Recovery of Multicomponent Shale Gas from Single Nanopores

Haiyi Wu, Yadong He, and Rui Qiao©

Department of Mechanical Engineering, Virginia Tech, Blacksburg, Virginia 24061, United States

Supporting Information

ABSTRACT: The adsorption of multicomponent gas mixtures in shale formations and their recovery are of great interest to the shale gas industry. Here we report molecular dynamics simulations of the adsorption of methane/ethane mixtures in 2 and 4 nm-wide nanopores and their recovery from these nanopores. Surface adsorption contributes significantly to the storage of methane and ethane inside the pores, and ethane is enriched inside the nanopores in equilibrium with bulk methane—ethane mixtures. The enrichment of ethane is enhanced as the pore is narrowed but is weakened as the pressure increases due to entropic effects. These effects are captured by the ideal adsorbed solution (IAS) theory, but the theory overestimates the adsorption of both gases. Upon opening the mouth of the nanopores to gas baths with lower pressure, both gases enter the bath. The production rates of both gases show only weak deviation from the square root scaling law before the gas diffusion front reaches the dead end of the pores. The ratio of the production rate of ethane and methane is close to their initial mole ratio inside the nanopore despite the fact that the mobility of pure ethane is smaller than that of pure methane inside the pores. Scale analysis and calculation of the Onsager coefficients for the transport of binary mixtures of methane and ethane inside the nanopores suggest that the strong coupling between methane and ethane transport is responsible for the effective recovery of ethane from the nanopores.

I. INTRODUCTION

Natural gas production from shale formations has received extensive attention recently and can potentially lead to a new global energy source. The U.S. Energy Information Administration shows that shale gas has provided the largest source of the growth in U.S. natural gas supply during the past decade, and its share is expected to grow continuously in the future. The composition of shale gas varies among shale reservoirs. While methane is always the most abundant component in shale gas, hydrocarbons with higher molecular weight are also present. In particular, ethane is the usually the second primary component in shale gas and can account for up to 15 vol % of the recovered gas. The effective extraction of these hydrocarbons, which are much more valuable than methane, is of great interest for the shale gas industry.

A distinguishing feature of shale formations is that, while they typically feature pores ranging from subnanometer to millimeters in size, their porosity is often dominated by nanopores smaller than 10–100 nm. As such, much of the gas is stored inside narrow nanopores as adsorbed gas and free gas. Based on extensive research on transport of fluids in nanopores, e.g., layering of fluids on pore walls modulates their transport through narrow pores, one can expect that the thermodynamic and hydrodynamic properties of gas inside such pores, which govern the extraction of gases from the pores, can be quite different from those of bulk gas. Tremendous progress has been made in understanding the adsorption and transport of single-component gas (usually methane) inside nanopores in the past decade, but those of multicomponent gas mixtures are much less well understood. Among the available studies, much attention is focused on the competitive adsorption of CH4/N2/CO2 gas species in shale, which plays an important role for enhancing shale gas recovery. However, a general understanding of the adsorption and transport of multicomponent shale gas under realistic reservoir conditions (e.g., considering CH4/C2H6 mixtures) is still limited.

Research on the adsorption and transport of multicomponent shale gas in nanopores can potentially benefit from the extensive works on the thermodynamic and transport properties of multicomponent gases in other industrial applications. For example, previous studies have attempted to predict the multicomponent adsorption via the extend Langmuir (EL) model and the ideal adsorbed solution (IAS) model. In addition to these theoretical models, previous simulations also indicated that there exits complex competition between the adsorbed species.

We are interested in understanding the storage of multicomponent gas mixtures in shale formations and their recovery during gas extraction using numerical simulation. Simulation of shale gas storage and recovery in shale formations appears extremely difficult because of the vast spectrum of pore sizes involved, the diverse surface chemistry of the pores, and the complicated connectivity between nanopores. Nevertheless, prior studies show that the overall shale gas recovery is mainly controlled by the transport of gas from the narrow pores to wide secondary fractures, and the essential features of shale gas adsorption and transport inside nanopores can be well captured by pore scale modeling with simple pore geometries, e.g., cylindrical or slit pores. In this work, we study the adsorption of CH4/C2H6 gas mixtures in single nanopores and their subsequent recovery from these pores using molecular dynamic (MD) simulation. In particular, we focus on the competitive adsorption of the two gases in the adsorbed phase of the nanopore and the gas recovery behavior of each species from the nanopore. Our work is inspired in part by prior

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studies on recovery of multicomponent shale gas,\textsuperscript{16,20,21,28} which have provided many useful insights into the behavior of adsorption and transport of gas mixture. Nevertheless, some important issues have not been addressed yet in these studies. For example, the competitive adsorption of CH\textsubscript{4}/C\textsubscript{2}H\textsubscript{6} gas inside nanopores and its impacts on shale gas storage is still not well understood, and a molecular view of the recovery of multicomponent gas mixture from nanopores is not yet available. By using MD simulations to study the gas recovery from nanopores, we will explicitly address these issues.

The rest of the paper is organized as follows: In section II, we introduce the model and methods for simulations of the equilibrium isotherm and dynamic extraction of CH\textsubscript{4}/C\textsubscript{2}H\textsubscript{6} mixture from single nanopores. In section III, we present our simulation results on adsorption isotherm (with a focus on the competition between methane and ethane in the adsorbed phase) and the recovery characteristics of the two gases (with a focus on the recovery of ethane). Finally, conclusions are drawn in section IV.

II. MD SIMULATION DETAILS

Figure 1 shows a schematic of the MD system. The system features a slit pore, a gas bath with a constant pressure, a piston, and the shale gas molecules inside the pore and the gas bath. The system is periodic in the y- and z-directions. The pore’s right end is permanently sealed. The center-to-center widths \( d \) of the investigated nanopores are 2 and 4 nm since pores with such small sizes are abundant in shale formations.\textsuperscript{6,20} Due to the finite size of the methane molecules and the wall atoms, the accessible pore width in the MD model \( d_\text{a} \) is about 1.62 and 3.62 nm, respectively. The pore length \( L \) is 202.5 nm. The left side of the gas bath is bounded by a piston plate (green plate in Figure 1), whose atoms move only in the x-direction. The constant gas pressure inside the gas bath is maintained by applying an external force \( F_{\text{ext}} = L_y L_z P_{\text{bath}} \) on the piston plate \( (L_y = 13.5 \text{ nm and } L_z = 12.8 \text{ nm are the plate’s width in y- and z-directions, respectively; } P_{\text{bath}} \text{ is the desired gas pressure inside the bath.}) \). All forces experienced by the piston plate are averaged and redistributed evenly on the atoms of the piston plate such that the plate remains planar.

Two types of simulations are performed: adsorption isotherm simulations and gas recovery simulations. For the adsorption isotherm simulations, initially methane and ethane molecules are placed in the gas bath and equilibrated; the pore is empty and sealed by “blocker” atoms at its mouth (see the black spheres in Figure 1). During the simulations, the pore is made open by removing the blocker atoms at its mouth, thus gas molecules move from the gas bath into the nanopore until a new equilibrium is reached. By using a gas bath that is sufficiently large and adjusting the number of gas molecules inside the gas bath by trial and error, the mole fraction of methane and ethane molecules inside the gas bath have been controlled to be 4:1 within a few percent. The amount of methane and ethane adsorbed inside the nanopores from the equilibrium run is used to determine the gas reserve inside the nanopores.

For the gas recovery simulations, the system is setup with pure methane in the gas bath and gas mixtures (CH\textsubscript{4}/C\textsubscript{2}H\textsubscript{6} molar ratio is 4:1) in the pore. Initially, the left end of the pore is sealed, and the system is equilibrated with higher pressure \( (P_0 = 200 \text{ bar}) \) in the pore and lower pressure in the bath \( (P_{\text{bath}} = 20 \text{ bar}) \). At \( t = 0 \), the blocker atoms at the left end of the pore are removed. Because the initial pressure inside the pore is higher than that in the gas bath, gas recovery is initiated.

Methane and ethane molecules are modeled using the TraPPE force field.\textsuperscript{30} Briefly, methane molecules are modeled as structureless, spherical molecules; ethane molecules are modeled as dumbbell-shaped molecules featuring two united atoms each representing one CH\textsubscript{3} site, and the two united atoms are separated by a bond length of 1.54 Å. While it is desirable to explicitly resolve all hydrogen atoms in the methane and ethane molecules, prior works show that the united atom approach taken here already allows accurate prediction of gas adsorption on planar walls well with lower computational cost.\textsuperscript{6,20} The intermolecular interactions between an atom “\( m \)” and another atom “\( j \)” are modeled using the Lennard-Jones (LJ) potential

\[
\phi_{mj} = 4 \epsilon_{mj} \left( \frac{\sigma_{mj}}{r} \right)^{12} - 2 \left( \frac{\sigma_{mj}}{r} \right)^{6}
\]

where \( \sigma_{mj} \) and \( \epsilon_{mj} \) are the LJ parameters for the pair (\( mj \)) and \( r \) is the distance between the two atoms. The piston plate is modeled as a square lattice of carbon atoms (lattice spacing 0.3 nm). We confirmed that the simulation results are independent of the piston mass. Following ref 31, the pore walls are modeled as semi-infinite slabs constructed from a FCC lattice oriented in the (111) direction and the lattice constant is 0.54 nm. To avoid the significant computational cost of explicitly simulating all wall atoms, only the innermost layer of the wall atoms (i.e., the layer in contact with gas molecules) are explicitly modeled. The wall atoms beneath this layer are treated collectively as an implicit slab, and gas molecules inside the system interact with
the implicit slab of wall atoms (the shaded region in Figure 1) via the LJ 9-3 potential

\[ \varphi_{\text{lw}} = \frac{2}{3} \sigma_{\text{lw}}^6 \epsilon_{\text{lw}} \left[ \frac{2}{15} \left( \sigma_{\text{lw}} / \delta_0 \right)^9 - \left( \sigma_{\text{lw}} / \delta_0 \right)^3 \right] \]

where \( \rho_w \) is the number density of the wall atoms in the implicit slab, \( \epsilon_{\text{lw}} \) and \( \sigma_{\text{lw}} \) are the LJ parameters for the interactions between the wall atoms and the gas molecules, and \( \delta_0 \) is the closest distance between a gas particle and the surface of the implicit slab. The LJ parameters for the gas molecules and the wall atoms are summarized in Table 1.\(^{30}\) The interaction parameters for the methane–ethane (i.e., CH\(_4\)–CH\(_3\)H) pair are determined from those for the CH\(_4\)–CH\(_4\) and CH\(_3\)H–CH\(_3\)H pairs using the Lorentz–Berthelot combination rule. With the structure of the wall and the LJ parameters for gas–wall interactions chosen here, the adsorption behavior of pure methane is found to be similar to those found in prior studies of methane adsorption in organic shale pores.\(^ {32}\)

Simulations are performed using the Lammmps code\(^ {13}\) with a time step size of 2 fs. The cutoff lengths for gas–gas interactions and gas–wall interactions are 1.4 and 1.5 nm, respectively. For each simulation, the volume of the system and the number of gas molecules inside the entire system are both kept constant. The pore wall atoms are fixed. We emphasize that, although the dimensions of the simulation box are fixed, the volume of the gas bath can change during gas recovery and adsorption isotherm simulations because its pressure is maintained by the piston (the simulation box is large enough in the \( x \)-direction so that the piston plate never protrudes out of the left boundary of the simulation box). The temperature of the gas molecules is maintained at 353 K using the Nose–Hoover thermostat. In each simulation, the density, pressure, and temperature of gas in both the nanopore and the gas bath are computed on-the-fly. Each simulation is repeated three times with different initial configurations to obtain reliable statistics. Specifically, before gas recovery simulations start, the gas molecules in the bath and in the pore are separated by the blocker atoms (see Figure 1) and the system is equilibrated for 2 ns. The configurations of the system at three time instants (1.0, 1.5, and 2.0 ns) of the equilibrium run are saved. We then perform three gas recovery simulations, in which the initial configuration is taken from those three equilibrium configurations just saved.

### III. RESULTS AND DISCUSSION

**Methane and Ethane Storage in Nanopores.** As is customary in studying material adsorption in porous materials, the storage of a gas species \( i \) inside a nanopore is measured per unit surface area of the pore using

\[ n_{i,\text{tot}} = \frac{1}{2} \int_{-d/2}^{d/2} \rho_i(z) \, dz \]

where \( \rho_i(z) \) is the density of the species across the pore, \( d \) is the accessible pore width, and the factor 1/2 appears because each slit pore has two walls. Because the gas density is strongly inhomogeneous across nanopores (see, for example, the methane density profile in the inset of Figure 2b), researchers often partition shale gas inside nanopores into adsorbed gas and free gas.\(^ {3,4}\) To delineate these two forms of gas and to help understand the storage of methane–ethane mixture in nanopores, we first examine the storage of pure methane and pure ethane in nanopores. Here, gas molecules in the density peaks next to the wall (shaded region in the inset of Figure 2b, hereafter referred to as the adsorption layer) are taken as adsorbed gas and said to belong to the adsorbed phase. The rest of the gas molecules inside the pore are considered as free gas and said to belong to the free phase. The adsorbed gas of species \( i \) is quantified using

\[ n_{i,\text{ads}} = \int_{-d/2}^{-d/2+\delta} \rho_i(z) \, dz = \int_{d/2-\delta}^{d/2} \rho_i(z) \, dz \]

\( \delta \) is the thickness of the adsorbed gas layer in the \( z \)-direction (taken as 4 Å here). In principle, \( \delta \) may vary with pore size and

<table>
<thead>
<tr>
<th>atom pair</th>
<th>parameters</th>
<th>( \sigma ) (nm)</th>
<th>( \epsilon/k_B ) (K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>methane–methane</td>
<td>0.3730</td>
<td>148.0</td>
<td></td>
</tr>
<tr>
<td>ethane–ethane(^ {6})</td>
<td>0.3750</td>
<td>98.0</td>
<td></td>
</tr>
<tr>
<td>methane–wall</td>
<td>0.3315</td>
<td>207.2</td>
<td></td>
</tr>
<tr>
<td>ethane–wall(^ {10})</td>
<td>0.3325</td>
<td>168.6</td>
<td></td>
</tr>
</tbody>
</table>

\(^{\text{a}}\)This is for the interactions between the CH\(_4\) sites of two different ethane molecules. \(^{\text{b}}\)This is for the interactions between a wall atom and one CH\(_3\) site of an ethane molecule.

Figure 2. (a) Isotherms of the storage of methane in 2 nm-wide nanopores in equilibrium with a pure methane bath and the contribution by the adsorbed and free gas to the total gas storage. (b) Isotherms of methane and ethane adsorbed on the walls of a 2 nm-wide nanopore when the pore is in equilibrium with pure methane or ethane gas baths. (inset) Density profile of methane across a 2 nm-wide slit pore when the pore pressure is 100 bar. Gas molecules in the region I (region II) are taken as the adsorbed (free) gas.

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pressure. However, examination of the gas density profiles in different pores at various pressures (see Figure S3 in the Supporting Information) show that, within the parameter space explored here, such variation is small and can be safely neglected.

Figure 2a shows the isotherm of the methane storage inside a 2.0 nm-wide pore and its contribution by free and adsorbed gases. We observe that the adsorbed methane accounts for more than 60% of all methane stored inside 2.0 nm-wide pores in the range of pressure investigated here, and their contribution increases as the pressure decreases. Similar trends are observed when pure ethane is stored inside the same pore (see Figure S1). However, as shown in Figure 2b, at the same pressure, the amount of ethane adsorbed on the walls of 2 nm-wide pores is larger than that of methane. This is expected because the adsorption of gas molecules on pore walls is driven by the attractive van der Waals forces between the gas molecules and the wall, and ethane molecules, which contain two CH₃ sites, have stronger van der Waals interactions with the wall than methane molecules. The stronger adsorption of ethane on pore walls is also observed in the 4 nm-wide pores (see Figure S2).

Next, we examine the gas storage inside nanopores in equilibrium with bath of binary methane—ethane mixtures. In all cases studied, the mole fraction of ethane in the gas bath is set to 0.2 ± 0.01. Figure 3a compares the storage of methane and ethane in a 2 nm-wide pore as a function of pressure (data for methane—ethane mixture storage in 4 nm-wide pore exhibit similar trend and are shown in Figure S4). The nonlinear increase of the gas storage with pressure, a signature of significant contribution of surface adsorption to gas storage, is similar to that of pure gas. Figure 3b shows that the molecular fraction of ethane inside the nanopores is higher than that in the bath, i.e., ethane is enriched in the pores. The enrichment is stronger in the 2 nm-wide pores. Figure 3c further shows that, in both 2 nm- and 4 nm-wide pores, the enrichment of ethane inside pores is mainly caused by its enrichment in the wall-adsorbed phase. The latter is expected based on the results shown in Figure 2b. However, Figure 3c shows that the ethane enrichment decreases as the pressure increases beyond ~100 bar. This means that the adsorption of methane on the pore walls becomes more competitive than ethane as pressure increases beyond 100 bar. To better understand this result, we examine the isotherm for the concurrent adsorption of methane and ethane on the walls of 2 nm-wide pores.

Figure 4 shows the isotherms for the adsorption of methane and ethane on the walls of 2.0 nm-wide pores. At P ≤ 100 bar, the adsorption of methane and ethane increases rapidly with pressure. As pressure increases further, the increase of the adsorption of both methane and ethane slows down. However, the slowdown is much more distinct for ethane and the adsorption of ethane reaches a plateau at P ~ 150 bar. In other words, ethane becomes less competitive in adsorbing on the pore walls than methane, and this leads to the reduced ethane enrichment at elevated pressure (see Figure 3c). The less competitive adsorption of ethane at high pressure can be understood as follows. At low to moderate pressure, the gas adsorption layer on the wall is loosely packed and gas—wall interactions play the most important role in determining gas adsorption. Since ethane molecules interact more strongly with the pore walls than methane molecules, they outcompete methane in adsorption on pore walls. As the pressure increases, the gas adsorption layer becomes more densely packed and the entropic effects become more important. Since ethane molecules are larger than methane molecules, their entropy is lower than that of the methane molecules at elevated pressure. Therefore, ethane molecules become less competitive in adsorption on pore walls than the methane molecules. We note that a similar effect has been predicted theoretically in prior studies of binary adsorption of methane—ethane in zeolites.34–36

We next investigate to what extent the isotherms computed using MD simulations can be predicted using existing theories. Among the available theories on multicomponent gas adsorption, the ideal adsorbed solution (IAS) theory24 is popular due to its simplicity and good predicting power. The
B. Methane and Ethane Recovery from Nanopores. In this section, we study the recovery of gas from a single nanopore (d = 2 or 4 nm) to a large gas bath. As detailed in section II, initially, each nanopore is filled with a methane/ethane mixture (CH₄:C₂H₆ molar ratio 4:1; total pressure 200 bar), and the gas bath contains pure methane. The pressure in the gas bath is maintained at 20 bar throughout the simulation. At t = 0, the blocker atoms at the pore mouth are removed to initiate the gas recovery.

To gain generalized insight into the kinetics of gas recovery using our MD simulations, we nondimensionalize the time using a characteristic time $t_c$. Since gas recovery from nanopore is generally considered as a diffusive process, following previous work, $t_c$ is chosen as

$$t_c = \frac{4L^2}{D_m^0/Kn^0}$$

where L is the nanopore length, $D_m^0$ and $Kn^0$ are the reference molecular diffusion coefficient and Knudsen number inside the pore prior to gas recovery. $D_m^0$ and $Kn^0$ depend on the molecular properties of the gas species and the pore size. For nanopores filled with multiple gas species, it is difficult to define $D_m^0$ and $Kn^0$, and thus $t_c$ uniquely. Here, we use the molecular properties of methane (cross-sectional area 0.45 nm²) to calculate the characteristic time $t_c$ since methane dominates the gas storage inside the nanopores. With the temperature and initial pressure inside the nanopores, we obtain a $D_m^0$ of $5.53 \times 10^{-8}$ m²/s. Furthermore, we compute a mean free path corresponding to the initial pressure inside the gas bath ($\lambda_0$) and a mean free path corresponding to the pressure inside the gas bath ($\lambda_f$). $\lambda_0$ is more relevant to the gas transport in the pore interior and at early stage of gas recovery; $\lambda_f$ is more relevant to gas transport near pore mouth. It is found that $\lambda_0 = 0.26$ nm and $\lambda_f = 1.9$ nm. Using the accessible width of the pore as the characteristic length scale, the corresponding Knudsen numbers are found to be $Kn^0 = 0.16$ and $Kn_f = 1.19$ for the 2 nm-wide pore. Note that the accessible width of a pore with a width of 2.0 nm (4.0 nm) measured between the center planes of the atoms in its two walls is 1.62 nm (3.62 nm) because the methane molecule has a diameter of ~0.38 nm.

We first examine the qualitative features of gas recovery process. Figure 5a shows the methane density profile across the 4 nm-wide pore at different x-positions at $t/t_c = 0.01$. We observe that, both near the pore wall and in the central portion of the pore, methane density decreases from their initial values (marked using a dashed line in Figure 5a), which suggests that the adsorbed gas and the free gas are recovered from the pore concurrently. The drop of the gas densities from their initial values becomes less significant as we move from the pore mouth toward the pore interior, which is indicative of the diffusive nature of the gas recovery process. Figure 5b shows the evolution of the cross-section averaged density profiles of methane and ethane along the pore length. After the gas recovery starts, the methane and ethane densities near the pore mouth drops quickly because of the gas exchange between the gas bath (where the gas density is lower) and the gas at the pore mouth. As gas recovery proceeds, gas density in the pore interior drops and a clear diffusion front moving toward the pore interior is observed. The diffusion front reaches the pore’s sealed end at $t/t_c = 0.01$, and the density of both methane and ethane decreases along the entire pore length after that. We note that the gas density at the pore mouth gradually decreases as gas recovery proceeds, which suggests that the gas exchange between the gas bath and pore mouth has a finite resistance. Such a resistance also manifests as concentration polarization in...
the gas bath. Indeed, as shown in Figure 5c, the densities of methane and ethane are nonuniform inside the gas bath, and a diffusion boundary layer in which the gas density varies sharply is visible near the pore mouth. The strength of concentration polarization, as indicated by the drop of gas density across the diffusion boundary layer, is comparable for methane and ethane. Because concentration polarization reduces the driving force for the gas transport from the pore to the gas bath, it slows down gas recovery. The nominal driving force for recovery of methane is larger than that for ethane because the maximal difference of the density of methane inside the pore and the gas bath (~4 nm⁻³ according to Figure Sb and c) is larger than that for ethane (~1 nm⁻³ according to Figure Sb and c). Because the strength of concentration polarization is similar for methane and ethane, concentration polarization likely slows down the recovery of ethane more notably than that of methane.

Next, we investigated the production rate of methane and ethane during the gas recovery process. The production rate of a gas species \( i \) is quantified as its flux out of a nanopore using

\[
Q_i(t) = -\frac{dN_i}{dt}
\]

where \( t = t/t_0 \) is the dimensionless time and \( N_i \) is the number of species \( i \) molecules inside the pore. The fluxes of methane and ethane from the 2 and 4 nm-wide pores are shown in Figure 6a and b, respectively. At earlier time (\( t \approx 0.01 \)), the fluxes of both methane and ethane follow a superdiffusive scaling law \( Q(t) \sim t^{-\alpha} \) with \( 0.4 < \alpha < 0.5 \), which differs from the \( Q(t) \sim t^{-0.5} \) for purely diffusive gas recovery. Similar deviation from the purely diffusive scaling law has been reported for the recovery of pure gas from nanopores, and is caused by the delayed removal of gas molecules adsorbed on the pore walls. At \( t \approx 0.04 \) (\( t \approx 0.01 \)), the production of methane and ethane from the 2 nm-wide (4 nm-wide) pore starts to deviate from the power law scaling behavior. This is consistent with the fact that the diffusion front reaches the pore’s sealed end at these times (see Figure 5b). Figure 6c shows the evolution of the ethane and methane flux ratio \( Q_e/Q_m \) with time. We observe that, during the entire gas recovery operation, \( Q_e/Q_m \) is mostly in the range of 0.19–0.22, which is very close to the initial ethane-to-methane ratio of 0.25 inside the nanopores. This result is somewhat surprising. Specifically, because of the stronger attraction of ethane to the pore walls (and consequently stronger adsorption on the pore walls) than methane (see Figure 3), the mobility of ethane molecules confined inside narrow pores is much smaller than that of the methane molecules. For example, we compute the total diffusion coefficient of methane (ethane) molecules confined inside 2 and 4 nm-wide pores filled with pure methane (ethane) at various pressures (see the Supporting Information) and find that the mean total diffusion coefficient of ethane molecules confined in these pores is 4–5 times smaller than that of the methane molecules at the same pressure (see Figure S5). In addition, as discussed earlier, the concentration polarization near the pore mouth likely slows down the recovery of the ethane more significantly than that of the methane. Therefore, one could have expected \( Q_e/Q_m \) to be much smaller than 0.25 during gas recovery. Below we seek to understand why this is not what we observed in the direct simulation of gas recovery from nanopores.

In principle, the composition of the gas recovered from pores filled with multiple gas species can be understood using several theoretical methods for describing multicomponent fluid transport, e.g., the Onsager formulism based on nonequilibrium thermodynamics and the Maxwell–Stefan formulism. Here we adopt the Onsager formulism, in which the transport of two species \( i \) and \( j \) in a mixture can be described using

\[
J_i = -L_i \nabla \mu_i - L_j \nabla \mu_j
\]

where \( J_i \) and \( \mu_i \) are the flux and chemical potential of species \( i \), respectively. \( L_i \) is the Onsager coefficient, which depends on the concentration and distribution of species \( i \) and \( j \) in the mixture, as well as the interactions between gas species. Rigorous calculations of the gas fluxes from nanopores initially filled with multicomponent gas is complicated by the fact that, because the gas density and composition evolves both temporally and spatially inside the pore, the Onsager coefficients also vary temporally and spatially inside the pore and thus difficult to compute accurately. Here we combine scale analysis and MD simulations to obtain a semiquantitative understanding of the composition of gas fluxes from pores during gas recovery operations.

Since the IAS theory predicts the isotherm of gas storage inside nanopores quite well (see Figure 4), it is reasonable to assume that methane and ethane are mixed ideally inside the pore. We thus approximate the fugacity of gas species \( i \) using its partial pressure and write the chemical potential of species \( i \) as

\[
\mu_i = \mu_{i,0} = k_BT \ln(P_i/P_{i,0})
\]

where \( \mu_{i,0} \) and \( P_{i,0} \) are the chemical potential and partial pressure of species \( i \) at a reference state, respectively. \( k_BT \) is the thermal energy. Further assuming that each gas inside the pore behaves ideally, the gradient of the chemical potential of the species \( i \) can now be written as

\[
\nabla \mu_i = \nabla \left( k_BT \ln(P_i/P_{i,0}) \right)
\]

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the pore mouth can be estimated as $P_{\text{nanopore}} = 1.4$ at pore widths $0.8$ and $0.2$. It is consistent with the facts that there are much more methane molecules in the pore than ethane molecules and the total diffusion coefficient of methane molecules confined in nanopores is larger than that of the ethane molecules (see Table S5). Importantly, we observe that the cross-correlation $L_{11}^0$ is even higher than the $L_{22}^0$ term. This indicates that the interactions between methane and ethane play a critical role in the transport of ethane in the nanopores.

Using the Onsager coefficients shown in Table 2 and eq 11, the ratios of the ethane and methane fluxes ($Q_i/Q_m$) during gas recovery from 2 and 4 nm-wide pores are estimated to be 0.219 and 0.220, respectively. These estimations are in good agreement with the results shown in Figure 6c. The relative large $Q_i/Q_m$ originates from the facts that $L_{22}^0$ is not much smaller than $L_{11}^0$ and $L_{11}^0$ is larger than $L_{22}^0$, both of which reflect the effective coupling between the transport of ethane and methane inside nanopores. In the above analysis, the effect of concentration polarization at the pore mouth is neglected. When such effect is taken into account by using the gas composition measured at pore mouth at representative time (e.g., at $t/t_c = 0.01$, see Figure 5c) to determine $P_{1,m}$ and $P_{2,m}$ in eq 11b, the $Q_i/Q_m$ predicted using eq 11a only decreases slightly (see the Supporting Information).

## IV. CONCLUSIONS

We use MD simulations to investigate the adsorption of binary mixture of methane and ethane in nanopores and the gas recovery from these pores to low pressure gas bath. Calculation of the binary adsorption isotherm of nanopores in equilibrium with bulk methane–ethane mixture shows that while both...
methane and ethane are stored as free gas and adsorbed gas inside the nanopores, the adsorption of ethane on the wall is stronger than methane. The stronger adsorption of ethane on pore walls compared to methane leads to an enrichment of ethane inside nanopores. Such an enrichment is more pronounced in narrow pores but is weakened as the pressure increases due to entropic effects. During gas recovery operations, free and adsorbed gas are extracted concurrently from the nanopores, and noticeable concentration polarization occurs near the pore mouth. Nevertheless, the production rates of both gases approximately follow the square root scaling law before the diffusion front reaches the sealed end of the nanopores. The ratio of the production rate of ethane and methane from the pores is only slightly smaller than their initial mole ratio inside the pores, which is attributed to the effective coupling of the transport of methane and ethane inside nanopores.

Our simulation results suggest that the storage of binary gas in narrow nanopores is affected strongly by the adsorption of different gas molecules on the wall, which depends on both the pore pressure and the nature of gas molecules. Since molecular simulations are computationally too costly for determining such adsorption in practical applications, molecular theories are needed. In this regard, we show that the classical IAS theory can predict the essential trends of binary gas adsorption on the pore walls, although it tends to overestimate the gas adsorption. Our simulations revealed that the recovery of binary gas mixture from single, narrow nanopores to gas baths is approximately a diffusive process and the coupling between the transport of gas species inside the nanopores plays an essential role in determining the composition of the recovered gas. These results lend support to the existing theories for gas recovery from nanopores and point to the need to develop effective models for predicting the coupling of the transport of different gas species confined inside nanopores.

Molecular simulations of gas recovery are limited to systems with pores many orders of magnitude shorter than in real shale formations, and this necessarily introduces some undesirable features such as an extremely large pressure gradient along pore length and from the pore opening to the gas bath during gas recovery. Nevertheless, the fact that the present and our earlier MD simulations capture the scaling law of gas recovery rate reported in field studies and continuum simulations suggests that these undesired features do not introduce significant artifacts into the simulation results. Therefore, MD simulations not only can be used to understand the transport and adsorption properties of gases in nanopores as demonstrated extensively in the past years, but can also be used as a powerful tool for exploring the essential physics of gas recovery process. In this study, we focus on the recovery of CH$_4$−C$_2$H$_6$ mixture from nanopores. The same method can be used for understanding the recovery of other gas mixtures. For example, since some shale formations contain a greater share of CO$_2$ than C$_2$H$_6$, it would be interesting to study the recovery of CH$_4$−CO$_2$ mixtures. The same method can also be used for studying the enhanced recovery of methane by injection of CO$_2$ in shale formations.

### ASSOCIATED CONTENT

1. Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.energyfuels.7b01013.

Isotherm of pure ethane in 2 nm-wide pores, isotherms of surface adsorbed pure methane and pure ethane in 4 nm-wide pores, isotherms of the surface adsorbed methane and ethane in 4 nm-wide pores, density profiles of methane and ethane molecules inside nanopores in equilibrium with baths with different pressures, the total diffusion coefficient of methane and ethane in 2 and 4 nm-wide nanopores, and the calculation of the ethane-to-methane flux ratio during gas recovery when concentration polarization is taken into account (PDF)

### AUTHOR INFORMATION

**Corresponding Author**

*E-mail: ruiqiao@vt.edu.*

**ORCID**

Rui Qiao: 0000-0001-5219-5530

**Notes**

The authors declare no competing financial interest.

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