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DOI:
10.1016/j.ijmultiphaseflow.2015.03.008

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Document Version
Peer reviewed version

Citation for published version (Harvard):
LAGRANGIAN PARTICLE TRACKING IN MECHANICALLY AGITATED POLYDISPERSE SUSPENSIONS: MULTI-COMPONENT HYDRODYNAMICS AND SPATIAL DISTRIBUTION

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Abstract
We investigated the local hydrodynamics and phase distribution of complex polydisperse suspensions ‘just-suspended’ by a down-pumping pitched blade impeller in a mechanically agitated vessel. The solid-liquid mixtures consisted of five size fractions of coarse glass particles in water, and the total concentration was varied within the range 2.4–23.6 vol% (5–40 wt%). The whole flow field and spatial distribution of the liquid phase and of each particle size fraction were resolved using the Lagrangian technique of positron emission particle tracking (PEPT). For the first time, it has been possible to conduct such a detailed ‘pointwise’ examination within an opaque polydisperse suspension of this type and complexity. For each component of the two-phase flow, results are presented in the form of maps of local velocity, solid concentration and time-averaged slip velocity, as well as plots of spatial suspension uniformity index. The accuracy of measurements is verified through six-component mass balance and mass continuity calculations.

Keywords: Lagrangian tracking; mixing; PEPT; polydisperse suspension; slip velocity; suspension uniformity.

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1. Introduction

Industrial processing of solid-liquid suspensions such as bio-slurries, minerals, food, crystallisation and catalytic mixtures, is commonly conducted in mechanically agitated vessels. A key aspect of such a process is the achievement of good mixing to ensure a good distribution of all the different phase components and enable an efficient interaction between them. The methods generally used for designing stirred vessels for solid-liquid mixing have tended to follow a global ‘black-box’ approach, but a more detailed description of the internal flow structure is needed to enable more rational rules to be established for equipment and process design. The advent of powerful measurement and modelling techniques in recent years has revivified interest in this field and increasingly more attempts are being devoted to better understand the mechanisms behind particle suspension and distribution in such systems (Kagoshima and Mann, 2006; Guha et al., 2007; Unadkat et al., 2009; Tamburini et al., 2009).

Homogeneous suspension exists when the solids are practically uniformly distributed throughout the vessel volume, i.e. there are no solids concentration gradients, and for a polydisperse system the particle size distribution should also be approximately the same everywhere. This condition is required when it is necessary to obtain uniform treatment of all the particles, when a suspension must be dispensed at a constant concentration for sampling, for further processing, or for packaging and the process result that the mixing has to achieve is that the discharge from the vessel should enable homogeneous filling of packaging lines, e.g. meat/vegetable pieces in a sauce or fruit particles in yoghurt. Another example would be in a crystalliser where a non-uniform solids distribution may cause high local supersaturation levels and hence non-uniform crystal growth which is undesirable. Therefore, knowledge of the local solid concentration and distribution in polydisperse systems, where multiple particle size fractions are suspended in a liquid phase, would be useful for optimum process design, development and scale-up.

Detailed experimental investigation of these phenomena has remained a major challenge for decades due to a lack of adequate measurement techniques capable of probing such opaque multiphase systems (Barigou, 2004). Among the various flow visualisation techniques, laser Doppler velocimetry (LDV) and particle image velocimetry (PIV) have become the most reliable to examine complex flow fields in optically transparent single-phase systems (Ducci and Yianneskis, 2006; Chung et al., 2007, 2009). Application to multi-phase studies, however, has been confined to dilute suspensions as these techniques fail completely in dense systems which are opaque.

Attempts at local measurements in these complex systems have been chiefly limited to the measurement of mean axial solid-concentration profiles at relatively low concentrations, using a single vertical conductivity or capacitance probe traverse or a withdrawal technique (Barresi and Baldi, 1987; Brunazzi et al., 2004). More recent experimental studies using electrical resistance tomography (ERT) demonstrated, albeit mainly qualitatively, how visualisation of gas, solid or liquid distribution can help improve understanding of mixing processes (Wang et al., 2000). However, none of these methods is capable of probing concentrated suspensions in detail to acquire quantitative information on the local flow dynamics of the different phase components or their individual 3-D distributions.
Rammohan et al. (2001) developed a more sophisticated computer automated radioactive particle tracking (CARPT) technique based on gamma-ray emissions. It has been argued for some time that Lagrangian data obtained from a single particle trajectory, where such data can be accurately determined, may unravel valuable mixing information which is not provided by Eulerian observations (Villermaux, 1996; Wittmer et al., 1998; Doucet et al., 2008; Guida et al., 2009; Barigou et al., 2009). The technique of positron emission particle tracking (PEPT) allows non-invasive probing of opaque fluids and within opaque apparatus by using a single small sub-millimetre positron-emitting particle as flow tracer (Parker and Fan, 2008). Being able to examine flow phenomena in three dimensions in dense opaque systems that could not be observed as effectively by using other techniques, PEPT is particularly useful for the study of non-transparent multi-phase or multi-component flows, where one component can be selectively labelled and its behaviour observed (Fangary et al., 2000, 2002).

A number of mixing studies have reported that in stirred solid-liquid systems, variation in the particle size produces significant effects on the fluid dynamics and phase distribution. For example, Virdung and Rasmussen (2007) using LDV measurements showed that the fluid velocity fluctuations due to turbulence in the liquid phase were higher in solid-liquid suspensions than in single phase flow, an effect that increased with particle size. Although their simulations involved particle sizes less than 0.1 mm, Altway et al. (2001) showed that the pattern of solid concentration contours for different particle sizes was qualitatively similar; however, smaller particles showed a more uniform distribution. Using larger particles with 0.1 mm < d < 0.5 mm, Špidla et al. (2005) experimentally confirmed by means of a conductivity probe that smaller solid particles generate a more homogeneous suspension. Their results also showed that the homogeneity of the solid-liquid suspension improved with increasing average solid concentration. We recently observed the same effect using coarse monosized or binary suspensions of glass particles (d ~ 1, 3 mm) (Guida et al., 2010a, 2011; Liu and Barigou, 2013).

In this paper, we report on the use of the Lagrangian PEPT technique to study the mixing of dense polydisperse mixtures consisting of glass beads of five different size classes suspended in water. We present for the first time extensive data for these polydisperse systems containing up to 40 wt% (23.6 vol%) solids. Detailed Lagrangian information is obtained on particle and fluid trajectories which is converted to give a ‘pointwise’ Eulerian description of the flow field as well as the spatial distribution of each of the liquid and solid components of the mixture.

2. Materials and methods
2.1 Experimental conditions
The mixing experiments were conducted in a fully-baffled flat-base Perspex vessel of diameter $T = 288$ mm, agitated by a down-pumping 6-blade 45° pitched-turbine (PBT) of diameter $D = T/2$, as illustrated in Fig. 1. The height of the suspension was set at $H = T$ and the impeller off-bottom clearance was $T/4$. The suspending liquid was an aqueous solution of NaCl of density 1150 kg m$^{-3}$ ~ NaCl was added to enable the water density to be matched to that of the resin PEPT tracer used to track the liquid phase ~ and the solid particles used were spherical glass beads of density 2485 kg m$^{-3}$. Five nearly-monomodal particle size fractions, ($d_p ~ 1.0$ mm; ~ 1.5
mm; \sim 2.0 \text{ mm}; \sim 2.5 \text{ mm}; \sim 3.0 \text{ mm}) were used to make polydisperse solid-liquid suspensions. The five size fractions were mixed in equal proportions with a total solid mass concentration, $X$, varying from 0 to 40 wt%, i.e. $X_1 = X_2 = X_3 = X_4 = X_5 = 0.2X$. Experiments were conducted under the ‘just-suspended’ regime corresponding to the minimum impeller speed for particle suspension $N_{js}$, which was visually determined in the transparent vessel according to the well-known Zwietering criterion (Zwietering, 1958), i.e. no particle remains stationary on the bottom of the tank for longer than 1-2 s. The experimental mixing conditions are summarized in Table 1.

### 2.2 Multi-component positron emission particle tracking

The multi-component two-phase flow field and spatial phase distribution in the mixing vessel were determined by the technique of positron emission particle tracking (PEPT), as illustrated in Fig. 1. PEPT uses a single positron-emitting particle as a flow tracer which is then tracked in 3-D space and time to reveal its full Lagrangian trajectory. The measurements can be analysed in two different ways: a Lagrangian-statistical analysis exploiting concepts such as residence time, circulation time and trajectory length distribution; or, a Lagrangian-Eulerian analysis used to extract local Eulerian quantities from the purely Lagrangian information contained in the tracer trajectory such as the three velocity components ($u_r$, $u_z$, $u_\theta$) of each phase component, its local occupancy or time-average concentration (Guida et al., 2012). PEPT has been validated by comparing its measurements with PIV in water, and has been shown to be an accurate and reliable technique (Pianko-Oprych et al., 2009). More details about the technique can be found in our earlier papers (Barigou, 2004; Guida et al., 2009; Barigou et al., 2009; Guida et al., 2010a, 2011, 2012).

In multi-component positron emission particle tracking, the full three-dimensional trajectory of each component must be resolved separately using different particle tracers. Currently, PEPT can most accurately track one particle at a time and, therefore, each component of the suspension was individually tracked in the suspension and its full trajectory separately determined in one of six successive experiments (liquid phase + 5 solid components). A neutrally-buoyant radioactive resin tracer of 600 \(\mu\)m diameter was used to track the liquid phase, and from each particle size fraction a representative glass particle was taken, was directly irradiated via a cyclotron and its motion tracked within the agitated slurry (Parker and Fan, 2008). The PEPT tracking time was 30 min in each case, long enough so that good data resolution was obtained in every region of the vessel. Furthermore, ergodicity of flow, the theoretical condition which guarantees that a flow follower is representative of all the solid or liquid phase it represents, could be safely assumed under such conditions. This is justified given that in such a system the probability of visit of a cell by the tracer particle is sufficiently uniform by virtue of the highly turbulent state of the suspension ($Re_{cap} = ND^2/\nu > 10^5$), and the open geometry of the vessel where all regions are more or less equally accessible by the tracer. Experimental confirmation of the ergodicity assumption has been discussed in previous papers (Guida et al., 2010a, 2010b, 2011).

### 3. Results and discussion

The measurements obtained from the 3-D Lagrangian tracking of the liquid phase and the five solid fractions were analysed and converted, for each component of the two-phase flow, to azimuthally-averaged Eulerian
maps of local flow velocity, spatial concentration distribution and time-averaged particle-liquid slip velocity, using a cylindrical grid consisting of \( n_c = 512 \) (16 by 32) cells of equal volume, as illustrated in Fig. 2. Cells near the wall have dimensions \( 4.49 \times 2.29 \text{ mm}^2 \), whilst the innermost cells have dimensions \( 4.49 \times 18.00 \text{ mm}^2 \). Six-component mass balance and continuity calculations were also conducted throughout the vessel to verify the accuracy of the PEPT measurements. In order to enable efficient consultation and comparison of data, results are presented in a concise format using sets of images judiciously produced by grouping together multiple graphs, greyscale contours, or vector plots. Finally, the uniformity of the suspensions was assessed by means of a uniformity index.

### 3.1 Local velocity fields

Maps of the local flow velocity are shown in Fig. 3 for the liquid phase and for the five solid fractions at conditions of suspension corresponding to the \( N_{js} \) values given in Table 1. The \( N_{js} \) value shown for each suspension concentration is the one which ensured that all particles were just suspended. It should be noted, however, that \( N_{js} \) was not significantly affected by particle size over the range of particle diameters investigated (1–3 mm). Using cylindrical coordinates, the radial-axial 2-D velocity maps were obtained by azimuthally averaging the 3-D velocity data and projecting them onto the 2-D radial-axial plane \((r, \theta)\). Whilst two-dimensional, these flow maps are, however, representative of the whole 3-D velocity field.

The observed flow field with a single flow loop is typical of a down-pumping PBT with a low off-bottom clearance. As expected, fluid velocities are highest in the impeller region (about \( 0.6u_{\text{tip}} \) near the edge of the impeller) and around the flow loop, but much lower in the upper parts of the vessel and near the centre of the loop.

For ease of comparison, all the velocity plots have been normalised by the impeller tip speed, \( u_{\text{tip}} \). The normalised plots for the liquid and solid components of the flow exhibit remarkable similarity regardless of particle size and solid concentration. The tendency of flow structures at high Reynolds numbers to present similar characteristics, is a phenomenon often referred to as Reynolds number similarity. In fully turbulent flow, viscosity effects vanish and dimensionless mean values such as velocities normalised by the impeller tip speed should be independent of Reynolds number. The values of the impeller Reynolds number (\( \text{Re}_{\text{imp}} = ND^2/\nu \)) are given in Table 1, and span the range \( 1.5\times10^5 - 2.4\times10^5 \). We have previously observed such a similarity in less complex suspensions of mono and binary composition (Liu and Barigou, 2013, 2014). Sample results plotted in Fig. 4, show the azimuthally-averaged radial distributions of the normalised velocity components for the liquid phase and two solid components (1 mm, 3 mm) at all solid concentrations investigated. Some minor disparities are observed, but the principle of Reynolds number similarity seems to hold for both liquid and solids at all solid concentrations.

### 3.2 Time-averaged slip velocity maps

The magnitude of the local time-averaged slip velocity vector was determined for each solid component \( j \), thus:

\[
s_j = \sqrt{(u_r^{(L)} - u_r^{(j)})^2 + (u_z^{(L)} - u_z^{(j)})^2 + (u_{\theta}^{(L)} - u_{\theta}^{(j)})^2}
\]  

(1)
where \((u_r, u_z, u_\theta)\) are the mean cylindrical velocity components for the liquid \((L)\) and solid component \((j)\). The effects of the mean solid concentration on the spatial distribution of the normalised total slip velocity for each solid component of the mixture are depicted in Fig. 5. The largest local slip velocities occur in the discharge region of the impeller and are of the order of \(\sim 0.10u_{tip}\). At the lower solid concentrations, the larger particles with more inertia display significantly more slip than the smaller ones. This effect, however, is barely noticeable at higher solid concentrations as particle-particle interactions become important and differences in velocity reduce through frequent particle collisions.

There are wide variations in the spatial distribution of \(s_j\). As a consequence of Eq. (1), which combines radial, axial and tangential contributions, the regions of maximum slip velocity occasionally correspond with the lower tip of the impeller blades (tangential dominance), with the discharge region of the impeller (radial-axial dominance), or with both areas when there is a more balanced contribution of the three cylindrical components of the slip velocity vector. Considering the individual components of the slip velocity vector, detailed analysis showed that by far the largest slip occurs in the axial \(z\) direction (data not shown). Nevertheless, some significant slip (up to \(\sim 0.10u_{tip}\)) was also observed in the \(r\) and \(\theta\) directions in small regions confined to the impeller discharge and below the impeller (data not shown). For all particle size fractions, there is a gradual but significant reduction in the slip velocity with increasing \(X\) in the upper parts of the vessel, but below the impeller and in the impeller discharge region substantial increases are observed.

### 3.3 Spatial solids distribution

In addition to location and velocity information, PEPT allows the particle occupancy distribution within the vessel to be determined. By dividing the vessel volume into a grid of \(n_c\) cells, occupancy can be obtained by calculating the fraction of the total experimental time, \(t_\infty\), spent by the tracer in each cell during the experiment. Such a definition establishes a mathematical identity between occupancy and probability of presence of the particle tracer, but undesirably makes occupancy highly dependent on the density of the grid so that as the number of cells increases occupancy tends to zero. If the cells are chosen to have equal volume, however, this problem is circumvented by using the ‘ergodic time’ defined as \(t_E = t_\infty / n_c\), instead of the total experimental time \(t_\infty\).

The ergodic time represents the time that the tracer would spend in any cell if the flow were ergodic, an asymptotic status in which the flow tracer has exactly an equal probability of presence everywhere within the system. Thus, the local occupancy, \(O_j\), can be defined as:

\[
O_j = \frac{\Delta t_j}{t_E}
\]

where \(\Delta t_j\) is the time that the tracer of the \(j\)-th component (solid or liquid) of the mixture spends inside a given cell. For the polydisperse systems examined in this work, the index \(j\) assumes values 1 to 6, where \(j = 1\) to 5 denotes the solid components, and \(j = 6\) denotes the liquid phase also designated by the symbol \(L\).
On the basis of such a definition, it can be shown that, in a multiphase system, occupancy, $O_j$, of component $j$ is directly related to its local phase volume concentration, $c_j$, via the relationship:

$$O_j = \frac{c_j}{C_j}$$  \hspace{1cm} (3)$$

where $C_j$ is the mean phase volume concentration of component $j$ in the vessel. Full details of the mathematical treatment can be found in our previous papers (Guida et al., 2010a, 2011).

Thus, using Eq. (3) the spatial distribution of the local concentration of each flow component was fully resolved in the polydisperse suspensions investigated. Azimuthally-averaged solid concentration maps are presented in Fig. 6. The highest local concentrations are several times greater than the mean solid concentration and are observed in a clearly defined mound of solids accumulating beneath the agitator as well as in small fillets in the corners. This effect is more pronounced at the lower solid loadings for which solids distribution in the upper parts of the vessel is poor. However, this maldistribution of solids improves substantially as $X$ increases for all particle fractions in the mixture, tending gradually towards a much more uniform state at $X = 40$ wt%. This seems to suggest, perhaps counter-intuitively, that in the ‘just suspended’ regime more concentrated suspensions tend towards a homogeneous state by virtue of their increased solids loading. The smaller particles, however, are significantly better dispersed than the larger ones at all the values of $X$ used here.

The local particle size distributions in the vessel are presented in the maps shown Fig. 7. The azimuthally- and vertically-averaged particle size distributions are shown above the local particle distribution maps. There are considerable concentration gradients in the vertical direction, an effect which is clearly captured in the radially-averaged particle size distributions shown to the right of the maps. Such an effect, as pointed out above, is much more pronounced at lower $X$ values. Horizontally, however, the vertically-averaged distributions above the maps are fairly flat except close to the wall in the upper parts of the vessel where the smaller particles seem to predominate somewhat.

Locally, at lower mean concentrations, the larger particles tend to represent the higher fractions beneath the impeller, but the local size distributions become increasingly more uniform as $X$ increases. However, the mass of solids is clearly still not quite homogeneously distributed in the volume of the vessel even at $X = 40$ wt%. It is well known that, in general, impeller speeds substantially higher than $N_{js}$ and, consequently, much higher power inputs are required to achieve a state of homogeneous suspension. Such an increase in power demand is even larger for (denser) faster settling particles as is the case here, and the power consumption could be several folds its value at $N_{js}$ (Oldshue, 1983).

The uniformity of a given component in the suspension can be quantified using the uniformity index, $\xi$, introduced in our recent works (Guida et al., 2009, 2010a), thus:
\[ \xi = \frac{1}{\sigma^2 + 1} = \frac{1}{n_c \sum_{i=1}^{n_c} \left( \frac{c_i - C}{C} \right)^2 + 1} \]  

(4)

where \( n_c \), as defined above, is the total number of cells in the PEPT measurement grid and \( i \) is the cell number. The uniformity of the suspension is at its minimum when \( \xi \rightarrow 0 \) (condition achieved for a theoretically infinite variance, \( \sigma^2 \), of the normalised local concentration); and the solids are uniformly distributed within the vessel volume when \( \xi = 1 \), i.e., the local solid concentration is equal to the average concentration everywhere in the vessel (\( \sigma^2 = 0 \)).

The effects observed in the particle-size-distribution maps displayed in Fig. 7, are clearly captured in Fig. 8 by the variations of \( \xi \). The uniformity index is considerably higher for the smaller particles and, for all particle sizes, it increases considerably as a function of \( X \), but the rate of increase reduces significantly at higher concentrations. The gap in homogeneity between small and large particles reduces gradually with increasing concentration from 70% at \( X = 5 \) wt% down to 12% at \( X = 40 \) wt%. For the smallest particles, \( \xi \) reaches 0.9 at \( X = 40 \) wt%.

### 3.4 Mass balance and mass continuity

The 'pointwise' measurements obtained from PEPT enabled the solids mass balance throughout the vessel to be accurately verified, i.e. the measured local values, \( c_i \), of the five solid volume concentrations should balance the experimental parameter \( C_j = V_j / V_T \), so that:

\[ \frac{1}{n_c} \sum_{i=1}^{n_c} c_i = C_j \]  

(5)

where \( V_j \) is the volume of component \( j \) in the vessel and \( V_T \) is the total suspension volume. The identity symbol '=' appears in Eq. (5) because the local concentration is obtained from Eq. (3), through the measurement of local values of the occupancy \( O_j \) whose average value is by definition identically equal to 1. The error on the mass balance for all components was ~ 1%.

Another important tool for checking the accuracy and reliability of flow data is mass continuity. The net mass flux through a volume bounded by a closed surface \( S \) should be zero, thus:

\[ \sum_{i} \mathbf{u}^{(i)} \cdot \Delta \mathbf{S} = 0 \]  

(6)

Calculations can be made by considering a closed cylindrical surface \( S \) with the same vertical axis, base and diameter as the tank but of a shorter height. Because the vector \( \mathbf{u} \) is zero over the external surface of the vessel or is parallel to it, the term \( \mathbf{u} \cdot \Delta \mathbf{S} \) is zero everywhere except in the horizontal plane across the tank, so that \( S \) can
be reduced to such a plane, \( S_h \), and Eq. (6) is reduced to a zero average of axial velocities across the horizontal plane considered.

Alternatively, a similar situation can be envisaged where \( S \) is a closed cylindrical envelope with the same vertical axis and height as the tank but with a smaller diameter; Eq. (6) then becomes a zero average of radial velocities across the lateral surface, \( S_r \), of the considered cylindrical volume. These two types of surfaces \( S_h \) and \( S_r \) were numerically introduced in the vessel and the continuity test was applied to all the sets of data collected in this study. The results are summarised in Fig. 9, showing a very good verification of the mass continuity as the velocity average across the given surface in each case is close to zero, generally less than 0.02\( u_{tip} \). The slight discrepancy is probably due to systematics in the intrinsic measurement error or in artefacts introduced by data processing, e.g. uncertainty associated with the location of the tracer, grid size for occupancy, smoothing of position data during velocity calculations etc.

**Conclusions**

The Lagrangian technique of PEPT has been successfully extended, for the first time, to the analysis of complex highly-concentrated solid-liquid flows of polydisperse suspensions consisting of five particle size fractions. Thus, it has been possible to determine for each component of the two-phase flow detailed distributions of the local velocity, spatial distribution, and time-averaged slip velocity, and to quantitatively assess the homogeneity of suspension of each particle size fraction relative to the other size fractions. The accuracy of the 3-D Lagrangian measurements has been ascertained through multi-component mass balance and continuity verifications throughout the vessel.

The measured local velocity distributions showed that the Reynolds number similarity holds for all components of the two-phase flow under all conditions investigated. The local time-averaged slip velocity varies widely within the vessel, is generally very different from the particle terminal settling velocity, and can reach considerable values greater than 0.10\( u_{tip} \). At the lower solid concentrations, the larger particles with more inertia display considerably more inter-phase slip than the smaller ones. This effect, however, reduces substantially with an increase in solid concentration and impeller speed as particle-particle interactions become important.

The uniformity of suspension improves as a function of mean solid concentration when suspensions are considered at the same ‘just suspended’ hydrodynamic regime corresponding to the \( N_{js} \) speed. Smaller particles exhibit considerably better suspension homogeneity. This study has shown that PEPT is a powerful non-intrusive Lagrangian technique which is ideally suited to the study of dense solid-liquid systems of complex composition. The technique can, therefore, be extended to the study of more challenging systems with wider particle size distributions, more complex rheologies and probably even higher solid concentrations.
Acknowledgements

Li Liu’s PhD research was funded jointly by a China Scholarship and an Elite scholarship from the University of Birmingham. We are grateful to Prof. D.J. Parker and J.F. Gargiuli of the Positron Imaging Centre at The University of Birmingham for their assistance with the PEPT experiments.

References


Figure and Table captions

Figure 1. Experimental set-up for studying the mixing of polydisperse suspensions by PEPT: 1 tank; 2 shaft; 3 baffle; 4 PBT; 5 γ-ray detectors; 6 PEPT particle tracer; 7 polydisperse glass particles.

Figure 2. Illustration of the cylindrical grid consisting of equal volume cells used for the analysis of the Lagrangian data, and a grid cell with multiple PEPT tracer visits.

Figure 3. Azimuthally-averaged local velocity maps of liquid and solid phase components at different solid concentrations: greyscale contours represent the total 3-D velocity \( u \); vectors represent the radial-axial 2-D velocity \( u_{rz} \).

Figure 4. Demonstration of Reynolds number similarity – Azimuthally-averaged radial distributions of the normalized velocity components for three mixture components: — \( X = 5 \) wt%; --- \( X = 10 \) wt%; ···· \( X = 20 \) wt%; ··· \( X = 40 \) wt%; \( Re_{imp} = 1.5 \times 10^5 \) – \( 2.4 \times 10^5 \) (see Table 1).

Figure 5. Azimuthally-averaged maps of local time-averaged slip velocity at different solid concentrations: greyscale contours represent the total 3-D velocity \( u \); vectors represent the radial-axial 2-D velocity \( u_{rz} \).

Figure 6. Azimuthally-averaged maps of local solid volume concentration at different solid concentrations.

Figure 7. Maps of azimuthally-averaged local particle size distribution at different solid concentrations: particle size fractions from left to right, \( d_p \sim 1.0 \) mm; \( d_p \sim 1.5 \) mm; \( d_p \sim 2.0 \) mm; \( d_p \sim 2.5 \) mm; \( d_p \sim 3.0 \) mm.

Figure 8. Variation of uniformity index for all solid components at varying \( X \).

Figure 9. Normalised radial and axial velocities averaged on surface \( S_r \) (of diameter 0.53\( T \)) and \( S_z \) (0.3\( H \) off the base), respectively: ○ 1.0 mm; ○ 1.5 mm; Δ 2.0 mm; × 2.5 mm; ● 3.0 mm; ■ liquid.

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<table>
<thead>
<tr>
<th>Concentration (wt%)</th>
<th>Solids Volume %</th>
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<tbody>
<tr>
<td>5</td>
<td>2.4</td>
</tr>
<tr>
<td>10</td>
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<tr>
<td>20</td>
<td>10.4</td>
</tr>
<tr>
<td>40</td>
<td>23.6</td>
</tr>
</tbody>
</table>

Radial position, $r/R$ (-) 
Axial position, $z/H$ (-) 

| $|u|/u_{tip}$ |
|------------|
| 0          |
| 0.5        |
| 1          |

$|u|/u_{tip}$ values for different concentration: 
- Liquid $d_p \sim 1.0$ mm 
- $d_p \sim 1.5$ mm 
- $d_p \sim 2.0$ mm 
- $d_p \sim 2.5$ mm 
- $d_p \sim 3.0$ mm
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<table>
<thead>
<tr>
<th>C (vol%)</th>
<th>X (wt%)</th>
<th>$N_{in}$ (rpm)</th>
<th>Re&lt;sub&gt;imp&lt;/sub&gt; (•)</th>
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<tr>
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<td>$1.5 \times 10^5$</td>
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<td>4.9</td>
<td>10</td>
<td>450</td>
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<td>20</td>
<td>510</td>
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<tr>
<td>23.6</td>
<td>40</td>
<td>612</td>
<td>$2.4 \times 10^5$</td>
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</table>
Highlights

- The technique of multi-component positron emission particle tracking is used
- New unique data are reported for the mixing of dense complex polydisperse suspensions
- 3-D local velocities and spatial distribution of each phase component are measured
- Maps of local particle size distribution and particle slip velocity are reported
- The uniformity of dispersion of each flow component is quantified