Chapter 5

Pebble-Bed Simulation¹

5.1 Introduction

5.1.1 Background

A worldwide effort is underway to develop more economical, efficient, proliferation resistant, and safer nuclear power [4]. A promising Generation IV reactor design is the uranium-based, graphite moderated, helium-cooled very high temperature reactor [49], which offers meltdown-proof passive safety, convenient long-term waste storage, modular construction, and a means of nuclear-assisted hydrogen production and desalination. In one embodiment, uranium dioxide is contained in microspheres dispersed in spherical graphite pebbles, the size of billiard balls, which are very slowly cycled through the core in a dense granular flow [97, 130]. Control rods are inserted in graphite bricks of the core vessel, so there are no obstacles to pebble flow.

The pebble-bed reactor (PBR) concept, which originated in Germany in the 1950s, is being revisited by several countries, notably China [110] (HTR-10 [58]) and South Africa [97] (PBMR [5]), which plan large-scale deployment. In the United States, the Modular Pebble Bed Reactor (MPBR) [130, 2] is a candidate for the Next Generation Nuclear Plant of the Department of Energy. A notable feature of MPBR (also

¹This chapter is based on reference [113], Analysis of Granular Flow in a Pebble-Bed Nuclear Reactor, published in Physical Review E in 2006. See http://pre.aps.org/ for more details.

present in the original South African design) is the introduction of graphite moderator pebbles, identical to the fuel pebbles but without the uranium microspheres. The moderator pebbles form a dynamic central column, which serves to flatten the neutron flux across the annular fuel region without placing any fixed structures inside the core vessel. The annular fuel region increases the power output and efficiency, while preserving passive safety. In the bidisperse MPBR, the moderator pebbles are smaller to reduce the permeability of the central column and thus focus helium gas on the outer fuel annulus. The continuous refueling process is a major advantage of pebble-bed reactors over other core designs, which typically require shutting down for a costly dismantling and reconstruction. The random cycling of pebbles through a flowing core also greatly improves the uniformity of fuel burnup.

In spite of these advantages, however, the dynamic core of a PBR is also a cause for concern among designers and regulators, since the basic physics of dense granular flow is not fully understood. Indeed, no reliable continuum model is available to predict the mean velocity in silos of different shapes [27], although the empirical Kinematic Model [90, 95, 94] provides a reasonable fit near the orifice in a wide silo [137, 84, 114, 28]. A complete statistical theory of dense granular flow is still lacking. The classical kinetic theory of gases has been successfully applied to dilute granular flows [115, 61, 108], in spite of problems with inelastic collisions [64], but it clearly breaks down in dense flows with long-lasting, frictional contacts [93, 28], as in pebble-bed reactors. Plasticity theories from soil mechanics might seem more appropriate [94], but they cannot describe flows in silos of arbitrary shape and often lead to violent instabilities [117, 106].

For now, experiments provide important, although limited, information about dense granular flows. Many experiments have been done on drainage flows in quasi-2D silos where particles are tracked accurately at a transparent wall [84, 83, 114, 28, 27]. Experimental studies of more realistic geometries for PBR have mostly focused on the porosity distribution of static packings of spheres [50, 119], which affects helium gas flow through the core [30, 141, 143].

As a first attempt to observe pebble dynamics experimentally in a reactor model,

the slow flow of plastic beads has recently been studied in 1:10 scale models of MPBR in two different ways [63]: the trajectories of colored pebbles were recorded (by hand) along a Plexiglas wall in a half-core model, and a single radioactive tracer pebble in the bulk was tracked in three dimensions in a full-core model. Very slow flow was achieved using a screw mechanism at the orifice to approximate the mean exit rate of one pebble per minute in MPBR. These experiments demonstrate the feasibility of the dynamic central column and confirm that pebbles diffuse less than one diameter away from streamlines of the mean flow. However, it is important to gain a more detailed understanding of pebble flow in the entire core to reliably predict reactor power output, fuel efficiency, power peaking, accident scenarios using existing nuclear engineering codes [131, 51].

5.1.2 Discrete-Element Simulations

Simulations are ideally suited to provide complete, three-dimensional information in a granular flow. Some simulations of the static random packing of fuel pebbles in a PBR core have been reported [38, 103], but in the last few years, large-scale, parallel computing technology has advanced to the stage where it is now possible to carry out simulations of continuous pebble flow in a full-sized reactor geometry using the Discrete Element Method (DEM). In this chapter, we present DEM simulations which address various outstanding issues in reactor design, such as the sharpness of the interface between fuel and moderator pebbles (in both monodisperse and bidisperse cores), the horizontal diffusion of the pebbles, the geometry dependence of the mean streamlines, the porosity distribution, wall effects, and the distribution of "residence times" for pebbles starting at a given height before exiting the core.

Our simulations are based on the MPBR geometry [130, 2], consisting of spherical pebbles with diameter d = 6cm in a cylindrical container approximately 10m high and 3.5m across. In this design there is a central column of moderating reflector pebbles, surrounded by an annulus of fuel pebbles. The two pebble types are physically identical except that the fuel pebbles contain sand-sized uranium fuel particles. Particles are continuously cycled, so that those exiting the container are reintroduced at the top of the packing. In order to efficiently maintain the central column, a cylindrical guide ring of radius $r_{\rm in} = 14.5d$ extends into the packing to z = 140d. Reflector pebbles are poured inside, while fuel pebbles are poured outside, and the guide ring ensures that two types do not mix together at the surface. Figure 5-1 shows the two main geometries that were considered; for much of this analysis, we have concentrated on the case when the exit funnel is sloped at thirty degrees, but since this angle can have a large effect on the pebble flow, we also consider the case of the when the funnel is sloped at sixty degrees. In both cases the radius of the opening at the bottom of the funnel is $r_{\rm exit} = 5d$.

In MPBR, as in most pebble-bed reactors, the drainage process takes place extremely slowly. Pebbles are individually removed from the base of the reactor using a screw mechanism, at a typical rate of one pebble per minute, and the mean residence time of a pebble is 77 days. Carrying out a DEM simulation at this flow rate would make it infeasible to collect enough meaningful data. However, the results of the previous chapters, and the experimental work of Choi [28], show that for granular drainage, altering the overall flow rate does not alter the geometry of the flow profile – the flow velocities are scaled by a constant factor. Furthermore, geometric properties of the flow, such as particle diffusion, are unaffected by the overall flow rate. We therefore chose to study a faster flow regime in which pebbles drain from the reactor exit pipe under gravity. Our results can be related directly to the reactor design by rescaling the time by an appropriate factor.

As well as the two full-scale simulations described above, we also considered a half-size geometry in order to investigate how various alterations in the makeup of the reactor would affect the flow. In particular, we examined a series of bidisperse simulations, in which the diameter of moderator particles in the central column was reduced. As explained in section 5.8, this has the effect of reducing the gas permeability of the central column, thus focusing the helium coolant flow on the hottest region of the reactor core, in and around the fuel annulus. The purpose of the simulations is to test the feasibility of the bidisperse PBR concept, as a function of the size ratio of moderator and fuel pebbles, with regard to the granular flow. It is not

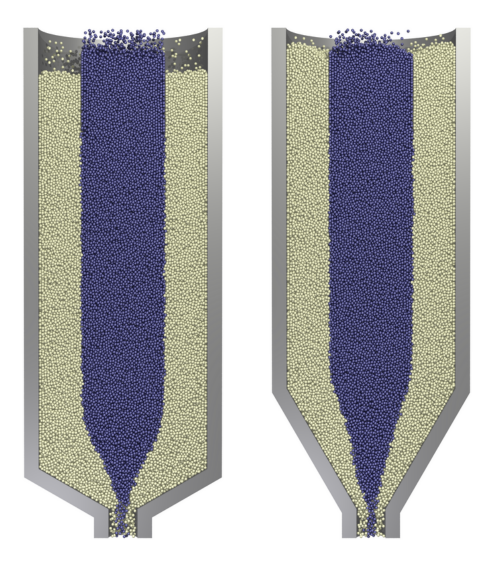


Figure 5-1: Snapshots of vertical cross-sections of the simulations for the two geometries considered in this report. We make use of a cylindrical coordinate system (r, θ, z) where z = 0 at the orifice. At the base of the container there is a small exit pipe of radius $r_{\text{exit}} = 5d$ that extends upwards to z = 10d. This connects to a conical funnel region, which has slope thirty degrees (left) or sixty degrees (right). The conical wall connects to a cylindrical wall of radius $r_{\text{out}} = 29d$, at z = 23.86d and z = 51.57dfor the thirty and sixty degree reactor geometries respectively. Particles are poured into the container up to a height of approximately z = 160d. A cylindrical wall at $r_{\text{in}} = 14.5d$ extends down into the packing to a height of z = 140d to keep the two types of pebbles mixing at the surface.

clear *a priori* under what conditions the dynamic column will remain stable with little interdiffusion of moderator and graphite pebbles.

To study this issue, we made a sequence of three runs using a half-size reactor geometry. (The smaller core size is needed since the number of smaller pebbles increases as the inverse cube of the diameter ratio.) The geometry is similar to that used above, except that the radius of the cylindrical container is decreased to 15*d*, with the guide ring at $r_{\rm in} = 7.5d$. The radius of the exit pipe is decreased to $r_{\rm exit} = 4d$. In the experiments, we keep the diameter of the fuel pebbles fixed at *d*, and use *d*, 0.8*d*, and 0.5*d* for the diameters of the moderator pebbles. The same geometry was also used to study the effect of wall friction, by making an additional run with the particle/wall friction coefficient $\mu_w = 0$.

This chapter is organized as follows. In section 5.2, we discuss the simulation technique that was used and briefly describe its implementation. This is followed with some basic analysis of the velocity profiles and a comparison to the Kinematic Model in section 5.3. We study diffusion around streamlines in section 5.4 and the distribution of porosity and local ordering in section 5.5. Next, in section 5.6 we examine the residence-time distribution of pebbles in the reactor, which is related to fuel burnup, and in section 5.7 we show that wall friction plays an important role. In section 5.8 we analyze the bidisperse PBR concept with half-size reactor simulations for a range of pebble-diameter ratios, focusing on the mean flow, diffusion, and mixing. We conclude in section 5.9 by summarizing implications of our study for reactor design and the basic physics of granular flow.

5.2 Models and Methods

Unlike all other simulations in this thesis, the DEM simulations presented here were carried out at Sandia Labs, since the MIT AMCL was too small to handle the number of particles used in the fully three dimensional system. For the monodispersed simulation, the spheres have diameter d = 6 cm, mass m = 210 g and interparticle friction coefficient $\mu = 0.7$, flowing under the influence of gravity g = 9.81 ms⁻². For the bidispersed systems, the moderator particles have diameter 0.8*d* or 0.5*d*. The particle-wall friction coefficient $\mu_w = 0.7$ except in one case where we model a frictionless wall, $\mu_w = 0.0$.

The initial configurations are made by extending the inner cylinder from 140d to the bottom of the container, adding a wall at the bottom of the container to stop particles from draining, and pouring in moderator pebbles into the inner cylinder and fuel pebbles between the inner and outer cylinders until the reactor was loaded. The bottom wall is then removed, the inner cylinder is raised to 140d, and particles are allowed to drain out of the container. As noted above, particles are recycled with moderator particles reinserted within the inner cylinder, and fuel particles between the inner and outer cylinders. All results presented here are after all the particles have cycled through the reactor at least once. The number of moderator and fuel particles was adjusted slightly from the initial filling so that the level at the top of the reactor is approximately equal. For the full scale simulation with a thirty degree outlet, the total number of pebbles is 440,000 with 105,011 moderator pebbles and 334,989 fuel pebbles, while for the sixty degree outlet, the total number of pebbles is 406,405 with 97,463 moderator and 308,942 fuel pebbles. For the former case, a million steps took approximately 13 hours on 60 processors on Sandia's Intel Xenon cluster.

For the bidispersed simulations the total number of pebbles is 130,044, 160,423, and 337,715 for the diameter of the moderator particles equal to d, 0.8d and 0.5d respectively. As the diameter of the moderator pebbles is decreased the number of particles required rapidly increases, since it scales according to the inverse of the diameter cubed.

A snapshot of all the particle positions is recorded every $5\tau = 0.39$ s. For the thirty degree reactor geometry we collected 1,087 successive snapshots, totaling 24.9Gb of data, while for the sixty degree reactor geometry, we collected 881 successive snapshots, totaling 18.7Gb of data. A variety of analysis codes written in Perl and C++ were used to sequentially parse the snapshot files to investigate different aspects of the flow. We also created extended data sets, with an additional 440 snapshots for the thirty degree geometry, and 368 snapshots for the sixty degree geometry, for examining long residence times in section 5.6.

5.3 Mean-Velocity Profiles

5.3.1 Simulation Results

Since we have a massive amount of precise data about the positions of the pebbles, it is possible to reconstruct the mean flow in the reactor with great accuracy. However care must be taken when calculating velocity profiles to ensure the highest accuracy. Initial studies of the data showed that crystallization effects near the wall can create features in the velocity profile at a sub-particle level, and we therefore chose a method that could resolve this.

By exploiting the axial symmetry of the system, one only needs to find the velocity profile as a function of r and z. The container is divided into bins and the mean velocity is determined within each. A particle which is at \mathbf{x}_n at the *n*th timestep and at \mathbf{x}_{n+1} at the (n + 1)th timestep, makes a velocity contribution of $(\mathbf{x}_{n+1} - \mathbf{x}_n)/\Delta t$ in the bin which contains its midpoint, $(\mathbf{x}_{n+1} + \mathbf{x}_n)/2$.

In the z direction, we divide the container into strips 1d across. However, in the r direction we take an alternative approach. Since the number of pebbles between a radius of r and $r + \Delta r$ is proportional to $r\Delta r$, dividing the container into bins of a fixed width is unsatisfactory, as the amount of data in bins with high r would be disproportionately large. We therefore introduce a new coordinate $s = r^2$. The coordinate s covers the range $0 < s < r_{out}^2$, and we divide the container into regions that are equally spaced in s, of width $1d^2$. The number of pebbles in each bin is therefore roughly equal, allowing for accurate averaging in the bulk and high resolution at the boundary.

This result yields extremely accurate velocity profiles in the cylindrical region of the tank. However, it fails to capture crystallization effects in the conical region: since the particles are aligned with the slope of the walls are averaged over a strip in

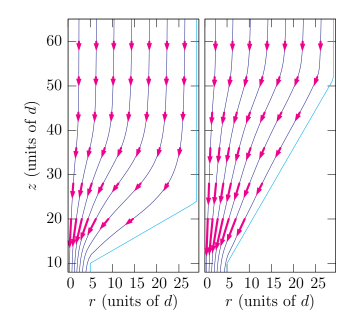


Figure 5-2: Computed streamlines of the mean flow in the 30° (left) and 60° reactor geometries. Arrows are proportional to the velocity vectors in selected horizontal slices.

z of width 1d, any effects are smeared out across several bins. We therefore scaled the radial coordinate to what it would be if the particle was in the center of the strip. Specifically, if the radius of the container is given by R(z), a particle at (r_n, z_n) is recorded as having radial coordinate $r_n R(z)/R(z_n)$. In the cylindrical region of the tank this has no effect, while in the conical region, it effectively creates trapezoidshaped bins from which it is easy to see crystallization effects which are aligned with the wall.

The streamlines of the mean flow are shown in Fig. 5-2 in the two geometries. Streamlines are computed by Lagrangian integration of the DEM velocity field, starting from points at a given height, equally spaced in radius. In each geometry, there is a transition from a nonuniform converging flow in the lower funnel region to a nearly uniform plug flow in the upper cylindrical region, consistent with the standard engineering picture of silo drainage [94]. In the wider funnel, there is a region of much slower flow near the sharp corner at the upper edge of the funnel. Our results for both geometries are quite consistent with particle-tracking data for quasi-2D silos of similar shapes [27] and half-cylinder models of the MPBR core [63], which provides

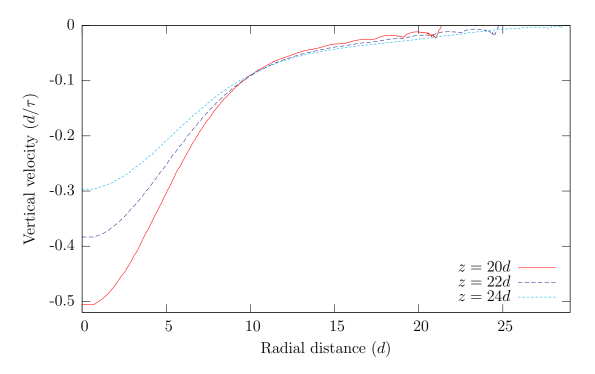


Figure 5-3: Velocity profiles for the thirty degree reactor geometry for several low cross-sections.

an important validation of our simulations.

We now look more closely at horizontal slices of the velocity field. Figure 5-3 shows several velocity profiles for the thirty degree case in the narrowing section of the container. As expected, we see a widening of the velocity profile as z increases. We can also see lattice effects, spaced at $\sqrt{3}d$ apart, due to to particles crystallizing on the conical wall section.

Figure 5-4 shows similar plots for several heights in the upper region of the container. At these heights, the velocity profile is roughly uniform across the container. However a boundary layer of slower velocities, several particle diameters wide, still persists. The average velocities of particles touching the boundary is between one half and two thirds that of particles in the bulk; it is expected that this behavior is very dependent on particle-wall friction; this issue is studied in more detail in section 5.7.

High in the container, results for the sixty degree geometry are very similar to the thirty degree case (and thus are not shown). However, as would be expected, a

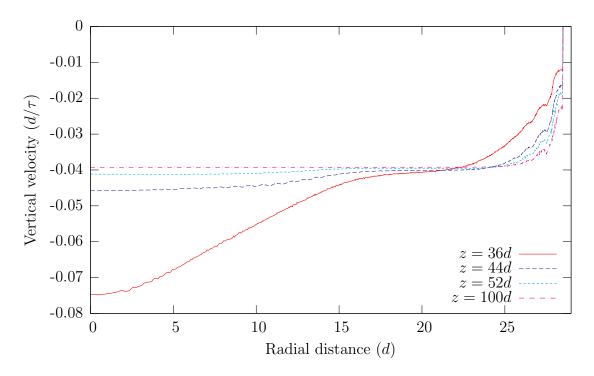


Figure 5-4: Velocity profiles for the thirty degree reactor geometry for several high cross-sections.

significantly different crossover from parabolic flow to plug-like flow in the lower part of the tank is observed, as shown in figure 5-6.

5.3.2 Comparison with the Kinematic Model

In section 2.3 we introduced the Kinematic Model, which is perhaps the only continuum theory available for the mean flow profile in a slowly draining silo. The vertical velocity v satisfies

$$\frac{\partial v}{\partial z} = b \nabla_{\perp}^2 v, \tag{5.1}$$

where the vertical coordinate z acts like "time". Boundary conditions on Eq. (5.1) require no normal velocity component at the container walls, except at the orifice, where v is specified (effectively an "initial condition"). As described in Appendix B, this boundary-value problem can be accurately solved using a standard Crank-Nicholson scheme for the diffusion equation.

Consistent with a recent experimental study of quasi-2D silos [27], we find reason-

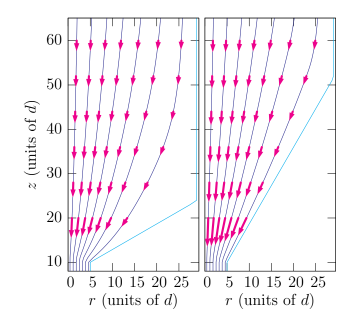


Figure 5-5: Streamlines of the mean flow in the 30° (left) and 60° reactor geometries for the numerical solution of the Kinematic Model. Arrows are proportional to the velocity vectors in selected horizontal slices.

able agreement between the Kinematic Model predictions and the DEM flow profiles, but the effect of the container geometry is not fully captured. In the converging flow of the funnel region, the streamlines are roughly parabolic, as predicted by the Kinematic Model and found in many experiments [137, 84, 114, 28, 27]. For that region, it is possible to choose a single value (b = 3d) to achieve an acceptable fit to the DEM flow profiles for both the 30° and 60° funnel geometries, as shown in figure 5-6.

In spite of the reasonable overall fit, the Kinematic Model has some problems describing the DEM results. It fails to describe the several particle thick boundary layer of slower velocities seen in the DEM data. In the original model, b depends only on the properties of the granular material, but we find that it seems to depend on the geometry; the best fit to the 30° DEM data is $b \approx 2.5d$, while the best fit for the 60° DEM data is $b \approx 3.0d$. Such discrepancies may partly be due to the boundary layers, since in the lower section of the container the conical walls may have an appreciable effect on the majority of the flow. We also find that the Kinematic Model fails to capture the rapid transition from converging flow to plug flow seen in the DEM data. This is shown clearly by comparing the streamlines for the Kinematic Model in figure

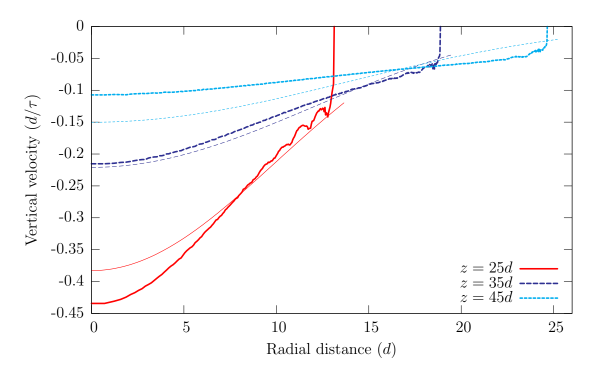


Figure 5-6: Velocity profiles for the 60° reactor geometry (heavy lines), with a comparison to the Kinematic Model for b = 3d (thin lines).

5-5 with those for DEM. Streamlines for the Kinematic Model are roughly parabolic, and no single value of b can capture the rapid change from downward streamlines to converging streamlines seen in DEM.

The difficulty in precisely determining b is also a common theme in experiments, although recent data suggests that a nonlinear diffusion length may improve the fit [27]. Perhaps a more fundamental problem with the Kinematic Model is that it cannot easily describe the rapid crossover from parabolic converging flow to uniform plug flow seen in both geometries our DEM simulations; we will return to this issue in section 5.5.

5.4 Diffusion and Mixing

Nuclear engineering codes for PBR core neutronics typically assume that pebbles flow in a smooth laminar manner along streamlines, with very little lateral diffusion [131, 51]. Were such significant diffusion to occur across streamlines, it could alter the

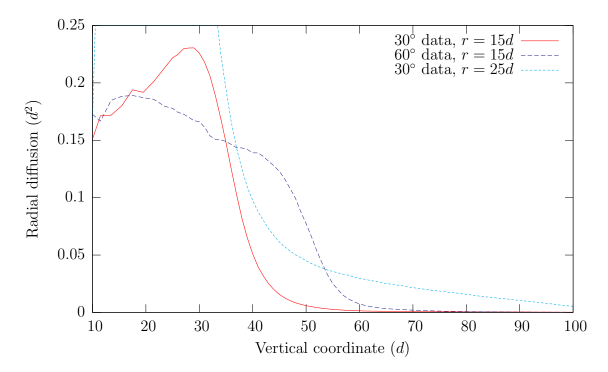


Figure 5-7: Radial diffusion of particles about streamlines of the mean flow as a function of height, z, in both reactor geometries for pebbles starting at z = 110d in an annulus of radius r = 15d, at the edge of the dynamic central column in MPBR. For the 30° geometry, we also show data for pebbles near the wall at r = 25d.

core composition in unexpected ways. In the MPBR design with a dynamic central column [2], diffusion leads to the unwanted mixing of graphite pebbles from the central reflector column with fuel pebbles from the outer annulus, so it must be quantified.

Particle-tracking experiments on quasi-2D silos [28] and half-cylinder MPBR models [63] have demonstrated very little pebble diffusion in slow, dense flows, but the observations were made near transparent walls, which could affect the flow, e.g. due to ordering (see below). Three-dimensional tracking of a radioactive tracer in a cylindrical MPBR model has also shown very little diffusion, at the scale of a single pebble diameter for the duration of the flow [63]. Here, we take advantage of the complete information on pebble positions in our DEM simulations to study core diffusion and mixing with great accuracy.

We collected extensive statistics on how much pebbles deviate from the mean-flow streamlines during drainage. Consistent with theoretical concepts [17], experiments have demonstrated that the dynamics are strongly governed by the packing geometry, so that diffusion can most accurately be described by looking at the mean-squared horizontal displacement away from the streamline, as a function of the distance dropped by the pebble (not time, as in molecular diffusion), regardless of the flow rate. Motivated by the importance of quantifying mixing at the fuel/moderator interface in the dynamic central column of MPBR, we focus on tracking pebbles passing through z = 110d with |r - 15d| < 0.16d. The variance of the r coordinate of the particles as they fall to different heights in z can be calculated. From this, we can determine the amount of radial diffusion, defined as the increase in the variance of r of the tracked particles from the variance at the initial height.

The diffusion data for both reactor geometries is shown in figure 5-7. We see that for large values of z in the cylindrical part of the container, the pebbles undergo essentially no diffusion; this is to be expected, since we have seen that in this area the packing is essentially plug-like, and particles are locked in position with their neighbors. However for lower values of z the amount of radial spreading begins to increase, as the particles experience some rearrangement in the region corresponding to converging flow. Note however that the scale of this mixing is very small, and is much less than a pebble diameter. The height where the amount of diffusion begins to increase is approximately z = 35d in the 30° geometry and z = 50d in the 60° geometry. In the 30° geometry, this transition is significantly above the height of the interface between conical and cylindrical walls, while in the 60° geometry, the transition is almost level with the interface. This suggests that while the container geometry may play a role in diffusion and velocity profiles, it is a lower-order effect. For very small values of z, there is a decrease in the variance of the radial coordinate, since the pebbles must converge on the orifice as they exit the container.

We applied a similar analysis for different initial values of r, and found very similar results over the range 0 < r < 25d. However, for particles close to the container boundary, very different behavior is observed, as shown by the third line in figure 5-7 for particles with |r - 25d| < 0.10d. In this region, the particles undergo rearrangement, and this causes a (piecewise) linear increase in the mean-squared displacement with distance dropped, which corresponds to a constant local diffusion length. There is also evidence of a sharp transition in the boundary-layer diffusion length, which increases significantly as pebbles pass the corner into the convergingflow region of the funnel.

5.5 Packing Statistics

5.5.1 Pebble Volume Fraction

Pebble-bed experiments [50, 119] and simulations [38, 103] of static sphere packings in cylinders have revealed that there are local variations in porosity near walls, at the scale of several pebble diameters, but there has been no such study of flowing packings, averaging over dynamic configurations. Similar findings would have important implications for helium flow in the core, since the local gas permeability is related to the porosity [30, 141, 143].

First, we study the distribution of local volume fraction (% of volume occupied by pebbles) throughout the container, averaged in time. (The porosity is one minus the volume fraction.) We make use of the Voronoi algorithm discussed in the previous chapter to examine the packing fraction in vertical cross sections through the containers. Figure 5-8 shows density snapshots for cross sections through the thirty degree and sixty degree reactor geometries, based on computing the local density at a particular point by averaging over the Voronoi densities of particles within a radius of 2.2d. Figure 5-9 shows density plots over the entire flow of the data, but using a smaller averaging radius of 0.8d. Many interesting features are visible, which corroborate our other results. High in the center of the container, we see that the local packing fraction is mostly close to 63%, suggesting that the plug-like region is in a nearly jammed and rigid state. This is consistent with our earlier reactor data showing nearly uniform plug flow with no significant diffusion or mixing.

We also observe two annular lines of lower density propagating down from the guide ring, which form due to wall effects on the guide ring itself (see below) and are advected downward. The fact that these subtle artifacts of the guide-ring constraints

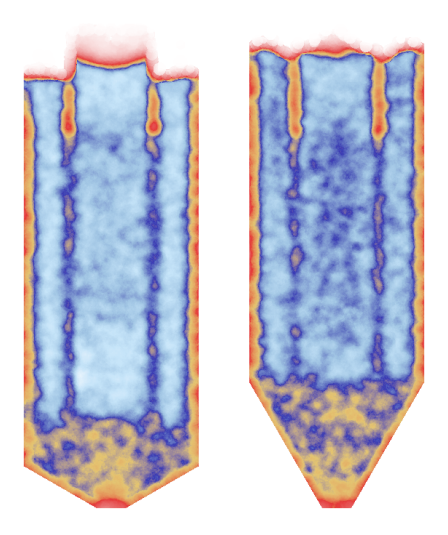


Figure 5-8: Plots of local volume fraction (1 - porosity) in a vertical cross section for the thirty degree reactor geometry (left) and the sixty degree reactor geometry (right), calculated using a Voronoi cell method. Volume fractions of 50%, 57%, 60%, and 63% are shown using the colors of red, yellow, dark blue, and cyan respectively. Colors are smoothly graded between these four values to show intermediate volume fractions. High in the bulk of the container, the packing fraction is approximately 63%, apart from in a small region of lower density at $r_{\rm in} = 14.5d$, corresponding to packing defects introduced by the guide ring. In both geometries a sharp reduction in density is observed in a region above the orifice, where particles in the parabolic flow region are forced to undergo local rearrangements.

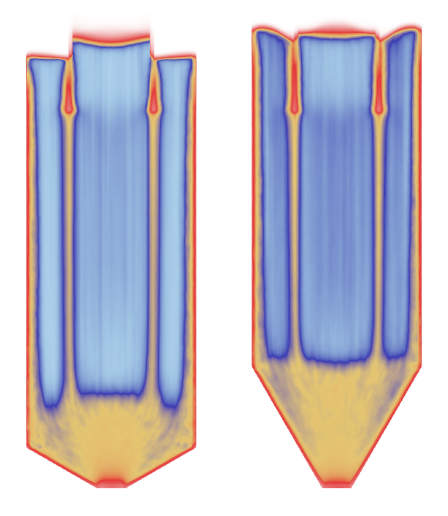


Figure 5-9: Time-averaged plots of the local volume fraction, using the same color scheme as figure 5-8.

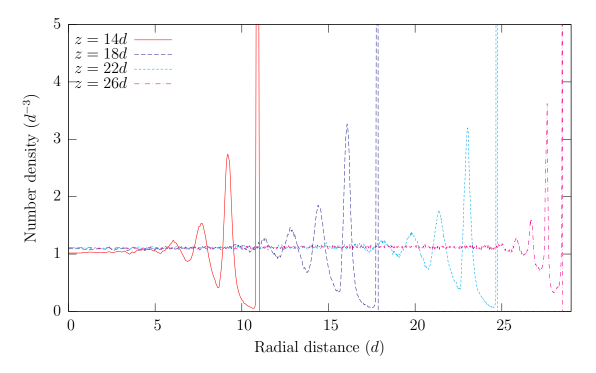


Figure 5-10: Plots of number density (defined as the number of particle centers per unit volume) in the 30° reactor geometry for several low cross sections.

are felt far down in the flow further demonstrates that very little diffusion or shearing occurs in the upper region. There are also similar lower-density regions along the walls, related to partial crystallization described in more detail below.

It is also clear in both geometries, especially the 30° model, that there is a fairly sharp transition between the upper region of nearly rigid plug flow and a less dense lower region of shear flow in the funnel. Similar features are in the velocity profiles described above, but the transition is much more sharp, at the scale of at most a few particles, for the local packing fraction. These sudden variations in material properties and velocities are reminiscent of shock-like discontinuities in Mohr-Coulomb plasticity theories of granular materials [94, 117]. It seems no such existing theory can be applied to the reactor flows, but our results suggest that plasticity concepts may be useful in developing a continuum theory of dense granular flow [65].

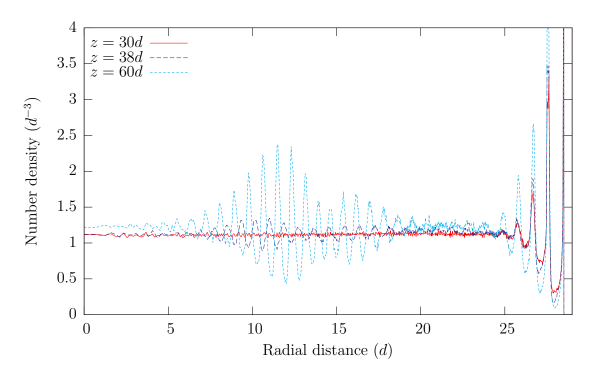


Figure 5-11: Number density plots in the 30° reactor geometry for several high cross sections.

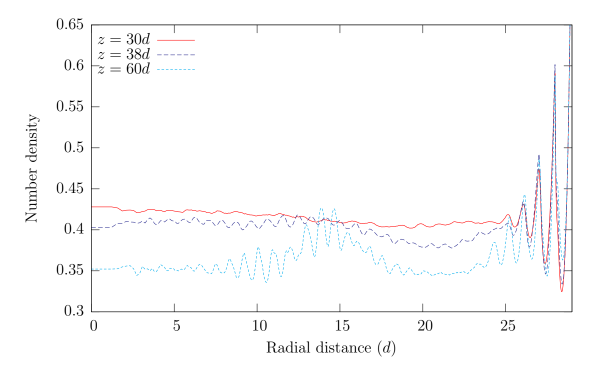


Figure 5-12: Horizontal profiles of porosity at different heights in the 30° reactor geometry.

5.5.2 Local Ordering and Porosity

As noted above, previous simulation studies of local ordering near walls have focused on static packings in simplified cylindrical geometries (without the funnel, outlet pipe, or guide ring) [38, 103], while we compute average statistics for slowly flowing packings in realistic full-scale reactor models. To take a closer look at ordering near walls, we study the number density profile in horizontal slices at different heights. The container is divided into bins in the same way as discussed previously and the number density in a bin is obtained by counting the number of times a particle center lies within that bin.

Figure 5-10 shows a sequence of number density profiles for several low values of z in the thirty degree reactor geometry. At all four heights, lattice effects are clearly visible and quite similar to those observed in experiments [50, 119] and other simulations [38, 103]. For the lowest three heights, these peaks are roughly $\sqrt{3}d$ apart, corresponding to particles crystallized against the conical wall, while for the highest value of z, these effects are roughly 1d apart, due to particles being crystallized against the cylindrical wall. The above graph also shows that in the middle of the container, no lattice effects are present.

However, this situation changes dramatically higher up in the container, as shown in Fig. 5-11. As z increases from 30d to 60d, the interior of the packing goes from being disordered to having a strong radial ordering, centered at around z = 12d. The reason for this ordering is due to the presence of the guide ring high in the container, which keeps the fuel and moderator pebbles separate. The ring, placed at $r_{\rm in} = 14.5d$ in the container, creates radial crystallization, which can then propagate very far downward, since the packing is plug-like for most of the cylindrical part of the reactor. At much lower heights, around z = 40d, this radial ordering is broken, as the particles are forced to reorganize once they enter the parabolic region of flow.

To make a direct connection with the modeling of gas flow, we show horizontal slices of the porosity at different heights in figure 5-12. The porosity is measured here by intersecting the spheres with annular cylindrical bins to compute the fraction of each bin volume not occupied by pebbles. The features noted above appear in the porosity and alter the local permeability, which enters continuum descriptions of helium gas flow in the core [30, 141, 143].

5.6 Residence-Time Distribution

5.6.1 Predictions of the Kinematic Model

The statistical distribution of fuel burnup is closely related to the distribution of pebble residence times in the reactor core, differing only due to nonuniform sampling of the neutron flux profile. Since the upper pebble flow is essentially a uniform plug flow, the distribution of residence times is the same (up to a constant time shift) as the distribution of waiting times for pebbles starting at a given height in the core to exit through the orifice, and we concentrate on these distributions in this section. However, we conclude by examining the residence times for particles to pass through the entire container, to investigate the effects of the guide ring and the outer walls.

We have seen that there is very little pebble diffusion, so fluctuations in the residence time are primarily due to hydrodynamic dispersion in the mean flow. We have also seen that the Kinematic Model gives a reasonable description of the mean flow profile in the conical funnel region, where most of the shear and hydrodynamic dispersion occur. Therefore, we can approximate the residence-time distribution by the distribution of times to travel along different streamlines of the mean flow, starting from different radial positions, r_0 , at a given height z_0 . Below we will compare such predictions, based on our numerical solutions to the Kinematic Model, to our DEM simulations for the two reactor geometries.

5.6.2 An Analytical Formula

We can obtain a simple, exact formula for the residence-time distribution in a somewhat different geometry using the Kinematic Model, as follows. The similarity solution to Eq. (5.1) for a wide, flat bottomed silo draining to a point orifice at z = 0

$$u(r,z) = -\frac{Qr}{2bz^2}e^{-r^2/4bz}$$
(5.2)

$$v(r,z) = \frac{Q}{bz}e^{-r^2/4bz}$$
 (5.3)

where u and v are the radial (horizontal) and downward velocity components and Q is a constant proportional to the total flow rate through the orifice. (This is just the classical Green function for the diffusion equation in two dimensions, where z acts like "time".) A slightly more complicated solution is also possible for a parabolic silo, but let us focus on the simplest case of Eqs. (5.2)-(5.3), which is a good approximation for a wide parabolic funnel, where the velocity near the walls is small, i.e. $R > \sqrt{4bz_0}$. A more detailed analysis is not appropriate here, since a simple analytical solution does not exist for the actual reactor geometry of a conical funnel attached to straight cylinder.

For the flow field in Eqs. (5.2)-(5.3), the trajectory of a Lagrangian tracer particle along a streamline is given by

$$\frac{dr}{dt} = u(r,z), \quad r(t=0) = r_0 \tag{5.4}$$

$$\frac{dz}{dt} = -v(r,z), \quad z(t=0) = z_0 \tag{5.5}$$

Combining these equations and integrating, we find that the streamlines are parabolae, $z/z_0 = (r/r_0)^2$, and that the residence time for a pebble starting at (r_0, z_0) is

$$\tau_0(r_0, z_0) = \frac{bz_0^2}{2Q} e^{r_0^2/4bz_0}.$$
(5.6)

Now we consider pebbles that are uniformly distributed at a height z_0 in a circular cross section of radius R in the flow field Eqs. (5.2)-(5.3. The probability distribution

for the residence time of those pebbles is

$$p(\tau|z_0, R) = \int_0^R \delta(\tau - \tau_0(r_0, z_0)) \frac{2\pi r_0 dr_0}{\pi R^2}$$
(5.7)

$$= \begin{cases} 0 & \text{for } \tau < \tau_{min}(z_0) \\ 4bz_0/R^2\tau & \text{for } \tau_{min} < \tau < \tau_{max} \\ 0 & \text{for } \tau > \tau_{max}(z_0, R) \end{cases}$$
(5.8)

where

$$\tau_{min} = \tau_0(0, z_0) = \frac{bz_0^2}{2Q}$$
(5.9)

$$\tau_{max} = \tau_0(R, z_0) = \frac{bz_0^2}{2Q} e^{R^2/4bz_0}$$
(5.10)

Once again, this solution is strictly valid for an infinitely wide and tall silo draining to a point orifice, and it is roughly valid for a parabolic funnel, $z/z_0 = (r/R)^2$, as an approximation of a conical funnel in the actual reactor geometry. We can further approximate the effect of a nearly uniform flow of speed v_0 to describe the upper cylindrical region by simply adding $(z - z_0)/v_0$ to the residence time for a starting point $z > z_0$.

Although this analysis is for a modified geometry, we will see that it captures the basic shape of the residence-time distributions from the DEM simulations in a simple formula (5.8). The probability density is sharply peaked near the shortest residence time, τ_{min} , corresponding to pebbles near the central axis traveling the shortest distance at the largest velocity. The longer distance and (more importantly) the smaller velocity at larger radial positions cause strong hydrodynamic dispersion, resulting a fat-tailed residence-time density which decays like 1/t, up to a cutoff τ_{max} .

5.6.3 Simulation Results

For the DEM reactor simulations, we calculate the distribution of times it takes for particles to drop from several different values of z_0 , adding in a weighting factor to take into account that shorter residence times are preferentially observed in the data

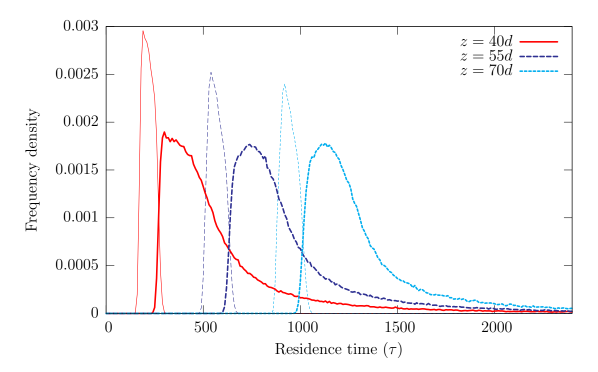


Figure 5-13: Residence-time probability densities for the time it takes particles to drop from a specific height z out of the container, for the 30° reactor geometry for fuel pebbles (heavy lines) and for moderator pebbles (thin lines).

set.

Since we are primarily interested in the radioactive burnup, we concentrate on the residence times for the fuel pebbles, but for comparison, we also report results for the moderator pebbles. Figure 5-13 shows the residence-time probability densities for pebbles starting at z = 40d, 55d, 70d to exit the container for the 30° reactor geometry. The distributions for the moderator pebbles are quite narrow, showing all particles exit over a short time window. In contrast, the distributions for the fuel pebbles exhibit fat tails, as expected qualitatively from the Kinematic Model approximation (5.8) for a parabolic geometry. A closer analysis of the data confirms that the longest waiting times are associated with pebbles passing close to the walls, especially near the corner between the conical and cylindrical wall sections, although there are no completely stagnant regions.

Figure 5-14 shows corresponding plots for the 60° reactor geometry. In general, the residence-time densities have similar shapes as for the 30° geometry, but they are

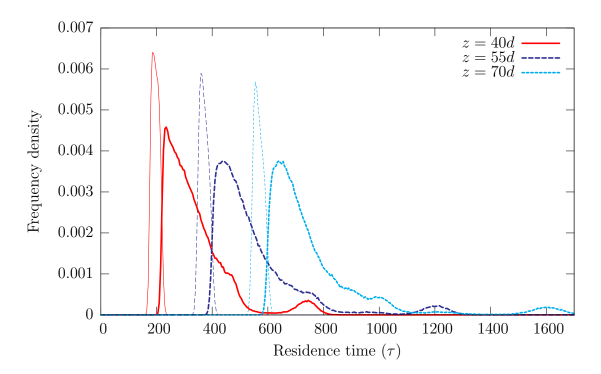


Figure 5-14: Residence-time probability densities for the time it takes particles to drop from a specific height z out of the container, for the 60° reactor geometry for fuel pebbles (heavy lines) and for moderator pebbles (thin lines).

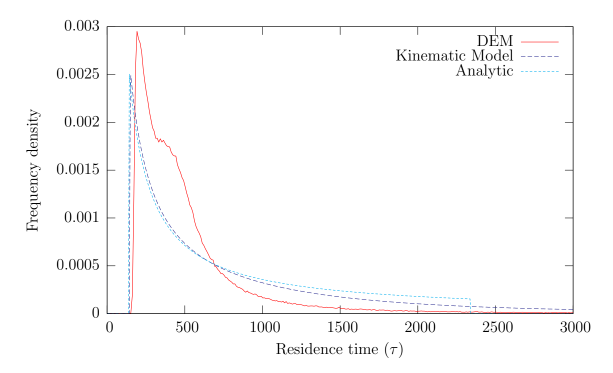


Figure 5-15: Comparison of the residence time distributions between DEM simulation, numerical solution of the Kinematic Model, and the analytic formula.

much narrower and exhibit a small secondary peak far into the tail. Examining movies shows that this extra peak is due to a boundary layer of particles, roughly one-pebble thick, touching the 60° conical wall sliding down at a speed lower than the nearby bulk. This extra source of hydrodynamic dispersion could not be easily captured by a continuum model for the mean flow. A simple way to eliminate it would be to replace add an outer annulus of moderator pebbles (controlled by another guide ring at the top), which would flow more slowly along the walls, leaving the fuel pebbles in a more uniform flow with smaller fluctuations. Another possibility would be to reduce the wall friction, which makes the flow more uniform, as discussed in the following section.

Figure 5-15 investigates the accuracy of the Kinematic Model in predicting the DEM residence-time distribution. The total residence-time distribution for both fuel and moderator pebbles to exit the reactor from z = 40d in the 30° geometry is shown, and is compared with two predictions from the Kinematic Model, one making use of the analytic formula (5.8), and one making use of the numerical solution of the velocity profile. We use of the value b = 2.5d and calibrate the total flow to match the total flow from the DEM data. Both the numerical solution and the analytic formula can roughly capture the overall shape of the DEM distribution, although neither achieves a good quantitative agreement, particularly in the tails. Since the analytic formula assumes all streamlines are parabolic, it fails to take into account the slow-moving particles that stay close to the wall, and it therefore predicts a cut-off in the residence time distribution which is much shorter than some of the observed residence times in the DEM simulation. The numerical solution of the Kinematic Model accounts for this and provides a better match, although it is clear that a model correctly accounting for the flow of pebbles near the container walls may be required in order to achieve high accuracy.

5.6.4 Residence times for the entire container

We also considered the distribution of times for the particles to pass through the entire container. While the flow in the upper part of the reactor is essentially plug-

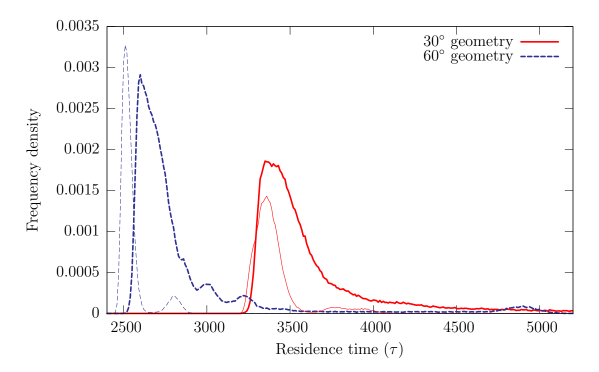


Figure 5-16: Distribution of times to pass through the entire container for fuel pebbles (heavy lines) and moderator pebbles (thin lines).

like, boundary effects near the container walls and on the guide ring can have an appreciable effect on the pebble residence times, which we study here. Since it takes a long time for particles to pass through the entire container we made use of the two extended data sets, consisting of 1,427 snapshots for the thirty degree geometry and 1,249 snapshots for the sixty degree geometry.

Figure 5-16 shows the time distributions for pebbles to pass through the entire container. Apart from a large positive time shift, the curves are similar in form to those in Fig. 5-13 and Fig. 5-14. However, for both geometries, we see second small peaks in the distributions for the moderator pebbles, corresponding to a slow-moving boundary layer of pebbles touching the guide ring. The sixty degree curve for the fuel pebbles also exhibits several undulations corresponding to multiple layers of pebbles crystallized against the outer wall, each moving at different speeds.

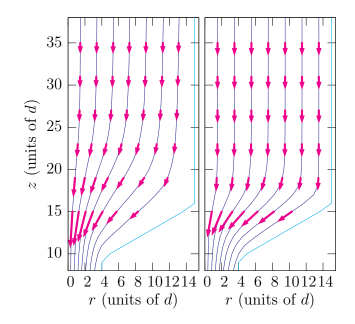


Figure 5-17: Streamlines for the half-size, monodisperse geometries with wall friction (left) and without wall friction (right). Arrows are proportional to the velocity vectors in selected horizontal slices.

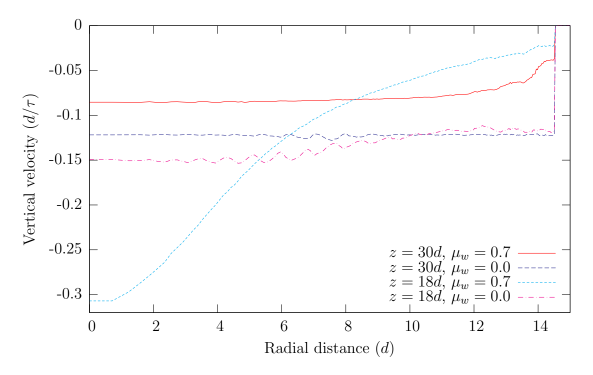


Figure 5-18: Comparison of velocity profiles for simulations with and without wall friction for two different heights.

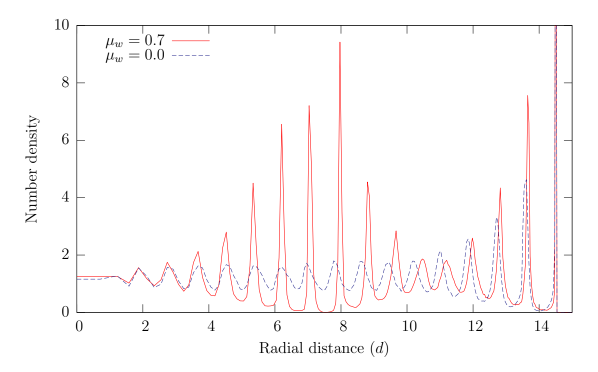


Figure 5-19: Comparison of number density profiles at z = 60d for simulations with and without wall friction.

5.7 Wall friction

The behavior of pebbles near the walls is of significant interest to reactor design, and to look into this further, we investigated the effect of wall friction by comparing two simulations runs in the half-size geometry, with wall friction coefficients $\mu_w = 0$ and $\mu_w = 0.7$. All other aspects of the simulation, including the interparticle interactions, were kept the same.

Figures 5-17 and 5-18 show comparisons of flow streamlines and velocity profiles respectively for the two simulations. We see that the $\mu_w = 0$ simulation results in a significantly larger flow speed, with a mass flow rate of $104m\tau^{-1}$, as opposed to $59.6m\tau^{-1}$ for $\mu_w = 0.7$. As would be expected, removing wall friction also removes the boundary layer of slower velocities at the wall, creating an almost perfectly uniform velocity profile high in the reactor. This also has the effect of increasing radial ordering effects, and we can see from figure 5-19 that the number density profile is more peaked close to the wall. Figure 5-19 also shows that the radial ordering created by the guide ring is also significantly enhanced. While this is due in part to the more plug-like flow allowing packing effects to propagate further down, it is also due to the frictionless guide ring initially creating radial ordering. Thus it may be possible to tune the material properties of the guide ring (or the roughness of its walls) to enhance or reduce the radial ordering effects.

Removing wall friction also has the effect of increasing radial ordering effects near the wall. Perhaps most surprisingly, removing wall friction results in a significant alteration of the flow in the *interior* of the packing, as shown by the two velocity profiles in figure 5-18 for z = 18d. While both velocity profiles must converge upon the orifice, we see that the velocity profile for the $\mu_w = 0.7$ case is significantly more curved than that for $\mu_w = 0$. This also has the effect of preferentially speeding up the relative flux of fuel pebbles: with wall friction, the fuel pebbles make up 71.5% of the total mass flux, but without wall friction, this increases to 74.7%.

5.8 Bidispersity

5.8.1 The Bidisperse PBR Concept

The two-pebble design of MPBR with a dynamic central moderator column has various advantages over a solid graphite central column (as in the revised PBMR design). For example, it flattens the neutron flux profile, while preserving a very simple core vessel without any internal structures, which would be subjected to extreme radiation and would complicate the granular flow. It also allows the widths of the moderator column and fuel annulus to be set "on the fly" during reactor operation, simply by adjusting the guide ring at the top.

A drawback of the dynamic moderator column, however, is its porosity, which allows the passage of the helium-gas coolant, at the highest velocity (along the central axis). In most PBR designs, high-pressure helium gas is introduced from a reservoir above the core, through holes in the graphite bricks which make up the core vessel. The gas then flows through the core and exits through holes in the graphite bricks of the conical funnel to another reservoir at high temperature. To improve the thermal

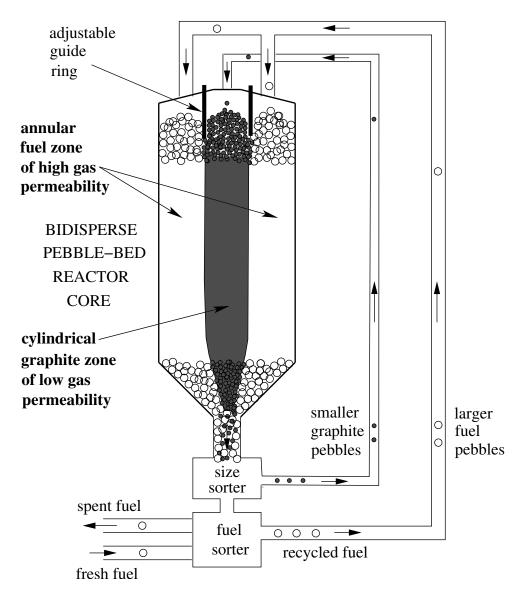


Figure 5-20: Schematic diagram of the pebble flow in a bidisperse MPBR design.

efficiency and power output, it would be preferable to focus the gas flow on the fuel annulus and the interface with the moderator column, where the most heat is generated. This is automatically achieved with a solid graphite column, but there is a very simple way to shape the gas flow in a similar way with a dynamic column, while preserving its unique advantages.

The idea is to make the graphite moderator pebbles in the central column smaller than the fuel pebbles in the outer annulus, as shown in Fig. 5-20. (This also helps with sorting of fuel and moderator pebbles as they exit the core.) In standard continuum models of flow in porous media [30, 141, 143], the permeability of the packing scales with the square of the pebble diameter (or pore size), so reducing the diameter of the moderator pebbles can greatly reduce the gas flow (e.g. by a factor of four for halfdiameter pebbles). This argument holds everywhere that the packing is statistically the same, in the monodisperse packings of the fuel annulus and the moderator column, which have the same porosity. At the interface between the two regions, we have seen in Figures 5-8 and 5-12 that the porosity is enhanced for a monodisperse core due to the guide ring, although a bidisperse interface will have different structure. In summary, if helium gas is introduced outside the guide ring in a bidisperse core, it can be made to pass almost entirely through the fuel annulus and the interface with the moderator column.

5.8.2 Simulation Results

The only question regarding the feasibility of the bidisperse core is the stability of the central column over time and the possibility of enhanced diffusion of the small moderator pebbles into the annulus of larger fuel pebbles. In other systems, such as rotating drums [96, 70, 69], vibrated buckets [121, 120], and draining silos [114], bidisperse granular materials display a tendency to segregate (rather than mix) during dynamics, but there is currently no general theory which could be applied to our reactor geometry. Therefore, our DEM simulations provide a useful means to address this important question.

Figure 5-21 shows snapshots of vertical cross sections for the three different bidis-

perse simulations that were run in the half-size geometry. As shown in the diagram, the central column remains stable and coherent in all three cases, and very little mixing between the two types of pebbles is visible. Figure 5-22 shows a comparison of the velocity profiles from the three simulations for two different heights. It is reassuring to see that the bidisperse simulations do not significantly differ from the monodisperse simulation, although we do see a slightly higher overall flow rate in the bidisperse systems: we see total mass flow rates of $59.6m\tau^{-1}$, $60.8m\tau^{-1}$, and $65.0m\tau^{-1}$ for the monodisperse, 0.8:1, and 0.5:1 simulations respectively.

The velocity profiles are slightly more curved in the bidisperse central core; this is particularly apparent in the 0.5:1 simulation. This leads to a small cusp in the velocity profile near the interface between the two types of particles which may lead to adverse mixing effects. The faster flow also leads to a significantly larger turnaround of the moderator pebbles. In the monodisperse system, the moderator pebbles comprise 28.5% of the total mass flux, but this is increased to 31.7% in the 0.8:1 bidisperse simulation, and 42.6% in the 0.5:1 bidisperse simulation.

To investigate the amount of mixing of the central column, we used a technique similar to that described in section 5.4. At z = 110d all moderator particles with r > 8d are marked, and their radial diffusion is then calculated as a function of z. The results are shown in figure 5-23: in the cylindrical section of the packing, there is very little difference between the three simulations, but in the area of convergent flow, we see that bidispersity leads to significantly more mixing. However, even for the 0.5:1 simulation, the scale of diffusion is still smaller than a single particle diameter, and essentially the central column remains stable.

Due to computational limitations, we were unable to investigate smaller size ratios in the reactor geometries, so we carried out simulations in a smaller container with a 0.3:1 size ratio (figure 5-24) and found dramatically different behavior: During drainage, the central column became unstable, and the small particles penetrated many particle diameters into the packing of larger particles. We expect that there is a fundamental crossover in behavior simply due to geometry of amorphous packings, when the moderator pebbles become small enough to pass through the gaps between

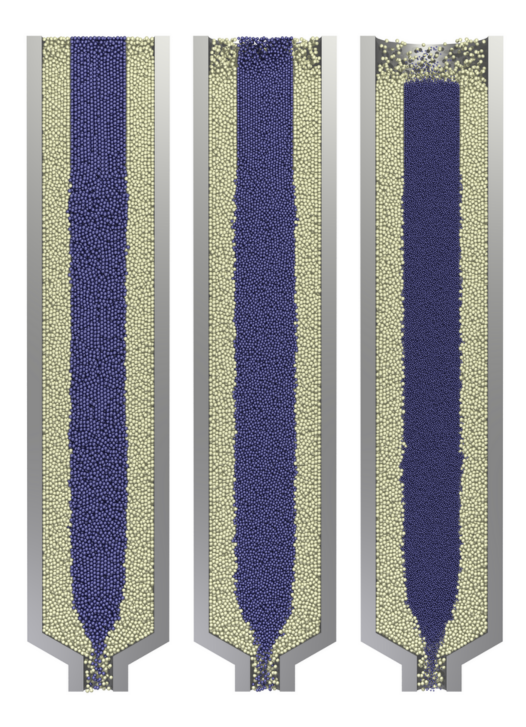


Figure 5-21: Snapshots of vertical cross-sections for the bidisperse simulations. From left to right, the moderator pebbles have diameters 1d, 0.8d, and 0.5d while the fuel pebbles are of constant size 1d.

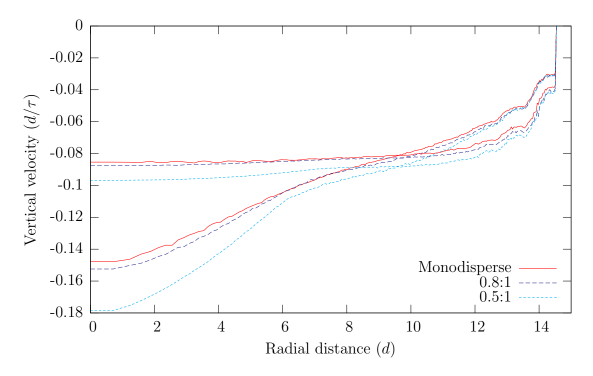


Figure 5-22: Comparison of velocity profiles for the three bidisperse simulations. The three flatter curves are calculated at z = 30d in the plug-like flow region while the other three were taken at z = 22d in the parabolic flow region.

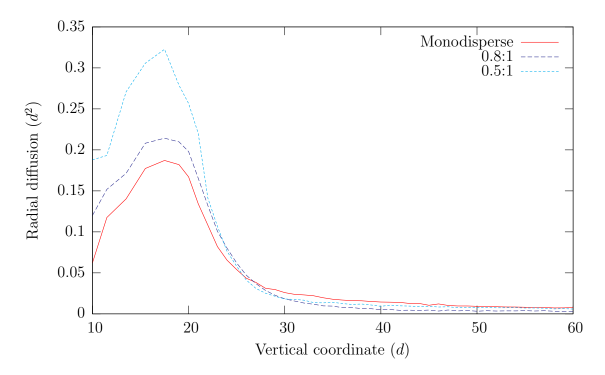


Figure 5-23: Comparison of particle diffusion for the three bidisperse simulations.

the densely packed fuel pebbles. An in-depth study of this phenomenon remains a subject of future work. For now, we can safely recommend a diameter ratio of 0.5:1, which reduces the dynamic central column's permeability by a factor of four without introducing any significant diffusion of moderator pebbles into the fuel annulus.

5.9 Conclusions

5.9.1 Pebble-Bed Reactor Core Design

Using DEM simulations, we have analyzed many aspects of granular flow in pebblebed reactor cores of direct relevance for design and testing. We close by summarizing some key conclusions.

The mean flow profile exhibits a smooth transition from a nearly uniform plug flow in the upper cylindrical region to a nonuniform, converging flow in the lower funnel region, consistent with recent experiments [27, 63]. There are no stagnant regions in the 30° and 60° conical funnels considered in this study, although the flow is slower near the corner at the top of the funnel, especially in the former case. Moreover, the wider 30° funnel has a boundary mono-layer of slower pebbles partially crystallized on the wall.

The only available continuum theory for such flows, the simple Kinematic Model [76, 90, 95, 94], gives a reasonable qualitative picture of the flow profiles, although it cannot capture discrete boundary-layer effects. As in other experiments on similar geometries [27], the Kinematic Model does not quantitatively predict the dependence of the flow profile on geometry. We suggest that it be used to get a rough sense of the flow profile for a given core geometry prior to (much more computationally expensive) DEM simulations and/or experiments.

We have quantified the degree of pebble mixing in the core. Although there is some horizontal diffusion in the funnel region, pebbles depart from the streamlines of the mean flow by less than one pebble diameter prior to exiting the core.

We have demonstrated that the "mixing layer" between the central moderator

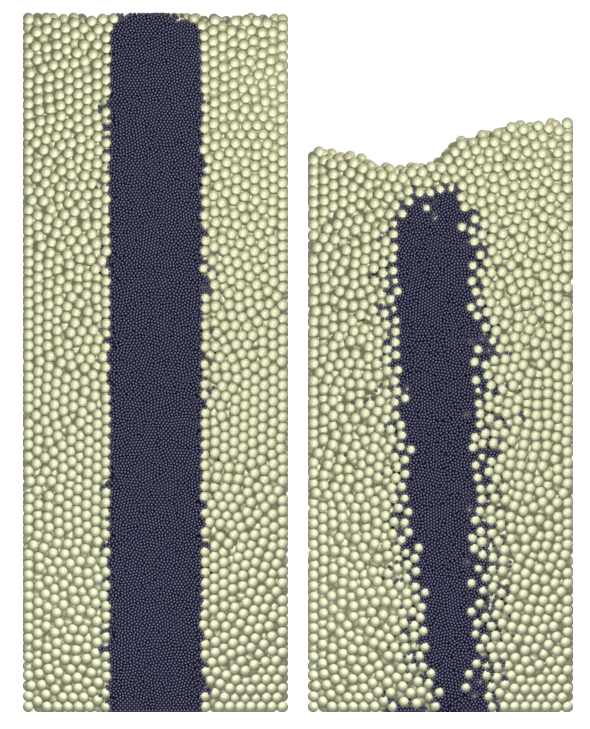


Figure 5-24: Simulation snapshots of a bidisperse drainage experiment with size ratio 0.3:1 in a rectangular silo of width 25d and depth 4d. The left image is taken before drainage taken place, and even at this stage, some of the smaller particles have penetrated into the packing of larger particles. During drainage (right) some of the small particles penetrate deep into the side packings. Small particles are also visible on the container base, having fallen through the gaps between the large particles.

column and the outer fuel annulus, which appears in prior models [51], can be reduced to the thickness of one pebble diameter by separating moderator and fuel pebbles with a guide ring at the ceiling (to eliminate mixing by surface avalanches), consistent with experiments on MPBR models [63]. We conclude that the dynamic central column of moderator pebbles is a sound concept, which should not concern regulators.

We have constructed Voronoi tessellations of our flowing packings to measure the profile of volume fraction (or porosity) and found some unexpected features which would affect coolant gas flow through the core. The bulk of the core, in the plug-flow region of the upper cylinder, has a volume fraction near the jamming point (63%), but there is a sharp transition to less dense packings (55 - 60%) in the funnel region, due to shear dilation. We also observe lower volume fractions in this range at the moderator/fuel interface in the upper cylinder, below the guide ring, and lower volume fractions (50 - 55%) against the walls. These narrow regions of increased porosity (and thus, increased permeability) would allow faster helium gas flow.

We have also studied local ordering in the flowing packings and find evidence for partial crystallization within several pebble diameters of the walls, consistent with previous experiments [50, 119] and simulations [38, 103]. Such ordering on the walls of the guide ring, then advected down through the core, is responsible for the increased porosity of the moderator/fuel interface.

We have varied the wall friction in our DEM simulations and observe that it can affect the mean flow, even deep into the bulk. Reducing the wall friction increases radial ordering near the walls and makes the flow profile more uniform.

Since diffusion is minimal, the probability distribution of pebble residence times is dominated by advection in the mean flow. Therefore, we have made predictions using the Kinematic Model, numerically for the conical-funnel reactor geometries, and analytically for a wide parabolic funnel. The model predicts a fat-tailed ($\sim 1/t$) decay of the residence-time density due to hydrodynamic dispersion in the funnel region.

Our DEM simulations predict that the 60° conical funnel results in a narrower residence-time distribution than the 30° funnel, which has more hydrodynamic dis-

persion. The steeper 60° funnel also exhibits a boundary layer of slower, partially crystallized pebbles near the wall which lead to an anomalous bump far in the tail of residence-time distribution. These results have important implications for nonuniformity in the burnup of fuel pebbles.

We have introduced the concept of a bidisperse core with smaller moderator pebbles in the dynamic central column than in the outer fuel annulus, in order to focus the helium gas flow on the fuel. Our DEM simulations demonstrate that there is negligible pebble mixing at the interface for diameter ratios as small as 0.5:1, for which the permeability of the moderator column is reduced by a factor of four. We conclude that the bidisperse MPBR design is sound and will produce a stable moderator-pebble column of greatly reduced gas permeability.

A natural next step would be to combine our full-scale DEM model for the pebble flow with existing computational approaches to reactor core physics [131, 51], which rely on pebble flow as an empirical input. More accurate studies of gas flow in the core could also be done, starting from our complete pebble packings, or the average quantities such as the porosity. With such computational tools, one should be able to reliably test and develop new reactor designs.

5.9.2 Basic Physics of Dense Granular Flow

We have noted a number of favorable comparisons between our simulations and experiments in similar geometries [63, 27, 50, 119], which provides further validation of the Discrete-Element Method as a realistic means of simulating granular materials. As such, it is interesting to consider various implications of our results for the theories of dense granular flow, since the simulations probe the system at a level of detail not easily attained in experiments.

Our conclusions about the Kinematic Model are similar to those of a recent experimental study [27]: The model describes the basic shape of the flow field in the converging region, but fails to predict the nearly uniform plug flow in the upper region with vertical walls or the precise dependence on the funnel geometry. It also cannot describe boundary-layers due to partial crystallization near walls or incorporate wall friction, which we have shown to influence the entire flow profile.

On the other hand, there is no other continuum model available for dense silo drainage, except for Mohr-Coulomb plasticity solutions for special 2D geometries, such as a straight 2D wedge without any corners [94], so it is worth trying to understand the relative success of the Kinematic Model for our 3D reactor geometries and how it might be improved. A cooperative microscopic mechanism for random-packing dynamics, based on "spots" of diffusing free volume, has recently been proposed, which yields the mean flow of the Kinematic Model as the special case of independent spot random walks with uniform upward drift from the orifice (due to gravity) [17]. Under the same assumptions, the spot model has also been shown to produce rather realistic simulations of flowing packings in wide silos (compared to DEM simulations) [112], where the Kinematic Model is known to perform well [137, 84, 114, 28]. This suggests that some modification of the spot dynamics, such as spot interactions and/or nonuniform properties coupled to mechanical stresses, and an associated modification of the Kinematic Model in the continuum limit, may be possible to better describe general situations.

From a fundamental point of view, perhaps the most interesting result is the profile of Voronoi volume fraction (or porosity) in our flowing random packings in Figure 5-8. Although the mean velocity in Figure 5-2 shows a fairly smooth transition from the upper plug flow to the lower converging flow, the volume fraction reveals a sharp transition (at the scale of 1-3 particles) from nearly jammed "solid" material in the upper region (63%) to dilated, sheared "liquid" material in the lower region (57-60%). The transition line emanated from the corners between the upper cylinder and the conical funnel. We are not aware of any theory to predict the shape (or existence) of this line, although it is reminiscent of a "shock" in the hyperbolic equations of 2D Mohr-Coulomb plasticity [94].

Our measurements of diffusion and mixing provide some insights into statistical fluctuations far from equilibrium. Consistent with the experiments in wide quasi-2D silos [28], we find that diffusion is well described geometrically as a function of the distance dropped, not time (as in the case of thermal molecular diffusion). As a clear

demonstration, there is essentially no diffusion as pebbles pass through the upper core, until they cross the transition to the funnel region, where the diffusion remains small (at the scale of one pebble diameter) and cooperative in nature. The behavior in the funnel is consistent with the basic spot model [17], but a substantial generalization would be needed to describe the transition to the upper region of solid-like plug flow, perhaps using concepts from plasticity theory [65].

We view silo drainage as a fundamental unsolved problem, as interesting and important as shear flow, which has received much more attention in physics. The challenge will be to find a single theory which can describe both shear cells and draining silos. Our results for pebble-bed reactor geometries may provide some useful clues.