Flow behavior of colloidal rod-like viruses in the nematic phase

M. Paul Lettinga,1 Zvonimir Dogic,2 Hao Wang,3 and Jan Vermant4

1IFF, Institut Weiche Materie, Forschungszentrum Jülich, D-52425 Jülich, Germany
2Rowland Institute at Harvard, Harvard University, Cambridge MA 02142, USA
3Department of Physics, North Dakota State University, Fargo, ND 58105, USA
4Department of Chemical Engineering, Katholieke Universiteit Leuven, de Croylaan 46, B-3001 Leuven, Belgium

The behavior of a colloidal suspension of rod-like fd viruses in the nematic phase, subjected to steady state and transient shear flows is studied. The monodisperse nature of these rods combined with relatively small textural contribution to the overall stress make this a suitable model system to investigate the effects of flow on the non-equilibrium phase diagram. Transient rheological experiments are used to determine the critical shear rates at which director tumbling, wagging and flow-aligning occurs. The present model system enables us to study the effect of rod concentration on these transitions. The results are in quantitatively agreement with the Doi-Edwards-Hess model. Moreover, we observe that there is a strong connection between the dynamic transitions and structure formation, which is not incorporated in theory.

I. INTRODUCTION

When subjected to shear flow liquid crystals can exhibit a variety of surprising phenomena, which arise because of the anisotropic shape of the constituent rods. Theoretically the behavior of a suspension of hard rods during shear flow can be described by the equation that governs the time development of their probability distribution function, as derived by Hess16 and by Doi and Edwards9. In the absence of a flow, the Doi-Edwards-Hess (DEH) theory reduces to the Onsager description of equilibrium nematic liquid crystals and can be used to describe the isotropic-nematic (I-N) phase transition of a hard rod suspension31. The rheological properties are predicted to be highly non-linear functions of the Péclet number (Pe), which is the ratio of shear rate $\dot{\gamma}$ over rotational diffusion constant $D_{\text{r}}$. This is not surprising as the Pe number can be much larger than unity when the rod-like molecules have large aspect ratios.

The nonlinear response of the rheological properties indicates that the shear flow distorts the equilibrium distribution of macromolecules or rods. The spatiotemporal microstructural changes during flow are even more complex. At low shear rates, the DEH theory predicts that the pseudo vector describing the average alignment of the rods, i.e. the “director”, undergoes a continuous “tumbling” motion in the plane defined by the velocity and the velocity gradient vectors. At high shear rates the director is predicted to align with the flow22,28. At intermediate shear rates it is possible to obtain multiple solutions to the Doi-Edwards-Hess equation, which are dependent on the initial orientation of the director10,12. For one stable solution called “wagging” the nematic director oscillates between two angles in the plane defined by the flow and the gradient of the flow. Other solutions such as kayaking and log-rolling are also possible, in which the director oscillates out of the flow-gradient plane at these intermediate shear rates15.

Experiments on polymeric liquid crystals have confirmed several predictions of the Doi-Edwards equation. Using a combination of rheological and rheo-optical measurements it was shown that nematic solutions of poly(benzyl-glutamate) tumble (PBG) at low shear rate and become flow aligning at high shear rates23. The existence of a wagging regime and a potential coexistence of wagging and log-rolling regimes at intermediate flow rates have also been revealed in experiments15,29. However, there remain significant difficulties when comparing experiments on polymeric liquid crystals (PLC) to theoretical predictions. One problem is that different levels of the microstructure may lead to different contributions to the stress tensor1. In addition to the molecular contribution to the stress tensor, textural aspects contribute. The latter include Frank elasticity contributions due the presence of spatial distortions of nematic director, and viscous interactions between ‘domains’. In addition, there is an indirect effect to the stress tensor as the defects disturb the orientation distribution function. These textural contributions to the total stress dominates the behavior at high concentrations and low shear rates29, making it difficult to accurately extract information about the concentration dependence of different flow transitions. The textural portion of the stress typically displays scaling of the transient rheological response with strain rather than with Pe number30. The strain scaling is a typical feature of materials where the time response is determined by an inherent length scale which in the case of PLC’s is set by the size of the large non-Brownian nematic domains24.

The DEH theory describes the flow behavior of a homogenous ensemble of rods but does not consider any polydomain effects. Therefore an ideal system for testing DEH theory should have small textural contributions. In this paper we use rod-like fd virus suspensions to access the concentration dependence of the transition of tumbling to wagging, and wagging to flow aligning. We show that the contribution of textural stress is very low, although the spatial distribution of directors still has to be accounted for. The main motive for using fd virus is the thorough understanding of its equilibrium behavior, which has been quantitatively described us-
ing the Onsager theory extended to take into account the semi-flexible nature of $fd$ as well as its surface charge $32$. Moreover, $fd$ has already successfully been used for (micro-)rheology experiments in the isotropic phase $14, 34$. The aim of the present paper is to make a comparison between the dynamic flow behavior of $fd$ suspensions and the available microscopic theoretical predictions of the DEH theory for a homogeneous system of colloidal rods under shear.

The paper is organized as follows. In section II we discuss the equation of motion of the orientational distribution function and the numerical method we use to solve it. The experimental details about sample preparation and measurements are given in section III. The results are discussed in five parts: the stationary viscosity of $fd$ suspensions, the concentration and shear rate dependence of the oscillatory response to a flow reversal, the relaxation after cessation of flow at high concentration and in situ microscopy under shear. In section IV the textural contribution to the stress tensor is investigated more detail. Finally we present a non-equilibrium phase diagram of shear and concentration dependence of different flow transitions.

II. THEORY

The distribution of an ensemble of rods can be described by the probability density function $P(\hat{1}_{1}, ..., \hat{1}_{n}, \vec{r}_{1}, ..., \vec{r}_{n})$ of the positions $\{\vec{r}_{i}\}$ and orientations $\{\hat{1}_{i}\}$ of the rods. Ignoring any spatial correlations, i.e. restricting to a monodomain, we have $P(\hat{1}_{1}, ..., \hat{1}_{n}, \vec{r}_{1}, ..., \vec{r}_{n}) = \tilde{\rho}P(\hat{1}_{1}, ..., \hat{1}_{n})$, where $\tilde{\rho} = N/V$ is the particle density. Therefore, the orientational probability density function, or orientational distribution function (ODF), fully characterizes the system. The time evolution of the ODF for a suspension of rods during flow is obtained by solving the equation of motion for the ODF, given by the $N$-particle Smoluchowski equation.

$$\frac{\partial P(\hat{1}, t)}{\partial t} = D_r \hat{\mathcal{R}} \cdot \left\{ \hat{\mathcal{R}}P(\hat{1}, t) + DL^2 \tilde{\rho}P(\hat{1}, t) \right\} \int d\hat{1}' P(\hat{1}', t)|\hat{1}' \times \hat{1}|$$

$$-\hat{\mathcal{R}} \cdot P(\hat{1}, t)\hat{1} \times (\hat{\Gamma} \cdot \hat{1})$$

(1)

where $\hat{\mathcal{R}}(\vec{r}) = \hat{1} \times \nabla_{\hat{1}}(\vec{r})$ is the rotation operator with respect to the orientation $\hat{1}$ of a rod. $D_r$ is the rotational diffusion of a rod at infinite dilution. Furthermore, $D$ is the thickness of the rods and $L$ is their length. $\hat{\Gamma} = \hat{\gamma} \hat{\Gamma}$ is the velocity-gradient tensor with $\hat{\gamma}$ the shear rate. Here we choose

$$\hat{\Gamma} = \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}$$

(2)

which corresponds to a flow $\vec{v}$ in the $x$-direction and its gradient $\nabla \vec{v}$ in the $y$-direction.

The concentration where the isotropic phase becomes unstable in the absence of shear flow can be calculated by solving the Smoluchowski equation at zero shear rate. This equation agrees with Onsagers approach to the I-N transition. Often the Maier-Saupe potential is used instead of the exact potential, which in fact corresponds to the first term of the Ginzburg-Landau expansion of the outer product in the exact potential given between the brackets in Eq. 1. Under flow conditions, a rich dynamics phase behavior is found as a function of shear rate and rod concentration. Marrucci and Maffettone were the first to solve the equation of motion of the ODF numerically, restricting themselves to two dimensions in order to reduce the computational effort $28$. They found that the director undergoes a tumbling motion with respect to the flow direction, resulting in a negative normal stress $N_1$. Larson expanded the ODF in three dimensions using spherical harmonics and truncated the expansion after checking for convergence $22$. This treatment predicts a transition from tumbling to “wagging” and finally to flow aligning state with increasing shear rates. A closure relation is frequently used for the interaction term on the right side of Eq. 1. This can greatly bias the results, see i.e. Feng et al $11$. The location of the flow transitions in the flow-concentration diagram is very sensitive to the choice of the closure, and no satisfactory closure has been found up till now.

In this paper we use a finite element method to numerically solve the equation of motion for the ODF, thus avoiding the use of any specific closure relation. As a typical diffusion-convection equation, this equation describes the diffusive-convective transport dynamics of an orientation of a homogeneous ensemble of thin rigid rods. A surface of a sphere is constructed on which a tip of the rod moves with respect to its center of mass. The equation for the probability of finding the tip of a rod in an area is determined by the transport fluxes on its boundaries due to (1) the Brownian diffusion (the first term in the brace brackets of Eq. 1), (2) the convection induced by the interparticle forces (the second term in the brace brackets of Eq. 1) and (3) the convection due to the imposed shear flow (the third term of Eq. 1).

To solve Eq. 1 numerically, a discretization scheme is used, and meshes on the surface of a unit sphere are constructed. For those operators inside the brace brackets which represent the transport fluxes we apply the central differences approximations. However, the rotation operator outside of the brace brackets needs to be discretized using the concept of transport fluxes through the boundaries of the mesh. In other words, the integral form of the Eq.(1) is invoked and applied to each of the mesh elements. To do this the identity, $(\hat{1} \times \nabla_{\hat{1}}) \cdot \vec{F} = \nabla_{\hat{1}} \cdot (\vec{F} \times \hat{1})$ is used in order to transform the angular transport flux of a rod to the translational transport flux of one tip of that rod. It differs from the conventional method of discretizing a differential equation where the operators are
written explicitly into the sum of the first- and the second
order derivatives and then the latter are approximated by
selected difference schemes. The advantage of the current
method is that, since neighboring meshes share bound-
aries, the fluxes leaving one mesh are always absorbed by
the surrounding meshes and vice versa. Therefore, there
is no loss and generation in the total amount of the ODF’s
as the computation proceeds (see Fig. 1). In practice a
$40 \times 80$ mesh was used on the surface of a unit sphere
with 40 equi-spaced grids in the polar angle and 80 equi-
spaced grids in the azimuthal angle in a spherical coor-
dinates. The right hand side of Eq. (1) is discritized on the
meshes according to the flux-conservative method men-
tioned above. A fourth order Adams’ predictor-corrector
method was invoked to follow the time evolution of the ODF. More details will be published in a forthcoming
paper.

The time-dependent ODF is now used to calculate the
time-dependence of three parameters characterizing the
flow behavior of a nematic phase: (1) $\theta$ describing the
angle between the nematic director and flow direction,
(2) the scalar magnitude of the director of defined by the
order parameter $P_2$ and (3) the total stress of an ensemble
of flowing rods. The angle and magnitude of the order
parameter are obtained from the order parameter tensor

$$ S = \int d\hat{u} \hat{u} \hat{u} P(\hat{u}, t). \tag{3} $$

The largest eigenvalue of the order parameter tensor $\lambda$,
characterizes the degree of alignment of rods with respect
to the director given by the corresponding eigenvector $\hat{n}$. The largest eigenvalue of $S$ is $1/3$ in the isotropic phase
and to 1 for a perfectly aligned nematic phase. Scalar
order parameter $P_2$ is defined as $P_2 = (3\lambda - 1)/2$.

The stress $\sigma_{12}$ is obtained from the deviatoric part of
the stress tensor derived by Dhont and Briels [10]:

$$ \Sigma_D = \eta_0 \dot{\gamma} + 3 \phi \kappa B T \left\{ S - \frac{1}{3} I + \frac{L}{D} \phi \Sigma_D^P + \frac{1}{6} P_{e r} [S^{(4)} : \hat{E} - \frac{1}{3} IS : \hat{E}] \right\}, \tag{4} $$

where

$$ \Sigma_D^P = \frac{8}{3\pi} \int d\hat{u} \int d\hat{u} \hat{u} \hat{u} \hat{u} \cdot (\hat{u} \times \hat{u}^\prime) \hat{u} \cdot \hat{u}^\prime P(\hat{u}, t) P(\hat{u}^\prime, t) \tag{5} $$

and

$$ S^{(4)} = \int d\hat{u} \hat{u} \hat{u} \hat{u} P(\hat{u}, t). \tag{6} $$

Here, $\phi = \frac{\pi}{8} L_D^4 \rho \kappa T$ is the volume fraction of rods, and
$P_{e r} = \dot{\gamma}/D_r$ the rotational Péclet number which is de-

The viscosity and stress response was measured using
an ARES strain controlled rheometer (TA instruments,
Delaware). A double wall Couette geometry was used
because of fairly low viscosity of the samples. Polarized
light microscopy images of $fd$ under shear flow were taken

FIG. 1: Flux conservation method used in discritizing Eq. (1)
The rod indicates the orientation of the director with respect
to the shear flow. The probability of finding a tip of one rod
in the shaded area of the unit sphere is determined by the
flux of the probabilities through the boundary of that area.

III. MATERIALS AND METHODS

The viscosity and stress response was measured using
an ARES strain controlled rheometer (TA instruments,
Delaware). A double wall Couette geometry was used
because of fairly low viscosity of the samples. Polarized
light microscopy images of $fd$ under shear flow were taken

The viscosity and stress response was measured using
an ARES strain controlled rheometer (TA instruments,
Delaware). A double wall Couette geometry was used
because of fairly low viscosity of the samples. Polarized
light microscopy images of $fd$ under shear flow were taken
P2

FIG. 2: Three plots showing the behavior of the angle of the nematic director $\theta$ (a), the magnitude of the nematic order parameter (b), and the average stress (c) as a function of strain after a start up of the flow. The dimensionless rod concentration is $\phi L / D = 4.5$. Data is obtained by numerically solving Eq. 1 using the finite element method. The rods are initially placed in the flow-gradient plane. For the stress calculation only the elastic contribution (Eq. 4) was considered.

using a Linkam CSS450 plate-plate shear cell.

The physical characteristics of the bacteriophage $fd$ are its length $L = 880 \text{nm}$, diameter $D = 6.6 \text{nm}$, persistence length of 2200 $\text{nm}$ and a charge per unit length of around 10$^{-7}$ $\text{e}^{-/\text{nm}}$ at pH 8.2. When in solution, $fd$ exhibits isotropic, cholesteric, and smectic phases with increasing concentration. $Fd$ forms a cholesteric phase while the DEH theory is valid for nematic structures. In practice nematic and cholesteric phase are locally almost identical and the free energy difference between these phases is very small. In this paper we do not distinguish between these two phases. The $fd$ virus was prepared according to standard biological protocols using XL1-Blue strain of E. coli as the host bacteria. The standard yields are approximately 50 mg of $fd$ per liter of infected bacteria, and virus is typically grown in 6 liter batches. The virus is purified by repetitive centrifugation (108 000 g for 5 hours) and re-dispersed in a 20 mM Tris-HCl buffer at pH 8.2.

A. $fd$ as a model hard rod system

The Onsager theory for hard rod dispersions predicts a first order phase transition between a disordered, isotropic phase and an orientationally ordered, nematic phase. Due to hard core athermal interactions considered in the Onsager model, the phase diagram is temperature independent and the rod concentration is the only parameter that determines the location of the I-N phase transition. The two points spanning the region of isotropic-nematic coexistence are called the binodal points. The spinodal point is located at a rod concentration higher than the lower binodal point and is determined by the following condition $\phi L / D = 4$. $Fd$ viruses are not true hard rods, due to surface charge and limited flexibility. As a consequence, their equilibrium phase behavior differs from the ideal hard rod case described by Onsager based theory, e.g. DEH. The finite flexibility of $fd$ viruses drives the concentration of the binodal points to a 30% higher value when compared to equivalent but perfectly stiff hard rods. In addition, flexibility also reduces the value of the order parameter of the coexisting nematic phase. For $fd$ the order parameter of the coexisting nematic is about 0.65 while Onsager theory for hard rods in equilibrium predicts the order parameter of 0.8. The effect of surface charge is to increase the effective diameter of the rod $D_{\text{eff}}$ and therefore the excluded volume interaction between charged rods. As a consequence the charge reduces the real concentration of the phase transition.

For the $fd$ suspension used, the binodal point at high rod concentration $c_{IN}$ occurs at 11 mg/ml. After taking the effects of flexibility and charge into account it was shown that the order parameter of the nematic solution of $fd$ is quantitatively described by the extensions of the Onsager theory to the semi-flexible case. Hence, even though $fd$ is flexible and charged, it can be used to quantitatively test predictions of the DEH theory. It is, however, a very difficult and until now unfulfilled task to incorporate charge and flexibility into a non-equilibrium equation of motion such as Eq. 1. Therefore in this paper we use data from reference to convert the measured concentration of $fd$ to the nematic order parameter of the sample. After that we compare experiments and theory at the same values of the order parameter.
A hesitation in the shear rate dependence of the viscosity was predicted theoretically by Larson [22]. It was argued that the transition from the tumbling regime to the wagging regime implies a broadening of the ODF which leads to higher dissipative stresses. The broadening of the ODF is illustrated in Fig. 2b. As can be seen in Fig. 2b, it is not straightforward that ODF broadening really has an effect on the stress. We calculated the time-dependent viscosity by numerically solving the equation of motion of the ODF for 20 different initial orientations of the director. From the time-dependent ODFs we calculated the viscosity using either only the elastic term or both elastic and viscous terms. The viscosity is averaged over all 20 traces and a tumbling period after the transient start up flows have died out. The results are scaled to the experiment using a typical concentration of 16; mg/ml for $\bar{\rho}$ in Eq. 4 and the value of $D^0_{\text{inf}} = 20 \text{ s}^{-1}$, taken for the rotational diffusion at infinite dilution [20, 32]. Fig. 4 the stationary viscosity decreases continuously with increasing shear rate and only shows a hesitation when the viscous contribution to the stress is not included. The shear rate where this hesitation occurs corresponds with the shear rate where the system nematic ordering is significantly reduced and the transition from tumbling to wagging takes place, as can be concluded from Fig. 2. Comparing the model predictions to the experiments it should be noted that the experimentally observed features are much more pronounced. Moreover, there is no real reason to leave out the viscous contribution although it does obscure the behavior we see in the experiment. Still, the maximum in the viscosity is interpreted as a signature of the transition from tumbling to a wagging state.

There are three observations to keep in mind when considering $fd$ in the nematic phase under shear flow, which all point to very low stresses in such systems when compared to polymeric liquid crystals. First, the viscosity of $fd$ in the nematic phase is two to three orders of magnitude lower than the viscosity of typical polymeric liquid crystals such as poly(benzyl glutamate) (PBG) [38], although the difference in solvent viscosity is only one order of magnitude. Second, the range over which the viscosity of $fd$ suspension varies is more limited with changing shear rate and rod concentration: the viscosity lies between 70 times the solvent viscosity for low shear rate and low rod concentration and 20 times the solvent viscosity for high shear rate and rod concentration. Moreover, the viscosity as calculated from the equation of motion of the ODF is of the same order as the measured viscosity. Third, polymer nematics exhibit negative first normal stress differences for certain shear rates as was first observed for PBG solution [18]. This is a direct consequence of the tumbling of the nematic director. Attempts have been made to measure the first normal stress difference for nematic $fd$ solutions but due to very low force the

![Figure 3](image_url)  
**FIG. 3:** Stationary viscosity as a function of shear rate for four different concentration of $fd$ virus at 11.5, 13, 16 and 25 mg/ml.

![Figure 4](image_url)  
**FIG. 4:** The theoretical time averaged viscosity at a dimensionless concentration of $\frac{D}{T} \bar{\phi} = 4.5$ with (□) and without (■) the viscous contribution of the rods, as calculated by solving the equation of motion of the ODF for 20 independent initial orientations of the director. The lines indicate the transition from tumbling to wagging, and from wagging to flow aligning as found from Fig. 2. The results are scaled to the experiment using a typical concentration of 16; mg/ml for $\bar{\rho}$ in Eq. 4 and $D^0_{\text{inf}} = 20 \text{ s}^{-1}$ for the rotational diffusion at infinite dilution [20, 32].

**IV. RESULTS**

**A. Stationary viscosity**

The measurements of a stationary viscosity as a function of the shear rate for different $fd$ concentrations are shown in Fig. 3. For the lowest concentrations of $fd$ the viscosity decreases continuously with shear rate except for a small hesitation at a shear rate of 10 s$^{-1}$. This hesitation is similar to what is observed for solutions of PBG at low concentration in solvent $m$-cresol [18, 30]. For $fd$ at intermediate concentrations, shear thinning becomes less pronounced, the hesitation shifts to higher shear rates and turns into a local maximum. For the highest $fd$ concentration almost no shear thinning is observed, only a pronounced peak in the viscosity. This shear thickening behavior has not been previously reported.

The measurements of a stationary viscosity as a function of shear rate for different $fd$ concentrations are shown in Fig. 3. For the lowest concentrations of $fd$ the viscosity decreases continuously with shear rate except for a small hesitation at a shear rate of 10 s$^{-1}$. This hesitation is similar to what is observed for solutions of PBG at low concentration in solvent $m$-cresol [18, 30]. For $fd$ at intermediate concentrations, shear thinning becomes less pronounced, the hesitation shifts to higher shear rates and turns into a local maximum. For the highest $fd$ concentration almost no shear thinning is observed, only a pronounced peak in the viscosity. This shear thickening behavior has not been previously reported.

The measurements of a stationary viscosity as a function of shear rate for different $fd$ concentrations are shown in Fig. 3. For the lowest concentrations of $fd$ the viscosity decreases continuously with shear rate except for a small hesitation at a shear rate of 10 s$^{-1}$. This hesitation is similar to what is observed for solutions of PBG at low concentration in solvent $m$-cresol [18, 30]. For $fd$ at intermediate concentrations, shear thinning becomes less pronounced, the hesitation shifts to higher shear rates and turns into a local maximum. For the highest $fd$ concentration almost no shear thinning is observed, only a pronounced peak in the viscosity. This shear thickening behavior has not been previously reported.
signals were too small to be measured.

B. Flow reversal experiments

In flow reversal experiments, the sample is first sheared at a constant shear rate in one direction until the steady state condition is reached. Subsequently, the direction of flow is suddenly reversed while keeping the magnitude of shear rate constant. Such experiments have been very useful in characterizing and understanding the dynamics of sheared liquid crystalline polymers [30]. In the present work, flow reversal experiments were performed covering a wide range of shear rates and \( \text{fd} \) concentrations. Typical flow reversal experiments are depicted in Fig. 5 for a \( \text{fd} \) concentration of 11.5 mg/ml which corresponds to \( c/c_{IN} = 1 \).05. At the lowest shear rates a damped oscillatory response is obtained which decays within few oscillations (Fig. 5a). Increasing the shear rate results in a more pronounced oscillatory response, which damps out relatively slowly. The oscillatory response in Fig. 5b is most pronounced at a shear rates of 12 s\(^{-1}\). At even higher shear rates, the damping again increases (Fig. 5c).

In order to quantitatively characterize the response to a nematic to a flow reversal, the data is fitted to a damped sinusoidal superimposed onto an asymptotically decaying function of the following form:

\[
\eta(t) = \eta_{stat} \cdot \left\{ 1 + A e^{-\frac{\dot{\gamma}}{\tau_d}} \sin(2\pi \frac{\dot{\gamma}t - \varphi}{P}) \right\} (1 - b \cdot g^{\dot{\gamma}t}). (7)
\]

This is an empirical choice, but each variable in the fit contains important information about the behavior of rods in shear flow. Fig. 6 shows the behavior of fit parameters as a function of the shear rate at few selected concentrations of \( \text{fd} \) virus. In this figure we indicate with vertical dashed lines the shear rates at which the steady state viscosity exhibits a local maximum for four different concentrations. Interestingly, these are exactly the same shear rates at which the damping constant \( \tau_d \) as well as the tumbling period \( P \) show a sharp increase. The asymptotic constant \( b \), on the contrary, shows a decrease. These features disappear for the highest \( \text{fd} \) concentration. Presumably the three regions showing different flow reversal behavior correspond to tumbling, wagging and flow aligning regime. This will be discussed in more detail in section V B. In the next section we first discuss the concentration and shear rate dependence of the tumbling period in the regime where rods exhibit tumbling flow behavior.

C. Tumbling period as a function of shear rate and rod concentration

DEH theory predicts that as the De (or Pe) number is increased, that the 'molecular' period of oscillation decreases with increasing shear rate in the tumbling regime [22]. This feature was never fully explored, since in most polymeric liquid crystals it was found that the tumbling period was strain scaling, implying that the response overlaps when the period is scaled with the applied shear rate and the stress is normalized by its steady state value. The strain scaling arises as a consequence of the presence of a large, non-Brownian, length scale in the sample that determines the time response, even at relatively high De (or Pe) numbers. This most probably is the domain size characterizing the nematic texture. The log-log plot of the tumbling period (\( T = P/\dot{\gamma} \)) as a function of the shear rate is shown in Fig. 7. Here the data are only shown for a low shear rate region which is associated with the tumbling regime. Strain scaling, if present, would give a slope of -1. However, as can be seen in the inset of Fig 7 the reciprocal indicating strain scaling is only approached and not reached at the highest rod concentration studied here.

The shear rate dependence of the tumbling period is compared to the theoretical prediction for the same rod concentration as well as the same order parameter, see Fig. 8. The reason for using the order parameter to assess the theoretical predictions was discussed at length.
FIG. 6: a) Steady state viscosity as function of the shear rate for fd virus at four different concentrations. All viscosity curves exhibit shear thinning at low shear rates followed by a local maximum in viscosity. b-d) Behavior of the parameters obtained from fitting the response of the shear flow reversal experiments to Eq. 7. The vertical lines indicate the local maximum in viscosity curves. The local maximum in the steady state viscosity curve corresponds to maximum of the tumbling period $P$ and damping constant $\tau_d$ and minimum of asymptotic constat $b$ in the flow reversal.

in section IIIA For purposes of comparison the order parameter was obtained form x-ray experiments and the value of $D_\text{inf}^0 = 20 \text{ s}^{-1}$. We emphasize that DEH theory is microscopic and that there are no adjustable parameters in the comparison between theory and experiments. Clearly there is a qualitative correspondence between theory and experiment, both showing a continuous decrease of the period. The quantitative correspondence, on the other hand, is limited. This is probably due to fact that texture, although not dominating the response, is still present. It will be shown later in section IVB that the shear rate and rod concentration dependence of a tumbling to wagging and wagging to flow-aligning transition agree much better with DEH theory.

The concentration dependence of the tumbling period is shown in Fig. 9. Here, theory and experiments are compared at a fixed shear rate at which the tumbling to wagging flow transition occurs. The tumbling period increases with increasing rod concentration (Fig. 9a) or, equivalently, increasing order parameter of the nematic phase (Fig. 9b). The increase of the tumbling period with increasing order parameter was already predicted using a linearized version of the DEH theory[21].

In conclusion, the absence of strain scaling of the tumbling period and the qualitative agreement between theory and experiment the tumbling period indicates that the response of the suspension of fd-virus is dominated by the molecular elasticity arising from the distortion of the ODF of particles.

FIG. 7: The dependence of the tumbling periods on the shear rate for different concentrations of the nematic fd. The figure shows that the tumbling period scales with a power low as a function of the shear rate. The inset shows the power law dependence of the tumbling period on the shear rate for different fd concentrations.

FIG. 8: The period of the oscillations (in units of strain) as a function of the Péclet number, where the shear rate is scaled with the rotational diffusion of fd at infinite dilution.
FIG. 9: The period of the oscillations (in units of strain) as a function of the dimensionless concentration (a) and order parameter (b). The shear rate was chosen at the point we identify with the tumbling to wagging transition for experiment (■) and exact theory (○).

D. Relaxation at high concentration

In order to measure the relative magnitude of the elastic texture contribution to the overall stress, relaxation experiments were performed. For polymeric liquid crystals like polybenzylglutamate (PBG) solutions in cresol, Walker et al. [39] showed that there are three different regimes of relaxation behavior, each of which is related to a distinct structural relaxation. There is a "fast" relaxation of the nematic fluid; a "slower" relaxation that exhibits scaling with the shear rate before the cessation of flow, which is due to the indirect contribution of the texture to the overall stress; and a "long-time" relaxation due to the reorganization of the texture on a supra-molecular level which will not be addressed here.

Stress relaxation experiments were performed in the low shear rate “tumbling” region, at shear rates smaller than those corresponding to the maximum in the viscosity. The sample used had a relatively high fd concentration of 25 mg/ml, corresponding with c/c_in = 2.3. Some typical responses to the cessation of flow are depicted in Fig. 10. The stress is normalized to its value before the cessation of flow, and the time axis is scaled by the shear rate. The fast component of the decay takes place at less than a tenth of a second, which is comparable to the response of the force re-balanced transducer and therefore not shown. The slow component of the stress relaxation scales when time is multiplied with the previous shear rate, but only from the point that the stress has decayed to less than 30 % of its original value, or less for higher initial shear rates. From Fig. 10 it can be concluded that the contribution to the stress for the highest concentration used and for low shear rates is 30 %. This is the absolute upper limit for the samples used in this paper. It should be noted that for PBG solutions 30 % it was found to be the lower limit [39].

E. In situ microscopy

The flow-induced changes of the liquid crystalline texture during steady state shear flow were studied using a plate-plate geometry in combination with a polarization microscope. Measurements were performed for fd concentrations of 14 mg/ml and 25 mg/ml. Typical images are shown in Fig. 11 for different shear rates. Interestingly the characteristic size of the "domains" was very large. Birefringent regions of up to half a millimeter were observed under static conditions. When the sample is subjected to shear flow, these domains will elongate and eventually disappear, at values of the shear rate which correspond to the maximum in the viscosity (see Fig. 6a). An important difference between the two concentrations is that the elongated domains merge into bands for high rod concentration, whereas for the low concentration the structure disappears before such bands are formed. Interestingly, this transition to a banded structure in the high concentration fluid takes place at a shear rate which is higher than the shear rate where the low concentration

FIG. 10: The stress relaxation after cessation of flow for fd at 25 mg/ml (c/c∗ = 2.3), varying the initial shear rate. The time is scaled by the initial shear rate. The stress is normalized by the stress before the cessation of flow.
V. DISCUSSION

When comparing the flow behavior of the polymeric nematic phase and the colloidal nematic phase of the dispersed \textit{fd} viruses, the most striking observation is the qualitative agreement between the two systems, despite the fact that \textit{fd} is an order of magnitude larger. The viscosity of the \textit{fd} nematic is much smaller and the rotational diffusion of \textit{fd} is much slower when compared to polymeric liquid crystals. Flow reversal experiments reveal typical transitions in the transient rheological behavior: damped oscillations occur at low shear rates changing to undamped oscillations at intermediate shear rate, which disappear if the shear rate is increased even further; the time scale of the oscillations of the stress transients is comparable. Also other well known phenomena like the formation of very large bands upon cessation of flow along the vorticity direction which have been studied in detail in polymeric systems [37], can also be observed here (data not shown).

Having established that \textit{fd} virus dispersions indeed undergo a tumbling motion under flow, the dynamic behavior of \textit{fd} suspensions can be rationalized on the basis of the microscopic theoretical predictions for a homogeneous system of rods under shear. Doing so, one important prerequisite needs to be fulfilled, namely that the dominating contribution to the stress is coming from the nematic fluid and not from the texture. It will be argued here that this indeed is the case. Having done so, we will be able to map out a phase diagram of the dynamic transitions from tumbling to wagging to flow aligning.

A. Textural evolution during flow

The word “texture” refers to disclination points and lines where the director of the nematic phase changes discontinuously, marking domains in the sample. When a system containing these domains and disclinations is subjected to shear flow, part of the dissipated energy is used to destroy these structures. Fig. 11 shows that the domains tend to elongate and align with the flow. Disclinations can also cause a direct contribution to the total stress resulting in a high viscosity and a very pronounced shear thinning behavior, typically referred to as region I [40]. Experiments on polymeric liquid crystals have revealed several features of the flow behavior of nematic liquid crystals which are attributed to the presence of texture in the nematic phase. Tumbling induces distortions in the director field and the defects arrest the tumbling, thereby inducing an elastic stress. The length scale over which this distortion occurs, i.e. the ‘domain’ length scale, is an inherent non-Brownian length scale, see ref. [3]. As a consequence, stress patterns during flow reversal will display strain scaling. Also the damping of

\begin{figure}[ht]
\centering
\includegraphics[width=\textwidth]{fig11.png}
\caption{The polarization images of the nematic \textit{fd} at 14 and 25 mg/ml for a range of different steady state shear rates. The dashed line in the bottom right image indicates the border between the structured and unstructured regions.}
\end{figure}
the oscillations is explained on the base of the presence of the polydomain structure, where e.g. the "friction" between the domains would lead to a damping of the oscillations. The scaling of the stress relaxation process after the flow is stopped with shear rate has been explained using the same arguments. From such an experiment the relative contribution to the total stress of a homogeneous nematic phase and the polydomain texture can be estimated since the relaxation dynamics of the nematic phase is much faster than that of polydomain structure.

The micrographs in Fig. 11 clearly reveal that texture under flow exists in nematic fd dispersions. Their contribution to the rheology is, however, far less prominent when compared to polymeric liquid crystals such as PBG. This we can infer from several observations. First, very moderate shear thinning is observed in the low shear rate regime for the low concentrations, which gradually disappears with increasing concentration (Fig. 6b). This is very similar to theoretical predictions for a homogeneous nematic phase (Fig. 2b in Ref. [4]). Also, the calculated and measured viscosity are of the same order of magnitude. In contrast, shear thinning can be fairly strong in the low shear rate region (Region I) where texture dominates the response and it will increase with increasing concentration, although also other microstructural features can contribute here. Second, the tumbling period is not strain scaling (Fig. 7), which could be due to either a smaller relative magnitude of the textural stress or due to the fact that we are not in a low enough Pe regime. Third, 'strain' scaling is recovered for the slow 'textural' relaxation process after the flow has been stopped. This experiment shows that at the highest rod concentrations used and at low shear rates the distortional textural contribution is about 30%. For most experiments done this value is probably significantly lower. So, where texture is important, even dominating the stress response for molecular LCPs, molecular elasticity is far more dominating the fd-virus. Though we just argued that the texture does not dominate the shear response of the system, this does not mean that the shear response is not influenced by texture. For one the oscillations we observe are still strongly damped, and the damping only decreases when the transition to the flow aligning state is reached (see the behavior of \( \tau_d \) in Fig. 6). Moreover, the presence of texture might explain the discrepancy in the behavior of the period of the oscillations between experiment and theory (Fig. 9). Most importantly, we know from microscopy that texture is present under shear (see Fig. 11). It should be noted, however, that the size of the polydomain structure of the fd dispersions is one order of magnitude bigger as compared to PBG, so that the density of disclination lines and points is about three orders of magnitude lower for fd. Note that the length scale of the texture during flow is still small compared to the dimension of the flow cell. Since the contribution of texture scales with the density of the disclinations, texture will be far more dominating for e.g. PBG than for fd, even when elastic constants are almost the same for the two systems.

B. Phase diagram of dynamical flow transitions

In this section the experimental results or combined and a non-equilibrium phase diagram of fd rods under shear flow is presented. The results for the four fit parameters plotted in Fig. 6 show clear transitions at well defined shear rates for all fd concentration. Although they only give an indirect proof of the transitions, they can be used to infer information about the flow transitions. For all fd concentrations (except for the highest one) the shear rate where the maximum viscosity is reached is identical with the shear rate where the period as well as the damping constant start to increase (indicated by the vertical dashed lines in Fig. 5). The microscopic observations are in fairly good agreement with the transitions inferred from the rheology. Upon approaching the tumbling to wagging transition from tumbling to flow aligning, the texture becomes too faint to resolve in the microscope and texture subsequently disappears upon reaching the FA region. For the high fd concentration, i.e. the sample showing shear banding (Fig. 11 last), one can identify a sharp transition from a structured to an unstructured region in the same micrograph. Since this picture was taken in the plate-plate geometry, there is a shear rate distribution across the image: the shear rate is increasing going from the left side to the right. A sharp spatial transition therefore also represents a sharp transition at a given shear rate. Although, due to the method of zero gap-setting, the value of the shear rate is not exactly known (±20%), one can still identify the shear rate where structure disappears as the shear rate where the viscosity reaches its local maximum (the down pointing triangles in Fig. 6a). For low fd concentration of (14 mg/ml) the structure disappears around the point where the viscosity reaches its local maximum, although the morphological transition for the lower concentration is less abrupt.

Fig. 12 shows the behavior of flow transitions as a function of shear rate for various fd concentrations. For the experiment we plotted the Péclet numbers where the viscosity shows a local maximum and where the damping constant reaches a maximum. The theoretical predictions for the tumbling to wagging and wagging to flow aligning transitions are obtained from the plots the angle of the nematic director \( \theta \) under flow, see Fig. 2. Similar to the method used in Fig. 5 the experimental concentration is scaled to the theoretical concentration in two different ways: effective concentration (Fig. 12a) and the order parameter \( P_2 \) (Fig. 12b). This figure was shown in a preliminary paper without a detailed explanation. The shear rate is rescaled to the Péclet number by using the rotational diffusion coefficient at infinite dilution. Fig. 12 allows us to draw some important conclusions. First, it is clear that scaling the concentration with the equilibrium
VI. CONCLUSIONS

Colloidal suspensions of rod-like \textit{fd} viruses are an ideal model system to study the behavior of the nematic liquid crystalline phase under shear flow. Flow reversal experiments show signatures for tumbling, wagging, and flow aligning behavior, very similar to the behavior found in polymeric liquid crystals. The rigid rod nature of the \textit{fd} suspension, possibly combined with a smaller relative textural contributions to the overall stress tensor make \textit{fd}-virus a suitable model system for the DEH theory. Important in this respect is that the overall viscosity is only one to two orders of magnitude higher than the solvent viscosity. Also it is important to note that stress relaxation experiments combined with the absence of strain scaling in flow reversal experiments suggest that there is only a limited contribution of textural aspects to the overall stress, even for the highest \textit{fd} concentration used in this work. The shear thickening of the viscosity observed for a range of \textit{fd} concentrations is as yet, unex-

order parameter gives better agreement when compared to the scaling by the dimensionless concentration. The fact that theory and experiment agree without using any fitting parameters ($P_2$ was obtained in a separate experiments \cite{12}) leads to the conclusion that the DEH theory describes the flow behavior of the \textit{fd} nematics quite well, as long as the effects of flexibility and charge of the experimental rods are included in the calculation of the order parameter. Less convincing agreement is obtained when comparing the experimental and theoretically calculated periods (Fig. \textbf{11}). The reason for this could be the remaining textural contribution to the overall stress which, although small, cannot be neglected. Since we deduce from Fig. \textbf{12}, that a dimensionless concentra-
tion of $\phi D/L = 4$ corresponds with a \textit{fd} concentration of 16 mg/ml, we used this number \textit{a posteriori} to scale the calculated molecular viscosity in Fig. \textbf{11}. The pure elastic contribution shows a very nice quantitative correspondence with the experimental data. Interestingly, when the viscous term is added, the theoretical viscosity is higher than the experimental viscosity, despite of the fact that no hydrodynamics is incorporated.

In the previous subsection it was argued that the influence of textural contribution to the stress tensor of \textit{fd} are relatively small, as compared to PLC’s. There are however strong indications that the dynamic behavior is influenced by the macroscopic bands which are formed for the samples at the highest concentrations used (see Fig. \textbf{11}). As can be seen in Fig. \textbf{12}, the typical features for the transition to wagging disappear: there is no increase in the damping constant, nor in the period of the oscillations. Moreover, the theory shows only a moderate hesitation of the stationary viscosity (which even disappears then the viscous term is added, Fig. \textbf{9}), whereas in experiments a local peak is observed which is more pronounced with increasing concentration. The microscopy pictures results that at high concentrations the systems finds another way to handle the distortion of the particle distribution at high shear rates by forming shear bands where the overall orientational distribution is alternating, as was already observed and partially explained for the polymeric systems \cite{22, 24, 25}. In the present work, the concentration dependence of the phenomenon at hand suggests that this merits further experimental as well as theoretical work. In this context one should not forget that we compar experiments on charged and semi-flexible \textit{fd} with theory for hard and stiff rods. It could well be that these factors also play an important role. It will be a major challenge especially to take the semi-flexibility into account in the equation of motion.

FIG. 12: Phase diagram of flow transitions for the nematic \textit{fd} phase as a function of dimensionless concentration (a) and order parameter (b). The experimental points indicate the Péclet numbers where the viscosity shows a local maximum (■) and where the damping constant $\tau_d$ (●) reaches a maximum. The theoretical points indicate tumbling to wagging (□) and wagging to flow aligning (○) transitions.
plained. The maximum in the viscosity occurs at the critical shear rate where the tumbling to wagging transition takes place. Microscopic observations show that at this shear rate the morphological features disappear, suggesting a strong connection between the dynamic transitions and structure formation.

The experimental results have been compared to a microscopic theory for rod-like molecules subjected to shear flow. A non-equilibrium phase diagram is constructed, describing the transitions from tumbling to wagging and from wagging to flow-aligning as a function of rod concentration and applied shear stress. When scaling the results to the concentration where the isotropic-nematic transition takes place, the experiment and theory show only a qualitative agreement, possibly due to the fact that the real rods are both semi-flexible and charged. However, when scaling the results using the order parameter, which is determined by the interactions between the rods, theory and experiment show an excellent agreement without using any fit parameters. Thus, it can be concluded that the Doi-Edward-Hess theory accurately captures the dynamic features of a hard rod system. Pd dispersions constitute such a hard rod system as long as flexibility and charge are properly taken into account, which can be simply achieved by using the order parameter to scale the data. More theoretical work is needed, however, to explain the clear connection between the observed band formation at high concentrations and the dynamic transitions, and to incorporate the effect of flexibility of the rods.

Acknowledgement

We thank Jan Dhont for many discussions and critical reading of the manuscript. Pier-Luca Maffettone is acknowledged for stimulating remarks. MPL is supported by the Transregio SFB TR6, “Physics of colloidal dispersions in external fields”. ZD is supported by Junior Fellowship at Rowland Institute at Harvard. The authors acknowledge support of the EU (6th FP) in the framework of the Network of Excellence “SOFTCOMP.”

[1] R. G. Larson. On the relative magnitudes of viscous, elastic and texture stresses in liquid crystalline PBG solutions. Rheol. Acta, 35 (2): 150-159, 1996.
[2] Ugaz V. M. and Cinader D. K. and W. R. Burghardt. Origins of region I shear thinning in model lyotropic liquid crystalline polymers. Macromolecules, 30 (5): 1527-1530, 1997.
[3] W. R. Burghardt and G. G. Fuller. Transient shear flow of nematic liquid crystals: Manifestations of director tumbling. J. Rheol., 34(6):959-992, 1990.
[4] W. R. Burghardt and G. G. Fuller. Role of director tumbling in the rheology of polymer liquid crystal solutions. Macromolecules, 24:2546, 1991.
[5] J. K. G. Dhont and W. J. Briels. Viscoelasticity of suspensions of long, rigid rods. Colloid Surface A, 213(2-3):131-156, 2003.
[6] Z. Dogic and S. Fraden. Smectic phase in a colloidal suspension of semiflexible virus particles. Phys. Rev. Lett., 78:2417, 1997.
[7] Z. Dogic and S. Fraden. Cholesteric phase in virus suspensions. Langmuir, 16:7820-7824, 2000.
[8] Z. Dogic and S. Fraden. Development of model colloidal liquid crystals and the kinetics of the isotropic-smectic transition. Phil. Trans. R. Soc. Lond. A., 359:997, 2001.
[9] M. Doi and S. F. Edwards. The Theory of Polymer Dynamics. Oxford, 1986.
[10] V. Faraoni, M. Grosso, S. Crescitelli, and P. L. Maffettone. The rigid-rod model for nematic polymers: An analysis of the shear flow problem. J. Rheol., 43:829, 1999.
[11] J. Feng, C. V. Chaubal, and L. G. Leal. Closure approximations for the doy theory: Which to use in simulating complex flows of liquid-crystalline polymers? J. Rheol., 42(5):1095-1119, 1998.
[12] M. G. Forest and Q. Wang. Monodomain response of finite-aspect-ratio macromolecules in shear and realted linear flows. Rheol. Acta, 42:20-46, 2003.
[13] S. Fraden. Observation, Prediction, and Simulation of Phase Transitions in Complex Fluids, volume 460 of NATO-ASI - Series C, pages 113-164. Kluwer Academic Publishers, Dordrecht, m. baus and l. f. rull and j. p. ryckaert, edition, 1995.
[14] C. Graf, H. Kramer, M. Deggelmann, M. Hagenbüchle, Ch. Johner, Ch. Martin, and R. Weber. Rheological properties of suspensions of interacting rodlike fd-virus particles. J. Chem. Phys., 98(6):4921-4928, 1993.
[15] M. Grosso, S. Crescitelli, E. Somma, J. Vermant, P. Moldenaers, and P. L. Maffettone. Prediction and observation of sustained in a shear liquid crystalline polymer. Phys. Rev. Lett., 90:098304, 2003.
[16] S. Hess. Fokker-planck-equation approach to flow alignment in liquid crystals. Z. Naturforsch., 31(a):1034-1037, 1976.
[17] M. N. Kawaguchi and M. M. Denn. A mesoscopic theory of liquid crystalline polymers. J. Rheol., 43(1):111-124, 1999.
[18] G. Kiss and R. S. Porter. Rheolog of concentrated solutions of poly(γ-benzyl-glutamate). J. of Polym. Sci., 65:193-211, 1978.
[19] Korn and Korn. Mathematical Handbook for Scientists and Engineers. Mc-Graw and Hill, 1968.
[20] H. Kramer, M. Deggelmann, C. Graf, M. Hagenbichle, C. Johner, and R. Weber. Electric birefringence measurements in aqueous fd virus solutions. Macromolecules, 25:4325-4328, 1992.
[21] N. Kuzuu and M. Doi. J. Phys. SOC. Jpn., 53:1031, 1984.
[22] R.G. Larson. Arrested tumbling in shearing flows of liquid crystal polymers. Macromolecules, 23:3983-3992, 1990.
[23] R.G. Larson. Roll-cell instabilities in shearing flows of ne-
matic polymers. *Liquid Crystals*, 37(2):175–197, J. Rheol.

[24] R.G. Larson and M. Doi. Mesoscopic domain theory for textured liquid crystalline polymers. *J. Rheol.*, 35(4):539–563, 1991.

[25] R.G. Larson and D. W. Mead. The ericksen number and deborah number cascades in sheared polymeric nematics. *Liquid Crystals*, 5(2):151–169, 1993.

[26] M.P. Lettinga and J.K.G. Dhont. Non-equilibrium phase behavior of rod-like viruses under shear flow. *J. Phys.: Condens. Matter* 16 (2004), 16:S3929S3939, 2004.

[27] G Marrucci and F. Greco. Flow behavior of liquid crystalline polymers. *Adv. Chem. Phys.*, 86:331, 1993.

[28] G Marrucci and P. L. Maffetone. Description of the liquid-crystallin phase at high shear rates. *Macromolecules*, 22:4076, 1989.

[29] J. Mewis, M. Mortier, J. Vermant, and P. Moldenaers. Experimental evidence for the existence of a wagging regime in polymeric liquid crystals. *Macromolecules*, 30(5):1323–1328, 1997.

[30] P. Moldenaers and J. Mewis. Transient behavior of liquid crystalline solutions of poly(benzylglutamate). *J. Rheol.*, 30(1):567–584, 1986.

[31] L. Onsager. The effect of shape on the interaction of colloidal particles. *Annals of the New York academy of science*, 51:62–659, 1949.

[32] K. R. Purdy, Z. Dogic, S. Fraden, A. Rühm, L. Lurio, and S. G. J. Mochrie. Measuring the nematic order of suspensions of colloidal fd virus by x-ray diffraction and optical birefringence. *Phys. Rev. E.*, 67:031708, 2003.

[33] J. Sambrook, E. F. Fritsch, and T. Maniatis. *Molecular Cloning: A Laboratory Manual*, chapter 4. Cold Spring Harbor Laboratory Press, 2 edition, 1989.

[34] F. G Schmidt, B. Hinner, E. Sackmann, and J. X. Tang. Viscoelastic properties of semiflexible filamentous bacteriophage fd. *Phys. Rev. E*, 62(4):5509–5517, 2000.

[35] J. Tang and S. Fraden. Isotropic-cholesteric phase transition in colloidal dispersions of filamentous bacteriophage fd. *Liquid Crystals*, 19(4):459–467, 1995.

[36] V. G. Taratuta, A. J. Hurd, and R. B. Meyer. Light-scattering study of a polymer nematic liquid crystal. *Phys. Rev. Lett.*, 55(2):246–249, 1985.

[37] J. Vermant, P. Moldenaers, J. Mewis, and S. J. Picken. Band formation upon cessation of flow in liquid-crystalline polymers. *J. Rheol.*, 38(5):1571–1589, 1994.

[38] J. Vermant, P. Moldenaers, S. J. Picken, and J. Mewis. A comparison between texture and rheological behaviour of lyotropic liquid crystalline polymers during flow. *J. Non-Newton. Fl. Mech.*, 53:1–23, 1994.

[39] L. M. Walker, M. Mortier, and P. Moldenaers. Concentration effects on the rheology and texture of pbm/m-cresol solutions. *J. Rheol.*, 40(5):967–981, 1996.

[40] L. M. Walker and N. J. Wagner. Rheology of region i flow in a lyotropic liquid-crystal polymer: The effects of defect texture) under shear and during relaxation. *J. Rheol.*, 38(5):1524–1547, 1994.