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Citation: The Journal of Chemical Physics 143, 044105 (2015); doi: 10.1063/1.4927084
View online: http://dx.doi.org/10.1063/1.4927084
View Table of Contents: http://scitation.aip.org/content/aip/journal/jcp/143/4?ver=pdfcov
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Monte Carlo simulation of dense polymer melts using event chain algorithms

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(Received 23 February 2015; accepted 9 July 2015; published online 23 July 2015)

We propose an efficient Monte Carlo algorithm for the off-lattice simulation of dense hard sphere polymer melts using cluster moves, called event chains, which allow for a rejection-free treatment of the excluded volume. Event chains also allow for an efficient preparation of initial configurations in polymer melts. We parallelize the event chain Monte Carlo algorithm to further increase simulation speeds and suggest additional local topology-changing moves (“swap” moves) to accelerate equilibration. By comparison with other Monte Carlo and molecular dynamics simulations, we verify that the event chain algorithm reproduces the correct equilibrium behavior of polymer chains in the melt. By comparing intrapolymer diffusion time scales, we show that event chain Monte Carlo algorithms can achieve simulation speeds comparable to optimized molecular dynamics simulations. The event chain Monte Carlo algorithm exhibits Rouse dynamics on short time scales. In the absence of swap moves, we find reptation dynamics on intermediate time scales for long chains.

I. INTRODUCTION

Polymer melts or polymer liquids are concentrated solutions of long chain molecules above their glass or crystallization temperature. In a dense polymer melt, long-range excluded volume interactions become screened and an individual polymer shows ideal behavior. Polymer melts exhibit a characteristic and complex dynamical and rheological behavior because of entanglement effects, which impede chain diffusion and give rise to reptation dynamics of polymer chains. The melt state is also most relevant for processing and manufacturing polymer materials.

In this paper, we introduce a novel Monte Carlo (MC) algorithm for the off-lattice simulation of a melt of flexible hard sphere polymer chains, which are connected by springs or tethers. This event chain (EC) algorithm allows for a much faster equilibration as compared to MC algorithms based on local moves.

The simulation of polymer melts by Molecular Dynamics (MD) or MC simulations is a challenging problem, in particular, for long chains at high density, where polymers in the melt exhibit slow reptation and entanglement dynamics. For chain molecules of length $N$, the entanglement time increases $\propto N^3$, which impedes the equilibration of long chain molecules in a melt if only local self-avoiding displacement moves of polymer segments are employed as in a typical off-lattice MC simulation. In order to reach equilibrium by such local moves, the system has to go through slow reptation dynamics on time scales between the Rouse and entanglement time.

In MD simulations, such reptation dynamics has been observed. In MC simulations, indications of reptation dynamics have been observed in lattice models or fluctuating bond lattice models. To our knowledge, reptation dynamics has not yet been observed in an off-lattice MC simulation so far, where equilibration is more difficult.

The dynamics of MC simulations depends on the MC moves that are employed. For local MC moves, the polymers obey Rouse dynamics on short time scales until entanglement effects eventually give rise to the crossover to reptation dynamics if MC moves obey the self-avoidance constraint. This means that the resulting MC dynamics can resemble the actual motion of coarse-grained polymers, although the MC dynamics is not explicitly based on a realistic microscopic dynamics. Local MC reptation moves (slithering snake moves) are used to initiate reptation dynamics and obtain faster equilibration of a polymer melt. MC simulations have the general advantage that also non-local or collective MC moves can be introduced, for example, chain-topology changing double-bridging moves, which speed up equilibration (such moves can also be combined with MD simulations to equilibrate the system). Dynamic properties, however, are no longer realistic if such topology-changing moves are employed. In particular, reptation dynamics will not occur if chain-topology changing moves are employed.

If polymers in a melt are modeled as bead-spring models with hard sphere beads, an additional simulation problem arises, in particular in MC simulations. At high segment or monomer densities, the mean free path of segments is limited and local MC displacement moves are restricted to very small step-sizes.

For hard sphere systems, non-local cluster moves represent a successful strategy to overcome the problem of slow MC equilibration in general by reducing rejection rates in the dense limit. In Ref. 22, the rejection-free event chain algorithm has been proposed, which coherently moves large clusters of polymer segments as in a typical off-lattice MC simulation. In order to reach equilibrium by such local moves, the system has to go through slow reptation dynamics on time scales between the Rouse and entanglement time.

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In MD simulations, such reptation dynamics has been observed. In MC simulations, indications of reptation
particles in the form of a chain, and a significant speed-up in the sampling of the hard sphere system has been shown. The EC algorithm can be generalized from athermal hard sphere systems to spheres with interaction potentials. In Ref. 25, we showed that the EC algorithm can be used for simulations of semiflexible bead-spring polymer systems. In this work, we adapt the EC algorithm for the MC simulation of dense polymeric melts consisting of flexible hard sphere polymers, verify the algorithm, and benchmark its performance.

The paper is structured as follows: in Section II, we present our EC based MC algorithm for hard sphere polymer melts. In order to further improve performance, we also introduce a parallelized version of the EC algorithm and a version employing local topology-changing “swap” moves. Furthermore, we show that EC moves can also be used to efficiently generate initial polymer configurations for the simulation, which are already representative of equilibrium configurations. In Section III, we verify our algorithm by a detailed comparison of equilibrium structural properties, such as the polymer shape and the end-to-end distance distribution, to simulation results from other MD and MC simulation techniques. Naturally, these results are not novel. Therefore, all details of this validation are presented in the Appendix. Finally, in Section IV, we benchmark the performance of serial and parallelized EC algorithms with or without swap moves against standard local MC schemes and against state of the art MD simulations (using the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) package). We use time-dependent mean-square displacements (MSDs) of polymer beads to monitor inter- and intrapolymer diffusions and use the intrapolymer diffusion to compare the performance of all algorithms in terms of a polymer relaxation time. The EC algorithm obeys Rouse dynamics on short time scales. Moreover, we show that in the absence of topology-changing swap moves and for long polymer chains, the EC algorithm exhibits reptation dynamics on intermediate time scales, before a crossover to chain diffusion on the longest time scales. We end with a conclusion and outlook.

II. EVENT CHAIN ALGORITHM FOR POLYMER MELTS

A very fundamental model for a self-avoiding flexible polymer is a bead-spring model, in which all beads interact via an excluded volume constraint, i.e., the polymers consist of hard impenetrable spheres, and the beads in one polymer are bonded with Hookean springs. The spring constant has to be sufficiently large as to enforce the impenetrability of polymers and avoid unphysically large bond stretching. In summary, we have a hard sphere interaction between all pairs of beads,

\[ V(r, r') = \begin{cases} \infty & \text{if} \ |r - r'| > \sigma \\ 0 & \text{else} \end{cases} \]

with the diameter \( \sigma \) of the hard spheres, and a harmonic stretching energy which, for a single polymer, can be written as

\[ H_{\text{bonds}} = \frac{k}{2} \sum_{i=1}^{N-1} (b_i - \sigma)^2. \]

Here, \( k \) is the spring constant, \( N - 1 \) is the number of bonds in a polymer (containing \( N \) beads), \( b_i \) is the length of the \( i \)-th bond, and the equilibrium length of the bonds coincides with the hard sphere diameter \( \sigma \) in our model. In our simulations, we chose bond stiffnesses \( k \sigma^2/b_kT = 30 \) such that thermal bond stretching remains weak with \( \langle b_i \rangle \approx 1.1 \sigma \). To simulate a polymer melt at a given density \( \rho \), we generate a system of \( M \) polymers in a cube of edge length \( L = 40 \sigma \), see Fig. 1. We employ periodic boundary conditions in all directions.

Alternatively, we also consider systems of hard sphere polymers bonded by tethers of maximal length \( b_{\text{max}} = 1.4 \sigma \) rather than springs.

In systems of dense hard spheres, standard Metropolis MC schemes based on local moves of individual spheres suffer from very slow sampling, as the move length is limited to roughly the mean free distance between spheres. This has been overcome by the introduction of suitable cluster moves, the so-called ECs. In an earlier work, we extended this approach to parallel computation and demonstrated how the EC algorithm can be applied to dense polymer systems. Because the EC algorithm moves dense regions of hard spheres or polymer beads coherently, it also mimics the essential features of the actual physical dynamics on a coarse time scale such as diffusion of polymer bundles. If the bonded beads in a polymer interact only via pair potentials, such as in the present example of hard sphere bead-spring polymers, a completely rejection-free simulation solely based on EC moves is possible.

In the EC algorithm, we first choose a total displacement \( \ell \), which is the same for all EC moves. For hard spheres, each EC move is constructed according to the following rule, which is also illustrated in Fig. 2(a).
1. Select the starting pivot bead for the EC move and a direction (which we call $v$) randomly. Initially, the remaining displacement is $x_{\text{max}} = \ell$.

2. Evaluate the largest possible displacement $x \leq x_{\text{max}}$ of the pivot bead in the chosen direction before it touches another bead. Move the pivot bead by $x$.

3. Continue the EC move at the new pivot bead, which is the hit bead. The remaining EC displacement $x_{\text{max}}$ is decreased by $x$.

4. Iterate by going back to step 2 until $x_{\text{max}} = 0$.

Then, the next EC move is started. The relevant computational step in the EC moves is the evaluation of the admissible displacement. In a system consisting of unbonded hard spheres, this is the distance to the bead hit first by the pivot bead while moving in the chosen direction.

The EC algorithm can be adapted to spheres with pairwise position-dependent interaction potentials. For each move of the pivot bead in an EC chain, an energy difference $\Delta E > 0$ is drawn according to the Boltzmann distribution. A displacement of the pivot bead that reduces the interaction energy is accepted (as in the standard Metropolis algorithm). A displacement increasing the energy is only partly executed, up to the point where the energy difference $\Delta E$ that has been drawn is reached or until the remaining EC displacement has been exhausted.

Now we consider the general situation that the pivot bead has several pairwise interaction energies. For each interaction partner $i$, the energy difference $\Delta E_i$ then defines a maximal displacement of the pivot bead $x_i \leq x_{\text{max}}(\Delta E_i)$. The largest possible displacement $x$ of the pivot bead is the minimum of all $x_i$, which shall be realized for an interaction partner $j$, i.e., $x = x_j = \min_i x_i$. The EC is then continued at bead $j$ as next pivot bead. For hard sphere interactions, this algorithm reduces to the standard EC collision rule.

Fig. 2(b) shows an example for hard sphere polymers bonded by springs. The attempted EC displacements of the green bead are in one of two classes: (i) the beads do not collide along the path (for an EC move in direction $v_1$) or (ii) the beads do collide along the path (for an EC move in direction $v_2$). In both cases, the energy stored in the bond reduces on the blue part of the trajectory, which, therefore, is always admissible, and increases on the red part, where the maximal admissible displacement is set by the “consumable” energy $\Delta E$ drawn from the Boltzmann distribution. Thus, the bond energy is only relevant for the maximal displacement if beads do not collide, because the other case is dominated by the hard sphere constraint. After displacing the green bead, the red bead becomes the pivot bead in both cases (i) and (ii).

We prefer to choose the direction $v$ of ECs randomly, which satisfies detailed balance. This can be relaxed, in principle, to other choices as discussed in Ref. 22 for hard sphere systems such that global balance is still satisfied. One particular simple choice, which can also be applied to the hard sphere polymers, is to start ECs only into three positive cartesian directions, which can gain a factor of approximately 2 in simulation speed (essentially by simplifications in the collision detection). This simplification is not efficient, however, if combined with the parallelization scheme discussed in Sec. II A, which decomposes the system into simulation cells and reflects the ECs on simulation cell boundaries rather than rejecting the whole EC move. For a cell decomposition with rectangular boundaries along cartesian directions, as it is usually used, EC moves started into cartesian directions will always reflect on themselves.

A. Parallelization

We use a parallelized version of this event-chain algorithm and refer to our earlier work for details of the parallelization. As discussed there, the parallelization requires a spatial decomposition of the system (which is changed in every sweep to ensure ergodicity) into simulation cells. This limits the displacement of each sphere to its respective simulation cell. For non-bonded hard spheres, this can be treated by reflection of non-admissible ECs at the cell boundaries. If a spring-triggered event occurs, where the bonded bead, which caused the event and would be the next pivot bead, is lying outside the current simulation cell, we proceed in a very similar manner, i.e., by reflection at the plane normal to the bond of the two participating spheres as illustrated in Fig. 2(c): the gray bead is rendered immotile because of the currently chosen spatial decomposition into parallel simulations cells. Therefore, the EC cannot be transferred to the gray bead at the occurrence of a spring “collision.” Then, the pivot bead (green) does not change, and the propagation direction is reflected as if there was a wall normal to the bond.

In this work, we use a spatial decomposition scheme different from a checkerboard partitioning: we use a rectangular tile-joint partition, where large tiles are separated by small...
The equilibration of polymer melts in simulations can be improved by generating initial configurations that are already representative of equilibrium configurations.\cite{21} Frequently used strategies consist in a slow compression of an equilibrated dilute solution\cite{9,10} or a "push-off" procedure, where the strongly repulsive steric interaction is switched on only after generating equilibrated configurations with a soft repulsive potential.\cite{21} For the hard sphere polymer melt, we propose an EC-based algorithm, which is conceptually similar to the slow push-off procedure in Ref. 21 for a Lennard-Jones melt.

The flexible polymers in the equilibrated melt are ideal but acquire an effective stiffness. The effective stiffness is characterized by a finite value of \( \langle \cos \theta \rangle \) (\( \theta \) being the bond angle).\cite{21} For a long ideal chain of bond length \( \sigma \), this results in a mean-square end-to-end distance \( \langle R^2 \rangle(N) = cN\sigma^2 \) with a parameter \( c \equiv (1 + \langle \cos \theta \rangle)/(1 - \langle \cos \theta \rangle) > 1 \),\cite{21} which depends on the short-range interaction between polymer beads. From long-run simulation data, we find \( c \approx 1.9 \) for a melt of long hard sphere chains.

In order to capture the effective stiffness already at the level of the initial configurations, we set up a system with randomly placed phantom polymers with vanishing hard sphere diameter and bond length \( \sigma \), which we grow as non-reversal random walks by restricting subsequent (unit) tangents to \( t_i \cdot t_{i+1} < \cos (\theta_{\text{max}}) \).\cite{21} For an otherwise uniform distribution of bond vectors, this leads to \( \langle \cos \theta \rangle = \cos^{2}(\theta_{\text{max}}/2) \). We choose \( \theta_{\text{max}} \) such that \( \langle R^2 \rangle/N \approx 1.90 \rho^2 \) holds in accordance with our long-run simulation data, see green and blue lines with small symbols in comparison to black line in Fig. 3.

We then introduce a finite excluded volume, but with a hard sphere diameter that is only a fraction of the target diameter \( \sigma \). This generates some "conflicts," i.e., overlapping spheres. We remove these conflicts by repeatedly starting EC moves into different directions from the overlapping spheres only, until the conflicting overlap has been removed. In these ECs, we ignore pre-existent overlaps so that the EC will not only be transferred to a bead the current pivot bead is not overlapping with. This procedure corresponds to locally "rattling" in the hard sphere system until enough space has been created around the overlapping bead to insert it. Once all conflicts for a given diameter are solved, we increase the diameter and continue iteratively until the target diameter \( \sigma \) is reached. The iterative growth of sphere diameters (which we call "slow push-off") due to conceptual similarity with Ref. 21 leads to a smaller change in the initial distribution of mean-square internal distances \( \langle R^2 \rangle(n) \) between two monomers with a chemical distance \( n \) along the chain (averaged over all chains), see curves with large symbols in comparison to corresponding curves with small symbols in Fig. 3. For comparison, we also generate initial configurations by a fast increase of \( \sigma \) (which we call "fast push-off" as in Ref. 21), see green curves in Fig. 3.

Configurations after the push-off should exhibit internal distances \( \langle R^2 \rangle(n) \) as close as possible to the equilibrium result as found by a long simulation run, see black line in Fig. 3. The initial configurations generated with slow push-off and the optimal value \( c \approx 1.9 \) (blue lines in Fig. 3) are indeed similar to the long-run simulation results. The fast push-off configurations (green lines) deviate with a maximum at intermediate \( N \), which takes a long time to equilibrate due to the slow reptation dynamics.\cite{21} Initial configurations generated from ideally flexible phantom chains (red lines) differ considerably.

Using the slow push-off we can initialize systems at (in principle) any geometrically possible density without resorting to configurations that are far from equilibrium (e.g., placing the beads on a lattice) in a reasonable amount of time (a couple of minutes wall time,\cite{27} for the system parameters below). Even for a very dense system with \( \eta = 0.63995 \), \( N = 120 \), and \( M = 275 \), we can generate initial configurations in \( O(100\, h) \) wall time. For such dense systems, however, these are only valid configurations, which are far from equilibrium because bonds are very elongated, and a thorough equilibration is still necessary.

C. Additional bead swapping

Typical conformations in a dense melt consist of highly entangled polymers. In the dense limit, the ECs become very long, i.e., the rather large displacement of an EC is distributed on a lot of very small displacements of many beads participating in the EC move. This results in a small collective translation of all beads participating in an EC cluster move with only small changes to the topology of entanglements.

Topology changing MC moves such as the double-bridging move\cite{20} can speed up equilibration in polymer melts significantly.\cite{9,10,21} Here, we improve sampling with EC moves further by introducing an additional swap MC move, which can locally change the topology of entanglements. In contrast to the double-bridging move, which changes bonds, the swap move changes topology by changing bead positions. For this purpose, we modify the EC move so that the EC does not directly transfer to the next bead upon hard sphere contact but, instead, a swap of the two touching spheres is proposed, see

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig3.png}
\caption{Mean-square internal distances \( \langle R^2 \rangle(n)/n\sigma^2 \) between two monomers with a chemical distance \( n \) along the chain for different initial condition generators and a long run simulation (thick black line). Small points are mean-square internal distances directly after setting up the phantom chains, corresponding large points after increasing the bead size to \( \sigma \), either by a "slow push-off" (red and blue curves) or by a "fast push-off" (green curve). The simulation parameters are \( N = 121 \), \( M = 500 \), \( L = 40\sigma \) (packing fraction \( \eta = 0.495 \)). We used ECs with a total displacement length \( f = 2\sigma \) for the push-offs.}
\end{figure}
The EC swap move is accepted according to the standard Metropolis algorithm. If the swap is rejected, the EC is transferred and the standard EC algorithm as described above is recovered. If it is accepted, the two beads are exchanged, and the EC continues with the same pivot bead. The example of a swap move in Fig. 4 shows a situation where it might be energetically favorable to swap beads. Note that in the absence of bonds, all beads become indistinguishable, and the EC algorithms with and without swapping are identical up to book-keeping differences.

The swap move is EC-specific: the EC automatically selects colliding pairs of beads for swapping; if the swap move is rejected, the EC move can continue without rejection of the entire EC move. Moreover, detailed balance is satisfied, and bead swapping can be included with very little computational overhead into the EC scheme. An analogous swap move in a standard MC algorithm needs to select pairs of beads such that the swap move has a reasonable acceptance rate (the particles have to be reasonably close). Moreover, the selection rule has to satisfy detailed balance (for example, simply proposing the nearest neighbor for swapping will lead to a violation of detailed balance). Therefore, there is no straightforward analogue of the EC swap move in a standard MC simulation with local moves.

Since the swap move locally changes topology and de-entangles polymers, the dynamics is no longer realistic if swap moves are applied. In particular, reptation dynamics is suppressed by swap moves (see numerical results below). On the other hand, this is the reason why swap moves can accelerate equilibration of the melt.

III. VALIDATION

In order to verify our algorithm, we address structural equilibrium properties of chains in a polymer melt by investigating their typical shape as characterized by the moment of inertia tensor and the distribution of end-to-end distance. These structural equilibrium quantities provide a detailed comparison across polymer melt simulation algorithms. The results of a comparison between different MC and MD simulation algorithms are shown in the Appendix. We find quantitative agreement between the EC algorithm and standard MC and MD algorithms, and agreement with previous MC simulation results and theoretical predictions, where available.

IV. PERFORMANCE AND DYNAMICS

For the comparison of the performance of different algorithms, we distinguish algorithms by whether they use the EC or standard Metropolis algorithm for (i) the hard sphere interactions and/or (ii) the bond spring interactions (“EC” for event chain and “MC” for standard Metropolis) and (iii) if the algorithm is executed parallelly (par) or serially (ser) and (iv) if the swap move is used (swap). Accordingly, we introduce a naming scheme for algorithms where, for instance, “EC-MC-par” refers to a parallelized simulation, where hard sphere interactions are handled by the EC, springs handled by standard Metropolis algorithm, and the swap move is not used.

We compare five different algorithms, namely EC-EC-par-swap, EC-EC-par, EC-EC-ser, MC-MC-par, and MC-MC-ser. This allows us to analyze the parallelization performance gains by comparing EC-EC-par-ser and check if we achieve the theoretical speed-up factor given by the number of processor cores. We do not parallelize the standard MC algorithm, because it was shown previously that strong scaling is achievable. The comparison of MC-MC-ser/EC-EC-ser gives the algorithmic speed-up by using the event chain algorithm. The comparison EC-EC/MC-par demonstrates the advantage of using the event chain on the pair potential, i.e., the bonds.

Additionally, we compare our results to those from MD simulations performed using the highly optimized LAMMPS package. As hard spheres cannot be used in a force-based MD simulation, we compare with beads that are interacting via the repulsive part of standard Lennard-Jones potentials, whereas the bonds remain Hookean springs. The identification of the effective hard sphere radius of such soft Lennard-Jones spheres has been subject of prior research. Our results show that identifying the Lennard-Jones length scale \( \sigma_{LJ} \) (defined by the zero of the full Lennard-Jones potential, \( V_LJ(\sigma_{LJ}) = V_LJ(\infty) = 0 \)) with the hard sphere diameter \( \sigma \) suffices for the purposes of this work. This comes with the advantage that we can use the same initial configurations (generated by our EC-based procedure) for the MD and MC evolutions.

A. Diffusional dynamics and algorithm speed

We compare the speed of different algorithms in terms of wall time. Since the simulations ran on different Central Processing Units (CPUs), all wall times were calibrated with short run simulations on the same workstation with four CPUs for comparable results.

We choose three different systems to investigate the influence of the occupied volume fraction \( \eta \) and chain length \( N \) on algorithm performance (we use the same systems for the validation of equilibrium properties in the Appendix):

1. System I: \( M = 400, N = 120, \) and \( \eta = 0.390 \);
2. System II: \( M = 500, N = 120, \) and \( \eta = 0.490 \);
3. System III: \( M = 250, N = 240, \) and \( \eta = 0.490 \).

FIG. 4. From left to right, we show EC moves with/without bead swapping. The currently selected pivot bead is highlighted by a red halo. The EC direction is along the vector connecting the middle beads of the two “polymers.” The initial EC (first column) is restricted by the hard sphere interaction of the middle beads, thus we use the Metropolis algorithm when the two beads touch (second column), to evaluate whether to swap beads (third column, upper row) or transfer the EC (third column, lower row). Finally, the remainder of the displacement is performed (fourth column).
This means the volume fraction $\eta$ increases from System I to System II, whereas the polymer length $N$ increases when going from System II to System III.

In the following, we will compare the performance of these algorithms by the inter- and intrapolymer diffusional behavior of polymer chains using time-dependent MSDs of polymer beads. For a chain with bead positions $\mathbf{r}_i$ ($i = 1, \ldots, N$) and center of mass $\mathbf{R} = \frac{1}{N} \sum_{i=1}^{N} \mathbf{r}_i$, we measure the MSD functions,$^{12,15}$

$$g_1(\Delta t) = \left( \langle (\mathbf{r}_{N/2}(t + \Delta t) - \mathbf{r}_{N/2}(t))^2 \rangle \right),$$

$$g_2(\Delta t) = \left( \langle (\mathbf{r}_{N/2}(t + \Delta t) - \mathbf{R}(t + \Delta t)) - (\mathbf{r}_{N/2}(t) - \mathbf{R}(t))^2 \rangle \right).$$

$g_1$ describes the diffusion of the middle bead including contributions from inter- and intrapolymer diffusion and $g_2$ the intrapolymer diffusion of the middle bead relative to the center of mass of the polymer. For both quantities, the average $\langle \ldots \rangle$, is an ensemble average and an average over time.

In a polymer melt, the time evolution is governed by a sequence of crossovers,$^{12}$

$$g_2(t) \sim \begin{cases} t^{1/2} & \text{for } t < \tau_e \\ t^{1/4} & \text{for } \tau_e < t < \tau_R \\ t^{1/2} & \text{for } \tau_R < t < \tau_d \\ t & \text{for } t > \tau_d \end{cases},$$

with three different crossover time scales: the entanglement time scale $\tau_e$, the Rouse time scale $\tau_R$, and the disentanglement time scale $\tau_d$. For all times scale $t > \tau_R$, reptation slows down the diffusional dynamics. The relative MSD $g_2$, exhibits the same regimes as $g_1$ but is insensitive to center of mass diffusion. For $t > \tau_d$, it approaches a plateau value given by the radius of gyration $R_g^2(N) = N^{-1} \sum_{i=0}^{N}(r_i^2 - \mathbf{R}^2)^2$.

Any simulation dynamics achieving equilibration of intrapolymer modes will reach the plateau in the relative MSD $g_2(t)$, beyond which intrapolymer fluctuations are equilibrated. We use the relaxation time to reach the plateau as a measure of simulation speed because it characterizes the equilibration performance of an algorithm on the scale of whole polymer chains. If the algorithm correctly describes the polymer melt dynamics on long time scales and exhibits Rouse, reptation, and chain diffusion dynamics as in Eq. (5), this relaxation time will coincide with the polymer disentanglement time $\tau_d$. In Fig. 5(a), we show the wall time evolution of $g_2(t)$ for different algorithms. Both MD (LAMMPS) and local MC dynamics follow Rouse dynamics with a $t^{1/2}$-behavior for short times.$^{11,13}$ Remarkably, we find such Rouse dynamics also for the cluster EC algorithm, even in the presence of swap moves. All algorithms approach a plateau in the relative MSD $g_2(t)$.

This also allows us to easily compare the performance of the algorithms and to determine a speed-up factor for each algorithm by rescaling time, i.e., shifting the double logarithmic curves such that the curves $g_2(t)$ coincide for long time scales close to the plateau. As a result, the single polymer relaxation time, which is identical to the disentanglement times $\tau_d$ if the algorithm exhibits all characteristic regimes of polymer melt dynamics, should be identical after rescaling. The resulting speed-up factors with respect to the standard local Metropolis algorithm MC-MC-ser are shown in Table I.

If we use these speed-up factors for a linear rescaling of the time, the data for both MSD functions $g_2(t)$ and $g_1(t)$ and from all algorithms collapse onto two “master curves” as shown in Fig. 5(b). The exception is the EC algorithm employing topology-changing swap moves. Because this collapse includes the MD algorithm, this provides evidence that both local MC dynamics and the cluster EC dynamics (in the absence of swap moves) evolve the system in a way that allows for an identification of “Monte Carlo time” (i.e., number of moves) with physical time.

For the comparison in Table I, we did not explicitly optimize the free simulation parameters like the total displacement length $\ell$ for an EC or the number of started ECs per

![FIG. 5.](image)

| Algorithm     | System I | System II | System III |
|---------------|----------|-----------|------------|
| MC-MC-ser     | 1        | 1         | 1          |
| EC-EC-par     | 9        | 7         | 7          |
| EC-MC-par     | 14       | 17        | 17         |
| EC-EC-par     | 31       | 25        | 28         |
| EC-EC-par     | 230      | 115       | 423        |
| LAMMPS (par)  | 330      | 625       | 770        |

TABLE I. Comparison of relative speed-up compared to the standard local Metropolis algorithm MC-MC-ser for different system parameters and algorithms. Both parallel EC and LAMMPS simulations were performed using four cores.
sweep in a parallelized simulation (see Ref. 25 for a detailed discussion). Nevertheless, it is obvious that all algorithms clearly outperform the standard MC algorithm. Without parallelization, the EC-EC-ser algorithm achieves speed-up factors up to 10 compared to the standard MC algorithm (MC-MC-ser). The parallelization gives an additional speed-up factor of 3.5...3.9 close to the theoretical limit of 4 given by the number of cores we used for the parallel simulation. We note that also standard MC algorithms could be parallelized such that this additional parallelization speed-up factor is not specific to the EC algorithms.

Despite these speed-up factors for the EC algorithm, the LAMMPS MD simulation is still the fastest algorithm. For the comparison in Table I, we used a parallel version of LAMMPS running on four cores. We note that LAMMPS is under development since the mid 1990s26 whereas our EC algorithm implementation, while adhering to general good practice rules for scientific computation, should still have room for optimization. In view of these preliminaries, the performance difference between the MD LAMMPS simulation and our fastest EC variant including swap moves seems very promising.

Table I also shows that the EC-MC algorithm gains some efficiency with respect to EC-EC algorithms with increasing η. In such dense systems, the springs are compressed to a value close to their rest length σ. Therefore, the rejection rate caused by the spring energy is rather low, such that the gain from the additional computational effort in the rejection-free treatment of springs is small in denser systems. Since the disentanglement time τd ∝ N^3 is strongly influenced by the chain length N, the efficiency of the swap algorithm increases with longer chains. MD performance does not decrease with density, whereas local MC and also EC performances depend on the displacement length ℓ, which decreases with density. This explains the performance differences if the density η is increased.

The speed-up factors in Table I characterize algorithm equilibration times based on the polymer disentanglement time τd. Alternatively, the autocorrelation time of the end-to-end vector can be used to characterize equilibration times.10,16,20 According to Ref. 31, these equilibration time scales are comparable for moves not changing the chain topology; the disentanglement time τd is to be preferred if topology-changing moves are employed that cannot relax density fluctuations (e.g., double bridging moves or our swap move).

B. Reptation dynamics

The good collapse onto master curves in Fig. 5(b) suggests that we can observe the same regimes of polymer melt dynamics in the EC simulation as in a MD simulation, at least if no swap moves are employed. Therefore, we investigated whether also a regime of reptation dynamics is observable with the EC algorithm.

The reptation regime for τc < t < τR is rather hard to observe in simulations of shorter chains, and one expects a slightly increased exponent g_1(t) ∝ t^x with 0.25 ≤ x < 0.4.32 For the chain lengths N = 120 used in Fig. 5(b), the intermediate reptation regime is not clearly visible. In lattice MC simulations, evidence for an intermediate reptation-like regime with a considerably slower increase than t^{1/2} has only been found in melts with long chains of length of N = 512.15

Therefore, we also simulated a smaller system with less (M = 20) but longer polymers (N = 500) at η = 0.298 and measured g_1(t) for EC-EC algorithms in comparison with MD simulations (using LAMMPS), see Fig. 6. For these longer chains, the MD simulations show a much more pronounced intermediate regime of slowed down dynamics. We find that this regime is increasing for stiffer polymer springs (k = 100k_B T/σ^2 as compared to k = 30k_B T/σ^2 in Fig. 6): with a small probability, chains can still cross via thermally activated bond stretching, which becomes less probable for stiffer springs. For stiff springs (k = 100k_B T/σ^2), the MD simulations exhibit a reptation regime with a time-dependence g_1(t) ∝ t^{0.3} close to t^{1/4} and in accordance with theoretical predictions and simulations in Ref. 32.

The parallelized EC-EC simulations show exactly the same dynamical regimes in the MSD function g_1(t) as the MD simulations, see Fig. 6. In each EC move, all beads that would collide successively during a short time interval in a MD simulation are displaced at once. This gives rise to a MC dynamics which is effectively very similar to the realistic MD. For the EC-EC simulations, we considered a spring constant k = 30k_B T/σ^2 and, instead of stiff springs, hard sphere polymers bonded by tethers of maximal length b_{max} = 1.4σ. In the tethered system, thermally activated bond crossing is absent similarly to a system with very stiff springs. Very similar to the stiff spring MD simulation, the tethered EC-EC simulation shows evidence of an intermediate reptation regime with g_1(t) ∝ t^{0.3}. To our knowledge, this is the first off-lattice MC simulation, where clear indications of reptation dynamics could be observed.

Fig. 6 also shows that the reptation regime is absent as soon as we employ additional disentangling swap moves in accordance with our expectation. Swap moves can thus be used to accelerate equilibration by effectively “switching off” the slow reptation dynamics.

V. CONCLUSION

We introduced novel efficient off-lattice MC algorithms for the simulation of dense polymer melts of hard sphere polymers, which are based on event chain cluster moves
previously known for hard sphere systems, see Fig. 2. These EC cluster moves allow for a rejection-free treatment of the excluded volume interaction in the polymer melt. We generalize the algorithm to also handle the spring interactions in polymer bond rejection-free.

In addition, we introduce an efficient procedure to generate initial configurations, which are representative of typical equilibrated configurations in polymer melts. Using EC “rattling,” we can generate initial configurations up to very high packing fractions (up to $\eta = 0.639\,95$).

We parallelize the event chain Monte Carlo algorithm and suggest additional local topology-changing swap moves, see Fig. 4, to further increase simulation speeds in melts.

We validated the EC algorithm by comparing equilibrium structural properties. In the Appendix, we show results for the polymer shape (Fig. 7) and the end-to-end distance distribution (Fig. 8), which are in quantitative agreement with other MD and MC simulation techniques.

We assessed the performance of the EC algorithm by measuring its equilibration speed using the relative mean squared displacement function $g_2(t)$ of a polymer bead in the middle of a polymer with respect to the polymer center of mass, see Fig. 5. This allows us to define a polymer relaxation time, which is specific to the algorithm and represents a measure for its equilibration speed. We find that the combination of EC moves and parallelization can increase MC simulation speeds by factors up to 30. If also swap moves are employed, MC simulation speeds become comparable to optimized MD simulations that we performed with the LAMMPS package for comparison.

Without swap moves, the dynamics of the EC algorithm appears to be very similar to MD. A simple rescaling of simulation times can collapse MD and EC simulation dynamics, see Figs. 5 and 6. The collective dynamics generated by the EC moves, which essentially displace all beads coherently that collide successively in a short time interval in a MD simulation, appears to be very similar to the MD collision dynamics.

Accordingly, in the absence of swap moves, the EC algorithm exhibits all dynamical regimes expected for polymer melts, i.e., Rouse, reptation, and chain diffusion dynamics. In particular, we can identify an intermediate reptation regime with a mean squared displacement function $g_2(t) \propto t^{0.3}$ close to $t^{1/4}$ in simulations of a system with long chains ($N = 500$), see Fig. 6. To our knowledge, this is the first off-lattice MC simulation, where reptation dynamics could be observed. If topology-changing swap moves are used, which disentangle polymer chain, reptation dynamics is absent in the EC algorithms.

Although we only presented results for the most simple case of a melt of flexible polymers with no interpolymer interaction other than excluded volume, the added value of EC algorithms should persist in more complex systems. For (bond) interactions that are not pair interactions, e.g., bending energies, rejection-free sampling in the way presented here is not possible. We have already shown in a previous work, however, that such bending energies can still be treated by proposing moves that are compliant with the hard sphere constraint by using ECs and then accepting (or declining) this move according to the standard Metropolis algorithm.

**ACKNOWLEDGMENTS**

We acknowledge financial support by the Deutsche Forschungsgemeinschaft (No. KI 662/2-1).

**APPENDIX: VALIDATION**

1. **Moment of inertia tensor**

The shape of a polymer in a dense melt is ellipsoidal rather than spherical, which can be shown by the distribution...
TABLE II. Comparison of shape descriptors from MC algorithms, LAMMPS, and theoretical expectations. For clarity, we only show the results from MC-MC-ser, EC-EC-swap-par, and LAMMPS algorithms; the other variants do not differ significantly.

| System | MC-MC-ser | EC-EC-swap-par | LAMMPS |
|--------|-----------|----------------|--------|
|        | I        | II             | III    | I     | II     | III    | I     | II     | III    |
| \((k^2)\) | 0.390    | 0.388          | 0.387  | 0.399 | 0.397  | 0.395  | 0.402 | 0.397  | 0.395  |
| \(<I_1 + I_2 - I_3>/\langle Tr(I)\rangle\) | 0.763    | 0.758          | 0.757  | 0.765 | 0.764  | 0.766  | 0.765 | 0.767  | 0.763  |
| \(<I_1 - I_2 + I_3>/\langle Tr(I)\rangle\) | 0.173    | 0.178          | 0.177  | 0.172 | 0.173  | 0.172  | 0.172 | 0.170  | 0.173  |
| \(<-I_1 + I_2 + I_3>/\langle Tr(I)\rangle\) | 0.0642   | 0.064 73       | 0.065 40| 0.062 84| 0.063 10| 0.062 51| 0.062 25| 0.063 02| 0.064 27|
| \(<4b^2 + 3c^2>/\langle Tr(I)\rangle^2\) | 0.653    | 0.621          | 0.629  | 0.640 | 0.638  | 0.672  | 0.646 | 0.643  | 0.633  | 0.667

2. Distribution of the end-to-end distance

The distribution of the end-to-end distance \(W(R)\) for an ideal chain with a mean-square end-to-end distance \(<R^2>(N) = cN\sigma^2\) (see Section II B for the definition of the stiffness parameter \(c\)) is approximately given by a Gaussian distribution:

\[ W(R)dR = 4\pi R^2 \left( \frac{3}{2pcN\sigma^2} \right)^{3/2} \exp\left( -\frac{3R^2}{2cN\sigma^2} \right) dR. \]

Therefore, the ideality of chains in a polymer melt can be tested by comparing simulation results for the distribution of the end-to-end distance with Gaussian expectation (A4), see Fig. 8. The agreement with the Gaussian expectation is indeed good, apart from an oscillating behavior at small distances \(R\). These oscillations can be explained by the influence of the pair correlation function \(g(r)\) characterizing the additional local liquid-like ordering of neighboring polymer beads. These oscillations are in quantitative agreement with \(g(r)W_L(R)dR\), where we determined the pair correlation \(g(r)\) of beads in the polymer melt numerically.

Also the agreement among the results for different simulation algorithms in Fig. 8 is very good. Only the standard serial MC-MC algorithm shows deviations because of its long equilibration times.
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