MONTE CARLO SIMULATION OF COAGULATION 
IN DISCRETE PARTICLE SIZE DISTRIBUTIONS 
I. BROWNIAN MOTION AND FLUID SHEARING 

H. J. Pearson, I. A. Valioulis and E. J. List 

W. M. Keck Laboratory of Hydraulics and Water Resources 
Division of Engineering and Applied Science 
California Institute of Technology 
Pasadena, California 91125 
U.S.A. 
August 1983
ABSTRACT

A method for the Monte Carlo simulation, by digital computer, of the evolution of a colliding and coagulating population of suspended particles is described. Collision mechanisms studied both separately and in combination are: Brownian motion of the particles, and laminar and isotropic turbulent shearing motions of the suspending fluid. Steady state distributions are obtained by adding unit size particles at a constant rate and removing all particles once they reach a pre-set maximum volume. The resulting size distributions are found to agree with those obtained by dimensional analysis (Hunt, 1982).
1. INTRODUCTION

In many fluid systems with a continuous size distribution of suspended particles the primary mechanism for the production of larger particles from smaller particles, over much of the size range, is coagulation, the process of collision and coalescence of particles. These coagulating particles can be solid or liquid with the suspending medium gaseous or liquid, for example: atmospheric aerosols, cloud water droplets, colloidal suspensions in water or emulsions of one liquid or another. The computations described in this paper are primarily concerned with suspensions of solid particles in water but the techniques used have general applications.

In describing the dynamics of continuous size distributions it is convenient to introduce the particle size distribution, \( n(v) \), defined by

\[
dN = n(v)dv
\]

so that \( dN \) is the number of particles per fluid volume whose sizes (volumes) lie in the range \( v \) to \( v + dv \). The collision rate, per unit volume of fluid, of particles of volumes \( v_i \) and \( v_j \) is given by the product of their respective concentrations and a collision function, \( \beta(v_i, v_j) \), representing the geometry and dynamics of the collision mechanism, so that

\[
\text{collision rate} = \beta(v_i, v_j) n(v_i) n(v_j) dv_i dv_j
\]

Then the change with time of the particle size distribution is given by the general dynamic equation (GDE)

\[
\frac{\partial n(v)}{\partial t} = I(v) + \frac{1}{2} \int_0^v \int_0^x \beta(v', v-v') n(v') n(v-v') dv' dv' - \int_0^\infty \int_0^\infty \beta(v, v') n(v) n(v') dv' \, Sv \quad (1)
\]
Here \( I(v) \) is a source of particles (through condensation, for example) and \( S(v) \frac{\partial n}{\partial z} \) is a particle sink resulting from particles sedimenting in the \( z \) direction at their Stokes' settling velocity, \( S(v) \). If we restrict attention to size ranges where the source term is negligible, and to homogeneous situations (so that spatial derivatives may be neglected) then (1) reduces to the coagulation equation

\[
\frac{\partial n(v)}{\partial t} = \frac{1}{2} \int_0^\infty \beta(v', v-v') n(v') n(v-v') dv' - \int_0^\infty \beta(v, v') n(v) n(v') dv'
\]  

(2)

The two terms on the right-hand side of (2) represent, respectively, the rate of gain of particles of volume \( v \) by coagulation of pairs of smaller particles, conserving volume, and the loss of particles, \( v \), due to their coagulation with particles of all sizes.

In writing equation (2) it is assumed that the suspension is completely mixed and that correlations between particles induced by the coagulation process can be ignored. For example, as particles of a given size coalesce, a local reduction in their number occurs, so fewer particles of this size remain for further coalescence. Such spatial inhomogeneities, which can be expressed as conditional probabilities for the collision of two particles (Gillespie, 1972) are ignored in (2) whose solution, thus, may not include all possible histories of particle growth.

A variety of techniques have been used to investigate equations (1) and (2) and an extensive literature has resulted (see Pruppacher and Klett, 1978 for a recent account). In this study we present a Monte Carlo method for the direct simulation of the processes of collision and coalescence of particles. Unlike the direct numerical solutions of equation (2), such as developed by Gelbard, Tambour and Seinfeld (1980),
the technique avoids approximations involved in writing (2). It also
does not require analytic forms of the collision functions \( \beta \) (except
for when Brownian motion is the collision mechanism, see Appendix A).
Direct comparison of the results of the Monte Carlo simulation method
and numerical integrations of (2) should therefore prove useful.
Furthermore, in addition to predicting the average particle size spectrum
the simulation method provides information on higher moments of properties
of the suspension, thereby enabling estimates of the uncertainty of
the size spectrum.

In order to confirm the appropriateness of the method and verify
the techniques used, it was considered sensible first to ignore particle
interactions such as those due to hydrodynamic and interfacial forces.
Clearly, this is unrealistic and only defensible as a first step toward
a more sophisticated model. However, there seemed little point in
proceeding to this enhanced model if the simulation method would not
reproduce the results of Hunt and Friedlander which were derived in the
absence of interparticle forces. A more complex model including also
the simulation of differential settling induced collisions is the
subject of a subsequent paper.

For particles to coagulate two processes are required: (a) a
mechanism to develop relative motion of the particles through the fluid
which will bring them into close proximity, and (b) short-range inter-
facial forces acting between the particles to bring about their
coalescence. Relative motion of particles in a fluid can be due to one
or a combination of the following:
1. Brownian or thermal motion.
2. Laminar or turbulent fluid shear or straining.
3. Particle inertia in turbulent flows.
4. Differential sedimentation of different size particles.

If hydrodynamic and interfacial forces between particles are ignored, relatively simple analytic estimates for $\beta$ are available for each of these collision mechanisms acting independently and these are summarized in Table 1. The table also includes the dimensional parameters that characterize the mechanisms and determine, in any given situation, the characteristic size of particles that they affect.

Note that all the collision functions depend on properties of the suspending fluid, the structure of its velocity field, and the size of the particles. However, only the functions for the final two collision mechanisms depend on a physical property of the particles: the difference between their density and that of the fluid. If the particle density excess ratio $(\rho_p - \rho_f)/\rho_f$ is small then sedimentation and inertia will only be important for larger particles. In a turbulent flow sedimentation will dominate inertial effects unless the characteristic acceleration $(\epsilon^3/\nu)^{1/2}$ is comparable with $g$, the gravitational acceleration. In this paper will be concerned only with the first two collision mechanisms.

For a coagulating system more than one collision mechanism can be important for a given size range of particles. However, if there is a particle size subrange in which the coagulation is dominated by only one collision mechanism, and this subrange is in a state of dynamic equilibrium, then the theory of Friedlander (1960a,b) and Hunt (1982) predicts the local size distribution given a constant flux of mass
through the particle size distribution. Aside from ignoring interparticle forces, the theory depends on two basic hypotheses: an equilibrium size distribution being established and non-interference of particles of a size characteristic of one collision mechanism with those of another collision mechanism.

Hunt's (1982) experimental results generally support the predictions of the theory for Brownian motion and laminar shear but are limited by uncertainty over the effects of the unsteadiness in the experiments due to particle sedimentation and loss from the system. In the present work these limitations are overcome by performing a computer "experiment" in which particle collisions are directly simulated by Monte Carlo techniques. The size evolution of a population of particles is followed. This allows the effects of each collision mechanism to be evaluated independently, and, by combining mechanisms, the hypothesis of non-interference of characteristic particle sizes to be tested. Collision rates as well as the approach to and the final form of an equilibrium size distribution are studied. The method could also be used to study the "aging" of an initially fixed number of particles as they collide and grow.

Monte Carlo simulations have been used by Nowakowski and Sitarski (1981) to model the collision function for Brownian coagulation of aerosol particles and by Husar (1971) and Gartrell and Friedlander (1975) to find solutions to the coagulation equation (2). In this paper we first briefly describe Hunt's theory and experimental results. Subsequent sections describe in detail the simulation techniques used to model Brownian, laminar shear and turbulent shear induced coagulation and the results obtained. The results are then compared with previous experiments
2. HUNT'S WORK

Friedlander (1960a,b) explained observed regularities in the size distributions of atmospheric aerosols by assuming that a state of dynamic equilibrium existed between production, coagulation and loss through sedimentation of particles. He then employed methods analogous to those developed by Kolmogorov for the analysis of turbulence spectra. If it is assumed that the size distribution in some subrange depends only on the particle volume, \( v \), the constant flux of particle volume through the size distribution, \( E \), and a dimensional parameter, \( C \), characterizing the sole dominant coagulation mechanism (see Table 1) so that

\[
n(v) = n(v, E, C),
\]

then the form of \( n(v) \) can be determined by dimensional analysis alone. This is analogous to postulating an inertial subrange of scales in which the turbulent energy spectrum is determined solely by the wavenumber and the flux of energy through the subrange (equal to the rate of energy dissipation by viscous stresses at the smallest scales).

(see, for example Monin and Yaglom, 1975, Ch. 21).

Hunt (1982) extended these ideas to hydrosols and compared the predictions of his theory with both laboratory and field measurements. In particular, he performed experiments on Brownian and laminar shear induced coagulation. His theory predicts the following size distributions for regions dominated by Brownian, shear and differential sedimentation coagulation.
Brownian

\[ n(v) = a_b (E/K_b)^{1/2} v^{-3/2}, \]  

(3)

Shear

\[ n(v) = a_{sh} (E/G)^{1/2} v^{-2}, \]  

(4)

Differential Sedimentation

\[ n(v) = a_{ds} (E/K_{ds})^{1/2} v^{-13/6}, \]  

(5)

He shows (Hunt 1982, Figure 1) that it is plausible, for a typical coagulating hydrosol, that these three mechanisms could dominate in regions of successively increasing particle size.

Hunt's measurements indicated that his system was in a quasi-dynamic equilibrium where size distributions taken at progressively later times were similar in shape but decreasing in magnitude. This unsteadiness was due to the overall particle concentration decreasing as a result of the larger aggregates settling to the bottom. Hunt measured the varying total suspended volume by light absorbance and used the computed rate of volume loss as an estimate for \( E \). He explains why this will be an overestimate for the quantity (see Hunt, 1982, for details), but it is still a useful approximation. The measured value of \( E^{1/2} \) can then be used to normalize size distributions (c.f. equations (3) - (5)) partially correcting for the effects of unsteadiness.

Hunt successfully collapsed much of his data at various times and for different experiments at different shear rates by normalizing the size distributions not just with \( E^{1/2} \), but with the ratio \((E/G)^{1/2}\) and
non-dimensionalizing the particle volume with the characteristic volume at which particles have both Brownian collisions and shear induced collisions at the same rate. This characteristic volume, found by putting $r_i = r_j$ in the expressions for the relevant collision rates in Table 1, is seen to be $v = \frac{\pi K_b}{3G}$, proportional to the ratio of the Brownian and shear parameters.

For some of the particle types tested the normalized volume distributions expressed as functions of non-dimensional size provide support for the relations (3) and (4) (see in particular Hunt 1982, Figure 5). However, as we have already noted, there are some reservations about the experiments, complicated as they are by instrumental difficulties and uncertainties about the effects of unsteadiness. Also, no single experiment exhibits a size distribution having regions with the equilibrium power laws corresponding to both Brownian and shear dominated mechanisms. One of the main aims of the present study, then, is to provide support or otherwise for Hunt’s results by means of a computer "experiment." This allows a genuine steady state to be set up and detailed probing of the interaction between Brownian and shear collision mechanisms.
3. COMPUTER SIMULATION

3.1 General Technique

The simulation proceeds by tracking the positions and sizes of a variable number, \( N \), of spherical particles (typically \( 50 < N < 600 \)). Whenever two particles collide they are coagulated to form a larger (still spherical) particle, conserving particle volume, thereby reducing \( N \) by one. The population of particles studied therefore consists of particles of unit volume, \( v_0 \), and integral multiples, \( v_i = i v_0 \), of the unit volume. In this paper the suffix \( i \) is used to denote properties of \( i \)-fold particles made up from \( i \) elemental particles. All lengths and times in the computer model are non-dimensionalized with the radius of the unit particle and the time step \( \Delta t \). The collision simulation algorithm is programmed for a digital computer.

The program can also function in a different mode in which collisions are counted but particles are not coagulated. On collision, one of the particles is randomly repositioned so as to avoid repeated collisions of the same pair of particles. This allows direct measurement of the collision function, \( \beta \), for any given mechanism.

Particle motions take place in a cubical box or "control volume" of side \( L \) and volume \( V \). (Figure 1 gives a schematic representation of this box and a definition of the rectangular coordinate system used.) Particle positions are denoted by \( \mathbf{r}(i) = (P_1(i), P_2(i), P_3(i)) \). The simulation employs what are essentially periodic boundary conditions, so that particles that have left the control volume at the end of a time step are replaced, for the next time step, by image particles that enter from the opposite side. This type of boundary condition is commonly employed in Monte Carlo simulations (see Alder and Wainwright,
and allows an infinite homogeneous system to be modeled approximately by a finite volume. Edge effects are reduced by allowing particles to interact with image particles just outside the control volume. The slight modifications to these boundary conditions required for laminar and turbulent shearing motions are described in §3.4 and §3.5 below.

In order to model a system in dynamic equilibrium, a fixed number $N_c$ of particles of unit volume are added to the population at random each time step and any particles that have reached a preset maximum volume $v_{\text{max}} = i_{\text{max}} v_o$ are removed from the population. (Typically, $i_{\text{max}} = 125$). The constant addition of small particles is a crude attempt to represent, indirectly, the particle volume flux $E$ into the size range from the collision of particles smaller than $v_o$. In the simulation

$$E = \frac{v_o N_c}{V \Delta t}.$$  

The removal of large particles is necessary to limit the total volume density of particles in the simulation. It can be physically justified as a crude representation of the loss of larger particles from a region by the combined action of sedimentation and vertical concentration gradients. The procedure of adding small particles and extracting large ones is consistent with the hypothesis that collisions between particles of similar size are more important and is justified a posteriori by the success of the simulation in reproducing Hunt's (1982) dimensional results.

The simulation starts either by generating a monodisperse population of particles randomly distributed over the control volume,
or by reading a set of particle positions and sizes from a preexisting file. This file is either a set of particles of given size distribution generated by an auxiliary program, or the end point of a previous simulation that is to be continued. Controlling parameters for the simulation run are either input manually or read from a file.

The particular methods for generating the particle displacements at each time step, \( \chi(i) = (Y_1(i), Y_2(i), Y_3(i)) \), and updating their positions between time steps are described in detail below in connection with each physical collision mechanism. Each particle is assumed to travel on a straight line path at constant speed during each time step. The algorithm used to detect particle collisions is described in §3.2 below.

At the end of every time step the particle size distribution is computed. After a prescribed number \( k \) of time steps, the size distribution, averaged over time \( t = k \Delta t \), is placed, along with the positions and sizes of all the particles, in a file in permanent computer storage. The particle positions and sizes are written over the previous copy to save storage space. The latest version is then always available to restart a run at a later time. The simulation continues until the required number of time steps have been completed.

Time averages are needed to provide reasonable particle size statistics as only a small number of particles are followed. Once a simulated system has reached a statistical steady state (dynamical equilibrium) then long time averages can be employed to produce well converged statistics. To follow the evolution of a rapidly changing system with any precision, it would be necessary to repeat the simulation many times and compute ensemble averages.
Most simulations were started with a monodisperse population of particles. The total volume of particles in the simulation increases continuously until the first particle grows by coagulation to $v_{\text{max}}$ and is removed. In order to have reasonable computational times the volume concentration, $\phi$, of suspended particles used in the simulations is larger than that occurring in many natural systems. (For example, typically $\phi$ is about 20 p.p.m. in Hunt's experiments but is about $10^3$ larger in the simulation runs).

The simulation requires the generation of relatively large numbers of (pseudo-) random numbers from both uniform and Gaussian distributions. First, a sequence of (pseudo-) random numbers distributed uniformly on the interval $(0,1)$ are generated by the standard congruence method. Random variates are then scaled to any required uniform distribution. Variates with Gaussian distribution are generated from this sequence by various algebraic manipulations and employing a six constant rational function approximation to the inverse of the Gaussian cumulative distribution function (details in Abramowitz and Stegun, 1964, §26.2.23 and §26.8).

3.2 Collision algorithm

Detecting which particles have collided at each time step is very costly in computer time and so an efficient method is needed. To this end the basic control volume is divided into cubic sub-cells. The cells are chosen to be as small as possible consistent with the constraint that any particle can only collide, during the next time step, with particles in the same cell or in the adjoining 26 cells. Each cell is given three integer coordinates that define its position in the control
For each particle the numbers of the cell it occupies are stored along with its actual position. The first stage in checking for collisions is to determine for each pair of particles whether they are in the same or adjoining cells. Only if this is so, are they considered candidates for a collision and a detailed calculation performed. Checking whether particles are in adjoining cells is performed by computationally fast integer arithmetic. Given two candidate particles, their relative initial position, $\mathbf{r}_p = \mathbf{p}(1) - \mathbf{p}(2)$, and displacement, $\mathbf{r}_y = \mathbf{y}(2) - \mathbf{y}(1)$, are computed (note the different ordering of particles). Then the condition for collision is that the vector $\mathbf{r}_y$ enters the sphere of radius $\sigma = r_i + r_j$ around the point $\mathbf{r}_p$, a simple geometrical test. This corresponds to following the motion of the two particles in a frame of reference moving with the (1) particle (see Figure 2 for schematic illustration).

A further advantage of the sub-cell system is that it allows for easy implementation of periodic boundary conditions. Particles in cells along any of the boundaries of the control volume are allowed to interact with particles in the requisite cells on the opposite side of the volume.

3.3 **Brownian motion**

The thermal impact of molecules cause suspended particles to perform random motion relative to the bulk fluid. In contrast to the recent work of Nowakowski and Sitarski (1981), the particles studied here are much larger than the molecular free-path in the fluid and so are in the continuum regime of Brownian motion. Also the time step, $\Delta t$, of the simulation is very much larger than the particle viscous
relaxation time, $t_r = 2r^2/9u$. Therefore, the relevant probability distribution function (p.d.f.) for the displacement, $Y$, of a particle during a time step is (Chandrasekhar, 1943)

$$W(Y) = \frac{1}{(4\pi D \cdot \Delta t)^{3/2}} \exp \left(-\frac{|Y|^2}{4D\Delta t}\right)$$

where $D$ is the diffusivity of the particle

$$D = kT/(6\pi \mu r) = \frac{K_b}{(6\pi \tau)}.$$

Each component of $Y$ has an independent Gaussian p.d.f.

$$W(Y_k) = \frac{1}{(4\pi D \Delta t)^{1/2}} \exp \left(-\frac{y_k^2}{4D\Delta t}\right) \quad k = 1, 2, 3$$

and this is used to replace the Brownian motion of the particles by a finite random walk. At each time step three independent random components of displacement are generated for each particle from the corresponding Gaussian distribution. The r.m.s. displacement in any direction, $\Delta x$, of an $i$-fold particle is

$$\Delta x_i = (2 \cdot D_i \cdot \Delta t)^{1/2}$$

where, $D_i = k_b/(6\pi \tau_i)$, is the particle diffusivity. $D_i$ can be obtained in terms of the diffusivity $D_0$ of an elemental particle by

$$D_i = D_0 \cdot i^{-1/3}.$$

Particle collisions are simulated on the basis of straight line trajectories during each time step. The question arises, therefore, of the validity of this as an approximation to Brownian induced coagulation. The r.m.s. displacement has been chosen correctly, but a particle of
mass $m$ undergoing Brownian motion actually travels along a tortuous path at r.m.s. speed $(kT/m)^{1/2}$. At first sight this suggests that the simulation would underpredict the collision rate. However, replacing Brownian motion by a finite random walk must change the pair distribution function, that is to say the probability distribution function for the spacing between any given pair of particles. So, while modeling Brownian motion by a finite random walk introduces inefficiency into the basic collision process it can compensate by increasing the probability that any pair of particles are found close together at the beginning of a time step. Here, "close together" means a separation on the scale of the r.m.s. steplength of the random walk. These matters are investigated in detail in Appendix A. Tests with the non-coagulating form of the program have shown that satisfactory collision rates for monodisperse populations of particles are obtained when the ratio $\Delta x/r$ is about 0.5. It is important to use the maximum possible time step in order to minimize computation times.

3.4 Laminar shear

The coagulating effects of a velocity gradient are investigated by imposing a uniform shearing motion on the control volume:

$$u_1 = G \cdot x_3$$

with $G$ the shear rate. The particles are assumed to move with the fluid so their displacement in any time step is just

$$\mathbf{Y}(i) = (Y_1(i),0,0); \quad Y_1(i) = G \cdot P_3(i) \cdot \Delta t$$

As stated in the introduction, in this paper we are ignoring hydrodynamic interactions between particles. The large body of work on particle
interactions in low Reynolds number flows (see e.g. Mason, 1976, for a review) shows that hydrodynamic forces will always come into play in a detailed analysis of collision dynamics. This is investigated in detail in a subsequent paper.

A uniform shearing motion, on average, moves a fraction of the particles out of the control volume at every time step. If they were replaced in the control volume according to simple periodic boundary conditions \( P_1 = P_1 - L \), whenever \( P_1 > L \) the simulation would be completely deterministic once initial positions had been chosen for the particles. Each particle would move in a straight line with fixed \( P_2 \) and \( P_3 \) coordinates. After a certain time all collisions between existing particles would cease as each particle would have swept out its own track through the control volume. In a real flow this would not occur as particles are continually meeting "new" particles. Therefore, in the simulation, when a particle leaves the volume it is replaced at a randomly chosen height \( P_3 \) on the other side of the control volume. The random value of the height \( P_3 \) must be chosen from a distribution that reflects the increasing flux of particles at larger values of \( P_3 \). This flux is proportional to \( P_3 \) and a uniformly distributed random variate may be converted to this linear p.d.f. by taking its square-root. This strategy leads to a further complication: particles may be replaced on top of one another, leading to spurious collisions. This is almost totally eliminated by checking for such particle overlaps at the end of each time step and randomly moving one of each overlapping pair. This may introduce a few further overlaps as no final check is made. An estimate of this number is available.
from the number of initial overlaps, which is recorded. This error is acceptable in the light of other approximations in the simulation. Overlaps are also introduced by the process of adding new elemental particles at each time step, whatever the collision mechanism. All types of overlaps are resolved simultaneously in the same manner.

3.5 **Turbulent shear**

We wish to simulate the coagulation of small particles by turbulent flow. The motion of suspended particle can be identified with the motion of an adjacent fluid particle provided that the time scale of the (fluid) particle acceleration is much greater than the particle relaxation time, \( t_r \), that is to say, if inertial effects are negligible, as will be the case here. Then for particles of radius smaller than the smallest scale of the turbulent motion (the Kolmogorov length scale, \( (\nu^3/\varepsilon)^{1/4} \)), coagulation rates are determined solely by the kinematics of the small scales of the turbulent flow field, in particular by the r.m.s. strain rate \( (\varepsilon/\nu)^{1/2}/15^{1/2} \). These small scales are very nearly isotropic (Batchelor, 1953).

Under these conditions, two particles separated by a distance smaller than the Kolmogorov length scale are subjected to a motion that can be decomposed into a rigid body rotation representing the local vorticity, and a locally uniform three-dimensional straining motion. The rigid body rotation component of the motion has no effect on the collisions of non-interacting particles and so only the straining motion (with symmetric velocity gradient tensor) is modeled. The straining motion will be uniform over length scales smaller than the Kolmogorov
micro-scale but there is no agreement as to the duration of this straining (Monin and Yaglom, 1975). Two time scales are important for the small scale straining: the rate of rotation of the principal axes of strain and the rate of change of the magnitude of the principal rates of strain. For turbulent flow at high Reynolds number the rate of change of the deformation fields of the small eddies is related to the Lagrangian time microscale $\alpha$ (Lumley, 1972). The time scale of the deformation field is $\lambda/u'$, where $\lambda$ is the Taylor microscale and $u'$ the r.m.s. fluctuating velocity. Corrsin (1963) approximates the ratio of the two as

$$\alpha u'/\lambda \approx (R_\lambda/30)^{1/2}$$

and since by definition

$$\lambda^2 = 15\nu \frac{u'^2}{\epsilon} \quad \text{and} \quad R_\lambda = \frac{u'\lambda}{\nu}$$

we have

$$\frac{\alpha}{(\nu/\epsilon)^{1/2}} \approx (R_\lambda/2)^{1/2}$$

which implies that the strain and vorticity fields of the small eddies remain constant for a time interval at least equal to the Kolmogorov time scale, $t = (\nu/\epsilon)^{1/3}$. This is just the inverse of the characteristic strain rate.

The effect of the rate of rotation of the principal axes of strain on the collision rate was investigated using the monodisperse, non-coagulating version of the simulation. The velocity gradient was simulated so that both the principal axes and principal rates of strain could be changed independently. The magnitude of the strain was kept
constant for a time interval equal to the Kolmogorov time scale. No statistically significant difference in the collision rate was found, whatever the time scale of rotation of the principal axes of strain. Therefore in the coagulation simulation both principal axes and rates of strain were varied at the same rate.

Assuming homogeneous, isotropic, unbounded turbulence with a Gaussian velocity gradient field, the elements of the rate of strain tensor were chosen randomly to satisfy (Hinze, 1959)

\[
\frac{\partial u_i}{\partial x_k} \frac{\partial u_j}{\partial x_\ell} = \begin{cases} 
\frac{1}{15} \frac{\varepsilon}{\nu} & \text{i=j=k=\ell} \\
- \frac{1}{30} \frac{\varepsilon}{\nu} & \text{j=\ell and i=k or i=\ell and j=k and i\neq j} \\
\frac{2}{15} \frac{\varepsilon}{\nu} & \text{k=\ell and i=j and i\neq k} \\
0 & \text{all other combinations}
\end{cases}
\]

subject to

\[
\frac{\partial u_i}{\partial x_i} = 0
\]

and kept constant for a time interval equal to the Kolmogorov time scale.

The simulation proceeds as in the case of laminar shear with particle displacements being given by the product of the time step (\(\Delta t\)) and the fluid velocity corresponding to the particle position. Now, however, as the motion is three-dimensional and stochastic, true periodic boundary conditions can be used. This corresponds to the control volume being surrounded by copies which are deformed with the original. Particles in the control volume at the end of one time step can then be used for the next. However, in preliminary simulations,
random fluctuations in the number of particles were found to cause trouble. To avoid the program halting because of too many or no particles left in the control volume the total number was adjusted at each time step according to

\[ N_{i+1} = N_i - N_{\text{COL}} + N_c \]

where \( N_{\text{COL}} \) is the number of collisions that had occurred during the time step, and \( N_c \) the number of elemental particles added. In order to satisfy the above condition, either particles were removed at random, or a particle whose volume had been chosen at random from the existing population was added at a random position. Finally, particle overlaps were resolved as explained in §3.4.

### 3.6 Multiple mechanisms

Simulations were performed in which the particle displacement was the linear sum of a fluid shearing and a Brownian component. The relative magnitude of the Brownian and shearing parameters could then be varied to investigate their interaction.

### 4. RESULTS

Figure 3 shows the effect of changing the r.m.s. steplength on collision rate in Brownian motion (see Appendix A for a discussion). There is some statistical scatter in the results but the general shape of the curve is correct. From these results a suitable time step can be chosen for simulations involving Brownian motion. Similar computations of collision rates in laminar and turbulent shear induced
coagulation were performed to check that they yielded the values given by Table 1. This, indeed, was found to be the case. The result for turbulent shear due to Saffman and Turner (1956) has been amended by a factor of $\frac{1}{2}$ from that in the original paper, correcting an algebraic error.

The development of a size distribution in a typical simulation starting with particles all of unit volume $v_0$ and undergoing Brownian induced coagulation is shown in Figure 4. The size distribution is non-dimensionalized according to equation (3) and plotted logarithmically against particle volume non-dimensionalized with the unit particle volume. The curves plotted are smoothed approximations to the actual data points, at $v = i \cdot v_0$, which are rather scattered. The upper portion of the data attains a slope of $-3/2$ once a range of about one decade in volume has been reached. Then, as particles of increasing size are formed, the slope of the size distribution remains the same, but its absolute level declines gradually. It reaches a statistically steady state once the first large particle is lost from the system. The final steady state for this set of parameters is shown in Figure 5, along with that for a run at a higher final volume concentration $\phi$ (this is obtained by adding more particles at each time step). The points plotted are actual data from the simulations, averaged over 1000 time steps. Even with this time averaging there is still some statistical scatter in the data, especially at the lower end of the size distribution where very small numbers of particles are actually involved. To further smooth the data in the region $v/v_0 = 20-100$, they have been averaged in groups of 5.
For both these runs $v_{\text{max}} = 125 v_o^*$, although the volume distribution is only plotted out to $v/v_o = 100$. Beyond this the data becomes erratic. The two sets of data are fully collapsed by the normalization used and very clearly exhibit the -2/3 power law expected from Hunt's (1982) theory. The intercept of the best fit line of slope -3/2 with the axis $v/v_o = 1$ gives the constant $a_b$ in equation (3).

Figure 6 is a comparison of the steady state size distributions for laminar shear at two volume concentrations differing by an order of magnitude. Again the data points are averaged over 1000 time steps, and are collapsed onto a slope of -2 by the normalization suggested by dimensional arguments. Similar results are shown for turbulent shear in Figure 7, where the inverse of the Kolmogorov time scale, $(\varepsilon/\nu)^{\frac{1}{2}}$, is used in place of $G$ in the normalization of the size distribution. Again, a -2 power law is achieved at steady state and the normalized results are independent of the flux of particle volume through the size range. Note, however, that the data points are slightly lower than in the case of laminar shear. This is simply a consequence of the collision functions given in Table 1: the expressions for laminar and isotropic turbulent shear are identical if $G$ is replaced by $1.72 (\varepsilon/\nu)^{\frac{1}{2}}$. With this scaling the data of Figures 6 and 7 collapse. This result strongly suggests the equivalence of laminar rectilinear shear and three-dimensional turbulent shear as coagulating agents.

It is gratifying that the results of the simulations, which do not assume forms for the collision functions, $\beta$, agree well with arguments suggested by the analytic estimates for $\beta$. 
The next series of simulation runs illustrate the effect that the ratio \( i_{\text{max}} = v_{\text{max}}/v_0 \) (i.e., the size range covered by the simulation has on final steady state size distributions in Brownian motion and laminar shear. Figures 8 and 9 give size distributions for the three cases \( v_{\text{max}}/v_0 = 27, 125, \) and \( 512 \). In all cases the relevant \(-3/2\) or \(-2\) power law prevails. For Brownian motion the results for \( v_{\text{max}}/v_0 = 125 \) and \( 512 \) are indistinguishable, while those for the smallest size range are slightly higher at the upper end of the size range. For laminar shear there is a slight but consistent decline in level with increasing size range. This reflects the extent to which the size distribution is affected by the collisions of the relatively small number of large particles. In laminar shear the collision function increased with the volume of the particles involved faster than in Brownian coagulation. Work on the effects of hydrodynamic interactions between particles on coagulation (see Adler, 1981 for most recent study) suggests that they act to reduce most the collision rate between particles of widely different sizes. This would probably result in weaker dependence of the level of the size distribution (the value of \( a_{\text{sh}} \)) on the size range covered by the simulation. Further work, with a more sophisticated simulation incorporating hydrodynamic interactions, will elucidate this point.

A consensus of the simulations performed gives the values,

\[
\begin{align*}
    a_b &= 0.2 \pm 0.04, \\
    a_{\text{sh}} &= 0.24 \pm 0.05
\end{align*}
\]

However, it is likely that accounting for hydrodynamic and interfacial forces will alter the values of these dimensionless constants.
So far all the results have been for simulations in which only one collision mechanism has been present. We now turn to cases where both Brownian motion and fluid shearing operate. A new normalization of the size distribution and volume variable is now required to collapse all the data. Following Hunt (1982) we define a non-dimensional volume

$$\chi = v^* (K_{sh}/K_b)$$

where $K_{sh}$ represents $G$ or $1.72 (c/v)^{1/2}$ and $K_b$ is as before. This is such that the collision rates due to Brownian motion and shear are equal for particles of size $\chi = 1$. Then if a normalized size distribution is defined by

$$n^*(\chi) = a_b \cdot \chi^{-3/2}$$

and

$$n^*(\chi) = a_{sh} \cdot \chi^{-2}$$

Results of three simulations each for laminar and turbulent shear with Brownian motion are plotted in this normalized form in Figure 10. Lines of slope $-3/2$ and $-2$ are drawn for comparison. There is some indication of a change in slope around $\chi = 1$ but it is not conclusive. Also, the constant $a_b$ and $a_{sh}$ obtained from the data in Figure 10 are the same (within statistical error) as those obtained from simulations with only one collision mechanism present, providing some support for the hypothesis of non-interference of mechanisms.
5. DISCUSSION

The main aims of this study have been:

1. to study the feasibility of a Monte Carlo simulation of both the collision function, $\beta$, and the coagulation equation (2) for the evolution of a population of particles to a steady state;

2. the investigation of Hunt's (1982) theory for the form of the resulting size distributions.

The simulation method described has proved most successful in modeling the coagulating powers of both Brownian and bulk shearing mechanisms and the development of steady state size distributions. This is in spite of the relatively restricted range of particle sizes that can be followed in any one computer run and the somewhat artificial strategy of adding new unit particles at each time step.

The results show that final steady state is rather insensitive to the size range covered, and that the size distribution at the upper end, (small particles), is not very disturbed by replacing the interactions of all small particles with the addition of unit particles at a constant rate. These observations are in accord with the striking success of dimensional analysis in predicting the observed size distributions. For dimensional analysis to be successful the dynamics of the coagulation process must be mainly "local" in size space so that further independent parameters (such as $v_o$ and $v_{max}$) are not important. We expect that accounting for hydrodynamic interactions between particles will decrease the dependence of the level of the size distribution, for given volume
flux, in shear-induced coagulation. Notice that the evolving populations of particles start to exhibit the relevant power-law over much of their size distribution long before a steady state is reached.

Hunt's further hypothesis that different collision mechanisms can act independently over separate size ranges has been partially confirmed. A slope of $-\frac{3}{2}$ is not very different from one $-2$ when there is scatter in the data! However, complete resolution of this point would require the simulation to cover a greater range of particle sizes. This is not feasible with the available computer storage. The perturbation analysis of van de Ven and Mason (1977), for the effect of weak shear on Brownian coagulation, suggests that when hydrodynamic interactions are considered the two mechanisms may not be strictly additive.

In conclusion, it can be said that, while simple in concept, and using acceptable computer resources, the simulation method has provided useful elucidation of Hunt's hypotheses and experimental results under carefully controlled conditions. Further work on the technique to include hydrodynamic interactions, interfacial forces and gravitational settling will be reported in a subsequent paper.

ACKNOWLEDGEMENTS

Financial support for this work was provided by NOAA/Sea Grant NA80AA-D-00120, NOAA Grant NA80RAD-0084 and a Mellon Foundation Grant to the Environmental Quality Laboratory at Caltech. The authors would also like to thank Dr. R.C.Y. Koh for advice on many aspects of the computer code and Dr. R. C. Flanagan for helpful discussions of the problem.
APPENDIX A

FINITE STEPLENGTH AND COLLISION RATE IN BROWNIAN MOTION

The theoretical collision function, $\beta$, for Brownian induced collisions between particles of radii $r_i$ and $r_j$ given in Table 1 was computed (see e.g. Chandrasekhar, 1949) by solving a diffusion equation for the pair distribution function, $w(s)$, where $s$ is the distance between the particles. In particular, the collision function is given by the asymptotic flux to the surface of a fixed sphere of radius $\sigma = r_i + r_j$, with a total diffusivity $D = D_i + D_j$. The "concentration", $w$, is held at zero at $s = \sigma$ and at units $s = \infty$. Initially, $w$ is uniform outside the sphere. Then at large times the pair distribution function is given by

$$w = 1 - \sigma/s \quad \text{(A.1)}$$

whence the required result:

$$\beta = 4\pi D(s^2 dw/ds)_{s=\sigma} = 4\pi D \sigma \quad \text{(A.2)}$$

If the actual pair distribution function in the finite steplength simulation was identical to that in (A.1), then the collision rate measured would be no larger than one-half of that in (A.2), however small the steplength. This result can be obtained either by careful evaluation of the expected collision probability from the algorithms used for generating particle displacements and detecting collisions, or by the following simple argument. In the limit of $\Delta x \ll \sigma$, i.e., very small r.m.s. steplength, but still with $\Delta t \gg t_r$, two particles must be so close at the beginning of the time step in which they collide that the curvature of their surfaces may be neglected. The problem then
reduces to that of the collision of a diffusing point with an adsorbing plane and we need only consider the component of the random walk perpendicular to the plane.

Consider now this one-dimensional problem. The particle is judged to have collided with the plane if its final position is on the far side of the plane. For any given final position on the far side of the plane there is a whole class of possible Brownian trajectories leading to it. Now each of these trajectories must cross the plane for the first time at some point. There will be an associated trajectory defined to be identical with the original until the first contact with the adsorbing plane and then the mirror image, in the plane, of the original. As the end-point of this associated trajectory lies on the near side of the plane it would not be judged a collision by the collision algorithm. Hence the 50 percent inefficiency.

However, for the same reason, the pair distribution function will not be identical in the theoretical and simulated cases. In the finite steplength case, $w$ will be larger within a distance of order $\Delta x$ of $s=\sigma$. This can compensate for the basic inefficiency of the collision algorithm. The actual form of $w$ for a given distribution of steplengths and hence the collision function could be computed by solving the relevant integral equation. This has not been done as yet, but the non-coagulating form of the simulation has been used to determine the collision rate for a monodisperse population of particles as a function of the mean steplength. The results of this "experimental" determination are shown in Figure 3. The ratio of measured collision rate to that predicted from (A.2) is
plotted against the ratio of r.m.s. displacement in any direction, $\Delta x$, and the particle radius $r$. The ratio is unity for $\Delta x/r$ about 0.6 and so $\Delta x$ is chosen accordingly in all the coagulation simulations.
REFERENCES


Findheisen, W. 1939 Meteor. Z. 56, 356.


Collision functions and characteristic dimensional parameters for various particle collision mechanisms. Values of $\beta$ are for collision mechanisms acting individually with no hydrodynamic or other interparticle forces.
FIGURE CAPTIONS

Figure 1. Schematic diagram of simulation box or "control volume" with cartesian coordinate system and representative particle at position $(P_1, P_2, P_3)$. Displacement of particle in current time step is $(D_1, D_2, D_3)$.

Figure 2. (a) Geometry for collision algorithm. (b) Viewed in frame of reference in which particle 2 is at rest.

Figure 3. Simulated collision rate of monodisperse particles undergoing Brownian motion as a function of r.m.s. displacement. The ratio between measured collision rate and theoretical rate is plotted against the ratio of step length to particle radius.

Figure 4. Development towards steady state of size distribution of initially monodisperse population undergoing Brownian induced coagulation. $D_0 = 0.222$, $E = 5.6 \times 10^{-5}$, $i_{\text{max}} = 125$; time: $--- t = 25$; $- - - - t = 50$; $- - - - t = 100$; $- - - t = 200$; $- - - t = 400$; $- - - t = 600$.

Figure 5. Steady state non-dimensional size distribution for Brownian motion. $D_0 = 0.222$, $i_{\text{max}} = 125$; $+ E = 5.6 \times 10^{-5}$, $\phi = 0.016$; $\Box E = 4.4 \times 10^{-4}$, $\phi = 0.043$.

Figure 6. Steady state non-dimensional size distribution for laminar shear. $+ G = 0.25$, $E = 1.4 \times 10^{-5}$, $i_{\text{max}} = 125$, $\phi = 0.013$; $\Box G = 1$, $E = 1.1 \times 10^{-3}$, $i_{\text{max}} = 125$, $\phi = 0.114$.

Figure 7. Steady state non-dimensional size distribution for turbulent shear. $K_{sh} = 0.5$, $i_{\text{max}} = 125$; $+ E = 5.7 \times 10^{-4}$, $\phi = 0.053$; $\Box E = 1.4 \times 10^{-5}$, $\phi = 0.008$. 
Figure 8. Comparison of steady state non-dimensional size distribution for Brownian motion for different $i_{\text{max}}$. $D_0 = 0.222$, $E = 5.6 \times 10^{-5}$; $\square i_{\text{max}} = 27$, $\phi = 0.011$; $\triangle i_{\text{max}} = 125$, $\phi = 0.016$; $\Delta i_{\text{max}} = 512$, $\phi = 0.0022$.

Figure 9. Comparison of steady state non-dimensional size distribution for laminar shear for different $i_{\text{max}}$. $\square G = 1$, $E = 1.1 \times 10^{-3}$, $i_{\text{max}} = 27$, $\phi = 0.049$; $\oplus G = 1$, $E = 1.1 \times 10^{-3}$, $i_{\text{max}} = 125$, $\phi = 0.057$; $\Delta G = 0.25$, $E = 1.4 \times 10^{-5}$, $i_{\text{max}} = 512$, $\phi = 0.014$.

Figure 10. Steady state normalized size distribution for (i) Brownian motion and laminar shear, (ii) Brownian motion and turbulent shear. (i) $E = 5.6 \times 10^{-5}$, $i_{\text{max}} = 125$; $\square K_{\text{sh}} = 0.25$, $\phi = 0.0081$; $\times K_{\text{sh}} = 0.125$, $\phi = 0.011$; $\Delta K_{\text{sh}} = 0.063$, $\phi = 0.013$.
(ii) $\oplus K_{\text{sh}} = 0.344$, $D_0 = 0.8$, $E = 1.7 \times 10^{-5}$, $i_{\text{max}} = 512$, $\phi = 0.0096$; $\Diamond K_{\text{sh}} = 0.215$, $D_0 = 0.222$, $E = 5.6 \times 10^{-5}$, $i_{\text{max}} = 125$, $\phi = 0.0077$; $\bigcirc K_{\text{sh}} = 0.043$, $D_0 = 0.222$, $E = 5.6 \times 10^{-5}$, $i_{\text{max}} = 125$, $\phi = 0.013$. 
Figure 1
Collision Rates for B.M.
Figure 8
Figure 10

\[ n(v)K_b^{-2}E^{-0.5}K_{sh}^{-1.5} = \frac{vK_{sh}}{K_b} \]