From creeping to inertial flow in porous media: a lattice Boltzmann–finite element study

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Abstract. The lattice Boltzmann method has been successfully applied for the simulation of flow through porous media in the creeping regime. Its technical properties, namely discretization, straightforward implementation and parallelization, are responsible for its popularity. However, flow through porous media is not restricted to near zero Reynolds numbers since inertial effects play a role in numerous natural and industrial processes. In this paper we investigate the capability of the lattice Boltzmann method to correctly describe flow in porous media at moderate Reynolds numbers. The selection of the lattice resolution, the collision kernel and the boundary conditions becomes increasingly important and the challenge is to keep artifacts due to compressibility effects at a minimum. The lattice Boltzmann results show an accurate quantitative agreement with finite element method results and evidence the capability of the method to reproduce Darcy’s law at low Reynolds numbers and Forchheimer’s law at high Reynolds numbers.

Keywords: discrete fluid models, flow in porous media, lattice Boltzmann methods
1. Introduction

Prediction of the transport properties of porous media, like the fluid permeability, defined by Darcy’s law, or heat conductivity, is of paramount importance in the chemical, mechanical, geological, environmental and petroleum industries. Flow situations in porous media are not restricted to the creeping flow regime, i.e., near zero Reynolds numbers where Darcy’s law applies [1]. Many important natural and industrial processes are characterized by large Reynolds numbers, where inertial effects also play a role. Examples include gas flow through a catalytic converter, groundwater flow, filtration processes and the flow of air in our lungs [2]. Hence, accurate simulation methods are needed to improve our understanding of these processes.

The lattice Boltzmann (LB) method has become an efficient tool [3]–[5] as an alternative to a direct numerical solution of the Navier–Stokes equation [6, 7] for simulating fluid flow in complex geometries such as porous media. Historically, the LB method was developed from the lattice gas automata [5, 8], replacing the number of particles in each lattice direction with the ensemble average of the single particle distribution function, and the discrete collision rule with a linear collision operator. In the LB method all computations involve local variables so that it can be parallelized easily [7]. Together with uniform grids and thus straightforward discretization, the LB method has become very popular in the field of porous medium flow simulations. With the advent of more powerful computers it has become possible to perform detailed simulations of finely resolved complex samples [3, 4, 7], [9]–[15].

In this work we investigate the accuracy of the LB method in flow regimes beyond Darcy’s regime. We focus on limitations of the method with respect to the lattice resolution and the selection of parameters. In particular, we address the requirement of reducing undesired compressibility effects by keeping the Mach number low, and how this influences the achievable maximum Reynolds numbers. In order to keep the compressibility as
To confirm the validity of the LB results, they are compared to finite element method (FEM) simulations that are performed with the commercial software package ANSYS. During recent decades, the FEM has been widely used to simulate fluid flow through porous media. It is known that the FEM can deal with complex pore geometries and boundary conditions, see for example [16]–[18]. Tezduyar et al [19] have developed the so-called deforming-spatial-domain/space–time (DSD/ST) procedure for flow problems with deforming interfaces using the so-called arbitrary Lagrangian–Eulerian (ALE) method and a space–time finite element method. This approach is based on fully resolved simulations around particles and is therefore computationally expensive in dense flows. For an overview of some finite element and finite difference techniques for incompressible fluid flow see [20] and for the efficiency of the solution algorithms see [21]. In recent studies, Yazdchi et al have related the macroscopic properties of porous media, namely permeability and inertial coefficients, to their microstructure and porosity [22, 23].

The remainder of this article is organized as follows. In section 2, an introduction to porous medium flow including laminar and weakly inertial flow, i.e. creeping flow, named Darcy’s flow, and Forchheimer’s flow is given. In section 3, the LB and FEM methods are introduced along with the description of the simulation setup. In section 4, we demonstrate quantitative agreement of LB and FEM simulations with theoretical predictions for fluid flow in porous media in the above mentioned regimes. Finally, we analyze and compare the results before we conclude.

2. Porous medium flow

Weak inertial flow, also named creeping flow (near zero Reynolds numbers) in porous media, can be described by Darcy’s law [24, 25]. It is defined by

$$\kappa = -\mu \frac{u_s}{\langle \nabla p \rangle_{x_2}}$$

(1)

where the porous medium parameter $\kappa$ (always positive) is called the permeability of the medium. $\langle \nabla p \rangle_{x_2}$ represents the average pressure gradient in the direction of the flow $x_2$, see figure 1, and $\mu$ represents the dynamic viscosity of the fluid. $u_s$ represents the superficial velocity defined by

$$u_s = \frac{1}{V_t} \int_{V_t} u_2 \, dV,$$

(2)

where $u_2$ is the fluid velocity in the flow direction. $V_t$ and $V_f$ represent the total volume and the fluid volume, respectively. The average velocity within the porous medium, $\langle u \rangle$, is related to the superficial velocity by $u_s = \varepsilon \langle u \rangle$, where $\varepsilon$ is the porosity of the medium. According to equation (1), Darcy’s law corresponds to a linear relation between the average pressure gradient $\langle \nabla p \rangle_{x_2}$ and the superficial velocity $u_s$, in the literature also referred to as the seepage velocity [26]. Darcy’s law was obtained empirically in 1846, but it can
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Figure 1. The computational domain with dimensions $X_1 \times X_2$ ($X_1 = 0.4X_2$). The sample is composed of 266 circles with radius $r$ representing 1.25% of $X_2$. They are randomly distributed assuring a minimum circle center distance $\Delta_{\text{min}}$ of 3.125% of $X_2$. Pure fluid chambers with lengths of 5% of $X_2$ are placed after and before the sample. The right inset shows a zoom of a typical unstructured, fine and triangular FEM mesh.

be derived from the continuous moment and mass balance assuming that the solid–fluid hydrodynamic interaction is proportional to the relative solid–fluid velocity [27].

When the Reynolds number is increased, inertia effects become relevant ($Re \approx O(1)$). The average pressure gradient $\langle (\nabla p)_{x_2} \rangle$ and the superficial velocity $u_s$ do not follow a linear relation anymore as was empirically shown by Forchheimer [28, 29]. For this flow regime, named Forchheimer’s regime, a quadratic term in $u_s$ is included to take the inertia effects into account. According to Massarani [27], Forchheimer’s law can be written as

$$\langle (\nabla p)_{x_2} \rangle = -\left(\frac{\mu}{\kappa} + \frac{\hat{\rho}_0 c}{\sqrt{\kappa} u_s} \right) u_s,$$  \hspace{1cm} (3)

where $c$ is a positive dimensionless parameter and $\hat{\rho}_0$ is the reference fluid density. The quadratic term in equation (3) describes a linear relationship between the solid–fluid hydrodynamic interaction and the relative solid–fluid velocity. Darcy’s law is recovered in the limit of $u_s \to 0$.

A transition from creeping flow to inertial flow has been reported in many studies, see for example the work of Koch and Ladd on flow in random arrays of cylinders [11] and spheres [12]. Also, the existence of a transition regime between the creeping and inertial regimes has been reported in the past [30]. This departure from Darcy’s regime is of the order of $u_s^3$ [22, 26], [31]–[33] and the relation between $\langle (\nabla p)_{x_2} \rangle$ and $u_s$ in this short interval is then given by

$$\langle (\nabla p)_{x_2} \rangle = -\left(\frac{\mu}{\kappa} + \frac{\gamma \hat{\rho}_0^2}{\mu} u_s^2 \right) u_s,$$  \hspace{1cm} (4)
where the cubic term in $u_s$ with positive parameter $\gamma$ represents the weak inertia correction.

3. Simulation methods

3.1. The lattice Boltzmann method

The integration of the Boltzmann equation on a regular lattice using a discrete set of velocities $c_i$ defines the lattice Boltzmann method. The lattice is defined by the spacing $\Delta x$, with the discrete velocity in units of $\Delta x/\Delta t$, where $\Delta t$ represents the timestep. Thus, the basic differential equation for the method is

$$f_i(x + \Delta t c_i, t + \Delta t) - f_i(x, t) = -\frac{\Delta t}{\tau} \left( f_i(x, t) - f_{eq}^i(x, t) \right),$$

(5)

where $f_i(x, t)$ represents the number of particles moving at position $x$ with velocity $c_i$ at discrete time $t$. The term on the right-hand side represents the collision operator as introduced by Bhatnagar, Gross and Krook (BGK) [34, 35], which approximates the collision through a linearization towards the equilibrium distribution function $f_{eq}^i(x, t)$ with a unique relaxation time $\tau$. This value is restricted to $\tau/\Delta t > 1/2$, assuring positive viscosity. If $\tau/\Delta t$ approaches $1/2$, numerical instabilities can arise [36]. This collision operator is often referred to as the ‘single relaxation time’ or BGK model. Under the assumption of very low Knudsen and Mach numbers ($Ma$), which assures small compressibility effects, $f_{eq}^i(x, t)$ is calculated by a second order Taylor expansion of the Maxwell distribution as [5]

$$f_{eq}^i(x, t) = \omega_i \frac{\rho}{\rho_0} \left( 1 + \frac{u \cdot c_i}{c_s^2} + \frac{(u \cdot c_i)^2}{2c_s^4} - \frac{u \cdot u}{2c_s^4} \right),$$

(6)

$\rho_0$ is a reference density. The coefficients $\omega_i$ are called lattice weights and are chosen to assure conservation of mass and momentum. They differ with lattice type, number of space dimensions and number of discrete velocities. For the cubic 3D lattice with 19 discrete velocities ($i = 0, \ldots, 18$) used in this work they are $1/3, 1/18$ and $1/36$ for the rest particles, the particles moving parallel to the $x_1$, $x_2$ or $x_3$, and the particles moving in diagonal directions, respectively [5, 37]. Even though the system of interest in this paper is intrinsically two-dimensional, we apply a three-dimensional implementation of the LB method and compute the flow in a very flat simulation domain with periodic boundary conditions in the $x_3$ direction. We do not expect this to have any influence on the simulation results, but it allows us to use our well tested implementation ‘LB3D’. The only disadvantage is higher computational costs, but we do not report on the amount of CPU time required for a given flow problem in this paper.

No-slip boundary conditions on the solid walls are implemented by mid-plane bounce back rules [38]. The BGK model is known to suffer from an artificial viscosity dependent slip at boundaries if these boundary conditions are used. An alternative approach for the collision operator, which reduces this well known drawback of the BGK model, is the multi-relaxation time (MRT) method [13, 14]. Here, the right-hand side of equation (5) is replaced by the expression

$$-\Delta t \left[ M^{-1} \cdot \mathbf{S} \cdot (\mathbf{m}(x, t) - \mathbf{m}_{eq}(x, t)) \right]_i,$$

(7)

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where $\mathbf{M}$ is a linear transformation chosen such that the moments
\[ m_i(x, t) = \sum_j M_{ij} f_j(x, t) \] (8)
represent hydrodynamic modes of the problem. We use the definitions given in [39], where $m_0$ is the fluid density, $m_2$ represents the kinetic energy, $m_i$ with $i = 3, 5, 7$ is the momentum flux and $m_i$ with $i = 9, 11, 13, 14, 15$ are components of the symmetric traceless viscous stress tensor. During the collision step the density and the momentum flux are conserved so that $m_i = m_i^{eq}$ with $i = 0, 3, 5, 7$. The non-conserved equilibrium moments $m_i^{eq}$, $i \neq 0, 3, 5, 7$, are assumed to be functions of these conserved moments and are explicitly given in [39]. $\mathbf{S}$ is a diagonal matrix, $\hat{S}_{ij} = \hat{s}_i \delta_{ij}$. The diagonal elements $\tau_i = 1/\hat{s}_i$ in the collision matrix are the relaxation times of the moments $m_i$. One has $\hat{s}_0 = \hat{s}_3 = \hat{s}_5 = \hat{s}_7 = 0$ because the corresponding moments are conserved. $\hat{s}_1 = 1/\tau_{\text{bulk}}$ describes the relaxation of the energy and $\hat{s}_9 = \hat{s}_{11} = \hat{s}_{13} = \hat{s}_{14} = \hat{s}_{15} = 1/\tau$ the relaxation of the stress tensor components. The remaining diagonal elements of $\mathbf{S}$ are chosen such that one has
\[ \hat{S} = \text{diag}(0, 1/\tau_{\text{bulk}}, 1.4, 0, 1.2, 0, 1.2, 0, 1.2, 1/\tau, 1.4, 1/\tau, 1.4, 1/\tau, 1/\tau, 1/\tau, 1.98, 1.98, 1.98) \] (9)
to optimize the algorithm performance [39, 40]. The two relaxation times $\tau$ and $\tau_{\text{bulk}}$ are restricted as well as for the BGK method to be $> \Delta t/2$, but allow the kinematic and bulk viscosity, respectively, to be defined. This multi-relaxation time scheme is commonly referred to as the ‘two relaxation time’ (TRT) method. An alternative TRT implementation can be found in [41, 42].

The macroscopic density $\rho(x, t)$ and velocity $u(x, t)$ are obtained from $f_i(x, t)$ as
\[ \rho(x, t) = \rho \sum_i f_i(x, t), \] (10)
\[ u(x, t) = \frac{\rho}{\rho(x, t)} \sum_i f_i(x, t) \mathbf{c}_i. \] (11)
The pressure is given by
\[ p(x, t) = c_s^2 \rho(x, t), \] (12)
where $c_s = 1/\sqrt{3}(\Delta x/\Delta t)$ is the speed of sound [5, 8]. The kinematic viscosity of the fluid, $\nu = \mu/\rho$, is a function of the discretization parameters $\Delta x$ and $\Delta t$ and the relaxation time $\tau$ [43, 44]. It is given by
\[ \nu = \frac{c_s^2 \Delta t}{2} \left( 2 - \frac{\tau}{\Delta t} \right). \] (13)

### 3.2. The finite element method

The velocity and pressure profiles through the system can be obtained from the solution of the conservation laws, namely, the continuity equation (conservation of mass) and the Navier–Stokes equations (conservation of momentum). In the absence of body forces, but assuming a constant density (i.e. incompressible flow) and steady state flow conditions, the
equations of conservation of mass and momentum for a Newtonian fluid are simplified to
\[
\nabla \cdot \mathbf{u} = 0, \quad \rho (\mathbf{u} \cdot \nabla \mathbf{u}) = -\nabla p + \mu \nabla^2 \mathbf{u}. \quad (14, 15)
\]

The FEM makes use of the variational formulations that allow the transformation of the above equations into a system of linear algebraic equations, which can be solved using a simple LU decomposition or iterative algorithms \[45, 46\]. Stable discretizations of the above equations are difficult to construct and it is known that the incompressibility constraint is not strongly enforced when using continuous polynomial shape functions for pressure. See \[21\] for a detailed theory and discussion. Langtangen et al. presented an overview of the most common numerical solution strategies, including fully implicit formulations, artificial compressibility methods, penalty formulations and operator splitting methods \[47\]. Using the conventional FEM scheme, we solve the above equations with the commercial software ANSYS \[48\]. The nonlinear solution procedure used in ANSYS belongs to a general class of semi-implicit methods for pressure linked equations (SIMPLE). In the flow domain, the steady state Navier–Stokes equations combined with the continuity equations are discretized into linear triangular elements. They are then solved using a segregated sequential solution algorithm. This means that element matrices are formed, assembled and the resulting system is solved using the Gaussian elimination algorithm for each degree of freedom separately. The number of iterations required to achieve a converged solution may vary considerably, depending on the number of elements, inertial contribution and the stability of the problem. Some more technical details are given in \[48\].

By knowing the fluid velocity field, the superficial velocity is then calculated from equation (2). Recently, using FEM simulations, Yazdchi and Luding \[22\] have shown that for both ordered and random fiber arrays, the weak inertia correction to the linear Darcy relation is third power in superficial velocity, up to small \(Re \approx 1 - 5\). When attempting to fit the data with a particularly simple relation, a non-integer power law performs astonishingly well up to moderate \(Re \approx 30\).

A typical unstructured, fine and triangular FEM mesh is shown in figure 1. The mesh size effect is examined by comparing the simulation results for different resolutions. The mesh refinement is carried out on the element level, meaning that a finer grid is overlaid on the coarse one. In order to be able to apply periodic boundary conditions in the \(x_1\) direction, see figure 1, we discretize the system such that we come up with the same number of nodes on the left and right boundaries. Periodic boundary conditions are applied by setting additional constraints, i.e. the same velocity, on these nodes.

3.3. Computational domain and simulation setup

The computational domain is presented in figure 1. The porous sample has a length of \(X_S\), representing 0.9 \(X_2\), and it is composed of 266 circles, randomly distributed with a radius \(r\) of 1.25\% of \(X_2\). A minimum distance \(\Delta_{\text{min}}\) of 3.125\% of \(X_2\) is imposed between the circle centers. The sample porosity then follows to be \(\varepsilon = 0.63730\).

For the LB method three different resolutions are used, namely low resolution (L), intermediate resolution (M) and high resolution (H), see table 1 for details. For the FEM method also three resolutions are used, corresponding to 22048, 49670 and
Table 1. Domain discretization for the LB simulations.

<table>
<thead>
<tr>
<th>Resolution</th>
<th>$X_1$</th>
<th>$X_2$</th>
<th>$X_S$</th>
<th>$X_C$</th>
<th>$r$</th>
<th>$\Delta_{\text{min}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low (L)</td>
<td>128</td>
<td>320</td>
<td>288</td>
<td>16</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td>Medium (M)</td>
<td>256</td>
<td>640</td>
<td>576</td>
<td>32</td>
<td>8</td>
<td>20</td>
</tr>
<tr>
<td>High (H)</td>
<td>512</td>
<td>1280</td>
<td>1152</td>
<td>64</td>
<td>16</td>
<td>40</td>
</tr>
</tbody>
</table>

98 2376 triangular elements, respectively. These meshes are referred to using the same abbreviations as for the LB simulations. However, one should keep in mind that the numbers of discretization elements used in the two methods cannot be compared easily since the LB simulations utilize a regular Cartesian lattice, while the FEM simulations are based on unstructured grids with locally varying resolution.

In the LB simulations a constant acceleration $g = ge_2$ drives the fluid using Guo’s method [49]. This acceleration defines the pressure gradient and acts on the fluid inside the sample, i.e. from $x_2 = X_C$ to $x_2 = X_C + X_S$. Together with this, on-site pressure boundary conditions are implemented on the inlet-plane $x_2 = 0$ and on the outlet-plane $x_2 = X_2$, setting a constant density value $\rho_0$ (reference density, i.e. $\sum f_i = 1$) on both planes. See equation (12) for the relation between the density and pressure within the LB method. These pressure boundary conditions are implemented using the method of Zou and He [50, 51]. Different values of the acceleration $g$ are used in order to study different Re regimes.

In our earlier work we proposed to use just on-site pressure boundary conditions at the inlet- and outlet-planes ($\sum f_i = 1 + \delta$ and $\sum f_i = 1 - \delta$, respectively), to impose a pressure drop of $2\delta$ [15]. The new setup presented here allows us to reduce compressibility effects inside the sample when the Reynolds number is increased. Periodic boundary conditions are applied in the $x_1$ direction.

In the FEM simulations, a pressure drop is imposed while the density is kept constant to drive the fluid. Further, we impose zero velocity on the surface of the fibers and also apply periodic boundary conditions in the $x_1$ direction.

4. Simulation results

We present a detailed analysis of the simulation of porous medium flow at low to moderate Reynolds numbers using the LB method. Special attention is given to its collision kernel, the discretization and the value of the relaxation time. To increase Re either the average flow velocity can be increased or the fluid viscosity can be decreased. However, the range of these parameters is limited: the lattice Boltzmann method is known to reproduce the Navier–Stokes equations in the low Mach number limit only, i.e., at high flow velocities compressibility artifacts can occur and render the results invalid. To validate the obtained data and to understand the impact of these limits on the precision of the LB results, we present a quantitative comparison with FEM results and theoretical predictions.

For the LB simulations 100 000 timesteps assure the steady state. The superficial velocity defined by equation (2) is calculated from the steady state LB data by

$$u_s = \frac{(\Delta x)^2}{X_1 X_2} \sum_{\mathbf{x} \in \mathcal{X}} u_2(\mathbf{x}), \quad (16)$$

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Figure 2. The normalized maximum fluid density as a measure for the effect of compressibility versus Reynolds number. Open symbols denote data obtained from LB-MRT simulations where the fluid is driven by an imposed pressure drop \( (p - BC) \) as suggested in [15]. When combining a fixed density at the inlet and outlet of the domain with a driving body force \( (p - BC + g) \), compressibility effects can be reduced substantially. Together with low values of the relaxation time \( \tau / \Delta t \), higher \( Re \) can be reached (solid symbols).

where \( X \) represents all fluid nodes in the simulation domain. The average pressure gradient is expected to be

\[
\langle (\nabla p)_{x_2} \rangle = -g \rho_0 .
\]  

(17)

The Mach and Reynolds numbers are defined by

\[
Ma = \frac{\langle u \rangle}{c_s},
\]  

(18)

\[
Re = \frac{\langle u \rangle r}{\nu}.
\]  

(19)

Figure 2 shows the relative maximum fluid density as a measure for the compressibility. The data are obtained from LB-MRT simulations of flow in the model geometry introduced above. While the open symbols represent data obtained using standard pressure drop boundary conditions (see [15]), the closed symbols describe data obtained with the newly proposed simulation setup combining pressure boundary conditions with a body force \( g \). In figure 2 the compressibility and thus the the relative maximum density in the system increases for higher \( Re \). Further, a clear reduction of compressibility effects is obtained when the newly proposed simulation setup is used.

Figure 2 also shows the effect of a reduced relaxation time (and thus viscosity) on the maximum reachable \( Re \) and the related compressibility effects. The compressibility of the fluid starts to become less important for lower viscosities and the combination of a low relaxation time \( \tau / \Delta t = 0.6 \) together with the combined pressure and body force boundary conditions leads to higher maximum Reynolds numbers. With this combination, \( Re \) of the order of 30 can be reached before, at large \( Re \), the fluctuations become too large for reliable analysis of the simulations.
Figure 3. Resolution and collision kernel \( (\tau/\Delta t = 0.6) \) analysis for permeability estimation versus \( Re \). The results show a dependence with the domain discretization, which is stronger when the BGK collision kernel is used.

Figure 3 shows the permeability estimate obtained from LB simulations using different discretizations and collision kernels. In our previous work it was shown for BGK simulations of 3D Poiseuille flow that when a relaxation time close to \( \tau/\Delta t = 0.8 \) is used, the dependence of the results on the discretization is minimal [14]. As is also demonstrated in the same article, finding such an optimal value for the relaxation time is impossible for more realistic samples such as a discretized Fontainebleau sandstone or an array of cylinders as used here: a strong dependence on the discretization appears when the BGK model is used, even though the relaxation time is set to \( \tau/\Delta t = 0.8 \). By using the MRT model this dependence can be decreased substantially. As can be observed in figure 3, the results of both collision kernels are qualitatively similar, i.e., both methods correctly reproduce the expected shape of the curve for comparable maximum \( Re \). One can see that in the case of the low resolution sample (L) the departure from Darcy’s regime (constant permeability) is to higher permeability values, which is unphysical since \( c \) in equation (3) is a non-negative parameter. In the case of intermediate (M) and high resolution (H), the departure is to lower permeability values, but only using the high resolution sample is it possible to simulate high-enough Reynolds numbers to analyze this phenomenon.

Figure 4 shows the permeability estimation for the LB-MRT method using different values for the relaxation time. Due the resolution used for this study (H) together with the MRT collision kernel it is not surprising that the results do not differ much quantitatively, but only by about 3%. The main difference is in the highest possible Reynolds numbers that can be reached by using a small value of the relaxation time \( \tau/\Delta t = 0.6 \), in accordance with figure 2. As mentioned above, numerical instability arises when the relaxation time approaches \( \tau/\Delta t = 0.5 \). For this reason the smallest value used in this work is \( \tau/\Delta t = 0.6 \).

For porous medium flow calculations using LB-MRT and \( \tau/\Delta t = 0.6 \), figure 2 shows that at high Reynolds numbers the compressibility is of the order of 30%. Furthermore, as we can see in figure 5 the superficial velocity remains low enough to keep Mach numbers below \( \approx 10^{-1} \). However, inside the sample there are zones with high velocity. Indeed, for the two higher Reynolds numbers simulated, on \( \approx 25\% \) and \( \approx 45\% \) of the fluid nodes the
Figure 4. Relaxation time analysis for permeability estimation versus $Re$ calculated using the LB-MRT method and the high resolution (H) computational domain. Different values for the relaxation time $\tau/\Delta t$ were used. The results only differ by $\approx 3\%$ and the highest $Re$ can be reached with the smallest value of $\tau/\Delta t = 0.6$.

Figure 5. The superficial velocity $u_s$ and maximal velocity $u_{y,max}$ inside the sample for LB-MRT and $\tau/\Delta t = 0.6$. The inset shows the percentage of fluid nodes with velocity $u_y$ higher than 20% of the speed of sound.

velocity is higher than 20% of the speed of sound, $c_s$, see figure 5. Such high velocities and strong compressibility which are present at high Reynolds numbers greatly question the validity of the results.

To investigate the validity of the results FEM simulations are performed as a benchmark for the LB simulations. In figure 6 the data obtained using both methods are plotted. One can see that the FEM results also show a dependence on the sample resolution. As stated above, in the case of the FEM the L, M and H consist of domains with 22,048, 49,670 and 982,376 elements. The resolution dependence is also demonstrated in the inset of the figure, where the permeability is shown as a function of resolution for a Reynolds number of $Re \approx 10^{-3}$. For simulations using $1 \times 10^6$ elements, there

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is a difference of $\approx 2\%$ between the LB and FEM results. This can be explained by several factors. Neither the FEM nor the LB results are fully converged with respect to the required number of elements, but performing a large number of simulations with substantially higher resolution is not feasible with the computational resources available, in particular since the final values are expected to change by not more than a few percent. Since the LB results are systematically lower than the FEM results, a possible explanation could be the loss of accuracy of LB due to the relatively small choice for the relaxation time $(\tau/\Delta t = 0.6)$, as demonstrated in figure 4. In any case, the deviation is small, and thus it can be concluded that the data in figure 6 show a qualitative and quantitative agreement when the Reynolds number increases. Figure 6 shows that for both methods the value of the permeability, beyond the validity of Darcy’s law, equation (1), drops consistently. Darcy’s law is limited to the weak inertia regime, i.e. low Reynolds numbers. To analyze the validity of the simulated data for high Reynolds numbers, we plot $u_s$ versus $\langle (\nabla p)_{xx} \rangle$ in figure 7 using the H data and $\tau/\Delta t = 0.6$ for LB-MRT. For both methods, at small velocity values (small $Re$), the flow follows Darcy’s prediction (constant slope) and equation (1) accurately fits the data. When the velocity increases the measured permeability departs from Darcy’s law and the best possible fit is obtained using equation (4), which confirms the existence of a transition regime with cubic velocity dependence. Only for higher velocities does the flow enter Forchheimer’s regime and can the data be fitted accurately by equation (3). The departure from the different flow regimes is clearly seen in the bottom panel of figure 7, where the relative error of the fits is plotted versus $Re$. It is calculated by

$$\epsilon(w) = \frac{w_{\text{fit}} - w_{\text{sim}}}{w_{\text{sim}}},$$

where the indices $\text{sim}$ and $\text{fit}$ represent the results obtained by simulation and from the fit, respectively. It can clearly be seen from both datasets that Darcy’s regime holds
Figure 7. Top: fit of the expressions that relate the superficial velocity $u_s$ and the average pressure gradient $\langle (\nabla p)_{x_2} \rangle$ for the porous medium flow. Bottom: the relative error of the fit presented above. The relative error shows that Darcy’s law fits the simulation results for $Re \ll 10^0$. Above this value the fit of the expressions given by equation (4) is better than the fit with Darcy’s law.

until $Re \approx 10^0$ and the inertia transition correction expression holds until $Re \approx 10^1$. Forchheimer’s regime describes the flow for higher $Re$.

5. Conclusions

We have presented an analysis of the calculation of the flow in porous media in different flow regimes using the LB method and compared our results to FEM simulations. Special attention has been given to discretization, selection of a collision kernel and choice of parameters. For the LB method a simulation setup combining a constant pressure at the inlet and outlet and an external acceleration acting inside the sample has been proposed in order to reduce compressibility artifacts at high $Re$. However, at higher $Re$ a substantial number of lattice nodes show flow velocities beyond 20% of the speed of sound and compressibility effects can clearly be observed. In order to clarify the validity of these results we have compared our data to FEM simulations and demonstrated a good
quantitative agreement for the full range of $Re$ studied. Furthermore, the data show good agreement with theoretical predictions, demonstrating that the range of $Re$ studied in this work is well accessible for the LB method and that compressibility effects only have a minor influence. The results accurately predict three different flow regimes. These are, firstly, Darcy’s regime for $Re \lesssim 10^9$, where the average pressure gradient $\langle \nabla p \rangle_{x_2}$ and the surface velocity $u_s$ obey a linear relationship. Secondly, a transient regime $10^9 \lesssim Re \lesssim 10^{10}$, where the flow is modeled by Darcy’s law plus an inertia correction term represented by a cubic term on $u_s$, see equation (4). Finally, for higher Reynolds numbers ($Re \gtrsim 10^{11}$) the porous medium flow follows Forchheimer’s prediction up to the limit of both simulation methods of $Re \approx 0$.

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