Extensional and shear flows, and general rheology of concentrated emulsions of deformable drops

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The rheology of highly concentrated monodisperse emulsions is studied by rigorous multidrop numerical simulations for three types of steady macroscopic flow, (i) simple shear ($\dot{\gamma}x_2$, 0 0), (ii) planar extension (PE) ($\dot{\Gamma}x_1$, $-\dot{\Gamma}x_2$, 0) and (iii) mixed $(\dot{\gamma}x_2, \dot{\gamma}\chi x_1, 0)$, where $\dot{\gamma}$ and $\dot{\Gamma}$ are the deformation rates, and $\chi \in (-1, 1)$ is the flow parameter, in order to construct and validate a general constitutive model for emulsion flows with arbitrary kinematics. The algorithm is a development of the multipole-accelerated boundary-integral (BI) code of Zinchenko & Davis (J. Fluid Mech., vol. 455, 2002, pp. 21-62). It additionally incorporates periodic boundary conditions for (ii) and (iii) (based on the reproducible lattice dynamics of Kraynik-Reinelt for PE), control of surface overlapping, much more robust controllable surface triangulations for long-time simulations, and more efficient acceleration. The emulsion steady-state viscometric functions (shear viscosity and normal stress differences) for (i) and extensiometric functions (extensional viscosity and stress cross-difference) for (ii) are studied in the range of drop volume fractions c = 0.45-0.55, drop-to-medium viscosity ratios $\lambda = 0.25-10$ and various capillary numbers Ca, with 100–400 drops in a periodic cell and 2000–4000 boundary elements per drop. High surface resolution is important for all three flows at small Ca. Large system size and strains $\dot{\gamma}t$ of up to several thousand are imperative in some shear-flow simulations to identify the onset of phase transition to a partially ordered state, and evaluate (although still not precisely) the viscometric functions in this state. Below the phase transition point, the shear viscosity versus Ca shows a kinked behaviour, with the local minimum most pronounced at $\lambda = 1$ and c = 0.55. The $\lambda = 0.25$ emulsions flow in a partially ordered manner in a wide range of Ca even when c = 0.45. Increase of λ to 3–10 shifts the onset of ordering to much smaller Ca, often outside the simulation range. In contrast to simple shear, phase transition is never observed in PE or mixed flow. A generalized five-parameter Oldroyd model with variable coefficients is fitted to our extensiometric and viscometric functions at arbitrary flow intensities (but outside the phase transition range). The model predictions compare very well with precise simulation results for strong mixed flows, $\chi = 0.25$. Time-dependent PE flow is also considered. Ways to overcome the phase transition and drop breakup limitations on constitutive modelling are discussed.

Key words: boundary integral methods, emulsions, rheology

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1. Introduction

An outstanding problem for any non-Newtonian liquid is in the formulation of a constitutive equation for the stress tensor valid in a broad class of kinematic conditions, not necessarily those encountered in rheological experiments (like shear flow, steady or oscillatory). Knowledge of such an equation would allow the solution of boundary-value problems of non-Newtonian hydrodynamics in complex geometries of technological interest. Obviously, rheological experiments alone, for all their value, are not sufficient to formulate a constitutive equation. The importance of microstructural analysis for general constitutive modelling was emphasized by Hinch & Leal (1975). Emulsions, i.e. dispersions of one immiscible liquid in another liquid, represent an important class of non-Newtonian liquids encountered in many applications, including the food and pharmaceutical industries, oil recovery and transportation. Early analytical theories (Schowalter, Chaffey & Brenner 1968; Frankel & Acrivos 1970; Barthés-Biesel & Acrivos 1973a,b) were able to produce general constitutive models for dilute emulsions of non-interacting drops limited by small deformations. Along these lines, Vlahovska, Loewenberg & Bławzdziewicz (2005) and Vlahovska, Bławzdziewicz & Loewenberg (2009) completed the $O(\varepsilon^3)$ theory for small deformations (where ε is the shape deviation from spherical relative to the drop size). However, the complexity of the analysis has restricted rheological applications so far, and no constitutive equation has resulted yet from the complete $O(\varepsilon^3)$ theory.

Outside this regime, numerical simulations for particular cases provide valuable insights into the emulsion rheology, although they have not led so far to a general constitutive model. Kennedy, Pozrikidis & Skalak (1994) calculated the deformation and rheological response of a single drop in a simple shear at finite drop deformations. Loewenberg & Hinch (1996) offered the first shear-flow dynamical simulations for random non-dilute monodisperse emulsions, with up to N = 12 drops (in a periodic cell) and up to 30% drop volume fractions. Their direct boundary-integral (BI) code had an $O(N^2 N_{\Delta}^2)$ scaling in the system size N and in the number N_{Δ} of triangular boundary elements per drop ($N_{\Delta} = 320$ in their work). Obviously, larger simulations (see below) require alternative approaches. Using the same implementation, Loewenberg (1998) presented additional calculations at 30% concentration and offered a mean-field criterion for drop breakup. This criterion was mostly confirmed in experiments with emulsion concentrations $c \leq 0.7$ and moderate drop-to-medium viscosity ratios λ (Jansen, Agterof & Mellema 2001), although not for higher concentrations and much smaller λ (Golemanov *et al.* 2008). This subject is reviewed at some length in Derkach (2009). For single drops in shear flow, breakup simulations are reported in several studies (e.g. Cristini et al. 2003).

Zinchenko & Davis (2002, hereafter referred to as paper I), offered the first large-scale dynamical simulations of monodisperse emulsions in shear flow by a multipole-accelerated BI algorithm (a development from the multidrop sedimentation code of Zinchenko & Davis 2000). It was possible to handle both matching ($\lambda = 1$) and contrast ($\lambda = 3$) viscosities, up to c = 0.55, N = 100-200, $N_{\Delta} = 1000-1500$ and up to $\sim 10^4$ time steps to systematically study the steady-state viscometric functions (emulsion viscosity and normal stress differences). An increased system size ($N \sim 100$) was particularly important to accurately describe the first normal stress difference at high concentrations, while high surface resolutions are paramount to accurately reach the limit of small capillary numbers *Ca*. This limit *Ca* \ll 1 is arguably the most difficult case in emulsion flow simulations at high *c*, due to geometrical blockages between nearly non-deformed drops. In retrospect, we have found all the results of paper I to be accurate, except for $\lambda = 1$ with $c \ge 0.45$ and small $Ca \le 0.05$, where they somewhat overestimate the emulsion viscosity and the first normal stress difference, due to still insufficient surface resolutions and shorter than necessary simulation times. As a consequence of this accuracy deterioration at small *Ca*, too sharp shear thinning was predicted (compared with the results of the present study which are far more accurate in this range). Using the same code as in paper I, Zinchenko & Davis (2004) even increased resolutions to $N_{\Delta} = 6000-9000$ for $\lambda = 1$ and c = 0.45-0.5 to refine the results at $Ca \leq 0.05$. However, the simulated strains were still limited to 25–45, thus leaving some of the small-*Ca* viscosity values in Zinchenko & Davis (2004) on the metastable level and, hence, slightly overestimated. Much larger strains are required to elucidate the difficult phenomenon of flow-induced 'phase transition' to a partially ordered emulsion in simple shear. Neither study (paper I or Zinchenko & Davis 2004) included interesting extreme viscosity ratios ($\lambda \ll 1$ or $\lambda \gg 1$).

Regarding potential applications to non-Newtonian hydrodynamics of emulsions, there appears to have been too much emphasis in the literature on simple shear. Neither steady nor oscillatory shear flow can mimic the history of deformation of a material element in realistic large-strain flows through complex geometries. 'Any reasonably abrupt change in geometry... will generate a flow with an extensional component and, in particular, flows through a sudden contraction or out of an orifice often lead to flow characteristics which cannot be predicted on the basis of shear viscosity alone.... The historical convention of matching only shear flow data with theoretical predictions in constitutive modelling may have to be rethought' (Barnes, Hutton & Walters 1989). In recent experiments (Rózanska et al. 2014), the Trouton ratio of the extensional and shear viscosities was variable and reported to reach 11 at 70 % emulsion concentrations (versus 3 for Newtonian liquids). Successful simulations of complex microstructures in extensional flows to large strains necessarily require that the lattice of periodic cells (each containing many drops, etc.) repeats itself after some strain period, so that cells could be rebuilt to continue simulation indefinitely. Such lattice self-reproducibility is obvious for shear (and has been heavily exploited) but not for extensional flows. Kravnik & Reinelt (1992) discovered that no lattice is reproducible in general extensional flows. An exception is planar extension (PE), where an infinite number of reproducible lattices exist, the 'critical' lattice being most convenient with minimum skewage during the flow. The Kraynik-Reinelt lattice geometry gained much popularity in molecular dynamics simulations. More recently, Hunt, Bernardi & Todd (2010) showed how to combine this geometry with simple matrix transformations to also provide periodic boundary conditions for mixed-flow simulations. In fluid mechanics, we are only aware of Ahamadi & Harlen's (2008) extensional flow simulations for 2D suspensions using the Kraynik-Reinelt lattice geometry. To the best of our knowledge, there are no extensional or mixed-flow simulations for emulsions.

Of course, in real complex flows the history of deformation of a material element is neither simple shear nor pure extension (uniaxial or PE). A recent novel approach (Martin, Zinchenko & Davis 2014) shows how to build a general constitutive model for a non-Newtonian liquid incorporating the results from both simple shear and PE. Namely, a generalized five-parameter traceless Oldroyd equation is assumed, with variable coefficients postulated to depend on one instantaneous flow invariant. In principle, these five coefficients can be found by simultaneously fitting the equation to three viscometric and two extensiometric (in PE) functions at arbitrary flow intensities. This general approach to constitutive modelling was tested in Martin *et al.* (2014) against exact BI results for dilute emulsions of non-interacting drops with small to large deformations, and found to be rather accurate in a broad class of kinematics, both Lagrangian-steady and unsteady.

The present paper seeks to explore in detail the steady-state rheological functions for concentrated emulsions in the two base flows – simple shear and PE – by large-scale dynamical simulations, and then use them simultaneously, in the manner of Martin et al. (2014) for a general constitutive model. Precise simulation results are also generated for mixed flows and used to validate this constitutive model. In § 2, the problem formulation and the general solution scheme for all three types of flow are discussed, including the implementation of periodic boundaries and the BI equations. The version of multipole acceleration, more flexible and efficient than in paper I, is outlined in §3. Miscellaneous features of the algorithm, discussed in §4, include control of surface overlapping and much more robust, compared with paper I, control of surface triangulations. These features have allowed for unlimited long-time simulations and accurate time averaging. In §5, the whole code is validated and tested for efficiency in the cases of PE and mixed flows. In §6, the steady-state extensiometric functions for PE (the effective viscosity and stress cross-difference) are calculated for c = 0.45-0.55, a broad range of viscosity ratios $\lambda = 0.25-10$ and up to drop breakup. The analysis of the viscometric functions for c = 0.45-0.55 in §7 is a major extension of the work performed in paper I. Not only have two new viscosity ratios $\lambda = 0.25$ and 10 been added, but we have also increased the simulated strains by 1.5–2 orders of magnitude (to hundreds for $\lambda \neq 1$ and thousands for $\lambda = 1$), the maximum system size to N = 400 and roughly doubled typical resolutions. These improvements are critical to shed light on the phenomenon of flow-induced partial ordering and related anomalous kinked behaviour of the viscometric functions, and evaluate the onset of phase transition for each value of c and λ . Still, some severe ergodic difficulties remain in the transition range, as discussed in §7. We also comment on the deep distinction between simple shear and PE simulations. In §8, the constitutive modelling scheme of Martin et al. (2014) is applied (with some improvements) to concentrated emulsions, based on the extensiometric and viscometric functions from §§6 and 7. In §9, the constructed constitutive model is validated against precise simulations for mixed flows, both rheologically 'strong' and 'weak'; an example of a time-dependent PE flow is also included in the validation. In the conclusions ($\S10$), we comment on possible ways to overcome the phase transition and drop breakup limitations on constitutive modelling.

All calculations have been performed as single core on several multicore PCs, with an i7 4960X 3.6 GHz CPU being the fastest. The full multiparameter study reported in this work required a large number (\approx 250) of independent runs. We view, therefore, the lack of code parallelization not as a drawback, since it was beneficial to simply run many calculations concurrently with 100% efficiency, avoiding communication bottlenecks typical of parallel computations.

2. Problem formulation and the general solution scheme

Consider an infinite set of three-dimensional deformable Newtonian drops of viscosity μ' freely suspended in a Newtonian medium of viscosity $\mu_e = \mu'/\lambda$ and subject to a steady macroscopic flow $\boldsymbol{u}_{\infty}(\boldsymbol{x}) = \nabla \boldsymbol{u}_{\infty} \cdot \boldsymbol{x}$ with a constant velocity gradient matrix $\nabla \boldsymbol{u}_{\infty}$. The inertial effects are assumed to be negligible on the microscale, and the Stokes equations apply to the fluid motion inside and outside the drops. An equivalent radius of non-deformed drops (assumed to be the same for all drops) is a_o , and the capillary number is defined as $Ca = \mu_e (2I_2)^{1/2} a_o/\sigma$ through the second invariant $I_2 = \text{tr}(\boldsymbol{E}_{\infty}^2)$ of the rate-of-strain tensor \boldsymbol{E}_{∞} for the macroscopic flow \boldsymbol{u}_{∞} . The system is surfactant-free with a constant interfacial tension σ . At

each time $t \ge 0$, the infinite drop system is obtained from the basic configuration of $N \gg 1$ drops with surface centroids $\mathbf{x}_1^c, \ldots, \mathbf{x}_N^c$ in the basic cell V by triply periodic continuation into the whole space with periods $\mathbf{e}_1, \mathbf{e}_2, \mathbf{e}_3$; the cell $V = \{\xi^i \mathbf{e}_i, 0 \le \xi^i < 1\}$ is the span of $\mathbf{e}_1, \mathbf{e}_2$ and \mathbf{e}_3 . Initially, at t = 0 the basis vectors $\mathbf{e}_1, \mathbf{e}_2, \mathbf{e}_3$ are $\mathbf{e}_1^o, \mathbf{e}_2^o, \mathbf{e}_3^o$. As time proceeds, the drop shapes and positions evolve, and so do the lattice vectors $\mathbf{e}_1(t), \mathbf{e}_2(t), \mathbf{e}_3(t)$. However, for proper long-time averaging of the results, it is imperative that the system remains spatially periodic for all times $t \ge 0$, and with limited skewage of V. The latter is achieved, when necessary, by systematic periodic cell restructuring, but is only possible for three types of flow:

simple shear:
$$\boldsymbol{u}_{\infty}(\boldsymbol{x}) = (\dot{\gamma}x_2, 0, 0), \quad Ca = \mu_e \dot{\gamma}a_o/\sigma;$$
 (2.1a)

planar extension (PE):
$$\boldsymbol{u}_{\infty}(\boldsymbol{x}) = (\dot{\Gamma}x_1, -\dot{\Gamma}x_2, 0), \quad Ca = 2\mu_e \dot{\Gamma}a_o/\sigma;$$
 (2.1b)

mixed flow:
$$u_{\infty}(x) = (\dot{\gamma}_m x_2, \dot{\gamma}_m \chi x_1, 0), \quad Ca = (1 + \chi) \mu_e \dot{\gamma}_m a_o / \sigma.$$
 (2.1c)

Here, $\dot{\gamma}$, $\dot{\Gamma}$, $\dot{\gamma}_m > 0$ and $-1 < \chi \leq 1$; the capillary number expressions are listed for future reference. We use the notation $\dot{\gamma}_m$ for mixed flow to distinguish it from the simple shear rate; $\chi = 1$ gives a flow equivalent to PE (Rallison 1981, 1984; Bentley & Leal 1986).

The main goal of the present work is to study the steady-state emulsion rheology in these three types of flow (2.1a-c), and use it for general constitutive modelling. The relevant non-dimensional rheological functions are defined through the emulsion stress tensor Σ and its deviatoric part $T = \Sigma - (1/3) \operatorname{tr}(\Sigma) I$ as

simple shear:
$$\mu_{sh} = \frac{\Sigma_{12}}{\mu_e \dot{\gamma}}, \quad N_1^{sh} = \frac{\Sigma_{11} - \Sigma_{22}}{\mu_e \dot{\gamma}}, \quad N_2^{sh} = \frac{\Sigma_{22} - \Sigma_{33}}{\mu_e \dot{\gamma}};$$
 (2.2a)

PE:
$$\mu_{pe} = \frac{\Sigma_{11} - \Sigma_{22}}{4\mu_e \dot{\Gamma}}, \quad N_{pe} = \frac{\Sigma_{11} + \Sigma_{22} - 2\Sigma_{33}}{4\mu_e \dot{\Gamma}};$$
 (2.2b)

mixed flow:
$$v_{ij} = \frac{I_{ij}}{\mu_e \dot{\gamma}_m (\chi + 1)}$$
. (2.2c)

Obviously, ν_{12} is interpreted as the non-dimensional mixed-flow viscosity, in addition to the similar quantities μ_{sh} and μ_{pe} .

2.1. Implementation of periodic boundaries

Simple shear. The reproducible lattice of points $\{m^i e_i\}$, where m^i are arbitrary integers, has long been known for shear flow (2.1*a*), and is extensively used in molecular dynamics and suspension flow simulations. It has also received applications in emulsion rheology studies (Loewenberg & Hinch 1996, paper I). At t = 0, the basic cell V is a cube $[0, L)^3$, with the side L, which we use as the length scale. For t > 0 the basis vectors $e_1 = (1, 0, 0)$ and $e_3 = (0, 0, 1)$ are unchanged. Once the strain $\dot{\gamma}t$ crosses a half-integer value, the lattice is restructured by taking $e_2 - e_1$ as the new e_2 -vector, to make, in general, $e_2 = (\gamma, 1, 0)$, where $\gamma = \dot{\gamma}t - [\dot{\gamma}t + 1/2] \in [-1/2, 1/2)$ and the brackets denote the integer part. Figure 1(*a*) (two-dimensional slice) shows the simple mapping to translate the drop centres from the old cell V to the new cell to continue the simulation. Such restructuring ensures the minimal possible skewage at all t > 0.

Planar extension. Far less obvious is the existence of reproducible lattices in PE found by Kraynik & Reinelt (1992). Following their results for a 'critical lattice'

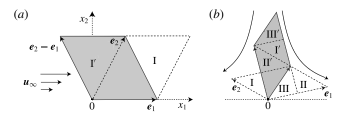


FIGURE 1. Mapping between the old and new (shaded) periodic cells at the restructuring moments for (*a*) simple shear and (*b*) planar extension. Region I is translated to I', region II to II', etc. The third dimension x_3 is not shown.

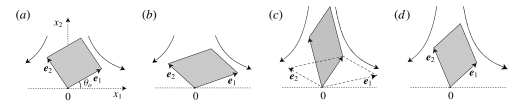


FIGURE 2. Dynamics of the computational cell in PE. At the Hencky strain of $\dot{\Gamma}t = \varepsilon_p/2$, the cell is restructured (the shaded area is the new cell) to continue the simulation. The third dimension is not shown; (a) $\dot{\Gamma}t = 0$, (b) $\dot{\Gamma}t = \varepsilon_p/4$, (c) $\dot{\Gamma}t = \varepsilon_p/2$, (d) $\dot{\Gamma}t = 3\varepsilon_p/4$.

(with minimal skewage), we choose a square with side *L* inclined at angle $\theta_o = \tan^{-1}[(\sqrt{5} - 1)/2] \approx 31.7^\circ$ to the x_1 -axis as the two-dimensional slice of the initial cubic periodic cell *V* (figure 2*a*). Again, *L* is used as the length scale. The initial lattice vectors then become $e_1^o = (\cos \theta_o, \sin \theta_o, 0)$ and $e_2^o = (-\sin \theta_o, \cos \theta_o, 0)$. Figure 2 shows how the lattice reproducibility is achieved in the Hencky strain interval of $\dot{\Gamma}t = \varepsilon_p$, with $\varepsilon_p = \ln[(3 + \sqrt{5})/2] \approx 0.9624$. By $\dot{\Gamma}t = \varepsilon_p/2$, the lattice dynamics $de_i/dt = diag(\dot{\Gamma}, -\dot{\Gamma}, 0)e_i$ brings the basic periodic cell *V* to a shape shown as the dashed-line parallelogram in figure 2(*c*). At this time moment, switching to $e_1 + e_2$ and $e_1 + 2e_2$ as the new lattice vectors and integrating the lattice dynamics further to $\dot{\Gamma}t = \varepsilon_p$ reproduces the initial cell shown in figure 2(*a*), due to the choice of θ_o and ε_p . In simulations, such a lattice restructuring must be carried out at all half-integer values of $\dot{\Gamma}t/\varepsilon_p = n + 1/2$, but there is no need to match (n + 1/2) precisely. The vectors e_1 and e_2 are replaced with $e_1 + e_2$ and $e_1 + 2e_2$ once $\dot{\Gamma}t$ crosses a half-integer value, to make, in general,

$$\boldsymbol{e}_1 = (\cos\theta_o \mathbf{e}^{\Gamma}, \sin\theta_o \mathbf{e}^{-\Gamma}, 0), \quad \boldsymbol{e}_2 = (-\sin\theta_o \mathbf{e}^{\Gamma}, \cos\theta_o \mathbf{e}^{-\Gamma}, 0), \quad (2.3a,b)$$

where

$$\Gamma = \dot{\Gamma}t - \varepsilon_p \left[\frac{1}{2} + \frac{\dot{\Gamma}t}{\varepsilon_p}\right] \in \left[-\frac{\varepsilon_p}{2}, \frac{\varepsilon_p}{2}\right).$$
(2.4)

Prior to a jump in e_1 and e_2 , mapping between the old and new periodic cells (figure 1b) is used to recalculate the new drop positions after the cell restructuring. Namely, drops with centroids in region I are translated by $e_1 + e_2$ to region I' of the new cell. Region II is translated by e_2 to II', region III – by $e_1 + 2e_2$ to III'.

Strong mixed flows. Kraynik & Reinelt (1992) noted that their reproducible lattice dynamics for PE can be generalized to other macroscopic flows with a constant

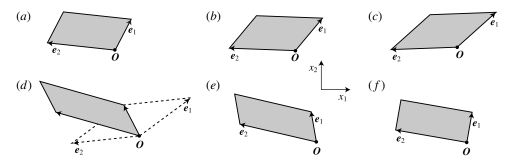


FIGURE 3. The geometry of the computational cell in a strong mixed flow ($\chi = 0.25$) at $\dot{\gamma}_m t/\varepsilon_p = 0$ (a), 1/2 (b), 2/3 (c), 1 (d), 4/3 (e) and 5/3 (f). In (d), the cell is restructured (the shaded area is the new cell) to continue the simulation.

velocity gradient matrix ∇u_{∞} , if a diagonalizing transformation

$$\nabla \boldsymbol{u}_{\infty} = \boldsymbol{\mathsf{S}} \operatorname{diag}(\dot{\boldsymbol{\Gamma}}, -\dot{\boldsymbol{\Gamma}}, 0) \boldsymbol{\mathsf{S}}^{-1}$$
(2.5)

exists between ∇u_{∞} and the velocity gradient matrix for PE, with a real-valued matrix **S**. Indeed, the required lattice vector dynamics $d\mathbf{e}_i/dt = \nabla u_{\infty} \cdot \mathbf{x}$ is reduced to the PE flow dynamics $d\mathbf{\tilde{e}}_i/dt = \text{diag}(\dot{\Gamma}, -\dot{\Gamma}, 0)\mathbf{\tilde{e}}_i$ for auxiliary vectors $\mathbf{\tilde{e}}_i = \mathbf{S}^{-1}\mathbf{e}_i$. The restructuring transformation $(\mathbf{\tilde{e}}_1, \mathbf{\tilde{e}}_2) \rightarrow (\mathbf{\tilde{e}}_1 + \mathbf{\tilde{e}}_2, \mathbf{\tilde{e}}_1 + 2\mathbf{\tilde{e}}_2)$ to the new basis vectors in the PE flow space at half-integer values of $\dot{\Gamma}t/\varepsilon_p$ begets the same transformation of the lattice vectors $\mathbf{e}_i = \mathbf{S}\mathbf{\tilde{e}}_i$.

For a strong mixed flow, the transformation (2.5) was found by Hunt *et al.* (2010) and used in sample molecular dynamics simulations. Their definition of mixed flows, though, differs from our (2.1*c*) in the choice of the coordinate axes (rotated by 45°). For our definition (2.1*c*) and $\chi > 0$ (characterizing 'strong' flows), the relation between $\dot{\Gamma}$ in (2.5) and $\dot{\gamma}_m$ is $\dot{\Gamma} = \dot{\gamma}_m \sqrt{\chi}$; the matrix **S** and its inverse take the forms

$$\mathbf{S} = \begin{pmatrix} (4\chi)^{-1/4} & -(4\chi)^{-1/4} & 0\\ (\chi/4)^{1/4} & (\chi/4)^{1/4} & 0\\ 0 & 0 & 1 \end{pmatrix}, \quad \mathbf{S}^{-1} = \begin{pmatrix} (\chi/4)^{1/4} & (4\chi)^{-1/4} & 0\\ -(\chi/4)^{1/4} & (4\chi)^{-1/4} & 0\\ 0 & 0 & 1 \end{pmatrix}. \quad (2.6a,b)$$

An arbitrary factor in **S** is chosen to make det(**S**) = 1, which preserves the cell volume in transformations between \tilde{e}_i and e_i . The side of the initial cubic box span on \tilde{e}_1^o , \tilde{e}_2^o and \tilde{e}_3^o (see figure 2*a*) is used as the length scale in our mixed-flow algorithm for $\chi > 0$. It can be derived from (2.3)–(2.6) that $e_1^2 + e_2^2 = (1/2)(\chi^{1/2} + \chi^{-1/2})$ ($\tilde{e}_1^2 + \tilde{e}_2^2$), indicating only a modest additional skewage of the periodic cell in mixed-flow simulations compared with PE simulations (except for very small χ with a stronger effect). For $\chi = 0.25$, the dynamics of the periodic cell V in the interval $0 \leq \dot{\gamma}_m t < 2\varepsilon_p$ is shown in figure 3; at $\dot{\gamma}_m t = 2\varepsilon_p$, the cell V returns to its initial shape. It can be noted that the formal mapping (2.6) between PE and mixed flow is only used to implement the periodic boundary conditions for the latter; of course, mixed-flow rheological simulations are irreducible to PE simulations.

Weak mixed flows. For $\chi < 0$ (when the transformation (2.5) does not exist), implementation of periodic boundaries is much simpler and does not require the computational cell restructuring. As for simple shear, we choose the cube $[0, L)^3$ as

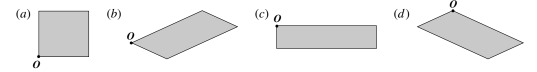


FIGURE 4. The geometry of the computational cell in a weak mixed flow ($\chi = -0.25$). The directions of the x_1 and x_2 coordinate axes are the same as in figure 3; (a) $\omega t = 0$, (b) $\pi/4$, (c) $\pi/2$, (d) $3\pi 4$.

the initial periodic cell and use *L* as the length scale. In contrast to simple shear and PE, though, the lattice vector dynamics $d\mathbf{e}_i/dt = \nabla \mathbf{u}_{\infty}$. \mathbf{e}_i does not lead to indefinite stretching/contraction (which would require cell restructuring), but gives the periodic solution instead:

$$\boldsymbol{e}_1 = (\cos \,\omega t, \, -|\chi|^{1/2} \sin \,\omega t, \, 0), \quad \boldsymbol{e}_2 = (|\chi|^{-1/2} \sin \,\omega t, \, \cos \,\omega t, \, 0), \quad (2.7a,b)$$

with $\omega = |\chi|^{1/2}\dot{\gamma}_m$. Figure 4 illustrates the degree of compactness of the periodic cell for $\chi = -0.25$; the selected shapes at $\omega t = 0$, $\pi/4$, $\pi/2$ and $3\pi/4$ also give the periodic cells at $\omega t = \pi$, $5\pi/4$, $3\pi/2$ and $7\pi/4$ respectively, through central symmetry about the origin **O**. Difficulties with bad aspect ratios would appear for much smaller $|\chi|$ (not considered herein); for the shear-flow limit $\chi = 0$, an alternative standard implementation of periodic boundaries (see above) is used instead.

2.2. Boundary-integral formulation

For all three flows (2.1), the time scale is defined as $(2I_2)^{-1/2}$, the velocity scale is $(2I_2)^{1/2}L$. The non-dimensional system of BI equations for the interfacial fluid velocity u(y) on drop surfaces S_{α} ($\alpha = 1, ..., N$), to be solved iteratively at each time step, is used in the form of paper I:

$$\boldsymbol{u}(\boldsymbol{y}) = \frac{(\lambda - 1)}{(\lambda + 1)} \left[2 \sum_{\beta=1}^{N} \int_{S_{\beta}} \boldsymbol{Q}(\boldsymbol{x}) \cdot \widetilde{\boldsymbol{\tau}}(\boldsymbol{x} - \boldsymbol{y}) \cdot \boldsymbol{n}(\boldsymbol{x}) \, \mathrm{d}S_{\boldsymbol{x}} + \langle \boldsymbol{u} \rangle_{\alpha} + 2 \sum_{\beta=1}^{N} \int_{S_{\beta}} (\boldsymbol{Q} \cdot \boldsymbol{n}) (\boldsymbol{x} - \boldsymbol{x}_{\beta}^{c}) \, \mathrm{d}S_{\boldsymbol{x}} \right] + \boldsymbol{F}(\boldsymbol{y}), \quad \boldsymbol{y} \in S_{\alpha}.$$
(2.8)

Here, the inhomogeneous term is

$$\boldsymbol{F}(\boldsymbol{y}) = \frac{2}{\lambda+1} \left[\boldsymbol{u}_{\infty}(\boldsymbol{y}) + \frac{2a}{Ca} \sum_{\beta=1}^{N} \int_{S_{\beta}} (k(\boldsymbol{x}) - \langle k \rangle_{\beta}) \boldsymbol{n}(\boldsymbol{x}) \cdot \boldsymbol{G}(\boldsymbol{x}-\boldsymbol{y}) \, \mathrm{d}S_{x} \right], \quad (2.9)$$

where $a = a_o/L$ is the non-dimensional non-deformed drop radius, $k(\mathbf{x})$ is the local mean of the principal surface curvatures, $\langle k_\beta \rangle$ is the average of k over S_β , $\mathbf{n}(\mathbf{x})$ is the outward unit normal at $\mathbf{x} \in S_\beta$ and $\mathbf{G}(\mathbf{x})$ is the triply periodic symmetric second-rank Green tensor. This tensor satisfies

$$\nabla^2 \boldsymbol{G}(\boldsymbol{x}) - \nabla \boldsymbol{q}(\boldsymbol{x}) = \nabla \cdot \boldsymbol{\tau}(\boldsymbol{x}) = \sum_{\boldsymbol{m}} \delta(\boldsymbol{x} - \boldsymbol{P}(\boldsymbol{m})) \boldsymbol{I}, \qquad (2.10)$$

where q(x) is the vector of pressures for G(x) and the summation is over all lattice points $P(m) = m^1 e_1 + m^2 e_2 + m^3 e_3$ with integer m^1 , m^2 , m^3 . The Ewald-like form for G(x) for a lattice with arbitrary periods e_1 , e_2 , e_3 follows from Hasimoto (1959) and is given in appendix B of paper I. It is inconvenient, though, that the associated third-rank stress tensor $\tau(x)$ is not periodic (and not symmetric), since it contains a linearly growing part from the pressure q(x). Following Zinchenko & Davis (2000) and paper I, we work instead with the periodic and fully symmetric tensor $\tilde{\tau}(r) = \tau(r) - rI$ in the double-layer integrals (2.8). Additionally, velocity fluctuations $Q(x) = u(x) - \langle u \rangle_{\beta}$ from the surface-averaged values $\langle u \rangle_{\beta}$ are used on S_{β} instead of u(x) to significantly reduce the integrand variations and thereby accelerate convergence in the most difficult multipole part (§ 3) of the algorithm. For the same reason, we work with $k(x) - \langle k \rangle_{\beta}$ instead of k(x) in the single-layer integrals (2.9). It is also convenient that Q(x) (unlike u(x)) is the same for all periodic images of S_{β} .

Once the interfacial velocity is found, the instantaneous volume-averaged stress tensor Σ in the emulsion can be expressed via surface integrals (e.g. Batchelor 1970; Kennedy *et al.* 1994). With $\mu_e(2I_2)^{1/2}$ as the stress scale, our non-dimensional stress for all three flows (2.1) is calculated as

$$\Sigma_{ij} = \text{I.T.} + \nabla_i (u_{\infty})_j + \nabla_j (u_{\infty})_i + \sum_{\alpha=1}^N \int_{S_{\alpha}} \left\{ \frac{2a}{Ca} [k(\mathbf{x}) - \langle k \rangle_{\alpha}] n_i (x - x_{\alpha}^c)_j + (\lambda - 1)(u_i n_j + u_j n_i) \right\} \, \mathrm{d}S, \quad (2.11)$$

where I.T. is an insignificant isotropic term; in deriving (2.11), the unit volume of the periodic cell has been taken into account. The tensor Σ is symmetric upon accurate surface discretization. From (2.11), all the rheological functions (2.2) are calculated.

3. Fast calculation of the boundary integrals

The present multipole-accelerated scheme for calculating the boundary integrals in (2.8) and (2.9) largely follows the logic of paper I, but it is more flexible and more efficient for the present set of problems, where considerable skewage of the periodic cell may occur, and some drops may become extremely elongated. This scheme, briefly outlined below, additionally incorporates some elements from our more recent codes (Zinchenko & Davis 2003, 2008, 2013) for multidrop–multiparticle systems.

3.1. Surface discretization and BI desingularizations

An unstructured mesh of $N_{\triangle} \gg 1$ flat triangles with $N_{\triangle}/2 + 2$ vertices (called nodes) is used to represent each drop surface, including the choices $N_{\triangle} = 2160$, 2880, 3840 and 6000 (in one simulation) used in the present calculations. Initial nearly uniform triangulations were prepared in a standard way from sphere tesselations with mesh refinements. As the drops move and deform, the quality of triangulations is maintained as described in §4 (quite differently from paper I). Surface normals and curvatures are calculated by the best paraboloid-spline method of Zinchenko & Davis (2000). Surface integrations were performed by a standard trapezoidal rule, with reassignment of triangle contributions to vertices for efficiency; the latter procedure follows Rallison (1981) and was used in much of our prior work. The double- and single-layer kernels in (2.8) and (2.9) are singular at all lattice points x - y = P(m) due to free-space contributions:

$$\boldsymbol{G}(\boldsymbol{r}) \simeq \boldsymbol{G}_o(\boldsymbol{r} - \boldsymbol{P}(\boldsymbol{m})), \quad \widetilde{\boldsymbol{\tau}}(\boldsymbol{r}) \simeq \boldsymbol{\tau}_o(\boldsymbol{r} - \boldsymbol{P}(\boldsymbol{m})), \quad (3.1a,b)$$

where

$$\boldsymbol{G}_{o}(\boldsymbol{r}) = -\frac{1}{8\pi} \left(\frac{\boldsymbol{I}}{r} + \frac{\boldsymbol{rr}}{r^{3}} \right), \quad \boldsymbol{\tau}_{o}(\boldsymbol{r}) = \frac{3}{4\pi} \frac{\boldsymbol{rrr}}{r^{5}}.$$
 (3.2*a*,*b*)

This behaviour necessitates efficient desingularizations of the boundary integrals prior to numerical solution. For a mesh node $y \in S_{\alpha}$ on a drop surface different from an integration surface S_{β} , the double-layer contribution from S_{β} is written as (paper I)

$$\int_{S_{\beta}} \boldsymbol{Q}(\boldsymbol{x}) \cdot \widetilde{\boldsymbol{\tau}}(\boldsymbol{x} - \boldsymbol{y}) \cdot \boldsymbol{n}(\boldsymbol{x}) \, \mathrm{d}S_{\boldsymbol{x}} \approx \sum_{\boldsymbol{x}_{j} \in S_{\beta}} \boldsymbol{Q}(\boldsymbol{x}_{j}) \cdot \widetilde{\boldsymbol{\tau}}(\boldsymbol{x} - \boldsymbol{y}) \cdot \boldsymbol{n}(\boldsymbol{x}_{j}) \Delta S_{j} - \sum_{\boldsymbol{m}} \boldsymbol{\varTheta}(\boldsymbol{m}, \boldsymbol{y}) \\ \times \boldsymbol{Q}^{*} \sum_{\boldsymbol{x}_{j} \in S_{\beta}} \boldsymbol{\tau}_{o}(\boldsymbol{x}_{j} + \boldsymbol{P}(\boldsymbol{m}) - \boldsymbol{y}) \cdot \boldsymbol{n}(\boldsymbol{x}_{j}) \Delta S_{j}.$$
(3.3)

Here, the summation is over all mesh nodes $x_i \in S_\beta$ and all integer vectors m, and ΔS_i is the surface area associated with node x_i . The barrier function Θ is designed to act only when y is close to a periodic image $S_{\beta} + P(m)$ of S_{β} . Namely, Θ is close to unity when the distance from y to the nearest node $x^* + P(m)$ on $S_{\beta} + P(m)$ is much less than a threshold h_o , and Θ gradually falls to zero as this distance approaches h_o . For the precise definition of Θ , see paper I; h_o is set to 0.3a in the present calculations. The choice of Q^* is crucial for successful large-strain simulations at high drop volume fractions and significant viscosity contrast λ . The simplest near-singularity subtraction $Q^* = Q(x^*)$ is insufficient to overcome divergence of BI iterations even for moderate $\lambda \neq 1$. The more robust (yet more complex) variational technique of paper I, which we follow herein, requires the subtracted quantity Q^* to minimize the Euclidean norm of the discrete free-space double-layer contribution from $S_{\beta} + P(m)$ to y after subtraction. When $\Theta \approx 1$, the subtracted term in (3.3) effectively alleviates the near-singular behaviour of the double-layer integrand (although it does not eliminate it completely). Of course, this subtracted term, helping calculations at finite triangulations, disappears in the limit $N_{\triangle} \rightarrow \infty$. The optimized way of calculating the subtracted term (Zinchenko & Davis 2006, 2008) had a relatively small cost in the present simulations.

The single-layer contribution in (2.9) from a surface $S_{\beta} \not\supseteq y$ is calculated as

$$\int_{S_{\beta}} f(\mathbf{x}) \mathbf{G}(\mathbf{x} - \mathbf{y}) \cdot \mathbf{n}(\mathbf{x}) \, \mathrm{d}S_{\mathbf{x}} \approx \sum_{\mathbf{x}_{j} \in S_{\beta}} f(\mathbf{x}_{j}) \mathbf{G}(\mathbf{x}_{j} - \mathbf{y}) \cdot \mathbf{n}(\mathbf{x}_{j}) \Delta S_{j} - \sum_{\mathbf{m}} \Theta(\mathbf{m}, \mathbf{y})$$
$$\times f^{*} \sum_{\mathbf{x}_{j} \in S_{\beta}} \mathbf{G}_{o}(\mathbf{x}_{j} + \mathbf{P}(\mathbf{m}) - \mathbf{y}) \cdot \mathbf{n}(\mathbf{x}_{j}) \Delta S_{j}, \qquad (3.4)$$

with the same barrier Θ as in (3.3), and $f(\mathbf{x}) = k(\mathbf{x}) - \langle k_{\beta} \rangle$ for brevity. We found the simplest choice of $f^* = f(\mathbf{x}^*)$ to be sufficient in the present calculations, although a more complex scheme was offered in paper I. The subtracted term in (3.4) completely eliminates near-singular behaviour of the integrand (thus helping calculations at finite triangulations), but, again, disappears in the limit $N_{\Delta} \to \infty$. Due to the barrier Θ , typically, only one periodic image of S_{β} contributes to the subtracted terms in (3.3) and (3.4), and so it was inexpensive to calculate these terms by direct summations.

For self-interactions $(y \in S_{\alpha} = S_{\beta})$, the contributions of S_{β} to the BIs in (2.8) and (2.9) are computed as

$$\int_{S_{\alpha}} \boldsymbol{Q}(\boldsymbol{x}) \widetilde{\boldsymbol{\tau}}(\boldsymbol{x} - \boldsymbol{y}) \cdot \boldsymbol{n}(\boldsymbol{x}) \, \mathrm{d}S_{x}$$

$$\approx \frac{1}{2} \boldsymbol{Q}(\boldsymbol{y}) + \sum_{\boldsymbol{x}_{j} \in S_{\alpha}} \boldsymbol{Q}(\boldsymbol{x}_{j}) \cdot [\widetilde{\boldsymbol{\tau}}(\boldsymbol{x}_{j} - \boldsymbol{y}) - \boldsymbol{\tau}_{o}(\boldsymbol{x}_{j} - \boldsymbol{y})] \cdot \boldsymbol{n}(\boldsymbol{x}_{j}) \Delta S_{j}$$

$$+ \sum_{\boldsymbol{x}_{j} \in S_{\alpha}} [\boldsymbol{Q}(\boldsymbol{x}_{j}) - \boldsymbol{Q}(\boldsymbol{y})] \cdot \boldsymbol{\tau}_{o}(\boldsymbol{x}_{j} - \boldsymbol{y}) \cdot \boldsymbol{n}(\boldsymbol{x}_{j}) \Delta S_{j} \qquad (3.5)$$

and

$$\int_{S_{\alpha}} f(\mathbf{x}) \mathbf{n}(\mathbf{x}) \cdot \mathbf{G}(\mathbf{x} - \mathbf{y}) \, \mathrm{d}S_{\mathbf{x}} \approx \sum_{\mathbf{x}_j \in S_{\alpha}} f(\mathbf{x}_j) \mathbf{n}(\mathbf{x}_j) \cdot [\mathbf{G}(\mathbf{x}_j - \mathbf{y}) - \mathbf{G}_o(\mathbf{x}_j - \mathbf{y})] \Delta S_j + \sum_{\mathbf{x}_j \in S_{\alpha}} [f(\mathbf{x}_j) - f(\mathbf{y})] \mathbf{n}(\mathbf{x}_j) \cdot \mathbf{G}_o(\mathbf{x}_j - \mathbf{y}) \Delta S_j. \quad (3.6)$$

For the second lines in (3.5) and (3.6) (with $x_j = y$ excluded), direct summations are always used. Unlike in (3.3), complete desingularization is achieved in (3.5).

3.2. Multipole-accelerated scheme

For the first lines in (3.3) and (3.4), it would be, of course, prohibitively expensive to use direct summations in dynamical simulations, with a cost of $O(N^2 N_{\Delta}^2)$ per time step and a large coefficient, due to iterations and the complex nature of $\tilde{\tau}$ and G. A highly efficient multipole-accelerated scheme is used instead. Each drop, when elongated, is divided into a few (not more than three) compact blocks; see paper I for the procedure and figure 2 therein. A compact drop coincides with its only block. As a result, all mesh nodes on all drops are partitioned into compact blocks $\mathcal{B}_1, \ldots, \mathcal{B}_{N_B}$ ($N_B \ge N$). A minimal spherical shell \mathcal{D}_{γ} with a centre \mathbf{x}_{γ}^o and radius d_{γ}^o is constructed around each block \mathcal{B}_{γ} with sufficient accuracy.

The next step is to partition the periodic kernels G(r) and $\tilde{\tau}(r)$ into the 'nearfield' and 'far-field' parts. In paper I, only the $G_o(r)$ and $\tau_o(r)$ parts of G(r) and $\tilde{\tau}(r)$ were qualified as near-field. Such partition causes difficulties and/or performance degradation when N is not too large (~100), periodic cells are significantly more stretched than in shear flow (see figures 2–4) and some drops are strongly deformed. A more flexible and robust scheme herein is to assign free-space contributions from lattice points to near-field, if their distance from the origin is within a given threshold R_{sub} :

$$\mathbf{G}(\mathbf{r}) = \sum_{|\mathbf{P}(\mathbf{m})| \leq R_{sub}} \mathbf{G}_o(\mathbf{r} + \mathbf{P}(\mathbf{m})) + \mathbf{G}_1(\mathbf{r}), \quad \widetilde{\boldsymbol{\tau}}(\mathbf{r}) = \sum_{|\mathbf{P}(\mathbf{m})| \leq R_{sub}} \boldsymbol{\tau}_o(\mathbf{r} + \mathbf{P}(\mathbf{m})) + \boldsymbol{\tau}_1(\mathbf{r}).$$
(3.7*a*,*b*)

The number of free-space contributions to be subtracted from $(\mathbf{G}, \tilde{\boldsymbol{\tau}})$ to form the farfield (\mathbf{G}_1, τ_1) depends, therefore, on both R_{sub} and the instantaneous lattice geometry. For $R_{sub} = 1.4$, which we found to be in the optimal range in the present simulations with N = 100-400 drops, there are 7–9, 7–13 and 9–13 subtractions for simple shear, PE and mixed-flow ($\chi = 0.25$) lattices respectively. The partition (3.7) with $R_{sub} = 1.4$ moves the singularities of $G_1(r)$ and $\tau_1(r)$ farther away from the origin, compared with the $R_{sub} = 0$ choice in paper I.

The free-space contributions of each block \mathscr{B}_{γ} ($\gamma = 1, ..., N_B$) to the summations in the first lines of (3.3) and (3.4) (i.e. with $G_o(x_j - y)$ and $\tau_o(x_j - y)$ in place of $G(x_j - y)$ and $\tilde{\tau}(x_j - y)$) are expanded in Lamb's singular series about the block centre x_{γ}^o . These expansions are generated to a sufficient order by the alternative algorithm of Zinchenko & Davis (2008, § 4.2 therein), which is ≈ 2.5 times faster for double-layer expansions than the original algorithm of Zinchenko & Davis (2000) used in paper I for this purpose.

Now, to calculate the sums in the first lines of (3.3) and (3.4) for $y \in \mathscr{B}_{\delta} \subset S_{\alpha}$, we consider, for each block $\mathscr{B}_{\gamma} \not\subset S_{\alpha}$, a periodically shifted block $\mathscr{B}_{\gamma}^{min} = \mathscr{B}_{\gamma} + P(\boldsymbol{m}_{\delta\gamma})$, which minimizes the centre-to-centre distance $|\boldsymbol{x}_{\gamma}^{o,min} - \boldsymbol{x}_{\delta}^{o}|$ (with $\boldsymbol{x}_{\gamma}^{o,min} = \boldsymbol{x}_{\gamma}^{o} + \boldsymbol{x}_{\delta}^{o}$) $P(m_{\delta \gamma})$). In what follows, the index min will supplement quantities related to $\mathscr{B}_{\gamma}^{min}$. According to (3.7) and periodicity of the kernels **G** and $\tilde{\tau}$, the contribution of \mathscr{B}_{ν} to either (3.3) or (3.4) can be written as the sum of free-space contributions from $\mathscr{B}_{\mathcal{Y}}^{\min} + \mathbf{P}(\mathbf{m})$ (with $|\mathbf{P}(\mathbf{m})| \leq R_{sub}$) plus the far-field contribution from $\mathscr{B}_{\mathcal{Y}}^{\min}$ using τ_1 and G_1 . The latter is calculated by a special form of Taylor double series for $\tau_1(x - y)$ and $G_1(x - y)$ in powers of $x - x_{\nu}^{o,min}$ and $y - x_{\delta}^{o}$ (Zinchenko & Davis 2000), which only requires knowledge of $G_1(r)$, the associated vector of pressures $q_1(r)$ and their high-order derivatives at $r = R_{\gamma\delta} = x_{\delta}^o - x_{\gamma}^{o,min}$ (appendix B of paper I). The optimized form of far-field expansions (appendix A of Zinchenko & Davis 2003) valid for any lattice geometry (with a unit cell volume) is used herein in favour of the algorithm from paper I. The necessary functions $G_1(r), q_1(r)$ and their derivatives are pretabulated (prior to iterations of (2.8)) to a sufficient order $n_{tab}(n_1, n_2, n_3)$ on a skewed uniform mesh $x = hn_1e_1 + hn_2e_2 + hn_3e_3$, with $h = 0.5/N_T$ and the integers n_i in the range $|n_1| \leq N_T$, $0 \leq n_2$, $n_3 \leq N_T$. Extension to negative n_2 or n_3 simply follows from the symmetry properties of $G_1(r)$ and $q_1(r)$. Since the singularities of $G_1(r)$ and $q_1(r)$ are far away from the origin, these functions are very smooth in the tabulation range $V_{tab} = \{\xi^i e_i, |\xi^i| \leq 1/2\}$, so only a small value of N_T (typically 7) and a small order n_{tab} of derivatives suffice. Still, due to the lattice time dependence, the tabulation is required at each time step. The most efficient Ewald-like forms for G_1 and q_1 follow from Hasimoto (1959), like in paper I. These forms, together with fast high-order differentiation techniques (appendix C of paper I), allow dynamical tabulation of G_1, q_1 and their high-order derivatives at a negligible cost. For a skewed lattice, the minimal vector $\mathbf{R}_{\gamma\delta}$ is not always in the tabulation range V_{tab} . In general, the periodic shift $P(m^*)$ is found to map $R_{\gamma\delta}$ on V_{tab} . The necessary values of G(r), $q_1(r)$ and their high-order derivatives at $r = R_{\gamma\delta}$ are easily linked to those at $r = R_{\gamma\delta}^*$ $R_{v\delta} + P(m) \in V_{tab}$, by adding/subtracting a few free-space contributions in an efficient manner (appendix D of paper I). In turn, the values of G_1, q_1 and their derivatives at $r = R_{\gamma\delta}^*$ are obtained from the nearest table node by Taylor expansions (typically of second or third order). Again, due to the smoothness of $\tau_1(r)$ and $G_1(r)$ near $r = R_{\gamma\delta}$, the far-field double series in powers of $x - x_{\gamma}^{min}$ and $y - x_{\delta}^{o}$ for the (τ_1, G_1) contribution to (3.3) and (3.4) are rapidly convergent, and just a few terms suffice.

If the shell $\mathscr{D}_{\gamma}^{\min} + P(m)$ around $\mathscr{B}_{\gamma}^{\min} + P(m)$ and the shell \mathscr{D}_{δ} have 'enough clearance' (see below), the free-space contributions from $\mathscr{B}_{\gamma}^{\min} + P(m)$ to (3.3) and (3.4) are evaluated at $y \in \mathscr{B}_{\delta}$ by first re-expanding Lamb's singular series for $\mathscr{B}_{\gamma}^{\min} + P(m)$ as Lamb's regular series near x_{δ}^{o} . All such re-expansions are greatly accelerated (Zinchenko 1994, Zinchenko & Davis 2000) by rotational transformations of spherical harmonics with Wigner functions, and the cumulative Lamb's regular series from all

blocks 'sufficiently separated' from \mathscr{B}_{δ} are then efficiently evaluated at all mesh nodes $y \in \mathscr{B}_{\delta}$.

If the block $\mathscr{B}_{\gamma}^{min} + P(m)$ is not 'sufficiently separated' from \mathscr{B}_{δ} , but node $y \in \mathscr{B}_{\delta}$ is 'well outside' the shell $\mathscr{D}_{\gamma}^{min} + P(m)$ (as defined below), Lamb's singular series for $\mathscr{B}_{\gamma}^{min} + P(m)$ are used directly to compute the contributions of $\mathscr{B}_{\gamma}^{min} + P(m)$ to (3.3) and (3.4). Only in rare cases, when y is too close to $\mathscr{D}_{\gamma}^{min} + P(m)$ from outside, or is inside $\mathscr{D}_{\gamma}^{min} + P(m)$, should we use direct summations for these contributions.

A separate scheme (also different from the one in paper I) is used for selfinteractions, i.e. to calculate the sums in the first lines of (3.5) and (3.6). It was observed in our PE and strong mixed-flow simulations in near-critical conditions that one (or a few) drop(s) can acquire extreme elongation (even comparable with the size of the periodic cell), still without necking, while all the other drops remain compact. To handle such cases in a simple and efficient manner, we replace, for self-interactions only, the uniform partition (3.7) by a variable partition with the parameter R_{sub}^{α} (instead of R_{sub}) now dependent on the drop elongation. Drop partitioning into blocks is not used in the self-interactions (3.5) and (3.6); we work with entire drops instead. Let ζ_o be the minimum distance from the origin to the lattice points $P(m), m \neq 0$, and d_{α} be the radius of the minimal spherical shell around S_{α} centred at \mathbf{x}_{α}^{c} . The convergence rate of the far-field expansion (with τ_1 and G_1 in place of $\tilde{\tau}$ and G in (3.5) and (3.6)) is conservatively measured by the progression exponent $2d_{\alpha}/\max(R_{sub}^{\alpha}, \zeta_{o})$. We require this quantity to stay within $(1 + 2a/\zeta_o)/2$, which determines R_{sub}^{α} . Upon the partition of **G** and $\tilde{\tau}$, the remaining free-space contributions from surfaces $S_{\alpha} + P(m)$ (with $0 < \mathbf{P}(\mathbf{m}) \leq R_{sub}^{\alpha}$) to (3.5) and (3.6) are handled by direct summations. For an extremely deformed drop, a fairly large number of free-space contributions need to be handled in this manner; still, the procedure remains fast compared with the rest of the code, since it is used for self-interactions only.

In BI simulations of emulsion flow through a packed bed (Zinchenko & Davis 2008, 2013) with ultrahigh surface resolution ($N_{\Delta} > 10\,000$) on solid boundaries, it was found to be useful to additionally divide each surface into a large number of 'patches' for multipole acceleration. This second level of mesh-node decomposition would not be beneficial in the present simulations with typically $N_{\Delta} \leq 3840$.

Economical truncation bounds. As in paper I, the vital feature of the present multipole-accelerated scheme is the 'economical truncation' of multipole expansions/ re-expansions to greatly boost the performance. It is taken into account, in nearly optimal fashion, that the rate of convergence of Lamb's free-space singular series for block $\mathscr{B}_{\gamma}^{\min} + \mathbf{P}(\mathbf{m})$ is a strong function of the clearance from the surrounding minimal spherical shell $\mathscr{D}_{\gamma}^{\min} + \mathbf{P}(\mathbf{m})$. Likewise, the rate of convergence of singular-to-regular free-space re-expansion between $\mathscr{B}_{\gamma}^{min} + P(m)$ and \mathscr{B}_{δ} strongly depends on the clearance between the surrounding shells $\mathscr{D}_{\gamma}^{\min} + \mathbf{P}(\mathbf{m})$ and \mathscr{D}_{δ} . In constructing the economical truncation bounds, we follow the technique of paper I, based, in part, on plausible estimations for the rate of decay of multipole coefficients. The only difference is that, according to the partition (3.7), there are now more free-space interactions qualified as 'near-field'. Accordingly, for blocks $\mathcal{B}_{\gamma}^{min}$ and \mathcal{B}_{δ} separated (centre-to-centre) by $\mathbf{R}_{\gamma\delta}$, the appropriate distance to affect the rate of convergence of far-field series contributions $\mathscr{B}_{\gamma}^{\min} \to \mathscr{B}_{\delta}$ is now $\zeta_{\alpha\beta} = \min |\mathbf{R}_{\gamma\delta} + \mathbf{P}(\mathbf{m})|$ (over $|P(m)| > R_{sub}$). Our truncation scheme, inevitably semi-intuitive, contains a single heuristic 'precision parameter' ε . Most importantly, as $\varepsilon \to 0$, all multipoles in near-field expansions/re-expansions and all terms in the far-field Taylor series are eventually included. However, for optimal performance, a threshold order k_o is set to

limit the use of multipoles in near-field operations. For a given ε , blocks $\mathscr{B}_{\gamma}^{min} + P(m)$ and \mathscr{B}_{δ} are qualified as 'sufficiently separated' if the multipole truncation order required for the block-to-block re-expansion is within k_o . Likewise, the node y is considered to be 'well outside' the shell $\mathscr{D}_{\gamma}^{min} + P(m)$ if the necessary truncation order for Lamb's singular series for $\mathscr{B}_{\gamma}^{min} + P(m) \rightarrow y$ free-space contribution does not exceed k_o . The value of $k_o = 25$ was about optimal in the present calculations. In the limit $\varepsilon \rightarrow 0$, though, the same non-multipole solution is reached regardless of k_o and other artificial parameters R_{sub} , R_{sub}^{α} . The correctness and high efficiency of our multipole-accelerated code are demonstrated in § 5.

4. Miscellaneous details of the algorithm

The generalized minimal residual method GMRES(k), with k = 5, is used to iterate the BI equations (2.8) at each time step; additional remarks in Zinchenko & Davis (2013) may be useful in this respect. The initial approximation to u(y) was constructed from the two preceding time steps to reduce the number of iterations. A version of 'passive mesh stabilization' (a family of techniques originated in Zinchenko, Rother & Davis 1997) is used to maintain the mesh quality in dynamical simulations. Namely, the mesh-node velocities $V_i = dx_i/dt$ to be used in the drop shape update are required to minimize a 'kinetic energy' function (separately on each surface), under the constraints on the normal velocity $V_i \cdot n(x_i) = u(x_i) \cdot n(x_i)$ from the BI solution. We use the newest (and most successful) form of this function ((3.1)-(3.4) of Zinchenko & Davis 2013), which both resists mesh triangle collapse and is moderately adaptive to surface curvature. In the runs with modest drop deformation, this technique alone, which maintains fixed mesh topology, has soft stability limitations on the time step and avoids any surface interpolations, was sufficient to maintain mesh quality for an indefinite time. Only for larger deformations did we have to use, still very rarely, two additional tools for the mesh control - (i) active mesh restructuring through minimization of the 'potential energy' function E and (ii) node reconnection (edge swapping). Our curvature-adaptive form of E ((3.5) of Zinchenko & Davis 2013) is more complex than the simplest physically motivated 'spring-like' form of Cristini, Bławzdziewicz & Loewenberg (2001); in particular, our form better helps us to prevent mesh-node crowding and mesh triangle collapse. Every 50-100 time steps, for each drop that has mesh triangles with compactness $C_{\Delta} < 0.09$, E-minimization iterations (Zinchenko & Davis 2013) are combined with recursive node reconnections. The resulting mesh changes are only accepted (and completed), though, if mesh improvement is observed after the first 200 iterations. The criterion for mesh improvement is the same as in Zinchenko & Davis (2013); the criterion for edge swapping mostly follows Cristini et al. (2001). It is the power of 'passive mesh stabilization', with the new kinetic energy function F, that makes the actual use of two other tools (i) and (ii) very infrequent, thus reducing the number of BI iterations - the main contributor to the computational cost. Only for an extremely deformed drop did we have to use (i) and (ii) more often (every 10 time steps). It should be noted also that our mesh strategy avoids more complex topological changes (node addition/subtraction) altogether.

Control of surface overlapping. As in paper I, van der Waals attractive forces between the drops are neglected compared with the hydrodynamical forces (which is justified, except for very small drops of colloidal size). In principle, under these conditions, lubrication layers between neighbouring drops would prevent surface contacts, if the problem could be solved exactly. In practice, though, elimination of all

surface overlaps at high emulsion concentrations would require exorbitant resolution and make the present simulations impossible. In paper I, small drop-drop overlaps were ignored altogether. This approach was mostly justified in the calculations of paper I, but is not in the present work with systematic runs in a much wider (1.5-2)orders of magnitude) time range; accumulation and growth of overlaps, however slow, would eventually destroy the solution. The control of surface overlapping was added to the present code as a major improvement over paper I; the principle is the same as in emulsion flow simulations through a packed bed (Zinchenko & Davis 2013). Let $\mathscr{R}(y)$ be the ray emanating from the mesh node $y \in S_{\alpha}$ in the direction of the outward normal to S_{α} , and $\delta(y)$ be the distance from y along $\mathscr{R}(y)$ to the nearest intersection with other drop surfaces (including periodic images). On each time step, all mesh nodes y entering the zone $\delta(y) < \delta_{min}$ are pushed back along -n(y) to make $\delta(\mathbf{y}) = \delta_{min}$; the default threshold δ_{min} is set to 0.001*a*. This *ad hoc* procedure is justified, because lubrication is 'almost resolved' due to high triangulations, so the above geometrical correction is required only for a very small portion (within 0.5-1%) of the total number of mesh nodes, and was not observed to compromise the overall accuracy (§ 6). To calculate $\delta(y)$, the nearest intersection of $\Re(y)$ with flat mesh triangles is first found, which selects a triangle and the corresponding surface. Finally, the intersection of $\mathscr{R}(y)$ with the best paraboloid locally fitting the selected surface near the selected triangle is used to determine $\delta(y)$. A plausible reason why it would be extremely difficult to eliminate all the remaining rare overlaps through the increase in the surface resolution is in the highly non-uniform thin film profile between nearly touching drops. It is well known from semianalytical studies of axisymmetrical thin film drainage between two drops under the action of a driving force that the surface clearance in the narrow transition zone from the film to the meniscus is much smaller than in the rest of the film.

The drop configuration is advanced by the simplest Euler scheme, with a variable non-dimensional time step

$$\Delta t = K_{\Delta t} C a \min(\Delta x, 0.7 \Delta x^*) / a, \qquad (4.1)$$

where $K_{\Delta t}$ is a numerical constant. The presence of Δx (the minimum spacing between directly connected mesh nodes) in (4.1) is in line with the usual Courant stability condition. The second term in (4.1) accounts for close drop interactions. Namely, Δx^* is the minimum distance between nodes $x_i \in S_{\alpha}$ and $x_i \in S_{\beta}$ on different surfaces (or their periodic images), but excluding pairs for which x_i is the node on S_{β} closest to x_i , or vice versa. This exclusion is allowed by near-singularity subtraction in the boundary integrals and prevents Δt from becoming prohibitively small. The criterion (4.1) was first used by Zinchenko & Davis (2005) in the two-drop problem, but with different factors $K_{\Delta t}$. In the present calculations for highly concentrated emulsions, $K_{\Delta t} \approx 2$ was universally applicable when $\lambda = 1$ and 3; for more extreme viscosity ratios $\lambda = 0.25$ and 10, smaller $K_{\Delta t}$ had to be used (≈ 1 and 0.65–1 respectively). With the rule (4.1) and c = 0.55 drop volume fraction, each time step required roughly 5–8 BI iterations for $\lambda = 0.25$ and 3, and $\approx 7-11$ iterations when $\lambda = 10$. The effect of time integration errors on the steady-state rheological functions (2.2) was generally well within statistical discrepancies, and so more expensive higher-order time-marching schemes were unjustified. As the only exception, time integration errors were observed to exceed statistical errors for a few weak mixed-flow runs $(\chi = -0.25)$ at the highest Ca; these were simply recalculated by the Euler scheme with much smaller $K_{\Delta t}$.

In some long-time simulations, a development of abnormal high curvature in a single node on just one drop was observed, with local corrugation of the otherwise

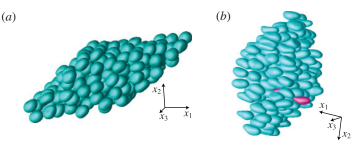


FIGURE 5. (Colour online) Snapshots of the dynamical simulations with c = 0.55, $\lambda = 3$, N = 200, $N_{\triangle} = 2160$ for (a) strong mixed flow ($\chi = 0.25$, Ca = 0.0375) and (b) PE (Ca = 0.088). Only N independent drops with centres in the periodic box V are shown. In (b), one drop develops an extreme elongation, while all other drops remain relatively compact; (a) t = 44.33, (b) t = 34.13.

smooth and compact drop shape. These events were extremely rare (not more often than every $O(10^4)$ time steps, even less frequently than in paper I due to improvements herein), but they still required treatment to continue the simulation to large times for averaging. Swelling techniques (Zinchenko & Davis 2008, 2013) provide a suitable and very accurate remedy. A spherical drop of maximum possible volume is first inserted inside the problematic ('parent') drop. The new drop is then expanded with deformation, until it very closely fits the parent drop (except in the very vicinity of the problematic node, of no significance). The new smooth shape replaces the parent drop to continue the simulation. This approach is superior to the ellipsoid fitting used in paper I for the same purpose. The swelling technique is also a flexible tool to change the surface resolution on demand for any chosen drop, since the new drop resolution (unrelated to that for the parent drop) can be set arbitrarily. Such remeshing was used in a few runs of §§ 5 and 6 to probe the evolution to drop breakup in emulsion flow.

The initial drop configuration for each run was a random arrangement of non-overlapping spheres (and their periodic images) well mixed by the Metropolis algorithm (with millions of Monte Carlo random displacements).

5. Algorithm testing and efficiency

Comprehensive validations of the multipole-accelerated BI code for shear-flow simulations were performed in paper I. However, since the algorithm has undergone many changes/improvements to be used in new applications herein, we have retested the new code and re-evaluated its efficiency. Figure 5 presents snapshots of our dynamical simulations with c = 0.55, $\lambda = 3$, N = 200, $N_{\triangle} = 2160$ and a default 'precision parameter' $\varepsilon = 10^{-3}a$ for (a) strong mixed flow ($\chi = 0.25$, Ca = 0.0375) and (b) PE (Ca = 0.088). For the non-dimensional times shown, the systems are at statistical steady state, as far as the rheology (2.2) is of interest (see below). As an alternative to our (fairly complex) multipole-accelerated code, much simpler (but very inefficient) direct summations were used to compute the first lines in (3.3)-(3.6)for the instantaneous configurations from figure 5. To this end, the smooth functions $\boldsymbol{G}(\boldsymbol{\xi}) - \boldsymbol{G}_{o}(\boldsymbol{\xi})$ and $\widetilde{\boldsymbol{\tau}}(\boldsymbol{\xi}) - \boldsymbol{\tau}_{o}(\boldsymbol{\xi})$ are tabulated on a 215³ mesh in $V_{tab} = \{\xi^{i}\boldsymbol{e}_{i}, |\xi^{i}| \leq 1/2\}$ to second derivatives. Mapping of $x_j - y$ on V_{tab} and using periodicity of **G** and $\tilde{\tau}$ allows us then to calculate $G(x_i - y)$ and $\tilde{\tau}(x_i - y)$ for every pair of nodes x_i , y using quadratic Taylor expansions from the table nodes and adding the free-space parts G_o and τ_{o} . The fine table size is chosen to exclude any appreciable numerical errors in this simple non-multipole calculation. The results obtained in this way are viewed as exact (for given discretizations (3.3)–(3.6)) and are supplemented with index *ex*. For the inhomogeneous term (2.9), only $\Delta F(y) = F(y) - 2u_{\infty}(y)/(\lambda + 1)$ is interesting to compare between the two methods, in order to exclude the macroscopic contribution from u_{∞} . Likewise, for the velocity field, the excess $\Delta u(y) = u(y) - u_{\infty}(y)$ must be compared between the two methods. For the single-layer test of the integrals (2.9), three measures are used to quantify the deviation between ΔF (for the multipole-accelerated scheme) and ΔF_{ex} (cf. paper I):

$$\delta_1(\boldsymbol{F}) = \frac{1}{\langle \Delta \boldsymbol{F}_{ex}^2 \rangle^{1/2}} \max_{\boldsymbol{x}_i} |\Delta \boldsymbol{F}(\boldsymbol{x}_i) - \Delta \boldsymbol{F}_{ex}(\boldsymbol{x}_i)|, \qquad (5.1a)$$

$$\delta_2(\boldsymbol{F}) = \frac{1}{N} \sum_{\alpha=1}^{N} \frac{1}{\langle \Delta \boldsymbol{F}_{ex}^2 \rangle_{\alpha}^{1/2}} \max_{\boldsymbol{x}_i \in S_{\alpha}} |\Delta \boldsymbol{F}(\boldsymbol{x}_i) - \Delta \boldsymbol{F}_{ex}(\boldsymbol{x}_i)|, \qquad (5.1b)$$

$$\delta_3(\boldsymbol{F}) = \langle (\Delta \boldsymbol{F} - \Delta \boldsymbol{F}_{ex})^2 \rangle^{1/2} / \langle \Delta \boldsymbol{F}_{ex}^2 \rangle^{1/2}.$$
 (5.1c)

Here, $\langle \cdots \rangle_{\alpha}$ and $\langle \cdots \rangle$ denote averaging over S_{α} and all surfaces respectively; in the conservative measure (5.1*a*), the maximum is taken over all nodes on all drops. In the double-layer test for the integrals (2.8), both the multipole-accelerated and the non-multipole solutions use the same F and the same initial approximation to u (both taken from our dynamical simulation with $\varepsilon = 10^{-3}a$), and one iteration of (2.8) is performed by both methods to compare Δu for the multipole-accelerated scheme with Δu_{ex} . The deviation between the two is quantified in the same way as in (5.1), with u in place of F everywhere. The mixed-flow and PE runs were also repeated with N = 100 drops and interrupted at t = 37.71 and 51.51 respectively, for a similar accuracy analysis.

The results are summarized in tables 1 and 2. In addition to $\delta_i(F)$ and $\delta_i(u)$, the CPU times (in seconds on a single 3.6 GHz PC core) to compute F (with all overheads including curvature/normal calculations, drop partitioning into blocks, singularity subtractions, etc.) and for one BI iteration of (2.8) are also given. In the PE test for N = 200, it was slightly better to limit the partition of each drop by two blocks (with 395 total), versus three blocks for N = 100 (and 243 total). For much larger systems, it would be optimal to disable partitioning altogether. In the mixed-flow test with smaller drop deformation, partitioning was almost inactive even at N = 100. In tables 1 and 2, all deviations $\delta_i(\mathbf{F})$ and $\delta_i(\mathbf{u})$ strongly correlate with the intuitive parameter ε and tend to zero as $\varepsilon \to 0$, which demonstrates convergence of our multipole-accelerated code to the direct summations code. The computational cost is weakly sensitive to ε . The code is particularly fast for small-to-moderate drop deformations (table 1 with the average drop length $\langle L_d \rangle = (2.35 - 2.37)a$), and degrades only modestly for larger deformations (table 2 with $\langle L_d \rangle / a = 3.01 - 3.19$). For comparison, the standard summation algorithm required 26-27 min for F, and \approx 35 min for one BI iteration, even at N = 100. Compared with the shear-flow algorithm from paper I, the present code is more flexible but also more efficient in less favourable conditions (more skewed periodic cells and larger drop deformations). For example, in the PE test with N = 100 ($\langle L_d \rangle = 3.19a$) and $\varepsilon = 10^{-3}a$, our CPU time of 9.5 s per BI iteration (table 2) is 140 times faster than by direct summations, even when the simplest linear interpolation is used in the latter for $\tilde{\tau} - \tau_{a}$ (and not including substantial tabulation time for the direct code). In the shear-flow test of paper I with Ca = 0.1, c = 0.55, $\lambda = 3$, N = 100, $N_{\triangle} = 1500$ and smaller deformations $(\langle L_d \rangle = 2.80a)$, the corresponding gain was 106-fold. The CPU time scaling in tables 1 and 2 is roughly linear in the number of drops N.

| | $N = 100 \ (t = 47.14)$ | | | $N = 200 \ (t = 44.33)$ | | | |
|----------------------------|--------------------------|----------------------|----------------------|--------------------------|----------------------|----------------------|--|
| | $\varepsilon = 10^{-2}a$ | $10^{-3}a$ | $10^{-4}a$ | $\varepsilon = 10^{-2}a$ | $10^{-3}a$ | $10^{-4}a$ | |
| $\delta_1(\boldsymbol{F})$ | $7.0 	imes 10^{-2}$ | $9.6 	imes 10^{-3}$ | 1.2×10^{-3} | $5.7 	imes 10^{-2}$ | $1.0 	imes 10^{-2}$ | $1.3 	imes 10^{-3}$ | |
| $\delta_2(F)$ | 2.3×10^{-2} | 3.4×10^{-3} | 4.5×10^{-4} | 2.2×10^{-2} | 3.2×10^{-3} | 4.2×10^{-4} | |
| $\delta_3(F)$ | 6.2×10^{-3} | 9.6×10^{-4} | 1.4×10^{-4} | 5.6×10^{-3} | $8.8 	imes 10^{-4}$ | $1.2 	imes 10^{-4}$ | |
| CPU | 7.0 | 8.1 | 9.6 | 15.3 | 17.9 | 21.3 | |
| Time (\mathbf{F}) | | | | | | | |
| $\delta_1(\boldsymbol{u})$ | 4.6×10^{-3} | $6.7 	imes 10^{-4}$ | $8.5 	imes 10^{-5}$ | 7.4×10^{-3} | 9.4×10^{-4} | 1.3×10^{-4} | |
| $\delta_2(\boldsymbol{u})$ | 1.8×10^{-3} | 2.4×10^{-4} | 3.6×10^{-5} | 1.5×10^{-3} | $2.6 	imes 10^{-4}$ | $2.6 	imes 10^{-5}$ | |
| $\delta_3(\boldsymbol{u})$ | 5.7×10^{-4} | $8.0 	imes 10^{-5}$ | 1.1×10^{-5} | 4.4×10^{-4} | 8.2×10^{-5} | 8.2×10^{-6} | |
| CPU | 4.7 | 5.7 | 7.0 | 10.9 | 13.3 | 16.6 | |
| Time (1 it.) | | | | | | | |

TABLE 1. Convergence of the multipole-accelerated solution to the standard $O(N^2 N_{\Delta}^2)$ solution as $\varepsilon \to 0$ in the mixed-flow tests ($\chi = 0.25$, Ca = 0.0375, c = 0.55, $\lambda = 3$, $N_{\Delta} = 2160$).

| | $N = 100 \ (t = 103.02)$ | | | $N = 200 \ (t = 34.13)$ | | |
|----------------------------|--------------------------|----------------------|----------------------|--------------------------|----------------------|----------------------|
| | $\varepsilon = 10^{-2}a$ | $10^{-3}a$ | $10^{-4}a$ | $\varepsilon = 10^{-2}a$ | $10^{-3}a$ | $10^{-4}a$ |
| $\delta_1(\boldsymbol{F})$ | $1.3 	imes 10^{-2}$ | 2.2×10^{-3} | $2.8 	imes 10^{-4}$ | $2.4 	imes 10^{-2}$ | $3.3 	imes 10^{-3}$ | $4.9 	imes 10^{-4}$ |
| $\delta_2(F)$ | 5.3×10^{-3} | $7.3 	imes 10^{-4}$ | $9.6 	imes 10^{-5}$ | 6.7×10^{-3} | $9.2 	imes 10^{-4}$ | $1.1 	imes 10^{-4}$ |
| $\delta_3(F)$ | 1.3×10^{-3} | $1.9 	imes 10^{-4}$ | 2.6×10^{-5} | 1.9×10^{-3} | $2.7 	imes 10^{-4}$ | 3.4×10^{-5} |
| CPU | 10.8 | 13.1 | 16.0 | 24.6 | 30.1 | 36.7 |
| Time (\mathbf{F}) | | | | | | |
| $\delta_1(\boldsymbol{u})$ | 9.6×10^{-3} | 1.1×10^{-3} | 1.5×10^{-4} | 6.9×10^{-3} | 1.0×10^{-3} | 1.1×10^{-4} |
| $\delta_2(\boldsymbol{u})$ | 2.6×10^{-3} | 3.6×10^{-4} | $4.8 	imes 10^{-5}$ | 2.5×10^{-3} | 3.7×10^{-4} | $4.6 	imes 10^{-5}$ |
| $\delta_3(\boldsymbol{u})$ | 6.1×10^{-4} | 8.7×10^{-5} | 1.1×10^{-5} | $5.8 	imes 10^{-4}$ | 9.2×10^{-5} | 1.1×10^{-5} |
| CPU | 7.6 | 9.5 | 11.9 | 18.3 | 23.1 | 29.0 |
| Time (1 it.) | | | | | | |

TABLE 2. Convergence of the multipole-accelerated solution to the standard $O(N^2 N_{\Delta}^2)$ solution as $\varepsilon \to 0$ in the planar-extension tests (Ca = 0.088, c = 0.55, $\lambda = 3$, $N_{\Delta} = 2160$).

The default value $10^{-3}a$ of the intuitive parameter ε in our dynamical simulations provides good accuracy for instantaneous configurations (tables 1 and 2), but is also tight enough to not affect time averaging of the rheological functions. In figure 6(*a*), the thin lines show the trajectories of the non-dimensional stresses (2.2*c*) v_{ij} for the N = 100 mixed-flow simulation from table 1 continued to larger times $t \sim 250$ (outside the range in the figure). Instead of $v_{12}(=\mu^*)$, we plot $\mu^* - 1$ to exclude the continuous-phase contribution; very small values of $v_{33} = -(v_{11} + v_{22})$ are not included. For comparison, the run was repeated with $\varepsilon = 10^{-4}a$ (thick blue lines) to $t \sim 260$. The two simulations are indistinguishable for small times $t \leq 8$, but then quickly randomize. However, the long-time averages

$$\langle \mu^* - 1 \rangle = 3.544 \pm 0.01, \quad \langle \nu_{11} \rangle = 0.628 \pm 0.01, \quad \langle \nu_{22} \rangle = -0.592 \pm 0.01 \quad (\varepsilon = 10^{-3}a),$$
(5.2*a*-*c*)
 \langle \mu^* - 1 \rangle = 3.537 \pm 0.01, \quad \langle \nu_{11} \rangle = 0.612 \pm 0.01, \quad \langle \nu_{22} \rangle = -0.572 \pm 0.01 \quad (\varepsilon = 10^{-4}a), (5.3*a*-*c*)

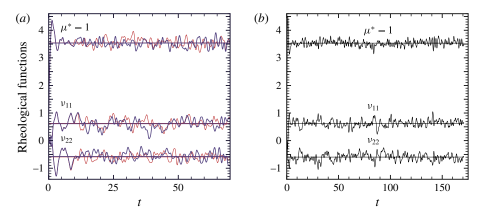


FIGURE 6. (Colour online) The effects of the precision parameter ε and the system size N on the rheological functions in the mixed-flow simulations with c = 0.55, $\lambda = 3$, $\chi = 0.25$ and Ca = 0.0375. In (*a*), the thin lines are for $\varepsilon = 10^{-3}a$ and the thick lines are for $\varepsilon = 10^{-4}a$, all for N = 100. In (*b*), $\varepsilon = 10^{-3}a$ and N = 200. The horizontal lines represent long-time averages.

obtained by excluding short transient parts ($t \le t_{cut} \approx 4$) of the trajectories and shown in figure 6(*a*) by horizontal lines (red thin and blue thick respectively), are in excellent agreement between the two simulations, within 0.6% of $\mu^* - 1$ (the main drop-phase contribution to the stress).

Figure 6(*b*) presents the trajectories of $\mu^* - 1$ and ν_{ij} for a larger system N = 200 ($\varepsilon = 10^{-3}a$) from table 1 and figure 5(*a*). The long-time averages 3.534 ± 0.01 , 0.628 ± 0.01 and 0.597 ± 0.01 for $\mu^* - 1$, ν_{11} and ν_{22} respectively (obtained with $t_{cut} \approx 8$ and shown by horizontal lines) are in excellent agreement with those for N = 100, $\varepsilon = 10^{-3}a$.

A similar accuracy analysis for PE flows with Ca = 0.088, c = 0.55, $\lambda = 3$ and $N_{\Delta} = 2160$ is shown in figure 7, including the trajectories of extensiometric functions (2.2b) $\mu_{pe} - 1$, N_{pe} and their long-time averages. The N = 100, $\varepsilon = 10^{-3}a$ simulation (thin red lines in figure 7a) had to be terminated at t = 105.6 due to extreme elongation of one drop (figure 8a) still without clear signs of breakup. For the same reason, the N = 100, $\varepsilon = 10^{-4}a$ simulation (thick blue lines in figure 7a) was stopped at t = 99.6. In both cases, though, most drops remain compact at the end of the simulation (the average drop length $\langle L_d \rangle$ being (3–3.15)a, versus L = 11a in figure 8a). Therefore, extreme elongation of individual drops still did not preclude the rheology (2.2b) from reaching a statistical steady state. The long-time averages

$$\langle \mu^* - 1 \rangle = 4.44 \pm 0.02, \quad \langle N_{pe} \rangle = 0.72 \pm 0.02 \quad (\varepsilon = 10^{-3}a),$$
 (5.4*a*,*b*)

$$\langle \mu^* - 1 \rangle = 4.43 \pm 0.01, \quad \langle N_{pe} \rangle = 0.73 \pm 0.01 \quad (\varepsilon = 10^{-4}a)$$
 (5.5*a*,*b*)

for the two simulations (obtained using $t_{cut} \approx 12$) agree very well within statistical errors.

The N = 200, $\varepsilon = 10^{-3}a$ PE simulation from figure 5(b) was only continued to t = 35.3 due, again, to extreme elongation of one drop (figure 8b). Here, the incipient breakup is more pronounced. The average drop length at t = 35.3 is only 3.03*a*, and the rheology is still at statistical steady state with the averages $\langle \mu^* - 1 \rangle = 4.46 \pm 0.02$, $\langle N_{pe} \rangle = 0.76 \pm 0.02$ (using $t_{cut} = 8$). These results are close to those for N = 100.

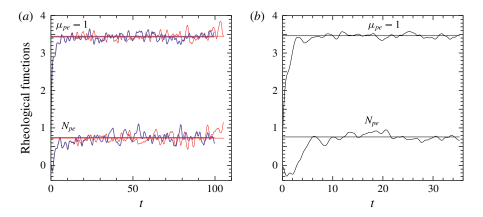


FIGURE 7. (Colour online) The effects of the precision parameter ε and the system size N on the rheological functions in the PE simulations with c = 0.55, $\lambda = 3$ and Ca = 0.088. In (a), the thin lines are for $\varepsilon = 10^{-3}a$ and the thick lines are for $\varepsilon = 10^{-4}a$, all for N = 100. In (b), $\varepsilon = 10^{-3}a$ and N = 200. The horizontal lines represent long-time averages.

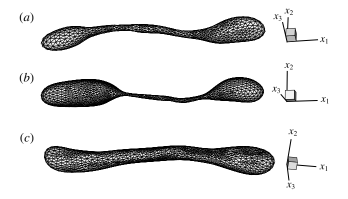


FIGURE 8. The final shape of the most deformed drop in the PE simulations with (a) Ca = 0.088, N = 100, (b) Ca = 0.088, N = 200 and (c) Ca = 0.1, N = 200, all for $\lambda = 3$ and c = 0.55. In (b) and (c), the initial resolution $N_{\triangle} = 2160$ has been refined to $N_{\triangle} = 3840$ for the chosen drop, shortly prior to its extreme deformation.

With Ca just slightly above 0.088, a major portion of the drops experience extreme elongation, so Ca = 0.088 is near-critical. It would be essential to extend rheological simulations to supercritical conditions, when multiple breakups eventually make the system polydisperse. Unfortunately, we currently do not see a prospect of such simulations, at least in PE and strong mixed flows. Emulsion flow through a packed bed was simulated by Zinchenko & Davis (2013), with up to ~60 primary and secondary breakups in an individual run, to observe the drop size evolution and the approach of the statistical steady state for macroscopic properties (phase permeabilities). In those simulations, solid obstacles acted as geometrical constraints limiting drop elongation, which greatly alleviated the numerical analysis of multiple breakups. No such constraints act strongly on drops in pure emulsion flows. The near-critical case in figure 8(b) is not representative of supercritical conditions. More typical is the scenario (at least for PE and strong mixed flows) when emulsion drops

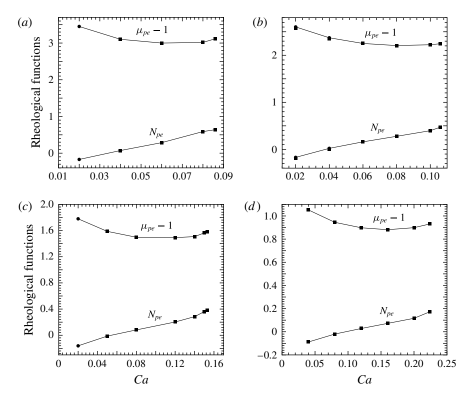


FIGURE 9. Steady-state extensiometric functions at c = 0.45 and N = 100 for $\lambda = 10$ (*a*), 3 (*b*), 1 (*c*) and 0.25 (*d*). Squares, $N_{\Delta} = 2160$; circles, $N_{\Delta} = 3840$. Lines connect the data deemed most accurate.

attain very large deformations still without any signs of incipient breakup (figure 8c). Each such case would need to be extended much further to approach pinch-off and fragment the drop, which is difficult to do repeatedly in a multidrop system. For this reason, the present simulations are only for subcritical conditions.

The above examples are for the 'normal' case, with fast relaxation to the statistical steady state and system-size independence of the results. Far more complex cases are discussed in § 7.

6. Extensiometric functions

In figures 9–11, our systematic calculations are summarized for the steady-state rheological functions $\mu_{pe} - 1$ and N_{pe} in PE. Figures 9(*a*-*d*) and 10(*a*-*d*) cover a broad range of viscosity ratios $\lambda = 10$, 3, 1 and 0.25 for both 45% and 55% drop volume fractions. These results are complemented by figure 11 for the intermediate case c = 0.5 (with $\lambda = 1$ only). Resolutions $N_{\Delta} = 2160-3840$ and system sizes N = 100-200 were used. The rightmost value of *Ca* on each graph is very close to critical for massive breakup to occur. Except for one shorter run (figure 7*b*), these near-critical simulations reached $t \sim 85-120$, with enough statistical accuracy to assess the steady-state $\langle \mu_{pe} - 1 \rangle$ and $\langle N_{pe} \rangle$. For other *Ca*, the runs were terminated at *t* from 120 to several hundred, to make the absolute statistical uncertainties in $\langle \mu_{pe} - 1 \rangle$ and $\langle N_{pe} \rangle$ as small as 0.005 (with 95% confidence) and distil the effects of the surface resolution and the system size.

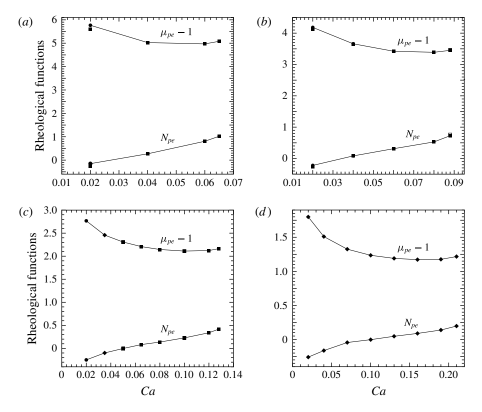


FIGURE 10. Steady-state extensiometric functions at c = 0.55 for $\lambda = 10$ (a), 3 (b), 1 (c) and 0.25 (d). Squares, $N_{\triangle} = 2160$; diamonds, $N_{\triangle} = 2880$; circles, $N_{\triangle} = 3840$. Solid symbols are for N = 100, open symbols (distinct from N = 100 results for only one data point in (b)) are for N = 200. Lines connect the data deemed most accurate.

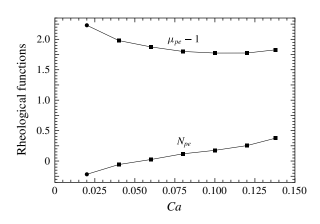


FIGURE 11. Steady-state extensiometric functions for c = 0.5 and $\lambda = 1$; system size N = 100. Squares, $N_{\triangle} = 2160$; circles, $N_{\triangle} = 3840$.

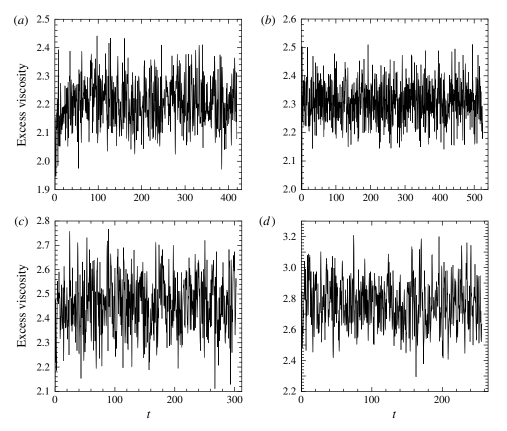


FIGURE 12. Trajectories of the excess viscosity $\mu_{pe} - 1$ in PE simulations for c = 0.55 and $\lambda = 1$ with (a) Ca = 0.065, N = 100, (b) Ca = 0.05, N = 200, (c) Ca = 0.035, N = 100 and (d) Ca = 0.02, N = 100.

An essential characteristic feature of all of our PE simulations (as opposed to the shear-flow runs discussed in § 7) is a short relaxation time to the statistical steady state, regardless of Ca, λ or c, so that cut-out values of $t_{cut} = 4-14$ were always enough for averaging. As an example, figure 12(a-d) gives the trajectories of $\mu_{pe} - 1$ for c = 0.55, $\lambda = 1$ and different capillary numbers. As Ca is progressively reduced from 0.065 to 0.02 for a fixed N = 100, the fluctuations grow significantly due to configurational blockages of less deformed drops, but relaxation to the statistical steady state remains fast. The smaller cross-difference N_{pe} (not shown in figure 12) has the same trends.

Increasing the surface resolution is most important at high emulsion concentrations, large viscosity ratio λ and small *Ca*, due to the increased role of lubrication effects. In the least favourable case c = 0.55, $\lambda = 10$, Ca = 0.02, the effect of N_{Δ} (2160 versus 3840) on $\langle \mu_{pe} - 1 \rangle$ is 3%; for $\langle N_{pe} \rangle$, this difference is 2% of the main stress $\langle \mu_{pe} - 1 \rangle$ (cf. the circles and squares in figure 10*a*). At the same *c*, *Ca* and $\lambda = 3$, these differences are only 1.3% and 1% respectively. At $\lambda = 3$, even smaller triangulation effects are observed for (i) c = 0.55, Ca = 0.04 and (ii) c = 0.45, Ca = 0.02-0.04 (cf. the solid squares and circles in figures 9*b* and 10*b*).

The slight difference between the results for N = 100 (solid squares) and those for N = 200 (open squares) at the near-critical Ca = 0.088 in figure 10(b) may be mostly statistical, due to the short averaging interval in the N = 200 run (see § 5 and figure 7b). In two other tests, for c = 0.55, $\lambda = 1$, and Ca = 0.05 and 0.1, carried out to $t \sim 500$ to greatly reduce the statistical errors, the effect of N (100 versus 200) on $\langle \mu_{pe} - 1 \rangle$ and $\langle N_{pe} \rangle$ did not exceed 0.2% of $\langle \mu_{pe} - 1 \rangle$. As a result of this analysis, the lines connecting the most reliable data in figures 9-11 are believed to accurately describe the extensiometric functions, resolution- and system-size-independent. Several physical trends in $\langle \mu_{pe} - 1 \rangle$ and $\langle N_{pe} \rangle$ (without the angular brackets in what follows) are interesting to discuss. The cross-difference N_{pe} is generally small compared with the excess viscosity $\mu_{pe} - 1$, but can reach 24% at the critical Ca. For small enough Ca, this cross-difference is negative (which is not a numerical artefact), and definitely remains negative in the limit $Ca \rightarrow 0$. The excess viscosity $\mu_{pe} - 1$, as a function of Ca, has a shallow minimum, so that the initial sharp tension thinning is followed by mild tension thickening (in contrast, a dilute emulsion only tension thickens in PE, Martin et al. 2014). In the phenomenological theory of non-Newtonian 'simple liquids' (e.g. Astarita & Marucci 1974), certain smoothness hypotheses are assumed and, as a result, the constitutive equation in the limit of slow flows is shown to be of Rivlin & Ericksen (1955) type. This general form was indeed confirmed by a small-deformation analysis for a single drop in linear flows (Schowalter et al. 1968; Frankel & Acrivos 1970; Barthés-Biesel & Acrivos 1973b). If the Rivlin–Ericksen theory were applicable to our concentrated emulsions of deformable drops, then, in particular, the viscosity μ_{pe} would approach its limiting value at Ca = 0 with an error of $O(Ca^2)$, and N_{pe} would vanish as O(Ca). Both phenomenological predictions, though, contradict our numerical results in figures 9-11, so a concentrated emulsion (of non-Brownian drops) is not a Rivlin-Ericksen fluid (in support of the views from paper I). This stark difference finds an easy explanation. In a flowing concentrated emulsion at $Ca \ll 1$, drops are nearly in contact, and a small-deformation analysis of such close interactions (hardly feasible in practice because of its extreme complexity) would be a problem of singular perturbations. Therefore, unlike for single drops, a regular shape expansion in Ca would no longer be possible, which should affect the behaviour of the rheological functions in the limit $Ca \ll 1$. However, it would be prohibitively expensive to extend the convergent results in figures 9-11 to even smaller *Ca* to further elucidate this issue.

7. Viscometric functions

A far more complex picture emerges in shear-flow simulations for concentrated emulsions, requiring a more difficult analysis.

7.1. Results for c = 0.45

Figure 13(*a*–*d*) summarizes our calculations for the non-dimensional steady state viscometric functions – the excess shear viscosity $\langle \mu_{sh} - 1 \rangle$ and normal stress differences $\langle N_1^{sh} \rangle$ and $\langle N_2^{sh} \rangle$ (defined in (2.2*a*)) at 45% drop volume fractions and different viscosity ratios λ . Resolutions $N_{\Delta} = 2160-3840$ and system sizes N = 100-200 were used, as specified in the caption. For $\lambda = 10$ (figure 13*a*) and Ca = 0.06-0.02, the runs were stopped at non-dimensional times (i.e. strains) t = 60-66, to average $\mu_{sh} - 1$, N_1^{sh} and N_2^{sh} with estimated absolute errors of 0.01–0.02. As an example of the difficulties of such $\lambda = 10$ simulations, the run N = 100, $N_{\Delta} = 3840$ for the smallest Ca = 0.02 in figure 13(*a*) took 160000 time steps with ≈ 10 BI iterations per time step. For the same λ and higher Ca = 0.08, 0.10 and 0.16, the runs could not be continued beyond t = 46, 42 and 27 respectively, due to divergence of the BI iterations. Obviously, our variational technique (§ 3.1)

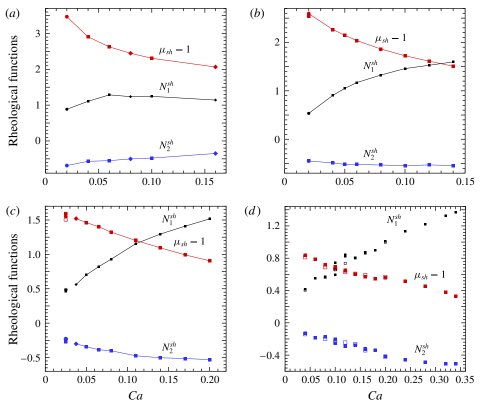


FIGURE 13. (Colour online) Steady-state viscometric functions at c = 0.45 for $\lambda = 10$ (*a*), 3 (*b*), 1 (*c*) and 0.25 (*d*). Squares, $N_{\Delta} = 2160$; diamonds, $N_{\Delta} = 2880$; circles, $N_{\Delta} = 3840$. Solid symbols are for N = 100, open symbols are for N = 200. Small-size symbols are used for N_1^{sh} to distinguish them from $\mu_{sh} - 1$ values. Lines connect the data points selected for constitutive modelling in §§ 8 and 9.

of the double-layer near-singularity subtraction, although highly beneficial, may be not powerful enough for concentrated emulsions at extreme viscosity ratios and large deformations. These shorter runs still gave the averaging errors likely within 0.01–0.02, since the increase of *Ca* reduces the statistical fluctuations. For all *Ca* in figure 13(*a*), relaxation to the statistical steady state is fast, and cut-out times $t_{cut} = 2-12$ sufficed. The results in figure 13(*a*) show significant shear thinning at the smallest *Ca*. The first normal stress difference $\langle N_1^{sh} \rangle$, smaller but comparable with $\langle \mu_{sh} - 1 \rangle$, is also a strong function of *Ca* when $Ca \ll 1$, but it levels off at larger drop deformations. The negative second normal stress difference is small compared with $\langle \mu_{sh} - 1 \rangle$.

When $\lambda = 3$ (figure 13*b*), relaxation to the statistical steady state remains fast for all capillary numbers considered. We used $t_{cut} = 2.5-5$ to exclude the transient regimes and integrated to $t \sim 100-300$ in each run to reduce likely statistical errors in $\langle \mu_{sh} - 1 \rangle$, $\langle N_1^{sh} \rangle$ and $\langle N_2^{sh} \rangle$ to $\pm 0.005-0.01$. Divergence of BI iterations was never encountered at this λ . At the smallest Ca = 0.02 (with the strongest surface resolution effect), the results were obtained for both $N_{\Delta} = 2160$ and 3840:

 $\langle \mu_{sh} - 1 \rangle = 2.539, \quad \langle N_1^{sh} \rangle = 0.540, \quad \langle N_2^{sh} \rangle = -0.446 \quad (N_{\triangle} = 2160), \quad (7.1a-c)$

$$\langle \mu_{sh} - 1 \rangle = 2.594, \quad \langle N_1^{sh} \rangle = 0.534, \quad \langle N_2^{sh} \rangle = -0.449 \quad (N_{\triangle} = 3840). \quad (7.2a - c)$$

The deviations between $\langle \mu_{sh} - 1 \rangle$ are only 2% and are negligible for $\langle N_1^{sh} \rangle$ and $\langle N_2^{sh} \rangle$. The triangulation effects should be much smaller for the other *Ca* values in figure 13(*b*). The qualitative trends of the viscometric functions are roughly the same as for $\lambda = 10$.

For $\lambda = 1$ (figure 13*c*), relaxation to the statistical steady state is fast ($t_{cut} = 3-4$) in all cases considered, except for Ca = 0.025, N = 200 when $t_{cut} = 45$ had to be excluded from the averaging. Even with fast relaxation in other runs, the statistical fluctuations grow dramatically as Ca = 0.025 is approached. These are signs of configurational 'phase transition' to a partially ordered state. The final simulation times were $T \sim 100$ for all $Ca \ge 0.11$, but 400–900 for all $Ca \le 0.065$. For the most difficult case Ca = 0.025, the resolution and system-size effects on the steady-state viscometric functions were explored:

$$\langle \mu_{sh} - 1, N_1^{sh}, N_2^{sh} \rangle = 1.591, 0.462, -0.269 \quad (N = 100, N_{\Delta} = 2160),$$
 (7.3)

$$\langle \mu_{sh} - 1, N_1^{sh}, N_2^{sh} \rangle = 1.549, 0.477, -0.226 \quad (N = 100, N_{\triangle} = 3840),$$
(7.4)

$$\langle \mu_{sh} - 1, N_1^{sh}, N_2^{sh} \rangle = 1.502, 0.488, -0.233 \quad (N = 200, N_{\triangle} = 3840).$$
 (7.5)

These effects are within 3% of the main drop-phase contribution $\langle \mu_{sh} - 1 \rangle$ to the stress. The increased system-size sensitivity (cf. the extensiometric functions in §6) is another sign of (incipient) phase transition, when a partially ordered emulsion finds it easier to flow with lower viscosity. This picture of eventual partial ordering is confirmed by the graphical analysis of drop configurations at Ca = 0.025, N = 200 and large times, with visible layers of drops sliding past each other along x_1 , and an approximately hexagonal structure in the x_2 , x_3 plane. No signs of order were detected for $Ca \ge 0.0375$.

The case of low viscosity ratio $\lambda = 0.25$ (figure 13d) stands out, by far, as the most difficult, since anomalies are observed in a wide range of capillary numbers. The simulation times were $T \sim 60-80$ for $Ca \ge 0.24$, but much larger (T = 200-800) for all smaller Ca. Rather fast relaxation $(t_{cut} \sim 10)$ to the statistical steady state for $Ca \ge 0.24$ dramatically slows down with Ca reduced to 0.16, and it remains slow $(t_{cut} \sim 100-200)$ for all smaller Ca. To illustrate this point, figure 14 presents the trajectories of $\mu_{sh} - 1$, N_1^{sh} and N_2^{sh} for (a) Ca = 0.16, (b) Ca = 0.12 and (c,d) Ca = 0.1, all at N = 100and $N_{\Delta} = \tilde{2}160$. For (*c*,*d*), two different well-mixed initial configurations of spherical drops were used; to avoid overlapping of data, the trajectories of $\mu_{sh} - 1$ are shown as insets. By excluding the transient parts ($t_{cut} = 125$, 150, 230 and 100 for (a-d) respectively), excellent averaging quality is achieved, with the absolute errors in $\langle \mu_{sh} - 1 \rangle$, $\langle N_1^{sh} \rangle$ and $\langle N_2^{sh} \rangle$ within 0.004 for (a), and within 0.002 for (b-d). However, the two Ca = 0.1 runs clearly belong to two different ergodic classes. While the effect of the initial configuration on $\langle \mu_{sh} - 1 \rangle$ is small (0.675 in c versus 0.694 in d), it is more noticeable for $\langle N_2^{sh} \rangle$ (-0.202 versus -0.253) and particularly large for $\langle N_1^{sh} \rangle$ (0.594 versus 0.747). In contrast, we did not find any appreciable effects of the initial drop configuration on the time averaging for Ca = 0.12 and 0.16 at N = 100. Ergodic difficulties akin to those at Ca = 0.1 are familiar in phase transition simulations for molecular systems by molecular dynamics or Monte Carlo methods. In principle, increasing the system size should merge separate ergodic classes inaccessible to each other, and make average results independent of the initial conditions. In practice, though, such system-size increase can make the averaging task much harder, if not insurmountable, since the system must now transition with enough frequency between metastable quasiergodic classes for good statistics. For N = 200, we successfully obtained steady-state viscometric functions at Ca = 0.04, 0.08, 0.1, 0.12, 0.14, 0.16

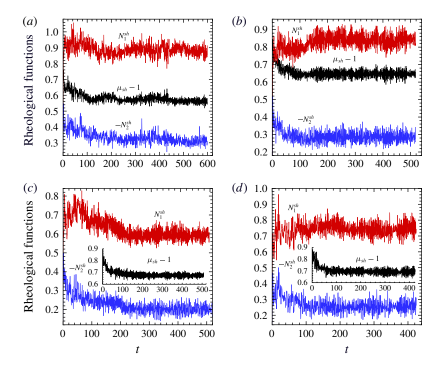


FIGURE 14. (Colour online) Trajectories of the viscometric functions for c = 0.45, $\lambda = 0.25$, N = 100, $N_{\triangle} = 2160$ at (a) Ca = 0.16, (b) Ca = 0.12 and (c,d) Ca = 0.1.

and 0.2 (although with larger statistical uncertainties of $\pm 0.01-0.02$ in $\langle N_1^{sh} \rangle$ and $\langle N_1^{sh} \rangle$) to compare with those for N = 100. The only large differences were found for $\langle N_1^{sh} \rangle$ at Ca = 0.1 and 0.12. From this analysis, $Ca \approx 0.1-0.12$ is believed to be the left end of the phase transition interval (while the right end is estimated as $Ca \approx 0.18$). The excess viscosity $\langle \mu_{sh} - 1 \rangle$ in figure 13(d) is the least sensitive to N and ergodic difficulties, and is reasonably accurate in the whole range of $0.04 \leq Ca \leq 0.34$ considered. In particular, the kink of the viscosity curve near Ca = 0.18 is not a numerical artefact.

Our results in figure 13 confirm the hypothesis (suggested in paper I, although with little support therein) that increasing λ shifts the phase transition in the emulsion shear flow to smaller capillary numbers. Indeed, the related ergodic difficulties, so pronounced at $\lambda = 0.25$, are not felt at $\lambda = 1$ until *Ca* is reduced to 0.025, and these difficulties are absent altogether for $\lambda = 3$ and 10 in the whole range of *Ca* considered. One reason for this shift is the increase in the drop deformation in the flow with higher λ (in the dilute limit, the trend would be the opposite for $\lambda > 1$). This larger deformation helps to eliminate geometrical blockages and merge different ergodic classes. Another reason is, presumably, more efficient mixing in the flow with higher λ acting the same way.

The classical model of a liquid of molecules interacting as hard spheres is known to exhibit phase transition to the solid state in the interval $0.494 \le c \le 0.545$ of volume fractions c (e.g. Hoover & Ree 1968; Hansen & McDonald 1976), with discontinuities of the first derivatives of pressure versus c at the end points. The present case of dynamical phase transition in emulsion shear flow is certainly different. Still, it

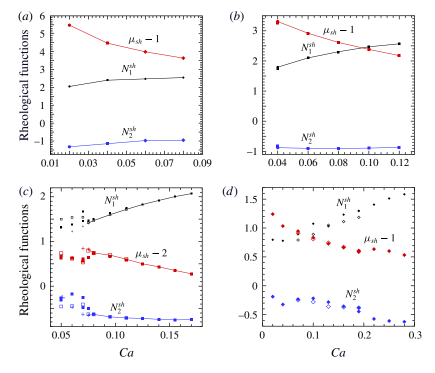


FIGURE 15. (Colour online) Steady-state viscometric functions at c = 0.55 for $\lambda = 10$ (*a*), 3 (*b*), 1 (*c*) and 0.25 (*d*). Squares, $N_{\Delta} = 2160$; diamonds, $N_{\Delta} = 2880$; circles, $N_{\Delta} = 3840$. Solid symbols are for N = 100, open symbols are for N = 200. Plus symbols are for N = 400 with $N_{\Delta} = 2160$. In (*c*), $\mu_{sh} - 2$ is presented (instead of $\mu_{sh} - 1$) to avoid data overlapping. Small-size symbols are used for N_1^{sh} to distinguish them from the viscosity values. Lines connect the data points selected for constitutive modelling in §§ 8 and 9.

is unexpected that we observe phase-transitional behaviour at significantly smaller volume fraction c = 0.45 without non-hydrodynamical attractive forces between the drops.

7.2. *Results for* c = 0.55

Our calculations of the steady-state non-dimensional viscometric functions $\langle \mu_{sh}-1\rangle$, $\langle N_1^{sh}\rangle$ and $\langle N_2^{sh}\rangle$ at c = 0.55 are summarized in figure 15(*a*-*d*). Again, at $\lambda = 10$ (figure 15*a*), proceeding to large simulation times is most prohibitive due to smaller time steps and a larger number of BI iterations. For this λ , the runs Ca = 0.02 and 0.04 were stopped at t = 63 and 74 respectively. At larger Ca = 0.06 and 0.08, though, eventual divergence of the BI iterations did not allow us to proceed beyond t = 35 and 45 respectively, even with the resolution $N_{\Delta} = 2880$. Still, the absolute statistical errors for all the data in figure 15(*a*) are expected to be within ± 0.02 . Relaxation to the statistical steady state was fast for all capillary numbers considered, and $t_{cut} = 2-5$ was sufficient to exclude the transient stages from averaging. As well as for c = 0.45 (figure 13*a*), the $\lambda = 10$ emulsion shows most pronounced shear thinning, but is also significantly more viscous with the drop volume fraction increased to 55 %.

For $\lambda = 3$ (figure 15b), the steady-state results could not be extended to Ca = 0.02 for an unexpected reason. In figure 16, the trajectories of the excess viscosity $\mu_{sh} - 1$

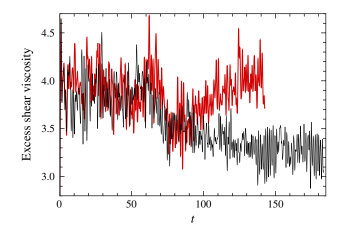


FIGURE 16. (Colour online) Two trajectories of the excess shear viscosity $\mu_{sh} - 1$ for c = 0.55, $\lambda = 3$, Ca = 0.02 and N = 100, starting from the same random configuration of spherical drops. Thin line, $N_{\triangle} = 2160$; thick line, higher resolution $N_{\triangle} = 3840$.

are shown for two simulations with Ca = 0.02 and N = 100. Both simulations start from the same random configuration of spherical drops, the thin (blue) line being for $N_{\Delta} = 2160$, the thick (red) line for $N_{\Delta} = 3840$. The two runs appear to quickly reach statistical steady states and closely agree, on average, up to $t \sim 65$. After $t \sim 70$, though, the lower-resolution run $N_{\triangle} = 2160$ abruptly settles to a different state with much lower viscosity, while the run $N_{\triangle} = 3840$ remains at the higher viscosity. We believe that the only explanation for this erratic behaviour and the unusually strong effect of the resolution is that the conditions are on the verge of phase transition, and a small numerical perturbation (change in N_{Δ}) brings the system to a different quasiergodic class after a long time. Graphical analysis of drop configurations confirms that the $N_{\Delta} = 2160$ run settles to a partially ordered state, unlike the other run. To make sure that such difficulties (compromising time averaging) do not appear for larger Ca in figure 15(b), we performed detailed testing of the effects of the initial conditions and programme parameters on the steady-state functions $\langle \mu_{sh} - 1 \rangle$, $\langle N_1^{sh} \rangle$, $\langle N_2^{sh} \rangle$ at Ca = 0.04. Two-level behaviour (akin to that in figure 16 for $N_{\Delta} = 2160$) was never observed in these test runs; the simulated times were T = 230-470, with always fast relaxation ($t_{cut} = 4$) to the statistical steady state. For one initial drop configuration, the $N_{\triangle} = 2160$ results (solid squares in figure 15b) at Ca = 0.04) are (3.254, 1.745, -0.817). Repeating the run with $N_{\Delta} = 3840$ gave (3.307, 1.793, -0.879) (solid circles in figure 15b). Tightening the 'precision' ε to $10^{-4}a$ in the $N_{\triangle} = 2160$ run yielded (3.267, 1.733, -0.852). Coarsening the barrier δ_{min} (which controls drop overlap, see § 4) to 0.003*a* resulted in (3.271, 1.743, -0.846). Finally, replacing the initial drop configuration in the $N_{\Delta} = 2160$ run gave (3.269, 1.770, -0.832). The deviations from the base results (shown by solid squares in figure 15b) are within 1.8% of $\langle \mu_{sh} - 1 \rangle$, the main drop-phase contribution to the stress, and often much less. Therefore (unlike for Ca = 0.02), the steady-state results for $Ca \ge 0.04$ are stable to the programme parameters and do not suffer from ergodic difficulties. For $Ca \ge 0.06$, shorter simulation times $T \sim 110-160$ were sufficient (with $t_{cut} = 4-5$). Although the results in figure 15(b) (and 15a) are for N = 100 only, any significant N dependence is not suspect; the latter could only result from phase transition and related ergodic difficulties.

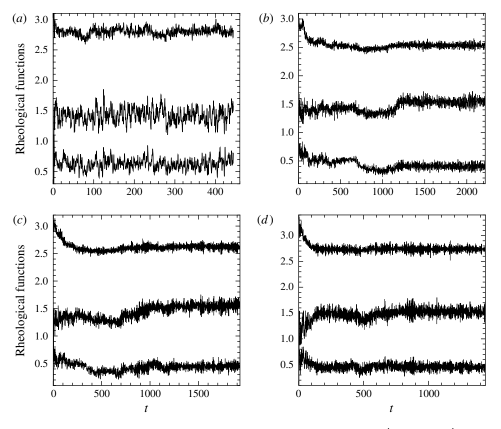


FIGURE 17. The trajectories of the viscometric functions μ_{sh} , N_1^{sh} and $-N_2^{sh}$ (top to bottom on each graph) for c = 0.55, $\lambda = 1$, N = 200 and $N_{\triangle} = 2160$; (a) Ca = 0.075, (b) Ca = 0.07, (c) Ca = 0.06, (d) Ca = 0.05.

When λ is decreased to 1 (figure 15c), the phase transition is shifted to much higher Ca; the ergodic difficulties in the transition range are severe and appear to be insurmountable, even when using N = 400 drops. Figure 17(a-d) shows the trajectories of $\mu_{sh}(t)$, $N_1^{sh}(t)$ and $-N_2^{sh}(t)$ at N = 200 and $N_{\Delta} = 2160$, when Ca is progressively decreased from (a) 0.075 to (d) 0.05. Here, we plot the full viscosity (instead of μ_{sh} – 1) to avoid data overlapping. Fast approach to the statistical steady state at Ca = 0.075 ($t_{cut} = 4$) changes to very slow relaxation ($t_{cut} \sim 1000-1200$) at Ca = 0.06and 0.07. Relaxation is still slow ($t_{cut} = 160$), although not as slow, at Ca = 0.05. The interval $Ca \approx 0.06-0.07$ is therefore believed to be phase-transitional. For $Ca \ge 0.075$, very good averaging quality is achieved with simulation times $T \sim 200-600$. The results are well convergent with the number of drops N = 100-200 when $Ca \ge 0.08$ (as tested for Ca = 0.08-0.11), but the system size has to be increased to N = 200-400at Ca = 0.075 for convergence. For $Ca \le 0.07$, all the runs were extended to much larger times for adequate averaging, but the results still show a strong dependence on the initial drop configuration (as tested for N = 100, cf. solid squares in figure 15c) and on the system size N = 100-400. Therefore, only the $Ca \ge 0.075$ portion of the results in figure 15(c) is trusted as unambiguous and is used for constitutive modelling in \S and 9.

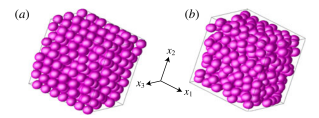


FIGURE 18. (Colour online) Snapshots from the shear-flow simulations at c = 0.55, $\lambda = 1$, N = 400 and $N_{\triangle} = 2160$ with (a) Ca = 0.05, t = 837 and (b) Ca = 0.06, t = 1155. The centres of N independent drops have been mapped into the cube $(0, 1)^3$; the same view is shown in (a) and (b).

It is very instructive to compare the transition from fast to slow relaxation in figure 17 with figure 12 for PE at the same c = 0.55 and $\lambda = 1$. For shear flow (figure 17), we observe this transition to occur when the average drop length $\langle L_d \rangle$ in the steady state (correlating with the drop deformation and the degree of configurational arrest) decreases from 2.42*a* at Ca = 0.075 to 2.36*a* at Ca = 0.07. In contrast, in the PE simulations (figure 12), $\langle L_d \rangle$ progressively decreases from 2.42*a* at Ca = 0.06a to much smaller, 2.15*a*, at Ca = 0.02, but the relaxation remains fast. This comparison reveals a profound difference between the two types of flow: in PE, phase transition does not occur regardless of how small Ca is, at least for drop volume fractions $c \leq 0.55$. The same is true for other viscosity ratios λ .

The small viscosity ratio $\lambda = 0.25$ at c = 0.55 (figure 15*d*) is another very challenging case in the shear-flow simulations. The transition from fast $(t_{cut} \sim 10)$ to slow $(t_{cut} \sim 100)$ relaxation is observed when $Ca \approx 0.19-0.22$, which estimates the right end of the phase transition interval. The simulations for figure 15(*d*) were performed to T = 72-75 at $Ca \ge 0.25$, and to $T \sim 200-400$ at each smaller $Ca \le 0.22$. There is no doubt in the accuracy of all steady-state viscometric functions for Ca > 0.22. In the transition range, though, strong effects of the initial drop configuration and/or the system size N = 100-200 on the average normal stress differences (especially $\langle N_1^{sh} \rangle$) are observed. In contrast, the excess viscosity $\langle \mu_{sh} - 1 \rangle$ is almost insensitive to these difficulties and is well convergent even in the transition range, at least for $Ca \ge 0.07$. Again, the viscosity curve has a kink at $Ca \approx 0.19$, near the upper end of the transition range.

Since the flow-induced partially ordered drop arrangement has an approximately hexagonal structure in the (x_2, x_3) plane, it would seem advantageous to adapt the shape of the periodic cell to this structure. Such algorithm modification, though, was not attempted in the present work for two reasons. The present version of the code heavily exploits the symmetry properties of the periodic Green function G(r) about the plane $r_3 = 0$, so an extension to an arbitrary geometry, with $e_2(t)$ non-orthogonal to e_3 , would not be straightforward. More importantly, while such an algorithm adaptation can alleviate ergodic and convergence difficulties in the simulations well below the transition point to order, it would not help near the transition with the coexistence of ordered and disordered states. Nor would such an adaptation help to identify the onset of transition. As an example, figure 18(a) shows a well-ordered flow achieved at a large strain t = 837 in the simulation with c = 0.55, $\lambda = 1$, N = 400 and Ca = 0.05. For Ca = 0.06, still below the right end of the transition interval, the emulsion is not nearly as ordered (figure 18b).

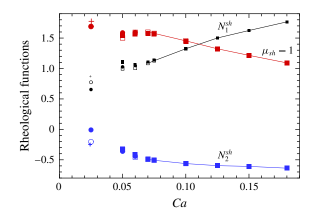


FIGURE 19. (Colour online) Steady-state viscometric functions at c = 0.5 and $\lambda = 1$. Squares, $N_{\triangle} = 2160$; circles, $N_{\triangle} = 3840$. Solid symbols are for N = 100, open symbols are for N = 200. Plus symbols are for N = 100 with $N_{\triangle} = 6000$. Small-size symbols are used for N_1^{sh} . Lines connect the data selected for constitutive modelling in §§ 8 and 9.

7.3. The intermediate case c = 0.5

At the intermediate volume fraction of c = 0.5, we have studied the steady-state viscometric functions for $\lambda = 1$ only (figure 19). For moderate $Ca \ge 0.1$, simulation times T = 80-190 were sufficient to provide high averaging accuracy. As Ca is decreased to 0.06–0.07, the statistical fluctuations sharply grow, necessitating much larger simulation times (up to $T \sim 3000$), but there is still no clear transient regime, so we used $t_{cut} = 3-5$ for all $0.06 \le Ca \le 0.18$. For all $Ca \ge 0.07$, the results are proven to be practically system-size-independent (compare the solid squares at N = 100with the open squares at N = 200 at Ca = 0.07, 0.075 and 0.1; indistinguishable for $\langle N_2^{sh} \rangle$). At Ca = 0.06, the agreement between the N = 100 and N = 200 results is less favourable for $\langle N_1^{sh} \rangle$ and $\langle N_2^{sh} \rangle$. At this *Ca*, there is also a comparable and modest resolution effect when N_{Δ} is increased from 2160 (solid squares) to 3840 (solid circles). Pronounced slow relaxation to the statistical steady state is first observed when Ca is decreased to 0.05. In the four runs for this Ca, the simulation times were T = 800-1900 with cutoffs $t_{cut} = 160-400$. Starting the N = 100, $N_{\triangle} = 2160$ runs from two different initial configurations of spherical drops gave practically indistinguishable results. Increasing the resolution to $N_{\triangle} = 3840$ (solid circles) increases the excess viscosity (from 2.55 to 2.58) and the magnitude of $\langle N_2^{sh} \rangle$ (from 0.32 to 0.36), but reduces $\langle N_1^{sh} \rangle$ (from 1.10 to 1.03). Increasing the system size to N = 200 (open squares) has the opposite effect of slightly reducing the viscosity (which we associate with the phase-transitional behaviour) and further reducing $\langle N_1^{sh} \rangle$; the second normal stress difference is not affected.

In the three most difficult runs at Ca = 0.025, (i) N = 100, $N_{\triangle} = 3840$, (ii) N = 200, $N_{\triangle} = 3840$ and (iii) N = 100, $N_{\triangle} = 6000$, our simulation times were limited to T = 900, 500 and 400 respectively, with still acceptable quality of averaging. There are only modest variations, if any, in the excess viscosity $\langle \mu_{sh} - 1 \rangle$ between the runs, but serious misconvergence in the normal stress differences.

Overall, the accuracy of our viscosity calculations in figure 19 is sufficient to state the existence of a shallow local minimum of $\langle \mu_{sh} - 1 \rangle$ versus *Ca* near *Ca* \approx 0.05–0.06 due to phase transition, followed by increase in the viscosity as the capillary number is further reduced. It is more difficult to argue about the behaviour of $\langle N_1^{sh} \rangle$ in the transition region due to larger uncertainties. Only the $Ca \ge 0.07$ part of the results from figure 19 is used in the constitutive modelling of §§ 8 and 9.

7.4. Comparison with paper I and other work

Paper I was the first to present large-scale shear-flow simulations and steady-state viscometric functions for concentrated emulsions. In addition to the results for $c \leq 0.4$ (not relevant to the present work), $\langle \mu_{sh} \rangle$, $\langle N_1^{sh} \rangle$ and $\langle N_2^{sh} \rangle$ were reported versus *Ca* for (i) c = 0.45 and 0.5 at $\lambda = 1$ and (ii) c = 0.45 at $\lambda = 3$. The viscosity values were also reported versus *Ca* for c = 0.55 at $\lambda = 1$ and 3, but it was not possible to even estimate the steady-state normal stress differences in either case due to the short simulation times and large statistical fluctuations.

In addition to substantially new calculations for the difficult viscosity ratios of $\lambda = 0.25$ and 10 at c = 0.45 and 0.55, the present results for $\lambda = 1$ and 3 stem from a far more systematic analysis, with much larger simulated strains for averaging, finer surface triangulations and (often) larger system size. In paper I with $N \sim 100$, the strains were limited to 25–45 for $\lambda = 1$ and only $\sim 15-25$ for $\lambda = 3$, while they are at least of the order of hundreds for all λ (and even thousands for $\lambda = 1$) in the present work. For $\lambda = 3$, the surface resolutions in paper I were limited to $N_{\Delta} \leq 1500$, compared with 2160–3840 used herein for all $\lambda \neq 1$. Finally, in the present work the maximum system size was extended to N = 400 for matching, and to N = 200 for contrast viscosities, twice as large as in paper I.

For moderate Ca (roughly when the average drop length $\langle L_d \rangle$ exceeds 2.6a), the present results for $\lambda = 1$ and 3 are fully consistent with those of paper I, within the statistical errors therein. In the most difficult range of small Ca (and $c \ge 0.45$), the results of paper I somewhat overpredict the first normal stress difference and (when $\lambda = 1$) the shear viscosity. The reason is twofold. First, Zinchenko & Davis (2004) used the same code as in paper I and in the same limited strain range, but with ultrahigh resolutions $N_{\Delta} = 6000-8640$ to recalculate the $\lambda = 1$ shear-flow results for c = 0.45-0.5 at Ca = 0.025 and 0.05. This change results, indeed, in noticeably smaller $\langle \mu_{sh} \rangle$ and $\langle N_1^{sh} \rangle$. Presumably, tiny surface overlaps (allowed by the code of paper I) favour some drop clustering (especially at small Ca and $\lambda = 1$ versus $\lambda = 3$), leading to higher emulsion viscosity and $\langle N_1^{sh} \rangle$, unless this artefact is eliminated by ultrahigh surface resolution. The present code, with explicit surface overlap control, achieves much faster convergence for the viscometric functions with respect to surface triangulations at small Ca. Second, the corrected viscosity values from Zinchenko & Davis (2004) are still slightly higher than the more accurate results herein (most noticeably, at c = 0.5, $\lambda = 1$ and Ca = 0.025, their work gives $\langle \mu_{sh} \rangle = 3.07$ versus the narrow range of 2.69–2.77 from our figure 19). This remaining discrepancy is obviously due to the relatively short simulation times in paper I and in Zinchenko & Davis (2004), so that averaging was, in fact, performed over metastable levels, well before the emulsion viscosity settles down to the true statistical steady state. Accordingly, the phase-transitional behaviour at c = 0.5 and $\lambda = 1$ could not be detected in those studies.

At c = 0.55 and $\lambda = 1$, the viscosity calculations of paper I are indicative of phase transition, but the results for $Ca \leq 0.05$ suffer again from limited triangulations and statistical errors due to short simulation times. For $Ca \geq 0.1$, good agreement with the present work is observed in the viscosity values, but paper I does not give the normal stress differences to compare with.

All of our shear-flow simulations in this section are for essentially subcritical conditions, away from drop breakup. When combining the extensiometric and

viscometric functions for general constitutive modelling (as suggested in the next section), early breakup in planar-extensional flow is the limiting factor (except for small λ), which was the main reason not to try extending the present shear-flow simulations to breakup. Incidentally, the results of paper I for $\lambda = 1$ and 3 come closer to breakup conditions, and they do not suffer in that range from limited triangulations or short simulation times.

8. A generalized Oldroyd model for constitutive modelling

A principal question is whether the rheological functions obtained for steady simple shear and PE can help in general constitutive modelling; that is to predict, with some accuracy, the stress tensor in (large-strain) flows with an arbitrary history of deformation of a material element. Knowing such a constitutive equation would allow the solution of boundary-value problems of non-Newtonian hydrodynamics in complex geometries of technological interest. Obviously, the information from shear flows only (steady or oscillatory) would not suffice for this purpose. In Martin et al. (2014), we suggested using a generalized Oldroyd model with five variable material parameters, found from simultaneously fitting the model to viscometric (in steady simple shear) and extensiometric (in PE) functions at arbitrary flow intensities. These parameters are postulated to depend on one carefully chosen instantaneous flow invariant. The approach was validated in a broad class of kinematics for a dilute emulsion of single deformable drops. For this particular system, the BI method was used to both prepare the viscometric and extensiometric functions at arbitrary Ca and provide exact results for the drop rheological response in a flow with arbitrary kinematics. Although drop breakup is a limitation, this approach was found to give, for finite drop deformation, far more accurate results than small-Ca expansions.

Below, we are trying to explore (although not as comprehensively) this new method of constitutive modelling in application to flowing concentrated emulsions. It is convenient, until otherwise stated, to return to dimensional variables. The macroscopic instantaneous deviatoric stress T for a material element of an emulsion can be written as $T = 2\mu_e E + \tau^d$ to single out the Newtonian continuous-phase contribution. Unlike the intrinsic stress τ^d in Martin *et al.* (2014), our excess stress (drop-phase contribution) τ^d is not normalized by the drop-phase volume fraction. Instead of u_{∞} and E_{∞} (cf. § 2), we use here simpler notations v and $E = (\nabla v + \nabla^T v)/2$ for the local macroscopic emulsion velocity v and the corresponding rate-of-strain tensor. The Oldroyd model can be formulated either for the full stress T or, equivalently, for the excess part:

$$\left(\boldsymbol{I} + \lambda_1 \frac{\mathscr{D}}{\mathscr{D}t}\right) \boldsymbol{\tau}^d - \mu_1 (\boldsymbol{\tau}^d \cdot \boldsymbol{E} + \boldsymbol{E} \cdot \boldsymbol{\tau}^d) + \frac{2}{3} \mu_1 (\boldsymbol{\tau}^d : \boldsymbol{E}) \boldsymbol{I}$$
$$= 2\eta \left[\boldsymbol{E} + \lambda_2 \frac{\mathscr{D} \boldsymbol{E}}{\mathscr{D}t} - 2\mu_2 \boldsymbol{E}^2 + \frac{2}{3} \mu_2 (\boldsymbol{E} : \boldsymbol{E}) \boldsymbol{I} \right].$$
(8.1)

The Jaumann or corotational derivative $\mathcal{D}/\mathcal{D}t$ can be calculated in fixed Cartesian axes for any tensor **A** as

$$\left(\frac{\mathscr{D}\boldsymbol{A}}{\mathscr{D}t}\right)_{ij} = \left(\frac{\mathbf{D}\boldsymbol{A}}{\mathbf{D}t}\right)_{ij} + \frac{1}{2}(\boldsymbol{\nabla}_i\boldsymbol{v}_k - \boldsymbol{\nabla}_k\boldsymbol{v}_i)A_{kj} + \frac{1}{2}(\boldsymbol{\nabla}_j\boldsymbol{v}_k - \boldsymbol{\nabla}_k\boldsymbol{v}_j)A_{ik}, \quad (8.2)$$

where D/Dt is the material derivative. Equation (8.1) can be derived from a very general multiparameter phenomenological model of Hand (1962), if his coefficients

 β_i for $i \ge 3$, α_j for $j \ge 6$ and α_3 are all set to zero. With these restrictions, the structural tensor **a** in Hand's model is linearly connected to **E** and **T**, and Hand's evolution equation for **a** results in (8.1). Oldroyd (1958) originally postulated an eightparameter model (with constant coefficients), which reduces to (8.1) if formulated for the deviatoric excess stress τ^d (i.e. with $I: \tau^d = 0$). However, to match the model (8.1), precisely and unambiguously, to the calculated viscometric and extensiometric functions at arbitrary flow intensities, the material parameters λ_1 , μ_1 , λ_2 , μ_2 and η must be allowed to be functions of one instantaneous flow invariant. In the dilute case, Martin *et al.* (2014) explored two choices for this invariant: (i) the second invariant **E**: **E** of the rate-of-strain tensor and (ii) the excess energy dissipation rate $\tau^d : \mathbf{E}$. Both versions, of course, satisfy the objectivity principle (and do not contradict Hand's model), but the less obvious choice (ii) was found to give a significantly more accurate and less restrictive model. Accordingly, here we follow the choice (ii) (which makes the model (8.1) implicit and only quasilinear in τ^d).

In the notations of § 2, the dimensional Oldroyd coefficients λ_i and μ_i are scaled with $\mu_e a_o/\sigma$, while η is scaled with μ_e ; the non-dimensional argument of all of these functions is $\zeta = a_o(\mu_e \tau^d : E)^{1/2}/\sigma$. Applying the model (8.1) to the shear flow (2.1*a*) and using the definitions (2.2*a*) of the viscometric functions, three non-dimensional equations are obtained:

$$\mu_{sh} - 1 = \frac{\eta \left[1 + Ca_{sh}^2 \left(\lambda_1 \lambda_2 - \frac{1}{3} \mu_1 \mu_2 \right) \right]}{1 + Ca_{sh}^2 \left(\lambda_1^2 - \frac{1}{3} \mu_1^2 \right)},$$
(8.3*a*)

$$N_{1}^{sh} = \frac{\eta C a_{sh} \left[2(\lambda_{1} - \lambda_{2}) + \frac{2}{3} \mu_{1} C a_{sh}^{2} (\mu_{1} \lambda_{2} - \mu_{2} \lambda_{1}) \right]}{1 + C a_{sh}^{2} \left(\lambda_{1}^{2} - \frac{1}{2} \mu_{1}^{2} \right)},$$
(8.3b)

$$N_{2}^{sh} = \frac{\eta C a_{sh} \left[\mu_{1} - \mu_{2} + \lambda_{2} - \lambda_{1} + \frac{1}{3} C a_{sh}^{2} (3\lambda_{1} - \mu_{1}) (\mu_{1}\lambda_{2} - \mu_{2}\lambda_{1}) \right]}{1 + C a_{sh}^{2} \left(\lambda_{1}^{2} - \frac{1}{3} \mu_{1}^{2} \right)}.$$
 (8.3c)

Here, the capillary number $Ca_{sh} = Ca$ is defined by (2.1*a*), but we add the subscript *sh* to stress its connection to the shear flow. Without confusion, $\mu_{sh} - 1$, N_1^{sh} and N_2^{sh} are used here and below for the steady-state macroscopic quantities (instead of $\langle \mu_{sh} - 1 \rangle$, etc.). In principle, for any given *c* and the viscosity ratio λ , the left-hand sides of (8.3*a*)–(8.3*c*) are known functions of Ca_{sh} through shear-flow simulations (§ 7). The non-dimensional Oldroyd coefficients λ_i , μ_i and η on the right-hand sides also depend on Ca_{sh} through $\zeta = Ca_{sh}\sqrt{\mu_{sh} - 1}$.

Likewise, by applying (8.1) to PE (2.1b) and using the definitions (2.2b) of the extensiometric functions we obtain two more non-dimensional equations:

$$\mu_{pe} - 1 = \frac{\eta \left(1 - \frac{1}{3}\mu_{1}\mu_{2}Ca_{pe}^{2}\right)}{1 - \frac{1}{3}\mu_{1}^{2}Ca_{pe}^{2}},$$
(8.4*a*)

$$N_{pe} = \frac{\eta C a_{pe}(\mu_1 - \mu_2)}{1 - \frac{1}{3}\mu_1^2 C a_{pe}^2}.$$
(8.4b)

Here, the relevant capillary number Ca_{pe} is given by (2.1b). The left-hand sides of (8.4*a*,*b*) are provided versus Ca_{pe} by the PE simulations (§ 6), while the Oldroyd coefficients on the right-hand sides depend on Ca_{pe} through $\zeta = Ca_{pe}\sqrt{\mu_{pe}-1}$. The algebra (8.3)–(8.4) is the same as in the dilute case (Martin *et al.* 2014), except that we use $\mu_{sh} - 1$ and $\mu_{pe} - 1$ for the excess (instead of intrinsic) viscosities, and our

definition (2.2b) of the cross-difference N_{pe} has an additional factor of four in the denominator.

To be able to use (8.3a-c) and (8.4a,b) simultaneously, the argument ζ must be the same for both flows, which gives the mapping between Ca_{sh} and Ca_{pe} :

$$Ca_{sh}\sqrt{\mu_{sh}-1} = Ca_{pe}\sqrt{\mu_{pe}-1} = \zeta.$$
(8.5)

Our numerical procedure for determining the Oldroyd coefficients works as follows. For every $\zeta \in [\zeta_{min}, \zeta_{max}]$ (see below for the choices of ζ_{min} and ζ_{max}), the corresponding values of Ca_{sh} and Ca_{pe} are found from (8.5) using local quadratic interpolations/splines for $Ca_{sh}\sqrt{\mu_{sh}-1}$ and $Ca_{pe}\sqrt{\mu_{pe}-1}$ versus Ca_{sh} and Ca_{pe} respectively. After that, the system (8.3) and (8.4), with the left-hand sides interpolated from our data for the viscometric and extensiometric functions (§§ 6 and 7), may be regarded as five nonlinear equations for five unknowns, λ_1 , μ_1 , λ_2 , μ_2 and η . It was easy to use a large number ($\sim 10^3 - 10^4$) of steepest descent iterations to localize the solution, followed by just a few necessary Newton–Raphson iterations. In the order of descending ζ , the initial guess is taken from the preceding converged solution (and arbitrary at $\zeta = \zeta_{max}$). Importantly, whenever iterations of (8.3) and (8.4) converge, the solution was found to be independent of an initial guess. To capture the complex behaviour of the Oldroyd coefficients and successfully reach small ζ (when appropriate, see below), the step in ζ must be very small, notwithstanding our scarce data in figures 9–11, 13, 15 and 19.

The values of ζ are naturally limited by our tabulated ranges for $Ca_{sh}\sqrt{\mu_{sh}-1}$ and $Ca_{pe}\sqrt{\mu_{pe}-1}$. An additional limitation on ζ_{min} is that the corresponding Ca_{sh} must exceed the phase transition range (if any). Precise calculations of viscometric functions (especially N_1^{sh}) are extremely difficult, if not impossible, in that range (§ 7). Moreover, the exact shear-flow results would still exhibit the inherent non-smooth kinked behaviour due to phase transition. It may be questionable to combine such data with smooth extensiometric functions (not suffering from phase transition) for a general constitutive modelling.

As in the dilute case (Martin *et al.* 2014), for $\lambda = 0.25$, an additional unfavourable factor limiting ζ_{max} is the non-monotonic behaviour of $\zeta(Ca_{sh})$ due to steep shear thinning at large Ca_{sh} . Namely, $\zeta(Ca_{sh})$ reaches a maximum while the corresponding Ca_{pe} is still subcritical. We did not include $\lambda = 0.25$ in the constitutive modelling discussed below. For all other $\lambda \ge 1$, ζ_{max} corresponds to the critical breakup condition in PE. We note finally that, even in the absence of the phase transition limitation, it was not always possible to reach the ζ_{min} prescribed by our tables for $\zeta(Ca_{pe})$ and $\zeta(Ca_{sh})$. In some cases, we had to stop at a slightly larger (but still small) value of ζ , when either the Newton iterations diverged or the solution became unacceptable $(\lambda_1 < 0)$.

Figures 20–22 present the Oldroyd parameters for c = 0.45-0.55 and λ from 1 to 10. The η -parameter is almost constant. The non-smoothness and small-scale wiggles of the λ_i and μ_i curves are due to the scarcity of our tables for μ_{sh} , etc. and the chosen manner of interpolation from them. However, the predictions of the Oldroyd model (8.1) with these coefficients (discussed in the next section) were found to be very insensitive to such details, obviously because the whole combination of five Oldroyd parameters matters. The sharp rise of some coefficients at small *Ca* (outside the range shown in figure 20*a*,*b*) is not quite as surprising. Even in the absence of the phase transition limitation, a well-defined limit $\zeta \rightarrow 0$ with finite λ_i and μ_i is not expected, because a flowing concentrated emulsion is not a Rivlin–Ericksen fluid.

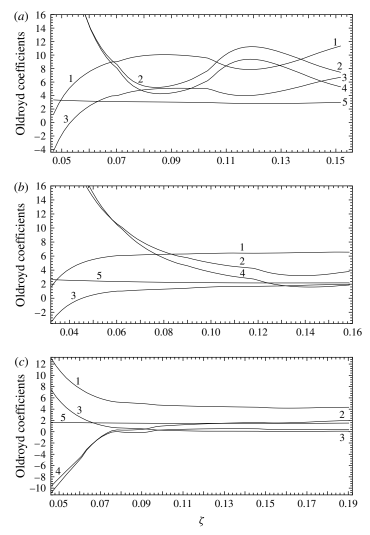


FIGURE 20. Generalized Oldroyd coefficients for concentrated emulsions with c = 0.45 and (a) $\lambda = 10$, (b) $\lambda = 3$ and (c) $\lambda = 1$. Lines 1–5 are for λ_1 , μ_1 , λ_2 , μ_2 and η respectively. Some lines are labelled twice for clarity.

9. Validation of constitutive modelling

For a given history of deformation of a material element, integration of the Oldroyd equation (8.1) requires an initial condition for τ^d at $t = t_0$. Obviously, knowledge of τ^d (or, equivalently, of a single microstructural tensor **a** in Hand's theory) does not completely characterize the instantaneous emulsion microstructure. For $\lambda = 1$, in particular, $\tau^d = 0$ can be obtained with an infinite number of statistically different initial configurations of non-deformed drops. In principle, more general constitutive equations could be attempted, with additional structural variables/tensors and possibly higher-order time derivatives of τ^d and/or **E**. Such models, though, do not necessarily represent a viable alternative to our constitutive modelling scheme, since these models would contain many more parameters/material functions problematic to determine from a finite set of flows. As usual, an acceptable behaviour of a multiparameter model in a broad range of conditions may be more difficult to achieve. It is not

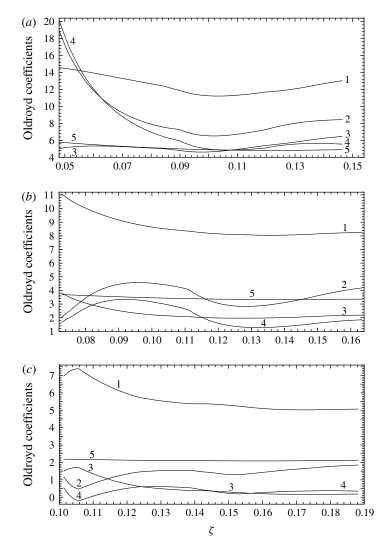


FIGURE 21. Generalized Oldroyd coefficients for c = 0.55 with (a) $\lambda = 10$, (b) $\lambda = 3$ and (c) $\lambda = 1$. Lines 1–5 are for λ_1 , μ_1 , λ_2 , μ_2 and η respectively. Some lines are labelled twice for clarity.

even clear to us how such more complex models would incorporate the details of the initial microstructure at $t = t_0$. Instead, the present work explores the idea of using only two steady canonic flows – PE and simple shear – for a robust and general constitutive modelling with only five material functions. An initial condition for τ^d is clearly artificial (as for all first-order constitutive equations of differential type, Astarita & Marucci 1974), and cannot replace the whole prehistory of the material element deformation prior to $t = t_0$. However, there are many potential applications of the generalized Oldroyd model (8.1) to non-Newtonian hydrodynamics, where the fading memory of the material makes the initial state insignificant. For example, in the problem of unbounded steady laminar flow of a non-Newtonian liquid past a body, there are two types of trajectories of a material element: (i) open trajectories arriving from infinity and (ii) closed trajectories in the recirculating zone(s). In both cases, an

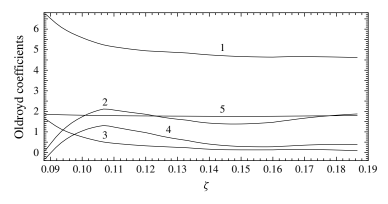


FIGURE 22. Generalized Oldroyd coefficients for c = 0.5 and $\lambda = 1$. Lines 1–5 are for λ_1 , μ_1 , λ_2 , μ_2 and η respectively.

initial condition for integrating (8.1) is imposed at $t = -\infty$ and does not affect the established regime at finite t (i.e. near the body) of only interest. In case (ii), the history of deformation of a material element and established rheological response are Lagrangian time-periodic. In case (i), they are aperiodic but still Lagrangian-unsteady. Obviously, dependence on the initial state would introduce much uncertainty, requiring details about the material preparation prior to simulation/experiments. Accordingly, we are only interested in validation tests for (8.1) where the initial state is sufficiently far in the past to not affect the results.

For dilute emulsions of single drops, the validation of the constitutive model (8.1), with the coefficients fitted to shear flow and PE, was performed (Martin *et al.* 2014) in a broad class of kinematic conditions, both Lagrangian-steady and unsteady. For concentrated emulsions, though, the only exact results that can be obtained for comparison with the model (8.1) are for steady mixed flows (2.1c), or their straightforward time-dependent generalizations (including particular cases of simple shear, $\chi = 0$, or PE, $\chi = 1$). These types of flow appear to be the only ones allowing for implementation of periodic boundaries (§ 2.1), and, hence, credible long-time simulations. Both steady and unsteady test flows are considered below.

9.1. General steady mixed flows

In contrast to shear flow, our mixed-flow simulations performed for $0.25 \le \lambda \le 3$ and $c \le 0.55$ have never exhibited phase transition, so simulation times $T \sim 100$ and the system size N = 100 suffice, combined with the moderate resolution $N_{\Delta} = 2160$ for $Ca \ge 0.0375$.

The model (8.1), with our variable Oldroyd coefficients (figures 20–22), is numerically integrated for the mixed-flow kinematics (2.1c) until the steady state. At the transient stage, the argument ζ is allowed to be somewhat outside the tabulated range [ζ_{min} , ζ_{max}] of the Oldroyd coefficients (with the extrapolation used), but we require it to be within [ζ_{min} , ζ_{max}] at the steady state. This requirement limits the range of simulated mixed-flow capillary numbers accordingly. The initial condition for (8.1) is either $\tau^d = 0$ or the converged solution at the nearest available *Ca*.

Strong mixed flows. For a strong mixed flow, $\chi = 0.25$, the model predictions (lines) are compared with exact simulation results (symbols) in figure 23(*a*,*b*). In addition to the excess mixed-flow viscosity $\mu^* - 1$, only the components v_{11} and v_{22} of the

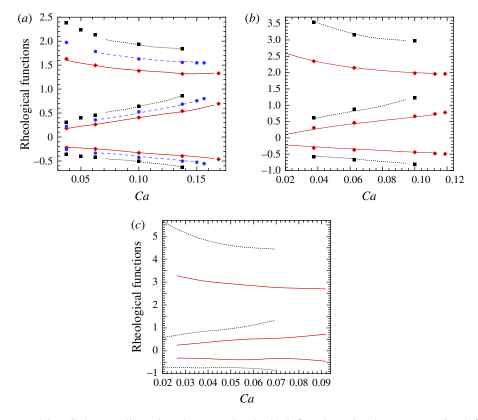


FIGURE 23. (Colour online) Steady-state rheological functions in the strong mixed flow with $\chi = 0.25$ and (a) $\lambda = 1$, (b) $\lambda = 3$, (c) $\lambda = 10$. Lines are from constitutive modelling, and symbols are exact BI simulation results. Diamonds and solid lines are for c = 0.45. Circles and long-dashed lines are for c = 0.5. Squares and short-dashed lines are for c = 0.55. In each graph, every three vertically aligned symbols of the same shape, and every three lines of the same pattern, represent $\mu^* - 1$, ν_{11} and ν_{22} from top to bottom.

non-dimensional stress (2.2c) are shown, since $v_{33} = -(v_{11} + v_{22})$ is not independent and is generally small. Apart from c = 0.55 (with the rightmost data points slightly subcritical), the other mixed-flow simulations (diamonds and circles) cover the whole range from small drop deformations to breakup. The corresponding model predictions (solid and long-dashed lines respectively) terminate at slightly smaller *Ca* due to the technical limitation of PE breakup on the argument ζ .

Another technical limitation – shear-flow phase transition – does not allow our model predictions for mixed flow to approach small *Ca*, namely when $\lambda = 1$, $c \ge 0.5$ and $\lambda = 3$, c = 0.55 (figure 23*a*,*b*). Despite these limitations, a very good agreement between the exact results and the model predictions suggests the success of the generalized Oldroyd model (8.1), with the variable coefficients fitted to simple shear and PE, for constitutive modelling of general flows. In the absence of the exact results at $\lambda = 10$, figure 23(*c*) provides our model predictions for mixed flow. In the conclusions, we discuss possible ways to remove/relax the current limitations on our constitutive modelling.

Weak mixed flows ($\chi < 0$). In the dilute limit, the above constitutive modelling scheme was also found to be very accurate for weak mixed flows, $\chi = -0.2$

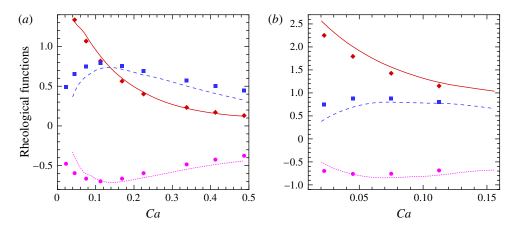


FIGURE 24. (Colour online) Steady-state rheological functions for a weak mixed flow $\chi = -0.25$, c = 0.45 at (a) $\lambda = 1$ and (b) $\lambda = 3$. Lines are from constitutive modelling and symbols are exact BI results: $\mu^* - 1$ (solid lines and diamonds), ν_{11} (long-dashed lines and squares), ν_{22} (short-dashed lines and circles).

(Martin *et al.* 2014). This is not quite the case for concentrated emulsions. In figure 24, for $\chi = -0.25$, c = 0.45 with (a) $\lambda = 1$ and (b) $\lambda = 3$, the agreement between the exact results and the model predictions is only approximate (except for the excess viscosity $\mu^* - 1$ in (a) with less discrepancy). For the exact results in the entire range of figure 24(*a*,*b*), the drop deformation remains small. Since a concentrated emulsion is not a Rivlin–Ericksen fluid and does not possess a well-defined limit of slow flows (with small drop deformations), it would be, indeed, difficult to expect accuracy of the model predictions for $\chi = -0.25$ based on the results for $\chi = 0$ (simple shear) and $\chi = 1$ (PE).

This moderate loss of accuracy for $\chi < 0$ is not perceived as a major drawback of our constitutive modelling scheme, since weak mixed flows are apparently less relevant to realistic non-Newtonian hydrodynamics in complex geometries. The trajectory of the displacement ξ between two close points of a continuum with velocity gradient ∇v is described by $d\xi/dt = \nabla v \cdot \xi$. Such trajectories are open (of hyperbolic shape) for strong mixed flows $\chi > 0$, and periodic (of elliptical shape) for $\chi < 0$. As a qualitative example of possible more general kinematics, consider a stationary flow past a macroscopic sphere at finite Reynolds number Re. Using, for simplicity, the Newtonian flow solution, the structure of the fluid trajectories past the sphere can be analysed up to Re = 60, at least qualitatively, by the two-term small-Re inner expansion for the stream function (Proudman & Pearson 1957; Van Dyke 1967). Figure 25(a) shows one closed trajectory in the eddy zone obtained in this manner for Re = 50. Along this pathline, ∇v is Lagrangian time-periodic. However, according to the Floquet theory for linear systems with periodic coefficients, periodicity of $\boldsymbol{\xi}(t)$ is not expected. Indeed, numerical integration of $d\boldsymbol{\xi}/dt = \nabla \boldsymbol{v}(t)\boldsymbol{\xi}$ along this pathline gives the open trajectory of $\xi(t)$ shown in figure 25(b). Therefore, even in the eddy zone, the history of deformation of material elements is qualitatively different from that for weak mixed flows.

9.2. Time-dependent PE flow

Our multidrop BI algorithm can be easily adapted to handle time-dependent simple shear, PE and mixed-flow simulations, if the time dependence is only in the scalar

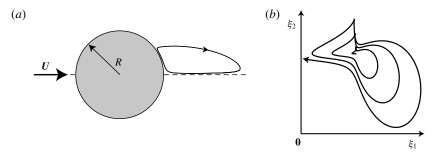


FIGURE 25. (a) Stationary Newtonian flow past a solid sphere at a finite Reynolds number $Re = UR/\nu = 50$; (b) a typical trajectory in the $\xi_3 = 0$ plane of the ξ -vector for a material element moving along the pathline shown in (a). The plane (ξ_1, ξ_2) is parallel to the meridian plane in (a), with ξ_1 along the ambient velocity U. The aspect ratio $\xi_1:\xi_2$ of the axes in (b) is 6.

prefactor $\varphi(t)$ before the constant velocity gradient tensor for either of the three flows. Returning to dimensional variables, consider an example of a time-dependent PE flow with the velocity gradient diag($\dot{\Gamma}, -\dot{\Gamma}, 0$) and the deformation rate $\dot{\Gamma}(t) = \langle \dot{\Gamma} \rangle \varphi(t)$, where $\langle \dot{\Gamma} \rangle > 0$ is constant. Assuming $\varphi(t) > 0$ for all t, a new monotonic variable \hat{t} is introduced through the integration of $\varphi(t) dt = d\hat{t}$. Then, the dynamics of the lattice vectors takes the usual form $d\boldsymbol{e}_i/d\hat{t} = diag(\langle \dot{\Gamma} \rangle, -\langle \dot{\Gamma} \rangle, 0)\boldsymbol{e}_i$ for steady PE, which enables the Kraynik–Reinelt scheme of periodic boundary implementation ($\S 2.1$); lattice restructuring is now made at all half-integer values of $\langle \dot{\Gamma} \rangle \hat{t} / \varepsilon_p$. Of course, the artificial variable \hat{t} is only used in the lattice dynamics and restructuring; in all the rest of the algorithm, the real time t is employed. A flow was considered with c = 0.45, $\lambda = 1$ and strong sevenfold non-harmonic oscillations in the deformation rate $\dot{\Gamma}(t)$, provided by $\varphi(t) = \text{const.}/(1 + 0.75 \sin 2\pi\omega t)$, with a numerical constant of order one (see below) to make the average of $\varphi(t)$ equal to one. Pure harmonic oscillations are avoided for the sake of generality. In order for this test to be relevant to real flow kinematics in complex geometries, the oscillation period $1/\omega$ should not be much smaller than $\langle \dot{\Gamma} \rangle^{-1}$, at least in the laminar regime. For example, the period of orbiting along the closed trajectory in figure 25 is $\sim R/U$, comparable with the inverse deformation rate; accordingly, we set $\omega = \langle \dot{\Gamma} \rangle$. With this choice, the above numerical constant in $\varphi(t)$ is found to be 0.6614. The capillary number $\overline{Ca} = 2\mu_e \langle \dot{\Gamma} \rangle a_e / \sigma$, based on the average deformation rate $\langle \dot{\Gamma} \rangle$, was set to 0.074.

Two non-dimensional rheological functions are defined for this flow through the drop-phase stress contribution τ^d and the average deformation rate,

$$\Delta \mu_{pe}^{\omega}(t) = \frac{\tau_{11}^d - \tau_{22}^d}{4\mu_e \langle \dot{\Gamma} \rangle}, \quad N_{pe}^{\omega}(t) = \frac{\tau_{11}^d + \tau_{22}^d - 2\tau_{33}^d}{4\mu_e \langle \dot{\Gamma} \rangle}.$$
 (9.1*a*,*b*)

These quantities quickly reach a periodic regime in our BI simulations, but small stochastic fluctuations around this regime remain, due to a finite number of drops; to reduce these additional fluctuations, a large system size N = 400 was used. Unlike for the steady-state results in §§ 5–7, time averaging would be irrelevant in this essentially unsteady flow test.

The initial evolution of $\Delta \mu_{pe}^{\omega}(t)$ in the N = 400 BI simulation to a periodic regime is shown in figure 26 by a solid line; as in the rest of the paper, the initial state was a well-mixed configuration of spherical drops. The results of the integration of the

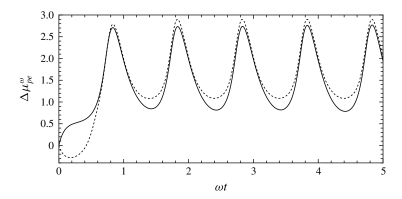


FIGURE 26. Evolution of the excess viscosity function to a periodic regime for time-dependent PE flow at c = 0.45 and $\lambda = 1$. The solid line is the exact result from the N = 400 BI simulation. The dashed line is from the generalized Oldroyd model (8.1).

generalized Oldroyd model (8.1) are also shown by a dashed line using our variable coefficients from figure 20(c) and the initial condition $\tau^d = 0$. The BI simulation and the model start to agree well before the periodic regime is reached, and the initial discrepancy at small times is not of much concern. At small times the initial drop configuration has an effect on simulations, and the BI results would first need to be ensemble-averaged over many initial conditions for a meaningful comparison. In view of the discussion at the beginning of this section, such averaging was not pursued since we are only interested in the long-time regime independent of the initial state; one initial drop configuration is sufficient to explore the long-time behaviour.

The long-time comparisons between the N = 400 BI simulation and various models are presented in figure 27, for both rheological functions (9.1). The two solid lines are from BI (top, $\Delta \mu_{pe}^{\omega}(t)$; bottom, $N_{pe}^{\omega}(t)$). The dashed lines are from the generalized Oldroyd equation (8.1) (top, $\Delta \mu_{pe}^{\omega}(t)$; bottom, $N_{pe}^{\omega}(t)$). The simplest alternative model is the quasisteady approximation using the steady-state PE results at the instantaneous deformation rate $\dot{\Gamma}(t) = \langle \dot{\Gamma} \rangle \varphi(t)$. Accordingly, the rheological functions (9.1) are approximated as $\langle \mu_{pe} - 1 \rangle \varphi(t)$ and $\langle N_{pe} \rangle \varphi(t)$, where the steady-state extensiometric quantities $\langle \mu_{pe} - 1 \rangle$ and $\langle N_{pe} \rangle$, as functions of the instantaneous capillary number (2.1b) $Ca = Ca\varphi(t)$, are interpolated from figure 9(c). This approximation is shown in figure 27 by two dotted lines (top, $\Delta \mu_{pe}^{\omega}(t)$; bottom, $N_{pe}^{\omega}(t)$). For some values of ωt , the quasisteady results could not be computed when Ca was outside the data range of figure 9(c). Regardless, the quasisteady approximation is seen to be very poor compared with the BI simulation, especially for $N_{pe}^{\omega}(t)$. The generalized Oldroyd model (8.1) behaves much better. For $\Delta \mu_{pe}^{\omega}(t)$, this model is practically indistinguishable from the BI results in broad ranges around maxima, and has very small phase errors near minima; only amplitude errors at minima are appreciable. For $N_{pe}^{\omega}(t)$, the Oldroyd model prediction is still acceptable, considering the small values of this cross-difference compared with $\Delta \mu_{pe}^{\omega}(t)$. Thus, remarkably, combining the steady-state results for PE with those for simple shear in our constitutive modelling scheme works to dramatically improve the predictions for time-dependent PE. This conclusion was confirmed by another test for a purely harmonic deformation rate $\dot{\Gamma}(t) = \langle \dot{\Gamma} \rangle (1 + 0.75 \sin 2\pi\omega t)$ at $\overline{Ca} = 0.1$.

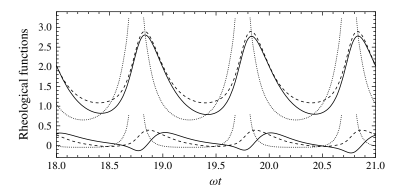


FIGURE 27. Rheological functions for time-dependent PE flow at c = 0.45 and $\lambda = 1$. The solid lines are exact results from the N = 400 BI simulation (top line, $\Delta \mu_{pe}^{\omega}$; bottom, N_{pe}^{ω}). The dashed lines are from the generalized Oldroyd model (8.1) (top, $\Delta \mu_{pe}^{\omega}$; bottom, N_{pe}^{ω}). The dotted lines are from the quasisteady PE flow approximation (top, $\Delta \mu_{pe}^{\omega}$; bottom, N_{pe}^{ω}).

10. Conclusions

We have studied the rheology of highly concentrated monodispersed emulsions of deformable drops by multidrop dynamical simulations for three types of steady macroscopic flows: (i) simple shear $v = (\dot{\gamma} x_2, 0, 0)$, (ii) planar extension (PE) $\boldsymbol{v} = (\dot{\Gamma}x_1, -\dot{\Gamma}x_2, 0)$ and (iii) mixed flow $\boldsymbol{v} = (\dot{\gamma}x_2, \dot{\gamma}\chi x_1, 0)$. The results for these flows are further used as a database to construct and validate the general constitutive model for concentrated emulsion flows with arbitrary kinematics. The algorithm is a further development of the original multidrop multipole-accelerated boundary-integral code for shear-flow simulations (Zinchenko & Davis 2002). Among the major improvements/additions, the present code incorporates periodic boundary conditions for PE and mixed flows (based on the reproducible lattice dynamics of Kraynik & Reinelt (1992) and matrix transformations of Hunt et al. (2010)), control of drop surface overlapping and much more robust (than in Zinchenko & Davis 2002) control of surface triangulations. These features allow for unlimited long-time simulations and accurate time averaging. The present version of the multipole acceleration is also more efficient and flexible, allowing for considerable skewage of the periodic cells and extreme drop elongation.

For shear flow, the steady-state viscometric functions (emulsion shear viscosity and normal stress differences) were studied for drop volume fractions c = 0.45-0.55, drop-to-medium viscosity ratios $\lambda = 0.25$, 1, 3 and 10, and different subcritical capillary numbers $Ca = \mu_e \dot{\gamma} a_0 / \sigma$, based on the ambient viscosity μ_e , non-deformed drop radius a_0 and surface tension σ . Compared with the initial effort (Zinchenko & Davis 2002), the present work includes two new more extreme viscosity ratios $\lambda = 0.25$ and 10. We have also extended the range of surface triangulations to typically $N_{\Delta} = 2000-4000$ boundary elements per drop (important for small *Ca*), and the system size to N = 200 (for $\lambda \neq 1$) and 400 (for $\lambda = 1$) drops in a periodic cell. Moreover, the simulated strains $\dot{\gamma}t$ were increased by 1.5–2 orders of magnitude (to hundreds for $\lambda \neq 1$ and thousands for $\lambda = 1$) to address the challenging phenomenon of dynamical phase transition (to a partially ordered state) in emulsion shear flows. The onset of this transition is quantified by a sharp increase in the relaxation time to the statistical equilibrium as *Ca* is reduced. In the simulated ranges $c \leq 0.55$ and $Ca \geq 0.02$, the $\lambda = 10$ emulsion does not show any phase transition. The $\lambda = 3$ emulsion flows in a disordered manner at c = 0.45, but starts to transition to ordered flow at the higher concentration c = 0.55 for *Ca* between 0.02 and 0.04. The $\lambda = 1$ emulsion can even transition at c = 0.45, but not until *Ca* approaches 0.025; at c = 0.55, the transition capillary number for such an emulsion is $Ca \approx 0.07$. Finally, the $\lambda = 0.25$ emulsion flows in a partially ordered manner in a wide range of $Ca \leq 0.18-0.22$ for c = 0.45-0.55. The shift of the phase transition to smaller capillary numbers, as λ is increased, is due to larger drop deformation and more efficient mixing. It is unexpected that our emulsions are found to transition even at c = 0.45 in shear flow; for comparison, molecular dynamics and Monte Carlo simulations for a classical liquid of hard spheres give higher transition values $c \ge 0.494$ (e.g. Hoover & Ree 1968; Hansen & McDonald 1976).

In the transition range, ergodic difficulties are severe, with noticeable effects of the system size N and the initial conditions on the steady-state viscometric functions, especially N_1 . Nevertheless, the accuracy of the calculations is sufficient to state the existence of the local minimum of the shear viscosity versus Ca; this non-smooth kinked behaviour is most pronounced for $\lambda = 1$ and c = 0.55. For the $\lambda = 0.25$ emulsion, the shear viscosity is almost insensitive to ergodic difficulties and is accurately calculated in a wide range, from large to small drop deformations. For such emulsions, though, the first normal stress difference can greatly exceed the drop-phase contribution to the shear stress, making them 'more elastic than viscous'.

The PE simulations behave very differently. The steady-state extensiometric functions (the effective viscosity μ_{pe} and cross-difference N_{pe}) were accurately computed in the same ranges of c = 0.45-0.55 and $\lambda = 0.25-10$ up to critical $Ca = 2\mu_e \dot{\Gamma} a_o / \sigma$ for drop breakup. High surface resolution $N_{\Delta} \sim 4000$ is paramount to accurately describe the range of $Ca \ll 1$. In the limit $Ca \rightarrow 0$, a concentrated emulsion is not a phenomenological Rivlin–Ericksen fluid, since close interaction of slightly deformed drops is a problem of singular perturbations. Our viscosity μ_{pe} versus Ca shows sharp tension thinning in this range, which then levels off for larger Ca with a shallow minimum. The cross-difference N_{pe} becomes significant for near-critical Ca. Most importantly, phase transition is never observed in our PE simulations. Relaxation to statistical equilibrium is always fast, with N = 100 sufficient to eliminate finite-size effects.

A natural question is whether the results obtained for steady simple shear and PE can help in general constitutive modelling for large-strain emulsion flows with arbitrary kinematics. Martin et al. (2014) offered a new general approach to constitutive modelling for non-Newtonian liquids, based on the traceless five-parameter Oldroyd model with variable coefficients. These five coefficients, depending on one instantaneous flow invariant ζ , must be found by simultaneously fitting the model to viscometric and extensiometric (in PE) functions at arbitrary flow intensities. Here, we applied this approach (already validated in Martin et al. 2014 for dilute emulsions of single drops) to concentrated emulsions. The Oldroyd model is written for the drop-phase contribution τ^d to the stress, and the chosen argument $\zeta = a_o(\mu_e \tau^d : E)^{1/2} / \sigma$ of the Oldroyd functions is related to the excess energy dissipation rate. The resulting constitutive model was validated for $\lambda = 1$ and 3 versus the exact simulation results in mixed flows at various flow intensities. For a strong flow $\chi = 0.25$, excellent agreement is observed. A modest loss of accuracy of the model for $\chi = -0.25$ is not a major drawback, since weak mixed flows ($\chi < 0$) are less relevant to non-Newtonian hydrodynamics in complex geometries, even in the eddy zones.

It is straightforward to extend the present multidrop BI algorithm with periodic boundaries to time-dependent large-strain flows of either simple shear, PE or mixed type, if this time dependence is only in the scalar prefactor before the constant velocity gradient. An example of PE flow with large periodic but non-harmonic oscillations in the deformation rate was considered for a concentrated emulsion c = 0.45, $\lambda = 1$ to further validate our approach to constitutive modelling. Of most interest was to find the long-time periodic stress behaviour independent of the initial conditions. Here, the steady-state PE results alone cannot adequately predict the rheological response. In contrast, combination of these results with those for simple shear in our constitutive modelling scheme works to dramatically improve the model predictions for this timedependent PE flow.

Still, our constitutive modelling for concentrated emulsions has limitations yet to be overcome. The range of capillary numbers below the phase transition point had to be excluded from the viscometric functions database. We found it virtually impossible to eliminate all the numerical uncertainties and accurately cover the whole range of Ca below the phase transition point for all three viscometric functions. Moreover, even in the absence of numerical errors, it may be questionable to combine such kinked data with smooth extensiometric functions (not suffering from phase transition) for a general constitutive modelling. This phase transition limitation restricts from below, sometimes very substantially, the range of flow intensities that can be simulated by our constitutive modelling. We anticipate the following approach to resolve this issue in future work. The fact that emulsions transition in shear flows but not in other flows (e.g. PE and mixed, as shown in this work) puts the relevance of simple shear, as a building block for general computational rheology of concentrated emulsions, into question. For future applications to non-Newtonian hydrodynamics in complex geometries, it would be much better to replace simple shear in the database by a mixed flow with a small $\chi > 0$ (say, 0.1–0.2). Such a modification would leave the basics of our constitutive modelling scheme intact, since mixed flow also has three rheological functions to match, and can still be accurately simulated for $\chi \sim 0.1$ –0.2, with not too much skewage of the periodic cells. The absence of phase transition in the database should largely eliminate the lower limitation on flow intensities in our constitutive modelling. Such a modification is also a way to overcome mapping difficulties (discussed in §8) and extend constitutive modelling to small viscosity ratios λ in future work. Of course, the modified constitutive modelling scheme would no longer be exact for pure simple shear, but this drawback is mostly irrelevant to non-Newtonian hydrodynamics in complex geometries. Interestingly, in the entirely different context of dilute suspensions of Brownian particles, Hinch & Leal (1976) also found simple shear to fall out from their modelling scheme, due to the lack of flow-induced particle orientation.

The upper limitation $\zeta \leq \zeta_{max}$ on our constitutive modelling is due to early drop breakup in steady PE. In a general flow with $\zeta > \zeta_{max}$, Lagrangian-steady or unsteady, drops will or will not break. Unfortunately, it is unclear, even in principle, how to extend this constitutive modelling to $\zeta > \zeta_{max}$ or even formulate the concept of the rheology of concentrated emulsions experiencing drop breakup. In some cases, it may work to simply extrapolate our computed Oldroyd coefficients beyond ζ_{max} . For example, at c = 0.45 and $\lambda = 1$, our Oldroyd coefficients (figure 20c) show large variations at small ζ (this variability is essential for the success of constitutive modelling in figure 23*a*). At larger ζ , though, these coefficients almost do not change, and it may be tempting to continue them as constants beyond ζ_{max} as an approximate constitutive modelling in the breakup range, at least for moderately supercritical conditions. We hope to probe this approach in future work. The concepts and techniques of the present work can be extended to the rheology of concentrated emulsions with an insoluble surfactant, the subject of our current research. Finally, our general constitutive modelling scheme should be applicable to other non-Newtonian liquids (suspensions, polymeric solutions, etc.).

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