




Self-propulsion of a calcium alginate surfer†

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A droplet of sodium alginate dripped into calcium chloride solution results in plate or boat shaped hydrogels. Both exhibit several minute-long self-propelled motion on the liquid surface without any extra fuel added, offering a new method to making active materials. By changing the initial concentrations, we are able to tune the transient dynamic activities from translational to rotational or stop-and-run motion. Dynamics are governed by osmotic pressure induced Marangoni effect, depending on the density difference and initial concentrations.

1 Introduction

Most living organisms move by self-propulsion, which is an essential tool for their survival.¹ The locomotion is prerequisite for doing tasks and therefore scientists are highly interested in designing inanimate active matter, that can move itself in a pre-planned manner.^{2,3} Several techniques are available to generate autonomous activities by consuming chemical energy⁴ or exploiting physical phenomena (electromagnetic,⁵ electrical energy,⁶ light^{7,8}). Further mechanisms have been proposed as well, such as Marangoni effect,⁹ electroosmosis,⁶ bubble ejection¹⁰ and dielectrophoresis.¹¹ Different names also distinguish artificial active objects depending on their characteristics. In general, active matter is called either motor¹² or machine¹³ reflecting on that the object powers itself by converting some kind of energy into kinetic energy. Gear¹⁴ and droplet¹⁵ are also used to mirror the physical design. Moreover, a surfer⁷ propels along the interface, whilst a swimmer^{16,17} moves in the bulk phase.¹⁸

Marangoni effect is an interfacial phenomenon, where thermal or chemical gradients cause surface tension gradient, which induces mass transport. A well-known example of this is simply based on the dissolution of camphor in water.^{19–22} In literature, reactive self-propulsion considered as a chemical reaction induced motion, but an extra component (usually a surfactant

or an organic solvent) always implemented in the reaction to create the surface tension gradient.^{23–25} In addition to the camphor dissolution, only a recent article reports on tension-lowering polyelectrolyte reactants, which self-generate motion without an additive.²⁶ In a different way, the concept of exclusively one chemical reaction is responsible for the motion was also imagined earlier by van't Hoff, by generating osmotic pressure gradient as the concept of truly exploiting “chemomechanical” forces.²⁷ The idea to convert chemical energy into mechanical motion in a form of an osmotic motor is supported by computational results.²⁸ Our purpose is to demonstrate experimentally that a simple reaction without additives can induce self-propulsion by exploiting osmotic pressure gradient through gelation.

In designing synthetic motile objects, their physical and chemical constructions are very important, because they affect the dynamics differently, resulting in translational, rotational, spinning or even oscillatory motions. This could mean applying liquids in the form of oil droplets,^{24,29} solid materials, like in the case of strings of camphor disks³⁰ or gel-based structures like enzyme-programmed alginate objects.³¹ Gel-based systems^{25,32–34} are especially favorable in technology-development due to their soft matter properties. Besides that, various applications are being explored, such as developing bio-actuators,³⁵ drug delivery carriers,^{17,36} sensors for toxicity detection³⁷ and environmental remediation of chemical spills.³⁸ More control can be achieved with the help of studying interactions between the self-propelled objects.³⁹

In this work, we introduce a new type of active material exploiting osmosis. We use the simple reaction between calcium and alginate ions without any additives.⁴⁰ While the gelation takes place, different types of motions are observed. We demonstrate, how the initial concentrations influence the dynamics along with the hydrogel shape.

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2 Experimental

Fig. 1 illustrates the experimental system. Self-propelled hydrogels were made in a glass Petri dish (i.d. = 6 cm) filled with 15 mL, 2–6 M calcium chloride solution. To make the surfer, one droplet of 0.3–1.5% w/V sodium alginate (ALG) solution was dripped (0.38 mL min⁻¹) into the electrolyte from 5 mm above, aiming for the center of the Petri dish. The dripping was implemented by using a peristaltic pump (Ismatec Reglo) installed with a Tygon tube (i.d. = 0.76 mm) ending in a scalp vein needle (i.d. = 0.4 mm). In order to minimize geometrical effects, we kept all experimental conditions constant. Trajectories of the hydrogel surfers were recorded from above and sidewise by taking images with 5 fps using a digital camera (Unibrain Fire-i 630c) equipped with a 24 mm focal length lens (Vivitar). For shadowgraph imaging, a 12 mm extension tube (Vivitar) was inserted between the lens and the camera. The Petri dish was placed above an LED light source and 0.1 mL of 0.4% congo red dye was added to the alginate solution to enhance the visibility.

Analytical grade CaCl₂·2H₂O (99–105%) and technical grade sodium alginate reagents from VWR International were used. Chemicals were dissolved in deionized water (Purite RO100). Alginate solution was first stirred overnight to properly dissolve the polysaccharide, and then kept in a fridge for a maximum usage of one week. Experiments were repeated at least three times at 23 ± 2 °C. Images were analyzed using Fiji or Wolfram Mathematica, and edited in Inkscape. Viscosity, density, droplet weight, surface tension and nuclear magnetic resonance (NMR) measurements are added in Tables S1–S3 in the ESI.†

3 Results and discussion

Phenomenon

When the dripped alginate droplet comes into contact with the calcium chloride solution, crosslinking occurs immediately. During the gelation, the guluronate block of the alginate chains crosslinks with calcium ions diffusing inward from the surrounding solution. As soon as gelation begins on the outside,

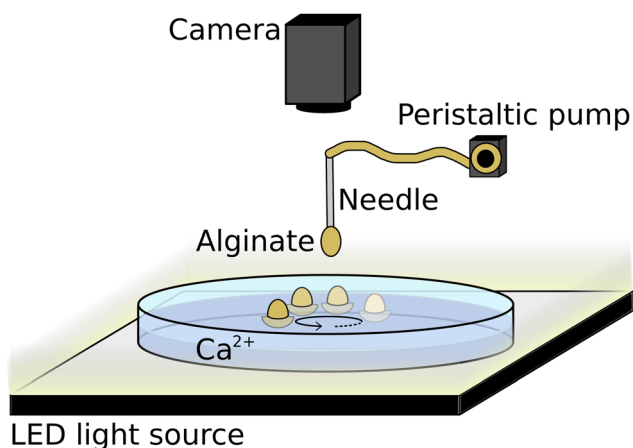


Fig. 1 Schematic diagram of the experimental setup.

the self-propulsion of the droplet starts because of the minute asymmetry in the shape and gel structure (Movie S1 in ESI†). Initially the floating droplet spreads out on the surface and takes up the shape of a plate. For smaller alginate concentrations (0.3 or 0.5% w/V), the droplet maintains this shape (see Fig. 2b). However, at greater alginate content (1 or 1.5% w/V), it transforms into a boat (see Fig. 2a). This transition is shown in Fig. 2c–h), where the entire droplet, both under and above the liquid surface, shrinks till it reaches the boat form. The bottom part of the droplet crosslinks immediately after contacting the CaCl₂ solution (Fig. 2c), while the upper part remains initially in a solution form and bulges higher (Fig. 2d–f). Later, it stops rising and shrinks as gelation also takes place inside (Fig. 2g and h). In contrast, plates (at lower alginate concentration) crosslink in a different way resulting in thin and flexible structures due to slower gelation. After contacting with the calcium chloride solution, the surfer only shrinks and becomes thinner. Along with the drastic shape transformations, particle image velocimetry (PIV) reveals that strong fluid flow evolves (see Fig. S3, S4 and Movies S2 and S3 in ESI†) in the calcium solution. These observations are inevitably the results of the arising osmotic pressure,⁴¹ whilst gelation also supports the shrinkage.⁴²

With shadowgraph imaging, we detect that the refractive-index changes in the immediate vicinity of the surfer when it

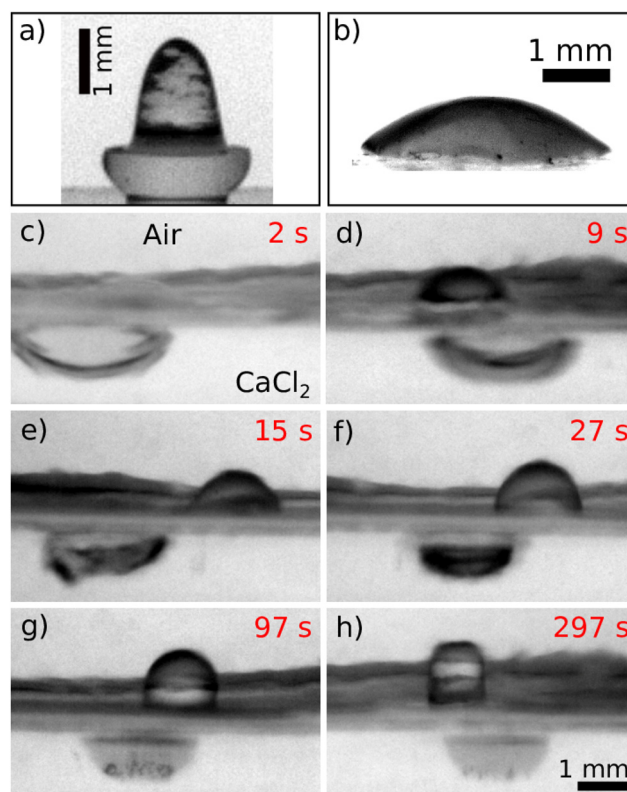


Fig. 2 A picture of the (a) boat and the (b) plate in a completely cross-linked state. (c)–(h) Side view of a 1.5% w/V ALG droplet in the 4 M CaCl₂ solution during self-propulsion. The curved wall of the Petri dish and the change in the refractive index of CaCl₂ solution and air cause the visual displacement between the upper part above the liquid and the lower submerged of the same boat.



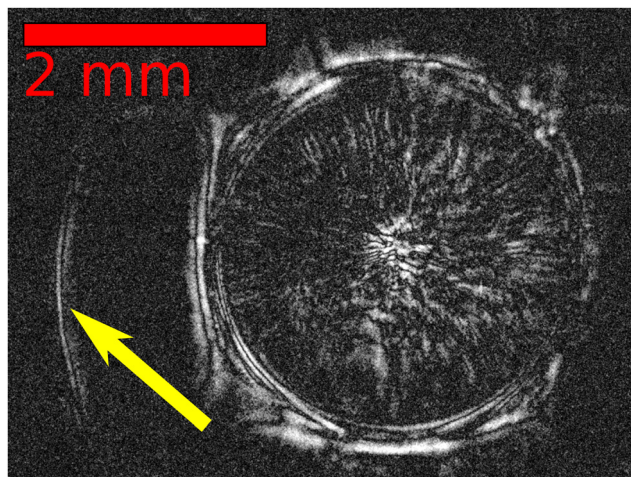


Fig. 3 Top view of a surfer (1.5% w/v ALG) after exerting water to the surrounding 4 M CaCl_2 solution using shadowgraph imaging. The yellow arrow points to the boundary of the diluted area.

moves (see Fig. 3). The only chemical that can be exerted out from the hydrogel, which is detected as it dilutes the surrounding calcium solution, is water.⁴³ The arising osmotic pressure exerts the water out which decreases the surface tension around (see Table S2 for the surface tension values, ESI[†]). The surfer appears symmetric, however, around its perimeter small fluctuations in the surface tension ($\delta\gamma$) exist. The magnitude can be estimated from the velocity on the surface (u) and the liquid viscosity (η) as $\delta\gamma \approx u\eta = 0.02 - 0.7 \text{ mN m}^{-1}$. This yields an upper limit on the Marangoni number as $Ma = \delta\gamma l / (\eta D) \approx 10^5$ where l is the liquid depth. Hence, motion is driven by surface tension gradient.

Our system does not include any additional “fuel” unlike in other published cases,^{24,25,30,34} only the gelation of two reactants is responsible for the self-propulsion as osmosis takes place.

Dynamics

The surfer position in the xy -plane of the Petri dish was gathered by an in-house imaging script using built-in macros in Fiji. First, the contour of the surfer was determined using Canny Edge Detector, then a circle with the corresponding radius was fitted and its center position (X , Y coordinates) was exported through Hough Circle Transform. Knowing the time delay (Δt) between two images, the instantaneous speed (v) of the object, defined as

$$v = \frac{\sqrt{(X_{t+\Delta t} - X_t)^2 + (Y_{t+\Delta t} - Y_t)^2}}{\Delta t}, \quad (1)$$

was determined. An example of a speed profile is illustrated in Fig. 4a and its related trajectory in Fig. 4b. This experiment shows all of the four separable periods (I–IV) which were observed during a surfer’s movement. However, only three phases can be distinguished based on the velocity profile: period I and period II are both part of the continuous phase followed by the intermittent phase (period III) before reaching the quiescent state (period IV). We also consider this as the dynamic transition from harmonic to relaxation-like motion.⁴⁴

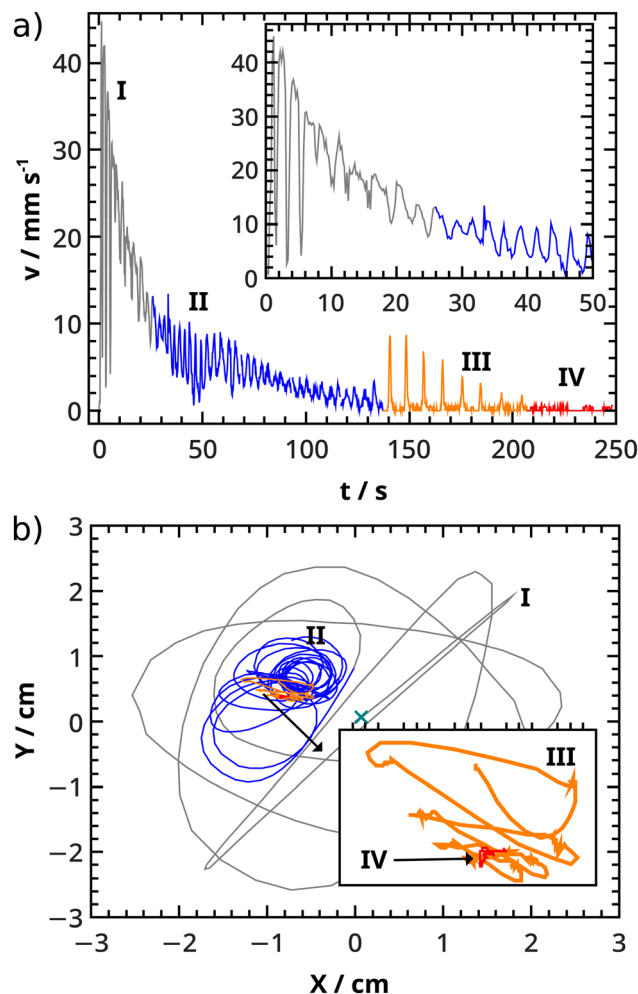


Fig. 4 (a) Change in speed (v) of a self-propelling surfer (6 M CaCl_2 , 0.5% w/v ALG) as a function of time (t). Four different forms of phases are distinguished: translational (I), rotational (II), stop-and-run (III) and quasi-equilibrium (IV). (b) Trajectory of the same surfer demonstrating its four pathways of the different movements.

Period I is characterized by translational movement, where the maximum speed of the surfer is reached in terms of its total activity time. In this section, the surfer begins its movement in a random direction heading to the Petri dish boundary from the center. The surfer and the dish wall deform the solution surface in the opposite direction: there is depression at the former and elevation at the latter. Hence, the capillary force represents repulsion between them,⁴⁵ which slows down the surfer (local minima in the gray regime in Fig. 4a and see Fig. S5 in ESI[†]) and reverses its direction. This motion is characterized by paths with a constant angle $\alpha = \text{atan2}(Y_{t+\Delta t} - Y_t, X_{t+\Delta t} - X_t)$ with π or $-\pi$ jumps representing a sharp reverse (see Fig. S6(A), ESI[†]). The turns are sharper in the earlier stages of the motion, but as the reaction proceeds, the velocity decreases and the shape transformation directly affects the repulsion, which leads to turn the surfer away less sharply. Between two turns, the surfer always returns to follow the rectilinear movement as much as possible. When the translational motion transforms



into smaller and continuous circles, the trajectory of period II shows rotational motions. This stage is characterized by constant $d\alpha/dt$, indicating a constant angular velocity (see Fig. S6(B), ESI†). Although period I and II are not distinguishable from each other in the speed profile, because they show the same harmonic decay, the following relaxation-like motion in period III is very distinct, as sharp velocity peaks of sudden movements repeatedly appear between rest states and thus this section referred as stop-and-run motion. The random values of α associated with the sudden movements indicates the stochastic nature of the behavior (see Fig. S6(C), ESI†). At this stage, the gelation took place already in the surfer to such an extent that it controls the release of water. Motion is possible only after sufficient amount of water is exerted, until which the surfer remains stationary. In period IV, the surfer is no longer active, thus no characteristic movement can be allocated either in the speed profile or in the trajectory.

With different initial concentrations, we created different dynamic behavior (see Fig. 5). We see trajectories rather moving linearly at 1–1.5% w/V alginate concentrations (Fig. 5a and b and Fig. S7 in ESI†). Nonetheless, parallel experiments show that trajectories stochastically located in the Petri dish in all of the different concentrations (see Fig. S7 in ESI†). Increasing the

CaCl₂ concentration creates higher surface tension gradient (see Table S2 in ESI†), hence gives more energy to the surfer (see Fig. 5a and b). However the linear motion of period I does not change due to the similar repulsive force between the wall and the surfer caused by the similar boat shape. We also see in Fig. 5b that turning points move further away from the wall over time, because the surface tension gradient (thus repulsion) changes along the crosslinking. At 0.5% w/V alginate concentration, repulsion decreases between the plate and the wall. The surfer manages to approach the wall for a longer time, yet soon diverts its direction and therefore the traveled route in period I resembles a star-shape (Fig. 5c and d). The repulsion between the wall and the surfer changes and proceeds similarly to the above mentioned alginate composition in a single experiment. At 0.3% w/V alginate, the surfer can touch the wall due to the small repulsion, so it can even follow the Petri dish boundaries (Fig. 5e). If we use high CaCl₂ concentration at this alginate amount, more repulsion exerted yields the turnings of the surfer (Fig. 5f).

The surfer accelerates the most in period I. It reaches its maximum velocity (v_{\max}) in this phase, where different maximum velocities are measured depending on the initial concentrations (see Fig. 6a). By increasing the alginate concentration, the maximum velocity decreases. The alginate concentration regulates the amount of exerted water, which sets the extent of the Marangoni effect (and thus the velocities). We observe this (Fig. 2c–h) when the surfer is distorted upwards into a boat at higher alginate concentrations, while it retains water unlike plates, which only releases water. It is also important to note, that plates (lower alginate concentrations) are more likely to approach the wall boundaries closer because of the smaller repulsion, which slows down the plates less.

We have found that the highest maximum velocities are reached in 4 M CaCl₂ solutions. To understand this, we discuss the increase and decrease in maximum velocity separately,

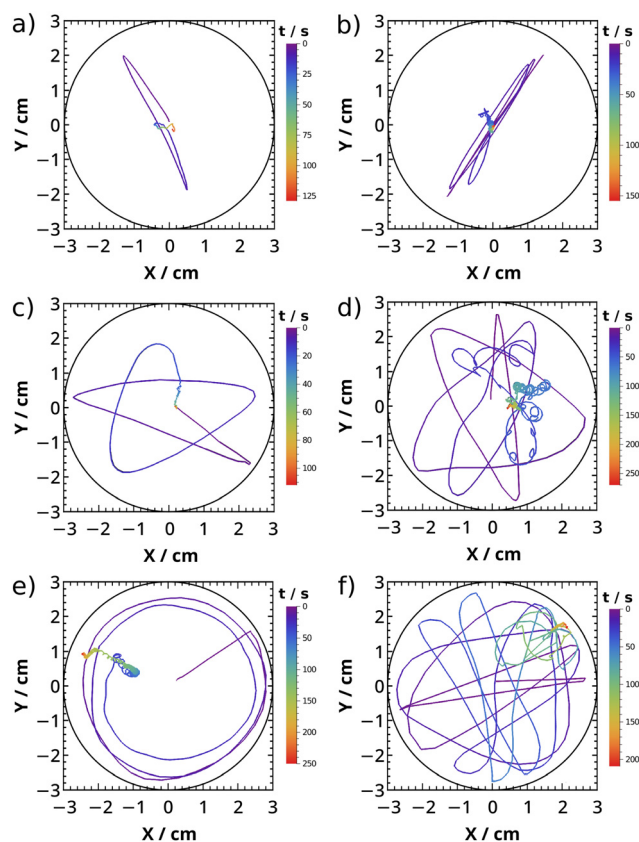


Fig. 5 Trajectory of a calcium alginate surfer supplemented with its temporal evolution (colourbar) at (a) 2 M CaCl₂, 1.5% w/V ALG, (b) 4 M CaCl₂, 1.5% w/V ALG, (c) 2 M CaCl₂, 0.5% w/V ALG, (d) 4 M CaCl₂, 0.5% w/V ALG, (e) 2 M CaCl₂, 0.3% w/V ALG and (f) 6 M CaCl₂, 0.3% w/V ALG. The black circle represents the Petri dish boundaries.

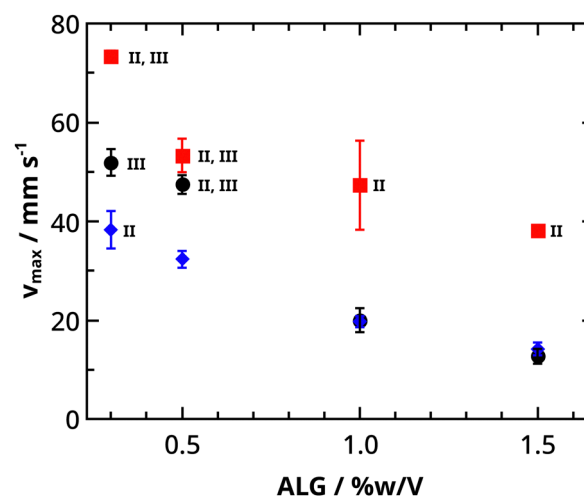


Fig. 6 Maximum velocity (v_{\max}) values in period I of the self-propelling calcium alginate surfers. The existence of characteristic rotational (II) and stop-and-run (III) motions is also marked. (♦ 2 M CaCl₂, ■ 4 M CaCl₂, ● 6 M CaCl₂).



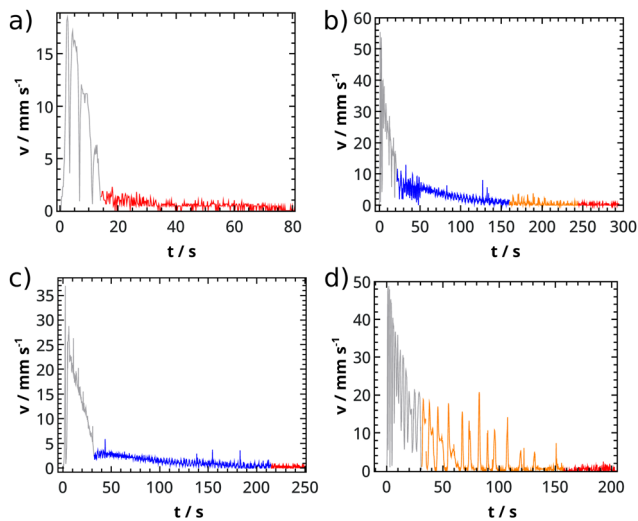


Fig. 7 Different velocity profiles of (a) 2 M CaCl_2 + 1% w/V ALG, (b) 4 M CaCl_2 + 0.5% w/V ALG, (c) 2 M CaCl_2 + 0.3% w/V ALG and (d) 6 M CaCl_2 + 0.3% w/V ALG.

along with the CaCl_2 concentration. At 2 M CaCl_2 , the density difference significantly decreases, which influences the surfer's depth of immersion. This affects the speed, which was already published about camphor boats: deeper immersion decreases velocity.⁴⁶ We have also observed, that fully immersed surfers do not move. At 6 M CaCl_2 , we suggest that the gelation is faster due to the high concentration, which restricts the exertion of water. For example, at 6 M CaCl_2 and 0.3% w/V alginate, dominantly stop-and-run motion is observed (see Fig. 7d), which indicates water regulation in a short period of time due to the high-rate gelation.

Occasionally the kinetic energy of period I is transformed into period II and/or III motions as indicated in Fig. 6. This transformation can occur by changing both CaCl_2 and alginate concentration, but randomly based on the durations of period I (see Fig. S8 in ESI[†]). From Fig. 6 we can also see that there are concentrations where only period I exists besides period IV (an example in Fig. 7a). When surfers have more energy, which is indicated by a higher maximum velocity value, they are able to make rotational motions (II). With even more energy, they are also able to make stop-and-run motion (III) in addition to period II (see Fig. 7b). Increasing the CaCl_2 concentration can make a complete transformation from period II to III (illustrated in Fig. 7c and d) through a transition state (see Fig. S9 in ESI[†]), where both types exist while the frequency of period II is altered with the changing spatial extent of the rotational motion (larger or smaller circles). If the alginate concentration is reduced, period III is more strongly present than period II (see Fig. 4a and 7d). By changing the initial concentrations, we are able to dissipate chemical energy in the four characteristic periods and observe different dynamical behavior.

4 Conclusions

We have created an active material exploiting osmosis in the gelling reaction of alginate and calcium ions. The appearing

osmotic pressure during the crosslinking exerts water from the surfer, which changes the surface tension around the surfer and thus forcing it to move due to Marangoni effect. We have not used any extra chemical to provide “fuel” for the motion, which offers a new method for transient self-propulsion based on hydrogels.

Two types of surfer shapes have been observed after complete crosslinking, where plates considered more active than boats. We have observed the complete dynamical transition from harmonic to relaxation-like motion with the more active surfers. Consecutively four periods (translational, rotational, stop-and-run, quasi-equilibrium) have been distinguished and different trajectories have been observed according to the initial concentrations. The occurrence of these motions (from linear to curvilinear to rotational) is also influenced by the shape of the surfer.

We have found that maximum velocities increase with decreasing alginate concentration, which depends on the water retention ability set by the alginate concentration. It also increases with the CaCl_2 concentration to a certain extent, because the density difference increases, which decreases the depth of immersion. Further increase in the CaCl_2 concentration results in smaller maximum velocities because of the greater gelation rate. We have also explained that the duration of the observed periods can be tuned by the initial concentrations. The study presented here advances energy-based motors and suggests a new way on how to design soft self-propelled surfers based on osmosis.

Data availability

Original data are available from the authors upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

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