Large Eddy Simulations of a Stirred Tank Using the Lattice Boltzmann Method on a Nonuniform Grid

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A nonuniform grid lattice Boltzmann technique previously described by He et al. [1] has been extended to simulate three-dimensional flows in complex geometries. The technique is applied to the computation of the turbulent flow in a stirred tank driven by a standard Rushton turbine. With the nonuniform grid approach, the total CPU time required for a simulation of the flow in a stirred tank can be reduced by roughly 75% and still provide the same spatial accuracy as would be obtained with a uniform high-resolution grid. Statistical results for the computed flow fields will be compared with experimental results (H. Wu and G. K. Patterson, Chem. Eng. Sci. 44, 2207 (1989)) and with simulations by J. G. M. Eggels (Int. J. Heat Fluid Flow 17, 307 (1996)) and J. J. Derksen and H. E. A. Van den Akker (AIChE J. 45, 209 (1999)). The results of the nonuniform mesh simulation are in reasonable agreement with the available experimental data and the results of previous simulations.

Key Words: lattice Boltzmann method; grid refinement; stirred tank; turbulence.

1. INTRODUCTION

Eggels [3] presented the results of large eddy simulations (LES) of a turbulent stirred tank flow using the lattice Boltzmann method (LBM). He showed phase-averaged velocity measurements and snapshots of the flow field in the tank. Derksen and Van den Akker [4] made a more detailed study using a similar approach. They collected statistics not only for the phase-averaged results but also for the phase-resolved data that can reveal the trailing vortex structures. They also presented the contours of the turbulent kinetic energy
and the phase-averaged energy dissipation rate. Comparisons were made between their phase-resolved results and LDA experimental data by Wu and Patterson [2].

Both the simulations and the experimental measurements reveal that the turbulence in the tank is highly inhomogeneous with the largest turbulence dissipation rates confined to a relatively small region near the impeller. These findings suggest that one could reduce the very large computation times needed for LES of stirred tanks by using nonuniform grids with the resolution concentrated in the regions of the tank where the turbulence is most intense. This motivated the present study, which involves an extension of previous work on nonuniform grids to the simulation of the time-dependent, three-dimensional turbulent flow in a stirred tank.

Several researchers have extended the LBM to nonuniform lattices. Nannelli and Succi [5] and Amati et al. [6] developed a finite-volume version of the LBM (FVLBM) that involves a nonuniform coarse lattice that contains several of the basic lattice cells. Filippova and Hänel [7] proposed a local grid refinement method based on the hierarchical grid refinement in conventional CFD methods. He et al. [1] proposed a method that He and Doolen [8] called the “interpolation-supplemented lattice Boltzmann equation model” (ISLBE). This method applies interpolation to update the particle distribution functions on the coarse portion of the grid after the propagation step. He et al. applied the ISLBE method to simulate the flow in a 2D symmetric channel with a sudden expansion on a nonuniform rectangular grid. Since the grid was coarse in one direction, one-dimensional quadratic interpolation was used to obtain a result with second-order accuracy. Later, He and Doolen [8] constructed a curvilinear coordinate system by applying the ISLBE method to simulate the vortex shedding behind a circular cylinder. Two-dimensional second-order upwind interpolation was applied in their simulation. Compared with the other nonuniform grid techniques, the ISLBE method is easy for programming, parallel computation, and incorporation of microscopic interactions in multiphase flow computations. It is valid for grids with arbitrary shapes and for both the BGK scheme and the Somers–Eggels scheme.

In the work reported in this paper, a nonuniform grid technique developed from the ISLBE method is applied to improve the efficiency of the large eddy simulation of a stirred tank; the LDA experimental results reported by Wu and Patterson [2] and Derksen et al. [9, 10] will be compared with the simulation results.

The nonuniform LBM simulations in this paper have several features that were not present in the previous ISLBE method. The simulations to be reported were performed on a three-dimensional FCHC lattice having 18 directions. A second difference is that the ISLBE scheme presented by He et al. [1] applies interpolation after the propagation step to redistribute the particle distribution functions, while the new scheme replaces the propagation step with interpolation. This results in a further improvement in computational efficiency. Finally, the computations are eliminated on grid points that lie outside the computational domain using an approach similar to that employed by Eggels [3] for uniform grids. It will be shown that in addition to reducing the memory requirements by roughly 65%, the total CPU time can be reduced by roughly 75% in the large eddy simulation of a stirred tank without significant sacrifices in accuracy.

The paper is organized as follows. Section 2 presents a concise review of the Somers–Eggels LBM scheme. The nonuniform grid technique is presented in Section 3. The implementation of boundary conditions with the nonuniform grid method is discussed in Section 4. Section 5 discusses the parallelization of the code. Section 6 discusses an “arbitrary computational domain” method in which the computations are skipped for points
lying outside the domain of physical interest. The simulation procedures for a stirred tank are presented in Section 7. The results of the simulations are presented and discussed in Section 8. Finally, the conclusions are presented in Section 9.

2. SOMERS–EGGELS LBM SCHEME

The lattice Boltzmann method (LBM) is a relatively new type of time-dependent Navier–Stokes solver in the incompressible flow regime. The fundamental idea of the LBM is to use a microscopic model of a many-particle system obeying the conservation laws of mass and momentum to simulate fluid flow (Frisch et al. [11]). Unlike conventional CFD schemes based on discretization of the Navier–Stokes equation or equivalent formulations, such as the streamfunction–vorticity equations, the LBM is based on microscopic models and finds solutions of a “mesoscopic” kinetic equation. Chen and Doolen [12] discuss it in detail.

The general form of the lattice Boltzmann equation (LBE) is

$$N_i(x + c_i, t + 1) = N_i(x, t) + \Omega_i(N(x, t)),$$  \hspace{1cm} (1)

where \( \Omega_i(N(x, t)) \) is the collision operator and \( N_i(x, t) \) is a single-particle distribution function, which can be considered as the probability of finding a particle with velocity \( c_i \) on lattice site \( x \) at time \( t \). The collision operator has to obey mass and momentum conservation,

$$\sum_i \Omega_i(N(x, t)) = 0,$$  \hspace{1cm} (2)

$$\sum_i c_i \Omega_i(N(x, t)) = F(x, t),$$  \hspace{1cm} (3)

where \( F(x, t) \) is the external force. The mass density \( \rho \) and fluid velocity \( u \) are related to \( N_i \) and \( c_i \) as follows:

$$\rho(x, t) = \sum_i N_i(x, t),$$  \hspace{1cm} (4)

$$\rho(x, t)u(x, t) = \sum_i c_i N_i(x, t).$$  \hspace{1cm} (5)

The LBM algorithm involves two main computational steps: the collision step and the propagation step. In the collision step, the particle distribution functions are changed by the collision. The postcollision particle distribution functions are calculated by

$$N_i^c(x, t) = N_i(x, t) + \Omega_i(N(x, t)),$$  \hspace{1cm} (6)

where \( N_i^c \) denotes the postcollision value of the particle distribution function. In the propagation step, the particles propagate to their neighbor sites. Thus, the distribution functions on a node are updated by their upstream nodes as follows:

$$N_i(x + c_i, t + 1) = N_i^c(x, t).$$  \hspace{1cm} (7)

It may be seen that Eqs. (6) and (7) imply Eq. (1).
The Somers–Eggels formulation of the LBM model is based on the 4D face-centered-hyper-cubic (FCHC) lattice introduced by d’Humieres et al. [13]. The velocities of the 4D FCHC lattice can be written
\[
c_i = \text{perm}(\pm 1, \pm 1, 0, 0),
\]
where “perm” means the coordinate permutation. The 18 velocities of the three-dimensional projection of the FCHC lattice are given by (see Fig. 1)
\[
c_i = \text{perm}(\pm 1, 0, 0), \text{perm}(\pm 1, \pm 1, 0).
\]

The differential form of Eq. (1) can be derived from the first-order Taylor expansions of \(N_i(x + c_i, t + 1)\),
\[
\frac{\partial N_i}{\partial t} + c_i \frac{\partial N_i}{\partial x} = \Omega_i(N),
\]
where \(c_{i\alpha}\) is the component of the \(i\)th velocity vector \(c_i\) in the Cartesian coordinate direction \(\alpha\). Repeated indices are understood to be summed over in Eq. (10) and subsequent equations except where noted. The index \(i\) in Eq. (10) is not summed over.

Assuming that the magnitude of the fluid velocity, \(u\), is much smaller than unity and the lattice gas is close to equilibrium, they obtained an expression for the collision operator \(\Omega_i\) in Eq. (1) by using an asymptotic expression for \(N_i\). The asymptotic result for \(N_i\) was obtained by Frisch et al. [11] using a multiple time-scale analysis together with an expansion of \(N_i\) around the equilibrium solution. The results are as follows:
\[
N_i = \frac{m_i \rho}{24} \left[ 1 + 2c_{i\alpha}u_\alpha + 3c_{i\alpha}c_{j\beta}u_\alpha u_\beta - \frac{3}{2}u_\alpha u_\alpha - 6\nu \left( c_{i\alpha} \frac{\partial c_{j\beta}u_\beta}{\partial x_\alpha} - \frac{1}{2} \frac{\partial u_\alpha}{\partial x_\alpha} \right) \right],
\]
\[
\Omega_i(N) = \frac{m_i \rho}{12} \left( c_{i\alpha} \frac{\partial c_{j\beta}u_\beta}{\partial x_\alpha} - \frac{1}{2} \frac{\partial u_\alpha}{\partial x_\alpha} \right) + \frac{m_i}{12} c_{i\alpha} F_\alpha.
\]
A summation over all directions $i$ of Eq. (10) with the mass conservation constraint on $\Omega_i$ given in Eq. (2) yields the continuity equation:

$$\frac{\partial \rho}{\partial t} + \frac{\partial \rho u_\alpha}{\partial x_\alpha} = 0.$$  (13)

From Eqs. (11) and (10) and from the symmetry relations (see Eggels and Somers [14]) with a summation over all velocity directions, one can obtain the following equation:

$$\frac{\partial \rho u_\alpha}{\partial t} + \frac{\partial \rho u_\alpha u_\beta}{\partial x_\beta} = -\frac{\partial p}{\partial x_\alpha} + \frac{\partial}{\partial x_\beta} \left[ \nu p \left( \frac{\partial u_\beta}{\partial x_\alpha} + \frac{\partial u_\alpha}{\partial x_\beta} \right) \right] - \frac{1}{2} \frac{\partial}{\partial x_\alpha} \left( \nu p \frac{\partial u_\beta}{\partial x_\beta} \right) + F_\alpha. \quad (14)$$

Equation (14) corresponds to the Navier–Stokes equation in the incompressible limit, with the following equation of state for the pressure:

$$p = \frac{1}{2} \rho \left( 1 - \frac{1}{2} u_\alpha u_\alpha \right). \quad (15)$$

The explicit dependence of the pressure on the velocity is unphysical. In single relaxation time schemes the dependence can be avoided by adding rest particles. However, in practice, it does not create serious problems as long as $|u_\alpha| \ll 1$.

To solve the LBE, Eggels and Somers [14] introduced a filter matrix $E_{ik}$ and solution vectors $\alpha_k^\pm$ by factorizing the asymptotic expression for $N_i$. The postcollision distribution function $N_i^+(x, t)$ is calculated as follows:

$$N_i^+(x, t) = \frac{m_i}{24} \sum_{k=1}^{n} E_{ik} \alpha_k^+ (x, t). \quad (16)$$

The vector $\alpha_k^-$ is calculated as

$$\alpha_k^-(x, t) = \sum_{i=1}^{n} E_{ki}' N_i(x, t), \quad (17)$$

where $E_{ki}'$ satisfies

$$\frac{m_i}{24} E_{ik} E_{ki}' = I. \quad (18)$$

The Somers–Eggels formulation computes the particle distribution function by multiplying the two matrices $E$ and $\alpha^\pm$. The details of the method may be found in Eggels and Somers [14].

### 3. NONUNIFORM GRID TECHNIQUE IN LBM

Figure 2 shows a cross section of a nonuniform grid based on the 18-node FCHC lattice. In the propagation step, particles with velocities in the fifth direction would move as indicated in the figure. On the fine portion of the grid, each particle will move to a neighboring lattice point. However, on the coarse portion of the grid, a particle will move to an interstitial position. It is not necessary to perform interpolation to update the single-particle distribution
function on a fine lattice point unless it is adjacent to a coarse portion of the grid because every node can be updated by its upstream nodes during the propagation step. However, on the coarse portion of the grid, no particle can reach a given lattice point in one time step since the length of the velocity vector is set by the fine grid. Thus, one must use interpolation to update the single-particle distribution function. This approach was developed by He et al. [1].

The nonuniform LBM simulations in this paper have several features that were not present in the ISLBE method developed by He et al. [1]. The simulations to be reported were performed on a three-dimensional FCHC lattice having 18 directions. This lattice is shown in Fig. 1. One can use one-dimensional interpolation on square or cubic grids if the coarse grid spacings have the same values in all the orthogonal coordinate directions. This is because the particles move along a straight line connecting the nodes in this case (see Fig. 2). Applying quadratic one-dimensional interpolation is the most efficient method for the three-dimensional nonuniform LBM scheme. A second difference is that the ISLBE scheme developed by He et al. applies interpolation after the propagation step to redistribute the particle distribution functions, while the new scheme replaces the propagation step with interpolation. In this paper, it will be shown that in addition to reducing the memory requirements by roughly 65%, the total CPU time can be reduced by roughly 75% in the large eddy simulation of a stirred tank.

The quadratic one-dimensional interpolation through three known points $f(x_k)$ [$k = 1, 2, 3$] is given by

$$y(x) = f(x_0) \frac{(x - x_1)(x - x_2)}{(x_0 - x_1)(x_0 - x_2)} + f(x_1) \frac{(x - x_0)(x - x_2)}{(x_1 - x_0)(x_1 - x_2)}$$

$$+ f(x_2) \frac{(x - x_0)(x - x_1)}{(x_2 - x_0)(x_2 - x_1)},$$

where $x$ is the point of interest.
It will be assumed in what follows that the coarse lattice spacing is an integer multiple of the fine grid spacing. The symbol $n_c$ denotes the ratio of the coarse grid spacing to the fine grid spacing.

One must consider three cases that arise in the implementation of interpolation in the nonuniform grid technique. These cases are shown in Fig. 3. In the coarse portion of the grid, quadratic one-dimensional interpolation can be applied to the three-dimensional cubic grid in our simulation because particles always propagate along a line parallel to a particular velocity. As shown in case 1 of Fig. 3, one can apply Eq. (19) to interpolate the postpropagation distribution function at the coarse grid node that is denoted by the square symbol because no particle can stream there in the propagation step. If $x$ represents the three-dimensional position vector of that node, then the position vectors of the three points

\[ (X - n_c c_i + c_i), (X), (X + n_c c_i + c_i) \]

denotes the nodes where interpolation is needed.

\[ (X - 2n_c c_i + c_i), (X - n_c c_i + c_i), (X), (X + c_i) \]

Case 2

\[ (X - n_c c_i), (X - n_c c_i + c_i), (X), (X + c_i) \]

Case 3

\[ (X - n_c c_i), (X - n_c c_i + c_i), (X), (X + c_i) \]

Case 3

\[ (X - n_c c_i), (X - n_c c_i + c_i), (X), (X + c_i) \]

Case 3

\[ (X - n_c c_i), (X - n_c c_i + c_i), (X), (X + c_i) \]

FIG. 3. Three cases in which quadratic one-dimensional interpolation is applied in the nonuniform grid technique.
that are denoted by circles are \( x - n_c \mathbf{c}_i + \mathbf{c}_i \), \( x + \mathbf{c}_i \), and \( x + n_c \mathbf{c}_i + \mathbf{c}_i \). The values of the postpropagation distribution function of the positions that are denoted by circles are already known after the propagation step. Therefore, one can interpolate the unknown value of the postpropagation function on the position denoted by the square from them. One may replace \( y(x), f(x_0), f(x_1), f(x_2) \) by the corresponding postpropagation distribution functions and rewrite Eq. (19) as

\[
N_i(x, t + 1) = a_1 N_i(x - n_c \mathbf{c}_i + \mathbf{c}_i, t + 1) + b_1 N_i(x + \mathbf{c}_i, t + 1) + c_1 N_i(x + n_c \mathbf{c}_i + \mathbf{c}_i, t + 1),
\]

where

\[
a_1 = \frac{n_c + 1}{2n_c^2}, \quad b_1 = \frac{n_c^2 - 1}{n_c}, \quad c_1 = \frac{1 - n_c}{2n_c^2}.
\]

Then, one can apply Eq. (7) to Eq. (20) and obtain an expression involving postcollision distribution functions:

\[
N_i(x, t + 1) = a_1 N^c_i(x - n_c \mathbf{c}_i, t) + b_1 N^c_i(x, t) + c_1 N^c_i(x + n_c \mathbf{c}_i, t).
\]

Case 2 of Fig. 3 shows a situation in which a particle crosses the boundary separating the coarse and fine grids and moves out of the coarse portion. In this case, applying the method used in case 1, one can rewrite Eq. (19) to update the postpropagation distribution function on the position denoted by the square; i.e.,

\[
N_i(x, t + 1) = a_2 N^c_i(x - 2n_c \mathbf{c}_i, t) + b_2 N^c_i(x - n_c \mathbf{c}_i, t) + c_2 N^c_i(x, t),
\]

where

\[
a_2 = \frac{n_c - 1}{2n_c^2}, \quad b_2 = \frac{2n_c^2 - 1}{n_c^2}, \quad c_2 = \frac{2n_c^2 - 3n_c + 1}{n_c^2}.
\]

For some nodes in a corner of the coarse portion, the situation will be like that shown in case 3 of Fig. 3. Applying the same method, one can rewrite Eq. (19) to update the postpropagation distribution function on the square’s position; i.e.,

\[
N_i(x, t + 1) = a_3 N^c_i(x - n_c \mathbf{c}_i - \mathbf{c}_i, t) + N^c_i(x - n_c \mathbf{c}_i, t) - a_3 N^c_i(x, t),
\]

where

\[
a_3 = \frac{1 - n_c}{n_c + 1}.
\]

Since the right hand sides of Eqs. (21) and (22) involve the postcollision distribution functions \( N^c_i \), one can replace the propagation step with an interpolation step by applying these equations after the collision step.

It is necessary to discuss the fine interface that connects the coarse grid and the fine grid (see Fig. 4). It will be understood in what follows that the interfaces belong to the fine grid. As shown in Fig. 4, the nodes on the fine interface can be updated by their upstream nodes
in the fine region, but they cannot be updated directly by their upstream nodes in the coarse region. Thus, on the nodes of the fine interfaces, interpolation will be needed to update the distribution functions with velocities that point to the fine region. Since this is the same as case 2, which was discussed previously, one can apply Eq. (22) to this situation. However, as may be seen from Fig. 4, not all of the nodes on a fine interface can be updated by this interpolation method. In such cases, the other fine-grid nodes on the interfaces will be interpolated from their neighboring nodes, which have been updated by interpolation from the neighboring coarse planes. Figure 5 shows a section of an interface and illustrates

\[\square\] denotes the nodes where interpolation is needed.

\[\circ\] denotes the updated nodes, which are used for interpolation.
the points that would be used to perform the two-dimensional interpolation that would be needed to update the distribution function.

4. BOUNDARY CONDITIONS

In the LBM, the physical boundary usually lies between lattice points. The neighboring lattice points that are outside the physical boundary constitute the “computational boundary” in this paper. For simplicity, the computational boundary will be referred to as the “boundary” in what follows.

As discussed in Section 2, the LBM has two main computational steps: the collision step and the propagation step. On the boundary nodes, however, no calculation is made for the collision step. The boundary nodes are used only for the implementation of boundary conditions. To implement boundary conditions, another computational step, the boundary condition step, is placed between the two main computational steps. The implementation of boundary conditions is performed in two steps.

1. In the boundary condition step, the particle distribution functions on the boundary are updated from the postcollision distribution functions on the neighboring interior grid nodes. Different boundary conditions require different updating rules.

2. In the subsequent propagation step, the distribution functions at the interior points are set equal to the values of the distribution functions at the upstream boundary points.

Therefore, different boundary conditions have significantly different effects on the interior nodes.

4.1. Boundary Conditions for the LBM

The bounce-back boundary condition imposes a no-slip boundary condition that requires the fluid locally to have the same tangential velocity as the wall. In the boundary condition portion of a time step, the postcollision distribution function at a boundary grid point with a velocity directed toward a neighboring interior point is set equal to the postcollision distribution function with the opposite velocity at the interior point. If the wall is stationary, the bounce-back boundary condition can be enforced on the boundary nodes by

\[ N_\xi^c(x_b, t) = N_{\xi^-}^c(x_{nb}, t), \]

where \( \xi \) and \( \xi^- \) represent opposite directions, \( c_\xi \) are the velocities moving into the computational domain, and \( c_{\xi^-} \) are the velocities moving out (see Fig. 6), \( N^c \) are postcollision

![FIG. 6. Bounce-back boundary condition.](image-url)
LES OF A STIRRED TANK USING LBM

particle distribution functions, $x_b$ is the position of a boundary node, and $x_{nb}$ is the interior node next to the boundary. It follows from their definitions that

$$x_{nb} = x_b + c_\xi.$$  

In the subsequent propagation step, the interior node next to the boundary will be updated by the boundary nodes as follows:

$$N_\xi(x_{nb}, t + 1) = N_\xi(x_b, t).$$  

Therefore, one can combine Eqs. (24) and (26) and obtain

$$N_\xi(x_{nb}, t + 1) = N_\xi(x_{nb}, t).$$  

If the wall is moving with velocity $u_b$, one has to modify Eq. (27) to enforce the moving-wall no-slip boundary condition. In the Somers–Eggels scheme, the equation for the moving-wall no-slip boundary condition can be obtained from Eq. (11) (see Eggels and Somers [14]):

$$N_\xi(x_{nb}, t + 1) = N_\xi(x_{nb}, t) + \frac{m_\xi(p(x_{nb}, t))}{6} [c_\xi \cdot u_b].$$  

The symmetric boundary condition is applied on a stress free surface. It is also called the free-slip boundary condition. In the boundary condition step, the postcollision distribution functions at boundary grid points with velocities directed toward a neighboring interior point are set equal to the postcollision distribution function with the symmetric velocity at the interior point. Figure 7 shows the implementation of the symmetric boundary condition. It can be expressed as

$$N_\xi(x', t) = N_\xi(x, t),$$  

where $\xi^s$ is the direction symmetric to $\xi$ and $x'$ are the boundary positions that are symmetric to $x$ across the physical boundary.

FIG. 7. Symmetric boundary condition.
4.2. Boundary Conditions for the Nonuniform Grid Technique

To implement the boundary conditions for coarse grids, it is helpful to discuss the imposition of these boundary conditions in the LBM. In this paper, the boundaries of both the coarse grid and the fine grid are fine and the distance between the boundary and the interior nodes next to the boundary is equal to a fine lattice spacing. This is shown in Fig. 8. As discussed previously in this section, the boundary conditions are performed in two steps. For the fine grid, no matter what updating rules are applied in the boundary condition step, the updated particles on the boundary nodes can always stream to the interior fine grid nodes. However, for the coarse grid, only some of the boundary nodes can be updated in the boundary condition step (see Fig. 9). As a result, for most updating rules, the updated particles on the boundary nodes can only stream to the interstitial positions of the coarse grid next to the boundary in the subsequent propagation step (see Fig. 10).

For the no-slip boundary condition, Eq. (27) shows that the postpropagation distribution at an interior node next to the boundary can be updated by the postcollision distribution function at the same position. The values of \( N_{c}^{\xi}(x_{nb}, t) \) are known for the coarse grid. Thus, interpolation is not needed in this case.

However, for the free-slip boundary condition, let us first consider the boundary condition step. Figure 9 shows that, when applying the free-slip boundary condition, the distribution

\[
\text{boundary condition step}
\]

function for velocity $c_i$ on a boundary node will be set equal to the distribution function for velocity $c_s$ on the neighboring interior node. In the propagation step, the updated boundary nodes update interstitial sites, as shown in Fig. 10. Thus, one needs to apply interpolation to update the coarse grid nodes.

- One may apply interpolation to the nodes that are denoted by squares on the neighboring coarse grid before the boundary condition step, as shown in Fig. 11.
- In the boundary condition step, the value of the distribution function at the interpolated coarse nodes will be assigned to the corresponding nodes on the boundary. (This is shown by the arrow ↑ in Fig. 11.)
- The boundary nodes will update coarse grid nodes in the propagation step.

5. PARALLEL COMPUTATION FOR THE NONUNIFORM GRID

To perform the parallelization, one needs to send border values of a process to neighboring processes and receive border values from neighboring processes in each time step because the computation in a process may need the border values of its neighboring processes. According to the definition of ghost points, “the elements of the array that are used to hold data from other processes” (see Gropp et al. [15]), one may refer to the border values of the neighboring processes as ghost points. Thus, in performing parallel computation, one may

![Diagram of interpolation](image-url)
need to transfer the data of some ghost point lines or areas, or ghost point zones in some three-dimensional cases.

In the uniform grid LES program described by Derksen and Van den Akker [4], the data transfer of some ghost point planes is applied in the message-passing step. However, in a three-dimensional parallel implementation in the coarse portion of the grid, one should transfer the data of some coarse ghost point zones instead of some ghost point planes. This is because, in the worst case, the interpolation for a border coarse grid node needs two coarse ghost points (see Fig. 12).

However, the cost of the data transfer operation is still small because the number of nodes in the coarse ghost point zone that contains two planes of the coarse grid is still less than that of a fine ghost point plane if \( n_c > \sqrt{2} \). Figure 13 shows a two-step process to transfer data. Dashed boxes indicate the ghost point zones; the data to be moved is shaded (see Gropp et al. [15]).
6. ARBITRARY COMPUTATIONAL DOMAIN METHOD

In many cases, the computational domain is a complicated geometry. If one uses a rectangular or cubic computational domain for the calculation in these cases, a large portion of the computational cost would be associated with points outside the region of physical interests.

Based on earlier work by Ladd [16], an arbitrary computational domain method was developed to solve this problem. The method is as follows:

1. Determine the functions that describe the boundaries of the computational domain.
2. Classify the lattice points that are inside the actual computational domain and outside the domain. (For example, one may introduce a symbol $\beta(x)$ to perform the classification by assigning the values of $\beta(x) = 1$ for the region of physical interest and $\beta(x) = 0$ for the points outside the region of physical interest.)
3. For some velocities, the particles on the interior nodes next to the boundary will move out of the interior region of physical interest. Test each velocity and record the information of the positions of these nodes, $x_{nb}$, the velocities $c_{-\xi}$ for which the particles on these nodes could move out of the actual computational domain, and the positions when they move out of the interior region of physical interest $x_{nb} + c_{-\xi}$, where the subscript $-\xi$ denotes the direction pointing from the neighboring interior nodes to the boundary. The subscript $nb$ indicates the position of the interior nodes next to the boundary.

All the operations above are performed in a separate program with the purpose of providing the necessary information for the main computational program to apply boundary condition and skip the lattice nodes outside the region of physical interest. Therefore, there is no additional cost to the main computational program. To illustrate how the main program can use the information above to apply boundary conditions, consider the no-slip boundary condition as an example. The key to applying boundary conditions is to enable the particles on the boundary nodes to stream to the neighbor interior nodes in the propagation step. It could be written as

$$N_\xi(x_{nb}, t + 1) = N_\xi(x_{nb} - c_\xi, t),$$

where the subscript $\xi$ denotes the direction pointing from the boundary to the neighboring interior nodes, where $-\xi$ always indicates the direction opposite to $\xi$, and where $x_{nb} - c$ denotes the corresponding upstream boundary nodes and is equal to $x_b$ (see Fig. 14).

To apply Eq. (30), one should obtain the value of $N_\xi(x_{nb} - c_\xi, t)$ before the propagation step. For the no-slip boundary condition, one may calculate $N_\xi(x_{nb} - c_\xi, t)$ in the previous boundary condition step as follows:

$$N_\xi(x_{nb} - c_\xi, t) = N_\xi(x_{nb}, t).$$

The quantities of $x_{nb} - c_\xi$, $x_{nb}$, and $-\xi$ are known from step 3. Thus, the main computational program can use the information to apply the no-slip boundary condition to complicated geometries correctly and skip the lattice nodes outside the region of physical interests by only calculating the nodes with $\beta(x) = 1$. 


7. SIMULATIONS

The flow simulation was performed with a modification of a program developed by Derksen and Van den Akker [4] to simulate a stirred tank. The program used by Derksen and Van den Akker makes use of a version of the LBM described by Eggels and Somers [14]. The program also makes use of the Smagorinsky [17] subgrid scale model to account for turbulent eddies with length scales smaller than the computational lattice spacing. Eggels [3] computed statistical results for the mean flow. Derksen and Van den Akker [4] computed time-averaged and phase-averaged results for the mean flow and turbulence statistics. These quantities have been computed in the present study to permit a comparison with the above results.

7.1. Computational Procedure

The LES program consists of six main computational steps in each time step.

1. Determine the elements of the solution vector $\alpha_k^-(x, t)$ from the postpropagation distribution function $N_i(x, t)$ of the last time step using Eq. (17).

2. Compute the elements of the solution vectors $\alpha_k^+(x, t)$ from $\alpha_k^-(x, t)$ (see Somers [18] and Eggels and Somers [14]).

3. Compute the postcollision distribution function $N_i^c(x, t)$ field from $\alpha_k^+(x, t)$ using Eq. (16).

4. Implement boundary conditions to update $N_i^c(x, t)$ on the boundary nodes.

5. In the message-passing step, send border values of $N_i^c(x, t)$ to neighbors and receive border values from neighbors in order to implement parallel computation.

6. In the propagation step, the postcollision distribution function $N_i^c(x, t)$ associated with the grid point at position $x$ moves to the grid point at $x + c_i$ to become $N_i(x + c_i, t + 1)$ for the next time step.

Most modifications have been made on the last three steps to perform a nonuniform LES. In the boundary condition step, the bounce-back boundary condition is imposed on the tank wall, baffle plates, and bottom of the tank, the free-slip boundary condition is applied on the top of the tank, and the adaptive force-field method (Derksen and Van den Akker [4]) is implemented on the impeller. The details of the implementation of boundary conditions for the nonuniform grid are discussed in Sections 4.1 and 4.2. The interpolation methods
discussed in Section 3 are applied in the propagation step to update the distribution functions in the coarse portion of the grid. The arbitrary computational domain method discussed in Section 6 is applied to set up the geometry of the tank wall and the baffles and to eliminate the computation of the nodes outside the tank. In the message-passing step, the method described in Section 5 is implemented to handle the message passing in the coarse portion.

7.2. Large Eddy Simulation

A direct simulation of a stirred tank with an industrially relevant Reynolds number (Re > 10^4) is not feasible because of the enormous numbers of grid points and time steps that would be required. However, in a large eddy simulation (LES), the effect that the small scales have on the flow is assumed to be universal and isotropic so that one may use a simple subgrid-scale model. Thus, the large eddy simulation can be performed with a relatively smaller resolution and fewer time steps. In this research, the standard Smagorinsky eddy viscosity model (Smagorinsky [17]) is applied. The model uses a subgrid-scale eddy viscosity \( v_t \),

\[
v_t = \lambda_{mix}^2 \sqrt{S^2},
\]

where \( \lambda_{mix} \) is the mixing length of subgrid-scale motion and \( S^2 \) the resolved deformation rate,

\[
S^2 = \frac{1}{2} \left( \frac{\partial u_\alpha}{\partial x_\beta} + \frac{\partial u_\beta}{\partial x_\alpha} - \frac{2}{3} \delta_{\alpha\beta} \nabla \cdot \mathbf{u} \right)^2,
\]

where \( \delta_{\alpha\beta} \) is the Kronecker delta. The ratio between the mixing length and the lattice spacing \( \Delta \) is set to a constant:

\[
c_s = \frac{\lambda_{mix}}{\Delta}.
\]

In the present study, \( c_s \) was chosen to be 0.12, which is within the range of commonly used values in shear-driven turbulence (see Piomelli et al. [19]). This value was selected to facilitate comparisons with the results obtained by Derksen and Van den Akker. In the coarse portion of the nonuniform grid, the values of the lattice spacing and the mixing length are twice as big as those in the fine portion. The total viscosity is \( \nu + v_t \) in the subgrid-scale model. Through the solution vectors of the Somers–Eggels LBM scheme, the stresses that are needed for the calculation of \( v_t \) can be obtained locally. Thus, the large eddy simulation does not destroy the locality of the lattice Boltzmann scheme.

7.3. Parameter Choices

The configuration of the stirred tank together with a sketch of the front and top view of the impeller is shown in Fig. 15. The Reynolds number is defined as \( Re = ND^2/\nu \), where \( N \) is the rotational speed of the impeller, \( D \) is the impeller diameter, and \( \nu \) is the kinematic viscosity of the flow. The four baffles are placed at the perimeter of the tank. The axial level \( z = 0 \) corresponds to the impeller disk plane.

Table I gives the values of the geometrical quantities, namely the tank height and diameter, the impeller diameter, the blade dimensions, the width of the tank baffles, and the diameter of the impeller shaft. In all cases, the quantities are made dimensionless in terms of the
lattice spacing. The values of the quantities in the simulations reported by Derksen and Van den Akker and by Eggels are also given in Table I.

The rotational speed, $N$, of the impeller is limited by the incompressibility condition in the LBM $|u|^2 \ll 1$. (In the lattice units used in this paper, the speed of sound is unity.) The impeller tip speed, $v_{\text{tip}} = \pi ND$, should be much smaller than unity. The Reynolds number can be varied by varying the kinematic viscosity. In this study, the Reynolds number was 29,000 because of the availability of experimental data (Wu and Patterson [2] and Derksen et al. [9, 10]) and simulation results (Derksen and Van den Akker [4]). Table II gives the values of the dimensionless physical parameters, namely the kinematic viscosity of the fluid, the density of the fluid, the impeller cyclic velocity, the impeller tip velocity, and the Reynolds number. The corresponding values for the Derksen–Van den Akker simulation and Eggels’ simulation are given for comparison.

In Derksen et al.’s and Eggels’ studies, low-resolution simulations were started from a quiescent state and continued until steady state was achieved. Then, the flow fields were interpolated to a higher resolution grid. Finally, high-resolution simulations were started from the interpolated flow fields. However, for the nonuniform simulation, since the coarse mesh covers most of the computational domain, one can perform the simulation directly from a quiescent state. According to the temporal monitoring of the velocity and the

### Table I

<table>
<thead>
<tr>
<th>Parameters for the Geometry of the Stirred Tank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tank height</td>
</tr>
<tr>
<td>Tank diameter</td>
</tr>
<tr>
<td>Impeller diameter</td>
</tr>
<tr>
<td>Length of blades</td>
</tr>
<tr>
<td>Width of blades</td>
</tr>
<tr>
<td>Width of tank baffles</td>
</tr>
<tr>
<td>Diameter of the impeller shaft</td>
</tr>
</tbody>
</table>
energy levels in a subsection of a stirred tank (Egels [3]), at least 30 impeller revolutions are needed to reach a quasi-steady state when starting from quiescent states. Thus, the nonuniform grid LES starts from a quiescent state and runs for 30 revolutions to develop a steady state. In the nonuniform grid simulation, the impeller makes a full revolution in 3000 time steps. Therefore the impeller tip velocity is much smaller than that of Derksen and Van den Akker’s and Eggels’s simulations. A small tip velocity will be beneficial for multiphase stirred tank simulations that will be performed in the future because large velocity fields will create numerical instability for multiphase simulations with the LBM. The simulation data were collected and statistically processed for 20 revolutions after reaching the steady state. This sampling period is smaller than that of Derksen et al.’s and Eggels’ simulations only because of the CPU time limits. Table III gives computational parameters for the grids that were considered in the present study, namely the number of time steps for a revolution, the number of revolutions to reach a steady state, the number of revolutions to collect the statistics, the numbers of grid points in the x, y, and z directions, the total number of grid points, and the ratio of the total number of grid points for the nonuniform simulation to those in the other simulations.

**TABLE II**

Parameters for the Physical Properties of the Flow

<table>
<thead>
<tr>
<th></th>
<th>Nonuniform grid simulation</th>
<th>Derksen et al.’s simulation</th>
<th>Eggels’ simulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kinematic viscosity</td>
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<td>$7.356 \times 10^{-5}$</td>
<td>$3.988 \times 10^{-5}$</td>
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<tr>
<td>Density</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Impeller cyclic velocity</td>
<td>$2\pi/3000$</td>
<td>$2\pi/1600$</td>
<td>$2\pi/1500$</td>
</tr>
<tr>
<td>Impeller tip velocity</td>
<td>0.084</td>
<td>0.12</td>
<td>0.17</td>
</tr>
<tr>
<td>Reynolds number</td>
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<td>29000</td>
<td>107000</td>
</tr>
</tbody>
</table>

**TABLE III**

Computational Parameters

<table>
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<th>Nonuniform grid simulation</th>
<th>Derksen and Van den Akker’s simulation</th>
<th>Eggels’ simulation</th>
</tr>
</thead>
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<tr>
<td></td>
<td>Low resolution</td>
<td>High resolution</td>
<td>Low resolution</td>
</tr>
<tr>
<td>Number of time steps for</td>
<td>3000</td>
<td>1000</td>
<td>750</td>
</tr>
<tr>
<td>a revolution</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of revolutions to</td>
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<td>20</td>
<td>30</td>
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<tr>
<td>reach a steady state</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of revolutions to</td>
<td>20</td>
<td>25</td>
<td>40</td>
</tr>
<tr>
<td>collect the statistics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of grid points in</td>
<td>92 in the coarse portion,</td>
<td>120</td>
<td>120</td>
</tr>
<tr>
<td>the axial direction (x-) 57 in the fine portion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of grid points in</td>
<td>120 in the coarse portion, 120</td>
<td>180</td>
<td>120</td>
</tr>
<tr>
<td>other directions (y-,z-) 240 in the fine portion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total number of grid</td>
<td>4,608,000</td>
<td>1,728,000</td>
<td>1,728,000</td>
</tr>
<tr>
<td>points</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normalized total number</td>
<td>2.667</td>
<td>1</td>
<td>3.375</td>
</tr>
<tr>
<td>of grid points</td>
<td></td>
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<td></td>
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TABLE IV

Computational Parameters

<table>
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<th>Uniform grid simulation</th>
<th>180(^\circ) grid</th>
<th>240(^\circ) grid</th>
</tr>
</thead>
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<tr>
<td>Total number of grid nodes</td>
<td>4,608,000</td>
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<td>13,824,000</td>
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<tr>
<td>Normalized number of grid nodes</td>
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<td>1.27</td>
<td>3</td>
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<tr>
<td>Normalized CPU time for a time step</td>
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</tr>
<tr>
<td>Normalized CPU time for a revolution</td>
<td>1</td>
<td>0.75</td>
<td>1.9</td>
</tr>
</tbody>
</table>

8. RESULTS

In the nonuniform simulation, one-dimensional quadratic interpolation is implemented in the coarse portion of the grid using the method in Section 4. Based on computational tests, the CPU time consumed on the interpolation for a coarse node is almost the same as that of the calculation for a fine node in the propagation step. This accounts for much of the efficiency of the interpolation method. Computational tests have also been made.

FIG. 16. An instantaneous velocity field in a vertical cross section of the stirred tank.
for uniform simulations with different resolutions to compare their CPU time. The uniform simulations with 180³ grid points and 240³ grid points represent the simulations of Derksen and Van den Akker [4] and Eggels [3], respectively. The results are shown in Table IV.

The general flow pattern for the nonuniform simulation of the stirred tank is illustrated in Figs. 16–19, showing instantaneous and phase-averaged velocity fields in a vertical cross section of the tank and in the horizontal plane z = 0 where the center of the impeller is located (see Fig. 15). In order to show the flow structure clearly, these plots are shown with only half of the resolution of the nonuniform simulation.

In Fig. 16, strong radial outflows as well as trailing vortices can be observed. The trailing vortices are produced near the edge of the impeller blades and sent out by the strong outflows toward the tank wall. It seems that the vortices can travel over a long distance until they come near the tank wall. One can also identify the shape of the shaft and the impeller. The continuous motion of flow passing through the interfaces of the coarse and fine grid suggests that the method for handling the interaction of the coarse grid and the fine grid is proper. Several trailing vortices can be observed in Fig. 17. Figures 18 and 19 show the mean flow.

**FIG. 17.** A velocity field of the turbulent flow in a horizontal plane that contains the middle of the impeller (z = 0).
With the exception of small vortices near the baffles, no vortices are seen in the mean flow. In Fig. 17, one can identify the baffles and the tank wall. The impeller is not easy to see due to the complexity of the flow field. However, one can still identify the position of the six blades and the impeller disk.

The trailing vortex structure that is developed by the impeller blades and advected into the bulk of the tank with the impeller outflow has high levels of turbulent activity and large velocity gradients. Hinze [20] and Prince et al. [21] argued that bubble breakup was probably caused by the eddies that were equal to or marginally smaller than the bubble size. Thus, this structure is very important for mixing and, possibly, bubble breakup in two phase flows. Derksen and Van den Akker [4] proposed the comparison of the phase-resolved results with the experimental data to study trailing vortices. In this paper, a comparison between phase-resolved statistical results of the nonuniform simulation and the experimental results by Derksen et al. [9, 10] is also made. Figure 20 shows that the agreement between the simulated and measured phase-resolved flow fields is good. To show the comparison clearly, the velocity fields are plotted with half the resolution of the simulation (i.e., the same resolution as the experimental measurements).

Figures 21 and 22 show the time-averaged results of the radial and tangential velocity components in the impeller outflow for various values of $2r/D$. The results reveal an
acceptable agreement with the experimental data of Wu and Patterson [2] and have an accuracy comparable to the results of Eggels [3] and Derksen and Van den Akker [4]. The maximum tangential velocities are in better agreement with experiment than the results of Derksen and Van den Akker. This is probably because of the higher resolution in the fine region.

The errors were calculated as follows:

\[ E_{rr} = \frac{|U_{20}^* - U_{10}^*|}{U_{20}^*}, \]  \hfill (35)

In the nonuniform grid simulation, the statistics collection interval was 20 revolutions. The statistical results are influenced by the limited length of the interval because the statistical error is significant for a short interval. The deviations of the simulation results from the experimental results for different radial positions are calculated according to

\[ D_v = \frac{|U_{sim}^* - U_{exp}^*|}{U_{max, exp}^*}, \]  \hfill (36)
FIG. 20. Phase-resolved flow fields in the vicinity of the impeller for $\theta = 10^\circ$, $\theta = 19^\circ$, $\theta = 31^\circ$, and $\theta = 40^\circ$. The flow fields on the top are the nonuniform simulation results. The bottom fields are experimental results by Derksen et al. [9, 10].
FIG. 20—Continued

θ = 31°

θ = 40°
### TABLE V
Deviation for Radial Velocity

<table>
<thead>
<tr>
<th>$2r/D$</th>
<th>Simulations</th>
<th>$2z/w$</th>
<th>$-1.6$</th>
<th>$-1.1$</th>
<th>$-0.55$</th>
<th>$0.0$</th>
<th>$0.55$</th>
<th>$1.1$</th>
<th>$1.6$</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.07</td>
<td>10 revolutions</td>
<td>8%</td>
<td>13%</td>
<td>12%</td>
<td>2%</td>
<td>3%</td>
<td>3%</td>
<td>10%</td>
<td>14.3%</td>
<td>7.3%</td>
</tr>
<tr>
<td></td>
<td>20 revolutions</td>
<td>7%</td>
<td>12%</td>
<td>15%</td>
<td>0%</td>
<td>1%</td>
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<tr>
<td></td>
<td>Eggels’ simulation</td>
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<tr>
<td></td>
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<td>1%</td>
<td>4%</td>
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</tr>
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<td>1.50</td>
<td>10 revolutions</td>
<td>2%</td>
<td>11%</td>
<td>19%</td>
<td>6%</td>
<td>9%</td>
<td>8%</td>
<td>0%</td>
<td>7.7%</td>
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<td>11%</td>
<td>0%</td>
<td>19%</td>
<td>9%</td>
<td>0%</td>
<td>6%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Eggels’ simulation</td>
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<td>6%</td>
<td>5%</td>
<td>9%</td>
<td>11%</td>
<td>8%</td>
<td>1%</td>
<td>6.2%</td>
<td></td>
</tr>
<tr>
<td>1.66</td>
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<td>19%</td>
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<td>21%</td>
<td>23%</td>
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<td>2%</td>
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<td>4%</td>
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<td>6%</td>
<td>10%</td>
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<tr>
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<td>6%</td>
<td>8%</td>
<td>10%</td>
<td>13%</td>
<td>0%</td>
<td>6.7%</td>
<td></td>
</tr>
</tbody>
</table>

**FIG. 21.** Comparison between simulations and experimental results for the mean radial velocity profiles in the impeller outstream (nu, nonuniform grid simulation; experiment, Wu and Patterson [2]; Eg, Eggels [3]).
FIG. 22. Comparison between simulations and experimental results for the mean tangential velocity profiles in the impeller outstream (nu, nonuniform grid simulation; experiment, Wu and Patterson [2]; DK, Derksen et al. [9, 10]).

where $D_v$ denotes the deviation, $U^*$ is the dimensionless velocity which is $U_r/U_{tip}$ for the radial velocity or $U_\theta/U_{tip}$ for the tangential velocity. The subscripts “sim”, “exp” and “max, exp” denote simulation result, experimental result, and maximum value of experimental results, respectively. Comparison of deviations between the statistics of 10 revolutions and 20 revolutions is shown in Tables V and VI, where $2r/D$ is the dimensionless radial position, $2z/w$ is the dimensionless axial position, “average” denotes the average of deviation, and the subscripts “20” and “10” denote the statistics for 20 revolutions and 10 revolutions, respectively. The results are given in Tables VII and VIII. The results suggest that there is a significant amount of statistical error in the results for 20 revolutions. This is especially true for positions near the tank wall. However, one may notice that the statistical results for 20 revolutions agree better with the experimental results. Therefore, the phase-resolved and phase-averaged results could be improved by increasing the statistics interval. As the resolution increases, the flow field tends to become more complex. Consequently, the statistics interval should be larger for a simulation with higher resolution. Eggels [3] suggested 40 revolutions for the statistics collection.
### TABLE VI
Deviation for Tangential Velocity

<table>
<thead>
<tr>
<th>(2r/D)</th>
<th>Simulations</th>
<th>(-1.6)</th>
<th>(-1.1)</th>
<th>(-0.55)</th>
<th>0.0</th>
<th>0.55</th>
<th>1.1</th>
<th>1.6</th>
<th>Average</th>
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<tbody>
<tr>
<td>1.07</td>
<td>10 revolutions</td>
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<td>6%</td>
<td>15%</td>
<td>15%</td>
<td>1%</td>
<td>3%</td>
<td>26%</td>
<td>9.4%</td>
</tr>
<tr>
<td></td>
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<td>15%</td>
<td>19%</td>
<td>1%</td>
<td>0%</td>
<td>24%</td>
<td>10.3%</td>
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<td></td>
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<td>7%</td>
<td>5%</td>
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</tr>
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### TABLE VII
Uncertainty for Radial Velocity

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<th>(2r/D)</th>
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### TABLE VIII
Uncertainty for Tangential Velocity

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9. CONCLUSION

In this paper, large eddy simulations for the complex flow field in a stirred tank driven by a Rushton turbine have been investigated. The results show that the lattice Boltzmann scheme with the nonuniform grid technique and the arbitrary computational domain method is efficient for turbulent flow simulations that demand large computational resources. The nonuniform grid technique can be implemented efficiently in three-dimensional simulations on parallel computer platforms. One-dimensional quadratic interpolation applied in the nonuniform grid technique does not reduce the efficiency of the technique. Therefore, the application of this technique makes it feasible for the simulation of a high-resolution stirred tank started from a quiescent state. The phase-averaged results are comparable in accuracy to the results obtained by Eggels [3] and Derksen and Van den Akker [4]. The phase-resolved flow fields are also well predicted by the simulation. The deviation of these results from the experimental data is probably attributable to the fact that the subgrid-scale model does not incorporate the effects of the tank wall and to the short time interval used to collect statistics. The arbitrary computational domain method introduced in this paper has been shown to be useful in complex geometries. With this method, one can efficiently implement bounce-back boundary conditions with complex geometries.

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REFERENCES