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Non-parametric wall model and methods of identifying boundary conditions for moments in gas flow equations

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Abstract

In this paper, we use Molecular Dynamics (MD) simulation method to study gas-wall boundary conditions. Discrete scattering information of gas molecules at the wall surface are obtained from collision simulations. The collision data can be used to identify the accommodation coefficients for parametric wall models such as Maxwell, Cercignani-Lampis scattering kernels. Since these scattering kernels are based on a limited number of accommodation coefficients, we adopt non-parametric statistical methods to construct the kernel to overcome these issues. Different from parametric kernels, the non-parametric kernels require no parameter (i.e. accommodation coefficients) and no predefined distribution. We also propose approaches to derive directly the Navier friction and Kapitza thermal resistance coefficients as well as other interface coefficients associated to moment equations from the non-parametric kernels. The methods are applied successfully to systems composed of CH\textsubscript{4} or CO\textsubscript{2} and graphite, which are of interest to the petroleum industry.

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INTRODUCTION

The study of transport properties in porous media plays an important role in many applications such as soil mechanics, geohydrology and the storage of nuclear waste. Along with the development of unconventional reservoirs (shale gas) extraction technology, like hydraulic fracturing, more attention has been paid to the transport of gas molecules in carbon pores. As a result, modeling the gas behavior and its interaction with the boundary is of significant interest.

Due to the size of the pore, gaseous molecules, here methane (CH$_4$) and carbon dioxide (CO$_2$) in this study, can travel with few collisions, resulting in high Knudsen number (Kn), a similar situation as the rarefaction effect. It is known that when Kn $> 0.01$, predictions based on the continuum Navier-Stokes-Fourier (NSF) equations and classical no-slip, no-jump conditions are no longer in agreement with experiences and atomistic simulation results [1, 2]. In order to capture these phenomena, more advanced continuum equations and boundary conditions are necessary [3, 4].

Unlike liquids where the friction and thermal resistance are characterized by layers of interacting molecules adsorbed at the wall [5–7], the gas molecules collide infrequently and their residence time near the wall can be neglected. The exchange of momentum and energy between the gas and the wall can be understood from ensemble of independent gas-wall collisions. In most cases, the collisions are usually modeled with scattering kernels based on several accommodation coefficients [8–14]. Other class of wall models for rough surfaces [15–17] were analytically derived from corrugation parameter and potential well depth. Although these parametric models are simple to code, they rely on many oversimplification hypotheses which cannot guarantee the accuracy of the collisions for the whole velocity range. These problems can have consequences on the boundary conditions at the continuum level and simulation results based on these scattering kernels.

The paper presents a systematic study of gas-wall collision models based on Molecular Dynamics (MD) simulations. The systems in consideration are composed of methane CH$_4$ (considered as monatomic gas) or carbon dioxide CO$_2$ (rigid linear molecule) interacting
with a graphite wall constituted of carbon atoms. By beaming independently gas molecules onto the surface and recording the reflected flux, we can determine the accommodation coefficients. More importantly, we can reconstruct numerically a non-parametric (NP) wall model. Different from parametric models in literature, this scattering kernel is able to capture the reflection process in a more realistic way. Originated from non-parametric statistics, the NP scattering kernel requires no parametrization (i.e. accommodation coefficients) and no predefined analytical form. Interestingly, it can be used to directly determine the parameters of any phenomenological boundary conditions, including those for NSF or moment equations. While these equations are valid for a limited range of Kn in the transition regime, the NP wall model can be directly implemented in particle methods like DSMC (Direct Simulation Monte Carlo) or MD to simulate flows at any Kn number. The development of gas-wall boundary conditions for continuum equations from a non-parametric kernel is the major contribution of the present work, which will be detailed in the subsequent sections.

II. STUDY OF GAS-WALL MODELS WITH MOLECULAR DYNAMICS METHOD

A. Scattering kernels

In kinetic theory, the state of monatomic gas at any location \( x \) at time \( t \) is entirely determined from the local number density \( n(x, t) \) and the probability density function \( f(x, c, t) \) of velocity \( c \). The evolution of the latter is governed by the Boltzmann equation and the boundary conditions

\[
c_z(nf)^+(c) = \int_{\Omega^-} B(c|c')|c'_z|(nf)^-(c') \, dc', \quad c' \in \Omega^-, \quad c \in \Omega^+. \tag{1}
\]

In the above expression, we assume that the boundary is normal to the \( z \) direction, and time \( t \) and space \( x \) variables are dropped for simplicity. Eq. (1) connects the incoming flux \( c'_z(nf)^-(c') \) and the outgoing flux \( c_z(nf)^+(c) \) via the scattering kernel \( B(c|c') \). The two velocities \( c \) and \( c' \) belong to dual half-spaces \( \Omega^- \) and \( \Omega^+ \) in \( \mathbb{R}^3 \), respectively, defined below

\[
c' \in \Omega^- = \mathbb{R}^2 \times \mathbb{R}^-, \quad c \in \Omega^+ = \mathbb{R}^2 \times \mathbb{R}^+. \tag{2}
\]
For fluid in equilibrium, the distribution of velocity is equal to the Maxwell-Boltzmann distribution

\[ f_{eq}(c) = f_M(c) = \frac{1}{\sqrt{2\pi \theta}} \exp \left[ -\frac{c^2}{2\theta} \right], \quad \theta = k_B T/m, \tag{3} \]

where \( k_B \) is the Boltzmann constant, \( m \) the atomic mass, and \( T \) the temperature. For rigid gas molecules, in addition to translational velocity of the center of mass, we must account for the rotational velocity \( \omega \). The scattering kernel must be replaced by \( B(\omega, c|\omega', c') \) and the probability density by \( f(\omega, c) \). The two half-spaces \( \Omega^- \) and \( \Omega^+ \) are also extended to include the rotational velocity \( \omega \), e.g \( \Omega^- = \mathbb{R}^5 \times \mathbb{R}^- \) for incident molecules and \( \Omega^+ = \mathbb{R}^5 \times \mathbb{R}^+ \) for reflected molecules. It is possible to include the orientation distribution in the scattering kernel but this will not be considered in the present work. At equilibrium, this density function is given by

\[ f_{eq}(\omega, c) = f_M(c)f_{\omega M}^\omega(\omega), \quad f_{\omega M}^\omega(\omega) = \frac{1}{\sqrt{2\pi \theta^d}} \exp \left[ -\frac{\omega^2}{2\theta^d} \right], \quad \theta^\omega = k_B T/I. \tag{4} \]

The quantity \( I \) represents the moment of inertia and the power \( d \) the rotational degree of freedom, \( d = 2 \) for linear molecules and \( d = 3 \) otherwise. It is noted that for linear molecules, the rotation around its proper axis is not considered.

The scattering kernel \( B(\omega, c|\omega', c') \) which is the probability of finding molecules bouncing with velocity \( (\omega, c) \) with given colliding velocities \( (\omega', c') \) can be determined by Molecular Dynamics collision simulation. Gas molecules are beamed at given velocities \( (\omega', c') \) onto the surface in consideration and the velocity distribution of reflecting molecules associated to \( (\omega', c') \) is recorded. Next the arriving velocities \( (\omega', c') \) are also varied to cover the incident velocity space. Generally, if the number of realizations is sufficiently large, we have a large set of discrete points which can represent the true probability density \( B(\omega, c|\omega', c') \).

We are also concerned about the use of the kernel as wall boundary conditions in other simulation methods (for example Molecular Dynamics, Direct Simulation Monte Carlo or Lattice Boltzmann). If we use the discrete form of \( B(\omega, c|\omega', c') \), output results must be obtained from the interpolation of known points. This method is accurate but less computationally convenient. The scattering kernel can be analytically modeled using some physical
parameters for example Tangential Momentum Accommodation Coefficients (TMAC) or Energy Momentum Accommodation Coefficients (EAC), etc... Some notable scattering models are Maxwell-Yamamoto (MY) [11], Cercignani-Lampis (CL) [9] etc... which can be used for atomistic gas flow simulations and accommodation coefficients can be used to derive velocity slip and temperature jump coefficients for NSF equations. To account for the special reflection mechanism of the anisotropic surface, one can use Dadzie-Meolans (DM) kernel [18] or anisotropic Cercignani-Lampis (ACL) kernel [14] with three different coefficients associated to the three directions $x, y, z$.

The MD collision point cloud can be fitted by analytical scattering models and the model parameters can be identified. However, the data can be scattered and there is no truly efficient fitting algorithm, for example, one can use the mean square of the difference between the two probability densities or methods based on accommodation parameters. We note that constant accommodation coefficients are only meaningful for analytical scattering kernel listed previously. For realistic gas surface interaction, those coefficients are usable in approximative sense and can oversimplify the true behavior.

B. Expressions for fluxes, average values and accommodation coefficients

Given molecular quantities $Q$ as function of velocities $\mathbf{c}, \omega$, the average value $\overline{Q}$ and the flux $\Phi_{Q}$ across a plane normal to $z$ can be computed as

$$\overline{Q} = \int Q(\mathbf{c}, \omega) f d\mathbf{c} d\omega, \quad \Phi_{Q} = n \int Q(\mathbf{c}, \omega) c_{z} f d\mathbf{c} d\omega. \quad (5)$$

Given the fact that all the physical quantities such as density $n$, temperature $T$, stress $\sigma$, velocity $v$, and heat flux $q$ are either average value or flux of molecular quantities, it is possible to investigate their relations at the boundary by examining the gas wall collisions. With respect to the wall normal to the $z$ direction, we define influx $\Phi_{Q}^{-}$ and outflux $\Phi_{Q}^{+}$ of atomic quantity $Q(\mathbf{c}, \omega)$ at the wall via the expressions

$$\Phi_{Q}^{-} = \int_{\Omega^{-}} |c_{z}| (n f)^{-} Q(\mathbf{c}, \omega) d\mathbf{c} d\omega, \quad \Phi_{Q}^{+} = \int_{\Omega^{+}} |c_{z}| (n f)^{+} Q(\mathbf{c}, \omega) d\mathbf{c} d\omega. \quad (6)$$
From an atomistic viewpoint, $\Phi_\pm Q$, $\Phi_Q^+$ can be computed by counting the number of atoms $N$ crossing the control plane in a given time $t$

$$\Phi_Q^\pm = \frac{1}{t} \sum_{\text{incident}} Q(c, \omega) = \frac{N}{t} \langle Q \rangle_i = \nu \langle Q \rangle_i, \quad \Phi_Q^\pm = \nu \langle Q \rangle_o,$$  \tag{7}$$

where the subscript $i$ stands for input (incident), $o$ for output (reflection), $\nu$ collision rate. Here the notation $\langle Q \rangle$ is the average of molecular quantities $Q$ that cross the control plane in Molecular Dynamics simulations. By breaking each relation in (5) into two integrals in half-spaces $\Omega^-$ and $\Omega^+$ as follows

$$nQ = \int_{\Omega^-} Q(c, \omega) |c_z| |n(f)| dcd\omega + \int_{\Omega^+} Q(c, \omega) |c_z| |n(f)| dcd\omega,$$

$$\Phi_Q = -\int_{\Omega^-} Q(c, \omega) |c_z| |n(f)| dcd\omega + \int_{\Omega^+} Q(c, \omega) |c_z| |n(f)| dcd\omega,$$  \tag{8}$$

and making use of (7) and (6), the flux $\Phi_Q$ and the average value $\overline{Q}$ at the wall can also be expressed as

$$n\overline{Q} = \Phi_Q^+ |c_z| + \Phi_Q^- |c_z| = \nu \langle Q/|c_z| \rangle_{i+o}, \quad \Phi_Q = \Phi_Q^+ - \Phi_Q^- = \nu \langle Q \rangle_o - \langle Q \rangle_i,$$  \tag{9}$$

with notation $\langle Q \rangle_{a\pm b} := \langle Q \rangle_a \pm \langle Q \rangle_b$. The relation between the average value $\overline{Q}$, and fluxes $\Phi_Q^+ |c_z|$ and $\Phi_Q^- |c_z|$ is useful because it is more convenient to compute $\overline{Q}$ with MD simulations. Choosing $Q = 1$ in (9) and noting that $\overline{Q} = \langle Q \rangle_i = \langle Q \rangle_o = 1$, we have the equalities

$$\nu = n \frac{1}{\langle 1/|c_z| \rangle_{i+o}}, \quad \Phi_1 = 0.$$  \tag{10}$$

Substituting $\nu/n$ from the first expression in (10) back into (9) for the general $Q$, we can derive that

$$\overline{Q} = \langle Q/|c_z| \rangle_{i+o}/\langle 1/|c_z| \rangle_{i+o}, \quad \Phi_Q = n \langle Q \rangle_{o-i}/\langle 1/|c_z| \rangle_{i+o}.$$  \tag{11}$$

We remark that the second relation of (10) is equivalent to the no atom accumulation condition at the wall, i.e the influx is equal to the outflux. If the leaving atoms are fully thermalized by the wall, the phase density $f^+$ should be replaced by the equilibrium distribution $f_{eq}(\omega, c)$ at the wall temperature $T_w$. The outgoing flux $\Phi_Q^+ w$ associated to this
distribution is given by the expression

$$\Phi_{Q_w}^+ = \int_{\Omega^+} |c_z| n f_{eq}(\omega, c) Q dcd\omega = \nu \langle Q \rangle_w,$$

(12)

where the subscript $w$ is for outgoing flux at the wall temperature $T_w$. Since $f_{eq}(\omega, c)$ is known from (4) and $\nu$ is estimated by setting $Q = 1$, we can compute the expected values for thermal wall $\langle Q \rangle_w$. They are functions of the reduced wall temperature $\theta_w = k_B T_w / m$ and given in Tab I. It is noted that for the special case where $d = 0$, the tabulated values are consistent with previous works for monatomic gas [19].

<table>
<thead>
<tr>
<th>Component</th>
<th>Velocity</th>
<th>Energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tangential $(x, y)$</td>
<td>$\langle c_x \rangle_w = 0$</td>
<td>$\langle c_x^2 \rangle_w = \theta_w$</td>
</tr>
<tr>
<td>Normal $(z)$</td>
<td>$\langle c_z \rangle_w = 1/2 \sqrt{2 \pi \theta_w}$</td>
<td>$\langle c_z^2 \rangle_w = 2 \theta_w$</td>
</tr>
<tr>
<td>Total</td>
<td>$\langle c \rangle_w = 3/2 \sqrt{2 \pi \theta_w}$</td>
<td>$\langle c^2 + \frac{I}{m} \omega^2 \rangle_w = (4 + d) \theta_w$</td>
</tr>
</tbody>
</table>

TABLE I. Expected values as functions of the reduced wall temperature $\theta_w = k_B T_w / m$. For CH$_4$, the rotation energy is neglected $d = 0$ and for CO$_2$, $d = 2$.

As a result, the accommodation coefficient of quantity $Q$ is equivalent to the expression

$$\alpha_Q = \frac{\Phi_Q - \Phi_{Q_w}^+}{\Phi_Q - \Phi_{Q_w}^-} = \frac{\langle Q \rangle_{i-o}}{\langle Q \rangle_{i-w}} \quad \text{or} \quad \langle Q \rangle_o = (1 - \alpha_Q) \langle Q \rangle_i + \alpha_Q \langle Q \rangle_w.$$  

(13)

The above expression which is independent of the collision rate $\nu$, is useful for the determination of the accommodation coefficients using MD method. Usually, the value for $\langle Q \rangle_w$ is known explicitly (see Tab I) and the coefficient can be computed based on Eq. (13). Most analytical wall models in literature are based on constant accommodation coefficients, which are independent of the input data $nf^-$. These assumptions may not be true for a general kernel $B(\omega, c|\omega', c')$ and this is the major disadvantage of using accommodation coefficients to model realistic surfaces.

When accommodation coefficients are not properly defined, different methods can be used to compute those coefficients and result differently. For example, in Ref.[19], the authors...
proposed using the least-squares formula

\[
\alpha_Q = 1 - \frac{\sum_{\text{collision}} (Q_i - \langle Q_i \rangle)(Q_o - \langle Q_o \rangle)}{\sum_{\text{collision}} (Q_i - \langle Q_i \rangle)^2},
\]

from the collision clouds. They found that results are very different from those obtained by Eq. (13).

These observations pose some problems on theories based on the existence of the constant accommodation coefficients for general surfaces. However, interface phenomena like slip velocity and temperature jumps do exist. Modeling those effects and identifying the parameters without using accommodation coefficients will be considered in the following.

C. Boundary conditions for Navier Stokes Fourier (NSF) equations

In this subsection, we present a new method to directly determine the macroscopic velocity and temperature jump coefficients via collision simulations. This completely avoids the intermediate modeling and simulations based on scattering kernels. As we know, all available analytical models have limitations. First, they only allow at most three accommodation coefficients. If we choose to model momentum accommodation effect along one direction, we have to sacrifice the energy accommodation along this direction. Accommodation effects for high order moment are also unavailable. Secondly, using constant accommodation coefficients, like most analytical wall models in literature can be a strong assumption. Numerical evidence in the latter section shows that in some cases, the true behavior deviates significantly from that hypothesis.

The approach proposed here is independent of scattering model and can be applied to any surfaces. It can also be extended to deal with general boundary conditions involving higher order moments. In slip regimes, the usual macroscopic boundary conditions for velocities \(v_x, v_y\) and reduced temperature jump \(\theta - \theta_w\) are given in the following forms

\[
v_k = -\frac{2 - \alpha_k}{\alpha_k} \frac{\sigma_{\text{hz}}}{nm \sqrt{2\theta/\pi}}, \quad \theta - \theta_w = -\frac{2 - \alpha_e}{2\alpha_e} \frac{q_z}{nm \sqrt{2\theta/\pi}}, \quad k = x, y,
\]

(15)
where \( \sigma_{xz}, \sigma_{yz} \) are the (minus) shear stress components and \( q_z \) the normal heat flux at the wall. Constants \( \alpha_k \) are the tangential accommodation coefficients associated to the tangential translational molecular velocities, and \( \alpha_e \) is the energy accommodation coefficient associated to its kinetic energy. The above equation where the thermal transpiration is neglected can be derived from the scattering models. In this paper, we propose a more general phenomenological form for the boundary conditions

\[
\beta_{1k} v_k = -\frac{\sigma_{kz}}{nm \sqrt{2\theta/\pi}}, \quad \beta_2 (\theta - \theta_w) = -\frac{q_z}{2nm \sqrt{2\theta/\pi}}, \quad k = x, y,
\]  \hspace{1cm} (16)

where \( \beta_{1x}, \beta_{1y} \) and \( \beta_2 \) are the dimensionless friction and Kapitza coefficients, depending on the gas-wall couple. It is clear that we recover the original equation if the coefficients \( \beta_{1k} \) and \( \beta_2 \) are connected to the accommodation coefficients \( \alpha_k \) and \( \alpha_e \) via the relation

\[
\beta_{1k} = \frac{\alpha_k}{2 - \alpha_k}, \quad \beta_2 = \frac{\alpha_e}{2 - \alpha_e}, \quad k = x, y,
\]  \hspace{1cm} (17)

It is noted that the two expressions in (16) can also be used for the cases where the accommodation coefficients are not constant.

Using Eqs. (7-11), we shall derive schemes to determine \( \beta_{1k} \) and \( \beta_2 \) from MD simulations. The velocity defined as \( v_k = \overline{c_k} \) (i.e \( Q = c_k \)) can be computed by the expression

\[
v_k = \frac{\langle c_k / |c_z| \rangle_{i+o}}{\langle 1 / |c_z| \rangle_{i+o}} \tag{18}
\]

The (minus) shear stress \( \sigma_{kz} = \Phi mC_k \) (i.e \( Q = mC_k \)) can also be computed in the following way

\[
-\sigma_{kz} = m\nu (\langle C_k \rangle_i - \langle C_k \rangle_o) = mn \frac{\langle c_k \rangle_i - \langle c_k \rangle_o}{\langle 1 / |c_z| \rangle_i + \langle 1 / |c_z| \rangle_o} = mn \frac{\langle c_k \rangle_{i-o}}{\langle 1 / |c_z| \rangle_{i+o}}, \tag{19}
\]

where \( C_k = c_k - v_k \) is the peculiar velocity. Comparing Eqs. (18,19) with (16), we can derive the interface coefficients \( \beta_{ik} \) via the expression

\[
\beta_{1k} = \frac{\langle c_k \rangle_{i-o} / \sqrt{2\theta/\pi}}{\langle c_k / |c_z| \rangle_{i+o}}. \tag{20}
\]
Next, the reduced temperature $\theta = \frac{C^2 + (I/m)\omega^2}{(3 + d)}$ and the heat flux $q_z = \Phi_{(mC^2 + I\omega^2)/2}$ are given by the equation

$$\theta = \frac{\langle(C^2 + (I/m)\omega^2)/|c_z|\rangle_{i+o}}{(3 + d)\langle1/|c_z|\rangle_{i+o}}, \quad -q_z = mn\frac{\langle(C^2 + (I/m)\omega^2)/2\rangle_{i-o}}{\langle1/|c_z|\rangle_{i+o}},$$

(21)

where $d$ is the number of rotation degrees of freedom of gas molecule, $d = 2$ for CO$_2$. Comparing Eqs. (21) with (16), we can calculate the Kapitza coefficient

$$\beta_2 = \frac{(3 + d)\langle(C^2 + (I/m)\omega^2)/2\rangle_{i-o}/(4\theta/\pi)}{\langle(C^2/|c_z|)\rangle_{i+o} - (3 + d)\theta_w\langle1/|c_z|\rangle_{i+o}/\sqrt{2\theta/\pi}}.$$  

(22)

We note that for monatomic gas, it is sufficient to remove the terms $I\omega^2$ and $d$ in the above expression and obtain

$$\beta_2 = \frac{3\langle(C^2/2\rangle_{i-o}/(4\theta/\pi)}{\langle(C^2/|c_z|)\rangle_{i+o} - 3\theta_w\langle1/|c_z|\rangle_{i+o}/\sqrt{2\theta/\pi}}.$$  

(23)

To facilitate the comparison between the numerical results, we normalize stress and heat flux computed by MD method with suitable quantities and rewrite the phenomenological law in the following way

$$\beta_{1k} = -\frac{\hat{\sigma}_{kz}}{\hat{v}_k}, \quad \beta_2 = \frac{-\hat{q}_z}{\Delta\hat{\theta}}, \quad k = x, y.$$  

(24)

Here, the hat notation is used for the normalized quantities,

$$\hat{\sigma}_{kz} = \frac{\sigma_{kz}}{nm\zeta^2}, \quad \hat{v}_k = \frac{v_k}{\zeta}, \quad \hat{q}_z = \frac{q_z}{2nm\zeta^3}, \quad \Delta\hat{\theta} = \frac{\theta - \theta_w}{\zeta^2}, \quad \zeta = \sqrt{2\theta/\pi}.  

(25)

D. Extension to 13 moments equations

Without being limited to NSF equations, the method presented previously can be applied to higher order model. Given any macroscopic boundary conditions in terms of moments, the present method can be used to derive the coefficients associated to boundary conditions. As an example, we consider the boundary conditions of R13 equations written for isotropic surfaces in dimensionless form[3, 20]

$$\beta_1 = -\frac{\hat{\sigma}_{xz}}{\hat{k}_1}, \quad \beta_2 = -\frac{\hat{q}_z}{\hat{k}_2}, \quad \beta_3 = -\frac{\hat{m}_{xxz}}{\hat{k}_3}, \quad \beta_4 = \frac{\hat{m}_{zzz}}{\hat{k}_4}, \quad \beta_5 = \frac{\hat{R}_{xz}}{\hat{k}_5},$$

(26)
\[ \hat{m}_{ijl} = \frac{m_{ijl}}{nm\zeta^3}, \quad \hat{R}_{ij} = \frac{R_{ij}}{nm\zeta^2}, \]
\[ \hat{\kappa}_1 = \sqrt{\frac{2}{\pi\theta}} \left( P v_x + \frac{1}{2} m_{xxz} + \frac{1}{5} q_x \right) / (nm\zeta^2), \]
\[ \hat{\kappa}_2 = \sqrt{\frac{2}{\pi\theta}} \left( 2P(\theta - \theta_w) - \frac{1}{2} P v^2 + \frac{1}{2} \theta q_{zz} + \frac{R}{15} + \frac{5}{28} R_{zz} \right) / (2nm\zeta^3), \]
\[ \hat{\kappa}_3 = \sqrt{\frac{2}{\pi\theta}} \left( \frac{1}{14} R_{xx} + \theta q_{xx} - \frac{1}{5} \theta q_{zz} + \frac{1}{3} P(\theta - \theta_w) - \frac{4}{5} P v^2 - \frac{R}{150} \right) / (nm\zeta^3), \]
\[ \hat{\kappa}_4 = \sqrt{\frac{2}{\pi\theta}} \left( \frac{2}{5} P(\theta - \theta_w) - \frac{1}{14} R_{zz} - \frac{3}{5} P v^2 - \frac{7}{5} \theta q_{zz} + \frac{R}{70} \right) / (nm\zeta^3), \]
\[ \hat{\kappa}_5 = \sqrt{\frac{2}{\pi\theta}} \left( P \theta v_x - \frac{11}{5} \theta q_x - \frac{1}{2} \theta m_{xxz} - P v^2 v_x + 6P v_x(\theta - \theta_w) \right) / (nm\zeta^4). \] (27)

The quantities \( P, R, R_{ij} \) and \( m_{ijl} \) are defined from the moments
\[ P = nm\theta + \frac{1}{2} \sigma_{zz} - \frac{1}{120} \frac{R}{\theta} - \frac{1}{28} \frac{R_{zz}}{\theta}, \quad R_{ij} = mnC^2(C_iC_j - C^2/3\delta_{ij}) - 7\theta q_{ij}, \]
\[ R = mn(C^4 - 15\theta^2), \quad m_{ijl} = mnC_iC_jC_l - C^2(C_i\delta_{jl} + C_j\delta_{il} + C_l\delta_{ij})/5, \]
\[ k = x, y, \quad i, j, l = x, y, z. \] (28)

These original boundary conditions are derived for Maxwell molecules and Maxwell scattering kernel where all the coefficients are identical \( \beta_1 = \beta_2 = \ldots = \beta_5 = \alpha/(2 - \alpha) \) with \( \alpha \) being the accommodation coefficient. Moments \( R, R_{ij} \) and \( m_{ijl} \) are connected to stress, heat flux, velocity, temperature and their derivatives via a regularization procedure [3, 21, 22].

Although the derivation conditions are rather restrictive, we shall assume that they are valid and determine the coefficients \( \beta_1, \beta_2, \ldots, \beta_5 \). We shall base directly on the moment definitions Eqs. (28) which are independent of the regularization methods and also relax the conditions that all coefficients \( \beta_1, \beta_2, \ldots, \beta_5 \) must be identical. We note that in Ref. [21], these authors already consider that \( \beta_1, \beta_2, \ldots, \beta_5 \) can be different and take empirical values allowing matching with a more accurate method. In these cases, those coefficients are used to fix the Knudsen layer effect that the R13 equation fails to capture completely. This empirical approach seems to be incompatible with the rigorous mathematical derivation of R13 equations.
E. Generation of pre-collision velocity

Molecular Dynamics simulation requires generating velocities of atoms that cross the control plane and collide with the wall. The (unnormalized) distribution of the latter is $|c_z| f^-(c, \omega)$ with $c_z < 0$ as seen in the previous section. In this paper, we use three types of distribution

- The Maxwell Boltzmann (MB) distribution

  For molecular gas $|C_z| f^-(c, \omega) = |C_z| f_{M}(C) f_{M}^\omega(\omega)$.  
  For monatomic gas $|c_z| f^-(c) = |c_z| f_{M}(C)$.  

The parameters of the distribution are the mean velocity $\mathbf{v}$ and the reduced temperature $\theta$. Using this distribution, we can model equilibrium system where the fluid is stationary $\mathbf{v} = 0$ and the temperature is uniform $\theta = \theta_w$ or non-equilibrium system by assuming that the gas adjacent to the wall is in local equilibrium with temperature and velocity different from the wall, i.e $\mathbf{v} \neq 0, \theta \neq \theta_w$.

- The Chapman-Enskog (CE) distribution [23, 24]

  $|c_z| f^-(c) = |C_z| f_{CE}(C)$.

In addition to temperature $\theta$ and mean velocity $\mathbf{v}$, there are also parameters associated to heat flux $q_k$ and shear stress $\sigma_{ik}$. This distribution is for non-equilibrium monatomic gas.

- The R13 distribution [20, 22]

  $|c_z| f^-(c) = |C_z| f_{R13}(C)$.

The last two distributions CE and R13 are for monatomic gases with expressions given in Appendix A. The generation of the input velocity is done via the Acceptance-Rejection approach. For example, in Ref. [25], a scheme to generate distributions in the form $f(c) = f_{M}(C)\Gamma(C)$ where $\Gamma(C)$ is a polynomial of $C$, is proposed. The distribution to be treated in this paper is slightly different since we are limited to the half-space $c_z < 0$ and there is a
function $|c_2|$ standing before $f_M(C)$ due to the flux definition (see Appendix A).

III. NUMERICAL SIMULATIONS AND RESULTS

A. Molecular Dynamics model

We study the collision of gas molecules, methane (CH$_4$) and carbon dioxide (CO$_2$) on a graphite wall (C atoms). The system contains two parts: the reservoir and the collision zone. The MD simulation is only done in the collision zone and the pre-collision velocity of gas molecules is generated directly at the stochastic boundary between the reservoir and the collision zone. Two graphite wall models will be considered. The first is a smooth wall model composed of 3 graphene layers with dimensions 17.04 Å × 17.22 Å (336 carbon atoms). The second is a rough model where a narrower band of graphene with surface 8.52 Å × 17.22 Å is added on the smooth surface (392 carbon atoms). Due to the surface geometry and symmetry, the scattering behavior of the smooth model is close to being isotropic and that of the rough model is anisotropic. We define the $x$ and $y$ directions as respectively “armchair” and “zigzag” directions, and the $z$ direction as the normal direction to the graphite plane. During the simulation, the lowest sheet is fixed and the second layer is maintained at constant temperature (350 K for CH$_4$ model and 600 K for CO$_2$ model) by Nose-Hoover thermostat with the relaxing temperature parameter equals to 100 time steps. The two final layers are free to interact with the gas molecules. We use periodic boundary conditions for $x$, $y$ directions, and we fix the height of box along $z$ direction. The gas-wall truncation distance is set to 15 Å from the upper layer, and the stochastic boundary is located at the truncation distance from the graphite wall. A simple sketch of the system and snapshots of MD simulations are shown in Fig. 1.

All MD simulations are performed with LAMMPS (Large-scale Atomic/Molecular Massively Parallel Simulator) package [26]. A typical simulation of $10^5$ collisions takes roughly 50 hours on an architecture of 92 Intel(R) Xeon(R) processors 2-3 GHz. The adaptive intermolecular reactive bond order (AIREBO) potential [27] is used for the interaction between the graphite carbon atoms. The CH$_4$ molecule is modeled as a united atom and its
FIG. 1. Simple sketch of the system. The stochastic boundary is indicated by the dashed line and the graphite wall is indicated by the solid line. Snapshots of MD simulations show the local orientation of the smooth (isotropic) and rough (anisotropic) systems.

interaction with graphite atoms is governed by the Lennard Jones (LJ) potential

\[ V = 4\epsilon \left[ \left( \frac{\sigma}{r} \right)^12 - \left( \frac{\sigma}{r} \right)^6 \right], \]

(32)

where \( r \) is the distance between two atoms under consideration and \( \sigma \) and \( \epsilon \) the parameters of the LJ model. Regarding the CO\(_2\) molecules, we do not consider the contribution of the internal degrees of freedom (bending/stretching) and use the rigid model [28]. The interaction of each site with the graphite atoms is also of LJ type with parameters taken from Ref. [5] (see Table II). After equilibrating the graphite system at the given temperature (10\(^6\) time steps of 1 fs), gas molecules are inserted one by one in the collision zone. Only after one collision event, i.e. a molecule interacts with the wall and goes out of the collision zone, another molecule is inserted in the zone from a random position at the stochastic boundary (for CH\(_4\) and CO\(_2\)) and with a random orientation (for CO\(_2\)). The residence time is considered negligibly small with respect to the flying time outside this zone and the velocities at the entrance and the outlet are collected (see Fig. 2).

Some comments can be made about the models for CH\(_4\) and CO\(_2\) used in the present work. Both rigid molecule models do not account for the vibrational internal degrees of freedom. The bending mode of CO\(_2\) associated with a wavenumber of 667 cm\(^{-1}\) [29] is the most concerned by an excitation due to collisions with the solid surface since this mode is the lowest energetic. The CO\(_2\) molecule collides with a surface at 600 K, i.e. \( k_B T = 417 \)
Using Boltzmann statistics, it can be estimated that only 20% of the CO\(_2\) molecules may be concerned by such an excitation. The lowest energetic vibrational mode of CH\(_4\) is the angle deformation mode associated with a wavenumber at 1306 cm\(^{-1}\) [29] and this molecule collides a surface at 350 K \((k_B T = 243 \text{ cm}^{-1})\). The Boltzmann statistics indicate that only 5% of the molecules would be excited. From these estimates and for simplification, CO\(_2\) and CH\(_4\) are kept rigid.

<table>
<thead>
<tr>
<th></th>
<th>(\sigma) [Å]</th>
<th>(\epsilon) [meV]</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH(_4) - C (Graphite)</td>
<td>3.550</td>
<td>5.547</td>
</tr>
<tr>
<td>C (CO(_2)) - C (Graphite)</td>
<td>3.059</td>
<td>2.418</td>
</tr>
<tr>
<td>O (CO(_2)) - C (Graphite)</td>
<td>3.197</td>
<td>4.091</td>
</tr>
</tbody>
</table>

TABLE II. LJ parameters for the interaction of CH\(_4\) and CO\(_2\) with the graphite surface.

FIG. 2. A collision showing the trajectory of a gas molecule.

We implement 3 simulation schemes to study gas-wall models and determine the model coefficients. They are different in terms of the gas state (equilibrium/non-equilibrium) and the associated velocity generator.

- **Batch average (BA) scheme**: We repeat the same incident velocity \((\omega', c')\) many times and record the reflected velocity \((\omega, c)\) which is a distribution. To generate samples, the incident velocity is taken from equilibrium distribution and the reflected velocity is aver-
By this way [30], we can examine the accommodation coefficients via their definition (Eq. (13)). In Fig. 3, We have plotted the input quantities $\langle Q \rangle_i$ against output quantities $\langle Q \rangle_o$, where $Q = c_x, c_y, c_z$ or $c^2 + I\omega^2/m$. In the ideal case, the data population should concentrate along a straight line and its slope corresponds to the constant accommodation coefficient $\alpha_Q$ (see section III.B for details).

- **Stochastic equilibrium (SE) scheme**: The reservoir is considered to be in equilibrium. The pre-collision velocity $(\omega', c')$ for each collision is generated using equilibrium distribution at zero mean velocity and at the same temperature as the graphite wall. By this way, we obtain numerical estimates of the density $B(\omega, c | \omega', c')$. The accommodation coefficients can be extracted using (14) (see section III.B for details).

- **Stochastic non-equilibrium (SN) scheme**: The non-equilibrium gas is considered. Depending on the problems, we use Maxwell-Boltzmann (MB), Chapman Enskog (CE) or R13 distribution as discussed in the previous section. The surface can be modeled atomistically (AM) as described from the beginning of the present section III.A. It can also be modeled statistically using the non-parametric (NP) model $B(\omega, c | \omega', c')$. The latter is constructed by the scattering results on the atomic model, which is detailed at the end of section III.B.

The simulation results will be analyzed using the theory we have proposed in the previous section.

**B. Determination of accommodation coefficients and construction of non-parametric wall model from collision data**

As mentioned previously, the accommodation coefficients are parameters based on Eq. (13). To verify this assumption, it is sufficient to study incident fluxes of constant velocity. Using BA scheme for the couples CH$_4$/CO$_2$-Graphite (smooth and rough surfaces), we set up 100 sampling groups and each group contains 500 collisions with the same incident velocity. These 100 incident velocities are drawn from equilibrium distribution at the same temperature as the wall. Then, we average the reflected values in the group for later anal-
Theoretically, if the ratio between the input and the output values are constant, the collision data will form a straight line. In Fig. 3, we find a strong correlation between the input and the output velocities. For the tangential velocity, despite some slight curvature the relation between the input and output is visibly linear for most of the data. However, the linear regression works less well for the normal velocity and especially for the kinetic energy. These data suggest that Eq. (13) is not valid for these cases and linear coefficients obtained by fitting (see Tab. III) are not representative.

Parametric studies based on varying the number of samples per input velocity from 20 to 1000 show that the scattering of energy data is always present. Given the fact that the results for $c_x$ and $c_y$ are clearly correlated, the energy data merit more detailed investigation to understand the origin of the deviation. We still use the same data and decompose the kinetic energy into tangential and normal components $c_n^2, c_t^2$. In Fig. 4(a) and (b), we can see that the tangential and normal kinetic energy data $\langle c_t^2 \rangle$ vs. $\langle c_t^2 \rangle$, $\langle c_n^2 \rangle$ vs. $\langle c_n^2 \rangle$ are strongly correlated. However, like the data $\langle c'_x \rangle$ vs. $\langle c_x \rangle$ and $\langle c'_y \rangle$ vs. $\langle c_y \rangle$ in Fig. 3, their slopes are different. There is a strong contrast between the tangential reflection $c_t^2$ which is more specular-like (energy mostly conserved after collision) and the normal reflection $c_n^2$ which is more diffusive-like (energy close to the walls’ after collision). On the other hand, in Fig. 4(c) and (d), the data $\langle c_t^2 \rangle$ vs. $\langle c'_n \rangle$ and $\langle c'_n \rangle$ vs. $\langle c_n^2 \rangle$ are scattered and the correlation is weak. It is suggested that the combination of different tangential and normal reflection behaviors can be responsible for the scattering of the total energy data $c^2$. As a final remark, the energy data scattering exists in literature models, especially for parametric models like ACL/CL or DM which are not based on a constant energy accommodation coefficient. Indeed, Fig 5 shows that the parametric kernel ACL/CL with suitable parameters has produced the same pattern, ie energy data scattering, as the MD results.

Next, with the SE scheme, we simulated $10^5$ collisions on graphite surfaces. The input velocities are taken from equilibrium distribution and results are presented in Fig. 6. From these figures we can analyze the correlation between incident and reflected velocities on graphite wall then we can compute the accommodation coefficients from Eq. (14) issued from [19]. It is noted that due to the equilibrium state, Eq. (13) takes the form $0/0$ and cannot be used to determine the accommodation coefficients in this case. We find that for
smooth graphite surface, the incident and reflected velocity data have significant correlation in $x$ and $y$ directions. Despite its crystalline nature, the surface behavior is isotropic and no visible difference is observed between directions $x$ and $y$. The accommodation coefficients calculated by the least-squares method [19] in Tab. III also confirm this remark. However, the influence of periodic roughness affects the anisotropy of the surface (second row figures) and the magnitude of the accommodation coefficients. We can see that the scattering is
FIG. 4. Detailed analysis of kinetic energy of CH\textsubscript{4} beamed on the smooth surface. The tangential kinetic energy component \(c_t^2 = c_x^2 + c_y^2\) and the normal kinetic energy component \(c_n^2 = c_z^2\) are used for analysis. Subfigures (a), (b), (c), (d) show the correlation between incident energy and reflective energy components.

FIG. 5. Energy data scattering observed from (a) MD simulations, (b) parametric ACL model with parameters \(\alpha_x = \alpha_y = 0.16\) (constant tangential momentum accommodation coefficient) and \(\alpha_z = 0.915\) (constant normal kinetic energy accommodation coefficient)

more diffusive along both directions but more pronounced for \(x\) direction. In all cases, the correlation between \(c_z\) and \(c'_z\) as well as the correlation between \(c^2\) and \(c'^2\) are very weak, and close to the diffusive wall.

We note that all analytical surface models in literature are based on the accommodation coefficients. Thus, we can construct scattering kernels which can serve as boundary conditions for atomistic method like MD or DSMC. The main advantage of these kernels is the simplicity in implementation but theirs drawbacks are their differences from the real surface behavior. This can be explained from the fact that they rely on the existence of the limited number of constant accommodation coefficients. To reconstruct \(B(\omega, c|\omega', c')\), we do not use any parameter and make no assumption on the distribution form except for the
FIG. 6. Collision data of incident and reflected velocities for the CH$_4$/CO$_2$-Graphite system. Both stochastic reservoir and graphite wall are maintained at 350 K for CH$_4$ and 600 K for CO$_2$. Columns (a), (b), (c) represent the velocity (nm/ps) along directions $x$, $y$, $z$, respectively, and column (d) the kinetic energy for CH$_4$ and CO$_2$ (the prefactor $m/2$ is removed for simplicity). The horizontal axis shows the incident values and the vertical axis the reflection values. The solid lines indicate the linear least square fit of incident and reflected values using Eq. (14). The diagonal dashed line and the horizontal dashed line indicate the zero accommodation case ($\alpha = 0$) and full accommodation case ($\alpha = 1$), respectively.

decomposition of each components

$$B(\omega, c|\omega', c') = \prod_{i=1}^{3} B_i(c_i|c_i') B_i^\omega(\omega_i|\omega_i').$$

(33)

The above relation reduces the realization of $B(\omega, c|\omega', c')$ to the realizations of independent
TABLE III. Accommodation coefficients computed by stochastic equilibrium (SE) simulation method using data of Fig. 6 and by batch average (BA) simulation method in Fig. 3: $\alpha_l$ coefficient is associated to the velocity $c_l$, $\alpha_e$ is the kinetic energy (the angular velocity of CH$_4$ is zero).

Without losing generality, we take the case of translation velocity $c_i$. The probability density function (PDF) of reflective velocity with given incident velocity $P(c_i|c'_i)$ can be determined by joint PDF of reflective-incident velocity $P(c_i, c'_i)$ and marginal PDF of incident velocity $P(c'_i)$ with relation:

$$B_i(c_i|c'_i) = P(c_i|c'_i) = \frac{P(c_i, c'_i)}{P(c'_i)}.$$  \hspace{1cm} (34)

The discrete collision data can be used to estimate the joint probability $P(c_i, c'_i)$ by histogram or kernel density estimation method. After determining $B_i(c_i|c'_i)$, we can use it to generate the outgoing velocities at any given incident velocities. This can be done via the use of conditional cumulative distribution function (CDF) $F(c_i|c'_i)$.

To illustrate the robustness of the non-parametric model, we take the case of anisotropic surface and plot the probability density of input and output velocities of different kernels together with the MD data in Fig. 7. The considered kernels are Dadzie-Meolans (DM) [12] and anisotropic Cercignani-Lampis (ACL) [31], and non-parametric kernel constructed numerically from MD simulations (see Appendix B). The accommodation parameters of
ACL kernel \( (\alpha_x \text{ associated to } c_x, \alpha_y \text{ to } c_y \text{ and } \alpha_z \text{ to } c^2_z) \) and DM kernels \( (\alpha_x \text{ and } \alpha_y \text{ are the same as ACL kernels and } \alpha_z \text{ is associated to } c_z) \) are determined by SE simulations using Eq. (14). One can find that the DM kernel that includes mirror-reflected mechanisms is very different from the true collision data, the probability density shows a strong discontinuity. The ACL kernel performs better but the non-parametric kernel is the most faithful to the MD data. Such differences can have significant influences on the results based on the kernel.

FIG. 7. Velocity probability density of MD simulations and from some scattering kernels: Dadzie-Meolans (DM) [12], Anisotropic Cercignani-Lampis (ACL) [31] and non-parametric (NP) kernel constructed from MD data. The MD data are from collision simulation of CH\(_4\) (350 K) at anisotropic atomic graphite wall (350 K). Columns \((a), (b), (c)\) represent the velocity \( \text{nm/ps} \) at directions \( x, y, z \), respectively, and column \((d)\) the kinetic energy for CH\(_4\) (the prefactor \( m/2 \) is removed for simplicity). The horizontal axis shows the incident values and the vertical axis the reflection values.
C. Direct computation of interface coefficients

The methods of determining the accommodation parameters like SE and BA depend on the existence of these constant coefficients. The BA gas beam experiments show that the postulates are rather restrictive (see e.g. Fig. 3, column d). Deriving those coefficients by fitting may correspond values in average sense without connection to the boundary conditions (16). Nevertheless, the slip and jump phenomena still exist and it is of interest to identify the coefficients associated to these effects. In this situation, one must make use of the non-equilibrium state of the gas near the wall. In principle, the more realistic the gas distribution is, the better interface coefficients we obtain. Before using more sophisticated distribution like Chapman-Enskog or R13 density, we shall examine the workability of a simpler distribution, MB at different temperatures and mean velocities. For $\beta_{1x}, \beta_{1y}$ associated to the friction coefficients, we use Maxwellian with the same temperature as the wall but non zero mean velocity. The latter is a vector lying in the bisector plane (making angle $\pi/4$ with respect to axis $x$ and $y$) with variable magnitude. For $\beta_2$ related to the Kapitza coefficient, we use Maxwellian with zero mean velocity but different temperatures. Specifically, the temperature ranges from 250 K to 450 K for CH$_4$, from 500 K to 700 K for CO$_2$ and mean velocities range from -0.05nm/ps to 0.05nm/ps for both CH$_4$ and CO$_2$. At this stage, we have two wall models:
- Atomic wall model (AM)
- Non-parametric wall model (NP)
for comparisons. To examine the connection between the quantities in the boundary models, results for the slip velocities, the temperature jump, stress and heat flux obtained by Eqs. (18-23) are plotted together in Fig. 8.

We see a clear linear relation between $-\hat{\sigma}_{kz}$ and $\hat{v}_z$ as well as between $-\hat{q}_z$ and $\Delta \hat{\theta}$ in Fig. 8. The slopes of fitted lines represent the value of the dimensionless friction coefficient $\beta_{1k}$ (subfigures $a$ and $b$) and the dimensionless thermal coefficient $\beta_2$ (subfigure $c$). The friction coefficient of the $x$ direction increases 8 times from smooth wall to rough wall, compared with the 3 times increase in the $y$ direction. It’s also interesting to see that the $\beta_2$
FIG. 8. Method of computing coefficients $\beta$s by non-equilibrium simulation, MB distribution and atomic wall model (AM). (a) $-\hat{\sigma}_{kz}$ and $\hat{v}_k$ of the isotropic graphite wall. (b) $-\hat{\sigma}_{kz}$ and $\hat{v}_k$ of the anisotropic graphite wall. (c) $-\hat{q}_z$ and $\Delta \hat{\theta}$ are calculated by incident and reflective velocities on isotropic and anisotropic graphite wall.

In terms of friction coefficient $\beta_{1k}$, the results of the non-equilibrium method with MB distribution are very close to those obtained with the methods based on equilibrium distribution. This is reasonable since the BA method also shows that the accommodation model works well for this case. For thermal coefficient $\beta_2$, the SN method seems to agree better with the coefficient derived from SE method than the BA method. This observation can be explained from the scattering data in the BA method, meaning that the theoretical definition of the thermal accommodation coefficient is no longer valid. In this case, an effective coefficient which reproduces the thermal jump effect can be determined.
TABLE IV. The $\beta$ coefficients of CH$_4$ and CO$_2$ computed by SN methods and compared with values obtained from accommodation coefficients (SE method, Tab. III) via formula $\beta_{1k} = \alpha_k/(2 - \alpha_k)$, $\beta_2 = \alpha_e/(2 - \alpha_e)$. MB: Maxwell-Boltzmann distribution, CE: Chapman-Enskog distribution, AM: atomic model for wall, NP: the wall is modeled by non-parametric scattering kernel $B(c|c')$, instead of atomic wall (as in MB-AM method). To increase the precision of the slip coefficients for smooth (isotropic) surface, we average values along $\beta_{1x}$ and $\beta_{1y}$.

D. Influences of non-equilibrium distributions and discussion

Essentially based on the same procedure as previous subsection, we study the influence of the near wall distribution and the surface models on the interface coefficients. We focus on the monatomic gas CH$_4$, NP wall model and the following non-equilibrium distributions: - Chapman-Enskog distribution - R13 distribution

To generate CE and R13 distributions, we use the Acceptance-Rejection approach described in Appendix A. In addition to temperature and mean velocity, the CE and R13 distributions require input fluxes and moments which are generally unknown. We carry out the following iterations:

- Step $n$
  - Generation of input distribution using CE (or R13) distribution with average moments of the previous steps $n - 1$.
  - Compute the output distribution using the kernel $B(c|c')$.
  - Calculate the average moments at the wall from input and output distributions.
  - Next step.

The loop stops when all the average moments converge (see e.g Fig. 9). The input
distribution at the first step \( n = 1 \) can be initialized with MB distribution as the previous SN/MB scheme, i.e all fluxes (heat fluxes, stress, etc...) are set zero. The latter quantities become non-zero after the initialization \( (n \geq 2) \) and we can effectively use the CE and R13 generator. Numerical tests show that while the average moments at convergence depend on the initialized values, the value of \( \beta \) coefficients are insensitive to them.

![Graphs showing convergence of different parameters](image)

**FIG. 9.** Convergence test of CE-NP kernel iteration for rough anisotropic surface. Subfigures (a): Friction coefficient \( \beta_{1x} \), (b) Gas temperature (c) Gas velocity.

About \( 10^7 \) collisions with scattering kernel are simulated. In Tab. IV, we find that all the methods yield results close to each other, especially for isotropic surface. For anisotropic surface and friction coefficient along \( x \), the method based on CE shows some discrepancies with the rest (although the coefficient is of the same range of order). The visible differences can be explained from the influences of heat flux and shear stress at convergence. By examining in detail the convergence of \( \beta_{1x} \) in Fig. 9, we find the final parameter is different from the first iteration one. It is suggested that the presence of the roughness perturbs considerably the phase density. Due to the realistic kernel \( B(c|c') \), the output is not necessarily of the same distribution class as the input. This raise questions on the use of CE distribution near the wall, especially the component \( c_x \) along the roughness directions. Another possible reason is that there may be a considerable coupling between different moments which must be taken into account in the phenomenological equations. The complete answer can only be found from flow simulations using the same surface model, but with different input distributions.

Next we consider another non-equilibrium distribution associated to R13 moment equations. The boundary conditions are originally derived for Maxwell scattering kernel with one accommodation coefficient (isotropic surface) and all \( \beta \) coefficients being identical. In
order to test the R13 generator, we use first the Maxwell kernel with $\alpha = 0.3$ and obtain interface coefficients. From Tab. IV, we find that these coefficients are overall in good agreement with the theory prediction $\beta = \alpha/(2 - \alpha) = 0.176$. Most of the computed coefficients are within less than 1% error from the analytical value. This is a good starting point to proceed with our graphite surface.

We use NP model for our graphite surface. The results show that coefficients $\beta_1, \beta_2$ agree with the computed values for NSF equations listed in Tab. V. Coefficients $\beta_3, \beta_4, \beta_5$ correspond to boundary conditions for higher order terms. As the overall remark, all coefficients $\beta$ are different, showing that the use of Maxwell kernel cannot capture correctly the boundary conditions at the wall. In this case, the present approach can provide an alternative and reliable solution for any surface.

<table>
<thead>
<tr>
<th>Surface</th>
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<th>$\beta_1$</th>
<th>$\beta_2$</th>
<th>$\beta_3$</th>
<th>$\beta_4$</th>
<th>$\beta_5$</th>
</tr>
</thead>
<tbody>
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<td>0.177</td>
<td>0.160</td>
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</tr>
<tr>
<td>Graphite</td>
<td>NP</td>
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<td>0.373</td>
<td>0.024</td>
<td>0.028</td>
<td>0.129</td>
</tr>
</tbody>
</table>

TABLE V. $\beta$ coefficients of R13 equations using Maxwell kernel ($\alpha = 0.3$) and non-parametric scattering kernel.

IV. CONCLUSIONS

It is known that gas flows at high Knudsen number are present in micro-nanopore underground. To accurately obtain macroscopic transport behavior (permeability for example), we must use relevant gas model and boundary conditions at the pore scale. The present paper is devoted to the construction of gas-wall interaction models and to the determination of boundary conditions for continuum equations such as Navier-Stokes-Fourier or R13 equations.

Based on MD simulation of independent collisions of CH$_4$ and CO$_2$ rigid molecules on graphite surface, we collect data of pre- and post-collision and numerically recover the scattering kernel for the gas-wall couples. Specifically, it can be used to construct non-parametric models capable of generating a distribution of post-collision velocity, given the pre-collision velocity. The effective accommodation coefficients can also be obtained from
the scattering kernel using a different fitting procedure.

In the general case, a method is proposed to directly compute the parameters of any phenomenological boundary conditions without using the concept of accommodation coefficients. The approach relies on the general scattering kernel issued from MD with suitable input gas distribution, which is similar to the theoretical method of Grad [24]. Such an approach is of interest since it can capture more accurately the distribution of reflection velocity. There are still some drawbacks to the current version which will be improved in the future, for example, the use of rigid gas molecule model and the assumptions of independence between the velocity components. Another interesting subject which has not been treated in this work is the Knudsen layer effect. We note that by simulating flows using non-parametric kernel, one can obtain correction coefficients to the microslip obtained by the present paper.

**APPENDIX**

**A. Generation of non-equilibrium distribution**

Generating pre-collision velocity of particle requires first the PDF of initial velocity. The Maxwell-Boltzmann distribution is used for a equilibrium dilute gas at temperature \( T \). In order to generate the pre-collision velocity, we rewrite the flux associated to the normalized MB distribution \( f^*_M(\tilde{C}) \) as:

\[
f^*_M(\tilde{C}) = 2\frac{|\tilde{C}_z|}{\pi} \exp\left(-\tilde{C}^2\right), \quad \tilde{C}_z < 0,
\]

where \( \tilde{C} = C/\sqrt{2\theta} \) and \( \theta = k_B T/m \). This distribution is constituted of three independent distributions: two normal distributions along \( x, y \) and a Rayleigh distribution along \( z \). Then we can calculate the flux associated to Chapman-Enskog distribution \( f^*_{CE}(\tilde{C}) \) by the relation:

\[
f^*_{CE}(\tilde{C}) = \Gamma_{CE}(\tilde{C}) f^*_M(\tilde{C}), \quad \tilde{C}_z < 0,
\]
\[ \Gamma_{CE}(\tilde{C}) = 1 + \left( \tilde{q}_x \tilde{C}_x + \tilde{q}_y \tilde{C}_y + \tilde{q}_z \tilde{C}_z \right) \left( \frac{2}{5} \tilde{C}^2 - 1 \right) + 2 \left( \tilde{\sigma}_{xy} \tilde{C}_x \tilde{C}_y + \tilde{\sigma}_{xz} \tilde{C}_x \tilde{C}_z + \tilde{\sigma}_{yz} \tilde{C}_y \tilde{C}_z \right) \]

\[ + \tilde{\sigma}_{xx} \tilde{C}_x^2 + \tilde{\sigma}_{yy} \tilde{C}_y^2 + \tilde{\sigma}_{zz} \tilde{C}_z^2, \]

(37)

with

\[ \tilde{\sigma}_{ij} = \frac{\sigma_{ij}}{p}, \quad \tilde{q}_i = \frac{q_i}{p \sqrt{\theta/2}}, \quad p = mn\theta; \quad i, j = x, y, z. \]  

(38)

Similarly, the R13 distribution can be calculated from MB distribution by the relation:

\[ f_{R13}^*(\tilde{C}) = \Gamma_{R13}(\tilde{C}) f_{M}^*(\tilde{C}), \quad \tilde{C}_z < 0, \]

(39)

where

\[ \Gamma_{R13}(\tilde{C}) = 1 + \varphi_{13}(\tilde{C}) + \varphi_{R1}(\tilde{C}) + \varphi_{R2}(\tilde{C}). \]

\[ \varphi_{13}(\tilde{C}) = \left( \tilde{q}_x \tilde{C}_x + \tilde{q}_z \tilde{C}_z \right) \left( \frac{2}{5} \tilde{C}^2 - 1 \right) + 2 \tilde{\sigma}_{xx} \tilde{C}_x \tilde{C}_z + \tilde{\sigma}_{yy} \tilde{C}_y^2 + \tilde{\sigma}_{zz} \tilde{C}_z^2, \]

\[ \varphi_{R1}(\tilde{C}) = \frac{1}{3} \left( \tilde{m}_{xxx} \tilde{C}_x^3 + \tilde{m}_{zzz} \tilde{C}_z^3 \right) + \tilde{m}_{xzz} \tilde{C}_x^2 \tilde{C}_z + \tilde{m}_{zzx} \tilde{C}_x \tilde{C}_z^2 + \tilde{m}_{zyy} \tilde{C}_y^2 \tilde{C}_z + \tilde{m}_{zyz} \tilde{C}_y \tilde{C}_z^2, \]

\[ \varphi_{R2}(\tilde{C}) = \left( \tilde{R}_{xx} \tilde{C}_x^2 + \tilde{R}_{yy} \tilde{C}_y^2 + \tilde{R}_{zz} \tilde{C}_z^2 + 2 \tilde{R}_{xy} \tilde{C}_x \tilde{C}_y \right) \left( \frac{\tilde{C}^2}{7} - \frac{1}{2} \right) + \frac{\tilde{R}}{30} \left( \tilde{C}^4 - 5 \tilde{C}^2 + \frac{15}{4} \right), \]

(40)

with

\[ \tilde{m}_{ijl} = \frac{m_{ijl}}{p \sqrt{\theta/2}}, \quad \tilde{R}_{ij} = \frac{R_{ij}}{p \theta}, \quad \tilde{R} = \frac{R}{p \theta}, \quad i, j, l = x, y, z. \]

(41)

Since the CE and R13 distributions have analytical PDF expression, we can generate random velocities corresponding to these distributions by the Acceptance-Rejection method. The specific steps are as follows:

1. Find \( B_{CE} = \max(|\sigma_{ij}|, |q_i|) \) for distribution CE, \( B_{R13} = \max(|\sigma_{ij}|, |q_i|, |m_{ijkl}|, |R_{ij}|, |R_{ij}|) \) for distribution R13 .

2. Set amplitude parameter \( A_{CE} = 1 + 30B_{CE} \) and \( A_{R13} = 1 + 60B_{R13} \).
3. Generate a normalized velocity $\hat{C}_{MB}$ obeying MB distribution (Eq. (35)), and a uniform random number $U(0,1)$.

4. If $\Gamma_{CE}(\hat{C}_{MB}) \geq AU$, accept $\hat{C}_{MB}$ as normalized CE distribution velocity $\hat{C}_{CE}$, and if $\Gamma_{R13}(\hat{C}_{MB}) \geq AU$, accept $\hat{C}_{MB}$ as normalized R13 distribution velocity $\hat{C}_{R13}$; else reject this velocity and return to step 3;

5. The real velocity is $c = \sqrt{2\theta \hat{C}_{MB}} + v$, $v$ is the mean velocity.
During the Acceptance-Rejection process, the function $\Gamma(\hat{C})$ can be negative. This velocity is rejected in this case.

**B. Method for generating non-parametric kernel**

Different from parametric scattering models, like CL or Maxwell, the non-parametric wall model is not specified a priori but is instead determined from data. Using the discrete incident-reflection velocities data, we can generate the reflective velocities at any given incident velocities. The implementation of Eq. (34) is as follows:

1. Discretize the velocity space $c_i'$ and $c_i$ to a series of velocity points with sufficient small interval. Choose a volume $\Delta c_i'$ and let us call class $c_i'$ the collisions in the volume centered at $c_i'$.

2. Use sliding window method to count the number of collision $\Delta N$ in class $c_i'$.

3. Calculate discrete CDF of every class $c_i'$ as $F(x|c_i') = \frac{\Delta N(c_i \leq x)}{\Delta N}$.

4. Generate a uniform random number $U$ between 0 to 1, then the reflective velocity is $c_i \sim F^{-1}(U|c_i')$.

The procedure is shown in Fig. 10. The reflective rotation velocity can be obtained by the same procedure.

**FIG. 10. Generation of velocity using non-parametric kernel derived from MD collision clouds**
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\( \langle c_x \rangle \) (nm/ps) \( \langle c_y \rangle \) (nm/ps) \( \langle c_z \rangle \) (nm/ps) \( \langle c^2 + \frac{1}{m} \omega^2 \rangle \) (nm\(^2\)/ps\(^2\))
(2a) $CH_4 - An_i$

(2b)

(2c)

(2d)
\[ \langle c^2 \rangle \left( \text{nm}^2 / \text{ps}^2 \right) \]

\[ \langle c'^2 \rangle \left( \text{nm}^2 / \text{ps}^2 \right) \]

Atomic

(a)
\begin{align*}
\langle c^2 \rangle &= \left( \frac{\text{nm}^2}{\text{ps}^2} \right) \\
\langle c'^2 \rangle &= \left( \frac{\text{nm}^2}{\text{ps}^2} \right)
\end{align*}
\( \beta_{1k} (iso) \)

\[ C_{H_4} \left( \frac{-\hat{\sigma}_{kz}}{\hat{v}_k} \right) \]

\( \beta_{1k} (ani) \)

\[ \hat{\sigma}_{kz} \]

\[ \Delta \hat{\theta} \]

Isotropic wall

Anisotropic wall
Isotropic wall
Anisotropic wall