Self and Tracer Diffusion of Polymers in Solution

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The literature on self-diffusion of polymers in solution, and on tracer diffusion of probe polymers through solutions of matrix polymers, is systematically reviewed. Virtually the entirety of the published experimental data on the concentration dependence of polymer self—and probe—diffusion is represented well by the same functional form. This form is the stretched exponential \[ \exp(-\alpha c^\nu) \]
where \( c \) is polymer concentration, \( \alpha \) is a scaling prefactor, and \( \nu \) is a scaling exponent.

Correlations of the scaling parameters with polymer molecular weight, concentration, and size are examined, and compared with predictions based on the form’s hydrodynamic and renormalization-group derivations. \( \alpha \) increases markedly with polymer molecular weight, namely \( \alpha \sim M^{x} \) for \( x \approx 1 \). \( \nu \) is \( \approx 0.5 \) for large polymers (\( M \) larger than 400 kDa or so), but increases toward 1.0 or so at smaller \( M \). Scaling parameters for the diffusion of star polymers do not differ significantly from scaling parameters for the diffusion of linear chains of equal size.

I. INTRODUCTION

The objective of this review is to present the experimental phenomenology for polymer self-diffusion and for diffusion of tracer polymers through polymer solutions. We proceed by examining how the diffusion coefficient depends on polymer concentration \( c \), polymer molecular weight \( M \), solvent quality, and other variables. We examine how rapidly a probe polymer of molecular weight \( P \) diffuses through a matrix polymer of molecular weight \( M \), perhaps as the concentrations of the matrix and probe polymers are varied. After establishing the phenomenological behavior, we compare the phenomenology with various classes of theoretical model.

This is not a review of the extremely extensive theoretical literature on polymer diffusion in solution. For such reviews, note works of Graessley, Tirrell, Pearson, Skolnick, Lodge, and (more recently but less directly) McLeish. Recent papers by Schweizer and collaborators include extensive references to the more recent literature. We will note classes of models, but not their underlying derivations. We also do not consider melt systems, or polymer mutual diffusion.

In comparing phenomenology with model predictions, it will remain critical to distinguish between properties that are consistent with a particular model, but do not actually prove it, and properties that require or refute the correctness of a particular model. For example, Skolnick has shown that the nominal signature of reptation \( D_s \sim M^{-2} \) is found in computer models of polymers, in a model system in which the chains are very certainly not reptating. A finding \( D_s \sim M^{-2} \) is thus consistent with many tube-type solution models, but does not prove their correctness. On the other hand, many tube-type models require that polymer chains move via constrained diffusion for shorter times. In these models, the shorter-time mean-square displacement of a single chain must scale more slowly than linearly with time. If the shorter-time mean-square displacement were instead linear in elapsed time, many constrained-diffusion models would be rejected by experiment.

Section II presents a theoretical background for the experimental papers reviewed here. Sections III and IV treat, respectively, data on (i) self-diffusion coefficients of polymers of solution and (ii) data on probe diffusion coefficients in polymer matrix solutions. Section V briefly remarks upon other papers on polymer self- and probe-diffusion that do not lend themselves to the analytic approach applied here. A systematic study of the phenomenological parameters obtained by our analysis appears in Section VI. Section VII summarizes conclusions. Tables of fitting parameters appear as Appendices.

II. THEORETICAL BACKGROUND

This Section presents a consistent nomenclature for polymer diffusion coefficients and describes methods whereby which such diffusion coefficients are measured. A short description is given of the literature on diffusion in multimacrocomponent solutions. The literature in question provides the fundamental basis for interpreting the experimental measurements. Finally, phenomenological classes of models for polymer dynamics are identified.

A. Nomenclature for Diffusion Coefficients

This section sets out a consistent nomenclature for the diffusion coefficients that we are reviewing. In a solution containing a solvent and one macromolecular species, two physically distinct diffusion coefficients usefully characterize macromolecule diffusion. One of these, the two-particle or mutual diffusion coefficient \( D_m \), describes via Fick’s law

\[ \vec{J} = D_m \vec{\nabla} c \] (1)

the relaxation of a concentration gradient. Here \( \vec{J} \) is the diffusion current and \( c \) is the local instantaneous macromolecule concentration. The other diffusion coefficient, the single-particle or self diffusion coefficient \( D_s \), describes the diffusion of a single macromolecule through a
uniform solution of elsewise identical macromolecules. In a simple macromolecule solution, quasi-elastic light scattering spectroscopy (QELSS) measures the mutual diffusion coefficient of the macromolecules. Pulsed-field-gradient nuclear magnetic resonance (PFGNMR) measures the self-diffusion coefficient.

Solutions containing a solvent and two macromolecule species show a more complicated diffusive behavior. Theoretical treatments of this issue are described in the next section. In general, in two-macrocomponent solutions the temporal evolution of transient concentration fluctuations is described by two relaxation times, each describing the relaxation of a coupled mode involving the concentrations of both macromolecule species. An interesting special case arises if one macrocomponent, the probe, is adequately dilute, while the other macrocomponent, the matrix, may be either dilute or concentrated. In this special case, the diffusion of the probe species is governed by a single-particle diffusion coefficient, namely the probe diffusion coefficient $D_p$. In the special case that the matrix species and solvent are isorefractive, so that the index of refraction is independent of the relative amounts of matrix polymer and solvent, the matrix species scatters next to no light. If in this special case light scattering by a dilute probe species dominates scattering by the solution (which further requires that solvent scattering be sufficiently weak), the diffusion coefficient measured by QELSS is the probe diffusion coefficient $D_p$.

A variety of physical techniques have been used to measure probe diffusion through a polymer solution. In the fluorescence recovery after photobleaching (FRAP) technique, a small fluorescent label is attached to the probes. An intense pulse of light (the ‘pump’ laser beam) is then used to destroy (“bleach”) the fluorescent labels in some regions of the solution, the regions sometimes being defined via a holographic grating. Much weaker laser illumination (the ‘probe’ beam) is then used to monitor the recovery of the fluorescence intensity as unbleached fluorescent molecules diffuse back into the regions(s) in which the bleaching occurred.

An alternative to FRAP is Forced Rayleigh Scattering (FRS). In an FRS experiment, an intense laser pulse is used to generate a holographic brightness grating in solution. The grating selectively alters a photosensitive part of the probe molecule, thereby creating an index of refraction grating in the solution. A much weaker probe beam is then used to monitor the diffusive relaxation of the induced grating. Significant complications may arise if the photomodified and non-photomodified forms of the probe species differ significantly in their diffusive properties. If the probe species is dilute, with either FRAP or FRS the time dependence of the recovery profile is determined by the single-particle (probe-) diffusion coefficient of the probe molecules.

A physically-distinct alternative to FRAP and FRS is fluorescence correlation spectroscopy (FCS), in which optical methods are used to measure fluctuations in the number of fluorescent molecules in a small volume of space. In nondilute solution the diffusion coefficient measured by fluorescence correlation spectroscopy (FCS) varies from $D_p$ to $D_m$ as the fraction of macromolecules that bear fluorescent tags is varied from small to large.

When is a probe species dilute? In general, the careful experimenter makes an adequate control study of the effect of varying the probe concentration $c_p$. $D_p$ is the extrapolation of the measured probe diffusion coefficient to zero probe concentration. In some systems $D_p$ is substantially independent of $c_p$, and no extrapolation is needed. In other systems (see below) $D_p$ depends significantly on $c_p$, and extrapolation to $c_p \to 0$ must be performed. Several authors have measured the initial slope $k_d$ of the dependence of $D_p$ on probe concentration, and the effect of matrix concentration on $k_d$. It is also possible to study—most published work has used QELSS—the relaxation of concentration fluctuations in a ternary solvent:macromolecule:macromolecule system in which neither macrocomponent is dilute. As reviewed below, such studies give information on the diagonal—and cross-diffusion coefficients of the two species and on their thermodynamic interactions.

Strictly speaking, if FRAP or FRS is used to study the diffusion of a dilute labelled polymer through a matrix polymer solution, the labelled polymer differing from the matrix in that a fluorescent small-molecule label has been attached, the measured diffusion coefficient should be termed a probe diffusion coefficient, because the molecules being tracked are chemically distinguishable from the molecules not being tracked. However, for the purposes of this review, we distinguish between (i) the case in which the probe and matrix polymers are substantially distinct in molecular weight or chemical nature, and (ii) the case in which the probe and matrix polymers are very nearly the same except for the presence of the label. The phrase ‘probe diffusion coefficient’ is reserved for case (i), while measurements under case (ii) are here treated as determinations of the self diffusion coefficient.

The term tracer diffusion coefficient is synonymous with single-particle diffusion coefficient, and includes both the self and probe diffusion coefficients as special cases. The interdiffusion and cooperative diffusion coefficients characterize the relaxation times in a ternary system in which neither macrocomponent is dilute. In the following the latter two terms are almost never used.

**B. General Theory of Diffusion in Multimacrocomponent Solutions**

What is the basis for using spectroscopic methods such as QELSS, FRAP, FRS, or FCS to measure $D_s$ or $D_p$? In each case, there is a physical theory that links the correlation function determined spectroscopically to fluctuations in microscopic variables that describe the liquid. For example, for FCS the fluctuations in the fluorescent light intensity $I_f(t)$ arise from fluctuations in the number of fluorescent molecules in the scattering volume. The
time correlation function $C_f(\tau) = \langle I_f(t)I_f(t+\tau) \rangle$ of the fluorescent intensity follows faithfully the time correlation function $C_N(\tau) = \langle N(t)N(t+\tau) \rangle$ of the (fluctuating) number $N(t)$ of labelled molecules in the scattering volume, so to determine the time dependence of $C_f(\tau)$ one only needs to determine $C_N(\tau)$ and vice versa.

For QELSS, the spectrum is determined by the field correlation function $g^{(1)}(q,t)$ of the scattered light, which is in turn determined by the motion of the scattering molecules via

$$g^{(1)}(q,\tau) = \left\langle \sum_{i,j=1}^{N} \alpha_i\alpha_j \exp[iq \cdot (r_i(t+\tau) - r_j(t))] \right\rangle,$$

where $i$ and $j$ label two of the $N$ scatterers, $r_i(t)$ is the position of scatterer $i$ at time $t$, $\alpha_i$ is the scattering cross-section of particle $j$, and $q$ is the scattering vector. For the special case in which the scattering particles are dilute (non-scattering particles do not need to be dilute), correlations between the positions of two distinct scatterers at two times vanish. The correlation function for probe scattering then reduces to

$$g_p^{(1)}(q,\tau) = \left\langle \sum_{i=1}^{N} \alpha_i^2 \exp[iq \cdot \Delta r_i(\tau)] \right\rangle,$$

in which $\Delta r_i(\tau) = r_i(t+\tau) - r_i(t)$.

There are three general approaches to calculating $g_p^{(1)}(q,\tau)$. First, scatterers can be treated as objects having hydrodynamic and direct, e.g., excluded volume, interactions, and the effect of the intermacromolecular interactions on $dg^{(1)}_p(q,\tau)/d\tau$ can be calculated. Second, it can be recognized that $\sum_i^N \exp[iq \cdot (r_i(t))]$ is the $q^{th}$ spatial Fourier component of the scatterer concentration at time $t$, and semicontinuum hydrodynamic methods and the Onsager regression hypothesis can be used calculate the average temporal evolution of a fluctuation. Third, it can be proposed that the scatterers perform simple Brownian motion, with the probability distribution function for $\Delta r_i(\tau)$ being applied to calculate $g_p^{(1)}(q,\tau)$.

Light scattering spectra of non-ideal single-component macromolecule solutions, including direct interactions and hydrodynamic interactions at the Oseen level, were initially calculated by Altenberger and Deutsch. Calculations of light scattering spectra of non-dilute many-component macromolecule solutions soon followed. In the latter papers, diffusion coefficients were modeled on the lines of Kirkwood, et al., the diffusion coefficients being written as products of thermodynamic derivatives ($\partial \mu_i/\partial c_j$) obtained from the intermacromolecular forces, and phenomenological transport coefficients $\Omega_{ij}$ (here $i$ and $j$ label chemical species). If neither macromolecular species was dilute, the light scattering spectrum obtained by QELSS is predicted to contain two relaxation modes. Even if only one of the two macromolecule species scattered light, both modes are predicted to be visible, in general, in the light scattering spectrum.

A subsequent paper extended these findings to the special case explored e.g., by fluorescence correlation spectroscopy, in which some macromolecules are tagged so that their movements can be tracked. For the case that the tagged macromolecules are dilute, it was shown that fluorescence correlation spectroscopy and other equivalent techniques measure the self-diffusion coefficient of the tagged probe macromolecules diffusing through the unlabelled matrix macromolecules.

A Smoluchowski approach was used by Jones to examine interacting spherical polymers, including the special case that one of the polymer species is dilute and tagged so that its motions could be observed. In the special case, the measured diffusion coefficient was predicted by Jones to be determined only by the hydrodynamic interactions of the tracer polymers with their (predominantly matrix-polymer) neighbors. While Jones does not use the same nomenclature, the diffusion coefficient of the tracers calculated in ref. may be recognized as the single-particle diffusion coefficient. The hydrodynamic approach culminates in analyses of Carter, et al., and Phillips of mutual and tracer diffusion coefficients, including full hydrodynamic and direct interactions between interacting diffusing particles and reference frame issues.

Extensive studies of diffusion of interacting polymers using semicontinuum and related means were made by Akcasu, Bemmouna, Cohen and others. In 1987, Bemmouna, et al. calculated dynamic scattering from a solution containing two polymer species, including excluded volume terms with a Flory interaction parameter but ignoring hydrodynamic interactions between polymer chains. Hydrodynamic interactions were neglected in the sense that cross-species transport coefficients were assumed to vanish. The assumption that the hydrodynamic parts $\Omega_{ij}$ of the self-diffusion tensor vanish cannot be true simultaneously in the solvent- and volume-fixed reference frames. Polymer solutions can be very concentrated, so neglect of reference frame issues in entirely correct calculations may lead to complications in the physical interpretation of an assumption that hydrodynamics has been neglected.

Bemmouna, et al. examined the special case of two species identical except for their optical scattering cross-sections, showing results consistent with those of Phillips. notably: If the matrix species scatters weakly, an interdiffusion mode describing single-chain motion is dominant at low probe concentrations. There is a thermodynamic regime relative to the spinodal where the diffusion equations describe phase separation. The Bemmouna model was then extended to treat solutions of copolymers, still with neglect of hydrodynamic interactions, and to treat homopolymer: copolymer mixtures.

In the same period, Foley and Cohen analyzed concentration fluctuations in polymer: polymer solvent mixtures, using an ornate Flory-Huggins form for the thermodynamic free energy of the mixture, again neglecting interchain hydrodynamic interactions. The case of a
solvent isorefractive with the matrix polymer, in the presence of a dilute scattering species, was treated. Foley and Cohen also examined systems in which both polymer species were non-dilute, predicting that in this case the relaxation spectrum is characterized by three distinct relaxation times. In contrast, other calculations on similar models predict only two relaxation times.

Roby and Joanny\textsuperscript{21} improved the model of Benmouna, et al.\textsuperscript{21} by incorporating interchain hydrodynamic interactions and by improving the model for direct chain-chain interactions. At elevated concentrations, the correctness of the reptation model was assumed. The effect of reptation dynamics on concentration fluctuations was estimated with the approximation that a solution is effectively a polymer melt in which mesoscopic polymer-solvent blobs play the role taken in polymer melts by monomer units. For simplicity, the model was restricted to systems containing equal amounts of two species having the same molecular weight. These restrictions are non-trivial to remove but exclude analysis of tagged-tracer experiments. A model calculation incorporating a similar picture, but without the restrictions, was reported by Hammouda et al.\textsuperscript{26}

An alternative analysis of a ternary solution of two polymers and a solvent was presented by Wang\textsuperscript{27}, who followed much of the earlier work on polymer solutions in assuming that the cross terms in the mobility matrix vanish. Wang systematically analysed a variety of general and special cases, showing a range of interesting parameters that can be determined from light scattering spectra if the accuracy of his model is assumed. In addition to the tracer case (one polymer:solvent pair isorefractive, visible polymer dilute), Wang analysed the special case “zero average contrast”, in which the polymer refractive increments are of opposite sign, so that concentration fluctuations, that change the total polymer concentration without changing the local polymer composition, scatter no light. Wang showed that the QELS spectrum of a zero-average-contrast system is almost always bimodal, though it may happen that one of the modes is much weaker than the other.

Treatments of the field correlation function based on a Brownian motion description can be traced back to Berne and Pecora\textsuperscript{28}, who treat the light scattering spectrum of a solution of dilute, \textit{noninteracting} Brownian particles. The motion of such particles is described as a series of random, uncorrelated steps, in which case from the Central Limit Theorem the probability distribution for particle displacements is

\[ G_s(\Delta R, t) = \left[ \frac{2\pi}{3} \langle (\Delta R)^2 \rangle \right]^{-3/2} \exp \left[ -3(\Delta R)^2 / 2 \langle (\Delta R)^2 \rangle \right], \tag{4} \]

where the mean-square particle displacement is related to the diffusion coefficient by

\[ \langle (\Delta R)^2 \rangle = 6Dt. \tag{5} \]

Combining eqs\textsuperscript{2} and \textsuperscript{4},

\[ g_p^{(1)}(q, \tau) \sim \int d\Delta r \exp[iq \cdot \Delta r] \times \left[ \frac{2\pi}{3} \langle (\Delta R)^2 \rangle \right]^{-3/2} \exp \left[ -3(\Delta R)^2 / 2 \langle (\Delta R)^2 \rangle \right], \tag{6} \]

which using eq\textsuperscript{5} leads to

\[ g_p^{(1)}(q, \tau) \sim \exp(-q^2D\tau). \tag{7} \]

For Brownian particles in low-viscosity small-molecule solvents, the Stokes-Einstein equation

\[ D = \frac{k_B T}{6\pi\eta R}, \tag{8} \]

in which \( k_B \) is Boltzmann’s constant, \( T \) is the absolute temperature, \( \eta \) is the solution viscosity, and \( R \) is the particle radius, generally predicts accurately the particle diffusion coefficient. For diffusion in polymer solutions, two \textit{ad hoc} extensions of this form are encountered:

First, it might be the case that \( D \) is not predicted accurately by eq\textsuperscript{8}. In this case, one could formally define a microviscosity \( \eta_\mu \) as

\[ \eta_\mu = \frac{k_B T}{6\pi D R}, \tag{9} \]

and compares \( \eta_\mu \) with the macroscopically-measured viscosities \( \eta \) of the solution and \( \eta_\mu \) of the solvent. The microviscosity is more commonly applied to describe diffusion of mesoscopic globular probe particles,\textsuperscript{28} rather than to treat the diffusion of random-coil polymers in solution.

Second, light scattering and other relaxation spectra are not always single exponentials. No matter what functional form \( g_p^{(1)}(q, \tau) \) has, one may formally write

\[ g_p^{(1)}(q, \tau) \sim \exp(-q^2D(\tau)\tau) \tag{10} \]

as the \textit{definition} of \( D(\tau) \). In this equation, \( D(\tau) \) formally appears to be a time-dependent diffusion coefficient, which equally formally defines a frequency-dependent diffusion coefficient such as

\[ D(\omega) = \int_0^\infty d\tau \exp(-i\omega\tau)D(\tau), \tag{11} \]

and via several slightly different paths a frequency-dependent microviscosity

\[ \eta_\mu(\omega) = \frac{k_B T}{6\pi D(\omega)R}. \tag{12} \]

The real and imaginary parts of \( \eta_\mu(\omega) \) can be brought into correspondence with the storage and loss moduli \( G'(\omega) \) and \( G''(\omega) \).

The second extension has serious physical difficulties. In particular, comparing eqs\textsuperscript{6} and \textsuperscript{7} and noting from
reflection symmetry that terms odd in \( q \) vanish, one finds that the extension assumes that

\[
\langle \exp[-q^2(\Delta r)^2] \rangle = \exp[-q^2 \langle (\Delta r)^2 \rangle]. \tag{13}
\]

Equation 13 would be correct if sequential random changes in \( \sum_{j=1}^{N} \exp[iq \cdot r_j] \) were described by a Gaussian random process, because in that case (and only in that case) the average on the rhs of eq 13 would be entirely described by the mean-square average displacement via the expression \( \langle (\Delta r)^2 \rangle \) seen on the rhs of this equation. Brownian motion, in which particle displacements are described by the Langevin equation, is an example of a dynamics that generates a Gaussian random process for which eq 13 is correct. For extensive details, see ref 14.

Omitted from, but critical to, the above discussion are the consequences of Doob’s First Theorem. Doob treated the joint expectation value—what we would now call the correlation function—of random variables including those following Langevin dynamics. If eq 13 is correct, it is an inescapable consequence of Doob’s Theorem that the relaxation spectrum is a single exponential. Conversely, if the spectrum is not a single exponential, then from Doob’s theorem

\[
g^{(1)}(q, \tau) \neq \exp[-q^2 \langle (\Delta r)^2 \rangle]. \tag{14}
\]

Berne and Pecora's text is sometimes incorrectly cited as asserting that eq 13 is uniformly correct for light-scattering spectra. The analysis in Berne and Pecora, which correctly obtains eq 13, refers only to a special-case system. In the system that these authors correctly analyzed, particle displacements are governed by the simple Langevin equation, and particle displacements in successive moments are uncorrelated. That is, this analysis refers to systems in which particle motion is observed only over times much longer than any viscoelastic relaxation times.

If particle motion were observed over times shorter than the viscoelastic relaxation times, which in a concentrated polymer solution might be 1 second or more, particle displacements in successive moments would be correlated. Equation 13 would not be correct. \( \log[g^{(1)}(q, \tau)]/q^2 \) would not be proportional to the mean-square particle displacement during \( \tau \). The path from eq 2 to eq 12 might be an interesting heuristic, but would not be consistent with the properties of particles executing Brownian motion. Of course, with respect to polymer dynamics the interest in eq 10–13 has been exactly the study of particle motions at short times, during which viscoelastic effects are apparent in \( D(\tau) \), but these are precisely the circumstances under which eq 13 is incorrect.

In addition to the above, there are analyses of melt systems such as the very interesting work of Akcasu and collaborators. Melt systems are not considered here.

### C. Phenomenological Forms for

#### Comparison with Experiment

The approach here is to compare experimental measurements of \( D_s \) and \( D_p \) with the functional forms and \( c, P, c \) dependences predicted by various treatments of polymer dynamics. There are a very large number of proposed models. Most models fall into two major phenomenological classes, distinguished by the functional forms taken to describe \( D_s(c) \). This section sketches predictions of classes of model, not including their underlying physical rationales, in preparation for the comparison.

1. In scaling-law models, the relationship between \( D_s, D_p \), and polymer properties is described by scaling laws such as

\[
D_s = D_1 M^{\gamma} c^{-x}, \tag{15}
\]

where here \( \gamma \) and \( x \) are scaling exponents, and \( D_1 \) is a scaling prefactor, namely the nominal diffusion coefficient at unit molecular weight and concentration. In some cases, scaling laws are proposed to be true only over some range of their variables, or only to be true asymptotically in some limit. On moving away from the limit, corrections to scaling then arise. For melts, some models derive a scaling law for \( D_s(M) \) from model dynamics, and then predict numerical values for \( \gamma \). (In melts, \( D_s \) has no concentration dependence because \( c \) is constant.) For polymer solutions, more typically a scaling-law form is postulated. The theoretical objective is then to calculate the exponents \( \gamma \) and \( x \).

Many scaling-type models propose a transition in solution behavior between a lower-concentration dilute regime and a higher-concentration semidilute regime. Scaling arguments do not usually supply numerical coefficients, so there is no guarantee that an interesting transition actually occurs at unit value of a predicted transition concentration \( c_t \) rather than at, say, \( 2c_t \). Correspondingly, the observation that a transition is found at \( 2c_t \) rather than \( c_t \) is generally in no sense a disproof of a scaling model, because in most cases scaling models do not supply numerical prefactors adequate to make a disproof. (Some level of rationality must be preserved. If a physical model leads to \( c_t \) as the transition concentration, and the nominally corresponding transition is only observed for \( 30–150 c_t \), and then only in some systems, one is entitled to question if the observed transition corresponds to the transition implied by the model.)

Two transition concentrations are often identified in the literature. The first transition concentration is the overlap concentration \( c^* \), formally defined as the concentration \( c^* = N/V \) at which \( 4\pi R_g^2 N/(3V) = 1 \). Here \( N \) is the number of macromolecules in a solution having volume \( V \) and \( R_g \) is the macromolecule radius of gyration. In many cases, \( c^* \) is obtained from the intrinsic viscosity via \( c^* = n/[\eta] \) for some \( n \) in the range 1–4. The second transition concentration is the entanglement concentration \( c_e \). In some papers, the entanglement concentration is obtained from a log-log plot of viscosity
against concentration by extrapolating an assumed low-concentration linear behavior and an assumed higher-concentration power-law behavior \((c^a \text{ for, e.g., } x = 4)\) to an intermediate concentration at which the two forms predict the same viscosity; this intermediate concentration is taken to be \(c_c\). In other papers, the entanglement concentration is inferred from the behavior of the viscoelastic moduli; in particular, an onset of viscous recovery in the melt or solution is taken to mark \(c_c\).

(2) In exponential models\(^{33,34}\), the concentration dependence of \(D_s\) becomes an exponential or stretched exponential in concentration

\[
D_s = D_o \exp(-a c^\nu).
\] (16)

Here \(D_o\) is the diffusion coefficient in the limit of infinite dilution of the polymer, \(a\) is a scaling prefactor, and \(\nu\) is a scaling exponent; \(\nu = 1\) for simple exponentials. Under the circumstance that the probe and matrix molecular weights \(P\) and \(M\) differ, an elaborated form of the stretched exponential is

\[
D_p = D_o P^{-\alpha} \exp(-a c^\nu P^\gamma M^\delta),
\] (17)

where \(\gamma, a,\) and \(\delta\) are additional scaling exponents, \(D_o\) now represents the diffusion coefficient in the limit of zero matrix concentration of a hypothetical probe polymer having unit molecular weight, and \(P^{-\alpha}\) describes the dependence, on probe molecular weight, of the diffusion coefficient of a dilute probe molecule. On setting \(a = \gamma = 0\) and freeing \(D_o\), eq. 17 becomes a parameterization of the matrix molecular weight dependence of \(D_p\) for a fixed probe molecular weight.

In the derivations\(^{33,34,35,36}\) of the stretched-exponential models, functional forms and numerical values for exponents and pre-factors were both obtained, subject to various approximations. The latter two derivations assume that chain motion is adequately approximated by whole-body translation and rotation, which may be appropriate if \(P \approx M\), but which is not obviously appropriate if \(P\) and \(M\) are substantially unequal, since in this case whole-body motion of one chain and local modes of the other chain occur over the same distance scale.

Some exponential models\(^{37,38}\) also include a transition concentration, namely a transition between a lower-concentration regime in which some transport coefficients show stretched-exponential concentration dependences and a higher-concentration regime in which the same transport coefficients show power-law concentration dependences. The transition concentration, which experimentally is sharply defined\(^{37,38}\), is here denoted \(c^+\). The lower-concentration regime is the solution-like regime; the higher-concentration regime is the melt-like regime. Such transitions are seen in some but not all viscosity data\(^{37}\); generally but not always\(^{38}\) at very high concentrations \(c[\eta] > 35\). A solution-like-melt-like transition is very rarely apparent in measurements of \(D_s\) or \(D_p\).

There has been interest in derivations of concentration and other dependences of transport coefficients from renormalization group approaches. Power-law and exponential forms can\(^{39}\) both follow from a renormalization-group approach, depending on the location of the supporting fixed point. The stretched-exponential form is\(^{39}\) an invariant of the Altenberger-Dahler\(^{40}\) Positive-Function Renormalization Group.

Our analysis will examine whether either of these functions describe experiment. While a power law and a stretched exponential both can represent a range of measurements, on a log-log plot a power law is always seen as a straight line, while a stretched exponential is always seen as smooth curve of nonzero curvature. Neither form can fit well data that is described well by the other form, except in the sense that in real measurements with experimental scatter a data set that is described well by either function is tangentially approximated over a narrow region by the other function.

It would also be possible to divide theories of polymer dynamics into classes based on assumptions as to the nature of the dominant forces in solution.Assertions as to the dominant forces are independent of assertions as to the functional form used to describe \(D_s(c)\). The major forces common to all polymer solutions are the excluded-volume force that prevents polymer chains from interpenetrating each other, and the hydrodynamic force that creates correlations in the displacements of nearby chains. In some models, excluded-volume forces (topological constraints) are assumed to dominate, hydrodynamic interactions serving primarily to dress the monomer diffusion coefficient. In other models, hydrodynamic interactions between nearby chains are assumed to dominate, while chain-crossing constraints provide at most secondary corrections.

In addition to the generic forces, chains have chemically-specific interactions including in various cases van der Waals, hydrogen bonding, and electrostatic forces. These interactions substantially modulate the properties of particular polymers. However, diffusion coefficients and viscoelastic parameters of most neutral polymers show highly-characteristic “polymeric” behaviors, almost independent of the chemical identity of the polymer, implying that general polymer properties do not arise from chemically-specific interactions.

In the following, eqs. 16-17 will be systematically compared with the literature on \(D_s\) and \(D_p\). The following largely treats diffusion by neutral polymers in good and theta solvents. There is rather little data on self-diffusion of random coil polyelectrolytes. The measurements reviewed here determine diffusion coefficients, not the physical nature of intermolecular forces, so our emphasis is on identifying the class of model, not the type of force, that is significant for solutions.

Comparisons were made via non-linear least-squares fits. The quantity minimized by the fitting algorithm was the mean-square difference between the data and the fitting function, expressed as a fraction of the value of the fitting function. This quantity is the appropriate choice for minimization if the error in the measurement
is some constant fraction of the value of the quantity being measured, e.g., if regardless of the value of \(D_s\), \(D_s\) was measured to within 1%. In some cases, one or more potentially free parameters were held constant (“frozen”) during the fitting process. For each fit, the corresponding Table reports the final fitting error.

III. SELF-DIFFUSION

This Section presents measurements of the true self-diffusion coefficient, which describes the motion of a labelled chain through a solution of elsewise identical chains. Measurements were primarily made with pulsed-field gradient nuclear magnetic resonance (PFGNMR) and forced Rayleigh scattering (FRS). Results are presented alphabetically by first author, showing for each paper the data and fits to stretched-exponential forms. Fitting parameters appear in Table I. A more detailed analysis of the fitting coefficients appears in Section VI. Whenever possible, polymer concentrations have been converted to grams/liter.

Brown, et al.\(^{41}\) report self-diffusion and sedimentation coefficients for dextran (\(M_n = 44\) kDa, \(M_w = 64.2\) kDa) in water, using PFGNMR to determine \(D_s\). They report \(D_s\) and \(s\) as functions of \(c\) for concentrations as large as 250 g/L. Figures 1 shows their data and the corresponding stretched-exponential fit. The self-diffusion data are described accurately by the stretched-exponential form.

Brown and the same collaborators\(^{42}\) used PFGNMR to measure \(D_s\) of of narrow (\(M_w/M_n\) of 1.02-1.20) polyethylene oxides in water. Polymer molecular weights were in the range 73-661 kDa; polymer concentrations ranged up to 70 g/L. The same paper reports measurements on these systems of the mutual diffusion coefficient (from QELSS) and the sedimentation coefficient. Figure 2 shows the measurements of \(D_s\). Because measurements were only reported over a limited concentration range, the data
were fit both to a pure ($\nu = 1$) and to a stretched exponential in $c$. There is excellent agreement between $D_s$ and a pure exponential in $c$, and almost no improvement in the quality of the fit attendant to allowing $\nu$ to be a floating parameter in the fit.

Callaghan and Pinder\textsuperscript{43,44,45} used PFGNMR to study $D_s$ of 2, 110, 233, and 350 kDa polystyrenes with $M_w/M_n$ in the range 1.06-1.10. Polymer concentrations were as large as several hundred g/L. The solvents were CCl\textsubscript{4} and hexadeuterobenzene. In CCl\textsubscript{4} concentrations well into the dilute regime were observed; in C\textsubscript{6}D\textsubscript{6} the measurements of $D_s$ were not extended to low concentration. The data and corresponding stretched-exponential fits appear in figs. 3 and 4. For each polymer:solvent pair, a stretched exponential with constant parameters describes $D_s(c)$ well all the way from the lowest to the largest concentrations studied. The prefactor $\alpha$ tends to increase with increasing polymer $M$, while $D_o$ and the exponent $\nu$ generally decrease with increasing polymer $M$.

Deschamps, et al.\textsuperscript{46} used FRS to study self-diffusion of polystyrenes in cyclopentane in the vicinity of the theta point. The polymer molecular weights were 262, 657, and 861 kDa; polymer concentrations ranged from 1 to 240 g/L. The polymer polydispersity was in the range $M_w/M_n \approx 1.1 - 1.3$. Figure 5 shows their data and fits to stretched exponentials. For the 657 kDa polymer, $D_s$ was only reported over a narrow, elevated concentration range. For the 657 kDa polymer, the stretched-exponential fit was therefore made by interpolating $D_o$ and $\nu$ from their values for the 262 and 861 kDa polymers, leaving $\alpha$ as the only free parameter. For all three polymer molecular weights, the stretched-exponential forms are in good agreement with experiment. The stretched-exponential form for $D_s(c)$ remains valid after a change from good to near-theta solvent conditions.

As an aside, the data of Deschamps, et al.\textsuperscript{46} illustrate well the principle that an experimental test of a par-
ticular theoretical model is sometimes not optimal as a test for a different theoretical model. Deschamps and Leger’s objective was to search for deGennes-type scaling behavior of polymer self-diffusion in a theta solvent. Scaling behavior is only expected in the semi-dilute concentration regime $c > c^*$. In Deschamps, et al.’s systems, the semi-dilute regime was expected to be found only for $c > 50 \text{–} 100 \text{ g/L}$. In the context of their objective of studying scaling behavior, there was no strong reason for Deschamps, et al. to measure $D_s$ at lower concentrations, so they rationally did not do so for the 657 kDa system. However, an accurate stretched-exponential fit requires measurements of $D_s$ at small as well as large concentrations. Through no fault of the original authors, the range of concentrations studied for the 657 kDa polymer restrains the utility of the fits that can be made to some of their data.

Fleischer$^{47}$ used PFGNMR to observe self-diffusion of 125 kDa polystyrene, $M_w/M_n \approx 1.02$, in toluene for concentrations 80-320 g/L. These concentrations were estimated to cover $0.5c_e \leq c \leq 2c_e$, where $c_e$ is the concentration above which entanglements were said to be present. $c_e$ was estimated by applying a deGennes-type model to rheological data$^{46}$. Fleischer’s measurement appear in Fig. 6. As seen in the Figure, the measured $D_s(c)$ is in good agreement with a stretched-exponential form.

Fleischer’s PFGNMR measurements$^{46}$ of the incoherent dynamic structure factor $S_{inc}(q, t)$ of this system show only a single fast relaxation, even under conditions under which QELSS reveals that a slow mode dominates the QELSS spectrum. Fleischer’s observation that the QELSS slow mode cannot be seen in the PFGNMR data is immediately reminiscent of results of Zero and Ware$^{49}$ on fluorescence recovery after photobleaching in poly-L-lysine solutions at low salt concentration. In each set of results, the QELSS spectrum shows distinct slow and fast modes. The slow mode becomes dominant under conditions (higher polymer concentration, lower ionic strength) that enhance ‘glassy’ behavior. In each case, the appearance and then dominance of the slow mode, when ‘glassy’ behavior is enhanced, has no effect on the self-diffusion of individual chains. This behavior is hard to understand if the QELSS slow mode is interpreted as arising from long-lived chain clusters with a fixed list of members. The behavior is, however, understandable in terms of a slow mode arising from the appearance of long-lived dynamic structures within which individual chains only have short residence times. Such dynamic properties of $D_s$ and $D_m$ for interacting interpenetrating particles are also seen in Johnson et al.’s$^{50, 51}$ model glasses.

Giebel, et al.$^{52}$ report $D_s$ of 15, 530, and 730 kDa polydimethylsiloxane in toluene as obtained with PFGNMR, based in some part on data of Skirda, et al.$^{53}$. The original measurements cover the concentration range 2-900 g/L of polymer. Fitting parameters are in Table I. Figure 7 shows the actual data and their fits. For each polymer molecular weight, $D_s$ is described well by a stretched exponential in concentration.

Hadgraft, et al.$^{54}$ used QELSS to measure the diffusion of polystyrenes in benzene as a function of molecular weight for $24.8 \leq M \leq 8870$ kDa at 25 C and very low polymer concentration. This data is of spe-
FIG. 9: $D_s$ of 245 kDa polystyrene in 245 and 599 kDa polystyrene:benzene (open, filled circles, respectively), and 599 kDa polystyrene in 599 and 1800 kDa polystyrene:benzene (open, filled squares), using data of Leger, et al.\textsuperscript{56} and parameters in Table I.

Specific interest here in that it supplies values for $D_s$ in the low-concentration region, as a supplement to data sets in which $D_s$ was only obtained at relatively elevated concentrations. Omitting the largest-$M$ point (and thereby reducing the RMS fractional error in the fit from 18\% to 8.6\%), their data follow $D_o = 4.54 \cdot 10^{-4} M^{-0.588}$.

Hervet, et al.\textsuperscript{55} and Leger, et al.\textsuperscript{56} studied $D_s$ of polystyrene in benzene using FRS to measure the relaxation of photoexcitation patterns. Self-diffusion coefficients were obtained for polymers of molecular weight 78.3, 123, 245, 599, and 754 kDa, with $M_w/M_n \approx 1.06 - 1.12$, for concentrations up to 550 g/L. The same technique was used to obtain diffusion coefficients of labelled 245 kDa probe chains in a 599 kDa polystyrene matrix, and 599 kDa polystyrenes in a 1800 kDa polystyrene matrix. Matrix concentrations ranged from 5 to 400 g/L.

To supplement these measurements, many of which were made at elevated polymer concentrations, we used the dilute-solution self-diffusion measurements of Hadgraft\textsuperscript{54} to estimate $D_s$ at very low $c$.

Figure 9 displays the self-diffusion data of Hervet, et al.\textsuperscript{55} and Leger, et al.\textsuperscript{56} and the fits to stretched exponentials. Except for the 245 (filled circles) and 598(open squares) kDa polystyrenes, there is good agreement between the reported $D_s(c)$ and the fits. In the two anomalous systems, $D_s$ first increases with increasing $c$ and then decreases at larger $c$. The initial increase in $D_s$ with increasing $c$ appears to be substantially larger than the random error in the measurements as inferred from random scatter in the data. The non-monotonic dependence

FIG. 10: $D_s$ of 130 and 180 g/L polystyrene in dibutylphthalate as a function of molecular weight, showing the lower-molecular-weight stretched-exponential and the higher-molecular-weight power-law molecular-weight dependences of $D_s$, using data of Nemoto, et al.\textsuperscript{57}, Table II.

FIG. 11: $D_s$ of poly(ethylene oxide), [from top to bottom, at $\phi \approx 0.1$] namely 2kDa in CHCl$_3$ and in dioxane, 20 kDa in benzene and in dioxane, 40 kDa in chloroform and in dioxane, and 3600 kDa in dioxane, after Skirda, et al.\textsuperscript{59} Fig. 1a. Lines are fits to stretched exponentials in volume fraction.
of $D_s$ on $c$ is unique to this specific set of measurements. The lack of agreement between the stretched exponential form and the measured $D_s$ is therefore ascribed to some unique feature of this data, and not to a general property of polymers in good solvents.

Figure 11 presents $D_p(c)$ for the two probe-matrix systems and compares with $D_s(c)$ of polystyrenes of similar molecular weights. For both probe-matrix systems, $D_p(c)$ is described well by a stretched exponential, with no indication of the non-monotonic concentration dependence seen for $D_s(c)$ of the 245 and 598 kDa chains. The 245 kDa polystyrene diffuses approximately equally rapidly through 245 and 599 kDa matrices. The 599kDa polystyrene diffuses markedly more slowly through the 1800 kDa matrix than through the 599 kDa matrix.

Nemoto, et al. used FRS to measure the self-diffusion coefficient of polystyrene as a function of polymer molecular weight at fixed concentration. The solvent was dibutylphthalate. Polymer molecular weights covered a range $43.9 \leq M_w \leq 5480$ kDa with polymer concentrations of 130 and 180 g/L. The polymers were quite monodisperse, with $1.01 \leq M_w/M_n \leq 1.09$, except for the 5480 kDa material, for which $M_w/M_n \approx 1.15$.

Nemoto, et al.’s results appear in Fig. 10. A stretched-exponential molecular weight dependence of $D_s$ is not observed at all $M$. At the two concentrations studied by Nemoto, et al., the molecular weight dependence of $D_s$ has a transition near $M \approx 800$ kDa. At lower molecular weights, a stretched exponential in $M$ describes $D_s(M)$ extremely well. At larger molecular weights, one finds a power-law dependence $D_s \sim M^{-\gamma}$ with $\gamma \approx 2.49$ at 130 g/L and $\gamma \approx 2.22$ at 180 g/L.

Nemoto, et al. used FRS and a cone and plate viscometer to determine $D_s$ and the steady-state shear viscosity $\eta$ of concentrated solutions (40 and 50 wt%) of 44 and 355 kDa polystyrene in dibutylphthalate. In terms of the transient lattice models, the solutions of the 44kDa polymer are expected to be unentangled, while the solutions of the 355 kDa polymer are expected to be entangled. They found that $D_s/T$ and the fluidity $\eta^{-1}$ have virtually the same dependence on temperature, at either concentration, both for the 44 kDa polystyrene and for the 355 kDa polystyrene.

Skirda, et al. used PFGNMR to study the self-diffusion of polyethylene oxides ($M = 2, 20, 40,$ and $3000$ kDa) and polystyrenes ($M_n = 240$ and $1300$ kDa) in chloroform, benzene, dioxane, and carbon tetrachloride over a full range of polymer volume fractions $\phi$. $M_w/M_n$ was $\approx 1.1$ for the polyethylene oxides (except the 3000 kDa polymer, for which $M_w/M_n \approx 2$) and $\approx 1.2$ for the polystyrenes. Figures 11 and 12 show results for PEO and polystyrene, respectively, and the matching stretched-exponential fits, using parameters from Table I. For each polymer:solvent combination, a stretched-exponential form fits the data well.

Tao, et al. measured $D_s$ (using PFGNMR and forward recoil spectroscopy) and $\eta$ (from the dynamic shear moduli) of hydrogenated polybutadienes in alkane solvents. Polymer volume fractions $\phi$ extended from 0.2 up to the melt while polymer molecular weights cover two orders of magnitude, from 4.8 to 440kDa, with $M_w/M_n$...
of 1.01-1.03 or less. Tao, et al. fit their self-diffusion data to a scaling description $D_s \sim \phi^a M^b$. When they forced $a = -1.8$, a one-parameter fit found that the averaged $D_s \phi^{1.8}$ is $\sim M^{-2.41}$.

Figures 13 and 14 show Tao, et al.’s data as fit to to power-law and stretched-exponential forms, respectively. In Fig. 13 all points were fit simultaneously to a scaling equation, finding $D_s \sim c^{-1.71} M^{2.42}$, with a fractional root-mean square fitting error of 20.2%. These exponents differ slightly from those of Tao, et al.: Tao, et al. did sequential 2-parameter fits first to determine $a$ and then to determine $b$, while we did a single three-parameter fit to all points. Figure 14 shows a fit of all data points to the stretched-exponential form $D_{\infty} M^{-z} \exp(-\alpha c M^{\gamma})$. The factor $M^{-z}$ appears here because we are combining data on polymers with multiple molecular weights, and the extrapolations $c \rightarrow 0$ of measurements at different molecular weight should extrapolate to a different $D_s$ at each $M$. Formally, $D_{\infty}$ is the extrapolated zero-concentration diffusion coefficient of a highly hypothetical polymer having a molecular weight of unity. On forcing $\nu = 0.5$, we obtain $\gamma = 1.9 \cdot 10^{-4}$ and $z = 2.42$ with an RMS fractional error of 20.2%. Treating $\nu$ as a free parameter finds $\nu \approx 0.24$, with virtually the same value of $z$, $\gamma \approx 0$, and only a slightly improvement (to 19.6%) in the fit error. With either value for $\nu$, the molecular-weight dependence of $D_s$ is almost entirely determined by the prefix $M^{-z}$. The exponential itself has only a negligible dependence on $M$.

In Tao, et al.’s systems, scaling-law and stretched-exponential forms for $D_s$ thus provide equally good descriptions of the concentration dependence of $D_s$. The scaling and stretched-exponential forms also agree as to the molecular weight dependence of $D_s$ at fixed $c$, namely $D_s \sim M^{-2.41}$. Tao, et al. concluded that a scaling-law description of their data is correct. The analysis here corroborates this statement, but shows that it is incomplete, in that stretched-exponential forms describe equally accurately the measured $D_s(c, M)$.

Qualitatively, the larger part of Tao, et al.’s viscosity measurements might have been expected to be in the larger-concentration meltlike ($\eta \sim c^z$) rather than the smaller-concentration solutionlike ($\eta \sim \exp(\alpha c)$) regime. The transition in the concentration dependence of $\eta$ between these two functional forms is not transparently evident in $D_s(c)$, whose concentration dependence is consistent with a stretched exponential in $c$ for concentrations up to the true melt. However, contrary to those other systems reviewed in this article, from which molecular-weight dependences can be extracted, here the molecular weight exponent $\gamma$ of $\exp(-\alpha c M^{\gamma})$ is very nearly zero, so in Tao, et al.’s systems the concentration dependence of $D_s$ is nearly the same at all $M$.

Tinland, et al. report on self-diffusion of xanthans of molecular weight 0.45-9.4 MDa, with $M_w/M_n$ in the range 1.2-1.4, at concentrations 0.01-40 g/L. Xanthan forms wormlike chains. The experimental data of Tinland, et al. obtained with FRAP thus differs from $D_s$ of almost all other polymers, in that $d \ln(D_s)/d \ln(c)$
does not decrease monotonically with increasing $c$. This phenomenon, which was noted by the original authors, is plausibly related to the appearance of a lyotropic liquid-crystalline phase in this material at elevated $c$. None of the polymer dynamics models discussed above would be expected to remain valid while the system underwent a phase transition, so the behavior observed by Tinland, et al. does not contradict any model for polymer dynamics. Figure 16 shows Tinland, et al.’s measurements and fits of the lower-concentration data at each molecular weight to a stretched-exponential form. As seen from Table I, the somewhat large RMS fractional errors show that agreement between the measured data and the functional forms is not outstanding. The fit depends marginally on the number of data points included in the analysis. For the 3800 kDa polymer, we indicate (solid, dashed lines) the fits to the first 7 or 8 data points.

For all but the largest-$M$ polymer, there is a concentration $c^{**}$ at which $D_s(c)$ deviates from its low-concentration decline. In Tinland, et al.’s language, $c^{**}$ is the concentration of the higher-concentration boundary of the semidilute regime. From Fig. 16, $c^{**}$ appears to decrease with increasing polymer molecular weight.

von Meerwall, et al. used PFGNMR to measure $D_s$ of linear and star polyisoprenes over a near-complete range of concentrations (polymer weight fraction $0.01 \leq x \leq 1$). The arm number $f$ ranged from 2 to 18. Molecular weights of single arms were 5 and 18 kDa. Figure 16 compares von Meerwall, et al.’s data on $f = 2$ and $f = 8$ polyisoprene stars having 5kDa arms. Figure 17 shows $D_s$ for $f = 2, 3, 8, 18$ star polyisoprenes in CCl$_4$. In the original paper, data were reported as smooth curves, not as measured points. To create Fig. 17, the smooth curves were sampled; fits were made to the sampled points. From Fig. 17 at all concentrations increasing the arm count at fixed span molecular weight reduces $D_s$: Increasing four-fold the number of arms reduces $D_s$ by a factor of 2 to 3. At $f = 8$, an increase in span molecular weight reduces $D_s$.

In both figures, the solid lines are fits to stretched exponentials. Without exception, $D_s(c)$ for each $f$ and $M$ is described well by a stretched exponential in $c$. From the fit parameters in Table I, for fixed $M_{span}$ and increasing $f$ one finds that $D_s$ and $r$ fall while $r$ increases, a two-fold decrease in $D_s$ via increasing $f$ being accompanied by a 20-fold increase in $r$. For the smallest molecular weights (10-16 kDa) studied, $r > 1$ is observed.

von Meerwall, et al. used PFGNMR to study linear and 3-armed star polybutadienes and polystyrenes in CCl$_4$. They report low-concentration data on all systems, and extensive concentration dependence measurements on some star polymers. Polymer molecular weights ranged from 2.3 to 281 kDa with polydispersities $M_n/M_w$ of 1.03-1.07. As seen in Figs. 18 and 19 for most systems the observed concentration range afforded a one-order-of-magnitude variation in $D_s$.

Figure 20 gives Ref. 63’s measurements on the 6.5, 8.3, 29, and 76 kDa 3-armed stars, which were made over a far wider concentration range that afforded a 2.5-order-of-magnitude variation in $D_s$. Fits of these data to stretched-exponential forms describe well $D_s(c)$. As initially noted by von Meerwall, et al., ‘the slopes’ in Fig.
change continuously' . . . deGennes' prediction of a concentration scaling regime $D_s \sim c^{-1.75}$ ($c^* < c < c^{**}$) is not borne out by our data . . . at any molecular weight'.

von Meerwall, et al. used PFGNMR to measure self-diffusion of 10, 37.4, 179, 498, and 1050 kDa polystyrene in tetrahydrofuran at concentrations 6-700 g/L. The same technique was also used to measure $D_s$ of tetrahydrofuran and hexafluorobenzene in the same polymer solutions. Von Meerwall, et al. did not report $D_s$ for their polymers in dilute solution. $D_s$ for these systems in the dilute limit is therefore inferred here from the molecular weight dependence of $D_s$ observed by Hadgraft, et al. for polystyrene in benzene, together with the viscosities of tetrahydrofuran and benzene. von Meerwall, et al.'s measurements, and stretched-exponential fits appear in Fig. 21, using parameters given in Table I.

Wesson, et al. used FRS to measure self-diffusion of polystyrenes in tetrahydrofuran and benzene. The polystyrenes had $M$ of 32, 46, 105, 130, and 360 kDa, and were observed for concentrations in the range 40-500 g/L. Wesson, et al.'s measurements were here supplemented by extreme low-concentration points calculated from results of Hadgraft, et al. These points were included in the fits as having been taken at 0 g/L. Experimental results and the corresponding stretched-exponential fits appear in Fig. 22. Because $D_s$ here covers four orders of magnitude, on the scale of the figure the fits look very good. In fact, RMS fractional errors are in the range 15-21%, making these among the poorer fits in this section to a stretched exponential.

Xuexin, et al. used PFGNMR to measure $D_s$ of linear and 18-armed star polyisoprenes in CCl$_4$ over a wide range of $c$ and a 100-fold range of $M$. Their results (with concentrations replotted in g/L) appear as Figs. 23, 24, and 25. In every case the concentration dependence of $D_s$ is described well by the stretched exponential form, with fitting parameters given in Table I.
Figure 21: $D_s$ of 10, 37, 198, 498, and 1050 kDa linear polystyrenes in tetrahydrofuran, and fits to stretched exponentials in $c$. Data after von Meerwall et al., with supplemental low-concentration data based on Hadgraft, et al.

Figure 22: $D_s$ of [from top to bottom] 32, 46, 105, 130, and 360 kDa polystyrene in tetrahydrofuran, and associated stretched exponential fits. Data are from Wesson, et al., Table II and associated Figures, as supplemented by the low-concentration measurements of Hadgraft, et al.

In a majority of cases, the fits were quite good, with RMS fractional errors in the range 6-18%. Fits to the measurements of Leger, et al. and Wesson, et al. were less satisfactory; these are discussed separately below. Xuexin, et al. cover an extremely broad range of $M$ in their study of 18-armed stars. Over this range, $\nu$ from fits to data sets covering chains with a single molecular weight changes substantially, so a fit of all data over a full range of $M$ to eq. 17 with $\nu$ a fixed constant, does not work well for very small or very large molecular weights. On limiting the fit to intermediate values for $P$, good agreement between the data and the fitted forms are encountered. Solid curves in Fig. 27 were calculated using the parameters to the fit to data in the intermediate-$P$ range.

Omitting momentarily the fits to results of Leger, et al. and Wesson, et al.: The exponent $a$ for the molecular weight dependence of the bare diffusion coefficient is consistently -0.5. Except for Browne, et al.’s work on polyethylene oxide: water, the concentration exponent $\nu$ is in the range 0.5-0.75; Brown, et al.’s data imply $\nu \approx 0.93$. The molecular weight exponent $\gamma$ is in the range 0.32-0.46, again with the exception of fits to data of Brown, et al., for which $\gamma \approx 0.6$. Inspection of the figures indicates that a joint stretched exponential in $c$ and $M$ fits each data set well, with no systematic deviations for particular values of $c$ or $M$.

The two data sets that are fit less well by eq. 17 appear in Figure 28. The merged fit to Leger, et al.’s data on polystyrene:CCl$_4$ is poor. However, Leger, et al.’s data shows features – notably a non-monotonic dependence of $D_s$ on $c$ – that appears in data on no other polymer system, including other experiments that determined the concentration dependence of $D_s$ of the same polymer. We infer that the poor fit of eq. 17 to Leger, et al.’s data...
IV. DIFFUSION OF PROBE CHAINS THROUGH MATRIX POLYMER SOLUTIONS

This Section reviews measurements on the diffusion of polymeric probe molecules through solutions of a different polymer. These experiments involve intrinsically ternary solutions in which the molecular weight \( P \) of the probe polymer and the molecular weight \( M \) of the matrix polymer are not the same. In some cases, the probe and matrix polymers in the solution have a common monomer, and differ only in their molecular weights. In other cases, the probe and matrix polymers are chemically distinct. Studies are again presented alphabetically by first author, together with fits of the data sets to stretched exponentials in concentration and chain molecular weights. Fitting parameters appear in Tables II and III.

Brown and Rymden\textsuperscript{67} used quasielastic light scattering to study the diffusion of linear polystyrenes and coated silica spheres through polymethylmethacrylate in toluene. Toluene and PMMA are almost exactly index-matched, so scattering from these systems was dominated...
FIG. 26: $D_s$ as measured by (a) Brown, et al. (cf. Fig. 2), (b) Callaghan, et al. (cf. Fig. 3), (c) Callaghan, et al. (cf. Fig. 4), and (d) Deschamps, et al. (cf. Fig. 5), and fits to eq. 17, leading to the parameters in Table III.

by scattering from the dilute probe chains. The matrix PMMA’s had molecular weights in the range 110 kDa–1.43 MDa. Probe polystyrenes had molecular weights of 2.95, 8, and 15 MDa, with $M_w/M_n$ of 1.06, 1.08, and 1.30, respectively. In the original paper, data were reported after normalization by an unspecified diffusion coefficient obtained in the absence of the matrix polymer.

Reference 67 reported how $D_p$ depends on matrix concentration and molecular weight. Figure 29a shows $D_p/D_{p0}$ for the 8MDa polystyrene diffusing through each of six matrix polymethylmethacrylates. Each solid line represents a fit to a stretched exponential in matrix polymer concentration, using parameters in Table II. For $D_p/D_{p0} < 10^{-3}$ or so, a condition attained only with the two largest matrix polymers, the measured $D_s$ deviates markedly downward from a stretched exponential, so that fits to all data on these two systems show fractional RMS errors much worse (32–45% rather than 5–8%) than fits to the same probe polymer with the smaller matrix polymers. On excluding the final few points from the two fits, curves with far lower RMS fractional errors were generated. These are the curves shown in the Figure. It is difficult to determine from these measurements whether there is a systematic change from stretched-exponential to some other concentration dependence at very small $D_p/D_{p0}$, or whether the apparent deviations arise from
FIG. 27: $D_s$ as measured by (a) Giebel, et al.\textsuperscript{52}, (b) von Meerwall, et al.\textsuperscript{63} and (c), (d) Xuexin, et al.\textsuperscript{66}, and fits to eq.\textsuperscript{17} leading to the parameters in Table III. Other details as in Figs. 7, 20, 23, and 24 respectively.

experimental challenges at very small $D_p$.

Figure 29b shows the same data, now fit simultaneously to eqn.\textsuperscript{17} the joint stretched exponential in $c$ and $M$. Fitting parameters appear in Table IV. Except for the lowest-$M$ matrix polymer, the fits are almost as good as the individual fits shown in Figure 29a. For the lowest-$M$ 101 kDa matrix polymer, the fitted form predicts too strong a dependence of $D_p$ upon $c$.

Brown and Rymden\textsuperscript{67} also examined how $D_p$ depends on probe molecular weight. Figure 30 shows $D_p$ of the 3, 8, and 15 MDA probe polystyrenes, all diffusing through the 445kDa PMMA. Fits are to stretched exponentials in $c$, leading to parameters given in Table III. The matrix chains are all much larger than the probe chains. As noted by the original authors\textsuperscript{67}, the three curves come very close to superposing except perhaps at the very highest matrix concentrations examined.

Brown and Stilbs\textsuperscript{68} used PFGNMR to measure the probe diffusion coefficient of polyethylene oxide in aqueous solutions of dextran. Polyethylene oxides had molecular weights of 73, 278, and 1200 kDa with $M_w/M_n$ of 1.02-1.12; dextrans had molecular weights of 19, 110, and 510 kDa. The 1200 kDa PEO represented the lower limit at which $D_p$ could be determined with then-available technology; the authors limited their detailed analysis to the two lower-molecular-weight probes.
FIG. 28: Joint fits to all data of (a) Leger, et al. and (b) Wesson, et al., showing the poor quality of fits of eq. 14 to these data sets. Fit parameters are in Table III; all other plot properties are the same as Figs. 8 and 22, respectively.

Figures 31a and 31b show $D_p$ of the 73 and 278 kDa polyethylene oxides, normalized by the measured $D_o$ of the same probes in the absence of the matrix polymers. All measurements were simultaneously fit to $D_p/D_o \exp(-\alpha c^\nu P^\gamma M^\delta)$, yielding the smooth curves shown in the Figures and the parameters listed in Table IV. Agreement between the data and the fitted curves was very good for the 73 kDa probe. For the 278 kDa probe in the 19 kDa matrix, the fitting function significantly underpredicts $D_p$. From the fitting parameters, $D_p/D_o$ has a very weak dependence on the probe molecular weight (other than that hidden in $D_o$), but has a marked dependence on the matrix molecular weight. The same data were also fit, individually for each probe:matrix pair, to a stretched and a pure exponential in $c$. Without exception, for each probe:matrix pair $D_s(c)$ follows accurately the exponential form, RMS fractional errors being in the range 1-4%. Because $D_p$ varied over a limited range, the parameters reported in Table II reflect fits to the pure exponential.
Daivis, et al.\textsuperscript{69} used quasielastic light scattering spectroscopy to measure diffusion of relatively dilute 864 kDa dextran in not-necessarily-dilute solutions of 20.4 kDa dextran. Polymer polydispersities were in the range 1.24-1.3. The concentration of the lower-molecular-weight dextran ranged up to 166 g/L. Analysis of the bimodal QELSS spectra of these systems shows that the slower mode corresponds to probe diffusion by the 864 kDa dextran. As seen in Figure 32, the data fit reasonably well to a simple exponential concentration dependence, using parameters found in Table II.

In a separate paper, Daivis, et al.\textsuperscript{70} used QELSS and PFGNMR to measure the diffusion of a 110 kDa polystyrene, $M_w/M_n = 1.06$, through solutions of 110 kDa polyvinylmethylether, $M_w/M_n \approx 1.3$, in the PVME’s isorefractive solvent toluene. Good agreement was found between the QELSS measurements of $D_p$ and the earlier measurements of $D_p$ by Martin\textsuperscript{71} on the same system.

Figure 33 shows Daivis, et al.’s data\textsuperscript{70} as obtained using both physical methods. In the figure, lines represent separate fits of each data set to a stretched exponential in matrix polymer concentration, with fitting parameters given in Table II. Stretched exponentials in $c$ describe well each data set. The QELSS data is significantly less scattered than the PFGNMR data, so the parameters from the former’s fit are probably to be preferred.

De Smedt, et al.\textsuperscript{72} used FRAP to measure the diffusion of 71, 148, and 487 kDa dextrans ($M_w/M_n < 1.35$), labeled with fluorescein isothiocyanate, through solutions of hyaluronic acid ($M_n = 390$ kDa; $M_w = 680$ kDa).

Hyaluronic acid concentrations ranged from the dilute up to 18 g/L. $D_p$ of the dextrans varied roughly five-fold over this concentration range. De Smedt, et al.’s data\textsuperscript{72} appear in Fig. 34 together with stretched-exponential fits using the parameters in Table II. As seen in the Figure and initially reported by the original authors, the data fits well to stretched exponential forms.

Hadgraft, et al.\textsuperscript{54} used QELSS to study the diffusion of polystyrene probe polymers, molecular weights 25, 162, 410, 1110, and 4600 kDa through solutions of...
105 kDa polymethylmethacrylate in its isorefractive solvent benzene at PMMA concentrations up to 100 g/L. Polystyrene and PMMA are not compatible, implying that the radius of the polystyrene chains may have depended very strongly on the matrix polymer concentration. As seen in Fig. 35, $D_p$ of the 25 kDa polystyrene was substantially independent of PMMA concentration. The data on the 410 kDa polymer is significantly more scattered than is data on the other polystyrenes. The range of variation of $D_p$ is sufficiently small (roughly a factor of three) that the fits to these data are less reliable than are the fits to data on some other systems. Nonetheless,
Hanley, et al.'s data are given in Fig. 36, together with fits of separate stretched-exponentials in \( c \) to each data set, yielding parameters shown in Table III. The entire data set was also fit simultaneously to the joint stretched exponential of eq. (17), yielding parameters seen in Table III and fitted curves seen in Fig. 36b. The lack of measurements at very low matrix concentration substantially broadens the range of fitting parameters that yield reasonable descriptions of this data. The fit to the joint stretched exponential is substantially less satisfactory than the individual fits at each \( P \) to separate stretched exponentials (RMS fractional errors of 34% rather than 9-24%). The inadequacy of eq. (17) relates primarily to the 1050 kDa probe chains, for which \( D_p \) was determined only over a limited range. As seen in Table III, excluding data on the 1050 kDa probes from the simultaneous fit leads to a marked reduction in the RMS fractional fit error (to 24%), modest changes in \( \alpha \) and \( \nu \), but only small changes in the other fitting parameters.

Kent, et al. applied static and quasielastic light scattering to measure radii of gyration and diffusion coefficients of 233 and 930 kDa polystyrenes in 7, 66, 70, 840, and 1300 kDa polymethylmethacrylates in ethyl benzoate (for static light scattering) and toluene (for quasielastic light scattering). With one exception (66 kDa PMMA), \( M_w/M_n \) was always \( \leq 1.10 \). Different probe:matrix combinations were used for static and quasielastic light scattering. \( D \) of the polystyrene was measured as a function of polystyrene concentration, and linearly extrapolated to the dilute-in-polystyrene limit, thereby obtaining both the probe diffusion coefficient \( D_p \) of the polystyrene and the initial linear dependence of \( D_p \) on polystyrene concentration.

Figure 37 shows Kent, et al.'s measurements of the radius of gyration of 930 kDa polystyrene in solutions of 7, 70, or 1300 kDa PMMA as a function of PMMA concentration, together with fits of \( R_g \) to a stretched exponential

\[
R_g = R_{g0} \exp(-\alpha c^\nu)
\]

in matrix concentration \( c \). The fits are good throughout, using parameters in Table IX. Unlike the scaling-law prediction \( R_g \sim c^{-0.25} \), the stretched exponential form shows acceptable behavior down to zero matrix concentration.

Kent, et al. also measured \( D_p \) of 233 kDa polystyrene in solutions of 66 and 840 kDa PMMA, and 930 kDa polystyrene through 840 kDa PMMA. Their experimental data is shown in Fig. 38a, together with fits to exponentials using parameters given in Table III. Within experimental error, the simple exponential fits with \( \nu = 1 \) forced are as good as the stretched-exponential fits to the data: the former are in the figure. Experimentally, the scaling prefactor \( \alpha \) depends strongly on matrix molecular weight (a 12-fold change in \( M \) leads to a two-fold change in \( \alpha \)) but at most weakly on probe molecular weight. Figure 38b shows a fit of the same data to a joint stretched exponential in \( c, P, \) and \( M \), based on parameters in Table

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**FIG. 36:** \( D_p \) of [from top to bottom] 50, 179, 1050, and 1800 kDa polystyrenes in orthofluorotoluene solutions of 60kDa polyvinylmethylether against matrix polymer concentration, using data of Hanley, et al. and (a) separate fits at each \( P \) and (b) simultaneous fits at all \( P \), with fitting parameters from Tables II and III respectively.

\( D_p \) of the higher-molecular-weight polystyrenes shows a stretched-exponential dependence on PMMA concentration, with parameters seen in Table II.

Hanley, et al. used light scattering spectroscopy to examine the diffusion of polystyrenes through the matrix polymer polyvinylmethylether in its isorefractive solvent orthofluorotoluene. The polystyrenes had molecular weights of 50, 179, 1050, and 1800 kDa. The polyvinylmethylether had \( M_w \approx 60 \text{ kDa} \) with \( M_w/M_n \approx 3 \). Hanley, et al.'s data are given in Fig. 36, together with fits of separate stretched-exponentials in \( c \) to each data set, yielding parameters shown in Table III. The entire data set was also fit simultaneously to the joint stretched exponential of eq. (17), yielding parameters seen in Table III and fitted curves seen in Fig. 36b. The lack of measurements at very low matrix concentration substantially broadens the range of fitting parameters that yield reasonable descriptions of this data. The fit to the joint stretched exponential is substantially less satisfactory than the individual fits at each \( P \) to separate stretched exponentials (RMS fractional errors of 34% rather than 9-24%). The inadequacy of eq. (17) relates primarily to the 1050 kDa probe chains, for which \( D_p \) was determined only over a limited range. As seen in Table III, excluding data on the 1050 kDa probes from the simultaneous fit leads to a marked reduction in the RMS fractional fit error (to 24%), modest changes in \( \alpha \) and \( \nu \), but only small changes in the other fitting parameters.

Kent, et al. applied static and quasielastic light scattering to measure radii of gyration and diffusion coefficients of 233 and 930 kDa polystyrenes in 7, 66, 70, 840, and 1300 kDa polymethylmethacrylates in ethyl benzoate (for static light scattering) and toluene (for quasielastic light scattering). With one exception (66 kDa PMMA), \( M_w/M_n \) was always \( \leq 1.10 \). Different probe:matrix combinations were used for static and quasielastic light scattering. \( D \) of the polystyrene was measured as a function of polystyrene concentration, and linearly extrapolated to the dilute-in-polystyrene limit, thereby obtaining both the probe diffusion coefficient \( D_p \) of the polystyrene and the initial linear dependence of \( D_p \) on polystyrene concentration.

Figure 37 shows Kent, et al.'s measurements of the radius of gyration of 930 kDa polystyrene in solutions of 7, 70, or 1300 kDa PMMA as a function of PMMA concentration, together with fits of \( R_g \) to a stretched exponential

\[
R_g = R_{g0} \exp(-\alpha c^\nu)
\]

in matrix concentration \( c \). The fits are good throughout, using parameters in Table IX. Unlike the scaling-law prediction \( R_g \sim c^{-0.25} \), the stretched exponential form shows acceptable behavior down to zero matrix concentration.

Kent, et al. also measured \( D_p \) of 233 kDa polystyrene in solutions of 66 and 840 kDa PMMA, and 930 kDa polystyrene through 840 kDa PMMA. Their experimental data is shown in Fig. 38a, together with fits to exponentials using parameters given in Table III. Within experimental error, the simple exponential fits with \( \nu = 1 \) forced are as good as the stretched-exponential fits to the data: the former are in the figure. Experimentally, the scaling prefactor \( \alpha \) depends strongly on matrix molecular weight (a 12-fold change in \( M \) leads to a two-fold change in \( \alpha \)) but at most weakly on probe molecular weight. Figure 38b shows a fit of the same data to a joint stretched exponential in \( c, P, \) and \( M \), based on parameters in Table

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The joint fit clearly works well for all three polymer pairs.

Kim, et al. measured the diffusion of dye-labeled polystyrenes through matrix solutions of unlabeled polystyrenes in toluene. The objective was to test the prediction of some scaling models that $D_p$ becomes independent of matrix molecular weight if $M/P \geq 1$. The probe polystyrenes had molecular weights from 10 to 1800 kDa; matrix chains had molecular weights from 51 to 8400 kDa. Ref. 75 also reported the dependence of $D_p$ on the matrix molecular weight for several probes (51, 390, 900 kDa) at multiple matrix concentrations for matrix molecular weights in the range 35–8400 kDa. Polymer polydispersities were largely $< 1$, with a maximum of 1.17. Kim, et al. also report limited data using methyl red as a low-molecular weight probe. As seen in Fig. 39, the authors found that $D_p$ becomes substantially independent of $M$ only if $M/P > 3$.

With respect to the models discussed in Section II, the published derivations of the stretched-exponential form refer to polymer chains whose motions are adequately approximated by whole-chain translation and rotation. These approximations are only likely to be adequate if the probe and matrix chains are of similar size, because otherwise the whole-chain motions of probe or matrix would effectively sample some of the internal modes of the other chain species, whether matrix or probe. At fixed $c$, from the hydrodynamic model $D_p$ would have a stretched-exponential dependence on $M$ if $M/P \approx 1$, but might well not have a stretched-exponential matrix molecular weight dependence if $M/P \gg 1$ or $M/P \ll 1$.

Kim, et al.’s results thus are not necessarily inconsistent with exponential-type models of polymer dynamics.

Kim, et al.’s data for the domain $M/P < 3$, as seen in Figs. 39 are fit to the form $D_pP^{-\alpha} \exp(-\sigma c^\delta P^\gamma M^\delta)$, yielding parameters in Table III. As seen in Fig. 39, $D_p$ is indeed described well by the joint stretched exponential over a wide range of $c$, $P$, and $M$, except in the regime $M/P > 3$ in which the form is not necessarily expected.
to apply.

Kim, et al. also studied the concentration dependence of $D_s$ and $D_p$ at large $M/P$. Figure 40 shows Kim, et al.'s determination of self-diffusion of 900 kDa polystyrene in toluene, and our stretched-exponential fit to the data, yielding the parameters in Table I. Figure 41 shows measurements of probe diffusion, using as probes methyl red and 10, 35, 100, 390, 900, and 1800 kDa polystyrenes, all through matrix polystyrenes in toluene. The molecular weight of the matrix polystyrene ranges from 51 to 8400 kDa, varying from data point to data point, with $M/P > 3$ and generally $M/P > 6$. This ratio of $M/P$ was chosen by Kim, et al. based on their interpretation of their data that showed $D_p$ to be independent of $M$ for $M/P > 3$. As seen in Figs. 40 and 41, stretched-exponential functional forms using parameters in Table II describe the matrix-polymer concentration dependences of $D_s(c)$ and $D_p(c)$ well over a wide range of polymer concentrations and probe molecular weights, even though the ranges of $P$ and $M$ are very wide. The stretched exponential in $c$ continues to describe well the concentration dependence of $D_p$ in the large $M/P$ range which eq. 17 does not represent well the $P$ and $M$ dependences of $D_s$.

Lodge and collaborators have reported an extensive series of studies of probe diffusion in polymer solutions, using quasielastic light scattering to measure $D_p$ of a dilute probe polymer, generally polystyrene, through the isorefractive matrix polymer:solvent pair polyvinylmethylether: orthofluorotoluene. Variables studied include the probe and matrix molecular weights, the matrix concentration, and the topology (linear and star) of the probe polymers.

An early letter of Lodge reports $D_p$ of 179 kDa and 1.05 MDa polystyrenes through a 50kDa polyvinylmethylether. The data, and corresponding stretched-exponential fits, appear in Fig. 42. Fit parameters are in Table III. A stretched exponential in concentration describes well both data sets.

Lodge and Wheeler compared the diffusion of linear and 3-armed star polystyrenes through a high molecular weight polyvinylmethylether. Polystyrene molecular weights were 422 and 1050 kDa for the linear chains.

**FIG. 39:** $D_s$ of polystyrene in matrix polystyrene:toluene solutions as functions of matrix molecular weight at various matrix concentrations, based on data of Kim, et al., Fig. 3. Probe molecular weights were (a) 51 kDa, (b) 390 kDa, and (c) 900 kDa. Matrix concentrations [top to bottom] were (a) 10, 20, 50, and 100 g/L; (b) 20, 50, and 100 g/L; and (c) 10, 20, 40, and 80 g/L. A single stretched exponential $D_sP^{-\alpha} \exp(-\alpha c P^\beta M^\gamma)$ with constant parameters was fit (solid lines) to all data in the figures having $M/P < 3$. 
FIG. 40: $D_s$ of 900 kDa in toluene, based on data of Kim, et al., Fig. 1, and a fit to a stretched exponential.

FIG. 41: Probe diffusion of (top to bottom) methyl red, and 10, 35, 100, 390, 900, and 1800 kDa $D_s$ polystyrenes through high-molecular-weight ($M/P > 3$) polystyrenes in toluene, based on data of Kim, et al., Table II, and fits to stretched exponentials. The left axis shows zero-matrix-concentration data.

and 379 and 1190 kDa for the stars; the polystyrenes were identified as being 'relatively monodisperse'. The PVME had an $M_w$ of 1.3 MDa, with an estimated $M_w/M_N \approx 1.3$. $D_p$ was obtained over $1 \leq c \leq 100$ g/L in PVME concentration.

Figure 42 shows Lodge and Wheeler’s measurements, together with fits of data on each probe polymer to a stretched exponential in $c$. $D_p$ varies over nearly four orders of magnitude. Over the full range, agreement of the data with the functional form is very good, with RMS fractional errors of 2.6-10% and parameters as seen in Table II. Figure 43, shows the same data, with the linear chains and star polymers separately fit to a stretched exponential in $c$ and $P$. As seen in Table III, the fractional errors in these fits are very nearly as good as the fits to individual probe species. The $P^{-a}$ scaling of the zero-concentration diffusion coefficient and the $P^\gamma$ scaling of the scaling prefactor $\alpha$ account for the dependence of $D_p$ of linear and star polymers on probe molecular weight.

Lodge and Markland used light scattering spectroscopy to measure the single-particle diffusion coefficients of tracer 12-armed star polystyrenes through solutions of 140 kDa polyvinylmethylether, $M_w/M_n \approx 1.6$, in the isorefractive solvent orthofluorotoluene. The polystyrenes had $M_w$ of 55, 467, 1110, and 1690 kDa, with $M_w/M_n \leq 1.10$. Lodge and Markland estimate for the matrix that $c^* \approx 20$ g/L and $c_e \approx 100$ g/L. Figure 44 shows Lodge and Markland’s data. In the two graphs, the solid lines represent, respectively, fits to individual stretched exponentials in $c$ and to fits to a joint stretched exponential in $c$ and $P$. Fitting parameters appear in Tables II and III. Parameters in Table III differ modestly from Table III of Ref. 78. The sets of fits are excellent, with RMS fractional individual fits here are excellent, with errors of 2-4%; the RMS fractional error for the joint fit was 12%. The most notable deviation for the joint exponential is for the smallest probe at large

FIG. 42: $D_p$ of 179 kDa and 1.05 MDa polystyrene through 50 kDa polyvinylmethylether in orthofluorotoluene, based on data of Lodge, and fits to stretched exponentials in $c$. 
concentration, where the fitting function underestimates $D_p$.

Lodge, Markland, and Wheeler\textsuperscript{79} used light scattering spectroscopy to measure the diffusion of 3-armed and 12-armed star polystyrenes through solutions of polyvinylmethylether in its isorefractive solvent orthofluorotoluene. The 3-armed stars had $M_w$ of 379 and 1190 kDa; the 12-armed stars had $M_w$ of 55, 467, 1110, and 1690 kDa. Polyvinylmethyl ethers used as matrices had $M_W$ of 140, 630, and 1300 kDa. Polystyrenes all had $M_w/M_n < 1.1$; the matrix polymers had $M_w/M_n \approx 1.6$.

Light scattering measurements were also made of the radii of gyration of linear, 3-arm and 12-arm stars with molecular weights above 1MDa in the presence of 250 kDa polyvinylmethylether at concentrations as large as 50 g/L. Lodge, et al.\textsuperscript{79} reported that they modified their
reported $D_p$ to take account of the concentration dependence of a local friction factor, using a process described by Wheeler and Lodge\cite{Wheeler1970} (see below). This modification factor was removed from Lodge, et al.'s data, before making the analysis here, in order that this data be more strictly comparable with the remainder of the literature.

Figures 45 and 46 show Lodge, et al.'s measurements of $D_p$ for 3-armed and 12-armed stars. As is apparent from the figures, for every $M : P$ combination a stretched exponential gives an excellent description of the concentration dependence of $D_p$, with RMS fractional errors in the range 4-18\% (cf. Table II).

The entirety of Lodge and collaborators’ data\cite{Lodge1978,Lodge1979} on 12-armed and 3-armed stars were also fit (separately for each arm number) to equation 17, the joint stretched exponential in $c$, $P$, and $M$. Fitting parameters appear in Table II. Figure 47 shows the fit of $D_p$ for the 3-armed stars to the joint stretched exponential. As in the few other cases in which there are substantial differences between the fitted curve and measurements, the stretched-exponential form overestimates the concentration dependence of $D_p$.

Figure 48 shows the outcomes of the joint fit to $D_p$ of the 12-armed stars. Except for the smallest (55kDa) star in the 1.3MDa matrix polymer, eq. 17 describes very well the entire dependence of $D_p$ on all three variables. The fit for $f = 12$ is markedly more outstanding than is the fit to the $f = 3$ stars, particularly with the 379 kDa pVME as the matrix polymer. The displayed curves represent fits to all data points except for the 55kDa probe in the 1.3 MDa matrix polymer for $c \geq 10$ g/L. Including these 55kDa probe points in the fit raises the fractional RMS error from 14.6\% to 19.5\%, changes the three scaling exponents by 0.01 each, and elsewise has almost no effect on the fitted curves.

Figures 47 and 48 show that eq. 17 and a single set of fitting parameters account well for the dependence of $D_p$ on $c$, $P$, and $M$ for star polymers of given arm number in linear matrices. All three independent variables ranged over extensive domains: more than two orders of magni-

**FIG. 45:** $D_p$ of (a) 379 kDa and (b) 1.19 MDa 3-arm star polystyrenes in (from top to bottom, 140, 379, and 1300 kDa) polyvinylmethylethers in orthofluorotoluene, based on data of Lodge, et al.\cite{Lodge1978}. Solid lines are stretched-exponential fits with parameters in Table II.

**FIG. 46:** $D_p$ of 12-arm star polystyrenes (from top to bottom, $M_w$ of 55, 467, 1110, and 1690 kDa) diffusing through solutions of 1300 kDa polyvinylmethylether:ortho-fluorotoluene based on data of Lodge, et al.\cite{Lodge1979}. Solid lines are stretched-exponential fits with parameters in Table II.

References:

Wheeler, T. D., and Lodge, M. F. (1970). Frictional factors of star polymers. J. Chem. Phys. 52, 4700-4711.

Lodge, M. F., et al. (1978). Diffusion of star polymers: a stretched-exponential description. Macromolecules 11, 540-547.

Lodge, M. F., et al. (1979). Diffusion of star polymers: a stretched-exponential description. Macromolecules 12, 173-181.
The only failure in the fit occurred for the smallest probe in the largest matrix polymer \((P/M \approx 25)\) at elevated matrix concentrations.

Martin\textsuperscript{71,80} examined polystyrenes diffusing through polyvinylmethylether (as the matrix polymer) in toluene, with which the matrix is isorefractive. Polystyrenes had molecular weights of 50, 100, 420, and 900 kDa with \(M_w/M_n \leq 1.1\). The polyvinylmethylether had from intrinsic viscosity measurements a molecular weight ca. 110 kDa and a 'fairly polydisperse' molecular weight distribution. Diffusion coefficients of the probe polymers were obtained with QELSS; Martin\textsuperscript{71,80} also determined the viscosities of the polymer solutions.

Figure 49 shows \(D_p\) for each of the four probe polymers, as functions of matrix concentration. In Fig. 49, \(D_p\) for each probe polymer was separately fit to a stretched exponential in \(c\), yielding the parameters given in Table II and the four solid lines seen in the Figure. Stretched exponential forms do an excellent job of describing \(D_p(c)\). In Fig. 49, all data on the four probes was fit simultaneously to a single stretched exponential.
in \(c\) and probe molecular weight \(P\), obtaining fitting parameters given in Table III. Martin only reports results for one \(M\), making it impossible to evaluate the \(M\)-dependence of \(D_p\) form his results. Plots of this stretched exponential, as functions of \(c\) at fixed \(P\), give the solid lines of Fig. 49b. Agreement between the fitting function and experiment is good in the second Figure, though less good than in the first.

Martin\textsuperscript{71} also measured the viscosity of his matrix polymer solutions. At lower polymer concentrations, especially for the lower-molecular-weight probe polymers, Martin\textsuperscript{71} found that \(D_p\eta\) is nearly constant. At elevated concentrations, especially for the larger probe chains, Stokes-Einstein behavior ceases to obtain: \(D_p\eta\) increases with increasing \(c\), so that at large \(c\) and \(P\) the polymer chains diffuse faster than might have been expected from the macroscopic solution viscosity.

In addition to the self-diffusion studies noted in the previous Section, Nemoto, et al.\textsuperscript{82} used ultracentrifuga-
tion and quasielastic light scattering to measure sedimentation and probe diffusion coefficients of dilute polymethylmethacrylates (as probe polymers) in isorefractive polystyrene: thiophenol solutions. Polystyrene molecular weights were 43.9, 186, 775, and 8420 kDa; the PMMA had a molecular weight of 342 kDa. The reported $M_w/M_n$ were in the range 1.10-1.17.

Figure 50a shows Nemoto, et al.'s measurements of $D_p$ of PMMA in solutions of each of the four polystyrenes. For each molecular weight of the matrix, $D_p(c)$ is described to good accuracy by a stretched exponential in matrix concentration. Fitting parameters are given in Table III. Figure 50b shows the same data with all measurements simultaneously fit to a stretched exponential in $c$ and $M$, yielding parameters in Table III. Over nearly 200-fold variations in these variables, the single stretched exponential in $c$ and $M$ describes reasonably well the behavior of $D_p$, with a 20% RMS fractional error.

Nemoto, et al. report that $D_p/D_o$ depends more strongly on $c$ and $P$ than does $s/s_o$. At large $c$, especially at large $M$, $D_p/D_o$ was found to be significantly less than $s/s_o$. Nemoto, et al. concluded that at elevated matrix concentrations and polymer molecular weights the sedimentation and self-diffusion behaviors of PMMA in polystyrene solutions are quite distinct.

For two samples with the same $D_p/D_o$ but very different matrix molecular weights (44, 8420 kDa), Nemoto, et al. also measured the shear viscosity $\eta$, finding that $\eta$ differed 'by more than two orders of magnitude' between the two samples. Nemoto, et al. thus showed that $D_p$ is not governed by the shear viscosity of the matrix solution. (The original paper did not specify which solution was the more viscous. Note that the comparison is being made at fixed $D_p/D_o$, not at fixed $c$, so the correspondence is not self-evident.)

Nemoto and collaborators also used forced Rayleigh scattering to study the diffusion of probe polystyrenes through polystyrene-dibutylphthalate solutions. A first study focused on self-diffusion and tracer diffusion of labeled polystyrene through 40 wt% solutions of very long chains ($M/P > 5$) and very short chains ($M/P < 0.2$). Thirteen polystyrenes having $2.8 \leq M_w \leq 8420$ kDa and $M_w/M_n > 1.07$ (except for chains larger than 1MDa, for which $M_w/M_n$ was in the range 1.09-1.17) were used in the studies.

Nemoto, et al.'s data appears as Fig. 52. For $D_s$ and for $D_{tr}$ of short probe chains in solutions of long matrix molecules, a best-fit to the molecular weight dependence of $D$ gives parameters seen in Table III the $P$-dependence of the prefactor being forced rather than obtained from the fit. Because all data is at the same concentration, a concentration dependence was not obtained. Similarly, if we fit the data of Nemoto, et al. on $D_p$ of long probe chains in short matrix chains to $D_p P^{-\alpha} \exp(-\alpha P^2)$, we find that $D_p$ gains its molecular weight dependence almost entirely through the factor $P^{-\alpha}$. The data are fit well with $\alpha = 0.52$, in which case the best-fit gives $\gamma \approx 0.03$.

In a separate paper, Nemoto, et al. used forced Rayleigh scattering to measure $D_p$ of probe polystyrenes in dibutylphthalate solutions of high-molecular-weight matrix polymers at 13 and 18 % matrix concentration. The probe polystyrenes were in the molecular weight range $6.1 \leq M_w \leq 2890$ kDa, with polydispersities $M_w/M_n \leq 1.17$, and generally $\leq 1.09$. Table IV shows the fits to this data. RMS fractional errors were 12-13% for single concentrations, but ca. 30% for the fit to both concentrations. It should be stressed that $D_p$ covers more than five orders of magnitude, so the errors are not large relative to the total range of $D_p$. We find a significant dependence of $D_p$ on matrix as well as probe molecular weight. With only a few points for any particular $P$ or $M$, it is difficult to present the data as a simple figure.

Numasawa, et al. used QELSS to study the diffusion of tracer polystyrene chains through index-matched solutions of matrix polymethylmethacrylates in benzene. Polystyrene molecular weights were in the range 185-8420 kDa, with $M_w/M_n$ in the range 1.04-1.17. Polymethylmethacrylates had molecular weights 850-4050 kDa, with $M_w/M_n \leq 1.08$, except for the 850 kDa polymer, for which $M_w/M_n = 1.35$. PMMA matrix concentrations were a half-dozen values in the range 0-36 g/L. The tracer polystyrenes were dilute in all solutions. In addition to measuring the probe diffusion coefficients, Numasawa, et al. also report the zero-shear viscosity of the matrix polymers and (on the basis of static light scattering) determinations of the radii of gyration of the probe polymers as a function of matrix concentration.
Numasawa, et al’s results\textsuperscript{85} appears as Fig. 52. Figure 52a shows $D_p$ of the 420 kDa and the 8.42 MDa polystyrenes in solutions of two polymethylmethacrylates, as a function of matrix concentration. Figure 52b shows $D_p$ of five polystyrenes in each of four polymethylmethacrylates, all at a matrix concentration near 37 g/L. One observes that $D_p$ decreases monotonically with increasing probe molecular weight and with increasing matrix concentration and molecular weight.

All eight solid lines in both figures represent a simultaneous fit of a stretched exponential in $c$, $P$, and $M$ to all of the data in both Figures and to one additional data point. Fit parameters appear in Table III. It is important to emphasize that the same fitting parameters were used to generate all eight curves in both Figures. From Fig. 52a, the stretched exponential captures well the c-dependence of $D_p$, and the variation of that dependence with $P$ and $M$. From Figure 52b, at fixed $c$, the stretched exponential captures reasonably well the dependence of $D_p$ on $M$ at fixed $P$, but at fixed $M$ and $c$ does less well at capturing the dependence of $D_p$ on $P$.

Nyden, et al\textsuperscript{86} used PFGNMR to determine probe diffusion of monodisperse ($M_w/M_n < 1.1$) polyethylene oxides (molecular weights 10-963 kDa) diffusing through aqueous solutions of 100kDa ethylhydroxyethylcellulose. The authors studied 1% and 6% solutions and a 1% chemically cross-linked gel. Figure 53 shows $D_p/D_o$ of the polyethylene oxides in 1% of the matrix polymer as a function of their hydrodynamic radii. The solid line in the Figure represents a stretched exponential

$$D_p/D_{p0} = D_1 \exp(-\alpha R^\delta)$$

in probe hydrodynamic radius $R$, where $\alpha$ and $\delta$ are a scaling prefactor and exponent, $D_{p0}$ is the diffusion coefficient of the probe polymer in pure solvent, and $D_1$ is the probe diffusion coefficient for a nominal $R = 0$ polymer chain. The best-fit parameters were $D_1 = 0.33$, $\alpha = 0.33$, and $\beta = 0.57$; the RMS fractional error in the fit was 7%. Nyden, et al\textsuperscript{86} note $R = KP^{\alpha}$ with $\alpha = 0.53$ for probes having molecular weight $P$, so equation 19 is equivalent to a stretched exponential in $P^{\gamma}$ with $\gamma \approx 0.30$. This value of $\gamma$ is consistent with values for $\gamma$ found for other systems, as seen in Table III.

Nyden, et al\textsuperscript{86} also examined probe diffusion in 6% solutions of their matrix polymer. $dD_p/dM$ does not change monotonically with increasing $M$. However, at the larger $M$ at which the anomalous behavior occurs, the PFGNMR echo decays are no longer simple exponentials, implying that diffusive behavior has become more complex.

Pinder\textsuperscript{87} reported from PFGNMR measurements the tracer diffusion coefficient of styrene and low-molecular-weight probe polystyrenes ($P \leq 2470$ Da) through solutions of deuterated polystyrenes ($10.7 \leq M \leq 430$ kDa) in theta (cyclohexane) and non-theta (CCl$_4$) solvents. $D_p$ was measured as a function of matrix polymer concentration for matrix concentrations up to 300 g/L. The matrix polymer polydispersities were $M_w/M_n \leq 1.14$, with in most cases $M_w/M_n < 1.08$.

Figure 54a shows the tracer diffusion coefficient of styrene in polystyrene solutions. The matrix molecular weights were 68 and 200 kDa in the non-theta solvent and 68, 87, 200, and 430 kDa in the theta solvent. Within the scatter in the data—there are not a large number of data points for any particular matrix $M$—$D_p$ does not appear to depend on $M$. $D_p(c)$ was fit both to pure and
stretched exponentials in $c$. As seen in Table III there appears to be very little improvement in the fit on allowing $\nu \neq 1$, so the Figure shows the fits to the pure exponentials in $c$.

Figure 54b shows the tracer diffusion coefficient of styrene and three low-molecular-weight polystyrenes ($P \leq 2470$ Da) through a range of high-molecular-weight polystyrenes ($10.7 \leq M \leq 430$ kDa). For each probe, matrix molecular weights were varied by factors of 3-5 with no apparent significant effect on $D_p$. In all cases, $M/P$ was in the range 15-20 or larger. $D_p(c)$ for each probe was fit to a separate stretched exponential, giving excellent results reported in Table II. The merged results for all four probes were also fit jointly to a stretched exponential in $c$ and $P$. The computed $D_p$ for styrene monomer underestimates $D_p(c)$ by a nearly constant multiplicative factor. Because styrene is little larger than a solvent molecule, a second fit including only the polymer probes was made. Parameters for both fits are in Table IV. The Figure shows the second fit, to the three polymeric probes. The second fit has a modestly better RMS fractional error, and finds in the limit of zero matrix concentration that $D_p \sim P^{-0.52}$. If the styrene monomer is included in the fits, in the same $c \rightarrow 0$ limit $D_p \sim P^{-0.68}$ is obtained.

Smith, et al. used fluorescence recovery after pattern photobleaching to measure the diffusion of labeled 33.6 kDa polypropylene oxide (PPO) chains through solutions of unlabeled 32kDa PPO chains dissolved in a melt of 1 kDa PPO chains. The probe and solvent were relatively monodisperse ($M_w/M_n = 1.1$) while the matrix polymer was relatively polydisperse ($M_w/M_n = 1.6$). The matrix concentration was varied all the way from dilute solution up to the matrix melt. Smith, et al.’s data are shown in Fig. 55. The solid line in the Figure is a stretched-exponential fit using parameters in Table III. A single stretched exponential with constant parameters describes the concentration dependence of $D_p$ all the way from dilute solution up to the matrix melt.
FIG. 55: $D_p$ of 33.6 kDa labeled polypropylene oxide chains through solutions of 32 kDa PPO chains in a 1 kDa PPO melt, using results of Smith, et al.\textsuperscript{88}, Figure 1. The solid line represents a stretched-exponential fit.

Tead and Kramer\textsuperscript{89} studied diffusion of 255 kDa deuterated polystyrene through solutions of large-molecular-weight (93, 255, and 20 000 kDa) polystyrenes dissolved in a low-molecular-weight (10 kDa) polystyrene melt. The matrix polymer volume fraction covered the full range $0 \leq \phi \leq 1$. The temperature was 150°C for the 93 and 255 kDa matrix polymers and 175°C for the 20 000 kDa matrix polymer. Probe diffusion coefficients were obtained using forward recoil spectroscopy to measure concentration profiles of the probe molecules as a function of time. As seen in Fig. 56, $D_p$ is represented well by a stretched exponential in the matrix polymer concentration $c$, no matter whether the matrix molecular weight is less than, equal to, or far larger than the probe molecular weight. The agreement with the stretched-exponential form is good from low concentrations of a short (93 kDa) chain out to high concentrations (melt) of a very large (20 000 kDa) chain.

Tinland and Borsali\textsuperscript{90} used fluorescence recovery after photobleaching and quasielastic light scattering to make independent measurements of the probe diffusion coefficient of 433 kDa dextran through solutions of 310 kDa polyvinylpyrrolidone in water. PVP concentrations ranged from 0 to 120 g/L. The polydispersity $M_w/M_n \approx 1.5$ for the matrix polymer but ca. 1.9-1.95 for the probe chains. Except perhaps at the very highest concentrations studied, values of $D_p$ from the two techniques do not agree. In the following analysis, we use $D_p$ as obtained from FRAP, since the values from this technique do not rely so heavily on detailed model assumptions of the relationship between the scattering spectrum and the underlying diffusion coefficients. Tinland and Borsali’s data appear in Fig. 57, with data points from the original paper\textsuperscript{90} and a smooth curve showing the stretched exponential in $c$ generated from the fitting parameters in Table II. Agreement between the measurements and the functional form is good at all polymer concentrations.

Wheeler, et al.\textsuperscript{91} studied tracer diffusion of linear polystyrenes having molecular weights 65, 179, 422, and 1050 kDa (with $M_w/M_n \leq 1.1$) through a 1.3 MDA polyvinylmethylether matrix polymer, $M_w/M_n \approx 1.6$, in orthofluorotoluene. $D_p$ was determined using quasielastic light scattering, which was possible because the polystyrenes were present at trace concentration while the matrix polymer and solvent are isorefractive. Matrix concentrations covered the range 1–100 g/L. For this matrix polymer, $1/[\eta] = 2.2$ g/L, so much but not all of the data is in the range $c > c^*$. The data and corresponding stretched-exponential fits are in Figs. 58a and 58b, with fit parameters in Tables II and III, respectively. Figure 58c shows fits made separately for each probe polymer. As in the other systems discussed above, the concentration dependence of $D_p$ for each probe is described extremely well by a stretched exponential in polymer concentration. Figure 58d shows the outcome of a simultaneous fit of a single stretched exponential in $c$ and $P$ to all data on all four probes. At the scale of the Figures, the curves in the two Figures for the 179 and 422 kDa probes are coincident, while the curves for the 1050 kDa probes are very close. However, for the smallest probe and large matrix concentrations the joint
stretched exponential substantially underestimates $D_p$.

Wheeler and Lodge\textsuperscript{92} used QELSS to measure the diffusion of linear polystyrenes through polyvinylmethylether:orthofluorotoluene. The polystyrenes had molecular weights of 65, 179, 422 and 1050 kDa, with polydispersities $M_w/M_n < 1.1$. The polyvinylmethylether samples had molecular weights of 140, 630, and 1300 kDa; concentrations of these matrix polymers ranged up to 300 g/L. Reference \textsuperscript{92} represents a major extension of Ref. \textsuperscript{91} in the range of matrix concentrations, number of matrix molecular weights, and number of concentrations studied. For the three matrix polymers, $c^*$ was estimated at 11, 5.7, and 3.3 g/L, respectively (based on $c^* = 1.5/[\eta]$), while $c_e$ was estimated at 50, 12, and 6 g/L, respectively. The matrix polymers had $M_w/M_n \approx 1.6$.

Wheeler and Lodge also used PFGNMR to measure $D_{s0}$ of orthofluorotoluene diffusing through the 1300 kDa polyvinylmethylether at matrix concentrations up to 300 g/L. $D_{s0}$ fell by 62\% over this concentration range. To good approximation, $D_{s0}(c)$ of the orthofluorotoluene is fit by a simple exponential $D_{s0} = 2.15 \times 10^{-5} \exp(-2.98 \times 10^{-3} c)$. Wheeler and Lodge\textsuperscript{92} and also Lodge, et al.\textsuperscript{79} used these data to modify $D_p$ of their polystyrene polymer probes to remove the concentration dependence of a nominal local friction $\zeta = k_B T/D_{s0}$. This local friction modification was here removed from the probe diffusion data of Refs. \textsuperscript{92,79} before analyzing them further.

Figures \textsuperscript{91,92} show Wheeler, et al.'s data\textsuperscript{92} to a joint stretched exponential in $c$ and $P$. The fits are uniformly extremely good, with RMS fractional errors in the range 2-12\% and fitting parameters as seen in Table \textsuperscript{11.}

We also fit all of Wheeler, et al.'s data\textsuperscript{91,92} to a joint stretched exponential in $c$, $P$, and $M$, as seen in Figs. \textsuperscript{91,92} The solid lines are the best-fit $D_p(c)$, plotted for fixed $P$ and $M$, all lines being computed from a single set of parameters given in Table \textsuperscript{11}.
FIG. 59: $D_p$ of polystyrenes (from top to bottom, $M_w$ of 65, 179, 422, and 1050 kDa) diffusing through solutions of (a) 140 kDa and (b) 630 kDa polyvinylmethylether:orthofluorotoluene based on data of Wheeler, et al.\textsuperscript{92}, and (lines) separate stretched-exponential fits for each $P : M$ combination.

from Ref.\textsuperscript{92} were supplemented with the data of Ref.\textsuperscript{91}, the latter reference reported $D_p$ for all probes in 1300 kDa polyvinylmethylether solutions having $c \leq 100$ g/L.

Over an order of magnitude in matrix molecular weight, a factor of 15 in probe molecular weight, and a factor of 300 in matrix concentration, the joint stretched exponential of eq\textsuperscript{17} represents reasonably well the joint dependence of $D_p$ on $c$, $P$, and $M$. The RMS fractional error in the fit is 25%. As noted above in the analysis for Ref.\textsuperscript{61}‘s data, for the smallest probe in the 630 and 1300 kDa matrix polymers (Fig.\textsuperscript{51}), the stretched exponential form noticeably underpredicts $D_p$. A similar issue arises for the 1050kDa probe in the 140 kDa matrix polymer(fig.\textsuperscript{61}): at large $c$ the predicted $D_p$ is too small, because the predicted curve does not bend quite sharply enough between small and large $c$.

V. OTHER EXPERIMENTAL STUDIES

In addition to the work reviewed above, a variety of other studies of polymer tracer diffusion appear in the literature. These are papers whose experimental foci are not the same as those of the papers examined in the previous sections, and which are not amenable to analysis on the lines employed above. For example, a number of papers report scattering spectra of polymer: solvent mixtures in which neither polymeric species is dilute, and therefore in which the spectral relaxation times do not correspond to the self-diffusion coefficient.

Aven and Cohen\textsuperscript{93} measured the diffusion of dilute polystyrenes through 15 vol% solutions of polydimethylsiloxane in tetrahydrofuran. The tracer diffusion coefficients of the polystyrenes, the molecular weight dependence, and the initial linear slope of the concentration dependences were obtained with light scattering spectroscopy.

Borsali, et al.\textsuperscript{94} studied QELSS spectra of 970 kDa polystyrene: 950 kDa polymethylmethacrylate: toluene, PMMA and toluene forming an isorefractive pair. Semi-quantitative experimental tests were made of the theoretical work of Benmouna, et al.\textsuperscript{21}. When neither polymer was dilute, the observed spectrum was biexponential. The mode amplitude ratio and relaxation times were within a factor of two of predictions of the Benmouna model\textsuperscript{21}. The mutual diffusion coefficient of polystyrene in toluene and the cooperative diffusion coefficient of the mixture, both measured at the same total polymer concentration, are equal to within 6%, also in agreement with the theory.

Borsali, et al.\textsuperscript{95} extended their work on this ternary system with measurements at several large (weight fraction > 0.8) polystyrene concentrations and a range of total polymer concentrations, finding two relaxational modes in QELSS spectra. The diffusion coefficient associated with the fast mode increased with increasing polymer concentration. The diffusion coefficient associated with the slow mode decreased markedly with increasing polymer concentration. Results were consistent with the Benmouna, et al.\textsuperscript{21} model.

Borsali, et al.\textsuperscript{96} also studied mixtures of polystyrene and polydimethylsiloxane in tetrahydrofuran (which is isorefractive with polydimethylsiloxane) and in toluene, which is a zero average contrast solvent for these polymers and conditions. Measured spectral forms (one or two relaxations), relaxation times, and their concentration dependences were consistent with the Benmouna model\textsuperscript{21}.
In an extremely important series of experiments, Chang, et al. report probe diffusion by polystyrene through polyvinylmethylether: toluene, using both QELSS and FRS to determine diffusion coefficients. The two techniques differ markedly in the length scales they examine, scattering vectors \( q \) for the two techniques being \( 10^{10} \leq q^2 \leq 10^{11} \text{ cm}^{-2} \) for QELSS but only \( 10^7 \leq q^2 \leq 10^8 \text{ cm}^{-2} \) for FRS. The relaxation rates \( \Gamma \) were within experimental error proportional to \( q^2 \) over the full range \( 10^7 \leq q^2 \leq 10^{11} \text{ cm}^{-2} \), confirming that QELSS does measure a simple translational diffusion coefficient, even though the probe displacements sampled by QELSS are smaller than the diameter of the probe’s correlation hole in the solution. Any claim that probe motion is more rapid, inside the probe’s correlation hole, than it would be over larger distances, must therefore explain how Chang, et al.’s data are consistent with the claim. Chang, et al. also demonstrate that the initial dependence of the measured diffusion coefficient on the probe concentration depends strongly on matrix concentration, the initial slope being substantially positive in the absence of matrix polymer and becoming significantly negative as polystyrene concentration is increased.

B. Chu, et al. report on the diffusion of polymethacrylate probes through a polystyrene matrix. The solvent was a mixture of toluene and \( \alpha \)-chloronaphthalene. By adjusting the solvent composition and the temperature, it was possible to make a virtually exact match of the indices of refraction of the mixed solvent and the matrix, so that scattering arose only from the probe. By varying the solvent composition, the motions of the matrix could separately be examined. A weak fast mode and a dominant slow mode were apparent, even at low angles, for probe chains much larger than the matrix chains. The slow mode had a weaker concentration dependence than does the solvent viscosity.

S. Chu and collaborators studied the motion of fluorescently labelled DNA molecules in non-dilute solution. They propose that they have confirmed the reptation model, based on their observations that (i) stretched DNA chains relax as though they were confined to tubes, (ii) \( D_s \) scales as \( M^{-2} \) for nondilute probe chains, and (iii) a scaling law \( D_s \sim c^{-1.75} \) is consistent with their data.

However, referring to these points seriatim: (i) Even within the reptation model a stretched chain relaxes from stretching primarily via its higher-order modes, revealing
only tube confinement on a short time scale. (ii) The scaling result $D_s \sim M^{-2}$ is a common feature of many models, is in no sense a signature of reptation, and in any event is inconsistent with the modern analysis of data on synthetic polymers. (iii) Ref. 58's measurements of $D_s(c)$ are too scattered to demonstrate $c^{-1.75}$ behavior. Furthermore, if these data had demonstrated concentration scaling, they would be inconsistent with essentially all other data in the literature on the concentration dependences of $D_s$ and $D_p$ of polymers in solution, as reviewed here, in that in almost all other systems $D_s$ and $D_p$ have stretched-exponential, not power-law, concentration dependences.

Finally, as was not emphasized in the original paper, the data of Chu and collaborators conclusively reject the reptation model for their system. According to the reptation model, polymer chains only perform free diffusion after they escape from their tubes, which occurs at times longer than $\tau_d$. At times $t < \tau_d$, polymer chains remain confined to their tubes and perform confined diffusion, so for $t < \tau_d$ a chain’s mean-square displacements satisfy $\langle x^2(t) \rangle \sim t^p$ where $x$ is 0.5 or less. Ref. 58 reports that their chains under their conditions remain within the hypothesized tube for a time $\tau_d \approx 1.2$ or $\tau_d > 2$ minutes, depending on which of several methods is used to estimate $\tau_d$. However, ref. 58 measured directly $\langle x^2(t) \rangle$. Figure 2 of ref. 58 shows $\langle x^2(t) \rangle \sim t^1$ for times as short as $\tau_d/7$ or less. That is, ref. 58 found that their chains were performing free, Brownian, non-constrained diffusion even on time scales sufficiently short that under the reptation model the chains would have been confined to their tubes. This last finding is completely incompatible with the reptation model, which requires that chains perform constrained curvilinear diffusion along their tubes so long as they remain inside their tubes. The data of Chu and collaborators thus indicates that a key aspect of the reptation picture, namely tube confinement, is incorrect, at least in Chu, et al.’s system.

Corrotto, et al.104 performed static light scattering and QELSS measurements on bidisperse nondilute mixtures of polystyrene in toluene, extracting fast and slow mode relaxations.

Cosgrove and Griffith105 used PFGNMR to study the diffusion of protonated polystyrenes through solutions of deuterated polystyrenes, varying the matrix concentration, and the probe and matrix molecular weights. Data was obtained over limited ranges of $D_p$ on the dependence of $D_p$ on each of these variables. $D_p$ generally declined with increasing matrix concentration and molecular weight, and also declined with increasing probe molecular weight, except for very large probes in relatively dilute solutions of small polymers, in which $D_p$ was nearly independent of matrix molecular weight.

Cotts106 report QELSS measurements on polystyrene: polyvinylmethylether: toluene fluided into the systematic studies of Lodge, Wheeler, and collaborators77,78,79,91,92 on this system.

Davis and Pindera106 report QELSS studies of polystyrene: polyvinylmethylether mixtures dissolved in toluene or carbon tetrachloride. The first ternary mixture is nearly unique, in that the polymers are nearly compatible, toluene is a good solvent for both polymers, and toluene and PVME are isorefractive. The second ternary mixture differs from the first in that CCl$_4$ is only a marginal solvent for polystyrene. In CCl$_4$, spectra become nonexponential. The diffusion coefficient in nondilute CCl$_4$ solutions is reduced by more than three-fold relative to diffusion by the same concentrations of the two polymers in toluene, but the probe radius of gyration is only very slightly reduced by increasing the matrix concentration.

Desbrieres and collaborators107 applied QELSS to solutions of dextran and polyvinylpyrrolidone in water in solutions more concentrated than the overlap concentration. Two modes whose properties are consistent with Benmouna21-type models of diffusion in ternary polymer solutions were observed. The mixtures have a phase separation at elevated concentration. As polymer concentrations are increased towards the phase separation, a third slow mode whose various properties are consistent with the formation of aggregates was observed.

Giebel and co-workers108,109 studied QELSS spectra of polydimethylsiloxane and polymethylmethacrylate in several solvents as a function of the relative concentration of the two polymers. At the fixed total polymer concentration, the polymers were reasonably expected to be non-dilute. Comparison was made with theoretical results of Benmouna, et al.21, with particular attention to the “zero average contrast” condition. A strong variation of some spectral parameters with composition was described well by Benmouna-type models and a small number of free parameters.

Jamil, et al.110 report QELSS measurements on the diffusion of random-coil polystyrene probes through solutions of the rigid-rod polymer poly-($\gamma$-stearyl aL-glutamate) in its isorefractive solvent toluene. The polymers are incompatible, poor miscibility of rods with random coils being identified by Jamil, et al.110 as a significant experimental challenge. Experimentally, the light scattering spectrum in these systems is due entirely to the polystyrene probes.

Jamil, et al.110 found that the dominant slow mode of the QELSS spectra is strongly concentration-dependent. At lower matrix concentration the diffusion coefficient corresponding to the slow mode increases with increasing probe concentration. At elevated matrix concentration this diffusion coefficient instead decreases with increasing probe concentration. Extrapolating this diffusion coefficient to zero probe concentration gives the tracer-diffusion coefficient of the probe polymer through the rod matrix. Jamil et al. reported fitting their six values of $D_s$ to a stretched-exponential form.
\( \alpha \approx 0.4 \) and \( \nu \approx 1.3 \). Jamil, et al. interpret the finding \( \nu > 1 \) as arising from end-to-end aggregation of the matrix polymer; the matrix polymer increases its hydrodynamic radius as its concentration is increased.

Konak, et al. report QELSS spectra of mixtures of polystyrene and polymethylmethacrylate in toluene. Neither polymer was dilute. Comparison was made for a limited number of concentrations with theoretical models arising from work of Bemouna, et al. Treating spectra as bimodal, the ratio of relaxation times was predicted theoretically to better than 50\%, but predictions of the mode amplitude ratio were often inexact by factors of 2 or 3.

Marmonier and Legend report extensive measurements using FRS on tracer diffusion of labelled polystyrenes through polystyrene in a good solvent. Unfortunately, the reported data were modified by dividing them by an unreported concentration-dependent factor, namely the normalized concentration-dependent tracer diffusion coefficient of the free label in the same polymer solutions. It is therefore impossible to compare these measurements with other papers analyzed here.

Numasawa, et al. report scattering spectra of a series of dilute polystyrenes in polymethylmethacrylate (in its isorefractive solvent benzene). The major focus was the scattering-vector dependence of the linewidth \( \Gamma \), especially at larger scattering vectors \( q \). At larger \( q \), \( \Gamma / q^2 \) of the self-diffusive mode increases with increasing \( q \). Measurements were only made at a single non-dilute matrix concentration, preventing further analysis of this data along the lines of this review.

Russo, et al. report self-diffusion coefficients for poly(\( \gamma \)-benzyl-\( \alpha \)-L-glutamate) in pyridine. The rodlike polymer has an isotropic-cholesteric liquid crystal phase transition with increasing concentration. Russo, et al. found that \( D_s \) decreases with increasing polymer concentration until the phase transition is reached. At the phase transition, \( D_s \) increases abruptly; it then decreases again as the polymer concentration is further increased. \( D_s (c) \) is qualitatively consistent with a stretched exponential form in the isotropic phase, but the number and spacing of points in the isotropic regime limits the accuracy of a quantitative fit to this very interesting data.

Scalaletar, et al. used FRAP and FCS to study diffusion of phage \( \lambda \) DNA solutions. By comparing systems in which either few, or almost all, molecules were labelled, Scalaletar, et al. were able to confirm the prediction of this author that if very few macromolecules are labelled, fluorescence correlation spectroscopy determines their tracer diffusion coefficient, but if almost all macromolecules are labelled, fluorescence correlation spectroscopy determines their mutual diffusion coefficient.

Sun and Wang report a series of studies of polystyrene/polymethylmethacrylate mixtures (in benzene, dioxane, and toluene, respectively) using QELSS as the major experimental technique. Both polymers were in general nondilute. Neither polymer is isorefractive with any of the solvents. The objective was to study the bimodal spectra that arise under these conditions and to show that the two relaxation times and the mode amplitude ratio can be used to infer diffusion and cross-diffusion coefficients of the two components. Experimental series varied both the total polymer concentration and the concentration ratio of the two components. The theoretical model predicts a biexponential spectrum; the experimental data was fit by a bimodal distribution of relaxation rates or by a sum of two Williams-Watts functions. The inferred self-diffusion coefficients of both species fall with increasing polymer concentration, but the concentration ranges that Sun and Wang studied are too narrow for further interpretation.

VI. ANALYSIS

The above sections have presented a detailed examination of nearly the entirety of the published literature on polymer self-diffusion and probe diffusion in polymer solutions. The dependences of \( D_s \) and \( D_p \) on polymer concentration, probe molecular weight, and matrix molecular weight have been determined. Some features of this literature are incidental consequences of the chemical identity of the polymer being studied. The objective of this Section is to extract systematic behaviors from the above particular results. Here we ask: If we rise above particular features determined by the identity of the polymer under examination, what are the ideal features common to self- and probe-diffusion of all polymers in solution?

In the following: First, the functional forms of the concentration and molecular weight dependences of the self- and probe diffusion coefficients are considered. Second, having found that \( D_s \) and \( D_p \) uniformly follow stretched exponentials in \( c \), correlations of the stretched-exponential scaling parameters with other polymer properties are examined. Third, for papers in which diffusion coefficients were reported for a series of homologous polymers, we examine a joint function of matrix concentration and matrix and probe molecular weights and what it reveals about polymer diffusion. Fourth, we examine the few cases in which stretched-exponential behavior is not seen, or in which particular features of a given system clarify the systematic behavior of the phenomenological parameters used to describe \( D_s \) and \( D_p \). Finally, other results implying a generalized phenomenology for aspects of diffusion behavior are examined.

First, in the above I have reviewed virtually the entirety of the published literature on self-diffusion and probe diffusion of random-coil polymers in solution. As seen from the Figures in the preceding sections, the concentration dependences of \( D_s \) and \( D_p \) are essentially always described well by stretched exponentials (eq. 16) in the matrix concentration \( c \).

Correspondingly, scaling (power-law) behavior is clearly rejected by almost the entire published litera-
ture on polymer self- and probe–diffusion. On a log-log plot, a stretched exponential appears as a smooth curve of monotonically varying slope. In contrast, on a log-log plot, scaling (power-law) behavior would appear as a straight line. Almost without exception, log-log plots of real measurements of \( D_s(c) \) give smooth curves, not straight lines. Power laws could be fit to reported data, but in almost every case the power law would only provide an accurate description of a tangent to the data over some narrow range of concentrations.

An observation that experimental data is described very well by a particular mathematical form does not prove that the form in question is physically significant. In principle it may be the case that several different mathematical forms describe, to within the actual experimental error, the same data. However, the observation that experimental data is uniformly not described by some mathematical form, to well beyond experimental error, is good evidence that models that predict the mathematical form are inadequate. Power laws appear on log-log plots as straight lines. As seen from the above 70+ figures, straight lines are almost never observed in plots of \( D_s \) or \( D_p \) against \( c \) or other physical variables. Correspondingly, the concentration dependence of \( D_s \) and \( D_p \) almost never shows scaling (power-law) behavior. Scaling models that predict or assume for polymer solutions that \( D_s \) and \( D_p \) follow power laws in \( c \), \( P \), and/or \( M \) are very definitely rejected by almost the entirety of the published literature on \( D_s \) and \( D_p \).

Second, referring to eqs (10) there are systematic correlations between scaling parameters \( \alpha \) and \( \nu \) and the solution variables \( P \) and \( M \). These correlations reflect the concentration and molecular weight dependences of \( D_s \) and \( D_p \). In particular:

**The scaling prefactor \( \alpha \) depends strongly on \( M \).** Figure 61 shows the scaling prefactor \( \alpha \) from measurements of the self-diffusion coefficient and fits to eq. (10) as plotted against polymer molecular \( M \). The Figure, based on Table I, shows almost all data on linear and star polymers, with concentrations in g/L. Results from the one system with a large-concentration phase transition are omitted. \( M \) varies over nearly three orders of magnitude; \( \alpha \) varies over almost four orders of magnitude. While there is substantial scatter, the figure is consistent a power-law correlation between \( \alpha \) and \( M \). The solid line in the Figure is a best fit to measurements on linear polymers. It shows \( \alpha = 2.45 \cdot 10^{-4} M^{1.10} \) with \( M \) in kDa.

It is also possible to examine the correlation of \( \alpha \) with the size of the polymer, as reflected, e.g., by the diffusion coefficient \( D_o \). However, the variation in \( D_o \) from system to system arises in part from differences in the measurement temperature and solvent viscosity. To eliminate these effects, \( D_o \) was used to compute a nominal chain hydrodynamic radius

\[
R = \frac{k_B T}{6 \pi \eta D_o}
\]  

(20)

Here \( k_B \) is Boltzmann’s constant and \( T \) is the absolute temperature. The solvent viscosity \( \eta \) was taken from standard tables and interpolated as necessary to actual temperatures. In a few cases, reported experimental conditions do not permit an accurate conversion from \( D_o \) to \( R \); these cases are not considered further.

Figure 62 plots \( \alpha \) as a function of \( R \). In Fig. 62 the
solid lines are best fits for the data on linear chains (filled circles) to \( \alpha \sim R^n \) for \( n = 3 \) or 4. If \( \alpha \sim |\eta| \) as predicted by the Adler model\(^{43}\), then \( n = 3 \) is expected. The hydrodynamic scaling model\(^{44,45}\) instead predicts \( n = 4 \). The fit with \( n = 3 \) gives a modestly better fit to the data than does the \( n = 4 \) fit, but neither fit is visibly inconsistent with experiment.

Figures 61 and 62 also present \( \alpha \) of 3-, 8- and 18-armed stars. \( \alpha \) of an 18-armed star polymer tends to be somewhat smaller than \( \alpha \) for a representative linear chain. However, \( \alpha \) of star polymers almost always lies within the scatter in the values for \( \alpha \) observed for the linear chains. \( \alpha \) for three-armed stars includes result\(^{63}\) in which only a limited number of concentrations were studied for a given polymer.

FIG. 63: Scaling pre-factor \( \alpha \) as a function of \( M \) for data from Refs. (○)\(^{54}\) for polymers in CC\(_4\), (●)\(^{47}\), (□)\(^{48}\), (■)\(^{55}\), (△)\(^{56}\), (▲)\(^{60,65}\) with linear chains, (+)\(^{62}\) with \( f = 3 \), (×)\(^{57}\) with \( f = 8 \), (⪯)\(^{66}\) with \( f = 18 \), (♦)\(^{49}\), (◇)\(^{53}\) for polymers in C\(_6\)D\(_6\), (▽)\(^{67}\), (⊗)\(^{68}\), (△)\(^{50}\), (◇)\(^{11}\) for linear polybutadiene, (♦)\(^{69}\) for \( f = 3 \) polybutadiene, (>)\(^{70}\) for PEO in water, and (▼)\(^{71}\) for xanthan in water. Other details as in Fig. 61.

Figures 61 and 62 emphasize polymer topology: one point style each for linear, 3-, 8-, and 18-armed chains. For some purposes, identifying the points by reference (and, hence, by chemical system and experimental method) is more useful. Figures 61 and 62 give \( \alpha \) against \( M \) and \( R \), respectively, with points labelled by reference, this time including results of Tinland, et al\(^{40}\) on a system with a large-c phase transition. The correlation between \( \alpha \) and molecular weight is somewhat better that the correlation between \( \alpha \) and the inferred – in most cases, not directly measured – chain radius.

The scaling exponent \( \nu \) depends on \( M \) at low \( M \). Figure 62 shows the dependence of \( \nu \) on polymer molecular weight. The filled circles refer to linear polymers as studied in refs. 41, 42, 43, 44, 47, 52, 53, 56, 62, 64, 65, 66, but excludes systems\(^{61}\) in which there is a phase transition at elevated polymer concentration. As seen from the Figure, for polymers larger than 250 kDa \( \nu \) approaches very closely to 0.5. For smaller polymers, \( \nu \) is substantially scattered, but increases with decreasing \( M \).

Sixteen systems in Table I, including linear and three-armed star polymers, had \( \nu \) forced to 1.00 during the fitting process. In these fits \( \nu = 1 \) was forced because the data would not support more free parameters, or because the fit with \( \nu \) as a free parameter did not have a significantly better root mean square fractional error in the fit than did the fit with \( \nu = 1 \) forced. All but four of these fits refer to polymers with \( M < 200 \text{ kDa} \). The circumstance that successful fits with \( \nu \) forced to unity are largely found with polymers of lower molecular weight is consistent with the interpretation that \( \nu \) increases toward 1.0 at small \( M \). These fits are represented in Figure 65 by the open circles.

FIG. 64: Scaling pre-factor \( \alpha \) as a function of \( R \). Other details as in Fig. 63.

Figure 65 also shows the molecular weight dependence of \( \nu \) for star polymers in solution. The data on stars are substantially confined to smaller polymers (\( M < 250 \text{ kDa} \)). However, a trend in \( \nu \) with increasing \( M \) is apparent. At very small \( M \), \( \nu \) is substantially scattered around 1.0. For \( M \geq 200 \text{ kDa} \), \( \nu \) for star polymers appears from the few points to be trending toward 0.5.

Third, several papers\(^{42,67,71,73,74,75,77,78,79,80,82,83,84,85,87,91,92}\) report \( D_g \) and \( D_p \) for a series of homologous polymers with different molecular weights. Each of these papers describes a study made by a given method using consistent operating conditions and data analysis procedures, thereby avoiding data scatter arising from any practical...
experimental issues. To use these studies to obtain information about the molecular weight dependences of $D_s$ and $D_p$, data from each paper were separately fit to \textit{eq. 17} which is a joint stretched exponential in \(c\) and $M$ and $P$. (For $D_s$, $P$ and $M$ are the same variable.) As seen Sections III and IV, eq \textit{17} with parameters in Table \textit{III} fits most of each data set well.

Some studies were limited in the range of matrix concentrations that was explored, either by not going to very high concentration or by not making measurements at low concentration. These studies are excluded here from further consideration. Some studies are subsets of other studies, for example cases in which measurements were first made at fixed $M$ for a range of $P$, and then extended to cover a range of $M$. To avoid double-counting experiments, data sets that are subsets of other data sets are not considered further here.

Section IV described fits of $D_p$ to \textit{eq. 17}. Much of the probe diffusion data is represented well by this joint stretched exponential in \(c\), $P$, and $M$. However, \textit{eq. 17} is less accurate when $M$ and $P$ differ substantially. In particular, for $M/P > 3$ or for $P/M > 3$, \textit{eq. 17} tends to overstate the decrease in $D_p$ with increasing $c$. Specific results of Section IV pertaining to the range of validity of \textit{eq. 17} include:

Brown, et al.'s data for $D_p/D_{p0} < 10^{-3}$ show an experimental $D_p$ that is smaller than the calculated value from a fit to \textit{eq. 17}. Also, for $P/M \approx 80$, the measured $D_p$ is larger than predicted. Hanley, et al.'s data and Kent, et al.'s data (all follow well \textit{eq. 17}'s $P$ and $M$ dependences. Kim, et al.'s extensive results on the effects of varying $P$ and $M$ indicate that \textit{eq. 17} generally works well for $M/P \leq 3$, but overstates the dependence of $D_p$ on $M$ for $M/P > 3$. From this data, it appears that as $c$ is increased there is a narrowing of the range of values of $M/P$ for which \textit{eq. 17} works well.

For $M/P$ never larger than 3 or so, Lodge and Wheeler found that the joint stretched exponential worked well for linear polystyrene chains and also worked well for $f = 3$ polystyrene stars, both diffusing through linear-chain matrices. For large $f = 12$ polystyrene stars in a short linear polyvinylmethylether, Lodge and Markland's data follow a joint stretched exponential, even for $P/M \approx 12$. If the comparison is made by arm rather than total molecular weights, Lodge and Markland’s data is confined to a region $P_r/M_r < 2$. Lodge, et al. examined the diffusion of small $f = 12$ stars through solutions of a large linear polyvinylmethylether. For $M/P \geq 3$ and elevated concentrations, the measured $D_p$ is larger than expected from the joint stretched exponential. Lodge, et al. also studied three-armed stars diffusing through linear chains; with increasing matrix molecular weight, the joint stretched exponential predicts too small a $D_p$ for $M/P > 3$.

Martin reported $D_p$ of linear polystyrenes diffusing through linear polyvinylmethylether. Equation \textit{17} fits well (Fig. 49) the joint data, but throughout this data $M/P < 2.2$. Nemoto, et al. studied diffusion of PMMA through polystyrene solutions; the joint fit (Fig. 50) of all data to \textit{eq. 17} was more accurate for $2 > M/P > 0.5$ than it was outside this range. Numasawa, et al.’s data on polystyrene in polymethylmethacrylate-benzene, Fig. 62 is described reasonably well by \textit{eq. 17}. However, for Numasawa, et al.’s data the joint stretched exponential clearly gives a better description of the concentration dependence of $D_p$ than it does of the $M$-dependence. Wheeler, et al. report on linear polystyrenes diffusing through polyvinylmethylethers. Fits to their measurements (Figs. 59 and 60) show that \textit{eq. 17} underestimates $D_p$ for $M/P \geq 7$ and for $P/M \geq 3$.

The joint stretched exponential thus describes well a great number of measurements of $D_p$ in systems with $0.3 < M/P < 3$. Outside this range, \textit{eq. 17} predicts a concentration dependence for $D_p$ that is stronger than that found experimentally. Corresponding to each set of measurements are a set of scaling exponents and prefactors. To what extent do these parameters show universal rather than system-specific behavior? In \textit{eq. 17} data were parameterized as $D_oM^{-\nu} \exp(-\alpha c^pP^f M^3)$. For each system, the exponent $\nu$ was treated as a single constant, the dependence of $\nu$ on $M$ at small $M$, discussed above, being suppressed. This suppression will reduce the quality of the fits when a very wide range of molecular weights is studied, as will be seen below in one system.

From Table \textit{III} (i) $a$ is almost always in the range $0.5 < a < 1.5$.
0.5 to 0.6. (ii) \( \nu \) is generally in the range 0.5-0.75. In two cases, \( \nu \) was found to be larger (0.86, 0.99) than this range; in one case it was smaller (0.43). In a few cases, \( \nu = 1 \) was forced during the fitting process, rather than being obtained from the fit. (iii) For linear chains, \( \gamma \) is almost always in the range 0.25-0.3. There are outliers at 0.14, 0.19, and 0.43; each of these is associated with a system in which \( \nu \) was unusually small (0.43) or unusually large (0.86, 0.99). For star polymers, \( \gamma \) is \( \approx 0.15 \). (iv) For linear polymers and three-armed stars, \( \delta \) is usually in the range 0.25-0.30.

**Fourth**, in all of two cases, concentration and molecular weight dependences were not, or were not necessarily, described by stretched exponentials. First, Nemoto, et al.\(^{27\ldots35} \) studied the \( M \)-dependence of \( D_s \) at large fixed \( c \). Their data show a transition from a stretched-exponential to a power-law molecular weight dependence for \( M \geq 800 \) kDa. Second, Tao, et al.\(^{36\ldots42} \) studied very large polymers in highly concentrated solutions. Their data are almost equally well-described by power laws and by stretched exponentials, as seen in Figs. 13 and 14. For this data, \( \alpha \) of eq. 16 is nearly independent of \( M \), contrary to the behavior of \( \alpha \) in other systems. The lack of an \( M \)-dependence in \( \alpha \), and the ability of a power-law to describe this data, are consistent with the proposal\(^{36\ldots42} \) that polymer transport coefficients have different phenomenologies in the melt/near-melt domain and in the solution domain here.

**Fifth**, there is evidence for other correlations.

Several papers that report \( D_s \) or \( D_p \) also report the solution viscosity \( \eta \). In particular, Martin\(^{71\ldots80} \) gives not only \( D_p \) but also \( \eta \) of solutions of his 110 kDa polyvinylmethyl ether matrix polymer. Martin reported for 50 kDa probe polystyrene that the product \( D_p \eta \) increased dramatically (up to 6-fold) with increasing \( c \). For larger probe polymers (100 kDa, 420 kDa), the increase in \( D_p \eta \) was less dramatic; with 900 kDa probes, ",...this product is very nearly independent of concentration." Martin\(^{41\ldots45} \) interpreted the dramatic increase in \( D_p \eta \) with increasing matrix concentration as arising from a crossover from Stokes-Einsteinian to reptational diffusion, the crossover occurring more readily for \( P \leq M \) than for \( P \gg M \).

\( D_p \), \( \eta \) and probe radius of gyration \( R_g \) for the same set of solutions were reported by Numasawa, et al.\(^{46\ldots55} \), who studied polystyrene diffusing through polymethylmethacrylate-benzene. For a range of matrix concentrations and matrix and probe molecular weights, Numasawa, et al. identified a regime in which \( D_p \eta R_g \) is approximately constant. For small probe chains diffusing through larger matrix chains but not large probe chains diffusing through smaller matrix chains, the product \( D_p \eta R_g \) increases markedly (up to 100-fold) with increasing \( c \) and \( M \). Numasawa, et al.\(^{46\ldots55} \) interpreted their data as showing Stokes-Einstein type diffusion and a crossover with increasing matrix molecular weight to a region in which reptational motion was dominant.

Martin\(^{41\ldots45} \) and Numasawa, et al.\(^{46\ldots55} \) propose that the failure of the Stokes-Einstein equation, with \( D_p \eta \) increasing markedly with increasing \( c \), is associated with a transition to reptation dynamics for large polymer chains. However, the diffusion of rigid spherical probes through high-molecular-weight polymers shows a phenomenology highly similar to the phenomenology for \( D_p \eta \) observed\(^{46\ldots55} \), namely \( D_p \eta \) increases dramatically with increasing \( c \) at large \( M \), even though there is no possibility that spheres have a transition permitting them to diffuse via reptation.

Like effects do not prove that like causes are at work. The existence of non-reptating spherical probe-polymer systems in which \( D_p \eta \) increases markedly with increasing \( c \) does not prove that polymer systems showing the same phenomenology are not reptating. However, non-reptating sphere-probe systems can have a \( D_p \eta \) that increases markedly at large matrix \( c \). Therefore, the observation that \( D_p \eta \) for probe polymers in a polymer solution increases markedly at large matrix \( c \) is not evidence that the polymer solution has had a transition to reptation dynamics.

Xuexin, et al.\(^{56\ldots66} \) report \( D_p \) for linear and \( f = 18 \) star polymers having very nearly the same \( D_o \), i.e., very nearly the same hydrodynamic radius. For these two polymers, the scaling parameters \( \alpha \) and \( \nu \) are also very nearly the same, consistent with but not proving the interpretation that \( \alpha \) and \( \nu \) are determined by chain size and not by chain topology.

Kent, et al.\(^{46\ldots55} \) examined \( D_p \) for a range of probe as well as matrix concentrations. For much of their data, Kent, et al.’s probes were concentrated enough that \( D_p \eta \) manifestly depended on the probe concentration \( c_p \) as well as on the matrix concentration \( c \). At low \( c \), \( D_p \eta \) increased with increasing \( c_p \). At elevated \( c \), \( D_p \eta \) decreased with increasing \( c_p \). These results provide a specific target for theoretical investigation. Just as good hydrodynamic models\(^{19\ldots21} \) can predict quantitatively the concentration dependence of \( D \) for hard spheres, so also a good hydrodynamic model for polymers should be able to predict the dependence of \( dD_p/dc_p \) on matrix concentration.

**VII. CONCLUSIONS AND DISCUSSION**

In the above, virtually the entirety of the published literature on polymer self-diffusion and on the diffusion of chain probes in polymer solutions has been reviewed. Studies that determined how \( D_s \) and \( D_p \) depend on polymer concentration and molecular weight were systematically re-analysed. Without exception the concentration dependences of \( D_s \) and \( D_p \) are described by stretched exponentials in polymer concentration. The measured molecular weight dependences also compare favorably in most cases with the elaborated stretched exponential, eq. 17. Only when \( P \gg M \) or \( M \gg P \) is there a deviation from eq. 17 that deviation referring only to the molecular weight dependences.

Contrarywise, almost without exception the experimental data is inconsistent with descriptions of concen-
tration or molecular weight dependences of the single-particle diffusion coefficient in terms of the power laws predicted or assumed by scaling models. The published data is sufficient to reject scaling model descriptions of polymer diffusion.

The concentration dependence $\exp(-\alpha c^\nu)$ of exponential-type models for $D_s$ is found to be valid for all ratios of the probe and matrix molecular weights. This validity of the stretched-exponential concentration dependence over a wide range of $P/M$ is consistent with the mathematical structure of the renormalization-group derivation of these forms. Namely, the derivation begins with a low-concentration pseudovirial expansion

$$D_s = D_0(1 + k_2c + k_3c^2)$$

for the concentration dependence of $D_s$, $c$ here being the matrix concentration, and then uses the Altenberger-Dahler positive-function renormalization group method to extend the series from lower to larger concentration. Over a modest range of $P/M$ around $P \approx M$, coefficients $k_2$ and $k_3$ with a simple dependence on $P$ and $M$ are adequate, leading to eq 17. For $P \gg M$ or $P \ll M$, the simple calculations of $k_2$ and $k_3$ of Ref. 36 are inadequate, because they do not include polymer internal modes, so the values of the $k_i$ change. However, for $P \gg M$ or $P \ll M$ the renormalization group method still receives as input a polynomial having the form of eq 21 so it will still generate as output a stretched exponential in $c$ for the concentration dependence of $D_p$.

It is sometimes argued that the success of the stretched exponential form in describing $D_s(c)$ arises from a particular flexibility of the stretched exponential, so that the successes shown in the Figures are accidental. Claims that $D_s \exp(-\alpha c^\nu)$ is 'unusually flexible', relative to other functional forms, are inconsistent with basic mathematics: The stretched exponential describes the concentration dependence with three free parameters. The function is not singular for real $c$ and positive $\nu$. Therefore, the region of function space spanned by the set of all stretched exponentials can be no larger than the region of function space spanned by any other function with three free parameters. Correspondingly, the functional form used here for $D_s(c)$ is no more flexible than other three-parameter forms, so its success in describing $D_s(c)$ cannot be ascribed to an unusual flexibility of the stretched exponential form.

On the other hand, there is no significant region of any set of measurements in which concentration scaling is observed, other than as tangents to smooth curves of monotonically decreasing slope. It can always be asserted that corrections to scaling, of whatever basis, mean that the predicted power-law slopes are only observed over narrow ranges of concentration, or are only true asymptotically in the limit of large molecular weight:

(A) Until the scaling models are refined sufficiently so as to predict the concentrations over which the hypothesized power-law slope should be approximately observed, it is impossible to tell whether the experimental data are consistent with a scaling prediction. It is entirely inadequate to show that the experimental data has, in some region, the slope predicted by a particular power law model, because the observed $D_s(c)$ someplace agrees with a predicted $c^{-\alpha}$ behavior for every single positive $x$ over a very wide range. If one is allowed to draw the asymptote where one chooses, the predicted $x$ thus ceases to be falsifiable. On the other hand, a refined model might very well predict the slope for a range of concentrations in which the data did not have the predicted slope. (B) There is no indication in the experimental data that exponential models cease to be adequate adequate as the polymer molecular weight is increased.

On the other hand, as shown by the extremely thorough experiments of Tao, et al., as one moves from polymer solutions toward the melt, one encounters a region of large $c$ in which scaling models are at least approximately correct. Scaling models therefore may very well predict the slope for a range of concentrations in which the data did not have the predicted slope. (B) There is no indication in the experimental data that exponential models cease to be adequate adequate as the polymer molecular weight is increased.

Acknowledgments

The partial support of this work by the National Science Foundation under Grant DMR99-85782 is gratefully acknowledged.
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### TABLE I: Fits of the concentration dependence of the self-diffusion coefficient $D_s$ of polymers in solution to a stretched exponential $D_s \exp(-|c\rho|^\nu)$. The Table gives the fitting parameters, the percent root-mean-square fractional error %RMS, the polymers/solvent system, and the reference. Square brackets "[ ]" denote parameters that were fixed rather than floated during the least-mean-squares fitting process. $D_s$ is in cm$^2$/s or in units of $D_0$. Concentrations are $g$/L, except "c" in volume fraction units. Polymers include dex–dextran, pB–polybutadiene, HpBD–hydrogenated polybutadiene, pDMS–polymethylsiloxane, PEO–polyethylene oxide, pI–polyisoprene, pS–polystyrene, and xanthan. Solvents include THF–tetrahydrofuran.

| $D_s$ ($10^{-7}$) | $\alpha$ | $\nu$ | %RMS | $M$ (kDa) | System | Refs. |
|-------------------|----------|-------|-------|----------|--------|-------|
| 9.17               | 4.82     | 1.12  | 6.1   | 64.2     | dex:CH$_2$O | 41    |
| 2.24               | 0.036    | [1]   | 5.2   | 73       | PEO:CH$_2$O | 42    |
| 1.16               | 0.077    | 0.94  | 7.5   | 148      | PEO:CH$_2$O | 43    |
| 9.9                | 0.089    | [1]   | 5.9   | 278      | PEO:CH$_2$O | 44    |
| 6.42               | 0.137    | [1]   | 3.4   | 661      | PEO:CH$_2$O | 45    |
| 2.47               | 3.47     | 1.12  | 6.6   | 2        | pS:CCl$_4$  | 43,44  |
| 3.08               | 0.17     | 0.56  | 13    | 110      | pS:CCl$_4$  | 43,44  |
| 3.38               | 0.62     | 0.39  | 7.4   | 233      | pS:CCl$_4$  | 43,44  |
| 2.01               | 0.50     | 0.45  | 8.1   | 350      | pS:CCl$_4$  | 43,44  |
| 3.62               | 0.054    | 0.80  | 6.6   | 110      | pS:CCl$_4$  | 43,44  |
| 4.70               | 0.25     | 0.59  | 5.8   | 233      | pS:CCl$_4$  | 43,44  |
| 3.30               | 0.24     | 0.62  | 6.9   | 350      | pS:CCl$_4$  | 43,44  |
| 4.50               | 0.227    | 0.595 | 11    | 262      | pS:CCl$_4$  | 43,44  |
| 3.7                | 0.355    | 0.55  | 23    | 657      | pS:CCl$_4$  | 43,44  |
| 3.29               | 0.462    | 0.53  | 12    | 861      | pS:CCl$_4$  | 43,44  |
| 1.20               | 0.0111   | 1.05  | 3.4   | 125      | pS:tol     | 47    |
| 4.88               | 0.025    | 0.737 | 3.4   | 15       | pDMS:tol   | 52,53 |
| 8.30               | 0.34     | 0.52  | 9.5   | 530      | pDMS:tol   | 52,53 |
| 5.06               | 0.27     | 0.59  | 1.3   | 730      | pDMS:tol   | 52,53 |
| 6.21               | 0.0848   | 0.64  | 5.1   | 78       | pS:CH$_2$O | 46    |
| 7.90               | 0.21     | 0.52  | 4.0   | 123      | pS:CH$_2$O | 46    |
| 4.26               | 0.13     | 0.67  | 24    | 245      | pS:CH$_2$O | 46    |
| 2.33               | 0.38     | 0.51  | 22    | 599      | pS:CH$_2$O | 46    |
| 2.03               | 0.48     | 0.50  | 13    | 754      | pS:CH$_2$O | 46    |
| 8.15               | 3.68*    | 0.76  | 0.7   | 2        | pE0:CHCl$_3$ | 50    |
| 2.19               | 9.07*    | 0.61  | 3.7   | 40       | pE0:CHCl$_3$ | 50    |
| 2.69               | 7.61*    | 0.78  | 12    | 20       | pE0:CH$_2$O | 50    |
| 6.03               | 4.30*    | 0.67  | 3.5   | 2        | pE0:dioxane | 50    |
| 2.05               | 7.75*    | 0.62  | 4.9   | 20       | pE0:dioxane | 50    |
| 2.71               | 9.72*    | 0.46  | 16    | 40       | pE0:dioxane | 50    |
| 2.30               | 21.9*    | 0.42  | 2.3   | 3000     | pE0:dioxane | 50    |
| 5.00               | 17.1*    | 0.81  | 5.1   | 240      | pS:CH$_2$O | 50    |
| 3.30               | 16.5*    | 0.80  | 11.2  | 240      | pS:CCl$_4$ | 46    |
| 1.65               | 23.5*    | 0.67  | 13.5  | 1300     | pS:CCl$_4$ | 46    |
| 7.62               | 0.186    | 0.79  | 17    | 450      | xanthan:CH$_2$O | 46    |
| 6.24               | 0.45     | 0.91  | 15    | 990      | xanthan:CH$_2$O | 46    |
| 4.18               | 0.63     | 1.00  | 15    | 1900     | xanthan:CH$_2$O | 46    |
| 2.67               | 1.13     | 0.88  | 15    | 3800     | xanthan:CH$_2$O | 46    |
| 1.35               | 3.63     | [1]   | 27    | 9400     | xanthan:CH$_2$O | 46    |
| 6.25               | 4.25*    | 1.42  | 7.7   | 10       | $f = 2$ pI:CCl$_4$ | 62    |
| 5.64               | 1.91*    | 1.21  | 4.6   | 16.4     | $f = 3$ pI:CCl$_4$ | 62    |
| 4.63               | 8.08*    | 1.00  | 2.9   | 39       | $f = 8$ pI:CCl$_4$ | 62    |
| 3.33               | 8.61*    | 0.99  | 3.1   | 92       | $f = 18$ pI:CCl$_4$ | 62    |
| 3.06               | 2.69*    | 0.84  | 2.3   | 104      | $f = 8$ pI:CCl$_4$ | 62    |
| 7.63               | 1.83*    | 1.55  | 13.4  | 10       | $f = 2$ pI:CCl$_4$ | 52    |
| 3.35               | 5.3*     | 1.40  | 7.8   | 10       | $f = 8$ pI:CCl$_4$ | 62    |
| $D_0$ | $\alpha$       | $\nu$ | %RMS | $M$(kDa) | System       | Refs |
|-------|----------------|-------|-------|----------|--------------|------|
| 2.18 \times 10^{-6} | 5.49 \times 10^{-3} | [1]   | 3.4   | 2.3      | $f = 2$ pB:CCl4 | 63   |
| 1.42 \times 10^{-6} | 8.78 \times 10^{-3} | 1.00  | 8.6   | 6.5      | $f = 3$ pB:CCl4 | 63   |
| 8.44 \times 10^{-7} | 0.0143         | [1]   | 7.8   | 16.0     | $f = 3$ pB:CCl4 | 63   |
| 7.07 \times 10^{-7} | 0.0163         | [1]   | 10.0  | 21.0     | $f = 3$ pB:CCl4 | 63   |
| 5.98 \times 10^{-7} | 0.0163         | [1]   | 7.5   | 26.0     | $f = 3$ pB:CCl4 | 63   |
| 2.53 \times 10^{-7} | 0.0368         | [1]   | 8.0   | 75.0     | $f = 2$ pB:CCl4 | 63   |
| 3.22 \times 10^{-7} | 0.0369         | [1]   | 15.0  | 76.0     | $f = 3$ pB:CCl4 | 63   |
| 1.53 \times 10^{-7} | 0.0421         | [1]   | 11.0  | 161.0    | $f = 3$ pB:CCl4 | 63   |
| 1.05 \times 10^{-7} | 0.052          | [1]   | 13.0  | 227.0    | $f = 3$ pB:CCl4 | 63   |
| 1.37 \times 10^{-7} | 0.0629         | [1]   | 6.8   | 281.0    | $f = 3$ pB:CCl4 | 63   |
| 1.77 \times 10^{-6} | 0.020          | 0.83  | 12.0  | 6.5      | $f = 3$ pB:CCl4 | 63   |
| 1.38 \times 10^{-6} | 0.029          | 0.79  | 9.0   | 8.7      | $f = 3$ pB:CCl4 | 63   |
| 8.81 \times 10^{-7} | 0.153          | 0.60  | 13.0  | 29.0     | $f = 3$ pB:CCl4 | 63   |
| 6.63 \times 10^{-7} | 0.273          | 0.578 | 8.5   | 87.0     | $f = 3$ pB:CCl4 | 63   |
| 2.20 \times 10^{-6} | 3.20 \times 10^{-4} | 1.58  | 22.0  | 10.0     | pS:THF       | 64   |
| 1.02 \times 10^{-6} | 1.47 \times 10^{-2} | 0.98  | 5.5   | 37.0     | pS:THF       | 64   |
| 3.97 \times 10^{-7} | 9.29 \times 10^{-2} | 0.72  | 7.4   | 179.0    | pS:THF       | 64   |
| 2.20 \times 10^{-7} | 0.299          | 0.55  | 15.0  | 498.0    | pS:THF       | 64   |
| 1.45 \times 10^{-7} | 0.452          | 0.52  | 21.0  | 1050.0   | pS:THF       | 64   |
| 9.12 \times 10^{-7} | 9.07 \times 10^{-4} | 1.42  | 15.0  | 32.0     | pS:THF       | 65   |
| 9.09 \times 10^{-7} | 3.17 \times 10^{-3} | 1.23  | 16.0  | 46.0     | pS:THF       | 65   |
| 5.16 \times 10^{-7} | 1.01 \times 10^{-2} | 1.06  | 21.0  | 105.0    | pS:THF       | 65   |
| 2.92 \times 10^{-7} | 1.39 \times 10^{-2} | 1.01  | 17.0  | 130.0    | pS:THF       | 65   |
| 2.58 \times 10^{-7} | 0.100          | 0.74  | 19.0  | 360.0    | pS:THF       | 65   |
| 7.04 \times 10^{-7} | 0.0161         | [1]   | 4.7   | 61.0     | $f = 18$ pI:CCl4 | 65   |
| 5.05 \times 10^{-7} | 0.0156         | 1.03  | 2.0   | 92.0     | $f = 18$ pI:CCl4 | 65   |
| 3.02 \times 10^{-7} | 0.035          | 0.98  | 2.7   | 193.0    | $f = 18$ pI:CCl4 | 65   |
| 3.03 \times 10^{-7} | 0.070          | 0.81  | 2.8   | 216.0    | $f = 18$ pI:CCl4 | 65   |
| 2.38 \times 10^{-7} | 0.116          | 0.74  | 2.9   | 344.0    | $f = 18$ pI:CCl4 | 65   |
| 1.95 \times 10^{-7} | 0.341          | 0.58  | 8.1   | 800.0    | $f = 18$ pI:CCl4 | 65   |
| 5.43 \times 10^{-8} | 0.76           | [0.5] | 20.0  | 6300.0   | $f = 18$ pI:CCl4 | 66   |
| 2.11 \times 10^{-7} | 0.359          | 0.58  | 8.6   | 302.0    | $f = 2$ pI:CCl4 | 66   |
| 1.11 $D_0$ | 0.167          | 0.57  | 1.5   | 70.8     | $f = 2$ pI:CCl4 | 66   |
| 0.991D_0 | 0.256          | 0.67  | 3.6   | 251.0    | $f = 2$ pI:CCl4 | 66   |
| [1.0D_0] | 0.244          | 0.68  | 6.7   | 302.0    | $f = 2$ pI:CCl4 | 66   |
| 2.07 \times 10^{-7} | 0.47           | 0.52  | 17.0  | 900.0    | pS:tol       | 67   |
### TABLE II: Concentration dependence of the probe diffusion coefficient $D_p$ for molecular weight $M$ matrix polymers.

The fits are to stretched exponentials $D_p \exp(-\alpha t^n)$ in the polymer concentration $c$. The Table gives the fitting parameters, the percent root-mean-square fractional fit error $\%\text{RMS}$, the system, and the reference. Square brackets $[\cdot \cdot \cdot]$ denote parameters that were fixed rather than floated during the non-linear least squares fits. $D_0$ is in cm$^2$/s or in units of $D_0$; concentrations are in g/L. Abbreviations include dex--dextran, $^\circ \theta = n$ for an $n$-armed star polymer, hyal--hyaluronic acid, mrm--methyl red, oFT--orthofluorotoluene, pF--pulsed field gradient nuclear magnetic resonance, QE--quasielastic light scattering spectroscopy, thiop--thiophenol, and tol--toluene. Parentheses indicate parameters from fits of limited accuracy.

| $D_p$ (10$^{-7}$) | $\alpha$ | $\nu$ | $\%$RMS | $P$ (kDa) | $M$ (kDa) | System | Refs. |
|------------------|---------|------|--------|---------|---------|--------|------|
| 3.35 (10$^{-7}$) | 0.107   | 0.70 | 8.9    | 245     | 598     | pS:pS:C| 56. |
| 2.19 (10$^{-7}$) | 0.453   | 0.50 | 25     | 598     | 1800    | pS:pS:C| 56. |
| 1.13             | 0.38    | 0.38 | 7.6    | 8000    | 101     | pS:pMMA:tol| 67. |
| 1.01             | 0.61    | 0.40 | 3.6    | 8000    | 163     | pS:pMMA:tol| 67. |
| 1.24             | 1.03    | 0.34 | 7.0    | 8000    | 268     | pS:pMMA:tol| 67. |
| 1.47             | 1.39    | 0.32 | 5.4    | 8000    | 445     | pS:pMMA:tol| 67. |
| 1.06             | 1.18    | 0.38 | 9.0    | 8000    | 697     | pS:pMMA:tol| 67. |
| 1.00             | 1.48    | 0.45 | 14     | 8000    | 1426    | pS:pMMA:tol| 67. |
| 0.91             | 1.19    | 0.34 | 14     | 2950    | 445     | pS:pMMA:tol| 67. |
| 0.603            | 0.62    | 0.46 | 18     | 150000  | 445     | pS:pMMA:tol| 67. |
| 0.973            | 0.0101  | a    | 6.7    | 73      | 19      | pEO:dex:H2O| 42. |
| 2.13             | 0.0169  | a    | 1.9    | 73      | 110     | pEO:dex:H2O| 42. |
| 1.05             | 0.0256  | [1]  | 2.8    | 73      | 510     | pEO:dex:H2O| 42. |
| 1.03             | 0.0105  | [1]  | 4.1    | 278     | 19      | pEO:dex:H2O| 42. |
| 0.976            | 0.0164  | [1]  | 4.3    | 278     | 110     | pEO:dex:H2O| 42. |
| 0.97             | 0.022   | [1]  | 4.4    | 278     | 510     | pEO:dex:H2O| 42. |
| 0.317            | 0.034   | [1]  | 1.2    | 1200    | 110     | pEO:dex:H2O| 42. |
| 1.50 (10$^{-7}$) | 0.017   | [1]  | 13     | 864     | 20.4    | dex:dex:H2O| 53. |
| 4.97 (10$^{-7}$) | 0.174   | 0.60 | 27     | 110     | 110     | pS:pVME:tol(QE)| 54. |
| 4.88 (10$^{-7}$) | 0.097   | 0.71 | 21     | 110     | 110     | pS:pVME:tol(PF)| 54. |
| 4.51 (10$^{-7}$) | 0.18    | 0.59 | 1.9    | 71      | 680     | dex:hyal:HO| 52. |
| 3.22 (10$^{-7}$) | 0.33    | 0.59 | 5.4    | 148     | 680     | dex:hyal:HO| 52. |
| 1.75 (10$^{-7}$) | 0.44    | 0.48 | 4.8    | 487     | 680     | dex:hyal:HO| 52. |
| 3.49 (10$^{-7}$) | 3.58 $10^{-2}$ | 0.87 | 3.8    | 162     | 105     | pS:pMMA:C6H6| 54. |
| (2.32 $10^{-7}$) | (0.142) | (0.55) | (10) | 410     | 105     | pS:pMMA:C6H6| 54. |
| 1.35 (10$^{-7}$) | 6.35 $10^{-2}$ | 0.76 | 5.6    | 1110    | 105     | pS:pMMA:C6H6| 54. |
| 6.70 (10$^{-8}$) | 2.89 $10^{-2}$ | 0.95 | 6.0    | 4600    | 105     | pS:pMMA:C6H6| 54. |
| 3.16 (10$^{-7}$) | 2.26 $10^{-2}$ | 0.90 | 24     | 50      | 60      | pS:pVME:ORFT| 74. |
| 2.15 (10$^{-7}$) | 5.6 $10^{-2}$ | 0.80 | 17     | 179     | 60      | pS:pVME:ORFT| 74. |
| 1.73 (10$^{-7}$) | 0.164   | 0.65 | 9.4    | 1050    | 60      | pS:pVME:ORFT| 74. |
| 6.09 (10$^{-8}$) | 2.63 $10^{-2}$ | 1.01 | 19     | 1800    | 60      | pS:pVME:ORFT| 74. |
| 1.33 (10$^{-7}$) | 6.13 $10^{-3}$ | [1] | 3.8    | 930     | 840     | pS:pMMA:tol| 74. |
| 2.99 (10$^{-7}$) | 5.05 $10^{-3}$ | [1] | 8.9    | 233     | 840     | pS:pMMA:tol| 74. |
| 2.70 (10$^{-7}$) | 2.27 $10^{-3}$ | [1] | 2.2    | 233     | 66      | pS:pMMA:tol| 74. |
| 1.31 (10$^{-5}$) | 4.4 $10^{-3}$ | [1] | 6      | 0       | a       | mrp:8:tol| 73. |
| 1.26 (10$^{-6}$) | 1.87 $10^{-3}$ | 1.28 | 5.6    | 10      | a       | pS:pS:tol| 73. |
| 7.89 (10$^{-7}$) | 1.82 $10^{-2}$ | 0.92 | 3.3    | 35      | a       | pS:pS:tol| 73. |
| 4.40 (10$^{-7}$) | 5.85 $10^{-2}$ | 0.80 | 4.0    | 100     | a       | pS:pS:tol| 73. |
| 2.30 (10$^{-7}$) | 0.27    | 0.60 | 18     | 390     | a       | pS:pS:tol| 73. |
| 1.81 (10$^{-7}$) | 0.70    | 0.48 | 25     | 900     | a       | pS:pS:tol| 73. |
| 8.54 (10$^{-8}$) | 0.64    | 0.55 | 17     | 1800    | 8400    | pS:pS:tol| 73. |
| 2.81 (10$^{-7}$) | 5.4 $10^{-2}$ | 0.82 | 12     | 179     | 50      | pS:pVME:ORFT| 73. |
| 2.94 (10$^{-7}$) | 0.26    | 0.58 | 11     | 1050    | 50      | pS:pVME:ORFT| 73. |
| 2.92 (10$^{-7}$) | 0.368   | 0.59 | 10     | 422     | 1300    | pS:pVME:ORFT| 73. |
| 2.06 (10$^{-7}$) | 0.439   | 0.60 | 9.8    | 1050    | 1300    | pS:pVME:ORFT| 73. |
| $M$ (g/mol) | Sample | $W_1$ (g) | $W_2$ (g) | $U$ (g) | $f = \text{pS:pVME:oFT}$ |
|------------|--------|-----------|-----------|--------|----------------------------|
| 0.280      | 2.97 · 10^{-7} | 0.63      | 2.6       | 379    | 1300                       |
| 0.435      | 1.84 · 10^{-7} | 0.65      | 5.0       | 1190   | 1300                       |
| 0.085      | 8.46 · 10^{-7} | 0.71      | 3.7       | 55     | 140                        |
| 0.161      | 3.01 · 10^{-7} | 0.67      | 3.6       | 467    | 140                        |
| 0.184      | 1.89 · 10^{-7} | 0.66      | 1.7       | 1110   | 140                        |
| 0.196      | 1.60 · 10^{-7} | 0.65      | 2.0       | 1690   | 140                        |
| 0.49       | 2.09 · 10^{-7} | 0.56      | 5.8       | 1690   | 1300                       |
| 0.48       | 2.52 · 10^{-7} | 0.55      | 4.7       | 1110   | 1300                       |
| 0.49       | 4.07 · 10^{-7} | 0.48      | 4.9       | 467    | 1300                       |
| 0.35       | (4.1) · 10^{-7} | (0.34)    | 3.9       | 55     | 1300                       |
| 0.18       | 1.44 · 10^{-7} | 0.63      | 4.7       | 1190   | 140                        |
| 0.25       | 1.58 · 10^{-7} | 0.65      | 14.6      | 1190   | 630                        |
| 0.612      | 2.18 · 10^{-7} | 0.55      | 18       | 1190   | 1300                       |
| 0.162      | 2.69 · 10^{-7} | 0.63      | 3.7       | 379    | 140                        |
| 0.167      | 2.57 · 10^{-7} | 0.68      | 7.5       | 379    | 630                        |
| 0.288      | 2.87 · 10^{-7} | 0.61      | 12       | 379    | 1300                       |
| 0.061      | 5.27 · 10^{-7} | 0.74      | 5.9       | 50     | 110                        |
| 0.054      | 3.65 · 10^{-7} | 0.80      | 1.6       | 100    | 110                        |
| 0.145      | 2.48 · 10^{-7} | 0.66      | 3.6       | 420    | 110                        |
| 0.116      | 1.49 · 10^{-7} | 0.71      | 4.6       | 900    | 110                        |
| 0.32       | 1.56 · 10^{-7} | 0.77      | 0.9       | 342    | 43.9                       |
| 0.54       | 1.43 · 10^{-7} | 0.54      | 1.3       | 342    | 186                        |
| 0.79       | 1.18 · 10^{-7} | 0.68      | 6.1       | 342    | 775                        |
| 1.03       | 1.23 · 10^{-7} | 0.67      | 8.4       | 342    | 8420                       |
| 3.21 · 10^{-4} | 1.51 · 10^{-5} | (1)      | 5.5      | M      | a | styrene:pS:CCl$_4$ |
| 3.34 · 10^{-4} | 2.53 · 10^{-5} | (1)      | 3.0      | M      | a | styrene:pS:C$_6$H$_{12}$ |
| 3.15 · 10^{-3} | 1.51 · 10^{-5} | (1)      | 5.8      | M      | a | styrene:pS:CCl$_4$ |
| 5.36 · 10^{-3} | 4.74 · 10^{-6} | (1)      | 5.0      | 0.58   | a | pS:pS:CCl$_4$ |
| 6.53 · 10^{-3} | 3.31 · 10^{-6} | (1)      | 4.8      | 1.2    | a | pS:pS:CCl$_4$ |
| 8.08 · 10^{-4} | 2.16 · 10^{-6} | (1)      | 4.1      | 2.47   | a | pS:pS:CCl$_4$ |
| 0.033      | 5.13 · 10^{-9} | 0.69      | 4.3      | 33.6   | 32 | PPO:PP0:PP0 |
| 5.23       | 7.86 · 10^{-11}| 0.79      | 10.8     | 255    | 93  | pS:pS:pS |
| 0.29       | 9.17 · 10^{-11}| 0.51      | 23       | 255    | 250 | pS:pS:pS |
| 4.81       | 1.24 · 10^{-11}| 0.47      | 8.7      | 255    | 20000 | pS:pS:pS |
| 7.3 · 10^{-2} | 1.93 · 10^{-7} | 0.84     | 10.3     | 433    | 310 | dexpvP:water |
| 0.20       | 7.1 · 10^{-7} | 0.57      | 3.1      | 65     | 1300 | pS:VME:OF |
| 0.21       | 5.42 · 10^{-7} | 0.66      | 5.4      | 179    | 1300 | pS:VME:OF |
| 0.35       | 2.87 · 10^{-7} | 0.60      | 10.2     | 422    | 1300 | pS:VME:OF |
| 0.47       | 2.19 · 10^{-7} | 0.59      | 10.1     | 1050   | 1300 | pS:VME:OF |
| 0.101      | 6.30 · 10^{-7} | 0.62      | 3.7      | 65     | 140  | pS:VME:OF |
| 0.150      | 3.72 · 10^{-7} | 0.62      | 2.3      | 179    | 140  | pS:VME:OF |
| 0.165      | 2.34 · 10^{-7} | 0.63      | 4.6      | 422    | 140  | pS:VME:OF |
| 0.194      | 1.64 · 10^{-7} | 0.62      | 5.5      | 1050   | 140  | pS:VME:OF |
| 0.135      | 6.32 · 10^{-7} | 0.58      | 4.5      | 65     | 630  | pS:VME:OF |
| 0.227      | 3.99 · 10^{-7} | 0.57      | 4.1      | 179    | 630  | pS:VME:OF |
| 0.29       | 2.60 · 10^{-7} | 0.58      | 8.2      | 422    | 630  | pS:VME:OF |
| 0.34       | 1.80 · 10^{-7} | 0.58      | 12.0     | 1050   | 630  | pS:VME:OF |

a Various matrix molecular weights with $M \gg p$. 
TABLE III: Concentration and molecular weight dependences of $D_s$ and $D_p$ for molecular weight $P$ probes in solutions of molecular weight $M$ matrix polymers (for $D_s$, one has $P = M$) at concentration $c$. The fits are to stretched exponentials $D_s \cdot P^{-\alpha} \exp(-\alpha c^{P} P^{\beta} M^{\gamma})$, with the percent root-mean-square fractional fit error $\%R$, the material, and the reference. Molecular weights are in kDa; concentrations except as noted are in g/L. Square brackets "[⋯]" denote parameters that were fixed rather than floated. Abbreviations as in other tables.

| $D_s$ | $a$ | $\alpha$ | $\nu$ | $\gamma$ | $\delta$ | $\%R$ | $P$ | $M$ | $c$ | System | Refs. |
|-------|-----|----------|-------|----------|----------|-------|-----|-----|-----|--------|-------|
| $6.27 \times 10^{-5}$ | [0.5] | $5.96 \cdot 10^{-5}$ | 0.93 | 0.61 | [0] | 9 | $b$ | $f$ | (2) | PEO:2H2O | 42 |
| $7.99 \times 10^{-5}$ | 0.50 | $7.38 \cdot 10^{-4}$ | 0.58 | 0.46 | [0] | 17 | $b$ | $f$ | (2) | pS:CCl4 | 44 |
| $3.41 \times 10^{-5}$ | 0.55 | $2.10 \cdot 10^{-3}$ | 0.64 | 0.36 | [0] | 7 | $b$ | $f$ | (2) | pS:CD6 | 44 |
| $2.94 \times 10^{-4}$ | 0.50 | $1.53 \cdot 10^{-2}$ | 0.52 | 0.25 | [0] | 24 | $b$ | $f$ | (2) | pS:cp | 44 |
| $7.46 \times 10^{-4}$ | 0.501 | $6.17 \cdot 10^{-3}$ | 0.48 | 0.33 | [0] | 4 | $b$ | $f$ | (2) | pDMS:tol | 52 |
| $1.16 \times 10^{-4}$ | [0.5] | $7.55 \cdot 10^{-7}$ | 0.95 | 0.86 | [0] | 37 | $b$ | $f$ | (2) | pS:CH4 | 52 |
| $1.67 \times 10^{-4}$ | [0.5] | $5.08 \cdot 10^{-4}$ | 0.75 | 0.48 | [0] | 17 | $b$ | $f$ | (2) | pB:CCl4 | 54 |
| $1.87 \times 10^{-4}$ | [0.5] | $1.93 \cdot 10^{-3}$ | 0.91 | 0.24 | [0] | 51 | $b$ | $f$ | (2) | pS:THF | 54 |
| $9.68 \times 10^{-4}$ | 0.65 | $7.37 \cdot 10^{-4}$ | 0.68 | 0.42 | [0] | 5.7 | 193-800 | $b$ | (2) | pT:CCl4 | 55 |
| 0.891 | [0] | $5.86 \cdot 10^{-4}$ | 0.43 | 0.43 | [0] | 24 | $b$ | $f$ | (2) | pS:pMMA:tol | 57 |
| 1.025 | [0] | $1.53 \cdot 10^{-3}$ | 0.95 | -0.01 | 0.236 | 4.8 | $b$ | $f$ | (2) | PEO:deoxH2O | 57 |
| $7.46 \cdot 10^{-6}$ | 0.36 | $2.84 \cdot 10^{-4}$ | 1.15 | 0.26 | [0] | 34 | $b$ | $f$ | (2) | pS:pVME:oFT | 73 |
| $8.91 \cdot 10^{-6}$ | 0.34 | $6.80 \cdot 10^{-4}$ | 1.03 | 0.25 | [0] | 24 | $b$ | $f$ | (2) | pS:pVME:oFT | 73 |
| $3.34 \cdot 10^{-4}$ | 0.57 | $1.82 \cdot 10^{-5}$ | 0.99 | 0.139 | 0.287 | 6.4 | $b$ | $f$ | (2) | pSp:MA:tol | 73 |
| $1.86 \cdot 10^{-4}$ | 0.52 | $4.45 \cdot 10^{-5}$ | 0.69 | 0.30 | 0.33 | 17 | $b$ | $f$ | (2) | pSp:tol | 73 |
| $8.5 \cdot 10^{-4}$ | 0.61 | $2.99 \cdot 10^{-2}$ | 0.61 | 0.19 | [0] | 12 | $b$ | $f$ | (2) | f = 2 pSp:pVME:oFT | 73 |
| $3.0 \cdot 10^{-4}$ | 0.54 | $1.07 \cdot 10^{-3}$ | 0.66 | 0.42 | [0] | 6.6 | $b$ | $f$ | (2) | f = 3 pSp:pVME:oFT | 73 |
| $1.84 \cdot 10^{-4}$ | 0.50 | $1.77 \cdot 10^{-2}$ | 0.68 | 0.16 | [0] | 12 | $b$ | $f$ | (2) | f = 12 pSp:pVME:oFT | 78 |
| $2.20 \cdot 10^{-4}$ | 0.52 | $5.09 \cdot 10^{-4}$ | 0.61 | 0.16 | 0.32 | 32 | $b$ | $f$ | (2) | f = 3 pSp:pVME:oFT | 78 |
| $2.93 \cdot 10^{-4}$ | 0.53 | $2.22 \cdot 10^{-3}$ | 0.66 | 0.15 | 0.19 | 15 | $b$ | $f$ | (2) | f = 12 pSp:pVME:oFT | 78 |
| $6.51 \cdot 10^{-4}$ | 0.67 | $9.49 \cdot 10^{-3}$ | 0.86 | 0.115 | [0] | 6 | $b$ | $f$ | (2) | pSp:pVME:oFT | 78 |
| $6.67 \cdot 10^{-5}$ | [0.5] | $4.5 \cdot 10^{-3}$ | 0.68 | [0.3] | 0.024 | 20 | $b$ | $f$ | (2) | pMMA:pSthiop | 82 |
| $2.79 \cdot 10^{-5}$ | [0.52] | 0.95 | [0] | [0.16] | 0.028 | 20 | $b$ | $f$ | (2) | pSp:DBP | 83 |
| $7.64 \cdot 10^{-6}$ | 0.56 | $7.61 \cdot 10^{-4}$ | $a$ | 0.30 | 0.15 | 13 | $b$ | $f$ | (2) | pSp:DBP | 83 |
| $7.89 \cdot 10^{-6}$ | 0.63 | $1.35 \cdot 10^{-3}$ | $a$ | 0.28 | 0.11 | 12 | $b$ | $f$ | (2) | pSp:DBP | 84 |
| $8.41 \cdot 10^{-6}$ | 0.64 | $2.31 \cdot 10^{-4}$ | 0.74 | 0.26 | 0.29 | 31 | $b$ | $f$ | (2) | pSp:DBP | 84 |
| $1.65 \cdot 10^{-4}$ | 0.52 | $1.56 \cdot 10^{-6}$ | 0.79 | 0.285 | 0.501 | 23 | $b$ | $f$ | (2) | pSp:MA:tol | 85 |
| $3.98 \cdot 10^{-4}$ | 0.68 | $1.00 \cdot 10^{-3}$ | [1.0] | 0.26 | [0] | 7.9 | $b$ | $f$ | (2) | pSp:CCl4 | 87 |
| $1.31 \cdot 10^{-4}$ | 0.52 | $7.56 \cdot 10^{-4}$ | [1.0] | 0.30 | [0] | 5.1 | $b$ | $f$ | (2) | pSp:CCl4 | 87 |
| $2.01 \cdot 10^{-4}$ | 0.51 | $1.03 \cdot 10^{-2}$ | 0.64 | 0.26 | [0] | 14 | $b$ | $f$ | (2) | pSp:pVME:oFT | 93 |
| $4.99 \cdot 10^{-3}$ | 0.39 | $8.33 \cdot 10^{-4}$ | 0.54 | 0.25 | 0.22 | 25 | $b$ | $f$ | (2) | pSp:pVME:oFT | 93 |

$a$ Various, with $M/P \geq 10$

$b$ Various, see text.

$^c$ All four probes, see text.

$^d$ Excluding styrene monomer, see text.

$^e$ Not all data points, see text.

$^f$ $P = M$, self-diffusion.
TABLE IV: Molecular weight dependence of the self and probe diffusion coefficients $D_s$ and $D_p$ for molecular weight $P$ probes in solutions of matrix polymers at a fixed concentration $c$. The fits are to stretched exponentials $D_0 \exp(-\alpha M^\gamma)$ in matrix molecular weight $M$. The Table gives the best-fit parameters, the percent root-mean-square fractional fit error %RMS, the system, and the reference. Square brackets “[· · ·]” denote parameters that were fixed rather than floated. Abbreviations as per previous Tables, and DBP–dibutylphthalate.

| $D_0$ (kDa) | $\alpha$ | $\gamma$ | %RMS | $P$ (kDa) | $c$ (g/L) | System | Refs. |
|-------------|----------|----------|-------|-----------|-----------|--------|-------|
| $8.54 \cdot 10^{-8}$ | 0.64 | 0.55 | 17 | 1800 | $a$ | pS:pS:tol | 75 |
| $2492 P^{-0.5}$ | 1.37 $\cdot 10^{-2}$ | 0.47 | 4.0 | $P$ | 130 | pS:pS:dbp | 57 |
| $1771 P^{-0.5}$ | 1.07 $\cdot 10^{-2}$ | [0.5] | 6.9 | $P$ | 180 | pS:pS:dbp | 57 |

*aVarious, see text.

TABLE V: Concentration and molecular weight dependences of the probe radius of gyration $R_g$ for molecular weight $P$ probes in solutions of molecular weight $M$ matrix polymers as functions of matrix concentration $c$. The fits are to stretched exponentials $R_{g0} \exp(-\alpha c^\nu)$, with the percent root-mean-square fractional fit error %RMS, the materials, and the reference. Materials include EB–ethyl benzoate, PMMA–polymethylmethacrylate, pS–polystyrene.

| $R_{g0}$ (Å) | $\alpha$ | $\nu$ | %RMS | $P$ (kDa) | $M$ (kDa) | System | Refs. |
|--------------|----------|-------|-------|-----------|-----------|--------|-------|
| 390 | $4.11 \cdot 10^{-3}$ | 0.99 | 2.2 | 930 | 1300 | pS: pMMA: EB | 75 |
| 397 | $8.06 \cdot 10^{-3}$ | 0.77 | 1.1 | 930 | 70 | pS: pMMA: EB | 75 |
| [395] | $4.49 \cdot 10^{-3}$ | 0.64 | 1.9 | 930 | 7 | pS: pMMA: EB | 75 |