Microscopic Description of Entanglements in Polyethylene Networks and Melts: Strong, Weak, Pairwise, and Collective Attributes

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Abstract:

We present atomistic molecular dynamics simulations of two Polyethylene systems where all entanglements are trapped: a perfect network, and a melt with grafted chain ends. We examine microscopically at what level topological constraints can be considered as a collective entanglement effect, as in tube model theories, or as certain pairwise uncrossability interactions, as in slip-link models. A pairwise parameter, which varies between these limiting cases, shows that, for the systems studied, the character of the entanglement environment is more pairwise than collective. We employ a novel methodology, which analyzes entanglement constraints into a complete set of pairwise interactions, similar to slip links. Entanglement confinement is assembled by a plethora of links, with a spectrum of confinement strengths, from strong to weak. The strength of interactions is quantified through a link ‘persistence’, which is the fraction of time for which the links are active. By weighting links according to their strength, we show that confinement is imposed mainly by the strong ones, and that the weak, trapped, uncrossability interactions cannot contribute to the low frequency modulus of an elastomer, or the plateau modulus of a melt. A self-consistent scheme for mapping topological constraints to specific, strong binary links, according to a given entanglement density, is proposed and validated. Our results demonstrate that slip links can be viewed as the strongest pairwise interactions of a collective entanglement environment. The methodology developed provides a basis for bridging the gap between atomistic simulations and mesoscopic slip link models.

1 Introduction

In the molecular description of structure-property relations of polymer melts and networks, one of the fundamental concepts is chain entanglement. When macromolecules interpenetrate, the term entanglements intends to describe interactions resulting from the an-

crossability of chains. In flexible polymer melts and rubbers these interactions alter in a universal manner their dynamical, flow, and deformation properties.

Molecular-based theories which aim to incorporate entanglements have first to define them. This task poses certain difficulties. In mathematical terms, entanglement interactions originate from topological constraints (TCs). Thus, in a many-chain theory, and in order to derive entanglement effects from ‘first principles’, a rigorous description invokes the use of topological invariants and certain approximations to make the problem tractable. In such theories, however, the range of problems that can be studied is limited by mathematical complexity.

In order to deal with entangled matter, several conceptual simplifications were introduced by Edwards,8,11–15 de Gennes,6,7 Doi and Edwards,8,9 and other pioneers in the field of polymer physics. The crucial simplification was the description of entanglement through a mean field and a single chain model. In tube model theories8,7,9,10 it is postulated that the entanglements generate a confining, (quadratic) mean field potential, which restricts lateral monomer fluctuations to a tube-like region surrounding each chain. In a perfect network, the potential attracts the monomers toward the centerline of the tube and chains are restricted to adopt conformations which are compatible with the tube constraint. In polymer melts, this confinement is not permanent, but leads to reptation,9 a one-dimensional diffusion of the chain along the tube.

The axis of the tube plays the role of a coarse-grained representation of the real chain, and it is called the primitive path (PP). It has a random-walk like conformation, like the real chain, but with a larger step length, which corresponds to the entanglement molar mass, M_e. The latter is determined from the plateau modulus of long chain polymer melts. The average tube diameter is not so well-defined; it is considered to be of the order of magnitude of the PP step length; it is often assumed that the two lengths are equal.

An alternative, discrete, localized version of the tube constraint is utilized in models employing slip-links,8,11–15 The tube is replaced by a set of slip links along the chain, which restrict lateral motion but permit chain sliding through them. The real chain is represented by the corresponding PP, which is a series of strands of average molar mass M_e connecting the links. The tube model provides a basis for working out analytically the linear and nonlinear viscoelasticity of polymer melts9,16–20 and networks10,15,21–24 at a molecular level. Slip link models19,20,24 also allow for analytic8,11–13,15,25,26 or stochastic simulation14,27–37 treatments with a molecular representation, at the single or many-chain level.

The concepts conveyed by the tube and slip-link models provide the prevailing microscopic view of entanglement. In stochastic simulations, the slip links are usually envisaged as binary entanglements, a form of discrete pairwise contacts which confine (link) the motion of two neighboring chains. The tube constraint, on the other hand, acts along the whole chain; locally, it is thought that confinement is applied collectively by many overlapping surrounding chains, and not by specific pairwise interactions. Any gain or
loss in the local pairwise interaction density does not necessarily affect the tube diameter or length, as it does the local entanglement density of slip-link models \(^1\) (see p 552 in ref 20).

A microscopic picture of entanglement is also suggested by scaling models relating the degree of entanglement to chemical structure. Notably, the most successful of them are based on similar ideas. A good correlation with polymer melt data is provided by the packing length \(^2\) model, which is based on the conjecture of Lin, \(^3\) and Kavassalis and Noolandi. \(^4,5\) The conjecture is in support of a collective \(^4,6\) argument: an entanglement results from a fixed number of polymer chains cohabiting the same volume. On the other hand, the scaling model \(^58,59\) of Colby and Rubinstein, which is in agreement with data from solutions in a theta solvent, is in support of a pairwise \(^10,34\) argument; a fixed number of binary contacts between chains gives rise to an entanglement strand.

Despite the success of tube and slip-link models in describing the linear (especially) and nonlinear rheology of polymer melts, as well as rubber elasticity, the nature of entanglements remains elusive. It would be desirable, therefore, to probe entanglements microscopically in order to correct, refine and support the perspective given above with microscopic information. Simulation is probably the method of choice, since we need access to some kind of topological information which is not directly accessible with spectroscopic methods.

The question, then, is how to define entanglements in a set of densely packed chain molecules. A unique definition might not exist, but a certain microscopic description has to be invoked to address the above topics. In this direction, several attempts have been made. Assuming that entanglements lead to persistent chain interactions, the detection of long-lived chain contacts \(^48–51\) was the target of many simulations. Although it was found that such contacts do exist, it is rather difficult to evolve this picture further. Additionally, the topological origin of entanglements is overlooked. It becomes prominent, however, in studies \(^52–57\) employing topological invariants. The latter can detect pairs of entangled and unentangled chains, and the ‘amount’ of winding between the chains. We have to note that, for open chains, true invariants do not exist. Moreover, the simplest invariant for closed chains, the Gauss linking number, can misclassify \(^2\) topological constraints. In any case, the provided information is at the global level of a chain. Though very useful, it is not clear how to put it into the context of current tube and slip-link model ideas. The latter require a somewhat localized description of entanglement, especially in connection with constraint release (CR) processes. In this respect, the use of a local topological measure may be more suitable.

Microscopic investigations of tube model ideas have greatly benefited from simulations which construct the PPs according to the Edwards definition, \(^5\) i.e., by holding chain ends fixed and removing all chain slack from the system. The first implementation \(^58,59\) was by Everaers et al., but soon thereafter other methods for shrinking the chains in continuous \(^80,61\) and lattice models \(^62,63\) appeared. There is strong evidence \(^61,64–68\) that the PP Kuhn segment coming out from these methods is in very good agreement with \(M_e\) estimations from plateau modulus measurements. The PP representation is also popular in slip-link simulations. Thus, it is hoped that these coarse graining methods could serve as a bridge \(^69,70\) between the different levels of description.

The properties of PPs have been studied in several systems, such as rubbers, \(^71\) melts, nanocomposites, \(^72–76\) grafted brushes, \(^77\) glasses, \(^78\) in equilibrium or under deformation \(^75\) and flow. \(^79\) Concepts and properties such as entanglement density scaling \(^58,59,80\), entanglement molecular weight \(^58,61,64–67,81\), CLF potential \(^62,81\), tube potential \(^82\), dilution exponent \(^63\), onset of entanglements \(^83,84\) and even the literal picture of a tube \(^65–89\) enclosing a chain and the associated tube survival probability, have been investigated. Despite the available microscopic information, the many chain nature of entanglement remains unclear. A possible reason is that most of the simulations follow tube model ideas and examine, with the help of PPs, single chain properties. Nevertheless, it is the many chain nature of entanglement which is elusive and is replaced by a mean field. Therefore, rather than examining single chain properties, which are the result of tube confinement, we can try to analyze the chain environment that gives rise to confinement. This is the strategy followed here.

In order to make the problem tractable, we have only examined systems with trapped entanglements; a polyethylene (PE) melt where all chain ends are grafted to other chains, and a perfect PE network formed by connecting cross-links initially situated at the sites of a diamond lattice. In this way, our results are free of complications related with CR processes. The latter can be addressed after a suitable microscopic description of entanglement has been given. In the overview of the paper given below, the terms tube and tube constraint are used as generic terms for entanglement constraints, and are not specifically related to tube model ideas.

As probes of the entanglement environment, we have used the PPs themselves. Exploiting the fact that they sample the uncrossability constraints of the surroundings, we show that, by mapping short molecular dynamics (MD) trajectories to PP trajectories, it is possible to detect all the pairwise chain interactions assembling the tube constraint. To verify that these interactions result from true topological constraints, we ensure that they satisfy a local topological criterion. The latter is based on the common idea of two concatenated rings that define a binary link. Therefore, entanglement constraints are decomposed into a large set of local links, which is more or less complete. Similar ideas have been explored, analytically, by Iwata and Edwards \(^90,91\).

By construction (chain shrinking), all these links are sampled inside the pervaded volume of a chain. The mechanism by which link sampling takes place is clarified in detail. While some of those links are continuously present, many of them are intermittent or ‘blinking’ links. They are not sampled continuously over a specific observation time. Less constraining (weak) links are sampled less frequently. This leads to a distribution of linking times, and thus the links assembling the tube constraint have a spectrum of confinement strengths, with maxima at the strong \(^92\) and weak part of the spectrum.

From this picture, we then develop a simple statistical model and we introduce a pairwise parameter, that can quantify the pairwise versus collective character of mean-field type links representing many uncrossability interactions. With the help of these tools we show quantitatively that:

- the entanglement environment consists of a plethora of pairwise interactions, so that it is inherently collective;
- there exist strong and weak pairwise uncrossability constraints;
- the majority of pairwise interactions are weak and confinement is mainly enforced by the strongest ones;
- weak topological interactions can be fully relaxed by stored length fluctuations, and they do not contribute to the low frequency storage modulus of an elastomer, or the plateau modulus of a melt;
- the average effect of the collective entanglement environment is to confine a chain through a much smaller number of mean-field type links, which have a dominant pairwise character;
- the pairwise parameter is chain length and entanglement density independent.

In order to be able to map the entanglement environment to specific pairwise interactions, we present a procedure for selecting the strongest links according to a given link density. The procedure...
introduces an auxiliary time scale which is self-consistently determined to be approximately one $\tau_e$. It is shown that strong links have the property that they are sampled at least once within this time interval. By following the space-time trajectories of selected links, under equilibrium, deformation, or flow, we could obtain useful mesoscopic information for the development of hierarchical slip-link, or single particle$^{93-95}$ simulation schemes. The mean square displacement of selected links is presented in the appendices.

The PPs employed in our study have been generated by the CReTA algorithm,$^{61}$ which applies a contour reduction (chain shrinking) scheme. Different schemes, based on time averaging,$^{96,97}$ have been proposed. However, the presented results are not based on specific properties of the PPs, except that they probe pairwise uncrossability constraints in the pervaded volume of a chain. Thus, our conclusions are fairly general and independent of the particular scheme employed for obtaining the PPs.

In order to make the article more readable and focus on the analysis of entanglement constraints, many of our results are presented in the Appendices. Our microscopic description is also supported by several explanatory videos$^{98}$ provided with the Supporting Information for this article.

2 Systems Studied

The systems analyzed are a PE network "rubber" and an end-grafted PE melt (see next section). The rubber system contains 64 tetrafunctional cross-links and 128 chains. The cross-links are initially situated at the sites of a diamond lattice and all chains are attached to cross-links at both ends. Thus, the network is perfect (absence of pendant chains, chain loops, etc.). Each chain consists of 201 united atoms, not counting the cross-links at its two ends.

The melt system consists of 16 linear chains of 500 united atoms each. In order to examine the chain length dependence of some of our results, we also analyzed a melt with eight chains and double the chain length, i.e., of 1000 united atoms. The entanglement density of these melt configurations is approximately the same. Whenever required, these systems will be denoted by (C500) and (C1000), but in general, our analysis will focus on the (C500) system.

The simulations were performed in the NPT ensemble, at temperature 450 K, pressure 1 Atm, by employing the united atom TraPPE$^{99,100}$ force field and 3D periodic boundary conditions. United atom beads correspond to CH$^2$ units, and carbon atoms, respectively. The rubber was equilibrated by the gradual push-off$^{101}$ method, discussed in Appendix I. The melt was equilibrated by standard connectivity altering$^{102,103}$ Monte Carlo algorithms. The shortest relaxation time related with the tube constraint, $\tau_c$, is 1.3 ns for the rubber, and 1.7 ns for the melt (C500). It was determined from the first crossover time in the mean square displacement (msd) of midchain monomers (Appendix II). The longest relaxation time of each system, $\tau_{ee}$, is determined from the msd crossover to an invariant regime (absence of long-range diffusion), and is 9.8 ns for the rubber and 11.5 ns for the melt (C500). The corresponding crossover times for the (C1000) melt are 1.6 and 19.4 ns. Details about the dynamics and the equilibration of the systems will appear elsewhere.$^{104,105}$

Both systems share, approximately, the same density,$^{106,107}$ $\rho \approx 0.78$ g/cm$^3$, and characteristic ratio, $C_v$. The latter is 8.5 for the rubber and 8 for the (C500) melt. Volumetric, structural,$^{102,103}$ and entanglement$^{61}$ properties of the melt system have been presented elsewhere. They are in very good agreement with corresponding experimental estimates. The rubber system is structurally similar to the melt, except for entanglement properties. Its $M_e$ was estimated$^{61}$ from the Kuhn segment of the PPs, and is about half that of the PE melt. The large entanglement density of the rubber is also reflected$^{104}$ in a comparatively high, entanglement-dominated, shear modulus for an elastomer, of 5.9 MPa. As is discussed in Appendix I, it is due to the short length of the precursor chains and the artificial preparation conditions of the network.

3 Methodology

For both systems, we generated a molecular dynamics (MD) trajectory, which is of length 35 $\tau_e$ for the rubber and approximately 50 $\tau_e$ for the melts, covering a few times the longest relaxation time of each system. Our purpose is to use the trajectory as an ensemble of states with invariant topology, representative of the entanglement constraints in the system. This means that under thermal motion all entanglements should remain trapped, a condition satisfied in the perfect network but not in the melt, due to the free motion of chain ends.

The melt topology can become invariant by introducing additional topological constraints. Our choice is to graft (bond) all chain ends to their nearest united atom bead along a different nearby chain. Thus, the melt system will be referred to as end-grafted melt (EG-melt). Effectively, the EG-melt becomes a perfect trifunctional network with as many cross-links as chain ends. The initial chains are grafted at both ends and have on average two randomly placed cross-links along their length (from the grafted ends of other chains).

In a MD simulation of this network, the conformations sampled by grafted chains are restricted by the same topological constraints present in the parent melt state, plus the constraints due to grafting. From such a network trajectory we store chain conformations by excluding all bonds inserted to impose grafting, i.e., the network trajectory is mapped to a melt system, with virtually free chain ends. Thus, starting from a melt configuration with a specific topology, we can generate thermally fluctuating chain conformations with approximately the same topology, by eliminating CR through grafting.

Our results for the EG-melt refer to this ‘perturbed’ MD trajectory. The perturbation preserves the average PP contour length, but it alters contour length fluctuations. Because of the presence of additional constraints, the grafted chains cannot sample certain Rouse modes with long length scales. However, the short and medium length scale modes, which are responsible for the sampling of the enclosing tube, are adequately sampled. In this respect, at short times, the dynamics of grafted chains are similar to those of melt chains with free chain ends. For example, at short times, the mean square displacement (see Appendix II) scales as $t^{0.5}$ and then displays a crossover to a regime scaling as $t^{0.28}$, a crossover indicative of tube confinement.

4 Reduction to Networks of Local Links

The generated MD trajectories were subjected to topological analysis, frame by frame, by using the CReTA$^{61}$ (Contour Reduction Topological Analysis) algorithm. CReTA reduces a dense system of polymer chains to the corresponding system of primitive paths (PPs), constructed as the shortest paths$^9$ under the same ‘topological constraints’ (TCs) as the original chains. By fixing chain ends in space and by prohibiting chain crossing, the contour lengths of all chains are simultaneously minimized (shrunk), until they become piecewise linear objects coming together at the nodal points of a network. During contour reduction, chain thickness is progres-
of ‘tightened’ entangled curves, consisting of consecutive beads, as semblance to binary slip links. The details of mapping a system can also be envisaged as local links that have a physical re-
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fused beads

sarily reduced. Finally, the PPs become very thin objects consist-
ing of fused beads which can be also mapped to the initial chain monomers. Explanatory videos and other details can be found elsewhere.61,98

Network nodes are placed where PPs are mutually blocked (en-
tangled) and contour reduction can no longer proceed (see network pictures in Supporting Information). They represent effective spatial localization points of the TCs, that each chain is subjected to. They can also be envisaged as local links that have a physical resemblance to binary slip links. The details of mapping a system of ‘tightened’ entangled curves, consisting of consecutive beads, as our system of PPs here, to a network with point-like nodes and rectilinear edges, will be described elsewhere.105 It will be shown that the mapping is very stable with respect to changes in the free parameters involved. A short account of the mapping is given in Figure 1, and in Appendix III.

5 Results and Discussion

5.1 Fluctuations in the Number of Local Links and Linked Chains (LCs)

Here, we examine the tube constraint at the level of the surrounding chains that create it. To this end, we define as $z_{\alpha}(t)$ the number of chains linked, instantaneously, to a specific chain $\alpha$, at a specific time $t$. As shown in Figure 2, all chains in this set are different. That is, if at time $t$ chain $\alpha$ is linked twice or more with chain $\beta$, at different places along their contours, then $\beta$ counts as one chain in $z_{\alpha}(t)$, and vice versa. We will refer to this time-dependent set as the mate chains, or Linked Chains (LCs) of $\alpha$. If, due to periodic boundary conditions, $\alpha$ is also linked with different ‘images’ of $\beta$ or of itself, then these images108 are considered as different chains in $z_{\alpha}(t)$. Entanglements or knots of $\alpha$ with itself (same image) are not considered. For the chain lengths studied here their occurrence is negligible.59,61 We also note that $z_{\alpha}(t)$ characterizes a chain’s entanglement environment by keeping track of the identities of linked chains at time $t$, in a form suitable for comparisons at different times.

In Appendix IV (Figure 16), we show that, when two chains are linked, the probability that they are coupled with more than one link is very small (less than 10%). Hence, the simplification of dealing with linked chains instead of individual links is justified. This choice has been made in order to facilitate the presentation and the interpretation of our results. We could as well work with individual links. Therefore, in the following sections we will use interchangeably the terms local links and linked chains for $z_{\alpha}(t)$, though in our calculations ($z_{\alpha}(t)$) refers to LCs and not individual links.

In Figure 3a we plot the distribution of $z_{\alpha}(t)$ reduced by its equilibrium time average $\langle z_{\alpha}(t) \rangle$. In contrast to what one would expect for a system with solely trapped entanglements, $z_{\alpha}(t)$ fluctuates around a time average value. In a corresponding slip-link system,
for long chains. Also see that the amplitude of these fluctuations seems to be smaller this issue is discussed in more detail. In the inset to Figure 3a, we restrict continuously two fluctuating PPs and exhibit a permanent character. As a result, the instantaneous number of links sampled by a PP, \(z_\alpha(t)\), fluctuates around an average value. In Appendices V and VI, and in the videos provided with the Supporting Information, this issue is discussed in more detail. In the inset to Figure 3a, we also see that the amplitude of these fluctuations seems to be smaller for long chains.

The chain average of \(z_\alpha(t)\) is defined as

\[
z(t) = \frac{1}{N_{ch}} \sum_{\alpha=1}^{N_{ch}} z_\alpha(t) ,
\]

where \(N_{ch}\) is the number of chains. In Figure 3b it is plotted as a function of reduced time, \(t/\tau_e\). We see that for both systems it is approximately constant. Its time average, \(\langle z(t) \rangle\), is 9.5 for the rubber, and 14.6, 25.3 for the EG-melt (C500) and (C1000), respectively. These values lead to a rubber entanglement density which is approximately twice that of the EG-melt.

The fluctuations around \(\langle z(t) \rangle\) are much smaller for the rubber because chain averaging is more effective in smoothing individual chain fluctuations. The rubber consists of eight (sixteen) times more chains than the EG-melt (C500) and (C1000) systems, respectively. In a very large system, \(z(t)\) would appear as a thin horizontal line, i.e., it would remain constant over time. Thus, despite the fluctuations in the number of links of individual chains, when viewing the system as a single chain (through \(\langle z(t) \rangle\)), we observe the behavior expected of a slip-link system.

### 5.2 Cumulative Number of Sampled links: A Complete Set

Because of the absence of CR, after a certain time the PP cannot sample any new links in its surroundings. This fact is exploited in order to gather all sampled links into a complete set of pairwise un-crossability interactions characterizing the entanglement environment.

To elucidate these ideas further, we introduce \(U_\alpha(t)\), the cumulative number of different chains which have been linked to a given chain, \(\alpha\), up to time \(t\). All chains in \(U_\alpha(t)\) are different. They were not necessarily linked to \(\alpha\) at all times between 0 and \(t\). Some of them may have been linked for a long (discontinuous or continuous) subinterval, some of them for just one instant, or for the whole time \(t\). As shown in Figure 2, \(U_\alpha(t+\Delta t)\) is larger than \(U_\alpha(t)\) by the number of mate chains which link to \(\alpha\), for the first time, at any instant between \(t\) and \(t+\Delta t\).

An informative graph of \(U_\alpha(t)\), together with individual ‘link trajectories’, is shown in Figure 4. In this figure, the systems are observed at equal time instants, spaced by 0.01 ns, over a total observation time (OT), \(T\), which covers a few times the longest relaxation time of each system. Specifically, \(T \approx 35 \tau_e\) for the rubber, \(T \approx 50 \tau_e\) for the EG-melts.

We see that some links ‘blinking’. They are not present during the entire OT. From the number of points in a trajectory we can estimate the corresponding linking time (LT), \(\tau\), i.e., the total time during which a particular link was active. In order to analyze the plethora of links we introduce the link persistence, \(\tilde{\tau} = \tau/T\), which...

\(\text{Figure 3.}\) (a) Distribution of \(z_\alpha(t)\) reduced by the corresponding time average \(\langle z_\alpha(t) \rangle\) (see text). Log-log coordinates in the inset. To obtain these results, a separate distribution is constructed for each chain first and then an average is taken over the distributions of all chains. This is done because the entanglement environment (number of entanglements) and related fluctuations may be different for each chain. Thus, we consider each chain as a subsystem, we analyze the fluctuations within each subsystem, and then we average over the subsystems. This is necessary in order to map the fluctuations to a single chain model. (b) Average number of links per chain at time \(t\).
of each link, with values $\alpha$ equal to time $t$. Here, it enumerates with unique integer indices, as those denoted by $m$ lines, continuous or segmented, correspond to individual 'link trajectories', to $\tau$, of reduced time, the total plotted length of the corresponding trajectory. The reduced linking formation, intermittent link trajectories with $\tilde{t}$, time, $\tilde{t}$, of reduced time, $\tilde{t}$ is a stepwise, increasing, integer valued function, shown by a solid line. $U(A(T))$ is a stepwise, increasing, integer valued function, shown by a solid line. It enumerates with unique integer indices, $i$, cumulatively, the links sampled by chain $\alpha$, up to time $t$. When a link, $i$, is found to be paired to $\alpha$ at time $t$, a point with coordinates $(i,t)$ is plotted. Therefore, horizontal lines originating at $\tilde{t}$, of the LCs sampled at $t = 0$, appear as $\tilde{t} \geq 0$. Note that, by construction, the number of points in a $t$-column is $\alpha(t)$. A subset of the links appears to be permanent, with $\tau \simeq 1$. Most of the LCs sampled at $t = 0$ have this property. Instantaneous sampling at any $t$ (a $t$-column in Figure 4), will have a large overlap with this subset, which are basically strong links. This is evident in individual $t$-columns, where most of the points are located on horizontal lines originating at $t = 0$. Note that, by construction, the number of points in a $t$-column is $\alpha(t)$. A striking feature of Figure 4 is that the majority of the $U_A(T)$ links have small, even vanishing $\tilde{t}$'s. Moreover, the number of links sampled over the whole OT, $U_A(T)$, is much larger than those sampled instantaneously, $\alpha(t)$, at any time. Furthermore, after a certain time, $\tilde{t}'$, most of the links with appreciable $\tilde{t}$ have been sampled. The links sampled for the first time at later times, $\tilde{t} > \tilde{t}'$, appear to have a $\tau \simeq 0$. The sampling mechanism that generates blinking links is relatively simple, and is demonstrated in Figure 5. The driving force for this mechanism are stored length fluctuations, as described in detail in Appendix V. In order to avoid confusion with constraint release (CR) mechanisms driven by chain end motion, we call this mechanism lateral link sampling (LLS). However, upon nonlinear deformation or flow, LLS could possibly act as a CR mechanism. A similar mechanism was implemented by Padding and Briels to enforce chain uncrussability in a coarse grained model of polyethylene. The chain average of $U_A(t)$ is defined as

$$U(t) = \frac{1}{N_{ch}} \sum_{\alpha=1}^{N_{ch}} U_A(t)$$

and is shown in Figure 6. It is estimated by averaging over multiple time origins, so that by construction $U(0) = \alpha(0)$ (since $U_A(0) = \alpha(0)$). We see that $U(t)$ increases rapidly and approaches a plateau which is much larger than the average number of links $\langle \alpha(t) \rangle$ per chain (see Figure 3b). This is more evident in the EG-melt (C500); the longer chains and the wider tube of this system is basically the reduced linking time of a link. As the observation time increases, $\tau$ becomes proportional to $T$, with a different prefactor for each link (see Supporting Information). Thus, for adequately long OTs, as the ones utilized here, the $T$'s are equilibrium $T$-independent quantities that characterize the ‘persistence’ of link trajectories with values $0 < \tau \leq 1$. A subset of the links appears to be permanent, with $\tau \simeq 1$. Most of the LCs sampled at $t = 0$ have this property. Instantaneous sampling at any $t$ (a $t$-column in Figure 4), will have a large overlap with this subset, which are basically strong links. This is evident in individual $t$-columns, where most of the points are located on horizontal lines originating at $t = 0$. Note that, by construction, the number of points in a $t$-column is $\alpha(t)$. A striking feature of Figure 4 is that the majority of the $U_A(T)$ links have small, even vanishing $\tilde{t}$'s. Moreover, the number of links sampled over the whole OT, $U_A(T)$, is much larger than those sampled instantaneously, $\alpha(t)$, at any time. Furthermore, after a certain time, $\tilde{t}'$, most of the links with appreciable $\tilde{t}$ have been sampled. The links sampled for the first time at later times, $\tilde{t} > \tilde{t}'$, appear to have a $\tau \simeq 0$. The sampling mechanism that generates blinking links is relatively simple, and is demonstrated in Figure 5. The driving force for this mechanism are stored length fluctuations, as described in detail in Appendix V. In order to avoid confusion with constraint release (CR) mechanisms driven by chain end motion, we call this mechanism lateral link sampling (LLS). However, upon nonlinear deformation or flow, LLS could possibly act as a CR mechanism. A similar mechanism was implemented by Padding and Briels to enforce chain uncrussability in a coarse grained model of polyethylene. The chain average of $U_A(t)$ is defined as

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and is shown in Figure 6. It is estimated by averaging over multiple time origins, so that by construction $U(0) = \alpha(0)$ (since $U_A(0) = \alpha(0)$). We see that $U(t)$ increases rapidly and approaches a plateau which is much larger than the average number of links $\langle \alpha(t) \rangle$ per chain (see Figure 3b). This is more evident in the EG-melt (C500); the longer chains and the wider tube of this system is basically the reduced linking time of a link. As the observation time increases, $\tau$ becomes proportional to $T$, with a different prefactor for each link (see Supporting Information). Thus, for adequately long OTs, as the ones utilized here, the $T$'s are equilibrium $T$-independent quantities that characterize the ‘persistence’ of link trajectories with values $0 < \tau \leq 1$. A subset of the links appears to be permanent, with $\tau \simeq 1$. Most of the LCs sampled at $t = 0$ have this property. Instantaneous sampling at any $t$ (a $t$-column in Figure 4), will have a large overlap with this subset, which are basically strong links. This is evident in individual $t$-columns, where most of the points are located on horizontal lines originating at $t = 0$. Note that, by construction, the number of points in a $t$-column is $\alpha(t)$. A striking feature of Figure 4 is that the majority of the $U_A(T)$ links have small, even vanishing $\tilde{t}$'s. Moreover, the number of links sampled over the whole OT, $U_A(T)$, is much larger than those sampled instantaneously, $\alpha(t)$, at any time. Furthermore, after a certain time, $\tilde{t}'$, most of the links with appreciable $\tilde{t}$ have been sampled. The links sampled for the first time at later times, $\tilde{t} > \tilde{t}'$, appear to have a $\tau \simeq 0$. The sampling mechanism that generates blinking links is relatively simple, and is demonstrated in Figure 5. The driving force for this mechanism are stored length fluctuations, as described in detail in Appendix V. In order to avoid confusion with constraint release (CR) mechanisms driven by chain end motion, we call this mechanism lateral link sampling (LLS). However, upon nonlinear deformation or flow, LLS could possibly act as a CR mechanism. A similar mechanism was implemented by Padding and Briels to enforce chain uncrussability in a coarse grained model of polyethylene. The chain average of $U_A(t)$ is defined as

$$U(t) = \frac{1}{N_{ch}} \sum_{\alpha=1}^{N_{ch}} U_A(t)$$

and is shown in Figure 6. It is estimated by averaging over multiple time origins, so that by construction $U(0) = \alpha(0)$ (since $U_A(0) = \alpha(0)$). We see that $U(t)$ increases rapidly and approaches a plateau which is much larger than the average number of links $\langle \alpha(t) \rangle$ per chain (see Figure 3b). This is more evident in the EG-melt (C500); the longer chains and the wider tube of this system is basically the reduced linking time of a link. As the observation time increases, $\tau$ becomes proportional to $T$, with a different prefactor for each link (see Supporting Information). Thus, for adequately long OTs, as the ones utilized here, the $T$'s are equilibrium $T$-independent quantities that characterize the ‘persistence’ of link trajectories with values $0 < \tau \leq 1$. A subset of the links appears to be permanent, with $\tau \simeq 1$. Most of the LCs sampled at $t = 0$ have this property. Instantaneous sampling at any $t$ (a $t$-column in Figure 4), will have a large overlap with this subset, which are basically strong links. This is evident in individual $t$-columns, where most of the points are located on horizontal lines originating at $t = 0$. Note that, by construction, the number of points in a $t$-column is $\alpha(t)$. A striking feature of Figure 4 is that the majority of the $U_A(T)$ links have small, even vanishing $\tilde{t}$'s. Moreover, the number of links sampled over the whole OT, $U_A(T)$, is much larger than those sampled instantaneously, $\alpha(t)$, at any time. Furthermore, after a certain time, $\tilde{t}'$, most of the links with appreciable $\tilde{t}$ have been sampled. The links sampled for the first time at later times, $\tilde{t} > \tilde{t}'$, appear to have a $\tau \simeq 0$. The sampling mechanism that generates blinking links is relatively simple, and is demonstrated in Figure 5. The driving force for this mechanism are stored length fluctuations, as described in detail in Appendix V. In order to avoid confusion with constraint release (CR) mechanisms driven by chain end motion, we call this mechanism lateral link sampling (LLS). However, upon nonlinear deformation or flow, LLS could possibly act as a CR mechanism. A similar mechanism was implemented by Padding and Briels to enforce chain uncrussability in a coarse grained model of polyethylene. The chain average of $U_A(t)$ is defined as

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and is shown in Figure 6. It is estimated by averaging over multiple time origins, so that by construction $U(0) = \alpha(0)$ (since $U_A(0) = \alpha(0)$). We see that $U(t)$ increases rapidly and approaches a plateau which is much larger than the average number of links $\langle \alpha(t) \rangle$ per chain (see Figure 3b). This is more evident in the EG-melt (C500); the longer chains and the wider tube of this system.
lead to a larger number of surrounding chains assembling the tube. In this system, \( U(t) \) does not seem to level off. It increases slowly, with a very small but nonzero decreasing rate. A true plateau with a slightly larger value is expected to exist (fixed tubes), at some longer \( t \).

However, it is not required to sample this regime, since the links sampled for the first time at such long times have a vanishing \( \tau \) (see Figure 4). Most probably, their sampling involves large, cooperative stored length fluctuations (Appendix V), which are not frequent.\(^{112} \) While these links are part of the entanglement environment of a chain, their confinement strength is negligible, and they can be safely considered as weak.

Therefore, for the systems studied, and for all practical purposes, we consider that an observation time of \( T \approx 40 - 50 \tau_\kappa \) is adequate for sampling all pairwise interactions playing a significant role in confinement. We also consider that the links contained in \( U_\alpha(T) \) form a complete set. This set applies to \( \alpha \) the reputed tube constraint. The state of entanglement of \( \alpha \), described by \( z_\alpha(t) \), can be projected, at any time, to the \( U_\alpha(T) \) links. Corresponding statements can also be made for \( U(T) \), which views the system as a single chain. In the next sections, we will refer to these complete sets with the symbols \( U_\alpha, U \), defined as

\[
U_\alpha = U_\alpha(T), \quad U = U(T).
\]

These are adequate representations of the full sets \( U_\alpha(\infty), U(\infty) \), which may have contributions from additional weak links that can be neglected.

### 5.3 Number and Weight Distribution of Link Persistences

We have seen that for the average single chain, the entanglement environment consists of a complete set of \( U \) different links. Here, we characterize this set through the distribution of the persistence of links. In Figure 7a we present the normalized probability distribution, \( p(\tilde{\tau}) \), discretized in intervals of \( \Delta \tilde{\tau} = 0.1 \). The fraction of the \( U \) links falling in each interval is \( p(\tilde{\tau})\Delta \tilde{\tau} \). We observe that \( p(\tilde{\tau}) \) has an absolute maximum at \( \tilde{\tau} \approx 0 \), and a second maximum at \( \tilde{\tau} \approx 1 \). Hence, the majority of the \( U \) links are weak, and the average link persistence, \( \tau_\kappa \), is quite small, namely, 0.30 for the rubber, and 0.24 for the EG-melt.

In the Supporting Information we present \( p(\tilde{\tau}) \) for several increasing \( T \), starting from \( T = \tau_\kappa \). It is shown that for the observation times discussed here the distribution becomes invariant. That is, we have increased \( T \), we would have obtained, practically, the same distribution. This is a necessary condition in order to consider \( U = U(T) \) as a complete set and \( T \)-independent. It is satisfied when sampling the entanglement environment for long OTs, where the linking time of any link becomes proportional to \( T \), so that the corresponding persistence, \( \tilde{\tau} = \tau/T \), acquires an equilibrium value. Additionally, at long \( T \)’s the weak links of the environment have been adequately sampled.

In order to associate the persistence of a link with a confinement strength, we have to note that in \( p(\tilde{\tau}) \) all links are equally weighted. Therefore, \( \tau_\kappa \) corresponds to the number-average link persistence. However, equal weighting hides the fact that the tube constraint is applied mainly by links with \( \tilde{\tau} \approx 1 \). In order to give greater importance to these links we need to estimate the weight average distribution of \( \tilde{\tau} \)’s, \( \bar{w}(\tilde{\tau}) \), which is defined as

\[
\bar{w}(\tilde{\tau}) = \frac{\tilde{\tau}}{\tau_\kappa} p(\tilde{\tau}) = \sum_{\tilde{\tau} = 1}^{\tilde{\tau}} \frac{\tilde{\tau}}{\tau_\kappa} z_\alpha(t) \left\{ \sum_{\beta = 1}^{\tau_\kappa} \frac{\tilde{\tau}}{\tau_\kappa} z_\alpha(t) \right\}
\]

and corresponds to the confinement strength of this link. The closer \( \tilde{\tau} \) is to unity, the stronger the confinement provided by a link. Similarly, \( \bar{w}(\tilde{\tau})d\tilde{\tau} \) is the fraction of confinement provided by the \( U \) links having a persistence between \( \tilde{\tau} \) and \( \tilde{\tau} + d\tilde{\tau} \). Note that \( p(\tilde{\tau}) \) and \( \bar{w}(\tilde{\tau}) \) view the system as a single chain.

The normalized weight distribution \( \bar{w}(\tilde{\tau}) \) is shown in Figure 7b.

---

**Figure 6.** Average number of different linked chains or links which make the tube constraint up to time \( t \). \( \tau_\kappa \) is 1.3 ns for the rubber and 1.7 ns for the EG-melt (C500).

**Figure 7.** (a) Normalized distribution of link persistences. Since all links are equally weighted, this distribution corresponds to the number distribution of \( \tilde{\tau} \)'s. (b) Normalized weight distribution of link persistences. Log-log coordinates in the insets. The observation time, \( T \), is 35\( \tau_\kappa \) for the rubber, and 50\( \tau_\kappa \) for the EG-melt (C500).
We see that confinement comes from, mainly, links with $\bar{\tau} \approx 1$. However, the form of $w(\bar{\tau})$ does not allow for a separation of links into strong and weak. This issue is discussed in detail in Section 5.7. The weight-average link persistence, $\tau_w$, is 0.76, for both systems. It is much larger than $\tau_n$ and it corresponds to the strength weighted average persistence of links.

In order to clarify this section further, it is useful to make an analogy with a corresponding slip-link system where all entanglements are trapped. In this case the links do not blink and all link trajectories are continuous. Then, assuming $Z$ links per chain, we have $z(t) = Z$, at any $t$, and therefore $U = \langle z(t) \rangle = Z$, for any observation time. For each link $\bar{\tau} = 1$ (permanent links), which leads to $\tau_n = 1$. Regarding $p(\bar{\tau})$, it would be zero for $\bar{\tau} \neq 1$, and unity for $\bar{\tau} = 1$ (delta function at $\bar{\tau} = 1$). Regarding $w(\bar{\tau})$, from Eq. (4) we have $w(\bar{\tau}) = p(\bar{\tau})$, and thus, $\tau_n = \tau_n = 1$. Concerning individual links, from Eq. (5) we find that all links have an equal confinement strength, $w = 1/Z$.

Finally, we note that the strength of a link could be associated with additional quantities. For example, a time average of suitably defined elastic interactions over the four strands making the link, the local friction between the mated chains, the real space fluctuations of the link, the monomer diffusion through a link, etc. At a first approximation, here, we consider that all these quantities are expressed indirectly through the link ‘persistence’. They could be extracted from atomistic simulations as long as individual link trajectories can be defined, as discussed in Section 5.6 and in Appendix II).

### 5.4 Collective Entanglement Environment, and Mean-Field Type Links

A useful relation can be obtained by examining Figure 4, again. Denoting by $\tau_{\alpha\beta}$ the LT of a link between chains $\alpha, \beta$, the following equation holds,

$$\sum_{\beta=1}^{U_\alpha} \tau_{\alpha\beta} = \int_{0}^{T} \tilde{z}_\alpha(t) dt \ ,$$

where $U_\alpha = U_{\alpha}(T)$. With reference to Figure 4, Eq. (6) states that in order to add up the LTs of the links sampled by $\alpha$, we can sum either segmented horizontal lines (left-hand side), or the number of points in vertical r-columns (right-hand side). In either case, the sums correspond to the total plotted length of the link trajectories, up to time $T$.

To proceed further we note that the average LT of a link between chains $\alpha, \tau_{\alpha\alpha}$, and $\langle z_\alpha(t) \rangle$, can be written as

$$\tau_{\alpha\alpha} = \frac{1}{U_\alpha} \sum_{\beta=1}^{U_\alpha} \tau_{\alpha\beta} = T \tau_{\alpha\alpha} \ ,$$

$$\langle z_\alpha(t) \rangle = \frac{1}{T} \int_{0}^{T} \tilde{z}_\alpha(t) dt \ .$$

Then, by dividing Eq. (6) with $U_\alpha T$, we find

$$U_\alpha \tau_{\alpha\alpha} = \langle z_\alpha(t) \rangle T \ .$$

Since $U_\alpha$ and $\tau_{\alpha\alpha}$ are independently distributed variables over the chains of the system (numerically checked), the chain average of Eq. (9) reads

$$U \tau_n = \langle z(t) \rangle T \ ,$$

that can also be written in the $T$-independent form

$$U \tau_n = \langle z(t) \rangle \cdot 1 \ ,$$

which shows that the average link persistence is $\bar{\tau}_n = \langle z(t) \rangle / U$, and that the $U$ links can be mapped to $\langle z(t) \rangle$ links with a persistence of unity. Eq. (10), Eq. (11), which are exact, provide a concise microscopic description of entanglements. It can be understood as follows.

By observing the system over a time $T$, which covers a few times its longest relaxation time, we found that for the average single chain the entanglement environment is made by $U$ different chains, with which the constrained fluctuating PP collides and forms binary links. Each link represents a pairwise uncrossability interaction, with a specific mate chain. The average linking time, $\tau_n$, is much smaller than $T$, because the majority of links are active only for very short times (linking links). The large number of these interactions shows that the entanglement environment is inherently collective. This description corresponds to the left-hand side of Eq. (10).

The right-hand side is basically a mean-field type average of the left-hand side, where the identities of the mate chains are mixed. It is understood as follows. For the average single chain, the entanglement environment can be viewed as $\langle z(t) \rangle$ links per chain which are active over the entire observation time, $T$. These links do not blink. Their linking time is $\tau = T$ (permanent links), so that their link persistence is $\bar{\tau} = 1$. However, the identities of their mate chains, and therefore their coordinates in real space and along the chain, are not specific. They are of mean-field type. Each one represents many pairwise interactions, and all together represent the complete set of the $U$ links on the left-hand side. This view leads to the foundations of the mean field picture employed in slip-link models.

The above discussion is also valid in the limit $T \to \infty$. In this case, the appropriate equation for the mapping of the collective entanglement environment to a small number of $\langle z(t) \rangle$ mean-field type interactions is Eq. (11). In stochastic slip-link models, these mean-field interactions are usually realized as binary slip links, though they are not necessarily pairwise. For example, it is possible to consider that slip links represent multichain interactions, as has been discussed by Nair and Schieber. \ref{33} The degree of multichain or pairwise character of these mean field interactions is examined in the next section.

### 5.5 Pairwise Parameter

To examine the mean field uncrossability interactions in more detail, we define $z_{\alpha}(t)|z_{\alpha}(0)$ as the number of links at $t = 0$, which also contribute to $z_{\alpha}(t)$ at time $t$, i.e., the ‘overlap’ between the sets contributing to $z_{\alpha}(0)$ and $z_{\alpha}(t)$. By definition, this number is smaller than or equal to $z_{\alpha}(0)$ (see Figure 2). By averaging $z_{\alpha}(t)|z_{\alpha}(0)$ over multiple time origins along an equilibrium trajectory, we obtain the autocorrelation function $\langle z_{\alpha}(t)|z_{\alpha}(0) \rangle$. Note that the links which contribute to $z_{\alpha}(t)|z_{\alpha}(0)$, for a different time origin, need not always be the same. The ratio

$$q_{\alpha}(t) = \frac{\langle z_{\alpha}(t)|z_{\alpha}(0) \rangle}{\langle z_{\alpha}(t) \rangle} \ .$$

is the percentage of links at time $t = 0$ which are also present at time $t$, irrespective of their state of entanglement with $\alpha$ at times between 0 and $t$. The corresponding average over all chains, $q(t)$, is plotted in Figure 8. It is defined as
The plateau indicates the pairwise character of the mean-field interactions that confine the average single chain. The squares denote corresponding predictions from the weight-average link persistence $\bar{\tau}_w(T)$, for observation times $(T/\tau_c) = 0, 1, 2, 5, 10, 15, \ldots, 50$ (see text).

$$q(t) = \frac{\sum_{\alpha=1}^{N\alpha} \langle z_{\alpha}(t) \mid z_{\alpha}(0) \rangle}{\sum_{\alpha=1}^{N\alpha} \langle z_{\alpha}(t) \rangle} = \frac{\langle z(t) \mid z(0) \rangle}{\langle z(t) \rangle}$$

We observe that $q(t)$ starts from unity ($t = 0$), the largest possible value, then decreases, and after a certain time reaches a plateau. For all systems, this time is around $t = 10\tau_c$. The decay of $q(t)$ is due to the mean-field nature of the $\langle z(t) \rangle$ links. The pairwise character of these links is quantified by the pairwise parameter, $q$, which is defined as the asymptotic value $q = q(t \to \infty)$. We find that $q$ is 0.76 for the rubber, 0.74 for the EG-melt (C500), and 0.81 for the EG-melt (C1000).

In principle, $0 < q \leq 1$, though it is obvious that it cannot become arbitrarily small. According to Eq. (13), when $q = 0$, the chains linked to a reference chain at times zero and $t$ would all be different to each other, something impossible in a system with fixed topology, $q = 1$ corresponds to a strictly pairwise character. The larger the deviation of $q$ from unity, the higher the multichain or collective character of the environment. Here, the mean field links have a strong pairwise character, supporting the employment of binary links in slip-link models. Moreover, it seems that the pairwise parameter is practically chain length and entanglement density independent, a very interesting result.

A simple model which relates $q$ to the weight-average link persistence, and therefore to the confinement strength of pairwise interactions, is presented below. The linking probability, $P_{\alpha \beta}$, that chain $\alpha$ is linked to chain $\beta$, can be estimated as the ratio of the linking time $\tau_{\alpha \beta}$, over the observation time $T$, i.e., $P_{\alpha \beta} = \tau_{\alpha \beta} / \tau$. The probability that these chains are not linked is 1. Obviously, $P_{\alpha \beta} = 1 - P_{\alpha \beta}$. The analysis below concerns long observation times, $T \to \infty$, where these probabilities become $T$-independent, with the convention $t \equiv T$.

The average number of different chains linked to chain $\alpha$ is

$$\langle z_{\alpha}(t) \rangle = \sum_{\beta=1}^{U_{\alpha}} P_{\alpha \beta} = \sum_{\beta=1}^{U_{\alpha}} \tau_{\alpha \beta}$$

On the same footing, it is meaningful to assume that the time correlation function defined in Eq. (12) can be written as

$$\langle z_{\alpha}(t) \mid z_{\alpha}(0) \rangle = \sum_{\beta=1}^{U_{\alpha}} P_{\alpha \beta} P_{\alpha \beta}$$

Then, the weight-average persistence of the links of chain $\alpha$ reads

$$\bar{\tau}_w,\alpha = \sum_{\beta=1}^{U_{\alpha}} w_{\alpha \beta} \bar{\tau}_{\alpha \beta} = \frac{1}{\langle z_{\alpha}(t) \rangle} \sum_{\beta=1}^{U_{\alpha}} P_{\alpha \beta} P_{\alpha \beta}$$

which leads to

$$\langle z_{\alpha}(t) \rangle \bar{\tau}_{w,\alpha} = \langle z_{\alpha}(t) \mid z_{\alpha}(0) \rangle \cdot 1$$

As in Eq. (9), $\langle z_{\alpha}(t) \rangle$ and $\bar{\tau}_{w,\alpha}$ are independently distributed variables over the chains of the system (numerically checked). Thus, the chain average of Eq. (17) reads

$$\langle z(t) \rangle \bar{\tau}_w = \langle z(t) \mid z(0) \rangle \cdot 1$$

From Eq. (13), Eq. (18), and by taking into account that $T \to \infty$ we then find

$$q = \bar{\tau}_w$$

Moreover, Eq. (18) can now be written as

$$q U \bar{\tau}_w = \langle z(t) \rangle \bar{\tau}_w$$

which is the ‘weighted’ form of Eq. (11).

A key result of this work is Eq. (18). In the latter, the $\langle z(t) \rangle$ mean-field links, of strength-weighted average link persistence $\tau_w$, are mapped to $\langle z(t) \mid z(0) \rangle$ links with a strength-weighted persistence of unity (shown explicitly in the equation).

Eq. (19) is in very good agreement with the data of Figure 8. Even for short observation times $T$, where $\tau_w$ is $T$-dependent (see Supporting Information), the relation $q(T) = \bar{\tau}_w(T)$ is able to follow the data, while in the asymptotic regime the agreement is very good. The small deviations are due to Eq. (15), which holds only approximately.

The above eqs when $\tau_w = \bar{\tau}_w = 1$, i.e., when the links do not ‘blink’, lead to $\langle z(t) \rangle = \langle z(t) \mid z(0) \rangle = U$, and $q = 1$. This is the case of a corresponding slip-link model, i.e., of a strictly pairwise entanglement environment (see Section 5.3). In practical terms, the meaning of the pairwise parameter, and of Eq. (13), is that the mean-field links can be realized as $q(z(t))$ effective binary links, with specific mate chain identities and coordinates.

### 5.6 Selecting the Strongest Pairwise Interactions

In Section 5.3, we have seen that the links have various strengths, which were categorized according to their link persistence, $\bar{\tau}$. However, a specific persistence scale separating the links into strong and weak does not seem to exist. The reason is that, for intermittent links, it is not only the linking time which qualifies a link, but also how this time is distributed over the observation time. The more uniformly it is distributed, the less weak is the link, as long as its persistence is appreciable. Therefore, a methodology for selecting the strongest links has to consider the idle time intervals, or time gaps, $\tau_g$, of the blinking links. These are the gaps in the ‘link trajectories’ of Figure 4.

In Figure 9 we present the normalized distribution of $\tau_g$ reduced
by $\tau_e$. The time gaps before the first, or after the last appearance of a link contribute to the distribution. We see that it follows a power-law decay (inset), and thus it is scale-free. The exponent is somewhat higher than -2. Almost all values are concentrated at $\tau_g \ll \tau_e$, and large time gaps are rare.

In order to select the strongest links, a possible strategy is to consider that, for idle time intervals less than or equal to a maximum $\tau_{g,\text{max}}$, the links are instead active. This way we actually ‘fill in’ certain time gaps of intermittent link trajectories, such as those in Figure 4. When all the time gaps of a trajectory are less than or equal to the specified $\tau_{g,\text{max}}$, then this link spans the whole observation time, and it can be selected as a link which is sampled at least once within this time interval. The idle times at the beginning and end of the observation time, if present, can be treated with a different $\tau_{g,\text{max}}$. Here, for simplicity, a common $\tau_{g,\text{max}}$ has been used.

The ratio of the average number of selected links per chain, $z_{\text{sel}}$, over $\langle z(t) \rangle$, for various values of $\tau_{g,\text{max}}$, is shown in Figure 10. In the regime of interest the ratio scales as $\langle \tau_{g,\text{max}}/\tau_e \rangle^{0.42}$. From this plot we can pick out an appropriate time scale for filling the gaps of link trajectories. Below we examine two possible choices, and a third one is examined in the next section.

Following the outcome of weighted averaging, $\tau_{g,\text{max}}$ can be adjusted so that we select as many links as the effective binary links, i.e., $z_{\text{sel}} = q \langle z(t) \rangle$, where $q$ is the pairwise parameter. In Figure 10, this case is shown with filled symbols, and the corresponding $\tau_{g,\text{max}}$ is approximately one $\tau_e$, for both systems.

The other choice is to select as many links as the average number of links per chain, which are mean-field type links (see Section 5.4). That is, $z_{\text{sel}} = \langle z(t) \rangle$. This scenario corresponds to a ‘realization’ of the mean field in terms of specific pairwise interactions and link coordinates. In Figure 10, the corresponding $\tau_{g,\text{max}}$ is estimated from the intersections of the dashed line with the data. It is 1.5$\tau_e$ for the rubber and 2$\tau_e$ for the EG-melt (C500).

The time scale of the maximum time gaps is very reasonable. The distribution of the persistence of selected links is shown in Figure 11. The peak at weak links has now disappeared. We also see that a smaller $\tau_{g,\text{max}}$ leads to fewer selected links, with larger persistencies. It also seems that the distributions are superposable. By adjusting $\tau_{g,\text{max}}$ to fit the effective binary links, contributions from links with $\tau < 0.45$ vanish.

To examine the effectiveness of this strategy, in Figure 12 we plot the trajectories of all the links selected under the condition $z_{\text{sel}} = q \langle z(t) \rangle$, with $\tilde{\tau} \simeq 0.45$. There are few of them. It is seen that they blink in such a way that they cover the whole observation time.

The above results show that the entanglement environment can be mapped to specific pairwise interactions, according to a given link density. The selection procedure sorts out the links according to their strength, and then selects interactions from the strong part of the confinement strength spectrum. A certain time scale (interval), $\tau_{g,\text{max}}$, is introduced, which is self-consistently determined to be of the order of one $\tau_e$. The selected links have the property that they are sampled at least once within this time interval.

The use of this time scale permits a controlled study with respect to a slip-link mapping of the system, and the development of simulation schemes which track the motion of entanglements at equilibrium, or under deformation and flow. For trapped entanglements, this is a straightforward procedure. First, the strong links can be selected by analyzing a short equilibrium MD trajectory. Then, it is possible to follow these links upon any kind of deformation. The whole method is relatively fast, as long as the necessary software exists. When CR is active, an additional scheme for dealing with the update of links would be required.

### 5.7 Link Density, Storage Modulus and Slip-Link Mapping

Rheological estimates of $M_e$ from plateau modulus measurements are in very good agreement with $M_e$ estimates from
the Kuhn segment of the PPs, as obtained from atomistic systems. However, this length scale is larger than the mesh length, \( M_{TC} \) of the networks presented here. Conformationally speaking, the larger link density is due to exponentially decaying orientational correlations along successive steps of the PP. A slowing down of Rouse modes conforming to the mesh length \( M_{TC} \) of the networks has also been observed.\(^{61,111}\)

Hence, a one-to-one mapping of our networks to slip-link models employing the rheological entanglement density, is not viable. However, given that the links here have various strengths, a way to resolve this issue is to consider the PP length as obtained from the networks, but for the slip-link motion under deformation, flow, or in equilibrium, to consider only a subset of very strong links. For example, in Appendix II we present the mean square displacement of the selected links \( z_{sel} = q(z(t)) \), which was constructed by averaging over their individual real space trajectories.

Then, a practical approach is to adjust \( \tau_{g,max} \) of the previous section, so that the selected link density matches the rheological entanglement density. Applying this scheme here, we found that only the very strong links, with a persistence \( \tau > 0.96 \), were selected (see below). The required \( \tau_{g,max} \) is very small, approximately 0.1\( \tau_e \), for both systems. It remains to be shown that the spatial statistics of these links provide a meaningful mesoscopic picture. This approach seems promising as a simple coarse graining scheme for bridging the gap between atomistic and slip-link models. We note that this scheme is self-consistent, since \( \tau_{g,max} \) is adjusted by the \( M_e \) obtained from the Kuhn length of PPs. Alternatively, \( \tau_{g,max} \) can be adjusted so that the selected link density matches a relaxation modulus obtained from simulation.

Evidently, the most important question is how the trapped links revealed here contribute to the low frequency storage modulus of an elastomer. For example, we have seen that the majority of the links blink instead of being continuously present (Section 5.2). The blinking is due to stored length fluctuations (Appendix V), which at the network level lead to thermal PP fluctuations and considerable idle times in certain link trajectories (Figure 4). Stored length fluctuations can ‘relax’ the constraining effect of these links, either fully or partially. Most probably, weak links can be fully relaxed so that they do not contribute to the modulus, while strong links can only partially relax. In this respect, the persistence of each link is indicative of its contribution to the low frequency modulus.

To investigate this issue further, we define the cumulative distribution function (CDF) of the persistence of links

\[
p_{cdf}(\tau) = \int_{\tau}^{1} p(\tau') d\tau' ,
\]

where \( p(\tau) \) is integrated backward, from the right to the left. \( p_{cdf}(\bar{\tau}) \) is the fraction of \( U \) links with a persistence \( \tau \leq \bar{\tau} \leq 1 \). We also introduce \( w_{cdf}(\bar{\tau}) \), the CDF corresponding to \( w(\tau) \), which is defined similarly to \( p_{cdf}(\tau) \). \( w_{cdf}(\bar{\tau}) \) is the fraction of confinement strength provided by links with a persistence \( \tau \leq \bar{\tau} \leq 1 \). Both distributions are shown in Figure 13. From the form of the CDFs we see that a \( \tau \)-scale separating the links into strong and weak is not obvious. However, we observe that the two \( w_{cdf}(\bar{\tau}) \) superpose, and that there are two regimes. From \( \bar{\tau} \approx 1 \) to \( \bar{\tau} \approx 0.95 \) the confinement strength provided by these links increases abruptly, while below \( \bar{\tau} \approx 0.95 \) it follows a smooth increase.

A way to examine Figure 13 with respect to relaxation effects is the following. In the inset we present a table with the required maximum time gap, \( \tau_{g,max} \), so that the number of selected links, \( z_{sel} \), matches certain link densities discussed in the article. \( Z = \langle L_{pp} \rangle / d - 1 \), where \( \langle L_{pp} \rangle \) is the average PP contour length, and \( d \) is the PP Kuhn length obtained by mapping the PPs to random walks (see Appendix II). \( Z \) corresponds to the number of ‘rheo-
logical’ entanglements, i.e., \( Z = (M/M_e) - 1 \), and is much smaller than \( \langle z(t) \rangle \).

It is meaningful to assume that links with a large \( \tau_{g, \text{max}} \) can relax to a larger degree than links with a small \( \tau_{g, \text{max}} \). For example, we can safely consider that the links with \( \tau_{g, \text{max}} \geq \tau_R \) is the longest monomer relaxation time in the system (Appendix II), do not contribute to the low frequency modulus. On the other hand, links with \( \tau_{g, \text{max}} < \tau_e \) can contribute to the modulus, but they can also partially relax, possibly to a different degree. Grossly speaking, we can assume that at low frequencies links with \( \tau_{g, \text{max}} > \tau_e \) become fully relaxed. We envisage that links which disappear and reappear, with idle times longer than \( \tau_e \), have been relaxed by stored length fluctuations.

To proceed further we need to associate a pair of \( (\tau_{\text{sel}}, \tau_{g, \text{max}}) \), to a link persistence. This can be done by associating \( \tau_{\text{sel}} \) to a specific \( \tau \), such that the fraction of the \( U \) links with \( \tau \leq \tau \leq 1 \) is \( \tau_{\text{sel}} \). That is, \( U_{\tau_{\text{eff}}} (= \tau_{\text{sel}}) \). For example, each symbol of Figure 13 is plotted at a \( \tau \) associated with a specific pair \( (\tau_{\text{sel}}, \tau_{g, \text{max}}) \) of the table in the inset. We see that the fraction of links with \( \tau_{g, \text{max}} < \tau_e \) is approximately 20%, with \( \tau > 0.65 \), for both systems (circle symbols). These links provide approximately 70% of the confinement strength, (see \( U_{\tau_{\text{eff}}} \) they are \( q_{\tau}(t) \)) in number, and they can potentially contribute to the low frequency modulus. In Figure 13 they are enclosed in a gray shaded area.

The rheological entanglement density corresponds to \( Z \) very strong links with \( \tau_{g, \text{max}} \sim 0.1 \tau_e \) and \( \tau > 0.96 \) (square symbols). These links provide only 30% to 40% of the confinement strength.

Interestingly, the full confinement strength is provided by links with \( \tau_{g, \text{max}} < \tau_e \) (diamond symbols). The above results demonstrate that weak links, though essentially trapped, do not contribute to the low frequency modulus, in contrast to common belief. It seems that stored length fluctuations can fully relax weak topological interactions. The role of these fluctuations in the viscoelasticity of elastomers and melts is central. Regarding the contribution of weak links to the plateau modulus of a melt, the above discussion is still valid.

### 5.8 Discussion

We have seen that the pairwise uncrossability constraints detected as kinks by chain shrinking methods, such as the Z1,60,61 CReTA 61 algorithms, or the primitive path analysis method,58,118 can be either strong or weak interactions, and not detected at all times (link blinking), even in the absence of CR. The entanglement environment is collective, and certain binary interactions are more important than others. Thus, under nonlinear deformation or flow, where some kind of entanglement network evolution takes place (due to CR), it is possible that changes in the number of detected kinks 75,118,119 do not reflect the actual microscopic changes of the entangled state of the system.

### 6 Summary and Conclusions

In molecular theories of rubber elasticity and polymer rheology, the microscopic entanglement constraints are replaced with a confining mean field, that restricts laterally chain dynamics and fluctuations. Here, we examine in detail at what level this mean field is a collective entanglement effect, as in tube model theories, or can be described through certain pairwise uncrossability interactions, as in slip-link models (see p. 552 in ref 20).

To this end, for the average single chain, the entanglement environment was expanded to a set of \( U \) local links. Each link represents a pairwise uncrossability interaction with a different neighboring chain, resembling a familiar slip-link. Since in our systems all entanglements are trapped, we claim that the \( U \) pairwise interactions form a complete set, and that the entangled state of the system, at any time, can be projected to them. Even if some interactions have been left out of the analysis, it is shown that they are too weak to be considered.

The number of the \( U \) interactions, or links, is very large, which means that the entanglement environment is inherently collective. However, their confinement strength varies from strong to weak, and the majority of them are weak. The plethora of these links is due to the fact that Primitive Paths (PPs) are fluctuating objects. Thus, as a PP fluctuates, it collides with the contours of other PPs and forms local links with them. This sampling mechanism reveals that weak links appear as intermittent, blinking links, because they are not continuously sampled. However, there also exist strong links which are sampled continuously, and appear as permanent constraints. The weak links can “disappear” for time intervals longer than \( \tau_e \), thus they do not contribute to the low frequency modulus of an elastomer, or the plateau modulus of a melt. Overall, the average effect of these interactions is to confine a chain through a much smaller number of mean-field type, permanent links, \( \langle z(t) \rangle \), which in principle represent many other pairwise interactions.

The multichain, or pairwise character of these mean-field links was quantified by defining a pairwise parameter, and it was shown that the pairwise character dominates. Moreover, for the systems studied the pairwise parameter is entanglement density and chain length independent. These results imply that the realization of a confining mean field with binary interactions, as in slip-link models, is not in contrast with a collective entanglement environment.

On this basis, we proposed a self-consistent scheme for mapping the entanglement environment to a slip-link model. The scheme discards weak links and selects the strongest binary interactions conforming to a given entanglement density. It is directly applicable to trapped entanglements, and to the analysis of MD deformation trajectories of bead-spring and atomistic polymer networks. From such studies, available slip-link network models can be refined, corrected, or extended, in a hierarchical manner. With suitable extensions,105 the tracking of constraint release events in polymer melt simulations, seems also possible. A list of symbols and abbreviations for this paper is given in Table 1.

### Appendix I: Construction of a Defect-Free Entangled PE Network

In flexible polymers, the degree of chain overlap is controlled120 by density and stiffness. Therefore, it seems unusual that the two systems examined here (rubber and end-grafted melt) have a different degree of entanglement, though the slightly larger \( C_w \) of the rubber would decrease120 \( M_e \), in comparison to the melt. Since both systems are well equilibrated, the difference must be due to the short subchains and the artificial preparation conditions of the network. The latter is constructed as follows. At the nodes of the diamond lattice we place carbon atoms. The lattice constant is then adjusted so that the carbon atoms are bonded with PE chains at full extension (all-trans state of dihedral angles). This is an unentangled state. Then, the network is let to relax at 1 Atm. Because of entropic elasticity and cohesive interactions it contracts to the melt density. At this stage the system is not equilibrated configurationally. To impose equilibration a phantom chain simulation at melt density (NVT ensemble) is performed. Bead overlaps are then eliminated by the gradual push-off104 method.

Entanglements are introduced dynamically after the chains have been linked to a network, by letting them interpenetrate at the phantom chain simulation stage. It seems that this procedure, together
with the short length of the chains, leads to a larger degree of chain overlap in comparison to the melt state. We expect that in networks of very long subchains, where the cross-link density vanishes, this effect would subside. It can also be avoided through random linking of a melt system, at the cost of introducing network defects. On the other hand, the fact that our results have been obtained with different entanglement densities makes our conclusions fairly general.

A different path to defect-free networks has been followed by Everaers, 52,121 It invokes constructing identical interpenetrating diamond networks, where the lattice constant is set to the root mean squared end-to-end distance of melt chains. The system consists of as many networks as are required to reach the melt density, which is held fixed. As in our case, equilibration is achieved by letting the networks interpenetrate at a phantom chain simulation stage. Chain overlaps are then eliminated through a gradual "push-off" stage. In comparison to our networks the chains can also entangle with chains from other identical networks.

**Appendix II: Mean Square Displacement and Characteristic Relaxation Times**

In Figure 14 we present the mean square displacement (msd), $g_1(t)$, of middle chain monomers (10% of each chain), reduced by $t^{0.5}$. Four regimes are discernible. The first one is unimportant. It is usually called ballistic. A second regime, scaling as $t^{0.5}$, corresponds to free Rouse motion within the tube. The next regime is indicative of tube confinement (restricted Rouse motion along a random-walk like tube), and it scales, approximately, as $t^{0.28}$, for both systems. The corresponding tube model prediction 7 is $t^{0.25}$. The last regime corresponds to an invariant msd. From the first crossover (between second and third regime) we estimate $\tau_2$, the time that a monomer ‘hits’ the tube for the first time. It is 1.3 ns for the rubber and 1.7 ns for the EG-melt. From the second crossover we estimate $\tau_R$, the longest monomer relaxation time of each system, which is 9.8 ns for the rubber and 11.5 ns for the EG-melt. This time corresponds to the Rouse time, $\tau_R$. Here, it is denoted with a prime, since the boundary conditions for a Rouse chain are different (free chain ends). For the (C1000) EG-melt, the corresponding crossover times are 1.6 and 19.4 ns (not shown).

In Figure 15 we present the msd of middle chain monomers together with the msd of links selected according to the pairwise parameter, i.e., $\zeta_{sel} = q(\zeta(t))$. The plateaus indicate the absence of reptation and long chain diffusion. We see that the selected links span much shorter distances than real chain monomers. Thus, we can say that the enclosing tube of PPs is a shrunk version of the alleged tube that restricts lateral motion of real chain monomers.

The bars plotted parallel to the plateaus correspond to estimations of certain link lengths related to entanglement constraints. $d$ is the tube diameter estimated from the Kuhn segment of the PP as $d = \langle R^2 \rangle / \langle L_{TP} \rangle$, where $\langle R^2 \rangle$ is the mean square end-to-end distance. $\langle L_{TC} \rangle$ is the mean square edge length of the link network. 61,83 $w_{TP}$ is an estimation of the mean square distance between PP contours from the inverse of the PP contour length density. It is defined as $w_{TP} = \langle V \rangle / \langle L_{TP} \rangle$, where $\langle V \rangle$ is the average volume. Note that the average distance between PP contours is smaller than the average distance between connected links along the PPs, $d_{TC}$. For example, in Figure 5, the distance between PP contours, $h$, is much smaller than the PP segments with average length $d_{TC}$.

For both systems, we see that the msd plateau of the monomers shows a better correlation with $d^2/4$, which corresponds to the squared radius of the tube, but is actually larger than that. This is not surprising, since real chain monomers can penetrate the tube walls, as parts of unentangled loops for example. In both systems we also observe a correlation between the squared tube radius and the mean edge length of the link network. The msd plateaus of selected links are much smaller than these quantities, which means that link fluctuations are restricted to the inner tube region along the tube axis. These plateaus are also larger than $w_{TP}$, especially in the EG-melt, which means that the fluctuating PPs can collide to each other, in accordance with the lateral link sampling mechanism.

**Appendix III: Binary Links, Local Knots, and Ternary Nodes**

As shown in Figure 1, each node links locally two chains, except from few ternary nodes (see below). Within this terminology, each node resembles a binary link that can be resolved into two TCs.
along two ‘mate chains’. Iwata and Edwards have discussed melt dynamics and rheology in terms of local knots which are conceptually similar to the ones presented here. The knots were defined at the real chain level by using as topological criterion a local form of the Gauss linking number. For open chains this criterion is not more rigorous than the one defined here, while it is computationally more complex.

In our networks about 5% of the nodes are ternary. That is, there exist situations where three chains, let us say α, β, and γ, are mutually constrained and form local links with each other (see videos). Detection of these cases is hard and needs special attention. They are mapped to nodes with six edges, which are further resolved into three links, αβ, αγ, βγ, so that all three chains are mutually coupled in pairs. With this choice, the mapping to a network leads exclusively to binary links, as in slip-link models. Information about ternary nodes is retrievable, though.

Upon dynamical evolution of the network, most of the ternary nodes tend to break up into consecutive binary nodes along the same chain, e.g., αβ, αγ, in the former example. The reverse procedure is also possible (see videos). A statistical analysis has shown that between successive links of a chain there exists an effective repulsion. Thus, we expect that, while ternary links certainly appear, they are transient. Interestingly, a repulsive interaction also emerges when enforcing detailed balance in the dynamical equations of the slip-link based primitive chain network model.

Appendix IV: Multiply Linked Chains

In Figure 16 we show the normalized probability, P(n), that a chain α forms n local links with a mate chain in α(t), instantaneously, at time t, (see also Figure 2). For the rubber and the short chain EG-melt P(n) is highly peaked at unity. Therefore, the simplification of dealing with LCs instead of individual links is justified. However, as chain length increases P(1) decreases. Thus, in systems with long chains the analysis should be made against individual links.

Appendix V: Lateral Link Sampling and CReTA Mapping

The large set of different chains sampled by the PP is rather surprising. Since the PPs are determined by contour reduction, then, by construction, the sampled chains have portions inside or at the boundaries of the pervaded volume of a reference chain. Moreover, they are not just simple chain contacts, since linked chains satisfy the local topological criterion of Figure 1, (although some very weak links could be considered as fluctuating chain contacts). The lateral link sampling (LLS) mechanism that takes place is simple and was demonstrated in Figure 5.

In order to provide more evidence for this mechanism, we first define the reduced curvilinear coordinates of a link, sα(t), sβ(t) between chains α, β. These are defined such that 0 ≤ s ≤ 1. That is, sα(t) = xα(t)/Lα(t), where xα(t) is the curvilinear coordinate of the link along chain α, and Lα(t) is the PP contour length. Likewise for sβ(t). Note that a local link can slide along PP contours, hence the time dependence of sα, sβ.

In Figure 17 we plot sα(t), sβ(t), for all the links that were first sampled at a time t > 0. In Figure 4, these links correspond to the trajectories with Uα(t) > Uα(0), (continuous or segmented trajectories above the column to the left of t = 0 in Figure 4). For each one of these links, we plot a single point that denotes sα(t), sβ(t), at the time the link was first sampled. Beyond some explainable heterogeneities, the homogeneous patterns support the LLS mechanism of Figure 5. The EG-melt data are less dense due to the smaller number of simulated chains.

Data concentration at the lower left corner of Figure 17b implies that many of the new links are formed between chain ends. This is explained by the higher probability of instantaneous link formation between PPs attached to the same tetrafunctional crosslink of a network. In the EG-melt, local links with sα ≈ 0 and sβ ≈ 0 are favored. This is an artificial effect, due to the less mobile chain ends because of grafting. The fluctuating PPs have higher probability to collide with certain PP segments which are relatively immobile, or attached to fixed points. In systems with very long chains these heterogeneities would disappear.

The physical origin of the LLS mechanism is the thermal motion of real chains and, more specifically, the thermal motion of stored length, or chain slack. The spatial distribution of this length is different at different time-snapshots of our systems. In slip-link models this ‘redistribution’ is taken into account by the thermal
In Figure 19 we present the calculation of \( q(t) \) for the (C500) melt, but this time analyzed as an end-constrained melt (EC-melt). That is, instead of grafting chain ends to nearby chains, the MD trajectory was generated by keeping all chain ends fixed. This difference has a strong effect on \( q(t) \) due to the presence of a CR mechanism, which can be called end-looping constraint release (EL-CR). A polymer chain, say chain A, can disentangle from a mate chain, say chain B, by developing a lateral unentangled hernia that passes over a chain end of B. That is, by hopping of an internal portion of A over one end of B, through Rouse motion (see Figure 7 in the work of Zhou).

As regards the mapping of a specific spatial distribution of stored length to a network of links by CReTA, this is not unique. CReTA minimizes the total contour length of the system. Assuming that there is a unique, 'global' minimum, the value of the objective function will hardly be changed if we remove some small length from a PP and add it to another. The stochastic nature of the reduction performed by CReTA is subject to this effect. At the network level, this will lead to small displacements of nodes, and therefore, possibly, to link sampling by the mechanism of Figure 5.

Thus, individual CReTA reductions of the same configuration (same spatial distribution of stored length), performed with different pseudorandom numbers, may sample, each time, a slightly different set of links, with practically the same shortest paths (PP conformations and length). This uncertainty is inherent to the reduction problem. Equivalently, one could say that the stored length can only be mapped to a small irregular volume around the centerline of the corresponding tube (see Figure 18), and not to a specific path. This volume encloses a subset of the links which define the tube constraint and has large overlap with the tube core. An individual reduction by CReTA, then, will sample a path line in this volume. The links of the subset which are inside the tube core will be sampled with higher probability. This kind of inherent 'noise' in the network mapping does not affect any of our conclusions. It is very small relative to the thermal noise driving the sampling of the links that assemble the tube constraint.

Appendix VI: Entanglement Constraints at the Level of Shortest Paths and Local Links

A full view of what happens at the link level requires the knowledge of the distances spanned by fluctuating PPs. A short account is given in Appendix II. A more detailed presentation will be given elsewhere. For the systems studied here, it suffices to say that, as time proceeds, the PP segments cover an increasing distance (through lateral fluctuations), until they have fully sampled their corresponding tubes. Because of the coarse graining procedure they span shorter distances than the real chain monomers. Thereby, their enclosing tubes are shrunk versions of the real chain tubes, and while they restrict lateral PP fluctuations they do not restrict lateral monomer fluctuations. We could say that a PP samples the domain around the centerline of the real chain tube. The picture that emerges is described in Figure 18.

Appendix VII: Constraint Release Effects on \( q(t) \)

In Figure 19 we present \( q(t) \) for the (C500) melt, but this time analyzed as an end-constrained melt (EC-melt). That is, instead of grafting chain ends to nearby chains, the MD trajectory was generated by keeping all chain ends fixed. This difference has a strong effect on \( q(t) \) due to the presence of a CR mechanism, which can be called end-looping constraint release (EL-CR). A polymer chain, say chain A, can disentangle from a mate chain, say chain B, by developing a lateral unentangled hernia that passes over a chain end of B. That is, by hopping of an internal portion of A over one end of B, through Rouse motion (see Figure 7 in the work of Zhou).
Larson). Here, we see that EL-CR reduces \( q(t) \) to a lower plateau value, which becomes discernible at \( t/\tau_e \approx 100 \). However, a safe estimation of this plateau would require a longer observation time.

Grossly speaking, the EC-melt can be viewed as the EG-melt with some links not being trapped, but subject to updating by EL-CR. This is similar to a network with dangling chains, i.e., with both trapped and temporary links. In such a system \( q(t) \) will decay toward a certain plateau, characteristic of the density of trapped entanglements. In a melt with free chain ends (exclusively temporary links), \( q(t) \) will eventually vanish. The plateau value for the EC-melt can be viewed as the EG-melt with some links not being trapped, but subject to updating by EL-CR. This is similar to a network with dangling chains, i.e., with both \( \text{trapped and temporary links} \). In such a system \( q(t) \) will decay toward a certain plateau, characteristic of the density of trapped entanglements. In a melt with free chain ends (exclusively temporary links), \( q(t) \) will eventually vanish. The plateau value for the EC-melt will be smaller than that of the EG-melt (smaller trapped entanglement density). However, the pairwise parameter is defined as the plateau of \( q(t) \) when a system is examined with all of its entanglements being trapped (fixed tubes). For this purpose, as explained in Section 3, CR is eliminated by grafting all free chain ends. Therefore, a plateau in \( q(t) \), in the presence of CR, does not correspond to a pairwise parameter for the entanglement environment. The decay of \( q(t) \), or the difference between the EC-melt and EC-melt plateau, can possibly provide some kind of CR information, but this issue was not examined here.

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**Supporting Information**

Entanglement network views of a PE melt (Figure S1), link trajectories for representative chains (Figure S2), number and weight distribution of link persistences, for increasing observation time \( T \) (Figure S3), Number-average and weight-average link persistence as a function of increasing \( T \) (Figure S4), distribution of unselected link persistences (Figure S5), magnified section of Figure 13 (Figure S6), and 15 explanatory videos for the material discussed in the document, with an associated description. This material is available free of charge via the Internet at http://pubs.acs.org.

**Table 1. List of Symbols and Abbreviations**

| Symbol | Description |
|--------|-------------|
| \( N_{ch} \) | Number of chains in the system |
| \( \alpha \) | Chain index, \( 1 \leq \alpha \leq N_{ch} \) |
| \( \tau_e \) | Entanglement time |
| \( \tau_R \) | Rouse time |
| \( M_e \) | Entanglement Molecular Weight |
| \( T \) | Observation Time (OT) |
| \( \tau \) | Linking Time (LT) |
| \( \bar{\tau} = \tau/T \) | Link persistence |
| \( z_\alpha(t) \) | Number of different links or Linked Chains (LCs) constraining \( \alpha \) at time \( t \) |
| \( U_\alpha(t) \) | Cumulative number of the different links sampled by \( \alpha \) up to time \( t \) |
| \( z_\alpha(t)|z_\alpha(0) \) | Subset of links constraining \( \alpha \) at both times, 0 and \( t \) |
| \( \langle z_\alpha(t)|z_\alpha(0) \rangle \) | \( z_\alpha(t)|z_\alpha(0) \) averaged over multiple time origins |
| \( q_\alpha(t) \) | Ratio of \( z_\alpha(t)|z_\alpha(0) \) over \( \langle z_\alpha(t)|z_\alpha(0) \rangle \) |
| \( z(t), U(t), q(t) \) | Chain averages of the corresponding quantities above. |
| \( q \) | Pairwise parameter, \( q = \lim_{t \to \infty} q(t) \) |
| \( p(\bar{\tau}) \) | Number distribution of link persistence |
| \( w(\bar{\tau}) \) | Weight distribution of link persistence |
| \( \bar{\bar{\tau}}_\alpha \) | Average persistence of links of chain \( \alpha \) |
| \( \bar{\tau}_n \) | Average link persistence |
| \( \bar{\tau}_w, \alpha \) | Weight average persistence of links of chain \( \alpha \) |
| \( \bar{\tau}_w \) | Weight average link persistence |
| \( \bar{\tau}_{\alpha\beta} \) | Persistence of a link between chains \( \alpha, \beta \) |
| \( w_{\alpha\beta}(\bar{\tau}_{\alpha\beta}) \) | Confinement strength of a link with persistence \( \bar{\tau}_{\alpha\beta} \) |
| \( P_{\alpha\beta} \) | Probability that chain \( \alpha \) is linked to chain \( \beta \) |
| \( \tau_{\alpha, \max} \) | Maximum time gap of link trajectories |
| \( z_{\text{sel}} \) | Number of selected links per chain |
| \( \text{PP} \) | Primitive Path |
| \( \text{TC} \) | Topological Constraint |
| \( \text{LC} \) | Linked Chain |
| \( \text{LLS} \) | Lateral Link Sampling |
| \( \text{CDF} \) | Cumulative Distribution function |
| \( \text{EG-melt} \) | End-Grafted melt |
| \( \text{EC-melt} \) | End-Constrained melt |
| \( \text{EL-CR} \) | End-Looping Constraint-Release |

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