Efficient Application of Continuous Fractional Component Monte Carlo in the Reaction Ensemble

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ABSTRACT: A new formulation of the Reaction Ensemble Monte Carlo technique (RxMC) combined with the Continuous Fractional Component Monte Carlo method is presented. This method is denoted by serial Rx/CFC. The key ingredient is that fractional molecules of either reactants or reaction products are present and that chemical reactions always involve fractional molecules. Serial Rx/CFC has the following advantages compared to other approaches: (1) One directly obtains chemical potentials of all reactants and reaction products. Obtained chemical potentials can be used directly as an independent check to ensure that chemical equilibrium is achieved. (2) Independent biasing is applied to the fractional molecules of reactants and reaction products. Therefore, the efficiency of the algorithm is significantly increased, compared to the other approaches. (3) Changes in the maximum scaling parameter of intermolecular interactions can be chosen differently for reactants and reaction products. (4) The number of fractional molecules is reduced. As a proof of principle, our method is tested for Lennard-Jones systems at various pressures and for various chemical reactions. Excellent agreement was found both for average densities and equilibrium mixture compositions computed using serial Rx/CFC, RxMC/CFCMC previously introduced by Rosch and Maginn (Journal of Chemical Theory and Computation, 2011, 7, 269−279), and the conventional RxMC approach. The serial Rx/CFC approach is also tested for the reaction of ammonia synthesis at various temperatures and pressures. Excellent agreement was found between results obtained from serial Rx/CFC, experimental results from literature, and thermodynamic modeling using the Peng−Robinson equation of state. The efficiency of reaction trial moves is improved by a factor of 2 to 3 (depending on the system) compared to the RxMC/CFCMC formulation by Rosch and Maginn.

1. INTRODUCTION

Substantial efforts have been made by scientists and engineers to study chemical reactions in nonideal environments.1−3 An optimal design and operation of many chemical processes relies on accurate information regarding reaction equilibria.4 Reaction equilibria vary as the operating conditions of a reactor change. As a result, an approach is required which can provide information regarding chemical equilibria for a wide range of operating conditions. In an ideal gas, chemical equilibria are determined by the difference between the standard Gibbs free energies of formation of reactants and reaction products.5 Due to interactions of the reacting molecules with surrounding molecules, the chemical equilibrium may significantly differ from the ideal gas situation as the medium formed by the surrounding molecules may stabilize reactant and reaction product molecules differently. It is not always possible to measure reaction equilibria experimentally. The main reasons for this are as follows: (1) Extreme conditions may not be accessible experimentally. (2) Kinetic limitation may prohibit reaching chemical equilibrium on accessible time scales. (3) Large-scale experimental screening of solvents for chemical reactions may not be feasible. Therefore, there is a demand for theoretical methods that can accurately predict reaction equilibria.
Molecular simulation is a natural tool for this as interactions between atoms and molecules are explicitly taken into account. One can perform Molecular Dynamics (MD) with a force field that can handle chemical reactions, e.g., DFT-based, Car−Parrinello, or ReaxFF based MD. The main disadvantage of these approaches is that reactions may not occur within the limited time scale of MD simulations. Therefore, advanced simulation techniques such as metadynamics or transition path sampling may be required. These types of simulation techniques are not considered further in this paper. One of the most commonly used approaches in molecular simulation is to simulate the reaction equilibria in the reaction ensemble (RxMC). In this approach, the chemical reaction is carried out by a Monte Carlo (MC) trial move. Beside thermalization (translation, rotation, etc.), trial moves are carried out in which reactants are removed and reaction products are inserted in the system, in such a way that an equilibrium distribution of reactants and reaction products is obtained. The mechanism and the transition state of the reaction are not considered as this approach is purely thermodynamic and reaction kinetics are not considered. As a result, the efficiency of this simulation technique is not affected by the height of the activation energy barrier of the reaction as reaction kinetics are not considered. The RxMC method requires the ideal gas partition functions of all reactant and reaction product molecules, a list of all possible chemical reactions in the system, and an appropriate force field accurately describing interactions between molecules. The ideal gas partition function can be obtained from Quantum Mechanics or standard thermochemical tables, e.g., the JANAF tables. Computing ideal gas partition functions using quantum packages is well established. However, due to lack of experimental data, it is not always straightforward to compute partition functions from QM software, especially for large molecules or ions. For a detailed review of RxMC techniques, the reader is referred to ref 4.

Just like many other ensembles that rely on sufficient molecule insertions/removals (e.g., the Gibbs ensemble and grand-canonical ensemble), RxMC struggles when insertions/removals of molecules are difficult, e.g., at low temperatures and high densities. During the past few years, significant progress has been made in Monte Carlo techniques for the insertion and deletion of molecules. There are two types of solutions to overcome low acceptance probabilities for molecule insertions/removals in RxMC: methods such as Configurational-Bias Monte Carlo (CBMC) or related methods that try to insert whole molecules in a single Monte Carlo trial move and methods based on the idea of expanded ensembles such as the Continuous Fractional Component Monte Carlo (CFCMC) method first developed by Shi and Maginn. The main advantage of the latter approach is that instead of inserting whole molecules in a single trial move, molecules are inserted gradually, so that the surrounding can easily adapt to the inserted/deleted molecules. This is particularly important at high densities. Therefore, CFCMC does not rely on the spontaneous occurrence of cavities in the system that are large enough to accommodate a large molecule. The CFCMC technique is frequently used for simulations that suffer from low acceptance probability of molecule insertions/removals. Applications of this approach include computation of the loading and enthalpy of adsorption of guest molecules in porous materials near the saturation loading, reaction equilibria of complex systems, and solubilities of small molecules in ionic liquids.

For more details on the challenges of Monte Carlo simulations in open ensembles, the reader is referred to refs 60−62. The combination of CFCMC in RxMC was first proposed by Rosch and Maginn from now on referred to as “parallel Rx/CFC”. Balaji et al. used parallel Rx/CFC to compute the equilibrium concentrations of the different species in CO2/monoethanolamine solutions for different CO2 loadings. In this method, fractional molecules of reaction products are gradually changed into whole reaction product molecules, while the fractional molecules of reactants are gradually removed and vice versa. This algorithm is shown schematically in Figure 1a. A key ingredient of parallel Rx/CFC is that the fractional molecules of both all reactants and reaction products are always present in the system. This CFCMC version of RxMC improves the acceptance probability of molecule insertions/removals significantly compared to the conventional RxMC algorithm. It does not allow direct calculation of chemical potentials, and it is not possible to directly check if the reaction is at equilibrium. Additional free energy calculations are needed to compute the chemical potential.
potentials of reactant and reaction product molecules. The fractional molecules of reactants and reaction products have to adapt to their surroundings simultaneously, which reduces the efficiency of the algorithm.

Recently, a more efficient approach to combine CFCMC with the Gibbs ensemble was introduced by us. Only one fractional molecule per component is present, which can be in either of the simulation boxes, and chemical potentials of all components are obtained without any additional calculations. Inspired by this, a new formulation for RxMC combined with CFCMC is introduced (denoted by serial Rx/CFC). The crucial difference with the parallel Rx/CFC is that either fractional molecules of reactants or reaction products are present in the system. The chemical potentials of reactants/reaction products are directly obtained without using Widom’s test particle insertion (or related) method. Therefore, one can directly check for the condition of chemical equilibrium.

This paper is organized as follows. In Section 2, the conventional RxMC ensemble and its combination with CFCMC are reviewed. Our formulation of RxMC with CFCMC (denoted by serial Rx/CFC) is introduced in Section 3. The partition function, types of trial moves, and computation of chemical potentials are also discussed in this section. Simulation details and systems are described in Section 4. In Section 5, the performance of serial Rx/CFC is compared to conventional RxMC and parallel Rx/CFC for Lennard-Jones (1J) molecules. We considered various model reactions and pressures for which ideal gas free energy changes are specified in advance. Our approach is also tested for the reaction of ammonia pressures for which ideal gas free energy changes are specified in advance. Our approach is also tested for the reaction of ammonia. Here, simplicity, we only consider systems with a single reaction, as extension to systems with multiple reactions is straightforward.\(^1\),\(^3\) The partition function for the constant pressure version of conventional RxMC equals\(^4,\)\(^5,\)\(^6\)

\[
Q_{\text{Conv,P}} = \beta P \sum_{N_i=0}^{\infty} \ldots \sum_{N_{i'}}^{\infty} \int dV \exp\left[-\beta PV\right] \exp\left[\sum_{i=1}^{S} \left(\beta \mu_i N_i + N_i \ln \frac{V_q}{\Lambda_i^3} - \ln N_i!\right)\right] \int ds \exp\left[-\beta U(s^{\text{new}})\right]
\]

where \(S\) is the number of components, \(\beta = 1/(k_B T)\), \(k_B\) is the Boltzmann constant, \(s\) are reduced coordinates, \(V\) is the volume of the simulation box, \(P\) is the pressure, \(N_{\text{total}}\) is the total number of molecules in the simulation box, and \(U\) is the total potential energy. Here, \(q_i, \mu_i, N_i\), and \(\Lambda_i\) are the ideal gas partition function excluding the translational part, the chemical potential, the number of molecules, and the thermal wavelength of component (molecule type) \(i\), respectively. The ensemble of eq 1 is subject to the constraints that the total number of atoms of each type is constant and that chemical reactions converting reactants into reaction products are in equilibrium. This sets limits on the values of \(\mu_i\). Sampling of configurations in this ensemble requires (1) sampling of the degrees of freedom of the interacting molecules (e.g., translation, rotation (for chain molecules), and sampling the internal configuration of molecules), (2) sampling the volume fluctuations, and (3) sampling of chemical reactions subject to the constraint that the total number of atoms of each component is constant, as well as that the reaction is at chemical equilibrium. The latter is obtained by performing reaction trial moves. The reaction trial move is attempted to remove randomly selected reactants and insert reaction product molecules, simultaneously. According to the partition function of conventional RxMC (eq 1), the probabilities of being in the old and new configurations for the reaction trial move in the forward direction are

\[
p_n = \frac{\beta P}{Q_{\text{Conv}}} \exp[-\beta PV] \exp\left[\sum_{i=1}^{S} \left(\beta \mu_i (N_i - \nu_i) + (N_i - \nu_i) \ln \frac{V_q}{\Lambda_i^3} - \ln (N_i - \nu_i)!\right)\right] \times \exp\left[\sum_{j=R+1}^{S} \left(\beta \mu_j (N_j + \nu_j) + (N_j + \nu_j) \ln \frac{V_q}{\Lambda_j^3} - \ln (N_j + \nu_j)!\right)\right] \exp[-\beta U_n]
\]

where \(\nu_i\) is the stoichiometric coefficient of component \(i\) in the reaction. Here, \(n\) and \(o\) denote the new and old configurations, respectively. We choose the convention that \(\nu_i\) is positive if component \(i\) participates in the reaction, and \(\nu_i\) is zero otherwise.

Here, \(R\) is the number of reactant components, and \(P\) is the number of reaction product components. As only systems with a single reaction where all components are either reactants or reaction products are considered here, one can write \(R + P = S\).
Therefore, the reaction product components are ranging from \( R + 1 \) to \( S \), with \( S \) being the total number of components. The summation \( \sum_{i=1}^{R} \) is a sum over all reactant types, and \( \sum_{R+1}^{S} \) is the sum over all reaction product types. Therefore, the ratio of the probabilities of being in the new and old configurations equals

\[
\frac{P_n}{P_o} = \exp \left[ - \beta \sum_{i=1}^{R} \mu_i \bar{n}_i - \sum_{i=1}^{R} \bar{n}_i \ln \frac{Vq_i}{N_i} + \sum_{j=R+1}^{S} \ln \frac{N_j!}{(N_j + \nu_j)!} \right] \times \exp \left[ \beta \sum_{j=R+1}^{S} \mu_j \bar{n}_j + \sum_{j=R+1}^{S} \bar{n}_j \ln \frac{Vq_j}{N_j} + \sum_{i=1}^{R} \ln \frac{N_i!}{(N_i - \nu_i)!} \right] \times \exp[-\beta \Delta U]
\]

hence \( \Delta U = U_n - U_o \) is the total change in the potential energy of the system. Reaction equilibrium implies \( \sum_{i=1}^{R} \mu_i \bar{n}_i = \sum_{R+1}^{S} \mu_j \bar{n}_j \). Consequently, the acceptance rule for the reaction trial move is\(^4,\)\(^3,\)\(^2\)

\[
\text{acc}(o \to n) = \min \left( 1, \left( \prod_{i=1}^{R} \frac{Vq_i}{N_i^\Delta} \right)^{\bar{n}_i} \times \left( \prod_{j=R+1}^{S} \frac{Vq_j}{N_j^\Delta} \right)^{\bar{n}_j} \times \exp[-\beta \Delta U] \right)
\]

(4)

Due to simultaneous insertion of the molecules in a single step, the efficiency of this algorithm can be very low at high densities. This is also the case when Configurational-bias Monte Carlo is used for inserting/deleting molecules.\(^26\)

In parallel Rx/CFC,\(^5,\)\(^5\) the conventional RxMC is expanded with fractional molecules of each component participating in the reaction (Figure 1a). The number of fractional molecules of each component is equal to its stoichiometric coefficient. Interactions of the fractional molecules are scaled with a coupling parameter \( \lambda \). The value \( \lambda = 0 \) corresponds to no interactions with the surrounding molecules (the fractional molecule acts as an ideal gas molecule), and \( \lambda = 1 \) corresponds to full interactions with the surrounding molecules (the fractional molecule has the same interactions as whole molecules of the same component). There are two coupling parameters per reaction, one for all reactants \( (\lambda_R) \) and one for all reaction products \( (\lambda_P) \). The coupling parameters for the fractional molecules of reactants and reaction products are constrained by \( \lambda_R + \lambda_P = 1 \). Attempts are made to change the coupling parameters by \( \lambda_{R,P} = \lambda_{R,P} + \Delta \lambda \) with \( \Delta \lambda \in [\Delta \lambda_{\text{min}}, \Delta \lambda_{\text{max}}] \). Due to the constraint \( \lambda_R + \lambda_P = 1 \), the coupling parameter of the fractional molecules of reaction products also changes according to \( \lambda_{P,R} = \lambda_{R,P} - \Delta \lambda \). When \( \lambda_{R,P} > 1 \) or \( \lambda_{P,R} < 0 \), an attempt is made to perform a chemical reaction. The acceptance rule for performing a chemical reaction in this ensemble is the same as eq 4. For more details, we refer the reader to the original publication by Maginn and Rosch.\(^5,\)

### 3. SERIAL Rx/CFC

#### 3.1. Partition Function.

In serial Rx/CFC, either fractional molecules of the reactants or reaction products are present in the system, in sharp contrast to parallel Rx/CFC where fractional molecules of both reactants and reaction products are always present (Figure 1b). Besides trial moves for thermalization and volume changes, there are three additional trial moves to facilitate the sampling of chemical reactions subject to the constraint that the total number of atoms of each component is constant, as well as chemical equilibrium. As derived in the Supporting Information, the partition function for the constant pressure version of the ensemble equals (not yet taking into account the conservation of atoms)

\[
Q_{\text{CFC,P}} = \beta P \sum_{N_{\text{tot}}=0}^{\infty} \sum_{N_{i}=0}^{\infty} \int \frac{dV}{V} \exp[-\beta PV] \exp \left[ \beta \sum_{j=1}^{R} \mu_j (N_j + \nu_j) + \sum_{j=R+1}^{S} (N_j + \nu_j) \ln \frac{Vq_j}{N_j^\Delta} - \sum_{j=1}^{R} \ln N_j! \right] \times \exp \left[ \beta \sum_{j=R+1}^{S} \mu_1 (N_j + \nu_1 (1-\delta)) + \sum_{j=R+1}^{S} (N_j + \nu_1 (1-\delta)) \ln \frac{Vq_j}{N_j^\Delta} - \sum_{j=1}^{R} \ln N_j! \right] \times \frac{1}{\lambda_1} \int ds_{\text{tot}} \exp[-\beta U_{\text{tot}}(s_{\text{tot}})] \left( \prod_{i=1}^{R} \int ds_{\text{frac}} \exp[-\beta U_{\text{frac}}(s_{\text{frac}}, \lambda)] \right) \times \left( \prod_{j=R+1}^{S} \int ds_{\text{frac}} \exp[-\beta (1-\delta) U_{\text{frac}}(s_{\text{frac}}, \lambda)] \right)
\]

(5)

where \( N_{\text{tot}} \) is the total number of whole molecules (regardless of the component type), and \( N_i \) is the number of whole molecules of component \( i \). When \( \delta = 1 \), fractional molecules of reactants are present in the simulation box \( (\nu_i \) fractional molecule for component \( i \)), and when \( \delta = 0 \), fractional molecules of reaction products are present. Here, a system with a single reaction is considered. Also, \( U_{\text{int}} \) is the total internal energy of whole molecules, and \( U_{\text{frac}} \) is the interaction energy of fractional molecules of component \( i \) with the other molecules, including other fractional molecules. The interactions of the fractional molecules with the surroundings are such that \( \lambda = 0 \) means no interactions and \( \lambda = 1 \) means full interactions, and the value of \( \lambda \) is restricted to \( \lambda \in [0,1] \).

Since fractional molecules are always distinguishable from whole molecules, the term \( N_i! \) only counts for whole indistinguishable molecules. The main difference between eq 5 and eq 1 is the integration over \( \lambda \) in eq 5. This is an immediate consequence of expanding the conventional RxMC with fractional molecules. In the Supporting Information, we show that for systems without intermolecular interactions (ideal gas
(phase) the partition functions of eq 5 and eq 1 are proportional. Therefore, these ensembles result in identical average numbers of molecules for each component, provided that fractional molecules are not counted when computing ensemble averages.

The fact that one should not count fractional molecules when computing the average number of molecules is in line with earlier studies in the Gibbs ensemble and in the grand-canonical ensemble.45,64

3.2. Trial Moves. In addition to Monte Carlo trial moves for thermalization and volume changes, there are three trial moves in this ensemble to mimic the chemical reaction. A detailed description of these trial moves and the derivation of the acceptance rules is provided in the Supporting Information.

3.2.1. Changing the Value of $\lambda$. This trial move is used to change the value of $\lambda$ either for reactants or reaction products, depending on the value of $\delta$ (Figure 2). The value of $\lambda$ is changed according to $\lambda_n = \lambda_o + \Delta \lambda$, in which $\Delta \lambda$ is a uniformly distributed random number between $-\Delta \lambda_{\text{max}}$ and $+\Delta \lambda_{\text{max}}$. Note that $\Delta \lambda_{\text{max}}$ can be different for reactants and reaction products. When the new value of $\lambda$ is not in the range $\lambda \in [0,1]$, this trial move is automatically rejected. In this trial move, the value of $\delta$, all positions of molecules, and the number of whole molecules and fractional molecules remain the same. By changing the value of $\lambda$, only the interactions between the fractional molecules and other molecules are changed. In the Supporting Information, it is shown that the acceptance rule for this trial move is

$$\text{acc}(o \rightarrow n) = \min[1, \exp[-\beta \Delta U]]$$

in which $\Delta U = U_n - U_o$ is the change in the total internal energy of the system.

3.2.2. Reaction for Fractional Molecules. In this trial move, fractional molecules of reactants/reaction products are removed, and fractional molecules of reaction products/reactants are inserted at random positions (Figure 3). In this trial move, essentially the value of $\delta$ is changed, so if $\delta_o = 1$ then $\delta_n = 0$ and vice versa. The number of whole molecules and also the value of $\lambda$ can be different for reactants and reaction products. When the new value of $\lambda$ is not in the range $\lambda \in [0,1]$, this trial move is automatically rejected. In this trial move, the value of $\delta$, all positions of molecules, and the number of whole molecules and fractional molecules remain the same. By changing the value of $\lambda$, only the interactions between the fractional molecules and other molecules are changed. In the Supporting Information, it is shown that the acceptance rule for this trial move is

$$\text{acc}(o \rightarrow n) = \min[1, \exp[-\beta \Delta U]]$$

in which $\Delta U = U_n - U_o$ is the change in the total internal energy of the system.
are constant. This trial move basically mimics a chemical reaction for the fractional molecules. Here, the acceptance rule for the forward reaction (reactants \(\rightarrow\) reaction products) is shown. The direction of the chemical reaction is defined by the value of \(\delta\) for the old configuration (if we have the fractional molecules of reactants or reaction products). In the Supporting Information, it is derived that the acceptance rule for converting the reactants into reaction products equals

\[
\text{acc}(o \rightarrow n) = \min \left[ 1, \prod_{i=1}^{R} \left( \frac{Vq}{\lambda_i^2} \right)^{-\nu_i} \prod_{j=R+1}^{S} \left( \frac{Vq_j}{\lambda_j' \lambda_j''} \right)^{\nu_j} \exp[-\beta \Delta U] \right]
\]  

(7)

Since the number of whole molecules remains constant during this move, the terms \(\prod_{i=1}^{R} \frac{N_i^o}{(N_i-o)_i}!\) and \(\prod_{j=R+1}^{S} \frac{N_j^o}{(N_j-o)_j}!\) are not present in eq 7. The acceptance rule for the reverse reaction (reaction products \(\rightarrow\) reactants) simply follows by swapping the labels of the reactants and reaction products. The acceptance probability for this trial move is large when \(\lambda\) is small. The reason for this is that fractional molecules have very limited interactions with the surrounding molecules, and therefore, the term \(\Delta U\) is nearly zero. For the limiting case of \(\lambda \downarrow 0\), the acceptance rule reduces to

\[
\text{acc}(o \rightarrow n) = \min \left[ 1, \prod_{i=1}^{R} \left( \frac{Vq}{\lambda_i^2} \right)^{-\nu_i} \prod_{j=R+1}^{S} \left( \frac{Vq_j}{\lambda_j' \lambda_j''} \right)^{\nu_j} \right]
\]  

(8)

### 3.2.3. Reaction for Whole Molecules

In this trial move, fractional molecules of reactants/reaction products are transformed into whole molecules of reactants/reaction products, while simultaneously, randomly selected whole molecules of reaction products/reactants are transformed into fractional molecules of reaction products/reactants. In this trial move, all molecule positions and the value of \(\lambda\) stay the same. The value of \(\delta\) is changed as follows: if \(\delta_o = 1\), then \(\delta_n = 0\) and vice versa. This trial move is illustrated in Figure 4 and can be seen as a reaction for whole molecules. In the forward reaction, whole molecules of reactants are transformed into fractional molecules, and fractional molecules of reaction products are turned into whole molecules. Essentially, the number of whole molecules of reactants is reduced, and the number of whole molecules of reaction products is increased, according to their stoichiometric coefficients. Trial moves are automatically rejected when there are not enough whole molecules to turn into fractional molecules. Here, the acceptance rule for the forward reaction (reactants \(\rightarrow\) reaction products) is shown. The direction of the reaction eventually depends on the value of \(\delta\) for the old configuration (if we have fractional molecules of reactants or reaction products). As derived in the Supporting Information, the acceptance rule for this trial move is

\[
\text{acc}(o \rightarrow n) = \min \left[ 1, \prod_{i=1}^{R} \frac{N_i!}{(N_i-o)_i!} \prod_{j=R+1}^{S} \frac{N_j!}{(N_j-o)_j!} \exp[-\beta \Delta U] \right]
\]  

(9)

in which \(\Delta U = U_o - U_n\) is the change in the total internal energy of the system including the interaction of fractional molecules as result of the trial move. The acceptance rule for the reverse reaction (reaction products \(\rightarrow\) reactants) simply follows by swapping the labels. Since the total number of whole and fractional molecules for each component remains constant, ideal gas partition functions are not present in eq 9. This trial move has a very high acceptance probability when the value of \(\lambda\) is close to 1. The reason for this is that fractional molecules have almost the same interactions as whole molecules, and therefore, the term \(\Delta U\) is nearly zero. For the limiting case of \(\lambda \uparrow 1\), the acceptance rule reduces to

\[
\text{acc}(o \rightarrow n) = \min \left[ 1, \prod_{i=1}^{R} \frac{N_i!}{(N_i-o)_i!} \prod_{j=R+1}^{S} \frac{N_j!}{(N_j-o)_j!} \right]
\]  

(10)
3.2.4. Volume Changes. This trial move is only used for the case where the temperature and external pressure are imposed. In this trial move, the volume of the simulation box is changed, while the number and relative coordinates of the whole molecules and fractional molecules stay the same. Here, the random walk is performed in V and not ln(V). The acceptance rule for this trial move is

\[
\text{acc}(o \rightarrow n) = \min \left[ 1, \left( \frac{V_n}{V_o} \right)^{\text{vol}} \right] \exp \left[ -\beta(\Delta U + P(V_n - V_o)) \right] \tag{11}
\]

3.2.5. Efficient Selection of Trial Moves. As discussed in the previous section, the reaction trial move for fractional molecules has a very high acceptance probability at low values of \( \lambda \), and the reaction trial move for the whole molecules has a very high acceptance probability at high values of \( \lambda \). In Figure 5, typical acceptance probabilities of these trial moves as a function of \( \lambda \) are shown. Therefore, one may attempt reaction trial moves for fractional molecules only at values of \( \lambda \) close to 0, and reaction trial moves for the whole molecules only at values of \( \lambda \) close to 1. In this way, each trial move is used where it is efficient, and the overall efficiency of the algorithm is improved. This is done as follows: one can define a switching point for the value of \( \lambda (\lambda_{\text{sw}}) \). The probabilities of selecting a trial move [thermalization, volume change, or changing the value of \( \lambda \) (Section 3.2.1)] are always constant. For selection of the remaining trial moves (Sections 3.2.2 and 3.2.3), one has a choice: selecting these with fixed probability or always selecting the reaction trial move for fractional molecules (Section 3.2.2) when \( \lambda < \lambda_{\text{sw}} \), or always selecting the reaction trial move for whole molecules (Section 3.2.3) when \( \lambda > \lambda_{\text{sw}} \). In the latter approach, the reaction trial moves are selected when they have a higher acceptance probability. Since the value of \( \lambda \) remains constant during either of these trial moves, the probabilities for selecting the trial moves also remain constant. Therefore, the condition of detailed balance is not violated.

3.3. Biasing the Probability Distribution \( p(\lambda, \delta) \). It is important to bias the probability distribution of \( p(\lambda, \delta) \) (\( \delta \) indicates whether fractional molecules of reagents or reaction products are in the simulation box) in such a way that the sampled probability distributions \( p(\lambda, \delta) \) is flat and that it is equally likely to have the fractional molecules of reagents (\( \delta = 1 \)) or reaction products (\( \delta = 0 \)). By using an adequate biasing function, one can overcome the problem of being "stuck" in free energy minima and can easily diffuse through the \( \lambda \) space. This is obtained by multiplying the statistical weight of each system state by a factor \( \exp [W(\lambda, \delta)] \). For parallel Rx/CFC, since fractional molecules of both reagents and reaction products are always present in the system, one would only need a one-dimensional weight function to obtain flat probability distribution of \( p(\lambda) \). It is important to note that in serial Rx/CFC the weight function \( W(\lambda, \delta) \) is a two-dimensional function that depends both on \( \lambda \) and the identity of the fractional molecules (\( \delta \)). By using this biased sampling, additional terms \( \exp [W(\lambda, \delta_n) - W(\lambda, \delta_f)] \) will be present in the acceptance rules of eqs 6, 7, and 9. For example, the acceptance rule for the trial move attempting to mimic a reaction for the fractional molecules (eq 7) will become

\[
\text{acc}(o \rightarrow n) = \min \left[ 1, \left( \prod_{i=1}^{R} \frac{V_q}{N_i} \right)^{-\frac{\delta}{2}} \prod_{j=R+1}^{S} \left( \frac{V_q}{N_j} \right)^{-\frac{\delta}{2}} \exp \left[ -\beta \Delta U + W(\lambda, \delta_n) - W(\lambda, \delta_f) \right] \right] \tag{12}
\]

To remove this bias, the Boltzmann averages of any observable \( X \) should be computed using

\[
\langle X \rangle_{\text{Boltzmann}} = \frac{\langle \exp[-W(\lambda, \delta)] \rangle_{\text{biased}}}{\langle \exp[-W(\lambda, \delta)] \rangle_{\text{biased}}} \tag{13}
\]

The weight function \( W(\lambda, \delta) \) can be obtained using the Wang–Landau algorithm or iteratively. To compute ensemble averages corresponding to the conventional RxMC while performing simulation with serial Rx/CFC, one should exclude the contribution of fractional molecules. By doing this, one can analytically show that for an ideal gas case the ensemble averages computed using the serial Rx/CFC and the conventional RxMC are identical (see Supporting Information). Including the contribution of the fractional molecules in the ensemble averages leads to small differences between the ensemble averages compute in the serial Rx/CFC and those computed in the conventional RxMC.

3.4. Free Energy Calculations. In serial Rx/CFC, chemical potentials can be computed without any additional computational efforts. As shown in the Supporting Information, one can show that
\[ \sum_{i=1}^{R} \nu \mu_i = -\frac{1}{\beta} \ln \left( \prod_{i=1}^{R} \left( \frac{q_i}{\Lambda_{i}\rho_i} \right)^{\nu_i} \right) = -\frac{1}{\beta} \ln \left( \frac{p(\lambda \uparrow 1, \delta = 1)}{p(\lambda \downarrow 0, \delta = 1)} \right) \]

where \( q_i \) is the ideal gas partition function of component \( i \) excluding the translational part. One can obtain the sum of the chemical potentials of reaction products in a similar way. Equation 14 allows for an independent check of reaction equilibria without any additional calculations (e.g., test molecules). The chemical potential of component \( i \) for a nonideal gas equals

\[ \mu_i = \frac{1}{\beta} \ln \left( \frac{\beta R \Lambda_i^3 q_i}{P_0} \right) + \frac{1}{\beta} \ln \left( \frac{y_i \rho_i}{P_0} \right) \]

in which \( \rho_i \) and \( y_i \) are the fugacity coefficient and mole fraction of component \( i \), respectively. Here, \( P_0 \) is the reference pressure (1 bar), and \( P \) is the pressure of the mixture. The first term on the right-hand side of eq 15 is the standard reference chemical potential \( \mu_i^{0}(T) \). Therefore, we have

\[ \sum_{i=1}^{R} \nu \mu_i = \frac{1}{\beta} \ln \left( \prod_{i=1}^{R} \left( \frac{\beta R \Lambda_i^3 q_i}{\nu \rho_i} \right)^{\nu_i} \right) \]

Combining this with eq 14 immediately leads to

\[ \prod_{i=1}^{R} q_i^{-\nu_i} = \frac{p(\lambda \uparrow 1)}{p(\lambda \downarrow 0)} \times \prod_{i=1}^{R} \left( \frac{\beta y_i P}{\rho_i} \right)^{\nu_i} \]

where \( \lambda \equiv \lambda \) when we have the fractional molecules of reactants (\( \delta = 1 \)). In this equation, \( p(\lambda \uparrow 1) \) is the probability that \( \lambda \) approaches 1, and \( p(\lambda \downarrow 0) \) is the probability that \( \lambda \) approaches 0. To compute the chemical potential of individual components, one should couple the interactions of different components in a smart way. The two limiting cases are well defined: at \( \lambda = 0 \), fractional molecules of reactants (or reaction products) do not interact, and at \( \lambda = 1 \), fractional molecules of reactants have full interactions with the surrounding molecules. However, for intermediate values of \( \lambda \), one has a choice. One can choose different paths to scale the interactions of fractional molecules from no interactions to full interactions. The interactions can be scaled atom by atom, or molecule by molecule, or in any other way. By scaling the interactions of the fractional molecules of only one of the components from no interactions to full interactions when the value of \( \lambda \) is changed from 0 to \( A \), one can write

\[ \nu \mu_i = -\frac{1}{\beta} \ln \left( \frac{q_i}{\Lambda_{i}\rho_i} \right)^{\nu_i} - \frac{1}{\beta} \ln \left( \frac{p(\lambda \uparrow 1, A \uparrow 1)}{p(\lambda \downarrow 0, A \downarrow 0)} \right) \]

The first term on the right-hand side accounts for the ideal gas part. The second term on the right-hand side accounts for the excess part of the chemical potential (due to interactions with surrounding molecules). Similar to eq 17, one can write for \( \nu_i \)

\[ q_i^{-\nu_i} = \frac{p(\lambda \uparrow 1, A \uparrow 1)}{p(\lambda \downarrow 0, A \downarrow 0)} \times \left( \frac{\beta y_i P}{\rho_i} \right)^{\nu_i} \]

When \( \nu_i > 1 \) and interactions of fractional molecules are scaled sequentially (one by one), fractional molecules that have interactions with surrounding molecules later (at higher values of \( \lambda \)) experience the effect of the fractional molecules which were inserted earlier (at lower values of \( \lambda \)). For sufficiently large system sizes, this will not affect the calculated chemical potentials. However, for small system sizes, there may be minor differences between the chemical potential of molecules that are inserted at lower values of \( \lambda \) and those inserted at higher values of \( \lambda \). Although these differences are expected to be small, one should be aware of them.

4. SIMULATION DETAILS

As a proof of principle, simulations are performed for different systems of LJ molecules. The ammonia synthesis reaction at various pressures (100–1000 bar) and temperatures (573–873 K) is also considered as a practically important application. For different systems of LJ molecules, simulations are performed at constant pressure and temperature using conventional ReaxMC, parallel Rx/CFC, and serial Rx/CFC. Various model reactions of LJ molecules are studied. For these simulations, all properties are defined in reduced units. LJ interactions are truncated and shifted at 2.5σ. The reduced temperature is set to \( T = 2.0 \), and simulations are always started with 400 molecules of component A. For simulations of LJ molecules using parallel Rx/CFC and serial Rx/CFC, the maximum molecule displacements, maximum volume change, and maximum change in the value of \( \lambda \) are set to achieve 50% acceptance for these trial moves. For simulations using serial Rx/CFC, the switching point for the value of \( \lambda \) is set to \( \lambda_{\text{eq}} = 0.3 \) (Section 3.2.4). In each Monte Carlo step, a trial move is selected at random with the following probabilities: 49.5% molecule displacements, 1% volume changes, and 49.5% reaction trial moves.

For the ammonia synthesis reaction, simulations are performed at constant pressure and temperature using serial Rx/CFC. All simulations start with a random configuration of 120 \( \text{N}_2 \), 360 \( \text{H}_2 \) molecules, and no ammonia molecules. Fractional molecules are added to this configuration. All molecules are rigid and interact only through LJ and electrostatic interactions. Force field parameters for \( \text{N}_2 \), \( \text{H}_2 \), and \( \text{NH}_3 \) are taken from the literature.67–70 The Wolf method is used to compute electrostatic interactions.1 Details regarding the force field parameters, scaling of the electrostatic interactions, and the Wolf method are provided in the Supporting Information. The ideal gas partition functions for this system are listed in Table S3 of the Supporting Information. In this table, a detailed comparison is made between ideal gas partition functions from experiments and QM computations using Gaussian.72 In each Monte Carlo step, a trial move is selected at random with the following probabilities: 33% molecule displacements, 33% molecule rotation, 1% volume changes, and 33% reaction trial moves. For the ammonia synthesis reaction, LJ interactions are switched on for \( \lambda \in [0,0.9] \), and electrostatic interactions are switched on for \( \lambda \in [0.9,1] \).

For all simulations using parallel Rx/CFC and serial Rx/CFC, the weight function is determined using the Wang–Landau algorithm.65 In serial Rx/CFC, the weight function \( W(\lambda, \delta) \) is determined such that the observed two-dimensional probability distribution \( p(\lambda, \delta) \) in the proposed ensemble is flat. Here, 200 bins are used to store the probability distribution of \( \lambda \) for reactants or reaction products. All simulations are started with 0.2 million Monte Carlo cycles to equilibrate the system, followed by 1 million production trial moves. The number of Monte Carlo steps per cycle equals the total number of molecules initially in the system, with a minimum of 20. LJ interactions are scaled according to.
In the conventional method and parallel Rx/CFC, there is only one type of reaction trial move. In contrast, serial Rx/CFC requires three types of trial moves for facilitating molecule transfers: 50% changes in the λ-space (Figure 2), 50% reaction for the fractional molecules (Figure 3) when λ < λ_{sw} or 50% reaction for the whole molecules (Figure 4) when λ > λ_{sw}. In serial Rx/CFC, the chemical potentials are computed from eq 14. The contribution of fractional molecules are excluded while computing ensemble averages.

To compare the efficiencies of the three methods, a fair way to define the efficiency of each method is required. In this work, the efficiency for all three methods is defined as the number of accepted trial moves (either forward or backward) resulting in a change in the number of whole molecules due to the reaction divided by the total number of reaction trial moves. For parallel Rx/CFC, this means the number of accepted λ trial moves that resulted in $\lambda_n > 1$ or $\lambda_n < 0$ divided by the total number of λ trial moves. For serial Rx/CFC, this means the number of accepted reaction trial moves for whole molecules (Figure 4) divided by the total number of all reaction trial moves, including changing the value of $\lambda_n$ reaction for fractional molecules, and reaction for whole molecules. It should be noted that reaction trial moves in serial Rx/CFC are computationally cheaper compared to parallel Rx/CFC. This is due to the reduction in the number of fractional molecules, and therefore, the number of interacting molecule pairs is reduced. Simulations performed using serial Rx/CFC require less CPU time compared to similar simulations when parallel Rx/CFC is used. The CPU time depends on the programming of the algorithms, the compiler, and CPUs used to perform the calculations. In this paper, different approaches are only compared in terms of efficiency and not the CPU time. This can be considered as the worst-case scenario for serial Rx/CFC.

5. RESULTS

To ensure that serial Rx/CFC is implemented correctly, the equilibrium composition for different reactions of LJ molecules are computed and compared for the three methods. The LJ parameters and partition function for LJ molecules used in this study are listed in Table 1. The equilibrium composition obtained with different methods at reduced pressures $P = 0.3, P = 1.0, P = 3.0$, and $P = 5.0$ are shown in Tables 2–5, respectively. Equilibrium compositions obtained for the three methods are the same for all reactions and conditions (Tables 2–5). This confirms the validity of the expressions used for the partition function and acceptance rules of serial Rx/CFC and indicates that this method is implemented correctly. The efficiencies of the three methods for different reactions are also shown in Tables 2–5. The conventional method has a very high efficiency for all reactions at the lowest pressure ($P = 0.3$). Since in this case the density of the system is very low and therefore interactions between the molecules are limited, there is almost no energy penalty for the reaction trial moves, and most of the attempts to perform reaction trial moves for whole molecules are accepted. Therefore, the method which attempts to directly replace the reactants with reaction products and vice versa has a high efficiency. For the conventional method, reaction trial moves for the whole molecules are the only reaction trial move, and this trial move is accepted with a high probability for the low pressure case. As a result, this method has high efficiencies for this case. In parallel Rx/CFC, many trial moves are spent diffusing through the entire λ-space, and less attempts are made to perform a reaction. Therefore, this method has the lowest efficiency for the low pressure case. Already at $P = 1.0$, the efficiency of the conventional method is much lower than its efficiency at $P = 0.3$. At higher pressures ($P = 3.0, P = 5.0$), the efficiency of the conventional method drops below 0.01 even for the simplest reaction ($A \rightleftharpoons B$). When the density is high, most of the reaction trial moves in the conventional method result in an overlap between the newly inserted molecules and molecules that are already in the system. Therefore, this move has very low acceptance probability. In this case, the efficiency of parallel Rx/CFC varies between 0.06 to 0.1, while the efficiency of serial Rx/CFC varies between 0.1 to 0.2 depending on the reaction. Due to the efficient use of the three trial moves in serial Rx/CFC, this method has a better performance compared to the conventional method and parallel Rx/CFC.

In Figure 6, the (unbiased) probability distributions $p(\lambda, \delta)$ for two different reactions ($A \rightleftharpoons B$ and $A \rightleftharpoons D + E$) and the weight functions to make $p(\lambda, \delta)$ flat are shown. For the reaction $A \rightleftharpoons B$, the probability distributions and the weight functions for the reactants and reaction products are identical, as A and B have a similar interaction with the surrounding molecules. For the reaction $A \rightleftharpoons D + E$, one reactant molecule is replaced with two product molecules. For this reaction, the interactions of the first product molecule are scaled from no interactions (ideal gas molecule) to full interactions (whole molecule) when the value of $\lambda$ is changed from 0 to $\frac{1}{2}$. For the second molecule, the interactions are scaled from no interactions (ideal gas molecule) to full interactions (whole molecule) when the value of $\lambda$ is changed from $\frac{1}{2}$ to 1. This can also be clearly seen in the shape of the probability distribution of $\lambda$ and the weight function of the reactant molecules. In this way, according to eq 18, one can obtain the excess chemical potential of the first reactant molecule using $p(\lambda_R \downarrow 0)$ and $p(\lambda_R \uparrow \frac{1}{2})$ and the excess chemical potential of the second reactant molecule using $p(\lambda_R \downarrow \frac{1}{2})$ and $p(\lambda_R \uparrow 1)$. The values obtained for the excess chemical potential of the first and second reactant molecules were very close to each other.

In Table 6, the sum of the total and excess chemical potentials times the stoichiometric coefficients are shown for the reactants and reaction products for different pressures and reactions. These values can only be directly computed in serial Rx/CFC according to eq 14. The data provided in Table 6 shows that for

\[
u_k(\tau, \lambda) = 4\lambda_k \left[ \frac{1}{\delta (1 - \lambda)^2 + (\frac{\tau}{\delta})^4} - \frac{1}{\delta (1 - \lambda)^2 + (\frac{\tau}{\delta})^4} \right]
\]

Table 1. Interaction Parameters (Lennard-Jones) and Partition Functions ($q/A^3$) for Different Molecule Types

| Molecule type | $\sigma$ | $\epsilon$ | $q/A^3$ |
|---------------|---------|-----------|---------|
| A             | 1.0     | 1.0       | 0.002   |
| B             | 1.0     | 1.0       | 0.002   |
| C             | 1.1     | 0.9       | 0.002   |
| D             | 1.0     | 1.0       | 0.02    |
| E             | 1.1     | 0.9       | 0.02    |
| F             | 1.0     | 1.0       | 0.02    |

"Note that there are several molecule types with exactly the same interaction parameters. This was done to show the effect of (in)distinguishability of the molecules in the reactions."
Table 2. Average Number of Molecules and Density at Equilibrium for Different Reactions for Different Methods

| Reaction | \(N_A\) | \(\langle N_{\text{Product 1}} \rangle\) | \(\langle N_{\text{Product 2}} \rangle\) | \(\rho_{\text{tot}}\) | Efficiency | Method |
|----------|----------|-----------------|-----------------|----------------|------------|--------|
| A ⇄ B    | 200.00(4)| 200.00(4)        | 0.433(0)        | 0.077          | Conventional |
| A ⇄ C    | 200.00(2)| 200.00(2)        | 0.432(0)        | 0.26           | Serial Rx/CFC |
| A ⇄ D    | 226.48(4)| 173.52(4)        | 0.392(0)        | 0.068          | Serial Rx/CFC |
| A ⇄ E    | 226.45(9)| 173.59(9)        | 0.391(0)        | 0.26           | Serial Rx/CFC |
| A ⇄ F    | 273.05(5)| 253.89(5)        | 0.433(0)        | 0.017          | Serial Rx/CFC |
| A ⇄ G    | 272.8(2) | 254.5(4)         | 0.430(0)        | 0.074          | Serial Rx/CFC |
| A ⇄ H    | 272.8(2) | 254.5(3)         | 0.431(0)        | 0.17           | Serial Rx/CFC |
| A ⇄ I    | 300.57(6)| 198.9(2)         | 0.395(0)        | 0.011          | Serial Rx/CFC |
| A ⇄ J    | 300.4(2) | 199.3(2)         | 0.393(0)        | 0.059          | Parallel Rx/CFC |
| A ⇄ K    | 177.73(5)| 222.27(5)        | 0.433(0)        | 0.017          | Serial Rx/CFC |
| A ⇄ L    | 177.4(3) | 222.6(3)         | 0.431(0)        | 0.075          | Parallel Rx/CFC |
| A ⇄ M    | 177.5(2) | 222.5(2)         | 0.431(0)        | 0.17           | Serial Rx/CFC |
| A ⇄ N    | 197.92(7)| 202.08(7)        | 0.401(0)        | 0.014          | Serial Rx/CFC |
| A ⇄ O    | 197.6(3) | 202.4(3)         | 0.399(0)        | 0.070          | Parallel Rx/CFC |
| A ⇄ P    | 197.6(2) | 202.4(2)         | 0.399(0)        | 0.17           | Serial Rx/CFC |

The efficiency is defined in Section 4. The reduced pressure and temperature are set to \(P = 0.3\) and \(T = 2.0\), respectively. Simulations are started with 400 molecules of type A. The interaction parameters of different molecules are listed in Table 1. The numbers between brackets denote the uncertainty in the last digit.

Table 3. Average Number of Molecules and Density at Equilibrium for Different Reactions for Different Methods

| Reaction | \(N_A\) | \(\langle N_{\text{Product 1}} \rangle\) | \(\langle N_{\text{Product 2}} \rangle\) | \(\rho_{\text{tot}}\) | Efficiency | Method |
|----------|----------|-----------------|-----------------|----------------|------------|--------|
| A ⇄ B    | 200.00(3)| 200.00(3)        | 0.162(0)        | 0.40           | Conventional |
| A ⇄ C    | 206.62(2)| 193.38(2)        | 0.155(0)        | 0.37           | Conventional |
| A ⇄ D    | 206.63(3)| 193.37(2)        | 0.154(0)        | 0.098          | Parallel Rx/CFC |
| A ⇄ E    | 192.59(6)| 414.8(2)         | 0.162(0)        | 0.26           | Serial Rx/CFC |
| A ⇄ F    | 202.75(5)| 394.5(1)         | 0.153(0)        | 0.21           | Serial Rx/CFC |
| A ⇄ G    | 202.35(6)| 395.3(2)         | 0.152(0)        | 0.086          | Parallel Rx/CFC |
| A ⇄ H    | 202.47(4)| 395.06(8)        | 0.152(0)        | 0.25           | Serial Rx/CFC |
| A ⇄ I    | 91.52(3) | 308.48(3)        | 0.162(0)        | 0.26           | Parallel Rx/CFC |
| A ⇄ J    | 91.22(9) | 308.78(9)        | 0.161(0)        | 0.097          | Parallel Rx/CFC |
| A ⇄ K    | 93.33(3) | 308.67(3)        | 0.161(0)        | 0.25           | Serial Rx/CFC |
| A ⇄ L    | 95.57(3) | 304.43(3)        | 0.156(0)        | 0.23           | Serial Rx/CFC |
| A ⇄ M    | 95.39(2) | 304.61(2)        | 0.155(0)        | 0.094          | Parallel Rx/CFC |

The efficiency is defined in Section 4. The reduced pressure and temperature are set to \(P = 1.0\) and \(T = 2.0\), respectively. Simulations are started with 400 molecules of type A. The interaction parameters of different molecules are listed in Table 1. The numbers between brackets denote the uncertainty in the last digit.

the reaction A ⇄ B, where the reactant and reaction products have identical LJ interactions, the values obtained for the chemical potentials of the reactants and reaction products are equal. Since molecules of A and B are identical (Table 1), this is exactly what is expected. This case is included because it is trivial and can serve as an additional check on the implementation and on convergence of the simulation. It is verified that the computed excess chemical potentials are identical to those obtained from Widom’s test particle insertion method in the conventional NPT ensemble at the same conditions (data not shown).

It is instructive to repeat this test case for systems with a full LJ potential. The use of tail correction is dictated by legacy use of some force field like TraPPE, as these were developed using Monte Carlo methodology. It tries to approximate the “full” Lennard-Jones interactions using a finite (small) cutoff to keep computations tractable. However, for many other system, especially for use in MD, the discontinuity such a cutoff would create in the forces is problematic. In the SI, it is also shown that for the case of LJ molecules (truncated interactions and using analytic tail correction) identical values for the excess chemical potentials are obtained from the serial Rx/CFC, Widom’s test particle insertion method in the conventional NPT, and EOS modeling using a full LJ potential.

Reaction equilibrium implies \(\sum_{i=R+1}^N n_i \mu_i = 0\). It can be clearly seen that this condition is satisfied for all reactions at all pressures within the error bars. This indicates that simulations have reached the condition of chemical equilibrium, and one can trust the results obtained from the simulations. Moreover, one
CFC simulations at different fractions of ammonia at equilibrium obtained from serial Rx/CFC compared with experimental results and results using equation approaches. This validates the applicability of serial Rx/CFC mixture compositions obtained using the three different methods. Excellent agreement is observed between the equilibrium computed using serial Rx/CFC simulations are in very good agreement with those calculated from equation of state modeling. No experimental data was found to compare with the results of thermodynamic modeling (using the PR equation of state) at different temperatures and pressures. For pressures below 600 bar, fugacity coefficients at very high pressures can be accurately estimated using the PR equation of state.

Table 4. Average Number of Molecules and Density at Equilibrium for Different Reactions for Different Methods

| Reaction | \(\langle N_A \rangle\) | \(\langle N_{Product 1} \rangle\) | \(\langle N_{Product 2} \rangle\) | \(\langle \rho_{tot} \rangle\) | Efficiency | Method |
|----------|-----------------|-----------------|-----------------|-----------------|-----------|--------|
| A ⇌ B    | 200.1(2)        | 199.9(2)        | 0.667(0)        | 7 × 10^{-3}     | Parallel Rx/CFC |
| A ⇌ C    | 199.9(4)        | 200.1(4)        | 0.665(0)        | 0.96            | Serial Rx/CFC |
| A ⇌ 2D   | 268.7(2)        | 131.3(2)        | 0.614(0)        | 5 × 10^{-3}     | Serial Rx/CFC |
| A ⇌ 2E   | 268.8(2)        | 131.2(2)        | 0.612(0)        | 0.76            | Serial Rx/CFC |
| A ⇌ D + F| 345.2(2)        | 109.5(4)        | 0.614(0)        | 0.20           | Serial Rx/CFC |
| A ⇌ D + E| 345.0(3)        | 110.0(5)        | 0.665(0)        | 0.066          | Parallel Rx/CFC |
| A ⇌ D + F| 293.1(6)        | 106.9(6)        | 0.665(0)        | 0.68           | Parallel Rx/CFC |
| A ⇌ D + E| 293.3(5)        | 106.7(5)        | 0.666(0)        | 0.11           | Serial Rx/CFC |
| A ⇌ D + E| 324.2(2)        | 75.8(2)         | 0.641(0)        | 2 × 10^{-4}    | Serial Rx/CFC |
| A ⇌ D + E| 324.2(5)        | 75.8(5)         | 0.638(0)        | 0.064          | Parallel Rx/CFC |

The efficiency is defined in Section 4. The reduced pressure and temperature are set to \(P = 5.0\) and \(T = 2.0\), respectively. Simulations are started with 400 molecules of type A. The interaction parameters of different molecules are listed in Table 1. The numbers between brackets denote the uncertainty in the last digit.

Table 5. Average Number of Molecules and Density at Equilibrium for Different Reactions for Different Methods

| Reaction | \(\langle N_A \rangle\) | \(\langle N_{Product 1} \rangle\) | \(\langle N_{Product 2} \rangle\) | \(\langle \rho_{tot} \rangle\) | Efficiency | Method |
|----------|-----------------|-----------------|-----------------|-----------------|-----------|--------|
| A ⇌ B    | 199.8(3)        | 202.3(3)        | 0.766(0)        | 1 × 10^{-3}     | Parallel Rx/CFC |
| A ⇌ C    | 200.1(4)        | 199.9(4)        | 0.764(0)        | 0.096          | Parallel Rx/CFC |
| A ⇌ 2D   | 298.5(5)        | 101.5(5)        | 0.718(0)        | 9 × 10^{-4}    | Serial Rx/CFC |
| A ⇌ 2E   | 298.5(8)        | 101.8(8)        | 0.716(0)        | 0.79           | Serial Rx/CFC |
| A ⇌ D + F| 298.6(4)        | 101.4(4)        | 0.718(0)        | 0.20           | Serial Rx/CFC |
| A ⇌ D + E| 324.2(1.4)      | 75.9(1)         | 0.659(0)        | 0.11           | Serial Rx/CFC |

The efficiency is defined in Section 4. The reduced pressure and temperature are set to \(P = 5.0\) and \(T = 2.0\), respectively. Simulations are started with 400 molecules of type A. The interaction parameters of different molecules are listed in Table 1. The numbers between brackets denote the uncertainty in the last digit.

Can directly compute the excess chemical potential of individual components according to eq 18.

To test the suitability of serial Rx/CFC simulations for practical systems and molecules with partial charges, the ammonia synthesis reaction \(\text{N}_2 + 3\text{H}_2 \rightarrow 2\text{NH}_3\) is considered. Equilibrium compositions obtained from serial Rx/CFC are validated with the RASPA software. In Figure 7, the mole fractions of ammonia at equilibrium obtained from serial Rx/CFC simulations at different temperatures and pressures are compared with experimental results and results using equation of state modeling (Peng–Robinson (PR) equation of state). Excellent agreement is observed between the equilibrium mixture compositions obtained using the three different approaches. This validates the applicability of serial Rx/CFC for systems including molecules with electrostatic interactions. In Figure 8, fugacity coefficients of ammonia at chemical equilibrium computed using serial Rx/CFC simulations are compared with the results of thermodynamic modeling (using the PR equation of state) at different temperatures and pressures. It is well known that cubic equations of state fail to provide accurate estimates for the fugacity coefficient at very high pressures. For pressures below 600 bar, fugacity coefficients computed using serial Rx/CFC simulations are in very good agreement with those calculated from equation of state modeling. No experimental data was found to compare with the values obtained for fugacity coefficients.
Figure 6. (a) and (c) Probability distributions $p(\lambda, \delta)$ for reactants ($\delta = 1$) and reactions products ($\delta = 0$) for reaction (a) $A \rightleftharpoons B$ and (c) $A \rightleftharpoons D + E$ at a reduced temperature $T = 2$ and constant reduced pressure $P = 3.0$. (b) and (d) Weight functions (in units of $k_BT$) to flatten the corresponding probability distributions of $\lambda$ and to ensure that it is equally likely to have fractional molecules of reactants and reaction products for reactions (b) $A \rightleftharpoons B$ and (d) $A \rightleftharpoons D + E$.

Table 6. Chemical Potentials of Reactants and Reaction Products for Different Reactions of the Lennard-Jones System at Different Pressures Obtained with Serial Rx/CFC$^a$

| Reaction       | $P$ | $\sum_{\text{reactants}} \rho \mu_{i}^{\text{reactants}}$ | $\sum_{\text{products}} \rho \mu_{i}^{\text{products}}$ | $\sum_{\text{reactants}} \rho \mu_{i}^{\text{excess}}$ | $\sum_{\text{products}} \rho \mu_{i}^{\text{excess}}$ |
|---------------|-----|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
| $A \rightleftharpoons B$ | 0.3 | $-0.344(9)$ | $7.036(9)$ | $-0.344(6)$ | $7.036(6)$ |
|               | 1.0 | 0.07(1) | 9.42(1) | 0.066(6) | 9.42(6) |
|               | 3.0 | 2.73(1) | 12.95(1) | 2.727(6) | 12.953(6) |
|               | 5.0 | 5.23(1) | 15.74(1) | 5.23(2) | 15.73(1) |
|               | 0.3 | $-0.265(8)$ | $7.101(8)$ | $-0.133(6)$ | $7.101(6)$ |
| $A \rightleftharpoons C$ | 1.0 | 0.25(1) | 9.66(1) | 0.79(1) | 9.67(1) |
|               | 3.0 | 2.887(6) | 13.541(6) | 4.32(1) | 13.53(1) |
|               | 5.0 | 5.35(2) | 16.53(2) | 7.51(2) | 16.52(2) |
|               | 0.3 | $-0.34(1)$ | 6.13(1) | $-0.68(1)$ | 6.12(1) |
| $A \rightleftharpoons 2D$ | 1.0 | 0.07(1) | 9.49(1) | 0.13(1) | 9.48(1) |
|               | 3.0 | 2.73(1) | 13.79(1) | 5.43(4) | 13.74(2) |
|               | 5.0 | 5.23(2) | 16.84(2) | 10.45(3) | 16.74(2) |
|               | 0.3 | $-0.240(9)$ | 6.253(8) | $-0.205(9)$ | 6.245(9) |
| $A \rightleftharpoons 2E$ | 1.0 | 0.247(8) | 9.791(7) | 1.553(9) | 9.772(9) |
|               | 3.0 | 2.81(1) | 14.08(1) | 8.45(3) | 13.98(1) |
|               | 5.0 | 5.25(3) | 17.02(3) | 14.78(8) | 16.67(6) |
|               | 0.3 | $-0.340(8)$ | 4.319(8) | $-0.68(1)$ | 4.32(1) |
| $A \rightleftharpoons D + F$ | 1.0 | 0.07(1) | 8.30(2) | 0.13(1) | 8.29(1) |
|               | 3.0 | 2.724(9) | 13.246(6) | 5.44(2) | 13.213(6) |
|               | 5.0 | 5.23(1) | 16.57(1) | 10.45(4) | 16.52(2) |
|               | 0.3 | $-0.270(7)$ | 4.418(7) | $-0.41(1)$ | 4.43(1) |
| $A \rightleftharpoons D + E$ | 1.0 | 0.220(9) | 8.578(9) | 0.96(1) | 8.57(1) |
|               | 3.0 | 2.809(9) | 13.573(9) | 7.04(2) | 13.528(6) |
|               | 5.0 | 5.27(2) | 16.80(2) | 12.69(6) | 16.90(3) |

$^a$The reduced temperature is set to $T = 2.0$. The interaction parameters of different molecules are listed in Table 1. The numbers between brackets denote the uncertainty in the last digit.
6. CONCLUSION
An improved formulation of the Reaction Ensemble combined with Continuous Fractional Component Monte Carlo is presented (serial Rx/CFC). The main difference between serial Rx/CFC and parallel Rx/CFC is that in serial Rx/CFC either the fractional molecules of the reactants or the fractional molecules of the reaction products are present in the system. In serial Rx/CFC, there are three trial moves to facilitate a chemical reaction: (1) changing the value of $\lambda$, (2) reaction for fractional molecules, and (3) reaction for whole molecules. As a proof of principle, serial Rx/CFC is compared to the conventional formulation of RxMC and parallel Rx/CFC for systems of LJ molecules at different reduced pressures. Moreover, equilibrium mixture compositions obtained for the ammonia synthesis reaction using serial Rx/CFC are compared with experimental results and mixture compositions computed using equation of state modeling. The equilibrium compositions obtained with serial Rx/CFC are in excellent agreement with those obtained from the conventional RxMC and parallel Rx/CFC. For the ammonia synthesis reaction, excellent agreement between the results of serial Rx/CFC and experimental measured mixture compositions was found as well. For systems at high pressures, the acceptance probability of the reaction trial move is improved by a factor of 2 to 3 (depending on the system under study) compared to parallel Rx/CFC. Serial Rx/CFC has the following advantages: (1) One directly obtains chemical potentials of all reactants and reaction products. These chemical potentials can directly be used as an independent check to ensure that chemical equilibrium is achieved. (2) Independent biasing is applied to the fractional molecules of reactants and reaction products; therefore, the efficiency of the algorithm is increased. (3) Changes in the maximum scaling parameter of intermolecular interactions can be chosen differently for reactants and reaction products. Serial Rx/CFC can be easily extended to molecules with intramolecular degrees of freedom. The trial moves of Figure 3 can be performed by inserting fractional molecules at random positions with random orientations. The internal configuration of the molecule can be generated randomly or using the Rosenbluth scheme. The trial moves of Figure 4 can be performed by keeping the internal configuration of the molecule the same as in the old configuration. For ergodic sampling, trial moves that attempt to change the internal configuration of flexible molecules should be added to the MC method.32 The serial Rx/CFC method could also be used for reactions involving ions. One can calculate the potential energy of a periodic system with a net charge by placing a dummy charge at the center of charges. Although it is difficult to interpret computed partial molar properties of ions (such as the chemical potential or the partial molar volume) by using serial Rx/CFC, one can still benefit from other advantages of the method such as efficient reaction trial moves.
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