UNIVERSITYOF BIRMINGHAM University of Birmingham

Epineuston vortex recapture enhances thrust in tiny water skaters

Rohilla, Pankaj; O'Neil, Johnathan N.; Bose, Chandan; Ortega-Jimenez, Victor M.; Choi, Daehyun; Bhamla, Saad

DOI: [10.1101/2024.06.17.599397](https://doi.org/10.1101/2024.06.17.599397)

License: Creative Commons: Attribution-NonCommercial (CC BY-NC)

Document Version Other version

Citation for published version (Harvard):

Rohilla, P, O'Neil, JN, Bose, C, Ortega-Jimenez, VM, Choi, D & Bhamla, S 2024 'Epineuston vortex recapture enhances thrust in tiny water skaters' bioRxiv.<https://doi.org/10.1101/2024.06.17.599397>

[Link to publication on Research at Birmingham portal](https://birmingham.elsevierpure.com/en/publications/71b6295c-606a-4139-9d7f-d133f7eef507)

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

•Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.

•User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?) •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

Epineuston vortex recapture enhances thrust in tiny water skaters

Pankaj Rohilla^{*a}, Johnathan N. O'Neil^{*a}, Chandan Bose^b, Victor M. Ortega-Jimenez^c, Daehyun Choi^a, and Saad Bhamla^{†a}

^a School of Chemical and Biomolecular Engineering, Georgia Institute of Technology, Atlanta, GA, USA ^bAerospace Engineering, School of Metallurgy and Materials, University of Birmingham, Birmingham, UK

^c School of Biology and Ecology, University of Maine, ME, USA

Vortex recapture underpins the exceptional mobility of nature's finest fliers and swimmers. Utilized by agile fruit flies and efficient jellyfish, this phenomenon is well-documented in bulk fluids. Despite extensive studies on the neuston—a vital fluidic interface where diverse life forms interact between air and water—neuston vortical hydrodynamics remain unexplored. We investigate epineuston (on water) vortical hydrodynamics in *Microvelia americana*, one of the smallest and fastest water striders, skating at 50 BL/s (15 cm/s) . Their middle legs shed counter-rotating vortices, re-energized by hind legs, demonstrating epineuston vortex recapture. High-speed imaging, particle imaging velocimetry, physical models, and CFD simulations show re-energization increases thrust by creating positive pressure at the hind tarsi, acting as a virtual wall. This vortex capture is facilitated by the tripod gait, leg morphology, and precise spatio-temporal placement of the hind tarsi during the power stroke. Our study extends vortex recapture principles from bulk fluids to the neuston, offering insights into efficient epineuston locomotion, where surface tension and capillary waves challenge movement. Understanding epineuston vortex hydrodynamics can guide the development of energy-efficient microrobots to explore the planet's neuston niches, critical frontlines of climate change and pollution.

¹ The unseen ballet of vortical forces orchestrates ² nature's most efficient swimmers and fliers [\[1–](#page-8-0)[7\]](#page-8-1). ³ These interactions, fundamental to minimizing energy expenditure and maximizing thrust, allow or-⁵ ganisms to utilize energy from their own or others' wakes $[1, 2, 8-10]$ $[1, 2, 8-10]$ $[1, 2, 8-10]$ $[1, 2, 8-10]$ $[1, 2, 8-10]$. Jellyfish boost thrust by capturing vortices during relaxation, creating high-pressure zones $[5,11]$ $[5,11]$. Fruit flies capture leading-edge vortices ⁹ during the fling motion, minimizing the energy re-10 quired to generate new vortices $[12, 13]$ $[12, 13]$. Fish exhibit ¹¹ such efficient wake capture that even dead fish can 12 swim upstream by resonating with oncoming Kármán 13 street vortices $[14, 15]$ $[14, 15]$.

 While these examples occur in bulk fluids, the neuston interface — a vital ecological niche — teems with life. From zooplankton, insects, and spiders to birds, reptiles, and plants, countless organisms in- teract at this boundary in marine and freshwater ecosystems [\[16–](#page-8-11)[27\]](#page-9-0). Despite the challenges of bal- ancing surface tension, drag, buoyancy, and capil- lary waves, no documented examples of vortex re- capture at this interface exist. Driven by curiosity about neuston vortical interactions, we reveal a vor- tex re-energization mechanism in Microvelia ameri-cana (Hemiptera, Veliidae).

²⁶ These millimeter-sized water walkers are ²⁷ epineustonic, living on the water surface and ²⁸ are one of the smallest and fastest on this ecological

niche (u_B ∼ 50 BL/s, Figure [2.](#page-3-0)d). Part of the 29 infraorder Gerromorpha, they are found in creeks ³⁰ and ponds worldwide and include over 200 species $\frac{31}{20}$ (Figure S1) $[28-31]$ $[28-31]$. Unlike most water striders that $\frac{32}{2}$ use elongated middle legs for rowing, Microvelia 33 employ all six legs to walk and run using a tripod $\frac{34}{4}$ gait $[16, 17, 32, 33]$ $[16, 17, 32, 33]$ $[16, 17, 32, 33]$ $[16, 17, 32, 33]$ $[16, 17, 32, 33]$ $[16, 17, 32, 33]$. Their unique morphology and $\overline{}$ kinematics enable them to recapture vortices shed ³⁶ from their middle legs, allowing them to speedily $\frac{37}{27}$ skate across the water surface. These amphibious 38 insects, whose ancestors were terrestrial and used a 39 tripod gait for movement on land, evolved to move ⁴⁰ on water while retaining this gait $[16, 34-37]$ $[16, 34-37]$. Using 41 high-speed imaging, particle imaging velocimetry, $\frac{42}{4}$ physical models, and CFD simulations, we describe ⁴³ the epineuston vortex interactions during the water 44 skating behavior of *Microvelia*.

RESULTS ⁴⁶

Skating on water. *Microvelia* possess dense hair 47 coverage on their bodies and legs (Figure [1.](#page-2-0)a) [\[38\]](#page-10-1). ⁴⁸ SEM analysis reveals a tarsal hair density of $\sim 15,000$ 49 hairs/mm² ($n = 3$), comparable to *Velia caprai* and so Gerridae. $[16, 39, 40]$ $[16, 39, 40]$ $[16, 39, 40]$ $[16, 39, 40]$. This dense coverage enables $\overline{}$ 51 Microvelia to maintain a Cassie-Baxter state $[41]$, $\overline{}$ limiting water infiltration and maintaining superhydrophobicity leading to dimples at air-water surface ⁵⁴ contact points (Figure [1.](#page-2-0)b). The low Weber num- ⁵⁵ ber, $We = \rho v^2 l / \sigma \ll 1$ (see Table S1) indicates that set

[∗]Equal contribution

[†]Corresponding author- \boxtimes saadb@chbe.gatech.edu

Figure 1: Behaviour and morphology of epineustonic *Microvelia americana* (a) Dorsal view of *Microvelia* americana with inset showing a SEM image of the dense hair coverage on middle leg tarsus (pseudo-colored). (b) Microvelia Sp. feeding on a trapped insect in a creek (Brunei), with legs deforming the water surface, forming dimples. (c) Size comparison showing M. americana's small body size relative to commonly found water striders, Gerridae. (d) Alternating tripod gait plot for M. americana locomoting on water surface, showing the gait cycle of each leg performing power (color filled boxes) and recovery strokes (empty boxes). (e) Snapshots showing the side view of M. americana walking on water. (f) Dynamics of M. americana on water, indicating short skating escape-sprints (∼2 s) and intermittent walking behavior over a time span 5 minutes.

⁵⁷ surface tension forces dominate over inertial forces ⁵⁸ in their interfacial locomotion, similar to other water ⁵⁹ striders like Gerridae [\[24\]](#page-9-7).

 Unlike water striders such as Gerridae that use a rowing gait, Microvelia employ an alternate tripod gait typical of terrestrial insects. In this gait, at least ϵ_{63} three legs – the front leg (FL), the contralateral mid- $_{64}$ dle leg (ML), and the ipsilateral hind leg (HL) – per- form a power stroke on water (Figure [1.](#page-2-0)d,e), while the other legs recover in air or sometimes on water

 $(SI \text{ Video } 1).$

To understand their epineustonic locomotion be- ⁶⁸ havior, we examine their dynamics over a 5-minute \qquad 69 period in the lab. During this time $Microvelia$ *ro* primarily engage in intermittent walking, spending $_{71}$ 99.6% of the time in this mode. However, they occasionally sprint as an escape response, skating a distance of \sim 30 mm in \sim 2 seconds (Figure [1.](#page-2-0)e). The 74 temporal trajectory of the middle and hind legs shows $\frac{75}{6}$ overlapping paths during this skating mode, indicat- ⁷⁶

Figure 2: Epineustonic kinematics of Microvelia. (a) Tarsal trajectories of middle and hind legs of Microvelia. The solid lines represent the power strokes, while the faded blue and red lines show the recovery strokes. The trajectories illustrate the time spent by the tarsi during movement. (b) Stroke amplitudes of the middle and hind legs (n = 15), illustrated with their tarsal tip trajectories relative to the motion of their respective shoulder joints. The middle legs exhibit larger stroke amplitudes ($\lambda_{ML} \sim 1.54 \pm 0.43$ mm and $\lambda_{MR} \sim 1.59 \pm 0.74$ mm) compared to the hind legs (λ_{HL} ~ 1.25 ± 0.46 mm and λ_{HR} ~ 1.17 ± 0.47 mm). (c) Stroke frequency (N = 3, n = 15) of the middle and hind tarsi, showing an average stroke frequency of $f \sim 30$ strokes/s. (d) Body speed of Microvelia on water and land (styrofoam) in cm/s (left Y axis) and BL/s (body lengths per second, right Y axis). The average maximum body speed on water is ~ 15 cm/s (~ 50 BL/s), compared to ~ 10 cm/s (~ 40 BL/s) on land. (e) Peak tarsi speeds of Microvelia on water. The middle legs achieve higher peak linear speeds during power strokes (∼ 17 cm/s) compared to the hind legs (~ 14 cm/s). This indicates that the middle legs act as the main hydrodynamic thrust propulsors, with higher acceleration ($\sim 2500 \text{ cm}^2/\text{s}$) compared to the hind legs ($\sim 2000 \text{ cm}^2/\text{s}$).

 π ing interfacial vortical interactions (Figure [2.](#page-3-0)a).

 During the skating mode, the middle legs of Mi - crovelia act as the main hydrodynamic thrust propul- sors [\[32,](#page-9-4) [33\]](#page-9-5). These legs exhibit a stroke ampli- tude 23% larger than the hind legs, while main- taining the same stroke frequency (Figure [2.](#page-3-0)b,c). This larger amplitude allows for greater displace- ment with each stroke, enhancing thrust. The mid- dle legs also achieve higher peak linear speeds during power strokes, 21% faster than the hind legs (Figure [2.](#page-3-0)e). This increased speed, coupled with greater ac- celeration—about 25% higher than that of the hind legs—indicates their dominant role as forceful thrust generators [\[33\]](#page-9-5).

91 Epineuston hydrodynamic interactions. Dur- ing the power stroke, the middle leg tarsi shed pairs of counter-rotating vortices (Figure [3.](#page-5-0)a, stage I). These vortices travel downstream, interacting with the hind tarsi, which enter the water at various spatio-temporal locations. The front tarsi generate weak vortices that dissipate without interacting with other tarsi (SI video II).

⁹⁹ The exact location and timing of the incident hind tarsi relative to the vortices dictate the outcome of these interactions. Favourable interactions re- sult in vortex re-energization, increasing the vortices' strength (Figure [3.](#page-5-0)a, Stage II and III). Body rocking and turning can misalign these interactions, altering the hind legs' angle of attack and leading to vortex annihilation or no interaction (Figure [3.](#page-5-0)b). Addition- ally, if Microvelia moves at high speed, its body can pass over the middle leg vortices before the hind legs can interact with them, emphasizing the importance of timing (Figure [3.](#page-5-0)f).

 We measure the circulation of vortex pairs gener- ated by the middle tarsi during re-energization until they dissipate after hind tarsi interaction. Circula-114 tion, $\Gamma = \int \int_s \omega \, dS$, where ω is the vorticity and S is the bounded area, measures the vortices' strength. As the middle leg initiates the power stroke (Figure [3.](#page-5-0)c, point 1), the vortices' circulation increases, peak- $\text{log at } \Gamma = 2 \text{ cm}^2/\text{s}$ (t = 71 ms), corresponding to the 119 maximum tarsal speed $(22 \text{ cm/s}, t = 70 \text{ ms})$. The middle leg then decelerates, reducing Γ as the vor- tices dissipate (point 3). The hind tarsi then enters the wake, re-energizing the vortices to enhance the circulation to a second, lower peak of $\Gamma = 1.6$ cm²/s $(t = 88.5 \text{ ms})$ due to a lower hind-tarsal speed of 17 $125 \, \text{cm/s}$ (Figure [3.](#page-5-0)c, point 4). This cycle ends with the hind tarsi completing their power stroke and dissipat-ing the vortices (SI video II, Figure S2).

128 Across 52 instances in 6 specimens, we observe that 129 60% of the interactions result in re-energization, 27% ¹³⁰ show no interaction, and the remainder lead to vortex annihilation (Figure $3.e$ $3.e$). We compare the normalized peak circulation before and after their hind 132 tarsal interaction $(\tilde{\Gamma} = \Gamma_2/\Gamma_1)$ with normalized body 133 speed ($\tilde{\mathbf{u}} = \mathbf{u}_B/\mathbf{u}_t$) and the time interval (Δt) be-tween strokes (Figure [3.](#page-5-0)d). Vortical re-energization 135 primarily occurs when the hind tarsi strike between ¹³⁶ the middle tarsi vortices with shorter Δt (typically 137 < 6 ms), during initial acceleration phase of the skat-ing sprint (Figure [3.](#page-5-0)f). At higher body speeds, longer $_{139}$ Δt , or due to body turning or rocking, the hind tarsi 140 miss the vortices resulting in no interaction. When $_{141}$ the hind legs skate across the pair of vortices rather $_{142}$ than slaloming between them, the interactions tend ¹⁴³ to weaken the vortices, leading to vortex annihila- ¹⁴⁴ tion (Figure [3.](#page-5-0)d). Collectively, this reinforces that $_{145}$ both the hind tarsi's entry position relative to the ¹⁴⁶ middle tarsi (angle-of-attack) and the inter-stroke in- ¹⁴⁷ terval play critical roles in determining the outcome ¹⁴⁸ of these interactions. ¹⁴⁹

Epineuston vortical recapture increases thrust 150 in *Microvelia*. Reconstructed pressure fields from 151 PIV-measured velocity fields reveal insights into vor- ¹⁵² tical interactions with the hind tarsi of *Microvelia* 153 (Figure [4.](#page-7-0)a). During vortex re-energization, a lo- ¹⁵⁴ cal pressure gradient forms from upstream to down- ¹⁵⁵ stream of the hind tarsi, generating the highest rel- ¹⁵⁶ ative pressure $(\Delta p \sim 5 \text{ Pa})$. In contrast, vortex an- 157 nihilation results in lower relative pressure $(\Delta p \sim 2$ 158 Pa, Figure S3), with cases of no interaction showing 159 similarly low pressure.

We calculate the total impulse by integrating the 161 relative pressure over time, $\mathbf{I} = \int_T \Delta p A dt$, where T is 162 the duration of the power stroke and A is the planar 163 area containing the tarsi and its wake (Figure S3). ¹⁶⁴ Normalizing the impulse, $\tilde{\mathbf{I}} = \int_T \Delta p dt / (\rho \bar{\mathbf{u}}_{t,n}^2 A)$, isolates the impact of hind tarsal interaction from tarsal 166 speed. Excluding the impulse from the middle tarsi 167 yields the relative impulse, $\tilde{\mathbf{I}}_r = \tilde{\mathbf{I}} / \tilde{\mathbf{I}}_{middle}$.

Our results show that vortex re-energization pro- ¹⁶⁹ duces a normalized impulse $(\tilde{\mathbf{I}}_r \sim 1.08)$, 34% higher 170 than vortex annihilation (\sim 0.81) and 15% higher $_{171}$ than no interaction (\sim 0.94) (Figure [4.](#page-7-0)b). This increased impulse results from enhanced fluid entrain- ¹⁷³ ment during re-energization, which raises pressure in 174 the tarsal plane. When hind tarsi step into the cen- ¹⁷⁵ ter of the vortex pair, they entrain more fluid mass ¹⁷⁶ due to the converging flow driven by the vortical mo- ¹⁷⁷ tion $[44]$, leading to increased pressure and greater 178 thrust.

The observed rise in normalized impulse during ¹⁸⁰ re-energization illustrates Microvelia's ability to har- ¹⁸¹ ness energy from its own wake, a phenomenon we ¹⁸² call 'Epineuston Vortex Recapture'. Typically, wakes 183 signifies lost energy to the environment. By step- ¹⁸⁴

Figure 3: Hydrodynamic interactions in *Microvelia's* epineustonic locomotion. (a) Stages of vortical shedding from the power strokes of the middle tarsus and their subsequent interactions with the hind tarsus. LHS: Flowfield streamlines visualization in Flowtrace [\[42\]](#page-10-6) and RHS: vorticity field generated in PIVlab [\[43\]](#page-10-7). Stage I - Vortices generated during the onset of the power stroke of the middle right tarsus, (II) Hind legs stepping into the vortices shed from the middle left tarsus, (III) re-energized vortices from the hind right tarsus; LHS shows the vorticity field corresponding to the frame on the right. (b) Illustrations represent the three different outcomes of vortical interactions based on the trajectory of the hind and middle tarsi. (c) Representative tarsal velocity profiles of the middle-right and hind-right tarsi of *Microvelia* walking on water and the corresponding circulation (filled circles) of the vortices for the case of vortex re-energization. (d) Effect of normalized body speed (relative to hind tarsi speed) on the circulation ratio of vortices originating from the middle legs pre- and post-interactions with the hind tarsi. (e) Percentage outcomes of the vortical interactions of the hind tarsi with vortices shed from the middle tarsi, and (f) Different vortical interactions within a single run on water in Microvelia.

 ping into vortices generated by its middle legs during previous strokes, Microvelia harnesses this energy to increase thrust production by the hind legs. This mechanism, driven by its tripod gait and interfacial movement, enables Microvelia to effectively generate thrust at the air-water interface.

 Physical models validate inter-stroke interval in epineustonic vortical interactions. To evalu-193 ate the effect of inter-stroke intervals (Δt) on vorti- cal interactions, we use a physical model. The model simulates Microvelia's middle and hind tarsi power strokes on water, varying Δt to alter the hind tarsi's angle of attack to the vortices shed by the middle legs. The Reynolds number of the model (∼18) is 199 within the range of *Microvelia* ($Re \sim 2-21$, see Ta- ble S1). The first arm generates a counter-rotating vortex dipole, which the second arm interacts with, $_{202}$ depending on Δt (Figure [4.](#page-7-0)c).

203 For large $\Delta t = -116$ ms, the first arm's vortices ²⁰⁴ dissipate before the second arm's entry, resulting in ²⁰⁵ no interaction (SI Video III, Figure S4). Reducing the ²⁰⁶ interval allows for re-energization, with the second ²⁰⁷ arm's vortices showing higher normalized circulation $_{208}$ $(\Gamma > 1)$ (Figure [4.](#page-7-0)d). However, at very short inter-²⁰⁹ vals $(-10 < \Delta t < 10$ ms), capillary waves generated ²¹⁰ by the arms disrupt the vortices, leading to annihila- $_{211}$ tion. The normalized circulation $(Γ)$ compares both ²¹² Microvelia and the physical model, revealing that op-²¹³ timal inter-stroke intervals enhance re-energization ²¹⁴ and thrust (Figures S4, S5). Vortical re-energization ²¹⁵ in *Microvelia* doesn't always imply $\Gamma > 1$ due to re-²¹⁶ duced hind legs' tarsal speed, imparting less energy $_{217}$ to the vortices (Figure [2.](#page-3-0)e).

 CFD analysis of thrust enhancement dur- ing vortex re-energization. We simulate thrust enhancement through vortex capture using 2D CFD 221 models of high aspect ratio $(AR = 20)$ rectangular plates undergoing prescribed rotation and transla- tion. Mimicking the physical model configuration, the first plate rotates counterclockwise, and the sec-225 ond rotates clockwise, starting with a time gap (Δt) , traversing the first plate's vortical wake (SI Video 4, Figure S5). These simulations evaluate the role of vortex re-energization on thrust via robotic arms' tra- jectory rather than mimicking the precise kinematics of Microvelia.

 Streamline analysis shows differences in flow ve-232 locity magnitudes for different Δt intervals (Figure [4.](#page-7-0)e). For $\Delta t = 0.2$ s, the second plate captures the first plate's wake, entering its recirculation region $_{235}$ closely (SI video 4). As a result, the vortex cores with the same sense of rotation from both plates co-align to increase the resultant circulation, augmenting the 238 propulsive force. In contrast, for $\Delta t = 0.5$ s, the second plate fails to interact effectively, leading to vortex ²³⁹ annihilation with lower flow velocities and thrust due ²⁴⁰ to the absence of effective wake capture.

To reinforce our findings, we present the tempo- ²⁴² ral evolution of the coefficient of thrust (C_T) for the 243 second plate, showing vortical re-energization, com- ²⁴⁴ pared to no interaction (Figure [4.](#page-7-0)f). The C_T , defined 245 as $2T/(\rho_f \mathbf{u_r}^2 A)$, where T is the thrust force, ρ_f is the 246 fluid density, \mathbf{u}_r is the relative linear velocity, and A 247 is the plate's projected area, illustrates the influence ²⁴⁸ of interaction on vorticity fields at different times. ²⁴⁹

For $\Delta t = 0.2$ s, C_T peaks at $t = 1.2$ s as vortex 250 dipoles from both plates interact, augmenting circu- ²⁵¹ lation. As the plates separate, C_T decreases, showing reduced wake interaction. Thrust enhancement ²⁵³ via wake capture correlates with changes in fluid im- ²⁵⁴ pulse, influenced by circulation and vortex core ve- ²⁵⁵ locities. This unsteady flow situation is consistent 256 with pressure data from *Microvelia* and the physical 257 model (Figure [4.](#page-7-0)a,b), where optimal stroke timing $_{258}$ increases entrainment and thrust, demonstrating en- ²⁵⁹ hanced thrust through vortex re-energization.

Conclusions and Outlook ²⁶¹

Our findings illuminate vortical interactions within ²⁶² the neuston, the dynamic water-air boundary that 263 supports diverse life forms. *Microvelia*, among one 264 of the smallest and fastest epineustonic animals, cre- ²⁶⁵ ate nearly 2D vortices due to their minute size and ²⁶⁶ weight, forming shallow dimples on the water surface $[45, 46]$ $[45, 46]$. Their alternating tripod gait, inherited 268 from terrestrial ancestors, enables versatile movement ²⁶⁹ across water, land, and duckweed $[17, 32, 47]$ $[17, 32, 47]$ $[17, 32, 47]$ $[17, 32, 47]$.

Although less energy-efficient than rowing gait, the 271 alternatig tripod gait excels in amphibious locomo- ²⁷² tion, providing *Microvelia* with a strategic advantage 273 in foraging and evading predators $[16, 24, 46]$ $[16, 24, 46]$ $[16, 24, 46]$ $[16, 24, 46]$. This $_{274}$ gait and leg proportions faciliate epineustron vortex ²⁷⁵ recapture combination, where hind leg tarsi boost the ²⁷⁶ circulation and fluid entrainment of vortices shed by ²⁷⁷ middle legs. This re-energization creates a positive ²⁷⁸ pressure at the hind tarsi, acting as a virtual wall ²⁷⁹ that augments thrust $[46]$. In other genera such as 280 Mesovelia, longer middle legs prevent effective vortex ²⁸¹ recapture, underscoring the critical role of leg size in ²⁸² this mechanism (Figure S7).

Epineuston vortex interactions hinge on the spatial ²⁸⁴ location, angle of attack, and trajectory of hind leg ²⁸⁵ tarsi, determining whether vortices are re-energized, ²⁸⁶ annihilated, or minimally interacted with. Our 287 robotic arm physical model and CFD simulations re- ²⁸⁸ inforce the impact of inter-stroke intervals on these ²⁸⁹ interactions. The data indicate that optimal timing ²⁹⁰

Figure 4: Quantifying epineuston vortical interactions through physical models and CFD analysis. (a) Temporal evolution of relative pressure (ΔP) and tarsal speed (v_t) of the hind leg. No-interaction and annihilation cases represent the the hind left tarsi, while re-energization corresponds to the hind right tarsi. (b) Normalized impulse for different types of vortical interaction. The semi-violin plot shows the distribution of the data as a jitter plot, while the box and whisker plot represent the median and the four quartiles (25%, 50%, 75%, and 100%) for Microvelia specimens (N=3) and strokes (n = 12). (c) Temporal evolution of the vortex circulation Γ for each robotic arm with varying Δt showing different vortical interaction outcomes. (d) Regime map of normalized circulation (Γ_2/Γ_1) for varying Δt . Γ_1 and Γ_2 represent peak circulation from the middle leg (or first arm) and hind leg (or second arm), respectively $(N=7, n=53)$. (e) CFD results showing effect of the time interval between plate movements (in quiescent fluid) on vortical interactions depicted by velocity magnitude contours. The second plate starts moving at $t = 0$ s with $\Delta t = 0.2$ s for re-energization and $\Delta t = 0.5$ s for annihilation. In snapshots at $t = 1$ s, arrows indicates the enhanced and reduced velocity field due to wake capture and wake annihilation respectively. (f) Temporal evolution of the coefficient of thrust (C_T) of the second plate for re-energization and no interaction. Snapshots show the interaction's impact on instantaneous vorticity fields at different times.

 and positioning of leg strokes enhance thrust through vortex re-energization, offering new insights into fluid dynamics at the air-water interface. Exploring mi- crovelia juvenile nymphs, multiphase CFD simula- tions, and turbulent flow regimes will further deepen our understanding of these interactions.

 By uncovering the physics behind the vortical re- capture in Microvelia, we extend similar mechanisms observed in jellyfish and fruit flies to the neuston [\[5,](#page-8-5) [11,](#page-8-6) [12\]](#page-8-7). Epineuston vortex recapture could inspire the development of efficient water-skating devices and amphibious robots, enhancing our exploration of the oceanic and freshwater neuston niches [\[26\]](#page-9-8).

304 **ACKNOWLEDGMENTS**

 The authors thank the members of the Bhamla Lab for their feedback and useful discussions. MSB ac- knowledges funding support from the NSF Grants CAREER 1941933 and PHY-2310691, and gift fund- ing from the Open Philanthropy Project. PR ac- knowledges the funding support from the Eckert Postdoctoral Fellowship, Georgia Tech. J.O. ac- knowledges funding support from the GT UCEM fel- lowshp program and the Herbert P. Haley fellowship ³¹⁴ program.

315 COMPETING INTERESTS

³¹⁶ The authors declare no competing interests.

317 REFERENCES

- ³¹⁸ [1] John O Dabiri. Optimal vortex formation as a ³¹⁹ unifying principle in biological propulsion. An-³²⁰ nual review of fluid mechanics, 41:17–33, 2009.
- ³²¹ [2] PF Linden and JS Turner. 'Optimal' vortex rings ³²² and aquatic propulsion mechanisms. Proceedings ³²³ of the Royal Society of London. Series B: Bio-³²⁴ logical Sciences, 271(1539):647–653, 2004.
- ³²⁵ [3] Erik J Anderson and M Edwin DeMont. The me-³²⁶ chanics of locomotion in the squid loligo pealei: ³²⁷ locomotory function and unsteady hydrodynam-³²⁸ ics of the jet and intramantle pressure. *Jour-* $_{329}$ nal of Experimental Biology, $203(18):2851-2863$, ³³⁰ 2000.
- ³³¹ [4] Michael S Triantafyllou, GS Triantafyllou, and ³³² DKP Yue. Hydrodynamics of fishlike swimming. ³³³ Annual review of fluid mechanics, 32(1):33–53, ³³⁴ 2000.
- ³³⁵ [5] John H Costello, Sean P Colin, John O Dabiri, ³³⁶ Brad J Gemmell, Kelsey N Lucas, and Kelly R

Sutherland. The hydrodynamics of jellyfish 337 swimming. Annual Review of Marine Science, 338 13:375–396, 2021. ³³⁹

- [6] Iztok Lebar Bajec and Frank H Heppner. Or- ³⁴⁰ ganized flight in birds. Animal Behaviour, ³⁴¹ 78(4):777–789, 2009. ³⁴²
- [7] John Roger Speakman and D Banks. The func- ³⁴³ tion of flight formations in greylag geese anser ³⁴⁴ anser; energy saving or orientation? Interna- ³⁴⁵ tional Journal of Avian Science, 140(2):280–287, ³⁴⁶ $1998.$ 347
- [8] Hao Liu, Shizhao Wang, and Tianshu Liu. Vor- ³⁴⁸ tices and forces in biological flight: Insects, ³⁴⁹ birds, and bats. Annual Review of Fluid Me- ³⁵⁰ $chanics, 56:147-170, 2024.$ ³⁵¹
- [9] You-Jun Lin, Sheng-Kai Chang, Yu-Hsiang Lai, ³⁵² and Jing-Tang Yang. Beneficial wake-capture ³⁵³ effect for forward propulsion with a restrained $\frac{354}{256}$ wing-pitch motion of a butterfly. Royal Society 355 open science, 8(8):202172, 2021. ³⁵⁶
- [10] Nils B Tack, Kevin T Du Clos, and Brad J Gem- ³⁵⁷ mell. Fish can use coordinated fin motions to ³⁵⁸ recapture their own vortex wake energy. Royal 359 Society Open Science, 11(1):231265, 2024.
- [11] Brad J Gemmell, Sean P Colin, and John H ³⁶¹ Costello. Widespread utilization of passive en- ³⁶² ergy recapture in swimming medusae. Journal of 363 Experimental Biology, $221(1)$:jeb168575, 2018. $\frac{364}{100}$
- [12] Fritz-Olaf Lehmann, Hao Wang, and Thomas 365 Engels. Vortex trapping recaptures energy in ³⁶⁶ flying fruit flies. Scientific Reports, $11(1):6992$, 367 $2021.$ 368
- [13] Michael H Dickinson, Fritz-Olaf Lehmann, and 369 Sanjay P Sane. Wing rotation and the 370 aerodynamic basis of insect flight. Science, ³⁷¹ $284(5422):1954-1960, 1999.$ 372
- [14] David N Beal, Franz S Hover, Michael S Tri- ³⁷³ antafyllou, James C Liao, and George V Lauder. ³⁷⁴ Passive propulsion in vortex wakes. Journal of 375 fluid mechanics, 549:385-402, 2006.
- [15] Liang Li, Máté Nagy, Jacob M Graving, Joseph $\frac{377}{277}$ Bak-Coleman, Guangming Xie, and Iain D ³⁷⁸ Couzin. Vortex phase matching as a strategy ³⁷⁹ for schooling in robots and in fish. Nature com-
s80 munications, 11(1):5408, 2020.
- [16] N Møller Andersen. A comparative study of loco- ³⁸² motion on the water surface in semiaquatic bugs 383

³⁸⁴ (insecta, hemiptera, gerromorpha). Vidensk. ³⁸⁵ Meddel. Natuirist. Foren. Kjobenhavn, 139:337– ³⁸⁶ 396, 2016.

 [17] Antonin JJ Crumiere, M Emilia Santos, Marie Sémon, David Armisén, Felipe FF Moreira, and Abderrahman Khila. Diversity in morphology and locomotory behavior is associated with niche \sum_{391} expansion in the semi-aquatic bugs. Current Bi- $ology, 26(24):3336-3342, 2016.$

- ³⁹³ [18] Jeffrey W. Shultz. Walking and Surface Film ³⁹⁴ Locomotion in Terrestrial and Semi-Aquatic ³⁹⁵ Spiders. Journal of Experimental Biology, ³⁹⁶ 128(1):427–444, 03 1987.
- ³⁹⁷ [19] S Tonia Hsieh and George V Lauder. Run-³⁹⁸ ning on water: Three-dimensional force gen-³⁹⁹ eration by basilisk lizards. Proceedings of the ⁴⁰⁰ National Academy of Sciences, 101(48):16784– ⁴⁰¹ 16788, 2004.
- ⁴⁰² [20] Glenna T Clifton, Tyson L Hedrick, and An-⁴⁰³ drew A Biewener. Western and clark's grebes ⁴⁰⁴ use novel strategies for running on water. The ⁴⁰⁵ Journal of Experimental Biology, 218(8):1235– ⁴⁰⁶ 1243, 2015.
- ⁴⁰⁷ [21] Victor M Ortega-Jimenez, Elio J Challita, Baek-⁴⁰⁸ gyeom Kim, Hungtang Ko, Minseok Gwon, ⁴⁰⁹ Je-Sung Koh, and M Saad Bhamla. Di-⁴¹⁰ rectional takeoff, aerial righting, and adhe-⁴¹¹ sion landing of semiaquatic springtails. Pro-⁴¹² ceedings of the National Academy of Sciences, ⁴¹³ 119(46):e2211283119, 2022.
- ⁴¹⁴ [22] S Wang and AM Ardekani. Swimming of a model ⁴¹⁵ ciliate near an air-liquid interface. Physical Re $view E, 87(6):063010, 2013.$
- $_{417}$ [23] Jonathan Voise and Jérôme Casas. The manage-⁴¹⁸ ment of fluid and wave resistances by whirligig ⁴¹⁹ beetles. Journal of The Royal Society Interface, $7(43):343-352, 2010.$
- $_{421}$ [24] David L Hu and John WM Bush. The hydrody-⁴²² namics of water-walking arthropods. Journal of ⁴²³ Fluid Mechanics, 644:5–33, 2010.
- ⁴²⁴ [25] Zexiang Huang, Hao Yang, Ke Xu, Jianing Wu, ⁴²⁵ and Jinxiu Zhang. Collecting differently sized ⁴²⁶ particles on water surface by maneuvering pedal ⁴²⁷ waves on the foot of the water snail pomacea ⁴²⁸ canaliculata. Soft Matter, 18(40):7850–7858, ⁴²⁹ 2022.
- ⁴³⁰ [26] Rebecca R Helm. The mysterious ecosys-⁴³¹ tem at the ocean's surface. *PLoS Biology*, $432 \hspace{1.5cm} 19(4):e3001046, 2021.$
- [27] Lanna Cheng and Himanshu Mishra. Why did ⁴³³ only one genus of insects, halobates, take to the ⁴³⁴ high seas? *PLoS Biology*, 20(4):e3001570, 2022. 435
- [28] Aidamalia Vargas-Lowman, David Armisen, ⁴³⁶ Carla Fernanda Burguez Floriano, Isabelle ⁴³⁷ da Rocha Silva Cordeiro, Séverine Viala, 438 Mathilde Bouchet, Marie Bernard, Augustin ⁴³⁹ Le Bouquin, M Emilia Santos, Alexandra ⁴⁴⁰ Berlioz-Barbier, et al. Cooption of the pteridine 441 biosynthesis pathway underlies the diversifica- ⁴⁴² tion of embryonic colors in water striders. Pro- ⁴⁴³ ceedings of the National Academy of Sciences, ⁴⁴⁴ 116(38):19046–19054, 2019. ⁴⁴⁵
- [29] Matthew R Pintar, Jeffrey L Kline, and Joel C ⁴⁴⁶ Trexler. The aquatic heteroptera (hemiptera) of $_{447}$ marshes in the florida everglades. Florida Entomologist, 104(4):307–319, 2021.
- [30] JT Polhemus. Water-striders (hemiptera: Ger- ⁴⁵⁰ ridae, veliidae, etc.). Marine insects. Elsevier, ⁴⁵¹ Amsterdam, pages 187–224, 1976.
- [31] Nils Møller Andersen. Microvelia polhemi, n. ⁴⁵³ sp. (heteroptera: Veliidae) from dominican am- ⁴⁵⁴ ber: The first fossil record of a phytotelmic water 455 strider. Journal of the New York Entomological 456 $Society, 107:135–144, 1999.$
- [32] Johnathan Nathaniel O'Neil, Kai Lauren Yung, ⁴⁵⁸ Gaetano Difini, Holden Walker, and M Saad ⁴⁵⁹ Bhamla. Tiny amphibious insects use tripod gait $_{460}$ for seamless transition across land, water, and ⁴⁶¹ duckweed. *bioRxiv*, pages $2024-04$, 2024 .
- [33] Johnathan Nathaniel O'Neil, Kai Lauren Yung, ⁴⁶³ Gaetano Difini, Pankaj Rohilla, and M Saad ⁴⁶⁴ Bhamla. Limb loss and specialized leg dynam- ⁴⁶⁵ ics in tiny water-walking insects. bioRxiv, pages ⁴⁶⁶ 2024–04, 2024. ⁴⁶⁷
- [34] Nils Møller Andersen. Phylogenetic inference as ⁴⁶⁸ applied to the study of evolutionary diversifica- ⁴⁶⁹ tion of semiaquatic bugs (hemiptera: Gerromor- ⁴⁷⁰ pha). Systematic Zoology, 28(4):554–578, 1979. ⁴⁷¹
- [35] Miguel Piñeirua, Anna Verbe, and Jérôme $_{472}$ Casas. Substrate-mediated leg interactions play ⁴⁷³ a key role in insect stability on granular slopes. ⁴⁷⁴ Physical Review E, 108(1):014903, 2023.
- [36] Antoine Humeau, Miguel Piñeirua, Jérôme Cras- 476 sous, and Jérôme Casas. Locomotion of ants 477 walking up slippery slopes of granular materials. Integrative Organismal Biology, 1(1):obz020, ⁴⁷⁹ $2019.$ 480

 [37] Pavan Ramdya, Robin Thandiackal, Raphael Cherney, Thibault Asselborn, Richard Benton, Auke Jan Ijspeert, and Dario Floreano. Climb- ing favours the tripod gait over alternative faster insect gaits. Nature communications, 8(1):14494, 2017.

- [38] Lanna Cheng. Marine and freshwater skaters: differences in surface fine structures. Nature, 242(5393):132–133, 1973.
- [39] Nils Møller Andersen. Fine structure of the body hair layers and morphology of the spiracles of semiaquatic bugs (insecta, hemiptera, gerromor- $_{493}$ pha) in relation to life on the water surface. Vidensk. Medd. Dansk Naturhist. Foren., 140:7-37, 1977.
- ⁴⁹⁶ [40] Cédric Finet, Amélie Decaras, Maria Rutkowska, Pascale Roux, Samuel Collaudin, ⁴⁹⁸ Pauline Joncour, Séverine Viala, and Abder- rahman Khila. Leg length and bristle density, both necessary for water surface locomotion, are genetically correlated in water striders. Proceedings of the National Academy of Sciences, 119(9):e2119210119, 2022.
- [41] ABD Cassie and S Baxter. Wettability of porous surfaces. Transactions of the Faraday society, 40:546–551, 1944.
- [42] William Gilpin, Vivek N Prakash, and Manu Prakash. Flowtrace: simple visualization of co- herent structures in biological fluid flows. Jour- nal of experimental biology, 220(19):3411–3418, 2017.

 [43] William Thielicke and Ren´e Sonntag. Particle image velocimetry for matlab: Accuracy and en- hanced algorithms in pivlab. Journal of Open Research Software, 9(1), 2021.

- [44] Michael Krieg and Kamran Mohseni. Modelling circulation, impulse and kinetic energy of start- ing jets with non-zero radial velocity. Journal of Fluid Mechanics, 719:488–526, 2013.
- [45] Robert B Suter. Spider locomotion on the water surface: biomechanics and diversity. The Jour-nal of Arachnology, $41(2):93-101$, 2013 .
- [46] Thomas Steinmann, Antoine Cribellier, and ₅₂₄ Jérôme Casas. Singularity of the water strider propulsion mechanisms. Journal of Fluid Me-chanics, 915, 2021.

[47] Stephen P Yanoviak and DN Frederick. Water 527 surface locomotion in tropical canopy ants. Journal of Experimental Biology, $217(12):2163-2170$, 529 $2014.$ 530