Light-switchable deposits from evaporating drops containing motile microalgae

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Light-switchable deposits from evaporating drops containing motile microalgae

In this study, we utilise the light-sensitivity of green microalgae to show that their tunable activity can force, completely suppress, and even direct the spatial structure of the coffee ring effect in an evaporating drop.

Image credit: Laura Quarto

Light-switchable deposits from evaporating drops containing motile microalgae†

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Deposits from evaporating drops have been shown to take a variety of shapes, depending on the physicochemical properties of both solute and solvent. Classically, the evaporation of drops of colloidal suspensions leads to the so-called coffee ring effect, caused by radially outward flows. Here we investigate deposits from evaporating drops containing living motile microalgae (Chlamydomonas reinhardtii), which are capable of resisting these flows. We show that utilizing their light-sensitivity allows control of the final pattern: adjusting the wavelength and incident angle of the light source enables forcing the formation, completely suppressing and even directing the spatial structure of algal coffee rings.

1 Introduction

Evaporation of a drop pinned to a surface induces capillary flows towards its edge, a phenomenon leading to ring-like deposits at the contact line.1 Ways to reverse this so-called coffee ring effect are much sought after, given its undesirability in various technical applications. Many strategies have been developed to counteract the formation of deposits on drop peripheries,2 such as the introduction of surface tension-driven Marangoni flows,3,4 capillary forces,5,6 prevention of contact line pinning by using hydrophobic7 or absorbing substrates,8,9 altering substrate elasticity,10 and even more complex strategies including the drying of frozen drops.11 Most commonly, the evaporation-induced stains are composed of either colloids or macromolecular solutes, in which the transport to the periphery is controlled by advection. Polymer solutions showing low Pe´clet numbers Pe = Re/D, for which diffusive transport, characterized by the diffusion coefficient D, dominates over advective flow, defined as the flow velocity v along the drop radius R, have shown to mitigate coffee rings in numerical and experimental studies.12,13 Other systems, which are presumably capable of overcoming advection controlled transport belong to the field of active matter.14,15 Active particles, such as biological cells, extract energy from their environment to fuel self-propulsion.16 Drying suspensions of drops of living motile bacteria have shown to lead to disk-like deposits,17 to alter coffee rings in colloidal suspensions,18 to create homogeneous patterns due to the auto-production of biosurfactants19 and also to induce complex wetting behaviors in micropillar arrays.20,21 While exhibiting a variety of possibilities, the capacity of these active entities to counter act (complex) internal flows in evaporating drops is limited. Due to their weak swimming power, most bacteria cells are subject to internal flows. Another class of active “particles” that potentially surpass this limitation are algae. Chlamydomonas reinhardtii (C. R.) emerged as a model system for the investigation of green microalgae,22 given their well understood interaction with light. Using the synchronous beating of their two flagella, C. R. steer either towards or away from a light source in a process referred to as phototaxis,23–26 and consequently develop high swimming velocities and large thrust.27–29 This model organism is even capable of light-induced adhesion to surfaces; it shows strong adhesion to substrates when exposed to blue light,30 independently of the chemical composition of the substrate.31 Considering these key characteristics, drops containing C. R. cells offer a promising experimental platform for studying the effect of activity on drying-induced pattern formation.

Using phase contrast microscopy coupled with particle tracking techniques, we show here that by manipulating the light conditions under which evaporation takes place, it is possible to control the final deposits left behind active drops containing C. R. cells. In contrast to bacteria, we find motile algae to counteract advection towards the contact line and undergo mixing within the drops. When dried under white light, C. R. show an increased adhesion on the substrate and an overall lower motility, which leads to a homogeneous final deposit. Using red light instead, increases the motility of the algae and causes them to aggregate at the contact line during

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the final stages of evaporation, which leads to the formation of a ring-like deposit. This is surprising, given the fact that the algae have sufficient thrust to easily withstand the advective flow that normally causes coffee rings. In fact we will see that, for the algae, the mechanism for the ring-deposition formation is completely different and can be explained in terms of confinement at the contact line related to a critical contact angle. We finally demonstrate that, by adjusting the incident angle of the light source, we can even maneuver the active particles in a desired direction and obtain a crescent-shaped deposit.

2 Results and discussion

To study the influence of motility and wavelength of the incident light on the deposits of drops containing either motile or non-motile microalgae, 0.3 μl drops of Chlamydomonas reinhardii (C. R.) suspended in commercial BG-11 medium (from Sigma) were deposited on glass microscope slides and left to dry at 20 °C and 50% relative humidity. C. R. are about 10 μm in diameter and we used a concentration of 2.57 × 10^3 ± 0.96 × 10^4 cells per μl. The aqueous drops exhibited an initial contact angle of θ = 40° and the contact line remained pinned during drying (see Materials and methods and Fig. S1, ESI†). We monitored the evaporation process using an inverted phase contrast microscope whilst illuminating the drops from above (see Materials and methods and Fig. S2, ESI†). The resulting patterns are summarized in Fig. 1. We begin by examining the final patterns left behind a sample containing non-motile algae cells, obtained by heating up the suspension to ≈50° prior to drop deposition. This allows a reference measurement containing inactive particles, as one would get with a regular non-Brownian dispersion of spherical particles. A dried drop containing these “passive” particles left behind a deposit resembling a coffee ring (Fig. 1a), which we quantified by calculating a rectangular intensity profile (Fig. S3, for the concentration dependence of this deposit see Fig. S4, ESI†). This is similar to the deposits from drying liquid drops containing inactive colloids, where capillary outward flows carry the solute towards the contact line.1

For motile algae, illumination with regular white light during the evaporation period produces a deposit where dried algae cells are homogeneously distributed (Fig. 1b and Fig. S3, ESI†). By implementing a longpass filter, excluding wavelengths below 650 nm (i.e. illumination with red light), we observed a ring-shaped pattern instead (Fig. 1c and Fig. S3, ESI†). From these observations we surmise that there are two key components to be considered relevant for the final pattern left behind a drying drop containing microalgae: the effect of the algal motility and the influence of the color of the light on the algal motility inside a drying drop.

To investigate these effects in detail, we used Particle Tracking Velocimetry (PTV), using the algae as particles, to dynamically determine flow fields in the drying drops containing either non-motile or motile cells. During evaporation, non-motile algae were being advected towards the contact line in a radially outward flow, as shown in Fig. 2a. In the latter stages of evaporation (>0.8t0), we observed an inward motion displacing the non-motile algae towards the drop center. This was found to be a consequence of capillary forces induced by the confinement of the particles at the drop edge.3,6,34 When the algae are motile, the picture provided by PTV is different. If illuminated with white light from above, the algae form multiple vortices,35–37 and “mix” within the drop, as shown in Fig. 2b. This effect is even more pronounced when illuminated with red light from above (Fig. 2c), suggesting an increased activity for this case. For both active cases we found the same inward motion in the later stages of evaporation as for the non-motile case, indicating independence of this effect on the motility. The PTV results show that the active algae resist the outward flows observed in the non-motile case and that their activity inside the drops increases when dried under red instead of white light.

To quantify C. R. activity under different light conditions, we tracked the algae directly at the glass surface. A snapshot of motile algae illuminated with white light from above is presented in Fig. 3a, along with a few selected trajectories. Here the algae move close to the surface, contrary to C. R. under red light (Fig. 3b), where most algae swim outside the viewing plane. The increased cell-surface interaction when illuminated with white light has been shown to originate from a light-switchable adhesiveness mediated by contact between the flagella of the algae and the substrate, which is regulated by a blue-light receptor.30 From the trajectories of the algae we calculated their mean square displacements MSD(τ) = ⟨|r(t) − r(t0)|^2⟩, from which we extracted their diffusion coefficients D as MSD(τ) = 4Dt for delay times τ > τc, where the critical delay time τc separates an initial ballistic regime from a diffusive-like behavior at longer timescales.38

The increased cell-adhesiveness when illuminated with white light reflects in the MSDs of the algae. The onset of the ballistic regime at τc is shifted to shorter delay times when illuminated with white light (τc ≈ 2 s, Fig. 3c) instead of red light (τc ≈ 8 s, Fig. 3d), and the diffusion coefficient found in the white light case, D ≈ 4.6 × 10^3 μm^2 s^{-1}, is significantly lower than the one found in the red light case, D ≈ 6.2 × 10^4 μm^2 s^{-1}. To evaluate the competition between the diffusivity of the algae and the rate
of advection due to the evaporative flux, we calculated the dimensionless Péclet number \( \text{Pe} = R \langle v(t) \rangle / D \), where \( R \) is the drop radius and \( \langle v(t) \rangle \) the averaged time-dependent velocity due to the evaporative flux, extracted from the flow fields governing the non-motile case in Fig. 2a. For the non-motile algae, we calculated \( D \) using the Stokes–Einstein equation \( D = 2k_b T/\eta d \) with the viscosity of the solvent \( \eta = 10^{-3} \) Pa s and the diameter of the algae \( d = 10 \) \( \mu \)m. The Péclet numbers as a function of time (Fig. 4a) reveal that, regardless of the light condition, the diffusivity of the motile algae dominates advection (\( \text{Pe} < 1 \)) and the red light case exhibits Péclet numbers an order of magnitude below the white light case, indicating a higher activity of the algae. The non-motile algae however show advection dominated evaporation (\( \text{Pe} > 1 \)), which is typical for evaporating drops of non-Brownian suspensions. From the PTV measurements and Péclet numbers, we inferred that active algae overcome advection and resist being dragged towards the contact line by the radially outward flux. We make this idea quantitative from the following considerations: first, the velocities of the active algae remain unchanged during evaporation, regardless of whether they face inwards or outwards with respect to the drop center (Fig. 4b). Secondly, we compare the drag force \( F_d \) exerted on the algae cells due to the capillary outwards flow to the dynamic propulsion force \( F_p \), arising from the beating of the flagella of the algae, which has been measured recently in vivo for motile \( C. \ R. \) and found to be \( 26 \pm 5 \) pN. Using Stokes’ law we find \( F_d = 3 \pi \eta d \langle v \rangle = 0.25 \pm 0.13 \) pN with \( \langle v \rangle \) being the maximum averaged velocity of the outward flow measured in the non-motile case (Fig. 4b). The ratio \( F_p/F_d \approx 10^2 \) indicates that the swimming force of individual \( C. \ R. \) cells is orders of magnitude higher than the force the algae experience due to drag in the evaporating drops.

The question remains why drops containing \( C. \ R. \) leave a ring-like deposit when dried under red light. Density maps obtained from the PTV measurements show that, under red light, active algae accumulate at the contact line in the later stages of evaporation (Fig. S5, ESI†). If illuminated with white light from above, the increased cell-substrate adhesiveness, and consequently low diffusivity, prohibits the algae to reach the contact line and the resulting deposit is thus homogeneous.
containing microalgae on a more wetting substrate (Fig. S6, ESI† we verified in a separate experiment involving deposition of drops line if the drop reaches below a critical contact angle,5,6,34,40 which consequence of algae getting confined and trapped at the contact

introduced in Fig. 1(c). We sugge st that these clusters form as a time (Fig. 5), at which they eventually form the ring-like pattern at

N

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within a cluster

f onwards,

of coffee ring manipulation, as neither the substrate nor the

meters of the light source, which is advantageous in the context

matter system can be influenced simply by adjusting para-

3 Conclusion

To conclude, the final patterns produced by drying drops containing Chlamydomonas reinhardtii (C. R.) microalgae vary significantly with their motility, which is tunable by light. If the algae are non-motile, the deposit resembles a coffee ring caused by radially outward capillary flows. Active algae, however, resist this drying induced flow and the deposits can be controlled by adjusting the color of the light source or its angle of incidence under which evaporation takes place. We surmise the variability in the shape of the deposits to originate from different swimming behaviors, which depend on the light condition. This shows that the coffee ring effect in this active matter system can be influenced simply by adjusting parameters of the light source, which is advantageous in the context of coffee ring manipulation, as neither the substrate nor the
sample composition have to be altered to achieve variations in the deposits. The findings presented here can be beneficial for a variety of (bio-)technological applications, which are especially important given the recent rise in algal technologies and applications.41

4 Materials and methods

4.1 Cell cultivation

*Chlamydomonas reinhardtii*, strain SAG 77.81, were grown axenically in BG-11 medium at 20 °C at a light intensity of \( \approx 18 \mu \text{mol (m}^{-2} \text{s}^{-1}) \). Samples from the culture were taken during its logarithmic growing stage and mixed with BG-11 medium to achieve a dilute suspension.

4.2 Substrate and sample preparation

As a substrate we used a partial-wetting glass slide obtained with a surface silanization technique. Prior to surface silanization, the glass slide was cleaned in a Zepto Electronic Diener plasma cleaner for 60 s. Then the glass slide was immersed in a mixture of 1 mL trichloro (octyl)-silane in 99 mL toluene. This procedure introduces a moderate degree of hydrophobicity, which we found to prevent algae cells from sticking to the substrate. The glass slide was cleaned with ethanol and deionized water prior use.

4.3 Experimental settings

The drying process of the microalgae drops was monitored using an inverted phase contrast microscope at 5x and 10x magnification. A 650 nm longpass filter was added for the experiments involving red light illumination. For the side light measurements, we positioned a collimated white light source at an incident angle of \( \approx 45^\circ \).

4.4 Tracking

For the tracking employed in this work, we used the python-based package trackpy.42,43 In the experiments involving the analysis of the whole drops we tracked particles within consecutive frames at 1 frames per second. The position vector is imposed a threshold to filter out non-moving particles. The position vector is imposed a threshold to filter out non-moving particles. The position vector is imposed a threshold to filter out non-moving particles. For the in-depth analysis of the cell-surface interaction we used an inverted phase contrast microscope at 5x magnification. A 650 nm longpass filter was added for the experiments involving red light illumination. For the side light measurements, we positioned a collimated white light source at an incident angle of \( \approx 45^\circ \).

\[
\omega = \frac{\partial u_y}{\partial x} - \frac{\partial u_x}{\partial y} \tag{1}
\]

For the in-depth analysis of the cell-surface interaction we extended the tracking time to 10 seconds recording at 10 frames per second. Within the tracking procedure we always imposed a threshold to filter out non-moving particles. We extracted the mean square displacement (MSD):

\[
\text{MSD}(t) = \langle |r(t + \tau) - r(t)|^2 \rangle \tag{2}
\]

where \( \langle \cdots \rangle \) indicates the time average.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


