Smoluchowski equations for linker-mediated irreversible aggregation†

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We developed a generalized Smoluchowski framework to study linker-mediated aggregation, where linkers and particles are explicitly taken into account. We assume that the bonds between linkers and particles are irreversible, and that clustering occurs through limited diffusion aggregation. The kernel is chosen by analogy with single-component diffusive aggregation but the clusters are distinguished by their number of particles and linkers. We found that the dynamics depends on three relevant factors, all tunable experimentally: (i) the ratio of the diffusion coefficients of particles and linkers; (ii) the relative number of particles and linkers; and (iii) the maximum number of linkers that may bond to a single particle. To solve the Smoluchowski equations analytically we employ a scaling hypothesis that renders the fraction of bondable sites of a cluster independent of the size of the cluster, at each instant. We perform numerical simulations of the corresponding lattice model to test this hypothesis. We obtain results for the asymptotic limit, and the time evolution of the bonding probabilities and the size distribution of the clusters. These findings are in agreement with experimental results reported in the literature and shed light on unexplained experimental observations.

I. Introduction

The self-assembly of micro- and nano-size particles mediated by linkers has been the focus of extensive experimental,1,9 theoretical10–12 and simulation4,10,11,13–17 studies in recent years, since the addition of a second species provides extra control of the aggregation process. Indeed the linkers to particles ratio,1,4,6,8,9 the strength of the linker-particle interactions,5,7 and the maximum number of linkers that can bond to one particle,7,8 may be used to tune the stability of the mixture, the extent and topology of the assembled structures, the equilibrium phase behavior and the rheological properties of the system.3,18 The linker-particle interactions are most often engineered using complementary biological macromolecules, such as complementary DNA strands5,8 or streptavidin and biotin.1,2,4 As a consequence, the bonds formed between particles mediated by linkers are extremely strong. These systems form several types of disordered and more or less open clusters, even at extremely low volume fractions.3,4,8 Theoretical studies have focused mostly on equilibrium properties.12,14 Some simulations have addressed the dynamics of these systems, either to get information about the early stages of aggregation4 or to try to predict the type of assembled structures at long times.15,16 An exception is the work reported in ref. 11, where a theory for the dynamics of aggregation of linker-particle binary mixtures was developed and studied.

The simplest theoretical description of the kinetics of aggregation is based on the Smoluchowski coagulation equation,19,20 two clusters of mass $i$ and $j$ merge at rate $K_{ij}$ to form a cluster of mass $i + j$. This description is general and flexible:21–23 its application to a particular type of aggregation is encoded in $K_{ij}$ -- the so called kernel. The choice of the appropriate kernel for a given aggregation process results from a wise combination of (intuitive) knowledge of the specificities of the clustering physical mechanism and the ability to draw relevant information from the solution of the resulting equation.24 The wide variety of kernels that have been studied22,23 depend on the size or mass of the aggregating clusters, but not on the presence of other components that promote aggregation. This includes the only proposal (to the best of our knowledge) to use Smoluchowski equations to study linker particle aggregation:11 the effect of linkers was subsumed into an effective functionality and a single time-scale for bonding was considered. These assumptions

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led to the use of the polymerization kernel that is appropriate for single particle aggregation.

The goal of this work is to develop a generalized Smoluchowski equation for linker-mediated aggregation where linkers and particles are explicitly taken into account. Inspired by the experimental works of ref. 1, 4 and 8, we will assume that the bonds between linkers and particles are irreversible and that clustering happens when there is an encounter between diffusive linkers and particles (limited diffusion aggregation). The choice of the kernel is dictated by analogy with single component diffusive aggregating systems (where the Brownian kernel is approximated by a constant kernel and by distinguishing clusters not only by their total size or mass, but by the number of particles and the number of linkers that it contains. We find that, even under these strong approximations, the dynamics depends on three relevant factors, all corresponding to controlable experimental quantities: (i) the ratio between the diffusion coefficients of the particles and linkers; (ii) the concentration or the relative number of particles and linkers and (iii) the maximum number of linkers that may bond to a single particle. This enables a direct comparison between the results of the theory and existing experimental results where the effect of these control parameters was investigated.

The paper is organized as follows. In Section IIA we describe and justify the choice of the kernel for linker-mediated aggregation and derive the corresponding Smoluchowski equations. We solve these equations in two ways: by simulating a lattice model of linkers and particles whose time evolution is determined by those equations – Section IIB – and analytically, using a scaling hypothesis that leads to an approximate analytical solution of the same equations – Section IIC. We report results for the asymptotic limit, for the time evolution of both the bonding fraction and the size distribution of the clusters, obtained from the simulations and from the approximate analytical theory, in Section IIIA–C. Some of the surprising experimental results reported in ref. 4 are revisited and explained in Section IIID. Finally, we discuss our results, draw some conclusions and elaborate on the possibility of generalizing the theory in Section IV.

II. Theory

A. Smoluchowski equations

We consider a mixture of \( N_p \) particles and of \( N_l \) linkers in a volume \( V \). Particles, linkers and clusters diffuse and bond irreversibly upon encounter. The linkers mediate the bonding of the particles: two particles become bonded when one of them bonds to a linker that is already bonded to another particle. Two bonded particles belong to the same cluster (and this defines a cluster as the set of particles that belong to it). Each linker bonds to a maximum of two particles and each particle bonds to a maximum of \( f \) linkers (\( f \) is often called the particle valence). As a consequence, each linker can be in one of three states: not bonded to any particle (state 0, free linker), bonded to one particle only (state 1) or bonded to two particles (state 2). \( N_i \) is the number of linkers in state \( i \) (so that \( N_0 + N_1 + N_2 = N_L \)). The linkers control the extent of aggregation of the particles, and to quantify this effect we use the parameter \( \phi \), the ratio between the actual number of linkers and the maximum number of linkers that could be bonded to all particles,

\[
\phi = \frac{N_L}{fN_p}
\]

The size \( i \) of a cluster is defined as the number of particles that belong to it. We assume that two particles that are bonded to each other are connected by only one linker (no double bonding) and that the clusters are tree like (no loops). Therefore, a cluster of size \( i \) contains \( i - 1 \) linkers in state 2, corresponding to the number of bonds needed to form it. Each cluster of size \( i \) contains also a number \( j \) of linkers in state 1 in the interval \([0;w_j]\), with \( w_j = (f-2)j + 2 \). Notice that a particle with no linkers bonded to it is a cluster of size \( i = 1 \) with \( j = 0 \).

It is useful to consider that a cluster of size \( i \) has a total of \( w_i \) bonding sites on its “surface”; if it contains \( j \) linkers in state 1, then \( j \) of these sites are occupied (by a linker) and \( w_i - j \) sites are unoccupied. Therefore each cluster is characterized by two integer numbers and thus a cluster \((i,j)\) consists of \( i \) connected particles and \( j \) linkers in state 1 or \( j \) occupied sites (see Fig. 1(a)). Two clusters will aggregate when one unoccupied site in one cluster bonds to an occupied site in the other.

We now propose a generalized Smoluchowski equation for the time evolution of the clusters, which takes into account two distinct aggregation mechanisms: bonding of free linkers to clusters \((i,j-1)\) and cluster-cluster aggregation. Let \( m_{ij} \) be the number of clusters \((i,j)\). The rate at which \( m_{ij} \) changes is,

\[
m_{ij} = K_{0,+} - K_{0,-} + K_{+} - K_{-},
\]

where:

1. \( K_{0,+} \) is the rate at which free linkers bond to clusters \((i,j-1)\) and form a \((i,j)\) cluster,

\[
K_{0,+} = K_0(i,j - 1) \frac{N_0 m_{i,j-1}}{V},
\]

2. \( K_{0,-} \) is the rate at which free linkers bond to \((i,j)\) clusters,

\[
K_{0,-} = K_0(i,j) \frac{N_0 m_{i,j}}{V},
\]

3. \( K_{+} \) is the rate at which two clusters bond to form a \((i,j)\) cluster,

\[
K_{+} = \frac{1}{2} \sum_{i_1+j_1=i} \sum_{i_2+j_2=j} K(i_1,j_1;i_2,j_2) \frac{m_{i_1,j_1} m_{i_2,j_2}}{V};
\]

4. \( K_{-} \) is the rate at which a cluster \((i,j)\) bonds to other clusters:

\[
K_{-} = \sum_{i=1}^{\infty} \sum_{j=0}^{w_i} K(i,j-1;i,j) m_{i,j-1} m_{i,j} \frac{1}{V};
\]

\( K_0(i,j) \) and \( K(i_1,j_1;i_2,j_2) \) are the kernels for the two types of aggregation. They are chosen to be the product of the Brownian...
A cluster \((i,j)\) consists of \(i\) particles (each can be bonded to a maximum of \(f\) other particles) and has \(j\) sites occupied (out of \(w_i = (f - 2i) + 2\), and \(w_i - j\) sites unoccupied; since the cluster is tree like, it has \(i - 1\) bonds. An occupied site is a linker in state 1 and a bond is a linker in state 2. In the example of the figure \(f = 4, i = 2\) and \(j = 3\), therefore \(w_i = 6\), there is 1 bond (or linker 2) and 3 unoccupied sites. (b) Diffusion. The constituents diffuse randomly: linkers with a diffusion coefficient \(D_P\) (or hopping rate \(H_L\) in the simulation) and clusters (of all sizes) with a diffusion coefficient \(D_L\) (or hopping rate \(H_L\) in the simulation). Only cases where \(D_L \geq D_P\) are considered. (c) Linker-cluster aggregation happens with probability \(p_0\) (given by eqn (9)) when a free linker encounters a cluster \((i,j)\). After bonding to the cluster, the free linker turns into a linker 1 while the cluster gains an occupied and loses an unoccupied site. The same bonding happens with equal probability when a cluster encounters a free linker. (d) Cluster-cluster aggregation happens with probability \(p\) (given by eqn (10)) when two clusters meet. The size of the new cluster is the sum of the sizes of the clusters that merged; as a consequence one occupied and one unoccupied sites (one in each cluster) disappear and a new bond forms. In the example of the figure, clusters \((1,2)\) and \((2,3)\) aggregate to form a cluster \((3,4)\).

The restrictions to aggregation expressed by eqns (9) and (10) incorporate in the dynamics the limited number of linkers that a particle can support in a simple way. In both cases, each aggregation event is considered to be instantaneous: the formation of a bond is much faster than all other processes and may be considered instantaneous compared to the other relevant time scales (diffusion limited aggregation). In addition, since we focus on the limit of low density of particles and linkers, the timescale of the linker-mediated aggregation is sufficiently large for the shape and orientation of each cluster to be uncorrelated between successive bonding events. As a consequence (as already expressed in (9) and (10)), every site of the “surface” of a cluster is equally likely to form the next bond. The relevant time scales are set by the diffusion coefficients of the particles, \(D_P\), and the free linkers, \(D_L\). We define as control parameter the ratio \(\Lambda\) of the diffusion coefficients,

\[
\Lambda = \frac{D_L}{D_P}
\]
We consider the limit where the linkers diffuse faster than the particles (i.e. $\Delta \geq 1$), since it is reasonable to assume that the linkers have a lower mass (and size) than the particles.

The generalized Smoluchowski equation that describes cluster formation is then given by eqn (2), within the approximations expressed by eqn (7)-(10). The description of the dynamics of aggregation is completed with the equation for the time evolution of the free linkers, i.e. the rate at which free linkers bond to clusters that, subject to the approximations described is,

$$N_0 = -\sum_{j=0}^{\infty} \sum_{i=0}^{\infty} K_{0i} = -2k_p z N_0 \sum_{i=0}^{\infty} p_0(i,j) n_{ij}. \quad (12)$$

The aggregation dynamics for clusters of size $i$ can be obtained from eqn (2) by using $m_i = \sum_{j=0}^{\infty} m_{ij}$ and taking into account that $\sum_{j=0}^{\infty} (K_{0i} + K_{0j} - K_{ij}) = 0$,

$$m_i = \frac{k_p}{V} \sum_{j=0}^{\infty} \sum_{l=0}^{\infty} p(i,j) m_{ij} + \sum_{j=0}^{\infty} \sum_{l=0}^{\infty} p(i,j) m_{ij} - 2k_p \sum_{j=0}^{\infty} \sum_{l=0}^{\infty} \sum_{l=0}^{\infty} p(i,j) m_{ij} \quad (13)$$

In the next two subsections we solve eqn (12) and (13) to obtain explicitly the time evolution of the aggregation process in the particle linker system, within the approximations adopted. First, we will perform kinetic Monte Carlo simulations of a lattice model with the same dynamics. Then, we will use a scaling hypothesis that leads to an (almost) analytical solution, by reducing eqn (12) and (13) to a system of two ordinary differential equations.

B. Numerical simulations

$N_p$ particles (or clusters $(1,0)$) and $N_t = \phi N_p$ free linkers are randomly distributed without overlap on a simple cubic lattice with $N_{latt}$ sites and periodic boundary conditions, each occupying one lattice site ($R_t = R_p$ in the simulations). The initial density of particles $\rho = N_p/N_{latt}$ is set to a low value ($\rho = 0.1$ for the simulations reported) and $f$ is chosen to be 6. The diffusive motion of the clusters and free linkers is described by kinetic Monte Carlo simulations. At each iteration, one cluster or one free linker is chosen and attempt a hop to a randomly chosen neighbouring lattice site. Free linkers are chosen with probability $H_t N_0 Q$ and clusters of size $i$ with probability $H_p m_i/Q$, where $N_0$ and $m_i$ are the actual numbers of free linkers and of clusters of size $i$, respectively, $Q$ is the total hopping rate,

$$Q = H_t N_0 + H_p \sum_i m_i, \quad (14)$$

and $H_t$, $H_p$ are two hopping rates that can be related to the diffusion coefficients of the linkers and particles (see below).

For a linker, depending on the state (occupied or empty) of the randomly chosen neighbouring site, the hop is:

- Accepted if it is not occupied;
- Rejected if it is occupied by another free linker;
- Rejected/accepted with probability $1 - p(i,j)/p_0(i,j)$ (given by eqn (9)) when it is occupied by a cluster ($i,j$); when accepted the cluster ($i,j$) is updated to a cluster ($i,j+1$), and the free linker is changed to a linker of type 1 contained in the new cluster.

Similarly, for a cluster ($i,j$), depending on the state of the randomly chosen neighbouring site, the hop is:

- Accepted if it is not occupied;
- Rejected/accepted with probability $1 - p(i,j)/p_0(i,j)$ (given by eqn (9)) if it is occupied by a free linker; when accepted the cluster ($i,j$) is updated to a cluster ($i,j+1$) and the free linker is changed to a linker of type 1 contained in the new cluster;
- Rejected/accepted with probability $1 - Q(i,j)/Q(i,j+1)$, when accepted the clusters ($i,j$) and ($i,j+1$) disappear by merging into a cluster ($i,j+1$).

After each hop that results in aggregation, the new cluster occupies the site to which the cluster/linker hops. Therefore each cluster/linker/particle occupies one lattice site only, and the number of occupied lattice sites decreases along the simulation.

The discrete hopping process is transformed into continuous diffusion by: (i) using the relations $H_t = 6 D_t a^2$ and $H_p = 6 D_p a^2$, where $a$ is the lattice constant; (ii) incrementing time, at each iteration, by a random number that follows a Poisson distribution with average value $1/Q$. The values of $N_0$, $m_i$, and $Q$ are updated as a result of the aggregation that follows from the encounter of the diffusive linkers and clusters. Notice that $H_t/H_p = D_t/D_p = A$.

Simulations were performed on a lattice of size $N_{latt} = 25^3$, with $N_p = \rho N_{latt} = 1562$ particles; $10^4$ samples were run for each pair $(A, \phi)$. For $A = 10$ and some values of $\phi$, runs were performed also on lattices of sizes $N_{latt} = 16^3$ and $32^3$ at the same density $\rho$, to check for finite size effects that turned out to be negligible.

C. Scaling hypothesis for the cluster size distribution

In this section we show that it is possible to approximate the full Smoluchowski description of eqn (13) by an analytically soluble equation, if a scaling hypothesis for $m_i$ is adopted. As demonstrated below, this scaling hypothesis is equivalent to assuming that, at any given instant, the fraction $j_{i,0}$ of occupied sites of a cluster ($i,j$) is independent of the size $i$ of the cluster and equals the ratio between the total number of occupied sites (or linkers 1) and the total number of sites of the clusters. Inspired by the fact that the kernel $p(i,j)$ in eqn (10) is a function of $j/w_i$, we assume the following Ansatz for the cluster size distribution $m_i$,

$$m_i(t) = m_i(0) \frac{g(j/w_i, t)}{\sum_j g(j/w_i, t)} \quad (15)$$
where \( g(x,t) \) is a time dependent scaling function. Furthermore, the moments of this function are calculated by replacing the sum by an integral,

\[
\sum_{j=0}^{w_i} j^k g(j/w_i, t) = w_i^{k+1} \int_0^{w_i} x^k g(x, t) \, dx.
\]  

(16)

A simple approximation for the time evolution of the cluster size distribution \( m_i(t) \) is then obtained by using eqn (15) and (16) in eqn (12),

\[
m_i(t) = \frac{2k_p}{V} q(1 - q) \left( \sum_{j=1}^{\infty} \frac{m_i m_{i-1}}{w_i} j \right),
\]

(17)

where,

\[
q = \int_0^{w_i} g(x, t) \, dx / \int_0^{w_i} g(x, t) \, dx
\]

(18)

Likewise, the time evolution of the free linkers is obtained by substituting eqn (15) and (16) in eqn (12),

\[
N_0 = -\frac{2k_p}{V} 2N_0(1 - q) \sum_{i=1}^{\infty} m_i.
\]

(19)

Notice that the quantity \( q \) has a simple physical meaning: the number of linkers in state 1 is \( N_1 = \sum_{i=1}^{\infty} j m_{ij} \), which may be written as,

\[
N_1 = q \sum_{i=1}^{\infty} w_i m_i.
\]

(20)

i.e., \( q \) is the ratio of the actual number of linkers in state 1 and its maximum value for a cluster size distribution \( m_i \). Moreover, if eqn (15) and (16) are used to replace the numerator in eqn (18), we obtain,

\[
q = \sum_{i=1}^{\infty} w_i m_{ij} / \sum_{i=1}^{\infty} w_i m_i
\]

(21)

which means that within the approximation the fraction of the linkers 1 bonded to clusters of size \( i \) at a given instant is independent of \( i \). It is also possible to determine the time evolution of \( N_2 \) using eqn (17) (and recalling that the clusters are tree like),

\[
\dot{N}_2 \equiv -\sum_{i,j} m_{ij} = \frac{2k_p}{V} q(1 - q)(N_p - N_2)^2.
\]

(22)

The time evolution of the number of free linkers \( N_0 \) can be expressed as (performing the sums in eqn (19)),

\[
\dot{N}_0 = -\frac{2k_p}{V} (1 - q) N_0 (N_p - N_2).
\]

(23)

The time evolution of the fraction of linkers in state \( i, p_i \equiv N_i/N_0 \), is then given by,

\[
\dot{p}_0 = -2p_0(1 - q)(1 - f \phi p_2)
\]

(24)

\[
\dot{p}_2 = \frac{q(1 - q)(1 - f \phi p_2)^2}{f \phi}
\]

(25)

III. Results

The dynamics of the model is obtained through simulations at several values of \( (\Delta, \phi) \) for \( f = 6 \) using the same initial conditions (free linkers at \( t = 0 \): \( p_0(0) = 1, p_2(0) = 0 \)). The results are compared with theoretical calculations based on the approximations discussed in the previous sections. This comparison tests the validity of those approximations. Notice that in the simulations \( R_i = k_p t \) so, in the theoretical calculations \( t = (1 + \Delta)/2 \) in eqn (24). In the following we present and discuss the
asymptotic regimes of the dynamics, the time evolution of the bonding probabilities, and the time evolution of the cluster size distribution.

A. Asymptotics

The asymptotic states of the model (i.e. the values of the probabilities $p_1$ and $p_0$ when $t \to \infty$) may be found by considering the aggregation rules and the nature of the clusters. A tree like cluster with $i$ particles has, by definition, $i - 1$ linker-mediated bonds. Thus, for the model under study, the actual number of linker-mediated bonds, $N_{2r}$, must obey $N_2 < N_P$ (or $p_2 < 1/(f \phi)$). Moreover, since each particle can bond to at most $f$ linkers the number of occupied sites is $N_1 \leq fN_P$ (or $p_1 \leq 1/\phi$). In Fig. 2 we plot the asymptotic values of the bonding probabilities $p_i$ and the bonding fraction $p_0 \equiv N_3/(fN_P/2) = 2\phi p_{2r}$, as a function of $\phi$ for $A = 10$. We find three different asymptotic regimes depending on the values of $\phi$:

1. $0 < \phi < \frac{1}{f}$: In this regime the total number of linkers is less than the total number of particles. In fact, the number of linkers is so low that all the linkers will establish bonds between the particles (i.e. all linkers 0 become linkers 2). The time taken to form these bonds depends on $A$ (as discussed below) but not the final state: $p_2 = 1$ and $p_1 = p_0 = 0$, which is a fixed point of eqn (24) and (25). Due to the tree-like character of the clusters, $N_2 < N_P$ or $p_2 \leq 1/(f \phi)$, and this regime will occur when $\phi < 1/f$. The asymptotic value of the bonding probability $p_0$ grows linearly with $\phi$ up to its maximum value $2/f$. These results are independent of $A$ (for simplicity only $A = 10$ is shown in Fig. 2).

2. $\frac{1}{f} < \phi < 1 - \frac{1}{f}$: In this regime, the number of linkers is larger than the total number of particles but smaller than $N_P(1 - f)$. The number of bonds between the particles takes its maximum value $p_2 = 1/(f \phi)$ and the bonding probability $p_0 = 2f/\phi$, is maximal and independent of $\phi$. Since $N_L > N_P$ or $\phi > 1/f$, the number of type 1 linkers is non-zero $N_1 = N_L - N_P$ or $p_1 = 1 - 1/(f \phi)$ (and $p_0 = 0$). This regime corresponds to the fixed point $p_0 = 0$, $p_2 = 1/(f \phi)$ of eqn (24) and (25), and will occur when $\phi < 1 - 1/f$; since $p_1 + p_2 \leq 1/(f \phi) + 1/\phi$. The asymptotic state is also independent of $A$, and is shown in Fig. 2 for $A = 10$. The same results are obtained for any other value of $A$. The constant bonding fraction $p_0 = 2/f$ implies that in this regime there is a single cluster, which contains all the particles, in the limit $t \to \infty$.

3. $\phi > 1 - \frac{1}{f}$: In this regime the asymptotic state is determined by the exhaustion of unoccupied sites in the clusters. At $t = 0$, the number of unoccupied sites is $fN_P$; since each linker 1 occupies one site and each linker 2 occupies two sites, the fraction of unoccupied sites at any time is $1 - \phi(p_1 + p_2)$, and vanishes when $p_1 + 2p_2 = 1/\phi$. The values of $p_1$ and $p_2$ which satisfy this condition depend on $A$. This can be seen by analysing two limiting cases. When $A \gg 1$, the $N_1$ linkers, free at $t = 0$, change to linkers 1 before any bonds between the clusters are formed; at this stage there are $N_L$ linkers 1 and $fN_P - N_1$ unoccupied sites that will then start to form bonds between the particles or clusters; at the end of this process, there are $fN_P - N_1$ bonds ($p_2 = 1/\phi - 1$) and $2N_L - fN_P$ linkers 1 ($p_1 = 2 - 1/\phi$), and the aggregation stops since all the sites are occupied. In the other limit, when $A \ll 1$, when a free linker bonds to an unoccupied site a bond between two clusters is also formed between the newly occupied site and an unoccupied one; this process lasts while $N_1$ remains below its maximum value $N_P$ ($p_1 = 1/(f \phi)$). Thereafter the remaining free linkers $N_L - N_P$ will bond with the $(f - 2)N_P$ still unoccupied sites; this second process ends when $N_1 = (f - 2)N_P$ (or $p_1 = (1 - 2/f)/\phi$), as all the sites are occupied. Notice that at this stage a number of linkers are still free as $N_0 = N_L - (f - 1)N_P$ (or $p_0 = 1 - (1 - 1/f)/\phi$) is non-zero. The asymptotic state for finite values of $A$ lies between these limits. This regime corresponds to the fixed point $q = 1$ of eqn (24) and (25). In Fig. 2 and 3(a) we show the asymptotic values of $p_i$ for $A = 10$ and $1 - 1/f < \phi < 1/f$, obtained from the simulations and the theory, which are in quantitative agreement. Fig. 3(b)-(d) illustrate this for other values of $A$ and reveal the same excellent agreement between the simulation and the theoretical results.

We conclude that the asymptotic numbers of bonds and clusters depend on the ratio of the diffusion coefficients $A$ only when $N_L > (f - 1)N_P$. Faster free linkers (larger $A$) promote the depletion of unoccupied sites decreasing the formation of bonds between the clusters. For intermediate values of $N_L$, i.e. $1/f < \phi < 1 - 1/f$, the model predicts the formation of a single cluster, containing all the particles, in the long time limit.

B. Time evolution of the bonding probabilities

We consider the dynamics of the aggregation at three different values of $\phi = 0.05$, $0.5$ and $0.95$ (representative of the three asymptotic regimes), and different diffusion ratios, $A = 1, 5, 10$ and $20$. We keep $f = 6$ fixed. At the two lowest values of $\phi$, the fraction of linkers of type 1, $p_1$, exhibits a maximum at early times. The value of this maximum increases and the time when it occurs decreases as $A$ increases (see Fig. 4(a) and 5(a)).
The presence of the maximum signals the change from an initial regime dominated by the formation of bonds between free linkers and unoccupied sites to a second regime where bonds between the particles form and the clusters merge. This transition is absent at $f = 0.95$ (see Fig. 6(d)): if the fraction of free linkers is sufficiently high bond formation between free linkers and unoccupied sites dominates at all times, due to the large decrease of unoccupied sites required for the formation of bonds between clusters. Fig. 4 and 5 show that, at low $f$, the time evolution of the bonding probabilities can be divided into an initial $\Delta$ dependent regime (up to $2\rho\kappa t \approx 10–20$, Fig. 4 and 5(a), (b)) and a late $\Delta$ independent regime (Fig. 4 and 5(c), (d)). By contrast, at large values of $f$ the dynamics depends on $\Delta$ at all times (see Fig. 6).

The theoretical results are validated by comparing them to the results of simulations (lines and symbols in Fig. 4–6). At low $f = 0.05$, initially, there is qualitative agreement for the time evolution of $p_1$ and $p_2$, including their dependence on $\Delta$ (Fig. 4(a) and (b)). At late times, there is agreement on the prediction that the dynamics does not depend on $\Delta$ (Fig. 4(c) and (d)). However, a quantitative discrepancy has been observed: while the theory predicts fast exponential decay to the asymptotic values, the simulation reveals a slower time evolution. This discrepancy between the simulation and the theoretical results (and also the observed in Fig. 6(c)) may be attributed to fluctuations in the distribution of the fraction of occupied sites in clusters with different sizes. Indeed, the analytical theory assumes that, at each instant, the fraction of occupied sites of a cluster is independent of its size (eqn (21)). This means that, under conditions where most of the clusters have either a few (low $f$) or many (high $f$) occupied sites, a strongly varying quantity is approximated by its average value, giving rise to the discrepancies observed in these regimes. At $f = 0.5$ (Fig. 5) the agreement between the theory and...
simulations is almost quantitative at all values of $\Delta$. Both predict a power-law decay of the evolution of the probabilities at late times, with an exponent $\approx -1$ (Fig. 5(c) and (d)). This feature results from the non-linearity of eqn (25), which for $1/f < \phi < 1 - 1/f$, and $t \to \infty$ reduces to,
\[ \dot{x} = -B(f,\phi)x^2, \] 
where $x = 1 - f\bar{p}_2$ and $B(f,\phi) = (f - 1 - f\phi)(f\phi - 1)/(f - 2)^2$. The solution of this equation yields $p_b(t) = p_b(t = \infty) = 1/(Bf\phi t)$. This power law decay was observed in simulations for other values of $\phi$ in the range $1/f < \phi < 1 - 1/f$, which strongly suggests that the present model in this range behaves as irreversible aggregation with a constant kernel.\(^{20}\)

The maximum number of bonds between particles (mediated by type 2 linkers) is reached asymptotically whenever $1/f < \phi < 1 - 1/f$ for any $\Delta$ as discussed in Section IIIA (see Fig. 2). In this regime, the system tends to form a single cluster that contains all particles. In order to investigate the conditions $(\phi,\Delta)$ under which the time required for the aggregation of this cluster is minimal, we analyse eqn (24) and (25) in the limits $t \to 0^+$ and $t \to \infty$, to obtain the behaviour of the bonding fraction $p_b$ in these limits. At $t = 0$, one has $\dot{\bar{p}}_b = 0$ and $\bar{p}_b = \phi(1 + \Delta)/f$, and therefore,
\[ \lim_{t \to 0} p_b(t) = \frac{\phi(1 + \Delta)}{f}. \] 
As a consequence, the initial growth of the clusters increases with $\Delta$ and $\phi$. The limit $t \to \infty$ may be obtained using eqn (31) as $p_b = 2f(1 - x)$ and thus,
\[ \lim_{t \to \infty} p_b(t) = \frac{2}{f} \left( 1 - \frac{1}{B(f,\phi)t} \right). \] 

The function $B(f,\phi)$ has a maximum at $\phi = 0.5$. Thus, for any $f$ and $\Delta$, the single cluster limit will be approached faster when $\phi = 0.5$. This behaviour is illustrated in Fig. 7(a), where the value of $p_b$ for $f = 6$ and $\Delta = 10$ is plotted for $\phi = 0.2, 0.5$ and 0.8, as obtained from the theory and simulations. Initially, the fastest growth is obtained for the largest $\phi$ ($\phi = 0.8$). However, the bonding fractions for $\phi = 0.8$ and $\phi = 0.5$ cross at $t \approx 5$ (in the units used in the figure) and the bonding fraction for $\phi = 0.5$ becomes the largest at later times. Fig. 7(b) represents this crossover time as a function of $\phi$ for different values of $\Delta$.

C. Cluster size distributions

In this section, we compare the results of simulations and theory for the cluster size distribution. Notice that this quantity can be measured in more realistic simulations and even in experiments.\(^{1,4}\) The total number of clusters is the zeroth moment, $M_0$, of the cluster size distribution $m_i$. It is, by definition, related to the number of linkers in state 2, $M_0 = N_p - N_s$, since the clusters are tree like. As a consequence, the mean cluster size, $\langle S \rangle = N_0/M_0$, can be calculated from $p_b$: $\langle S \rangle = 1/(1 - f\bar{p}_2)$. In Fig. 8(a) and (c), $\langle S \rangle$ is represented for $\Delta = 10, f = 6$. $\langle S \rangle$ grows with time, but the extent of this growth depends on $\phi$. At $\phi < 1/f$ it reaches the asymptotic value $\langle S \rangle = 1/(1 - f\phi)$ – see Fig. 8(a). At $\phi > 1 - 1/f$, the asymptotic value of $\langle S \rangle$ depends on $\Delta$. In both cases, the growth of large clusters is strongly limited as we observed $\langle S \rangle < 2$ at all times. By contrast, when $1/f < \phi < 1 - 1/f$, there is no finite asymptotic limit for $\langle S \rangle$, as shown in Fig. 8(c). The mean cluster size grows monotonically with time; the dependence on $\phi$ is limited to the time taken to reach a given cluster size, which has a minimum at $\phi = 0.5$ (at times larger than the crossover described in Fig. 7). The results in Fig. 8(a) at $\phi = 0.05$ and 0.95 reveal that the agreement between simulations and theory is good while those in Fig. 8(c) at $\phi = 0.5$ and 0.2 show that the theory slightly overestimates $\langle S \rangle$ at long times.

The cluster size distribution was determined using simulations for $\Delta = 10, f = 6$ at several values of $\phi$ and different times. The inverse of $M_2$, the second moment of these distributions, divided by $N_p$, is represented in Fig. 8(b) and (d), as a function
of \((1 - f p_b / 2) / (1 + f p_b / 2)\) to facilitate the comparison with the theoretical prediction. A behaviour similar to that of \((S_i)\) is observed: limited growth at \(\phi = 0.05\) and 0.95 and an asymptotic divergence at \(\phi = 0.2\) and 0.5. The theory overestimates \(M_i\) in all cases except at \(\phi = 0.5\).

The fraction of clusters of a given size \(i\), \(m_i/M_0\), is represented in Fig. 9 at several values of \(\phi\) and three different times, corresponding to \(p_b \approx 0.5p_b(\infty), p_b \approx 0.75p_b(\infty)\) and \(p_b \approx p_b(\infty)\). The theoretical results are obtained using eqn (30) for the fraction of clusters of size \(i\),

\[
m_i = \frac{1 - f p_b}{2} \left(1 + f p_b/2\right)^{-1},
\]

where \(p_b = 2\phi p_2\). At the initial stage of growth, the cluster size distributions are exponential for every \(\phi\) and the agreement between eqn (34) and the results of simulation is excellent. As time progresses, we find that this distribution is still exponential for large \(i\), at \(\phi = 0.05\) and \(\phi = 0.95\) (see Fig. 9(a) and (b)), but the agreement between theory and simulations is poor. The theory predicts an exponential distribution for all sizes and largely underestimates the frequency of large clusters. These differences are compatible with the results reported in Fig. 8(a) and (b); theory and simulation agree for the number of small clusters and for \((S_i)\) (which is dominated by the contribution of the more abundant small clusters); on the other hand, as large clusters have a more significant contribution to \(M_i\), the broader distributions observed in the simulations yield a larger \(M_i\) than that predicted by the theory.

There is a remarkable agreement between the cluster size distributions obtained by simulations and theory at \(\phi = 0.5\) (see Fig. 9(c)). At this value of \(\phi\) the asymptotic value of \(p_b\) is \(2f/\phi\), and thus the distribution eqn (34) becomes a very slow varying function of \(i\) for values of \(p_b\) close to \(2f/\phi\). Both theory and simulation exhibit a very broad cluster size distribution (almost uniform in the simulations) at the highest value of \(p_b\) in Fig. 9(c). To test if this agreement is obtained at other values of \(\phi \in [1/f; 1 - 1/f]\), the results at \(\phi = 0.2\) are shown in Fig. 9(d).

Exponential distributions are obtained at low and intermediate values of \(p_b\) and for large cluster sizes only when \(p_b\) is close to \(2f/\phi\). The theory underestimates both the number of very small and very large clusters. Still, the broadness of \(m_i/M_0\) is reached asymptotically in both theory and simulations.

In summary, both the theory and the simulations predict two different behaviours for the cluster size distribution: (i) at high and low \(\phi\), the distribution will be dominated by very small clusters and the growth of large clusters is rather limited; (ii) at intermediate values of \(\phi\) the distribution will evolve to almost uniform, with a large number of small and large clusters alike.

### D. Comparison with experiments

The theoretical framework developed here can be tested by comparison with experiments reported in ref. 4, where the assembly kinetics of dilute binary mixtures of streptavidin-coated large particles (\(P\)) and biotin-coated small ones \((L)\) was studied using Dynamic Light Scattering. From these measurements the time evolution of the number of clusters and the number of linkers \((i.e. m(t)\) for different values of \(i\) and \(N_0(t)\)) was obtained. The number of particles, \(N_P\), was fixed and samples with different number of linkers, \(N_L\), were investigated. A combination of geometrical parameters with the analysis of experimental results led to the estimate \(f = 65\) as the maximum number of linkers that can bond to a particle. The diffusion coefficients of particles and linkers were measured in an independent experiment and \(D_P/D_L \approx 5\) was found.

The analysis of the experimental results for \(m_1(t)\) (Fig. 5 of ref. 4) shows that, in the initial stages of growth, the rate at which monomeric particles disappear has a non-monotonic dependence on \(N_L\); at a given instant, \(|m_1|\) was found to be maximal for an intermediate value of \(N_L\). The prediction of our theory for \(m_1(t)\) is obtained by solving eqn (17) for \(i = 1\),

\[
m_1 = -2m_1 q_1(1 - q_1)(1 - f b p_2),
\]

together with eqn (24) and (25), using the experimental parameters \(\lambda = 5\) and \(R_P/R_L = 1/\lambda\) in (24)), \(f = 65\), and \(f b = N_L/N_P\). Fig. 10(a) shows the results of this calculation for the experimental values \(N_L/N_P = 2, 80\), and \(200\): the faster variation is obtained first for \(N_L = 200\) and then for \(N_L = 80\). This non trivial behaviour is elucidated in Fig. 10(b): \(|m_1|\) is represented as a function of \(N_L/N_P\) at three different times in the initial stages of growth \((2\rho k_B t = 0.1, 0.2\) and 0.3\) and is in line with the non-monotonic experimental observation depicted in Fig. 5(b) of ref. 4.

Notice that the numerical simulations reported in ref. 4 were not able to reproduce the behaviour depicted in Fig. 10, since they were limited by technical reasons to extremely short times. By contrast, in the theory described here, all the parameters (time included) can be varied freely, providing a simple tool for interpreting and predicting experimental results.

In particular the theory describes the long time behaviour reported in ref. 4. The asymptotic regime \((\text{limit } t \to \infty)\) of the kinetics was analysed by calculating the fraction of linkers bonded to particles (using the measurement of the number of
developing a generalized Smoluchowski theoretical framework in which linkers and particles are explicitly taken into account. Smoluchowski equations were solved using simulations and an approximated theory obtained via a scaling argument. Simulations have shown that the dynamics and the cluster size distributions are determined by \( \phi \) (the ratio between the actual number of linkers and the maximum number of linkers to which particles can bond or total number of bonding sites) and by \( A \) (the ratio between the diffusion coefficients of linkers and clusters) in a non-trivial way. In particular, \( \phi \) alone determines the asymptotic state of the system when \( \phi < 1 - 1/f \), while for \( 1 > \phi > 1 - 1/f \) (i.e. for a large number of linkers) the asymptotic state will depend also on \( A \). The cluster size distributions obtained at both low \( \phi \) and high \( \phi \) are exponential and exhibit finite and small moments at all times. On the other hand, for \( \phi \in [1/f; 1 - 1/f] \) the distribution becomes very broad asymptotically, with a diverging second moment. In this interval, the initial growth of the clusters is faster at larger values of \( \phi \) and \( A \); however, after a crossover time that depends on \( A \), the system with \( \phi = 0.5 \) exhibits always the fastest growth and the largest clusters (at a given time).

The results of the approximate analytical theory – eqn (24) and (25) – are in line qualitatively (and in some cases quantitatively) with the simulation results. This comparison tests the validity of the assumption that the fraction of unoccupied sites of a cluster is independent of its size. This hypothesis works quantitatively at the initial stages of growth but fails drastically at low \( \phi \) and high \( \phi \) \( \phi \) by failing to describe correctly the cluster size distributions at long times. This means that, at least at low and high \( \phi \), there is a relevant non-trivial time evolution of the dependence on the cluster size of the fraction of unoccupied sites that is not captured by the analytical theory. The theory makes the strong prediction that the cluster size distribution (34) depends on time only through the bonding probability \( p_b \), and the comparison with the simulations shows that this is valid for values of \( \phi \in [1/f; 1 - 1/f] \). Any extension of this theory would have to include a explicit \( \phi \) and cluster size dependence on the fraction of occupied sites.

The cluster size distribution (34) of the linker-mediated aggregation is equal to that obtained for a constant kernel, and therefore the model proposed in this work, with dynamics determined by the kernel defined in eqn (7) and (8), turns out to be similar to the model of irreversible aggregation with a constant kernel. In particular, the absence of percolation (i.e. the formation of a giant cluster in a finite time) and the asymptotic broad distribution at large times reinforce the similarity with the linker-mediated aggregation model in the range \( \phi \in [1/f; 1 - 1/f] \).

We found that the analytical theory explains some experimental results of ref. 4. In particular, the simplicity of the theory allowed to vary both \( f_0 = N_L/N_P \) and \( f \), enabling the exploration of length and time scales that are not accessible through detailed numerical simulations.

We focused on the study of limited linker-particle aggregation with two diffusion time scales. One could extend this study...
by adding more realistic ingredients to the theory such as: (i) clusters with more general structures (not just tree like), and even loop formation, since it seems to be particularly important, at least in equilibrium conditions, when the linkers are polymer like;\(^27\) (ii) other kernels, namely a generalized polymerization kernel, which may be amenable to analytical treatment and compared with the approximation developed in ref. 11; (iii) correlations in the distribution of unoccupied sites, etc. However, our study shows that this type of theories, regardless of the strong simplifying assumptions, do contribute to a deeper understanding of the dynamics of linker-mediated aggregation.

Conflicts of interest

There are no conflicts to declare.

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