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Mobilité interfaciale dans les films polymères vitrifiables : est-elle déterminée par le mouvement collectif ou l'énergie intramoleculaire ?

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Interfacial Mobility in Glass Forming Thin Films : is it Determined by Collective Motion or by Intra-Molecular Energies ?

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Chapter 1

Introduction

This thesis is about the temperature dependence of dynamics in polymer melts, and how this mobility may change when polymer melts are confined to thin films. Let us start by discussing what polymer melts are about.

Polymer melts are systems that contain many macromolecules in form of a liquid [1, 2]. In the simplest case, macromolecules are linear chains comprised of N monomeric repeat units (number N is usually called a degree of polymerization or, simply, the chain length). The chains can be up to $N \lesssim 10^5$ long making them quite big objects with the gyration radius $R_{\rm g} \lesssim 1 \ \mu {\rm m}$, even though the size of a monomer is only ~ 5 Å. Such a large span of length scales is pronounced in the structural and dynamical properties of polymer melts. Being liquids, they show an amorphous dense packing of the monomers on a local scale and low compressibility on a macroscopic scale [3]. On the intermediate scales (up to $\sim R_{\rm g}$) the structure of the melt is determined by the intra- and intermolecular contributions. The former manifests in the fact that long polymers are self-similar fractal objects [2, 4] that have the ability to adopt different conformations. Such a chain structure allows the other chains to penetrate inside the volume it occupies leading to on average \sqrt{N} other chains inside the pervaded volume by one chain. These neighboring chains screen the intrachain excluded volume interactions which in a dilute solution would lead to swelling of the polymer [1, 2, 4]. Thus, to a good approximation the chains in a melt have a random-coil-like conformation with the size scaling as $R_{\rm g} \sim \sqrt{N}$ with the chain length. The dynamics of the melt also varies with the length scale: the (longest) relaxation time of a polymer τ_N is much larger than the monomer relaxation time τ_1 . For moderate N it is commonly described by the Rouse model [1, 2, 4, 5], that predicts $\tau_N = \tau_1 N^2$. Due to the slow relaxations, polymer melts are more viscous than low-molecular-weight liquids. Moreover, since the chains cannot cross each other, a network of topological constraints appears and persists during significantly long times when the chain length exceeds a certain value $N_{\rm e}$. These entanglements slow down the dynamics of the melt even more. The so-called "reptation" model describes this effect and predicts $\tau_N = \tau_1 N^3 / N_{\rm e}$ [1, 2, 4]. In addition to slowing down the relaxation in the time window $\tau_1 < t < \tau_N$, the increase of the chain length leads to a pronounced viscoelastic behavior of the melt even at high temperatures.

Upon cooling to low temperatures, polymer liquids undergo a transition to a solid state, which is either semicrystalline or glassy (amorphous) depending on the structure of the chain. Only polymers with side groups having regular tacticity [2] (e.g. isotactic/syndiotactic polystyrene) or without side groups (ex. polyethylene) can form crystalline lamellae. Still, the full crystallization is almost never achieved and the lamellar sheets exist along with amorphous regions [6, 7]. For atactic (homo)polymers (e.g. atactic polystyrene) or random copolymers (*cis-trans* polybutadiene, etc.) the crystalline phase does not exist. Upon cooling these polymer melts are in general good glass formers [8, 9]. Similarly to nonpolymeric glass-forming liquids, with decreasing the temperature towards a characteristic value $T_{\rm g}$, all structural relaxation times drastically increase over about 14 orders of magnitude (from ~ 1 ps at high T to ~ 100 s at $T_{\rm g}$) in a super-Arrhenius fashion. Below $T_{\rm g}$ the relaxation is so slow, that the system is indistinguishable from an amorphous solid. On the contrary, static two-point correlation functions, such as the static structure factor S(q), change mildly and below $T_{\rm g}$ the system has the structure of a "frozen" liquid, that has stopped to flow [10]. In addition to slowing down, another characteristic feature of the glassy dynamics is a two-step relaxation of the time correlation functions $C(t) = \langle A^*(t)A(0) \rangle^{-1}$ of the structure-sensitive variables A, with the first step attributed to the weekly temperature dependent microscopic relaxation and the second step corresponding to structural relaxation. The latter one is stretched over large time windows and has a strong dependence on temperature. Typical experimentally observed quantities that show this behavior are the coherent/incoherent density fluctuations measured in dynamic neutron scattering experiments [11]. These effects are also observed as broadening of the dominant loss peak in the frequency domain when measuring the relaxation of the

 $^{{}^{1}}A^{*}$ denotes the complex conjugation and the angular brackets $\langle \rangle$ denote the thermal average.

local electric dipoles in dielectric spectroscopy measurements [12].

Even though the behavior of glass forming polymer melts outlined above has been observed over many decades already [10], the description of the glassy dynamics at the present time still has rather an empirical character and the mechanisms of the glass transition phenomenon are not yet fully understood. An overview of theoretical perspectives on the glass formation can be found in Refs. [13, 14]. The situation is even more complicated for the polymer glasses in confinement to the nanoscale dimension, that have extensive industrial application (coatings, semiconductors, polymer nanocomposites) [15]. With the current trend to the miniaturization, it is important to understand the effects of confinement on mechanical stability. In particular, many studies on $\sim 10-100$ nm thick polymer films reported significant shifts in the average $T_{\rm g}$ with respect to the bulk value (see [10] and references therein and Fig. 1.1). These shifts were found to depend on the geometry of the film (capped/supported/freestanding), the chemical structure of the polymer and even the experimental technique employed in the investigation. Interpretation of a huge amount of data seems to reach a consensus in the fact, that these shifts are the average effect of local $T_{\rm g}$ gradients created by the confinement. In particular, the presence of a free surface for the supported and freestanding films is commonly assumed to speed up the dynamics of the close-by polymer layer leading to a decrease of the local T_g . The presence of a substrate for the capped/supported geometries is considered to have either a slowing down (increase of $T_{\rm g}$) or a speeding up (decrease of $T_{\rm g}$) effect depending on the nature of its interaction with the polymer and the roughness of the substrate. Yet, there is still an ongoing debate whether the results of the non-equilibrium and equilibrium experimental techniques can in principle be compared, since they might probe different properties [16].

While it is relatively straightforward to measure the film averaged properties using a wide amount of experimental techniques (ellipsometry, calorimetry, dielectric spectroscopy etc.), it is extremely challenging to access experimentally the dynamical gradients created by the surface effects, especially on a scale of the monomer size relevant to the glass transition process. Insights on the magnitude of the surface effects and how deep into the film the dynamics is perturbed (along with the general mechanisms of the structural relaxation) can be obtained from the molecular dynamics computer simulations [17]. Moreover, they allow to measure many different observables on the same sample providing



Figure 1.1: Number of publications vs. shift in $T_{\rm g}$ reported therein for the thinnest studied layer of polystyrene (the figure is taken from Ref. [16]). The collection is restricted to molecular weights in the range of 102–104 kg/mol and thicknesses between 5 and 50 nm; the employed experimental method and type of support are indicated. In total 92 publications are included (see [16] and references therein). The number of publications on each type of support is indicated in brackets after the type of support; 5 papers employing more than one type of support are recorded in each corresponding category. The increment in the y-axis representing one publication is always one which means that for a publication covering several methods the representative area is split into as many parts.

a link between different experimental techniques and allowing to interpret the vast variety of results.

Concerning the modelling of polymer systems, multiple approaches exist that differ in the level of chemical details. The most generic approach grasps only the basic features of polymers: chain connectivity, excluded volume interactions and, possibly, some chain stiffness [18, 19]. The polymers are modeled as chains of beads (connected by springs), each of them representing a part of the real chain, coarse-grained to the scale of a Kuhn segment [2]. The size of the segment is the only intrinsic length scale in the system. Being thus independent of chemical details, these generic bead-spring models are used to study universal properties of polymers [20, 21] including the mechanisms of glass formation [22– 24]. The more thorough quantitative comparison to real polymers requires a more detailed approach in the simulations using atomistic models [25, 26] that treat every atom present in the polymer (all-atom models) or combine few of them (typically carbon-hydrogen groups) together into united atoms (united atom models). In this case the details of chemical interactions matter, and in addition to the generic features mentioned above, the models include intramolecular potentials for the torsional angles and even sometimes the long range electrostatic interactions.

At the present moment most of the state of the art realistic simulations are able to follow the dynamics only up to few microseconds of real time. The bead-spring models win additional few orders of magnitude due to their simplicity. Although this time scale is much smaller than the relaxation time near $T_{\rm g}$ (which is about 100 s), simulations are appreciated [27] because they can provide a tomographic picture of the glass former and allow direct access to its properties on the relevant length scale. This ability is heavily used in the current research on the glass transition in general [28] and has also been exploited for systems in confinement [10, 18, 24, 29–31].

In general the simulation results corroborate the experimental findings [24, 31, 32]. Smooth, repulsive or weakly attractive substrates or free surfaces enhance the relaxation relative to the bulk. Whereas strongly attractive substrates cause the retarding effect. These interface-induced perturbations smoothly transition from the boundaries to the interior of the confined liquid. The range of these gradients grows on cooling so that the perturbations can propagate across the entire film for sufficiently low T and strong confinement. Contributions from all layers in these gradients determine the average behavior of the films and so the shift of $T_{\rm g}$. Qualitatively such a behavior is observed in the simulations using both realistic and coarse-grained models. However, the quantitative description obtained from the realistic models has been more and more often reported to disagree with the results extracted from generic simulations. The reasons of the disagreements have been attributed to the importance of the intramolecular forces, that are present in the realistic models in addition to the bead-spring-like interactions [33]. In particular, extensive molecular dynamics simulations have recently been carried out on glassy polymer films using an atomistic model for polybutadiene (PBD) embedded between two graphite walls [31, 34–36]. By calculating the dielectric response the simulations find, in qualitative agreement with experiments of the group of F. Kremer [37, 38], that confinement has no effect on the film dynamics, except in a layer of about 1 nm thickness near the substrate. It is argued that this result is divergent relative to the bead-spring

models and is caused by the dominance of intramolecular barriers. Since such barriers are present in all real polymers, references [31, 33, 35, 36] represent an important warning sign, challenging simulation efforts based on the generic models and the unifying physical picture emerging from these simulations for the glassy dynamics of polymers.

Clearly, this warning sign calls for a systematic study of how the intramolecular barriers and substrate interactions influence the glassy dynamics in atomistic simulations. Our work is focused on one particular type of the intramolecular interactions, the torsion (dihedral) barriers, that control the conformations of a polymer chain.

We start from simulations using the chemically realistic united atom model of PBD [26, 39] (described in Chapter 2) and make a crossover to the model where all the dihedral barriers were switched off, complementing previous studies of how they influence the structure and the glassy dynamics of the PBD melt [40–42] (Chapter 3). Then we employ these models in the simulations of supported PBD films (Chapter 4). Comparing between them will reveal how the interplay of intermolecular packing, intrachain barriers and surface effects influences the glass-forming properties of the PBD film. Chapter 5 provides a synthesis of the key results and conclusions of this work. The French summary of the thesis follows this conclusion chapter. Finally, a few appendices provide further technical details.

Chapter 2

Simulation model and computational details

2.1 Introduction

The computational technique used in our work to simulate polybutadiene (PBD) is classical molecular dynamics (MD) simulations [17]. In its essence, the algorithm integrates Newton's equations of motion for all particles in the studied system: the positions and velocities of the particles at the next timestep are calculated from the positions and velocities at the precedent timestep using the interaction potentials defined by the user. This set of interaction potentials along with the values of their parameters is commonly referred to as the force field. The velocities and/or simulation domain dimensions can be coupled to additional dynamic variables in order to model the effects of a thermostat and/or a barostat [43]. This determines the thermodynamic ensemble in which the coordinates and velocities are generated. The simulations last until a sufficient number of timesteps has been reached. During a simulation the coordinates and velocities of the particles are saved at selected timestep intervals into the so-called simulation trajectory, which is used during the post-analysis to compute the quantities of interest.

Our simulations were conducted using the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) [43, 44], which is a high performance computer code that handles parallel computing, provides extensive force field customization and a variety of tools for efficient computation of physical quantities during ongoing simulations. Additional benefits of using LAMMPS were the broad expertise and plenty of trajectory analysis tools that were developed at the research site.

There are 4 versions of the polymer model used in this work:

- The Chemically Realistic Chain (CRC) model is a modified version of the quantum chemistry-based united-atom model (from here on it is referred to as the original model) of 1,4-polybutadiene [26] developed by G. D. Smith and W. Paul in 1998 for MD simulations of polymer melts. The original model was tuned to reproduce the structure and dynamics of bulk PBD very well by extensive comparison with experimental data [39, 45, 46]. Later it was also used for simulations of confined films [30, 31, 35, 36, 47] using the GROMACS computer code [48]. However, due to the specifities of the simulations explained later in this chapter, this model cannot be used directly for the supported film geometry having free interfaces. Also, the direct transfer of some of the model features into LAMMPS is not possible. The CRC model developed by the author incorporates the modifications to account for these issues while maintaining the ability to reproduce the equation of state, the dynamics and the structure of polybutadiene well.
- The CRC2 and CRC4 models are derived from the CRC model by reducing the dihedral potentials by a factor of 2 and 4 correspondingly.
- The Freely Rotating Chain (FRC) model is also obtained from the CRC model by completely disabling the dihedral potentials.

Two types of polymer systems were studied in this work: a rather small system $(\sim 5 \times 10^3 \text{ united atoms})$ with periodic boundary conditions that models a bulk PBD and a quite large supported PBD film ($\sim 10 \text{ nm}$ thick with $\sim 10^5 \text{ united atoms})$ on top of a graphite substrate. All 4 models were used for the simulations of bulk PBD. In the case of films, only the CRC and the FRC models were employed, since such simulations are much more time and storage space consuming.

The following Section 2.2 presents the definition of the models, and Section 2.3 describes the simulation methodology. The parameters and comparison to the original CRC model are discussed in Section 2.4. The last Section 2.5 gives some information about technical details of processing the simulation trajectories.

2.2 The united-atom model of polybutadiene

A polybutadiene polymer chain has 2 types of carbon-hydrogen groups (Fig. 2.1a): CH and CH₂, that are represented by united atoms (UAs) with the masses $m_{\rm CH_2} = 14.027$ u and $m_{\rm CH} = 13.019$ u. A monomer consists of 4 UAs that form either a *cis* or a *trans* configuration (Fig. 2.1b). A polymer chain contains $N_{\rm m} = 29$ monomers or $N = 4N_{\rm m} =$ 116 UAs (only one chain length was studied in this work). For simplicity, no distinction between CH₂ groups within the chain and CH₃ groups at both ends of the chain is made. Our PBD chains are random copolymers with 55% *trans* and 45% *cis* monomers.

The force field of PBD is composed of four potentials associated with bond stretching, bond angle bending, torsional rotations and pair interactions.

The stretching of the bond length ℓ is modeled by a harmonic potential

$$U_{\rm bond}(\ell) = K_{\rm bond}(\ell - \ell_0)^2 ,$$
 (2.1)

where K_{bond} is (half) the force constant and ℓ_0 is the equilibrium bond length. There are 3 types of bonds in the system: CH-CH, CH₂-CH and CH₂-CH₂. They are plotted in



Figure 2.1: (a) Chemical notation of the 1,4-polybut adiene. (b) Schematic illustration of a 1,4-polybut adiene chain, which contains 116 $\rm CH_2$ and CH united atoms combined into 29 monomers.

Fig. 2.2 with the parameters given in Table 2.1 1 .

In the simulations the bond angle θ is calculated by the scalar product of the two consecutive bond vectors. Computationally, it is therefore convenient to express the bending potential in terms of $\cos \theta$. We used the harmonic in $\cos \theta$ bending potential

$$U_{\text{bend}}(\theta) = K_{\text{bend}}(\cos\theta - \cos\theta_0)^2 , \qquad (2.2)$$

with the K_{bend} being (half) the bending stiffness constant and θ_0 the equilibrium angle. There are 2 types of bond angles in the system that correspond to different possible combinations of the consecutive pair of bond vectors along the PBD chain: CH₂-CH-CH and CH-CH₂-CH₂ (cf. Fig. 2.3 and Table 2.2). The scale of the potentials in Fig. 2.3 is quite big with respect to the value of the thermal energy $k_{\rm B}T \approx 0.7$ kcal/mol at the highest studied temperature. Thus the fluctuations of the bond angle only happen in a small region around the equilibrium values. We will discuss this later in Section 2.4.2.

The torsion (or dihedral) angle ϕ is the angle between two intersecting planes defined by three successive bonds along the chain backbone. The associated potential $U_{\text{tor}}(\phi)$ accounts for steric interactions between UAs separated by these three bonds. Since $U_{\text{tor}}(\phi)$ is an even and periodic function of ϕ (0 °≤ ϕ ≤ 360°), it can be expressed as an expansion in powers of cos ϕ . We employ the Ryckaert–Bellemans function [49] in our simulations



Figure 2.2: The bond stretching potential Eq. (2.1).

Table 2.1: Parameters for the bond potential of Eq. (2.1).

Bond length	$K_{\rm bond}$	ℓ_0
type	$(\text{kcal/mol}Å^2)$	(Å)
CH-CH	2500	1.34
CH ₂ -CH ₂	2500	1.53
CH-CH ₂	2500	1.50

¹The high value of the force constant is a compromise between the experimentally observed bond vibrations and the constrained bonds of the original model, see also Section 2.4.



Table 2.2: Parameters for the bending potential of Eq. (2.2). They are identical to the original model [26].

Bond angle	$K_{\rm bend}$	θ_0
type	(kcal/mol)	(deg)
CH ₂ -CH-CH	68.1475	125.896
$CH-CH_2-CH_2$	66.5925	111.652

Figure 2.3: The bond angle bending potential Eq. (2.2).

using the CRC/CRC2/CRC4 models

$$U_{\rm tor}(\phi) = \sum_{n=1}^{6} A_n(\cos \phi)^{n-1} . \qquad (2.3)$$

There are 5 types of dihedral potentials in the system, that are depicted in Fig. 2.4 with the parameters $\{A_n\}_{n=1,\dots,6}$ given in Table 2.3. We have adopted the same naming as in Ref. [26].

The double bond *cis* and *trans* potentials (Fig. 2.4b) determine the rotation along the double (CH-CH) bond axis, and therefore, the configuration of the PBD monomer. Per definition, there is only one minimum, that occurs at $\phi_{\min} = 0^{\circ}$ (or 360°) for the *cis* and at $\phi_{\min} = 180^{\circ}$ for the *trans* potential.

The β torsional potential determines the rotation along the axis of the CH₂-CH₂ bond between two adjacent monomers. It has a shape familiar from hydrocarbon chains, such as polyethylene [2]. There are three minima, the primary being the *trans* state at $\phi = 180^{\circ}$ and the secondary minima being the *gauche-plus* and *gauche-minus* states at $\phi \simeq 180^{\circ} \pm 120^{\circ}$

The α cis and α trans potentials correspond to the CH-CH₂ bond that is next to the double bond. Qualitatively, the shape of $U_{\rm tor}(\phi)$ for these dihedrals is the mirror image of the β potential, having minima at $\phi = 0^{\circ}, 360^{\circ}$ and $\phi \simeq 180^{\circ} \pm 60^{\circ}$.

The *cis* and *trans* potentials (both double bond and α) are randomly assigned to the monomers in a PBD chain with the proportion of 45% *cis* and 55% *trans*.

The values of the parameters $\{A_n\}_{n=1,\dots,6}$ in Table 2.3 correspond to our chemically



Figure 2.4: (a) Schematic illustration of the torsion (or dihedral) angles in a fragment of a PBD chain. (b-d) Torsional potential $U_{tor}(\phi)$ (in kcal/mol) for the different torsion angles ϕ illustrated in panel (a).

Table 2.3: Coefficients A_n (in kcal/mol) of the torsional potential of Eq. (2.3) in the CRC model for different dihedral angles of PBD indicated in Fig. 2.4.

Torsion angle	A_1	A_2	A_3	A_4	A_5	A_6
$\alpha \ (trans)$	0.385179	3.466610	1.050490	-3.86178	-0.320148	-0.720332
$\alpha \ (cis)$	1.160540	0.360163	0.680313	1.48069	-0.640294	-3.041420
β	-3.261500	-4.942280	0.340155	5.58258	0.280129	2.000930
db $(trans)$	24.011000	5.316470	0.100537	27.54850	-0.160949	-8.914360
db (cis)	24.011000	-5.316470	0.100537	-27.54850	-0.160949	8.914360

realistic chain (CRC) model of PBD. As it was mentioned in Section 2.1, we also study model variants with reduced torsional potential by replacing $U_{tor}(\phi)$ for all dihedral angles with $\lambda U_{\text{tor}}(\phi)$ where $\lambda = 0, 1/4, 1/2, 1$. We refer to these models as:

$$\lambda = 0 : \text{ freely rotating chain (FRC) model},$$

$$\lambda = \frac{1}{4} : \text{CRC4 model},$$

$$\lambda = \frac{1}{2} : \text{CRC2 model},$$

$$\lambda = 1 : \text{CRC model}.$$
(2.4)

The rescaled $(\lambda = \frac{1}{2}, \frac{1}{4})$ or disabled $(\lambda = 0)$ torsional potentials is the only difference between the models. The FRC model was already introduced and studied in Refs. [40, 41]. Due to the absence of torsional potential the model enables free rotation around the double bonds, thereby eliminating the difference between *cis* and *trans* conformers. Within the FRC model PBD is therefore no longer a random copolymer of *cis* and *trans* monomers, but a regular copolymer of CH and CH₂ units. While it would have been possible to preserve the random *cis/trans* copolymer character by maintaining the dihedral potentials for the double bonds only [50], we have adopted the FRC model here because it resembles extensively studied glass-forming bead–spring models with intramolecular forces resulting only from bond-length and bending potentials² [33, 51–54]. The CRC4 and CRC2 models are introduced because they interpolate between the FRC and CRC models.

Pair interactions between united atoms of a chain separated by four or more bonds and between united atoms of different chains are modeled by a 12-6 Lennard–Jones (LJ) potential with an additional switching function S(r) commonly used in the GROMACS code [55], but also implemented in LAMMPS (see pair_style lj/gromacs command [43]). The potential reads:

$$U_{\text{pair}}(r) = \begin{cases} 4\epsilon \left[\left(\frac{\sigma}{r}\right)^{12} - \left(\frac{\sigma}{r}\right)^6 \right] + S(r) & \text{if } r < r_{\text{c}} ,\\ 0 & \text{if } r \ge r_{\text{c}} , \end{cases}$$
(2.5)

with

$$S(r) = \begin{cases} C & \text{if } r < r_1 \ , \\ \frac{A}{3}(r - r_1)^3 + \frac{B}{4}(r - r_1)^4 + C & \text{if } r_1 \le r < r_{\rm c} \ , \end{cases}$$
(2.6)

²Though, the bond length of the FRC model is much shorter than in the bead-spring models.

and

$$\begin{split} A &= \frac{-3U'_{\rm term}(r_{\rm c}) + (r_{\rm c} - r_{1})U''_{\rm term}(r_{\rm c})}{(r_{\rm c} - r_{1})^{2}} , \\ B &= \frac{2U'_{\rm term}(r_{\rm c}) + (r_{\rm c} - r_{1})U''_{\rm term}(r_{\rm c})}{(r_{\rm c} - r_{1})^{3}} , \\ C &= -U_{\rm term}(r_{\rm c}) + \frac{1}{2}(r_{\rm c} - r_{1})U'_{\rm term}(r_{\rm c}) \\ &- \frac{1}{12}(r_{\rm c} - r_{1})^{2}U''_{\rm term}(r_{\rm c}) . \end{split}$$
(2.7)

where $U_{\text{term}}(r_{\text{c}})$ stands either for the repulsive term, $4\epsilon(\sigma/r)^{12}$, or for the attractive term, $-4\epsilon(\sigma/r)^6$, of the LJ potential, σ is the zero-crossing distance $(U_{\text{pair}}(r=\sigma)=0)$, ϵ is the value of the minimum of the potential at $U_{\text{pair}}(r=2^{\frac{1}{6}}\sigma)=-\epsilon$. Prime and double prime symbols denote respectively the first and second derivatives with respect to r. The switching function obeys the following boundary conditions:

$$\begin{split} S'(r_1) &= S''(r_1) = 0 , \\ S(r_c) &= -U_{\rm term}(r_c) , \\ S'(r_c) &= -U'_{\rm term}(r_c) , \\ S''(r_c) &= -U''_{\rm term}(r_c) . \end{split} \tag{2.8}$$

For $r < r_1$ Eqs. (2.5) and (2.6) imply that $U_{\text{pair}}(r)$ is just a shifted LJ potential, while for $r_1 \leq r < r_c$ the LJ potential and its 1st and 2nd derivatives are progressively smoothed to zero due to the form of S(r) and the boundary conditions.

We choose identical values, $r_1 = 9$ Å and $r_c = 12$ Å, for all pairs of UAs. The LJ parameters used in the present study are summarized in Table 2.4 and the corresponding potentials are plotted in Fig. 2.5

In the case of supported film, the united atoms of PBD interact with a substrate, that is modeled using the crystalline lattice of graphite atoms with mass $m_{\rm CH_2} = 12.011$ u. The LJ parameters for the graphite were taken from the work [30], which in turn has taken the values reported in [56]. They are also given in Table 2.4 and Fig. 2.5.

Only from the pair interactions point of view PBD corresponds to a binary mixture (see Section 3.3). The values in Table 2.4 show that PBD deviates from the standard Lorentz–Berthelot mixing rule [3]. By contrast to the Lorentz rule, PBD is a nonadditive mixture, since $\sigma_{ij} \neq (\sigma_{ii} + \sigma_{jj})/2$, and has enhanced mixing ability relative to the Berthelot rule because $\epsilon_{ij} > \sqrt{\epsilon_{ii}\epsilon_{jj}}$. In simple liquids and metallic alloys both features are favorable

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for better glass-forming ability [57, 58]. In the case of the interaction of PBD with the graphite, the Lorentz-Berthelot combining rules were used.

Our force field is different from the one used in the original works [26, 30, 31, 35, 36, 39, 45–47], that used the GROMACS code, in two aspects. In these works the bonds were constrained and the plain LJ potential truncated at $r_c = 9$ Å with long-range tail corrections to energy and pressure was used for pair interactions. The reasons for these differences are discussed in Section 2.4.

2.3 The simulation methodology

2.3.1 General information

We perform molecular dynamics (MD) simulations of PBD with the Large Scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) [17, 43, 44]. The classical equations of motion are integrated by the rRESPA multi-timescale integrator with two hierarchical levels. The outer level integrates the pair interactions with a timestep $\delta t_{outer} = 1$ fs, while the inner level integrates the bond-length, bending and dihedral potentials with a four times smaller timestep $\delta t_{inner} = 0.25$ fs. We combine the rRESPA integrator with the Nosé–Hoover thermostat and barostat to control temperature T and pressure p in NPT simulations and with the Nosé–Hoover thermostat when carrying out canonical NVT simulations at constant volume V (LAMMPS parameters [43]: Tdamp = 1000 fs, Tchain = 3; Pdamp = 10000 fs, Pchain = 3). During the simulations of the PBD melt the



Figure 2.5: The Lennard–Jones potential Eq. (2.5).

Table 2.4: Parameters of the Lennard–Jones potential Eq. (2.5) for the different types of interaction sites: $i, j = CH_2, CH, GRA$.

Pair type	ϵ_{ij}	σ_{ij}
	(kcal/mol)	(Å)
$CH \leftrightarrow CH$	0.114999	3.38542
$\mathrm{CH}_2\leftrightarrow\mathrm{CH}_2$	0.107639	4.00904
$\mathrm{CH}_2\leftrightarrow\mathrm{CH}$	0.116723	3.79256
$GRA \leftrightarrow GRA$	0.055640	3.40000

pressure equal to p = 0 atm was imposed under the NPT conditions and the corresponding volume V was imposed under the NVT conditions. In this way, the melt can be compared to the simulations of the supported films in contact with vacuum at the free surface.

The initial configurations at T = 353 K for the bulk and the film systems were taken from the work [36], which employed the implementation of the realistic model (with all dihedrals) in GROMACS code [55]. These configurations were converted to LAMMPS input files by means of a custom-made script using a conversion 1 kcal/mol = 4184 J/mol for the energy units. The bulk system was used to establish the parameters of our models in LAMMPS and compare to the original GROMACS simulations (see Sections 2.3.2 and 2.4). The film systems were simulated using our models from the beginning. We provide a detailed map of the conducted simulations in Figs. A.1 and A.2 in Appendix A.

The simulations were conducted on the High Performance Computing (HPC) Center of the University of Strasbourg [59] and the Jean Zay (HPE SGI 8600) supercomputer at the Institute for Development and Resources in Intensive Scientific Computing (IDRIS, Orsay) [60].

2.3.2 Bulk details

The bulk system contains $N_c = 40$ PBD chains in a cubic box of about 50 Åin size (Fig. 2.6). Periodic boundary conditions were applied in all directions. The initial configuration at T = 353 K was converted from GROMACS to LAMMPS format by means of a custom-made script, that properly converts the units and replaces constrained bonds by the harmonic potential.

In order to verify if the model was transferred from GROMACS to LAMMPS correctly, our first simulations were conducted using the truncated LJ potential and tail corrections used in both codes. This allowed us to make a comparison to the original GROMACS simulations, which used such a potential in the first place. From now on, we refer to this model as the 'original model'.

Fig. A.1 presents a detailed map of the simulations using the original model. Aiming to compare with the works [40, 41], two working temperatures were chosen: T = 353 K and T = 273 K and the equilibration time was set to 100 ns. At each working temperature, the following simulation protocol was adopted (Fig. 2.7).

• Starting from the initial configuration, the equilibration is performed under the NPT



Figure 2.6: The bulk system in a periodic simulation box.

conditions over 100 ns. This duration suffices to relax the volume, since we find the volume relaxation to last no longer than 30 ns for the studied temperatures. The final 70 ns are then used to determine the equilibrium volume V(T) as the time average over the time series of the volume.

- In the subsequent deformation stage we start from the instantaneous volume of the final configuration of the NPT run and isotropically change the size of the simulation box until the equilibrium volume V(T) is reached (fix deform command in LAMMPS [43]). The deformation is carried out over a short time of 10 ps.
- Due to the fast deformation, it is necessary to relax residual stresses possibly created by the quenching of the box volume, even though the deformation is small (≤ 0.1 Å). This is done over another 100 ns of the NVT run at fixed V(T), which suffices to stabilize the average pressure at p = 0 atm.
- After equilibration we perform a NVT production run that lasts at least 1 μ s (10⁹ timesteps).

At T = 353 K the initial (t = 0) configuration is the one converted from the work [36]. In order to obtain the initial configuration at T = 273 K, the system was continuously cooled down to T = 113 K at imposed pressure p = 0 at according to the ramp

$$T(t) = T_0 - \Gamma t , \qquad (2.9)$$

with the cooling rate $\Gamma = 20$ K / 6 ns = 3.33 K/ns (Fig. 2.7). Then the configuration that corresponded to the time point with T = 273 K was used as the starting one (t = 0) for constant temperature NPT equilibration.

In order to check the density of PBD, the NPT equilibration at T = 353 K was continued up to 1 μ s and the configurations at t = 200 ns, 400 ns, 600 ns, 800 ns, 1 μ s were used to start 9 additional cooling runs: 4 runs with $\Gamma = 3.33$ K/ns and 5 runs with $\Gamma = 0.83$ K/ns. The t = 200 ns point was also used to launch a cooling run of the system where all dihedral potentials were disabled³. This is motivated by the results of Honnell *et al.* [61], who derived that the pressure equation for chain molecules does not explicitly depend on the angular potentials, and the works [40, 41], where the dynamics without the dihedrals was reported to be orders of magnitude faster than with the dihedrals while the structure is identical. Thus, we expect the density at p = 0 atm to be indistinguishable between the realistic model and the model without the dihedrals within the fluctuations at studied temperatures. We also use this argument later for the initial configurations of our CRC2, CRC4 and FRC models.

From one of the slower cooling runs, we then followed the protocol described above for an additional set of working temperatures: T = 293, 253, 240, 225 K and launched the constant temperature NPT equilibrations. At the same time we set up the simulations



Figure 2.7: Sketch of the bulk simulation protocol consisting of a cooling run according to Eq. (2.9) followed by the equilibration and production runs. Starting from the melt configuration stored during the cooling run at a given T (here T = 273 K), equilibration comprises three steps: 1) NPT relaxation over 100 ns with determination of the equilibrium volume V(T) as the time average over the final part of the time series of the volume, 2) the instantaneous volume of the final configuration of the NPT run is rapidly deformed over 10 ps toward V(T), 3) NVT relaxation at V(T) over another 100 ns. Equilibration is followed by a production run which lasts at least 1 μ s depending on temperature.

 $^{^{3}}$ This is not yet our FRC model, since the pair interactions are still described by the truncated LJ potential with the tail corrections.

of supported films and developed the modified pair potential to yield equivalent result in inhomogeneous systems (free surface!) without tail correction (see Section 2.4.3 for detailed explanation). We used the system at T = 273 K to launch the runs with modified pair interactions to develop our CRC model. The aim of the modifications was to match the densities of the systems at the studied temperatures without having to resort to the tail corrections. The new pair potential was verified to recover desired density by simulating a system at T = 353 K and performing the same set of cooling runs using our CRC model.

Due to the indistinguishable densities and for time saving purposes, we used the equilibrium volumes V(T) obtained using the original model to continue our simulation protocol and performed the NVT equilibrations and the production runs using already our CRC/CRC2/CRC4/FRC models. As a last check, we also performed long (1 μ s) NPT runs at each temperature using all four models to make sure that the densities are indeed the same.

Additionally, the system at T = 213 K was simulated using all our models in a similar manner as the higher temperatures, with an exception that the NPT equilibration has been performed using our CRC model. In addition, the systems at T = 153, 113 K were simulated using the FRC model following the protocol described above..

2.3.3 Film details

The initial film configuration was taken from the work [36], where the film comprising 720 PBD chains (83520 UAs) was confined between two layers of graphite atoms at the bottom of the simulation box and one layer at the top with periodic boundary conditions in all directions. In order to make the film supported on one side with free surface on the other side, we have added about 10 nm of vacuum above the top graphite layer, then moved the whole system of PBD and graphite up by 3.4 Å (the distance between the graphite layers). Finally the top layer of graphite was moved to the bottom of the simulation box, so all three layers are together and separated by 3.4 Å. A snapshot of the box is presented in Fig. 2.8.

All film simulations are under NVT conditions. The presence of the free surface of the film in contact with vacuum effectively corresponds to the simulations at zero pressure. The periodic boundary conditions were used in all directions. In the lateral dimensions the simulations box is commensurable with the (0,0,1) graphite crystalline surface. During



Figure 2.8: The film system in a simulation box.

the simulations all graphite atoms are fixed at their equilibrium positions in the crystal lattice. In the z-dimension, the 10 nm of vacuum is sufficient for the united atoms of PBD not to interact with their periodic images.

The simulations of the films were performed using our CRC and FRC models. During the conversion of the initial configuration, the pair interactions were replaced by the ones from our models. The film simulation protocol is similar to the bulk one and is presented schematically in Fig. 2.9

- The initial configuration at T = 353 K is equilibrated over 100 ns.
- Then the cooling run is performed according to the ramp given by Eq. (2.9) with the cooling rate $\Gamma = 0.83$ K/ns.
- The cooling configurations that corresponded to the time points of the working temperatures T = 273, 253, 240, 225, 213 K were used as starting ones for the constant temperature equilibration and production runs.

In fact, there is one long constant temperature run at each working temperature. The equilibration and production parts are distinguished simply by the time point, from which we start measuring the quantities of interest. The same is valid for the systems at T = 353 K: the configuration after 100 ns of equilibration using the CRC model was copied to launch the cooling (and also the FRC equilibration) and the run was continued. The constant temperature runs lasted 1 μ s in total in the case of the CRC model. The FRC

runs at the same working temperatures lasted 700 ns. As in the bulk, the two additional temperatures T = 153 K and T = 113 K were performed using the FRC model, that lasted also 1 μ s. The detailed map of our simulations is given in Fig. A.2 in Appendix A.

2.4 Model parameters and previous works

2.4.1 General information

As it was mentioned previously, our CRC model has several differences in the force field with respect to the model used in recent works [30, 31, 34–36, 47]. These works were focused on polymer films and, in turn, also employed an adapted version of the original potential functions [26, 39] for implementation in the GROMACS code. Since our study is also focused on films, we prioritized the recent version of the model [30, 31, 34–36, 47] when implementing our simulations in LAMMPS for both bulk and film systems. We have also inherited the values for the timestep $\delta t = 1$ fs and the thermostat damping time Tdamp = 1000 fs. For the damping time of the barostat, we chose a ten times larger value Pdamp = 10000 fs as it is suggested by the LAMMPS documentation [43]. The default (in LAMMPS) number of chained thermostats and barostats was used: Tchain = 3, Pchain = 3.



Figure 2.9: Sketch of the film simulation protocol consisting of a cooling run according to Eq. (2.9) followed by the equilibration and production runs.

2.4.2 Intra-chain interactions

In order to verify if the intra-chain interactions were transferred from GROMACS to LAMMPS correctly, we set up rather short simulations of the bulk PBD using the original GROMACS scripts obtained from the authors of the work [36] using the same initial configuration and compare them to our LAMMPS simulations. Following the results of [62], our objective was not to obtain an exact match of the energies and particle movements during both simulations (it is impossible due to the difference in the software features discussed later), but rather to get a qualitative agreement of the mean values of the energy terms and a quantitative agreement of the representative structural and dynamical properties of PBD. At the moment of performing the comparison the pair interactions had not been modified yet. The truncated LJ potential with tail corrections was used with the same parameters in both MD engines.

The main difference in the intra-chain interactions between our LAMMPS simulations and the original GROMACS simulations is in the bond potential. All bonds between united atoms of PBD were constrained to their equilibrium values ℓ_0 in GROMACS. LAMMPS does not allow to constrain the backbone bonds of a chain in order to achieve good parallel performance. However, we wanted to stay close to the original simulations with $\ell = \ell_0$, therefore we used a harmonic bond potential instead with a large value for the force constant $K_{\text{bond}} = 2500 \text{ kcal/(mol Å}^2$ (cf. Eq. (2.1) and Table 2.1). This value is a compromise between spectroscopic data [63] and not too long computation times. With this value one finds $\sqrt{\langle (\ell - \ell_0)^2 \rangle} = \sqrt{k_{\rm B}T/(2K_{\rm bond})} \approx 0.01$ Å for T = 353 K, the highest studied temperature. Thus, bond length fluctuations are less than 1% of ℓ_0 . The estimation of the smallest bond oscillation time in the system as the period of oscillation of a bond between two CH united atoms (the atoms with the smallest mass) results in $\tau_{\rm bond} = 2\pi \sqrt{m_{\rm CH}/(2K_{\rm bond})} \approx 16$ fs, which is only 16 GROMACS timesteps. Thus we used a rRESPA integrator with two hierarchical levels as it was explained previously in Section 2.3.1: the outer level integrates the pair interactions with a timestep $\delta t_{\text{outer}} = 1$ fs, while the inner level integrates the bond-length, bending and dihedral potentials with a four times smaller timestep $\delta t_{\text{inner}} = 0.25$ fs.

Fig. 2.10 presents the bond length distribution of all bond types obtained in our LAMMPS simulations at T = 353 K using two values of the force constant in the harmonic bond potential: 2500 kcal/(molÅ²) (our final value) and 1000 kcal/(molÅ²) (in between



Figure 2.10: Bond length distributions measured during the simulations using the force constants $K = 1000 \text{ kcal/(molÅ}^2)$ (red squares) and $K = 2500 \text{ kcal/(molÅ}^2)$ (blue circles). The vertical lines mark the equilibrium values of ℓ_0 . The black solid lines are the sums of Eq. (2.10) calculated for each bond type and normalized by the number of the corresponding bonds in the system.

the values converted from the experimental bond stretching frequencies $\nu_{\text{C-C}} \approx 1000 \text{ cm}^{-1}$ and $\nu_{\text{C=C}} \approx 1650 \text{ cm}^{-1}$ [63]). For the latter, the peaks at $\ell_0 = 1.5$ Å and $\ell_0 = 1.53$ Å are merged into one peak in between. Even though this is probably closer to the experiment, our intention at this point was to introduce as little deviation to the model as possible and to reduce the influence on the other potentials for better comparison with the constrained system. We have compared the distribution measured in the simulations to the Boltzmann distribution

$$p_{\text{bond}}(\ell) = \frac{\exp(-U_{\text{bond}}(\ell)/k_{\text{B}}T)}{\int_{0}^{\infty} \mathrm{d}\ell \, \exp(-U_{\text{bond}}(\ell)/k_{\text{B}}T)}$$
(2.10)

and obtained an excellent agreement, which was expected because the bond potential is the strongest in the system.

As for the bending angle potentials, our LAMMPS simulations and recent GROMACS works used a harmonic in $\cos \theta$ bending potential given by Eq. (2.2) and Table 2.2, whereas the initial version of the model [26] employed

$$U_{\rm bend}(\theta) = \frac{1}{2}k_{\theta}(\theta - \theta_0)^2 . \qquad (2.11)$$

Assuming θ to be close to the equilibrium bond angle θ_0 the bending stiffnesses K_{bend} and



Figure 2.11: Bending angle distributions measured during the LAMMPS (red open squares) and GROMACS (blue filled circles) simulations. The vertical lines mark the average equilibrium values of θ_0 . The black solid lines are the sum of Eq. (2.13) calculated for each bending angle type and normalized by the number of corresponding angles in the system.

 k_{θ} are related to one another by ⁴

$$K_{\text{bend}} = \frac{k_{\theta}}{2\sin^2 \theta_0} \ . \tag{2.12}$$

In this way, K_{bend} was determined from k_{θ} of Ref. [26]. K_{bend} is smaller than K_{bond} by a factor of about 37, implying larger fluctuations of θ than for ℓ . Still, the fluctuations remain very small because $\sqrt{\langle (\theta - \theta_0)^2 \rangle} \approx \left[k_{\text{B}} T / (2K_{\text{bend}} \sin^2 \theta_0) \right]^{1/2} \approx 5^{\circ}$ even for T = 353 K.

Fig. 2.11 presents bending angle distributions measured in our LAMMPS and GRO-MACS simulations along with the Boltzmann prediction

$$p_{\rm bend}(\theta) = \frac{\exp(-U_{\rm bend}(\theta)/k_{\rm B}T)}{\int_0^{\pi} \mathrm{d}\theta \ \exp(-U_{\rm bend}(\theta)/k_{\rm B}T)} \ . \tag{2.13}$$

The data agree very well with the latter verifying the correct transfer of the bending angle potential from GROMACS to LAMMPS. The bending potential is also a hard variable in the system.

The torsion potential has been transferred from GROMACS to LAMMPS exactly. In both cases it is given by Eq. (2.3) and Table 2.3, while the initially developed model [26,

⁴To get Eq. (2.12) we insert the identity $\cos \theta - \cos \theta_0 = -2 \sin[(\theta + \theta_0)/2] \sin[(\theta - \theta_0)/2]$ into Eq. (2.2) and assume deviations of θ from θ_0 to be small.

39] utilized

$$U_{\rm tor}(\phi) = \frac{1}{2} \sum_{n=1}^{6} k_n \left[1 - \cos(n\phi) \right] \ . \tag{2.14}$$

The coefficients $\{A_n\}$ in Table 2.3 were obtained by fitting Eq. (2.3) to Eq. (2.14) for each type of dihedral angle (the details are described in Appendix B).

The double-bond dihedral potentials are much stronger than the α and β dihedral potentials. This ensures that the double-bond torsion angles stay close to the planar average equilibrium state during the simulations. It also implies that a monomer initially prepared in a *cis* or *trans* state will remain in this state for all times of current simulations with a length of few microseconds. In the initial configurations, that we received from the work [36], the potentials were randomly assigned to the monomers in a chain with the ratio of 45% cis and 55% trans configurations. The deviations from the planar configurations are possible, but quickly become penalized energetically due to large values of $U_{tor}(\phi)$ relative to the thermal energy that is $k_{\rm B}T\simeq 0.7$ kcal/mol at T=353 K⁵. The potential energy of the inflection points at $\phi = 90^{\circ}, 270^{\circ}$ corresponds to about $12\,000\,\mathrm{K}$ and that of the maxima at $\phi = 180^{\circ}$ for *cis* or $\phi = 0^{\circ}$, 360° for *trans* to about 24 000 K. These temperatures are much larger than the temperatures studied $(T \leq 353 \,\mathrm{K})$ so that the corresponding angles are never adopted in the simulation⁶. We can estimate typical fluctuations around this state by a harmonic expansion of $U_{\rm tor}(\phi)$ around $\phi_{\rm min}$. This expansion provides an excellent approximation up to $U_{\rm tor}(\phi \approx \phi_{\rm min} \pm 20^\circ) \approx 2.7 \,\rm kcal/mol$ with a force constant $k_{\phi} = 0.0133 \,\mathrm{kcal/(mol \, deg^2)}$, implying that $\sqrt{\langle (\phi - \phi_{\min})^2 \rangle} = \sqrt{k_{\mathrm{B}} T / k_{\phi}} \approx 7^{\circ}$ for T = 353 K. Deviations from the equilibrium state are therefore small, though larger than for ℓ and θ . The double bonds, together with the bond lengths and bond angles, are therefore stiff degrees of freedom.

The main source of chain flexibility and conformational rearrangements in PBD stems from torsional rotations around the β bond and the α *cis* and α *trans* bonds. This is the case because the maximum potential energy of these dihedrals is an order of magnitude smaller than for the double bonds, as seen from Fig. 2.4. The energy difference between *trans* and *gauche* states of the β potential is about 0.41 kcal/mol $\approx 0.58k_{\rm B}T$ at 353 K and the barrier at $\phi \simeq 180^{\circ} \pm 60^{\circ}$ is about 3.84 kcal/mol $\approx 5.48k_{\rm B}T$ at 353 K. The barriers

 $^{^5 \}rm We$ utilize the value of Boltzmann constant defined in the LAMMPS code, $k_{\rm B}=0.0019872067~\rm kcal/(mol~K).$

 $^{^{6}}$ When testing the reduced torsions, we found some flipping events at 1/10 of the initial height of the dihedral barriers.



Figure 2.12: Torsion angle distributions measured during the LAMMPS (red open squares) and GROMACS (blue filled circles) simulations. The black solid lines are the sum of all torsion types in Eq. (2.15). The separate distributions will be discussed in Section 3.1.1

between the minima of the α potential are smaller than those of the β bond, implying that torsional transitions are more easily possible about these dihedrals.

Fig. 2.12 presents the complete torsion angle distributions measured in our LAMMPS and constrained GROMACS simulations along with the Boltzmann prediction

$$p_{\rm tor}(\phi) = \frac{\exp(-U_{\rm tor}(\phi)/k_{\rm B}T)}{\int_0^{2\pi} {\rm d}\phi \ \exp(-U_{\rm tor}(\phi)/k_{\rm B}T)} \ . \tag{2.15}$$

The LAMMPS and GROMACS distributions agree with each other very well indicating a correct implementation of the dihedrals in our model. Both of them, however, differ from the Boltzmann distributions: the dihedral angles are redistributed from 0° and 360° to 120° and 240°. We examine this in detail in Section 3.1.1. According to the work [26], these deviations come from the influence of the intra-chain pair interactions between united atoms separated by four and more bonds on the α *cis* dihedral. In fact, in Ref. [26] the potentials were optimized to account for an effective torsional potential created by the presence of the pair interactions and to reproduce the correlations between the consecutive torsions.

As the last check of the model implementation we compared the structure and the dynamics of the PBD obtained from our LAMMPS simulations to the constrained GROMACS simulations. As a representative structural quantity we used the melt and single chain structure factors S(q) and F(q) (Fig. 2.13):

$$S(q) = \frac{1}{N_{\rm c}N} \sum_{i,j=1}^{N_{\rm c}} \sum_{a,b=1}^{N} \langle \exp[-i\vec{q}(\vec{r}_{a,i} - \vec{r}_{b,j})] \rangle \ , \tag{2.16}$$

$$F(q) = \frac{1}{N_{\rm c}N} \sum_{i=1}^{N_{\rm c}} \sum_{a,b=1}^{N} \langle \exp[-i\vec{q}(\vec{r}_{a,i} - \vec{r}_{b,i})] \rangle , \qquad (2.17)$$

where N_c is the number of chains, N the number of atoms in a chain, \vec{q} the wavevector, $\vec{r}_{a,i}$ the position of atom a of the chain i. For the S(q) the discrete \vec{q} are used, which are commensurable with the simulation box (the components of \vec{q} are multiples of $2\pi/L$ with L being the box size). The dynamics was compared using the mean-square displacement of united atoms $g_0(t)$ (Fig. 2.14)

$$g_0(t) = \frac{1}{N_{\rm c}N} \sum_{n=1}^{N_{\rm c}N} \langle [\vec{r}_n(t) - \vec{r}_n(0)]^2 \rangle, \qquad (2.18)$$

where $\vec{r}_n(t)$ the position of the *n*-th united atom at time *t*.

We have also compared our results to the data published in Ref. [41], which is an older work than Ref. [36], from which we received the initial configurations and GROMACS scripts. The data are in a very good agreement at the studied temperatures.

2.4.3 Pair interactions

As it was already mentioned in the previous chapters, the originally developed simulations of bulk PBD [26, 39] and the recent works on the confined films [31, 35, 36] utilized the plain Lennard-Jones potential truncated at $r_{\rm c} = 9$ Å for pair interactions

$$U_{\text{pair}}(r) = \begin{cases} 4\epsilon \left[\left(\frac{\sigma}{r}\right)^{12} - \left(\frac{\sigma}{r}\right)^6 \right] & \text{if } r < r_{\text{c}} ,\\ 0 & \text{if } r \ge r_{\text{c}} , \end{cases}$$
(2.19)

with the parameters for different types of united atoms given in Table 2.5. To account for missing interactions for distances r > 9 Å, the long range tail corrections [17, 64, 65] to pair energy and pressure were applied. We have also used the truncated LJ potential with tail corrections when setting up our first simulation of bulk PBD in LAMMPS. In practice,



Figure 2.13: Comparison of the melt (a) and the single chain (b) structure factors obtained form our LAMMPS simulations (red open squares) and constrained GROMACS simulation (blue filled circles) at T = 273 K. The black line is the data scanned from Ref. [41].



Figure 2.14: Comparison of the mean-square displacement of united atoms of PBD obtained form our LAMMPS simulations (red open squares) and constrained GROMACS simulation (blue filled circles) at T = 273 K and T = 353 K. The black line is the data scanned from Ref. [41].

these corrections are constant values calculated in the beginning of the simulation⁷, which are added to the values of pair energy and pressure at each timestep. The corrections to the energy, in fact, do not affect the dynamics of the system, because the forces between the particles are proportional to the gradient of the potential, which eliminates the constant. The pressure correction, however, has a direct influence on the dynamics under the NPT conditions, because it enters the barostating routine. This modifies the volume of the simulation box, and therefore, the density according to the desired pressure. For the bulk

 $^{^{7}}$ see Eq. (5) of Ref. [64] for the analytical expressions of the corrections

system under the NVT conditions, the influence of the corrections comes only via the value of the fixed volume. Enabling/disabling the tail corrections in the bulk NVT simulations at the same fixed volume does not change the dynamics of the simulated system whatsoever.

The situation is different for the simulations of the films, which are NVT simulations only. In the confined geometry the density can be adjusted by varying the distance between the walls. The works [31, 35, 36] used this method to match the density of PBD in the middle of the film to the bulk value. In the supported films, however, the density is determined by the interactions, and since the tail corrections have no effect in the NVT simulations, the value in the middle of the film corresponds to the one of the bulk simulated under the NPT conditions without the corrections. This issue does not allow us to continue using the truncated LJ potential with tail corrections, because we would not be able to directly compare the bulk and the film simulations under the same conditions. Yet, the original model has been validated against experiments [39, 45, 46, 66, 67], a salient feature we want to keep. Fig. 2.15a presents the bulk density of PBD obtained from the cooling runs at zero pressure using our LAMMPS simulations with the truncated LJ potential and corrections enabled. We compared our results to various experimental and simulation references, that we were able to find (see Appendix C for an overview). The data in the literature are for PBD of various compositions and molecular weight under different pressure conditions. Thus, it only gives a range of densities to compare with.

Due to the presence of the tail corrections, our data lay nicely within the references. Disabling them leads to a very significant decrease of the density in the NPT simulations at T = 353 K due to quite small $r_c = 9$ Å, as it is shown in Fig. 2.15b. Increasing r_c up to 14 Å results in densities closer to the experimental ones, but also leads to more than twice longer CPU times.

In order to avoid the tail corrections completely, we decided to apply the switching function S(r) to the LJ potential, as it is described in Section 2.2 (cf. Eqs. (2.5) to (2.7)). The inner cutoff in Eq. (2.5) was kept at $r_1 = 9$ Å, so that the shape of the potential at $r < r_1$ is kept as in the original model. The outer cutoff was set at $r_c = 12$ Å. The switching function S(r) smoothes the potential and its 1st and 2nd derivatives to zero over a distance of about one particle diameter 9 Å < r < 12 Å.

The modified pair potential Eq. (2.5) is of finite range and is still of shorter range than the original model with tail corrections. This difference affects the density of the melt, which is smaller for the modified model than for the original model and moreover does not fit into the interval of reference data from the literature (Fig. 2.15). Therefore, it is necessary to adjust the potential parameters in order to reproduce the desired density. To this end, we choose to leave the zero-crossing distance σ unchanged but to enhance ϵ by a factor $\lambda > 1$. This choice is motivated by the generally accepted point of view [3] that the local structure of (simple) liquids at high density is dominated by the packing of the molecular repulsive cores (i.e. by σ), whereas the attractive interactions (of strength ϵ) determine the cohesive energy and along with that, thermodynamic properties, such as the density or compressibility.

To determine λ we proceed in the following way. For the original model with truncated LJ potential and tail corrections we carry out an NPT simulation to get the reference value of the density at pressure p = 0 atm for two temperatures, T = 273 K and 353 K. As described in Section 2.3.2, we impose this density in a subsequent NVT simulation and calculate the average pair energy, $E_{\text{pair}}^{\text{ref}}(T)$, for the original model. For the modified model, Eqs. (2.5), (2.6) and (2.7), we also perform an NVT simulation, with the same density and



Figure 2.15: (a) Comparison of the PBD density $\rho(T)$ obtained from our LAMMPS simulations upon cooling the system in NPT conditions using a truncated LJ potential with tail corrections with the data from literature. Five independent cooling runs were averaged to obtain the density of PBD during our simulations (pink circles). (b) The dependence of PBD density $\rho(T)$ at T = 353 K on the cutoff distance $r_{\rm c}$ and tail corrections.
Table 2.5: Lennard–Jones energy parameters for the different types of united atoms $i, j = CH_2, CH$. Second column: ϵ_{ij} of the original model [26, 47]. Third column: ϵ_{ij} for the shifted and smooth model obtained from the original model by multiplying ϵ_{ij} by $\lambda = 1.149462$. Fourth column: σ_{ij} , which is the same for both models.

Pair type	original ϵ_{ij}	$\epsilon_{ij}^* = \lambda \epsilon$	σ_{ij}
	(kcal/mol)	$(\dot{k}cal/mol)$	(Å)
$CH \leftrightarrow CH$	0.1000460	0.114999	3.38542
$\overline{\operatorname{CH}_2\leftrightarrow\operatorname{CH}_2}$	0.0936436	0.107639	4.00904
$\operatorname{CH}_2\leftrightarrow\operatorname{CH}$	0.1015460	0.116723	3.79256

 ϵ as for the original model, and determine again the average pair energy $E_{\text{pair}}(T)$. The NVT simulations were performed in LAMMPS only and consisted of an equilibration run of 10 ns followed by a production run of another 10 ns to measure the pair energy and pressure. At both T = 273 K and 353 K the run length of 10 ns allows the UAs to displace on average over a distance of at least 2 particle diameters. As expected due to the shorter range of the modified model, we find $|E_{\text{pair}}| < |E_{\text{pair}}^{\text{ref}}|$. Since $U_{\text{pair}}(r)$ is proportional to ϵ , we define λ by

$$\lambda = \frac{E_{\text{pair}}^{\text{ref}}}{E_{\text{pair}}} \,. \tag{2.20}$$

At T = 273 K this gives $\lambda = 1.149462$. We utilize this value for all pairs ij of united atoms $(i, j = CH_2, CH)$, that is, we replace ϵ_{ij} of the original model by $\lambda \epsilon_{ij}$. The resulting values of ϵ_{ij} are given in the third column of Table 2.5 (they are the same as in Table 2.4). To verify the estimated λ we performed NVT simulations at T = 273 K and at T = 373 K with the new values of ϵ_{ij} . For both temperatures the densities and pressures agree with the target values of the original model.

We have also performed an inverse cross-check by imposing the p = 0 atm and comparing the density of PBD obtained from the cooling runs at two rates $\Gamma = 3.33$ K/ns and $\Gamma = 0.83$ K/ns using the original and the modified potential. The results are presented in Fig. 2.16.

All curves superimpose well at T > 225 K, so our new model is able to represent the density of PBD. Depending on the cooling rate Γ , the curves bend around $T \approx 200$ K, which suggests a glass transition. The modified model appears to bend slightly earlier upon cooling the system, which is expected due to the increased ϵ .

In addition, we have compared our results to the density obtained from the first simulations where all the dihedrals had been disabled, which also superimposed well in the



Figure 2.16: The density of PBD obtained from the cooling runs using the original and the modified pair interactions. The purple line represents the cooling at the rate $\Gamma = 3.33$ K/ns using the original model. This is the same data as in the Fig. 2.15. The green line is the data obtained from a slower cooling rate of $\Gamma = 0.83$ K/ns. The black and the red lines are from the simulations using the modified model. The data described above were obtained by averaging over 5 independent cooling runs (the simulation details are given in Appendix A). The blue line corresponds to a single cooling run using the original model without the dihedrals.

region of the working temperatures. This model has been already studied in Refs. [40, 41], in which much faster dynamics of the PBD was reported for the system without dihedrals while the structure was observed to be unchanged. Thus, even though our cooling run without dihedrals starts from the configuration obtained from the system with dihedrals, we expect all perturbations that may occur by doing so (if any) to be relaxed during first few nanoseconds of the run. This allowed us to omit the NPT equilibration step for the models with modified dihedrals, and to use the densities obtained using our CRC model for the NVT simulations of our models with reduced and disabled dihedral barriers. We will continue this discussion in Section 3.2 where we analyse the cooling runs using all our models.

2.5 Details of sampling and analysis of the trajectories

In order to compute the quantities of interest there are trajectory files written during the simulation, in which the coordinates of all particles in the simulated system (usually called "a frame") are saved at selected timesteps. Our simulations write two types of such



Figure 2.17: The details of the linear (a) and logarithmic (b) spacing of the trajectory files written during our simulations using LAMMPS.

files. The first type contains frames written at timesteps that are equidistantly spaced (Fig. 2.17a). These files are used for calculating ensemble averages of the static properties. The sequence of frames is firstly divided into equal blocks, and the quantities are computed and averaged within each block separately. The average of the results over all blocks is then performed along with the standard deviation of mean, that is usually used as the estimation of error bars. The second type of files, which we call "logarithmic" trajectory, also has blocks of frames, but the frames are spaced within each block exponentially at short times, such that there is an equal distribution of points when plotting dynamic quantities logarithmically in time (Fig. 2.17b). These files are used for calculating the time averages with the resolution of the time windows varying over several decades: these "logarithmic blocks" produce the short time average, and the first frame of each block is used for the long time average as in the files of the first type.

The thermodynamic quantites in the simulations were calculated by LAMMPS itself and writted into separate log files. The various static and dynamic quantities presented in this work were calculated using trajectory analysis tools developed in the Theory and Simulations of Polymers (TSP) team at Institute Charles Sadron and the library written by the author in C++ programming language (available at [77]). The post-analysis, figures and this manuscript itself were done using R programming language for statistical computing and the available packages [78–82].

The bulk simulations were carried out using LAMMPS build of 7 Sep 2016 at the High Performance Computing (HPC) Center of the University of Strasbourg (~ 4 million hours) [59]. The film systems were simulated using LAMMPS build of 29 Oct 2020 on the Jean Zay (HPE SGI 8600) supercomputer at the Institute for Development and Resources in Intensive Scientific Computing (IDRIS, Orsay; ~ 2.9 million hours) [60]. Examples of

the LAMMPS input scripts are presented in Appendix D.

Chapter 3

Bulk systems analysis

This chapter presents the results obtained from the simulations of the bulk systems. We begin with the analysis of the conformational properties in Section 3.1. Then the density and the glass transition are discussed in Section 3.2. Afterwards the static structure of the PBD is analyzed in Section 3.3. The obtained results allow us to check the thermodynamic consistency of the simulations and analyze the heat capacity in Section 3.4. In Section 3.5 we apply the Rouse model to describe the static properties of PBD and give the first insights on the dynamics. A more detailed analysis of the dynamics is made in Section 3.6. Finally, we discuss our results in Section 3.7.

3.1 Conformational properties

3.1.1 Distributions of the torsion angle

A first insight into the influence of reducing the torsional potential on the properties of PBD may be provided by the distribution of the dihedral angle, $p_{\lambda}(\phi)$, where λ identifies the different models as defined in Section 2.2. If the dihedral angle was independent of the other potentials, $p_{\lambda}(\phi)$ would be solely determined by λU_{tor} ,

$$p_{\lambda}(\phi) = \frac{\exp(-\lambda U_{\rm tor}(\phi)/k_{\rm B}T)}{\int_{0}^{2\pi} \mathrm{d}\phi \, \exp(-\lambda U_{\rm tor}(\phi)/k_{\rm B}T)} \,. \tag{3.1}$$

It is therefore instructive to compare Eq. (3.1) with the distribution obtained from the MD simulations for a given dihedral. Such a comparison is shown in Fig. 3.1. As seen from Fig. 3.1a, the distribution of the double bond *trans* dihedral angle from the MD for the



Figure 3.1: Distributions $p_{\lambda}(\phi)$ for different types of dihedrals angles at T = 353 K. (a) Log-linear plot of $p_{\lambda}(\phi)$ for the double bond of the *trans* conformer. (b) Linear plot of $p_{\lambda}(\phi)$ for the β bond between two monomers. (c) Linear plot of $p_{\lambda}(\phi)$ for the α bond adjacent to the double bond of a *trans* monomer. (d) Linear plot of $p_{\lambda}(\phi)$ for the α bond adjacent to the double bond of a *cis* monomer. In all panels the symbols represent the MD results and the full lines show the results of Eq. (3.1). The colors mark the simulation models: CRC ($\lambda = 1$) – red, CRC2 ($\lambda = 1/2$) – green, CRC4 ($\lambda = 1/4$) – purple, FRC ($\lambda = 0$) – blue. The black dashed horizontal line indicates the uniform distribution $p_{\lambda}(\phi) = 1/360$ expected for the FRC model from Eq. (3.1)

CRC model is in excellent agreement with Eq. (3.1). This is not unexpected because the dihedral associated with the double bond is a "hard" conformational variable which is kept close to the equilibrium value by its stiff torsional potential. Reducing this potential could change the situation. However, the agreement between the simulated distribution and Eq. (3.1) is still good for both the CRC2 and CRC4 models (Fig. 3.1a). While progressive softening of the torsional potential enhances deviations from the *trans* configuration, the population at $\phi = 0^{\circ}$ or 360° vanishes, even for the CRC4 model despite the decrease of the

potential barrier of the *cis* state by a factor of 4. This implies that cis/trans isomerization is precluded not only for the CRC model, but also for the CRC2 and CRC4 models. Since the torsional potential for the double bond of the *cis* conformer is only shifted by 180° relative to that of *trans* conformer, the results reported here for the *trans* conformer are found for the *cis* conformer, too (not shown).

The difference between the *cis* and *trans* conformers disappears when the torsional potential is switched off. Then, Eq. (3.1) predicts a uniform distribution, $p_{\lambda=0}(\phi) = 1/(2\pi)$. The simulated distribution for the FRC model deviates from this prediction (Fig. 3.1a). The probability of finding ϕ near 180° is increased relative to the uniform distribution, while it is decreased near $\phi = 0^{\circ}$ or 360°. These deviations stem from intrachain LJ interactions between united atoms separated by four and more bonds, which contribute to the conformational energies and thereby create an effective torsional potential. Such an influence of the LJ potential is expected from the force field parametrization developed in Ref. [26], where intramolecular pair interactions between consecutive torsional states (second-order effects between consecutive pairs of torsions and third-order effects involving three consecutive torsions).

For the FRC model this also implies that similar deviations between the simulated and the uniform distributions must be observed for the other dihedrals of the β , α *cis* and α *trans* bonds. This is illustrated in Figs. 3.1b to 3.1d. As in the case of the double bonds, we see that the probability of adopting torsional angles is enhanced near 180° and supressed near $\phi = 0^{\circ}$ or 360° for the FRC model. Deviations between the MD results for the dihedral distributions and the predictions from Eq. (3.1) are also visible for the CRC, CRC2 and CRC4 models. While Figs. 3.1b and 3.1c show that the deviations are small for the β and α *trans* bonds, they are large for the α *cis* bond (Fig. 3.1d), in particular for torsional states near $\phi = 0^{\circ}$ or 360° which are suppressed in the MD simulation, contrary to the prediction of Eq. (3.1). Fig. 3.2 shows a configuration of the *cis* monomer and the UAs adjacent to it, where all dihedral angles were fixed to $\phi = 0^{\circ}$. This geometry results in a distance of only 1.457 Å between the outer UAs. Since these UAs are separated by more than 4 bonds along the chain, the pair interaction between them takes place. According to Eq. (2.5) and Table 2.4, the pair energy at this distance is equal to $U_{\text{pair}} \approx 10^5$ kcal/mol ($\approx 10^7$ K), which is so high, that the α *cis* dihedrals at $\phi = 0^{\circ}$ are never adopted during



Figure 3.2: Fragment of PBD chain comprising 6 UAs. 4 inner UAs make a monomer of PBD. The bond lengths and bending angles are fixed to their equilibrium values according to Tables 2.1 and 2.2. All dihedral angles were fixed to $\phi = 0^{\circ}$ representing the ideal *cis* configuration.

the simulations and are redistributed around the other minima of the potential. This effect is commonly referred to in the literature as the 'steric hindrance' [83, 84] or 'pentane interference' [85, 86]. These intrachain correlations beyond those of single torsions are clearly important for the conformational properties of PBD [26].

3.1.2 Bond correlations and internal distances

Let \vec{r}_n denote the position of the *n*th united atom and $\vec{l}_n = \vec{r}_{n+1} - \vec{r}_n$ its bond vector (n = 1, ..., N). For the united atom m = n + s that is separated by *s* bonds from *n*, we define the mean-square internal end-to-end distance $R_e^2(s)$ and the bond correlation function $P_1(s)$,

$$R_{\rm e}^2(s) = \left\langle (\vec{r}_{n+s} - \vec{r}_n)^2 \right\rangle, \quad P_1(s) = \frac{\left\langle \dot{\ell}_{n+s} \cdot \dot{\ell}_n \right\rangle}{l^2} , \qquad (3.2)$$

where $l^2 = \langle \vec{\ell}_n^2 \rangle$ is the mean-square bond length of the PBD chain. Here $\langle ... \rangle$ denotes the thermal average (over all configurations and chains of the system) and also the average over all possible pairs n, m = n + s. (Hence, the statistics will deteriorate for $s \to N - 1$.) Since $\vec{r}_{n+s} - \vec{r}_n = \sum_{i=n}^{n+s-1} \vec{l}_i$, $R_e^2(s)$ and $P_1(s)$ are related to each other by

$$R_{\rm e}^2(s) = l^2 s + 2l^2 \sum_{k=1}^{s-1} (s-k) P_1(k) .$$
(3.3)

Both quantities are of considerable theoretical importance [2, 87, 88] and have often been studied in polymer melt simulations [21, 88–90]. Therefore, we also determine them here.

As an example for the typical behavior of $P_1(s)$ found for all models in the interval

213 K $\leq T \leq$ 353 K, Fig. 3.3 shows the results of the CRC model at T = 353 K. If bond correlations were only caused by the bond angle, one would expect from the freely rotating chain model theory an exponential decay $P_1(s) = |\langle \cos \theta \rangle|^s$ [2]. Clearly, this prediction (dashed line in Fig. 3.3) must underestimate $P_1(s)$ for PBD. It only accounts for correlations between consecutive pairs of bonds $[P_1(s = 1)]$, but not for correlations between more distant pairs along the chain backbone $[P_1(s > 1)]$, resulting from dihedral and interchain pair interactions. To fit these more distant correlations we use an exponential function $P_1(s) = A \exp(-s/\bar{s})$ with A and \bar{s} as adjustable parameters. This gives a good description of the MD data for $2 \leq s \leq 25$ (full line in Fig. 3.3). Fig. 3.3 shows that $P_1(s \approx 25) \sim 10^{-3}$, which is the noise level of our MD results. Since the statistical accuracy does not allow one to explore bond correlations for large $s \to N - 1$, the best adapted parameterization of the present data is given by

$$P_1(s) = \begin{cases} \alpha & \text{for } s = 1 \ , \\ A \exp(-s/\overline{s}) & \text{for } s > 1 \ . \end{cases}$$
(3.4)

where we defined $\alpha = -\langle \cos \theta \rangle$ (> 0). We have analyzed all models in the temperature interval 213 K $\leq T \leq 353$ K with Eq. (3.4). The directly measured values for α and the fit results for A and \bar{s} are compiled in Appendix E.

Insertion of Eq. (3.4) into Eq. (3.3) gives

$$R_{\rm e}^2(s) = l^2 s \left[C_{\infty} - \frac{2}{s} \left(\alpha - A\overline{\alpha} + A \frac{\overline{\alpha} - \overline{\alpha}^{s+1}}{(1 - \overline{\alpha})^2} \right) \right] , \qquad (3.5)$$

where $\overline{\alpha} = e^{-1/\overline{s}}$ (< 1) and C_{∞} is the characteristic ratio of the infinitely long chain,

$$C_{\infty} = \lim_{N \to \infty} \frac{R_{\rm e}^2(N-1)}{(N-1)l^2} = 1 + 2(\alpha - A\overline{\alpha}) + \frac{2A\overline{\alpha}}{1-\overline{\alpha}} . \tag{3.6}$$

The characteristic ratio determines the effective bond length $b_{\rm e} = \sqrt{C_{\infty}} l$ [1] and the temperature coefficient κ of the mean-square end-to-end distance $R_{\rm e}^2(N-1)$ of the chain [91],

$$\kappa = 1000 \,\frac{\mathrm{d}\ln R_{\rm e}^2(N-1)}{\mathrm{d}T} = 1000 \,\frac{\mathrm{d}\ln C_{\infty}}{\mathrm{d}T} \,. \tag{3.7}$$

The factor of 1000 is introduced in this definition because κ is typically of the order of 10^{-3} K^{-1} [26, 92].



Figure 3.3: Log-log plot of $P_1(s)$ versus the number of bonds s $(1 \le s \le N-1)$ for the CRC model (red circles) at T = 353 K. The dashed line indicates the exponential decay based on the freely rotating chain model theory, $|\langle \cos \theta \rangle|^s$ [2]. The solid line represents the fit result to $A \exp(-s/\bar{s})$ for $2 \le s \le 20$, yielding A = 0.7507 and $\bar{s} = 3.625$. \bar{s} is indicated by a vertical dotted line. For s > 25 the statistical accuracy of the data deteriorates: $P_1(s)$ can become negative, explaining the gap in the log-log plot for $30 \le s \le 100$. The associated 'noise' does not allow an analysis of $P_1(s)$ for $\bar{s} \ll s \le N-1$.

Fig. 3.4a depicts the ratio $R_e^2(s)/s$ as a function of s for all models at T = 353 K. This ratio starts from the mean-square bond length l^2 (= 2.158 Å²)¹, then increases first steeply for small s and eventually levels off when s approaches N - 1. We find that the dependence $R_e^2(s)/s$ on s can be well fitted by Eq. (3.5) for all models and temperatures (see Appendix E for a compilation of the fit results). The full line in Fig. 3.4a gives an example for the FRC model. By repeating this analysis at all studied temperatures we obtain the T dependence of C_{∞} (Table 3.1). The results are plotted as $\ln C_{\infty}$ versus T in Fig. 3.4b. This plot format approximately rectifies the data, yielding negative values for κ in the range $-0.64 \,\mathrm{K}^{-1} \leq \kappa \leq -0.52 \,\mathrm{K}^{-1}$ for all models studied. A negative value for κ implies that the chains tend to expand on cooling.

The results for κ may be compared with experimental values [93] and Rotational Isomeric State (RIS) calculations [26, 94]. These studies show that the thermal coefficient is sensitive to the stereochemical composition of PBD. While pure *cis*-PBD has positive κ , negative values are obtained for the pure *trans*-form. For mixed stereoirregular mi-

¹The stiff bond length potential of the PBD model implies that the bond length is Gaussian distributed around the equilibrium bond length ℓ_0 , leading to a *T* independent $\langle \vec{\ell}^2 \rangle = \ell_0^2$. Out of N-1 bonds, a PBD chain has N/4 CH=CH bonds of length $\ell_0 = 1.34$ Å, N/4 - 1 CH₂-CH₂ bonds of length $\ell_0 = 1.53$ Å, and N/2 CH–CH₂ bonds of length $\ell_0 = 1.50$ Å. The weighted average of ℓ_0^2 gives $l^2 = 2.158$ Å².



Figure 3.4: (a) Plot of the mean-square internal distance $R_e^2(s)$ divided by the number of bonds *s* versus *s* ($1 \le s \le N-1$) for the CRC (red circles), CRC2 (green triangles), CRC4 (purple pluses) and FRC models (blue crosses). All data refer to T = 353 K. The (black) dashed line shows a fit to Eq. (3.5) for the FRC model (not shown for other models). The horizontal dotted lines show respectively the mean-square bond length $l^2 = 2.158$ Å² and for the FRC model the square effective bond length $b_e^2 = 11.58$ Å². (b) Temperature dependence of the characteristic ratio C_{∞} from Eq. (3.6) for the CRC (red circles), CRC2 (green triangles), CRC4 (purple pluses) and FRC models (blue crosses). The plot format, $\ln C_{\infty}$ versus *T*, is motivated by Eq. (3.7) when assuming κ to be constant. The full line presents a linear fit to the FRC data, yielding $\kappa = -0.52$ K⁻¹. The dashed line shows the fit result to the CRC model, leading to $\kappa = -0.64$ K⁻¹. The dotted line indicates the experimental result, $\kappa = -0.10$ K⁻¹ (obtained for 298 K $\leq T \leq 373$ K), from Table VIII of Ref. [93].

crostructures similar to our simulation model, RIS calculations [94] mostly lead to small negative values of κ (~ -0.1 K⁻¹), in resonance with experimental results (where PBD also contains about 9% of vinyl groups in addition to *cis* and *trans* units) [93]. From this comparison we can conclude that our simulation results are in qualitative accord

Table 3.1: Characteristic ratio C_{∞} from Eq. (3.6) for the various models studied [Section 2.2] at temperature T.

$T(\mathbf{K})$	CRC	CRC2	CRC4	FRC
213	6.2	6.0	5.9	5.8
225	5.8	5.9	5.9	5.7
240	5.8	5.8	5.9	5.7
253	6.1	5.9	5.7	5.6
273	5.7	5.7	5.7	5.5
293	5.7	5.7	5.6	5.5
353	5.5	5.5	5.5	5.4

with available literature data ($\kappa < 0$), but the extent of the chain extension on cooling is stronger in the simulation.

Finally, a remarkable observation from the previous analysis is the weak influence of the torsions. Even when switching off all dihedral potentials the conformational properties of the resulting FRC model are very close to those of the CRC model. This observation agrees with the original works [40, 41]. Qualitatively, we can rationalize this feature by invoking the hindered rotation model that expresses C_{∞} as [2]:

$$C_{\infty} = \frac{1+\alpha}{1-\alpha} \frac{1-\langle \cos \phi \rangle}{1+\langle \cos \phi \rangle} \,. \tag{3.8}$$

If the torsions were uniformly distributed for the FRC model, we would have $\langle \cos \phi \rangle = 0$ and switching off of the torsional potential would give a big effect on C_{∞} . However, Fig. 3.1 shows that the distribution of the dihedral angles is not uniform for the FRC model due to the intrachain pair interactions, leading to a finite value of $\langle \cos \phi \rangle$. Therefore, we determined $\langle \cos \phi \rangle$ for all models. The results are summarized in Table 3.2, together with C_{∞} from Eq. (3.8) (using the values of α from Appendix E). We see that the values of C_{∞} from Eq. (3.8) for the different models are close to one another at a given *T*. This is consistent with the trend shown in Table 3.1. Even at a quantitative level the characteristic ratio from the hindered rotation model is close to the results from Table 3.1. This implies that the dominant contribution to chain stiffness stems on average from local correlations involving three bonds along the chain backbone in PBD.

Table 3.2: C_{∞} at temperature T from the hindered rotation model [Eq. (3.8)]. The values for $\langle \cos \phi \rangle$ were determined from the simulation.

	CRO	C	CRC	22	CRC	24	FRO	C
$T(\mathbf{K})$	$\langle \cos \phi \rangle$	C_{∞}						
213	-0.323	5.6	-0.275	5.0	-0.253	4.8	-0.246	4.7
225	-0.312	5.4	-0.270	4.9	-0.249	4.7	-0.244	4.7
240	-0.305	5.3	-0.264	4.9	-0.246	4.7	-0.242	4.7
253	-0.302	5.3	-0.261	4.9	-0.243	4.7	-0.240	4.6
273	-0.292	5.2	-0.255	4.8	-0.242	4.6	-0.237	4.6
293	-0.285	5.1	-0.250	4.8	-0.235	4.6	-0.234	4.6
353	-0.267	4.9	-0.238	4.6	-0.226	4.5	-0.227	4.5

3.2 Density

The comparison of the bulk density of simulated PBD to experimental data has been already presented in Section 2.4.3 in case of the original PBD model. Our CRC model essentially reproduces the same density, which fits within the experimental and simulation data from the literature (see Figs. 2.15 and 2.16 and Appendix C for an overview). In this section we complement the discussion by comparing the density obtained from cooling the PBD down to 3 K using all our models, comment on the equilibrium values at the working temperatures and analyze the glass transition.

Fig. 3.5 presents the mass densities $\rho_{\rm m}(T) = m/V(T)$ (*m* is the mass of PBD in the simulation box of volume *V*) of PBD obtained in our work. Different symbol shapes denote the runs, from which the data were extracted, and the colors mark the models. The empty circles represent the data obtained from 5 independent cooling runs at the



Figure 3.5: Main panel: Density of the bulk PBD. The empty colored circles represent the data from the cooling runs. The black triangles correspond to the equilibrium volumes, obtained from 100 ns NPT equilibration and imposed during the NVT runs for all models. The colored crosses are the data from 1 μ s long NPT runs. The solid black lines represent the fits using Eq. (3.10) with the parameters form Table 3.3. Inset: The fraction of the densities from the cooling (circles) and NVT runs (triangles) divided by the corresponding density from the long NPT runs for each model.

rate $\Gamma = 0.83$ K/ns, that were, at first, averaged together at each timestep, and then smoothed so that each point is an average of the raw data over a neighboring region of $\Delta T = 1$ K. The curves overlap at high temperatures before the bending occurs at different temperatures depending on the model. Clearly, such a bending suggests the glass transition that depends on the strength of the dihedrals. We will discuss this later in this section. The overlapping data reveals the mild influence of the dihedrals on the density of PBD in the liquid state.

Honnell et al have studied the equation of state for chain molecules in 1987 [61] and derived that the pressure does not depend on the angular potentials explicitly². It is determined by the inter- and intrachain contributions of the site-site correlations, that include only the pair and the bond terms:

$$\begin{pmatrix} \frac{p\beta}{\rho_{\rm c}} \end{pmatrix} = N - \frac{2\pi\beta\rho_{\rm c}}{3} \sum_{\alpha,\gamma} \int_{0}^{\infty} g_{\rm inter}^{\alpha,\gamma}(r) \frac{dU_{\rm pair}}{dr} r^{3} dr - \frac{4\pi\beta}{3V} \sum_{i=1}^{N-1} \int_{0}^{\infty} g_{\rm intra}^{i,i+1}(r) \frac{dU_{\rm bond}}{dr} r^{3} dr - \frac{4\pi\beta}{3V} \sum_{\alpha=1}^{N-4} \sum_{\gamma=\alpha+4}^{N} \int_{0}^{\infty} g_{\rm intra}^{\alpha,\gamma}(r) \frac{dU_{\rm pair}}{dr} r^{3} dr ,$$

$$(3.9)$$

where $\rho_{\rm c}$ is the number density of chains, N is the number of sites in a chain, $g_{\rm inter}^{\alpha,\gamma}(r)$ is the probability to find the sites α and γ , that belong to different chains, at a distance r; $g_{\rm intra}^{\alpha,\gamma}(r)$ is the probability to find the sites α and γ , that belong to the same chain, at a distance r.

Indeed, the pressure components reported by LAMMPS (Fig. 3.6a), that correspond to the contributions arising directly from the bending and torsional potentials are equal to zero for all models. However, the angular potentials may still influence the site-site correlations $g_{inter}^{\alpha,\gamma}(r)$ and $g_{intra}^{\alpha,\gamma}(r)$. In order to estimate these correlations we have calculated the total radial distribution function g(r) and its inter/intrachain contributions for all models assuming all sites to be equivalent. They are presented in Figs. 3.6b to 3.6d. The curves, that correspond to the intermolecular part, overlap (Fig. 3.6c), so the 1st integral term in Eq. (3.9) is also same for all models. The bonds are very stiff and do not feel the presence of the dihedrals. They result in very high peaks around the equilibrium bond lengths

²The main argument of the derivation is the fact, that the angles do not change when rescaling the volume of the system. Therefore, the derivatives of the angular potentials with respect to the volume vanish. Our simulations employed an isotropic barostat, so this statement holds.

3.2. DENSITY

(they are truncated in Fig. 3.6d), which bring the same contribution to the 2nd integral term in Eq. (3.9) for all models. The intramolecular part has visible differences between the models at 3 Å $\leq r \leq 5$ Å (Fig. 3.6d). However, the distances $r \leq 4$ Å correspond to the correlations only up to the 3rd neighbor along the chain ($\sqrt{R_e^2(s=3)} \approx 3.5$ Å in Fig. 3.4a). These interactions are excluded in the simulations. The intramolecular part of the pair potential is taken into account starting form the 4th neighbor, which corresponds to the distances $r \gtrsim 4$ Å ($\sqrt{R_e^2(s=4)} \approx 4.5$ Å). In this region the data for the $g_{intra}(r)$ differs noticeably only close to the peak. We expect these differences to be negligible with



Figure 3.6: (a) Contributions to the pressure arising from the different potential terms calculated in LAMMPS during the simulations at T = 353 K. The data were averaged over 10 ns of a NVT run. (b-d) The total (a), intermolecular (c) and intramolecular (d) radial distribution functions obtained from the NVT simulations of 1 μ s in length at T = 353 K. The colours denote different models: CRC - red, CRC2 - green, CRC4 - purple, FRC - blue.

respect to the other terms of Eq. (3.9). Thus, with sufficient amount of confidence, we can assume the pressure, and therefore, the resulting density of PBD to be the same for all studied models in the melt state.

Consequently, the values of the equilibrium volume obtained from the NPT runs using the CRC model were employed for the NVT simulations of all models at all working temperatures. These data are shown in Fig. 3.5 as black triangles. As a post factum check, we have compared our data to the densities extracted from the 1 μ s long NPT runs for each model (colored crosses), that were launched after the NVT runs. These values of the density are obtained with much better averaging statistics and still agree nicely with the rest of the data within the size of the symbol. The inset of Fig. 3.5 shows that the differences between the cooling densities, the densities imposed in the NVT runs and the ones after long NPT runs are less than 1%.

As it was mentioned above, the cooling curves change their slope at different temperatures depending on the model, which suggests an influence of the dihedrals on the glass transition temperature $T_{\rm g}$. There are several ways to extract the $T_{\rm g}$ from the evolution of the density upon cooling. A straightforward approach often used in the experiments and simulations is to estimate the $T_{\rm g}$ from the intersection point of the linear glassy and liquid regions [54, 95, 96]. Even though the choice of the linear regions is ambiguous, this method works reasonably well for the experimental data, because the experimental cooling rates of the order of $\Gamma \sim 0.001$ -1 K/s [97–99] result in sharply pronounced changes of the slope over only few Kelvins. Still, the ambiguity in the method leads to the problem of reproducibility of the results. In order to solve this problem Dalnoki-Veress et al proposed an alternative method to determine the $T_{\rm g}$ [100]. Assuming the thermal expansion coefficients to be constant in the liquid and in the glass (which correspond to the slopes M and G in the density curves), interpolating across the transition region by a hyperbolic tangent function and integrating the resulting equation, they obtained

$$\rho(T) = w\left(\frac{M-G}{2}\right) \ln\left[\cosh\left(\frac{T-T_{\rm g}}{w}\right)\right] + (T-T_{\rm g})\left(\frac{M+G}{2}\right) + \rho_{\rm g}, \qquad (3.10)$$

with $\rho(T)$ being the density at the temperature T, w the width of the glass transition region, M and G the slopes $d\rho(T)/dT$ in the melt and glass regions respectively, T_g the glass transition temperature, ρ_g the density at $T = T_g$. We employed this method in our simulations, which, due to significantly faster cooling rate $\Gamma = 0.83$ K/ns are expected to result in much broader transition region. Even by a visual estimation, our data in Fig. 3.5 bends over about $w \approx 30\text{-}50$ K, while the typical experimental values are $w \approx 2$ K [100]. The situation is difficult for the FRC model, where it is not clear if the glassy linear regime is pronounced. While accounting for the width of the transition explicitly, there is still an ambiguity in choosing the temperature interval $(T_g^* - \Delta T, T_g^* + \Delta T)$ over which the fit is applied³. In order to investigate the influence of the choice of ΔT , we have attempted to fit our smoothed data using Eq. (3.10) over a set of ΔT ranging from tens of Kelvins to the highest possible value. The results are presented in Fig. 3.7, where the values of the fitted parameters are plotted against ΔT . The lower bound of ΔT is limited by the amount of smoothing, since it is meaningless to apply the fit over a region of the fluctuation of the data. According to Fig. 3.7, the T_g , ρ_g and the slopes M and G converge to the corresponding plateaus, but the transition width parameter w appears to be the most sensitive to ΔT .

In order to obtain the final values of the parameters, we fixed the $T_{\rm g}$, $\rho_{\rm g}$, M and G to their plateau values and refitted the data for the highest ΔT in order to obtain an 'averaged' w. These data are presented in Table 3.3 and as dashed lines in Fig. 3.7. In case of the FRC model this resulted in slightly higher value of w with respect to the ΔT -dependent fit. The fitted curves are presented in Fig. 3.5 as solid black lines.

The most important fitting parameter is, of course, the glass transition temperature $T_{\rm g}$. The value for the CRC model is reasonably higher than the experimental $T_{\rm g} \approx 175$ K (for high molecular weights) [98, 101] again due to much faster cooling rate. Weakening the dihedrals significantly shifts the $T_{\rm g}$ to lower values. The value for the FRC model qualitatively agrees with the results of the coarse-grained simulations, where typical values of $T_{\rm g}$ are of the order of a fraction of $\epsilon/k_{\rm B}$ (≈ 55 K). [19, 54].

³The choice of the initial value of T_g^* for the fit interval is arbitrary. We observed no significant influence on the results of the fits after choosing the value of T_g^* visually or iterating after the first fitting attempts.



Figure 3.7: The values of the fitting parameters of Eq. (3.10) depending on the fitting interval ΔT . The colours denote different models: CRC - red, CRC2 - green, CRC4 - purple, FRC - blue. The dashed lines represent the final values of the parameters.

model	$T_{\rm g}$ (K)	$ ho_{ m g}~({ m g/cm^3})$	$M \times 10^{-4} \ (\text{g/cm}^3 K)$	$G \times 10^{-4} \; (g/cm^3 K)$	w (K)
CRC	192	0.967	-6.6	-2.0	50
CRC2	143	1.004	-6.9	-2.2	45
CRC4	108	1.032	-7.0	-2.5	34
FRC	43	1.089	-8.5	-4.0	31

Table 3.3: The final values of the fit parameters of Eq. (3.10)

3.3 Structure factors, compressibility

3.3.1 Lennard–Jones interactions: Comparison of polybutadiene with the Kob–Andersen binary mixture

From the viewpoint of the Lennard–Jones (LJ) interactions 1,4-*cis-trans* polybutadiene (PBD) is a binary mixture. Thus, it is interesting to compare PBD to models of glass-forming binary mixtures, an extensively studied example being the Kob–Andersen (KA) mixture [58, 102–104].

For PBD the CH_2 united atom (UA) is the big particle and CH united atom the small particle. Following commonly employed notations we refer to the big particle as the A particle (= CH_2) and to the small particle as the B particle (= CH_2).

For PBD the LJ interaction parameters are

$$\begin{split} \epsilon_{\rm AA} &= 0.107639 \, \frac{\rm kcal}{\rm mol} \,, \quad \sigma_{\rm AA} = 4.009 \, \rm \mathring{A} \,, \\ \epsilon_{\rm BB} &= 0.114999 \, \frac{\rm kcal}{\rm mol} \,, \quad \sigma_{\rm BB} = 3.385 \, \rm \mathring{A} \,, \\ \epsilon_{\rm AB} &= 0.116723 \, \frac{\rm kcal}{\rm mol} \,, \quad \sigma_{\rm AB} = 3.793 \, \rm \mathring{A} \,, \end{split}$$
(3.11)

while for the KA mixture they are [102]

$$\epsilon_{AA} = 1.0, \ \sigma_{AA} = 1,$$

 $\epsilon_{BB} = 0.5, \ \sigma_{BB} = 0.88,$
(3.12)
 $\epsilon_{AB} = 1.5, \ \sigma_{AB} = 0.8.$

These parameters imply that the particle size ratios $\delta_{\sigma} = \sigma_{\rm BB} / \sigma_{\rm AA}$,

$$\delta_{\sigma}^{\text{PBD}} = 0.844 \quad \text{and} \quad \delta_{\sigma}^{\text{KA}} = 0.88 \;, \tag{3.13}$$

are nearly the same for PBD and the KA mixture. Moreover, both systems deviate from the Lorentz rule [3], $\sigma_{ij} \neq (\sigma_{ii} + \sigma_{jj})/2$, and are thus "nonadditive". Following Eq. (8) of Ref. [57] we can quantify deviations from additivity by

$$\sigma_{ij} = \frac{1}{2} \left(\sigma_{ii} + \sigma_{jj} \right) \left[1 - f_{\sigma} |\sigma_{ii} - \sigma_{jj}| \right] \,. \tag{3.14}$$

This modified mixing rule implies that identical particles (i = j) have unaltered diameters, whereas the cross term of unlike particles deviates from additivity for $f_{\sigma} \neq 0$. If $f_{\sigma} > 0$, the small and large particles overlap more than they could for additive systems, while they overlap less for $f_{\sigma} < 0$. For PBD and the KA mixture we find

$$f_{\sigma}^{\text{PBD}} = -0.0416 \text{ \AA}^{-1} \quad \Rightarrow \quad f_{\sigma}^{\text{PBD}} \sigma_{\text{AA}} \simeq -0.1668 ,$$

$$f_{\sigma}^{\text{KA}} = 1.24 .$$
(3.15)

Compared to the KA mixture, the overlap between unlike particles is decreased for PBD, but the deviation from nonadditivity is fairly weak—weaker by a factor of about 7 relative to the KA mixture.

We can compare the LJ energy parameters of both models in a similar way. First, the LJ energy ratio $\delta_{\epsilon} = \epsilon_{\rm BB}/\epsilon_{\rm AA}$ have the following values

$$\delta_{\epsilon}^{\text{PBD}} \simeq 1.036 \quad \text{and} \quad \delta_{\epsilon}^{\text{KA}} = 0.5 \;.$$
(3.16)

For PBD the LJ energy between like particles is almost the same. This is not unexpected given that $A (= CH_2)$ and B particles (= CH) are chemically very close.

For simple mixtures the interaction energy is often modeled by the Berthelot rule [3], $\epsilon_{ij} = \sqrt{\epsilon_{ii}\epsilon_{jj}}$. It is convenient to describe deviations from this rule by an equation similar to Eq. (3.14), i.e.

$$\epsilon_{ij} = \sqrt{\epsilon_{ii}\epsilon_{jj}} \left[1 + f_{\epsilon} |\epsilon_{ii} - \epsilon_{jj}| \right].$$
(3.17)

As before, the modified mixing rule does not change the energy parameter of identical particles, but implies stronger attractions between unlike particles relative to the Berthelot rule, if $f_{\epsilon} > 0$. Stronger attractions implies enhanced mixing ability [58, 105]. Applying Eq. (3.17) to PBD and the KA mixture we get

$$f_{\epsilon}^{\text{PBD}} \simeq 6.674 \,\frac{\text{mol}}{\text{kcal}} \Rightarrow f_{\epsilon}^{\text{PBD}} \epsilon_{\text{AA}} \simeq 0.718 \,,$$

$$f_{\epsilon}^{\text{KA}} \simeq 2.243 \,.$$
(3.18)

Thus, mixing is enhanced for both models. However, as for the LJ diameters, the deviations from the standard mixing rule (Berthelot rule) is stronger for the KA system than for PBD—by a factor of about 3. Again, this difference just reflects the similarity of the A

and B particles in the case of PBD.

The KA mixture is produced by blending A and B particles which by themselves crystallize at low temperature. The nonadditive cross interactions in the KA system are specifically designed to suppress demixing and ensuing crystallization. Good glass-forming properties are thus expected. Nonetheless, composition fluctuations at various length scales are unavoidable in any mixed system. In the cold liquid these fluctuations can lead to regions containing just a single component. If such a region is large enough and long-lived enough, it will nucleate a crystal of this component. Exactly this ordering mechanism has recently been described for the KA mixture [58]. Due to this instability in favor of crystal nucleation by composition fluctuations Ref. [58] concludes that the KA mixture "has a fatal flaw as a glass former".

These findings for simple liquids invite to explore composition fluctuations also for PBD. This chapter is a first step in this direction by focusing on the partial static structure factors and the related Bhatia–Thornton structure factors. We introduce these structure factors below and apply them to PBD. A spin-off of this analysis is a temperature dependence of the isothermal compressilibity, which we also determine by a volumefluctuation method.

3.3.2 Partial static structure factors: Background for Binary Mixtures of Simple Liquids

We consider a binary mixture consisting of N particles with $N_{\rm A}$ particles of type A and $N_{\rm B}$ (= $N - N_{\rm A}$) particles of type B. The composition of the mixture is characterized by $x_{\rm A} = N_{\rm A}/N$, the number fraction of A particles, and $x_{\rm B} = N_{\rm B}/N = 1 - x_{\rm A}$, the number fraction of B particles.

Let us introduce the collective static structure factor,

$$S(q) = \frac{1}{N} \left\langle \rho(\vec{q}) \rho(-\vec{q}) \right\rangle , \qquad (3.19)$$

which is defined in terms of the coherent density fluctuations for wave vector \vec{q} ,

$$\rho(\vec{q}) = \sum_{j=1}^{N} \exp\left(\mathrm{i}\vec{q} \cdot \vec{r}_{j}\right) \quad (\text{for } q \neq 0), \tag{3.20}$$

where $\langle ... \rangle$ denotes the canonical average and \vec{r}_j is the position of particle $j \ (= 1, ..., N)$. For spatially homogeneous and isotropic systems, the structure factor depends only on the modulus of the wave vector, $q = |\vec{q}|$. In the limit $q \to 0$, S(q) is small because the fluctuations of the particle number relative to the average $\langle N \rangle \ (= N)$ are weak in a dense system (at constant chemical potential),

$$S(q \to 0) = \frac{\langle N^2 \rangle - \langle N \rangle^2}{\langle N \rangle} \ll 1.$$
(3.21)

For a system consisting only of one particle type we also have [3] $S(q \to 0) = k_{\rm B} T \rho \kappa_T$ where T is the temperature, ρ is the particle number density ($\rho = N/V$ with V the volume of the system) and κ_T is the isothermal compressibility.

Further insight into structural properties of a binary mixture can be obtained from the partial static structure factors

$$S_{\alpha\beta}(q) = \frac{1}{N} \left\langle \rho_{\alpha}(\vec{q})\rho_{\beta}(-\vec{q}) \right\rangle \quad (\alpha, \beta = A, B)$$
(3.22)

defined by the partial density fluctuations

$$\rho_{\alpha}(\vec{q}) = \sum_{j_{\alpha}=1}^{N_{\alpha}} \exp\left(\mathrm{i}\vec{q}\cdot\vec{r}_{j_{\alpha}}\right),\tag{3.23}$$

where $\vec{r}_{j_{\alpha}}$ is the position of particle j_{α} of species α . As $\rho_{\rm A}(\vec{q}) + \rho_{\rm B}(\vec{q}) = \rho(\vec{q}) \equiv \rho_{\rm n}(\vec{q})$, the collective structure factor can be expressed as

$$S(q) = S_{\rm nn}(q) = S_{\rm AA}(q) + S_{\rm BB}(q) + 2S_{\rm AB}(q). \tag{3.24}$$

While $S_{\alpha\beta}(q)$ characterize spatial correlations between like or unlike particles, S(q) describes number-number (nn) correlations (hence the notation $S = S_{nn}$). Eq. (3.24) is not the only physically significant linear combination of the partial structure factors. Since composition (or concentration) fluctuations $\rho_c(\vec{q})$ are defined by $\rho_c(\vec{q}) = \rho_A(\vec{q}) - x_A \rho(\vec{q}) = x_B \rho_A(\vec{q}) - x_A \rho_B(\vec{q})$, the structure factor

$$S_{\rm cc}(q) = \frac{1}{N} \langle \rho_{\rm c}(\vec{q}) \rho_{\rm c}(-\vec{q}) \rangle$$
$$= x_{\rm B}^2 S_{\rm AA}(q) + x_{\rm A}^2 S_{\rm BB}(q) - 2x_{\rm A} x_{\rm B} S_{\rm AB}(q)$$
(3.25)

represents composition-composition (cc) correlations, and the structure factor between ρ_n and $\rho_c,$

$$S_{\rm nc}(q) = \frac{1}{N} \langle \rho_{\rm n}(\vec{q}) \rho_{\rm c}(-\vec{q}) \rangle$$

= $x_{\rm B} S_{\rm AA}(q) - x_{\rm A} S_{\rm BB}(q) + (x_{\rm B} - x_{\rm A}) S_{\rm AB}(q),$ (3.26)

describes number-composition (nc) correlations. The structure factors $S_{nn}(q)$, $S_{cc}(q)$ and $S_{nc}(q)$ are often referred to as Bhatia–Thornton structure factors [106]. They have been studied extensively for metallic alloys [107, 108] or colloidal suspensions [109].

For binary mixtures a deviation between $S(q \to 0)$ and $k_{\rm B}T\rho\kappa_T$ is expected from the work of Bhatia and Thornton [106] and also from the Kirkwood–Buff theory for multicomponent solutions [110]. For $q \to 0$ the Bhatia–Thornton structure factors are related to the thermodynamic properties of the binary mixture:

$$S(q \to 0) = k_{\rm B} T \rho \kappa_T + \delta^2 S_{\rm cc}(q \to 0), \qquad (3.27)$$

$$S_{\rm cc}(q \to 0) = \frac{N k_{\rm B} T}{(\partial^2 G / \partial x_{\rm A}^2)_{p,T,N}},\tag{3.28}$$

$$S_{\rm nc}(q \to 0) = -\delta S_{\rm cc}(q \to 0), \qquad (3.29)$$

where G is the Gibbs free energy, p the pressure and

$$\delta = \rho(v_{\rm A} - v_{\rm B}) \tag{3.30}$$

is a volumetric factor given by the partial molar volumes

$$v_{\rm A} = \left. \frac{\partial V}{\partial N_{\rm A}} \right|_{p,T,N_{\rm B}} \quad \text{and} \quad v_{\rm B} = \left. \frac{\partial V}{\partial N_{\rm B}} \right|_{p,T,N_{\rm A}} \,.$$
 (3.31)

Eq. (3.27) shows that in a mixture fluctuations of the total particle number, $S(q \to 0)$, do not only stem from compressibility effects—that is, from the volume response of the system to a pressure variation—but also from composition fluctuations and their coupling to the number density. Since thermodynamic stability requires $(\partial^2 G/\partial x_A^2)_{p,T,N} > 0$, we have in general $S_{cc}(q \to 0) > 0$. This implies that the second term in the right-hand-side of Eq. (3.27) is positive, if $\delta \neq 0$ or $v_A \neq v_B$. The molar volumes of the two species can be calculated via the Kirkwood–Buff theory from the partial structure factors in the limit $q \rightarrow 0$ [110],

$$ov_{\rm A} = \frac{x_{\rm A} S_{\rm BB}(0) - x_{\rm B} S_{\rm AB}(0)}{x_{\rm A}^2 S_{\rm BB}(0) + x_{\rm B}^2 S_{\rm AA}(0) - 2x_{\rm A} x_{\rm B} S_{\rm AB}(0)} , \qquad (3.32)$$

$$ov_{\rm B} = \frac{x_{\rm B}S_{\rm AA}(0) - x_{\rm A}S_{\rm AB}(0)}{x_{\rm A}^2 S_{\rm BB}(0) + x_{\rm B}^2 S_{\rm AA}(0) - 2x_{\rm A}x_{\rm B}S_{\rm AB}(0)},$$
(3.33)

and similarly the theory also allows to express the compressibility as

$$k_{\rm B} T \rho \kappa_T = \frac{S_{\rm AA}(0) S_{\rm BB}(0) - [S_{\rm AB}(0)]^2}{x_{\rm A}^2 S_{\rm BB}(0) + x_{\rm B}^2 S_{\rm AA}(0) - 2x_{\rm A} x_{\rm B} S_{\rm AB}(0)} , \qquad (3.34)$$

where $S_{\alpha\beta}(0)$ is an abbreviation for $S_{\alpha\beta}(q \to 0)$.

3.3.3 Polybutadiene: Partial static structure factors, compressibility

Fig. 3.8 depicts S(q) and its decomposition $S_{AA}(q) + S_{BB}(q)$ and $2S_{AB}(q)$ according to Eq. (3.24). Panel (a) shows the results for the CRC model at T = 353 K, while panel (b) compares the CRC and FRC models. Additionally, panel (b) plots the data versus $q\sigma_{AA}$, i.e. in LJ units for comparison with simple liquids. From panel (b) we see that the



Figure 3.8: Panel (a): Collective static structure factor at T = 353 K for the CRC model and its decomposition into partial structure factors according to Eq. (3.24): $S_{AA}(q) + S_{BB}(q)$ and $2S_{AB}(q)$. Panel (b): Same as above but for both the CRC model (lines) and the FRC model (circles). In addition, the q axis is shown as $q\sigma_{AA}$ to facilitate comparison with models where results are given in LJ units.

results of the CRC and FRC models agree with each other (on the scale of the figure). In very good approximation, spatial correlations encoded in the partial structure factors are unaffected when switching off the torsional potential. However, by zooming into the data some differences become visible, in particular at small q, as we will see when discussing the Bhatia–Thornton structure factors (cf. Fig. 3.10 and Fig. 3.11). Here we focus on the qualitative behavior of the partial structure factors and compare this behavior with that of simple binary mixtures.

Fig. 3.8 shows that the PBD melt tends toward the behavior of an ideal mixture for $q \to \infty$. For an ideal mixture spatial correlations vanish, implying $S_{AB}(q \to \infty) = 0$ and $S_{AA}(q \to \infty) + S_{BB}(q \to \infty) = x_A + x_B = 1$. For large q, say $q \gtrsim 8 \text{ Å}^{-1}$ or $q\sigma_{AA} \gtrsim 32$, the structure factor S(q) is thus dominated by correlations between identical particles. The sum $S_{AA}(q) + S_{BB}(q)$ is positive for all q, whereas $S_{AB}(q)$ oscillates around 0 and remains positive for $q < q^*$ with q^* being the position of the first peak of S(q). This behavior for small q is different from the one typically found for simple binary mixtures [108, 111–113]: For simple liquids $S_{AB}(q)$ is negative for $q \approx q^*$ and goes through a minimum when q decreases. This minimum outweighs the positive contribution of $S_{AA}(q) + S_{BB}(q)$, leading to a dip in S(q) before S(q) increases toward a plateau as $q \to 0$. Such a dip is not observed for the PBD melt. Here S(q) continuously decreases—toward the compressibility plateau $S(q \to 0) = k_B T \rho \kappa_T$, as Fig. 3.9 indicates and will be discussed further.

Using the partial static structure factors and $x_{\rm A} = x_{\rm B} = 1/2$ (PBD has two CH₂ UAs and two CH UAs per monomer) we calculate $S_{\rm cc}(q)$ and $S_{\rm nc}(q)$ from Eqs. (3.25) and (3.26). Fig. 3.9 shows these structure factors together with S(q) at T = 353 K for the CRC and FRC models. Due to the small values of $S_{\rm cc}(q)$ and $S_{\rm nc}(q)$ the figure zooms on ordinate values below 1, thereby truncating the maxima of S(q). We find that $S_{\rm cc}(q)$ is positive for all q. For $q > q^*$, $S_{\rm cc}(q)$ oscillates around $x_{\rm A}x_{\rm B}$ (= 0.25)—the value expected for an ideal (equimolar) mixture—while $S_{\rm nc}(q)$ oscillates around 0 and is roughly in antiphase to $S_{\rm cc}(q)$ for large q. Both $S_{\rm cc}(q)$ and $S_{\rm nc}(q)$ exhibit a maximum for q near q^* . Qualitatively similar oscillations are also found for simple binary mixtures [107, 109, 114], reflecting the (nonuniversal) local composition-composition and number-composition correlations in these systems.

However, qualitative differences between simple binary mixtures and PBD occur in the limit $q \to 0$. While $S_{\rm cc}(q)$ and $S_{\rm nc}(q)$ tend to finite values for simple liquids [107, 109, 114], the structure factors (appear to) vanish for PBD. Vanishing of $S_{cc}(q \to 0)$ implies that there are no (thermodynamic) composition fluctuations at large scales. Such fluctuations are prevented by chain connectivity: At large scales, the character of the binary mixture disappears for PBD; PBD is a homopolymer.

What is the consequence of $S_{\rm cc}(q \to 0) = 0$ in view of the general relations of Eqs. (3.27) to (3.29)? From Eq. (3.27) we conclude that $S(q \to 0) = k_{\rm B} T \rho \kappa_T$ and from Eq. (3.29) that $S_{\rm nc}(q \to 0) = 0$, provided the volumetric factor δ [cf. Eq. (3.31)] does not diverge. Although it is physically clear that δ cannot diverge, Fig. 3.10 tests this point by parametrizing the q dependence of $\rho v_{\rm A}(q)$ and $\rho v_{\rm B}(q)$ by a polynomial fit function:

$$\rho v(q) = \rho v_0 + A_v q^2 + B_v q^4 , \qquad (3.35)$$

where ρv_0 , A_v and B_v are fit parameters. The values of δ are obtained using Eq. (3.30). They are given in Table 3.4.

Similarly, we parameterize the q dependence of $S_{cc}(q)$ and the q dependent compressibility for small q:

$$S_{\rm cc}^{\rm fit}(q) = A_{\rm cc}q^2 + B_{\rm cc}q^4$$
, (3.36)



Figure 3.9: Bhatia–Thornton structure factors at T = 353 K for the CRC model (symbols with dotted lines) and the FRC model (filled circles): $S_{\rm nn}(q) = S(q)$, $S_{\rm cc}(q)$ and $S_{\rm nc}(q)$. The horizontal dashed lines represent the large-q limits: $S_{\rm nn}(q \to \infty) = 1$, $S_{\rm cc}(q \to \infty) = x_{\rm A}x_{\rm B}$ and $S_{\rm nc}(q \to \infty) = 0$. The horizontal full line indicates $k_{\rm B}T\rho\kappa_T$ obtained from Fig. 3.11 by an extrapolation to $q \to 0$.

$$\left[k_{\rm B}T\rho\kappa_T(q)\right]^{\rm fit} = k_{\rm B}T\rho\kappa_T + A_\kappa q^2 + B_\kappa q^4 , \qquad (3.37)$$

where A_{cc} , B_{cc} , $k_{\rm B}T\rho\kappa_T$, A_{κ} and B_{κ} are adjustable parameters. The simulation results for $k_{\rm B}T\rho\kappa_T(q)$ are determined by inserting the q dependent partial structure factors into Eq. (3.34).

Fig. 3.11 shows the results of this analysis. From panel (b) we see that $S_{cc}(q)$ for the CRC and FRC models agree with each other and can be well described by Eq. (3.36). Using then $S_{cc}^{fit}(q)$ and the results for δ from Fig. 3.10, the difference for $S_{cc}(q)$ between the CRC and FRC models can be understood [cf. dashed lines in Fig. 3.11b]. Similarly,



Figure 3.10: Partial molar volumes (times density) at T = 353 K for the CRC model (open symbols) and the FRC model (filled circles). The wave vector dependence of $\rho v_A(q)$ and $\rho v_B(q)$ is defined by inserting the q dependent partial structure factors into Eq. (3.32) and Eq. (3.33). The dashed lines indicate the fits using Eq. (3.35) with estimated $q \rightarrow 0$ values, leading to the volumetric factor δ , defined in Eq. (3.30), for the CRC and FRC models.

Table 3.4: The values of the volumetric factor δ obtained from the partial molar volumes.

$T(\mathbf{K})$	$\delta_{ m CRC}$	$\delta_{ m CRC2}$	$\delta_{ m CRC4}$	$\delta_{ m FRC}$
213	0.829	1.001	1.025	1.178
225	0.905	1.012	1.045	1.176
240	0.960	1.021	1.058	1.224
253	0.865	1.017	1.099	1.217
273	0.999	1.061	1.106	1.228
293	1.029	1.078	1.102	1.260
353	1.250	1.187	1.228	1.377

Fig. 3.11a shows that the q dependent generalization of the compressibility is well described by Eq. (3.37). Within the error bars the simulation results for $k_{\rm B}T\rho\kappa_T(q)$ for both models are the same so that no separate fit was attempted.

Using $[k_{\rm B}T\rho\kappa_T(q)]^{\rm fit}$, $\delta_{\rm CRC}$ and $S_{\rm cc}^{\rm fit}(q)$ in Eq. (3.27), the solid line in Fig. 3.11a reveals that a good description of the simulation data for S(q) is obtained for the CRC model. Due to the extra term $\delta_{\rm CRC}^2 S_{\rm cc}^{\rm fit}(q)$ in Eq. (3.27), the compressibility plateau is less visible in S(q)than in $k_{\rm B}T\rho\kappa_T(q)$, for the simulated range of q values. Apparently, $k_{\rm B}T\rho\kappa_T(q)$ together with Eq. (3.37) is a more convenient method (with respect to straightforward extrapolation $S(q \to 0)$) to determine the (reduced) compressibility $k_{\rm B}T\rho\kappa_T$ from canonical simulations of binary-mixture-like polymer systems. Here this analysis gives $k_{\rm B}T\rho\kappa_T = 0.188$ at T = 353 K for the CRC model (and other models considering a single fit in Fig. 3.11a).

An alternative—and often employed—method to determine the compressibility is to monitor the volume fluctuations in NPT simulations (as a special case of the general theory of the ensemble dependence of fluctuations [115, 116])

$$\kappa_T = -\frac{1}{V} \frac{\partial V}{\partial p} \Big|_T = \frac{\langle V^2 \rangle - \langle V \rangle^2}{k_{\rm B} T \langle V \rangle} \,. \tag{3.38}$$

We employed this method to determine $k_{\rm B}T\rho\kappa_T$ from 1 μ s long NPT runs using all our models of PBD.



Figure 3.11: Panel (a): Zoom of the small q region for S(q) (circles) and $k_{\rm B}T\rho\kappa_T(q)$ (triangles) at T = 353 K for the CRC model (red) and FRC model (blue). The wave vector dependence of $\kappa_T(q)$ is defined by using the q dependent partial structure factors in Eq. (3.34). The dashed line shows a polynomial fit, denoted by $[k_{\rm B}T\rho\kappa_T(q)]^{\rm fit}$, in order to estimate the compressibility κ_T by extrapolation to $q \to 0$. The resulting value $k_{\rm B}T\rho\kappa_T = 0.188$ is indicated as a horizontal dotted line. Inserting $[k_{\rm B}T\rho\kappa_T(q)]^{\rm fit}$, $\delta_{\rm CRC} = 1.25$ from Fig. 3.10 and the fit result $S_{\rm cc}^{\rm fit}(q)$ from panel (b) into Eq. (3.27) gives the solid (black) line. Panel (b): Zoom of the small q region for $S_{\rm cc}$ (circles) and $S_{\rm nc}$ (triangles) at T = 353 K for the CRC model (red) and FRC model (blue). The solid (black) line shows a polynomial fit to $S_{\rm cc}$. The dashed lines show Eq. (3.29) when using $S_{\rm cc}^{\rm fit}$ and the volumetric factors $\delta_{\rm CRC}$ and $\delta_{\rm FRC}$ obtained from Fig. 3.10.

Fig. 3.12 shows the results for all the models (circles). The triangles presents the compressibility from Fig. 3.11. For comparison two additional results from the literature

are included in Fig. 3.12. The crosses depict MD data for *cis*-1,4 PBD from Ref. [117] and the squares experimental data taken from Ref. [118]. We find reasonable agreement between these literature and the present results with the volume fluctuations data being the most accurate. The data obtained from the partial structure factors appears to deviate significantly, even though we expect them to be more precise than a straightforward extrapolation of $S(q \rightarrow 0)$. Of course, our data for the structure factors at low q are still quite noisy even with the averaging statistics corresponding to several μ s of the simulations. However, we believe there can still be a room for improvement in the choice of the fitting function (cf. Eqs. (3.35) to (3.37)). This function was inspired by the Taylor expansions of the structure factors up to 4th order in q in Eqs. (3.28) and (3.32) to (3.34). While visually the fits look good, the post-factum look on the fitting parameters revealed that the coefficients at the q^4 term are either of the same order or even larger than the ones at the q^2 term (the values are given in Appendix F). Clearly, this suggests that the Taylor expansion does not work in this region of q. There was no clear temperature dependence of the corresponding length scales observed as well.

Fig. 3.12 plots the compressibility as $1/\sqrt{k_{\rm B}T\rho\kappa_T}$ versus 1/T. This plot format for the data was suggested by Schweizer and Saltzmann based on the Polymer-Reference-Interaction-Site theory, yielding [118]

$$\frac{1}{\sqrt{k_{\rm B}T\rho\kappa_T}} = -A + \frac{B}{T} \,. \tag{3.39}$$

Eq. (3.39) was not claimed to be "quantitatively reliable nor rigorous, but merely suggestive of how experimental data for the dimensionless compressibility might be organized" (cf. Appendix A in [118]). However, Eq. (3.39) was found to be practically relevant, since it linearizes the experimental data for numerous polymers, including PBD (cf. Figs. 4 and 5 in [118]). We reproduce the (scanned) PBD data and the fit to Eq. (3.39) from [118] in Fig. 3.12.

More recently, Mirigian and Schweizer derived Eq. (3.39) from a low-temperature expansion of a van-der-Waals (vdW) model for the equation of state [119]. Eq. (3.39) enters the mapping of a hard-sphere fluid to molecular liquids [119] and polymer melts [120]. This mapping underlies the elastically cooperative nonlinear Langevin equation (ECNLE) theory for the T dependence of the α relaxation time [119, 120] and extensions of the ECNLE theory to polymer films [32, 121, 122].



Figure 3.12: Temperature dependence of the compressibility. The data are plotted as suggested by Eq. (3.39). The orange squares are experimental results for PBD scanned from Fig. 4 of Ref. [118]. The dashed line shows Eq. (3.39) with parameters A = -0.435 and B = 987 K taken from Table I of Ref. [118]. The circles show the results for the CRC model obtained from volume fluctuations [Eq. (3.38)]. As the experimental results, they are linearized and can be fit to Eq. (3.39) (not shown). The solid line indicates the fit result to the Long–Lequeux theory obtained from a joint adjustment of the CRC data for the monomer density (inset) to Eq. (3.40) and for the compressibility to Eq. (3.41). The results for $k_{\rm B}T\rho\kappa_{T}$, found for all models are indicated by triangles. The orange crosses present the MD results from Ref. [117] for *cis*-1,4-PBD at 1 atm (systems with 128 C-atoms per chain and 32 chains), which were calculated by scanning the data from Fig. 4 and Fig. 7 of [117].

Interestingly, essentially the same vdW approach as employed by Schweizer and coworkers was proposed before by Long and Lequeux [123] to model the pressure-volumetemperature (PVT) behavior of polymer melts. Starting from a vdW-like ansatz for the free energy and imposing equilibrium conditions Long and Lequeux get the following expressions for the density

$$\rho = \frac{\rho_0}{2} \frac{1 + \sqrt{1 - T/T_c}}{1 + \beta (1 - \sqrt{1 - T/T_c})/2}$$
(3.40)

and the (inverse) compressibility

$$\frac{1}{\kappa_T} = \frac{k_{\rm B} T \rho_0}{2\gamma^2} \left[1 + \sqrt{1 - \frac{T}{T_{\rm c}}} \right]^3 \times \left[\frac{1}{\left(1 - \sqrt{1 - T/T_{\rm c}}\right)^2} - \frac{T_{\rm c}}{T} \right] \,. \tag{3.41}$$

Here ρ_0 is the maximum close-packing density and T_c the temperature at which air would

become a good solvent for the polymer. Clearly, this (hypothetical) temperature should be very high so that $T_{\rm g} \leq T \ll T_{\rm c}$ is the temperature regime where the theory is expected to hold. The other two parameters, β and γ , are numbers introduced to account for the incompressibility of the monomers.

A low-*T* expansion of Eqs. (3.40) and (3.41) gives back Eq. (3.39) with $B = 4T_c/\gamma$ and $A = (5 - \beta)/2\gamma$. Moreover, Ref. [123] showed that Eqs. (3.40) and (3.41) provide a consistent description of *T* dependence of ρ and κ_T for supercooled polymer melts down to the glass transition. Therefore, we fit Eqs. (3.40) and (3.41) to ρ and κ_T of all our models. As shown in Fig. 3.12, the fits give good agreement, yielding values for the parameters $(\rho_0, T_c, \beta, \gamma)$ that are in reasonable accord with literature results for other hydrocarbon polymers, such as polystyrene or poly(isobutylene) [123].

Table 3.5: The values of the fit parameters of the Long-Lequeux theory for the density and compressibility.

model	$T_{\rm c}~({\rm K})$	$\rho_0~({\rm g/cm^3})$	β	ρ_0/γ^2
CRC	920.751	1.104	1.251	3.902896e + 27
CRC2	1054.873	1.109	1.692	2.773495e+27
CRC4	997.521	1.108	1.493	3.190466e + 27
FRC	1087.512	1.108	1.851	2.516277e+27

3.4 Energies and heat capacity

Let x denote the microstate of the PBD melt containing N_c polymers with N united atoms each. The hamiltonian is written as

$$\mathcal{H}(x) = K(x) + U_{\text{tot}}(x) , \qquad (3.42)$$

where K is the total kinetic energy and U_{tot} is the total potential energy consisting of valence terms (bond, bond angle, dihedral) and pair (Lennard–Jones) interactions.

As explained in Section 2.3.2, the simulations are carried out in two steps: First, NPT relaxation to get the average volume $V = V(T, p, N_c N)$ corresponding to temperature T and pressure p (= 0). Second, NVT production runs by imposing the volume $V(T, p, N_c N)$.

During these canonical production runs the internal energy U is determined by

 $U = \langle \mathcal{H}(x) \rangle$. However, since $V = V(T, p, N_c N)$, we have (for simplicity the dependence on $N_c N$ is suppressed in the equations below)

internal energy
$$U = U(T, V(T, p)) = U(T, p)$$
, (3.43)

enthalpy
$$H = U(T, p) + pV(T, p)$$
. (3.44)

For p = 0 the enthalpy is given by U(T, p).

In the following sections we discuss the contributions to U(T, p) from the potential energy of PBD before we turn to heat capacity.

3.4.1 Hard variables: Bond length

The bond length ℓ is subjected to the harmonic bond potential defined by

$$U_{\rm bond}(\ell) = K_{\rm bond}(\ell - \ell_0)^2 , \qquad (3.45)$$

where K_{bond} is (half) the force constant and ℓ_0 the equilibrium bond length. The parameters for these constants are given in Table 2.1.

In Section 2.4.2 we observed an agreement of the bond length distribution obtained from the simulations with the one for independent bonds (i.e. with the distribution determined only by Eq. (3.45)). This allows us to assume that the bonds are independent of each other and also independent of the potential energies for bond angles, torsion angles and pair interactions. Then, the partition function for a bond reads

$$Z_{\text{bond}}(\beta) = 4\pi \int_0^\infty \mathrm{d}\ell \,\ell^2 \exp\left[-\beta K_{\text{bond}}(\ell-\ell_0)^2\right]$$

$$\simeq 4\pi \ell_0^2 \sqrt{\frac{\pi}{\beta K_{\text{bond}}}} , \qquad (3.46)$$

where $\beta = 1/(k_{\rm B}T)$. The last approximate equality was obtained by utilizing Eq. (3.462/7) from Ref. [124] and the fact that the dimensionless quantity $(k_{\rm B} = 1.9872067 \times 10^{-3} \text{ kcal/(mol,K)})$

$$\beta K_{\text{bond}} \ell_0^2 = \frac{2500}{1.9872067 \times 10^{-3}} \frac{\ell_0^2}{T}$$
(3.47)

is large because $\beta K_{\rm bond} \ell_0^2 \simeq 6500$ even for the smallest bond length, $\ell_0 = 1.34$ Å, and the

highest temperature studied, T = 353 K. From Eq. (3.46) we get the average energy per bond

$$\langle U_{\rm bond} \rangle = -\frac{\partial \ln Z_{\rm bond}(\beta)}{\partial \beta} \simeq \frac{1}{2} k_{\rm B} T , \qquad (3.48)$$

as expected for a one-dimensional harmonic oscillator according to the equipartition theorem.

PBD has three types of bonds:

 1st bond (b1) = CH-CH double bond: There is one such bond per monomer. So the total number of b1 bonds is given by

$$N_{\mathrm{b1}} = \# \text{ of monomers} = n = \frac{N}{4}$$
 ,

where $n \ (= 29)$ is the number of monomers per chain and $N \ (= 116)$ is the number of united atoms per chain.

2nd bond (b2) = CH-CH₂ bond: There are two such bonds per monomer. So the total number of b2 bonds is given by

$$N_{\mathrm{b}2} = 2n = \frac{N}{2} \; . \label{eq:Nb2}$$

• 3^{rd} bond (b3) = CH₂-CH₂ bond: This bond links consecutive monomers. So the total number of b3 bonds is given by

$$N_{\rm b3} = n - 1 = \frac{N}{4} - 1 \; .$$

When adding up the energies for all bonds in the melt and using Eq. (3.48), we find for the average extensive bond energy (E_{bond}) of the system

$$E_{\rm bond}(T, p, N_{\rm c}N) \simeq N_{\rm c}N\left(1 - \frac{1}{N}\right)\frac{1}{2}k_{\rm B}T\,,$$
 (3.49)

where $N_{\rm c}$ (= 40) is the number of chains in the system.

Fig. 3.13 shows the *T* dependence of the intensive bond energy $e_{\text{bond}} = E_{\text{bond}}/(N_c N)$. Two data sets are compared: results from cooling runs with rate = 0.83 K/ns and results for $T \ge 213$ K after NPT and NVT equilibration. We see that both data sets agree well for $T \ge 213$ K. Moreover, the main panel of Fig. 3.13 suggests that Eq. (3.49) gives an



Figure 3.13: Intensive bond energy $e_{\rm bond}$ (= $E_{\rm bond}/(N_{\rm c}N)$) versus temperature T for the CRC and FRC models. Results from continuous cooling runs with rate $\Gamma = 0.83$ K/ns (colored circles) are compared to equilibrium data obtained by NPT and NVT equilibration for 213 K $\leq T \leq 353$ K (colored crosses). The colors mark CRC (red) and FRC (blue) models. The black dashed line depicts Eq. (3.49). The vertical dotted line indicates $T_{\rm g} = 192$ K of the CRC model as determined by fitting Eq. (3.10) to the density. Inset: Plot of the ratio $e_{\rm bond}/[(1-1/N)k_{\rm B}T/2]$ versus T using the same data as in the main panel.

excellent description of the T dependence not only for the equilibrated polymer liquid, but also for the glass down to lowest T studied. The inset provides a more critical test of this point. If Eq. (3.49) was valid for all T, the ratio $e_{\text{bond}}/[(1-1/N)k_{\text{B}}T/2]$ should fluctuate around 1. While this is the case for $T \gtrsim 200$ K, the ratio slightly, but systematically, increases as $T \rightarrow 0$, leading to a larger bond energy than expected from Eq. (3.49). However, this deviation from equilibrium is much weaker than for the other potential energies, as we will see below.

3.4.2 Hard variables: Bond angle

An approach analogous to that of the bond length can be used to estimate the average energy of the bond angle. The bond angle θ is subjected to the potential

$$U_{\text{bend}}(\theta) = K_{\text{bend}}(\cos\theta - \cos\theta_0)^2 , \qquad (3.50)$$

where the values for K_{bend} and θ_0 are given in Table 2.2. As for the bond length, the agreement of the bond angle distribution obtained from the simulations (see Section 2.4.2) with the ones for intependent angles (i.e. with the distribution determined only by Eq. (3.50)) justifies the assumption of the angles being independent of each other and of the other variables. The partition function of an angle is then given by

$$Z_{\text{bend}}(\beta) = \int_0^{\pi} d\theta \sin \theta \exp\left[-\beta K_{\text{bend}}(\cos \theta - \cos \theta_0)^2\right]$$

$$= \frac{1}{2}\sqrt{\frac{\pi}{q}} \left[\operatorname{erf}\left([1 - x_0]q\right) + \operatorname{erf}\left([1 + x_0]q\right)\right],$$
(3.51)

where $\operatorname{erf}(x)$ is the error function, $q = \sqrt{\beta K_{\text{bend}}}$ and $x_0 = \cos \theta_0$. With the values from Table 2.2 one finds $\operatorname{erf}([1 \pm x_0]q) \simeq 1$ and so

$$Z_{\text{bend}}(\beta) \simeq \sqrt{\frac{\pi}{\beta K_{\text{bend}}}}$$
 (3.52)

For the average energy of a bond angle this implies

$$\langle U_{\rm bend} \rangle = -\frac{\partial \ln Z_{\rm bend}(\beta)}{\partial \beta} \simeq \frac{1}{2} \, k_{\rm B} T \,. \tag{3.53}$$

The PBD model has two types of bond angles:

• 1^{st} angle $(a1) = CH_2$ -CH-CH angle: There are two angles per monomer, leading to

$$N_{\rm a1}=2n=\frac{N}{2}$$

for the total number of a1 angles.

 2nd angle (a2) = CH-CH₂-CH₂ angle: A single monomer has no a2 angle, two monomers have 2 a2 angles, three monomers have 4 a2 angles, etc. So the total number of a2 angles is given by

$$N_{\rm a2} = 2(n-1) = \frac{N}{2} \left(1 - \frac{4}{N}\right) \; . \label{eq:n2}$$

Adding up the energies for all angles in the system and using Eq. (3.53), we get for


Figure 3.14: Intensive bond angle energy $e_{\text{bend}} (= E_{\text{bend}}/(N_c N))$ versus temperature T for the CRC and FRC models. Results from continuous cooling runs with rate $\Gamma = 0.83$ K/ns (colored circles) are compared to equilibrium data obtained by NPT and NVT equilibration for $213 \text{ K} \leq T \leq 353 \text{ K}$ (colored crosses). The colors mark CRC (red) and FRC (blue) models. The black dashed line depicts Eq. (3.54). The vertical dotted line indicates $T_{\rm g} = 192 \text{ K}$ of the CRC model as determined by fitting Eq. (3.10) to the density. Inset: Plot of the ratio $e_{\rm bend}/[(1-2/N)k_{\rm B}T/2]$ versus T using the same data as in the main panel. For $T \to 0$ the ratio increases to about 1.6 for the FRC model and to about 5.7 for the CRC model.

the average extensive bond-angle energy

$$E_{\rm bend}(T, p, N_{\rm c}N) \simeq N_{\rm c}N\left(1 - \frac{2}{N}\right)\frac{1}{2}k_{\rm B}T.$$
 (3.54)

This is the same result as for the bond length, bearing the (small 1/N) finite-chain length correction.

Fig. 3.14 shows the results for the bond angle; they are similar to those for bond length. The intensive energies e_{bend} (= $E_{\text{bend}}/(N_{\text{c}}N)$) from the cooling runs and equilibrium simulations agree with each other for $T \ge 213$ K and also with Eq. (3.54). However, for $T \to 0$ systematic deviations from Eq. (3.54) are clearly visible and much larger than for the bond length. The onset of these deviations occur in the glass transition zone around T_{g} , as illustrated for the CRC model.

3.4.3 Soft variables: Dihedral angles

As a start to the analysis of the dihedral angles, we can make the same assumption as for the bond lengths and bond angles: we take the dihedral angles to be independently distributed according to the Boltzmann weight of their potential energies. Section 3.1.1 compares the predicted distributions to those from the MD simulations and concludes the following :

• Double bond *cis* and *trans* dihedrals

The simulated distributions agree with the predictions from the Boltzmann weights for the CRC model and the reduced barrier models (Fig. 3.1a). This is expected, since the torsions of the double bonds are in fact hard variables. Their potentials are very stiff, allowing only for weak vibrations around the planar configuration at all temperatures studied. The vibrations contribute $k_{\rm B}T/2$ per double bond to the average dihedral energy.

• α trans and β dihedrals

The potential energies of these dihedrals have pronounced minima but the barriers between them are more than an order magnitude smaller than well depth of the double bond dihedrals. Due to this softening of the potentials some deviations from the predicted distributions must be expected. Indeed, Fig. 3.1b show that the predictions from the Boltzmann weights are not as good as for the double bonds. Still, the agreement is semiquantitative.

• α cis dihedrals

Fig. 3.1d show strong deviations between predicted and simulated distributions. In particular, the simulated distributions tend to 0 for $\phi \to 0^{\circ}$ and $\phi \to 360^{\circ}$, whereas the predicted distributions have a maximum for these limiting angles. Clearly, the assumption of independent dihedrals fails here. Rotation around the α *cis* bond leads to strong steric hindrance due to nonbonded pair interactions for $\phi \to 0^{\circ}$ and $\phi \to 360^{\circ}$, which suppresses the occurrence of these limiting angles. This influence of the nonbonded interactions is similar to the "pentane effect" for polyethylene [91].

A harmonic expansion around the minima of the dihedral potentials cannot be sufficient to describe the T dependence of the average torsion energy $E_{tor}(T, p, N_c N)$ in the polymer liquid. Anharmonic effects need to be taken into account. A possible scheme



Figure 3.15: Intensive energy of the dihedral angles $e_{\rm tor}$ (= $E_{\rm tor}/(N_{\rm c}N)$) versus T for the CRC model. Results from continuous cooling runs with rate $\Gamma = 0.83$ K/ns (red circles) are compared to equilibrium data obtained by NPT and NVT equilibration for 213 K $\leq T \leq$ 353 K (red crosses). The black dashed line depicts a fit to the equilibrium data with Eq. (3.55) (fit interval: 213 K $\leq T \leq$ 353 K), yielding $A_{\phi} = -1.231379$ kcal/mol, $B_{\phi} = 1.393 \times 10^{-3}$ kcal/(mol K) and $C_{\phi} = -4.644 \times 10^{-7}$ kcal/(mol K)². In the glassy phase the black solid line indicates the fit result to $e_{\rm tor}(T) = A_{\phi}^{\rm g} + B_{\phi}^{\rm g}T$ with $A_{\phi}^{\rm g} = -1.104315$ kcal/mol, $B_{\phi}^{\rm g} = 6.408 \times 10^{-4}$ kcal/(mol K) (fit interval: 3.5 K $\leq T \leq$ 120 K). The vertical dotted line indicates $T_{\rm g} = 192$ K as determined by fitting Eq. (3.10) to the density. Inset: Plot of $(e_{\rm tor} - A_{\phi})/(B_{\phi}T + C_{\phi}T^2)$ versus T to examine the quality of the fit with Eq. (3.55).

to include anharmonicity is to expand $E_{tor}(T, p, N_c N)$ in powers of T:

$$E_{\rm tor}(T, p, N_{\rm c}N) = N_{\rm c}N \Big[A_{\phi}(p) + B_{\phi}(p)T + C_{\phi}(p)T^2 + \mathcal{O}(T^3) \Big] .$$
(3.55)

This equation looks like a Taylor expansion around T = 0, where the coefficients $A_{\phi}(p)$, $B_{\phi}(p)$, $C_{\phi}(p)$, ...then depend only on the remaining variable p; it is inspired by similar approaches used in the potential energy description of supercooled liquids and glasses (cf. Sect. 6.2 of [125]).

Fig. 3.15 shows that including only the first anharmonic correction (C_{ϕ}) in Eq. (3.55) suffices to give an excellent fit to the simulation data in the polymer liquid (cf. inset of Fig. 3.15). Numerically, $|C_{\phi}T|$ is by about a factor of 10 smaller than B_{ϕ} , implying that the anharmonic correction is relatively weak. The fit result for B_{ϕ} (= 1.393 × 10^{-3} kcal/(mol K)) has a reasonable value. To see this, let us calculate the energy in harmonic approximation by considering only those dihedrals with pronounced minima (double bonds, α trans and β). Each such dihedral contributes $k_{\rm B}T/2$ to the energy. Since there are n = N/4 monomers per chain and the fraction of α trans dihedrals is $f_{\rm trans} = 0.55$, we have for one chain

- *n* double bonds having 1 minimum,
- n-1 beta dihedrals with 3 minima,
- $2nf_{\rm trans}$ alpha-*trans* dihedrals with 3 minima. Then, the harmonic contribution to $E_{\rm tor}(T,p,N_{\rm c}N)$ reads

$$\begin{split} N_{\rm c} \frac{N}{4} \Big[1 + 3(1 - 4/N) + 6 f_{\rm trans} \Big] \frac{1}{2} k_{\rm B} T &= N_{\rm c} N B_{\phi}^{\rm approx} \, T \\ \Rightarrow \quad B_{\phi}^{\rm approx} &= 1.788 \times 10^{-3} \, \rm kcal/mol \, K \; , \end{split}$$

which is close to the fit value for B_ϕ from Fig. 3.15.

The impact of the glass transition is clearly visible from the main panel of Fig. 3.15. Below 213 K the cooling process is too fast for the melt to stay at equilibrium. Accordingly, the dihedral energy at low T is larger than the extrapolated liquid curve. For $T \rightarrow 0$ a harmonic approximation provides an excellent description of $E_{tor}(T)$ (cf. solid line in Fig. 3.15). However, the fitted ground state energy, $A_{\phi}^{\rm g} = -1.104315$ kcal/mol, is much larger than the one expected from torsional potential, which is given by the energy minimum at 180° of the β dihedral (= -5.28281 kcal/mol). This exemplifies the nonequilibrium character of the glass created by the cooling process.

3.4.4 Soft variables: Pair interactions

From the viewpoint of the pair interactions the PBD model corresponds to a binary Lennard–Jones (LJ) mixture (Section 3.3.1). It is therefore interesting to survey in the literature how the excess energy of LJ and other simple liquids has been described.

In this respect, an influential theory was developed by Rosenfeld and Tarazona (RT) [126]. The RT theory starts from an accurate approximation for the excess free energy functional of densely packed hard spheres (so-called fundamental measure functionals) and uses thermodynamic perturbation theory to extend the theory to continuous ("soft") potentials. First-order perturbation theory around $\eta = 1$ packing fraction gives for the excess internal energy (E^{ex}) of a three-dimensional classical bulk fluid with N particles

the following constant volume (particle density ρ) and constant temperature expression (cf. Eq. (34) in Ref. [126]):

$$\frac{E^{\rm ex}(\rho,T)}{N} = A_{\rm RT}(\rho) + B_{\rm RT}(\rho)T^{3/5} + \cdots \ . \label{eq:expansion}$$

The leading $T^{3/5}$ term is expected to be accurate for large densities (near freezing) and predominanly repulsive interactions (see the detailed discussion in [126]). It implies that the excess isochoric heat capacity ($C_V^{\text{ex}} = \partial E^{\text{ex}} / \partial T|_{V,N}$) increases with decreasing temperature as $C_V^{\text{ex}} \propto T^{-2/5}$.

A detailed simulation study of the *T* dependence of C_V^{ex} was carried out in Ref. [127] for 18 model liquids with different stoichiometric compositions, molecular topologies and interactions. It was found that the RT expression $C_V^{\text{ex}} \propto T^{-2/5}$ provides a better approximation for liquids with strong correlations between equilibrium fluctuations of virial and potential energy, i.e., for so-called "Roskilde-simple" liquids [128]. The Kob–Andersen binary LJ mixture [58, 102–104] is a Roskilde-simple liquid (for $\rho = 1.2$ in LJ units) [127]. It is therefore tempting to test whether the following ansatz,

$$E_{\text{pair}}(T, p, N_{\text{c}}N) \simeq N_{\text{c}}N \Big[A_{\text{RT}}(p) + B_{\text{RT}}(p)T^{3/5} \Big] ,$$
 (3.56)

can provide a good description for the pair interactions of the PBD model.

Fig. 3.16 shows that the description is indeed excellent. The fit to Eq. (3.56) was carried out for the equilibrated polymer liquid in the interval $213 \text{ K} \leq T \leq 353 \text{ K}$ (dashed line), where the pair energies of the CRC and FRC models nearly coincide. Interestingly, the extrapolation of the fit to low T (solid line) agrees with the FRC results from the cooling run down to the temperature regime where the glass transition of the FRC model occurs (see also the inset in Fig. 3.16 supporting the quality of the fit).

The impact of the glass transition is clearly visible in the main panel of Fig. 3.16. For the CRC model we find, as for the dihedral angles, that the cooling process is too fast for melt to stay at equilibrium below 213 K. Accordingly, the pair energy at low T is larger than the extrapolated liquid curve. For $T \rightarrow 0$ a harmonic approximation provides again an excellent description of the energy (cf. dash-dotted line in Fig. 3.16), as expected for a solid.



Figure 3.16: Intensive LJ energy e_{pair} (= $E_{\text{pair}}/(N_c N)$) versus T for the CRC and FRC models. Results from continuous cooling runs with rate $\Gamma = 0.83$ K/ns (colored circles) are compared to equilibrium data obtained by NPT and NVT equilibration for 213 K $\leq T \leq 353$ K (colored crosses). The green dashed line depicts the result of a fit of Eq. (3.56) to the equilibrium data in the interval 213 K $\leq T \leq 353$ K. The fit yields $A_{\text{RT}} = -1.8340 \,\text{kcal/mol}$ and $B_{\text{RT}} = 0.0223 \,(\text{kcal/mol} \,\text{K})^{3/5}$. The thin full line shows the extrapolation of this fit to lower T. In the glassy phase of the CRC model the dash-dotted line indicates the fit result to $e_{\text{pair}}(T) = A_{\text{pair}}^{\text{g}} + B_{\text{pair}}^{\text{g}}T$ with $A_{\text{pair}}^{\text{g}} = -1.409614 \,\text{kcal/mol}$, $B_{\text{pair}}^{\text{g}} = 5.286 \times 10^{-4} \,(\text{kcal/mol} \,\text{K})$ (fit interval: $3.5 \,\text{K} \leq T \leq 120 \,\text{K}$). The vertical dotted line indicates $T_{\text{g}} = 192 \,\text{K}$ for the CRC model as determined by fitting Eq. (3.10) to the density. Inset: Plot of the ratio E_{pair} divided by Eq. (3.56) versus T to examine the quality of the fit of the equilibrium data by Eq. (3.56).

3.4.5 Heat capacity of the polymer liquid: results for the CRC model

The thermodynamic relation between the (extensive) heat capacities at constant pressure C_p and at constant volume C_V is given by

$$C_p = C_V + T V \frac{\alpha^2}{\kappa_T} = C_V + N_{\rm c} N \frac{T \alpha^2}{\rho \kappa_T} , \qquad (3.57)$$

,

where α is the thermal expansion coefficient,

$$\alpha = \left. \frac{1}{V} \frac{\partial V}{\partial T} \right|_{p, N_{\rm c} N}$$

 κ_T the isothermal compressibility,

$$\kappa_T = \left. -\frac{1}{V} \frac{\partial V}{\partial p} \right|_{T, N_c N}$$

and $\rho = N_c N/V$ is the number density of united atoms in the system (we recall that N = 4n where n is the number of monomers per chain).

If we consider the heat capacity as the ability of a material to take up energy and to statistically distribute this energy over all degrees of freedom, it is natural to report the specific heat $(c = C/(N_c N))$ divided by k_B because this quantity can be interpreted as the number of degrees of freedom per particle (united atom) contributing to the heat capacity at temperature T.

Fig. 3.17 shows such a plot for the CRC model. The specific heat at constant pressure c_p was obtained from the fluctuations of the enthalpy H = U + pV in the NPT runs (circles)⁴. Alternatively, c_p can also be calculated by

$$c_p = \frac{1}{N_{\rm c}N} \left. \frac{\partial H}{\partial T} \right|_{p,N_{\rm c}N} = \frac{1}{N_{\rm c}N} \left. \frac{\partial U}{\partial T} \right|_{p,N_{\rm c}N} \,, \tag{3.58}$$

where $U(T, p, N_c N)$ is the internal energy. The pV term of H does not contribute because the NPT simulations are performed at p = 0. The internal energy is given by

$$U = E_{\rm kin} + E_{\rm bond} + E_{\rm bend} + E_{\rm tor} + E_{\rm pair}$$
(3.59)

with $E_{\rm kin}$ being the (average extensive) kinetic energy,

$$E_{\rm kin} = \langle K \rangle = \frac{3}{2} N_{\rm c} N k_{\rm B} T \, . \label{eq:kin}$$

Adding $E_{\rm kin}$ and the energies for the bond length [Eq. (3.49)] and bond angle [Eq. (3.54)] gives the following contribution to the specific heat

$$\frac{1}{N_{\rm c}N} \left. \frac{\partial}{\partial T} (E_{\rm kin} + E_{\rm bond} + E_{\rm ang}) \right|_{p,N_{\rm c}N} = \left[\frac{5}{2} \left(1 - \frac{3}{5N} \right) \right] k_{\rm B} = 0.00494232 \, [\rm kcal/mol/K] \,.$$
(3.60)

⁴The enthalpy $H(x) = \mathcal{H}(x) + pV(x)$ depends on the microstate x of the system. The specific heat is given by $c_p = (\langle H^2 \rangle - \langle H \rangle^2)/(N_c N k_B T^2)$. However, the NPT simulations were carried out at p = 0 pressure so that $H(x) = \mathcal{H}(x)$ and c_p can be calculated from the fluctuations of the hamiltonian \mathcal{H} only.



Figure 3.17: Specific heat at constant pressure c_p and constant volume c_V versus T from equilibrium simulations of the CRC model. The symbols present the results from enthalpy fluctuations for c_p (circles) and energy fluctuations for c_V (squares). The solid line indicates c_p obtained when inserting Eqs. (3.60) to (3.62) into Eq. (3.58). The dashed line shows c_V calculated from Eq. (3.64) as discussed in the text. The horizontal dotted line indicates the sum of Eq. (3.60) and B_{ϕ} from Eq. (3.61), yielding the constant 3.188 $k_{\rm B}$. When adding Eq. (3.62) to this constant one obtains the dash-dotted line. The difference between the circles and the dash-dotted line shows the contribution of the T dependent part of the dihedrals. Inset: Comparison to experiment. The circles and solid line are the results for c_p from the main panel. The red dashed line shows Eq. (3.65) (divided by 4R with Rbeing the gas constant) representing the experimental data from Ref. [129]. The green squares depict the results from Eq. (3.67) using the experimental data (divided by 4R) for the solid state from Fig. 9 of [129] as a proxy to account for contributions from quantum mechanical vibrations in the experimental system.

For the "soft" dihedral and pair interactions we find from Eq. (3.55) and Eq. (3.56)

$$\frac{1}{N_{\rm c}N} \left. \frac{\partial E_{\rm dih}}{\partial T} \right|_{p,N_{\rm c}N} = B_{\phi} - 2|C_{\phi}|T\,, \tag{3.61}$$

$$\frac{1}{N_{\rm c}N} \left. \frac{\partial E_{\rm pair}}{\partial T} \right|_{p,N_{\rm c}N} = \frac{3}{5} B_{\rm RT} T^{-2/5} . \tag{3.62}$$

By adding Eq. (3.60), Eq. (3.61) and Eq. (3.62) the solid line in Fig. 3.17 is obtained, which agrees with the result from the enthalpy fluctuations. Of course, this agreement is expected for systems at thermal equilibrium. Here it illustrates the internal consistency of the simulation, i.e., of the thermostating/barostating methods and the length of the equilibration and production runs.

3.4. ENERGIES AND HEAT CAPACITY

The splitting of the internal energy into its contributions according to Eq. (3.59) allows to obtain further insight into the T dependence of c_p . With the numerical value for B_{ϕ} from Fig. 3.15 the first term of Eq. (3.61) adds a constant contribution to the specific heat of about $0.7k_{\rm B}$. Together with Eq. (3.60) this implies an overall constant contribution of about $3.188k_{\rm B}$ to the specific heat, resulting from the kinetic energy and hard variables (bond length, bond angle, and B_{ϕ}). This baseline accounts for the lion's share of the specific heat (dotted line in Fig. 3.17), while the T dependent parts from Eq. (3.61) and Eq. (3.62) represent less than one $k_{\rm B}$ of c_p . These parts are dominated by the pair interactions, as seen from dash-dotted line in Fig. 3.17.

The specific heat at constant volume can be calculated from the fluctuations of the hamiltonian \mathcal{H} in the NVT simulations

$$c_V = \frac{1}{N_{\rm c}N} \frac{1}{k_{\rm B}T^2} [\langle \mathcal{H}^2 \rangle - \langle \mathcal{H} \rangle^2] \;. \tag{3.63}$$

The squares in Fig. 3.17 show the result of this calculation. As expected, $c_V < c_p$. Alternatively, c_V can be obtained from c_p via Eq. (3.57)

$$c_V = \frac{1}{N_{\rm c}N} \left. \frac{\partial H}{\partial T} \right|_{p,N_{\rm c}N} - \frac{T\alpha^2}{\rho\kappa_T} \,. \tag{3.64}$$

We know the first term. The second term can be calculated independently from the simulations or from the model proposed by Long and Lequeux [123]. In a previous chapter we showed that the Long-Lequeux model provides a very good description of the T dependence of ρ and κ_T . Therefore, we employ this model here. With this choice the right-hand side of Eq. (3.64) can be calculated analytically. The dashed line in Fig. 3.17 shows that the result of this calculation agrees with Eq. (3.63), as it should be for thermally equilibrated systems. Here this agreement illustrates again the internal consistency of the MD simulations.

3.4.6 Heat capacity of the polymer liquid: comparison with experiment

The inset of Fig. 3.17 compares the simulation results for c_p (circles with solid line) with the experimental heat capacity of liquid 1,4 PBD (dashed line) [129]. The

experimental data refer to *cis*-PBD and *trans*-PBD. Both polymers crystallize, but have identical c_p in the liquid phase. To parametrize the *T* dependence of liquid PBD Ref. [129] recommends the following equation for the specific heat per monomer:

$$c_p = 52.63 + 0.178 T \, [\text{J/(mol K)}]$$
 (3.65)

Since the experimental c_p is reported per monomer, the dashed line in the inset shows Eq. (3.65) divided by 4R with R being the gas constant. We see that the order of magnitude is the same for both simulation and experiment, but the T dependence is different: The simulated c_p increases upon cooling, whereas the experimental c_p decreases.

How can this difference be explained? A comprehensive discussion of the thermal analysis of polymers is provided in the book by Wunderlich [130]. Section 2.3.9 of [130] reports that the heat capacity of many liquid polymers increases linearly with temperature. Eq. (3.65) thus represents a typical behavior found in experiment. The linear *T* dependence is interpreted as resulting from the superposition of three contributions [131]:

$$C_p = C_{\rm vib} + C_{\rm conf} + \frac{TV\alpha^2}{\kappa_T} \;, \tag{3.66}$$

where $C_{\rm vib}$ denotes the vibrational heat capacity at constant volume and $C_{\rm conf}$ the contribution to C_p due to conformational rotations of the chain backbone. The analysis for several hydrocarbon polymers in [131] shows that, even in the liquid, the major contribution to C_p comes from $C_{\rm vib}$. As explained in [129–131], $C_{\rm vib}$ can be calculated from the vibrational spectrum of a polymer by considering two modes of vibration: "skeletal modes", involving torsion vibrations, bond-angle vibrations as well as collective vibrations along the chain backbone, and "group modes", involving vibrations of small groups of atoms, e.g. stretching vibrations of the CC and CH bonds, bending vibrations of the angle in CH₂, etc. (see Fig. 2.48 on p. 123 in [130] and Table 4 in [129]). To calculate $C_{\rm vib}$ both modes are treated quantum mechanically, based on the Debye and Einstein theories for the heat capacity. The Debye and Einstein theories predict the heat capacity to increase toward the classical limit with increasing *T*. Applications reveal that this classical limit is not yet reached in the experimentally studied *T* regime of the polymer liquid (cf. [131] and p. 138 in [130]). Therefore, $C_{\rm vib}$ still increases with *T*. Although $C_{\rm conf}$ and $TV\alpha^2/(\rho\kappa_T)$ are found to decrease upon heating, this decrease is weaker than the increase of $C_{\rm vib}$. Therefore, the sum of the three contributions still leads to an increase of C_p with T, which can be fit by linear T dependence.

Clearly, our classical united-atom model must fail to reproduce this behavior: Hydrogen atoms are not explicitly taken into account and quantum mechanical effects are absent. In our case, $c_{\rm vib}$ is a constant given by the classical result as a multiple of $k_{\rm B}$.

If quantum mechanical skeletal and group vibrations are really the cause of the discrepancy between the simulated and experimental c_p , can we approximately correct for these missing effects? As an attempt to do so, we scanned the heat capacity data for the solid state from Fig. 9 of [129] and used them as a proxy for the vibrational contribution $(c_{\rm vib}(T))$ to replace the terms $c_{\rm vib} + \frac{T\alpha^2}{\rho\kappa_T}$. The resulting prediction for the experimental specific heat then reads

$$c_p \approx c_{\rm vib}(T) + \left[\frac{3}{5} \frac{B_{\rm RT}}{T^{2/5}} - 2|C_\phi|T\right] \ , \eqno(3.67)$$

where the term [...] stems from Eqs. (3.61) and (3.62). This term replaces C_{conf} in Eq. (3.66) because it accounts for contributions to c_p from dihedral rotations and pair interactions in the polymer liquid.

The squares in the inset of Fig. 3.17 show that Eq. (3.67) is indeed reasonable. Using $c_{\rm vib}(T)$ from experiments inverts the *T* dependence originally found from the simulation, now yielding semiquantitative agreement with experiment (squares in the inset). Our findings are similar to those published recently by Honguy et al [132], where an agreement of the heat capacity extracted from MD simulations with the experimental data was achieved after taking into account the corrections for quantum effects.

3.5 Rouse theory analysis

3.5.1 Introduction

Being one of the simplest theoretical polymer models, the Rouse model [5] is still extensively used for studying polymer systems in experiments (typically neutron scattering [133–135]) and simulations [23, 40, 41, 136–139]. The model considers a polymer chain as a sequence of beads connected by harmonic springs, that is surrounded by a (structureless) Langevin thermostat imposing in addition to temperature T also a friction coefficient (both parameters are coupled to a random force via the fluctuation-dissipation theorem) [1]. It is expected to provide a suitable description of the properties of unentangled polymer melts on length scales larger than the monomer size and on the time scales larger than the associated monomer relaxation time. Yet, since the excluded volume and chain stiffness are not taken into account, deviations from real polymers are also expected at small scales [136].

Analytically the Rouse model provides a framework of describing the motion of a chain using the discrete Rouse modes $\vec{X}_p(t)$, defined as [5]

$$\vec{X}_{p}(t) = \frac{1}{N} \sum_{n=1}^{N} \vec{r}_{n}(t) \cos\left[\frac{(n-\frac{1}{2})p\pi}{N}\right] (p=0,\dots,N-1)$$
(3.68)

with $\vec{r}_n(t)$ being the position of *n*-th bead at time *t*, *N* the number of beads in a chain, *p* the mode index. Eq. (3.68) is a decomposition of the equations of motion of the polymer into the normal coordinates (modes), where the *p*-th mode describes the part of the chain that includes (N-1)/p segments. The inverse transformation reads

$$\vec{r}_n(t) = \vec{X}_0(t) + 2\sum_{p=1}^{N-1} \vec{X}_p(t) \cos\left[\frac{(n-\frac{1}{2})p\pi}{N}\right] (n=1,\ldots,N) \;. \tag{3.69}$$

The modes are theoretically considered to be orthogonal with each other [140], so that the properties of the polymer are given only by their autocorrelations⁵

$$C_{pp}(t) = \frac{\langle \vec{X}_p(t) \vec{X}_p(0) \rangle}{\langle \vec{X}_p^2(0) \rangle} .$$

$$(3.70)$$

The functions $C_{pp}(t)$ allow direct observation of the chain relaxations on the corresponding length scales. Also, many static and dynamic observables, which are relevant in the experiments and simulations, can be expressed in terms of C_{pp} , so by knowing them one can fully characterize the polymer within the framework of the Rouse model.

As for the PBD, the original CRC and FRC models have been already studied within the framework of the Rouse model in Refs. [40, 41] with $\vec{r}_n(t)$ in Eq. (3.68) being the position of united atom n at time t of the PBD chain. In the following sections we compare

⁵In practice the cross-correlations $\langle \vec{X}_p \vec{X}_{q \neq p} \rangle$ are of the order of 1% of the autocorrelations $\langle \vec{X}_p^2 \rangle$, so they are usually neglected. Our calculations and the results reported in Ref. [40] show that this is indeed the case for PBD.



Figure 3.18: Main panel: Log-log plot of the static Rouse modes autocorrelations C_{pp} . The colored empty symbols denote the results obtained from our simulations at different temperatures. The filled black circles represent the data scanned from Fig. 3a in Ref. [40]. The vertical dashed lines correspond to p that are multiples of the number of chemical monomers in our PBD chains $N_{\rm m} = 29$. Inset: Zoom-in of the data in the main panel at p > 27.

and extend this analysis using our models across wider range of temperatures. We also relate our results to the literature data for common polymers.

3.5.2 Static Rouse modes

The static Rouse modes autocorrelations are defined as

$$C_{pp} = \langle \vec{X}_p(0) \vec{X}_p(0) \rangle = \langle \vec{X}_p^2(0) \rangle .$$
(3.71)

Ref. [40] reported no noticeable differences in C_{pp} between the models and temperatures up to p = 40. Indeed, the results of our calculations appear to overlap with the data scanned from Fig. 3a in Ref. [40], as presented in Fig. 3.18. The visible deviations appear for larger p.

Interestingly, for large p one observes three spikes in the data, which are present for all models and temperatures. A closer look on the data (inset in Fig. 3.18) revealed, that they correspond exactly to the values of p = 29, 58, 87. Those values are in fact multiples of the number of chemical monomers in our PBD chains $N_{\rm m} = 29$ (each of them consists of 4 UAs resulting in $N = 4N_{\rm m} = 116$ UAs per chain). Thus, the deviations of the values of C_{pp} at p = 29, 58, 87 from the overall trend can be explained by the fact that the cosine transform in Eq. (3.68) catches the periodicity in the PBD chain comprised of repeating sets of 4 UAs.

The theoretical prediction for the C_{pp} in the ideal Rouse chain reads [5]

$$C_{pp} = \frac{b_{\rm e}^2}{8N} \frac{1}{\sin^2(\frac{\pi p}{2N})} = \frac{C_{\infty}l^2}{8N} \frac{1}{\sin^2(\frac{\pi p}{2N})} \,. \tag{3.72}$$

However, due to the chain stiffness and excluded volume effects, only the first few modes follow such a scaling even in the bead-spring simulations [23, 140]. Although the corrections that account for these effects were proposed [141] and applied to the original CRC and FRC models of PBD in Ref. [40], we are able to use our parametrization of the conformational properties in Section 3.1.2 (cf. Eqs. (3.4) and (3.5)), that include the correction due to the stiff bending angle potential and an effective angle created by the dihedrals. By expressing the positions of UAs in Eq. (3.68) as the sums of bondvectors and plugging the result into Eq. (3.71) one obtains (after a long calculation that, due to its length, we do not reproduce in this thesis):

$$C_{pp} = \frac{l^2}{8N} \left[\frac{C_{\infty}}{\sin^2(\frac{\pi p}{2N})} - \frac{4A\overline{\alpha}\frac{1+\overline{\alpha}}{1-\overline{\alpha}}}{1-2\overline{\alpha}\cos(\frac{\pi p}{N}) + \overline{\alpha}^2} - 4(\alpha - A\overline{\alpha}) + O(N^{-1}) \right].$$
(3.73)

Fig. 3.19 tests our parametrization. The scaling of the axes is inspired by Eq. (3.72), so that for $\frac{p}{N} \to 0$, the value of $\frac{2(\pi p)^2}{N}C_{pp} \to l^2C_{\infty} = b_e^2$ as it is shown in the inset. We present the data at T = 353 K, but the result is valid for all studied temperatures. Using the values of the parameters reported in Appendix E, our predictions for the C_{pp} are in a good agreement with the simulation data. The differences between the models are very small, since the parameters also do not differ significantly. This shows once more that the static properties of PBD on the scales from the size of the statistical segment to the size of the chain are mildly sensitive to the presence of the dihedrals.



Figure 3.19: Main panel: Log-linear plot of the rescaled static Rouse modes autocorrelations C_{pp} . The colored symbols denote the results obtained from our simulations at T = 353 K. The colored lines with black dots are the results of Eq. (3.73). Inset: Zoom-in of the data from the main panel at $\frac{p}{N} \rightarrow 0$. The colored horizontal dashed lines correspond to the values of b_e^2 for each model.

3.5.3 Dynamic Rouse modes

The dynamic Rouse modes autocorrelations are defined in Eq. (3.70). The Rouse model predicts them to decay exponentially [5]

$$C_{pp}(t) = \exp\left[-\frac{t}{\tau_p}\right] \tag{3.74}$$

with $\tau_p = \tau_{\rm R}/p^2$ and $\tau_{\rm R} = \tau_{p=1}$ being the Rouse time, which is the longest relaxation time of the polymer chain corresponding to the mode of the largest length scale (p = 1). However, it is known from extensive simulation studies, that due to the excluded volume, chain stiffness and "viscoelastic hydrodynamic interactions" [142, 143], the $C_{pp}(t)$ deviates from simple exponential decay in Eq. (3.74) (especially for large p) and, instead, can be well described by the stretched exponential (Kohlrausch–Williams–Watts function) [23, 137–139]

$$C_{pp}(t) = A_p \exp\left[-\left(\frac{t}{\tau_p}\right)^{\beta_p}\right].$$
(3.75)

In addition, for a polymer melt close to the glass transition a two-step relaxation was reported [23]. Depending on the mode number, the emergence of a plateau between the initial short-time decay and the final long-time decay can be observed. Ref. [41] documented that this is the case for the original CRC model of PBD. By contrast, the original FRC model showed faster, single stretched exponential relaxation.

Fig. 3.20 presents the functions $C_{pp}(t)$ for p = 1, 10, 20, 50, 80, 100 for our models of PBD at T = 213 K. The first modes look like single decays with the relaxation times significantly faster when reducing the strength of the dihedrals (Figs. 3.20a and 3.20b). In case of the CRC2 and CRC models, with increasing the mode number p the two step decays gradually develop with the intermediate plateaus becoming more pronounced when decreasing the temperature (Figs. 3.20c to 3.20f). The curves also become more stretched. In case of the CRC4 and FRC modes the relaxations are quite fast, so the exponential relaxations are merged with the initial relaxation.

In order to characterize the relaxations quantitatively, at first, we extracted the relaxation times τ_p of all modes as points, where $C_{pp}(t=\tau_p)=1/e$. The result is plotted in Fig. 3.21. The axes are rescaled to test if the behavior of τ_p follows the p^{-2} prediction of Eq. (3.74). As it can be seen from the figure, only the first few modes, that correspond to large length scale relaxations have p^{-2} scaling. For the smaller length scale modes deviations are observed, which can be explained by the fact that in our simulations the chains are more complicated (stiffness, dihedrals, excluded volume etc.) than the Rouse model considers. Another observation is, that depending on the model, the times are shifted up systematically for all modes with respect to the data for the FRC model. With lowering the temperature the shifts become dramatic and span over many orders of magnitude. For the lowest studied temperatures T = 225,213 K, the first modes of the CRC model have not even reached 1/e over the simulation time of several μ s. Such an increase of the relaxation times is typically observed for glass forming systems upon decreasing the temperature. It is usually explained in the bead-spring model simulations [23] as a collective effect, where a particle becomes trapped within a cage created by its nearest neighbors. While such picture pertains for the FRC model, in case of the CRC(2,4) models, the trapping has been demonstrated to originate from the dihedral barriers ("intramolecular caging") [41]: when a particle cannot overcome the barrier it participates in caging another particle.

In order to characterize the emergence of the plateaus and the stretching, we attempted to fit the $C_{pp}(t)$ using Eq. (3.75). However, having 4 models, 7 working temperatures and



Figure 3.20: Dynamic Rouse modes autocorrelations $C_{pp}(t)$ for p = 1, 10, 20, 50, 80, 100. The colored symbols represent the data obtained from the simulations at T = 213 K. The colored lines with black dots are the fits using Eq. (3.76). The horizontal dashed lines correspond to the value 1/e.

115 modes means 3220 fits to perform! Another problem is that a unique subset of the data should be used for each fit due to the differently emerging plateaus for each model. In order to overcome these issues we use an automated fitting with the following approach:

• A typical fitting situation is presented in Fig. 3.22a. At first, the values of the relaxation times have been extracted previously in Fig. 3.21 as points where $C_{pp}(t =$



Figure 3.21: Relaxation times τ_p of the dynamic Rouse modes correlations extracted as points, where $C_{pp}(t = \tau_p) = 1/e$. The colors mark the simulation models and the symbols denote different temperatures.

 τ_p) = 1/e (horizontal dashed line in Fig. 3.22a). They were fixed in all the fits. The small p modes that had not relaxed over a time of the simulation at low temperatures were omitted from the fit.

• Since there is a prefactor A_p , Eq. (3.75) has to be corrected, so that the definition of the relaxation times $C_{pp}(t = \tau_p) = 1/e$ holds

$$C_{pp}(t) = A_p \exp\left[-\left(\ln A_p + 1\right) \left(\frac{t}{\tau_p}\right)^{\beta_p}\right].$$
(3.76)

- The values of $C_{pp}(t)$ were limited in time from above at $t < 10\tau_p$ in order to omit the noisy tails of the autocorrelation.
- For long times $t > 2 \times 10^6$ fs, where the data is spaced linearly, only the time points selected in logarithmic fashion were used in order to make the fit less biased by the tails, that are densely populated with data (black symbols in Fig. 3.22a).
- We are interested only in the second step of the relaxation. Since the initial relaxation appear to overlap for all our models, we can filter it out by considering only the points where the differences between the models start to occur. In practice this is done by determining the time point, where the difference in the data between the models⁶ $\Delta C_{pp}(t) = C_{pp}(t) C_{pp}^{ref}$ (dotted colored lines in Fig. 3.22a) exceed a

⁶For the CRC(2,4) models $\Delta C_{pp}(t) = C_{pp}(t) - C_{pp}^{\text{ref}}$ models the data of the FRC model were taken as a reference. For the FRC model, the difference between the CRC4 and FRC was used.

chosen threshold of, for example, $\Delta C_{pp}(t) > 0.05$ (horizontal black dotted line in Fig. 3.22a).



Figure 3.22: Panel (a): A typical stretched exponential fit of the dynamic Rouse modes. The colored symbols represent the data of the 13th mode obtained in our simulations at T = 213 K. The same black symbols are the subset of the data fed into the fitting routine. The solid colored lines with black dots are the results of the fit. The horizontal black dashed line corresponds to the value of 1/e. The colored dotted lines are the differences between the models $\Delta C_{pp}(t)$. The horizontal black dotted line represents the threshold of 0.05 used to determine the time, where $\Delta C_{pp}(t) > 0.05$. Panels (b-c): The values of the fitting coefficients A_p and β_p obtained from fitting the Rouse modes correlations with Eq. (3.76). The symbols correspond to different temperatures and the colors represent different models.

• Since the values of parameter $A_p > 1$ in Eq. (3.75) have no physical meaning, they

were constrained to unity from above.

With this fitting procedure we can fit the data for all p. However, since the fitting is automatic, the results should not be taken for granted. The resulting values of the fitting coefficients are presented in Figs. 3.22b and 3.22c. The inspection of the fits revealed that our approach worked well for all models at $p \leq 10$ $(N/p \geq 10)$ and at all p for the the CRC and CRC2 models, where the intermediate plateaus are reasonably developed. This is demonstrated by the values of the amplitude A_p different from unity in Fig. 3.22b. There are pronounced minima in the values A_p , that increase in depth depending on temperature. At the same time, the stretching exponent β_p is weakly dependent on the temperature and decreases with the mode number p starting from $\beta \approx 0.8$ down to $\beta \approx 0.5$, which means that the relaxation is slower than the exponential (the curves are stretched). While the decrease from $\beta \approx 0.8$ at small p agrees with the results reported in the literature [137, 138], our data is qualitatively different at large p, where the bead-spring simulations report a minimum of $\beta \approx 0.5$ at $N/p \approx 3$.

In principle, for $N/p \gtrsim 10$ it would possible to fix $\beta_p \approx 0.8$ for all T and obtain a fit of similar quality. This fixings implies time-temperature superposition, an important principle in glass physics and polymer physics. However, for $N/p \lesssim 10$, the principle appears to become progressively violated with decreasing N/p. With s = N/p, $N/p \lesssim 10$ corresponds to $s \lesssim 16$. According to Fig. 3.4, the regime $s \lesssim 16$ corresponds to chain segments where deviations from the Gaussian chain model, $R_{\rm e}^2(s) = b_{\rm e}^2 s$, become very prominent.

For the CRC4 and the FRC models, the stretched exponential, does not actually represent well the shape of the relaxation at $p \gtrsim 10$ (as it can be seen in Figs. 3.20c to 3.20f). In fact, the values of p, from which the fits fail, can be observed in Fig. 3.22b where the amplitudes A_p come back to unity after the initial decrease. This means that the fitting routine reached the upper bound in A_p , so the stretching parameter β_p was the only adjustable variable for the fit⁷. While for the CRC4 model β_p has a similar decreasing trend as of the CRC2 and CRC models, it is qualitatively different for the FRC model. In the case of the FRC model, the stretching exponent shows a minimum of $\beta_p \approx 0.5$ at $N/p \approx 3$. Interestingly, similar minima are reported in the bead-spring model simulations [137, 138]. In general, the failure of the stretched exponential for the CRC4

⁷This discussion is also motivated by the fact, that when we tried to discard the upper bound of A_p during the fits, we obtained the curves with $A_p > 1$, which has no physical meaning.

and FRC models can be explained by the fact that the studied region of temperatures $213 \leq T \leq 353$ K is quite far from the corresponding glass transition temperatures $T_{\rm g}$. We can observe such a temperature dependence in Fig. 3.22b by the fact that the amplitudes A_p for these models reach the cap of unity earlier for higher temperatures.

3.6 Mean-square displacements

Having seen in the previous section, that the dihedral barriers in PBD lead to an enormous change in the relaxation times of the Rouse modes, it is important to investigate how they influence the motion of the united atoms. In the simulations this motion can be observed directly by looking at their mean-square displacements (MSD). In particular, it is interesting to look at

$$g_0(t) = \frac{1}{N} \sum_{n=1}^{N} \left\langle \left[\vec{r}_n(t) - \vec{r}_n(0) \right]^2 \right\rangle \,, \tag{3.77}$$

which describes the MSD averaged over all united atoms of a chain;

$$g_1(t) = \left\langle \left[\vec{r}_{N/2}(t) - \vec{r}_{N/2}(0) \right]^2 \right\rangle \,, \tag{3.78}$$

which is the MSD of the innermost united atom of a chain;

$$g_2(t) = \frac{1}{N} \sum_{n=1}^{N} \left\langle \left[\vec{r}_n(t) - \vec{R}_{\rm cm}(t) - \vec{r}_n(0) + \vec{R}_{\rm cm}(0) \right]^2 \right\rangle \,, \tag{3.79}$$

which describes an averaged MSD of the united atoms with respect to the chain's center of mass $\vec{R}_{\rm cm}(t)$;

$$g_3(t) = \left\langle \left[\vec{R}_{\rm cm}(t) - \vec{R}_{\rm cm}(0) \right]^2 \right\rangle \,, \tag{3.80}$$

which is the MSD of the chain's center of mass itself;

$$g_4(t) = \left\langle \left[\vec{r}_{\text{end}}(t) - \vec{r}_{\text{end}}(0) \right]^2 \right\rangle,$$
 (3.81)

which describes the MSD of the united atoms at the ends of the chain.

Fig. 3.23 presents the $g_{0-4}(t)$ obtained from the simulations using the CRC model at T = 353 K. The united atoms at the ends of a PBD chain (orange diamonds in the figure)

show slightly faster dynamics than the innermost united atoms (blue squares). Yet, their influence is mild, since the contribution of many inner UAs dominates in the averaged dynamics (red filled circles). While at short time scales the local motion of the UAs is of importance (green filled triangles), at long times, being connected in chains the UAs cannot explore the distances beyond the chain's size. This is demonstrated by the plateau in the $g_2(t)$ (horizontal dotted line). At long times the diffusion of the chain as a whole prevails, so all the data in Fig. 3.23 coincide with the $g_3(t)$ (violet crosses).

Having already analyzed the Rouse modes for all our models, we, at first, examined how well they can actually describe our simulation data. By inserting Eq. (3.69) into Eq. (3.79) and assuming only the orthogonality of the Rouse modes, one obtains

$$g_2^{\rm R}(t) = 4 \sum_{p=1}^{N-1} C_{pp} \left[1 - C_{pp}(t) \right] \ . \tag{3.82}$$

This result is represented by a solid line in Fig. 3.23. It is nearly identical to the $g_2(t)$ obtained directly from the simulations, verifying that the properties of PBD can be indeed described by the orthogonal Rouse modes.

Typically, for unentangled polymers at the temperatures far above the glass transition



Figure 3.23: The mean-square displacements $g_0(t)$ (red filled circles), $g_1(t)$ (blue empty squares), $g_2(t)$ (green filled triangles), $g_3(t)$ (violet crosses) and $g_4(t)$ (orange empty diamonds) obtained from the simulations using CRC model at T = 353 K. The dark gray line is the $g_2(t)$ obtained from the Rouse modes correlations Eq. (3.82). The horizontal black dotted line correcponds to twice the gyration radius R_g of the PBD chain.

temperature $T_{\rm g}$, the MSD shows three distinctive regimes [2]. At short times, the initial motion of particles just after the beginning of the measurement is referred to as ballistic, since it is determined by Newtonian dynamics with $g_0(t) \sim t^2$. In the long times regime the particles diffuse with $g_0(t) \sim t$. The intermediate regime is characteristic of polymer systems. It shows a behavior of $g_0(t) \sim t^x$ with the exponent x < 1 and, therefore, is called subdiffusive. The value of x is typically between $0.5 < x < 0.75^8$. With lowering the temperature, the situation becomes slightly different. In particular, the ballistic regime is followed by an emerging plateau, that shifts the subdiffusive regime to longer times. Such a plateau is common for the glass forming systems: it indicates the trapping of the particles in a cage, that leads to the dynamical arrest.

Fig. 3.24 presents the MSD averaged over all united atoms of a chain $g_0(t)$ for PBD simulated using all our models at different temperatures as empty colored symbols. The corresponding Rouse predictions are given by the colored lines with black dots

$$g_0^{\rm R}(t) = g_2^{\rm R}(t) + g_3(t). \tag{3.83}$$

The general behavior of the curves follows the description above with the regimes denoted in the figure. However, the data for different models shows significantly different temperature dependences. At our working temperatures 213 < T < 353 K the FRC model demonstrates the fastest motion. The curves are very close to each other and have clearly distinct regimes. With introducing the dihedrals and making the barriers higher the dynamics becomes slower, so that the diffusive regime cannot be reached over few μ s of the simulations. In the case of the CRC model we observe the caging plateaus that make a dramatic effect on the dynamics with lowering the temperatures. As it was mentioned in the previous section, the presence of the dihedrals imposes constraints on the movement of the particles when the thermal fluctuations are comparable to the barrier height. For PBD those are the α and β dihedrals. When a particle becomes trapped within the barriers, it also cannot anymore make room for the rearrangements of its neighbors. Considering many such particles, this leads to the structural caging. Since the height of the dihedral barriers is larger than pair energy barriers, one observes the emergence of the cage at higher temperatures when comparing to the FRC model. We have performed

⁸The lower limit of x = 0.5 is in fact a result of the ideal Rouse chains. However by introducing chain stiffness, the value of x increases with the x = 0.75 being the limit of the stiff chains [2].



Figure 3.24: Log-log plots of the mean-square displacement $g_0(t)$ of united atoms for the CRC [panel (a), red], CRC2 [panel (b), green], CRC4 [panel (c), purple] and FRC [panel (d), blue] models. The colored symbols represent the data obtained from the simulations at different temperatures. The legend in the plot for the FRC model is also valid for other models. The colored solid lines with black dots are the predictions using the Rouse modes Eqs. (3.80), (3.82) and (3.83). The horizontal dashed line corresponds to the value σ^2 .

additional simulations of PBD using the FRC model at the temperatures T < 213 K and observed the cage effect appearing at the temperature 100 K below our working range (filled symbols in Fig. 3.24d).

The structural relaxation is typically described quantitatively by extracting the relaxation times from the MSD as points where $g_0(t = \tau) = \bar{\sigma}^2$ (horizontal dashed lines in Fig. 3.24) with $\bar{\sigma}$ being the average size of a particle (united atom in our case). With the values from Table 2.4, $\bar{\sigma} \approx 3.7$ Å. Fig. 3.25 plots these relaxation times in the form of the relaxation map, where the decimal logarithm $\log \tau$ is plotted against the inverse temperature 1000/T. When comparing the data at our working range of temperatures 213 < T < 353 K (colored symbols in the figure), one observes a super-Arrhenius increase of the relaxation times in case of the CRC(2,4) models. The data for FRC model appears

rather Arrhenius at these temperatures. Performing the simulations at lower temperatures allowed to see the increase in the relaxation times faster than Arrhenius. We have fitted our data using the Vogel-Fulcher-Tammann (VFT) equation in the logarithmic form

$$\log \tau(T) = \log \tau_{\infty} + \frac{B}{T - T_0} \tag{3.84}$$

with $\log \tau_{\infty}$, *B* and T_0 as adjustable parameters. The results are plotted in Fig. 3.25 as solid colored lines. These fits were used to extrapolate to lower temperatures and determine the glass transition temperature as points, where $\log \tau (T = T_g^{VFT}) = 100$ sec. The resulting values are presented in Table 3.6 along with the fitting parameters. Comparing to the cooling data (also given in Table 3.6 as T_g^{cool}), the T_g^{VFT} obtained from the equilibrium simulations are expected to be lower considering the fast cooling rate employed in the



Figure 3.25: The relaxation maps of PBD simulated using all our models (colored symbols). The solid colored lines are the extrapolations to lower temperatures using Eq. (3.84).

Table 3.6: The values of the fitting parameters of Eq. (3.84) and the glass transition temperatures $T_{\rm g}^{\rm VFT}$ obtained by extrapolating the fits to 100 sec. The values obtained from the cooling runs are also given for reference.

model	$\log \tau_{\infty} \ (ps)$	$B(\mathbf{K})$	T_0 (K)	$T_{\rm g}^{\rm VFT}$ (K)	$T_{\rm g}^{\rm cool}$ (K)
CRC	-1.2	641.14	119.2	161.3	192
CRC2	-0.3	316.11	116.8	139.0	143
CRC4	-0.3	351.70	72.6	97.1	108
FRC	-0.1	293.62	41.0	61.8	43

simulations. While this is the case for the CRC(2,4) models, the $T_{\rm g}^{\rm VFT}$ for the FRC model is about 1.5 times larger than the corresponding $T_{\rm g}^{\rm cool}$. Yet, this inconsistency rather reflects the poor quality of the extrapolation (cf. Fig. 3.27), than having a physical meaning. The VFT equation works well at the temperatures close to $T_{\rm g}$, but reaching the relaxation times at these temperatures requires much longer simulations than few μ s.

As an aside, we compared our results for the CRC model to various experimental references and original model simulations (Fig. 3.26). Our data fits within the experiments, confirming that our simulations represent the dynamics of PBD well in the studied temperature interval.

3.7 Discussion

By now we have adapted the united atom chemically realistic (CRC) model of 1,4polybutadiene to be able to perform the simulations of the bulk systems and systems with a free surface in LAMMPS code. We have performed extensive bulk simulation and studied



Figure 3.26: Comparison of the relaxation map of PBD simulated using the CRC and FRC models (filled symbols) with experimental and simulation references from the literature. The empty symbols represent the scanned data points, the solid lines are recreated VFT fits. Ref. [50] are the data from old simulations using the original CRC model. Refs. [144, 145] are the dielectric spectroscopy data. Ref. [146] are the data based on the measurements of the monomeric friction coefficient. Ref. [36] are dielecric relaxation data obtained from the recent simulations using the original CRC model. Ref. [147] are the combined data from dielectric spectroscopy (low T) and NMR (high T) for PBD of M = 1450 g/mol and M = 2020 g/mol (in our simulations M = 1569 g/mol).

the structure and dynamics of PBD over a wide range of temperatures 213 K < T < 353 K. Our results fit within the experimental references for PBD of similar composition and molecular weight.

We have also studied the influence of the dihedral barriers on the properties of PBD by performing the simulations with the reduced (CRC2 and CRC4 models) and completely disabled barriers (FRC model). We observed minor changes in the structure of the PBD melt with modified/disabled dihedrals, which could not be seen in the similar studies carried out earlier [40, 41]. By contrast, the dynamics of PBD is orders of magnitude faster for these models, which also leads to significant shifts of the glass transition temperature T_g to lower values. By the analysis of the local relaxation of the united atoms, we attribute these differences in the dynamics to the fact, that the trapping of the UAs between the dihedral barriers shifts the onset of the structural caging to higher temperatures.

Our results challenge the theories that try to predict the dynamics of the glass forming systems from their structure. In particular, the mode coupling theory (MCT) has been extensively discussed in the previous works [40, 41, 148]. Below we discuss the more recent elastically collective nonlinear Langevin equation (ECNLE) theory [119, 120, 149]. The theory considers a polymer melt as a system of hard spheres at volume fractions determined from the temperature dependence of the dimensionless compressibility. An energetic dynamic barrier can be calculated from the structure of the system, so that the time of crossing describes the local dynamics. In order to account for the collective rearrangements, the additional elastic barrier is introduced (which is also calculated using the dynamic free energy). Since the theory is non-linear, we expect the small differences in structure to cause significant differences in the dynamics. The theory was successfully applied to predict the segmental relaxation, glass transition temperature and fragility of many polymers, including PBD.

We have put a lot of effort into describing the structure of PBD simulated with our models and indeed observed small differences in the structural properties between the models. The two important quantities here are the characteristic ratio C_{∞} and the dimensionless compressibility $k_{\rm B}T\rho\kappa_T$. The ECNLE theory assumes C_{∞} to be temperature independent and uses it to map the compressibility of the melt to the one of the hard spheres [120]. We have implemented the numerical solution of the ECNLE theory following Refs. [119, 120, 149] and performed the calculations of the relaxation times plugging into the theory the C_{∞} at T = 353 K from Section 3.1.2 and the linearized in 1/T dimensionless compressibility using the Long-Lequeux fits from Section 3.3.3 for our CRC and FRC models. Fig. 3.27 present the results of the calculations of the relaxation times, that correspond to the hopping across the dynamic barrier only (without the additional contribution from the elastic barrier). Even though the differences in the structure between the models were taken into account in the calculations, the data overlaps. Interestingly, for both models, the dynamics at high temperature (where the barrier is very small) we observe an agreement with the MD data of the FRC model. However, with decreasing the temperature, even without the elastic contribution the ECNLE theory already overestimates the relaxation times of the FRC model. This means that in the studied range, the mapping of temperature to the volume fraction via the renormalized dimensionless compressibility is not valid for the FRC model. At the moment the reasons for this disagreement are not clear and further research is required. This is the main reason why we decided not to expose the ECNLE theory, its implementation, and the details of its application to our simulation data in this thesis (despite the large amount of work spent on this project).



Figure 3.27: The relaxation times obtained in our work (filled symbols with solid lines) compared to the predictions of the ECNLE theory (empty symbols) and the experimental reference (crosses).

Chapter 4

Film systems analysis

This chapter presents the results obtained from the simulations of the film systems. We begin our analysis with determining the thickness and studying its temperature dependence upon cooling in Section 4.1. Then in Section 4.2 we relate the obtained results to the layer-resolved investigation of the dynamical properties. Finally, we discuss our findings in Section 4.3.

4.1 Density profiles and thickness

Let the z-axis be perpendicular to the film plane (Fig. 2.8). Then the density profile along the z direction can be defined as

$$\rho(z) = \frac{1}{A} \left\langle \sum_{i=1}^{NN_c} \delta(z - z_i) \right\rangle \,, \tag{4.1}$$

where A is the area of the cross-section of the simulation box in the xy plane.¹ In practice, such a calculation is performed by introducing finite (yet, small enough in thickness) slabs along the z-direction and counting how many particles are in each slab volume. Fig. 4.1 presents the density profiles of our films simulated using the CRC and FRC models at different temperatures. By looking at the shape of the curves, we can get an idea on the influence of some surface effects. In particular, the peaks on the left-hand side of the figures suggest that the presence of the wall leads to layering of united atoms next to it, which is typically found in liquids in contact with an impenetrable substrate [18].

¹The periodic boundary conditions are employed in the simulations in x, y directions.

We compared this part of our density profiles to the results of Ref. [36], where the PBD was simulated in confinement between two graphite walls using the original CRC model (dashed black line in Fig. 4.1a). An excellent agreement was obtained, which is expected considering that our modified CRC model was tuned to reproduce the same density as the original one.

At the free surface, one observes a gradual decrease of the density down to zero, which is commonly attributed to the presence of capillary waves [150]. The width of the decrease region is determined by the wave amplitudes, which are temperature dependent and limited by the system size. The plateau in the middle of the films corresponds to the value of the bulk density, as marked by the horizontal dotted lines in Fig. 4.1. From the analysis of the bulk systems, we know that they have almost identical densities, so the plateaus for the CRC and the FRC models overlap. Interestingly, they also overlap at the wall and at the free surface, meaning that surface effects on the density appear to be mildly influenced by the dihedrals.

Measuring the density profile of the film allows us to determine the thickness of the films using the Gibbs dividing surface (GDS) method [3]. In order to do this, at first, the mean density $\bar{\rho}$ at the center of the film z_c is calculated by averaging $\rho(z)$ over a wide enough region $z_c - \Delta z/2 \le z \le z_c + \Delta z/2$. The choice of the center point z_c and the averaging region is arbitrary as long as the mean density can be well defined. In our case, the procedure was automatic with z_c being the middle of the non-zero data of the density profile and $\Delta z = 30$ Å. The positions of the upper $z_+^{\rm G}$ and lower $z_-^{\rm G}$ Gibbs dividing surfaces are calculated as

$$z_{\pm}^{\rm G} = z_{\rm c} + \frac{1}{\bar{\rho}} \int_{z_{\rm c}}^{\pm \infty} \rho(z) \mathrm{d}z \;.$$
 (4.2)

Qualitatively, the integration in Eq. (4.2) means that starting from a middle point z_c the position of the GDS is propagated to the left (or to the right) proportionally to the fraction of the density profile at position z to the mean value. The positions z_+^{G} and z_-^{G} are marked in the Fig. 4.1 as vertical dotted lines. The thickness of the film is then obtained as

$$h = z_+^{\rm G} - z_-^{\rm G} \ . \tag{4.3}$$

Since the density of PBD increases upon cooling, the thickness decreases, as it is observed from lower positions of z_{+}^{G} with decreasing the temperature in the Fig. 4.1. In



Figure 4.1: The density profiles of the films obtained from the simulations using the CRC (red) and FRC (blue) models at different temperatures. The dashed black line is the data scanned from Ref. [36], where the PBD film confined between graphite was simulated using the original CRC model at T = 353 K. The vertical dot-dashed lines are the positions of the graphite layers. The vertical dotted lines are the positions of the Gibbs dividing surfaces for the CRC model. The horizontal dotted lines correspond to the density of the bulk systems at the same temperatures.

principle, the z_{-}^{G} also shifts with decreasing the temperature, since the first peak (that contributes the most to the integration in Eq. (4.2)) increases in height. However, such an influence is actually quite small (~ 0.1 Å), so it is ignored.

Using the GDS method, the thickness can be monitored not only in the equilibrium simulations, but also upon cooling the films. Fig. 4.2 presents the dependence of thickness on temperature h(T) obtained from the cooling simulations at the rate $\Gamma = 0.83$ K/ns (colored circles)² and from the equilibrium simulations (colored crosses). Since the film system is much bigger than the bulk, the averaging statistics is better and small differences

²On the contrary to the bulk systems, it was not possible to perform multiple cooling runs for the same film systems and average over them. Only the smoothing was applied in order to obtain a data point per Kelvin.

in density are now more visible: the thickness of the CRC model is systematically slightly lower than for the FRC model at high temperature.

As it is usually observed in ellipsometry experiments [151], h(T) changes slope when the system is undergoing the glass transition. Yet, our cooling rate $\Gamma = 0.83$ K/ns is much faster than typically employed in the experiments, which leads to a broad transition region. We have extracted the $T_{\rm g}$ from the cooling data using a similar approach as during the analysis of the bulk densities. The thickness h(T) was fitted with:

$$h(T) = w\left(\frac{M-G}{2}\right)\ln\left[\cosh\left(\frac{T-T_{\rm g}}{w}\right)\right] + (T-T_{\rm g})\left(\frac{M+G}{2}\right) + h_{\rm g}, \qquad (4.4)$$

with w being the width of the glass transition region, M and G the slopes dh(T)/dT in the melt and glass regions respectively, $T_{\rm g}$ the glass transition temperature and $h_{\rm g}$ the thickness of the film at $T = T_{\rm g}$. In order to account for the dependence on the size of the fitting region $(T_{\rm g}^* - \Delta T, T_{\rm g}^* + \Delta T)$, we have done the fits over a set of ΔT ranging from tens of Kelvins to the highest possible value. Comparing to the bulk density data, the data for the film thickness have lower quality. Thus, the influence of the fitting region is more pronounced in the values of M and G, while the $T_{\rm g}$ and $h_{\rm g}$ are much less sensitive to the choice of ΔT (Fig. 4.3). The final iteration of the fits with $T_{\rm g}$, $h_{\rm g}$, M and G fixed to



Figure 4.2: The temperature dependence of the film thickness obtained by the GDS method from the simulations using the CRC (red) and the FRC (blue) models. The circles represent the data obtained from the cooling runs, and the crosses are the averaged values from the constant temperature simulations. The solid black lines are the fits using Eq. (4.4) with the parameters from Table 4.1.

their values at the highest possible ΔT leads to the values presented in Table 4.1 and the fitting curves represented by black solid lines in Fig. 4.2. We obtained an exact match of the glass transition temperature $T_{\rm g} = 43$ K for the FRC model with the bulk simulations. In case of the CRC model, the resulting value $T_{\rm g}^{\rm film} = 206$ K is slightly higher than the bulk value $T_{\rm g}^{\rm bulk} = 192$ K. Still, the film thickness data in the transition region is quite noisy even after smoothing, so it is hard to argue about reasons of possible differences. Moreover, refitting the CRC data with the $T_{\rm g}$ fixed to the bulk value resulted in the fitting curve of the same quality (not shown) with comparable values of the parameters. We consider them in agreement with each other.

Our $T_{\rm g}$ values are, of course, the film averaged values, that result from the contribution



Figure 4.3: The values of the fitting parameters of Eq. (4.4) depending on the fitting interval ΔT . The colours denote different models: CRC - red, FRC - blue. The dashed lines represent the final values of the parameters.

model	$T_{\rm g}$ (K)	$h_{\rm g}$ (Å)	M (Å/K)	G (Å/K)	w (K)
CRC	206.9	94.0	0.0766	0.0132	71.1
FRC	43.0	82.8	0.0704	0.0295	7.7

Table 4.1: The final values of the fit parameters of Eq. (4.4)

of the bulk-like region in the middle of the film and the surface effects. The fact that they are basically the same as in the bulk for both models may suggest that either the surface effects are very weak, or the influence of the graphite wall and the free surface average out. We have learned from the analysis of the bulk systems, that disabling the dihedrals leads to significant speed up of the dynamics of PBD while preserving its structure, which explains the presence of the shift of the $T_{\rm g}$ to lower values. It is interesting now to investigate the local dynamics of PBD across the film in order to understand the influence of the dihedral barriers on the surface effects, and consequently, on the $T_{\rm g}$.

4.2 Mean-square displacements and dielectric relaxation

Our study of the dynamics in the films utilizes two methods. The first one is essentially the same as employed for the bulk systems: the simulations allow direct observation of the mean-square displacements of united atoms $g_0(t)$ on different time scales. The second mimics the dielectric spectroscopy technique commonly employed in experiments (see [152] and references therein), where the dynamics is analyzed by measuring the dielectric response of the system to the applied electric field alternating with different frequencies. It is inspired by the previous works [31, 35, 46], where dielectric relaxation of PBD was studied in the simulations using the original CRC model. The fluctuation-dissipation theorem relates the experimentally relevant complex dielectric permittivity ϵ^* to the correlation of the total dipole moment of the system \vec{M} , which was shown to be equal to the sum of independent chain contributions, which, in turn, are the sums of the independent dipoles of the *cis* groups of PBD³ (Fig. 4.4) [46, 153]. Thus, the dielectric spectroscopy experiments on PBD probe the relaxation on the segmental scale. The studied sample is placed between the electrodes, that act like walls confining the film, so the relevant quantity extracted

 $^{^{3}}$ We refer the reader to the works [35, 46] for the theoretical and implementation details, since (as we will show later) our calculations are identical.



Figure 4.4: Schematic representation of the dipole moment of the *cis* group of PBD. The CH_2 united atoms are treated together. The hydrogen atoms from the CH groups are separated and placed in their equilibrium positions. The charges are given in the figure in the units of electronic charge $1 e = 1.60217733 \times 10^{19}$ C along with the equilibrium bond lengths. Such a configuration results in a small non-zero dipole moment depicted as an orange arrow. Since the dihedral angle for the *cis* monomer is strongly localized, the dipole is considered in plane of the monomer. Due to the stiff bonds and bending angles it can also be considered perpendicular to the double bond.

from the simulation trajectories is the autocorrelation of the perpendicular to the film component of the dipole moment (DACF) of the *cis* groups of the PBD:

$$C^{z}(t) = \frac{\langle M^{z}(t)M^{z}(0)\rangle}{\langle M^{z}(0)M^{z}(0)\rangle}.$$
(4.5)

This is an important limitation, since the relaxations in different directions may be different and it is the averaged dynamics that matters for the glass transition in the films. Thus, in our simulations we also study the autocorrelation of in-plane components $C^{x,y,xy}(t)$ as well as the total autocorrelation of a dipole moment vector

$$C(t) = \frac{\langle \dot{M}(t)\dot{M}(0)\rangle}{\langle \vec{M}^2(0)\rangle}.$$
(4.6)

At first, Fig. 4.5 presents the comparison of the total mean-square displacements of united atoms $g_0(t)$ and the total DACF C(t) obtained from the bulk simulations (black solid lines) and from the film simulations (colored solid lines) at selected temperatures averaged over the whole simulation box. The in-plane and perpendicular components will be discussed below. In case of the CRC model, the averaging over the correlations of the *cis* dipole moments was performed for C(t). In case of the FRC model, there is no distinction between the *cis* and *trans* monomers, because all the dihedral potentials are disabled. Therefore, one expects the dipole moment of each monomer to be equal to zero. However, due to the excluded volume, there is still a preferred range of the dihedral angles (see Section 3.1.1) which leads to a non-zero value of \vec{M} . Thus, the DACF for the FRC model was obtained by averaging the correlations of the dipole moments of all monomers. We use the time domain representation in order to be able to relate the relaxation regimes with the regimes of the MSD, and because performing the Fourier-transform of the data would require either a different approach to the spacing of the simulation trajectories, or enormous interpolation efforts.

The film averaged data of the MSD are slightly faster than the bulk curves at the corresponding temperatures. The DACF curves appear to decay slightly slower at long times in the films. Refs. [31, 47] attributed such additional slowing down to the adsorption-desorption kinetics of the chains at the graphite wall.

We are interested in the time scales of the structural relaxation. For the bulk systems, they were extracted as points, where a united atom has displaced over a distance of its own size $g(t = \tau) = \bar{\sigma}^2$. While for the MSD this definition is consistent between the CRC and FRC models, the same relaxation times correspond to different regimes in the dielectric relaxation between the models. Omitting the short time region of the DACF t < 100 fs, the data for the CRC model are single decays, which allows to make a rough approximation of the structural relaxation times to the decay of $C(t=\tau) \approx 0.2$. Meanwhile, the FRC model exhibits two-step relaxations and the structural relaxation times correspond to the second step with $C(t = \tau) \approx 0.01$. Of course, the two discussed quantities cannot be directly related: the MSD represents the motion of individual united atoms, whereas the DACF describes the reorientation of a group comprised of several united atoms. In addition, in the simulations using the realistic model each individual dipole persists at all times, so only their orientations contribute the most to the correlation. In case of the FRC model, the non-zero dipole moment is the result of the average excluded volume effect, so the reorientation of the united atoms within the monomer are of importance, instead of the reorientation of the monomer as a whole.

Having compared the overall film dynamics to the bulk, it is interesting now to look at the relaxations in the different regions of the films. To do so, we have divided the films into slabs and calculated the MSD and the DACF in each of them. The layout of the slabs is presented in the Fig. 4.6. The bottom of the first slab coincides with the topmost layer
of the graphite. The slabs have width equal to $2\bar{\sigma}$ and overlap with the neighboring slabs over a distance $\bar{\sigma}$ for better statistics in the data. A united atom or a dipole belongs to a slab if its position (center of mass of a corresponding monomer in case of a dipole) is



Figure 4.5: Panels (a-b): The mean-square displacements $g_0(t)$ obtained from the bulk simulations (black solid lines) and film simulations (colored solid lines) using the CRC (red) and FRC (blue) models at different temperatures averaged over all united atoms in the simulation box. Panels (c-d): The dielectric relaxation C(t) obtained from the bulk simulations (black solid lines) and film simulations (colored solid lines) using the CRC (red) and FRC (blue) models at different temperatures averaged over all dipoles in the simulation box. The vertical dotted lines correspond to the structural relaxation times extracted from MSD curves. The horizontal dotted lines correspond to the values C = 0.2(CRC) and C = 0.01 (FRC), obtained as averaged values of the DACF at the times where $g_0 = \bar{\sigma}^2$.



Figure 4.6: Schematic illustration of the geometry of the slabs in the PBD film.

inside that slab in the beginning of the correlation time window⁴.

Fig. 4.7 presents the xy (in film plane) components and Fig. 4.8 presents the z (perpendicular to the film) components of the MSD and DACF for slabs 1-5, 12, 14, 16, 23-27 at T = 240 K. The slabs are marked by the color spectrum with the blue and red sides corresponding to the slabs next to the wall and the free surface respectively. The in-plane data for both MSD and DACF show the slowing down of the dynamics at the wall and speeding up at the free surface with respect to the middle of the film. The slabs in the middle show bulk-like behavior. For the perpendicular to the film components, a similar behavior is observed for the MSD and the first step of the DACF. At long times, however, the MSD curves should be limited by the distance from the slab to the farthest edge of the film. The second steps of the dielectric relaxation are qualitatively different. It appears, that the DACF relaxes more slowly at the free surface as well. This suggests, that there are additional constraints on the orientation of the PBD monomers at the free surface, that result in preferred direction of the dipoles. Since both surfaces contribute in the same way to the averaged supported film dynamics, the long-time differences of the box-averaged DACF (short dashed black lines) with respect to the bulk curves (long dashed black lines) are understood (see also Figs. 4.5c and 4.5d). We have compared our results for $C^{z}(t)$ with the previous work on simulations using the original model of PBD confined between graphite walls. Wider film slabs of about 1.2 nm were analysed

⁴Considering the width of the slabs $2\bar{\sigma}$, this method yields nearly identical results for the structural relaxation to checking the positions in the beginning and the end of the correlation time window, or even at each time step within the time window. For large time windows the additional checks deteriorate the averaging statistics.

there. An excellent agreement is obtained between the $C^{z}(t)$ in the middle of the film and between our 2nd thin slab from the wall and the 1st thick slab from the work [35].

As it has been already mentioned, in the dielectric spectroscopy experiments, the electrodes act as walls for the film, so it is usually the perpendicular component of the dielectric relaxation that is measured. It is interesting to study if there are preferred orientations of the dipoles in the film slabs at the wall and at the free surface, since the experiments would not see the contributions from the dipoles that are oriented in-plane. Fig. 4.9 presents a layer-resolved comparison of the projections of the dipole moment vector onto different directions. The limits of the y axis are fixed for each model to highlight the temperature dependence of the surface effects. One immediately notices the difference in the magnitude of the dipole moment between the models. Our calculations show on average $|\vec{M}| \approx 0.33$ D (1 Debye = 0.2081943 eÅ) for the CRC model and $|\vec{M}| \approx 0.23$ D for the FRC model. The values for the CRC model also show a slight increase with decreasing the temperature, while the data for the FRC model appears to be independent of temperature. At high temperatures, all projections for each model are identical within the error bar of the measurement. There are small hints of a polarization in z direction with preservation of the total magnitude at the wall, and an increase of the magnitude without pronounced polarization at the free surface. Unfortunately, with lowering the temperature our data suffers from insufficient statistics (or equilibration). While in the middle of the film all three projections appear to be identical (again within the error bar of the calculation), it is not possible to interpret physically the influence of the confinement.

Similarly to the bulk analysis, we have extracted the structural relaxation times from the MSD data in each slab in the parallel and perpendicular to the film directions as points where $g_0^{xy}(t=\tau) = \frac{2}{3}\bar{\sigma}^2$ and $g_0^z(t=\tau) = \frac{1}{3}\bar{\sigma}^2$ respectively (the factors 2/3 and 1/3 are introduced to account for different number of degrees of freedom with respect to the bulk data). They are plotted in Fig. 4.10 as a function of the position of the slab. Comparing the data at the same temperatures between the CRC and FRC models we observe a similar behavior to what we have seen in the bulk: the dihedrals set the total order of magnitude of the relaxation time, which is then getting modified by the surface effects. The modifications become dramatic with decreasing the temperature. Yet, visually they penetrate no more than ~ 20 Å inside the film. The slowing down at the wall and



Figure 4.7: The parallel to the film components of the MSD (a-b) and DACF (c-d) obtained from the simulations using the CRC and FRC model at T = 240 K. The colored lines correspond to the fim slabs with blue side of the spectrum marking the slabs at the wall and the red side - at the free surface. The black long dashed line represents the bulk data and the black short dashed line is the in-plane component of the film-averaged data. In case of the MSD, the bulk curve is multiplied by 2/3 to account for different number of degrees of freedom. The horizontal dotted line represents the value $\frac{2}{3}\bar{\sigma}^2$.



Figure 4.8: The perpendicular to the film components of the MSD (a-b) and DACF (c-d) obtained from the simulations using the CRC and FRC model at T = 240 K. The colored lines correspond to the fim slabs with blue side of the spectrum marking the slabs at the wall and the red side - at the free surface. The black long dashed line represents the bulk data and the black short dashed line is the perpendicular component of the film-averaged data. In case of the MSD, the bulk curve is multiplied by 1/3 to account for different number of degrees of freedom. The horizontal dotted line represents the value $\frac{1}{3}\bar{\sigma}^2$. The purple lines show the results of the original work [35], where wider slabs were studied.



Figure 4.9: Projections M_x, M_y, M_z of the dipole moment obtained in each slab in the film plotted agains the slab coordinate. The colours denote different models used in the simulations: CRC - red, FRC - blue. The shapes mark different directions: x - squares, y - circles, z - triangles. The colored solid lines are the rescaled total magnitudes $\frac{|\vec{M}|}{\sqrt{3}}$. The limits of the y axis are fixed for each model to highlight the temperature dependence of the surface effects.

the speed up at the free surface must be averaging out, so the overall film dynamics (as we reported above) is quite similar to the bulk. Which also explains similar glass transition temperatures to the ones obtained for the bulk systems.

Let us focus on the relaxation time gradients at the wall and at the free surface of the films. Figs. 4.11 and 4.12 show these data separating between relaxations in xy and zdirections and Fig. 4.13 presents a superposition of the relaxation time profiles and the density profiles. Additionally, we have fitted the data with a commonly used empirical "double-exponential" relation (see [154, 155] and references therein)

$$\tau(z,T) = \tau_{\rm b}(T) \exp\left[-A(T) \exp\left(-\frac{z}{\xi}\right)\right] \,, \tag{4.7}$$

with $\tau_{\rm b}(T)$ being the relaxation time in the middle of the film at temperature T,⁵ and A, ξ are adjustable parameters. The results of the fits at selected temperatures are presented in Fig. 4.13 as colored lines and the values of the fitting parameters are plotted in the Fig. 4.14. Comparing this ample amount of data one can extract the following features:

• First of all, aligning the data so that the positions of the upper GDS coincide (Fig. 4.11) leads to an impression that the relaxation time gradients at the free surface are determined only by the time in the middle of the films and are the same for both studied models. This is seen the best comparing the CRC model at 353 K (red squares) and the FRC model at 225 K (blue triangles) in Fig. 4.11. Such an observation suggests that the scaling of the fitting parameters in Eq. (4.7) with $\tau_{\rm b}(T)$ should be the same between the two models. Such scaling has already been observed before in [155]. Figs. 4.14a and 4.14b show that this is the case for the relaxation in the *xy* direction. In the *z* direction the dependencies of the fit parameters are slightly different between the models, however these fits are definitely influenced by the closest points to the upper GDS which deviate from the overall behavior of the data (especially for the FRC model; see e.g. Figs. 4.10 and 4.11b)⁶. Furthermore, it is important to note that the gradients at the free surface increase with lowering

⁵We used the value averaged over 5 slabs in the center of each film as $\tau_{\rm b}(T)$. The bulk relaxation times sometimes slightly disagree with the values in the middle of the films, which, when used in the fits, lower their quality. We believe the disagreement comes from the statistical errors and insufficient equilibration rather than having a physical origin, since the films should be thick enough to have a bulk-like region in the middle.

⁶The origin of these artifacts is not clear. Excluding these points from the fits for all data resulted in overestimating the gradients for all models and temperatures.

the temperature as it is demonstrated in Figs. 4.14e and 4.14f, which is exactly the opposite than the behavior of the density profiles. As it has been already mentioned in Section 4.1 a gradual decrease of the density down to zero at the free surface is commonly attributed to the presence of capillary waves [150], that emerge from a competition between the thermal energy and the surface tension. Their height, and therefore the width of the region of the decrease of the density is proportional to temperature. Moreover, Fig. 4.13 shows that this width is significantly smaller than the length over which the relaxation times are modified. Similar results are reported from simulations of supported films using a bead-spring polymer model [24] (Fig. 4.15). All these observations suggest that at the free surface the profile of the dynamics and the density profile cannot be related. Especially considering the fact that the decrease of the density profile next to the upper GDS results from the presence of vacuum in between the waves. The local density nearby an arbitrary particle can be in fact not very different from the bulk. Assuming that this is the case, then there is no change in properties of the melt with respect to the bulk-like middle of the film that could be responsible for the change in the dynamics. The existence of the gradient of the relaxation times can be attributed to the presence of the free surface itself and the fact that the cage that surrounds a given particle is open, which significantly speeds up the relaxation. This effect gradually damps towards the interior of the film.

• The situation at the wall side of the film is ambiguous. At a first glance, it seems from Fig. 4.13 that the relaxation times data follows the envelope of the density profiles. Since the density profiles are identical between the models at the same temperature, one would expect the same scaling of the fit parameters with temperature. However, Figs. 4.14g and 4.14h show that this is not the case. While the amplitudes of the relaxation time gradients can be discussed, the values of the decay length are very scattered. Moreover, only the amplitudes of the gradients of the dynamics in z direction can probably be averaged to the same constant value for both models at all temperatures. The data in the xy direction actually show the same scaling with $\log_{10}(\tau_{\rm b})$ as for the free surface (empty symbols in Fig. 4.14c). Another observation that is important to note comes from the comparison of the gradients in the xy direction with previously mentioned results from the simulations using a coarse-



Figure 4.10: The natural logarithm of the structural relaxation times extracted from the MSD data at different slabs in the film. The symbols denote different temperatures and the colors mark different models.



Figure 4.11: Subset of the data from Fig. 4.10 for z > 70 Å (next to the free surface of the film). Panel (a) presents the relaxation times in xy direction; panel (b) presents the relaxation times in z direction. The legend is the same as in Fig. 4.10. The data were shifted so that the positions of the Gibbs dividing surfaces at the free surface of the film coincide at z = 110 Å for all models and temperatures (vertical dotted line).



Figure 4.12: Subset of the data from Fig. 4.10 for z < 50 Å (next to the wall). Panel (a) presents the relaxation times in xy direction; panel (b) presents the relaxation times in z direction. The legend is the same as in Fig. 4.10.



Figure 4.13: Structural relaxation times extracted from MSD data in film slabs. Filled symbols correspond to the perpendicular to the film direction, empty symbols correspond to the parallel to the film component. Colors mark different models. Colored lines are the results of the fits using Eq. (4.7). The density profiles are given for comparison as black solid lines. Vertical black dotted lines are the positions of the Gibbs dividing surfaces.



Figure 4.14: The values of the fitting coefficients in Eq. (4.7) for the relxation time gradients at the free surface (a-b,e-f) and the wall (c-d,g-h) of the PBD supported films simulated using the CRC (red) and FRC (blue) models. The empty symbols represent the in-plane relaxations, the same filled symbols are the relaxation in the perpendicular to the film direction.



Figure 4.15: The profile of the relaxation times obtained from Ref. [24] where the polymer film supported on a structureless wall was simulated using a bead-spring polymer model.

grained model (Fig. 4.15). In particular, both films exhibit the layering of the particles next to the attractive wall that is visible on the density profiles. However, the data from Ref. [24] show a speed up of the relaxation at the wall, not the slowing down as in our case for both models. This disagreement can be explained by the differences in the properties of the substrate. In our case, the substrate is modeled by particles in a crystal lattice that make the wall very attractive and rough. In the work [24] the substrate is structureless (smooth) and is modeled using a weakly attractive potential that acts only in z direction. Work [29] studied the influence of the wall roughness on the dynamics at the substrate also by means of simulations using a bead-spring polymer model for the film and reported that it is sped up at the smooth walls and slowed down at the rough walls. As for our results, it remains unclear how this effect is influenced by the presence of the dihedrals.

4.3 Discussion

We have performed molecular dynamics simulations of the polybutadiene films supported by a graphite wall using the chemically realistic (CRC) united-atom model and the freely rotating (FRC) model, where all the dihedral potentials were disabled. We observed no significant differences in the density profiles of the film simulated using both models. The glass transition temperatures extracted from the dependence of the film thickness on temperature upon cooling agree with the values obtained from the bulk systems.

The overall film dynamics observed in the mean-square displacements of united atoms and the dielectric relaxation is mildly different from the bulk dynamics. The structural relaxation times extracted from the MSD data correspond to different regimes in the dielectric relaxation between the CRC and the FRC models. Qualitatively different behavior of the dielectric relaxation between the parallel and perpendicular to the film components was observed: the in-plane relaxation is slower at the wall and faster at the free surface, while the perpendicular to the film relaxation is slowed down at both interfaces.

The layer resolved analysis of the structural relaxation times extracted from the MSD data showed significant influence of the interfaces with respect to the middle of the film. However, the effect of the wall and the free surface qualitatively appear to average out when contributing to the total $T_{\rm g}$ of the film. Analysis of the relaxation times gradients at the interfaces showed that at the free surface they nearly identical in the xy direction between the CRC and FRC models at the temperatures with the same relaxation times in the bulk-like middle of the film. The numerical results for the z direction at the free surface and both directions at the wall are not conclusive due to difficulties of interpreting the behavior of the gradient decay length ξ . A visual inspection suggests that these gradients are different from each other.

Our data for the CRC model agree with the results reported in the previous studies of confined PBD [31, 35, 36] in the fact, that the dynamics is modified in a region of about 1 nm next to the graphite wall. We report slightly deeper perturbations at the free surface. Refs. [31, 35, 36] argued that the perturbation of around 1 nm into the film are the result of the interplay between the dihedral barriers present in the realistic model and the density changes created by the interfaces. They should lead to much smaller shifts in the glass transition temperature in the confined geometry with respect to what is typically observed in the widely used coarse-grained bead-spring polymer models (which do not have the dihedrals). Our FRC model mimics the bead-spring models, but it has a realistic length scale and shorted bond length. The comparison between the CRC and FRC models agrees with this hypothesis for the relaxation in the perpendicular to the film direction (which was the only one discussed in Refs. [31, 35, 36] and only for the films confined between walls), since there were no similarities found in the gradients between the models. However, we report a different behavior of the relaxation time gradients in the parallel to the film direction, where the identical gradients were obtained between the CRC and FRC films, that have the same order of magnitude of the relaxation time in the middle.

Additionally, we were able to compare the in-plane relaxation gradient obtained from our CRC model simulations at T = 240 K ($T_g = 206$ K) to the same data obtained from the bead-spring model simulations of the supported film at T = 0.48 ($T_g = 0.392$) in [24, 150]. Both of them have $T/T_g \approx 1.2$. Fig. 4.16 presents this comparison. For all data, z = 0 is the position of the Gibbs dividing surface. The x axis is rescaled by the size of the particle $\bar{\sigma}$. Unfortunately, we do not have the data for the FRC model at the same T/T_g , so the lowest available temperature T = 113 K and also T = 240 K are presented for comparison. A remarkably good agreement is observed between the data for the CRC and the bead spring model. As expected, the gradient data for the FRC model is below due to the higher ratio of T/T_g .



Figure 4.16: The natural logarithm of the structural relaxation times extracted from the MSD data at different slabs in the film. The symbols denote different temperatures and the colors mark different models.

Chapter 5

Conclusions and Outlook

We now give a summary of the key results of our work and provide an answer to the title of the thesis "Is the interfacial mobility in glass-froming polymer films determined by collective motion or intramolecular energetics?"

In order to study the influence of realistic torsional constraints on the properties of glass forming polymers we have adapted the united atom model of 1,4-polybutadiene (PBD) *cis-trans* random copolymer (developed previously from quantum chemistry calculations and validated against experiments [39, 45, 46]) for classical molecular dynamics simulations in the LAMMPS code [17, 43, 44]. Four versions of the model were introduced varying the scale of all torsional (dihedral) potentials in the system:

- Chemically realistic chain (CRC) model with the unmodified dihedrals potentials (which provide the distinction between the *cis* and *trans* monomers).
- CRC2 and CRC4 models with the dihedral potentials reduced by a factor of 2 and 4 correspondingly.
- Freely rotating chain (FRC) model with all dihedral potentials switched off. Here the distinction between the *cis* and *trans* groups is lost.

Such an interpolation provides a crossover to the widely used generic bead-spring polymer models, in which the realistic chemical details (including the dihedrals) are coarse-grained. Our FRC model resembles the bead-spring models with an important exception, that it has much shorter bond length with respect to the size of the particles. The most used bead-spring model of Kremer-Grest [20, 23, 24] type has a single length scale that determines both the size of the beads and the length of the bonds between them.

Our simulations were performed in the bulk using all 4 models and in the confined

geometry (supported film on graphite) using the CRC and the FRC models in the temperature range $213 \le T \le 353$ K.

Note that in order to simulate supported films (including a free surface), it is not possible to use tail corrections for the pair-potential. The parameters of the original model had to be adapted to yield the same equation of state with a finite range pair-potential. The extensive chapter on bulk results is on the one hand a validation of these modified model parameters, and on the other hand a new look at the PBD results. Indeed, today's increased computer power enables us to obtain much longer trajectories.

5.1 Bulk systems

The simulations of the bulk system revealed that reducing/disabling the dihedral potentials leads to little change in the conformational properties of PBD (discussed in Section 3.1.2). The analysis of the characteristic ratio C_{∞} (Fig. 5.1) agrees with the results of previous works [40, 41] and its temperature dependence shows a similar (though slightly larger) chain extension upon cooling with respect to the experimental results on PBD of comparable composition [93]. The fact that the characteristic ratio of the FRC is almost identical to the CRC is certainly specific to PBD and makes this polymer an ideal model system to study the influence of the dihedral barriers on the dynamics, without changing the structure of the melt.

The study of the temperature dependence of the bulk density (Section 3.2) of PBD showed that the dihedrals have a negligible influence in the high temperature region (Fig. 5.2). For a small bulk system the densities at $T \gtrsim 213$ K are basically indistinguishable between the models. However, they have a significant influence on the glass transition temperature $T_{\rm g}$. The Dalnoki-Veress fits to the data obtained from the cooling runs using all models indicate significant shifts of the $T_{\rm g}$ with weakening and disabling the dihedrals: $T_{\rm g} \approx 192$ K for the CRC model $\rightarrow 143$ K for the CRC2 model $\rightarrow 108$ K for the CRC4 model $\rightarrow 43$ K for the FRC model.

A lot of efforts were put into investigating the influence of the dihedrals on the structure and dynamics of bulk PBD in order to understand such significant shifts of the $T_{\rm g}$. In particular, the analysis of the static liquid structure factors S(q) revealed no noticeable changes in the structural correlations in PBD when modifying the strength of the



Figure 5.1: Temperature dependence of the characteristic ratio C_{∞} from Eq. (3.6) for the CRC (red circles), CRC2 (green triangles), CRC4 (purple pluses) and FRC models (blue crosses). The plot format, $\ln C_{\infty}$ versus T, is motivated by Eq. (3.7) when assuming κ to be constant. The full line presents a linear fit to the FRC data, yielding $\kappa = -0.52 \text{ K}^{-1}$. The dashed line shows the fit result to the CRC model, leading to $\kappa = -0.64 \text{ K}^{-1}$. The dotted line indicates the experimental result, $\kappa = -0.10 \text{ K}^{-1}$ (obtained for 298 K $\leq T \lesssim 373$ K), from Table VIII of Ref. [93].



Figure 5.2: Density of the bulk PBD. The empty colored circles represent the data from the cooling runs. The black triangles correspond to the equilibrium volumes, obtained from 100 ns NPT equilibration and imposed during the NVT runs for all models. The colored crosses are the data from 1 μ s long NPT runs. The solid black lines represent the Dalnoki-Veress fits using Eq. (3.10) with the parameters form Table 3.3.

dihedrals (see inset in Fig. 5.3). Backed up by more detailed analysis in analogy to works on the Kob-Andersen binary mixture glass former¹ [58, 102–104], partial static structure factors and the related Bhatia–Thornton structure factors [106] (Section 3.3), this result

¹Our model of PBD has two types of united atoms, which allows application of the same approach.



Figure 5.3: Temperature dependence of the compressibility. The data are rectified as suggested by Eq. (3.39). The orange squares are experimental results for PBD scanned from Fig. 4 of Ref. [118]. The dashed line shows Eq. (3.39) with parameters A = -0.435 and B = 987 K taken from Table I of Ref. [118]. The circles show the results for the CRC model obtained from volume fluctuations [Eq. (3.38)]. As the experimental results, they are linearized and can be fit to Eq. (3.39) (not shown). The solid line indicates the fit result to the Long–Lequeux theory obtained from a joint adjustment of the CRC data for the monomer density (top left inset) to Eq. (3.40) and for the compressibility to Eq. (3.41). The results for $k_{\rm B}T\rho\kappa_T$, found for all models are indicated by triangles. The orange crosses present the MD results from Ref. [117] for *cis*-1,4-PBD at 1 atm (systems with 128 C-atoms per chain and 32 chains), which were calculated by scanning the data from Fig. 4 and Fig. 7 of [117]. The bottom right inset shows the static structure factors S(q), that overlap for CRC and FRC models.

leads to indistinguishable compressibilities between the simulated models. An alternative approach using the Long-Lequeux theory [123] combines the more precise data of the density and compressibility obtained from the volume fluctuations extracted from long NPT simulations. This demonstrates some influence of the dihedrals, but the differences are still extremely small (Fig. 5.3). Our results are comparable to the experimental [118] and simulation [117] data on PBD from the literature.

As an aside, the gathered data on the density and compressibility allowed to verify the internal consistency of our simulations by comparing the heat capacity at constant volume C_V and constant pressure C_p (Section 3.4). Focusing on the results obtained from the simulations using the CRC model, it was shown that the thermodynamic relation between them holds:

$$C_p = C_V + T V \frac{\alpha^2}{\kappa_T} = C_V + N_c N \frac{T \alpha^2}{\rho \kappa_T} , \qquad (5.1)$$

where $\rho = N_c N/V$ is the numeric density of the system (N_c and N are the number of chains and the number if united atoms per chain correspondingly, V is the volume), $\alpha = \frac{1}{V} \frac{\partial V}{\partial T}|_{p,N_cN}$ is the thermal expansion coefficient. However, the data showed a qualitative difference with respect to typical experimental results: while the simulated C_p increases upon cooling, the experimental C_p decreases [129]. We showed that this disagreement can be solved by considering the quantum mechanical corrections of the vibrational contribution to the C_p , which are absent in our classical modelling approach. Our findings are similar to those published recently by Honguy et al. [132], where an agreement of the heat capacity extracted from MD simulations with experimental data was achieved after taking into account the corrections for quantum effects.

A consolidated framework for describing the structure and dynamics of polymers is provided by the Rouse model [1, 5]. In particular the static and dynamic correlations of the Rouse modes $\vec{X_p}(t)$ are of interest. They were investigated for our models of PBD in Section 3.5. As it is expected from our results above and from previous works [40, 41], the influence of the dihedrals on the static Rouse mode correlations $C_{pp} = \langle \vec{X}_p(0)^2 \rangle$ is mild. In fact, the C_{pp} are completely determined by the conformational properties of PBD (the parametrization has been proposed in the main text) in the studied range of temperatures carrying weak temperature dependence as well (Fig. 5.4a). Contrary to the statics, the dynamic Rouse mode correlations $C_{pp}(t)=\langle \vec{X_p}(t)\vec{X_p}(0)\rangle$ were found to be dramatically dependent on the dihedrals. When the dihedral barriers become comparable to the thermal energy, they impose constraints on the particle movements shifting the caging effect, and thus the glass transition, at much higher temperatures than expected from the collective packing constraints without the dihedrals. The slowing down of the dynamics when approaching the $T_{\rm g}$ from above exposes itself via the characteristic feature of the glassy dynamics: the two step decay of the correlation functions with the second step being stretched with respect to the ordinary exponential decay (Fig. 5.4b).

The onset of caging appearing at higher temperatures was confirmed by the analysis of the mean-square displacements (MSD) of united atoms obtained from the simulations using all our models (Section 3.6). Our studied region of temperature $213 \leq T \leq 353$ K



Figure 5.4: Panel (a): Log-log plot of the static Rouse modes autocorrelations C_{pp} . The colored empty symbols denote the results obtained from our simulations at different temperatures. The filled black circles represent the data scanned from Fig. 3a in Ref. [40]. Panel (b): Dynamic Rouse modes autocorrelations $C_{pp}(t)$ for p = 20 at T = 213 K. The colored symbols represent the data obtained from the simulations. The colored lines with black dots are the stretched exponential fits using Eq. (3.76).

is close to the $T_{\rm g}$ of the CRC model, so the caging plateaus are clearly pronounced in between the ballistic and subdiffusive regimes of the MSD with decreasing the temperature. With weakening the dihedrals, the same region of temperatures is becoming farther from the corresponding values of $T_{\rm g}$ for the CRC2, CRC4 and FRC models, where the cage effect does not occur yet. Going to lower temperature revealed the caging behavior for the FRC model (Fig. 5.5a). The relaxation map of the structural relaxation times extracted from the MSD data shows a super-Arrhenius increase with decreasing temperature for all models, which is common to all glass-forming systems (Fig. 5.5b). It also illustrates the significant slowing down of the structural relaxation in PBD, that depends on the height of the dihedral barriers.

Having summarized our findings above, we now revisit some critical points of our study of the bulk PBD.

• First of all most of our analysis of the structural properties is biased by the fact that we considered the density to be indistinguishable between the models. We have neglected the minor differences in the pair correlations that arise from the dihedrals (which is reasonable for our small bulk system). It would be interesting to study a bigger system to improve the statistics on subtle packing effects and the resolution



Figure 5.5: Panel (a): Log-log plot of the mean-square displacement $g_0(t)$ of united atoms. The colored symbols represent the data obtained from the simulations for different models. The colored solid lines with black dots are the predictions using the Rouse modes Eqs. (3.80), (3.82) and (3.83). Panel (b): The relaxation map of PBD simulated using all our models (colored symbols). The solid colored lines are the extrapolations to lower temperatures using a Vogel-Fulcher-Tammann fit, Eq. (3.84).

of static structure factors at small scattering vectors.

- Another interesting point is that in our crossover from the realistic model to the generic models by disabling all the dihedrals, the distinction between the *cis* and *trans* monomers of PBD is lost. In Section 3.4 it was shown that dihedrals that keep such a distinction (associated with the rotation along the double-bond axis of the PBD monomer) are the "hard" variables and the dynamics is determined by the other "soft" dihedrals. A deeper insight on the dynamical properties of PBD can be obtained keeping the "hard" dihedrals unmodified and disabling only the "soft" ones ².
- In a different direction, the speed up of the dynamics using weaker dihedrals while preserving the structure of the polymer melt can be turned into an advantage as a new equilibration method. In principle, it should be possible to use the weaker dihedral configuration to equilibrate the large-scale structure of the system faster. Then, as a second step one can reintroduce the realistic potential to equilibrate the local structure. Using this trick, it should be possible to obtain equilibrated configurations of the CRC model in the supercooled regime around T_g which would be impossible to obtain by brute force dynamics.

²It is probably redundant to reduce only the "soft" dihedrals. Our results show that when reducing all of them by factor a of 4, the double bond dihedrals preserve their property of being "hard" variables effectively making the change from cis to trans impossible.

٠ The last, but not least remark can be made when relating our results to the common theories that describe the glassy dynamics, such as the mode-coupling theory [40, 41, 148] and the more recent elastically collective nonlinear Langevin equation (ECNLE) theory [119, 120, 149]. Being inspired from and tested on the simple glass forming liquids, they attempt to relate the dynamics with the structure of the system. The structure of the liquids is described by the spatial correlations, which are typically treated using central symmetry and averaged isotropically. While this is completely reasonable for the liquids, where the pair correlations are the most important, it is not clear whether this averaging can be applied for the systems with three- and/or four-point potentials, such as the dihedral potentials. The qualitative description, of course, should hold, since the purely structural constraints in liquids and the energetic constraints in polymers effectively lead to the same caging behavior. Yet, the different energy scales between the dihedral barriers and pair interaction barriers result in the cage effect appearing at higher temperatures for polymers. Therefore, it appears that an important challenge is to incorporate intrachain barriers into a liquid-state-theory based approach.

5.2 Film systems

The simulations of the PBD films supported on a graphite wall showed that disabling the dihedral potential does not lead to visible changes in the density profile of the films. The profiles overlap not only in the bulk-like middle of the film, but also in the region next to the wall and at the free surface (Fig. 5.6a). Extracting the thickness of the films using the Gibbs dividing surface (GDS) method revealed a tiny difference: the thickness of the films simulated using the CRC model are a fraction of an Ångström lower than the ones for the FRC model (Fig. 5.6b). We attribute this to the better averaging statistics in the films (they are ~ 18 times bigger than the bulk systems), so the influence of the dihedrals on the density should become more visible (though it was not detected by a visual inspection of the density profiles).

Upon cooling the films, the temperature dependence of the thickness show a change of slope (Fig. 5.6b) which suggests the glass transition at different temperatures depending on the presence of the dihedral potentials. Extracting the values of $T_{\rm g}$ for both models using the Dalnoki-Veress approach revealed that they agree with the bulk values within the precision of our small systems and fast cooling rates.

In order to study the reasons for such an agreement of the $T_{\rm g}$ the dynamical properties across the film were studied in a layer-resolved fashion. Two characteristic quantities were chosen: the mean-square displacements (MSD), that describe the dynamics of the united atoms, and the autocorrelation of the dipole moment vector (DACF) associated to each monomer (with the perpendicular to the film component being experimentally relevant). A consistent picture in the time window of the structural relaxation was observed between both quantities in case of the CRC model: the dynamics is slowed down at the graphite wall (which agrees with the previously reported results of Refs. [31, 35]) and sped up at the free surface in both parallel (xy) and perpendicular (z) to the film directions. While the same behavior is observed in case if the FRC model for the xy direction, the structural relaxation times in z direction extracted from the MSD correspond to a different regime in the DACF, which is slower at both interfaces (the hints of a similar slowing down was also observed for the layers at the free surface for the CRC model, but at longer times). The early appearance of such a relaxation step was attributed to the fact, that in the FRC model there are no dihedral potentials to preserve the configuration of the monomer. So the dipole moment vector autocorrelation (calculated in the same way as for the CRC



Figure 5.6: Panel (a): The density profiles of the films obtained from the simulations using the CRC (red) and FRC (blue) models at different temperatures. The vertical dot-dashed lines are the positions of the graphite layers. The vertical dotted lines are the positions of the Gibbs dividing surfaces for the CRC model. The horizontal dotted lines correspond to the density of the bulk systems at the same temperatures. Panel (b): The temperature dependence of the film thickness obtained by the GDS method from the simulations using the CRC (red) and the FRC (blue) models. The solid black lines are the fits using Eq. (4.4) with the parameters from Table 4.1.

model) is sensitive not only to the reorientation of the monomer (as it is for the CRC model), but also to the rearrangements of the united atoms within the monomer.

Further analysis of the structural relaxation times extracted from the MSD for both models is summarized in Fig. 5.7. It appears, that the slowing down effect of the wall and the speeding up at the free surface average out, which explains the agreement of the values of the total $T_{\rm g}$ with the the bulk. Moreover, the surface effects are mildly affected by the presence of the dihedrals themselves. They act with respect to the total order of magnitude of the dynamics, which is set by the dihedrals across the whole system. The quantitative analysis of the relaxation time gradients by fitting the data using the widely assumed empirical double-exponential form [154, 155]

$$\tau(z,T) = \tau_{\rm b}(T) \exp\left[-A(T) \exp\left(-\frac{z}{\xi}\right)\right] \,, \tag{5.2}$$

showed a similarity for the relaxation in the parallel to the film direction between the CRC and FRC models at the temperatures that have the same order of magnitude of the relaxation in the bulk-like region in the middle of the film. Being at different temperatures (densities), these results suggest that at the free surface the gradients of the dynamics in the xy direction can be related to the fact that at the free surface the cage surrounding each particle is open, which facilitates the relaxation. For the relaxation in the perpendicular to the film direction, the gradients are different between the models and temperatures suggesting a complex interplay between the confinement effects, the density and the dihedrals.

After having summarized our results for the films, let us also review the critical points in them.

- In general, a significant amount of our data suffers from insufficient statistics and lack of time for more detailed analysis. Moreover, the slowing down of the dynamics with decreasing the temperature barely allows to reach the structural relaxation time within the time of the simulations. This is the most crucial at the wall, where the dynamics is even slower. Therefore, more simulation time would be required to improve our analysis.
- As a second point, we would like to come back to the discussion of the overlapping density profiles. In principle, it is possible to express the density profile using pair



Figure 5.7: The natural logarithm of the structural relaxation times extracted from the MSD data at different slabs in the film. The symbols denote different temperatures and the colors mark different models.

distribution functions similarly to Eq. (3.9)³, by introducing the external potentials, that act on the system mimicking the surface effects. Such a description would, however, require knowledge of the radial distribution function in confinement. It

 $^{^{3}\}mathrm{In}$ fact, Eq. (8.1.34) of Ref. [156] proposes such an expression.

would be interesting to study its behavior at the interfaces and how it is influenced by the dihedrals.

- Another remark concerns the dielectric relaxation for the FRC model. We have already noted, that disabling the dihedrals leads to loosing the distinction between the *cis* and *trans* monomers of PBD. It would be interesting to study the dipole autocorrelation in the model where these "hard" dihedrals are preserved and only the "soft" ones are disabled. Alternatively, it is common in the simulations using the generic models to consider the dipole moment to be perpendicular to the bonds in the chain backbone. Applying this approach for the double bonds of PBD may help to understand better the segmental dynamics and the influence of the dihedrals.
- It is interesting to speculate what happens if the simulations could be pushed to temperatures near the experimental $T_{\rm g}$. While the range of the surface-induced changes of the dynamics is weak in the temperature interval of the current simulations, covering only a few nanometers, it might be expected to increase upon cooling. This low-T, strongly supercooled regime is the focus of the ECNLE theory which then predicts much more long-ranged gradients. If this was true, then in thin supported films the gradients from the substrate and the free interface should interfere, erasing the bulk-like behavior in the center of the film and leading, perhaps, to a continuous decrease of the local relaxation time from high values at the substrate to low values at the free interface. It would be interesting to check whether such a highly inhomogeneous situation could be observed in simulations, e.g. by studying thinner films or by harnessing a more efficient equilibration method based on reduced dihedral barriers.
- A complex interplay between the temperature dependence of the film thickness, the depth of the perturbation of the dynamics by the surface effects, as well as their direction and amplitude determines the film-averaged glass transition temperature. Thus, a comparison between the realistic and the coarse-grained approaches to the dynamics of polymers in confinement should be reviewed with a well defined mapping between the length and time scales, and the temperature.

Finally, we would like to address the question raised in the title of the thesis: Is the interfacial mobility in glass forming polymer film determined by collective motion or intramolecular energetics? Such a general question is very difficult to answer: a vast amount of experimental and simulation data (including ours) suggest that both factors are intimately intertwined. Moreover, the terms "intramolecular energetics" and "collective motion" themselves cover a wide range of different effects, that vary even depending on the studied polymer. Thus, the better question to ask would be how their interplay affects the dynamics? In our work we have focused on one of the intramolecular contributions – the energetic constraints on the torsional (dihedral) angle – for a polybutadiene polymer. We showed that their presence affects the collective caging effect shifting it to higher temperature, which leads to strongly non-Arrhenius glassy dynamics. While this coupling determines the structural relaxation in the system, it has an extremely small effect on the density (for PBD!). At the interfaces of the films, there is an additional factor, that comes into play – the surface effects that are determined by the type and the properties of the confinement. They are intertwined with the energetics and the collective motion influencing each other and, thus, the interfacial mobility in the films.

Résumé en français

Mobilité interfaciale dans les films polymères vitrifiables : est-elle déterminée par le mouvement collectif ou l'énergie intramoleculaire ?

Nous proposons une étude systématique de l'influence des barrières intramoléculaires et de l'interaction avec un substrat sur la dynamique vitreuse, et ceci par des simulations numériques chimiquement réalistes. Notre travail se concentre sur un type particulier d'interactions intramoléculaires: les barrières de torsion (dièdres), qui contrôlent majoritairement les conformations d'une chaîne polymère, contrairement les énergies de liasons et de valence.

Afin d'étudier l'influence de contraintes de torsion réalistes sur les propriétés des polymères vitreux, nous avons adapté un modèle atomes unifiés du copolymère aléatoire *cis-trans* du 1,4-polybutadiène (PBD) pour des simulations de dynamique moléculaire classique avec le code LAMMPS [17, 43, 44]. Ce modèle a été développé précédemment à partir de calculs numériques de chimie quantique et validé par rapport à des expériences [39, 45, 46]. Ce travail de doctorat fait suite aux travaux récents étudiants le PBD confiné entre deux parois de graphite. Le but est d'étudier le comportement avec une surface libre, ainsi que l'influence des potentiels dièdres. Notre modèle se décline dans en quatre versions , où nous faisons varier l'échelle de tous les potentiels de torsion (dièdres) dans le système :

- Modèle de chaîne chimiquement réaliste (CRC) avec les potentiels dièdres non modifiés (qui fournissent la distinction entre les monomères *cis* et *trans*).
- Modèles CRC2 et CRC4 avec les potentiels dièdres réduits d'un facteur 2 et 4 respectivement.
- Modèle FRC (Freely rotating chain) avec tous les potentiels dièdres désactivés. Ici, la distinction entre les groupes *cis* et *trans* n'a pas lieu.

Une telle interpolation permet de faire le lien avec les modèles génériques de type

billes-ressorts, largement utilisés, dans lesquels les détails chimiques réalistes (y compris les dièdres) sont confondus dans une seule interaction à "gros grains". Notre modèle FRC ressemble à un modèle billes-ressorts, à une exception près : la longueur des liaisons est beaucoup plus courte par rapport à la taille des particules. Le modèle billes-ressort le plus utilisé de type Kremer-Grest [20, 23, 24] a une seule échelle de longueur qui détermine à la fois la taille des billes et la longueur des liaisons entre elles.

Nos simulations ont été effectuées en volume en utilisant les 4 modèles et dans une géométrie confinée (film supporté sur graphite) en utilisant les modèles CRC et FRC dans la gamme de températures $213 \le T \le 353$ K.

Il est à noter que pour simuler des films supportés (incluant donc une surface libre), il n'est pas possible d'utiliser des corrections de queue ("tail corrections") pour prendre en compte la portée infinie du potentiel de paire. Les paramètres du modèle original ont dû être adaptés pour donner la même équation d'état avec un potentiel de paire à portée finie. Le chapitre sur les résultats en phase volumique est d'une part une validation de ces paramètres de modèle modifiés, et d'autre part un nouveau regard sur les résultats PBD. En effet, la puissance informatique accrue d'aujourd'hui nous permet d'obtenir des trajectoires beaucoup plus longues.

Systèmes en phase volumique

Les simulations du système volumique ont révélé que la réduction ou désactivation des potentiels dièdres entraîne peu de changements dans les propriétés conformationnelles du PBD (discuté en Section 3.1.2). L'analyse du rapport caractéristique C_{∞} (Fig. 5.8) est en accord avec les résultats de travaux antérieurs [40, 41] et sa dépendance de la température montre une extension de chaîne similaire (bien que légèrement plus importante) lors du refroidissement par rapport aux résultats expérimentaux sur le PBD de composition comparable [93]. Le fait que le rapport caractéristique du FRC soit presque identique à celui du CRC est certainement spécifique au PBD et fait de ce polymère un système modèle idéal pour étudier l'influence des barrières dièdres sur la dynamique, sans changer la structure du système.

L'étude de la dépendance en température de la densité apparente (Section 3.2) du PBD a montré que les dièdres ont une influence négligeable dans la région des hautes



Figure 5.8: Dépendance en température du rapport caractéristique C_{∞} , Eq. (3.6), pour les modèles CRC (cercles rouges), CRC2 (triangles verts), CRC4 (plus violet) et FRC (croix bleues). Le format du graphique, $\ln C_{\infty}$ en fonction de T, est motivé par l'Éq. (3.7) en supposant que κ est constant. La ligne pleine présente un ajustement linéaire aux données de la FRC, donnant $\kappa = -0, 52 \text{ K}^{-1}$. La ligne en pointillé présente le résultat de l'ajustement au modèle CRC, conduisant à $\kappa = -0, 64 \text{ K}^{-1}$. La ligne en pointillés indique le résultat expérimental, $\kappa = -0, 10 \text{ K}^{-1}$ (obtenu pour 298 K $\leq T \lesssim 373$ K), d'après le Tableau 8 de la Réf. [93].

températures (Fig. 5.9). Pour un petit système massif, les densités à $T \gtrsim 213$ K sont indiscernables entre les modèles. Cependant, elles ont une influence significative sur la température de transition vitreuse T_g . L'ajustement, selon la formule de Dalnoki-Veress, aux données obtenues à partir des cycles de refroidissement, en utilisant tous les modèles, indiquent des décalages significatifs de la température de transition vitreuse (T_g) avec l'affaiblissement ou la désactivation des dièdres : T_g d'environ 192 K pour le modèle CRC \rightarrow 143 K pour le modèle CRC2 \rightarrow 108 K pour le modèle CRC4 \rightarrow 43 K pour le modèle FRC.

Nous avons consacrée de nombreux efforts à l'étude de l'influence des dièdres sur la structure et la dynamique du PBD massif afin de comprendre ces changements significatifs du $T_{\rm g}$. En particulier, l'analyse des facteurs de structure statique S(q) n'a révélé aucun changement notable des corrélations structurelles dans le PBD lors de la modification de l'amplitude des dièdres (voir l'encart dans la Fig. 5.10). Etayé par une analyse plus détaillée en analogie avec les travaux sur le mélange binaire vitrifiable de Kob-Andersen⁴ [58, 102–104], des facteurs de structure statique partiels et des facteurs de structure connexes de Bhatia–Thornton [106] (Section 3.3), ce résultat conduit à des compressibilités

 $^{^4\}mathrm{Notre}$ modèle de PBD possè de deux types d'atomes unifiés, ce qui permet d'appliquer la même approche.



Figure 5.9: Densité du PBD en phase volumique. Les cercles colorés vides représentent les données des cycles de refroidissement. Les triangles noirs correspondent aux volumes d'équilibre, obtenus à partir de l'équilibration NPT de 100 ns et imposés pendant les simulations NVT pour tous les modèles. Les croix colorées représentent les données des cycles NPT longs de 1 μ s. Les lignes noires pleines représentent les ajustements de type Dalnoki-Veress utilisant l'Éq. (3.10) avec les paramètres du Tableau 3.3.

indiscernables entre les modèles simulés. Une approche alternative utilisant la théorie de Long-Lequeux [123] combine les données plus précises de la densité et de la compressibilité obtenues à partir des fluctuations de volume extraites de simulations longues en NPT. Cela démontre une certaine influence des dièdres, mais les différences restent extrêmement faibles (Fig. 5.10). Nos résultats sont comparables aux données expérimentales [118] et de simulation [117] sur le PBD issues de la littérature.

Par ailleurs, les données recueillies sur la densité et la compressibilité ont permis de vérifier la cohérence interne de nos simulations en comparant la chaleur spécifique à volume constant C_V et à pression constante C_p (Section 3.4). En se concentrant sur les résultats obtenus à partir des simulations utilisant le modèle CRC, il a été démontré que la relation thermodynamique entre C_p et C_V prend la forme habituelle :

$$C_p = C_V + T V \frac{\alpha^2}{\kappa_T} = C_V + N_{\rm c} N \frac{T \alpha^2}{\rho \kappa_T} \; , \label{eq:cp}$$

où $\rho = N_c N/V$ est la densité particulaire du système (N_c et N sont le nombre de chaînes et le nombre d'atomes unifiés par chaîne respectivement, Vest le volume), $\alpha = \frac{1}{V} \frac{\partial V}{\partial T}|_{p,N_cN}$ est le coefficient de dilatation thermique. Cependant, les données ont montré une différence



Figure 5.10: Dépendance en température de la compressibilité. Les données sont rectifiées comme suggéré par l'Éq. (3.39). Les carrés oranges représentent les résultats expérimentaux pour le PBD numérisé à partir de la Fig. 4 de la Réf. [118]. La ligne en pointillés indique l'Éq. (3.39) avec les paramètres A = -0, 435 et B = 987 K tirés du Tableau I de Réf. [118]. Les cercles montrent les résultats pour le modèle CRC obtenu à partir des fluctuations de volume [Éq. (3.38)]. Comme les résultats expérimentaux, ils sont linéairement dépenant en 1000/T et peuvent être ajustés à l'Éq. (3.39) (non montré). La ligne continue indique le résultat de l'ajustement à la théorie de Long–Lequeux obtenu à partir d'un ajustement conjoint des données CRC pour la densité des monomères (encart supérieur gauche) à l'Éq. (3.40) et pour la compressibilité à l'Éq. (3.41). Les résultats pour $k_{\rm B}T\rho\kappa_T$, trouvés pour tous les modèles sont indiqués par des triangles. Les croix oranges présentent les résultats MD de la Réf. [117] pour le *cis*-1,4-PBD à 1 atm (systèmes avec 128 atomes de carbone par chaîne et 32 chaînes), qui ont été calculés en scannant les données des Fig. 4 et Fig. 7 de la Réf. [117]. L'encart en bas à droite montre les facteurs de structure statique S(q), qui se chevauchent pour les modèles CRC et FRC.

qualitative par rapport aux résultats expérimentaux typiques : alors que le C_p simulé augmente lors du refroidissement, le C_p expérimental diminue [129]. Nous avons montré que ce désaccord peut être résolu en considérant des corrections issues de prédictions de mécaniques quantiques à la contribution vibrationnelle au C_p , qui sont absentes dans notre approche de modélisation classique. Nos résultats sont similaires à ceux publiés récemment par Honguy et al [132], où un accord entre chaleur spécifique extraite des simulations MD et les données expérimentales a été obtenu après avoir pris en compte les corrections pour les effets quantiques précédement évoqués.

Un cadre consolidé pour décrire la structure et la dynamique des polymères est fourni par le modèle de Rouse [1, 5]. En particulier, les corrélations statiques et dynamiques des modes de Rouse $\vec{X_p}(t)$ sont intéressantes. Elles ont été étudiées pour nos modèles de PBD dans la Section 3.5. Comme on peut s'y attendre à partir de nos résultats ci-dessus et des travaux précédents [40, 41], l'influence des dièdres sur les corrélations des modes de Rouse statiques $C_{pp}=\langle \vec{X_p}(0)^2\rangle$ est faible. En fait, les C_{pp} sont complètement déterminés par les propriétés conformationnelles du PBD (la paramétrisation a été proposée dans le texte principal) dans la gamme étudiée de températures : une faible dépendance en température est observée (Fig. 5.11a). Contrairement à la statique, les corrélations dynamiques des modes de Rouse $C_{pp}(t) = \langle \vec{X_p}(t) \vec{X_p}(0) \rangle$ se sont avérées dépendre fortement des dièdres. Lorsque les barrières dièdres deviennent comparables à l'énergie thermique, elles imposent des contraintes sur le mouvement des particules, déplaçant l'effet de cage, et donc la transition vitreuse, à des températures beaucoup plus élevées que celles attendues à partir des contraintes collectives d'encombrement sans les dièdres. Le ralentissement de la dynamique lors de l'approche à la $T_{\rm g}$ par le haut se montre via une caractéristique de la dynamique vitreuse : la décroissance en deux étapes des fonctions de corrélation, la deuxième étape étant étirée par rapport à la décroissance exponentielle ordinaire (Fig. 5.11b).

L'apparition du phénomène de cage à des températures plus élevées a été confirmée par l'analyse des déplacements carrés moyens (MSD) des atomes unifiés obtenus à partir des simulations utilisant tous nos modèles (Section 3.6). La région de température étudiée, $213 \leq T \leq 353$ K, est proche de la température $T_{\rm g}$ du modèle CRC, de sorte que les plateaux de cage sont clairement prononcés entre les régimes balistique et subdiffusif du MSD avec la diminution de la température. En affaiblissant les dièdres, la même région de températures s'éloigne des valeurs de $T_{\rm g}$ correspondantes pour les modèles CRC2, CRC4 et FRC, où l'effet de cage ne se produit pas encore. Le passage à une température plus basse a révélé le comportement de cage pour le modèle FRC (Fig. 5.12a). Le diagramme de relaxation obtenu à partir des temps de relaxation structurelle extraite des données MSD montre une augmentation super-Arrhenius avec la diminution de la température pour tous les modèles, ce qui est commun à tous les verres "fragiles" (Fig. 5.12b). Il illustre également le ralentissement significatif de la relaxation structurelle dans le PBD, qui dépend de la hauteur des barrières dièdres.


Figure 5.11: Panneau (a) : Tracé log-log des autocorrélations des modes de Rouse statiques C_{pp} . Les symboles vides colorés indiquent les résultats obtenus à partir de nos simulations à différentes températures. Les cercles noirs remplis représentent les données scannées de la Fig. 3a dans la Réf. [40]. Panneau (b) : Autocorrélations des modes de Rouse dynamiques $C_{pp}(t)$ pour p = 20 à T = 213 K. Les symboles colorés représentent les données obtenues à partir des simulations. Les lignes colorées avec des points noirs sont les ajustements exponentiels étirés en utilisant Éq. (3.76).



Figure 5.12: Panneau (a) : Tracé log-log du déplacement quadratique moyen $g_0(t)$ des atomes unifies. Les symboles colorés représentent les données obtenues à partir des simulations pour différents modèles. Les lignes solides colorées avec des points noirs sont les prédictions utilisant les modes de Rouse, Éqs. (3.80), (3.82) et (3.83). Panneau (b) : La diagramme de relaxation du PBD simulée pour tous nos modèles (symboles colorés). Les lignes colorées pleines sont les extrapolations à des températures plus basses à l'aide d'un ajustement Vogel-Fulcher-Tammann, Éq. (3.84).

Après avoir résumé nos résultats ci-dessus, nous allons maintenant revenir sur certains points de notre étude du PBD en phase volumique.

• Tout d'abord, la plupart de notre analyse des propriétés structurelles est biaisée

par le fait que nous avons considéré que la densité était indiscernable entre les modèles. Nous avons négligé les différences mineures entre les corrélations de paires qui découlent des dièdres (ce qui est raisonnable pour notre petit système massif). Il serait intéressant d'étudier un système plus grand pour améliorer les statistiques sur les effets subtils de l'empaquetage, et la résolution des facteurs de structure statique aux petits vecteurs de diffusion.

- Un autre point intéressant est que dans notre passage du modèle réaliste aux modèles génériques en désactivant tous les dièdres, la distinction entre les monomères *cis* et *trans* de PBD est perdue. Dans la Section 3.4, il a été montré que les dièdres qui conservent une telle distinction (associée à la rotation le long de l'axe de la double liaison du monomère PBD) sont des variables "dures" et que la dynamique est déterminée par les autres dièdres "souples". Un aperçu plus approfondi des propriétés dynamiques du PBD peut être obtenu en gardant les dièdres "durs" non modifiés et en ne désactivant que les dièdres "mous".⁵
- Dans une autre direction, l'accélération de la dynamique en utilisant des dièdres plus faibles tout en préservant la structure du polymère fondu peut être transformée en avantage, dans le sens d'une nouvelle méthode d'équilibration. En principe, il devrait être possible d'utiliser des dièdres plus faibles pour équilibrer plus rapidement la structure à grande échelle du système. Ensuite, dans un deuxième temps, on peut réintroduire le potentiel réaliste pour équilibrer la structure locale. En utilisant cette astuce, il devrait être possible d'obtenir des configurations équilibrées du modèle CRC dans le régime surfondu autour de T_g , ce qui serai impossible à obtenir par une dynamique par force brute.
- La dernière remarque, mais non la moindre, peut être faite en mettant en relation nos résultats avec quelques théories réppondus qui décrivent la dynamique vitreuse, telles que la théorie de couplage de modes [40, 41, 148] et la plus récente version basée sur l'équation de Langevin non linéaire incluant des effets élastique et collectif (ECNLE) [119, 120, 149]. Inspirés et testés sur des liquides simples formant un verre, elles tentent de relier la dynamique à la structure du système. La structure des liquides est décrite par les corrélations spatiales, qui sont typiquement traitées en

⁵Il est probablement redondant de réduire uniquement les dièdres "mous". Nos résultats montrent qu'en les réduisant tous par un facteur 4, les dièdres de double liaison conservent leur propriété de variables "dures" rendant effectivement impossible le passage d'un état cis à un état trans.

utilisant la symétrie centrale et une moyenne isotrope. Bien que cela soit tout à fait raisonnable pour les liquides, où les corrélations de paires sont les plus importantes, il n'est pas clair si cette moyenne peut être appliquée aux systèmes avec des potentiels à trois et/ou quatre points, tels que les potentiels dièdres. La description qualitative, bien sûr, devrait être valable, puisque les contraintes purement structurelles dans les liquides et les contraintes énergétiques dans les polymères conduisent effectivement au même comportement de cage. Pourtant, les différentes échelles d'énergie entre les barrières dièdres et les barrières d'interaction de paires font que l'effet de cage apparaît à des températures plus élevées pour les polymères. Par conséquent, il semble qu'un défi important consiste à intégrer les barrières intra-chaîne dans une approche basée sur la théorie de l'état liquide.

Film polymère supporté

Nos simulations des films PBD supportés par une paroi en graphite ont montré que la désactivation du potentiel dièdre n'entraîne pas de changements visibles dans le profil de densité des films. Les profils se superposent non seulement au milieu du film, mais aussi dans la région proche de la paroi et à la surface libre (Fig. 5.13a). L'extraction de l'épaisseur des films à l'aide de la méthode GDS (Gibbs dividing surface) a révélé une différence minime : l'épaisseur des films simulés à l'aide du modèle CRC est inférieure d'une fraction d'Ångström à celle du modèle FRC (Fig. 5.13b). Nous attribuons cela aux meilleures statistiques de moyennage dans les films (ils sont ~ 18 fois plus grands que les systèmes en phase volumique), de sorte que l'influence des dièdres sur la densité devrait devenir plus visible (bien qu'elle n'ait pas été détectée par une inspection visuelle des profils de densité).

Lors du refroidissement des films, la dépendance de la température de l'épaisseur montre un changement de pente (Fig. 5.13b), ce qui suggère la transition vitreuse à une température différente selon la présence des potentiels diédraux. L'extraction des valeurs de $T_{\rm g}$ pour les deux modèles en utilisant l'approche Dalnoki-Veress a révélé qu'elles sont en accord avec les valeurs en phase volumique dans la limite de précision de nos petits systèmes et des vitesses de refroidissement rapides.

Afin d'étudier les raisons d'un tel accord des $T_{\rm g}$ entre volume et film, les propriétés

dynamiques à travers le film ont été étudiées d'une manière résolue en couche. Deux quantités typiques ont été choisies : les déplacements carrés moyens (MSD), qui décrivent la dynamique des atomes unifiés, et l'autocorrélation du vecteur du moment dipolaire (DACF) associé à chaque monomère (la composante perpendiculaire au film étant expérimentalement pertinente). Une image cohérente dans la fenêtre temporelle de la relaxation structurelle a été observée entre les deux quantités dans le cas du modèle CRC : la dynamique est ralentie au niveau de la paroi du graphite (ce qui est en accord avec les résultats précédemment rapportés dans les références [31, 35]) et accélérée au niveau de la surface libre à la fois parallèlement (xy) et perpendiculairement (z) à l'orientation du film. Alors que le même comportement est observé dans le cas du modèle FRC pour la direction xy, les temps de relaxation structurelle dans la direction z extraits du MSD correspondent à un régime différent dans le DACF, qui est plus lent aux deux interfaces (l'indication d'un ralentissement similaire a également été observée pour les couches à la surface libre pour le modèle CRC, mais à des temps plus longs). L'apparition précoce d'une telle étape de relaxation a été attribuée au fait que, dans le modèle FRC, il n'y a pas de potentiels dièdres pour préserver la configuration du monomère. Ainsi, l'autocorrélation du vecteur du moment dipolaire (calculée de la même manière que pour le modèle CRC) est sensible



Figure 5.13: Panneau (a) : Les profils de densité des films obtenus a partir des simulations utilisant les modeles CRC (rouge) et FRC (bleu) à différentes températures. Les lignes verticales en pointillés sont les positions des couches de graphite. Les lignes pointillées verticales sont les positions des surfaces de division de Gibbs pour le modèle CRC. Les lignes pointillées horizontales correspondent à la densité des systèmes en phase volumique aux mêmes températures. Panneau (b) : La dépendance en température de l'épaisseur du film obtenue par la méthode GDS à partir des simulations utilisant les modèles CRC (rouge) et FRC (bleu). Les lignes noires pleines sont les ajustements utilisant Éq. (4.4) avec les paramètres du Tableau 4.1.

non seulement à la réorientation du monomère (comme pour le modèle CRC), mais aussi aux réarrangements des atomes unifiés au sein du monomère.

Une analyse plus approfondie des temps de relaxation structurelle extraits du MSD pour les deux modèles est résumée dans la Fig. 5.14. Il apparaît que l'effet de ralentissement à la paroi et l'accélération à la surface libre se compensent, ce qui explique la concordance des valeurs de $T_{\rm g}$ total avec le volume. De plus, les effets de surface sont légèrement affectés par la présence des dièdres eux-mêmes. Ils agissent par rapport à l'ordre de grandeur total de la dynamique, qui est fixé par les dièdres dans l'ensemble du système. L'analyse quantitative des gradients de temps de relaxation en ajustant les données à l'aide de la forme empirique double-exponentielle largement admise [154, 155] :

$$\tau(z,T) = \tau_{\rm b}(T) \exp\left[-A(T) \exp\left(-\frac{z}{\xi}\right)\right] \;, \label{eq:tau}$$

a montré une similarité pour la relaxation dans la direction parallèle au film entre les modèles CRC et FRC aux températures qui ont le même ordre de grandeur de la relaxation dans la région au milieu du film. Ces résultats, obtenus à différentes températures (densités), suggèrent qu'à la surface libre, les gradients de la dynamique dans la direction xy peuvent être liés au fait qu'à la surface libre, la cage entourant chaque particule est ouverte, ce qui facilite la relaxation. Pour la relaxation dans la direction perpendiculaire au film, les gradients sont différents entre les modèles et les températures suggérant une interaction complexe entre les effets de confinement, la densité et les dièdres.

Après avoir résumé nos résultats pour les films, passons également en revue quelques points critiques de ceux-ci.

- En général, une grande partie de nos données souffre de statistiques insuffisantes et du manque de temps pour une analyse plus détaillée. De plus, le ralentissement de la dynamique en diminuant la température permet à peine d'atteindre le temps de relaxation structurelle dans le temps des simulations. Ceci est le plus crucial au niveau de la paroi, où la dynamique est encore plus lente. Par conséquent, un temps de simulation plus long serait nécessaire pour améliorer notre analyse.
- En second lieu, nous aimerions revenir sur la discussion concernant la superposition des profils de densité. En principe, il est possible d'exprimer le profil de densité



Figure 5.14: Le logarithme naturel des temps de relaxation structurelle extraits des données MSD à différentes profondeures du film. Les symboles indiquent les différentes températures et les couleurs les différents modèles.

à l'aide de fonctions de distribution par paire de manière similaire à l'Éq. (3.9),⁶ en introduisant des potentiels externes, qui agissent sur le système en imitant les effets de surface. Une telle description nécessiterait cependant la connaissance de la

 $^{^6\}mathrm{En}$ fait, l'Eq. (8.1.34) de la Réf. [156] propose une telle expression.

fonction de distribution radiale sous confinement. Il serait intéressant d'étudier son comportement aux interfaces, et comment il est influencé par les dièdres.

- Une autre remarque concerne la relaxation diélectrique pour le modèle FRC. Nous avons déjà noté que la désactivation des dièdres conduit à la perte de la distinction entre les monomères *cis* et *trans* du PBD. Il serait intéressant d'étudier l'autocorrélation des dipôles dans le modèle où ces dièdres "durs" sont préservés et où seuls les dièdres "mous" sont désactivés. Alternativement, il est courant dans les simulations utilisant les modèles génériques de considérer que le moment dipolaire est perpendiculaire aux liaisons du squelette de la chaîne. L'application de cette approche pour les doubles liaisons du PBD peut aider à mieux comprendre la dynamique segmentale et l'influence des dièdres.
- Il est intéressant de spéculer sur ce qui se passerait si les simulations pouvaient être poussées à des températures plus proches des $T_{\rm g}$ expérimentale. Bien que la portée des changements de la dynamique induits par la surface soit faible dans l'intervalle de température des simulations actuelles, couvrant seulement quelques nanomètres, on pourrait s'attendre à ce qu'elle augmente lors du refroidissement. Ce régime à faible T, fortement surfondu, est au centre de la théorie ECNLE qui prédit alors des gradients beaucoup plus étendus. Si cela était vrai, alors dans les films minces supportés, les gradients du substrat et de l'interface libre devraient interférer, effaçant le comportement de masse au centre du film et conduisant, peut-être, à une diminution continue du temps de relaxation local des valeurs élevées au substrat aux valeurs basses à l'interface libre. Il serait intéressant de vérifier si une telle situation très inhomogène peut être observée dans les simulations, par exemple en étudiant des films plus fins ou en exploitant une méthode d'équilibration plus efficace basée sur des barrières dièdres réduites.
- Une interaction complexe entre la dépendance de la température de l'épaisseur du film, la profondeur de la perturbation de la dynamique par les effets de surface, ainsi que leur direction et leur amplitude, détermine la température de transition vitreuse moyenne du film. Ainsi, une comparaison entre les approches réalistes et à gros grains de la dynamique des polymères en confinement devrait être entreprise à nouveau avec une correspondance bien définie entre les échelles de longueur et de temps, et de la température.

Enfin, nous aimerions aborder la question soulevée dans le titre de la thèse : La mobilité interfaciale dans un film polymère vitrifiable est-elle déterminée par le mouvement collectif ou par l'énergie intramoléculaire ? Il est très difficile de répondre de manière franche à une question aussi générale : un grand nombre de données expérimentales et de simulation (y compris les nôtres) suggèrent que les deux facteurs sont intimement liés. De plus, les termes "énergie intramoléculaire" et "mouvement collectif" couvrent eux-mêmes un large éventail d'effets différents, qui varient même en fonction du polymère étudié. Par conséquent, la meilleure question à poser serait de savoir comment leur interaction affecte la dynamique. Dans notre travail, nous nous sommes concentrés sur l'une des contributions intramoléculaires - les contraintes énergétiques sur l'angle de torsion (dièdre) - pour un polymère de polybutadiène. Nous avons montré que leur présence affecte l'effet de cage collectif en le déplaçant vers une température plus élevée, ce qui conduit à une dynamique vitreuse fortement non-Arrhenius. Alors que ce couplage détermine la relaxation structurelle du système, il a un effet extrêmement faible sur la densité (pour le PBD !). Aux interfaces des films, un facteur supplémentaire entre en jeu : les effets de surface qui sont déterminés par le type et les propriétés du confinement. Ils sont intimement liés à l'énergétique et au mouvement collectif qui s'influencent mutuellement et, par conséquent, à la mobilité interfaciale dans les films.

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Appendix A

Simulation map

See next page.

This appendix gives a "genealogical tree" to trace the history of the simulated systems, i.e. from which parent system start configurations were taken. Each box in the figures below represents a simulation. A short description of the type of the simulation and its length are given inside each box. The boxes with white background denote the simulations using the original truncated LJ potential with tail corrections for pair interactions. The boxes with colored background are the simulations using the shifted and smoothed LJ potential. The colors mark different models: CRC - red, CRC2 - green, CRC4 - purple, FRC- blue. The arrows show the sequences of the simulations. If an arrow starts from the side of the box, then the last configuration of that run was used to start the next one. If an arrow starts from the bottom of the box, the configuration at the time point specified next to the arrow was used to start the next run.











Figure A.1: The schematic map of the simulations of the bulk PBD.



Figure A.2: The schematic map of the simulations of the PBD supported films.

Appendix B

Torsional potential

The original model of Ref. [26] represents the torsional potential $U_{\rm tor}(\phi)$ by the function

$$U_{\rm tor}(\phi) = \frac{1}{2} \sum_{n=1}^{6} k_n \left[1 - \cos(n\phi) \right] \,, \tag{B.1}$$

where the torsion angle lies in $0^{\circ} \leq \phi \leq 360^{\circ}$ and the energy coefficients k_n are given in Refs. [26, 39, 157]. When determining the torsion angles from the Cartesian coordinates of the united atoms one gets the cosine of ϕ . It is therefore computationally expedient to expand Eq. (B.1) in powers of $\cos \phi$. This gives [158]

$$\begin{aligned} U_{\rm tor}(\phi) &= \frac{1}{2}k_1 + k_2 + \frac{1}{2}k_3 + \frac{1}{2}k_5 + k_6 \\ &+ \left(-\frac{1}{2}k_1 + \frac{3}{2}k_3 - \frac{5}{2}k_5 \right)\cos\phi \\ &+ \left(-k_2 + 4k_4 - 9k_6 \right)\cos^2\phi \\ &+ \left(-2k_3 + 10k_5 \right)\cos^3\phi \\ &+ \left(-4k_4 + 24k_6 \right)\cos^4\phi \\ &- 8k_5\cos^5\phi - 16k_6\cos^6\phi \;. \end{aligned}$$
(B.2)

The GROMACS code provides such a representation via the Ryckaert–Bellemans function [49] (which we write here in the form employed by LAMMPS ¹)

$$U_{\rm tor}(\phi) = \sum_{n=1}^{6} A_n(\cos \phi)^{n-1} . \tag{B.3}$$

¹https://docs.lammps.org/dihedral_nharmonic.html.

Table B.1: Coefficients A_n from Eq. (B.4) as obtained from the k_n of Ref. [157] and utilized in prior work with GROMACS [31, 35, 36, 47, 159]. A_n are given in units of kJ/mol as employed in GROMACS, while LAMMPS uses kcal/mol (1 kJ/mol = 4.184 kcal/mol).

Torsion angle	A_1	A_2	A_3	A_4	A_5	A_6
$\alpha \ (trans)$	1.61159	-14.50430	4.39523	16.15770	-1.33950	3.01387
α (cis)	4.85568	-1.50692	2.84643	-6.19521	-2.67899	12.72530
β	-13.64610	20.67850	1.42321	-23.35750	1.17206	-8.37189
db $(trans)$	100.46200	0.00000	-100.46200	0.00000	0.00000	0.00000
db (cis)	100.46200	0.00000	-100.46200	0.00000	0.00000	0.00000

Comparison of Eq. (B.3) with Eq. (B.2) allows to determine A_n in terms of k_n . However, since Eq. (B.3) stops at $\cos^5 \phi$, the implementation in GROMACS sets $k_6 = 0$ [31, 35, 36, 47, 159]. This choice gives

$$A_{1} = \frac{1}{2}k_{1} + k_{2} + \frac{1}{2}k_{3} + \frac{1}{2}k_{5}$$

$$A_{2} = -\frac{1}{2}k_{1} + \frac{3}{2}k_{3} - \frac{5}{2}k_{5}$$

$$A_{3} = -k_{2} + 4k_{4}$$

$$A_{4} = -2k_{3} + 10k_{5}$$

$$A_{5} = -4k_{4}$$

$$A_{6} = -8k_{5}.$$
(B.4)

The values for A_n resulting from k_n of Ref. [157] are given in Table B.1.

Since k_6 vanishes for the double bonds in the original model [26, 39, 157], the choice $k_6 = 0$ is insubstantial. Differences from the original model may occur for the other dihedrals where $k_6 \neq 0$. However, graphical comparison of Eq. (B.1) with Eqs. (B.3) and (B.4) reveals that these differences are negligible for the α trans and β bonds (not shown), while they are a bit larger for the α cis bond. Even in this worst case, differences remain small, as illustrated by Fig. B.1a. Therefore, Eqs. (B.3) and (B.4) represent an excellent approximation to the original model and we utilize the potential parameters of Table B.1 for β , α cis and α trans bonds.

When comparing the double bond potentials, we noticed, that in the recent works using GROMACS [31, 35, 36, 47, 159] there are different sets of parameters $\{A_n\}$ for the *cis* and the *trans* double bonds, which does not agree with the Table B.1. As seen from Table B.1, Eqs. (B.3) and (B.4) propose the same $U_{tor}(\phi) = A_1(1 - \cos \phi)$ for the double



Figure B.1: (a) Torsional potential for the α *cis* bond. The solid line shows Eqs. (B.3) and (B.4) with the parameters from Table B.1. The dashed line represents the original model, Eq. (B.1), with the parameters from Smith et al. [26, 39, 157]. (b) Torsional potential for the double bonds. The dashed line depicts Eqs. (B.3) and (B.4) with the parameters from Table B.1. The solid lines present the results from Eq. (B.3) with the coefficients A_n from Table 2.3 of the main text for the *cis* (db cis, red line) and *trans* (db trans, blue line) configuration of the monomers.

bond of the *cis* and *trans* monomer. This function is shown as a dashed (green) line in Fig. B.1b. The same $U_{\rm tor}(\phi)$ is used for the *cis* and *trans* monomer because the barrier at $\phi = 90^{\circ}, 270^{\circ}$ corresponds to about 12 000 K and is much higher than the temperatures of a few hundred Kelvin typically studied. In practice, this implies that a monomer initially prepared in a *cis* or *trans* state will remain in this state for all times of current simulations with length of a few microseconds. We believe that the authors of the works [31, 35, 36,47, 159 have preferred to distinguish between the *cis* and *trans* states by turning the potentials to having single minima at $\phi = 0^{\circ}$ (or 360°) and $\phi = 180^{\circ}$, rather than two. In this way, even if a possibility of a rare escape from the barrier occurs, the *cis* monomer will still remain *cis*, and the *trans* monomer will still remain *trans*. In practice, this was achieve by turning the barrier at $\phi = 90^{\circ}, 270^{\circ}$ into an inflection point. For the *cis* monomer the composite function is: $U_{\text{tor}}(\phi) = A_1(1 - \cos \phi)$ for $\phi \leq 90^\circ$, $U_{\text{tor}}(\phi) = A_1(1 + \cos \phi)$ for $90^{\circ} < \phi < 270^{\circ}$ and $U_{\rm tor}(\phi) = A_1(1 - \cos \phi)$ for $\phi \ge 270^{\circ}$. For the trans monomer the composite function is: $U_{\rm tor}(\phi) = A_1(1 + \cos \phi)$ for $\phi \le 90^\circ$, $U_{\rm tor}(\phi) = A_1(1 - \cos \phi)$ for $90^{\circ} < \phi < 270^{\circ}$ and $U_{\rm tor}(\phi) = A_1(1 + \cos \phi)$ for $\phi \ge 270^{\circ}$. In the program, it is simpler to fit these composite functions to Eq. (B.3). The resulting values for the coefficients A_n are given in Table 2.3 of the main text. The fit functions are shown as solid lines in Fig. B.1b; they are the same as in Fig. 2.4 of the main text.

Appendix C

PBD density references

See next page.

ID	Ref	Experimantal/ Simulation	Polybutadiene type	Molecular weight	Pressure (atm)	Notes
1	[36]	Simulation	1,4, 45% cis / 55% trans	$1.6 imes 10^3$	1 atm	The same as in our work
2	[68]	Simulation	1,4,100% cis	$2.6 imes 10^3$	1 atm	
3	[69]	Experimental	1,4, 40% cis / 50% trans / 10% vinyl	?	?	
4	[70]	Experimental	pure cis	13.5×10^3 ?	?	
5	[71]	Experimental	1,4,100% cis	?	1 atm	
6	[45]	Simulation	1,4, 40% cis / 50% trans / 10% vinyl	$1.6 imes 10^3$	1 atm	
7	[72]	Experimental	1.4, 9% cis / 20% trans / 71% vinyl	13×20^3	?	
8	[73]	Simulation	1.4, 40% cis / 53% trans / 7% vinyl	$7 imes 10^3$?	Single chain simulation
9	[74]	Experimental	1.4, 34% cis / 58% trans / 8% vinyl	1.42×10^3	?	
10	[75]	Experimental	1.4, 36% cis / 55% trans / 9% vinyl	$\sim 2 \times 10^5$	0 atm	
11	[76]	Experimental	100% cis	$\sim 2-3\times 10^5$	0 atm	

 Table C.1:
 Description of the reference data of density for 1,4-polybutadiene.

Appendix D

LAMMPS input scripts

Bulk system. To run: \$LAMMPS -in inputScript.lmp -var sID packageId

units	real
atom_style	full
pair_style	lj/gromacs 9.0 12.0
bond_style	harmonic
angle_style	cosine/squared
dihedral_style	nharmonic
read_data	init_conf.dat
<pre># write_restart</pre>	NVTequil.restart.*
special_bonds	lj/coul 0.0 0.0 0.0
pair_modify	tail no mix arithmetic
neighbor	2.0 bin
neigh_modify	delay 0 every 1 check yes
timestep	1
fix	f2 all nvt temp 353.0 353.0 1000.0
thermo	100
thermo_style	custom step temp pe epair ebond eangle edihed ke vol press
run_style	respa 2 4 bond 1 angle 1 dihedral 1 pair 2
dump	logdump all custom 10000000 conf_dump_NVT_log-\${sID}.dat id mol xu yu zu
compute	clog all logscale 200000 2
# weed = 36	
variable	varlog equal c_clog
run	0
dump_modify	logdump every v_varlog
dump	dCONF all custom 20000 conf_dump_NVT-\${sID}.dat id mol xu yu zu
run	1000000
write_restart	NVTequil.restart.*

Film system. To run: $\Lambda MMPS$ -in inputScript.lmp -var sID packageId

processors	* * 1			
units	real			
atom_style	full			
pair_style	lj/gromacs 9.0 12.0			
bond_style	harmonic			
angle_style	cosine/squared			
dihedral_style	nharmonic			
read_data	supported_conf.dat			
#read_restart	PBDW.restart.*			
special_bonds	lj/coul 0.0 0.0 0.0			
pair_modify	tail no mix arithmetic			
neighbor	2.0 bin			
neigh_modify	delay 0 every 1 check yes			
timestep	1			
group	walls type 3			
group	polymers type 1 2			
fix	f1 polymers nvt temp 225.0 225.0 1000.0			
thermo	100			
thermo_style	custom step c_f1_temp pe epair ebond eangle edihed ke vol press			
run_style	respa 2 4 bond 1 angle 1 dihedral 1 pair 2			
dump	logdump all custom 1000000 conf_dump_log-\${sID}.dat.gz id mol xu yu zu			
compute	clog all logscale 100000 2			
# weed = 34				
variable	varlog equal c_clog			
run	0			
dump_modify	logdump every v_varlog			
compute	c_bins polymers chunk/atom bin/1d z lower 0.1 units box			
variable	v2x atom vx*vx			
variable	v2y atom vy*vy			
variable	v2z atom vz*vz			
fix	f_chunk polymers ave/chunk 1 1000000 1000000 c_bins			
	v_v2x v_v2y v_v2z density/number density/mass temp			
	file chunk_analysis-\${sID}.dat			
dump	dCONF all custom 10000 conf_dump-\${sID}.dat.gz id mol xu yu zu			
run	1000000			
write restart	PBDW.restart.*			

Appendix E

Bond correlations and internal distances

The mean-square internal end-to-end distance and the bond correlation function are defined by \vec{r}

$$R_{\rm e}^2(s) = \langle (\vec{r}_{n+s} - \vec{r}_n)^2 \rangle \,, \quad P_1(s) = \frac{\langle \vec{\ell}_{n+s} \cdot \vec{\ell}_n \rangle}{l^2} \,, \tag{E.1}$$

where \vec{r}_n is the position vector of the *n*th united atom (n = 1, ..., N), $\vec{l}_n = \vec{r}_{n+1} - \vec{r}_n$ the bond vector and $\langle ... \rangle$ the average over all configurations and chains of the system and over all possible pairs n, m = n + s of a chain. A viable parameterization of MD results for $P_1(s)$ is

$$P_1(s) = \begin{cases} \alpha & \text{for } s = 1 \ , \\ A \exp(-s/\overline{s}) & \text{for } s > 1 \ , \end{cases}$$
(E.2)

where $\alpha = -\langle \cos \theta \rangle$ (> 0). Using

$$R_{\rm e}^2(s) = l^2 s + 2l^2 \sum_{k=1}^{s-1} (s-k) P_1(k) , \qquad (E.3)$$

one gets

$$R_{\rm e}^2(s) = l^2 s \left[C_{\infty} - \frac{2}{s} \left(\alpha - A\overline{\alpha} + A \frac{\overline{\alpha} - \overline{\alpha}^{s+1}}{(1 - \overline{\alpha})^2} \right) \right] , \qquad (E.4)$$

where $\overline{\alpha} = e^{-1/\overline{s}}$ (< 1) and C_{∞} is the characteristic ratio of the infinitely long chain,

$$C_{\infty} = \lim_{N \to \infty} \frac{R_{\rm e}^2(N-1)}{(N-1)l^2} = 1 + 2(\alpha - A\overline{\alpha}) + \frac{2A\overline{\alpha}}{1-\overline{\alpha}} . \tag{E.5}$$

The characteristic ratio determines three often discussed length scales of a polymer: the effective bond length $b_{\rm e}$ [1],

$$b_{\rm e} = \sqrt{C_{\infty}} l \;, \tag{E.6}$$

the Kuhn length [92]

$$l_{\rm K} = C_{\infty} l \;, \tag{E.7}$$

and the persistence length $l_{\rm p}^{-1}$,

$$l_{\rm p} = \frac{1}{2} \, (C_\infty + 1) l \; . \tag{E.8}$$

For the CRC, CRC2, CRC4 and FRC models the following tables compile the values of α obtained from the simulations, the fit results for $A, \bar{s}, \bar{\alpha}$, and the resulting predictions for $C_{\infty}, b_{\rm e}, l_{\rm K}$ and $l_{\rm p}$.

Table E.1: Results for the CRC model at different T for $\alpha = -\langle \cos \theta \rangle$, the fit parameters $A, \bar{s}, \bar{\alpha} = e^{-1/\bar{s}}$, and the predictions for C_{∞} , $b_{\rm e}$, $l_{\rm K}$ and $l_{\rm p}$ using $l^2 = 2.158$ Å².

$T(\mathbf{K})$	α	A	\overline{s}	$\overline{\alpha}$	C_{∞}	$b_{\rm e}$ (Å)	$l_{\rm K}$ (Å)	$l_{\rm p}$ (Å)
213	0.480	0.769	4.025	0.780	6.213	3.661	9.126	5.297
225	0.480	0.833	3.542	0.754	5.807	3.540	8.530	4.999
240	0.480	0.816	3.581	0.756	5.790	3.534	8.505	4.987
253	0.480	0.751	3.985	0.778	6.058	3.615	8.898	5.183
273	0.480	0.776	3.680	0.762	5.749	3.522	8.444	4.956
293	0.480	0.777	3.620	0.759	5.668	3.497	8.325	4.897
353	0.481	0.751	3.625	0.759	5.549	3.460	8.150	4.810

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The expression for $l_{\rm p}$ given in Eq. (E.8) follows from the definition $l_{\rm p} = l \sum_{s=0}^{\infty} P_1(s)$ [91] and the relation between $R_{\rm e}^2(N-1)$ and $P_1(s)$ according to Eq. (E.3).
$T(\mathbf{K})$	α	A	\overline{s}	$\overline{\alpha}$	C_{∞}	$b_{\rm e}$ (Å)	$l_{\rm K}$ (Å)	$l_{\rm p}$ (Å)
213	0.480	0.722	4.060	0.782	5.999	3.598	8.811	5.140
225	0.480	0.730	3.930	0.775	5.866	3.558	8.617	5.043
240	0.480	0.730	3.850	0.771	5.758	3.525	8.458	4.963
253	0.480	0.719	3.971	0.777	5.866	3.557	8.616	5.042
273	0.480	0.720	3.850	0.771	5.706	3.509	8.382	4.925
293	0.480	0.723	3.800	0.769	5.651	3.492	8.300	4.884
353	0.481	0.716	3.700	0.763	5.486	3.440	8.058	4.763

Table E.2: Results for the CRC2 model at different T for $\alpha = -\langle \cos \theta \rangle$, the fit parameters $A, \bar{s}, \bar{\alpha} = e^{-1/\bar{s}}$, and the predictions for C_{∞} , $b_{\rm e}$, $l_{\rm K}$ and $l_{\rm p}$ using $l^2 = 2.158$ Å².

Table E.3: Results for the CRC4 model at different T for $\alpha = -\langle \cos \theta \rangle$, the fit parameters $A, \bar{s}, \bar{\alpha} = e^{-1/\bar{s}}$, and the predictions for C_{∞} , $b_{\rm e}$, $l_{\rm K}$ and $l_{\rm p}$ using $l^2 = 2.158$ Å².

$T(\mathbf{K})$	α	A	\overline{s}	$\overline{\alpha}$	C_{∞}	$b_{\rm e}$ (Å)	$l_{\rm K}$ (Å)	$l_{\rm p}$ (Å)
213	0.479	0.697	4.125	0.785	5.945	3.581	8.732	5.100
225	0.479	0.699	4.095	0.783	5.918	3.573	8.693	5.081
240	0.480	0.695	4.075	0.782	5.868	3.558	8.619	5.044
253	0.480	0.701	3.955	0.777	5.745	3.521	8.438	4.954
273	0.475	0.697	3.945	0.776	5.699	3.506	8.371	4.920
293	0.480	0.704	3.840	0.771	5.610	3.479	8.240	4.854
353	0.481	0.695	3.815	0.769	5.530	3.454	8.122	4.795

Table E.4: Results for the FRC model at different T for $\alpha = -\langle \cos \theta \rangle$, the fit parameters $A, \bar{s}, \bar{\alpha} = e^{-1/\bar{s}}$, and the predictions for C_{∞} , $b_{\rm e}$, $l_{\rm K}$ and $l_{\rm p}$ using $l^2 = 2.158$ Å².

$T(\mathbf{K})$	α	A	\overline{s}	$\overline{\alpha}$	C_{∞}	$b_{\rm e}$ (Å)	$l_{\rm K}$ (Å)	$l_{\rm p}$ (Å)
213	0.480	0.692	4.035	0.780	5.800	3.537	8.519	4.994
225	0.480	0.693	3.975	0.778	5.726	3.515	8.410	4.939
240	0.480	0.695	3.925	0.775	5.674	3.499	8.334	4.902
253	0.480	0.699	3.848	0.771	5.592	3.474	8.214	4.842
273	0.480	0.696	3.825	0.770	5.548	3.460	8.150	4.809
293	0.481	0.699	3.800	0.769	5.532	3.455	8.125	4.797
353	0.482	0.699	3.675	0.762	5.370	3.404	7.887	4.678

Appendix F

Partial structure factor fits

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	$T\left(\mathbf{K}\right)$	$\rho v_{\rm A,0}$	$A_{v,\mathrm{A}}$	$B_{v,\mathrm{A}}$	$ ho v_{\mathrm{B},0}$	$A_{\rm cc}$	$B_{\rm cc}$	$k_{\rm B}T\rho\kappa_T$	A_{κ}	B_{κ}
	213	1.415	0.154	-0.132	0.585	0.034	0.077	0.084	-0.079	0.356
_	225	1.452	0.122	-0.133	0.548	0.050	0.004	0.078	0.038	0.093
-	240	1.480	0.100	-0.134	0.520	0.037	0.055	0.086	0.080	0.049
-	253	1.433	0.201	-0.183	0.567	0.040	0.050	0.101	-0.008	0.275
-	273	1.500	0.116	-0.160	0.500	0.041	0.045	0.114	0.017	0.211
_	293	1.515	0.120	-0.172	0.485	0.045	0.037	0.130	0.023	0.213
-	353	1.625	0.125	-0.229	0.375	0.043	0.051	0.188	0.002	0.325

Table F.1: The values of the fit parameters for the partial structure factors in case of the CRC model (Eqs. (3.35) to (3.37)). The values $A_{v,B} = -A_{v,A}$ and $B_{v,B} = -B_{v,A}$



Mobilité interfaciale dans les films polymères vitrifiables : est-elle déterminée par le mouvement collectif ou l'énergie intramoleculaire ?

Résumé

L'influence des barrières énergétiques de torsion intramoléculaires sur la structure et la dynamique du copolymère aléatoire cis-trans du 1,4-polybutadiène (PBD) a été étudiée au moyen de simulations classiques de dynamique moléculaire. Le PBD en phase volumique et un film de ~10 nm d'épaisseur supporté par une paroi de graphite ont été simulés à l'aide d'un modèle chimiquement réaliste d'atomes unifiés, qui comporte des potentiels de torsion (dièdre). Les simulations ont également été effectuées avec des potentiels dièdres réduits, ainsi que sans potentiel dièdre. Il est démontré que la présence des dièdres renforce l'effet de cage dans la masse, ce qui ralentit la dynamique de plusieurs ordres de grandeur, déplaçant ainsi la température de transition vitreuse vers des valeurs plus élevées par rapport au modèle sans dièdres. En revanche, la densité et la structure du PBD en phase volumique ne sont pratiquement pas affectées par les dièdres. De la même manière, les dièdres déterminent l'ordre de grandeur de la dynamique dans les films. Leur désactivation n'entraîne pas de changements notables du profil de densité des films, non seulement dans la région centrale, semblable au volume, mais aussi aux interfaces. La dynamique a été étudiée par une analyse à résolution de couche des déplacements carrés moyens (MSD) et de l'autocorrélation du moment dipolaire des monomères (relaxation diélectrique). L'extraction des temps de relaxation structurelle a montré que le confinement imposé par la paroi de graphite et la surface libre est le facteur dominant dans la modification des gradients de mobilité interfaciale par rapport à l'ordre de grandeur global de la dynamique fixé par les barrières de torsion à travers tout le film PBD. Mots clés : films polymères supportés, polybutadiène, simulations de dynamique

Mots clés : films polymères supportés, polybutadiène, simulations de dynamique moléculaire, transition vitreuse, relaxation diélectrique, gradients de relaxation, barrières intramoléculaires.

Résumé en anglais

The influence of the intramolecular torsional energetic barriers on the structure and dynamics of 1,4-polybutadiene (PBD) *cis-trans* random copolymer was studied by means of classical molecular dynamics simulations. Bulk PBD and a ~10 nm thick film supported on a graphite wall were simulated using a chemically realistic united-atom model, which has torsional (dihedral) potentials associated to each bond in the polymer chain. The simulations were also performed with reduced, as well as without dihedral potentials. It is demonstrated that the presence of the dihedrals enhances the cage effect in the bulk, which slows down the dynamics by several orders of magnitude, thus shifting the glass transition temperature to higher values with respect to the model without the dihedrals. Meanwhile the density and the structure of bulk PBD are essentially unaffected by dihedrals. In the same way, the dihedrals determine the order of magnitude of the dynamics in the films. Disabling them does not lead to noticeable changes of the density profile of the films not only in the middle, bulk-like region, but also at the interfaces. The dynamics was studied by layer-resolved analysis of the mean-square displacements (MSD) of united atoms and the monomer dipole moment autocorrelation (dielectric relaxation). Extracting the structural relaxation times showed that the confinement imposed by the graphite wall and the free surface are the dominating factor in modifying the interfacial mobility gradients with respect to the overall order of magnitude of the dynamics set by the torsional barriers across the whole PBD film. **Keywords**: supported polymer films, polybutadiene, molecular dynamics simulations, glass transition, dielectric relaxation, relaxation gradients, intramolecular barriers.