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Discrete Salt Crystallization at the Surface of a Porous Medium

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Efflorescence refers to crystallized salt structures that form at the surface of a porous medium. The challenge is to understand why these structures do not form everywhere at the surface of the porous medium but at some specific locations and why there exists an exclusion distance around an efflorescence where no new efflorescence forms. These are explained from a visualization experiment, pore-network simulations and a simple efflorescence growth model.

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Salt crystallization is an important issue in building physics because of the severe damages caused by the crystallization process [1]. Salts can enter stone and masonry by several routes, the simplest of which is the capillary rise of ground water. As the water wicks up into a wall, it also evaporates. When the evaporation front is within the porous medium, salt crystals will precipitate inside the porous medium if the solution becomes supersaturated; this is called subflorescence. When the liquidvapor interface remains at the porous medium surface, supersaturation leads to precipitation at the surface or efflorescence. Although less damaging than subflorescence, efflorescence is an important issue for the conservation of old paintings and frescoes and can be also a problem in soil physics [2] or in relation with the underground sequestration of CO_2 [3].

In order to develop a better understanding of efflorescence formation, we performed the simple wicking experiment sketched and described in Fig. 1 [4]. Results are shown in Fig. 2. As can be seen, the localization of the efflorescence white spots varies with δ (δ is the distance between the porous medium surface and the hollow cylinder entrance in Fig. 1). The other striking phenomenon is the discrete nature of efflorescence. Efflorescence does not cover the entire porous medium surface but forms individual and isolated growing structures. Hence an important fraction of the surface remains free of efflorescence.

We now provide explanations to the observations, discussing first what happens before the occurrence of the first crystal at the porous medium surface.

It is essential to note that the evaporation process induces a flow within the porous medium. Since the menisci stay at the sample surface (the medium remains saturated), the evaporation rate J_m from a meniscus is balanced by a liquid flow rate q_m (due to capillarity) toward the meniscus,

$$q_m = J_m / \rho_\ell, \tag{1}$$

where ρ_{ℓ} is the liquid density.

As a result, there exists a velocity field **u** within the pore space with an average velocity \bar{U} directed toward the surface given by $\bar{U} = J/A/\rho_{\ell}/\varepsilon$ where $\varepsilon (\approx 0.36)$ is the porous medium porosity (see caption in Fig. 3 for the definition of J and A). Owing to the existence of the velocity field, the ions are transported by advection to the evaporation surface. As a consequence a salt peak builds up and salt concentration gradients develop. But as soon as a concentration gradient develops diffusion tends to level off the accumulation. Hence there is a competition between advection, which transports ions to the top of the sample and thereby causes accumulation, and diffusion. Using the initial concentration C_0 , the wick height L and \bar{U}

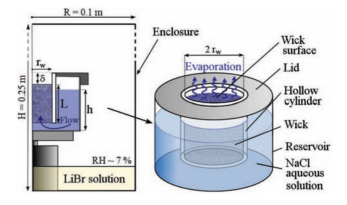


FIG. 1 (color online). Sketch of wicking-evaporation experiment. A pack of 1 mm glass beads ($d_b \approx 1$ mm) in a 50 mm long hollow cylinder of radius $r_w = 19$ mm forms a porous medium. The packing of height *L* is in contact at its bottom with a NaCl near saturated aqueous solution ($C = C_0 = 25$ g NaCl/100 g solution; the saturation concentration is $C_{\text{sat}} = 26.4$ g NaCl/100 g solution). The liquid level *h* in the reservoir is at any time such that the wick remains fully saturated owing to capillary effects. The system is set in a cylindrical enclosure of controlled temperature ($T \approx 22$ °C) and relative humidity (thanks to a LiBr saline solution, RH $\approx 7\%$). The evaporation rate at the wick surface and the hollow cylinder entrance.

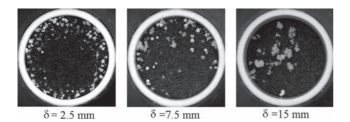


FIG. 2. Efflorescence discrete structures (top view of wick surface). The white spots are efflorescence. The projected vertical surface area of the white spots onto the wick surface is about the same in the three images.

as characteristic concentration, length and velocity, respectively, the dimensionless equation governing the dissolved salt transport within the pore space reads [5],

$$\operatorname{Pe}\left(\frac{\partial C'}{\partial t'} + \frac{\mathbf{u}}{\bar{U}} \cdot \nabla C'\right) = \Delta C', \qquad (2)$$

where $C' = C/C_0$ and $t' = t/t_{ref}$ with $t_{ref} = L/\overline{U}$; Δ is the Laplacian operator and Pe is the Peclet number characterizing the competition between advection and diffusion

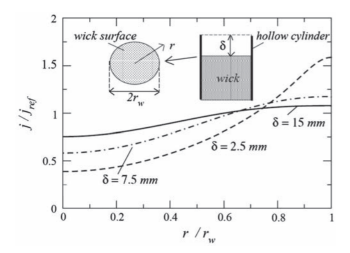


FIG. 3. Distribution of evaporation flux j at the porous medium surface. The reference flux j_{ref} is equal to J/A where J is the evaporation rate and A is the porous medium surface area: $J = 2.7 \times 10^{-8} \text{ kg/s}, 2.37 \times 10^{-8} \text{ kg/s}$ and $2.02 \times 10^{-8} \text{ kg/s}$ for $\delta = 2.5, 7.5$ and 15 mm, respectively. The evaporation rate is measured from weighing the set-up and also computed solving numerically the vapor transport equations in the enclosure with a finite element code [4,11]. A good agreement is found with the measurements provided that free convection effects are included in the computations. The flux distributions are deduced from the numerical computations. The crucial point is that the evaporation flux distribution at the surface varies with δ . When δ is small enough ($\delta = 2.5$ mm), the evaporation flux is significantly higher at the periphery (somewhat similarly as in the classical coffee ring problem [5]) whereas for a sufficiently large δ (δ = 15 mm) the evaporation flux is much more uniform $(i(r, \delta) \approx$ $J(\delta)/A$) over the surface. One can refer to [11] for more details.

effects; $Pe = \frac{\bar{U}L}{D_s}$ ($D_s = 1.3 \times 10^{-9} \text{ m}^2/\text{s}$ is the diffusion coefficient for the dissolved salt). For our experiment, this gives $Pe \approx 2$, 1.6 and 1.4 for $\delta = 2.5$, 7.5 and 15 mm, respectively.

As a result of the significant advection, the ion distribution is characterized by a narrow region of high salt concentration adjacent to the porous medium surface [6], whose size increases with time but remains narrow (on the order of 2–3 bead diameters). This explains why crystallization is observed at the surface and not inside the porous medium. Crystallization occurs when the saturation concentration C_{sat} is reached (supersaturation effects are negligible for NaCl [7]), and C_{sat} is first reached at the sample surface owing to the advection effect.

Efflorescence preferentially appears in regions where evaporation is the greatest (thus in the surface periphery when $\delta = 2.5$ mm) since the greater the evaporation flux, the greater the underlying velocities in this region as expressed by Eq. (1). When $\delta = 15$ mm the evaporation flux is much more uniform at the surface (see Fig. 3), and there is no peripheral preferential location of efflorescence spots anymore.

To explain now the discrete nature of efflorescence formation, it is essential to notice that the disordered nature of the random packing induces spatial fluctuations in the velocity field **u** within the pore space. The effect of this spatially random velocity field is illustrated solving numerically the transient transport problem represented by Eq. (2) using a 3D pore-network approach [8]. The porous medium is represented as a simple cubic network (of regular spacing d_b) of pores interconnected by tubes (referred to as bonds). The pore volumes and the bond radii are randomly distributed in the ranges $[0.3-0.426]d_b^3$ and $[0.09-0.21]d_b$ respectively, using Gaussian probability density functions, in accordance with known data for random packings of monodisperse spherical beads [9].

The velocity and concentration fields in the network (initial concentration C_0) are computed solving problems equivalent to resistance network problems [4]. An uniform pressure is imposed at the interface between the network and the bottom reservoir. The imposed flow rate in each surface bond (bond in contact with the top surface of network) is given by Eq. (1) and deduced from the data shown in Fig. 3 ($J_m(r, \delta) = j(r, \delta)d_b^2$). The concentration C_0 is imposed at the bottom of network. A salt zero flux condition is imposed at the surface menisci (the salt cannot leave the liquid phase before the crystallization onset). As a result of network disorder, the average velocity in the bonds varies randomly. This randomness induces in turn spatial fluctuations in the salt concentration within the network and therefore at the network surface.

The numerical simulations reveal that the structure of the salt concentration field at the pore-network surface remains unchanged (except at the very beginning of this transient process). Hence the location of each local

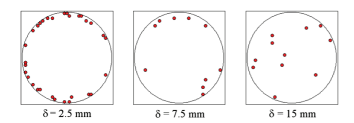


FIG. 4 (color online). Pore-network simulation of salt concentration maxima distribution at porous medium surface. The red dots correspond to the menisci where the reduced salt concentration C^* is greater than 0.8.

concentration extremum at the surface is always the same for a given realization of network and a given evaporation flux distribution. In others terms, the spatial distribution of the reduced concentration $C^*(t) = \frac{C(t) - C_{\min}(t)}{C_{\max}(t) - C_{\min}(t)}$ at network surface, where C_{\max} and C_{\min} are the surface maximum and minimum concentration at time *t*, is independent of time *t*. The simulation results are presented in Fig. 4, which shows an obvious correlation between the locations of concentration maxima (which correspond to the places where crystallization must start) at wick surface and the distribution of crystallization spots in Fig. 2. Thus the key role played by advection on the salt distribution in the pore space combined with the spatial fluctuations of the velocity field

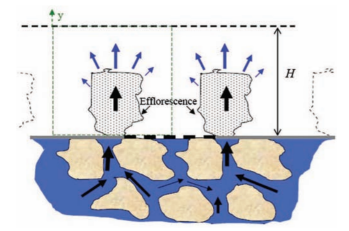


FIG. 5 (color online). Efflorescence is porous and pumps by capillarity the liquid solution traveling in the underlying porous medium [10]. Efflorescence grows owing to the salt deposition occurring preferentially in efflorescence upper regions where evaporation is higher. As a result of efflorescence growth, evaporation from surface pores located between two efflorescence structures is screened (the evaporation flux becomes negligible) and the crystallization concentration is not reached in these screened pores. By contrast, the liquid flow rate in a surface pore in contact with efflorescence increases. Compared to flow before the onset of crystallization, this induces a reorganization of the flow structure within the upper layer of porous medium, which explains why the efflorescence continues to grow forming discrete and isolated structures.

in the porous medium explains both the peripheral localization (small δ) and the discrete nature of efflorescence onset.

A puzzling question is why the efflorescence continues to grow forming isolated structures leaving surface pores between efflorescence structures free of efflorescence. To explain this phenomenon it is crucial to note that the salt structures forming efflorescence are porous [10]. Evaporation takes place at the outer surface of each structure, which consequently pumps by capillarity the aqueous solution contained in the underlying porous material.

Hence, as sketched in Fig. 5, the efflorescence structures first act as sinks at the porous surface (the dissolved salt is directed toward the efflorescence). The second effect is a screening effect, i.e., the fact that the evaporation flux becomes negligible in the porous surface regions located between the efflorescence structures. This can be illustrated through a simple growth model in two-dimensions. The 60×100 square cells computational domain corresponds to the rectangle bounded by a green thin dashed line in Fig. 5. Thus the underlying porous medium is not considered. In the gas region, we solve numerically the vapor concentration transport equation assuming a quasisteady purely diffusive transport ($\Delta c_v = 0$) with the boundary condition: $c_v = 0$ at y = H, $c_v = 1$ at y = 0 (porous medium surface) and at the efflorescence boundary (which

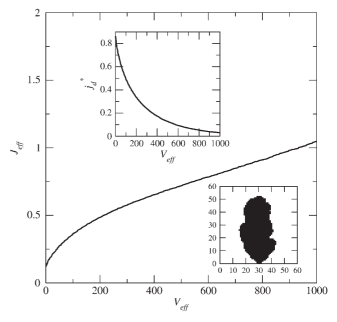


FIG. 6. Efflorescence pumping effect: evolution of evaporation rate J_{eff} from one efflorescence structure as a function of structure size V_{eff} (in number of cells). The screening effect is illustrated in the inset top, which shows the evolution of dimensionless mean evaporation flux between two efflorescence structures (along the thick dashed line in Fig. 5) as a function of structure size (the reference flux is the flux in the absence of efflorescence); an example of efflorescence computed shape is shown in the inset bottom.

initially is formed by two square cells located in the middle of the line y = 0). Spatially periodic boundary conditions are imposed in the lateral direction. The "evaporation rate" at each edge forming the efflorescence contour is then computed ($J_{ed} = -\nabla c_v$.**n** where **n** is the unit vector outwardly normal to the considered edge). The efflorescence grows locally proportionally to J_{ed} . Some disorder is introduced assuming that the porosity of each efflorescence cell varies randomly (details will be presented elsewhere). The results are summarized in Fig. 6 and illustrate the screening effect (top inset) and the local flow rate enhancement ($\propto J_{eff}$) explaining the efflorescence discrete growth (Fig. 5).

In summary, the quantitative theory of efflorescence formation and growth is yet to come but we believe that the phenomenology described in this Letter opens up the route in this direction.

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