I. INTRODUCTION

The standard atomistic approach to tribology, i.e. the study on friction dissipation and wear, usually focuses on indestructible “crushproof” particles, typically atoms or molecules [1,9]. Recent works [10,13] have brought colloids into the realm of tribology, by letting repulsive hard-core particles interact with a periodic “correlation potential”, generated by means of optical forces, and driven by viscous drag. The conditions of that experiment match the single indestructible particle paradigm. Colloids and, more generally, soft matter systems would however allow one to investigate situations where complex objects, carrying an internal structure, under the stress produced by external interactions and driving, can alter their internal structure both in shape and even in the number of the component sub-particles. Thus one can expect the presence of a varied set of regimes of response of the system to the external perturbation. In the present work, we address precisely this last situation by means of molecular-dynamics (MD) simulations of a system forming micro-phases. Micro phase formation can take place under a number of conditions on the interparticle interactions and on the state variables in three dimensions [14,19] as well as in two [20,23]. The microphases can be disordered [15,18,21,22] as well as ordered in a crystalline-like state or other patterned structures like stripes and lamellae [17,21,25] or even as an ordered bicontinuous state [26] in which both components of a two-component system span the space. Examples of micro-phase forming systems are block-polymers [24,25] or hard colloids with competing interactions, e.g. short range attraction-long range repulsion like for some colloid-polymer mixtures [15,19,18,20,22]. Certain soft-matter systems like star polymers or dendrimers in a good solvent can interpenetrate each other to a large extent so that the effective interaction between the centers of mass of two such entities has a soft-core character. Also such systems can form microphases in which the colloids spontaneously aggregate into clusters, and the clusters are organized in a crystalline state [27,28]. Rheological properties of cluster fluid phase [29] and of cluster crystal phase [30] have been investigated but no study has yet addressed the tribology of such systems.

In the present work, we simulate a system of interacting colloidal particles, whose mutual interaction potential generates the spontaneous formation of clusters. In particular, we concentrate on a 2D geometry with repulsive interactions, like in many experiments carried out with colloids [10,21,35], but the concepts introduced here may be relevant for vortexes in superconductors [36] too. The novelty compared to these experiments is the adoption of an interparticle interaction which supports the spontaneous formation of cluster phases. Cluster dynamics has been investigated in the past [37,39]: in these works clustering was forced by an external potential, while in the present work clustering is the intrinsic result of the particle-particle interaction, and is retained even when any external potentials are turned off.

Section [11] introduces and motivates the model. As a first step, in Sec. [11] we characterize the $T = 0$ equilibrium states of the model free from external interactions. We study the ground-state energy of several regular arrangements of clusters formed by $n$ particles (with $n$ from 1 to 6, that we will refer to as $n$-clusters) as a function of the mean colloids density, obtaining a $T = 0$ phase diagram. Then Sect. [IV] reports the investigation of the dynamics of the resulting arrays of clusters interacting with a lattice-matched external periodic potential and an homogeneous pulling force. Specifically, we report the mobility curves for these cluster systems, as a function of the intensity of the pulling force, for a few values of the amplitude of the corrugation potential. We characterize in detail the different sliding regimes, with entire clusters advancing, or only parts of them. The main outcome of this research is that the clusters internal dynamics is indeed affecting significantly the depinning force and the overall mobility.
II. THE MODEL

A. The Interparticle Interaction

Spontaneous cluster formation can be the result of competitive interactions $[15, 16, 18, 20–22]$, but clusters can also form due to soft potentials $[23, 27, 28]$. Here we adopt an interaction of the latter type. We consider the following form for the pairwise interaction potential energy:

$$\phi_{cc}(r) = \frac{A_0}{r^6} + \frac{B_0}{r^3}.$$  \hspace{1cm} (1)

This interaction, depicted in Fig. 1, consists of two terms: a soft-core repulsive interaction, and a weak hard-core repulsion. The $B_0 = 0$ (purely soft-core) model has been fully characterized in two dimensions in Ref. $[23]$: its phase diagram consists in a high-temperature low-density regular fluid phase, and a low-temperature high-density cluster phase. The reason for the cluster phase is that when the inter-particle distance decreases under $r \lesssim R_c$, the potential energy $\phi_{cc}(r)$ flattens out to a nearly constant energy, with the result of producing a quite small repulsive force. As a result, in one dimension a particle in between two other particles kept at fixed positions, when the distance between these fixed particles is not too large, may be better off energetically by coming closer to either of them, rather than remaining at the middle point. This suggests the mechanism whereby, in any dimensionality and at large enough density, a spontaneous symmetry breaking leading to clustering can be energetically favorable against a uniform fluid phase. A detailed mathematical criterion has been developed for the formation of cluster phases in the case of soft-core potentials in terms of their Fourier transform becoming negative in a range of finite wave vector $[40]$. At high $T$, entropic effects and fluctuations tend to favor the uniform fluid phase against cluster formation. In the opposite $T \to 0$ extreme, for $B_0 = 0$ no mechanism keeps the colloids apart, and so the clustering tendency would reach the extreme limit of collapsed point-like clusters with all colloids of a cluster sitting exactly at the same position. To prevent this singular behavior we add the hard-core $B_0$ term of Eq. (1) for whose coefficient we adopt a relatively small value $B_0 = 5 \times 10^{-5} A_0$ $[41]$ in order to perturb the phase diagram as little as possible, at least in the not-too-high-density region that we are interested in.

We take the repulsive-potential characteristic distance $R_c$, and characteristic energy $A_0/R_c^6$, and the mass $m$ of the colloidal particles as fundamental units. In the following we express all physical quantities in terms of suitable model units, as listed in Table I.

B. The Equations of Motion

In simulations, we let the entire system evolve following the standard Langevin dynamics in two dimensions $[42]$, under the effect of an external periodic corrugation potential $U_{ext}(r)$ and a constant driving force $F$ applied to each particle. The equation of motion for the $j^{th}$ particle is:

$$m \ddot{\mathbf{r}}_j = F \dot{\mathbf{r}}_j - \mathbf{R}_j + \sum_{k \neq j} N \phi_{cc}(|\mathbf{r}_k - \mathbf{r}_j|) + U_{ext}(\mathbf{r}_j) + \xi_j.$$  \hspace{1cm} (2)

Here $\eta$ is the coefficient of viscous friction associated to the fluid where the colloids move; we generate an overdamped dynamics by adopting a large $\eta = 28 \eta_0$. The $\xi_j$ terms are Gaussian-distributed random forces with amplitude $\sigma = \sqrt{2m\eta k_B T/\Delta t}$, where $\Delta t$ is the simulation time step: they simulate the collisions of the molecules in the fluid with the colloids, thus generating...
the appropriate Brownian dynamics. Together, the viscous and the random-force terms in Eq. (2) represent a standard Langevin thermostat \[10\]. In simulations, the inter-particle potential \( \phi_{cc} \) is cut off smoothly (vanishing potential and potential derivative) at a distance \( R_{\text{cutoff}} = 5 R_c \) \[11\].

The periodic external potential \( U_{\text{ext}}(r) \) simulates the effects of friction against a crystalline surface. In the laboratory, this interaction has been realized by means of a modulated light field constructed by laser interference, so that its spacing and intensity that can be tuned with considerable freedom \[10\]. In our simulations, we assume a corrugation potential of the form:

\[
U_{\text{ext}}(r) = V_0 W(r) .
\]

Here \( V_0 \) is the amplitude of the spatial modulation and \( W(r) \) is a 2D periodic function with hexagonal symmetry, period \( a_{\text{pot}} \), and unit amplitude. Its explicit form is:

\[
W(r) = -\frac{2}{9} \left[ \frac{3}{2} + 2 \cos \left( \frac{2\pi r_x}{a_{\text{pot}}} \right) \cos \left( \frac{2\pi r_y}{\sqrt{3}a_{\text{pot}}} \right) \right] + \cos \left( \frac{4\pi r_y}{\sqrt{3}a_{\text{pot}}} \right) .
\]

This function is illustrated in Fig. 2.

Following the \( r_y = 0 \) path in the \( x \)-direction a particle goes through alternating minima and saddle points of \( W(r) \). This is the energetically less costly trajectory that a colloid can follow when pushed by an \( x \)-directed force. Along this path, the external potential is a simple sinusoidal oscillation

\[
U_{\text{ext}}(r_x, 0) = -V_0 \left[ \frac{5}{9} + \frac{4}{9} \cos \left( \frac{2\pi r_x}{a_{\text{pot}}} \right) \right] .
\]

The resulting energy barrier separating the saddle points from the minima is therefore \( \frac{2}{9} V_0 \). In this situation the static friction of an isolated particle at \( T = 0 \), i.e. the minimum force needed to push that particle across the potential corrugation when inertial effects are negligible, equals the largest value of \( \frac{\partial}{\partial r_x} U_{\text{ext}}(r_x, 0) \), namely:

\[
F_{1s} = \frac{8\pi V_0}{9a_{\text{pot}}} .
\]

In the simulations and in the relative graphs we express all applied forces by comparison to the static friction \( F_{1s} \) of the single colloid, expressed in Eq. (6). For the lattice spacing we take \( a_{\text{pot}} = 1.5 R_c \) and select values of the particles density that favor arrays of clusters whose spacing matches precisely this separation, as discussed in Sect. \[11\].

C. The Cell and the Initial Condition

We simulate the 2D model in a parallelogram-shaped supercell with periodic boundary conditions (PBC), thus fixing the average density. To respect the hexagonal symmetry that the arrays repulsive colloids (or clusters thereof) tend to adopt, we use a supercell generated by two primitive vectors of equal length \( L \) and forming an angle of 60° with each other. For \( V_0 = 0 \) we are free to adopt an arbitrary cell size: in the static simulations of Sect. \[11\] we will vary \( L \) freely in order to evaluate the energetics as a function of the colloid density, and therefore determine the \( T = 0 \) phase diagram of the free model.

For \( V_0 \neq 0 \), this supercell needs to match the symmetry of the corrugation potential: we must stick to \( L \) given by an integer multiple of \( a_{\text{pot}} \). We settle for a compromise size \( L = 12 a_{\text{pot}} \), small enough to guarantee simulations involving a not-too-large number of particles (and therefore manageable simulation times), but large enough for a fair averaging over fluctuations and allowing for independent and possibly asynchronous movements of individual clusters. Accordingly, the cell contains 144 corrugation minima in an area \( A = \sqrt{3}L^2/2 \).

As discussed below, we find that a cluster-cluster spacing 1.5\( R_c \) as optimal for favoring a well-defined n-cluster phase. In the present work we focus on a lattice-matched condition between the cluster-cluster spacing and the spacing \( a_{\text{pot}} \) between adjacent minima of the corrugation. Therefore in all dynamical simulations of Sect. \[14\] we settle for fixing \( a_{\text{pot}} = 1.5 R_c \), and therefore a supercell side \( L = 18 R_c \), thus generating a supercell area \( A \approx 280.6 R_c^2 \). In this supercell, the potential has 144 wells, each initially hosting a single particle or a \( n = 2 \) to 6-particles cluster. We simulate a total \( N = 144 n \) particles, which thus sample the following discrete values for the average density:

\[
\rho = \frac{N}{A} = n \frac{2}{\sqrt{3}a_{\text{pot}}^2} = n \frac{8}{9\sqrt{3} R_c^2} .
\]

For the initial condition, we start off with distances \( \delta = 0.32 R_c \) between the colloids within each cluster suggested by the inflection of the inter-particle potential,
III. GROUND-STATE ENERGETICS FOR THE FREE MODEL

While at finite temperature the number of colloids in each cluster fluctuates, at $T = 0$, for every given density $\rho$, one expects that an hexagonal lattice of clusters with the same number $n$ of particles represents a local minimum of the total repulsive energy. We explore the density range from $\rho = 0.4 R_c^{-2}$ to $4 R_c^{-2}$. In the absence of corrugation potential ($V_0 = 0$) and driving force ($F = 0$), for every $n$-cluster and every density (i.e. every lattice spacing of the cluster crystal), we execute a full relaxation by means of a damped dynamics, for at least a total time $2000 \tau_0$, but extending the relaxation for longer time until the residual kinetic energy is safely below $10^{-8} E_0$. This method leads us very close to the precise ground state. Depending on $n$ and $\rho$, the system may remain trapped in a local minimum characterized by some residual energy associated to the cluster’s orientational order, see Fig. 3. Arrows identify the densities adopted for the driven dynamics studied in Sect. IV.

By running several energy minimizations starting from different cluster phases, we can follow the evolution of the ground-state energy as a function of the density $\rho$. For each configuration we evaluate the total colloid-colloid repulsive energy per particle, divide it by the number of colloids in the supercell, and obtain the energy per particle. Figure 4 displays the curves of the total energy per particle for each $n = 1$ to 6, as a function of $\rho$. We see that at each given density, several cluster configurations represent competitive local energy minima. However, at each density one of them is usually lowest in energy. Crossings between successive curves identify the coexistence density values where the $n$-cluster and the $(n + 1)$-cluster have the same stability, and can therefore coexist. By carrying out a finer numerical analysis and a Maxwell construction one could also identify density ranges characterized by the phase coexistence of homogeneous domains consisting of clusters of different $n$. We need not go into this detail: based on the obtained rough phase diagram, we identify the densities corresponding to stable $n$-cluster states, characterized by a cluster-cluster spacing $1.5 R_c$, and we adopt them for the driven simulations of Sect. IV.
Figure 5. Small portions of the $T=0$ configurations in a $V_0=1.5 E_0$ corrugation, under the action of a $F=0.6 F_{1s} < F_{dep}$ pulling force, i.e. these are static configurations prior to depinning, for: (a): $n=2$; (b): $n=3$; (c): $n=4$; (d): $n=5$ clusters.

IV. DRIVEN DYNAMICS

Starting from the equilibrated state obtained as discussed in Sect. III, we investigate the dynamics under the competing effects of the external periodic corrugation potential, which tends to immobilize the clusters at its minima, and the lateral driving force $F$ which, if there was no corrugation, would tend to establish a sliding state at speed $v = F/\eta$. As expected of a commensurate, and even lattice-matched configuration, a static pinning threshold is always present. When $F$ is slowly raised from 0 up to this pinning threshold, after an initial transient allowing for the cluster rearrangement, the steady sliding speed remains null. However, nontrivial rearrangements of the clusters are observed well in advance of depinning: the driving force clears all orientational orderings characteristic of the ground states. The clusters are forced to specific orientations, which are displayed in Fig. 5. These ferro-orientational $[43, 44]$ arrangements are made energetically advantageous by the anisotropic energy landscape present at the regions in between the minimum and the saddle point of the corrugation potential of Fig. 2 where $F$ pushes the clusters. Precisely this force-induced ferro-orientational cluster arrangement observed for all $n$ prior to depinning $[45]$ makes the orientational details of the free-model ground state (e.g. Fig. 3) irrelevant for the dynamics we address in the present work.

The most straightforward dynamical indicator for a driven model of this kind is the average mobility, i.e. the ratio of the average $\hat{x}$-directed velocity to the driving force, which we report for different numbers $n$ of colloids per cluster. If the corrugation potential was removed ($V_0 = 0$), then for any nonzero force, the mobility would equal the free mobility $\eta^{-1}$, which equals $0.0357 \eta_0$ with the present choice of the damping parameter. The addition of the corrugation leads to smaller values of mobility.

To evaluate the mobility, for each $2 \leq n \leq 5$ we execute three sequences of simulations, with the following amplitudes of the periodic external potential: $V_0 = E_0$, $V_0 = 1.5 E_0$, and $V_0 = 2 E_0$. The $T=0$ mobility of the $n=2$-cluster lattice as a function of the driving force $F$ expressed as a fraction of the depinning force $F_{1s} = 8\pi V_0/(9\eta_{pot})$ for an isolated particle, for corrugation amplitudes $V_0/E_0 = 1$, $1.5$, and $2$. The numerical error affecting the mobility due to averaging over noninteger numbers of periods is less than 0.3% of the free mobility $\eta^{-1}$. The labels along the curve identify different sliding regimes, with groups of $m/n$ particles leaving one well of $U_{ext}(r)$, and moving to the next well at the right. Arrows identify transitions from a sliding regime to the one occurring at larger force. Synchro/async regimes refer to whether all well-to-well particle jumps occur simultaneously across the cell or they do not. In this specific $n=2$ case, even without specification, all motions are “1D”, meaning that colloids advance along horizontal lines through the minima of $W(r)$.

Figure 6. The $T=0$ mobility of the $n=2$-cluster lattice as a function of the driving force $F$ expressed as a fraction of the depinning force $F_{1s} = 8\pi V_0/(9\eta_{pot})$ for an isolated particle, for corrugation amplitudes $V_0/E_0 = 1$, $1.5$, and $2$. The numerical error affecting the mobility due to averaging over noninteger numbers of periods is less than 0.3% of the free mobility $\eta^{-1}$. The labels along the curve identify different sliding regimes, with groups of $m/n$ particles leaving one well of $U_{ext}(r)$, and moving to the next well at the right. Arrows identify transitions from a sliding regime to the one occurring at larger force. Synchro/async regimes refer to whether all well-to-well particle jumps occur simultaneously across the cell or they do not. In this specific $n=2$ case, even without specification, all motions are “1D”, meaning that colloids advance along horizontal lines through the minima of $W(r)$.
Figure 7. Same as Fig. 6 but for the $n = 3$ cluster lattice.

Figure 8. Same as Fig. 6 but for the $n = 4$ cluster lattice.

Figure 9. Same as Fig. 6 but for the $n = 5$ cluster lattice. All observed dynamical modes are synchronized here. See text for a description of the very fast hop and 4/5 sync altern regimes.

Figure 10. The static friction, or depinning threshold, $F_{\text{dep}}$, expressed as a fraction of the single-colloid depinning force $F_{1s}$ (also relevant for the depinning of the $n = 1$ no-cluster phase), for $V_0/E_0 = 1, 1.5$, and $2$, as a function of the number $n$ of colloids per cluster.

For each sequence, we increase the applied lateral force in a range proportional to $V_0$, namely $0.55 F_{1s} \leq F \leq 1.3 F_{1s}$ in small steps $\Delta F = 0.015 F_{1s}$. At each step, the average velocity is evaluated by averaging over a total time $1400 t_0$ after an initial transient time $600 t_0$. Figures 9 and 10 report the resulting mobilities of the cluster lattices for the investigated numbers $n$ of colloids per cluster as a function of the applied driving force $F$. In the initial small-$F$ simulations the driving force is insufficient to extract the colloid clusters from the potential wells of $U_{\text{ext}}$: thus a null mobility is obtained. Beyond a certain static friction threshold $F_{\text{dep}}$, the colloids unpin and start to advance, resulting in a finite mobility, which grows progressively approaching the free-particle value $\eta^{-1}$. The small “adiabatic” increase in $F$ allows us to evaluate $F_{\text{dep}}$ with a relatively small uncertainty, and to identify the features of the particle motion as mobility increases after depinning.

Consider the depinning force $F_{\text{dep}}$. Although its values can be read directly from Figs. 9 and 10 for ease of comparison Fig. 10 reports them as a function of $n$. A first observation is that the clustering of the colloids causes a strong decrease of $F_{\text{dep}}$ compared to the value $F_{\text{dep}} = F_{1s}$ of a single colloid, by about $20\%$ for $n = 2$ and $3$ up to $33\%$ for $n = 5$. The static friction force peaks for $n = 3$. Above $n = 3$ the depinning force tends to decrease as $n$ increases, as expected due to the growing size of the clusters.

The $n = 2$ clusters mark an exception, as they depin at smaller force than $n = 3$. Also, in the comparison of the depinning force for three values of $V_0/E_0$, we see that depinning for $n = 2$ occurs at a smaller $F_{\text{dep}}/F_{1s}$ for $V_0 = E_0$ than $V_0 = 2E_0$, contrary to all other cluster sizes. The reason for the special behavior of the $n = 2$
Figure 11. The whole-cluster (5/5) advancement observed in the $n = 5, V_0 = E_0$ simulation driven at $F/F_1 = 1$, see Fig. [9]. Panels (a) - (d) represent subsequent snapshots, separated by $\Delta t = 2 t_0$. A particle is labeled with a vertical bar for ease of identification in the successive snapshots.

Figure 12. Successive snapshots illustrating the 2/5 dynamics observed in the $n = 5, V_0 = 2 E_0$ simulation driven at $F/F_1 = 0.7$, see Fig. [9]. Five particles are labeled for ease of identification in the successive snapshots. The time interval $\Delta t = 2 t_0$, the same as in Fig. [11].

cluster lays in its elongated conformation under traction, see Fig. [5]. As a result, in the pulling direction $\hat{x}$ the $n = 2$ cluster extends over a longer distance $\approx \delta$ than the size $\approx \delta \sqrt{3}/2$ of $n = 3$. For this reason, when the cluster center of mass approaches the inflection point of the corrugation potential (the point beyond which the retaining force begins to taper off), the rightmost particle has moved forward more for $n = 2$ than for $n = 3$.

After depinning, the mobility, on top of an overall smooth increase as a function of $F$, exhibits sharp changes in slope, Figs. [6] - [9]. These changes mark transitions between different sliding patterns, which can be examined by monitoring the advancement of the individual colloids. The following section describes precisely these sliding regimes.

A. Sliding regimes

Under driving, the internal structure of the clusters makes them advance either as a single object, or less trivially with structural decompositions and recompositions. Depending on the corrugation amplitude $V_0$ and on the number of colloid per cluster we observe (or not) several of these decomposition-recomposition phenomena as the lateral force is increased. For convenience, we indicate the observed dynamical patterns on the mobility curves of Figs. [6] - [9] where the switching between patterns can induce visible mobility anomalies. We use $k/n$ labels indicating dynamical patterns which persist for an entire force interval. The $k/n$ labels in the mobility curves indicate that the advancement from one well to the next occurs with $k$ particles out of a cluster of $n$ jumping ahead, joining the next cluster, and leaving $n - k$ particles behind. Arrows point at transitions from one kind of dynamical behavior to the next.

As a general rule, a small corrugation amplitude $V_0$ tends to favor the clusters advancing as a whole, namely in “$n/n$” modes. In contrast, for larger amplitude $V_0$ the corrugation competes favorably with the energy advantage of the clusters to remain entire, and as a result for larger $V_0$ we do observe nontrivial decomposition-recomposition patterns. These decompositions can occur synchronously across the entire supercell (in this case they are labeled synchro) or asynchronously (async label). Figures [11] and [12] report successive snapshots illustrating the two main kinds of dynamical patterns, as marked on the mobility plots. Specifically, Fig. [11] illustrates the most basic 5/5 advancement mode: whole cluster advancement from one corrugation well to the next, as observed for weak corrugation $V_0 = E_0$. In contrast, Fig. [12] illustrates an advancement mode involving synchronous decompositions and recompositions of the $n = 5$ clusters, with 2 particles abandoning each cluster and jumping to the next corrugation well while leaving 3 particles behind at each step. Examples of similar patterned movements involving different numbers of particles are reported in the Supplementary Material [46], as digital videos. Figure [13] illustrates the effect of these patterned movements on the $x$-coordinate of the center of mass of the colloids. As the particle hoppings are synchronized across the entire sample, the center-of-mass position $x_{cm}$ exhibits relatively fast jumps during the synchronized
in asynchronous dynamics as a consequence of averaging no (or very small) steps are observed in the overall center-of-mass coordinate. The asynchronous jumps can occur in a regular sequence at different rows or with apparently chaotic hoppings as for example in the video c14.2.71V2.avi provided as Supplementary Material [46]. Note that any regularity in asynchronous motions (and possibly the overall synchronization itself when present) is favored by the specific supercell periodic boundary conditions adopted in simulation: as such, the synchronized or regular patterns may be considered artifacts of the simulated model.

This zero-temperature dynamics, often dominated by unique regular or chaotic attractors [17, 48] does not represent a realistic condition that one could put in relation to actual experiments. Indeed, in the more realistic finite-temperature simulations of Sect. [49] below, irregular asynchronous depinning occurs in most cases, due to the Brownian forces randomly anticipating or retarding the advancement of individual particles or clusters.

By cycling the driving force \( F \) up and down, we investigated whether these dynamical systems retain some kind of dynamics memory, i.e. the sliding state is affected not only by the competition of the applied confinement potential and driving, but also by the initial sliding configuration. Indeed in these \( T = 0 \) simulation we do find some small difference of the boundaries between specific dynamical modes between the decreasing-\( F \) path and the increasing-\( F \) path. To rule out non-physical underdamping effects, we verified that such hysteretic effects persist even under a doubled damping rate \( \eta = 56 \gamma_0 \), indicating that a competition among multiple dynamical attractors is indeed present in this model.

In the present study the adopted \( \hat{x} \)-directed orientation of the driving force tends to drive the colloids straight across the saddle points of the corrugation potential. By exploring other more general pushing directions, depending on the angle formed by the force with the \( \hat{x} \)-direction, the colloids would be driven toward the corrugation maxima too. This kind of investigation has been carried out in the past for different models or experimental setups, typically finding locking to energetically and/or dynamically favorable advancement directions [45, 49, 50]. We expect similar directional-locking phenomena in the present model. However, possible novel regimes realized by the current cluster model may include concurrent multiple decomposition paths for the clusters, with several particles leaving simultaneously a potential well to reach different neighboring ones. We leave the investigation of these regimes to future research.

### B. Thermal Effects

We simulate finite temperature by restoring the random forces \( \xi \) in the equations of motion (2), and therefore sampling the canonical ensemble. The starting point of the \( T > 0 \) simulations is the appropriate \( T = 0 \)
Figure 14. The scattering intensity $S(q)$ obtained assuming that each colloid acts as structureless point-like scatterer. The simulations refer to $n = 4$, at four temperatures expressed in units of energy (i.e. assuming $k_B = 1$): (a) $T = 0$ – the ordered state in the static condition depicted in Fig. 5, (b) $T = 0.01 E_0$ – rotational melting of the cluster lattice, (c) $T = 0.15 E_0$ – crystal lattice phase just below structural melting, and (d) $T = 0.17 E_0$ – a liquid phase just above melting. The $T > 0$ simulations of panels (b-d) are carried out for the free model ($V_0 = 0, F = 0$). $S(q)$ is averaged over 30 snapshots widely spaced in time along an equilibrium simulation.

Static configuration (a perfect cluster lattice) obtained in Sect. 11. In the simulated scattering intensity reported in Fig. 14, the modulation of the Bragg peaks reflects the structure factor of the orientationally ordered lozenge-shaped clusters at $T = 0$, see Fig. 5. In the finite-temperature simulations of the free model ($V_0 = 0, F = 0$), it is straightforward to verify that even a low temperature, such as $T = 0.01 E_0$, induces the melting of the orientational order of the clusters, as seen in the diffuse rings at integer multiples of $2\pi/\delta \approx 20 R^{-1}$ reported in Fig. 14. Further raising temperature leads rapidly to more disordering of the cluster crystal, and eventually to melting, which occurs between $T = 0.15 E_0$ and $T = 0.17 E_0$, see Fig. 14.d.

According to this analysis, we focus the investigation of the driven model on moderately-low temperature $T \leq 0.05 E_0$. In the simulations under driving, as we did in the $T = 0$ protocol, successive runs at larger and larger driving force $F$ are started from the final configuration of the previous step in $F$, which is increased "adiabatically" by $\Delta F = 0.01 F_{1s}$ at each step. For each step, the simulation duration is $500 \tau_0$; the first 30% is dropped to prevent transient effects, and the remainder is used for the determination of the mobility, providing a fair averaging over thermal fluctuations.

Strictly speaking, at finite temperature there is no static friction, because thermal fluctuations would lead to a diffusive displacement of the layer even for $F = 0$, and to a slow systematic drift in the force direction for $F > 0$, if one had the patience to wait long enough for these thermally-activated rare events to occur. However, at the relatively small considered temperatures $T \ll V_0$, the rate of well-to-well thermal-assisted hopping is extremely small, and as long as the applied force is small $F \ll F_{1s}$, we observe no such hoppings at all for the duration of our simulations. This allows us to define $F_{\text{dep}}$ even for $T > 0$, as the threshold force beyond which a significant center-of-mass displacement is observed in the applied force direction within the simulation duration. For forces immediately before this threshold, the mobility is not precisely null, but it is really tiny, with overall advancements of the entire system by far less than one lattice spacing $a_{\text{pot}}$ over the entire simulation time. We note also that all particle well-to-well displacements occur within horizontal rows: although a-priori possible, we never observe any particle abandoning their initial X-oriented row and transferring to a well in the next row. These cross-row hoppings would of course become frequent for larger $T/V_0$. We do however observe particle jumps within rows, leading to a limited but nonzero random concentration of $n - 1$ and $n + 1$ "defective" clusters in the pre-depinning quasi-immobile state.

The mobility resulting in the finite-temperature simulations is exemplified in Fig. 15. This figure exhibits the following characteristic features, common to analogous simulations that we carried out for clusters of sizes $n = 2$ to $5$, and reported in Appendix: (i) The depinning threshold $F_{\text{dep}}$ decreases for increasing temperature, be-
cause thermal fluctuations help the particles anticipate the barrier hopping, thus activating sliding even when the driving force is nominally insufficient to overcome the barrier. (ii) For $F \gtrsim F_{\text{dep}}$, immediately above the threshold, thermal fluctuations tend to favor the sliding state, as shown by the higher mobility proving a sort of thermlubric effect \[51, 59\]. However, depending on $n$ and the corrugation amplitude $V_0$, for forces larger than the $T = 0$ depinning threshold $F_{1s}$, increasing temperature can lead to either a slightly decreasing mobility, as in Fig. 15, or essentially $T$-independent mobility, quite similar to the values obtained for $T = 0$, see Appendix A.

The main effect of finite temperature on the cluster dynamics is to suppress the $T = 0$ synchronized advancements, especially those associated to the fractional-advancement modes. As a result, at $T > 0$ the center-of-mass advancement is generally smoother, with no, or at least substantially smeared, steps like those occurring for the $T = 0$ fractional sliding regimes reported in Fig. 13. Nonetheless, some amount of approximate synchronization is preserved in the special case of whole-cluster hopping, e.g. at $T > 0$ the 3/3 synchro pattern of Fig. 7, even extends down to smaller $F$ than for $T = 0$. Relatedly, the sharp transitions between different sliding modes observed at $T = 0$, at finite $T$ are replaced by smooth crossovers.

\textbf{V. DISCUSSION AND CONCLUSIONS}

In the present work we introduce and study a driven model for friction at the micro scale where the protagonists are spontaneously formed clusters, rather than simple structureless objects \[12, 60-76\] or structured but unbreakable units \[77, 78\], as usually considered in previous investigations. We focus on a lattice-matched situation, with the external corrugation periodicity matching exactly the equilibrium spacing between the clusters. In the language of the Frenkel-Kontorova model \[3, 79-82\], this is the typical situation which maximizes the pinning effect of the corrugation, leading to a finite static friction regardless of how small the corrugation amplitude may be. We do find indeed a finite static friction in our simulations of the current model, with clustering affecting the static-friction threshold and the sliding mechanisms for driving force exceeding this threshold.

For small corrugation amplitude, the clustering tendency is dominating the dynamics, so that clusters tend to advance as a individual objects, with the partial exception of single-particle hopping induced by thermal fluctuations near the depinning threshold. In contrast, at larger corrugation amplitude, the stability of clusters is challenged when the driving force moves them in regions where the curvature of the external corrugation potential is negative: this leads to cluster decomposition and partial hoppings. In this regime, the internal structure of the clusters decomposes in several ways as a function of the driving force, thus producing a quite rich dynamics with complicate fractional advancement patterns for clusters of various size. We investigate the impact of these phenomena on the mobility of the system. At $T = 0$ we observe and characterize well-defined regimes of sliding of entire clusters or of decomposition-recomposition processes, with deformation of the clusters. These processes can take place in a synchronous or in asynchronous way across the system. Thermal effects tend to destroy such distinct regimes: we observe a gradual change of sliding under increasing driving force, already for temperature as low as ~ 5% of the temperature at which the transition from a cluster crystal to a uniform fluid occurs.

The phenomenon of cluster decomposition/recomposition under driving is not expected to be specific of the interparticle interaction adopted here. We therefore predict that experiments carried out in conditions where spontaneous clustering occurs should observe this kind of dynamical behavior.

In the future we plan to extend the investigation to lattice-mismatched configurations. In incommensurate conditions, for increasing corrugation amplitude $V_0$ there is room for an Aubry-type transition from a “superlubric” dynamics characterized by "solitonic" sliding modes \[79, 80, 83-88\]. A similar condition could be realized by taking for the substrate a symmetry other than hexagonal, as e.g. in Ref. \[72, 99, 101\]. In the small-$V_0$ superlubric ground state, a finite fraction of clusters is supposed to sit near one of the maxima of the incommensurate corrugation potential, rather than all in a potential well as in the fully commensurate setup of the present paper. The novelty for a cluster-supporting condition is that, as $V_0$ is raised, two competing effects may arise: either an entire cluster near the maximum slides

Figure 15. The mobility, relative to the value $\eta^{-1}$ of free colloids, as a function of the driving force, relative to the $T = 0$ depinning threshold $F_{1s}$ of isolated colloids, at three relatively small but finite temperatures, expressed in units of energy $E_0$. The $T = 0$ solid curve is the same as in Fig. 7. Simulations are carried out for $n = 3$-particle clusters, for corrugation $V_0 = 2E_0$. Statistical fluctuations account for error bars (not drawn) on mobility mostly smaller than $5 \times 10^{-3} \eta^{-1}$. 

\[\frac{\eta}{F}, \frac{V_0}{F_1s} \]

\[T = 0, T = 0.01 E_0, T = 0.02 E_0, T = 0.05 E_0\]

\[n = 3, V_0 = 2 E_0\]
down toward a nearby potential well, thus leading to a relatively standard Aubry-type transition \[94, 97\], or alternatively it decomposes under the action of divergent corrugation forces, leading to a structurally inequivalent ground state. Which of these mechanisms turns in earlier is open for investigation. These competing possibilities are likely to lead to a rich phase diagram.

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**Appendix A: Thermal effects on mobility**

Figures 16 - 19 report simulated mobilities as a function of the driving force. The simulations are carried out at the values of the average particle density given by Eq. (7), for \(n = 2, 3, 4, 5\), compatible with clusters of \(n\) particles arranged in a regular hexagonal lattice with spacing 1.5 \(R_e\). Three different values of corrugation amplitude are considered, namely: \(V_0 = E_0\), \(V_0 = 1.5 E_0\), and \(V_0 = 2 E_0\). Each figure compares the \(T = 0\) mobility (also shown in the Figs. 16-19) with the homologous quantity obtained in finite-temperature simulations, carried out at three comparably small temperatures: \(T = 0.01 E_0\), \(T = 0.02 E_0\), and \(T = 0.05 E_0\).

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Figure 18. Same as Fig. 16, but for clusters formed by \( n = 4 \) particles.

Figure 19. Same as Fig. 16, but for clusters formed by \( n = 5 \) particles.

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