Microstructure evolution in magnetorheological suspensions governed by Mason number

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The spatiotemporal evolution of field-induced structures in very dilute polarizable colloidal suspensions subject to rotating magnetic fields has been experimentally studied using video microscopy. We found that there is a crossover Mason number (ratio of viscous to magnetic forces) above which the rotation of the field prevents the particle aggregation to form chains. Therefore, at these high Mason numbers, more isotropic clusters and isolated particles appear. The same behavior was also found in recent scattering dichroism experiments developed in more concentrated suspensions, which seems to indicate that the dynamics does not depend on the volume fraction. Scattering dichroism experiments have been used to study the role played by the volume fraction in suspensions with low concentration. As expected, we found that the crossover Mason number does not depend on the volume fraction. Brownian particle dynamics simulations are also reported, showing good agreement with the experiments.

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I. INTRODUCTION

Magnetorheological (MR) suspensions are essentially suspensions of magnetizable particles immersed in a nonmagnetic fluid. These complex fluids show a unique ability to undergo rapid, nearly completely reversible, significant netic fluid. These complex fluids show a unique ability to

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biomedical applications

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ices such as controllable shock absorbers, electromagnetic

clutches and brakes, control valves, and artificial joints [2,3],
as well as other microfluidic devices which involve lower
concentrate magnetorheological suspensions especially for
biomedical applications [4,5]. On the other hand, and from a
more fundamental point of view, the relationship between
microscopic structure and mechanical and optical properties
of these systems has excited considerable interest in the last
decade [6–8].

Most of the studies to date have focused on the optical
response of MR suspensions to unidirectional magnetic fields
[8–13]. However, in the past years their dynamical response
to rotating magnetic fields has been a new research area of
interest. Indeed, the rotating field configuration closely
resembles the simple shear configuration appearing in most of
the practical applications of MR fluids. Moreover, the rotating
field experiments have a much simpler laboratory imple-
mentation than a simple shear experiment, which requires
precise mechanical motion. Thus, pioneering experimental
studies were reported on magnetic holes (nonmagnetic mi-
crospheres in a ferrofluid) [14,15], magnetic microdroplets
[16], and MR suspensions [17–20] subject to rotating mag-
netic fields. These systems show a rich dynamics depending
on the value of the rotating field frequency. The response of
a pair of magnetic particles immersed within a fluid was first
experimentally reported by Kashevsky and Novikova [17].
Later on, some theoretical works described the different ro-
tation modes of systems formed by few magnetic particles
under rotating fields [18]. At very high rotating field frequen-
cies, these modes became extremely complicated and chaotic
states were found. Colloidal suspensions subjected to high-
frequency rotating fields have been recently studied [21–25].
They reported aggregation of particles into two-dimensional
sheetlike structures aligned in the field plane.

The dynamics of semidilute MR fluids (volume fraction
\(\phi \approx 0.02\)) under rotating magnetic fields has been recently
studied using scattering dichroism [26,27]. In these suspen-
sions, dichroism is caused by the polarization-dependent
scattering from oriented aggregates [28] and, provided that
there is no lateral aggregation, the scattering dichroism is
proportional to the total number of aggregated particles, \(N_a\)
[27]. We found that the field-induced chains rotate synchro-
nously with the field but lag behind by a frequency-
dependent phase angle. Qualitatively similar results were ob-
tained in Ref. [29] in a system of magnetic microdroplets.
Furthermore, a more relevant result arises from Ref. [27]: the
Mason number, \(Ma\) (ratio of viscous to magnetic forces),
governs the chain dynamics under rotating fields. This di-

mensionless parameter has been defined with different pro-
portionality factors in the literature [24,30,31]. Here we use
the following definition:

\[
Ma = \frac{12 \eta_\alpha \nu}{\mu_0 \mu_s M^2},
\]

where \(\eta\) is the solvent viscosity, \(\nu\) is the rotating field fre-

quency, \(\mu_0\) and \(\mu_s\) are the vacuum and solvent magnetic
permeability, respectively, and \( M \) is the particle magnetization. Here the proportionality factor has been chosen to be in agreement with the dimensionless frequency obtained from the simulations analysis (Sec. III). A change in behavior of the dichroism and the phase lag was found above a crossover Mason number \( Ma \sim 1 \), where the viscous forces dominate and inhibit the aggregation process.

In this paper, we report on an experimental video microscopy study of the aggregation of magnetizable particles and the subsequent chain orientation dynamics under rotating magnetic fields. This technique allows us to directly visualize the dynamics of very dilute suspensions (\( \phi \sim 10^{-6} \)). Note that the scattering dichroism technique will not be useful to study very dilute suspensions \( \phi < 0.001 \) due to the small dichroism signal obtained. We used different rotating frequencies which correspond to Mason numbers below and above 1, i.e., around the value where both magnetic and hydrodynamic interactions are comparable. This range of Mason numbers is similar to the one used in our previous scattering dichroism experiments in more concentrated suspensions [26,27]. We find that the average size of the aggregates decreases with \( Ma \). Furthermore, we find a transition in the total number of aggregated particles at a crossover Mason number close to \( Ma \sim 1 \), in agreement with our previous work. Therefore, this result seems to indicate that the dynamics does not depend on the volume fraction, at least between \( \phi \sim 10^{-4} \) and \( \phi = 0.02 \). In order to verify this result, we analyze the role played by the volume fraction in the dynamics using scattering dichroism experiments. As expected, we found that the crossover Mason number does not depend on the volume fraction, which points out the generality of this behavior. Our experimental findings have been corroborated through Brownian particle dynamics simulations of “hard” spheres with induced dipolar interactions and Stokes friction against the solvent. The simulations are in agreement with the experiments. We also developed numerical simulations to analyze the thermal effect on the dynamics.

II. EXPERIMENTS

Two experimental setups were used to study the dynamics governing MR suspensions subject to rotating magnetic fields. First, we performed video microscopy experiments on very dilute suspensions (\( \phi \sim 10^{-6} \)) to directly visualize their dynamics. Second, to study the role played by the volume fraction, scattering dichroism technique was needed since it allows the use of suspensions with moderate concentration. So we performed scattering dichroism experiments on suspensions with different volume fractions ranging from \( \phi = 0.001 \) to \( \phi = 0.02 \).

A. Experimental materials and procedure

1. Magnetic suspensions

To prepare our samples we used an aqueous suspension M1-070/60 of super-paramagnetic microspheres supplied by Estapor with a solid content of 10% in weight. The particles in the suspension contain magnetite crystals (\( \text{Fe}_3\text{O}_4 \)) of small diameter (1–20 nm) dispersed in a polymeric matrix.

Table I. Properties of the magnetizable microspheres.

<table>
<thead>
<tr>
<th>Particle properties</th>
<th>M1-070/60</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean diameter (( \mu m ))</td>
<td>1.24</td>
</tr>
<tr>
<td>Magnetic content</td>
<td>61%</td>
</tr>
<tr>
<td>Saturation field (emu/g)</td>
<td>51.9</td>
</tr>
<tr>
<td>Surface group content (( \mu \text{eq/g} ))</td>
<td>117</td>
</tr>
</tbody>
</table>

The surface of the microspheres contains carboxylic acid (-COOH) groups with an added surfactant coating layer of sodium dodecyl sulfate (SDS) to stabilize the dispersions. The particles’ physical properties are detailed in Table I. We have characterized the average magnetic properties of the particles by measuring their magnetization curve using a vibrating sample magnetometer (VMS-Lakeshore 7300). We observed that under sufficiently low magnetic fields these particles exhibit super-paramagnetic behavior with virtually no hysteresis or magnetic remanence. Due to their small average diameter and density (\( \rho \sim 1.3 \text{ g/ml} \)) the sedimentation time is long enough to neglect gravitational effects. The sedimentation velocity was estimated according to the Stokes’ law to be \( v_s \sim 0.06 \mu \text{m/s} \).

For video microscopy experiments we diluted the suspension M1-070/60 with Milli-Q ultrapure water with the same SDS concentration as the original suspension in order to avoid particle aggregation. The volume fraction achieved is \( \phi \sim 10^{-4} \). To analyze the volume fraction effect through scattering dichroism experiments, we prepared suspensions with 50% glycerol and different volume fractions ranging from \( \phi = 0.02 \) to \( \phi = 10^{-3} \).

2. Magnetic field generation and sample cell

The video microscopy experiments requires optical transparency, so the fluid sample was sandwiched between two circular quartz windows with inner diameter 6.5 mm. These windows were held in place by a delrin attachment designed to prevent evaporation of the solvent and separated by an annular delrin spacer \( e = 100 \mu \text{m} \) thick along the light path (axis \( Z \)). The video microscopy images were taken at an area in the cell less than 1 mm in the vicinity of the center. The sample is placed at the center of two orthogonal pairs of coils that generate a rotating magnetic field in the plane of the images (\( XY \)). The rotating magnetic field was achieved by applying sinusoidal electric signals to the two orthogonal pairs of coils by means of two Kepco BOP20-10M power amplifiers, driven by two HP-FG3325A function generators referenced to one another at a phase difference of 90°. The function generators allowed for control of both the amplitude and the frequency of the rotating magnetic field. These coils are housed in a temperature controlled aluminum cylinder to prevent heating effects. All experiments were performed at a temperature of \( T = 282 \pm 1 \text{ K} \) on the sample. The coil’s relative positions and their dimensions were optimized to obtain the smallest possible spatial variation of the field over the sample (smaller than 3% across the whole sample). This minimizes possible local changes of concentration due to
migrates of the magnetic particles under the effect of field gradients. In Table II are summarized the experimental conditions for each experiment.

3. Video microscopy setup and procedure

We illuminated the top side of the sample from the direction perpendicular to the field plane using a white light source (American Optical H-80). To amplify the image we used a (Navitar 12X) zoom system composed of a 2X adapter that attaches the zoom to a charge-coupled-device (CCD) video camera (Sanyo VDC-3825), a 12X zoom, and a 2X attachment that joins the zoom to a 10X microscope objective placed close to the bottom side of the sample. This group of lenses has the advantage of combining large zoom range, high resolution, and long working distance (33 mm). This zoom system has the capability of allowing the field of view to be altered by changing the image resolution. The resolution could be varied between 0.34 and 4.2 μm/pixel for fields of view between 170 and 2100 μm, respectively. The CCD camera is connected to a S-VHS VCR (Panasonic AG 1975) which records images at 30 frames/sec. We subsequently digitalized single frames of 640 × 480 pixels with 256 gray levels on a computer at fixed time intervals depending on the field rotational frequency. To analyze the images we used IGOR Pro image-processing software from WaveMetrics, Inc. The images had a dark background over which the particles appear as clear areas. In the image analysis the pixel area for each aggregate was calculated by applying a gray level threshold that was kept constant for a given experiment.

A typical experimental run would begin with an homogeneous sample and then we applied the rotating magnetic field for 365 sec. Although the zoom magnification was varied for each experiment to adapt the field of view, a resolution that allowed us to see individual particles was always used. Then, the temporal evolution of the total number of aggregated particles was determined. The average rotation angle of the chains was also computed by analyzing the Hough transform of the images [32,33].

4. Scattering dichroism setup and procedure

The optical train used to measure linear dichroism consists of a He-Ne laser, a polarizer (that sets the angle for the polarization), a photoelastic modulator (set at π/4 displacement with respect to the polarizer), and a quarter wave plate (set parallel to the polarizer). The laser beam goes through the sample cell and the transmitted light is detected by a photodiode. The signal from the photodiode is sent to a dc amplifier to recover the dc component and two phase lock-in amplifiers (Priceton Applied Research, EG&G-128A) set at once and twice the photoelastic modulator frequency. The outputs of both lock-ins and the dc amplifier output are then digitized using a 16-bit resolution A/D data acquisition board (National Instruments, PCI-MIO-16X-E-10). A full description of this experimental technique can be found in Refs. [26,34]. With this optical setup we can simultaneously measure the temporal evolution of the dichroism \( n''(t) = n''_1 - n''_3 \), i.e., the difference between the extinction of the incident light with polarization parallel and perpendicular to the long axis of the aggregates (being \( n_1 = n'_1 - i n''_1 \) the refractive index in the direction parallel to the long axis of the aggregates). We can also measure the temporal evolution of the orientation angle of the aggregates \( \theta'' \), i.e., the average angle difference between the reference angle of the optical train and the long axis of the aggregates. By comparing \( \theta'' \) with the temporal evolution of the magnetic field direction, given by \( \omega t' \), we may define the phase lag between the field and the aggregates as \( \alpha(t) = \omega t' - \theta''(t) \).

The measurement procedure required, first, to set the values of frequency and amplitude of the magnetic field, second, to measure the time evolution of \( \Delta n''(t) \) and \( \theta''(t) \) until a steady state is reached, \( \Delta n''_i \) and \( \theta''_i \), and third, to wait the time necessary for the particles to break apart and to return to their initial state. Therefore, all measurements are independent, because there is no continuous sweep in either field frequency or amplitude.

B. Video microscopy results

In Fig. 1 video microscopy images show the temporal evolution of the structures induced by a magnetic field rotating clockwise with frequency \( f = 0.001 \) Hz and amplitude \( H = 1.55 \) kA/m (\( M_a = 0.0012 \)). When the field is first applied \(( t = 0 \) s), a magnetic dipole is induced in each particle in the direction of the field. At this low volume fraction \(( \phi \sim 10^{-4} \) the dipolar interaction induces particle aggregation into linear chains that follow the magnetic field direction. We have seen that the chains rotate synchronously with the field as was reported in Ref. [35]. The average chain length initially increases until it finally reaches a steady state [35]. Note that the aggregation kinetics under rotating magnetic fields is very different than the kinetics reported under unidirectional magnetic fields where power-law behavior governs the long-time aggregation regime [12,13]. Under rotating magnetic fields, the final size of the aggregates is roughly determined by the competition between the magnetic dipolar interaction and the hydrodynamic forces on the chains [36]. We observe that rotating chains undergo a process of dynamic chain growth and fragmentation that has a frequency twice that of the field frequency as was reported in Ref. [35].

We would like to emphasize that lateral aggregation of chains has not been observed at the low volume fractions reported in the video microscopy experiments as occur in unidirectional magnetic fields [9]. In the case of rotating fields, we have observed partial overlapping of chains at low rotational frequencies. This phenomenon is originated by a different mechanism than the lateral aggregation of chains.
under unidirectional fields since it is due to contact of close neighboring chains during their rotation when the distance between their centers is smaller than the average length of the chains. We observe that partial overlapping of chains always occurs in the plane perpendicular to the image since in this configuration the structure formed by two overlapping chains presents smaller inertia moment and is able to easily follow the field. Two dimensional sheets of several particles thick aligned in the plane of the field were reported by Martin et al. \cite{21,25,37} for high volume fraction suspensions under high-frequency rotating fields.

In the limit of low rotational frequencies, and for this low value of the magnetic field, aggregation and fragmentation processes are continuously observed between neighboring chains. Chains approach each other and, for a short time, they keep attached together by their ends and rotate as one. The phase lag of the longer formed chain respect to the field direction increases compared to the phase lag of the shorter chains. Hydrodynamic forces on the chains cause them to break up into smaller segments in order to remain oriented with the field \cite{38}.

In Fig. 3 video microscopy images show the behavior of the structures induced for different rotational frequencies. All the images have been recorded after applying the field during $t=300 \text{ s}$. We can see that the length of the induced aggregates, $L$, decreases when increasing the frequency. The same behavior $L \propto \text{Ma}^{-1/2}$ is also obtained for a chain model in the case of rotating magnetic fields \cite{35,36}. However, for $\text{Ma} \gg \text{Ma}_c \sim 1$, this low amplitude magnetic field is not strong enough for the structures to remain aligned in the field direction. Hence, at high frequencies ($f = 1 \text{ Hz}$ in Fig. 3, which corresponds to $\text{Ma}=1.2$) even two particle chains may break apart, therefore increasing the number of isolated particles; moreover, other more isotropic structures appear, as for example disklike structures formed by 3–5 particles. Similar

![FIG. 1. Video microscopy images at different times showing the structures induced by a magnetic field rotating clockwise with frequency $f = 0.001 \text{Hz}$ and amplitude $H = 1.55 \text{kA/m}$. Field of view: $58 \times 43 \mu \text{m}^2$; suspensions with $\phi \sim 10^{-4}$.](image1.png)

![FIG. 2. Details of the S shape (reversed in this case) developed by one chain just before the breakup. The field rotates counterclockwise with frequency $f = 0.01 \text{Hz}$ ($\text{Ma}=0.012$). Field of view: $14.4 \times 14.2 \mu \text{m}^2$; suspensions with $\phi \sim 10^{-4}$.](image2.png)
structures were found for ER fluids subjected to high-frequency rotating fields [39]. Therefore the total number of aggregated particles should decrease when increasing Mason number. In Fig. 3 (on the right) we see the dependence of the normalized total number of aggregated particles with Mason number, which shows the expected behavior, in agreement with our previous work [27]. A deeper insight in the behavior of $N_a$ (see Fig. 3) reveals that the crossover Mason number seems to be shifted to values lower than 1. We should say at this point that the criteria we followed to compute $Ma_c$ is to consider the value at which a change in behavior is observed. Usually, this change is not abrupt but occurs in a range of values, therefore a precise determination of $Ma_c$ cannot be made without some ambiguity. A possible responsible mechanism for the shift observed on $Ma_c$ to smaller values is the formation of small structures displayed in the direction perpendicular to the image plane at high rotating frequencies (close to $Ma = 1$). From the image analysis, this fact leads to a decrease of the normalized total number of aggregated particles, and therefore it could explain the mentioned curve shift.

In summary, we have seen that the behavior of the very dilute MR suspensions is the same as found in our previous work on more concentrated suspensions, that is, there is a crossover Mason number ($Ma_c \sim 1$) above which the rotation of the field prevents the particle aggregation process into chains from taking place, and other more isotropic structures appear. This result seems to indicate that this scenario is basically independent of the volume fraction.

C. Scattering dichroism results

In Fig. 4 we plot the variation of the dichroism (left) and the phase lag (right) with rotational frequency for decreasing values of $\phi$. We see, in agreement with our previous works for larger $\phi$ [26, 27], that dichroism shows two distinct regions for frequencies below and above a crossover frequency $f_c$. Below this crossover frequency, the dichroism is essentially independent of frequency. However, once this frequency is surpassed, the dichroism strongly decreases with frequency following a power-law behavior with an exponent $-1$. The phase lag also show two different responses de-
transition around the crossover Mason number as the volume fraction increases is observed, which could be observed at the left part of Fig. 4. At low volume fractions the rupture of the dimers leads to the abrupt change in dichroism behavior. However, at high volume fractions the mean distance between chains can be very small, so before the dimers break apart, they have a higher probability to join together with neighboring dimers forming more isotropic structures. Therefore, the collision of dimers around the crossover Mason number may lead to a broadening of the transition. Although it is not plotted here, a nice collapse of the curves corresponding to the phase lag is also found.

III. NUMERICAL SIMULATIONS AND DISCUSSION

A. Equations of motion

A complete algorithm to simulate MR fluids would include Brownian motion, multipolar magnetic interaction forces between the particles, local field corrections to the applied field, hydrodynamic friction forces (lubrication and long range), excluded-volume repulsive forces, and so on. However, our approach has been to consider only those contributions which we believe are essential to capture the physics of the problem. Therefore, in order to interpret the experimental results, we developed two-dimensional (2D) Brownian particle simulations of “hard” spheres with induced dipolar interactions and Stokes friction against the solvent. Similar approaches have been previously used [25,40–42]. We consider a monodisperse suspension of $N$ spherical particles of diameter $2a$ suspended in a fluid of viscosity $\eta$ and subjected to a rotating magnetic field of amplitude $H$ and angular frequency $\omega$. Taking into account that the aggregation takes place in the plane of the magnetic field rotation [17], we simplified our calculation by developing 2D simulations in this plane. Two fundamental length scales characterize the formation of chains. The first one is the so-called “thermomagnetic distance” $R_1$, which is the distance at which the magnetic energy corresponding to the dipolar interaction between two particles aligned in the field direction and with parallel dipole moments equals the thermal fluctuation energy. This length scale turns out to be $R_1 = 2a\lambda^{1/3}$, where $\lambda$ is a dimensionless parameter calculated as the ratio between magnetic and thermal energies,

$$\lambda = \frac{W_m}{k_B T} = \frac{\mu_0 \mu_r m^2}{16 \pi a^3 k_B T},$$

where $m = (4 \pi/3) a^3 M$ with $M$ the particle magnetization, $k_B$ is the Boltzmann constant, and $T$ is the temperature. The physical meaning of $R_1$ is that particles that are separated by a distance larger than $R_1$ do not “feel” the magnetic interaction due to each other, because the energy of thermal fluctuations is larger than the dipolar interaction energy. The second length scale is the average initial interparticle distance, which can be estimated as $R_0 \sim 2a/\phi^{1/3}$ [13]. Actually, the values of $\lambda$ that correspond to the experimental results reported here are $\lambda = 213$ for the video microscopy experiments with very dilute suspensions and $\lambda = 75$ for the scat-
tering dichroism experiments with more concentrated suspensions. In these experiments, since $R_1$ is typically $4\leq R_1 \leq 7.5$, and $R_0$ is in the range $4\leq R_0 \leq 20$, we will include the Brownian motion of the particles on the evolution of the structures. As a first approximation, we will neglect hydrodynamic interaction, and the only solvent effect that we will consider is the viscous force represented as a Stokes force acting on each particle. We can typically neglect the inertial term in the equation of motion because the viscous drag term dominates. Therefore, the equation of motion for the $i$ particle will contain the sum of the following forces [25]:

$$
\frac{d\vec{r}_i}{dt} = \frac{1}{\gamma} \sum_{j \neq i} \vec{F}_d(\vec{r}_{ij}) + \frac{1}{\gamma} \sum_{j \neq i} \vec{F}_r(\vec{r}_{ij}) + \vec{F}_B,
$$

(3.2)

where $\vec{r}_i$ is the position of the $i$ particle, $\vec{F}_d$ is the magnetic force, $\vec{F}_r$ is the repulsive force, and $\vec{F}_B$ is the Brownian force. Moreover $\gamma = 6\pi \eta a$ is the friction coefficient and $\vec{r}_{ij}$ is the vector between the centers of mass of particles $i$ and $j$. The dipolar force over particle $i$ will be the sum of the dipole-dipole forces exerted by all of the other particles. The dipole-dipole force between particles $i$ and $j$ is given by

$$
\vec{F}_d(\vec{r}_{ij}) = \frac{3 \mu_0 \mu_r m^2}{4 \pi \vec{r}_{ij}} \left\{ \left[ 1 - 5 (\vec{m} \cdot \vec{r}_{ij})^2 \right] \vec{r}_{ij} + 2 (\vec{m} \cdot \vec{r}_{ij}) \vec{m} \right\},
$$

(3.3)

where we take $\vec{m}$ to be aligned always with the field direction. Note that as we are using one-particle Stokes’ hydrodynamics, a repulsive force $\vec{F}_r$ must be included to avoid particles from overlapping. This force is calculated from Ref. [43]:

$$
\vec{F}_r(\vec{r}_{ij}) = A \frac{3 \mu_0 \mu_r m^2}{4 \pi (2a)^4} \exp[-B(r_{ij}/(2a) - 1)] \vec{r}_{ij},
$$

(3.4)

where we set $A = 2$ and $B = 10$, so that for two particles that are in mechanical contact the repulsive force exactly balances the attractive dipolar magnetic interaction. Similar repulsive forces have been used in previous works [21,43]. The Brownian force $\vec{F}_B$ is a stochastic force with zero mean, $\langle \vec{F}_B(t) \rangle = 0$, and $\delta_t$ correlated, $\langle F_B(t) F_B(0) \rangle = 2D \delta_t$, where $D$ is the translational diffusion coefficient $D = k_B T/\gamma$. As usual [1,21,44], we can make the particle evolution equation (3.2) dimensionless, using the particle diameter $2a$ as length scale, so that $r' = r/(2a)$, and a time scale $\tau' = 12^2 \eta a/(\mu_0 \mu_r M^2)$, such that $t' = t/\tau$. This temporal scale leads to a dimensionless rotational frequency equal to $\Omega = \omega \tau$. This dimensionless frequency is the well-known Mason number with the definition used in Eq. (1.1), $\Omega = \Omega_m$.

**B. Simulation procedure**

According to the experimental analysis, we have used only two control parameters, the Mason number $M_a$ and the particle volume fraction $\phi$. The simulations being 2D, we have set the initial average interparticle distance so that it corresponds to the volume fraction in the experiment with the different suspensions.

The numerical integrations have been performed using a time step of 0.005, which has proved to be small enough to avoid significant overlapping errors when the particles come into close contact. We have developed simulations using 400 and 100 particles, and the simulations are carried out in a square box of the proper size to adjust the initial average interparticle distance. Periodic boundary conditions were applied. The particles are set initially at random positions avoiding overlapping, and all of the results reported here are statistical averages over different realizations corresponding to different initial spatial distributions of the particles, typically ten runs. Each simulation is run long enough for the steady state to be reached, and then the quantities to be compared with the experimental results are averaged during the last period of rotation of the field.

The comparison between experimental and simulation results deserves some comments. First, the chains are recognized by means of a criterion according to which two particles are aggregated when the distance between their respective centers of mass is smaller than 1.1 times the particle diameter. Second, using Mie’s theory for light scattering by nonspherical objects, we can estimate the scattering dichroism generated from a chain $j$ formed by $N_j$ particles in the forward direction ($\varphi = 0$). Actually the scattering dichroism is proportional to the number of particles in the chain, $N_j$. Assuming that the scattering dichroism produced from a set of chains is the incoherent sum of the scattering dichroism produced by each chain, the total dichroism is proportional to the total number of aggregated particles, $N_a$. Hence, we will compare the experimental value of the scattering dichroism with the value of $N_a$ obtained in the simulations computed as

$$
N_a = \sum_j N_j,
$$

(3.5)

where the sum is performed for $j$ such that $N_j > 1$. We have computed the dimensionless average chain length (average number of particles in a chain) $L$ in the usual way, i.e., if a chain labeled as $j$ is formed by $N_j$ particles, then

$$
L = \frac{\sum_j N_j}{\sum_j 1}.
$$

(3.6)

The phase lag of a given chain ($\alpha_j$) is the angular difference between the orientation of the magnetic field and the unit vector parallel to the long axis of the chain. For quasi-linear chains, we have determined the chain orientation vector by calculating the eigenvalues of the chains’ inertia tensor, $I^\text{max}_j, I^\text{min}_j$. Then, the eigenvector corresponding to $I^\text{min}_j$ gives the orientation of the long axis of the chain.

The computations described so far consider straight chains and, obviously, do not take into account the shape of
FIG. 6. Particle position in the XY plane at an arbitrary time for different dimensionless rotating frequencies or Mason numbers.

the clusters. However, video microscopy results show that very long chains deviate significantly from straight lines, and, moreover, at high frequencies smaller and more isotropic clusters may appear. To assess the effect of these chain form anomalies in our simulations results, we calculated the same quantities, \( L \) and \( N_a \), using a weight function that takes into account the shape of the cluster. Actually we used \( W_j = N_j s_j \), where \( s_j \) is a shape factor with value 1 for the case of a straight chain, and 0 for a symmetric cluster,

\[
s_j = \frac{(I_j^{\text{max}})^{1/2} - (I_j^{\text{min}})^{1/2}}{(I_j^{\text{max}})^{1/2} + (I_j^{\text{min}})^{1/2}}.
\]

(3.7)

We did not observe appreciable changes in the simulations results using this shape correction. Therefore, we present here the results corresponding to the case without the shape factor.

C. Simulation results

1. Very dilute suspension

We have studied the dynamics of very dilute suspensions with the conditions of the microscopy experiments, i.e., volume fraction of particles \( \phi = 10^{-4} \) which corresponds to an initial average separation between particles equal to \( R_0/(2a) \sim 16 \). The particle diameter is \( 2a = 1.24 \mu \text{m} \), the magnetization is \( M = 2.82 \text{ kA/m} \) (for \( H = 1.55 \text{ kA/m} \)), and the viscosity of the solvent (water, in this case) at \( 10 \degree \text{C} \) is \( \eta = 1.307 \times 10^{-3} \text{ Pa.s} \). Using these values we obtain a time scale equal to \( \tau = 0.019 \text{ s} \) and \( \lambda = 213 \). We numerically solved Eq. (3.2) for a system with \( N = 400 \) magnetic particles. For each dimensionless rotational frequency \( \Omega \) or Mason number \( \text{Ma} \), we calculate the total number of aggregated particles, \( N_a \), and the average length \( L \).

In Fig. 6 we plot the particle positions in the \((X,Y)\) plane for different dimensionless frequencies at an arbitrary time. Several qualitative observations can be made in Fig. 6. First, the size of the structures becomes smaller as the rotational frequency increases. Second, at high \( \text{Ma} \), the tendency to form chains aligned with the field is not clearly apparent, and more isotropic clusters appear to be preferred; moreover, a higher number of isolated particles appears. We plot in Fig. 7 the distribution of lengths \( N_j \) and phase lags \( \alpha_j \) corresponding to the images shown in Fig. 6 at two different Mason numbers. At low Mason numbers \( \text{Ma} = 0.3 \), the distribution of \( N_j \) presents a maximum placed around \( N_j \sim 3 \) while the phase lag follows a narrow distribution around \( \alpha_j \sim 5 \degree \). On the other hand, at high Mason numbers \( \text{Ma} = 1.6 \), the maximum of the distribution of \( N_j \) is set at \( N_j = 1 \), which means that there are many isolated particles in the suspension, being responsible to the decrease of the total number of aggregated particles, \( N_a \). At this high Mason number, a broader distribution of \( \alpha_j \) appears, and the maximum shifts to higher values.

Some of these observations can be made more quantitative by studying the average chain length. The dimensionless average chain length \( \bar{L} \) versus \( \text{Ma} \) in a log-log plot is shown in Fig. 8(a). The average chain length follows a power-law behavior with an exponent approximately equal to \(-0.5\). This behavior agrees well with the one predicted by the chain model developed for ER fluids subject to steady shear flows [38], and for MR suspensions subject to rotating fields [36]. Figure 8(b) shows the total number of aggregated particles, \( N_a \), versus \( \text{Ma} \) in a log-log plot. Two different regions appear, in agreement with the experimental results. First, there is a plateau zone at low \( \text{Ma} \), in which \( N_a \) varies very little with \( \text{Ma} \). Second, a strongly decreasing response appears for \( \text{Ma} \gtrsim 1 \), with an apparent exponent close to \(-1\), although the region in which this power-law behavior could be studied in the simulations is rather small.

The picture that emerges from these results is the following: the average chain length decreases monotonously with

FIG. 7. Distribution of lengths \( N_j \) (left) and phase lags \( \alpha_j \) (right) at the same arbitrary time of Fig. 6 at a Mason number \( \text{Ma} = 1.6 \) (solid line) and \( \text{Ma} = 0.3 \) (dotted line).

FIG. 8. (a) Average length \( L \) and (b) normalized total number of aggregated particles \( N_a / N \) vs Mason number. Suspension with \( \phi \sim 10^{-4} \).
The purpose of this work was to study the spatiotemporal evolution of the induced structures in very dilute polarizable...
colloidal suspensions subject to rotating magnetic fields by means of video microscopy experiments. This technique allows us to directly visualize the dynamics and needs of very dilute suspensions ($\phi \sim 10^{-4}$). The average chain length decreases when increasing frequency. We also found that there is a crossover Mason number above which the rotation of the field prevents the particle aggregation to form clusters. Therefore, at these high Mason numbers, more isotropic clusters and isolated particles appear. The same behavior was also found in previous scattering dichroism experiments on more concentrated suspensions.

In order to study the role played by the volume fraction, we carried out scattering dichroism experiments in suspensions with low concentration ($\phi \sim 0.001–0.02$). Dichroism shows two distinct regions, above and below $M_a$. The crossover Mason number does not depend on the volume fraction which points out the generality of this behavior. We can conclude that the results obtained here (the Mason number governing the dynamics of chain rotation, the limits to the rotational dynamics come from both the mechanical instability of the chains due to shear and the formation of more isotropic structures) can be extrapolated to higher volume fractions, provided they are below a threshold volume fraction above which sheetlike structures will form [21].

Thermal particle dynamics simulations are also reported, showing good agreement with the experiments. We theoretically analyzed the thermal effect on the dynamics observing a decrease of the crossover Mason number for $\lambda < 50$.

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