively agrees with experimental observations, where individual worms are capable of moving the small blob (Fig. 4C and Movie S4). Taken together, we posit that the locomotion in small blobs can emerge through differential activities of the individuals in the front (puller) and rear (wiggler).

**Phototactic Robophysical Blobs.** Here we develop an active physical model (a “robophysical” model) (39, 40) of the worm blob to validate our hypothesis that an entangled robotic collective can exhibit emergent locomotion through two central principles: mechanical interactions (entanglements) and function (gait) differentiation. The advantage of a robophysical model is that it enables us to examine critical parameters that cannot easily be observed in the biological system and serves as a laboratory platform for testing the individual–environment interactions that could lead to real-world swarming robot collectives.

The robophysical blobs consist of six three-link, two-revolute joints, with planar, smart, active particles (smarticles) (38) equipped with two light sensors as shown in Fig. 5A. Based on the light intensity sensed by these optical sensors, we program robots with three autonomous behaviors as a function of their arm angles \(\alpha_1\) and \(\alpha_2\) (Fig. 5B and Movie S5): 1) For low light intensity (<200 lux), robots execute a wiggle gait, where \{\(\alpha_1, \alpha_2\) = \[−\pi/4, −\pi/4\], \(\pi/4, \pi/4\)\] such that the robot arms swing up and down (out of phase with each other) \(45°\) from the centerline of the body (Fig. 5D). 2) For high light intensity (>800 lux), robots follow a crawl gait, where \(\alpha_1\) and \(\alpha_2\) follow a sequence as shown in Fig. 5B. 3) For intermediate light (200 to 800 lux), robots can either hold their arms straight (rigid) with \(\alpha_1; \alpha_2 = 0\) or let their arms freely move (flexible) and deform based on neighboring interactions. We note that only the crawl gait leads to forward motion (symmetry breaking) toward
Collective Dynamics in Robophysical Blobs. We first demonstrate that strong physical entanglement among individuals is critical for collective locomotion. In previous work (38), robots achieved collective mobility only when they were forced to weakly entangle by an external ring structure. Here, we test the same robots—without any external ring—and since there is little traction (or entanglement) among individuals, it is not possible to create a cohesive robotic blob. Thus, when the robots undergo phototaxis through purely crawl or wiggle gaits, no collective locomotion is observed (SI Appendix, Fig. S14 and Movie S5). To achieve strong entanglements while allowing for ready disentanglement—reflecting the capabilities of the worm blobs—we add a mesh and L-shape pins on the robot arms (Fig. 5C). The size (width = 0.5 cm, length = 1 cm), shape, and orientation of the pins are chosen so that the arms easily attach to and detach from the mesh and form an entangled robotic blob mimicking the worm blobs (see Fig. 5D). For the rest of the robophysical experiments, we use these self-entangling and disentangling robots.

Next, we investigate the collective phototaxing behavior of the robophysical blobs in a quasi-2D confined arena (length = 60 cm, width = 5 cm). We cover the bottom surface of the arena with a mesh to increase substrate friction and place a light source at one end (SI Appendix, Fig. S15A). We expect that the leading robot (closer to the light source, high light intensity) would receive the most light and achieve a crawl gait, while the rearward robots (farther away from light, low light intensity) would execute a wiggle gait. The middle robots that receive intermediate light intensity would keep their arms rigid, to create a cohesively moving robotic blob. Surprisingly however, we observe that the robotic blob slowly disentangles and the robots move individually toward the light, rather than moving as a group (Fig. 64). By tracking the arm positions of each robot in a blob using an automated tracking software (DeepLabCut) (57), we observe that although the crawler robots and wiggler robots behave as expected, the intermediate light robots through their rigidity diminish effective force transmission within the blob, leading to disentanglement (SI Appendix, Fig. S16). Once the blob has disentangled, these intermediate robots also change to a crawling gait (no longer shaded from light in the blob) and move individually toward the light source (Movie S5).

To address this issue of disentanglement, we simply switch off a subset of the intermediate light intensity robots to allow them to freely deform their arms (flexible) in response to neighboring robot arm movements (Fig. 6B). Now, the inactive robots (not powered) act as passive and compliant “binders,” enabling the blob to move as a collective, similar to their biological counterparts (Fig. 6C, Right; SI Appendix, Fig. S15; Movie S5 and Table 1). We note that the remaining robots are active (powered) and change their gaits according to the light intensity as described previously (if >900 crawl, <900 wiggle).

Further analysis of the robophysical blob dynamics reveals little change in connectivity (defined as the total number of robots that individual robots touch) between the robots in the “flexible” case (8.7 ± 9.0%), while in the “rigid” case the robots go from highly connected to spread out (20.1 ± 25.1%) as shown in SI Appendix, Fig. S18. We also note that the mean displacement of the entire group is quite similar in both cases (flexible, 10.5 ± 4.9 mm; rigid, 9.5 ± 6.0 mm; Fig. 6D); however, the mean displacement differs significantly for individual robots in the rigid case as only one to two robots successfully make it all of the way to the light source as a unit (Fig. 6F). By tracking all robots, we also plot a summary of the distribution of the vertical arm positions with time (Δy) over time (Fig. 6E). When all or most of
Fig. 6. Collective locomotion of a physically entangled robotic blob. (A) Snapshots from the experiment where all of the robots are active and change their gaits from wiggling (<200 lux) to crawling (>800 lux) according to light intensity (Fig. 5B). At intermediate light intensity (200 to 800 lux) the robots kept their arms straight (rigid), i.e., $\alpha_1 = \alpha_2 = 0$ (Movie S5). (B) The robots with red dots are inactive (unpowered) and their arms can move flexibly to enable them to act as binders for the robotic blob. The remaining robots changed their gaits according to light intensity thresholds described previously (Fig. 5B and Movie S5). (C) Space–time overlap heatmaps of robot positions ($x$-axis), revealing collective motion in the flexible case. Color represents the normalized density of the robots. (D) Mean displacement of the individual robots in a run for the flexible ($n = 17$ trials) and rigid ($n = 11$) cases. $\Delta y = y(t) - y(0)$ with respect to initial arm position of the robots in a blob for four different gaits (blue, all crawl; orange, half crawl; yellow, rigid; purple, flexible). Each dataset has 10,000 data points randomly chosen from two to five experiments. Inset shows schematic illustration of $\Delta y$ measurement. (E) Mean displacement of all of the robots per cycle in a run for the all crawl ($n = 6$ trials), semicrawl ($n = 15$ trials), flexible ($n = 17$ trials), and rigid ($n = 11$) cases (see Table 1 for the details). The activity level is defined as how much the robots move their arms on a vertical plane.

the robots crawl, the distribution is flattened, resulting in spreading out or disentangling behavior. In the flexible case, we see a narrow distribution, revealing that only a few robots crawl, thus leading to collective transport (Fig. 6F).

We additionally conduct experiments to further reinforce that heterogeneous gait differentiation is critical for collective movement. Since only the crawl gait leads to forward movement (the wiggle gait does not break symmetry), we program all robots to execute only a crawl gait. A purely homogeneous crawl gait leads to disentanglement, where the robots move individually rather than collectively, and only a few robots ultimately reach the end of the arena ($3.73 \pm 2.5$ mm; SI Appendix, Fig. S17A; Movie S5; and Fig. 6F). To create a partial gait heterogeneity, we switch off the robots in the rear (three of six inactive), while the front robots execute only a crawl gait. We observe a modest improvement, as the robotic collective does move together now (mean displacement per cycle $6.3 \pm 5.7$ mm; Fig. 6F). A summary of these results is presented in Table 1. Taken together, these results support that a strong physical entanglement and heterogeneous gait differentiation are necessary for collective mobility in the robotic blob.

Finally, to test whether gait differentiation leads to energetic benefits, we measured the power usage of a single robot while performing wiggle and crawl gaits with no additional load on the arms or body. We find that the wiggle gait ($240 \pm 50$ mW) utilizes approximately half the power of the crawl gait ($480 \pm 150$ mW; SI Appendix, Fig. S15D). Although our single-robot measurements suggest that gait differentiation could lead to overall lower
collective energy consumption, further work needs to be done to measure the in situ power consumption of individual robots under different conditions. We anticipate that by harnessing gait differentiation, future entangled robotic swarms could improve their energy efficiency in addition to exhibiting adaptive and decentralized swarming locomotion.

Concluding Remarks
We performed a functional study on a biological system consisting of a multitude of worms, i.e., a worm blob, to investigate the fundamental mechanisms behind the emergent physical adaptability, mechanofunctionality, and locomotion of the entangled collective. We studied systematically how the worm blob reacts to different environmental stresses depending on the type, history, and intensity of the associated perturbations. We found that a worm blob that moves through the use of physical entanglement and differentiation can protect against desiccation collectively by reducing the size of a blob, which allows emergent behavior even in the absence of centralized control. To validate our hypotheses on the collective locomotion of the worm blob, we used a robophysical model of a blob consisting of six small, three-link robots that can entangle to each other with the help of mesh-covered pinned arms. Our model enabled us to investigate other parameters such as the gait, activity, and flexibility of the individuals that are challenging to directly test in the living system.

For robotics, the creation of a coherent swarm of simple robots has long been a goal, and our robophysical blob is part of an emerging trend in leveraging mechanics and physics to perform collective tasks in a decentralized way (37, 38) rather than the traditional algorithm-based and centrally controlled approach to swarms (31, 58–64). Typically in robotics, the complex physical interactions between individual robots are either simplified or ignored in idealized theoretical models. Through our robotic blobs, we show the rich behavior that exists between stochastic local interactions between individuals that leads to emergent functionalities of pulling, reducing substrate friction, and even force transmission in a dynamically cohesive robotic blob. These principles can be harnessed in the design of emergent swarm robots for real-world applications.

For biology, the worm blobs hold exciting potential to inspire adaptive active materials as well as advance our understanding of emergent biomechanics of living collectives (16–20). We anticipate that by harnessing gait differentiation, future entangled robotic swarms could improve their energy efficiency in addition to exhibiting adaptive and decentralized swarming locomotion.