PIFE-PIC: PARALLEL IMMERSED-FINITE-ELEMENT PARTICLE-IN-CELL FOR 3-D KINETIC SIMULATIONS OF PLASMA-MATERIAL INTERACTIONS∗
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Abstract. This paper presents a recently developed particle simulation code package PIFE-PIC, which is a novel three-dimensional (3-D) Parallel Immersed-Finite-Element (IFE) Particle-in-Cell (PIC) simulation model for particle simulations of plasma-material interactions. This framework is based on the recently developed non-homogeneous electrostatic IFE-PIC algorithm, which is designed to handle complex plasma-material interface conditions associated with irregular geometries using a Cartesian-mesh-based PIC. Three-dimensional domain decomposition is utilized for both the electrostatic field solver with IFE and the particle operations in PIC to distribute the computation among multiple processors. A simulation of the orbital-motion-limited (OML) sheath of a dielectric sphere immersed in a stationary plasma is carried out to validate PIFE-PIC and profile the parallel performance of the code package. Furthermore, a large-scale simulation of plasma charging at a lunar crater containing 2 million PIC cells (10 million FE/IFE cells) and about 520 million particles, running for 20,000 PIC steps in about 109 wall-clock hours, is presented to demonstrate the high-performance computing capability of PIFE-PIC.

Key words. immersed-finite-element, particle-in-cell, parallel domain decomposition, plasma-material interactions

AMS subject classifications. 35R05, 65N30, 65Y05

1. Introduction. Particle modeling of plasma dynamics has emerged as one of the most appropriate algorithms for first-principle-based modeling of many plasma-material interaction (PMI) problems. One of the fundamental phenomena in plasma-material interactions is surface charging. When an object is immersed in a plasma, its surface will collect charge from the plasma until it reaches an equilibrium surface potential determined by the current balance condition. Many plasma-material interaction problems involve multiple objects with complex geometries, therefore the interface conditions between the plasma and object need to be accurately resolved.

Being one of the most popular kinetic methods for collisionless plasma simulations, the Particle-in-Cell (PIC) method [9] models the charged particles as macro-particles and tracks the motions of particles in the electrostatic/electromagnetic field. The electric potential in a PIC simulation domain is governed by the second-order elliptic partial differential equations (PDEs) with discontinuous dielectric coefficients and non-homogeneous flux jumps across the material surface interface. Numerical methods based on structured meshes, especially Cartesian meshes, are particularly desirable in these simulations because they enable efficient particle tracking and save computing time in particle-mesh interactions.

The immersed-finite-element (IFE) method is a finite element method (FEM) for solving interface problems using interface-independent meshes such as Cartesian meshes. The main idea of IFE is to adjust approximating functions locally to accommodate the physical interface conditions. An IFE method can achieve optimal convergence on an interface-independent mesh with the number and location of the degrees-of-freedom isomorphic to the standard FEM on the same mesh. The first IFE method was introduced by Z. Li in [44] for solving one-dimensional (1-D) elliptic interface problems with piecewise linear polynomials. Since then, the IFE method has been extended to higher-order approximations [2, 11, 12, 21], higher-dimensional elliptic interface problems [18, 20, 45, 31, 46, 48, 59], and other interface problems with other PDE models [1, 3, 4, 19, 34, 47].

Over the past decade, the IFE method has been successfully used together with PIC in plasma particle simulations [3, 14, 38, 39, 42]. Recently, a non-homogeneous IFE-PIC algorithm has been developed for particle simulations of plasma-material interactions with complex geometries while maintaining the computational speed of the Cartesian-mesh-based PIC [15, 25, 28, 30, 49, 50]. To the best of our knowledge, most existing IFE-PIC algorithms are serial. The non-parallel algorithms have limitations in their capability to handle large-scale particle simulations and their efficiency in using multiple processors at the algorithm level.

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For a typical large-scale 3-D PIC simulation, millions to billions of particles are tracked in the computation domain that contains millions of elements. With the availability of multi-processor computational facilities, the call for parallel IFE-PIC algorithms is urgent.

The goal of this paper is to develop and test a new Parallel IFE-PIC package for particle simulations of electrostatic plasma-material interactions, namely, PIFE-PIC. We utilize a 3-D domain decomposition technique for both field-solve and particle-push procedures of the PIC model. The computations are distributed into multiple subdomains which can be handled independently by multiple processors. The key is how to efficiently exchange the information between these subdomains. In this work, neighboring subdomains have a small overlapping (“guard cells”) region which will be used as a common region to interchange the PDE solutions and the particle data. Extensive numerical experiments show that our PIFE-PIC scheme significantly outperforms the serial IFE-PIC scheme. Although it maintains a similar accuracy as the serial IFE-PIC computational scheme, the high parallel performance dramatically reduces the computational time for problems of practical interests. Hence, large-scale kinetic simulations of plasma-material interactions can be carried out much more efficiently.

The rest of this paper is organized as follows. In Section 2, we describe the details of 3-D domain decomposition for both IFE (field-solve) and PIC (particle-push) procedures of PIFE-PIC. In Section 3, we present a code validation using a 3-D sheath problem of a dielectric sphere immersed in a stationary plasma. Section 4 presents a parallel efficiency test of the PIFE-PIC code for strong scaling. Section 5 presents an application of PIFE-PIC to simulations of lunar surface charging at a crater. Finally, a summary and conclusion are given in Section 6.

2. Parallel IFE-PIC Algorithms.

2.1. Overview of PIC and IFE-PIC. PIC is a widely-used kinetic particle simulation method for plasma dynamics [10, 36]. In PIC, charged particles of plasma species are represented by a number of simulation particles (also referred to as macro-particles or super-particles) distributed “freely” in the entire computation domain, while the field quantities such as electric potential are discretized on a mesh (thus the name “particle-in-cell”). The kernel of PIC method is the “PIC loop” which includes four essential steps: scatter, field-solve, gather, and particle-push (Figure 2.1). Within a PIC loop, quantities carried by the simulation particles are weighted onto the mesh nodes (“scatter”) to form the right-hand side (RHS) term of the PDE for the solution of the electrostatic/electromagnetic field (“field-solve”), which is in turn interpolated at particle positions (“gather”) to update the velocity and position of the particles (“particle-push”). Such data exchange between particles and field quantities will iterate for a desired number of steps (or till a convergence criterion is met) to obtain the self-consistent solution of both particles and fields.

For problems of plasma-material interactions, the mathematical model is an interface problem including the electrostatic/electromagnetic field problem in a self-consistent solution to the corresponding plasma

![Fig. 2.1. Four essential steps in a PIC loop.](image)
dynamics problem (Figure 2.2(a)), together with the appropriate interface conditions between the plasma region and the material region (Figure 2.2(b)). For electrostatic problems presented in this work, we consider

\[
-\nabla \cdot (\varepsilon \nabla \phi(X)) = \rho(X), \quad X = (x, y, z) \in \Omega^+ \cup \Omega^-,
\]

\[
\phi(X) = g(X), \quad X \in \Gamma_D,
\]

\[
\frac{\partial \phi(X)}{\partial n_{\Gamma_N}} = p(X), \quad X \in \Gamma_N.
\]

Here, \( \Omega \in \mathbb{R}^3 \) is assumed to be a cuboidal domain, which is divided into two subdomains \( \Omega^+ \) and \( \Omega^- \) by an interface surface \( \Gamma \) such that \( \overline{\Omega} = \overline{\Omega}^- \cup \overline{\Omega}^+ \cup \Gamma \). \( \Gamma_D \) and \( \Gamma_N \) are the Dirichlet and Neumann boundaries such that \( \partial \Omega = \Gamma_D \cup \Gamma_N \). The vector \( n_{\Gamma_N} \) is the unit outward normal of \( \Gamma_N \). See the sketch in Figure 2.2(a). The functions \( \rho, g, \) and \( p \) are the source term, Dirichlet boundary function, and Neumann boundary function, respectively. The electric field \( E = -\nabla \phi(X) \) is discontinuous across the interface \( \Gamma \) with the following jump conditions imposed:

\[
[\phi(X)]_{\Gamma} = 0,
\]

\[
[\varepsilon \frac{\partial \phi(X)}{\partial n_{\Gamma}}]_{\Gamma} = q(X),
\]

where the jump \([\cdot]_{\Gamma}\) is defined by \([w(X)]_{\Gamma} := w^+(X)_{\Gamma} - w^-(X)_{\Gamma}\). The vector \( n_{\Gamma} \) is the unit normal of \( \Gamma \) pointing from \( \Omega^- \) to \( \Omega^+ \). The material-dependent coefficient \( \varepsilon(X) \) is discontinuous across the interface. Without loss of generality, we assume it is a piecewise constant function defined by

\[
\varepsilon(X) = \begin{cases} 
  \varepsilon^-, & X \in \Omega^- \\
  \varepsilon^+, & X \in \Omega^+
\end{cases}
\]

where \( \min(\varepsilon^+, \varepsilon^-) > 0 \).

In many applications of scientific and engineering interest, the shape of the interface \( \Gamma \) is usually non-trivial. Traditionally, when solving field problems involving complex-shaped objects, an unstructured body-fitting mesh is employed to improve accuracy (Figure 2.3(a)). However, a structured mesh, such as Cartesian mesh (Figure 2.3(b)), is more advantageous in kinetic PIC modeling of plasma dynamics from the perspective of computing speed and efficiency, although, it has been limited to problems with relatively simple geometries due to accuracy considerations inherited from finite-difference-based schemes. To solve this dilemma while taking into account both accuracy and efficiency, the immersed-finite-element particle-in-cell (IFE-PIC) method was developed to handle complex interface conditions associated with irregular geometries while maintaining the computational speed of the Cartesian-mesh-based PIC. The detailed IFE formulation and IFE-PIC steps are archived in [30] and the flowchart of the serial IFE-PIC algorithm is shown in Figure 2.4. Over the past few years, the IFE-PIC method has matured to successfully model plasma dynamics problems arising from many space applications, such as ion thruster grid optics [40, 41], ion propulsion plume-induced contamination [43, 60, 26], charging of lunar and asteroidal surfaces [25, 30, 28, 13, 29, 27, 63], and dust transport dynamics around small asteroids [64].
2.2. 3-D Domain Decomposition in PIFE-PIC. In our proposed PIFE-PIC algorithm, the 3-D computational domain is decomposed along each dimension using the Message Passing Interface (MPI) architecture (Figure 2.5). The domain is first decomposed into cuboid blocks with the same PIC mesh resolution. Each subdomain is handled by a processor for both field-solve and particle-push procedures of
the PIC method. Two overlapping PIC cells (“guard cells”) in each dimension are used in PIFE-PIC (Figure 2.5).

Therefore, the boundaries of each subdomain are either on the global boundary or in the interior of its neighboring subdomains. Local IFE mesh is then generated for each subdomain. By virtue of the IFE formulation, PIC and IFE can use different mesh resolutions. In PIFE-PIC, PIC mesh is globally uniform. However, IFE mesh could be globally non-uniform but still locally uniform within each subdomain. The data interaction between IFE and PIC meshes of different resolutions is described in detail in [43]. Figures 2.6 and 2.7 illustrate the 2-D and 3-D views of the domain decomposition and different resolutions.

2.3. Parallel Algorithm for IFE Field Solver. For the parallel electrostatic field solver, Dirichlet-Dirichlet domain decomposition with overlapping cells is used to distribute the subdomains among multiple MPI processes [6]. For each subdomain, the IFE solver is the same as the sequential IFE method with Dirichlet boundary conditions [32, 33]. These Dirichlet boundary conditions are imposed at the boundaries of the subdomain, which are interior for the neighboring subdomains (Figure 2.8, left). Therefore, the field solution at respective neighboring subdomains are used as Dirichlet boundary conditions for each subdomain. Within each field-solve step, inner iterations are performed such that the solutions of the overlapping cells are exchanged and updated as the new Dirichlet boundary conditions for the respective neighboring subdomains. It is noted here that since PIFE-PIC uses 3-D domain decomposition, such MPI data exchange will be carried out at guard cell nodes on “surfaces” (+/- neighbor in one direction, such as Rank 1 and Rank 2 in Figure 2.8), “edges” (+/- neighbor in two directions, such as Rank 3 and Rank 6 in Figure 2.8), and “vertices” (+/- neighbor in three directions). We denote this level of iteration as the “Domain Decomposition Method (DDM)” iteration. The relative error $e_{\text{rel}}$ of DDM is defined with the $L^2$ norm as below:

\[
e_{\text{rel}} = \frac{\|\phi_{\text{new}} - \phi_{\text{old}}\|_{L^2}}{\|\phi_{\text{old}}\|_{L^2}}
\]

where $\phi_{\text{new}}$ and $\phi_{\text{old}}$ denote solutions at the new and old steps in the DDM iteration, respectively.
2.4. Parallel Scheme for PIC Procedures. In PIFE-PIC, simulation particles belonging to the same subdomain are stored together on the processor that solves the electrostatic field of the same subdomain (Figure 2.8, right). In this sense, “particle quantities” and “field quantities” of each subdomain are handled by the same processor. Each processor (MPI rank) handles its own particles belonging to its domain without guard cells (see Figure 2.5). In particle-push, particles crossing the inner boundaries are sent to the corresponding rank based on their destination positions. Note that such particle motion includes similar cases as data exchange for field-solve, which are “crossing one surface” (+/- neighbor in one direction, such as Rank 1 and Rank 2 in Figure 2.8), “crossing an edge (two surfaces)” (+/- neighbor in two directions, such as Rank 3 and Rank 6 in Figure 2.8), and “crossing a vertex (three surfaces)” (+/- neighbor in three directions).
MPI data exchange among neighboring subdomains within DDM iteration. The thick edges (black and red) represent the boundaries of each subdomain without guard cells. Left: for field-solve operations: at inner boundaries with guard cells, the nodes at a certain subdomain's boundary (e.g., Rank 1’s boundary nodes) are also interior nodes of its neighboring subdomain (e.g., Rank 2). Therefore, the field quantities stored on interior nodes of Rank 2 are sent to Rank 1 and used as Dirichlet boundary nodes. Since PIFE-PIC has 3-D domain decomposition, such MPI data exchange will be carried out at guard cell nodes on “surfaces” (+/- neighbor in one direction, such as Rank 1 and Rank 2), “edges” (+/- neighbor in two directions, such as Rank 3 and Rank 6), and “vertices” (+/- neighbor in three directions, not shown on this 2-D illustration). Right: For particle-push operations: each processor handles its own particles belonging to its domain without guard cells (see Figure 2.5). In particle-push, particles crossing the inner boundaries are sent to the corresponding rank based on their destination positions. Note such particle motion includes similar cases as data exchange for field-solve, which are “crossing one surface” (+/- neighbor in one direction, such as Rank 1 and Rank 2), “crossing an edge (two surfaces)” (+/- neighbor in two directions, such as Rank 3 and Rank 6), and “crossing a vertex (three surfaces)” (+/- neighbor in three directions, not shown on this 2-D illustration). For charge-weighting, contributions from all neighboring subdomains are summed together at respective inner boundary nodes.

2.5. Flowchart for PIFE-PIC. Figure 2.9 shows the flowchart of PIFE-PIC. The steps in red color are major steps involving MPI operations associated with domain decomposition. In total, there are three levels of iteration in PIFE-PIC. The first level is the matrix-solving iteration which uses the preconditioned conjugate gradient (PCG) algorithm (PCG level). The second one checks the relative error in the iterations of the domain decomposition method (DDM level). The third one tracks the solution of each PIC step (PIC level).

3. Code Validation. We apply the PIFE-PIC code to simulate the charging of a small dielectric sphere immersed in a collisionless and stationary plasma in the orbital-motion-limited (OML) sheath regime. Successful validations of the serial IFE-PIC against analytic OML solutions are presented in earlier work [30, 28].

3.1. Problem Description and Simulation Setup. We consider a stationary, collisionless hydrogen plasma of equal ion and electron temperatures \(T_i = T_e\). The analytic expressions for ion and electron densities in the plasma are given by the revised OML theory [58, 16]. Therefore, the analytic potential profile near the sphere can be numerically solved from Poisson’s equation in spherical coordinates.

3.1.1. Computation Domain and Mesh. In the simulation, we use a computation domain of a \(5 \times 5 \times 5\) Debye cube with a globally uniform PIC mesh with the size of \(h = 0.1\lambda_D\) in all dimensions, where
Fig. 2.9. Flowchart of PIFE-PIC.

\( \lambda_D \) is the Debye length of the plasma. The entire simulation domain has \( 50 \times 50 \times 50 = 125,000 \) PIC cells which is \( 125,000 \times 5 = 625,000 \) tetrahedral FE/IFE cells as each cuboid PIC cell is partitioned into 5 tetrahedral FE/IFE cells in 3-D IFE-PIC \([30, 28]\). The IFE mesh size is also globally uniform and the same as that of the PIC mesh. The dielectric sphere is centered at \((0, 0, 0)\) with a radius of \( R_s = 0.401 \). Due to symmetry in all three dimensions, only \( 1/8 \) of the sphere is included in the domain. The entire domain is partitioned into \( 5 \times 5 \times 5 \) subdomains with each subdomain computed by one MPI process. Figure 3.1 shows the 3-D IFE mesh and setup used in the simulation.

3.1.2. Field Setup. At \( X_{\text{max}}, Y_{\text{max}}, \) and \( Z_{\text{max}} \) boundaries, the potentials are set to 0 as the reference potential. At \( X_{\text{min}}, Y_{\text{min}}, \) and \( Z_{\text{min}} \) boundaries, zero-Neumann boundary conditions are applied due to symmetry (Figure 3.1(b)). The relative permittivity of the sphere is set to 4. The floating potential of the sphere is calculated from the non-homogeneous flux jump condition at the sphere surface.

3.1.3. Particle Setup. The simulation is carried out using the realistic ion-to-electron mass ratio of \( m_i/m_e = 1836 \). Particles are pre-loaded into the domain before the initial field solution, and injected into the domain at \( X_{\text{max}}, Y_{\text{max}}, \) and \( Z_{\text{max}} \) within each PIC step. Particles hitting the \( X_{\text{min}}, Y_{\text{min}}, \) and \( Z_{\text{min}} \) boundaries are reflected due to symmetry. Particles hitting the \( X_{\text{max}}, Y_{\text{max}}, \) and \( Z_{\text{max}} \) are absorbed and removed from the simulation. The normalized time step size was set to be 0.01. There were 125 particles \((5 \times 5 \times 5)\) per species, per cell being loaded/injected into the domain.

3.2. Simulation Results. The simulation of the validation case finished in about 2 hours for a total of 50,000 PIC steps on the Foundry cluster provided by the Center of High-Performance Computing Research at Missouri University of Science and Technology. The computing nodes are configured with Dell C6525 nodes each having four node chassis with each node containing dual 32-core AMD EPYC Rome 7452 CPUs with 256 GB DDR4 RAM and six 480GB SSD drives in RAID 0. All other simulations presented in this work were also carried out on the same cluster.

For this test case, the maximum number of PCG iterations was set to 60 with a tolerance (for relative residual) of \( 1 \times 10^{-6} \), the max number of initial DDM iterations (solving the initial electrostatic field before
main PIC loop starts) was set to 150 and the max number of DDM iterations at each PIC iteration step was set to 50 with a tolerance of $1 \times 10^{-2}$. The simulation was set to run 50,000 PIC steps.

3.2.1. Initial Field Solution. The initial field solution (the zeroth PIC step) took about 100 DDM iterations which are more than what is needed at each step of the main PIC loop, to converge in terms of the relative error $1 \times 10^{-2}$. The idea of setting a relatively larger DDM iteration number is to obtain a better initial field for the main PIC loop. Since the initial field was solved only once, the extra DDM iterations contributed little to the overall wall-clock time of the entire simulation.

3.2.2. Solution History of Main PIC Loop. Figure 3.2 shows the field solution convergence history including the max absolute PCG residual and max DDM relative error as a function of PIC steps in the main PIC loop. A few phenomena are observed here:

1. For most PIC steps, PCG took about 45-50 iterations to reach the tolerance of $1 \times 10^{-6}$. The “max” in the vertical axis stands for “max among all subdomains” (first plot);
2. The maximum PCG absolute residual of the matrix solver has been maintained below $1 \times 10^{-6}$ (second plot);
3. At early PIC steps, most DDM steps took about tens of iterations to converge below $1 \times 10^{-2}$, while later on as PIC approaches steady state, most DDM steps were able to converge within about 10~15 iterations (third and forth plots).

Figure 3.3 shows the global particle number history. At the steady state, there are approximately $1.56 \times 10^7$ particles in the entire global domain. It is also shown that the numbers of particles reached steady state at normalized simulation time of about $\hat{t} = 125$.

3.2.3. Comparison with Analytic Solution. Figure 3.4 shows the comparison between PIFE-PIC simulation results against analytic solution for the OML sheath problem as well as a 3-D potential contour. The potential profile agrees very well with the analytic solution, as also shown in earlier work with the serial IFE-PIC [28, 30].

3.3. Performance Profiling. Table 3.1 shows the detailed timer profile of PIFE-PIC on the validation simulation for the entire 50,000 PIC steps: total wall-clock time and percentage of total wall-clock time for each main component of the PIFE-PIC procedures, namely, “gather”, “particle-push”, “particle-push-comm” (adjust particles at local boundaries and communication among subdomains), “scatter”, “field-solve”, “field-solve-philic” (communication among subdomains and update local potential boundary conditions), and “other” (including particle injection at global boundaries and particle collection at objects). The percentage breakdown shows the procedures of ‘particle-push’ and ‘field-solve’ took the majority of the computing (wall-
Fig. 3.2. Field convergence history of the code validation test case, PCG absolute residual and DDM relative error. The green line on max DDM relative error plot is the DDM tolerance.

clock) time. The computing time of ‘particle-push’ essentially depends on the number of simulation particles in the domain, which also affects the accuracy and smoothness of the source term for Poisson’s equation. Therefore, in practical PIC simulations, large numbers of particles are preferred when computing resources are available. The computing time of ‘field-solve’ essentially depends on 1) the size of each subdomain
Fig. 3.3. Global particle history of the code validation test case. “ns1” represents the number of Species #1 particles which is the number of electrons, while “ns2” represents the number of Species #2 particles which is the number of ions. “ntot” is the total number of particles (electrons plus ions).

Fig. 3.4. Validation of OML sheath solution: PIFE-PIC vs. analytic and 3-D potential contours.

(number of mesh cells and nodes) and 2) the number of DDM iterations. The size of each subdomain can be determined by the domain decomposition configurations, while the number of DDM iterations is affected by the DDM relative error tolerance and the max number of DDM iterations. In the following section, we vary the size of each subdomain and max number of DDM iterations to investigate the strong scaling performance of PIFE-PIC.

4. Parallel Efficiency: Strong Scaling. For most large-scale problems of practical interests, the problem size is usually determined by the physical phenomena to be resolved. Therefore, to test the parallel
Table 3.1

| Computing step                                      | Wall-clock time (s) | Percent of total wall-clock time (%) |
|-----------------------------------------------------|---------------------|--------------------------------------|
| Total wall-clock time                               | 6672.31             | 100.00                               |
| Total gather time                                   | 744.88              | 11.16                                |
| Total particle-push time                            | 3153.67             | 47.26                                |
| Total particle-push-comm (AdjustOuter local) time*  | 1652.67             | 24.77*                               |
| Total scatter time                                  | 48.48               | 3.72                                 |
| Total field-solve time                              | 2129.57             | 31.92                                |
| Total field-solve-phibc (Update Phi BC) time**      | 310.79              | 4.66**                               |
| Total other time                                     | 95.71               | 5.93                                 |

* Included in the ‘particle-push time’
** Included in the ‘field-solve time’

The speedup is defined as $S = T_s / T_p$, where $T_s$ is the serial runtime and $T_p$ is the parallel runtime on $p$ MPI processes. The strong scaling parallel efficiency is then defined as $E = S / p = T_s / (p \cdot T_p)$. We chose two groups of configurations to test the parallel efficiency:

- Group I: Using at most 10 DDM iterations per main-loop PIC step;
- Group II: Using at most 6 DDM iterations per main-loop PIC step.

The timer data was taken over all 20,000 PIC steps. Table 4.1 lists the domain decomposition configurations for each test case. Table 4.2 lists the total wall-clock time, speedup, and parallel efficiency of each case for both Group I and Group II.

Table 4.1

| # of subdomains | DD Configurations | Size of smallest subdomain (cells) | Size of biggest subdomain (cells) |
|-----------------|-------------------|-----------------------------------|----------------------------------|
| 1 (serial)      | 1×1×1             | 100×100×100                       | 100×100×100                      |
| 64              | 4×4×4             | 25×25×25                          | 25×25×25                         |
| 80              | 4×4×5             | 20×20×20                          | 25×25×25                         |
| 100             | 4×5×5             | 20×20×20                          | 25×25×25                         |
| 125             | 5×5×5             | 20×20×20                          | 20×20×20                         |
| 180             | 5×6×6             | 15×15×15                          | 20×20×20                         |
| 216             | 6×6×6             | 15×15×15                          | 17×17×17                         |

Figure 4.1 plots the percentage of total wall-clock time to show the performance of PIFE-PIC across different number of processors for Group I and Group II. The percentage of total wall-clock time breakdown is fairly consistent across all parallel configurations. It also shows the majority of the wall-clock time was always consumed by the field-solve step.

5. Application to Lunar Crater Charging. In this section, we apply PIFE-PIC to simulate the plasma charging at a lunar crater under average solar wind (S.W.) conditions to demonstrate the large-scale simulation capability of PIFE-PIC. In the following, we will first briefly describe the lunar surface charging problem, then introduce the setup of the simulation, and finally present the results and discussion.

5.1. Problem Description. The problem considered is solar wind plasma charging near the lunar surface, specifically, near the lunar craters at the terminator region for lunar exploration missions. The Moon is directly exposed to the solar radiation and various space plasma environments which directly interact with
Table 4.2
Strong Scaling Test Results

| # of sub-domains | Total time $T_1$ (min) | Speedup $S_1$ | Efficiency $E_1$ (%) | Total time $T_{II}$ (min) | Speedup $S_{II}$ | Efficiency $E_{II}$ (%) |
|------------------|------------------------|--------------|----------------------|---------------------------|-----------------|------------------------|
| 1 (serial)       | 12,509                 | 1            | 100                  | 12,509                    | 1               | 100                    |
| 64               | 196                    | 63.78        | 99.66                | 177                       | 70.54           | 110.22                 |
| 80               | 160                    | 78.42        | 98.03                | 141                       | 88.81           | 111.01                 |
| 100              | 149                    | 83.89        | 83.89                | 117                       | 106.87          | 106.87                 |
| 125              | 142                    | 88.26        | 70.61                | 109                       | 114.76          | 91.81                  |
| 180              | 102                    | 123.24       | 68.47                | 77                        | 162.93          | 90.51                  |
| 216              | 93                     | 134.01       | 62.04                | 70                        | 178.16          | 82.48                  |

the lunar surface. A direct consequence of such interactions is surface charging. Observations have found that the potential of the sunlit surface is typically a few tens of volts positive with respect to ambient due to photoelectron emission, while that of the surface in shadow can be hundreds to thousands of volts negative because of the hot electron flux from ambient plasma that can dominate the charging process [8, 17, 22, 23, 24, 55, 57, 62]. Both solar illumination and plasma flow can have a substantial influence on lunar surface charging. At the lunar terminator, the rugged surface terrain, such as that near a crater, generates localized plasma wakes and shadow regions which can lead to strong differential charging at the surface [7, 54, 61]. Both the localized plasma flow field and the charged lunar surface are expected to have substantial influence on the charging of spacecraft/landers/rovers/habitats for future surface missions.

The lunar surface is covered by the lunar regolith layer which separates the solid bedrock from the plasma environment. The regolith layer in most areas is about 4 to 20 meters thick [52, 56]. A complete model of plasma charging on the lunar surface needs to explicitly take into account the properties of the regolith layer, such as permittivity, layer thickness, and the lunar electrical ground.

The serial version of IFE-PIC method has been successfully applied to simulations of lunar plasma charging [29]. In order to illustrate the high performance computing capability of the PIFE-PIC package in this paper, we apply PIFE-PIC to a much larger scale parallel simulation with a larger simulation domain including a lunar crater and much more simulation particles. The plasma environment is chosen to be the average solar wind and photoelectron parameters at the lunar surface [61], as shown in Table 5.1. It is noted here that the Debye length of photoelectrons at 90° Sun elevation angle (1.38 m) is used as the reference length to normalize spatial dimensions in PIFE-PIC.

Table 5.1
Average solar wind and photoelectron (at 90° Sun elevation angle) parameters

|                      | Number density $n$, cm$^{-3}$ | Drifting velocity $v_d$, km/s | Thermal velocity $v_t$, km/s | Temperature $T$, eV | Debye length $\lambda_D$, m |
|----------------------|-------------------------------|-------------------------------|------------------------------|---------------------|----------------------------|
| S.W. electron        | 8.7                           | 468                           | 1453                         | 12                  | 8.73                       |
| S.W. ion             | 8.7                           | 468                           | 31                           | 10                  | N/A*                       |
| Photoelectron        | 64                            | N/A*                          | 622                          | 2.2                 | 1.38                       |

* N/A denotes “not applicable”

5.2. Simulation Setup.

5.2.1. Lunar Crater Geometry and Simulation Domain. In PIFE-PIC, the geometry of the lunar crater is realized through an algebraic equation describing the surface terrain in the form of $z = z(x, y)$ where $z$ denotes the surface height. For the lunar crater considered here, the shape is realized by a few characteristic parameters such as inner-rim radius, outer-rim radius, depth, rim height, etc. (Figure 5.1(a)) according to the Lunar Sourcebook [35]. The specific diameter of a real lunar crater can be measured through NASA Jet Propulsion Laboratory’s website, Moon Trek [53]. The crater considered in this study has these characteristic dimensions: inner-rim radius $10.5 \times 1.38 = 14.49$ m, top-rim radius $20.2 \times 1.38 = 27.88$ m, outer-rim radius $30.9 \times 1.38 = 42.64$ m, and top height $6.7 \times 1.38 = 9.25$ m. Details of the approach to set up the lunar crater
The simulation domain has 200×100×100 = 2 million PIC cells (10 million tetrahedral FE/IFE cells) including half of the lunar crater due to symmetry with respect to the X-Z plane at y = 0 (Figure 5.1(a)). Each PIC cell is a 1.38×1.38×1.38 cube. In physical units, the domain size is approximately 276 m by 138 m by 138 m. At the Z\textsubscript{min} boundary, the simulation domain includes a layer of the lunar bedrock with a thickness of \( L_{\text{bedrock}} = 4.5 \times 1.38 = 6.21 \) m. On top of the bedrock is a layer of dielectric regolith with a thickness of \( L_{\text{regolith}} = (9.5 - 4.5) \times 1.38 = 6.9 \) m. The relative permittivities of the lunar regolith layer and the bedrock are taken to be \( \epsilon_{\text{regolith}} = 4 \) and \( \epsilon_{\text{bedrock}} = 10 \), respectively [35]. 3-D domain decomposition of 8×4×4 (total 128 MPI processes) is used to run the simulation (Figure 5.1(b)).

5.2.2. Particle and Field Boundary Conditions. Particles representing solar wind ions and electrons are pre-loaded and injected into the domain with an angle of 10° towards the surface in the X-Z plane (Figure 5.1(a)). Particles representing photoelectrons are generated at the sunlit regions according to the local sunlight index. At the global X\textsubscript{min}, X\textsubscript{max}, Y\textsubscript{max}, and Z\textsubscript{max} domain boundaries, ambient solar wind particles are injected. Particles hitting the global Y\textsubscript{min} boundary are reflected due to symmetry. Particles hitting the lunar surface are collected and their charges are accumulated to calculate surface charging.
Dirichlet boundary condition of \( \Phi = 0 \) is applied at the \( Z_{\text{max}} \) boundary (the unperturbed solar wind), whereas Neumann boundary condition of zero electric field is applied on all other five domain boundaries. The PCG max iterations was set to 150 with a tolerance of \( 1 \times 10^{-6} \) (for relative residual). The max number of DDM iteration for initial field solution was set to 800 and the max number of DDM iteration for each step within the main PIC loop was set to 200 with a tolerance of \( 1 \times 10^{-3} \). The simulation ran for 20,000 PIC steps.

5.3. Convergence History. The run took about 109 hours to finish 20,000 PIC steps with the time step size of 0.05 (total simulation time till \( \hat{t} = 1,000 \)). Figure 5.2 shows the convergence history of the lunar crater charging simulation including the max absolute PCG residual and max DDM relative error and particle number histories. It is shown that the field solution residuals and relative errors started to level off near PIC step of 10,000 (\( \hat{t} = 500 \)), and at steady state, the entire domain had about 520 million particles. The results presented below are taken at \( \hat{t} = 1,000 \).

5.4. Surface Charging Results. Figure 5.3 illustrate the density contours of solar wind ions, solar wind electrons, photoelectrons, and total space charge near the crater. The solar wind ion and electron density contours clearly exhibit a localized plasma wake formed by the crater rim. The photoelectron density contours clearly exhibit the lack of photoemission in the shadow region. The total space charge density contours show the non-neutral regions associated with the wake caused by the crater rim.

Figure 5.4 illustrates the potential contours of the domain and near the crater. It is shown, for the average solar wind conditions considered here, the surface potential in the sunlit region of the crater is charged to about \( 16 \times 2.2 \approx 35 \) V while the surface in the shadow region is charged to about \(-24 \times 2.2 \approx -53 \) V. It is noted as this length scale is on the order of tens of meters, such differential surface charging will affect the lunar surface activities for exploration missions, such as the risk of discharging/arcing and horizontal/vertical transport of lofted charged lunar dusts.

6. Summary and Conclusion. In this paper, we presented a most recently developed 3D parallel immersed-finite-element particle-in-cell method, namely, PIFE-PIC, for kinetic particle simulations of plasma-material interactions especially electrostatic surface charging. PIFE-PIC is based on the serial non-homogeneous electrostatic IFE-PIC algorithm, which was designed to handle complex interface conditions associated with irregular geometries while maintaining the computational speed Cartesian-mesh-based PIC. 3D domain decomposition is used in both field-solve and particle-push procedures of PIC to distribute the computation among multiple processors. A validation case of 3-D OML sheath of a dielectric sphere im-
mersed in a stationary plasma was carried out and results agreed well with the analytic solution. A series of strong scaling tests were performed to profile the parallel efficiency for a problem of fixed size which has 1 million PIC cells (5 million tetrahedral FE/IFE cells), about 54 million particles, and running 20,000 PIC steps on the Foundry cluster at Missouri University of Science and Technology. Parallel efficiency up to approximately 110% superlinear speedup was achieved.

An application of PIFE-PIC to a larger problem, solar wind plasma charging at a lunar crater, is presented to show the capability of PIFE-PIC for practical problems of science and engineering interest.

Fig. 5.2. The lunar crater simulation convergence history.
The lunar crater charging simulation has 2 million PIC cells (10 million tetrahedral FE/IFE cells), about 520 million particles, and running for 20,000 PIC steps. The simulation finished in about 109 wall-clock hours with domain decomposition of $8 \times 4 \times 4 = 128$ MPI processes. This demonstrates that PIFE-PIC can be utilized to carry out realistic large-scale particle simulations of plasma-material interactions routinely on supercomputers with distributed memory.
(a) Potential contours showing the differential charging near the lunar crater.

(b) Zoom-in view of the potential contours near the lunar crater.

Fig. 5.4. Potential contours of lunar crater charging. The potential values are normalized by 2.2 V and the spatial dimensions are normalized by 1.38 m.
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