

Evaluating Mixture Adsorption Models Using Molecular Simulation

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The design of adsorption-based separation processes using novel adsorbents requires reliable data for the adsorption of fluid mixtures on candidate adsorbents. Due to the difficulty of generating sufficient data across possible operating conditions, process designs generally rely on interpolation of pure-component data using a model, most commonly ideal adsorbed solution theory (IAST), and related theories. There are many cases where IAST fails to provide an adequate description of mixture adsorption, usually due to the fact that practical adsorbents do not have uniform surfaces. We have evaluated the use of a segregated version of IAST, where competition is assumed to occur at isolated adsorption sites. This simple modification can provide the correct description of adsorption across a large range of pressures using ideal isotherm models. We also demonstrate the importance of identifying multiple sites even for weakly adsorbing components to provide the correct behavior at high pressure. © 2013 American Institute of Chemical Engineers AIChE J, 59: 3054–3064, 2013

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Introduction

Separating fluid mixtures via adsorption onto porous materials is a common process technology used in a wide variety of industrial contexts, including the separation of air, scrubbing solvent vapors from exhaust, and separation of xylene isomers.¹ The design of adsorption-based processes requires reliable thermodynamic data for the fluid-adsorbent system over a range of potential process conditions. Collecting these data for the adsorption of mixtures of gases are particularly challenging due to the complexity of the experiments and interpretation of the resulting data. This difficulty is mitigated in practice by the use of a model, such as ideal adsorbed solution theory (IAST), to interpolate mixture data from pure-component data, which is simpler to collect.² Although IAST has been shown to provide good predictions for a wide variety of fluid-adsorbent systems, there are numerous cases where it has been shown that the predictions are inaccurate.^{3–10} The IAST model assumes ideal behavior of the adsorbed phase and that all components have access to the same uniform surface. This second assumption is

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frequently violated in practical adsorbents, and variety of approaches have been proposed to correct for the inaccuracies it introduces, which will be discussed in more detail later.^{3,6–8,10–18} Recent efforts in synthesizing and predicting new materials have led to a plethora of potential new adsorbents, including metal–organic frameworks (MOFs), zeolitic imidazolate frameworks (ZIFs), and novel zeolites. These materials are very promising for several important gas separation applications.¹⁹ It is possible to introduce specific sites in these materials that selectively adsorb one of the components in preference to the others giving them a heterogeneous pore space by definition. From a practical point of view, it is important to reliably predict the mixture adsorption isotherms. This requires a better understanding how the structure of these materials influences the selection of the correct model for the adsorption of mixtures.

The large number of new porous materials that have been demonstrated in recent years introduces the practical problem of evaluating such a large number of materials for a given separation. For example, Deem and coworkers²⁰ have developed a database of millions of hypothetical zeolite structures. Efforts by many researchers have revealed a wide array of MOFs with diverse structures and compositions.²¹ The nature of MOF chemistry suggests that there are potentially millions of materials when taking into account all possible combinations of metal nodes and organic linker molecules.^{19,22–24} When organized into databases, there is

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the possibility of screening very large sets of materials simultaneously to find those best suited for a particular application.^{18,21,25}

An active area of research is the development of highthroughput screening techniques to identify the most promising materials for desired gas separations. In the absence of sufficient experimental data, direct computer simulations in the grand canonical ensemble²⁶ can provide the data over a wide range of conditions for microporous materials like zeolites,²⁷ MOFs,²⁸ and ZIFs.²⁹ Both the experimental and the computational approaches focus on obtaining the pure-component isotherms and rely on IAST to accurately predict the mixture isotherms.

In this work, we develop a systematic approach to incorporate these chemical inhomogeneities in the IAST. This approach is based on the simple notion that one should divide the pore space into different regions and apply IAST on each region separately. Interesting, variations of this idea have appeared several times in the literature but, as we will demonstrate in this work, it can only be used successfully with a molecular understanding of the adsorption behavior of the pure-component isotherms. We illustrate with several practical examples that incomplete information about the pure-component isotherms, namely data at insufficiently high pressures where weakly adsorbing components saturate adsorption sites, can explain the problems associated with typical application of IAST.

At this point, we emphasize that we rely on molecular simulations to provide us with accurate data on both the pure components and the mixture isotherms. From a purely conceptional point of view, there is no fundamental difference for IAST, in predicting mixture isotherms, whether these pure-component data are obtained from molecular simulations or experiments. Of course, only if our simulations are representative for the experimental system, our conclusions also hold for the experimental conditions.

One of the main conclusions of this work is that a reliable prediction of the mixture isotherm requires knowledge on the number of adsorption site and which component preferentially adsorbs on which site. For many systems, this can be reliably obtained from the pure-component isotherms. However, in some, and usually the most interesting, cases this molecular information is lacking. We also demonstrate how molecular simulation can be used to address these questions. Given that the assumption of a uniform surface is often violated in practice, simulations provide a molecule-scale picture that can be used to suggest the siting of molecules in the adsorbent's pores. These strong sites have important implications for the best model to interpolate the data. Simulations can also predict adsorption at very high pressures that may be infeasible in practice. Accurate information about the saturation loading of weakly adsorbed components may only be available at these high pressures and can help correctly parameterize the pure-component isotherms.

Adsorbed Solution Theories

IAST

Due to the importance of making reliable predictions of adsorption isotherms for fluid mixtures, many models have been proposed that attempt to predict the adsorbed phase composition and loading based on pure-component data and, in some models, properties of the fluid mixture. Most practical models rely on some variant of the IAST of Myers and Prausnitz.² This approach is popular due to the fact that it relies solely on the isotherms of the pure components in the

fluid mixture and is amenable to rapid solution using numerical techniques. IAST has been used extensively to describe adsorption from multicomponent mixtures in zeolites and other adsorbents, including mixtures of alkanes, CO₂, CH₄, and N_2 .^{30–33} Extensions to IAST include the real adsorbed solution theory³ and the non-ideal AST,¹¹ which attempts to account for nonideal behavior of the adsorbed phase by correlating activity coefficients from binary adsorption data or assumptions about the distribution of sites, respectively. Other approaches have attempted to deal with problems surrounding the assumption of a perfectly uniform surface. One key modification these approaches make to IAST is that the surface is assumed to be composed of independent adsorption sites, and any mixture model should be applied separately for each site. For the sake of clarity, this concept will be called segregated IAST (SIAST) when applied in the context of this work, and the terminology of other authors will be adopted when addressing their work.

The concept of modeling adsorption as independent processes at distinct sites or volumes has been applied many times before to improve mixture adsorption predictions relative to IAST. Valenzuela et al.¹² proposed what they termed heterogeneous IAST where each component is assumed to have a distribution of adsorption sites characterized by an adsorption energy. IAST is applied to each site separately based on a fitted energy difference between sites. Moon and Tien⁶ took the analysis a step further and proposed that matching the sites of different adsorbates based on a site-matching criteria that accounts for correlation of the adsorption of different components on the same site. Eiden and Schlünder^{7,8} applied a similar approach to the analysis of dichloromethane and benzene adsorption on activated carbons. Calling the approach multiphase IAST, they assumed a pore filling mechanism and applied IAST to separate sites based on a difference in saturation volumes between components. Cerofolini and Rudznski¹³ categorized these approaches as patchwise theories, as they treat the surface of the adsorbent material as composed of a set of discrete areas or volumes where a model was applied to each individual patch, but there was no interaction between different patches. One drawback of these approaches from a practical standpoint is that they require additional assumptions about the system, usually in the form of the distribution site energies. These distributions must also be parameterized, which can be difficult given the nature of the inverse problem.¹⁴ Recent work by Ritter et al.¹⁵ attempted to avoid this issue using a sum of competitive Langmuir sites approach, which they term a dual-process Langmuir (DPL) isotherm, using the parameters from fitting a dual-site Langmuir isotherm to the pure-component data. Their approach also permits accounting for nonideality in the system by applying the sitematching approach of Moon and Tien by simply swapping the parameters of a particular component between terms of the isotherm. Unfortunately, there is no clear method for deciding when this switch is appropriate other than comparing with experimental mixture data. A disadvantage of these approaches is that they potentially require a large quantity of detailed adsorption data at a variety of conditions that may not be easy or cheap to generate and often are specific to the material being studied.

In more recent work, several authors have attempted to use molecular simulations to evaluate the usefulness of IAST and similar approaches for predicting mixture properties in zeolites and MOFs. Cessford et al.¹⁶ have provided a comprehensive overview of the topic in their recent article, and

we will only address a few highlights here. Krishna and coworkers³⁴ have evaluated the use of IAST to predict mixtures of alkanes in the zeolite MFI and CO₂:CH₄ mixtures in cage-type zeolites.¹⁰ In this latter article, the authors observed that IAST failed to give a good description of the mixture at higher pressures and ascribed this to a segregation of the two components due to preferential adsorption at either the center of the cages or the windows between cages. Jee and Sholl¹⁷ suggested that using a segregated model for CO₂:CH₄ adsorption in zeolite DDR improves predicted isotherms relative to IAST, but is still inaccurate at high total fugacity. Van Heest et al.¹⁸ performed an extensive screening of MOFs for separating noble gas mixtures and noted that accounting for regions that would be inaccessible to one component of the mixture was important for correctly predicting adsorption selectivity. A key aspect of these studies is that they were able to correlate specific structural features of the adsorbents to the inaccuracy of IAST.

SIAST

While the IAST model does not rely on a specific description of the molecular system, it is usually interpreted in terms of a hypothetical uniform surface for the adsorption of the components of the mixture. Due to the atomic scale heterogeneity of nanoporous materials, it is useful to view the materials as having multiple "surfaces" where molecules can adsorb. Each of these surfaces can have a separate equilibrium, which we assume can be described with an ideal model. Strictly speaking, for microporous adsorbents like zeolites, the concept of spreading pressure is not applicable, because there is no well-defined surface.^{11,35} However, the idea of accessible surface area or pore volume is still a useful concept for diagnosing failures of IAST. If the components of the mixture can access different parts of the material, one component may be able to diffuse into cavities not accessible to other components and is, thus, no longer competing with the other components for the same adsorption sites. In recent work on the screening of zeolites for the separation of CO₂ and N₂, we discovered a class of materials that had particularly strong adsorption sites for CO₂.²⁵ These were often small pockets that could accommodate a single CO₂ molecule, and the arrangement of the framework atoms around the pocket provides a site with a particularly favorable interaction energy for the molecule. Although N₂ is not formally excluded from these pockets, they are more likely to be occupied by CO₂ when the two gases are adsorbing competitively. Materials of this sort typically exhibit an isotherm with a plateau or step at intermediate loading that can be accurately correlated with the dual-site Langmuir isotherm equation. The loading at the plateau corresponds to the density of strong adsorption sites in the unit cell.

To more accurately describe the total loading of a system that may have spatially distinct adsorption sites, we propose that applying IAST to distinct sites can provide more accurate predictions in many cases. Instead of considering the available pore volume to be one continuous space, the volume may be subdivided into natural regions where separate competition processes take place. For materials where it is possible to identify segregated sites, it is possible to model the mixture adsorption as the sum of separate competitions, which in turn may be modeled using IAST. Figure 1 shows how this might be extended to an arbitrary number of adsorbed phases. Equation 1 relates the isotherm of component i at site j to the surface potential.

$$\psi_{ij} = \int_0^{f_{ij}} n_{ij}(f) dlnf \tag{1}$$

In Eq. 1, the surface potential for adsorption of component i at site j is related to an adsorption isotherm representing only site j. In this work, we will assume that each site can be described by a single-site Langmuir isotherm

$$n_{ij}(f) = \frac{m_{ij}b_{ij}f}{1 + b_{ij}f} \tag{2}$$

giving the following expression for the surface potential

$$\psi_{ij}\left(f_{ij}^{\circ}\right) = m_{ij}\log\left(1 + b_{ij}f_{ij}^{\circ}\right) \tag{3}$$

where m_{ij} and b_{ij} are the saturation loading and Langmuir constant for adsorption, respectively, of component *i* at site *j*. The condition for equilibrium may be expressed as Equation 4 for competitive adsorption between *N* components at site *j*.

$$\psi_{1j} = \psi_{2j} = \dots = \psi_{Nj} \tag{4}$$

As we treat the adsorbed phase as composed of independent phases, each is related to the gas phase independently.

$$fy_i = x_{ij} f_{ij}^{\circ} \left(\psi_{ij} \right) \tag{5}$$

There is an independent mass balance on each phase.

$$\sum_{i=1}^{N} x_{ij} = 1$$
 (6)

The total loading in phase *j* is given by

$$n_{ij} = \left[\sum_{i=1}^{N} \frac{x_{ij}}{n_{ij} \left(P_{ij}^{\circ} \right)} \right]^{-1}$$
(7)

and loading of component i in phase j is given by

$$n_{ij} = x_{ij} n_{tj} \tag{8}$$

The total loading of component i is simply the sum of the loadings in the individual phases.

$$n_i = \sum_{j=1}^M n_{ij} \tag{9}$$

While this method is general, experimental isotherms may not cover a large enough range of pressures to accurately determine the saturation loadings for all components at each site. We are also neglecting any sort of accounting for the nonideality of the absorbed mixture that may exist at high density. Except at the highest densities, the interactions felt by molecules in nanoporous materials are dominated by the atoms of the material framework any contribution to nonideality from the components themselves is likely small in comparison.

To compare traditional IAST with the current approach, the dual-site Langmuir isotherm model was fitted to the purecomponent isotherms using nonlinear least-squares regression.

$$n_i(f) = \frac{m_{i1}b_{i1}f}{1 + b_{i1}f} + \frac{m_{i2}b_{i2}f}{1 + b_{i2}f}$$
(10)

For materials with strongly segregated sites for adsorption, the saturation at the first site was fixed to the same value for

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Figure 1. Schematic diagram of the phases in the SIAST model.

A gas phase (white) is in equilibrium with M independent adsorbed phases (shades of gray). The whole system is at a constant temperature. The gas phase is characterized by a total pressure P and its composition, usually expressed as the mole fraction of all components y_i . Each adsorbed j phase is characterized by the loading of each component i, n_{ij} in equilibrium with the gas phase. Each adsorbed phase is in equilibrium separately with the gas phase.

both CO_2 and N_2 , reflecting the fact that only one molecule of any component can occupy one of these sites at a given time.

Methods

To investigate different models of adsorption, grand canonical Monte Carlo (GCMC) simulations were used to generate pure-component and mixture isotherms for mixtures of CO_2 and N_2 in different zeolites and CO_2 and C₃H₈ in the zeolite MOR. These were treated as the reference data that allow us to compare the predictions of different models. Simulations in the grand canonical ensemble involve specifying the temperature and volume of an adsorbent system and the chemical potentials of all the adsorbing components.²⁶ The simulation box was defined by a tiling of one or more crystallographic unit cells of a given material. All framework atoms were held fixed. Trial configurations were generated by attempting to move, rotate, reinsert, or remove an existing molecule or adding a new molecule to the simulation box. The energy of all interactions was described using the 12-6 LennardJones potential plus an electrostatic term. The Ewald summation technique was used for the electrostatic interactions. All short-range interactions were cut off and shifted to zero at 12.0 Å. Interaction parameters and partial atomic charges were taken from different sources, depending on the system under study. CO_2 and N_2 adsorption in siliceous zeolites were modeled using the parameters of García-Pérez et al.³⁶ C₃H₈ adsorption in MOR was parameterized using the force field of Dubbeldam et al.³⁷ For simulations in aluminosilicate zeolites, CO_2 and sodium interaction parameters were taken from García-Sánchez et al.³⁸ N₂ interaction parameters in these materials were developed as a part of this work and are reported in the Supporting Information.

Results and Discussion

We have broken down the analysis into a three separate cases based on the degree of segregation between adsorption sites and the relative preference of each adsorbing component for each of the adsorption sites. In one extreme case of segregation, the adsorption sites may be spatially separated

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Figure 2. Comparison of isotherms computed via GCMC and different mixture adsorption models in pure silica PCOD8286959 at 300 K.



from one another and diffusion between sites requiring crossing a barrier. Each of these sites can represent a different free energy for each component, and components will occupy the sites in the order that represents the lowest free energy for the component. In this work, we have considered materials with two distinct adsorption sites, and we distinguish whether a given mixture exhibits a normal site preference, that is, components occupy sites in the same order, or reverse site preference, where one component occupies sites in the reverse order that the other component does. This parallels the site correlation concept used by Moon and Tien⁶ and Ritter et al.¹⁵ We also consider the case where sites are not isolated from one another and evaluate the use of competitive isotherm models.

Materials with isolated sites, normal site preference

GCMC simulations were performed for pure CO2 and pure N₂ in hypothetical zeolites PCOD8286959 and PCOD8200029 from Deem's database of hypothetical materials²⁰ at 300 K. These pure-component isotherms are shown in Figures 2A and 4A, respectively. The isotherms were simulated from very low fugacity (on the order of 10 Pa) up to 10^9 Pa fugacity. The very high fugacities are required to ensure that the plateaus in the isotherm are accurately represented. In the case of the more weakly adsorbing N_2 , fugacities above 107 and 106 Pa for PCOD8286959 and PCOD8200029, respectively, were required for the isotherm to approach a saturation loading. All pure-component isotherms were fitted using the dual-site Langmuir isotherm (Eq. 10) with the constraint that the saturation loading of the first adsorption site (m_{i1}) was fixed to be identical for both CO2 and N2 (fitted parameters can be found in the Supporting Information). Previous work²⁵ has indicated that adsorption in these materials occurs first at small localized pockets in the material that can accommodate only a single molecule. Figures 3 and 5 show accumulated snapshots of CO₂ and N₂ locations in PCOD8286959 and PCOD8200029, respectively, taken from the GCMC simulations of the mixture at 10⁹ Pa. In PCOD8286959 (Figure 3), large main channels (purple circle) run parallel to the c crystal axis, and small pockets (orange circle) connect between channels in the *b* direction. The density of these sites is set by the crystal structure of the material and hence should be identical for both components. Although N_2 does not exhibit a pronounced step in the isotherm, because it does not have as strong a preference for this site as CO₂ does, this constraint is required to ensure the correct description of the total loading of each site in the mixture model. In PCOD8286959 (Figure 3), the molecules can adsorb in two separate pore systems, one running parallel the *c* crystal axis (orange circle and arrow) and the other running parallel to the *b* axis (purple circle and arrow). The sizes of these two channels are slightly different, with CO₂ preferring to adsorb in the channels parallel to the *b* axis, as shown by the predominance of blue points in Figure 3.

Figures 2B and 4B show adsorption isotherms for a 14% CO₂, 86% N₂ mixture at 300 K predicted using GCMC, the SIAST model, and the original IAST model in PCOD8286959 and PCOD8200029, respectively. The original IAST data were generated by integrating Eq. 1 directly with Eq. 10 for the two components. For both materials, the models both agree with the GCMC data at very low fugacity. At these low fugacities, adsorption is occurring in the linear or Henry's law regime of the isotherm. Each component is in effect adsorbing independently and is completely described by its Henry coefficient. Both the segregated and traditional IAST model agree up to fugacities commensurate with the first saturation loading in the pure CO_2 isotherm. Above these fugacities, the N₂ loading predicted by traditional IAST is lower than the GCMC results. In the case of PCOD8286959 (Figure 2B), the IAST-predicted N₂ loading matches the GCMC results at fugacities above 10^7 Pa, whereas in PCOD8200029, the IAST-predicted N2 loading is consistently 10 times lower than the GCMC isotherm. For both of these materials, the SIAST model provides excellent agreement with the GCMC results at all fugacities.

The effect of using the traditional IAST model is that the step in the CO_2 isotherm induces an effective step in the N_2 isotherm when using IAST, because it imposes a competitive adsorption process between CO_2 adsorbing at the stronger adsorption sites and N_2 adsorbing at the weaker adsorption sites. The adsorption of N_2 at the weaker sites in the

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Figure 3. Snapshots of CO₂ (blue) and N₂ (red) center-of-mass configurations in zeolite PCOD8286959 during simulation at 10₉ Pa fugacity and 300 K.

Framework silicon and oxygen atoms are drawn in dark gray, and the centers-of-mass were collected from 10,000 samples during the GCMC simulation. The purple and orange circles indicate the main channels and the small pockets of the material, respectively. Due to the affinity of CO_2 for the small pocket, N_2 is effectively excluded and only adsorbs in the main channel. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

framework can occur simultaneously and independently of adsorption at the stronger sites due to the fact that there is likely a barrier between adsorption at these distinct sites. The fact that CO_2 is more likely to adsorb at the stronger sites first frees up the rest of the volume of the material where it would otherwise adsorb to provide adsorption sites for N₂. We also note that the relative saturation loadings of the two adsorption sites do not appear to matter. If we compare the fitted isotherm parameters the case of PCOD8286959 (Figure 2), the first saturation loading is 0.4229 mol kg⁻¹, compared to 8.811 mol kg⁻¹ for the second saturation loading of CO2. The same values for the two sites in PCOD8200029 (Figure 4) are 1.510 mol kg^{-1} and 0.7354 mol kg⁻¹, respectively. The first site accounts for 4.6% of the total saturation in PCOD8286959 but over 67% of the total saturation loading in PCOD8200029. Although one site in the material may provide the majority of all adsorption sites, it is important to partition them correctly, especially to predict mixture adsorption at high fugacity where both sites will likely be occupied to some extent.

Materials with isolated sites, reverse site preference

One set of data that has been used to evaluate different mixture adsorption models was the adsorption of CO2:C3H8 on H-MOR. Valenzuela et al., Moon and Tien, and Ritter et al. applied their approaches and showed that different segregated models improved predictions for the system relative to IAST.^{6,12,15} We computed isotherms for pure CO₂, pure C₃H₈ (Figure 6A) and a 50:50 CO₂:C₃H₈ mixture (Figure 6B) adsorbing in the pure-silica form of MOR at 300 K. The MOR topology has well-known side pockets along the main pores of the material, and this is reflected by the step in the isotherms for both components. Figure 7 shows center-ofmass configurations of CO_2 and C_3H_8 recorded during GCMC simulations at low (10³ Pa) and high (10⁹ Pa) fugacity of both pure components and the mixture described earlier. The purple circle indicates the main channels of the material, running parallel to the c crystal axis and the orange circles indicate the side pockets along these channels. As can be seen in the pure CO₂ snapshots, CO₂ is most likely to



Figure 4. Comparison of isotherms computed via GCMC and different mixture adsorption models in pure silica PCOD8200029 at 300 K.



be found in the side pockets at low fugacity, then as those are saturated, stars to fill the main channels. The reverse is true for C_3H_8 . We note that the second saturation for C_3H_8 does not appear until the fugacity goes above 10^8 Pa, whereas CO_2 begins to saturate its second site below 10^6 Pa fugacity. As these components have opposite preferences, at low fugacity, C_3H_8 occupies the main channels whereas CO_2 resides in the side pockets. At high fugacity, however, CO_2 has saturated the side pockets and competes for the main channels with C_3H_8 , which can be seen by the relatively few red points in the mixture snapshots for this condition.

Figure 6B compares the predictions of IAST, SIAST, and the DPL model with the results of the GCMC simulations. The dual-site Langmuir parameters for both pure components were fit without restriction. Although we did not constrain them as in the previous case, we observed that the first saturation loading for CO_2 (4.165 mol kg⁻¹) was comparable to the second saturation loading for C_3H_8 (4.993 mol kg⁻¹). As shown by Ritter et al., reversing the sense of the two sites in the isotherms of CO₂ and C₃H₈ provided the best predictions of experimental mixture compositions on H-HOR.¹⁵ In our application of the SIAST and DPL models, we also made this switch, so that the first site in the CO₂ isotherm competes with the second site in the C₃H₈ isotherm and vice versa. Traditional IAST does not provide for this reversal, as the entire isotherm is integrated simultaneously and provides the poorest description of the data. SIAST and the DPL model provide good agreement with the GCMC results at low fugacity. At high fugacity, the DPL model predicts a constant composition after the fugacity increases beyond 10^6 Pa. This is expected for competitive isotherm models, since the limit for this form of isotherm as fugacity or pressure approaches infinity is simply the saturation



Figure 5. Snapshots of CO₂ (blue) and N₂ (red) center-of-mass configurations in zeolite PCOD8200029 during simulation at 10₉ fugacity and 300 K.

Framework silicon and oxygen atoms are drawn in dark gray, and the centers-of-mass were collected from 10,000 samples during the GCMC simulation. The pores of the material consist of two orthogonal pore systems, indicated by the purple and orange arrows and circles. (A) and (B) show the views down the *c* and *b* crystallographic axes, respectively. The pores are small enough in the channel indicated by orange that CO_2 prefers to site there and tends to exclude N₂. Both components are accommodated in the slightly larger channels indicated by purple. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]



Figure 6. Comparison of isotherms computed via GCMC and different mixture adsorption models in pure silica MOR at 300 K.

(A) Pure-component isotherms of CO₂ (blue) and C_3H_8 (red) with lines indicating the fit of the dual-site Langmuir model (Eq. 10). (B) Mixture adsorption isotherms for a 50% CO₂ (blue), 50% C_3H_8 mixture. Symbols indicate GCMC results, solid lines represent the SIAST model, dashed lines represents the traditional IAST model, and densely dotted lines are predictions of the DPL model of Ritter et al.¹⁵ [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

loading of the component. The GCMC simulations predict that the loading of C_3H_8 will eventually decrease. The replacement of C_3H_8 by CO_2 in the pores should be expected at very high fugacity due to the entropic effect of smaller molecule CO_2 filling the available volume more effectively than C_3H_8 . The SIAST model provides a good fit to the GCMC results for CO_2 over the entire fugacity range, and C_3H_8 is well described up to about 10^7 Pa total fugacity. Above this fugacity, the predicted C_3H_8 loading decreases more rapidly than observed via GCMC. This may be due to other features in the isotherm that would be observed by simulating the pure components at higher fugacity then we did and would require an additional site in the isotherm to describe.

The SIAST and DPL model required that the second site in the C_3H_8 isotherm was matched with the first site of the CO_2 isotherm to provide accurate predictions. This is due to the fact that CO_2 readily enters the side pockets along the



Figure 7. Snapshots of CO_2 (blue) and C_3H_8 (red) center-of-mass configurations in zeolite MOR during simulation at low (10³ Pa) and high (10⁹ Pa) fugacity and 300 K.

Framework silicon and oxygen atoms are indicated in gray, and all views look parallel to the *c* crystal axis. The centers-of-mass were collected from 10,000 samples during the GCMC simulations. The purple and orange circles in the top left figure show the main channels and side pockets of the material, respectively. While both components can occupy either site, CO_2 tends to adsorb in the pockets first, while C_3H_8 only enters the pockets at high fugacity. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

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Figure 8. Comparison of isotherms computed via GCMC and different mixture adsorption models in aluminosilicate NaFAU (13X) at 300 K.

(A) Pure-component isotherms of CO₂ (blue) and N₂ (red) with lines indicating the fit of the dual-site Langmuir model (Eq. 10). (B) Mixture adsorption isotherms for a 14% CO₂ (blue), 86% N₂ mixture. Symbols indicate GCMC results, solid lines represent the SIAST model, and the dashed line represents the traditional IAST model. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

channels of MOR, whereas C3H8 first adsorbs in the main channels and enters the side pockets at high pressure. This behavior can readily be seen in snapshots of the molecular configurations recorded during the GCMC simulations. As only one molecule can occupy these side pockets at a time, it is appropriate that the loading of the first site in the CO_2 isotherm and second site in the C3H8 isotherm are comparable. The nonideal nature of this system arose solely from this difference in siting behavior and is not a necessarily the result of nonideality in the CO₂:C₃H₈ fluid mixture. Unfortunately, the matching of the correct spatial placement of the sites is not easy to ascertain from simply fitting the purecomponent isotherms. The complementary site-matching or perfect negative correlation described by Moon and Tien and Ritter et al., respectively, accounts for this ordering, but requires either additional parameterization or evaluation against experimental data to work. Information about the siting of molecules is a direct result of GCMC simulations and can provide a direct method for determining the correct pairing of sites when applying SIAST with no additional assumptions.

Materials where sites are not isolated

Another important class of materials for gas separations are aluminosilicate zeolites. The presence of aluminum atoms in the framework creates a charge imbalance that is compensated by the presence of cations in the pores of the material. These cations can be exchanged by washing the materials with solutions of metal salts, and a mixture of different cations can be used to modify the adsorption properties of the materials for a specific purpose. Recent studies on the common type X and type A zeolites suggest that they may be selective adsorbents for separating mixtures of CO₂ and N₂.³⁹⁻⁴¹ They typically exhibit a very large selectivity for CO2 over N2 due to the stronger interactions with the larger quadrupole moment of CO₂. We simulated adsorption of CO2, N2, and a 15%:85% CO2:N2 mixture in zeolite 13X (NaFAU). This material has a Si:Al ratio of approximately 1.24:1. Adsorption of these gases occurs in the supercages of the material, which are connected by 7.4 Å windows. The relatively large distances in these materials suggest that there are not volumes where an adsorbed molecule is segregated from the other pores space and can exclude other molecules. Figure 8A shows the adsorption isotherms for the pure gases at 313 K. Details of the simulation are given in the Supporting Information.

The CO_2 isotherm exhibits two-site behavior that can be well correlated using the DSL isotherm. Due to the weak adsorption of N₂ at ambient conditions, it is difficult to estimate the saturation loading from typically experimental isotherms. As seen in Figure 8A, it would take on the order of 10^8 Pa fugacity to observe saturation loading of N₂. In the absence of the required saturation loading, it is often assumed that the saturation loading for one component eventually takes the same value as the other component and a single-site isotherm. This assumption, in and of itself, is not a bad approximation for this system. At the highest fugacity simulated (10^9 Pa) , the loading of CO₂ and N₂ agree well (Figure 8A). However, it is possible to fit a dual-site model to the N2 data with a high level of confidence. Figure 8B compares the application of traditional IAST using a dualsite model for CO₂ and either a single-site or dual-site model for N₂. We have also included the dual-process model of Ritter et al. to demonstrate the utility of using competitive models for describing this system.

Figure 8B shows that the application of IAST with a dualsite model for both components improves the prediction of mixture isotherms for N₂ in NaX compared to using a single-site isotherm for N₂. The CO₂ isotherm is not strongly influenced by the choice of isotherm model for N₂, reflecting the fact that the stronger adsorption of CO₂, reflected by its larger adsorption constants, means that it will always be the preferred molecule to adsorb at a given site. Although it is the more likely component to adsorb at either site, CO₂ will tend occupy the strongest site first, as this site represents the lowest free energy for the molecule. It is important that the isotherm for N₂ also reflects this choice of adsorption site, even if the preference between the sites is not as strong as

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the other component. When CO₂ adsorbs at its lowest free energy site, it will tend to compete with N2 to a lesser extent at the higher free energy site, and consequently the N₂ adsorbs to a greater extent than would otherwise be expected. If a single-site isotherm is used to describe N₂ adsorption, the predicted mixture contain less nitrogen, because CO₂ will always be preferred at a given adsorption site and N₂ does not have the option at adsorbing at another site that represents a relatively high free energy for CO_2 . In contrast to the materials considered in the sections earlier, there is not a spatial segregation of adsorption sites, and both the traditional IAST and the DPL model provide a similar description of adsorption in these materials. Correct modeling of the mixture adsorption requires that each pure component is represented by a model that represents all possible sites where it may adsorb.

Conclusions

We have evaluated the prediction of adsorption isotherms for mixtures of gases in nanoporous materials using IAST on segregated adsorbed phases. This approach can provide a better description of adsorption in these materials, especially when there are distinct adsorption sites with different saturation behaviors. It is not necessary to directly treat the potential nonideality of the adsorbed mixture, because most of the nonideality is due to the differing preferences for the mixture components for different adsorption sites. We think simplicity of this model makes it sufficiently cheap to calculate for use in the screening of databases of materials for separations or for use in other computational modeling that would benefit from a computationally cheap description of mixture adsorption. High-pressure isotherm data are important to accurately fit saturation loadings. Any model of a pure component used to predict mixture adsorption should describe any multiple-site behavior, even if the preference for different sites is weak. This is especially important when interpreting isotherms where good estimates of the saturation loadings of all components are not available.

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