

Localized fluidization burrowing mechanics of *Ensis directus*

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Keywords: burrowing, bivalves, fluidization, particle image velocimetry, soil mechanics

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SUMMARY

Muscle measurements of *Ensis directus*, the Atlantic razor clam, indicate that the organism only has sufficient strength to burrow a few centimeters into its habitat soil, yet razor clams burrow to over 70 cm. In this paper, we show that the animal uses motions of its valves to locally fluidize surrounding soil and reduce burrowing drag. Substrate deformations were measured using Particle Image Velocimetry (PIV) in a novel visualization system that enabled us to see through soil and watch *E. directus* burrow *in situ*. PIV data, supported by soil and fluid mechanics theory, show that the contraction motion of *E. directus*' valves locally fluidizes the surrounding soil. Particle and fluid mixtures can be modeled as a Newtonian fluid with an effective viscosity based on the local void fraction. Using these models, we demonstrate that *E. directus* is strong enough to reach full burrow depth in fluidized soil, but not in static soil. Furthermore, we show that *E. directus*' method of localized fluidization reduces the amount of energy to reach burrow depth by an order of magnitude compared to penetrating static soil and leads to a burrowing energy that scales linearly with depth rather than depth squared.

1 INTRODUCTION

2 There are many examples of animals that live in particulate substrates and have adapted
3 unique locomotion schemes. The sandfish lizard (*S. scincus*) wiggles its body from side
4 to side in order to effectively swim through sand (Maladen et al., 2009). Smaller
5 organisms, such as nematodes (*C. elegans*), have been observed to move quite efficiently
6 via an undulatory motion in saturated granular media (Jung, 2010a; Wallace, 1968). Clam
7 worms (*N. virens*) have been observed to burrow in gelatin, a material with similar
8 properties as elastic muds, by propagating a crack (Dorgan et al., 2005). Throughout this
9 paper, we use the terms “soil” and “substrate” to refer to the media in which burrowing
10 organisms live, regardless of whether they are granular or cohesive. We specify that these
11 substrates are composed of small particles (from clay to coarse sand) with the term
12 “particulate.”

13
14 Numerous soft-bodied organisms that live in particulate substrates saturated with a pore
15 fluid use a two-anchor system to burrow: one section of the animal expands to form an
16 anchor while another section contracts and extends to progress forward in the burrow;
17 once extension is exhausted, the roles of each section are reversed (Dorgan et al., 2005;
18 Fager, 1964; Holland and Dean, 1977; Jung, 2010b; Shin et al., 2002; Stanley, 1969;
19 Trueman, 1966a; Trueman, 1966b; Trueman, 1967; Trueman, 1975). In this paper, we
20 show that the Atlantic razor clam (*Ensis directus*), which burrows via the two-anchor
21 method, uses motions of its valves to create a pocket of fluidized substrate around its
22 body in order to reduce drag forces and burrowing energy expenditure.

23
24 Trueman measured ≈ 10 N as the maximum pulling force *E. directus* can exert to pull its
25 valves into soil (Trueman, 1967). Using a blunt body with a similar size and shape as *E.*
26 *directus*, pressed into the animal’s habitat soil on a mud flat in Gloucester, MA, we
27 measured that 10 N of force should enable *E. directus* to submerge to approximately 1-2
28 cm (Fig. 9A, further described in the Discussion section). In reality, razor clams can dig
29 to 70 cm (Holland and Dean, 1977). Since drag force scales linearly with depth for a
30 body moving through a granular medium (Robertson and Campanella, 1983), muscle
31 force measurements indicate that *E. directus* is approximately 75 X too weak to reach full

1 burrow depth in static soil. Thus, the animal must manipulate the surrounding
2 environment to reduce drag.

3
4 *E. directus* is enclosed by two, long, slender valves that hinge along an axis oriented
5 longitudinally to the animal. A foot, which is a dexterous, soft organ, resides at the
6 bottom of the valves. *E. directus* burrows by using a series of valve and foot motions to
7 draw itself into the substrate (Fig. 1). We estimate an upper bound of the mechanical
8 energy per depth for the animal to advance its valves downwards by adapting pedal
9 strength, valve displacement, hinge stiffness, and mantel cavity pressure from (Trueman,
10 1967) and summing the expended mechanical energy and attained displacements for each
11 burrowing motion: valve uplift (0.05 J, 0.5 cm), valve contraction (0.07 J, 20°), valve
12 penetration (0.20 J, -2.0 cm), combine for a total of 0.21 J/cm. Re-expansion of the
13 valves is accomplished through elastic rebound of the hinge ligament and thus requires
14 no additional energy input by the animal. To put the magnitude of these energetics in
15 perspective, *E. directus* can travel over a half kilometer through soil on the energy stored
16 in a AA battery (Energizer, 2009).

17 18 **MATERIALS AND METHODS**

19 ***E. directus* collection**

20 *E. directus* specimens used for our research were collected from natural stocks in Orleans
21 and Gloucester, Massachusetts under the appropriate research permits issued by the
22 state's fish and game department. Once specimens were harvested and brought back to
23 the lab, they were held in a chilled, saltwater commercial lobster tank (Stark Products 50-
24 gallon lobster tank, College Point, NY, USA).

25 26 ***Blunt body drag measurement***

27 The device shown in Fig. 2 was used to measure drag force on *E. directus*' valves in
28 static soil. The device is constructed from a pair of *E. directus* valves bonded to an
29 aluminum rod with urethane casting compound (McMaster-Carr #8690K1, Princeton, NJ,
30 USA). The valves are positioned in their natural orientation and the cross-sectional area
31 of the rod is less than the frontal area of the valves, as to minimize its effect on drag. A

1 fish scale (Jennings Ultrasport 30, www.jscale.com) attached to a pushing handle is
2 mounted to the top of the rod to measure insertion force. The device was operated by
3 slowly pushing it into soil (as to avoid inertial effects) and recording force at each depth
4 increment marked on the rod.

6 ***Visualization tank***

7 The adage “clear as mud” is used to describe the difficulty of visually investigating
8 burrowing animals *in situ*. To surmount this challenge and view *E. directus* burrowing
9 motions, as well as deformations in the soil surrounding the animal, we developed the
10 visualizer shown in Fig 3A. The visualizer is essentially a backlit ant farm, or Hele-Shaw
11 cell, which is commonly used in fluid mechanics experiments to measure flow in two
12 dimensions (Kundu and Cohen, 2004). The front viewing pane is adjustable forward and
13 aft via a lead screw and bellowed side walls, allowing tank and animal thickness to be
14 matched. Specimens were oriented vertically in the visualizer with their valves
15 orthogonal to the tank walls (Fig. 3B). This forced a plane strain condition while allowing
16 for soil deformation in the principal directions of the animal’s burrowing motions.

18 The substrate used in the visualizer is composed of 1 mm diameter, optically clear soda
19 lime glass beads (Potters Industries A-100, Malvern, PA, USA). This substrate was
20 chosen because its particle size and density ($\rho = 2.52 \text{ g/cm}^3$) are close to that of coarse
21 silica sand (Terzaghi et al., 1996), one of the substrates in which razor clams live
22 (Holland and Dean, 1977). Although smaller substrate particles are more common in *E.*
23 *directus* habitat, we found through experimentation that soda lime glass beads less than 1
24 mm in diameter did not provide adequate light transmission for visualization.

26 Heat dissipation was an important consideration in the visualizer design, as the halogen
27 lamps generate sufficient heat to melt the polycarbonate walls of the tank (and kill the
28 animals) within minutes if not actively cooled. Chilled and oxygenated water supplied
29 from the commercial lobster tank used to hold specimens is circulated through the
30 visualizer (Fig, 3C). This water is mixed with hot water flowing out of the visualizer and
31 gravity-fed into a pump. The pump sends water to a reservoir with a float valve that

1 maintains fill level near the top of the visualizer. Two outlets leave the reservoir: one into
2 the top of the visualizer and one directly back to the lobster tank. The flow rate through
3 the visualizer is 10 gal/min, which yields a downwards bulk flow velocity of ≈ 0.08 cm/s.
4 This is 15 X slower than *E. directus*' uplift velocity (Trueman, 1967) and was therefore
5 assumed to not interfere with burrowing motions.

6
7 Because the flow resistance through the substrate within the visualizer is much greater
8 than the resistance of the "short circuit" line from the reservoir to the lobster tank, the
9 portion of heated water that enters the supply stream at the pump is small compared to
10 chilled water from the lobster tank. As a result, we were able to maintain temperature in
11 the visualizer to within 2° C of the lobster tank, which was set to 10° C.

12 13 ***Valve kinematics tracking***

14 *E. directus* specimens placed in the visualizer were filmed with a high-definition video
15 camera (Sony HD Handycam, Tokyo, Japan) at 30 frames/second. During some tests, the
16 animals were stimulated to dig by lightly touching their siphons. In most tests, *E. directus*
17 burrowed naturally without any applied stimulus. Each test was composed of one
18 complete burrowing cycle (Fig. 1). The data presented in this paper represents 23
19 burrowing cycles performed by three separate animals at a depth ranging from 0.5 to 1
20 body lengths.

21
22 Kinematics of *E. directus* were tracked using the following process: a) frames of each
23 digging cycle were converted to 720 X 480 pixel grayscale images (Fig. 4A); b) each
24 image was converted to black and white in Matlab (The MathWorks, Inc., Natick, MA,
25 USA), with a threshold chosen to accentuate the contrast between dark areas of the valves
26 and light areas of the foot and substrate; c) the images were filtered to remove all
27 connected components consisting of fewer than 50 pixels, which removed dark areas that
28 do not correspond to the valves; and d) the remaining points were divided into four
29 quadrants that approximate the top, bottom, left, and right sides of the valves using a
30 rectangular orthogonal regression algorithm (Arbenz, 2008). The size and position of this

rectangle was then used to identify the width and depth of the clam in each video frame of each test (Fig. 4A).

Particle Image Velocimetry analysis

To measure substrate deformations around burrowing *E. directus*, opaque particles were interspersed in the visualizer substrate at a concentration of seven percent and were tracked using MatPIV Particle Image Velocimetry (PIV) software (Sveen, 2004). Raw video from each of the 23 burrowing cycle tests was converted to grayscale and cropped to show only the substrate to one side of *E. directus* (Fig. 4A). This removed tracking artifacts generated by the motion of the valves and the shadow projected by the animal's body. The substrate to the side of *E. directus* that exhibited the greatest displacement was used for PIV analysis (Fig. 6, causes for asymmetry discussed in Results section). Each video frame was then converted to a 720 X 480 pixel image. Adjacent images were processed with MatPIV in three iterations with interrogation windows of 64 X 64 pixels, 32 X 32 pixels, and 16 X 16 pixels, all with 50% overlap.

Soils' mechanical properties largely depend on void fraction, ϕ , which is the fractional amount of the substrate's volume occupied by voids (liquid or gas). PIV data was used to calculate the instantaneous void fraction field around *E. directus* with the following method. The displacement field, $\partial\delta_i$, between subsequent video frames, was calculated using the velocity, v , measured via PIV, and time, dt , between frames.

$$\partial\delta_i = v_i dt = \frac{\partial\delta_i}{\partial t} dt . \quad (1)$$

Displacements between each video frame were summed to find the total horizontal and vertical displacement fields. At the point of valve contraction (Fig. 1D), immediately before valve penetration (Fig. 1E), the average horizontal displacement of the soil adjacent to the animal for all tests was found to be 9.2 ± 4.9 (mean $\pm$ s.d.) times greater than the vertical displacement. As a result, horizontal soil displacement was considered to be the most important factor in the animal's manipulation of the substrate. To reduce PIV

noise, horizontal displacement as a function of horizontal position was determined by averaging the rows of the horizontal displacement field (Fig. 4B).

Volumetric strain in an element of soil is

$$e = \frac{V^t - V_0^t}{V_0^t}, \quad (2)$$

where V^t is the current and V_0^t is the initial volume of the element. A volumetric strain field is calculated by summing the principal strain fields, ϵ_i . As a result of the disparity between measured horizontal and vertical displacements in our PIV data, and because the visualizer imposed a plane strain condition, we approximated the strain field around

burrowing *E. directus* as $e \approx \epsilon_x = \frac{\partial \delta_x}{\partial x}$. A smooth function for δ_x was required to evaluate this derivative without generating excessive noise; for each burrowing test, an exponential function was fit to the horizontal displacement data (Fig. 4B). Only tests with fits of $R^2 > 0.9$ were used in our analysis, which yielded the 23 data sets reported in this work.

To calculate changes in void fraction at a point, the same control volume must be used before and after deformation, in this case the initial volume of the differential element,

V_0^t . The initial void fraction is then $\phi_0 = 1 - \frac{V_0^s}{V_0^t}$ and the final void fraction is $\phi = 1 - \frac{V^s}{V^t}$,

where the amount of solids remaining in V_0^t after deformation, V^s , is $V^s = V_0^s \frac{V_0^t}{V^t}$.

Combining Eq. 2 with the previous relations yields

$$e = \frac{V_0^s - V^s}{V^s} = \frac{\phi - \phi_0}{1 - \phi} \Rightarrow \frac{e + \phi_0}{1 + e} = \phi, \quad (3)$$

1 an expression for the current void fraction, ϕ , given the initial void fraction, ϕ_o , and the
2 volumetric strain.

3
4 With Eq. 3, temporal changes in void fraction around burrowing *E. directus* were
5 calculated from PIV data (Fig 4B). The only initial condition required was the initial void
6 fraction, which was determined to be 0.38 by measuring the volume of the experimental
7 setup and the volume of the pore fluid.

9 RESULTS

10 Figure 5 shows valve kinematics during five consecutive burrowing cycles for one of the
11 *E. directus* specimens tested. Each stage of the burrowing behavior, corresponding to
12 Figs. 1C-F, can be seen. Digging progress per cycle is small but apparent. Each stage of
13 the digging motion does not happen discretely; contraction (Fig. 1D) continues during
14 penetration (Fig. 1E) and the valves tend to rise during expansion (Fig. 1F).

15
16 Figure 6 shows temporal changes in substrate displacement during one *E. directus*
17 burrowing cycle. Valve uplift (Fig. 6B) and contraction (Fig. 6C) result in a local soil
18 disturbance. Magnitude of the particle motion on either side of the animal was rarely
19 symmetric. This result is expected if *E. directus* lies at a slight angle relative to vertical;
20 substrate on the uphill side of the animal will preferentially deform by sliding
21 downwards, rather than the substrate on the downhill side of the animal deforming by
22 sliding upwards (Terzaghi et al., 1996). This behavior is consistent with our observations.
23 Soil deformations caused by *E. directus*' valve motions occurred on a timescale smaller
24 than that required for the soil to collapse and fill around the animal. The natural failure
25 wedge that occurs in this condition can be seen in Fig. 6D, with the matching overlaid
26 failure angle, θ_f , which was calculated from the soil's friction angle. The relationship
27 between friction and failure angle is explained in the Discussion section.

28
29 Void fraction values calculated from PIV displacement data taken at the point of valve
30 contraction (Fig. 1D) that immediately precedes valve penetration (Fig. 1E) show that
31 these burrowing motions unpack and locally fluidize the substrate proximate to *E.*

directus (Fig. 7). Void fraction at incipient fluidization for the substrate was assumed to be ≈ 0.41 , which is consistent with the level measured by Wen and Yu for substrate particles of similar size, shape, and density (Wen and Yu, 1966). Table 1 reports the horizontal displacement functions that were fit to the horizontal displacement PIV data in order to calculate changes in void fraction and the location of the fluidized zone. These functions were used to extrapolate data (gray dotted lines in Fig. 7) for the shadow regions between the animal's body and the cropped video frames (Fig. 4A).

Table 1: Horizontal soil displacement fit functions and location of fluidized zone

Animal	Test #	A	B	R^2	$\left. \frac{x}{R_E} \right _{\phi=0.41}$
a	1	-1.44	-0.0041	0.96	NA
a	2	-1.73	-0.0041	0.96	NA
a	3	-2.72	-0.0053	0.97	NA
b	1	-5.83	-0.0118	0.99	1.90
b	2	-4.80	-0.0086	0.99	NA
b	3	-4.72	-0.0046	0.92	NA
b	4	-3.77	-0.0063	0.98	NA
b	5	-7.80	-0.0055	0.90	2.41*
b	6	-7.82	-0.0060	0.99	NA
b	7	-5.12	-0.0061	0.99	NA
b	8	-7.17	-0.0055	0.98	NA
c	1	-5.25	-0.0121	0.98	1.84
c	2	-8.45	-0.0103	0.98	3.04
c	3	-7.25	-0.0103	0.98	2.84
c	4	-11.92	-0.0066	0.92	4.53
c	5	-13.43	-0.0061	0.94	4.41
c	6	-10.72	-0.0058	0.95	3.20
c	7	-5.96	-0.0051	0.99	NA
c	8	-4.95	-0.0062	0.98	NA
c	9	-6.64	-0.0053	0.99	NA
c	10	-6.22	-0.0059	0.99	NA
c	11	-7.95	-0.0057	0.98	1.48*
c	12	-6.21	-0.0051	0.99	NA

A and B correspond to the fitting function $\delta_x = Ae^{Bx}$; R^2 between fit and PIV data; fluidization radii marked as NA have $\phi < 0.41$ at $x/R_E = 1$; asterisk indicates extrapolated data at fluidization point.

DISCUSSION

Fluidization due to E. directus valve contraction

As *E. directus* contracts its valves, it reduces the level of stress acting between the valves and the surrounding soil. At some stress level, the imbalance between horizontal and vertical stress causes the soil adjacent to the animal to fail. Continued valve contraction draws pore water towards the animal, which mixes with the failed soil to create a region of localized fluidization.

The stress state in soil at equilibrium and failure can be represented as Mohr's circles (Hibbeler, 2000) on a normal versus shear stress plot (Fig. 8). Failure occurs when the internal shear stress equals the shear strength. Graphically, this is represented when circle b in Fig. 8 is tangent to the failure envelope formed by the soil's friction angle, ϕ . The stress state in Fig. 8 relates to a cohesionless soil; for a cohesive soil, the same failure analysis can be used, but the failure envelope will be shifted vertically by half the soil's cohesive strength.

When *E. directus* first starts to contract its valves, it brings the soil to a state of active failure (Terzaghi et al., 1996). At this point the soil will tend to naturally landslide downward at a failure angle, θ_f , illustrated by inset b in Fig. 8. The failure angle is the transformation angle between the principal stress state and the stress state at the tangency point between the circle and failure envelope. This angle can also be seen in Fig. 8 by connecting the tangency point on the failure envelope, the horizontal failure effective stress, σ'_{hf} , and the principal stress axis, and is given by $\theta_f = \frac{\pi}{4} + \frac{\phi}{2}$. Effective stresses, indicated with a prime, represent the actual stress between soil particles, neglecting hydrostatic pressure of the pore fluid.

We observed that valve uplift and contraction occur on a smaller timescale than that required to form a landslide failure wedge in the substrate around burrowing *E. directus* (Figs. 6B-D). This gives the animal time to move through the fluidized soil before it

1 collapses. The calculated failure angle in the visualizer substrate, determined via the
 2 friction angle (measured as 25° for 1 mm soda lime glass beads), aligns with the failure
 3 wedge angle observed in the PIV data (Fig 6D).

4
 5 Although bulk landslide movement of the soil does not occur when *E. directus* initiates
 6 valve contraction, the resulting imbalance in stresses brings the soil to a state of failure,
 7 creating a failure surface at angle θ_f . The existence of this failure surface is crucial
 8 because it creates a discontinuity in the substrate surrounding *E. directus*, where the soil
 9 particles within are free to move and the soil particles beyond remain stationary. As *E.*
 10 *directus* contracts its valves, it reduces its own body volume; this change in volume must
 11 be compensated by pore fluid drawn into the failure zone. Increasing the pore fluid while
 12 retaining the same number of particles within the failure surface results in a net increase
 13 in void fraction, creating a fluidized zone around *E. directus*.

14
 15 The importance of a soil failure surface around contracting *E. directus* is demonstrated by
 16 investigating whether motion of pore fluid flow alone could create fluidization. The
 17 Reynolds number of the pore fluid adjacent to *E. directus*' valves, calculated from valve
 18 velocity, particle diameter, and the pore fluid density and viscosity, varies between 0.02
 19 and 56, depending on particle size (0.002 to 2 mm) (Holland and Dean, 1977; Terzaghi et
 20 al., 1996; Trueman, 1967), animal size (10 to 20 cm) (personal observations), and valve
 21 contraction velocity ($v \approx 0.011$ to 0.028 m/s) (Trueman, 1967). As this range mostly falls
 22 within the regime of Stokes drag (Kundu and Cohen, 2004), the characteristic time for a
 23 particle to reach valve velocity during contraction can be estimated by conservation of
 24 momentum, given by

$$m_p \frac{dv_p}{dt} = 6\pi\mu_f d_p (v_v - v_p) \Rightarrow t_{char} = \frac{d_p^2 \rho_p}{36\mu_f}, \quad (4)$$

26
 27 where m_p is the mass of the particle, v_v is the valve contraction velocity, v_p is the particle
 28 velocity, and t_{char} is the time constant of the differential equation governing velocity
 29 change.

This analysis yields timescales of 0.075 s for 1 mm soda lime glass beads in water. These timescales are considerably less than the ≈ 0.2 s valve contraction time measured by Trueman (Trueman, 1967) and the contraction time of 1.46 ± 0.58 s (mean $\pm$ s.d.) measured in our tests. Furthermore, 1 mm particles represent an upper bound for *E. directus* habitat; the timescale mismatch is even greater for smaller substrate particles. As such, soil particles surrounding *E. directus* can be considered inertialess and are advected with the pore fluid during valve contraction.

The presence of a failure surface breaks the coupling between particles and pore fluid movement; particles within the failure wedge can freely move and fluidize, while particles outside remain static. Without the wedge, an initially uniform particle distribution would remain uniform as substrate particles follow the pore fluid flow, which is incompressible and governed by $\nabla \cdot v = 0$. No divergence in the flow field implies no divergence between particles, and thus no unpacking.

Fluidization due to E. directus uplift motion

A fluidized bed is created when fluid flows upwards (against gravity) through a substrate and the resulting pressure gradient, neglecting hydrostatic pressure, supports the weight of the particles. A common example of a fluidized bed is quicksand, where fluid pressure beneath the soil surface is great enough to induce upwards flow and levitate the soil. The bulk fluid flow velocity (as opposed to the flow velocity within the void spaces between particles) to induce fluidization is the same as the settling velocity of a fluidized bed at incipient fluidization. Wen and Yu derived two expressions for the void fraction at incipient fluidization, ϕ' , from tests of different size, density, and shape particles (Wen and Yu, 1966):

$$\frac{1 - \phi'}{\psi^2 (\phi')^3} \cong 11, \quad \frac{1}{\psi (\phi')^2} \cong 14 \quad (5)$$

where ψ is the particle shape factor ($\psi \approx 1$ used for round particles).

Using a series of empirically-derived correlations, we can relate ϕ' to the velocities required for fluidization. Richardson and Zaki (Richardson and Zaki, 1954) reported that the relationship between settling velocity of particles in fluid, v_s , to a single particle's terminal velocity in an infinite fluid, v_t , depends on the void fraction raised to the power n .

$$v_s = v_t \phi^n. \quad (6)$$

To find n , we use the correlation determined by Khan and Richardson (Khan and Richardson, 1989), which relates the void fraction exponent and Archimedes number, Ar , a dimensionless number that corresponds to the motion of fluids due to differences in density:

$$\frac{4.8 - n}{n - 2.4} = 0.043 Ar^{0.57}, \quad Ar = \frac{g d_p^3 \rho_f (\rho_p - \rho_f)}{\mu_f^2}, \quad (7)$$

where g is the gravitational constant, d_p is the particle diameter, ρ_f is the density of the fluid, ρ_p is the density of the particle, and μ_f is the fluid viscosity. The particle terminal velocity can be expressed as $v_t = \frac{Re_{p,t} \mu_f}{\rho_f d_p}$, where $Re_{p,t}$ is the Reynolds number for the particle at terminal velocity. Finally, the Reynolds number can, in turn, be calculated regardless of whether the flow is dominated by viscous or inertial effects using a correlation to the Archimedes number, developed by Gibilaro (Gibilaro, 2001).

$$Re_{p,t} = \left[-3.809 + \left(3.809^2 + 1.832 Ar^{0.5} \right)^{0.5} \right]^2. \quad (8)$$

Combining Eqs. 5-8 enables us to compute the settling velocity, and thus the required bulk upward fluid flow velocity, at the incipient fluidization void fraction. The fluidization velocity at $\phi' = 0.41$ for 1 mm soda lime glass beads is 1.35 cm/s. *E. directus*' uplift velocity measured by Trueman is 1.25 cm/s (Trueman, 1967). The peak uplift velocity during the uplift stroke (shown by segment C on Fig. 5A) measured in our experiments where there was a clearly distinguishable upward jerk before valve penetration was 1.30 ± 0.89 cm/s (mean $\pm$ s.d., $n = 18$). This means *E. directus*' uplift motion induces a velocity in the pore fluid on the order of the velocity required to fluidize the substrate below the animal. Furthermore, 1 mm is near the upper bound of particle size in *E. directus* habitat; using the same uplift velocity, *E. directus* can easily fluidize smaller particles. For example, spherical 0.5 mm soda lime glass beads will reach $\phi' = 0.41$ at an upward flow velocity of 0.5 cm/s.

Burrowing drag reduction due to localized fluidization

Moving through fluidized, rather than static, soil provides considerably less drag resistance for *E. directus*, as stresses – normally concentrated in force chains (Cates et al., 1998) – cannot be transferred between particles. The model in Fig. 9B depicts the flow fields and drag force experienced by *E. directus* while moving through a fluidized substrate. We chose to use a 3D model for this analysis, rather than a 2D model based on our visualizer, because: a) *E. directus* lives in three dimensions; b) the blunt body data in Fig. 9A was gathered in real, 3D ocean substrate; and c) the size of the fluidized zone in Fig. 7 is consistent with other 3D measurements and calculations (Winter, 2011). The flow field around burrowing *E. directus* is composed of summed Couette (u_C) and Poiseuille (u_P) flows (Kundu and Cohen, 2004) in an annulus (Fig. 9B), with

$$u_C = -V \frac{\ln\left(\frac{r}{R_B}\right)}{\ln\left(\frac{R_E}{R_B}\right)} \quad (9)$$

and

1

$$u_p = \frac{1}{4\mu_{eff}} \left(\frac{-dp}{dz} \right) \left[R_B^2 - r^2 + \frac{R_B^2 - R_E^2}{\ln\left(\frac{R_E}{R_B}\right)} \ln\left(\frac{R_B}{r}\right) \right], \quad (10)$$

2

3 where V is the penetration velocity of *E. directus*, R_E is the animal's radius, R_B is the
 4 burrow radius, r is the radial coordinate from the center of animal, μ_{eff} is the effective
 5 viscosity of the liquid and particle mixture, p is the pressure driving the Poiseuille flow,
 6 and z is the axial coordinate. From conservation of mass, the volume per unit time
 7 displaced by *E. directus* must equal the volumetric flow rate of liquid/particle mixture
 8 pushed upwards past the animal's body: $\pi V R_E^2 = \int_{R_E}^{R_B} (u_c + u_p) 2\pi r dr$. Evaluating this
 9 integral yields an expression for the pressure differential driving the Poiseuille flow.
 10 Once this differential is known, the total force acting on *E. directus* can be found by
 11 summing the forces caused by the pressure differential and skin friction:

12

$$F_D = \pi R_E^2 \frac{-dp}{dz} L + 2\pi R_E L \mu_{eff} \left. \frac{d}{dr} [u_c + u_p] \right|_{R_E}. \quad (11)$$

13

14

15 In order to evaluate Eq. 11, the effective viscosity of the liquid/particle mixture, μ_{eff} ,
 16 must be determined. Figure 9A juxtaposes *E. directus*' pulling capabilities in static
 17 substrates, determined through tests with the blunt body in Fig. 2, and drag forces
 18 calculated with Eq. 11 using the effective viscosity models in Table 2, which depend on
 19 void fraction at incipient fluidization.

20

21

22

23

24

1 **Table 2: Effective viscosity models depending on void fraction**

	Expression	Reference
$\mu_{eff} =$	$\mu_f \left(\frac{9}{8} \right) \frac{\left(\frac{1-\phi}{1-\phi'} \right)^{\frac{1}{3}}}{1 - \left(\frac{1-\phi}{1-\phi'} \right)^{\frac{1}{3}}}$	(Frankel and Acrivos, 1967)
	$\mu_f \left(1 - \frac{1-\phi}{1-\phi'} \right)^{-2.5\phi'}$	(Krieger and Dougherty, 1959)
	$\mu_f \left[1 + \frac{1.25(1-\phi)}{1 - \frac{1-\phi}{1-\phi'}} \right]^2$	(Eilers, 1941; Ferrini et al., 1979)
	$\mu_f \left(1 - \frac{1-\phi}{1-\phi'} \right)^{-2}$	(Maron and Pierce, 1956)

2

3

4 As the void fraction at incipient fluidization in *E. directus* habitat was not known, we
5 estimated $\phi' = 0.399$, which is the average of the expressions in Eq. (5 with round
6 particles. Variation in burrow radius corresponds to the zone in Fig. 7 where the mean
7 plus one standard deviation in void fraction is above the incipient fluidization threshold
8 ($1 < r/R_E < 3$). The fluidized void fraction was taken to be the largest mean value in Fig.
9 7, $\phi = 0.411$. The radius used for the animal was $R_E = 1.6$ cm. Maximum downwards
10 velocity (10 cm/s) and pulling force of the animal (10 N) were measured by Trueman
11 (Trueman, 1967). Although *E. directus* is too weak to move through static soil, Fig. 9A
12 demonstrates that moving through fluidized soil lowers drag forces to within the animal's
13 strength capability.

14

15 ***Energetic savings from burrowing through fluidized substrate***

16 Data adapted from Trueman's measurements of burrowing *E. directus* (Trueman, 1967)
17 compared to the blunt body energetics calculated from force data in Fig. 9A show that

moving through fluidized, rather than static, soil reduces the amount of energy *E. directus* has to expend to reach full burrow depth by an order of magnitude (Fig. 10). In an idealized fluidized medium, the drag force on a body does not depend on depth. Moving through a packed, static particulate medium, as in the case of the blunt body, requires pushing force that increases linearly with depth (Robertson and Campanella, 1983). As a result, burrowing energy, calculated by $E = \int F dz$, scales linearly with depth for *E. directus*, rather than depth squared for the blunt body.

ACKNOWLEDGEMENTS

This work was funded by the Battelle Memorial Institute, Bluefin Robotics, and the Chevron Corporation. The authors would like to thank C. Becker, M. Bollini, D. Dorsch, J. Kao, K. Ray, and D. Sargent for their experimental and analytical assistance.

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15
16

FIGURE LEGENDS

Figure 1: *E. directus* burrowing motions. Dashed line indicates depth datum. White arrows denote valve and foot motions. Panels are snapshots of live *E. directus* performing burrowing motions. Red silhouette denotes valve geometry in expanded state, before contraction. A) Start of burrowing cycle. B) Foot extends downward. C) *E. directus* pushes on foot, causing upstroke of valves. D) Valve contraction, which pushes blood into the foot, inflating it to form a terminal anchor. E) Contraction of the foot, which draws valves downward. F) Expansion of the valves, at the start of the next burrowing cycle.

Figure 2: Blunt body drag measurement device. The end of the device is made from real *E. directus* valves in their natural configuration. Pushing it into soil gives a measure of drag forces experienced by *E. directus* without localized fluidization. Force measurements are read from the fish scale.

Figure 3: Burrowing visualization system. A) Visualization tank filled with 1 mm soda lime glass beads. Viewing panes are adjustable via lead screws to match tank and animal width, forcing a plane strain condition. Labeled regions: FV, float valve; A, accumulator; LS, lead screw; L, 500 W halogen light; B, bellowed sidewall; and S, substrate. B) Schematic of *E. directus*' orientation within the visualizer. Green arrows denote principal directions of valve movement. C) Flow diagram of the visualizer's cooling and recirculation system. Chilled water, fed by gravity from a commercial lobster tank (LT), is mixed with warm water from the visualizer and fed into a pump. Water is pumped into the accumulator, which maintains level in the visualizer via a float valve. The outlet from the accumulator feeds both the visualizer and a "short circuit" line back to the lobster tank. As the resistance through the visualizer, R_V , is much more than the short circuit line resistance, R_L , nearly all of the water flowing from the pump and into the visualizer is fresh from the lobster tank.

Figure 4: Clam tracking and soil unpacking from PIV data. A) Grayscale image from raw video showing *E. directus*' valves and foot. Red box denotes boundary of the valves,

determined by tracking software. Yellow box outlines the area of substrate investigated with PIV. Soil and valve displacements were measured in reference to the animal's initial valve position, shown by the white coordinate system. B) Example of soil displacement and unpacking after valve contraction (Fig. 1D) and before valve penetration (Fig. 1E). Horizontal displacement from PIV data, δ_x (green line), have been averaged over the height of the interrogation window to produce a function that varies only in horizontal position, x . Dashed line is an exponential fit to the data. Blue line is the void fraction in the soil, ϕ , calculated from the exponential fit. All lengths have been non-dimensionalized by *E. directus*' initial radius, R_E .

Figure 5: Measured burrowing kinematics. A) Vertical and radial valve positions, δ_E and r_E respectively, of an *E. directus* specimen during five consecutive burrowing cycles. Sections of the burrowing cycle that correspond to Figs. 1C-F are labeled on the third cycle. Dashed lines show initial positions. Lengths have been non-dimensionalized by *E. directus*' initial radius, R_E .

Figure 6: Soil movement around burrowing *E. directus*. Horizontal and vertical displacement fields from PIV data, δ_x and δ_y respectively, are plotted together as displacement magnitude, non-dimensionalized by *E. directus*' initial radius, R_E . Data are overlaid on original video frames used for PIV. The animal's body is masked from the data. Color bar spans 0.001 to 0.07. Timeline shows progression of burrowing events. A) Initiation of burrowing cycle. B) Completion of valve uplift (Fig. 1C). C) Valve contraction (Fig. 1D), immediately before valve penetration (Fig. 1E). D) Moment when failure wedge fully forms, occurring after retraction of foot and downwards pull on valves (to Fig. 1E). Predicted failure wedge angle, θ_f , calculated from substrate friction angle and shown with white dashed line.

Figure 7: Localized fluidization around burrowing *E. directus*. Each curve corresponds to void fraction in the substrate, ϕ , as a function of horizontal position, x , for 23 burrowing tests taken at the point of valve contraction (Fig. 1D), immediately before valve

penetration (Fig. 1E). Position has been non-dimensionalized by *E. directus*' initial radius, R_E . Solid lines calculated from PIV horizontal displacement data (red, animal a; black, animal b; blue, animal c). Gray dotted lines are extrapolations through the shadow zone between the animal's valves and the section of soil investigated with PIV (Fig. 4A). Gray bars are average void fraction for all tests (including extrapolated data) in characteristic lengths away from the animal. Black dashed line is incipient fluidization void fraction. These data show that soil proximate to *E. directus* becomes fluidized during valve uplift and contraction.

Figure 8: Induced soil failure by valve contraction. Mohr's circle of equilibrium (a) and active failure (b) stress states. Inset b shows the formation of a failure surface at angle θ_f and the tendency of soil to naturally landslide during valve contraction, indicated by red arrows. Symbols: ϕ , friction angle; τ , shear stress; σ , normal stress; σ'_{h0} and σ'_{v0} , horizontal and vertical effective stress, respectively, at equilibrium; σ'_{hf} and σ'_{vf} , horizontal and vertical effective stress, respectively, at failure.

Figure 9: Drag forces on burrowing *E. directus*. A) Drag force as a function of depth for static and fluidized soil. Gray bar is *E. directus*' maximum pulling force measured by Trueman (Trueman, 1967). Data points from penetration tests in *E. directus* habitat using the blunt body in Fig. 2 (error bars $\pm$ s.d., $n = 13$). Drag force ranges in fluidized substrate calculated using $1 < R_B/R_E < 3$ and the following liquid/particle viscosity models: F&A (Frankel and Acrivos, 1967); K&D (Krieger and Dougherty, 1959); E+F (Eilers, 1941; Ferrini et al., 1979); M&P (Maron and Pierce, 1956). The plot shows that although *E. directus* is not strong enough to move through a static substrate, drag forces experienced in fluidized substrates are within its strength capabilities. B) Model of liquid/particle flow around *E. directus* used to predict fluid drag. Total flow around animal is sum of Couette (u_C) and Poiseuille (u_P) flow fields. Variables: F_D , pulling force, which is equal to drag force; R_B , burrow radius; R_E , *E. directus* radius; L , *E. directus* length; V , *E. directus* velocity.

1 Figure 10: Energetic savings achieved via localized fluidization burrowing. The blunt
2 body in Fig. 2 requires an order of magnitude more energy to reach full burrow depth
3 than real *E. directus* (error bars $\pm$ s.d., $n = 13$). Using localized fluidization, burrowing
4 energy scales linearly with depth, rather than depth squared for moving through static
5 substrate.

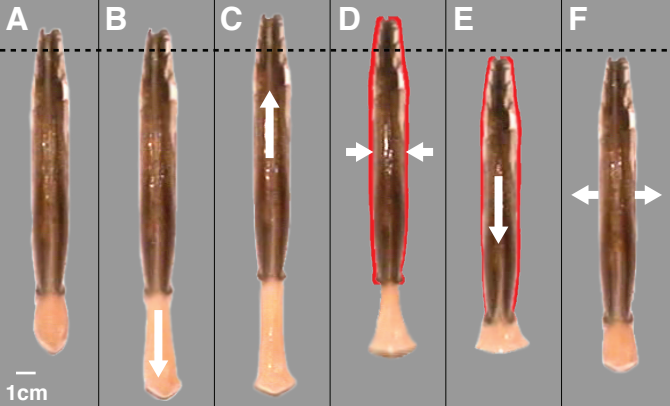


Figure 1

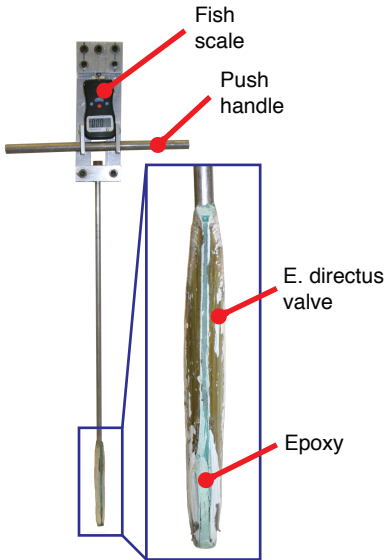


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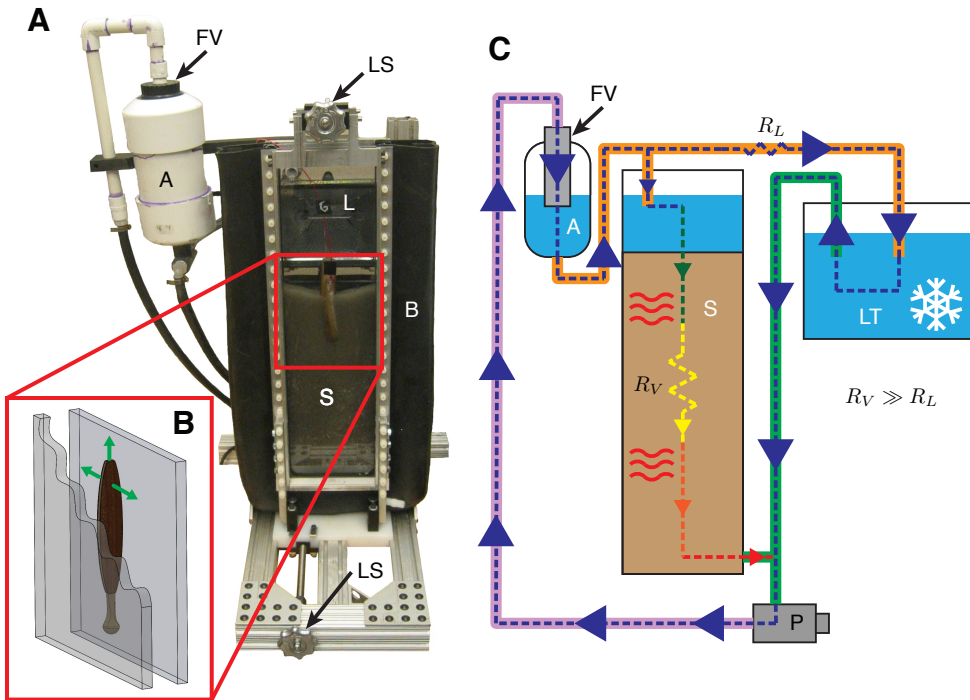


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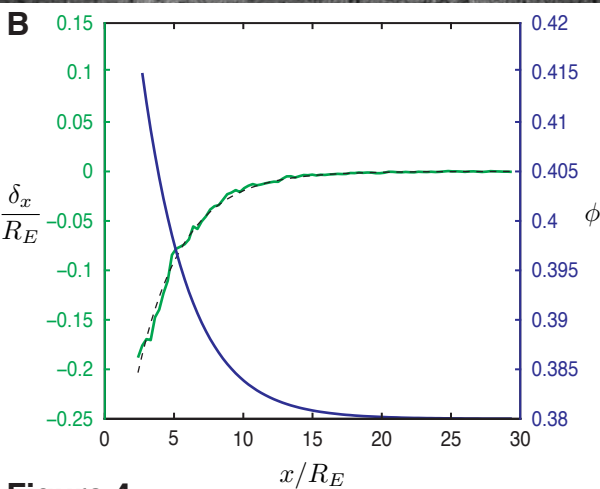
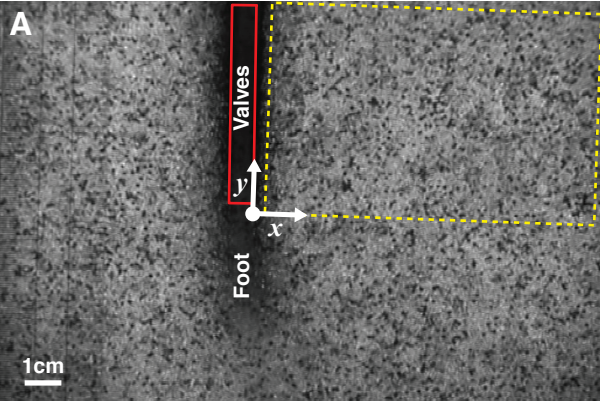


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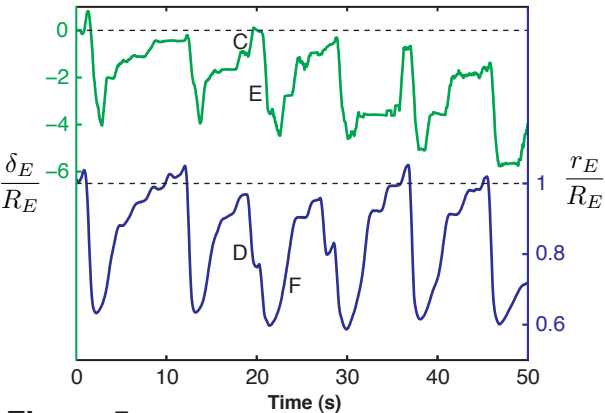


Figure 5

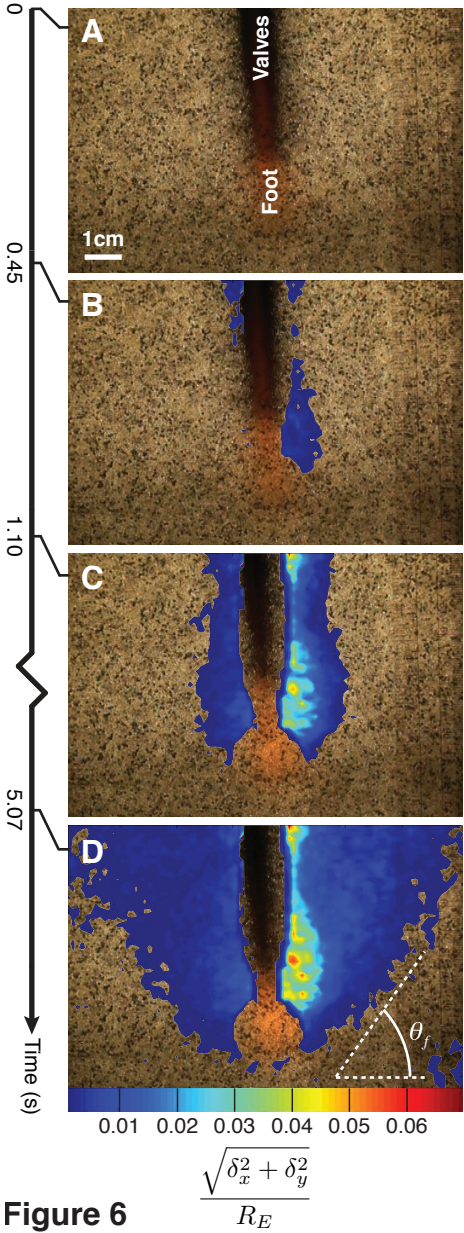


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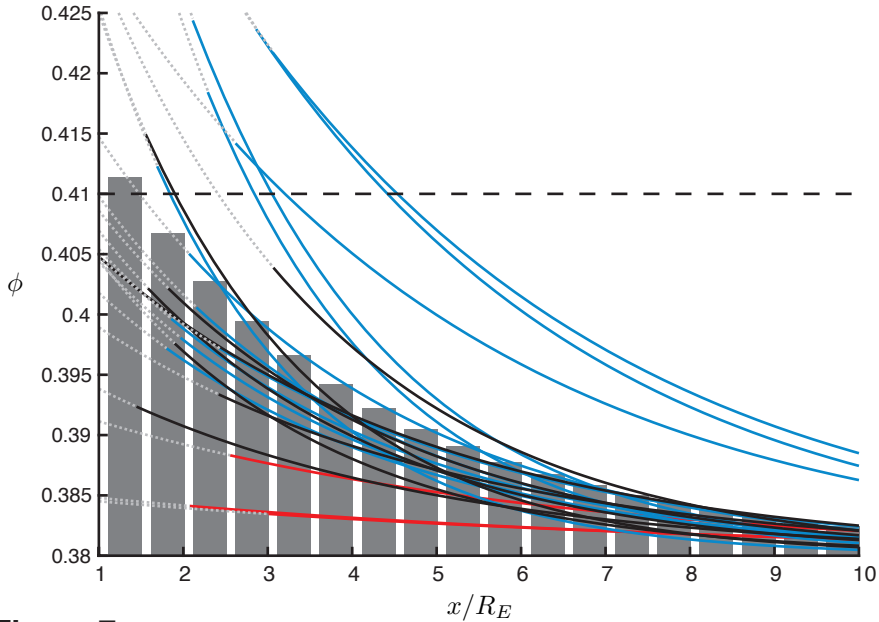


Figure 7

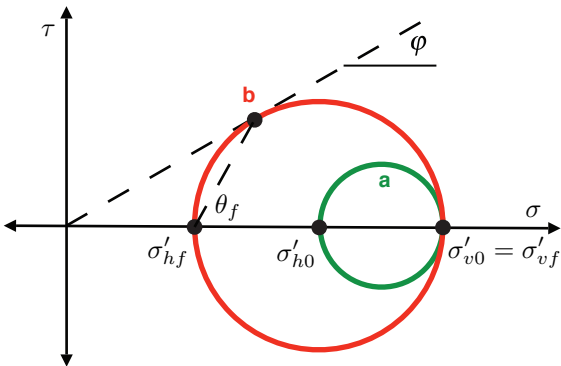
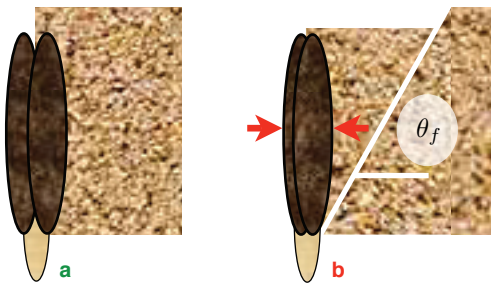


Figure 8

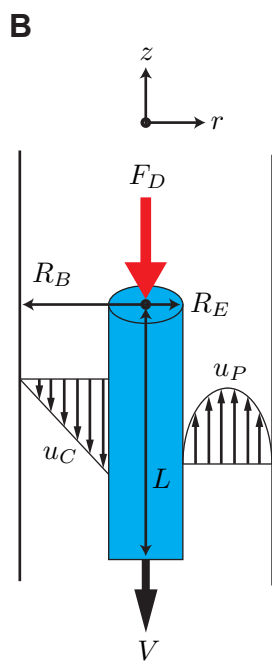
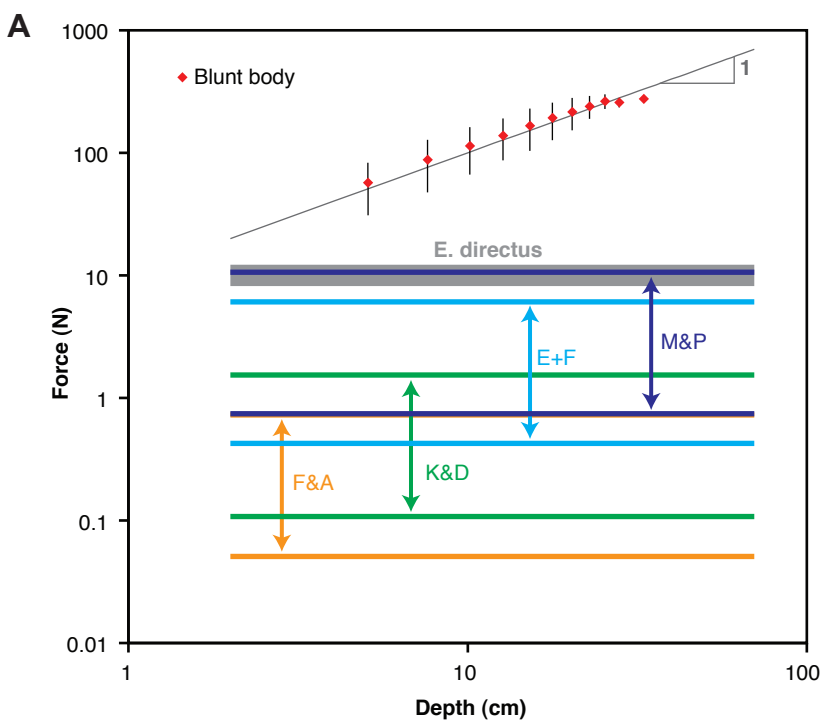


Figure 9

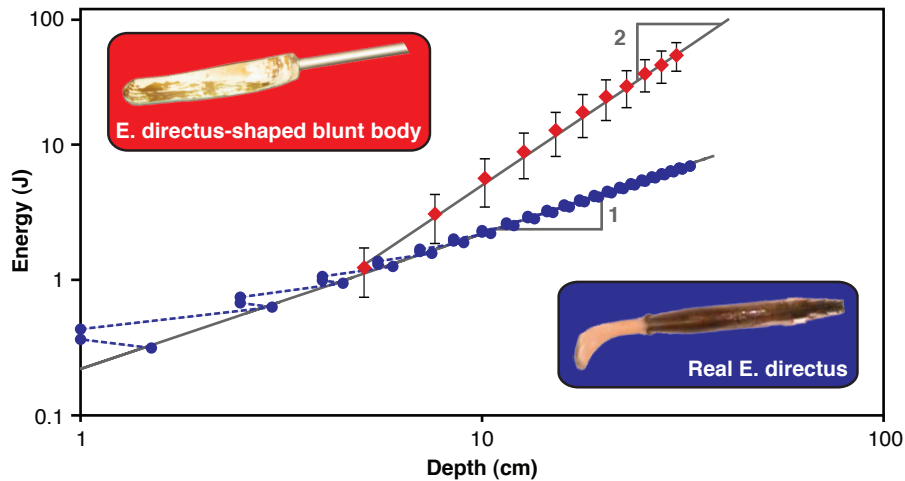


Figure 10