Pore-blockade times for field-driven polymer translocation

Henk Vocks\textsuperscript{1}, Debabrata Panja\textsuperscript{2}, Gerard T Barkema\textsuperscript{1,3} and Robin C Ball\textsuperscript{4}

\textsuperscript{1} Institute for Theoretical Physics, Universiteit Utrecht, Leuvenlaan 4, 3584 CE Utrecht, The Netherlands
\textsuperscript{2} Institute for Theoretical Physics, Universiteit van Amsterdam, Valckenierstraat 65, 1018 XE Amsterdam, The Netherlands
\textsuperscript{3} Instituut-Lorentz, Universiteit Leiden, Niels Bohrweg 2, 2333 CA Leiden, The Netherlands
\textsuperscript{4} Department of Physics, University of Warwick, Coventry CV4 7AL, UK

Received 6 November 2007, in final form 18 December 2007
Published 15 February 2008
Online at stacks.iop.org/JPhysCM/20/095224

Abstract
We study pore-blockade times for a translocating polymer of length $N$, driven by a field $E$ across the pore in three dimensions. The polymer performs Rouse dynamics, i.e., we consider polymer dynamics in the absence of hydrodynamical interactions. We find that the typical time for which the pore remains blocked during a translocation event scales as $\sim N^{1+2\nu/(1+\nu)}/E$, where $\nu \simeq 0.588$ is the Flory exponent for the polymer. We show, in line with our previous work, that this scaling behavior stems from polymer dynamics in the immediate vicinity of the pore—in particular, the memory effects in the polymer chain tension imbalance across the pore. This result, like numerical results from several other groups, violates the lower bound $\sim N^{1+\nu}/E$ suggested earlier in the literature. We discuss why this lower bound is incorrect and show, on the basis of the conservation of energy, that the correct lower bound for the pore-blockade time for field-driven translocation is given by $\eta N^{2\nu}/E$, where $\eta$ is the viscosity of the medium surrounding the polymer.

1. Introduction
Molecular transport through cell membranes is an essential mechanism in living organisms. Often, the molecules are too long, and the pores in the membranes too narrow, to allow the molecules to pass through as a single unit. In such circumstances, the molecules have to deform themselves in order to squeeze—i.e., translocate—themselves through the pores. DNA, RNA and proteins are such naturally occurring long molecules \cite{1–5} in a variety of biological processes. Translocation is also used in gene therapy \cite{6, 7}, and in delivery of drug molecules to their activation sites \cite{8}. Consequently, the study of translocation is an active field of research: as a cornerstone of many biological processes, and also due to its relevance for practical applications.

More recently, translocation has found itself at the forefront of single-molecule-detection experiments \cite{9–11}, as new developments in the design and fabrication of nanometer-sized pores and etching methods may lead to cheaper and faster technology for the analysis and detection of single macromolecules. In these experiments, charged polymeric molecules, suspended in an electrolyte solution, are initially located on one side of a membrane. The membrane is impenetrable to the molecule except for a nanometer-sized pore. Between the two different sides of the membrane, a DC voltage difference is then applied, which drives the molecule through the pore. When the molecule enters the pore, it affects the electrical resistivity of the circuit, leading to a dip in the electric current supplied by the voltage source. The magnitude and the duration of these dips have proved to be very effective in determining the size and the length of the molecule. The usage of protein pores (modified $\alpha$-haemolysin, mitochondrial ion channel, nucleic acid binding/channel protein etc) and the etching of specific DNA sequences inside the pores \cite{6, 12} have opened up promising new avenues of fast, simple and cheap technology for single macromolecule detection, analysis and characterization, perhaps even allowing DNA sequencing at the nucleotide level.

The subject of this paper is (charged) polymer translocation in three dimensions through a narrow pore in an otherwise impenetrable membrane placed at $z = 0$, as the polymer is driven by a DC voltage across the pore. Our interest is in the
Figure 1. Snapshot of a translocating polymer in a two-dimensional projection of our three-dimensional system. Across the pore of size unity a voltage difference $2\,V$ is applied. The monomer located within the pore is labeled $s$.

scaling behavior for the typical pore-blockade time during a translocation event with polymer length $N$. In practice, the electric field due to the applied voltage decays rapidly with increasing distance from the pore, and for simplicity it is often assumed that only those polymer segments residing within the pore feel the driving force due to the field. For our theory and simulations too, we consider a polymer which only experiences a force acting on its monomers that reside in the pore, as illustrated in figure 1.

To substantiate our theoretical analysis we use extensive Monte Carlo simulations with a three-dimensional self-avoiding lattice polymer model. For the voltage difference across the pore we choose

$$V(z) = \begin{cases} +V & (z \leq -1) \\ 0 & (z = 0) \\ -V & (z \geq 1). \end{cases}$$

Thus, during translocation through the pore, the energy gained by each monomer carrying a charge $q$, in dimensionless units, is given by $\Delta U = 2qV/k_BT$. From now on, favoring notational simplicity, we choose both $q$ and $k_BT$ to be unity. Since we also choose the lattice spacing to be unity in our simulations, the strength of the electric field acting on each monomer within the pore is given by $E = V$.

Details of the lattice polymer model used in this paper can be found in [13, 14]: the polymer moves through a sequence of random single-monomer hops to neighboring lattice sites. These hops can either be ‘reptation’-moves, along the contour of the polymer, or Rouse moves, in which the monomer jumps ‘sideways’ and changes the contour. The definition of time used throughout this paper is such that each monomer attempts a ‘reptation’-move as well as a ‘sideways’-move with rate unity. There is no explicit solvent in our analysis, i.e., the polymer performs Rouse dynamics.

Our conventions to study this problem, all throughout this paper, are the following. We place the membrane at $z = 0$. We fix the middle monomer (monomer number $N/2$) of a polymer of total length $N$ at the pore, apply the voltage as in equation (1) and thermalize the polymer. At $t = 0$ we release the polymer and let translocation commence. We define the typical time when the polymer leaves the pore as the dwell time $\tau_d$: it scales with $N$ in the same way as the pore-blockade time in a full (field-driven) translocation event.

This problem has recently been studied in [15], in which a lower bound $\propto N^{1+\nu}/E$ has been argued for $\tau_d$. This lower bound was derived in the limit of unimpeded polymer movement, i.e., for an infinitely wide pore, or equivalently, in the absence of the membrane. In [15] the authors also suggested that the dynamics of translocation is anomalous (see also [16] in this context).

In the recent past, some of us have been investigating the microscopic origin of the anomalous dynamics of translocation. We have set up a theoretical formalism, based on the microscopic dynamics of the polymer, and showed that the anomalous dynamics of translocation stem from the polymer’s memory effects, in the following manner. Translocation proceeds via the exchange of monomers through the pore: imagine a situation when a monomer from the left of the membrane translocates to the right. This process increases the monomer density in the right neighborhood of the pore, and simultaneously reduces the monomer density in the left neighborhood of the pore. The local enhancement in the monomer density on the right of the pore takes a finite time to dissipate away from the membrane along the backbone of the polymer (similarly for replenishing monomer density on the left neighborhood of the pore). The imbalance in the monomer densities between the two local neighborhoods of the pore during this time causes an enhanced chance of the translocated monomer to return to the left of the membrane, thereby giving rise to memory effects. The ensuing analysis enabled us to provide a proper microscopic theoretical basis for the anomalous dynamics. Further theoretical analysis then led us to the conclusion that in the case of unbiased translocation, i.e., when the polymer is not subjected to an external force, the dwell time scales with length as $\tau_d \sim N^{2+\nu}$ [13, 17, 18], both in two and three dimensions. Our approach based on the polymer’s memory effects also works beautifully for pulled translocation, for which a force $F$ is applied at the head of the polymer: we showed that if $FN^\nu$ is sufficiently large, then the dwell time scales as $\tau_d \sim N^{2+\nu} \sim (FN^\nu)^{-1}$ [19]. In this work, we push ahead with the same formalism to demonstrate that it reveals the physics of field-driven translocation too, thus providing a unified underlying theoretical basis for translocation, based on the theory of polymer dynamics.

Returning to the lower bound for the scaling of the dwell time with polymer length $N$ for field-driven translocation as proposed in [15], we note that subsequent numerical studies did not immediately settle the scaling for $\tau_d$ with $N$, including the one by the authors of [15] themselves. In table 1 we present a summary of the existing numerical results on the exponent for the scaling of $\tau_d$ with $N$ for field-driven translocation. All results quoted are for self-avoiding polymers in the absence of hydrodynamical interactions in the scaling limit.

More recently, this lack of consensus prompted three of us to investigate the issue of field-driven translocation in two dimensions, via a proxy problem, namely, polymer translocation in three dimensions out of strong planar confinements [18]. We showed that the actual lower bound for $\tau_d$ for field-driven translocation is given by $\eta N^{2\nu}/E$, where $\eta$ is the viscosity of the surrounding medium. This inequality
is derived from the principle of conservation of energy: it was shown in [18] that although the presence of the memory effects suggests that the scaling of \( \tau_d \) could behave as \( N^{(1+2\nu)/(1+\nu)} \), since \((1+2\nu)/(1+\nu) < 2\nu \) in two dimensions, conservation of energy overrides the memory effects in the polymer—high precision simulation data suggested, in accordance with those of [15, 21] that the actual scaling of \( \tau_d \) for field-driven translocation in two dimensions is given by \( \tau_d \sim N^{2\nu} \). In three dimensions \( 2\nu < (1+2\nu)/(1+\nu) \), implying that in three dimensions \( \tau_d \sim N^{(1+2\nu)/(1+\nu)} \), which is the central result of this paper.

This paper is organized in the following manner. In section 2 we derive the lower bound \( N^{2\nu} \) for \( \tau_d \) for field-driven translocation. In section 3.1 we discuss a method to measure the polymer’s chain tension in the imbalance of the polymer’s chain tension at the pore. In section 3.2 we analyze the memory effects in the imbalance of the polymer’s chain tension at the pore. In section 4 we discuss the consequence of these memory effects on the translocation velocity \( v(t) \), and obtain the scaling relation of \( \tau_d \) with the polymer length \( N \). We end this paper with a discussion in section 5.

2. Lower bound for \( \tau_d \) for field-driven translocation

As noted in section 1, a lower bound for the dwell time \( \tau_d \sim N^{1+\nu}/E \) has been proposed in [15]. The underlying assumption behind this result is that, with or without an applied field, the mobility of a polymer translocating through a narrow pore in a membrane will not exceed that of a polymer in bulk (i.e., in the absence of the membrane). This mobility is then obtained under two more assumptions for the behavior of a polymer under a driving field:

(i) To mimic the field acting on a translocating polymer, the field on the polymer in bulk has to act on a monomer whose position along the backbone of the polymer changes continuously in time. As a result, there is no incentive for the polymer to change its shape from its bulk equilibrium shape, i.e., the polymer can still be described by a blob with radius of gyration \( \sim N^{\nu} \) in the appropriate dimension.

(ii) The polymer’s velocity is proportional to \( DE \), where \( E \) is the applied field, and \( D \) is the diffusion coefficient scaling as \( D \sim 1/N \) for a Rouse polymer.

Of these two assumptions, note that (ii) is obtained as the steady state solution of the equation of motion of a Rouse polymer, in bulk, with uniform force and vanishing internal forces, see for instance [25] (equation VI.10). We have already witnessed in many occasions [13, 15–17, 19, 24, 26, 27] that the dynamics of translocation through a narrow pore is anomalous (subdiffusive), as a consequence of the strong memory effects discussed in the previous section, and also that these memory effects are so strong that the velocity of translocation is not constant in time [18, 19]. The anomalous dynamics and the memory effects are crucial ingredients that question the validity of the lower bound \( N^{1+\nu} \) for \( \tau_d \) for field-driven translocation.

It is however possible to derive a lower bound for \( \tau_d \) for field-driven translocation, based on the principle of conservation of energy. Consider a translocating polymer under an applied field \( E \) which acts only at the pore. By definition, the \( N \) monomers of the polymer translocate through the pore in a time \( \tau_d \). The total work done by the field in this time \( \tau_d \) is then given by \( E N \). During translocation, each monomer travels over a distance of order \( R_g \), leading to an average monomer velocity \( v_m \sim R_g/\tau_d \). The rate of loss of energy due to the viscosity \( \eta \) of the surrounding medium per monomer is given by \( \eta v_m^2 \). For a Rouse polymer, the frictional force on the entire polymer is a sum of frictional forces on individual monomers, leading to the total energy loss due to the viscosity of the surrounding medium during the entire translocation event scaling as \( \sim N \eta R_g^2 v_m^2/\tau_d \). This loss of energy must be less than or equal to the total work \( E N \) done by the field, which yields us the inequality \( \tau_d \geq \eta R_g^2/\nu \). This inequality must be less than or equal to the total work \( E N \) done by the field, which yields us the inequality \( \tau_d \geq \eta R_g^2/\nu \).

3. Memory effects in the chain tension perpendicular to the membrane

A translocating polymer can be thought of as two segments of polymers tethered at the pore, while the segments are able to exchange monomers between them through the pore. In [17] we developed a theoretical method to relate the dynamics of translocation to the imbalance of chain tension between these two segments across the pore. The key idea behind this method is that the exchange of monomers across the pore responds to \( \phi(t) \), this imbalance of chain tension; in its turn, \( \phi(t) \) adjusts to \( v(t) \), the transport velocity of monomers across the pore. Here, \( v(t) = \dot{s}(t) \) is the rate of exchange of monomers from one side to the other.

The memory effects discussed in section 1 in terms of relaxation of excess monomers (or the lack of monomers) in the immediate vicinity of the pore translates immediately to that of the imbalance of the chain tension across the pore—local accumulation of excess monomers reduce the chain tension, while local lack of monomers enhance it. Quantitatively speaking, in the presence of memory effects, the chain tension imbalance across the pore \( \phi(t) \) and the velocity of translocation \( v(t) \) are related by

\[
\phi(t) = \phi(t=0) + \int_0^t dt' \mu(t - t') v(t')
\]  

\[5\] For unbiased translocation, \( E N \) in this argument is to be replaced by the difference in free energy (or entropy) of a threaded polymer, corresponding to \( s = N/2 \), and the translocated polymer, corresponding to \( s = N \). This leads to the inequality \( \tau_d \geq \eta N^{1+2\nu} \).
via the (field-dependent) memory kernel $\mu(t)$, which could be thought of as time-dependent ‘impedance’ of the system. Using the Laplace transform, this relation could be inverted to obtain $v(t) = \int_0^\infty d' a(t - t') [\Phi(t - a) - \Phi(t')]$, where $\Phi(t)$ can be thought of as the ‘admittance’ of the system. In the Laplace transform language, these are related to each other as $\Phi(\Phi) = a^{-1}(k)$, where $k$ is the Laplace variable representing inverse time [13, 17–19].

3.1. Chain tension perpendicular to the membrane

Measuring chain tension directly is difficult. We therefore use a method developed earlier [18, 19] to monitor the chain tension near the pore.

By definition, the chain tension imbalance $\Phi(t)$ is the difference of the chain tensions on the right and the left side of the pore: $\Phi(t) = \Phi_R(E, t) - \Phi_L(E, t)$. Both $\Phi_R(E, t)$ and $\Phi_L(E, t)$ are functions of the applied electric field $E$ across the membrane. Note, from the applied potential $(1)$, that the field $E$ acts on the monomers at site $z = -1$ towards the pore, while it acts on those at site $z = 1$ away from the pore. Using the convention that $E < 0$ (resp. $E > 0$) implies a field acting towards (resp. away from) the membrane, we have

$$\Phi(E, t = 0) = \begin{cases} \Phi_R(t = 0) & \text{ (} E < 0 \text{) } \\ \Phi_L(t = 0) & \text{ (} E > 0 \text{) } \end{cases}$$

(3)

Now consider a different problem, where one end of a polymer is tethered to a fixed membrane, yet the number of monomers are allowed to spontaneously enter or leave the tethered end, under the effect of an electric field $E$. Then, following the methodology described in [18, 19], we have

$$\Phi(E, t = 0) = k_B T \ln \frac{P_+}{P_-}$$

(4)

where $P_+$ (resp. $P_-$) is the probability that the left (or the right) polymer segment has one monomer less (resp. one extra monomer).

Note that even for $E = 0$, as already stressed in [19], there is nonzero chain tension $\Phi_0$ at the pore, due to the presence of the membrane. A polymer’s free energy close to a membrane is higher than its free energy in bulk. In other words, the membrane repels the polymer, and as a result, for a polymer with one end tethered to a membrane, the monomers close to the membrane are more stretched than they would be in the bulk.

For a translocating polymer equation (4) cannot be used, so to compute $\Phi_R(t)$ and $\Phi_L(t)$ one needs a suitable proxy. In the cases of unbiased translocation [13, 17], translocation with a pulling force [19] and translocation out of planar confinements [18], we have seen that the center-of-mass distance of the first few, say 4 to 5 monomers from the membrane provides an excellent proxy for $\Phi$. In this paper we follow the same line. The average distance $\langle Z^{(4)}(t = 0) \rangle$ is plotted as a function of the chain tension $\Phi(E, t = 0)$ for various values of $E$ in figure 2. This figure shows that under an applied field, $\Phi(E, t = 0)$ is a reasonably linear function well-proxied by $Z^{(4)}$. The positive curvature seen in figure 2, i.e., the deviation from linearity, is seen only for $E > 0$. We believe that this is partly due to the saturation of $Z^{(4)}$. (By definition, in our lattice model the distance of the center-of-mass of the first 4 monomers from the membrane cannot exceed $(1 + 2 + 3 + 4)/4 = 2.5$.)

3.2. Memory effects in the chain tension

From equation (2), the behavior of the memory kernel $\mu_R(t)$ for the polymer segment on the right side of the membrane can be obtained with a sudden injection of $p$ extra monomers through the pore, corresponding to an impulse current $v(t) = p \delta(t)$. Physically, $v(t) = p \delta(t)$ with $p > 0$ (resp. $p < 0$) means that we tether a polymer of length $N$ halfway at the pore at $t \rightarrow -\infty$, let it thermalize until $t = 0$, and then inject $p$ extra monomers at the tethered end of the right (resp. left) segment at $t = 0$. We then ask for the time-evolution of the mean response $\langle \delta \Phi_R(t) \rangle$, where $\delta \Phi_R(t)$ is the shift in chemical potential for the right segment of the polymer at the pore. This means that for the translocation problem (with both right and left segments), we would have $\Phi(t) = \delta \Phi_R(t) - \delta \Phi_L(t)$, where $\delta \Phi_L(t)$ is the shift in chemical potential for the left segment at the pore due to an opposite input current to it.

In earlier works [13, 17], using $v(t) = p \delta(t)$ for a polymer of length $N$ tethered halfway at the pore as described in the above paragraph, three of us showed that for unbiased polymer translocation, i.e., for $E = 0$, this mean response, and hence $\mu(t)$ takes the form $\mu(t) \sim t^{-\alpha} \exp[-t/\tau_{\text{Rouse}}(N/2)]$ (note that for $E = 0$ there is a trivial symmetry between the right and the left segment of the polymer, hence $\mu_R(t) = \mu_L(t) \equiv \mu(t)$).

When the electric field is applied at the pore, and the same monomer injection method is used to probe the memory kernels $\mu_R(t)$ and $\mu_L(t)$, we expect the above arguments to hold again: since the field is applied very locally at the base of the tethered polymer segments, it does not destroy the broader structure of the polymer. However, we do expect to see deviations from the $t^{-\alpha} \exp[-t/\tau_{\text{Rouse}}(N/2)]$ at short times. Indeed, we have confirmed this picture—for various field
strengths we tracked $\langle \delta \Phi_R(t) \rangle$ and $\langle \delta \Phi_L(t) \rangle$ by measuring the distance of the average center-of-mass of the first 4 monomers from the membrane, $(Z^{(4)}(t))$, in response to the injection of extra monomers near the pore at $t = 0$. Specifically we consider the equilibrated right and left segments of the polymer, each of length $N/2 = 200$ (with the middle monomer threaded at the pore), adding 5 extra monomers at the tethered end of the right and the left segment each at $t = 0$, corresponding to $|p| = 5$, bringing the length of each segment up to $N/2 + |p|$. Using the proxy $\langle Z^{(4)}(t) \rangle$ for both segments we then track $\langle \delta \Phi_R(t) \rangle$ and $\langle \delta \Phi_L(t) \rangle$, denoting them by values $E > 0$ and $E < 0$ respectively in figure 3. The deviations from the expected power-law $t^{-(1+\nu)/(1+2\nu)}$ at short times and the $\text{exp}[-t/\tau_{\text{Rouse}}(N/2)]$ at long times makes the precise identification of the power-law $t^{-(1+\nu)/(1+2\nu)}$ difficult. Nevertheless, there is an extended regime where this power-law can be identified reasonably clearly, yielding us $\mu_R(t) = \mu_R(\nu) \equiv \mu_R \approx t^{-(1+\nu)/(1+2\nu)} \text{exp}[-t/\tau_{\text{Rouse}}(N/2)]$.

4. Scaling behavior of $\tau_d$ with $N$

The memory kernel we obtained in section 3 can be termed as the ‘static memory kernel’, as it is obtained under the condition that before the injection of the extra monomers both segments were thermalized. When the applied field is not too strong, we can expect the static memory kernel to yield the scaling of translocation velocity with time, in the following manner.

An inverse Laplace transform of equation (2) yields us

$$v(t) = \frac{\phi(t)}{k \mu(k)} - \frac{\phi(0)}{\mu(0)},$$

where $k$ is the Laplace variable representing inverse time. Thereafter, using the power-law part of $\mu(t) \sim \frac{1}{t^{\nu + 1}}$, demonstrated that $\phi(\nu) \sim \phi(1)$ reduces to a constant very quickly: $E = 0.05$ (circles), 0.15 (squares), and 0.25 (triangles). To generate these averages 16,000 individual polymers were unthreaded for each value of $E$.

$t^{-(1+\nu)/(1+2\nu)}$, i.e., $\mu(k) \sim k^{(1+\nu)/(1+2\nu)-1}$, and Laplace-inverting equation (5), we get

$$v(t) = \int_0^t \text{d}t' \left( t' \right)^{\nu} \left[ \phi_{t=0} - \phi(t') \right].$$

If $\phi(t)$ goes to a constant $\neq \phi_{t=0}$, then equation (6) reduces to

$$v(t) \sim t^{-1/(2\nu)},$$

where $[s(t) - N/2]$ is the distance unthreaded after time $t$, the $N/2$ appears in equation (7) as $s(0) = N/2$.

In figure 4 we show the behavior of $[\phi_{t=0} - \phi(t)]$ by means of the proxy variable $(Z^{(4)}(0) - Z^{(4)}(t))$, for $E = 0.05, 0.15,$ and 0.25 respectively, where $Z^{(4)}$ is the difference between the $Z^{(4)}$ values of the right and left segment of the polymer, i.e., $Z^{(4)}(t) = Z^{(4)}_R(t) - Z^{(4)}_L(t)$. Indeed the quantity $[\phi_{t=0} - \phi(t)]$ approaches a constant rather quickly. We also note that the relation between this constant and the applied field $E$ is almost linear.

For strong fields, there is no a priori reason that the dynamics can still be described by the static memory kernel instead of a suitably replacing ‘dynamic memory kernel’, but we find that the scaling $s(t) \sim t^{(1+\nu)/(1+2\nu)}$ is obeyed for fairly strong fields as well: in figure 5 we plot the average time $\langle t \rangle$ to unthread a distance $s$ to show this scaling. Note the strong finite-size effects for the scaling behavior as shown by the deviation from the $t^{(1+2\nu)/(1+\nu)}$ for larger values of $s$. The presence of such strong finite-size effects indicates that without the aid of $s(t)$ versus $t$ curves, determining the scaling of $t_d$ with $N$ will almost certainly lead to erroneous identification of the scaling laws—we believe that these finite-size effects are responsible for the wide range of existing numerical scaling.
results, as summarized in table 1. Nevertheless, figure 5 shows that these finite-size effects do not increase linearly with $N$, leading us to the scaling for $t_d$ as

$$t_d \sim N^{(1+2\nu)/(1+\nu)}/E,$$  \hspace{1cm} (8)

which is obtained from the condition that $s(t_d) = N$. For the above analysis to hold, the dwell time must be less than $t_{\text{Rouse}}$, which equation (8) confirms. Note that the $E$-dependence of equation (8) is only numerically obtained from figure 5. Note also that the curves in figure 5 for $E = 0.05$ tend to ‘sag’ a bit. We attribute this to our numerically inspired definition of $s(t)$, as the mean time to unthread a distance $s$, as opposed to, e.g., the numerically less favorable measure of distance $(s(t))$, i.e., the monomer which is most likely to reside in the pore at time $t$. At small fields, the polymer has ample time for fluctuations, pushing the time of first arrival up. Numerically, for $E = 0.15$ and 0.25, the exponent $\partial(\log s(t))/\partial(\log t)$ is found to be 0.73±0.02, in agreement with the theoretical value $(1+\nu)/(1+2\nu)$. The sagging and finite-size effects discussed above cause the apparent exponent to be slightly larger, ranging from 0.74 to 0.79, for $E = 0.05$.

With decreasing field strength, especially in the range where the thermal fluctuations are comparable to the work done by the field to translocate the entire polymer, given by $EN \simeq k_B T = 1$, one should obtain a crossover from the above scaling (8) to $t_d \sim N^{2+\nu}$ for unbiased translocation [13, 17]. This suggests that if $t_d/N^{2+\nu}$ is plotted as a function of $EN$, then one should obtain a scaling collapse; i.e., there exists a scaling function $f$ such that $t_d = N^{2+\nu} f(EN)$. However, $EN$ as a scaling variable is simply numerically inconsistent with figure 5 and equation (8). Instead $t_d = N^{2+\nu} f(E, N)$ is the proper description of the situation, with $f(E, N)$ approaching a constant for $E \to 0$, $N \to \infty$ and $f(E, N)$ behaving as $E^{-1} N^{-\nu-1/(1+\nu)}$ for $E \sim O(1)$ and $N \to \infty$. Note that $E$ in this paragraph should be interpreted as the dimensionless quantity $qV/(k_B T)$.

To demonstrate the scaling behavior of equation (8) for $E \simeq O(1)$, we plot $t_d/N^{2+\nu}$ as a function of $EN^{\nu/(1+\nu)}$ in figure 6. Keeping in mind that this way of plotting the data does not necessarily yield a data collapse at small but nonzero $E$, as discussed above, we also plot several data points for small $E$, in order to demonstrate that for $E \to 0$ our results in this paper are consistent with that of unbiased translocation [13, 17, 18].

5. Discussion

In this paper, we studied polymer translocation in three dimensions through a narrow pore in an otherwise impenetrable membrane, as the polymer is driven by a field $E$ across the pore. The polymer performs Rouse dynamics, i.e., we considered polymer dynamics in the absence of hydrodynamical interactions. We found that the typical time the pore remains blocked during a translocation event, for moderate field strengths scales as $N^{(1+2\nu)/(1+\nu)}$, which is the Flory exponent for the polymer. In line with our previous works, we showed that this scaling behavior stems from the polymer dynamics at the immediate vicinity of the pore—in particular, the memory effects in the polymer chain tension imbalance across the pore [13, 17–19]. We also showed that our results in this paper are consistent with that of unbiased translocation [13, 17, 18] in the limit $E \to 0$.

The above results for finite $E$, along with the numerical results by several other groups, violate the lower bound $N^{1+\nu}/E$ suggested earlier in the literature [15]. We also discussed why this lower bound is incorrect and showed, based on conservation of energy, that the correct lower bound for the pore-blockade time for field-driven translocation is given by $\eta N^{2\nu}/E$, where $\eta$ is the viscosity of the medium surrounding the polymer. Our theoretical analysis has been supported by high precision computer simulation data, generated with a three-dimensional self-avoiding lattice polymer model.
Having worked out the physics of field-driven polymer translocation in the absence of hydrodynamical interactions, it is worthwhile to reflect on the scaling of pore-blockade times as a function of the polymer length $N$ in the presence of hydrodynamical interactions. Hydrodynamical interactions will modify the memory kernel $\mu(t)$—changing it from $t^{-(1+\nu)/(1+2\nu)} \exp(-t/\tau_{\text{Rouse}})$ to $t^{-(1+\nu)/(3\nu)} \exp(-t/\tau_{\text{Zimm}})$ [13, 17], where $\tau_{\text{Zimm}}$ is the Zimm relaxation time, scaling as $N^3\nu$ in good solvent for a polymer of length $N$. This implies that the pore-blockade time will behave as $N^{3\nu/(1+\nu)}$ under the influence of hydrodynamical interactions. In this context we note that the scaling of the pore-blockade time has been experimentally measured to scale as $N^{1.2\pm0.07}$ [11]. In the scaling limit $3\nu/(1+\nu) \simeq 1.11$, the value for $\nu$ suggested in [11] is $0.611 \pm 0.016$, for which $3\nu/(1+\nu) \simeq 1.14 \pm 0.02$, a bit closer to $1.26 \pm 0.07$. For a physical explanation of the scaling of the pore-blockade times with polymer length, the authors of [11] arrived at an answer $2\nu$ using a macroscopic view of the translocating polymer, assuming that the translational velocity of the center-of-mass of the untranslocated part is constant in time, and (implicitly) that the memory kernel is a $\delta$-function. Our analysis in this paper, as well as in [13, 17–19] based on memory effects, therefore, casts serious doubts on the physical interpretation of [11]: as we have repeatedly shown that the velocity of translocation is not uniform in time, and the same part of the polymer visits the pore a multitude number of times. Although so far our work has not incorporated hydrodynamical interactions explicitly, it is difficult to imagine that introducing hydrodynamical interactions will mysteriously wipe out the entire memory effects in the polymer.

Acknowledgments

Ample computer time on the Dutch national supercomputer facility SARA is gratefully acknowledged.

References

[1] Dreiseikelmann B 1994 Microbiol. Rev. 58 293
[2] Henry J P et al 1989 J. Membr. Biol. 112 139
[3] Akimaru J et al 1991 Proc. Natl Acad. Sci. USA 88 6545
[4] Goerlich D and Rappaport T A 1993 Cell 75 615
[5] Schatz G and Dobberstein B 1996 Science 271 1519
[6] Szabó I et al 1997 J. Biol. Chem. 272 25275
[7] Hanss B et al 1998 Proc. Natl Acad. Sci. USA 95 1921
[8] Tseng Y-L et al 2002 Mol. Pharm. 62 864
[9] Nakane J J, Akeson M and Marziali A 2003 J. Phys.: Condens. Matter 15 R1365
[10] Kasianowicz J I et al 1996 Proc. Natl Acad. Sci. USA 93 13770
Henrickson E et al 2000 Phys. Rev. Lett. 85 3057
Meller A et al 2001 Phys. Rev. Lett. 86 3435
Akeson M et al 1999 Biophys. J. 77 3227
Meller A et al 2000 Proc. Natl Acad. Sci. USA 97 1079
Meller A and Branton D 2002 Electrophoresis 23 2583
[11] Storno A J et al 2005 Nano Lett. 5 1193
[12] Szabó I et al 1998 FASEB J. 12 495
Horowka S et al 2001 Proc. Natl Acad. Sci. USA 98 12996
Howorka S, Cheley S and Bayley H 2001 Nat. Biotechnol. 19 636
[13] Panja D, Barkema G T and Ball R C 2006 Preprint cond-mat/0610671
[14] van Heukelum A and Barkema G T 2003 J. Chem. Phys. 119 8197
van Heukelum A et al 2003 Macromolecules 36 6662
Klein Wolterink J et al 2005 Macromolecules 38 2009
Klein Wolterink J and Barkema G T 2005 Mol. Phys. 103 3083
[15] Kantor Y and Kardar M 2004 Phys. Rev. E 69 021806
[16] Chuang J et al 2001 Phys. Rev. E 65 011802
[17] Panja D, Barkema G T and Ball R C 2007 J. Phys.: Condens. Matter 19 432202
[18] Panja D, Barkema G T and Ball R C 2008 J. Phys.: Condens. Matter 20 075101
(Panja D, Barkema G T and Ball R C 2007 Preprint 0710.0147)
[19] Panja D and Barkema G T 2008 Biophys. J. at press
(Panja D and Barkema G T 2007 Preprint 0706.3969)
[20] Luo K et al 2006 J. Chem. Phys. 124 114704
Huopaniemi I et al 2006 J. Chem. Phys. 125 124901
[21] Cacciatore E and Luijten E 2006 Phys. Rev. Lett. 96 238104
[22] Wei D et al 2006 J. Chem. Phys. 126 204901
[23] Milchev A, Binder K and Bhattacharya A 2004 J. Chem. Phys. 121 6042
[24] Dubbeldam J L A et al 2007 Europhys. Lett. 79 18002
[25] de Gennes P-G 1979 Scaling Concepts in Polymer Physics (Ithaca, NY: Cornell University Press)
[26] Metzler R and Klafter J 2003 Biophys. J. 85 2776
[27] Dubbeldam J L A et al 2007 Phys. Rev. E 76 010801(R)