Kinetic interfaces of patchy particles

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Received 24 September 2014, revised 26 November 2014
Accepted for publication 4 December 2014
Published 29 April 2015

Abstract
We study the irreversible adsorption of patchy particles on substrates in the limit of advective mass transport. Recent numerical results show that the interface roughening depends strongly on the particle attributes, such as, patch–patch correlations, bond flexibility and strength of the interactions, uncovering new absorbing phase transitions. Here, we revisit these results and discuss in detail the transitions. In particular, we present new evidence that the tricritical point, observed in systems of particles with flexible patches, is in the tricritical directed percolation universality class. A scaling analysis of the time evolution of the correlation length for the aggregation of patchy particles with distinct bonding energies confirms that the critical regime is in the Kardar–Parisi–Zhang with quenched disorder universality class.

Keywords: kinetic roughening, adsorption on substrates, absorbing phase transitions

(Some figures may appear in colour only in the online journal)

1. Introduction
The nonequilibrium evolution of growing interfaces has attracted many experimental and theoretical studies [1–15]. One of the most popular theoretical approaches considers kinetic discrete models to describe particle aggregation on substrates. Albeit simple, these models are expected to contain the relevant physics [16–18]. In particular, the ballistic deposition model (BD) [19, 20], is considered the prototype for irreversible aggregation on substrates. In BD, the growth is driven solely by the sequential addition of particles to the aggregate, which stick to the first particle they touch, without subsequent rearrangement. From the simple rules of ballistic deposition a complex structure emerges with a nontrivial porous bulk structure (see e.g. [16, 19, 21]) and a kinetically rough interface in the Kardar–Parisi–Zhang universality class [16, 22].

Inspired by recent advances in the production of patchy particles we have proposed a stochastic model to study their aggregation on substrates [23], which in the limit of advective mass transport is a generalized version of BD. Patchy particles are colloids with functionalized surfaces, with new features such as selective and directional particle–particle interactions, control over the valence and the possibility of forming permanent electrical dipoles [24–32]. Studies of the irreversible aggregation on substrates reveal a nontrivial dependence of the bulk and surface properties on the mechanism of mass transport [33], on the strength of the patch–patch interactions [34, 35] and on the spatial-patch distribution [36].

Here, we focus on the scaling properties of the growing interface in the limit of advective mass transport. In this limit, we have found new absorbing phase transitions depending on the patch–patch interactions [33] and a crossover in the universality class of the interface depending on the relative strength of the patch–patch interactions [35]. These findings have been discussed previously in the context of functional colloids. Here we revisit these transitions and investigate their scaling properties in the framework of kinetic discrete models of interfacial growth.

The paper is organized in the following way. In section 2 we describe the model and recall some definitions. The two absorbing transitions are discussed in sections 3 and 4. The crossover of the universality class of the interface is discussed in section 5. Some final remarks and future perspectives are provided in section 6.

2. Model
In the ballistic deposition model [16, 18, 37] particles are sequentially released from a position above the interface,
chosen uniformly at random and move vertically towards the substrate sticking irreversibly to the first particle they touch. To account for the directionality of the interactions, the excluded volume interaction between particles and the short-ranged patch–patch attraction, characteristic of patchy colloids, we proposed a generalized version of this model in [23, 33], which we describe below.

To access larger-system sizes, let us consider a two-dimensional system of patchy particles (disks) of unit diameter $\sigma$ with an initially empty flat (linear) substrate of length $L$. As in the ballistic deposition model, we iteratively generate a horizontal position, chosen uniformly at random above the interface, to release a particle and follow its ballistic trajectory downwards until the particle collides either with the substrate or with another particle. Collisions with the substrate always result on adsorption of the particle at the collision point with a random orientation.

While in the ballistic deposition model two particles stick together upon collision, in the case of patchy particles the success of bond formation depends on the relative orientation between the particles. The $n$ patches are located on the surface of the particle and for each patch we define an interaction range around it. The interaction range accounts for the extension of the patch and the range of the patch–patch interaction and it is characterized by a single parameter $\theta = \pi/6$, representing the maximum angle with the center of the patch (see figure 1(a)). Two patches bond in an irreversible way, a process we call binding, if their interaction ranges partially overlap. Thus, stochastically, if the new particle collides within the interaction range of a particle already on the substrate, it binds to it with a probability $p = A_p/A$, where $A = \pi \sigma$ is the surface area of the incoming particle and $A_p$ is the extension of this area covered by the interaction range of all patches [23]. If binding is not successful the particle is removed from the system and a new one is released from the top.

Inspired by chemical or DNA mediated bonds [38–40], we consider highly directional and very strong bonds between patches. Thus, in the case of successful binding, the binding

\[ \delta < \delta_{\text{min}} \]

of the incoming particle is selected uniformly at random among its patches. The position and orientation of the incoming particle is then adjusted such that the binding occurs along the center of the two bonding patches (see figure 2).

### 3. patch–patch correlations

The dependence of the interface of patchy particle aggregates on the spatial arrangement of the patches was studied in [36]. In that study we considered three-patch particles and investigated the dependence on the opening angle, showing that growth is suppressed below and above a minimum and a maximum opening angles, with two absorbing phase transitions between thick and thin adsorbed film regimes. In this section, we investigate the nature of those transitions.

As shown schematically in figure 1(a), the particles are disks with three patches: one reference patch and two adjustable ones. The spatial arrangement of the patches is characterized by the opening angle $\delta$, from the reference patch to the adjustable ones. For simplicity, we define the units of $\delta$ as $\pi$ rad. As the particles are sequentially added, a network of patchy particles grows away from the substrate (see figure 3) but, its growth is only sustained for $\delta_{\text{min}} < \delta < \delta_{\text{max}}$.

For $\delta < \delta_{\text{min}}$, the angle between the patches is such that all patches are in the same hemisphere. Thus, the patches of particles in the aggregate are pointing most likely towards the substrate and are not accessible to new incoming particles. When there are no more patches available to establish new bonds, the growth is suppressed. A systematic finite-size study for the threshold value gives $\delta_{\text{min}} = 0.468 \pm 0.001$ [36].

This threshold is above that expected from purely geometrical

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arguments, revealing strong collective effects \[36\]. To analyze the transition to the absorbing state at \(\delta_{\min}\), we define as the order parameter \(r\) the fraction of successful binding attempts. Figure 4, shows the dependence of \(r\) on \(\delta\) for different system sizes. Clearly, in the thermodynamic limit, \(r\) vanishes at \(\delta_{\min}\) and it grows continuously with \(\delta\). As shown in the inset, a data collapse is obtained for the finite-size scaling consistent with the directed percolation (DP) universality class in two dimensions \[18, 41, 42\].

For \(\delta > \delta_{\max}\), the distance between the adjustable patches is such that, if one particle binds to one of these patches it shields the access of a new particle to the other. Consequently, branching is suppressed and only chains grow away from the substrate. These chains are locally tilted and their growth direction (given by the available patches), fluctuates while growing. Eventually, the growing tip of the chain points down and its growth is suppressed. The absorbing state occurs when all tips are either pointing down or covered by other and its growth is suppressed. The absorbing state occurs as a function of \(\delta\), for three different substrate lengths \(L\) namely, 512, 1024 and 2048. The inset shows the finite-size scaling universality class. We considered \(\beta_{DP} = 0.58\) and \(\gamma_{DP} = 0.73\). Results are averages over \(\{400, 200, 100\}\) samples for \(L = \{512, 1024, 2048\}\).

4. Bond flexibility

For simplicity, in the previous section the position of the incoming particle is adjusted such that the center of the colloids and of their patches is perfectly aligned. However, in reality, one expects some degree of flexibility around this optimal orientation \[38, 44\]. A simple strategy to account for flexibility was proposed in \[36\], which takes advantage of the stochastic nature of our model. The idea is still to consider rigid and irreversible bonds but, at a binding event, the orientation of the bond deviates by an angle \(\gamma\) from the optimal orientation (see figure 6). The value of \(\gamma\) is drawn randomly from a Gaussian distribution of zero mean and dispersion \(F\theta\), where \(F\) is the flexibility, truncated at \(F\theta\). The sense of rotation of \(\gamma\) is always from the center of the patch to the point of collision.

Results for different values of the flexibility are summarized in the diagram shown in figure 7. The active region does widen with the flexibility but there is always a lower and an upper thresholds. For the range of flexibilities considered, the transition at \(\delta_{\min}\) is always continuous and in the DP universality class, as shown in figure 8. The threshold \(\delta_{\min}\) decreases with \(F\) but it is always significantly higher than 1/3, the one predicted from purely geometrical considerations.

For the second transition, the threshold \(\delta_{\max}\) increases with \(F\) and converges towards unity at large flexibilities. In fact, the threshold value is well approximated by,

\[
\delta_{\max}(F) = \delta_{\max}(0) + \frac{F\theta}{\pi},
\]

obtained from geometrical arguments \[36\], corresponding to the dashed line in figure 7. The nature of the transition also changes with \(F\). Figure 9(a) shows the dependence of the order parameter \(r\) on \(\delta\) close to the second transition, for three different values of \(F\). The numerical results suggest that even for \(F = 0.075\), the transition is still discontinuous (see also the histogram of the order parameter in figures 9(b) and (c)) but the size of the jump decreases as \(F\) increases. For \(F > 0.2\)
the numerical results clearly suggest a continuous transition in the DP universality class (see figure 10). The inset of figure 10 suggests that the size of the jump decreases with the distance to $F = 0.1$ as a power law with exponent $0.25$. This suggests, in turn, that the nature of the transition changes from discontinuous to continuous at a tricritical flexibility $F_{tc} \approx 0.1$, in the tricritical directed percolation universality class [45, 46].

As the tricritical point is approached from lower flexibilities, the fraction of successful binding attempts goes to zero at $\delta_{\max}$. This is intimately related to the film rate of growth close to the threshold. For a discontinuous transition, the rate of growth is always non-zero in the active phase.

5. Weak and strong bonds

Patchy particles with distinct patch–patch interactions yield interesting bulk properties at equilibrium [47–56] and novel nonequilibrium interfaces [34, 35]. The typical strategy is to consider $2AnB$ particles, with two strong $A$- and $n$ weak $B$-patches. The $A$-patches are in the poles, while the $B$-patches are equally spaced along the equator (see figure 1(b)). A generalization of our model to account for these two energy scales was proposed in [34, 35], based on dissimilar binding probabilities that can be formally related to the activation energy of binding. At a collision event with partial overlap between the interaction range of two patches $i$ and $j$, the binding is successful with a binding probability $P_{ij}$ that depends on the type of patch pair $ij$. In particular, since $A$-patches are stronger than $B$-patches, we consider $P_{AA} > P_{BB} = P_{AB}$. Without loss of generality, we consider $P_{AA} = 1$ and define the sticking coefficient $r_{AB} = P_{AB}/P_{AA}$. Lower sticking coefficients favor chain growth over branching.

In [35] we have shown that a crossover of the interfacial roughening from the Kardar–Parisi–Zhang (KPZ) to the KPZ with quenched disorder (KPZQ) is observed when $r_{AB}$ is sufficiently small, i.e. when the $A$-patches are significantly stronger than the $B$-patches. For $r_{AB} \ll 1$, the strong $A$-patches, promote growth along the poles, favoring the aggregation of $AA$ chains. These chains are likely to extend over long lateral regions. These long chains expose their $B$-patches and shield the access to any underlying $A$-patch, thus
the probability of binding there is significantly lower. This is expected to have a similar effect to that of quenched noise [35].

Note that the KPZ universality class is very robust [22], while KPZQ is only observed at the critical depinning transition [1, 57–63]. Nevertheless, for patchy particles with weak and strong bonds one finds remarkably an entire critical region of $r_{AB}$ where KPZQ is observed. This is likely due to the balance of two competing mechanisms that keep the system at criticality. As $r_{AB}$ decreases, the probability of binding to $B$-patches decreases and the growth of longer $AA$-chains increases. The longer these chains, however, the larger is the number of $B$-patches available for bonding, which compensates the decrease in the binding probability.

Recent experiments on aggregation of ellipsoids at the edge of an evaporating drop suggest that, for sufficiently large major–minor axis aspect ratio the interface is also in the KPZQ universality class [64]. Together with these experiments, a discrete model was proposed to argue that KPZQ was driven by a colloidal Matthew effect [64]. However, numerical simulations of the same model by Nicoli et al [65] suggest a different interpretation of the theoretical results. Oliveira and Reis performed a careful statistical analysis of the correlation length and concluded that the differences observed in the growth and roughness exponents are due, instead, to a crossover to columnar growth [66]. Although the results for the interface of the simple models appear to be settled the nature of the interface of the experimental system is still an open problem [64].

In order to proceed and to discard the possibility of columnar growth in the model of patchy particles with strong and weak bonds, we performed the analysis proposed in [66]. Figure 11 shows a section of the aggregate and height profile for one sample with $r_{AB} = 0.01$. The time evolution of the height profile reveals no evidence of columnar growth. A more quantitative analysis is illustrated in figure 12 where we have plotted the scaling of the position $r_0$ of the first zero of the autocorrelation function with time. This position is expected to scale as,

$$ r_0 \sim t^{1/z}, $$

where $z$ is the dynamic exponent. In spite of the strong finite-size effects, the size scaling of the slope (inset of the
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We reviewed our recent results on the kinetic roughening of

figure) indicates that the growth is consistent with the KPZQ

universality class.

6. Final remarks

We reviewed our recent results on the kinetic roughening of interfaces of patchy particles and showed that the scaling of the interface depends strongly on the patch–patch correlations, the bond flexibility and the strength of interactions. For particles with patch–patch correlations and bond flexibility we found two absorbing phase transitions that are, in general, of different nature. While the first transition is always continuous in the directed percolation universality class the second is either continuous or discontinuous depending on the bond flexibility. A scaling analysis of the size of the jump close to the tricritical flexibility suggests that the tricriticality is in the tricritical directed percolation universality class. For particles with weak and strong bonds we analyzed the kinetics of particles with two types of patches (A and B). When the strength of the A and B patches is similar, the interface roughness is in the Kardar–Parasi–Zhang universality class. However, for A-patches significantly stronger than the B-patches, the interface is in the universality class of Kardar–Parasi–Zhang with quenched disorder. This critical universality class is observed for an entire range of the relative strength of the interactions.

Beyond their theoretical interest, our findings have three consequences with practical applications. First, we have shown that the roughness of the interface can be controlled by the spatial distribution of patches or the relative strength of their interactions. Second, we revealed that sustained growth is only possible for certain patch arrangements. Third, the existence of an extended region of the parameter space where the critical KPZQ universality class is observed opens the possibility for an experimental realization of such systems using patchy colloids with weak and strong bonds.

As a follow up, it would be interesting to analyze the experimental observations reported in [64] in the light of our findings. For simplicity, we have considered bonds to be irreversible within the time scale of interest. In spite of being a reasonable approach for the systems considered here, for sufficiently long timescales or at higher temperatures, an adsorbed colloid might detach and rebind. It would be interesting to analyze the stability and aging of the interfaces when reversibility is considered.

Acknowledgments

We acknowledge fruitful discussions with A G Yodh and T J Oliveira, as well as financial support from the Portuguese Foundation for Science and Technology (FCT) under contract nos. EXCL/FIS-NAN/0083/2012, PEst-OE/FIS/UI0618/2014 and IF/00255/2013.

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