



Cite this: *Soft Matter*, 2021, 17, 7554

Received 19th April 2021,  
Accepted 10th July 2021

DOI: 10.1039/d1sm00564b

[rsc.li/soft-matter-journal](http://rsc.li/soft-matter-journal)

## Agile reversible shape-morphing of particle rafts†

Kyungmin Son,<sup>a</sup> Jeong-Yun Sun<sup>b,c</sup> and Ho-Young Kim<sup>a</sup>\*

Materials that transform shapes responding to external stimuli can bring unprecedented innovations to soft matter physics, soft robotics, wearable electronics, and architecture. As most conventional soft actuation technologies induce large deformations only in a preprogrammed manner at designated locations, the material systems capable of agile reversible deformations without prescribed patterns are strongly desired for versatile mechanical morphing systems. Here we report a morphable liquid interface coated with dielectric particles, or a particle raft, which can reversibly change its topography under an external electric field. The rafts change from flat floors to towers within seconds, and the morphed structures are even capable of horizontal translation. Our experiments and theory show that the raft deformation is driven by electrostatic attraction between particles and electrodes, while being modulated by electric discharge. A broad range of materials serving as electrodes, *e.g.*, human fingers and transparent polymers, suggests this system's diverse applications, including the human-machine interface and the three-dimensional physical display.

## Introduction

Shape-morphing structures, aimed to provide enhanced dynamic functionality to robotics,<sup>1–3</sup> flexible electronics,<sup>4–6</sup> biomedicine,<sup>7,8</sup> and smart architecture,<sup>9,10</sup> have employed either highly stretchable materials or geometrical modifications of surfaces (*e.g.*, folds and cuts). Stimulus-responsive soft materials – such as elastomers<sup>11–15</sup> and hydrogels<sup>16–21</sup> – can undergo a great degree of volume change, which can lead to a transformation of structures embedding those materials. Origami<sup>22–26</sup> and kirigami,<sup>27–29</sup> respectively, employing folds and cuts of surfaces, can convert two-dimensional objects into three-dimensional structures. However, the deformation patterns and locations should be prescribed in these schemes, implying that the structures are set to be transformed only into a designated shape. Although ferrofluids can undergo large deformations without prescribed patterns in response to an external magnetic field,<sup>30–32</sup> it is nearly impossible to achieve localized stable shape equilibrium due to the Rosensweig instability<sup>33</sup> and interfacial instabilities induced by magnetic attraction.<sup>34,35</sup> Therefore, a material system capable of elaborate but stable large deformation without prescribed

patterns is strongly called for to enhance the versatility of morphing systems.

A promising material system capable of such dramatic shape change is a liquid interface coated with particles, which behaves as a two-dimensional solid with superior deformability. The elasticity of this composite interface is owing to capillary forces providing inter-particle cohesion,<sup>36</sup> and its fluidity comes from the underlying liquid. Depending on whether particles cover the free surface of a liquid pool or a liquid drop, the interfacial system is called a particle raft<sup>36</sup> or a liquid marble.<sup>37</sup> Particle rafts can buckle and wrinkle under compressive stress,<sup>36</sup> fracture when in contact with surfactants,<sup>38</sup> and encapsulate overlying fluids upon sinking.<sup>39</sup> Liquid marbles can be divided,<sup>40</sup> merged,<sup>41</sup> and deformed into different shapes under static and dynamic loadings.<sup>37</sup> However, no attempts have been made yet to achieve controllable and reversible shape-morphing of these particle-coated interfaces.

Here, we explore the deformability of the particle raft to give rise to dramatic interfacial deformations of extraordinary accuracy, speed, and controllability without prescribed patterns, which has been impossible with conventional liquid-gas or solid-fluid interfaces. Utilizing an electric field, we experimentally show that the particle raft can be locally upheaved to a tall mound, pulled to a tower, and translated horizontally. We theoretically explain the physical principles behind the morphing behaviors, which are essential for the precise control of the raft deformations. As a broad range of materials can serve as electrodes, our shape-morphing technology can find versatile applications, including in three-dimensional human-machine interfaces.

<sup>a</sup> Department of Mechanical Engineering, Seoul National University, Seoul 08826, Korea. E-mail: [hyk@snu.ac.kr](mailto:hyk@snu.ac.kr)

<sup>b</sup> Department of Materials Science and Engineering, Seoul National University, Seoul 08826, Korea. E-mail: [jysun@snu.ac.kr](mailto:jysun@snu.ac.kr)

<sup>c</sup> Research Institute of Advanced Materials (RIAM), Seoul National University, Seoul 08826, Korea

† Electronic supplementary information (ESI) available. See DOI: 10.1039/d1sm00564b



## Experimental

The particle raft consists of hollow glass spheres (Cospheric HGMS-0.09) densely covering a 5 mm thick film of castor oil (Sigma-Aldrich 83912). Glass particles are sprinkled over the oil film until they completely fill the interface, and the excess particles on the particle raft are removed using a gentle wind. The mean diameter and density of glass spheres are 108  $\mu\text{m}$  (standard deviation of 9.2  $\mu\text{m}$ ) and 90  $\text{kg m}^{-3}$ , respectively, and the exterior surface of the particles are smooth with a surface roughness less than 2  $\mu\text{m}$ . The castor oil has a density of 960  $\text{kg m}^{-3}$ , a relative permittivity of 4.7, and an electric conductivity of 45  $\text{pS m}^{-1}$ . It has a high polarity being composed of approximately 90% of an unsaturated hydroxyl 18-carbon fatty acid. Its contact angle with borosilicate glass and soda-lime glass was measured to be 35° and 37°, respectively, which are significantly higher than the contact angle of silicone oil with glass (9° to 10°). Therefore, glass particles gently scattered on the castor oil float on the air–oil interface without being completely wet while those on the silicone oil are completely wet even without forced immersion, as schematically compared in Fig. 1(c) and (d). The liquid film sits on a grounded stainless-steel plate, and a 10 mm diameter circular stainless steel disk electrode is located above the liquid–air interface.

As a means of controlled stimulation to the soft interface, we employed an electric field through the experimental setup shown in Fig. 1(a). The electric potential difference,  $\phi$ , was applied between the bottom ground plate and a metallic disk above the liquid. The electrostatic force per volume,  $\mathbf{f}$ , acting on dielectric media is caused by free charges in the media,  $\rho_e \mathbf{E}$ , and dipole force due to an abrupt jump in polarizability,<sup>42</sup>  $E^2 \nabla \epsilon / 2$ , where  $\rho_e$  is the volume density of free charge,  $\mathbf{E}$  is the electric field with a magnitude of  $E$ , and  $\epsilon$  is the permittivity of the medium. Because none of the leaky dielectric liquid (castor oil) and dielectric particles (hollow glass microspheres) employed in our experiments have a free charge ( $\rho_e = 0$ ) except at surfaces, the force acting on either the particle raft or the uncoated oil comes only from the interface where the permittivity changes ( $\nabla \epsilon \neq 0$ ) and thus surface charges can build up. We note that, although interfacial deformation by electro-dipping force is observed at the nonpolar fluid and water interface,<sup>43,44</sup> the dipole field effect is negligible at the air–castor oil interface due to the small difference in permittivity.

## Results and discussion

### Electrostatically induced deformation of interfaces

A static interface of a leaky dielectric oil slightly bulges under a weak electric field thanks to the charge accumulated at the air–oil interface as shown in Fig. 1(b) ( $t = 0$  s).<sup>45</sup> The short hump is destabilized upon the electric field exceeding a critical value. It is attributed to a quick surface charge build-up by charge convection in the leaky dielectric liquid. The electric field thus formed generates a strong electric shear force tangential to the tilted interface that exceeds the stabilizing forces of gravity and surface tension, accelerating the liquid toward the electrode to

form a jet (Fig. 1(b),  $t > 0$ ).<sup>46</sup> When dielectric particles covering the liquid interface are completely wetted due to forced immersion, the composite interface is also destabilized under a high electric field to generate a jet, as shown in Fig. 1(c). It is because the liquid layer still covering the entire surface including particles allows surface charge transfer that generates electric shear stress.

On the contrary, a raft of dry dielectric particles floating on the leaky dielectric liquid does not exhibit such instability under a high electric field. Rather, the particle raft reaches a static equilibrium as shown in Fig. 1(d) ( $t = 3.5$  s). Because surface charge convection does not occur along the insulating particles at the interface, the electric shear stress generated is relatively small. In addition, the layer of particles can support the anisotropic shear stress,<sup>36</sup> which consequently suppresses shear stress driven flow (see Appendix B, Fig. 9 for particle packing on the raft). Thus, the particle raft behaves like a perfectly insulating medium where the deformation is only driven by electric normal stresses caused by the polarization of the medium. The temporal evolution of the peak height of the three interfaces in Fig. 1(b)–(d), measured after the electric potential  $\phi$  reaches 10 kV, is plotted in Fig. 1(e). Although the humps rise initially toward the electrode at similar rates regardless of the interface types, only the raft of partially wetted particles reaches a steady height, uncovering the remarkable capability of the electrostatic scheme to induce agile but stable shape morphing. Fig. 1(d) ( $t > 7$  s) also shows that the raft returns to the original flat floor when the electric potential is removed, revealing the reversibility of the deformation. This corresponds to the most fundamental raft morphing mode, *i.e.* upheaving (see Movie S1, ESI†).

The steady raft height under constant electric potential and electrode height implies the balance of the forces acting on the raft. However, this balance contradicts the conventional understanding of the movement of a dielectric medium between capacitor plates. As a liquid of a higher permittivity occupies a larger portion between the electrodes, the capacitor's increased charge due to the increased polarization density strengthens the electric field. The resulting increase in the force acting on the medium should accelerate the interface to displace the less polarizable medium and touch the initially distant electrode, as was observed in Fig. 1(b) and (c), and reported for polymer–air interfaces<sup>47</sup> in a similar setting.

### Analysis of upheaving particle raft

Our particle raft with a dry interface experiences a similar initial bulging process to that of a polarizable medium in the capacitor. However, when the electric field between the particles and the electrode exceeds the dielectric strength of air, electric discharge takes place. Ions are drifted from the electrode to the surface of insulating glass particles, thereby offsetting the opposite charges originating from polarization. Thus, the electric field is reduced in the air gap, allowing the weakened electrostatic force to be balanced with the gravitational and capillary forces resisting the raft deformation. The process leading to force equilibrium of an upheaved raft is illustrated in Fig. 2(a).





**Fig. 1** Electrostatically induced deformation of interfaces. (a) Schematic of the experimental setup. (b) and (c) A castor oil (b) and a raft of fully wet hollow glass spheres on castor oil (c) develop jets under a strong electric field perpendicular to the unperturbed interface. (d) A raft of partially wet hollow glass spheres on castor oil is upheaved to an equilibrium height without instability, such as ejection, detachment, and interfacial jetting. The upheaved raft due to a high electric potential ( $\phi$ ) descends when the potential is removed, to finally return to a flat interface. (e) The changes in the height of the humps of each interface corresponding to (b), (c), and (d) as a function of time under an electric potential of 10 kV. The time is set to be zero when the ascent speed of each interface is  $0.1 \text{ mm s}^{-1}$ . In all three interface conditions, the top electrode is 6.3 mm above the unperturbed interface.

The electric discharge was evidenced by the electric current through the air gap, which was measured *via* a current meter (Keithley 2450, Tektronix) connected to the bottom electrode. As shown in Fig. 2(b), the current rose abruptly from approximately 0.2 to 2 nA when the bulging raft approached the top electrode. We note that if a conductive, instead of leaky dielectric, liquid is used, much stronger discharge occurs, which destabilizes the electric field and thus prevents stable transformation of the raft into a mound.

We can predict the fate of the raft as we increase the electric potential  $\phi$  or the distance of the top electrode from the unperturbed free surface  $h$ , based on a regime map for the steady-state raft response in Fig. 2(c). We start the raft's morphing with a short hill formed under a weak electric field corresponding to a large electrode distance and a low potential (the lower right regime, indicated as A). To increase the electric field, we either increase the potential at a fixed electrode distance (path I) or decrease the electrode distance at a fixed potential (path II). When the regime boundary denoted as a green line is met in the path, the raft experiences a strong uplift, which may continue to eventually result in contact of the

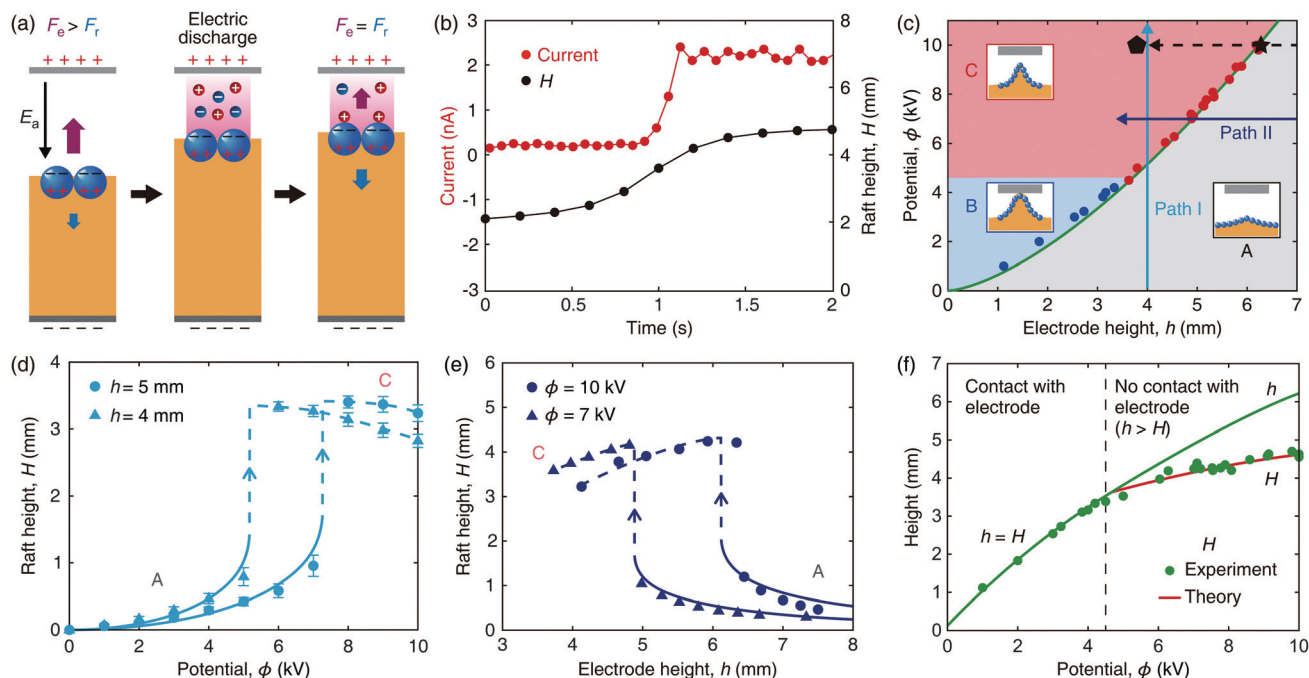
raft and the top electrode (regime B) or stop at a stable height (regime C) (see Appendix B, Fig. 10). Fig. 2(d) and (e) show the raft response to the change of  $\phi$  (path I) and  $h$  (path II), respectively, as we move from regime A to C.

In order to theoretically explain the raft behavior, we first analyze the height  $H$  of a short hill in regime A, which is determined by the electric field between the electrodes. The electric field is related to the raft height through the balance of the electrostatic force with gravitational and capillary forces:

$$\frac{1}{2}E^2\Delta\epsilon = \rho g\hat{H} + \gamma_p\nabla^2\hat{H}, \quad (1)$$

where  $\Delta\epsilon$  is the jump of the permittivity between castor oil ( $\epsilon_c$ ) and air ( $\epsilon_a$ ),  $\rho$  is the density of castor oil,  $g$  is the gravitational acceleration,  $\gamma_p$  is the equivalent surface tension coefficient of the particle raft, and  $\hat{H}$  is the raft height field with the maximum of  $H$  in the center. Here, we do not consider the solid viscoelastic properties of the raft because the lateral capillary forces, the origin of elastic behavior in dilatational rheology, are negligible due to nearly imperceptible deformation of the interface caused by the low density of the particles.<sup>48</sup> However, when the particle





**Fig. 2** Mechanism of electro-morphing of the particle raft. (a) Equilibration of forces on the particle raft. When the raft approaches the electrode, electric discharge takes place, and consequently the electrostatic force  $F_e$  (purple arrow) decreases to be equal to the resisting force  $F_r$  (blue arrow), thereby achieving force equilibrium. (b) Experimentally measured electric current through the air during particle raft deformation under the potential difference  $\phi = 10$  kV with the distance of the top electrode from the unperturbed interface  $h = 6.4$  mm. (c) Regime map of the shape of a particle raft (5 mm thick castor oil coated with hollow glass spheres) in response to  $\phi$  with  $h$ . The raft forms a small hump in regime A, touches the electrode in regime B, and forms a stable mound without touching the electrode in regime C. The green line is plotted by the approximate model,  $\phi = \eta(\rho g/\Delta\epsilon)^{1/2}h^{3/2}$  with  $\eta = 1.2$ . The blue and red points represent the experimental boundaries of A–B and A–C, respectively. (d) The change of raft height from the unperturbed interface,  $H$ , with the increase of  $\phi$  at fixed  $h$  corresponding to path I. (e) The change of  $H$  with the decrease of  $h$  at fixed  $\phi$  corresponding to path II. In (d) and (e), the solid lines are theoretical predictions of our elaborate model given in eqn (A3) in Appendix A, and the dotted lines provide a guide to the eye. (f) The circles correspond to the experimentally measured height of the raft at the green regime boundary of (c). The red line, the theoretical prediction of  $H$  by taking  $A_1/A_2 = 5/6$  in eqn (2), matches the experimental data.

layer is thick, the viscoelastic properties of the wet granular media should be additionally considered, which falls beyond the scope of this work.

As we cross the regime boundary in Fig. 2(c), the electrostatic force increases faster than the resisting forces due to capillarity and gravity, so that the raft rise cannot be stopped as in regime A. This boundary, or the conditions provoking the abrupt jump of  $H$  in Fig. 2(d) and (e), can be obtained by considering the force balance given by eqn (1):  $(\phi/h)^2\Delta\epsilon \sim (\rho g + \gamma_p/l_c^2)H$ , where we estimate the magnitudes of  $E$  and  $\nabla^2 H$  as  $\phi/h$  and  $H/l_c^2$ , respectively, with the capillary length  $l_c = [\gamma_p/(\rho g)]^{1/2}$ . Taking  $H \sim h$ , the potential at the regime boundary is scaled as  $\phi = \eta(\rho g/\Delta\epsilon)^{1/2}h^{3/2}$  with  $\eta$  being an empirical prefactor, implying that the potential to upheave the raft into regime C increases like  $h^{3/2}$ . We present a mathematically more elaborate model for the raft height in regime A and the regime boundary in Appendix A (see also Fig. 6 for the model used to analyze a non-uniform electric field). We see that the regime boundaries predicted by the current approximate analysis and the elaborate model are fairly close (Fig. 7), verifying our force argument leading to the raft upheaval.

Fig. 2(d) and (e) show that our elaborate model (eqn (A3) in Appendix A) can predict the variation of  $H$  with  $\phi$  and  $h$  before

the drastic uplift occurs (solid lines). After the sudden upheaval, the raft height is rather decreased with the increase of  $\phi$  and the decrease of  $h$  (broken lines). Such a repulsive behavior of the top electrode against the raft in regime C arises because the raft height is adjusted to maintain the electric field in the air gap at the dielectric strength of the air. Therefore, the raft height is maximized along the green regime boundary in Fig. 2(c) in the upheaving mode.

We now predict the maximum height of the raft, which is reached as a result of electric discharge. The above model is not suitable for the height of the upheaved raft owing to severe deformation of the interface and different force balance caused by discharge. We note that the electric field in the air gap has reached the dielectric strength of air,  $E_s = 3$  kV mm<sup>-1</sup>, when the raft is in equilibrium by discharge. We build a one-dimensional capacitance model [see Appendix B, Fig. 11(a)] where the electric field in the air gap is  $E_a = \phi_a/(h - H)$  with  $\phi_a$  being the potential difference across the air gap. The capacitance in the air and castor oil, respectively, given by  $C_1 = \epsilon_a A_1/(h - H)$  and  $C_2 = \epsilon_c A_2/(h_c + H)$ , interrelate *via* charge conservation:  $\phi_a C_1 = (\phi - \phi_a)C_2$ , where  $A_1$  and  $A_2$  are the effective areas of the capacitors, and  $h_c$  is the oil layer thickness. By equating  $E_a$  with  $E_s$ , we find the raft height at the boundary of regimes A





and C as the following:

$$H = \frac{(h - \phi/E_s)\epsilon_c/\epsilon_a + h_c A_1/A_2}{\epsilon_c/\epsilon_a - A_1/A_2}. \quad (2)$$

Fig. 2(f) plots the height of raft  $H$  along the green regime boundary in Fig. 2(c). Below the critical values of  $\phi$  and  $h$  (boundary between regimes A and B), the raft touches the top electrode, so that  $h = H$ . Beyond the critical point, we find that the theoretically predicted maximum raft height of eqn (2) is in good agreement with the experiment.

### Shape control of the particle raft

Once we generate a tall mound by reaching the aforementioned regime boundary, we can further control the raft morphology by moving the top electrode either upward or horizontally. When the electrode slowly moves upward, the mound is vertically pulled to turn to a tower, as shown in Fig. 3(a) and (b), which we term the remote pulling mode (see Movie S1, ESI<sup>†</sup>). We note that the already upheaved tall tower can keep standing even in regime A of Fig. 2(c). This is because once the raft has entered regime C, the force balance governed by the electric discharge can be maintained even when we follow path II backward to re-enter regime A. We note that repeating the remote pulling mode at the same location may leave a small trace as a result of local particle accumulation. This weakly hysteretic behavior, characteristic to viscoplastic systems, is associated with the plastic transition by the formation of intense force chains in large deformation.<sup>49</sup>

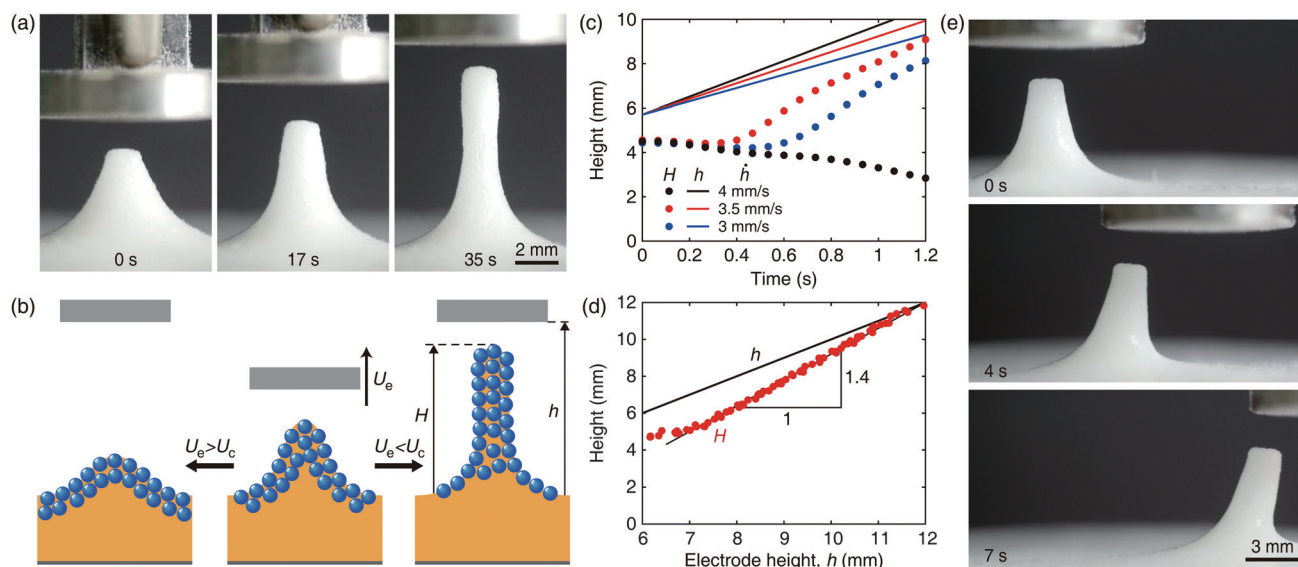
Fig. 3(c) plots the temporal evolution of the mound height,  $H$ , for different electrode ascent speeds  $U_e$ , revealing that the

raft can follow the electrode rising with a speed lower than the critical speed  $U_c \approx 4 \text{ mm s}^{-1}$  (see the ESI<sup>†</sup>). The slight decrease of the raft height in the initial stages for  $U_e = 3$  and  $3.5 \text{ mm s}^{-1}$ , as shown in Fig. 3(c), is because particles initially gathered toward the ascending electrode squeeze liquid out while narrowing the tower. The particle aggregate then grows vertically following the electrode with a nearly constant width.

The vertical growth rates of the particle aggregates are higher than the rising speed of the electrode as seen by greater slopes of the solid lines than the dotted lines in Fig. 3(c). Hence, the tower cannot be elongated indefinitely but rather reaches a maximum height at which it touches the electrode. To mathematically understand this remote pulling mode, *i.e.* the relationship between the electrode height  $h$  and the raft height  $H$ , we model the system as a three-layer structure of oil, particle aggregate, and air [see Appendix B, Fig. 11(b)]. In the one-dimensional series connection model of the three capacitors, the potential drops in air ( $\phi_a$ ), particle aggregate ( $\phi_p$ ), and oil ( $\phi_c$ ) are related as  $\epsilon_a A_1 \phi_a / (h - H) = \epsilon_p A_2 \phi_p / H = \epsilon_c A_3 \phi_c / h_c$  through charge conservation. Here,  $\epsilon_p$  is the permittivity of the particle aggregate,  $A_1$ ,  $A_2$ , and  $A_3$  are respectively the area of the equivalent capacitors of air, particle aggregate, and oil. The relation combined with  $\phi = \phi_a + \phi_p + \phi_c$  leads to an equation for  $H$ :

$$H = \frac{h + \epsilon_a A_1 h_c / (\epsilon_c A_3) - \phi / E_s}{1 - \epsilon_a A_1 / (\epsilon_p A_2)}. \quad (3)$$

The sensitivity of the raft height to the electrode height change is quantified by  $H' = dH/dh$ . Assuming  $A_1 \ll A_3$ ,  $A_2 \ll A_3$ , and  $A_1/A_2$  to be constant, we find  $H' = 1/[1 - \epsilon_a A_1 / (\epsilon_p A_2)]$  which is constant. We indeed see in Fig. 3(d) that the raft height



**Fig. 3** Morphology control of the particle raft. (a) Images of a raft tower pulled vertically following the electrode that ascends from  $h = 6.4$  to  $10 \text{ mm}$  at a speed of  $0.1 \text{ mm s}^{-1}$ . (b) Schematics of a particle raft that either collapses or grows depending on whether the electrode ascends at a speed higher or lower than the critical speed  $U_c$ . (c) Temporal evolution of the hill height (solid lines), which starts from  $H = 4.5 \text{ mm}$  with  $h = 5.7 \text{ mm}$ , for different ascent speeds of the electrode,  $U_e = 3, 3.5, \text{ and } 4 \text{ mm s}^{-1}$ . The particle raft grows upward following the electrode when the ascent speed is  $3$  and  $3.5 \text{ mm s}^{-1}$ , but it collapses when  $U_e = 4 \text{ mm s}^{-1}$ . The dotted lines indicate the elevation of the electrodes,  $h$ . (d) The change of the raft height  $H$  as the electrode rises at  $U_e = 0.1 \text{ mm s}^{-1}$ . (e) Horizontal translation of a tower following the electrode horizontally moving at a speed of  $2 \text{ mm s}^{-1}$ . In (a) and (c)–(e), the electric potential  $\phi$  is  $10 \text{ kV}$ .



increases at a constant slope and eventually equals the electrode height. The maximum height of the particle tower  $H_t$  is obtained by taking  $H = h = H_t$  in eqn (3):  $H_t = \phi \epsilon_p A_2 / (E_s \epsilon_a A_1)$ , implying that it increases with the potential difference. An empirically found maximum height  $H_t = 12$  mm for  $\phi = 10$  kV allows us to find the ratio  $\epsilon_p A_2 / (\epsilon_a A_1)$  to be 3.5. Then we get the theoretical slope  $H' = 1.4$ , which agrees well with the experimental results in Fig. 3(d).

Utilizing the attractive force between the upheaved particle raft and the moving electrode, we devised another morphing mode in the horizontal, rather than vertical, direction. The mound or tower originally built by the vertical deformation of the raft was translated horizontally following the electrode, as shown in Fig. 3(e), which is referred to as the horizontal translation mode (see Movie S1, ESI†). Such translation of the raft structure occurred when the electrode moved at a speed under a critical value, which is determined by the balance of the electrostatic attraction and the shear stress that resists the translational motion. We observed that the maximum horizontal velocity of the electrode that the tower could steadily follow was  $6.5 \text{ mm s}^{-1}$ . Above the critical speed, the shear stress acting on the raft becomes so large that the particle tower elongates toward the top electrode and finally collapses.

Fig. 4(a)–(d) show that versatile control of the upheaved raft shape is possible by combining the upheaving mode, following either path I or II in Fig. 2(c), and the remote pulling mode. A relatively slender and tall tower in Fig. 4(b) can be constructed by pulling a slender mound in Fig. 4(a) that has been upheaved under the critical condition in Fig. 2(c). Starting from Fig. 4(a), we can build a relatively wide and tall tower as shown in Fig. 4(d), by first decreasing the electrode height  $h$  following path II (from black pentagram to pentagon) in Fig. 2(c) to reach a state in Fig. 4(c). Then, the wide mound is pulled upward to attain a state in Fig. 4(d). The widening of the mound

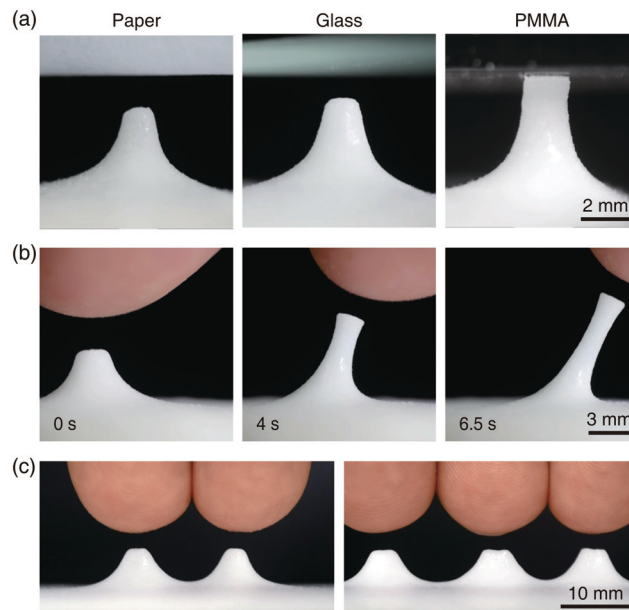


Fig. 5 Electro-morphing with electrodes of diverse materials. (a) Response of particle rafts to grounded top electrodes of low conductivities. The potential of 10 kV is applied to the bottom electrode. (b) A bare finger can upheave and translate the raft. (c) Upheaving of multiple mounds by sensing multiple fingers.

as shown in the process from Fig. 4(a)–(c) is attributed to the enlarged raft region affected by a high electric potential. Using the same principle, we can further increase the mound width by employing a top electrode of a larger diameter, as shown in Fig. 4(e), where the electrode diameter has been doubled in comparison with that in Fig. 4(c) with the other conditions kept identical. Then, a fairly wide and tall tower can be built by pulling the mound in Fig. 4(e) to reach a state in Fig. 4(f) (see the ESI†).

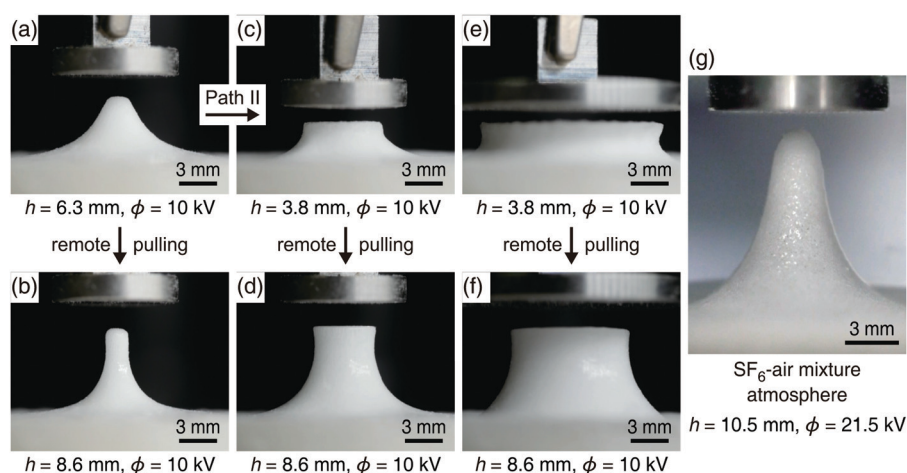


Fig. 4 Various shape control schemes of a raft mound. (a) A raft mound upheaved at a critical potential of 10 kV with a constant top electrode height  $h = 6.3$  mm. (b) A raft tower pulled by the ascent of the top electrode from (a). (c) A raft mound whose width has been increased as the electrode height  $h$  decreased to 3.8 mm from (a), following path II in Fig. 2(c). (d) A wide tall tower pulled by the ascent of the top electrode from (c). (e) A raft mound lifted by a top electrode of 20 mm diameter, twice that of those in (a)–(d). (f) A raft tower pulled by the ascent of the top electrode from (e). (g) A raft mound upheaved at a critical potential of 21.5 kV with a constant top electrode height  $h = 10.5$  mm in the mixture of  $\text{SF}_6$  and air. Movie S2 (ESI†) shows all of the shape control processes in real-time.



To increase the height of an upheaved raft in the upheaving mode alone, we replaced air with a mixture of SF<sub>6</sub> and air that can prevent unwanted electric discharge at a greater electric potential owing to its high dielectric strength (see the ESI<sup>†</sup>). The result is shown in Fig. 4(g), where the particle raft has been upheaved to the height of  $H = 9.55$  mm by a top electrode located at  $h = 10.5$  mm with  $\phi = 21.5$  kV. Such high altitude of the electrode is impossible in normal air because electric breakdown occurs before such a high potential is reached.

### Electro-morphing with electrodes of diverse materials

Our electro-morphing scheme is not restricted to highly conductive metallic electrodes. Rather, materials with moderately low conductivity can be employed as a top electrode. This is because the force balance of the upheaved raft requires only a very low current resulting from an electric discharge, as low as 2 nA (Fig. 2(b)). We indeed see in Fig. 5(a) that particle rafts can be upheaved by employing materials of poor conductivity ( $10^{-10}$ – $10^{-12}$  S m<sup>-1</sup>) like paper and glass, as a top electrode. Using an electrode of even lower conductivity ( $<10^{-14}$  S m<sup>-1</sup>), such as poly(methyl methacrylate) (PMMA), the raft rises with a significant delay but without stopping, as shown in the third panel of Fig. 5(a), because electric discharge can hardly be maintained due to slow charge transfer.

The top electrode can be replaced by a floating electrode, *i.e.* an object with high capacity for charge storage. Fig. 5(b) and (c) show the electro-morphing of the raft with single and multiple bare fingers moving freely above the interface. This is because living tissues of animals and humans have the required high capacity for charge storage.<sup>50</sup> This suggests that our particle raft system can serve as a human-machine interface as well as a sensor or actuator driven by bioelectricity (see Movie S3, ESI<sup>†</sup>). An array of electrodes that upheave the particle raft precisely where desired can construct three-dimensional (3D) topography out of a flat floor (see Movie S3, ESI<sup>†</sup>), suggesting another application of the system as a 3D physical display.

## Conclusions

We have shown that a raft of partially wet dielectric particles can respond to an external electric field to upheave and form a tall stable mound. The particle raft's transformation arises when the electrostatic force exceeds the capillary and gravitational forces, but the uprise is stabilized by the electric breakdown of the surrounding gas. The mound can be further changed in height or width and translated by a varying electric field mediated by the electrode motion. Our theoretical models explain and predict the raft height achieved by the upheaving and the remote pulling modes. Versatile control of the raft shape is enabled by combining such fundamental morphing modes, employing electrodes of different sizes and materials, or changing gas environments. The remarkable shape-morphing capability of our soft composite material system allows us to envision its diverse applications in soft robotics, biomedicine,

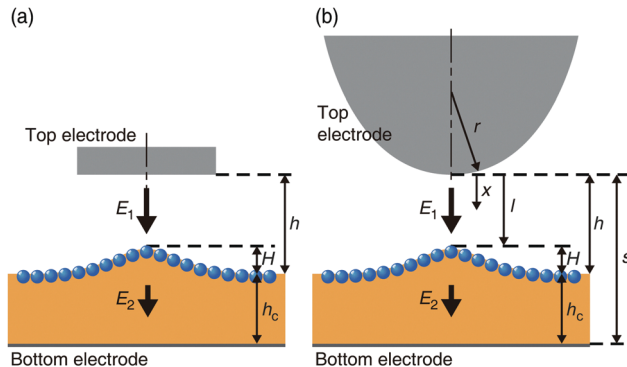


Fig. 6 'Top parabola – bottom plane' model to analyze the non-uniform electric field. (a) Schematic of the experimental setup, where the particle raft lies between a plane and a circular disk electrode. (b) Schematic of a particle raft between a plane electrode and a parabolic tip electrode with radius of curvature  $r$ . The major axis of the tip electrode is perpendicular to the plane. Its apex is at the same distance from the unperturbed interface as the circular disk electrode in (a).

and smart architecture with ever-evolving sophisticated control technologies of the external electric field.<sup>51</sup>

## Appendix A: Theoretical modeling of electro-morphing

### Theoretical modeling of particle raft height in regime A

To model the upheaving behavior of a particle raft under electrostatic forces, we need to analyze the non-uniform field created by a circular disk top electrode and a plane bottom electrode. We base our analysis on a 'top parabola – bottom plane' electrode system (Fig. 6), for which the analytical solution of  $E$  with homogeneous medium between the electrodes is given by  $E(x) = 2\phi/[(r + 2x)\ln(2s/r + 1)]$  with  $s$  being the distance between the parabola tip of curvature  $r^{-1}$  and the bottom electrode.<sup>52</sup> Assuming that the fluid interface does not distort the electric field, we estimate  $E_1 = k_1\phi/[(r + 2x)\ln(2s/r + 1)]$  for air and  $E_2 = k_2\phi/[(r + 2x)\ln(2s/r + 1)]$  for liquid, where  $k_1$  and  $k_2$  are to be determined by the following two conditions. First, continuity of electric displacement across the fluid–fluid interface at  $x = l$ , given by  $\epsilon_a E_1(l) = \epsilon_c E_2(l)$ , yields  $\epsilon_a k_1 = \epsilon_c k_2$  where  $l$  is the distance between the parabola tip and the fluid–fluid interface. Second, the line integral of the electric field from the tip to the plane results in the total potential difference,  $\int_0^l E_1 dx + \int_l^s E_2 dx = \phi$ . The two conditions give  $k_1 = 2\phi\ln(2s/r + 1)/\{\ln[(r + 2l)/r] + (\epsilon_a/\epsilon_c)\ln[(r + 2s)/(r + 2l)]\}$  and  $k_2 = (\epsilon_a/\epsilon_c)k_1$ , which yields the electric field in the air  $E_1$ , and the field in the liquid  $E_2$  as

$$E_1(x) = \frac{2\phi}{r + 2x} \left[ \ln \frac{r + 2l}{r} + \frac{\epsilon_a}{\epsilon_c} \ln \frac{r + 2s}{r + 2l} \right]^{-1} \quad (0 < x \leq l), \quad (\text{A1})$$

$$E_2(x) = \frac{2\epsilon_a\phi}{\epsilon_c(r + 2x)} \left[ \ln \frac{r + 2l}{r} + \frac{\epsilon_a}{\epsilon_c} \ln \frac{r + 2s}{r + 2l} \right]^{-1} \quad (l \leq x < s). \quad (\text{A2})$$

The electric field is related to the raft height through the balance of the electrostatic force with gravitational and





capillary forces:  $E^2 \Delta \varepsilon / 2 = \rho g \hat{H} + \gamma_p \nabla^2 \hat{H}$ , where  $\Delta \varepsilon$  is the jump of the permittivity between castor oil ( $\varepsilon_c$ ) and air ( $\varepsilon_a$ ),  $\rho$  is the density of castor oil,  $g$  is the gravitational acceleration,  $\gamma_p$  is the equivalent surface tension coefficient of the particle raft, and  $\hat{H}$  is the raft height field with the maximum of  $H$  in the center. We estimate  $\nabla^2 \hat{H}$  as  $H/l_c^2$  with the capillary length  $l_c = [\gamma_p/(\rho g)]^{1/2}$ , which allows us to write  $H \approx \beta E_1^2$  with  $\beta \sim \Delta \varepsilon/(\rho g)$ . In the real system with the disk electrode at the top,  $s$  and  $l$  are replaced by  $h + h_c$  and  $h - H$ , respectively. Then the total electric potential  $\phi$  is given by

$$\phi(H, h) = nH^{1/2}[r + 2(h - H)] \times \left[ \ln \frac{r + 2(h - H)}{r} + \frac{\varepsilon_a}{\varepsilon_c} \ln \frac{r + 2(h + h_c)}{r + 2(h - H)} \right], \quad (\text{A3})$$

where the prefactor  $n$ , scaled as  $\sim \beta^{-1/2}$ , can be empirically determined.

### Theoretical boundary of regimes A and C for strong uplift of a particle raft

The abrupt increase of hump height due to a slight increase of  $\phi$  as we cross the regime boundary between A and C *via* path I is mathematically represented by  $\partial H/\partial \phi \rightarrow \infty$  or  $\partial \phi/\partial H = 0$ . As we take path II, the uplift of the raft at the regime boundary corresponds to  $\partial H/\partial h \rightarrow \infty$  or  $\partial h/\partial H = 0$ . Differentiating eqn (A3) with respect to  $H$  with  $h$  fixed gives

$$\frac{\partial \phi}{\partial H} = \frac{1}{2} n H^{-1/2} \left\{ \left[ \ln \frac{r + 2(h - H)}{r} + \frac{\varepsilon_a}{\varepsilon_c} \ln \frac{r + 2(h + h_c)}{r + 2(h - H)} \right] \times (r + 2h - 6H) - 4H \left( 1 - \frac{\varepsilon_a}{\varepsilon_c} \right) \right\}. \quad (\text{A4})$$

If we differentiate  $h$  with respect to  $H$  with  $\phi$  fixed in eqn (A3), we get

$$4H \frac{\partial h}{\partial H} \left\{ \left[ \ln \frac{r + 2(h - H)}{r} + \frac{\varepsilon_a}{\varepsilon_c} \ln \frac{r + 2(h + h_c)}{r + 2(h - H)} \right] + \left( \frac{\varepsilon_a}{\varepsilon_c} \right) \times \left[ \frac{r + 2(h - H)}{r + 2(h + h_c)} - 1 \right] + 1 \right\} + \left[ \ln \frac{r + 2(h - H)}{r} + \frac{\varepsilon_a}{\varepsilon_c} \ln \frac{r + 2(h + h_c)}{r + 2(h - H)} \right] (r + 2h - 6H) - 4H \left( 1 - \frac{\varepsilon_a}{\varepsilon_c} \right) = 0. \quad (\text{A5})$$

Substituting  $\partial \phi/\partial H = 0$  and  $\partial h/\partial H = 0$  into eqn (A4) and (A5), respectively, leads to the same equation, indicating that the conditions for  $\partial \phi/\partial H = 0$  and  $\partial h/\partial H = 0$  for eqn (A3) are equivalent.

Eqn (A4) gives  $H = H^*$ , where  $\partial \phi/\partial H = 0$ , in the following implicit form:

$$\left[ \ln \frac{r + 2(h - H^*)}{r} + \frac{\varepsilon_a}{\varepsilon_c} \ln \frac{r + 2(h + h_c)}{r + 2(h - H^*)} \right] (r + 2h - 6H^*) - 4H^* \left( 1 - \frac{\varepsilon_a}{\varepsilon_c} \right) = 0. \quad (\text{A6})$$

Substituting  $H^*$  obtained in eqn (A6) into eqn (A3) allows us to find the theoretical regime boundary. By taking  $r = 7$  mm and  $n = 1.6 \times 10^7$  kg m<sup>1/2</sup> A<sup>-1</sup> s<sup>-3</sup>, the theoretical curve denoted by a

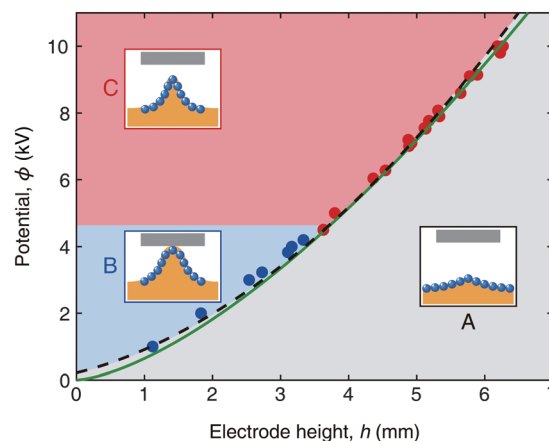


Fig. 7 Comparison of theoretical regime boundaries. The green solid line is plotted from the scaling analysis,  $\phi = \eta(\rho g/\Delta \varepsilon)^{1/2} h^{3/2}$ , by taking the prefactor  $\eta$  as 1.2. The black dotted line is the boundary obtained by eqn (A3) taking  $\partial \phi/\partial H = 0$ .

dotted line in Fig. 7 agrees well with the experimental measurement results. Here, the value of the prefactor  $n$  makes good sense as  $n \sim \beta^{-1/2} \sim 10^7$  kg m<sup>1/2</sup> A<sup>-1</sup> s<sup>-3</sup>.

### The dynamics of raft upheaving

Considering that the ascent speed of the raft in upheaving mode,  $\dot{H}$ , is less than 10 mm s<sup>-1</sup>, the Reynolds number of the upheaving flow,  $Re = \rho \dot{H} l_c / \mu_c$ , is less than 0.1, where we assume the capillary length as the characteristic length. The value of  $Re$  indicates that the inertial effect is negligible in the upheaving flow. Then the force balance equation is written as

$$\frac{1}{2} E^2 \Delta \varepsilon - \rho g \hat{H} - \gamma_p \nabla^2 \hat{H} + \mu_c l_\mu \nabla^2 \hat{H} = 0, \quad (\text{A7})$$

where  $\mu_c$  is the viscosity of castor oil and  $l_\mu$  is the thickness affected by viscosity. If we use eqn (A1) for the electric field and estimate the magnitudes of  $\nabla^2 \hat{H}$  and  $\nabla^2 \dot{H}$  as  $H/l_c^2$  and  $\dot{H}/l_c^2$ , respectively, eqn (A7) becomes

$$E_1^2 \Delta \varepsilon / 2 - c_1 \rho g H - c_2 \mu_c \dot{H} = 0, \quad (\text{A8})$$

with prefactors  $c_1$  and  $c_2$ . This approximated differential equation describes the change in raft height over time.

Fig. 8(a) shows the experimental height change (raft of mode I in Movie S1, ESI<sup>†</sup>) and the theoretical prediction obtained from eqn (A8) with fitting parameters of  $c_1 = 1.5$  and  $c_2 = 5.6 \times 10^3$  m<sup>-1</sup>. The experimental and theoretical curves exhibit a similar trend except for the late stages where the electric discharge effect newly comes in. The speed slows down after the initial rapid ascent and then accelerates again when the raft reaches a certain height. This change in speed is explained by the change in the force acting on the raft as a function of the raft height, which is shown in Fig. 8(b). Up to a height of 2 mm, the decrease in the difference between the electrostatic upheaving force and the other resisting forces slows the ascent. After the height of 2 mm, the difference between the two forces increases dramatically, which rapidly increases the ascent







**Fig. 8** The dynamics of raft upheaving. (a) Comparison of theoretically predicted and experimentally measured height change of the raft as a function of time at a critical potential of 10 kV with a top electrode height  $h = 6.3$  mm. (b) The changes of forces acting on the raft as a function of raft height. Electrostatic force is  $E_1^2 \Delta \epsilon / 2$  and the resisting force including gravitational and capillary force is  $c_1 \rho g H$  with a prefactor  $c_1 = 1.5$ .



**Fig. 9** Local surface coverage of the particle raft. (a) Particle distribution on a flat raft. (b) Top view of the raft upheaving and returning to the flat interface. A transparent ITO (indium tin oxide) electrode is used to observe the raft from the top.

speed. In the experiment, the equilibrium state is reached by an electric discharge after such a jump.

## Appendix B: Supplemental figures

For a particle raft to behave like a perfectly insulating medium, particles must be tightly packed at the interface. Fig. 9(a) shows the uppermost particle layer of the particle raft. Densely packed

layers of particles indicate that the raft can both suppress the surface charge convection and support significant shear stress. Fig. 9(b) also shows that the particle packing deteriorates little during the upheaving and returning process. We neither observed a significant increase in the inter-particle spacing nor particles expelled from the outermost layer during the morphing process, which implies that the particle layer well forms a two-dimensional solid.



**Fig. 10** Particle raft deformation at the regime boundary of Fig. 2(c). The upheaved raft touches the electrode at low  $h$  and  $\phi$  (boundary of regimes A and B). On the other hand, at high  $h$  and  $\phi$  (boundary of regimes A and C), the rafts stop without touching the electrode.





**Fig. 11** Analysis of the equilibrium height of particle rafts using 1D capacitance models. (a) 1D capacitance model with two capacitors arranged in series, each containing air and castor oil as dielectric materials. This model predicts the height of the upheaved raft. (b) 1D capacitance model with a series connection of three capacitors, each containing air, particle aggregate, and oil as dielectric materials. This model predicts the height of the particle aggregate grown by the remote pulling mode.

Following the regime map of Fig. 2(c), Fig. 10 shows the particle rafts reaching equilibrium after deformation at the regime boundary conditions. Under low voltage conditions, the low electric field between the upheaving particle raft and the electrode prevents electric discharge, causing the raft to touch the electrode. At 5 kV or higher, electric discharge occurs and the raft stops without touching the electrode. The gap between the raft and the electrode increases as the voltage increases, which is described in eqn (2) as  $h - H$ .

Fig. 11 presents the equivalent one-dimensional (1D) capacitance model used to predict the maximum height of the upheaved raft. We used a two capacitor series-connected model for the raft deformed by upheaving, and a three-capacitor connected model for predicting the height of particle aggregate grown long by remote pulling. The resistances in the capacitance models can be neglected until the electric field in the air gap reaches the dielectric strength because the current is negligible for very high air resistance ( $R_1$ ) in Fig. 11(a) and (b).

## Author contributions

H.-Y. K. conceived and supervised the research. K. S. conducted the experiments, and K. S., J.-Y. S., and H.-Y. K. analyzed the results. K. S. and H.-Y. K. prepared the manuscript and figures, and all of the authors contributed to the compilation and review of the manuscript.

## Conflicts of interest

H.-Y. K. and K. S. are inventors on the patent application (#10-2020-0170605, Republic of Korea) submitted by SNU R&DB Foundation that covers dielectric fluid and particle composition, and electrical operation methods for the fabrication of three-dimensional deformation systems.

## Acknowledgements

This work was supported by the National Research Foundation of Korea (grant no. 2018-052541) via SNU-IAMD.

## References

- 1 Y. Kim, H. Yuk, R. Zhao, S. A. Chester and X. Zhao, *Nature*, 2018, **558**, 274–279.
- 2 G. Gu, J. Zuo, Z. Ruike, X. Zhao and X. Zhu, *Sci. Robot.*, 2018, **3**, eaat2874.
- 3 S. J. Park, M. Gazzola, K. S. Park, S. Park, V. Di Santo, E. L. Blevins, J. U. Lind, P. H. Campbell, S. Dauth, A. K. Capullini, F. S. Pasqualini, S. Ahn, A. Cho, H. Yuan, B. M. Maoz, R. Vijaykumar, J.-W. Choi, K. Deisseroth, G. V. Lauder, L. Mahadevan and K. K. Parker, *Science*, 2016, **353**, 158–162.
- 4 H. Fu, K. Nan, W. Bai, W. Huang, K. Bai, L. Lu, C. Zhou, Y. Liu, F. Liu, J. Wang, M. Han, Z. Yan, H. Luan, Y. Zhang, Y. Zhang, J. Zhao, X. Cheng, M. Li, J. W. Lee, Y. Liu, D. Fang,



- X. Li, Y. Huang, Y. Zhang and J. A. Rogers, *Nat. Mater.*, 2018, **17**, 268–276.
- 5 M. Zarek, M. Layani, I. Cooperstein, E. Sachyani, D. Cohn and S. Magdassi, *Adv. Mater.*, 2016, **28**, 4449–4454.
- 6 M. Ma, L. Guo, D. G. Anderson and R. Langer, *Science*, 2013, **339**, 186–189.
- 7 B. Gao, Q. Yang, X. Zhao, G. Jin, Y. Ma and F. Xu, *Trends Biotechnol.*, 2016, **34**, 746–756.
- 8 S. Fusco, M. S. Sakar, S. Kennedy, C. Peters, R. Bottani, F. Starsich, A. Mao, G. A. Sotiriou, S. Pané, S. E. Pratsinis, D. Mooney and B. J. Nelson, *Adv. Mater.*, 2014, **26**, 952–957.
- 9 A. Holstov, G. Farmer and B. Bridgens, *Sustainability*, 2017, **9**, 435.
- 10 S. Poppinga, C. Zollfrank, O. Prucker, J. Rühle, A. Menges, T. Cheng and T. Speck, *Adv. Mater.*, 2017, **30**, 1703653.
- 11 E. Sifert, E. Reyssat, J. Bico and B. Roman, *Nat. Mater.*, 2019, **18**, 24–28.
- 12 E. Hajiesmaili and D. R. Clarke, *Nat. Commun.*, 2019, **10**, 183.
- 13 J. H. Pikul, S. Li, R. T. Hanlon, I. Cohen and R. F. Shepherd, *Science*, 2017, **358**, 210–214.
- 14 H. Koemer, G. Price, N. A. Pearce, M. Alexander and R. A. Vaia, *Nat. Mater.*, 2004, **3**, 115–120.
- 15 T. P. Russell, *Science*, 2002, **297**, 964–967.
- 16 A. Nojoomi, K. Arslan, K. Lee and K. Yum, *Nat. Commun.*, 2018, **9**, 3705.
- 17 A. Cangialosi, C. Yoon, J. Liu, Q. Huang, J. Guo, T. D. Nguyen, D. H. Gracias and R. Schulman, *Science*, 2017, **357**, 1126–1130.
- 18 S.-J. Jeon, A. W. Hauser and R. C. Hayward, *Acc. Chem. Res.*, 2017, **50**, 161–169.
- 19 A. S. Gladman, E. A. Matsumoto, R. G. Nuzzo, L. Mahadevan and J. A. Lewis, *Nat. Mater.*, 2016, **15**, 413–418.
- 20 M. C. Koetting, J. T. Peters, S. D. Steichen and N. A. Peppas, *Mater. Sci. Eng., R*, 2015, **93**, 1–49.
- 21 I. Tokarev and S. Minko, *Soft Matter*, 2009, **5**, 511–514.
- 22 Y. Liu, B. Shaw, M. D. Dickey and J. Genzer, *Sci. Adv.*, 2017, **3**, e1602417.
- 23 L. H. Dudte, E. Vouga, T. Tachi and L. Mahadevan, *Nat. Mater.*, 2016, **15**, 583–588.
- 24 P. M. Reis, F. L. Jimenez and J. Marthelot, *Proc. Natl. Acad. Sci. U. S. A.*, 2015, **112**, 12234–12235.
- 25 S. Felton, M. Tolley, E. Demaine, D. Rus and R. Wood, *Science*, 2014, **345**, 644–646.
- 26 Z. Y. Wei, Z. V. Guo, L. Dudte, H. Y. Liang and L. Mahadevan, *Phys. Rev. Lett.*, 2013, **110**, 215501.
- 27 T. Castle, D. M. Sussman, M. Tanis and R. D. Kamien, *Sci. Adv.*, 2016, **2**, e1601258.
- 28 D. M. Sussman, Y. Cho, T. Castle, X. Gong, E. Jung, S. Yang and R. D. Kamien, *Proc. Natl. Acad. Sci. U. S. A.*, 2015, **112**, 7449–7453.
- 29 G. P. T. Choi, L. H. Dudte and L. Mahadevan, *Nat. Mater.*, 2019, **18**, 999–1004.
- 30 J. V. I. Timonen, M. Latikka, L. Leibler, R. H. A. Ras and O. Ikkala, *Science*, 2013, **341**, 253–257.
- 31 I. Torres-Díaz and C. Rinaldi, *Soft Matter*, 2014, **10**, 8584–8602.
- 32 W. Wang, J. V. I. Timonen, A. Carlson, D. M. Drotlef, C. T. Zhang, S. Kolle, A. Grinthal, T. S. Wong, B. Hatton, S. H. Kang, S. Kennedy, J. Chi, R. T. Blough, M. Sitti, L. Mahadevan and J. Aizenberg, *Nature*, 2018, **559**, 77–82.
- 33 M. D. Cowley and R. E. Rosensweig, *J. Fluid Mech.*, 1967, **30**, 671–688.
- 34 U. Banerjee and A. K. Sen, *Soft Matter*, 2018, **14**, 2915–2922.
- 35 L. Wang, F. Li, M. Kuang, M. Gao, J. Wang, Y. Huang, L. Jiang and Y. Song, *Small*, 2015, **11**, 1900–1904.
- 36 D. Vella, P. Aussillous and L. Mahadevan, *Europhys. Lett.*, 2004, **68**, 212–218.
- 37 P. Aussillous and D. Quéré, *Nature*, 2001, **411**, 924–927.
- 38 D. Vella, H.-Y. Kim, P. Aussillous and L. Mahadevan, *Phys. Rev. Lett.*, 2006, **96**, 178301.
- 39 M. Abkarian, S. Protire, J. M. Aristoff and H. A. Stone, *Nat. Commun.*, 2013, **4**, 1–8.
- 40 P. Aussillous and D. Quéré, *Proc. R. Soc. A*, 2006, **462**, 973–999.
- 41 Y. Xue, H. Wang, Y. Zhao, L. Dai, L. Feng, X. Wang and T. Lin, *Adv. Mater.*, 2010, **22**, 4814–4818.
- 42 T. B. Jones, J. D. Fowler, Y. S. Chang and C. J. Kim, *Langmuir*, 2003, **19**, 7646–7651.
- 43 K. D. Danov, P. A. Kralchevsky and M. P. Boneva, *Langmuir*, 2004, **20**, 6139–6151.
- 44 M. G. Nikolaidis, A. R. Bausch, M. F. Hsu, A. D. Dinsmore, M. P. Brenner, C. Gay and D. A. Weitz, *Nature*, 2002, **420**, 299–301.
- 45 D. A. Saville, *Annu. Rev. Fluid Mech.*, 1997, **29**, 27–64.
- 46 R. T. Collins, J. J. Jones, M. T. Harris and O. A. Basaran, *Nat. Phys.*, 2008, **4**, 149–154.
- 47 E. Schaeffer, T. Thurn-Albrecht, T. P. Russell and U. Steiner, *Nature*, 2000, **403**, 874–877.
- 48 M. M. Nicolson, *Proc. Cambridge Philos. Soc.*, 1949, **45**, 288–295.
- 49 E. Jambon-Puillet, C. Josserand and S. Protière, *Phys. Rev. Mater.*, 2017, **1**, 042601.
- 50 W. D. Greason, *J. Electrostat.*, 2003, **59**, 285–300.
- 51 J. B. Pendry, D. Schurig and D. R. Smith, *Science*, 2006, **312**, 1780–1782.
- 52 A. A. Fridman and L. A. Kennedy, *Plasma Physics and Engineering*, Taylor and Francis, New York, 2004.

