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Mechanisms and scale-up potential of 3D solar interfacial-evaporators†

James H. Zhang, Rohith Mittapally, Abimbola Oluwade and Gang Chen*

Evaporation fluxes from porous evaporators under sunlight have been reported to exceed the solar-thermal limit, determined by relating the incoming solar energy to the latent and sensible heat of water, for applications in desalination and brine pond drying. Although flat two-dimensional (2D) evaporators exceeding the solar limit imply a non-thermal process, tall three-dimensional (3D) solar evaporators can exceed it by absorbing additional environmental heat into its cold sidewalls. Through modeling, we explain the physics and identify the critical heights in which a fin transitions from 2D to 3D evaporation and exceeds the solar-thermal limit. Our analyses illustrate that environmental heat absorption in 3D evaporators is determined by the ambient relative humidity and the airflow velocity. The model is then coarse-grained into a large-scale fin array device on the meters scale to analyze their scalability. We identify that these devices are unlikely to scale favorably in closed environment settings such as solar stills. Our modeling clearly illustrates the benefits and limitations of 3D evaporating arrays and pinpoints design choices in previous works that hinder the device's overall performance. This work illustrates the importance in distinguishing 2D from 3D evaporation for mechanisms underlying interfacial evaporation exceeding the solar-thermal limit.

Broader context

Over the last eight years, many groups have reported that evaporation rates from porous materials under sunlight can exceed the solar-thermal limit calculated by relating the incoming solar energy with the latent and sensible heat of water. There are broadly two classes of materials that are super solar-thermal: 2D and 3D materials. We have shown previously that super solar-thermal 2D evaporators imply a non-thermal evaporation process. In contrast, 3D evaporators can exceed the solar-thermal limit by absorbing environmental energy, which is still a thermal process. Here, we conduct extensive modeling on 3D thermal evaporators to clearly delineate 3D solar interfacial evaporators from 2D evaporators, explain the underlying mechanisms, and evaluate their scalability in large devices for desalination and brine pond drying. Our work pinpoints that 3D evaporators see the most benefit only in open environments under large bulk airflow conditions because environmental heat absorption is governed by the ambient humidity and vapor transport kinetics. This work lays a foundation for understanding the mechanisms underlying the super solar-thermal evaporation limit, and points to directions in exploring their potential applications.

Introduction

Passive solar evaporation to separate water from dissolved minerals has the potential to be a low capital cost and green method to produce clean water, harvest critical minerals, and treat wastewater ponds. The high latent heat of water relative to the solar flux strongly limits the evaporation rate of water. Using interfacial solar evaporating materials with capillary wicking abilities has been shown to be an especially promising

strategy due to its ability to concentrate the solar energy in a thin interfacial region near the evaporating surface.^{1–7} Under standard one sun insolation of 1000 W m^{-2} and assuming all solar energy is used for evaporation, one arrives at a maximum evaporation flux of around 1.45 to $1.49 \text{ kg m}^{-2} \text{ h}^{-1}$, which we call the solar-thermal limit. Machine learning has been applied to model solar evaporation devices as well to maximize their performances.^{8,9}

However, many studies have reported evaporation fluxes that exceed this solar-thermal limit.^{6,10–19} For 2D solar-driven interfacial evaporating materials in which the solar absorbing area is nominally the same as its evaporating area, reports have demonstrated evaporation fluxes beyond the solar-thermal limit by 2 to 4 times.^{6,10–13} Our recent work has shown that such high evaporation fluxes imply that water evaporates in the

Department of Mechanical Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, Massachusetts, 02139, USA.
E-mail: gchen2@mit.edu

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form of clusters, *i.e.* super solar-thermal, because no region of the evaporation system is below the ambient temperature and the reduced latent heat hypothesis is incorrect.^{20,21} Although the details of this phenomenon are still under investigation, our group has interpreted such super solar-thermal evaporation as arising from photons directly cleaving off water molecular clusters, which we called the photomolecular effect.^{10,20,22–24}

For tall evaporators, which we call 3D solar interfacial-evaporators and will quantitatively define them later, the evaporating surface area is much larger than its solar absorbing area due to its extended surface similar to fins used in heat transfer devices.^{15,19,25–29} It has been well appreciated in the field that these structures can exceed the solar-thermal limit due to the structure absorbing additional environmental energy along its sidewalls not exposed to direct sunlight.¹⁶ The reported evaporation fluxes normally range between 3 and 5 times the solar-thermal limit^{27,29} and sometimes even up to 7 times using forced convection,¹⁸ based on the projected top cross-sectional area of the absorber. Theoretical studies suggest that water can capillary rise up to meters in height in microporous materials,³⁰ and water has been observed to rise to 100 cm in soil.³¹ The maximum capillary flux in the 3D evaporator needs to be able to sustain the fins' total evaporation rate,¹⁹ with a past work demonstrating 3D evaporators up to 60 cm tall.¹⁷ The high evaporation fluxes have attracted lots of materials development and lab prototype testing. However, very few reports have analyzed the scale-up potential of 3D evaporators beyond a few fins on the decimeter device scale nor have they mechanistically studied the physics of the device.^{16,18,32–34} Experiments from Chen *et al.*¹⁷ showed that as the array of extended surfaces increase in number, the evaporation flux per structure decreases, illustrating the challenges of scaling up 3D solar evaporators. Yang *et al.*³⁵ hypothesized that the vapor from the solar absorbing region might re-condense in the evaporative cooling region due to the vapor concentration difference, leading to degraded performance with taller fin heights inside the condenser chamber. Although many works have analyzed 3D evaporators through heat transfer equations,^{15,16,19,29} very few have explicitly considered vapor transport which drives the environmental heat absorption.^{18,36} Currently, there lacks a systematic study to help guide these discussions for 3D solar-interfacial evaporators that considers both heat and mass transfer kinetics. No criteria have been established to clearly distinguish 2D from 3D effects, which is important for studying mechanisms behind solar-interfacial evaporation exceeding the solar-thermal limit.

In this work, we will systematically analyze the performance of 3D solar interfacial-evaporators, starting from a single fin and extend to scaled-up systems under both forced and natural convective conditions. First, a simplified model will be constructed to illustrate the performance of a single 3D solar evaporator and reveal the underlying physics. Criteria will be established to demarcate when the evaporating structures can be treated as 3D and when it will reach or exceed the solar-thermal limit. Then, the model will be coarse-grained to a large solar device to study its scale-up potential under forced convective

conditions. Our model illustrates that environmental heat input can only occur if the ambient air is below 100% relative humidity and the rate of environmental heat input depends on the airflow velocity and the humidity. Our analysis shows that despite the high performance of a single and a few fins, the performance of large-scale 3D solar evaporator arrays will degrade significantly in low airflow regimes and in closed environments, such as in solar still devices, because of limitations in vapor transport kinetics.

Results and discussion

Mechanisms and performance of a single fin

The performance of a 3D solar evaporator is driven by the balance between evaporative cooling effects from vapor transport and heat transport kinetics from the ambient environment. We will consider a cylindrical shaped 3D solar evaporator, often called a pin shaped fin, evaporating into a large ambient reservoir. In a typical laboratory setting, the fin is inserted into a larger diameter container of water (Fig. 1(a)). The fin is placed into an ambient reservoir at temperature T_∞ , ambient humidity RH, and characteristic bulk airspeed u_∞ . The ambient humidity and temperature determine the ambient vapor mole fraction $c_{v,\infty}$. For now, we assume the excess surface area of the container is covered and the solar beam spot size exactly matches the fin's cross-sectional area with light intensity $\dot{q}_{\text{sun}}''$. The foam thickness leads to an additional thickness of the fin t_{base} not exposed to the ambient air and in contact with a water reservoir underneath at temperature T_{bot} . The top surface becomes elevated in temperature T and has a high saturated vapor mole fraction $c_{v,s}$. The sidewalls that are not exposed to direct sunlight drops below the ambient temperature due to evaporative cooling effects, leading to heat absorption from the environment to sustain further evaporation. The solar-thermal limit for 2D evaporation can be calculated by assuming all solar energy is used to heat up and evaporate water, *i.e.*,

$$\dot{m}_{\text{sun,max}}'' = \frac{\dot{q}_{\text{sun}}''}{h_{\text{fg}} + c_p \Delta T_s} \quad (1)$$

where h_{fg} is the latent heat of water, c_p is the specific heat of water, and ΔT_s is the temperature rise of the surface. The above expression gives values between 1.45 and 1.49 kg m⁻² h⁻¹ depending on the evaporating surface temperature of the solar absorbing region.

Single fin model and metrics. We constructed an ideal model for a single fin, including solar heating on top surface, heat conduction along the fin, and evaporative, radiative, and convective heat exchange with the ambient (see the Methods section). Unless otherwise noted, the performances are calculated using a base case scenario in which the extended surface has a diameter (D) of 2.5 cm, a height (H) of 10 cm, a thermal conductivity (k_f) of 0.3 W m⁻¹ K⁻¹, a blackbody emissivity (ε) of 0.95, and evaporating into an ambient at (T_∞) 23 °C, 30% relative humidity (RH), and an external sidewall convective heat transfer coefficient (h_{conv}) of 5 W m⁻² K⁻¹. We set t_{base} to 2 cm and the water convective heat transfer coefficient h_{bot} at the bottom of the fin to 100 W m⁻² K⁻¹. We will show later that the



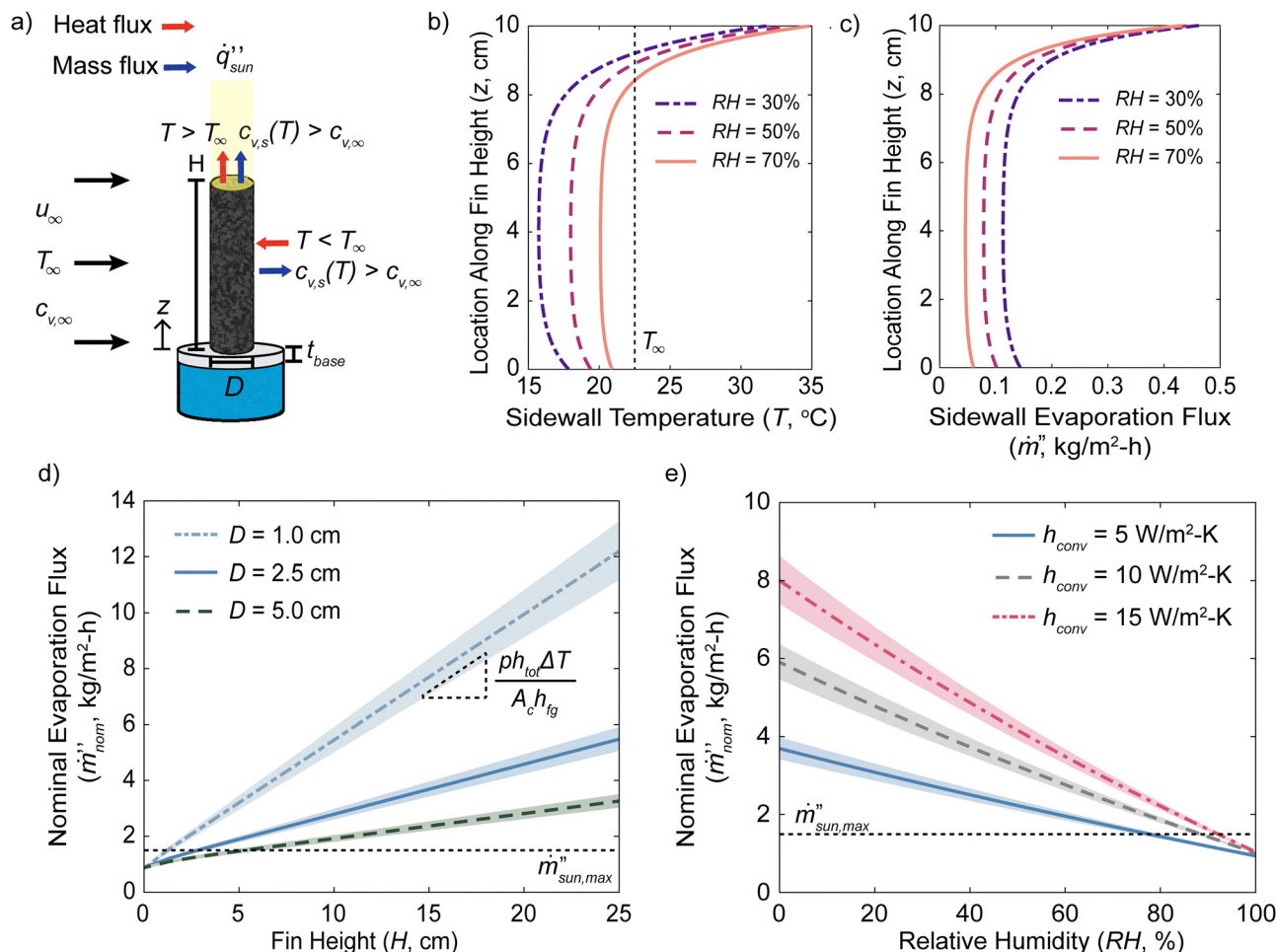


Fig. 1 Performance and mechanism of a single 3D evaporating fin. (a) Diagram of heat transfer, mass transfer, and typical testing geometry in laboratory test. T is the temperature and $c_{v,s}$ is the saturated mole fraction of vapor at the specified temperature. Unless stated otherwise inside the figure, the studied base case is a fin with a height (H) of 10 cm, a diameter (D) of 2.5 cm, and a thermal conductivity (k_f) of $0.3 \text{ W m}^{-1} \text{ K}^{-1}$ evaporating into an ambient at (T_{∞}) 23°C , 30% relative humidity (RH), and a convective sidewall heat transfer coefficient (h_{conv}) of $5 \text{ W m}^{-2} \text{ K}^{-1}$. Predicted sidewall (b) temperature profile and (c) local evaporation flux for different RH. (d) Predicted nominal evaporation flux's dependence on fin geometry. (e) Predicted nominal evaporation flux as a function of RH and h_{conv} . The shaded regions in (e) and (f) are variations in the nominal evaporation flux if the ambient temperature changes by $\pm 10\%$ relative to its value in °C.

predicted performances are strongly independent of these two values. The full model is described in the Methods section. Table 1 lists all variables used in the model and their descriptions. The diameter and heights were chosen due to the similarity in size with many previously published works.^{15,16,29,36} The chosen thermal conductivity value was used because previous evaporators are commonly based on carbonized or organic materials,^{16–18,29,32} which would have thermal conductivities on the order of $0.1 \text{ W m}^{-1} \text{ K}^{-1}$.³⁷ Water has a thermal conductivity of about $0.6 \text{ W m}^{-1} \text{ K}^{-1}$ and the thermal conductivity of the wetted material would be somewhere in the middle. The emissivity was chosen to match that of water due to the assumption that the surface is wetted. Organic materials typically have blackbody emissivity values of about 0.9, which do not deviate significantly from water.³⁷ Previous works have conducted experiments in laboratory humidity values ranging from 20 to 50% and temperatures close to 23°C .^{15–18,28,29} We will analyze the results in a forced convection setting with bulk air velocity u_{∞} in crossflow to

the cylinder to decouple the heat transfer coefficients from the surface temperatures but expect similar results for natural convection (Supplementary Note S1, see the ESI†). For a given convective condition, the heat transfer coefficients h are interrelated with the vapor mass transfer coefficients g_m because of similarities in the boundary layers above the surface.³⁸ The diameter of the fin and the bulk airspeed are interrelated with the convective coefficients (see the Methods section).

We will define the nominal evaporation flux, $\dot{m}_{\text{nom}}''$, as the total evaporation rate of the fin normalized to only the top projected cross-sectional area A_c . The nominal evaporation flux is the metric commonly reported in previous literature to characterize their performance and governs the solar absorption area when light is incident only on the top surface.

$$\dot{m}_{\text{nom}}'' = \frac{\int \dot{m}'' dA_{\text{wetted-surface}}}{A_c} \quad (2)$$



Table 1 Variable list and descriptions in the fin model

Symbol	Description
Greek symbols	
β	Fin parameter.
ΔT	Temperature difference between the flat region and ambient.
$\Delta \dot{q}_{\text{env}}$	Total heat exchange from the airflow to the device.
ε	Emissivity of the wetted fin.
θ	Zenith angle of sunlight
σ	Stefan-Boltzmann constant.
Roman symbols	
A_{base}	Device area in one control volume, equal to $S_{\text{f}}S_{\text{t}}$.
A_{c}	Cross-sectional area of the fin.
A_{dev}	Total device footprint area.
C_{g}	Molar density of air.
c_{p}	Specific heat capacity.
$c_{\text{v},\infty}$	Mole fraction of vapor in ambient air.
$c_{\text{v},\text{s}}$	Saturated mole fraction of vapor.
D	Diameter of fin
D_{v}	Diffusion coefficient of vapor in air.
g_{m}	Convective mass transfer coefficient of vapor in air.
H	Height of the fin.
$H_{\text{cr},2\text{D}}$	Critical height to absorb environmental heat.
$H_{\text{cr},\text{th}}$	Critical height to nominally evaporate at $\dot{m}''_{\text{sun,max}}$.
h_{bot}	Convective heat transfer coefficient of water underneath.
h_{conv}	Sidewall convective heat transfer coefficient in air.
h_{fg}	Latent heat of evaporation of water.
h_{tot}	Combined radiative and sidewall heat transfer coefficient.
k_{f}	Thermal conductivity of the wetted fin.
L	Total length of the device.
$\dot{m}''$	Evaporation flux of water.
$\dot{m}''_{\text{dev}}$	Evaporation flux normalized to device footprint area.
$\dot{m}''_{\text{nom}}$	Evaporation flux normalized to top area.
$\dot{m}''_{\text{sun,max}}$	Solar-thermal limit for 2D evaporation.
N_{cols}	Number of columns of fins in the device.
N_{rows}	Number of rows of fins in the device.
Nu	Average Nusselt number.
$\dot{n}_{\text{air}}$	Molar flowrate of air.
p	Outer perimeter of fin's cross-section.
Pr	Prandtl number.
$\dot{q}$	Rate of heat transfer
$\dot{q}_{\text{sun}}$	Solar flux on top surface.
Re	Reynolds number.
RH	Relative humidity of ambient air.
Sc	Schmidt number.
Sh	Average Sherwood number.
S_{l}	Longitudinal spacing between each row of fins.
S_{t}	Transverse spacing between each column of fins.
T	Temperature.
T_{∞}	Temperature of ambient air.
T_{bot}	Temperature of water reservoir.
t_{base}	Fin base thickness.
u_{∞}	Bulk airflow velocity.
u_{max}	Maximum airspeed in the device from venturi effect.
W	Total width of the device.
x	Axis along the device width.
y	Axis along the device length.
z	Axis along fin's height.
Subscripts	
base	Relating to fin array's base plate.
bot	Relating to water reservoir.
conv	Relating to fin's sidewall area.
i	Relating to i th row in the device.
top	Relating to fin's top face.

Later, we will further discuss evaporation flux normalized to the device footprint area.

Performance of a single fin. Fig. 1(b) illustrates the predicted surface temperature along the fin's sidewalls for different RH values, illustrating that the constructed model captures the physics of 3D evaporators shown in Fig. 1(a). The top evaporating surface is above the ambient temperature due to solar absorption. About 1.5 cm below the hot top surface, the fin drops below the ambient temperature from evaporative cooling. In the middle region along the sidewall, the temperature profile is flat, a common observation in previous experiments.^{15–18,29} The flat temperature value is strongly determined by the ambient humidity. Near the bottom of the extended surface, the temperature increases again due to the heat transfer with the water reservoir beneath the fin. Fig. 1(c) shows the local evaporation flux along the fin's sidewall for the same given conditions. The local sidewall evaporation flux from the flat temperature region is between 73% and 89% smaller than the hot solar absorbing region near the top because the saturated vapor concentration decreases rapidly with the surface temperature. The solar absorbing “hot” top region has much faster evaporation kinetics than the evaporatively cooled “cold” regions along the middle region.

Due to the many variables involved in the model, we have also conducted a sensitivity analysis from the chosen base case by calculating the percentage change in the nominal evaporation flux per percentage change of the variable to understand the most influential terms (Fig. S1, see the ESI†). The most sensitive variables are the fin diameter (−0.93), the ambient temperature (0.67), the fin height (0.64), the solar intensity (0.33), the ambient humidity (−0.31), and the bulk airspeed (0.27). The results are less sensitive to the blackbody emissivity (0.10) and very insensitive to the fin's thermal conductivity (−0.0009). From this analysis, we can appreciate which variables are the most important for single fin performance.

Using the model, we can predict the nominal evaporation flux's dependence on the fin height and diameter (Fig. 1(d)). The variations in the nominal evaporation flux with respect to ambient temperature are shown by the shaded region if the ambient temperature changed by $\pm 10\%$. Due to the convective heat and mass transfer coefficients' dependence on both fin diameter and bulk airspeed, we varied the bulk airspeed for the different diameters to maintain a constant sidewall convective heat transfer coefficient of $5 \text{ W m}^{-2} \text{ K}^{-1}$ (see the Methods section).

The model predicts that the nominal evaporation flux increases linearly with the fin height due to the increased evaporating surface area assuming no dry out occurs. In a real system, there will be a limit to the height of the fin based on the capillary water pumping in the designed fin. The reason why the nominal evaporation flux increases linearly with fin height is explained with the temperature profiles found previously. As the fin gets taller, it has minimal effects on the temperature profiles near the hot solar-absorbing top and the cooler bottom exchanging heat with the water reservoir below. These features are determined by the balance of the fin's heat conduction along its length with the heat and vapor exchange with the



ambient reservoir at the respective ends. Increasing the fin's height only increases the length of the flat temperature profile region in the middle. The energy balance in the flat temperature region is a simple relationship between the evaporative cooling from vapor transport kinetics and the combined convective and radiation heat transfer coefficient with the ambient.

$$g_{m, cov} C_g h_{fg} (c_{v, s}(T) - RH c_{v, s}(T_{\infty})) = h_{tot} (T_{\infty} - T) \quad (3)$$

where C_g is the molar density of air and h_{tot} is the combined convective and radiative heat transfer coefficients. Eqn (3) illustrates that the properties of the flat temperature region is independent of the fin's properties beyond its blackbody emissivity and driven by the air side boundary layers.³⁸ At high airflow external velocities when radiation becomes negligible relative to convection, this temperature should approach the wet-bulb temperature.

From Fig. 1(c), it is evident that the sidewall evaporation flux due to environmental heat flux is about 10 times lower than the maximum solar-thermal rate and the enhanced nominal evaporation flux is due to the cross-sectional area normalization. The wet-bulb temperature is determined by the ambient temperature and relative humidity and it is the lowest temperature the surface can achieve due to evaporative cooling (Fig. S2a, see the ESI†). Using this, we can estimate what the convective sidewall heat transfer coefficient needs to be for the fin's sidewalls to evaporate at the solar-thermal limit due to environmental heat input (Fig. S2b, see the ESI†). It can be seen that except for very high humidity values above 90%, the required sidewall convective heat transfer coefficient is on the order of $100 \text{ W m}^{-2} \text{ K}^{-1}$. The radiation heat flux contribution was neglected because its linearized heat transfer coefficient will only be about $5 \text{ W m}^{-2} \text{ K}^{-1}$, which is much smaller than the findings in Fig. S2b (ESI†). For a fin diameter of 2.5 cm, this would require an extremely high airspeed at around 17 m s^{-1} .

The performance of 3D evaporators and its environmental heat absorption is governed by the ambient humidity near the fin and the external airflow velocity. If the ambient is near 100% RH, eqn (3) will simplify to the zero solution and no environmental heat absorption can occur. Using this knowledge, we can estimate that the slope governing the increase in the nominal evaporation flux with the fin height is

$$\frac{dm''_{nom}}{dH} = \frac{p h_{tot} \Delta T}{A_c h_{fg}} \quad (4)$$

where ΔT is the temperature difference between the flat region and the ambient from eqn (3) and p is the perimeter of the fin. Linear growth in the nominal evaporation flux with height up to the fin dry-out was also observed in a previous study.³⁴

Due to the strong dependence of the nominal evaporation flux on the ambient humidity and convective conditions, we further plot the nominal evaporation fluxes as a function of these two variables (Fig. 1(e)). Similar to Fig. 1(d), the effects of ambient temperature variations are included using the shaded region as well. As the sidewall convective heat transfer coefficient, and consequentially the convective mass transfer coefficient, increases from 5 to $15 \text{ W m}^{-2} \text{ K}^{-1}$, the nominal

evaporation fluxes increase by over 2 times at 10% RH and are above the solar-thermal limit due to environmental heat absorption. However, as the ambient humidity increases to 100% RH, all curves predict that the performance will degrade to below the solar-thermal limit for 2D structures and converge towards similar values. At 100% RH, the 3D evaporator will always behave like a 2D evaporator regardless of its height. The structure will continue to evaporate under this condition, albeit at a very low rate compared to the solar-thermal limit, because the solar absorbing top region will be at an elevated temperature compared to the environment. This will cause the saturated vapor mole fraction at the surface to be higher than the vapor mole fraction at 100% RH in the ambient air. In this scenario, the fin's sidewall temperature can never drop below the ambient.

It is important to note that there may be some ambiguity in the definition of what constitutes a wetted-surface area in eqn (2). Many previous 3D interfacial evaporators are highly porous with surface roughness that will cause the surface area to be higher than that of a smooth cylinder. Similar to our previous analysis on 2D evaporators, the key comparison is the characteristic size of the surface roughness to the vapor concentration boundary layer thickness.²⁰ Using the studied conditions from Fig. 1, we can estimate that the vapor concentration boundary layer thickness will be on the order of mm to cm.³⁸ If the surface roughness is significantly smaller, such as on the order of microns, the vapor does not have kinetics to escape from the enhanced surface area into the far field because the macroscopic boundary layer is the rate limiting resistance. If the characteristic size is comparable with the boundary layer, then the fin's perimeter, cross-sectional area, and wetted surface area in eqn (2) should account for the enhanced surface area.

Critical heights for 3D evaporators

There is a critical height in which the evaporator will transition from a 2D to a 3D evaporator that can absorb environmental heat. If the water reservoir beneath the fin has the same temperature as the ambient, we can identify this critical height, $H_{cr, 2D}$, as the point in which the fin's temperature profile will drop below the ambient temperature due to evaporative cooling and start to absorb environmental heat (Fig. 2(a)). The shaded regions represent the variations in the critical heights predicted if the ambient temperature changes by $\pm 10\%$. When the fin behaves like a 2D evaporator, the temperature profile will monotonically decrease from the hot evaporating top surface to the ambient bottom temperature. This suggests that the $H_{cr, 2D}$ will be related to the fin parameter β .

$$\beta = \sqrt{\frac{h_{conv} p}{k_f A_c}} \quad (5)$$

When the convective heat transfer coefficient is at $5 \text{ W m}^{-2} \text{ K}^{-1}$ and the ambient humidity is 50%, the non-dimensional critical height is about 0.85. Using the base case scenario for a diameter of 2.5 cm, this corresponds to a height of about 1.15 cm. For structures with much smaller cross-section areas,



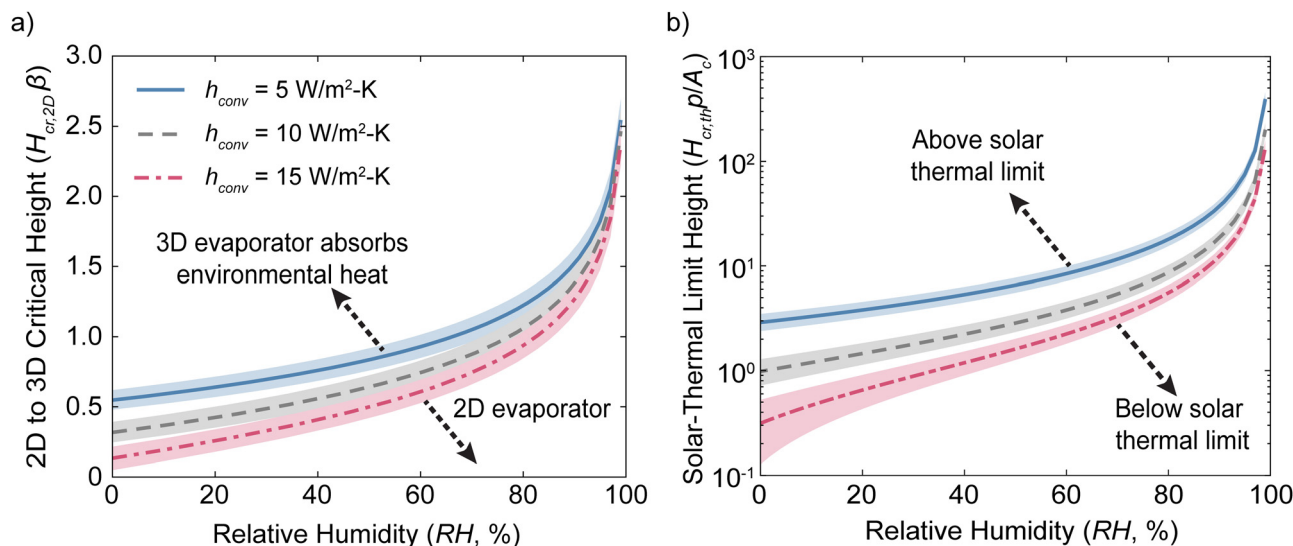


Fig. 2 Critical heights of 3D evaporators. (a) Height $H_{cr,2D}$, normalized to the fin parameter β , is when a 2D evaporator will behave like a 3D evaporator and absorb environmental energy. (b) Height $H_{cr,th}$, normalized to the fin's aspect ratio, is when a 3D evaporator will absorb enough environmental heat to exceed the solar-thermal limit. Each curve corresponds to a different convective condition. The plots were produced using the base-case scenario values described above unless otherwise stated in the figure. The shaded regions in (a) and (b) are variations in the nominal evaporation flux if the ambient temperature changes by $\pm 10\%$ relative to its value in $^{\circ}\text{C}$.

this critical height is significantly lower.²⁷ Fins with heights below $H_{cr,2D}$ can only lose heat to the environment whereas fins taller than $H_{cr,2D}$ can absorb heat from the environment. As the RH approaches 100%, $H_{cr,2D}$ diverges and can never absorb environmental heat. Higher convective heat transfer coefficients decrease $H_{cr,2D}$ because the evaporative and convective cooling will remove the solar heat more rapidly. The critical height is not just a function of the nondimensional parameters due to the nonlinearity of the saturated vapor molar concentration dependence on temperature.

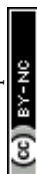
From the discussion on 2D to 3D evaporator transition, it will be meaningful to further identify how tall a fin needs to be, $H_{cr,th}$, for the nominal evaporation to reach the solar-thermal limit (Fig. 2(b)). $H_{cr,2D}$ doesn't determine the critical height needed to exceed $\dot{m}_{\text{sun,max}}''$ because the fin first needs to absorb sufficient environment heat to compensate for the heat lost from convective and radiative cooling at the top solar absorbing region. Rather than normalizing to the fin parameter β , $H_{cr,th}$ is normalized to the fin's geometric aspect ratio p/A_c because the flat-temperature middle region that absorbs environmental energy is independent of the fin's properties if dry-out doesn't occur. As the ambient humidity increases, $H_{cr,th}$ increases exponentially because the environmental heat absorption and evaporation are determined by the ambient humidity. For the same reason with $H_{cr,2D}$, $H_{cr,th}$ will also diverge as the ambient humidity approaches 100%. Many previous experiments have fins with fin aspect ratios Hp/A_c on the order of 10 and testing in environmental humidity below 50%.^{15–19,29} Fig. 2(b) suggests that these will achieve super solar-thermal evaporation fluxes.

We have also studied the robustness of the critical heights predicted by the model to variations in the base case parameters (Fig. S3, see the ESI[†]). The curves illustrate the changes

in the critical heights predicted if the thermal conductivity k_f (Fig. S3a and b, ESI[†]), fin diameter D (Fig. S3c and d, ESI[†]), water reservoir convective heat transfer coefficient h_{bot} (Fig. S3e and f, ESI[†]), and the base thickness t_{base} (Fig. S3g–h, ESI[†]) changed by $\pm 50\%$ of the base values used. The predicted heights to reach the solar-thermal limit $H_{cr,th}$ is very independent of these variables because the rate of environmental heat absorption is governed only by the airside properties, as shown in eqn (3). The predicted height when 2D evaporators transition to 3D evaporators $H_{cr,2D}$ has a weak dependence on the fin's thermal conductivity and diameter. $H_{cr,2D}$ is independent of the water reservoir heat transfer coefficient and the base thickness. The weak dependence on the thermal conductivity is due to the nonlinearity in the model introduced by the evaporative cooling from vapor transport. The weak dependence of $H_{cr,2D}$ on the fin diameter is due to the convective heat and mass transport on the top cross-sectional surface of the fin. Although the sidewall convective heat transfer coefficient is kept the same, the top cross-sectional's convective coefficients change slightly due to the different scaling with the fin diameter, causing $H_{cr,2D}$ to shift slightly as well.

Performance of scaled-up array in outdoor setting

In a scaled-up array of 3D fins for outdoors experiments, many conditions are significantly different from the laboratory setting (Fig. 3). The total solar energy and solar absorbing area of the device vary with the time of day due to the changing solar zenith angle θ . The fins will absorb additional sunlight along its sidewalls, cast a shadow, and shade the projected fin's area behind it.¹⁹ For small devices where fins have much taller heights than the device length, the device will absorb significantly more solar energy than the device area footprint, A_{dev} .



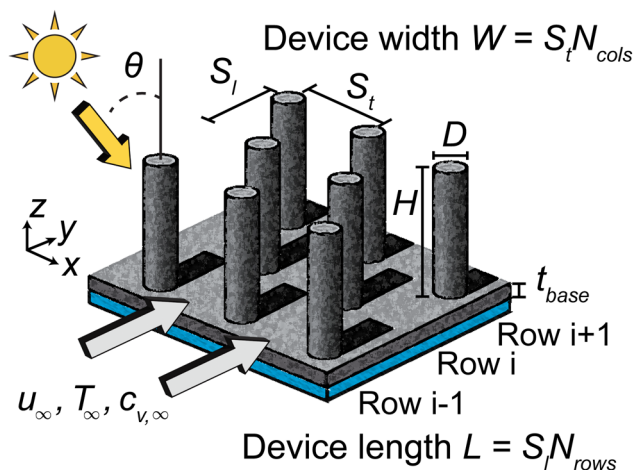


Fig. 3 Diagram of scaled-up fin array device in outdoor setting. The time of day will change the solar absorbing area of the device due to the zenith angle θ . The total device length L and width W depend on the number of fin columns and rows in the array. The geometric dimensions of the fins and air movement in crossflow to the fins are illustrated as well. The ambient air conditions are set by its temperature T_∞ , vapor mole fraction $c_{v,\infty}$, and bulk air speed u_∞ .

This is because the shaded area will extend beyond the device's footprint. The device's footprint area can be related to the device length and width as

$$A_{\text{dev}} = (S_l N_{\text{rows}})(S_t N_{\text{cols}}) \quad (6)$$

where N_{rows} is the number of rows of fins in the y -axis and N_{cols} is the number of columns in the transverse x -axis, S_l is the spacing length between rows, and S_t is the spacing width between columns. The total length of the device L is simply $S_l N_{\text{rows}}$. This additional area that absorbs sunlight will become negligible as the device is scaled-up in size where $L \gg H$ because this is only an edge effect of the backmost row of fins, leading to the total solar energy incident on the device to be approximately equal to

$$\dot{q}_{\text{sun}} = \dot{q}_{\text{sun}}'' A_{\text{dev}} \quad (7)$$

In many previously reported outdoor devices, the base of the device is a non-evaporating and non-absorbing insulating material such as white foam instead of an interfacial evaporator. We will show later that this configuration will degrade the true performance of the overall device because a significant amount of solar energy will be wasted when it can be used to further evaporate water. Finally, we define a device evaporation flux that is normalized to the device footprint area to give a representative metric for scaled-up devices that consider the solar absorption of the entire structure.

$$\dot{m}_{\text{dev}}'' = \frac{\int \dot{m}'' dA_{\text{wetted-surface}}}{A_{\text{dev}}} \quad (8)$$

This metric is only accurate either in the condition when $L \gg H$ or when $\theta = 0$, with the latter representing the best

scenario, and hence our discussion leads to an upper limit in evaporation fluxes.

Forced convection device model. The transport kinetics of vapor through the device determines the performance of the fin array because the environmental heat absorption depends on the air remaining below 100% RH. As the air flows through the fin array device, it will progressively become more humid and diminish the performance of the fins further downstream. If the device becomes too large, the hotter humid air can cause vapor recondensation on the sidewalls and decrease its overall evaporation rate. These effects suggest that there will be an optimum fin sizing and spacing given a total device length L , solar intensity $\dot{q}_{\text{sun}}''$, external bulk airspeed u_∞ , and reservoir conditions. However, the large mismatch in length scales between the boundary layer thickness and the device size make FEA simulations of an entire fin array very computationally expensive.

Using the single fin model, we developed a coarse-grained model based on a control volume analysis of the airflow (see the Methods section for details). The list of variables included are shown in Table 1 above. In this model, we draw a series of control volume along the y -axis of the array. Each control volume along the y -axis has dimensions S_t and S_l such that only one fin is inside. We assume that air only flows along the y -axis such that each fin in the same row behaves identically. We will consider a fin with the same characteristics as the single fin analysis. The transverse spacing S_t is $4D$ (10 cm) and the longitudinal spacing S_l is $2D$ (5 cm). This will cause the extended surface to have about 85% more evaporating surface area than the base plate around it in each control volume. The incoming air is initially at 23 °C and 30% RH. For solar absorption calculations, we will assume that the zenith angle is 0° and that the incoming solar intensity is 1000 W m⁻², all absorbed where the solar ray lands (on the top of the fin and the base plate around the fin). The total area, A_{base} , that absorbs sunlight in each control volume would simply be the product of the two array spacings. This should correspond to the maximum evaporate rate since in this case, the side wall is not heated directly by the solar radiation. In addition to the total device evaporation flux $\dot{m}_{\text{dev}}''$, we will further define a local device evaporation flux to illustrate the performance of each row of fins, denoted by row index i .

$$\dot{m}_{\text{dev},i}'' = \frac{\int \dot{m}_i'' dA_{\text{wetted-surface}}}{A_{\text{base}}} \quad (9)$$

where A_{base} is equal to $S_t S_l$. These parameters will depend on the local air temperature T_i and humidity RH_{*i*} at the specified row of fins in the device.

Performance of the device. We can now analyze the detailed airflow properties to better understand the mechanisms of the device. Fig. 4(a) illustrates the local air humidity along the length (y -axis) of the device with 50 rows in width (2.5 m) for both solar and dark conditions. The air's RH initially increases rapidly and then its growth decays further downstream in the device. The RH doesn't increase as rapidly under dark



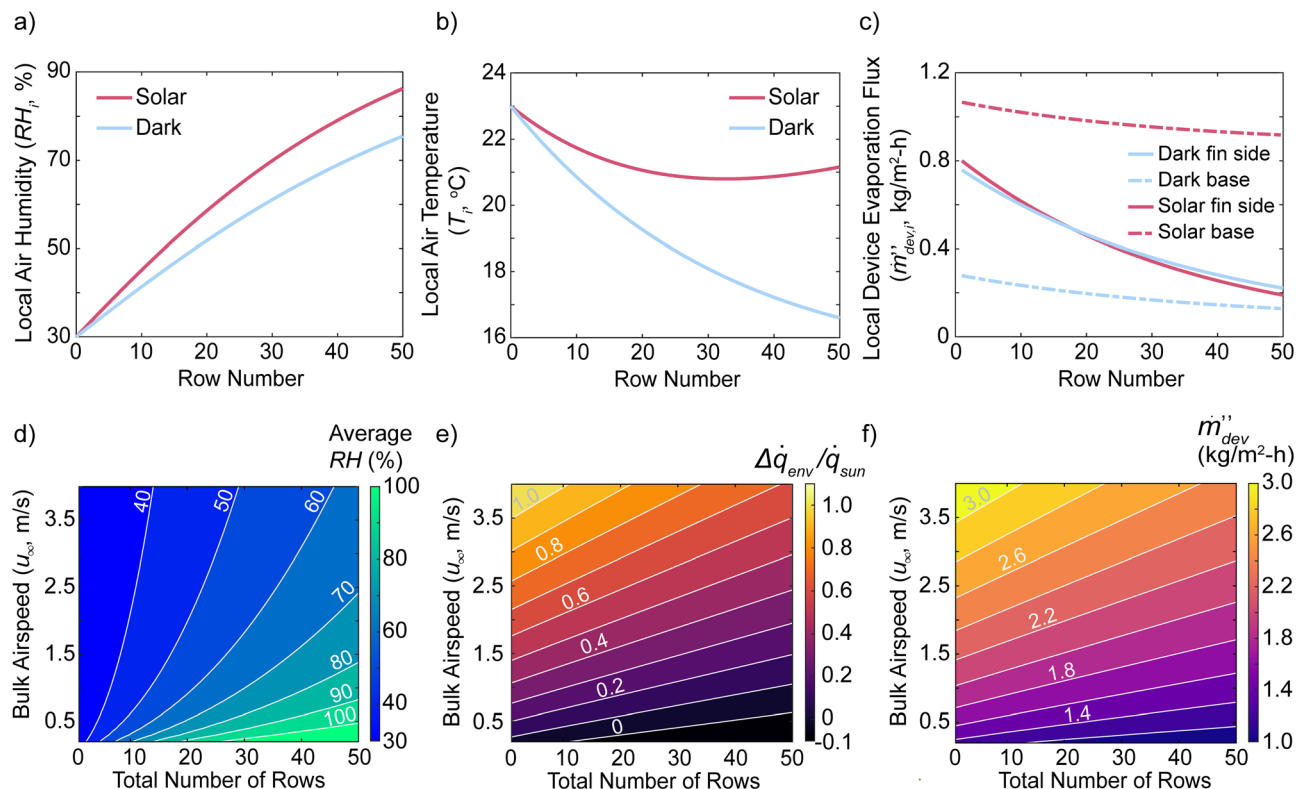


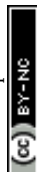
Fig. 4 Heat and vapor exchange with airflow through the device for large scale systems. (a)–(c) Properties and performance of a device 50 rows (2.5 m) long with an external bulk airspeed at 1 m s^{-1} . Local air (a) relative humidity and (b) temperature for dark and solar conditions as a function of row number. (c) Contributions from the base plate and the fins' sidewalls to the local device evaporation flux. (d)–(f) Performance of devices with different lengths and subjected to different bulk airspeeds. (d) Average relative humidity of air throughout the entire device. (e) Ratio of total environmental heat exchange to total solar absorption. (f) Total device evaporation fluxes. The fins have a diameter of 2.5 cm, a height of 10 cm, a thermal conductivity of $0.3 \text{ W m}^{-1} \text{ K}^{-1}$, a transverse fin spacing of 10 cm, and a longitudinal fin spacing of 5 cm. The incoming air is initially at 23°C and 30% RH. The variable lists are included in Table 1 above.

conditions as under solar conditions. Environmental heat exchange between each row and the airflow will cause the local air temperature to change as well (Fig. 4(b)). For the solar condition, the air temperature initially drops, reaches a local minimum, and then increases afterwards. For dark conditions, the local air temperature monotonically decreases.

The reason for these trends is revealed in Fig. 4(c), illustrating the contribution of the solar absorbing base plate area and the fin's sidewall contributions to the local device evaporation flux per row. For fins under both dark and solar conditions, the rows near the leading edge of the device contribute a significant amount of evaporation due to the low humidity in the air. In fact, their performances are almost identical because their evaporation fluxes are primarily determined by 3D environmental heat absorption effects. This causes the local air temperature to cool down due to the fins' sidewall environmental heat absorption. For the base plate near the leading edge, it will heat the air due to convective heat loss under solar conditions and cool the air under dark conditions due to an evaporative cooling effect like the fins' sidewalls. Further downstream inside the device, the humidity will increase and the fins' sidewall evaporation becomes less effective, leading to the local fin sidewall evaporation fluxes to degrade rapidly. The base

plate downstream under solar conditions continues to both evaporate into and heat up the air due to its enhanced temperatures. Under dark conditions, the base area's local device evaporation flux also decreases rapidly downstream. The balance between the fins' sidewall heat gain and the base area's heat loss coupled to the changing humidity and local evaporation fluxes create the air temperature minimum under solar conditions. In the initial rows of the device, the local device evaporation flux exceeds the solar-thermal limit due to the low humidity. Further downstream as the air saturates, the local device evaporation flux degrades to below the solar-thermal limit.

These calculations illustrate that the common procedure in previous works, in which the base area is covered with a non-evaporating and non-absorbing material such as white foam, will degrade the device performance significantly. The fins will only have a high device evaporation flux if the local air RH is below 100%. In contrast, the base plate will continue to evaporate water because it heats up from solar absorption, leading to higher saturated vapor molar concentration and enhanced evaporation kinetics. Not utilizing the total solar energy on the entire device will lead to large reductions in performances and this behavior is evident if the device



performance is normalized to the entire area footprint rather than only the fin's cross-sectional area.

Our single fin analysis shows that environmental heat gain can only occur if the air flow inside the device is below 100% humidity. Fig. 4(d) illustrates the average humidity of the air across the entire device as a function of its total number of rows (length) and external airspeeds under solar conditions. At low airspeeds, the air will rapidly saturate as the device gets wider. At a bulk airspeed of 2.5 m s^{-1} , the average RH is 70% if the device has 50 rows (2.5 m) in length. This scaling relation is because the rate at which air is replaced in the device scales linearly with the bulk airspeed and the rate of evaporation scales sub-linearly with the bulk airspeed due to the Sherwood number relationship (see the Methods section below). From this analysis, we can appreciate that larger devices and greater environmental gain can be achieved only if the external bulk airspeed is high and the ambient air is dry. Fig. 4(e) shows the total environmental heat gain $\Delta\dot{q}_{\text{env}}$ relative to the total solar energy $\dot{q}_{\text{sun}}$ the device absorbs. At lower airspeeds and longer device lengths, the device will lose heat to the environment ($\Delta\dot{q}_{\text{env}} < 0$) because the hot base loses heat faster than the sidewalls can absorb heat. This is because the humid air cannot be replaced by dry air fast enough to enhance the fins' sidewall evaporation. The corresponding device evaporation fluxes are illustrated in Fig. 4(f), which can exceed the solar-thermal limit under certain conditions. For a given bulk airspeed, there is an optimum number of fins and length the device can be scaled to. Due to the diminished performance of the fins as the device gets larger, the device performance will degrade and eventually be below the solar-thermal limit. If the device is too long, water vapor will evaporate from the hot base and recondense on the fins' sidewalls further downstream and decrease the device performance even further. From this analysis, the airflow velocity and ambient humidity dictate the potential scale-up of 3D evaporating arrays.

Performance of the device under natural convective conditions

The above discussion made it clear that the enhanced nominal evaporation flux of the extended fin due to environmental heat input depends on the humidity of its surrounding air. Under natural convective conditions, the airflow speed and its effects on vapor transport are strongly coupled to temperature and vapor concentration gradients due to induced buoyancy effects.^{38,39} Surfaces hotter than the ambient environment will cause the air to rise upwards and form a plume. Surfaces colder than the ambient will flow downwards. If the base area is an interfacial solar absorbing material, it will become hotter than the ambient. If the base plate is reflective like white foam, the temperature will be slightly elevated, and the diffuse solar reflections from the foam base will cause the bottom region of the evaporating fin to become hotter.

The airflow velocities will depend on the geometry and the boundary layer that forms over the entire device. The natural convection velocities for single fins in these geometries are on the order of cm s^{-1} . For scaled-up devices, the airflow velocity will decrease due to the thicker boundary layers it forms

(Fig. 5(a)). The fins near the edge of the device will have the highest heat transfer coefficients and airflow velocities due to it being adjacent to the air reservoir. The interior airflow velocities will be much lower, and it will decrease as the device's transverse width and length (x - and y -axis) increase. The sluggish airflows can lead to the hot vapor near the fins solar absorbing top and base to recondense on the colder sidewalls, as hypothesized in a previous experimental study.³⁵ This will strongly limit the scale-up potential of these evaporating fins under natural convective conditions.

Closed system devices can't absorb environmental heat.

These challenges are evident in closed environment devices, such as solar still desalination devices (Fig. 5(b)). Enhanced performance from environmental heat absorption only occurs if the air is below 100% RH. Furthermore, it relies on continuous air supply from the environment because the air will continue to cool down and become more humid as the fins absorb environmental heat as shown in Fig. 4. In a closed environment system, there is no net airflow and only internal circulation patterns will form. The condenser should be the coldest temperature in the system and will be at 100% RH near its surface. The hot solar evaporating surfaces are at an elevated temperature and its corresponding 100% RH as well. At steady-state, the air inside of a simple solar still device must be close to the saturated condition because it is bound by these two conditions. Unless the device's lateral dimension is so small that a non-condensing sidewall of the enclosure remains below the ambient temperature and transfers heat from outside, the only other heat inputs into the device are from the brine reservoir below and solar absorption. Thus, large-scale conventional solar stills relying on internal circulating natural convection cannot exceed the solar-thermal limit.

FEA simulation of recondensation. To further illustrate these challenges, we used transient finite element analysis (FEA) to simulate natural convective evaporation in a 1D straight fin geometry with sunlight normal to the z -axis ($\theta = 0^\circ$). The highlighted yellow regions indicate the solar absorbing interfaces. This simulation will be representative of a very large system in which the interior airflow velocities become small. Fig. 5(c) illustrates the temperature profiles and the natural convection flow patterns that form after 10 minutes of simulation time if the base evaporates water as well. Fig. 5(d) illustrates the vapor molar density profile and diffusion vapor fluxes for the same case. Fig. 5(c) shows an internal natural convection flow pattern forming between the fins. The base plate's solar absorbing region is not as hot as the fin's tips due to heat exchange with the brine reservoir beneath it. The fin's sidewalls are at a lower temperature. However, it can be seen clearly in Fig. 5(d) that all the water vapor that evaporates from the base plate recondenses on the fin's sidewalls. A significant amount of vapor from the fin's top surface circulates and recondenses at the fin's sidewalls as well. This recondensation problem is still illustrated in the case of Fig. 5(e) and (f), which show the same profiles for the case in which the base plate does not evaporate any water. The hot tips form a high concentration vapor region at 100% RH that blocks off the sidewall's evaporation



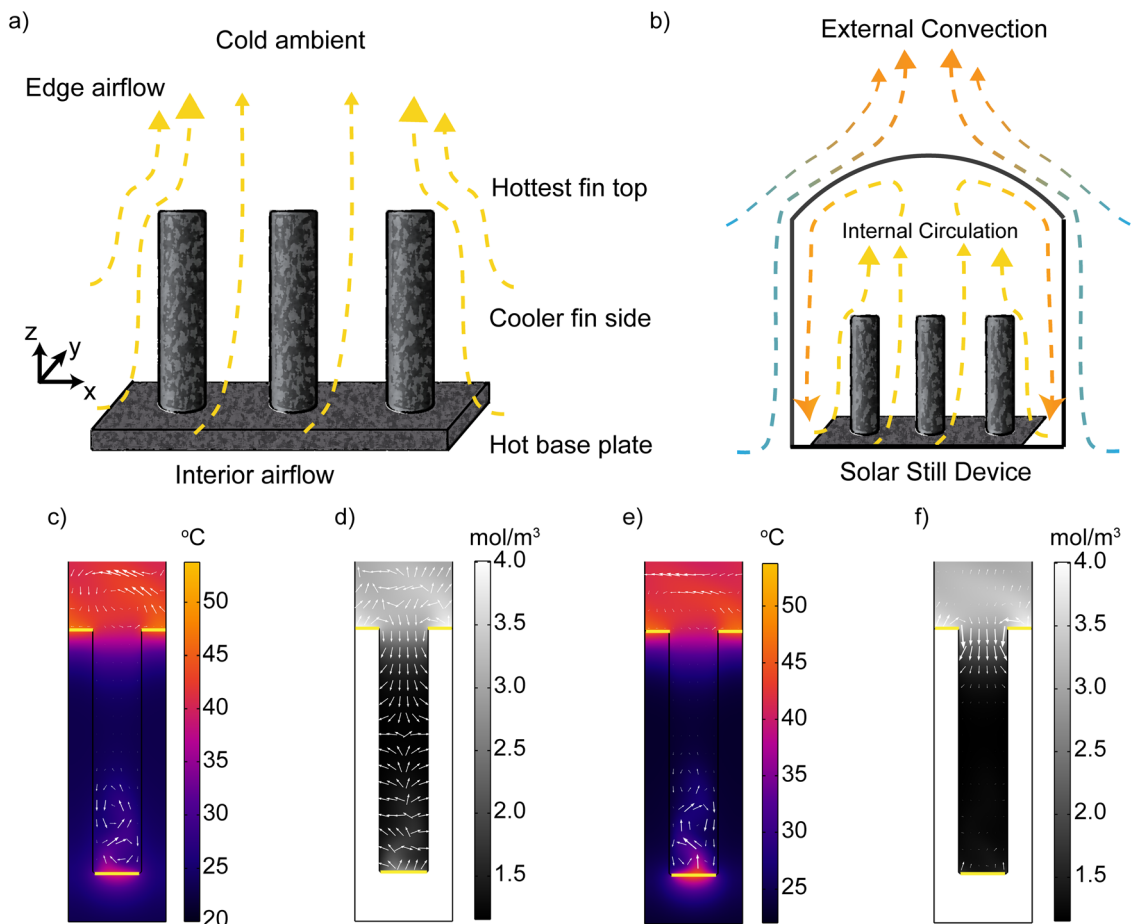


Fig. 5 3D fin performances under natural convective conditions. Hypothesized natural convection flow patterns in an (a) open and (b) closed environment such as a solar still device. Transient FEA simulated snapshots of 1D fin geometry for the case in which (c) and (d) the base evaporates water and (e) and (f) the base doesn't evaporate after 10 minutes of simulated time. (c) and (e) illustrate the natural convective flow patterns in white arrows and the temperature profiles. (d) and (f) illustrate the vapor molar density and the vapor diffusion gradients in white arrows. The outlined yellow regions illustrate the solar absorbing areas.

flow pathway, leading to the fin's sidewalls to only recondense water vapor. The low airflow velocities and incompatible device geometries with environmental heat absorption mechanisms lead to poor performances in scaled-up arrays under natural convection.

Regime map of 3D device performances

Summarizing the above discussions, we illustrate a regime map in Fig. 6 showing the required height to reach the solar-thermal limit, simulated performances of different forced convective conditions from Fig. 4(f), and the region in which conventional solar still devices operate in. Unlike in Fig. 2(b), an extra area ratio pre-factor A_c/A_{base} is included to account for the device absorbing more solar radiation than just the fin's cross-sectional area. This allows us to map the device performances back to the single fin regime map. The predicted device evaporation fluxes for some scenarios in Fig. 4(f) are mapped into Fig. 6, showing agreement with the solar-thermal limit curves corresponding to their respective convective heat transfer coefficients. The data corresponding to a bulk airspeed of 0.4 m s^{-1} was used because the convective heat transfer

coefficient is close to $15 \text{ W m}^{-2} \text{ K}^{-1}$, allowing easy comparisons with the calculations made in Fig. 1 and 2. The exact device performances will depend on the geographical and environmental conditions the device is placed in, as evident from the variation in the device performance due to the bulk airspeed as shown in Fig. 4(d) and (f). Furthermore, the predicted device evaporation fluxes are much lower than the single fin nominal evaporation flux performances predicted in Fig. 1(e), illustrating that the nominal evaporation flux is not an accurate performance metric in array devices. The total device length for the scaled-up array is also reproduced in the top x-axis to illustrate the scale-up potential of these devices relative to the chosen bulk airflow velocity. Since the longitudinal spacing between rows is 5 cm, the reported devices extend up to 2 meters in length. The figure shows that to achieve evaporation beyond the solar-thermal limit, the device length and the fin density can't be too high. For a given environment, there is an optimum fin spacing and size to maximize its performance.

The region in which most reported conventional solar still devices operate are shown by the gray box. Due to the device



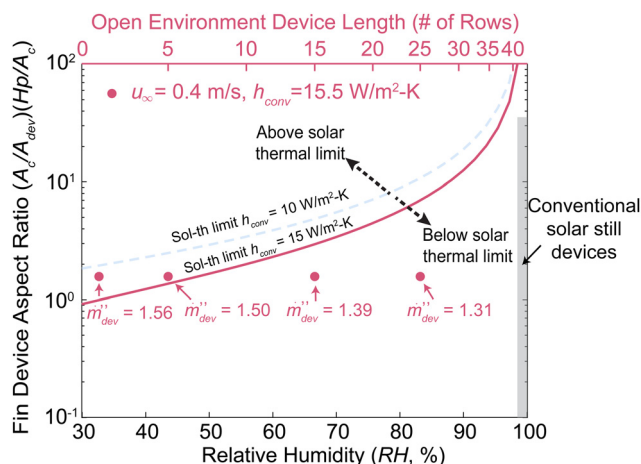


Fig. 6 Regime map of device performances. Lines represent single fin performances equal to the solar-thermal limit from Fig. 2(b). The extra area pre-factor in the y-axis accounts for the extra solar irradiation on the base of the device. Example forced convection device evaporation fluxes from Fig. 4(f) are reproduced to illustrate fin device aspect ratio's ability to predict super solar-thermal evaporation fluxes. The length of the devices corresponding to the average RH in the device with bulk airflow $u_{\infty} = 0.4 \text{ m s}^{-1}$ is reproduced as well on the top x-axis. The distance between each row is 5 cm. The operating regions of conventional solar still devices are outlined by the gray box. Note that the conventional solar still region is not related to the top axis.

utilizing natural convection with low induced airflow velocities, the heat transfer coefficients are unlikely to exceed $10 \text{ W m}^{-2} \text{ K}^{-1}$. The air inside of the device at steady state must be close to the saturated condition because of the working principle of the device. The regime map clearly illustrates that conventional solar still devices can't utilize environmental heat input due to the lack of net airflow through the device, the high device RH, and the low airflow velocities. The enhanced outdoor solar still performances reported previously are likely due to a combination of changing solar view factors and improper area normalizations.

Conclusions

3D solar interfacial-evaporation structures have been experimentally demonstrated to be able to evaporate water exceeding the solar-thermal limit due to environmental heat absorption. Although many lab-scale devices have been built and tested, there exist no criteria distinguishing 2D from 3D structures and very few attempts to test and analyze their scale-up potential for large scale deployment.

We have systematically modeled and explained the physics underlying the performance of 3D evaporating structures. First, the commonly reported nominal evaporation flux and the device evaporation fluxes are distinguished from each other due to the different area normalizations. A single fin model was constructed and was successfully able to reproduce the flat temperature profiles observed in experiments as well as the enhanced nominal evaporation fluxes. The model predicts that the nominal evaporation flux scales linearly with the fin height

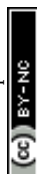
and that the environmental heat input critically relies on the humidity to be below the saturation point. Using the model, we have identified the non-dimensional critical heights in which a fin will first begin to absorb environmental heat and it absorbs enough environmental heat to nominally evaporate at the solar-thermal limit.

The single fin model was extended to a coarse-grained fin array to study the scale-up performance of these devices by coupling with the vapor and heat transport on the air side as well. Using this model, we have highlighted the potential and limitations in increased device performances by adding fin structures in forced convection situations. The model was able to illustrate the difference in performance between solar and dark evaporation, the environmental heat absorption, and the non-monotonic local air temperature profiles. The model pinpointed that the common procedure of using a foam insulating base plate is ill-advised because the base area can contribute a significant amount of evaporation flux due to enhanced kinetics from solar absorption. The model illustrates that the greatest gain in performance can occur if the external airflow velocity is high so that it can enhance the convective mass transfer coefficients and quickly replenish the humid air inside of the device with dry air. The device length can only be extended up to when the air becomes fully saturated with vapor. Afterwards, device performance will degrade significantly because hot vapor from the solar absorbing regions can recondense on the cooler fins' sidewalls.

We have identified that the enhanced benefits from 3D evaporating fins are unlikely to translate in large devices for natural convection open environment conditions due to the low airflow velocities. We have also illustrated that environmental heat absorption cannot be the mechanism for enhanced performance under natural convection closed environment conditions, such as in solar still devices, because the air internally is close to 100% RH. Instead, these small-scale devices have misleading improved performances due to the changing solar view factors on the fin and improper area normalization. Through FEA simulations, we have further clarified and illustrated the recondensation effect under natural convective conditions due to the hot vapor region that forms from the evaporating tip, blocking off the vapor transport pathway from the sidewalls.

We have compiled these findings into a regime map to illustrate the performances of single fins, example simulated studies of scaled-up forced convection devices in open environments, and the regime in which conventional solar stills operate in. An extra area ratio pre-factor allows us to approximately map back the overall device performance to the predicted performances of a single fin.

We conclude this paper by emphasizing the fundamental difference between 2D and 3D solar interfacial-evaporators with evaporation fluxes surpassing the solar-thermal limit. As shown in this paper, a 3D evaporator can absorb heat from the environment and achieving evaporation fluxes exceeding the solar-thermal limit.^{15,16} In this case, the evaporation is still thermally driven. No matter if the evaporators are 2D or 3D,



if nowhere in the system is below the ambient temperature, it is impossible to exceed solar-thermal limit based on purely thermal processes because all the heat used for evaporation must come from the solar energy. For 2D evaporators, it is difficult to achieve below ambient temperature unless there is a very high air flowrate.²⁰ Thus, most 2D evaporators tested under natural convection conditions with evaporation fluxes exceeding the solar-thermal limit imply a non-thermal evaporation process, such as *via* the photomolecular effect that we have discussed in several publications.^{10,22–24} It is important to point out that the solar absorbing top region of the fin could also evaporate through a non-thermal process as well. We hope this work, together with our investigations on 2D evaporators, will provide stimulus for future understanding of the mechanisms of solar interfacial evaporators exceeding the solar-thermal limit, which will enable better applications.

Methods

Single fin model

We make the following key assumptions in constructing an ideal performance model for a single fin: (1) we will ignore the detailed capillary flow of liquid brine in the 3D evaporator, (2) the external surface of the evaporating structure is always fully wetted, (3) the device will have a mechanism to reject salt accumulation during evaporation to prevent salt crystallization, and (4) the properties of the brine solution can be approximated using the properties of pure water. These assumptions are made to allow a system level modeling of the device while having results that are generalizable to different materials system. Assumption 1 holds true because the net mass flux of brine is small due to the low evaporation fluxes of 1 to 10 kg m^{−2} h^{−1}. Assumption 2 requires that the 3D solar evaporator has enough capillary pumping to continuously supply water without drying out and would depend ultimately on the material's pore structure, surface energy, and height of the fin. Assumption 3 is the least rigorous assumption due to the low diffusivities of ions and water relative to the evaporation flux and would place limits on the device geometry, leading to these results becoming a best-case scenario. Assumption 4 neglects the small change of the colligative properties such as vapor pressure depression due to the presence of salts. Thus, assumptions 2–4 govern the “ideal” performance of a 3D evaporator and the relaxation of each assumption will lead to performance degradation.

With such approximations, the differential energy balance equation can be written as the sum of conductive heat transfer along the fin and the convective, evaporative, and radiative heat exchange with the ambient reservoir.

$$0 = A_c k_f \frac{d^2 T}{dz^2} - p [h_{\text{conv}}(T - T_\infty) + \sigma \varepsilon (T^4 - T_\infty^4) + h_{\text{fg}} g_{\text{m,conv}} C_g (c_{\text{v,s}}(T) - \text{RH} c_{\text{v,s}}(T_\infty))] \quad (10)$$

where T is the local fin temperature, T_∞ is the ambient temperature, RH is the ambient relative humidity, z is the axis along the height of the fin, A_c is the cross-sectional area of the

extended fin, k_f is the wetted material's thermal conductivity, p is the perimeter, h_{conv} is the sidewall convective heat transfer coefficient, h_{fg} is the latent heat of evaporation, $g_{\text{m,conv}}$ is the sidewall convective mass transfer coefficient, $c_{\text{v,s}}$ is the saturated molar vapor fraction at a given temperature, σ is the Stefan–Boltzmann constant, and ε is the blackbody emissivity of the wetted fin. A full list of variables used is given in Table 1 above. The boundary conditions of this equation are two Robin style boundary conditions governing the heat fluxes. At the top interface, it will exchange heat and vapor with the ambient environment and absorb sunlight.

$$\frac{\dot{q}(z=H)}{A_c} = \dot{q}_{\text{sun}}'' - h_{\text{top}}(T - T_\infty) - \sigma \varepsilon (T^4 - T_\infty^4) - h_{\text{fg}} g_{\text{m,top}} C_g (c_{\text{v,s}}(T) - \text{RH} c_{\text{v,s}}(T_\infty)) \quad (11)$$

The subscript top denotes the convective coefficients on the top cross-section of the fin. The heat and mass transfer correlations are described using the heat and mass transfer analogies for crossflow on a cylinder (for sidewall) and laminar flow on a flat plate (for top surface). For crossflow, we will use the Churchill correlation.^{38,40} The corresponding Nusselt numbers Nu, and their relationships to the convective transfer coefficients are

$$\text{Nu}_{\text{conv}} = 0.3 + \frac{0.62 \text{Re}_D^{1/2} \text{Pr}^{1/3}}{(1 + (0.4/\text{Pr})^{2/3})^{1/4}} \left(1 + \left(\frac{\text{Re}_D}{282000} \right)^{5/8} \right)^{4/5} \quad (12)$$

$$\text{Nu}_{\text{top}} = 0.664 \text{Re}_D^{1/2} \text{Pr}^{1/3} \quad (13)$$

$$h = \frac{\text{Nu} k_{\text{air}}}{D} \quad (14)$$

where Pr is the Prandtl number of air, Re is the Reynolds number, and k_{air} is the thermal conductivity of air. We used the same correlations to describe the mass transfer of vapor in air. The mass transfer coefficient can be found analogously using eqn (12)–(14) by replacing h with g_{m} , Nu with the Sherwood number Sh, Pr with the Schmidt number Sc, and k_{air} with the diffusion coefficient of water vapor in excess of air D_v . The bottom heat transfer coefficient, h_{bot} , governs the convective heat transfer between the brine pond reservoir and the evaporating fin's bottom interface. The true value should depend on the temperature differences and the convective flow patterns inside the reservoir. For simplicity, we will set h_{bot} to 100 W m^{−2} K^{−1} and the brine pond temperature T_{bot} to the same as the ambient T_∞ at 23 °C. To account for the foam inserts used in the base in laboratory tests or the base plate thickness in outdoor devices, we will assume that there is an additional t_{base} of 2 cm of wetted material not exposed to the ambient environment. The bottom Robin boundary condition is thus

$$\frac{\dot{q}(z=0)}{A_c} = \frac{(T_{\text{bot}} - T)}{1/h_{\text{bot}} + t_{\text{base}}/k_f} \quad (15)$$



A system of non-linear equations is formed that relates the heat fluxes at each local cross-section of the fin at a given iteration n .

$$\bar{f}_n = \bar{A}\bar{T}_n + \bar{b}_n + \bar{c} \quad (16)$$

where $\bar{f}_n$ is the function vector that becomes zero at the equilibrium temperature profile, the $\bar{A}$ matrix has constant terms that couple linearly with the temperature vector $\bar{T}_n$, the $\bar{b}_n$ vector has terms that are non-linearly dependent on the temperature, and the $\bar{c}$ vector holds constant terms. The Jacobian can then be calculated by taking the multivariable derivative of each term in eqn (16).

$$\bar{J}_n = \bar{A} + \frac{\partial \bar{b}_n}{\partial \bar{T}} \quad (17)$$

The newly guessed temperature profile can then be calculated using the Jacobian.

$$\bar{T}_n^* = \bar{T}_n - \bar{J}_n^{-1} \bar{f}_n \quad (18)$$

The new temperature profile is then iterated by mixing part of the newly guessed temperature $\bar{T}_n^*$ and the old temperature $\bar{T}_n$ using a mixing term λ .

$$\bar{T}_{n+1} = \lambda \bar{T}_n^* + (1 - \lambda) \bar{T}_n \quad (19)$$

Coarse-grained scaled-up fin model

A coarse-grained model is developed using the single fin results to build the scaled-up model for fin arrays. Further simplifications are imposed to make the model computationally tractable. The first assumption is that due to the complex view factors and temperature profiles internal to the pin fin array, we will neglect radiation heat exchange. This effect will cause the temperature difference between the hot and cold evaporating surfaces to be larger, however we expect this effect to become small as the external airspeed and convective heat transfer coefficients increase. The second assumption is that there is a clear separation between air flow that is “internal” (flowing through the pin fin array) and a free stream “external” air flow on the tips of the fin. This separation will clearly define the boundary condition for the fins’ tips and the internal flow that will progressively become more humid. This simplification will likely over-predict the evaporation flux from the top cross-sectional area because this region will couple with the internal flow and become more humid locally as well. The third assumption is that due to the various pins dispersed throughout the array, it will induce local mixing and allow us to describe the heat and mass transfer using the bulk-averaged properties of the airflow at each control volume. The fourth assumption is that the air always flows in the same direction and that the array is large enough in the transverse direction (x -axis) so that there are minimal edge effects. The final assumption is that the temperature of the base plate in each control volume can be estimated independently of its neighbors using a 1D heat and mass transfer resistance network of a

flat plate correlation in which there are no pin fin array. This assumption becomes more valid as the spacing between each fin becomes large relative to the fin’s characteristic size and the base plate thickness.

The model works by doing a control volume analysis of air flowing downstream along the device and coarse graining the total heat and mass transfer exchange between it and the local row. At the inlet of the i th-row pin fin array, we denote the local air temperature T_i and vapor mole fraction $c_{v,i}$. The total molar flowrate of “internal” air is $\dot{n}_{\text{air}} = C_g u_{\infty} S_t H$. Žukauskas correlations for tube bundles are used to describe heat transfer between the fins and the air.^{38,41,42}

$$\text{Nu}_{\text{conv}} = \text{Pr}^{0.36} \text{fn}(\text{Re}_{D,u_{\text{max}}}) \quad (20)$$

$$\text{fn}(\text{Re}_{D,u_{\text{max}}}) = \begin{cases} 0.71 \text{Re}_{D,u_{\text{max}}}^{0.5}, & \text{Re}_{D,u_{\text{max}}} < 1180 \\ 0.35 \text{Re}_{D,u_{\text{max}}}^{0.6}, & \text{Re}_{D,u_{\text{max}}} \geq 1180 \end{cases} \quad (21)$$

where u_{max} is³⁸

$$u_{\text{max}} = u_{\infty} \frac{S_t}{S_t - D} \quad (22)$$

due to the venturi effect. The corresponding Sherwood numbers and mass transfer coefficients are found using the heat and mass transfer analogy again. To describe the heat (h_{base}) and mass transfer ($g_{\text{m,base}}$) coefficients between the base plate at temperature $T_{\text{base},i}$ and the air, the flat plate correlations in eqn (13) is used with a characteristic size of S_t . The base temperature and evaporation fluxes are then calculated by using a 1D heat and mass transfer resistance network at the evaporating surface.

$$0 = \dot{q}_{\text{sun}}'' - h_{\text{fg}} C_g g_{\text{m,base}} (c_{v,s}(T_{\text{base},i}) - c_{v,i}) - h_{\text{base}} (T_{\text{base},i} - T_i) - \frac{(T_{\text{base},i} - T_{\text{bot}})}{1/h_{\text{bot}} + t_{\text{base}}/k_f} \quad (23)$$

As the airflows over each row, the air’s temperature and vapor content will change based on the evaporation and heat exchange.

$$c_{v,i+1} = c_{v,i} + \frac{\rho g_{\text{m,conv}} C_g \int_0^H (c_{v,s}(T) - c_{v,i}) dz}{\dot{n}_{\text{air}}} + \frac{(S_t S_l - A_c) g_{\text{m,base}} C_g (c_{v,s}(T_{\text{base},i}) - c_{v,i})}{\dot{n}_{\text{air}}} \quad (24)$$

$$T_{\text{air},i+1} = T_{\text{air},i} + \frac{\rho h_{\text{conv}} \int_0^H (T - T_i) dz}{\dot{n}_{\text{air}} c_p} + \frac{(S_t S_l - A_c) h_{\text{base}} (T_{\text{base},i} - T_i)}{\dot{n}_{\text{air}} c_p} \quad (25)$$

where T is the temperature along the fin. Heat conduction and vapor diffusion along the air profile is neglected due to the high Peclet numbers involved in the study. The profiles are then solved by forward marching in space along the array length using the updated humidity and temperatures of the air.



FEA model

We have modeled natural evaporation from the 1D fin array using transient simulations in a 2D geometry. Due to the large-scale nature of these devices, we have modeled the behavior of the interior fins, which should dominate the overall device behavior. Periodic boundary conditions are imposed in the perpendicular directions for heat transport, vapor transport, and momentum equations. The top of the domain is set to be 10 times taller than the fin and set as an open boundary condition. Air is fully simulated around the fin structure. The general governing equations for mass, momentum, energy, and vapor transport in the air are

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \vec{u}) = 0 \quad (26)$$

$$\rho \frac{\partial \vec{u}}{\partial t} + \rho (\vec{u} \cdot \nabla) \vec{u} = -\nabla P + \nabla \cdot \left(\mu \left(\nabla \vec{u} + (\nabla \vec{u})^T \right) - \frac{2}{3} \mu (\nabla \cdot \vec{u}) I \right) + \rho \vec{g} \quad (27)$$

$$\rho c_p \left(\frac{\partial T}{\partial t} + \vec{u} \cdot \nabla T \right) = \nabla \cdot (k \nabla T) \quad (28)$$

$$\frac{\partial c_v}{\partial t} + \nabla \cdot (c_v \vec{u}) = \nabla \cdot (D_v \nabla c_v) \quad (29)$$

where ρ is the material density, t is the time, μ is the dynamic viscosity, P is the gauge pressure, $\vec{g}$ is the gravity vector, and $\vec{u}$ is the velocity vector. Using the weakly compressible mode, air's density is coupled with its temperature to induce natural convective flows. Only heat conduction is modeled inside the array structure due to the low flowrates of the liquid involved. The material's thermophysical properties are set to a thermal conductivity of $0.3 \text{ W m}^{-1} \text{ K}^{-1}$, a specific heat capacity of $2000 \text{ J kg}^{-1} \text{ K}^{-1}$, and a density of 1200 kg m^{-3} .²⁰ The fin was set to 10 cm tall, the base plate set to 2 cm in thickness, the width of the fin set to 2 cm, and the gap between the fins is set to 2 cm. The bottom of the baseplate is set to a Robin heat boundary condition through a heat transfer coefficient of $100 \text{ W m}^{-2} \text{ K}^{-1}$ interacting with a reservoir temperature of 23°C . The air surrounding the fin is initially set to 23°C and 30% RH. The top of the fins and the top of the base plate absorbs 1000 W m^{-2} to simulate solar absorption with a zenith angle of 0° . At the fin and air interface, a boundary evaporative cooling term is included that couples the vapor transport solved from eqn (29) with the heat equation in eqn (28) using the latent heat of evaporation. The simulation is then forward marched in time to produce the simulation snapshots.

Author contributions

J. H. Z. and G. C. developed the concept. J. H. Z. prepared the models. J. H. Z., R. M., A. O., and G. C. wrote the paper. G. C. directed the overall research.

Data availability

The modeling methods and formula have been fully included in the Methods section.

Conflicts of interest

There are no conflicts to declare.

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