Wavelets and fast summations for particle simulations of gravitational flows of miscible drops

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Abstract

For sedimentation of miscible drops through quiescent liquid of the same viscosity, a recent paper [J. Fluid. Mech. 447 (2001) 299] has shown the effectiveness of computer simulations based upon a swarm of point forces in tracking coalescence, mixing and rupture. Robustness of the approach was offset by the slow O(N 2 p) summations needed to calculate the mutual viscous interactions between all N p particles.

Motivated by applications of wavelets to linear operators, this paper develops a conceptually simple scheme for dramatically accelerating the simulations. After lumping the particles together into N local clusters (three-dimensional “pixels”), a Haar discrete wavelet transform (DWT) is used to “compress” the “bitmapped” six-dimensional image of pixel–pixel hydrodynamic interactions. Depending upon the criterion of accuracy, the numerically observed scaling of the operation count seems to be either O(N) or O[(N(logN)α] . The DWT also works without modification for hydrodynamic wall effects, where the kernel is not purely translational and therefore fast convolutions (FFT) cannot be used.

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1. Introduction: a swarm of point forces

This article describes the use of wavelets to substantially accelerate particle simulations of low-Reynolds-number drop flows. As a physical point of reference, within the realm of common experience and having also geophysical analogs in mantle plumes (Manga, 1997; Manga, Stone, & O’Connell, 1993) consider the rising (or sedimentation) of a miscible drop of ink (or suspension of glass beads) through a quiescent bath of solvent (Fig. 1). The resulting nonlinear shape instability has been described and studied extensively (Adachi, Kiriyama, & Koshioka, 1978; Arecchi, Buah-Bassuah, Francini, Pérez-García, & Quercioli, 1989; Joseph & Renardy, 1993; Kojima, Hinch, & Acrivos, 1984; Machu, Meile, Nitsche, & Schaflinger, 2001; Schaflinger & Machu, 1999; Thomson & Newall, 1885) and includes

(i) entrainment of solvent and formation of a tail; (ii) intermediate mushroom and ring shapes; and (iii) an eventual cascade of successive breakups. All of these phenomena are observed in the theoretically simplest case of inertialess flow with equal viscosity of the ink and solvent.

The work of Machu et al. (2001) formalized an analogy between a miscible drop of homogeneous liquid and a swarm of point forces (cf. Adachi et al., 1978; Kojima et al., 1984; Nitsche & Batchelor, 1997), and solved numerically a (nonlinear) dynamical system for the Stokeslet positions:

\[
\frac{d\mathbf{r}_i}{dt} = \sum_j G(r_i - r_j) \delta V
\]

with \( G \) the (dimensionless) disturbance field produced by a unit point force acting in the direction of gravity,

\[
G(r) = (|r|^{-1} I + |r|^{-3} r r) \cdot e_g
\]

and \( \delta V \) the equivalent volume of ink associated with one particle. The resultant error in the local liquid velocity scales with the number \( N_p \) of particles like \( N_p^{-1/2} \log N_p \).
The boundary integral method (Cristini, Bławzdziewicz, & Loewenberg, 1998, 2001; Davis, 1999; Koh & Leal, 1999; Pouzikids, 1990, 1992; Rallison & Activos, 1978; Zinchenko, Rother, & Davis, 1999) is very widely used in low-Reynolds-number drop dynamics because of the efficient premise of two-dimensional discretization. But when interfaces wind around, rupture or coalesce, even sophisticated adaptive surface gridding algorithms (Cristini et al., 1998, 2001) can get bogged down. For simulating the bell-mushroom-ring-breakup sequence described above, a volumetric method such as Machu et al. (2001) seems to be necessary. The purpose of this article is to develop a conceptually and practically simple scheme for speeding up the $O(N^2)$ summations of particle interactions. Within the equal-viscosity assumption, particle swarms then emerge as an attractive approach to low-Reynolds-number drop dynamics.

2. Lumping particles together

A common starting point for accelerating particle simulations in a wide variety of physical contexts is to lump nearby particles together within cells or pixels (Hockney & Eastwood, 1988); see Fig. 2. At the crudest, force-monopole level of approximation, one counts the total number $\phi_{lmn}$ of gravitational point forces in each pixel $(l,m,n)$, and then replaces the superposition of all of their locally distributed hydrodynamic disturbance fields with a single, central point force based upon the summed weight. From a six-dimensional table $F_{ijklmn}$ of all pixel-pixel interactions $(i,j,k) \leftrightarrow (l,m,n)$, the flow velocity $v_{ijklmn}$ at the center of each pixel $(i,j,k)$ is obtained from the summation

$$v_{ijklmn} = \sum_{lmn} F_{ijklmn} \phi_{lmn} \delta V$$

where the self-interaction $F_{ijklmn}$ is taken as the Stokes settling velocity of an equivalent sphere (radius roughly optimized to 0.295 times the pixel dimension in numerical trials), as an approximation of the corresponding weakly singular integral. [See comments on “desingularization” in Philips (2003, pp. 84–85).] Individual particle velocities in the swarm can then be interpolated between values in the pixel table $v_{ijklmn}$. Eq. (3) could be renumbered (somewhat inconveniently) to look like a matrix multiplied by a vector.

Refinement of this scheme, by evaluating near particle interactions from neighboring pixels directly (Greengard & Rokhlin, 1987; Hockney & Eastwood, 1988) is not necessary because of the fundamentally global nature of gravitational Stokes flow. Integrated over volume, the $1/r$ decay of the Stokeslet means that local structure of linear dimension $\varepsilon$ contributes only at order $\varepsilon^2$ to the local flow field that moves and deforms it: the main contribution is actually from everywhere else in the drop. Thus, local pixel refinement is actually superfluous, and coarse pixels can resolve comparatively fine structures. In particular, accurate resolution of the fuzzy, miscible interface is not necessary. For the drop shown in Fig. 2, the intrinsic quadrature errors for the $8 \times 8 \times 8$ and $16 \times 16 \times 16$ pixel grids are surprisingly low: sampled at the centers of all pixels, rough bounds on the maximum deviation in velocity are 0.59 and 0.24%, respectively, scaled to the maximum velocity. Fig. 3 shows that, until this intrinsic error is reached (horizontal asymptote),
Fig. 2. Lumping of a spherical swarm of 18,850 particles into pixels, illustrated in a sectional view (1 pixel deep) for two levels of discretization. (a) $8 \times 8 \times 8$ pixel grid; (b) $16 \times 16 \times 16$ pixel grid.

Fig. 3. Difference $\varepsilon_{\text{rms}}$ between the particle sum (1) and the pixel sum (3) as a function of number $n = N_p/N$ of particles per pixel, at two fixed pixel discretizations. The two velocities are computed at the centers of all pixels, and $\varepsilon_{\text{rms}}$ is the root-mean-square discrepancy, normalized as a percentage of the maximum velocity norm.

3. Wavelets

Leaving aside higher multipoles (which can be incorporated in a straightforward if tedious fashion) the main premise of this article is to exploit a discrete wavelet transform (DWT), which (i) applies, via simple matrix operations, directly to the fine pixel grid; and (ii) automatically compresses the summations without any explicit tree-code data structuring. This approach is suggested generally by applications of wavelets (and multiresolution algorithms) to linear operators (Beylkin, Coifman, & Rokhlin, 1991; Harten & Yud-Shalom, 1994) and more specifically, by the acceleration of matrix multiplications with the DWT (Press, Teukolsky, Vetterling, & Flannery, 1992).

The DWT is essentially a hierarchy of “filtering” operations applied to successively coarser samplings of a pixel image $\phi_{ijk}$ or $F_{lmn}$. Each index the filter operates by forming linear combinations of nearby elements in the ordered list given to it, which serve to extract equal amounts of “smooth” versus “detail” information. At each stage the filter is applied only to the smooth half of the output from the previous stage, in what is called a “pyramidal algorithm” (Press et al., 1992). There are two main points. First, all levels in the DWT pyramid can be encapsulated in the action of a single matrix $W$, which can be stored in packed form to exploit its sparseness (predominance of zero elements); see Fig. 4. Second, the matrix $W$ has the property that most elements of the resultant image in the wavelet basis, $\hat{\phi}_{IJK} = \sum_{ijk} W_{II} W_{JJ} W_{KK} \phi_{ijk}$ (4), are so small that they can be discarded without seriously degrading the image upon reconstituting it with the inverse DWT. Wavelets are particularly good for compressing localized detail, where spectral decompositions become uniformly fuzzy at the same level of truncation (Press et al., 1992). Our illustrative calculations are based on Haar wavelets (Kaiser, 1994; Walker, 1999) which are the simplest.

In some sense, the coarser averages at higher levels of the DWT pyramid are analogs of larger clusters in FMM. In other words, the pyramidal algorithm represents a kind of automatic, generic tree-code, independent of the specific operator (physics) involved. Reduction of the whole pyramid to a single matrix operation makes the DWT transparent in its
One can map the six-dimensional array of pixel–pixel interactions \( (i, j, k) \) to a highly sparse array for faster summations:

\[
\tilde{F}_{ijk} = \frac{1}{|x_i - x_j|}
\]

In the absence of wall effects [translational kernel \( G(r - q) \)], the six-dimensional sum can be reduced to triple sum:

\[
\tilde{G}_{ijk} = \sum_{a,b,c} A_{abc} \tilde{A}_{ijk} \tilde{A}_{abc} \tilde{G}_{abc}
\]

written in terms of an intermediate three-dimensional array \( A \) that is constructed from the wavelet matrix \( W \):

\[
A_{abc} = \sum_{ij} W_{ai} W_{bj} W_{ck}
\]

Depending upon both the kernel and the wavelet basis, one may or may not be able to exploit a theoretically justified pattern of truncation in the wavelet-space array \( \tilde{F}_{ijk} \) to discard most of its elements a priori, without having to first calculate them and then check their magnitude (Beylkin et al., 1991). This aspect will be investigated in a future paper. The important point is that the truncated table of wavelet coefficients \( \tilde{F}_{ijk} \) needs to be calculated only once for all nonzero entries in the array. (a) 8 pixels; (b) 16 pixels; (c) 32 pixels; (d) 64 pixels.

### 4. Scaling of the operation count

Our drop simulations were limited to the 16 × 16 × 16 pixel array illustrated in Fig. 2b. As will be seen below, this resolution is not fine enough for the asymptotic scaling of the operation count to emerge clearly. Thus, we shall extrapolate the scaling with reference to one- and two-dimensional analogs.

Focusing on the \( 1/r \) radial decay of the Stokeslet, which is here the determining feature, one recognizes a dimensional progression of analogous kernels, of which the 1D version is of the Calderon-Zygmund type (Beylkin et al., 1991):

\[
K^{(1)}(x, \xi) = \frac{1}{|x - \xi|} \tag{10}
\]

\[
K^{(2)}(x, y, \xi, \eta) = \frac{1}{\sqrt{(x - \xi)^2 + (y - \eta)^2}} \tag{11}
\]

\[
K^{(3)}(x, y, z, \xi, \eta, \zeta) = \frac{1}{\sqrt{(x - \xi)^2 + (y - \eta)^2 + (z - \zeta)^2}} \tag{12}
\]

The two and three-dimensional cases are similar; the major qualitative difference is between one dimension and two, where a singular line (\( \xi = \xi \)) gets replaced by a singular point \([\xi, \eta] = (x, y)\). However the pixels may be renumbered into linear order, this feature destroys the simple diagonal dominance of the corresponding matrix discretization of \( K^{(2)} \), which is now an array of diagonally dominant sub-blocks.

As an example of the scaling of the operation count for wavelet compression applied to the kernels \( K^{(1)}(x, \xi) \) and \( K^{(2)}(x, y, \xi, \eta) \), respectively, analogous mound-shaped signal functions in one and two dimensions,

\[
\phi^{(1)}(\xi) = \frac{1}{1 - 3\xi^2 + 2|\xi|^3}, \quad |\xi| \leq 1
\]

\[
\phi^{(2)}(\xi, \eta) = \phi^{(1)}\left(\sqrt{\xi^2 + \eta^2}\right) \tag{13}
\]

are “run through” these operators to produce the corresponding output signals \( \phi^{(1)}(x) \) and \( \phi^{(2)}(x, y) \):

\[
\phi^{(1)}(x) = \int_{-1}^{1} K^{(1)}(x, \xi) \phi^{(1)}(\xi) d\xi \tag{14}
\]

\[
\phi^{(2)}(x, y) = \int_{-1}^{1} \int_{-1}^{1} K^{(2)}(x, y; \xi, \eta) \phi^{(2)}(\xi, \eta) d\xi d\eta \tag{15}
\]
Residing conceptually between $K^{(1)}$ and $K^{(2)}$ is a bilinear functional:

$$\int \int K(x, \xi, \eta) f(\xi) g(\eta) \, d\xi \, d\eta.$$  

For $K^{(1)}$ and $K^{(2)}$ theorems on $O(N)$ summations are available (Beylkin et al., 1991; Harten & Yadal-Shalom, 1994).

For a given discretization of the $N$ domain (represented by the total number $N$ of pixels, irrespective of dimension), various truncation levels were applied to the DWT of the corresponding kernel array $K^{(1)}_{ij}$ or $K^{(2)}_{il}$, and the maximum error $\varepsilon_{\text{max}}$ of the discrete output $\psi^{(1)}_d$ or $\psi^{(2)}_d$ was recorded as a function of the total number $M$ of retained wavelet coefficients, which directly reflects the operation count in the summations. Plotted logarithmically, these $\varepsilon_{\text{max}}$ versus $M$ data were smoothed by least-squares linear regression (Fig. 5), and the linear fit was then used to interpolate the number of wavelet coefficients needed to achieve a given accuracy. Results for various resolutions in both one and two dimensions are combined in Fig. 6, which shows the scaled operation count $M/N$ or $M/[N \log N^{1/2}]$ as a function of the number $N$ of pixels. Holding the error fixed, the scaling in both one and two dimensions seems to be $O(N)$ [Fig. 6(a)]; it is certainly bounded by $N \log N^{1/2}$. Fixed error is not entirely realistic as a criterion, because the whole purpose of higher resolution is to diminish the discretization error in the integrals (14) and (15). A safe overestimate comes from taking $\varepsilon_{\text{max}} \propto N^{-1/2}$. This assumption (i) is more stringent than the $O(N^{-1/2} \log N)$ error scaling of the particular discretization (1) of a liquid drop; and (ii) assumes that $N$ is increased proportionally to $N_p$, which is unnecessary according to Fig. 3. From Fig. 6(b) we see that $M = O(N \log N^{1/2})$. The same asymptotic scalings would be expected to apply in 3D simulations as well.

5. Simulations of drop dynamics

Since the 16 pixel resolution that was computationally tractable for 3D problems lies well below the asymptotic regime of the operation count (Fig. 6), we shall here be content with the reasonable dimensional extrapolation of the previous section, and simply exhibit typical acceleration factors with reference to two test problems. Shown in Fig. 7 are results of computer simulations whereby the configurational evolution of two buoyant, miscible drops is modeled with swarms of particles. The smaller, leading drop flattens out

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**Fig. 5.** Error $\varepsilon$ in the output signals $\psi^{(1)}_d$ and $\psi^{(2)}_d$ as a function of the number $M$ of retained wavelet coefficients in the truncated kernel arrays $K^{(1)}_{ij}$ and $K^{(2)}_{il}$, at fixed, 16 pixel resolution. (a) 1D; (b) 2D.

**Fig. 6.** Scaling of the operation count $M$ with the number of pixels $N$ for wavelet compression of the 1D and 2D kernels defined by Eqs. (10) and (11). Numbers by the points indicate the linear pixel resolution. (a) Fixed accuracy $\varepsilon_{\text{max}} = 1\%$; (b) increasing accuracy $\varepsilon_{\text{max}} \propto N^{-1/2}$. 

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into a thin lens and coats the larger trailing drop, which becomes elongated (Manga & Stone, 1993). The wavelet-based calculations for the dots on the right-hand side of each graph were based upon a swarm of 18,850 particles lumped within a $16 \times 16 \times 16$ pixel grid, and used 3.22% of the wavelet coefficients $F_{i,j,k}$. Three thousand time steps took less than 3 min, and these could be used for any simulation. Note particularly that the thin lens shape and narrow gap between the two drops are accurately resolved by pixels which are of similar dimensions, owing to the global nature of the hydrodynamic interactions; cf. Eqs. (1) and (2).

Figure 8 shows the flattening and distention into an oblate shape of a miscible drop sedimentating toward a horizontal, no-slip surface. The same description as for Fig. 7, except for the particle interactions. Left (sparser) side: direct summation (1) for 16,755 particles. Right (denser) side: wavelet method for 16,755 particles. The original list of about 50 million wavelet coefficients was truncated to 3.94%. (a) Initial configuration, for which the drop-wall gap is 1.75 drop radii; (b) $t = 30$ (6000 time steps).

6. Concluding remarks

Widely used in physics (Carrier et al., 1988; Greengard, 1994; Greengard & Rokhlin, 1987; Hockney & Eastwood, 1988; Rokhlin, 1983) the simplicity and robustness of particle methods makes them useful for simulating low-Reynolds-number flows of drops at unit viscosity ratio that involve complex interfacial evolutions such as coalescence and rupture (Machu et al., 2001; Nitsche & Schaflinger, 2001). For the case of infinite Bond number, this article has illustrated how wavelets remove the bottleneck in computational speed, with only inconsequential losses of accuracy.

The pyramical structure of the DWT, which is vaguely analogous to the tree-coding of fast multipole methods, is encapsulated in one simple matrix multiplication, and may be useful for other particle interaction laws as well: molecular dynamics (Greenspan, 1990; Moseler & Landman, 2000; Murad & Law, 1999), dissipative particle dynamics (Clark, Lal, Ruddock, & Warren, 2000; Hoogerbrugge & Koelman, 1992; Jones, Lal, Ruddock, & Spenley, 1999), and smoothed particle hydrodynamics (Monaghan, 1992; Nitsche & Zhang, 2002).
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