Rheology and morphology of model immiscible polymer blends with monodisperse spherical particles at the interface

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Synopsis

We show that the addition of solid particles to droplet–matrix blends of immiscible polymers induces massive changes in the rheology and the flow-induced structure even at loadings as low as 0.1 vol. %. Experiments were conducted using blends of polyethylene oxide (PEO, dispersed phase), polyisobutylene (PIB, continuous phase), and 470 nm monodisperse silica particles with two different surface wettabilities. Rheological experiments were conducted under molten conditions, while the morphology was characterized at room temperature using scanning electron microscopy. We are able to image the morphology at both lengthscales: The >20 μm lengthscale of the dispersed phase, as well as the submicron lengthscale of the particles. Rheological experiments along different trajectories in the ternary particle/PEO/PIB composition diagram reveal that addition of ~1 vol. % particles that are preferentially wetted by the PIB induces a large increase in steady shear viscosity, severe shear-thinning, and yield-like behavior. However if the particles are equally wetted by PEO and PIB, these effects are greatly diminished. Remarkably, addition of very low loadings (~0.1 vol. %) of particles reduces the viscosity under some conditions regardless of wettability. These rheological changes are interpreted in terms of three observations from morphological studies: That particles greatly enhance coalescence at low volume loadings, that particles jam the interface at higher loadings, and that particles bridge across drops and glue them together into large clusters. The first two of these effects occur regardless of particle wettability, whereas the last occurs only with particles that are preferentially wetted by the continuous phase. © 2013 The Society of Rheology. [http://dx.doi.org/10.1122/1.4801757]

I. INTRODUCTION

The structure and rheology of blends of immiscible polymers have been examined for more than three decades. The effects of the various parameters (volume fraction of the two polymers, relative viscosity and viscoelasticity of the two phases, flow strength, and compatibilizers) on the structure and rheology have all been studied in detail. There is now a fairly comprehensive understanding of flow-induced structural evolution and of the corresponding structure–rheology relationships in these systems, and much of this research has been reviewed in books and review articles [Utracki and Shi (1992); Tucker and Moldenaers (2002); Bucknall and Paul (2009)].
In recent years, there has been significant research on ternary systems composed of two immiscible homopolymers, and a small fraction (few percent) of solid particles that adsorb to the interface between the homopolymers. Much of that research has been reviewed recently [Fenouillot et al. (2009)]. Some of the early research examined the preferential localization of carbon black at the interface between immiscible homopolymers [Sumita et al. (1991); Gubbels et al. (1994); Soares et al. (1995)]. One noteworthy result from these experiments was that high electrical conductivity could be realized at very low particle loadings if the particles were located at the interface of a bicontinuous structure. There have been several more recent studies of nanoclays [Gelfer et al. (2003); Ray and Okamoto (2003); Ray et al. (2004); Si et al. (2006); Fang et al. (2007); Hong et al. (2007); DeLeo et al. (2011a)], fumed silica [Vermant et al. (2004); Thareja and Velankar (2006); Elias et al. (2008); Vermant et al. (2008); Tong et al. (2010); Liu et al. (2012)], and other particles [Li et al. (2009); Hwang et al. (2012)] localizing at the interface between immiscible polymers. Many of these studies showed that addition of particles improved dispersion, e.g., in the case of droplet–matrix blends, addition of particles reduced the drop size. In this regard, the role of particles is analogous to conventional compatibilizers—block copolymers that are adsorbed at the interface between polymers—which also reduce drop size. Accordingly, such particles may be regarded as particulate compatibilizers. With conventional compatibilizers, the improved dispersion often results from their ability to prevent coalescence [Sundararaj and Macosko (1995); Van Puyvelde et al. (2001)]. Analogously, particles were also found to suppress coalescence, e.g., fumed silica particles were able to stop flow-induced coalescence at loadings of \( \sim 1\% \) [Vermant et al. (2004)]. Later research by the same group [Vandebril et al. (2010)] suggests that the interfacial rheology of the particles plays a significant role in coalescence suppression and morphological stabilization.

Beyond improving dispersion however, adding particles to two-phase polymer blends can induce other remarkable effects. The first is that the effect of particles on coalescence is not symmetric: A given particle may be able to prevent coalescence in an A-in-B droplet/matrix blend but not in a B-in-A blend. This was noted by Vermant [Vermant et al. (2004)] with fumed silica particles. Later experiments by our group using a wider variety of particles showed even more extreme asymmetry: A given particle type could either reduce or increase drop size as compared to the particle-free blend depending on which phase was the continuous phase, i.e., particles have the ability to both hinder as well as actively promote coalescence [Thareja et al. (2010)]. Such asymmetric effects are also known with conventional block copolymer compatibilizers [Van Hemelrijck et al. (2005); Martin and Velankar (2007); DeLeo et al. (2011b); Gong and Leal (2012)].

A second remarkable effect due to particle addition is interfacial jamming. Since particles adsorb almost irreversibly at the interface [Binks (2002)], if the interfacial particle coverage is sufficiently high, they can jam the interface giving nonspherical drops. This phenomenon is well-known in small-molecule systems [Asekomhe et al. (2005); Subramaniam et al. (2005); Monteux et al. (2007); Cheng and Velankar (2009); Pawar et al. (2011)], but has also been noted in polymer blends with clay nanoparticles [Si et al. (2006)]. Interfacial jamming has also been proposed as a way to stabilize a bicontinuous morphology: The essential idea is to create a bicontinuous morphology with sufficiently high particle coverage at the interface that structural coarsening can be arrested [Stratford et al. (2005)]. One approach to realizing such a “bijel” morphology is to induce spinodal decomposition of a ternary particle/polymer/polymer system [Gam et al. (2011)]. One may also achieve the initial bicontinuous morphology by blending a two-phase system under suitable conditions and allowing it to coarsen until the interface jams [Cheng (2009)].
A third surprising observation is that addition of particles can sometimes generate a string-like morphology: In droplet–matrix blends in shear flow, droplets were found to be highly deformed along the flow direction [Cheng (2009); Tong et al. (2010)].

Finally droplet–matrix blends with added particles can also show the unusual phenomenon of particle bridging when a single particle adsorbs simultaneously on two drops at once [Horozov and Binks (2006)]. When several such particles bridge across drops, they glue drops together into clusters with major structural and rheological consequences [Thareja and Velankar (2006); Thareja and Velankar (2008); Vermant et al. (2008); Walker et al. (2011); Nagarkar and Velankar (2012)].

In summary, numerous structural changes have been noted due to particle addition: Coalescence suppression or promotion in droplet–matrix blends, jamming due to interfacial crowding, and bridging. Nevertheless, as compared to the comprehensive understanding of particle-free binary polymer blends, our understanding of ternary polymer/particle systems remains poor. In particular, the dependence of flow-induced structure on particle loading, particle size, and particle wettability is all poorly understood. Most glaringly, the relationships between the rheology and the structure are not well understood at all. Indeed, this deficiency is not limited to particle-containing polymeric systems: More generally, rheology of ternary systems composed of two fluids (even oil/water) and one particulate species is poorly characterized. In contrast to the vast literature on particle–matrix suspensions and droplet–matrix emulsions, there is relatively little knowledge about what happens when particles and droplets are both present. The small-amplitude oscillatory behavior has been measured in several cases and there have been attempts to apply the Palierne emulsion model to the frequency response behavior of these systems [Vermant et al. (2004); Elias et al. (2008); Thareja and Velankar (2008)]. But beyond linear viscoelastic behavior, there is very little information; even the most basic issue of dependence of viscosity on the ternary composition is not known. As we shall show in this paper, the dependence of viscosity on composition shows remarkable trends that do not appear to have been recognized previously.

The approach in this paper is to examine model ternary systems composed of rheologically simple bulk fluids and spherical monodisperse particles. The literature on binary blends [Almusallam et al. (2000); Grizzuti et al. (2000); Tucker and Moldenaers (2002); Janssene et al. (2003)] shows enormous success in elucidating flow-induced structure and rheology relationship using polymers (often polydimethylsiloxane, PDMS and polyisobutylene, PIB) that are nearly Newtonian liquids at room temperature. Since the polymers are liquid at room temperature, long experiments are possible without thermal degradation. Since the bulk fluids are Newtonian, all non-Newtonian effects can be ascribed unambiguously to interfacial phenomena. More recently, similar success has been achieved in ternary systems composed of two immiscible polymers and a compatibilizer [Van Hemelrijck et al. (2004); Martin and Velankar (2007); DeLeo et al. (2011b); Gong and Leal (2012)]. It is therefore reasonable to attempt the same approach (using rheologically simple room temperature-liquid systems) to particle-containing blends, and indeed some of the work cited above took this approach. However elucidating the flow—structure—rheology relationships requires characterizing the structure, and this poses major challenges in a system that is liquid at room temperature. The two-phase structure is often on the order of 5–100 μm in size and can be characterized by in situ optical microscopy using a transparent shear cell—at least as long as light scattering is not too severe. However, the particle-scale structure is usually on the 0.1–5 μm scale and is difficult to characterize. Cryo-electron microscopy has been used for this purpose but it is experimentally challenging [Vermant et al. (2004); Binks et al. (2005); Madivala et al. (2009); Vandebril et al. (2010)]. Confocal imaging has been conducted successfully in
oil/water systems, but it requires fluorescent labeling of the particles and/or polymers, and refractive index matching the polymers and particles; furthermore, it can work only with particles that are roughly 1 μm or larger [Dinsmore et al. (2002); Tarimala and Dai (2004); Lee et al. (2012)]. Particle-scale characterization by transmission electron microscopy (TEM) or scanning electron microscopy (SEM) has been conducted with polymer blends that are solid at room temperature (see many of the citations in the second paragraph of this paper), but in such cases the polymers and particles used are fairly complex and do not permit the clean experimental interpretation possible in model systems.

We have developed a new experimental system that addresses these challenges. It is composed of polyethylene oxide (PEO) and PIB as the bulk phases, and monodisperse spherical silica as the particles [Nagarkar and Velankar (2012)]. The melting temperature of PEO is relatively low (65°C) allowing flow experiments to be conducted at 80°C—a temperature at which thermal degradation is slow. The molecular weight of each component is relatively low and hence both fluids are nearly Newtonian under experimental conditions. Upon cooling, the PEO can be crystallized, thus quenching the morphology; removal of the PIB in a selective solvent then allows the structure to be examined by SEM. This system preserves most of the advantages of the room temperature-liquid blends, but offers the additional advantage of allowing structural characterization on the scale of particles. In addition, since the particles are spherical and monodisperse, the complications associated with the high aspect ratios of nanoclays or the complex structure of fumed silicas can be avoided. Finally, the surface wettability and hence contact angle can be varied by chemical modification of the particles. Specifically, modification of the particles with octadecyltrichlorosilane (OTS) made them preferentially wetted by the PIB phase, whereas modification with dichlorodimethylsilane (DCDMS) made them roughly equally wetted by both PDMS and PIB [Nagarkar and Velankar (2012)].

In a previous publication [Nagarkar and Velankar (2012)] using this experimental system, we reported the following observations: (1) Particles promote coalescence regardless of wettability; (2) OTS-modified particles bridge across drops and create a bridged network of drop clusters; (3) at low particle loadings (~0.1 vol. %), the OTS-modified particles are not distributed uniformly on the interface; instead they are heavily concentrated in the bridged region between the drops; (4) at higher particle loadings (~1 vol. %), the surface coverage of OTS-modified particles on the interface is much higher, and bridging and jamming both occur; and (5) the corresponding blends show gel-like behavior in small-amplitude oscillatory experiments. Figure 1 summarizes the physical picture of the morphology that resulted from these observations. The SEM images supporting this physical picture were provided previously, and a limited set of images will be shown later in this paper (Fig. 9). The goal of this paper is to describe the nonlinear rheology of these systems. We will also highlight the differences between blends containing the OTS-modified particles described previously, and blends containing DCDMS-particles.

II. MATERIALS AND METHODS

The materials and methods are identical to those used previously. Briefly, the blends are composed of PEO, PIB, and monodisperse Stober silica particles of diameter 470 nm. The surface of the particles was modified with OTS or DCDMS by methods described previously. Figure 2 shows the SEM images of the particles adsorbed at the PEO/PIB interface, and the large difference in wettability is obvious. The values of contact angle (measured through the PEO phase) of roughly 150° and roughly 90° were calculated previously [Nagarkar and Velankar (2012)].
The blend preparation protocol was identical to that used previously. Blends were designated as Ex-y where “E” represents the dispersed phase (PEO), x is the weight percent of PEO (15, 25, 35, or 45) on a particle-free basis, and y is weight percentage of particles in the blend rounded to the nearest 0.01%. Most of the results in this paper refer to blends with OTS-modified particles, and hence the sample designation does not mention the silane used. In Sec. III E, blends containing DCDMS-modified particles are discussed, and in the corresponding discussion, we will specify the surface modifier explicitly. Finally, for convenience we will use the terms “OTS-particles” instead of “OTS-modified particles,” and “OTS-blends” instead of “blends containing OTS-modified particles.”

The compositions of the various blends studied here are shown in the composition diagrams of Fig. 3. We will discuss the rheological properties along each of the three traces in the ternary composition diagram: (1) The dotted blue line which corresponds to raising

**FIG. 1.** Schematic of effect of OTS-particles in polymer blends. (A) is a particle-free sample and particle concentration rises going from (B) to (D). (B) At low particle loadings, particles are preferentially adsorbed in the bridging region, and drop size is larger than in the particle-free blend. (C) Increased particle loading leads to a heavily bridged structure. (D) Further increase in particle loading reduces drop size.

**FIG. 2.** High resolution images of particles adsorbed on the surface of PEO drops. Images were taken after the PIB was removed in a selective solvent. (A) OTS-modified and (B) DCDMS-modified particles.
particle loading while keeping the weight ratio of the drop phase to the continuous phase fixed, (2) the horizontal green line which varies drop loading while holding the particle loading fixed, and (3) the dashed red “diagonal” line which varies the continuous phase loading. The last trajectory corresponds to raising the particle loading and the drop loading in a fixed proportion. We emphasize that the particle loadings are low, typically under 1 vol. %, and addition of particles to the matrix phase did not have a measurable effect on the bulk rheology. Thus, all the effects of particle addition can be attributable to interfacial phenomena.

The blends were sheared in a controlled stress rheometer (AR 2000, TA Instruments) at 80 °C using a cone and plate geometry with 1° cone and a diameter of 40 mm. The shear history for the most of the data presented in this paper is shown in Fig. S1 (Supplementary Material). Briefly, the sample was presheared at a stress of 250 Pa, and

FIG. 3. (A) Ternary composition diagram. (B) Magnified view of the bottom left corner of the composition diagram. Experiments are conducted along the three lines indicated. Black points correspond to compositions tested.
then shearing was continued at a stress of 50 Pa for 2000 strain units. At the end of shearing, small-amplitude oscillatory experiments were conducted. These were discussed previously [Nagarkar and Velankar (2012)] and will not be mentioned further in this paper. The experiments of Sec. III D used a different shear history, which will be explained separately. The morphology of the blends was examined as described previously. Briefly, the blends were cooled to –35 °C in the rheometer to freeze the PEO drops, the continuous phase removed with n-octane, and the frozen PEO drops examined by SEM.

III. RESULTS: RHEOLOGY OF BLENDS CONTAINING PARTICLES

A. Steady shear viscosity of blends containing OTS-modified particles

Upon applying a fixed stress to the blends, the viscosity shows a complex transient with time (discussed in Sec. III B) before reaching a steady value. These steady shear viscosities are discussed in this section. Independently mixed batches of E35-1.75 blend were examined four times and showed reproducibility to within 15%. Several other samples were examined two times and showed comparable reproducibility.

Figures 4(A) and 4(B) illustrate the effect of particle loading on the steady shear viscosity of the E35-y and E15-y blends, respectively, at two different stress levels. Focusing first on the particle-free blends, the viscosities of E35-0 and E15-0 are indicated by the horizontal lines. In both these blends, the viscosity at 250 Pa is slightly lower than at 50 Pa, indicative of shear-thinning. Such shear-thinning has been previously attributed to the orientation of the interface along the shear direction [Choi and Schowalter (1975); Doi and Ohta (1991)] and this will be discussed further in Sec. IV.

With addition of OTS-particles following the dotted blue line trajectories in Fig. 3, the following trends may be noted for both E15-y and E35-y blends. First, at the lower stress level of 50 Pa, the viscosity of the blends increases sharply over that of the particle-free blend. The magnitude of the increase is large: As little as 3.5 wt. % (roughly 1.6 vol. %) of particles increase the viscosity of the E35 blends roughly twofold. A comparable particle loading in either phase does not cause a measurable viscosity increase, and indeed Einstein’s equation predicts that the viscosity would rise by less than 5%. Second, the particle-containing blends are severely shear-thinning: Raising the stress from 50 to 250 Pa reduces the viscosity of the E35-3.5 by almost twofold—a much greater decrease than for the particle-free blend E35-0. Third and most remarkable, at the higher stress level of 250 Pa, the trend in viscosity with particle loading is not monotonic: Blends with a very low particle loading (e.g., E35-0.35) are less viscous than the corresponding particle-free blend (E35-0), but at higher particle loadings, the viscosity increases to above that of the particle-free blend. Section III E will show that such behavior occurs with the DCDMS-blends as well.

Figure 4(C) traces the viscosity of the blends as the continuous phase fraction is reduced, which corresponds to the dashed red trajectory in Fig. 3. The viscosities of the particle-free blends (corresponding to the lower edge of the ternary diagram in Fig. 3) are in excellent agreement with the Phan-Thien–Pham model [Phan-Thien and Pham (1997)]. Similar good agreement was shown previously [Martin and Velankar (2007)] for particle-free blends. For the particle-containing blends, the viscosity increases sharply. This is not surprising in light of Figs. 4(A) and 4(B): Since drop loading and particle loading each raise the viscosity, increasing both of them proportionately (as happens along the dashed red trajectory in Fig. 3) may be expected to raise the viscosity as well.

In Sec. IV A we will discuss the morphology of blends and note that for the E35-0.07 and E35-0.17 blends at low stress (the points circled in Fig. 4(A)), the drop size is
FIG. 4. Dependence of viscosity on particle loading at (A) 50 Pa stress, and (B) 250 Pa stress. (C) Dependence of viscosity on fraction of the continuous phase.
especially large because the particles promote coalescence. Some of these drops are on the order of the rheometer gap size. Accordingly, these two viscosity values may be affected by confinement effects.

B. Transient creep viscosity of blends with OTS-modified particles

Figures 5(A)–5(C) show the time-evolution of the viscosity during the last creep step of the shear history along each of the three trajectories in the ternary composition

**FIG. 5.** Viscosity during creep at 50 Pa. (A) Effect of particle loading for E35-y blends, (B) effect of drop loading at two fixed particle loadings, and (C) effect of continuous phase loading.
diagram. In all cases, at long times the viscosity reaches an approximately invariant value, and it is these values that were reported in Fig. 4. However Fig. 5 shows that this steady viscosity is not always approached in a monotonic fashion; in many cases, the viscosity shows an overshoot at short shearing times before leveling off. Figure 5(A) shows that for the E35-y blends, the magnitude of the overshoot increases significantly with particle loading if drop loading is held fixed. Figure 5(B) compares blends at different drop loadings while particle loading is held fixed, i.e., along the horizontal line trajectories in Fig. 3. Comparing E35-1.75 vs E15-1.75 for instance, it is clear that reducing drop loading at fixed particle loading reduces the magnitude of the overshoot. Finally Fig. 5(C) shows that reducing the continuous phase loading (dashed red line trajectories) significantly increases the magnitude of the overshoot. Incidentally, when the same results of Figs. 5(A)–5(C) are plotted as a function of strain (not shown), the peak in viscosity appears at strains slightly under 1 strain unit.

Viscosity overshoots are often indicative of breakdown of some large-scale structure in the fluid [Mujumdar et al. (2002)], and in this case, presumably the structure of particle-bridged drop clusters. This will be discussed further later in the paper. In this context however, we have conducted additional experiments on the E35-0.35 blend in which the sheared sample was allowed to stand for extended periods ranging from 30 s to 32 min before initiating creep. There was no significant difference in the overshoot behavior, i.e., under quiescent conditions, there do not appear to be significant changes in the transient creep viscosity. Referring to Fig. 1, this suggests that quiescent annealing does not induce significant changes in drop size or in cluster formation. Previously we have observed [DeLeo and Velankar (2008)] viscosity overshoots in similar blends, but with reactive compatibilizer rather than particles, but in that case, the magnitude of the overshoot increased significantly with quiescent rest time.

C. Recovery after cessation of shear for blends with OTS-modified particles

The elastic recoil after cessation of shear was measured after each of the creep steps. Figure 6 shows the recovery vs time after the very last shearing step at 50 Pa. Figure 6(A) shows the effect of particle loading at a fixed dispersed phase loading of 35%. The particle-free blend E35-0 shows an ultimate recovery of roughly 0.15, somewhat lower than the recovery observed previously for similar systems [Wang and Velankar (2006a, 2006b)]. Upon addition of particles [dotted blue trajectory of Fig. 3(A)], two trends are evident: (1) Particles greatly increase the ultimate recovery, and (2) particles greatly increase the time needed for full recovery. Indeed in some cases, we are not confident that full recovery was completed within the 3 min of the recovery step.

Figure 6(B) shows the effect of continuous phase loading (dashed red line from Fig. 3) on the elastic recoil. As the fraction of continuous phase decreases (i.e., the drop and particle loadings both increase in tandem), the creep recovery increases significantly.

The recoil for the E15-y blends is difficult to measure since it is very small; furthermore since these blends have fairly low viscosities, the recoveries are affected significantly by the residual torque of the rheometer. For this reason, we are unable to compare the E35-y and E15-y blends at a fixed particle loading (horizontal green line trajectory in Fig. 3).

The experimental protocol of Fig. S1 also gives recovery after cessation of shear at 250 Pa. These results are not shown in detail; however, Fig. S2 illustrates the main trend for some selected blends: The recovery is far faster at the higher stress level. In particle-free blends, such faster recovery can be interpreted readily: Previous models
VINCKIER ET AL. (1999); WANG AND VELANKAR (2006a)] for the recovery of droplet–matrix blends show that the retardation time depends on the mean drop size. Thus, the faster recovery of E35-0 at 250 Pa is consistent with a smaller drop size: Upon reducing the stress to 50 Pa, flow-induced coalescence raises the drop size, and slows the recovery. The behavior of the particle-containing blends is more difficult to interpret since it must involve deformation and recovery of the drops which are clustered together. Nevertheless, the faster recovery of particle-containing blends at higher stress level is consistent with having smaller drops at higher stress, and SEM images (Fig. 9) presented later do show that the individual drops in the clusters reduce in size at higher stress.

D. Stress ramp experiments

Sections III A and III B made two observations regarding blends with OTS-modified particles: (1) particle addition makes the blends highly shear-thinning (Fig. 4), and
(2) the viscosity transient during creep shows an overshoot at short shearing times (Fig. 5). Both observations indicate that the blends may possess a yield stress, and this was tested directly by subjecting the blends to a shear history of Fig. S3. The idea underlying this shear history is to create an initial morphology by shearing for an extended period at a fixed stress (either 500 Pa or 50 Pa), and then examining the structural breakdown by a stress ramp. Figure S3 shows that the ramp consists of increasing the stress continuously from 0.5 Pa to 500 Pa in a logarithmic fashion over 10 min. This same shear-and-ramp protocol was applied three times in succession to verify reproducibility. Below, we will only show the results of the first stress ramp at 500 Pa. The latter two stress ramps show good reproducibility, albeit with slightly lower viscosity prior to the yield point.

Figure 7(A) shows the results of these stress ramps for the E35-y blends. As expected the particle-free blend E35-0 shows a low viscosity at all stress levels with a slight decrease in viscosity indicative of shear-thinning as mentioned in Sec. III A. With increasing particle loading, the viscosity at low stress increases sharply; however, the shear-thinning becomes severe as well. At the highest particle loading examined here, the curves show characteristics of a yield-stress fluid. For the E35-1.75 blend, the apparent yield stress is roughly 10–20 Pa.

Before proceeding further, we must note that the high viscosities measured at the low stresses during the ramp are not completely reliable; the AR2000 rheometer measures the rotational velocity of the cone, and if the strain rate is very low (owing to the very low stresses early in the ramp), there may be some errors in the viscosity measurement. Thus the viscosities at stresses below the apparent yield stress are high, but there may be some errors in their absolute values.

Figure 7(B) shows the effect of drop loading at fixed particle loading on the stress ramp behavior; it is clear that the yield-stress behavior becomes weaker with lower drop loading, and blends with 15% drops do not show yield stress-like behavior.

Finally, Fig. 7(C) shows the effect of continuous phase loading on the stress ramp. As expected, the results are a combination of increasing particle loading and increasing drop loading. Accordingly the stress yield stress-like behavior becomes more prominent as the continuous phase loading is decreased.

All these stress ramp results can also be plotted in the form of viscosity vs strain (rather than viscosity vs stress), and in such cases, the yielding appears at a strain on the order of 1 strain unit (values ranging from about 0.3 to 2.3 strain units were noted for various samples). The apparent yield strain reduces with increasing drop loading and increasing particle loading.

Finally, the shear protocol of Fig. S3 also allows a comment on the effect of prestress on the stress ramp behavior by comparing ramps imposed after a prestress of 500 vs 50 Pa. A limited set of results is shown in Fig. S4. The following trends may be noted in samples that show yield-like behavior: (1) The viscosity below the yield point is generally higher at after 50 Pa prestress than after 500 Pa (however as noted above, these values may not be entirely accurate); (2) the value of yield stress is not highly sensitive to the prestress level: In fact across all the samples in Fig. 7 and Fig. S4, apparent yield stresses are always in the 10–25 Pa range; and (3) above the yield point, viscosity is nearly independent of prestress. Thus, tentatively we may conclude that the apparent yielding behavior depends only weakly on prestress.

We must note one caveat on stress ramps: These experiments were conducted with a smooth cone-and-plate geometry. Yielding phenomena can sometimes be accompanied by slip behavior, and some influence of wall slip on the results cannot be ruled out.
FIG. 7. Dependence of viscosity on stress in stress ramp experiments following the protocol of Fig. S2. (A) Effect of particle loading for E35-y blends, (B) effect of drop loading at two particle loadings, and (C) effect of continuous phase loading.
E. Effects of particle wettability: Rheology of blends with DCDMS-modified particles

In Sec. IV, we will interpret the results of the rheology in terms of structural characteristics of the blends. Specifically, we will comment on the major role played by particle bridging. As a prelude to that discussion, it is useful to examine the rheology of a blend in which particles adsorb at the interface but cannot bridge. Previously we had shown that DCDMS-modified particles adsorb roughly symmetrically at the interface, and therefore are incapable of bridging [Nagarkar and Velankar (2012)]. This section briefly describes the rheology of the corresponding blends. These experiments have only been performed on the E35-y series of blends (dotted blue trajectory in Fig. 3) with particle loadings up to 3.5 wt. %.

Figure 8(A) shows that the viscosity of the blends with the DCDMS-modified particles is significantly lower than of blends with the OTS-modified particles. Once again, the viscosity behavior is not monotonic with particle addition: The viscosity reduces at low particle loading as compared to the particle-free blend, before rising again at higher particle loading.

Figure 8(B) shows that the blend with DCDMS-modified particles shows a slight viscosity overshoot during creep experiments; however, the magnitude of the overshoot is substantially smaller than for the blends containing OTS-modified particles.

Finally Fig. 8(C) shows that in the stress ramp experiment, the blends containing DCDMS-modified particles show modest shear-thinning, but not the extreme yield-like behavior of the blends containing OTS-modified particles.

IV. DISCUSSION

Before proceeding, we will summarize the chief rheological consequences of adding interfacially active particles to the blends. It is noteworthy that in all cases, the volume fraction of particles was low, usually less than 1 vol. %, and the particles do not affect the bulk rheology of the phases significantly. Thus, all the effects listed below can be unambiguously attributed to the interfacial activity of the particles. These effects are as follows:

1. At very low particle loadings, both particle types can reduce the viscosity of blends, especially at higher stress levels. At higher particle loadings, the viscosity rises again. The OTS-particles can induce large increases in viscosity.
2. Addition of OTS-particles makes the blends highly shear-thinning, and these blends show apparent yield-like behavior.
3. During startup of steady flow, OTS-blends show an overshoot in viscosity during fixed-stress creep experiments.
4. OTS-particles make the blends more elastic, at least as judged by elastic recoil upon cessation of shear.
5. The effects of particles depend severely on wettability; blends containing DCDMS-modified particles show rheological behavior that is different—even qualitatively—from blends containing OTS-modified particles.

A. Morphology

We now discuss the microstructural origins of these rheological effects. Figure 9(A) shows the structure of the PEO drops (after they have been solidified) for the E35-1.75 blend. The drops are obviously nonspherical, and at higher magnification [Fig. 9(B)],
FIG. 8. Effect of particle wettability on rheological properties. (A) Steady shear viscosity at 50 Pa stress, (B) viscosity transient during steady creep, and (C) viscosity dependence on stress during stress ramp.
many drops are seen to have facets or shallow craters. While images such as Fig. 9(A) provide an accurate picture of drop sizes and shapes, they do not provide an accurate description of the morphology quenched from molten conditions. We showed previously [Nagarkar and Velankar (2012)] that in fact the morphology upon quenching consists of these same drops heavily clustered together due to particle-bridging. The shallow craters or facets in Fig. 9(B) correspond to regions that had been bridged. Most of these clusters are disrupted during selective solvent extraction of PIB, although occasional clusters are still visible. Even higher magnification at particle-scale resolution [Fig. 9(C)] reveals a tight packing of particles. Figure 9 refers to the E35-1.75 sample; similar imaging at much lower particle loading (not shown) reveals that the drops are far larger, sometimes comparable to the rheometer gap. Furthermore, at low particle loadings, there are not sufficient particles to cover the entire surface: Instead, the particles are preferentially located in the bridging regions leaving much of the drop surfaces particle-free. Finally, while our previous research did not discuss the effect of stress, a comparison of Fig. 9(A) (after 50 Pa stress) and Fig. 9(D) (after 500 Pa stress) shows a significant decrease in drop size as stress increases. Similar SEM images across a much wider range of compositions [Nagarkar and Velankar (2012)] led to the physical picture of Fig. 1. To summarize, addition of OTS-modified particles has two morphological consequences. The first is that the OTS-modified particles can bridge across PEO drops. Bridging occurs at all drop loadings and has profound effects on the morphology; most importantly, it glues together the drops into clusters. The second (not evident from Fig. 9, but shown previously) is an increase in flow-induced coalescence: Even at particle loadings as low as 0.07 wt. % (the lowest loading examined), particles were able to greatly increase the drop size.

**FIG. 9.** SEM images of E35-1.75 with OTS-modified particles. (A)–(C) After shearing at 50 Pa stress for 2000 strain units. (D) After shearing at 500 Pa stress for 2000 strain. The scalebars on the top left of each image are 200 µm in (A) and (D), and 10 µm in (B) and (C). Note that (A) and (D) are nearly the same magnification and the difference in drop size is readily apparent.
Our previous article did not present structural characteristics of blends containing DCDMS-modified particles in detail, and we will do so here. Figure 10(A) shows the dispersed phase from E35-1.75-DCDMS sample after being subjected to a stress of 50 Pa. The morphology is altogether different from Fig. 9. First, the size-scale of this structure is significantly larger than the 300 μm gap at the edge of the cone-plate geometry. It is no longer clear whether there are discrete drops in the sample; many of the drops appear to have coalesced together giving the appearance of a bicontinuous structure. The facets or shallow craters evident in the OTS-blends are not evident. At much higher magnification [Fig. 10(B)], the particle surface appears to be tightly packed with particles. It must be emphasized that this sample, which has only ~33 vol. % of dispersed phase, is not likely to be bicontinuous in reality; Fig. 10 is just one portion of the dispersed phase and not representative of the overall morphology.

It is of interest to examine the dependence of the morphology of the DCDMS blends at other blend compositions. However, the large size-scale of the morphology in Fig. 10 is likely to be heavily influenced by the gap size, and indeed under these conditions, the SEM images are not highly reproducible: In some cases, we see long string-like phases rather than a morphology that resembles bicontinuity. Therefore, we have conducted additional experiments examining the effect of particle loading and drop loading, but at a higher stress level of 500 Pa, at which the size-scale of the morphology is smaller, and therefore less affected by (although still not altogether independent of) the rheometer gap.

Figure 11 shows the effect of particle loading on E35-y DCDMS-blends quenched after shearing at 500 Pa. The particle-free blend E35-0-DCDMS [Fig. 11(A)] shows a morphology composed of small spherical drops with the largest drops being much less than 100 μm in diameter. Addition of 0.35 wt. % particles [Fig. 11(B)] greatly increases the drop size showing that the DCDMS particles strongly promote coalescence. This was reported in our previous research as well. At this loading, particle-scale images [Fig. 11(B1)] show particles on the interface, but the coverage is far short of complete.
FIG. 11. Morphology of E35 blends with DCDMS-modified particles at various particle loadings after shearing at 500 Pa. (A) is the particle free blend, whereas particle loading increases from (B) to (D). Right column of images is at higher magnification. Scalebars are 500 µm in the left column and 5 µm in the right column.
At a higher particle loading [Figs. 11(C) and 11(C1)], a string-like morphology appears, and the interface appears jammed with particles. To our knowledge, this is the first demonstration of a flow-induced interfacially jammed blend morphology. With an even higher particle loading [Figs. 11(D) and 11(D1)], the size-scale of the morphology reduces significantly and the aspect ratio of the drops reduces sharply although they are still not spherical.

These images suggest the following physical picture (Fig. 12) for the effect of DCDMS particles on the structure of the blends. At low particle loadings, the particles greatly promote coalescence. Since the particle loading is too low to cover the interface completely, drops are spherical. The drop size is chiefly controlled by a steady state balance between flow-induced breakup and flow-induced coalescence. As particle loading increases, the surface coverage increases to the point where the drops are forced to take on a nonspherical shape so that the larger surface area required for accommodating the particles can be maintained with the same drop volume. We speculate that under these conditions, the morphology can be either string-like or bicontinuous depending on the stress level. Finally, at even higher particle loading, the continuing need to accommodate all the particles on the interface forces a decrease in the size-scale of the dispersed phase.

Finally, we discuss the reasons for the promotion of flow-induced coalescence. As mentioned in the introduction, we had previously noted instances of particle-induced coalescence in polymer blends [Thareja et al. (2010)], and had speculated that the “bridging–dewetting” mechanism was involved. Consider two drops colliding when the upper drop already has a particle adsorbed at the equilibrium contact angle in the region of the thin film separating the drops [Fig. 13(A)]. During the collision, the particle bridges across the thin film and adsorbs on both drops simultaneously; at this instant, the lower contact line is not at the equilibrium contact angle [Figs. 13(B) and 13(D)], and therefore must recede across the particle. Whether this particle bridge is stable or not

**FIG. 12.** Schematic of the effect of adding DCDMS-modified particles in polymer blends. (A) is the particle-free sample, and the particle concentration rises going from (B) to (D). (B) At low particle loadings, the interface is not fully covered and the particles promote coalescence. At higher particle loadings [(C) and (D)], the interface is jammed and drop size reduces.
depends on how far the contact line recedes, i.e., on the wettability of the particle. If the particle is preferentially wetted by the drop phase [Fig. 13(B)], the lower contact line recedes across the particle until the two contact lines meet (C) and coalescence occurs. If the particle is preferentially wetted by the continuous phase, \( \theta > 90^\circ \) is desired. The contact line recedes until the desired contact angle is reached (E) and a stable bridge is formed.

In our previous research [Thareja et al. (2010)], the contact angles of the particles were unknown and hence bridging–dewetting was proposed as a hypothesis. In the present case however, the contact angles of the particles are known from the particle-scale SEM images (Fig. 2), and hence the bridging–dewetting mechanism may be examined more critically. The DCDMS-particles are roughly equally wetted by the two phases, and hence bridging–dewetting can indeed explain the coalescence promotion. In contrast, the OTS-particles are preferentially wetted by the continuous phase and hence stable bridging is expected, and indeed bridged clusters were observed at all compositions. Thus in the case of OTS-particles, bridging–dewetting, at least in the simplistic form of Fig. 13, cannot readily explain the promotion of coalescence. Indeed, our observations of coalescence promotion seem to contradict previous reports that particles wetted by the continuous phase bridge across drops and prevent coalescence [Ashby et al. (2004); Stancik and Fuller (2004); Horozov and Binks (2006)]. We speculate that in our OTS-particle case, the accelerated coalescence results from the fact that the region outside the bridged monolayer is particle-free and hence susceptible to coalescence; accordingly, the perturbations in this region during applied flow are sufficient to rupture the thin film separating...
the drops. Coalescence is promoted chiefly because bridging offers a means of holding the drops in close proximity. Thus flow-induced coalescence can be suppressed only at higher surface coverages when the entire drop surface—even the region outside of the bridged monolayer—is nearly covered with particles. In contrast, quiescent coalescence can be suppressed even when the region outside the bridged monolayer is particle-free [Horozov and Binks (2006)]. This explanation is speculative; other potential explanations, such as a change in the van der Waals attraction between the drops due to particle adsorption, cannot be ruled out.

B. Rheology

Figures 1 and 12 allow a clear interpretation of the rheology of the ternary blends. We will focus first on most extreme rheological consequence of addition of particles—the yield-like behavior. Since the blends with DCDMS-modified particles do not show such behavior, we may attribute such behavior to the fact that the OTS-modified particles bridge drops into clusters. While our structural characterization is not able to obtain the morphology in situ after flow, the fact that a yield stress exists suggests that the clusters form a percolating network. Indeed similar yield behavior was noted [Lee et al. (2012)] due to particle bridging in oil/water systems, albeit at far higher particle loadings (8–17 vol. % as compared to ~1 vol. % in this paper). In that paper, owing to the higher particle loading, the morphology appeared similar to a polyhedral foam. We may also draw a strong analogy to the highly nonlinear viscosity vs stress relationship in wet granular systems. The most familiar example of this is the yield stress developed by sand by adding a small amount of water; the water drops form menisci [Fig. 14(B)] that connect together the particles into percolating clusters, allowing sand castles to be constructed. Similar yielding behavior has been examined in particle suspensions containing a small amount of a second fluid that partially wets the particles [Cavalier and Larche (2002); Koos and Willenbacher (2011)]. Such wet granular systems or suspensions in pendular or capillary states are also ternary fluid/fluid/particle systems similar to the polymer blends here, but in a different region of the ternary composition diagram (Fig. 3). Indeed, as illustrated in Fig. 14, our particle-bridged drops may be considered the exact opposite of menisci-bridged particles, and it is not surprising to see similar rheological behavior.

In addition to the yield stress, another notable difference between the DCDMS and the OTS-blends is the overshoot of viscosity during startup of shear. Overshoots are commonly seen across a wide variety of systems, and previous articles [e.g., Groot and Agterof (1994); Whittle and Dickinson (1997); Carrier and Petekidis (2009);
Koga et al. (2009) have cited examples of a variety of systems, e.g., molten polymers, hard sphere suspensions, fiber suspensions, attractive suspensions, associating polymers, emulsions, and foams, which show such viscosity behavior, although most of this literature conducts experiments at a specified rate rather than stress. Such overshoots can often be attributed to the breakdown of some associated structure prior upon applying flow, and indeed, the point of maximum viscosity can be regarded as a definition of the yield point [Moller et al. (2006)]. Our own group has seen viscosity overshoots in polymer blends with reactive compatibilizer, a system in which optical microscopy showed formation of large drop clusters [DeLeo and Velankar (2008); DeLeo et al. (2011b)] qualitatively similar to those seen here.

Finally, there has been at least one article [Herzig et al. (2007)] showing that particle-stabilized bicontinuous morphologies (bijels) show a yield stress as well. In that case, the yield-like behavior cannot be attributed to bridging since the particles are roughly symmetrically adsorbed on the interface analogous to the DCDMS-modified particles. Instead, the yielding behavior was attributed to the fact that the bicontinuous structure allows the interfacial particles to form a percolating network. Analogously, DCDMS-blends might also show yield-like behavior if a bicontinuous jammed morphology were to be realized, e.g., by shearing blends with roughly equal volume fraction of PEO and PIB. We have not conducted experiments in this region of parameter space.

One last aspect of the rheology worthy of comment concerns the nonmonotonic behavior of viscosity with particle loading: In Fig. 5, we pointed out that the viscosity of blends at low particle loadings (e.g., E35-0.35) was lower than of the corresponding particle-free blend (E35-0), especially at high stress. This is true for the blends containing DCDMS-particles as well [Fig. 8(A)], and hence cannot be attributed to bridging-induced clusters. We believe that this unusual behavior is attributable to flow-induced coalescence. Specifically, it is well-known that the steady shear viscosity of blends depends on the orientation of the drops, which in turn depends on capillary number $Ca$, which is the ratio of the viscous stress to the Laplace pressure [Frankel and Acrivos (1970); Choi and Schowalter (1975)]. At small $Ca$ values, the drops remain nearly spherical and the orientation is negligible. With increasing $Ca$, the drops deform and orient along the flow direction. These changes in orientation induce shear-thinning as viscosity reduces from its zero shear value to a lower value that depends on the capillary number. Since a low level of particle addition was shown to greatly increase the drop size, we suggest that the correspondingly high capillary numbers reduce the viscosity to values below that of the particle-free blend. As the particle loading increases, the drop size (and hence $Ca$) reduces, and the viscosity increases again. Certainly, the more complex effects of particles—bridging and stabilization of nonspherical drops—would also contribute to the viscosity increase as particle loading increases.

V. CONCLUSIONS

In summary, we have examined the effect of small (~1 vol. %) of particles to model polymer blends with a droplet–matrix morphology. We find that the rheological changes due to particle addition include large increases in the viscosity, highly shear-thinning (or yield-like) behavior, and overshoots in viscosity during startup of steady shear. We interpret the results in terms of the morphological changes—the bridging and jamming phenomena that occur due to the OTS-particles, and the jamming phenomena due to DCDMS particles. Both particle types are able to promote flow-induced drop coalescence at low particle loadings, which is unexpected in light of previous results that bridging particles stabilize drops against coalescence.
In a qualitative sense, the foremost conclusions of this paper are (1) particles affect the morphology and rheology of polymer blends even at low volumetric loadings as low as 0.1 vol. %, and (2) the effects of morphology and rheology depend severely on the wettability of the particles. These results are likely to apply not just to polymer blends, but also to small-molecule systems (oil/water) although in that case charge interactions will likely play a significant role as well. It is remarkable that even though particle/liquid suspensions and liquid/liquid emulsions have been studied for decades, the rheology of ternary particle/liquid/liquid systems remains poorly characterized.

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References


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