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Molecular dynamics study of the swelling and osmotic properties of compact nanogel particles†

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Owing to their great importance in materials science and other fields, we investigate the solution and osmotic properties of uncharged compact nanogel particles over a wide range of solvent quality and particle concentration by molecular dynamics (MD) simulations. We characterize the osmotic pressure by estimating the second and third virial coefficients, and by extension, we identify the θ -point where the second virial coefficient vanishes. Calculations of the structure factor indicate that these particles are similar to macrogels in that the particle-like scattering profile disappears at moderate concentrations. We also find that improving the solvent quality enhances the spatial segmental uniformity, while significant heterogeneous structure arises near the θ -point. Well below the θ -point where the second osmotic virial coefficient vanishes, these heterogeneous structures become less prevalent as the particles tend to collapse. We also investigate the degree of swelling and structure of compact nanogel particles with a variable excluded volume interaction and gel particle concentration. The osmotic modulus and the scaling exponents in good and θ -point conditions of these gels are characteristic of interacting randomly branched polymers, i.e., "lattice animals".

I. Introduction

Nanogel particles are typically, but not always, hydrogel materials with a high capacity for swelling in water, depending on the extent of cross-linking and polymer excluded interactions within and between them. 1-3 They are increasingly becoming essential building blocks of modern (bio)-manufacturing, 4-6 because they are often highly bio-compatible, due to their hydrophilic nature. The high loading capacity for guest molecules and their unique physical properties offer distinct advantages over other types of nanomaterials for biomedical applications. Moreover, they are highly customizable since they can be composed of a variety of naturally occurring or synthetic polymers, and their size, softness, and degradability can be fine-tuned by varying the chemical composition. For example, nanogel particles can be designed to exhibit multi-stimuli responsiveness in targeted therapy of cancer⁷ or to provide targeted delivery of short interfering RNA (siRNAs), a gene-regulating tool to increase the efficacy of chemotherapy drugs.8 Owing to the growing number of applications of nanogel particles, there is a need to

understand these particles better to enable optimization of

targeted drug-delivery vehicles, nanogel particles have been also used in bacterial bio-production, 9 as self-healing materials, 10 in enhanced oil recovery,11 coatings12,13 and sensors.14 Another emerging role of nanogel particles is by embedding them within a polymer network to improve performance for the desired application. Specifically, the articular cartilage of the knee and hip exhibits compromised load-bearing capacity when its compressive resistance is reduced. Moreover, articular cartilage experiences large-amplitude repetitive loading during walking or jumping, so a rapid recovery of the bulk tissue is crucial to normal function. Deviation from the physiological material performance in articular cartilage is associated with degenerative conditions such as osteoarthritis. 15-18 Recently, Horkay and Basser¹⁹ have utilized nanogel particles to develop a synthetic composite hydrogel that mimics functional properties of articular cartilage. 19 As a prerequisite to studying composite gels, it is prudent to develop descriptions of the two primary components, the network and the nanogel particles. In this paper, we study the swelling and osmotic properties of compact nanogel particles in dilute and concentrated solutions in the absence of a polymer matrix. The inclusion of the polymer matrix is integral for creating the composite gel model, but it also increases the complexity of the model. We plan for this to be the subject of a subsequent paper. It is then essential to understand the behavior

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their properties for specific applications. While nanogel particles are predominately being used as

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of the nanogel particles in the absence of a polymer matrix as the natural step toward our goal.

Our previous study of polymers having a network topology indicated that the apparent exponents describing the mass scaling of these polymers under good solvent conditions generally depend on the density of branch points in the network and the fraction of bonds cut in the network. 20,21 This led us to define two distinct general classes of networks. First, the 'ideal' or 'compact nanogel particles' have a network topology defined by a lattice-like structure, resulting in a more symmetric particle-like geometry. Second, the 'open nanogel particles' have a polymer network that is highly defective so that the polymer network forms an anisotropic fractal-like structure. The mass scaling of the 'open gels' was found to conform rather well with randomly branched polymers ('lattice animals'), but the situation was found to be more complicated for the perfect nanogel particles whose size did not seem to conform with the expectations of randomly branched polymers.²¹ In particular, the apparent mass scaling exponents describing the average size of these nanogel particles depended on the mesh spacing in the network. The radius of gyration mass scaling exponent when the mesh size was small was found to be consistent with that for the swollen linear chains composing the network in isolation ($\nu \approx 0.588$) with increasing M while the apparent exponent was found to be near ($\nu \approx 1/3$) with increasing branched points in each direction. An examination of the structure factor of these closed gel nanoparticles exhibits a mass scaling similar to self-avoiding linear polymers at high scattering wavevector. However, a cross-over to a wavevector scaling is consistent with a higher effective fractal dimension at larger length scales. 20,21 It is apparent from these observations that the structures simulated cannot be described as being 'fractal' objects, but they are rather hybrid structures having properties in common to fractals, epitomized by linear polymer chains, and symmetric particles having uniform density. It is difficult from the limited size simulations to deduce the asymptotic mass scaling in the limit of large polymer mass from the study of individual network polymer chains, but additional insights into these mass scaling characteristics can be obtained from the study of the swelling of these structures as solvent quality is varied and from the concentration dependence of the osmotic pressure under variable solvent conditions. We will show that there are slow cross-over effects on the ν exponent, as found even in linear polymer chains when the polymers are semi-flexible, suggesting there is an appreciable rigidification of polymer network when the mesh spacing is small.

This paper studies the swelling and osmotic properties of compact nanogel particles in dilute and concentrated solutions. We have already developed methods for generating nanogel particles, and we have studied the structure and conformational properties in athermal dilute solutions. 20,21 We use these methods to construct gels composed of nanogel particles and characterize their structure, osmotic pressure, and osmotic modulus. In addition, we estimate the virial coefficients of compact nanogel particles and compare them with star polymer solutions. We also investigate the influence

of solvent quality on the degree of swelling in both dilute and concentrated nanogel particle concentrations.

The paper is organized as follows. Section II contains details of the simulation model and methods. The results are presented in Section III, where we investigate the influence of solvent quality in the swelling of nanogel particles in dilute and concentrated solutions. We also investigate the scattering profiles of these gels and the calculate the osmotic modulus. Finally, we draw our conclusions in Section IV.

II. Methods and models

We employ a bead-spring model suspended in an implicit solvent, in which we have adapted a model developed previously for studying the swelling behavior of linear polymer chains in solution. All the nanogel particles are assigned the same mass m, size σ , and strength of interaction ε ; we set ε and σ as the units of energy and length.

The expression describing the non-bonded interactions operating between all pairs of beads contains three terms. First we write the purely repulsive Weeks-Chandler-Andersen (WCA) potential,22 which is a Lennard-Jones potential cut and shifted at the position of the minimum, $r_{\min} = 2^{1/6}\sigma$:

$$V_{\text{WCA}}(r) = \begin{cases} 4\varepsilon \left[\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^{6} \right] + \varepsilon & r \le r_{\text{min}} \\ 0 & r > r_{\text{min}} \end{cases}$$
 (1)

To represent the attractive interactions, the WCA potential is shifted in the range $0 \le r \le r_{\min}$ by a square-well (SW) potential,

$$V_{\rm SW}(r) = \begin{cases} -\lambda \varepsilon & 0 < r \le r_{\rm min} \\ 0 & r > r_{\rm min} \end{cases}$$
 (2)

where the well-depth parameter λ allows for a tuning of the effective monomer-monomer attractive interaction strength or the "solvent quality". We again note that we follow previous work²³ in the use of this model coarse-grained interaction potential. To interpolate the potential smoothly between $-\lambda \varepsilon$ at $r = r_{\min}$ and 0 at a cut-off distance $r_{\text{cut}} > r_{\min}$, we add the term

$$V_{\cos}(r) = \begin{cases} \frac{1}{2} \lambda \varepsilon \left[\cos(\alpha r^2 + \beta) - 1 \right] & r_{\min} < r \le r_{\text{cut}} \\ 0 & r > r_{\text{cut}} \end{cases}$$
 (3)

 α and β satisfy the conditions $\alpha r_{\min}^2 + \beta = \pi$ and $\alpha r_{\text{cut}}^2 + \beta = 2\pi$. The cosine form of the potential also means that $dV_{\cos}/dr = 0$ at

 $r = r_{\rm cut}$. We choose $r_{\rm cut} = 3\sigma/2$, for which α and β become, $\alpha =$

$$\frac{4\pi}{9-4\sqrt[3]{2}}$$
 and $\beta=2\pi-\frac{9}{4}\alpha$. The final non-bonded potential is $V(r)=V_{\rm WCA}+V_{\rm SW}(r)+V_{\rm cos}(r)$, see Fig. 1. The parameter λ controls the depth of the potential well at $r=r_{\rm min}$, and provides a convenient measure of the solvent quality. In a good solvent, the effective bead–bead interactions are purely repulsive; this

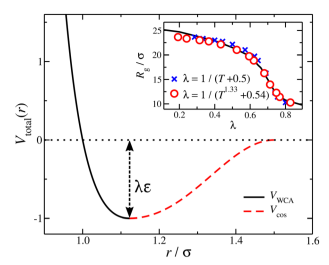


Fig. 1 The non-bonded, bead-bead interaction potential V(r) with $\lambda = 1$, showing the contributions from $V_{WCA}(r) + V_{SW}(r)$ (black) and $V_{cos}(r)$ (red). Inset: Average radius of gyration of a nanogel particle $via \lambda$ variation (black line) and temperature T variation (symbols). Two relations between λ and Tare presented.

corresponds to $\lambda = 0$. In a poor solvent, the bead-bead interactions are attractive, and this behavior can be modeled with $\lambda = 1$; the parameter λ also corresponds roughly to the Lennard-Jones potential well, which has been used frequently in the past in coarse-grained modeling of neutral polymers both in solution and in the melt.24,25

In simple models of uncharged polymer solutions, a variable excluded volume interaction strength is normally described theoretically by the ratio of the strength of the nearest-neighbor interaction (e.g., the depth of a square potential) divided by temperature as in the classical Flory-Huggins theory of polymer solutions and polymers in the melt state where the the dimensionless energetic well-depth parameter (normally denoted as χ) is estimated from fitting the model calculations based on such idealized intermolecular potentials to experiment.26 It is wellknown that this approach must often be modified by adding a constant "entropic" term to the γ interaction parameter so that the T dependence is more complicated than indicated by the original Flory-Huggins theory neglecting monomer and solvent structure, as well as the topology and bending rigidity of the molecules. A general, albeit complex extension of the Flory-Huggins model has been developed which addresses these specific monomer effects^{27,28} which explains this more complex temperature phenomenology of polymer-polymer interactions under high polymer polymer concentrations where mean field theory reasonably applies. Correspondingly, we may expect some deviation from the simple dependence of solvent quality on the ratio $\lambda \varepsilon/T$ that one may infer from the Lennard-Jones or the square-well like potential alone. We address this complication in describing the reduced interaction strength so that obtain a unified description of the variation of solvent quality by varying either λ or T.

The benefit of using the depth of the potential interaction instead of *T* for the control of the solvent quality is that a wider range of "solvent quality" can be explored by molecular dynamics simulation. A "map" between the λ and T can be constructed by matching an appropriate quantity, here we used the radius of gyration for a nanogel particle, see inset of Fig. 1. A first order approximation results in $\lambda = (T + 0.5)^{-1}$, which evidently athermal solvents having $\lambda \to 0$ require $T \to \infty$; an improved description is found with $\lambda = (T^{1.33} + 0.54)^{-1}$. In other words, athermal or very "good" solvents require high temperature values that are prohibited in a molecular dynamics simulation. Moreover, high temperature simulations of some complexity models, such as polymers, have the potential of not sampling appropriately the configurational space due to deviation between the configurational temperature and thermodynamic temperature.²⁹ The segments along a chain are connected with their neighbors via a stiff harmonic spring, $V_H(r) = k(r - l_0)^2$, where $l_0 = 0.99\sigma$ is the equilibrium length of the spring, and $k = 2000\varepsilon/\sigma^2$ is the spring constant.

A perfect compact gel that is composed of star polymers placed in a square or in a cubic lattice and with two or more of their free ends bonded with the free ends of the neighboring stars, the number of branched points (or star polymers) in each direction is labeled as N_x , N_y , and N_z , see Fig. 2.²⁰ The repeating structural unit of the polymer network studied here is a branched structure that is identical to a regular star polymer. Other polymeric structures and/or other lattices could be utilized but these are out of the scope of the current study. A regular star polymer has a core particle, which is connected with the free end of f chains (or arms) composed of Msegments. Thus, the total number of interaction centers per star polymer is $M_{w,star} = fM + 1$. The molecular mass of a nanogel particle is $M_{\rm w} = (N_x N_y N_z) M_{\rm w,star}$. We use the quantity $N_{\rm b}$

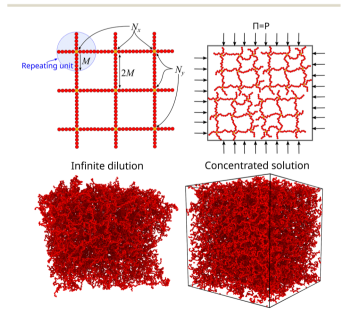


Fig. 2 Schematic of the molecular architecture of the nanogel particle. Typical screenshots of a equilibrated molecular configuration of compact nanogel particle at infinite dilution (the simulation box is so large that is not visible) and fixed osmotic pressure conditions, where the nanogel particles fill the simulation box and interacting through periodic boundaries.

Paper

to characterize the number of branched points in each direction. We focus on nanogel particles having $N_b = N_x = N_y = N_z$. Every star polymer unit at the interior of the nanogel is fully bonded with its neighbors and thus the only dangling polymer chains are located at the exterior of the nanogel structure. Nanogel particles are randomly placed in a simulation box without overlaps. For highly dilute conditions, we consider a single nanogel particle in a simulation box several times larger than the size of the nanogel particle. For concentrated solutions, we considered a small number of nanogel particles ranging from one to eight. Unless stated otherwise, we mainly focus on compact nanogel particles having $N_b = 10$, f = 4, and M = 25.

The systems in dilute concentrations were equilibrated at constant temperature $k_{\rm B}T/\varepsilon$ = 1.0, maintained by a Nosé-Hoover thermostat. In concentrated solutions, the systems were equilibrated at constant temperature $k_{\rm B}T/\varepsilon = 1.0$ and constant pressure maintained by a Nosé-Hoover thermostat and barostat. Typical simulations equilibrate for 5000τ and data is accumulated over a 150 000 τ interval, where $\tau = \sigma(m/\varepsilon)^{1/2}$ is the MD time unit; the time step used was $\Delta t/\tau = 0.005$.

The osmotic pressure of the system is calculated from the virial equation $\Pi = \rho kT + W/V$, where the internal virial W is calculated from the sum of a pair virial function $w(r_{ii}) =$ $r_{ii}dU(r_{ii})/dr_{ii}$. Due to the presence of non-vanishing intramolecular contributions, a small constant is added in Π to ensure that the correct limit is reached at vanishing small segmental densities.

III. Results and discussion

We first discuss the two main approaches to identifying the θ -point in compact nanogel particles MD simulations where the second virial coefficient vanishes. The first approach involves taking advantage of the invariance of the mass scaling of the size of the polymer, and the second approach uses the virial equation to estimate the second virial. Once the θ -point is identified, then we develop a universal description for the size of the compact nanogel particles. Following this analysis, we calculate the form factor and structure factor for these structures and characterize the structure of these gels as the θ -point is approached. Finally, we utilize our model of compact nanogel particles to estimate the osmotic modulus over a wide range of molecular characteristics and solvent conditions.

Compact nanogel particles in super-dilute concentrations

We initiate our discussion on the swelling of compact nanogel particles in super-dilute concentrations. We define the segmental density $\rho = N_{gel}M_w/V$, where N_{gel} is the number of nanogel particles and V is the volume of the simulation box. We note the local segmental density inside the nanogel particle $\rho_{gel} = M_w/V_{nanogel}$, where V_{nanogel} is the volume occupied by nanogel particle, does not decrease towards vanishing small values as $\rho \to 0$ due to the polymer crosslinking. For the purposes of the current study, we assume that the volume of the nanogel particle is approximately,

$$V_{\rm nanogel} \approx \frac{4\pi}{3} R_{\rm g}^3$$
.

At high temperatures, polymer chains in solution³⁰⁻³² and, by extension, the gels,³³ swell due to the dominance of repulsive interactions, while at low temperatures, the polymer chains collapse because the attractive interactions dominate over the repulsive interactions resulting in chain collapse conformations. At intermediate temperatures, near the θ -temperature where attractive interactions compensate the repulsive inter-polymer interactions, the chains adopt nearly 'ideal' conformations because the attractive and repulsive interaction contributions cancel each other, but the scaling of the polymer size under this ideal condition depends on polymer topology, in general. Linear chain melts and concentrated solutions have practically ideal chain conformations between monomers because the surrounding chains almost completely screen the interactions between the monomers.

There are two main approaches to identifying the θ -temperature for a polymeric system. The first one takes advantage of the size mass invariance that polymers exhibit at the θ -temperature, provided one knows the Flory exponent, ν_{θ} appropriate to the kind of polymer involved. The approach is relatively straightforward, requiring the estimation of the size of the polymer at different solvent quality solvents and molecular masses. The radius of gyration R_{g} is typically used as a measure of polymer size for this purpose. As the temperature of the system is increased, the polymer (assuming that it is neutral) will swell. This type of calculation is repeated for polymers of different molecular masses. The θ -temperature can be identified by the location where $R_{\rm g}/M_{\rm w}^{\nu_{ heta}}$ remains fixed for all molecular masses. The advantage of this approach is that it is relatively straightforward to implement computationally. However, its disadvantage is that it requires a priori knowledge of ν_{θ} , which cannot be assumed to be the same as polymers having a linear topology and this definition does not generally coincide with the condition at which the second virial coefficient vanishes for finite-sized polymers. The second approach is the estimation of θ -temperature by the calculation of the second virial, and we will discuss this approach in Subsection IIIC.

The scaling regions are clearly distinguishable in Fig. 3. In the first region $R_{\rm g}/M_{\rm w}^{\nu_{\theta}}$ increases as $M_{\rm w}$ increases corresponding to a good solvent regime, and in the second region $R_{\rm g}/M_{\rm w}^{\nu_{\theta}}$ decreases as Mw increases corresponding to a poor solvent regime. In between these two regimes all $R_{
m g}/M_{
m w}^{
u_{
m g}}$ curves for different values of $M_{\rm w}$ cross, suggesting that $R_{\rm g}/M_{\rm w}^{\nu_{\theta}}$ becomes independent of $M_{\rm w}$ variation. This crossover point defines the θ -point. Additionally, our high fidelity results provide a clear identification of a single point where $R_{
m g}/M_{
m w}^{
u_{ heta}}$ remains invariant of $M_{\rm w}$ corresponding to $\lambda_{\theta} \approx 0.7$, see Fig. 3. While the same mass scaling exponents for each solvent quality condition are found in both linear chains and our compact nanogel particles, the location of the θ -point is significantly different; for linear chains, $\lambda_{\theta, \text{chains}} = 0.646$. The branched nature of the polymer network evidently also influences the θ -point.

Knowledge of the conditions at which the θ -conditions emerge provide an important reference point to describe the changes in the size of a polymer over the variation of the solvent

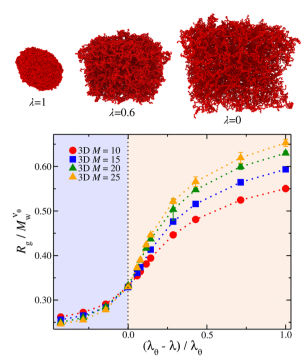


Fig. 3 Average radius of gyration, $R_{\rm g}$, of isolated nanogel particles normalized by the scaled molecular mass, $M_{\rm w}^{\nu_{\theta}}$, where the ν_{θ} = 2/5 is the Flory exponent at the θ -point, as a function of the reduced solvent quality parameter, $(\lambda_{\theta}-\lambda)/\lambda_{\theta}$, where λ_{θ} = 0.7. The highlighted regions outline the good solvent conditions for $(\lambda_{\theta} - \lambda)/\lambda_{\theta} > 0$ and poor solvent conditions for $(\lambda_{\theta} - \lambda)/\lambda_{\theta} < 0$. The uncertainty estimates correspond to two standard deviations. Typical equilibrated molecular configurations of compact nanogel particle at infinite dilution at different solvent conditions are also presented.

quality. There is no rigorous theory for the swelling of these networks, and we first offer a highly simplified mean field model of the swelling of the polymer networks. Inspired by previous modeling of the swelling of polymer brushes and gels;³⁴ The basic idea is that the extent of polymer swelling ultimately saturates, and we model the swelling by a tanh function which varies sigmoidally and inherently exhibits this saturation tendency. Indeed, we find $R_g/R_{g,q}$ for that compact nanogel particle having M and variable N_b follows this empirical functional form rather well. In particular, $R_g/R_{g,\theta}$ for compact nanogel particles can be described by the empirical functional form,

$$\frac{R_{\rm g}}{R_{\rm g,\theta}} = 1 + c_0 \tanh\left[\frac{\tau M^{\varphi}}{c_1}\right] \tag{4}$$

where the c_0 , c_1 , and φ are fitting parameters, and τ is the reduced temperature $\tau = (T - T_{\theta})/T$. The results of this approximation, which a direct extension of the scaling relation for the dimensions of macroscopic gels and grafted polymer brushes,³⁴ are shown in Fig. 4. We also used the following relation, $[(\lambda_{\theta} - \lambda)/$ $\lambda_{\theta}^{\gamma} \sim (T - T_{\theta})/T$, where γ is a fitting parameter. This simple approximant clearly captures the overall trend in the nanogel particle swelling rather well. We note that for the good and the poor solvent regimes, a different set of values for these

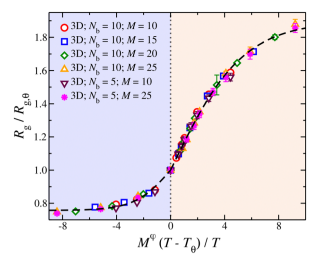


Fig. 4 Average radius of gyration, $R_{\rm g}$, of isolated compact nanogel particles normalized by R_q at θ -point, as a function of the reduced temperature, $\tau = (T - T_{\theta})/T$ and chain length, M, scaled with an exponent φ . The highlighted regions outline the good solvent conditions for $\tau > 0$ and poor solvent conditions for τ < 0. The uncertainty estimates correspond to two standard deviations. The dashed line is a fit of eqn (4) and the values of the parameters are presented in Table 1.

parameters were found, and their values are presented in Table 1. This simple approximation cannot apply to the case of highly variable M since the different mass scaling between the θ -solvent, and good solvent limits will make the gap between the theta and good solvent limits progressively larger with increasing M. This same situation arises in linear polymer chains, but where the good solvent and θ -solvent scaling exponents are different from branched polymers. Future work will describe the size of compact²⁰ and open²¹ nanogel particles that are more suitable when considering an extensive range of polymer molecular mass. We next briefly show that a simple extension of the renormalization (RG) group theory to describe the swelling of our nanogel particles appears to be promising for describing this type of crossover above the θ -point.

In Fig. 5, we show a comparison to the first order renormalization group expression for $R_{\rm g}^{\ \ 2}/R_{{\rm g},\theta}^{\ \ 2}$, 35,36

$$R_{\rm g}^2/R_{\rm g,q}^2 = (1 + z/u^*)^{(2\nu - 2\nu\theta)/\phi} [1 + b(z/u^*)/(1 + z/u^*)]$$
 (5)

for linear polymers with the swollen and θ -point exponents fixed by values appropriate for the present polymers, i.e., $\nu = 1/2$ and $\nu_{\theta} \approx 2/5$ in a good solvent and θ -conditions. The 'crossover exponent' φ for the binary excluded volume interaction equals, $\varphi = 2 - d\nu_{\theta}$, where d is the spatial dimension, and the observed value of φ is close to the predicted value $\varphi = 0.8$ obtained by taking ν_{θ} = 2/5. In the fit in Fig. 5 to eqn (5), the generalized

Table 1 Parameters and their values for universal description of the radius of gyration of compact nanogel particles, see egn (4)

Parameter	$\tau > 0$	$\tau < 0$
c_0 c_1 φ	0.89 5.31 0.8	0.24 2.64 0.8

Paper Soft Matter

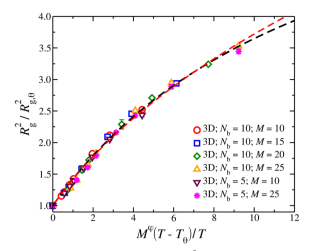


Fig. 5 Average square radius of gyration, R_q^2 , of isolated compact nanogel particles normalized by R_q^2 at θ -point, as a function of the reduced temperature, $\tau = (T - T_{\theta})/T$ and chain length, M, scaled with an exponent φ . The dashed line is a fit of eqn (5) with ν = 1/2 (black line) and ν = 0.588 (red line). The uncertainty estimates correspond to two standard deviations.

dimensionless excluded volume parameter parameter is taken to scale as $z = \tau M^{\varphi}$ and and b is a fitting parameter with values equal to b = 4.16 for $\nu = 1/2$ and b = 2.66 for $\nu = 0.588$, respectively. As discussed by Douglas, 37 there are two extensions of Wiener path model to describe polymer networks where the one dimensional time coordinate of the Wiener path describing linear polymers is replaced by a 'multi-dimensional time' variable that describes the topological structure of the polymer network. One of these models, the Wiener sheet model, leads to the same critical properties as randomly branched polymers, e.g., a critical dimension of 8 for binary excluded volume interactions and 6 for ternary interactions, $\nu =$ 1/4 in the absence of excluded volume interactions, etc., while the other network model has an 'infinite critical dimension' since the average radius of the polymer network only increases logarithmically with the polymer mass in the absence of excluded volume interactions, a well known property of the James and Guth model of polymer networks. 38,39 We expect that the development of a perturbative treatment of excluded volume interaction in 'Wiener sheet' polymers, and a subsequent renormalization of the perturbation theory series expansion in z should lead to an expression of the form eqn (5), but this type of expansion has not yet been attempted. However, there have been perturbative treatments of binary excluded volume interactions and a corresponding RG analysis of randomly branched polymers; see discussion in ESI.†

There has been some disagreement about which model is most appropriate in describing random networks which arise in a wide range of physical problems. We note that Parisi⁴⁰ and Cates⁴¹ have made arguments favoring the properties of the first type of network model. Numerical simulations of random surfaces having a free boundary in three dimensions 42,43 appear to closely accord with ν in three dimensions being exactly 1/2 under self-avoiding conditions, as suggested for our network polymer simulations (Fig. 5).

The crossover function for the swelling of the nanogel particles in eqn (5) provides a reasonable reduction of a swelling data and interpolates between the mass scaling before for the nanogel particles in the good solvent limit where R_g scales as $R_{g} \sim M^{1/2}$ and the observations of the present paper for nanogel particles at their θ -point where $R_{g,q}$ scales as, $R_{g,q} \sim$ $M^{0.4}$. We should note that in a previous paper we initially estimated the exponent ν for the closed gel polymers to have a value consistent with linear polymer chains in a good solvent ($\nu = 0.588$), but we now appreciate the uncertainties in this type of exponent estimation for the inherently limited size of the polymeric structures that we can simulate. Accordingly, we can obtain some insight into this apparent exponent by considering the crossover expression eqn (5) with ν assumed to equal our former estimate, i.e., $\nu = 0.588$. We see in Fig. 5 that this exponent also describes our swelling data for smaller values of z, but the cross-over curve is better described by $\nu = 1/2$ for large z. Minimally, this underscores the uncertainty in the estimation of ν and to the need for further investigation of this quantity for much larger networks. Similar trends were observed in the formation of clusters of branched polymers.⁴⁴ Below we approach this problem from another direction through the scaling of the osmotic pressure with polymer concentration where we find compelling evidence for $\nu = 1/2$ in a good solvent from the scaling of the osmotic pressure of solutions of these polymers in the semi-dilute concentration regime. At any rate, the extension of the linear chain RG crossover description of chain swelling to network polymers seems promising, and we plan to develop this type of crossover description further in the future.

Continuum perturbative models of polymers with excluded volume interactions should only apply for relatively high mass polymers under conditions where chain semi-flexibility and other effects related to the detailed functional form of the pair interaction does not influence the asymptotic scaling. 45-47 Even short random flight polymer chains do not closely resemble Gaussian chains because of short range correlations and chain semi-flexibility makes the approach to this asymptotic scaling slower. The limitations of the two parameter model of polymer excluded volume observed for linear polymers should apply also to randomly branched polymers; see discussion in ESI.†

B. Size of nanogel particles in concentrated solutions

Now that we understand the swelling behavior of compact nanogel particles under infinite dilution conditions, we focus on concentrated solutions where the nanogel particles fill the interstitial space resembling a mesh of polymer chains. Similar to linear polymer solutions, we can define three characteristic concentration regimes: (i) the dilute solution, in which the individual polymers do not overlap, (ii) the semi-dilute solution where polymers are strongly overlapping but the volume fraction of polymers are still small, and (iii) the concentrated solutions. The crossover between dilute and semi-dilute regimes is typically defined by the overlap concentration defined as $\rho/\rho^* = 1$, where $\rho^* = \frac{4\pi}{3} R_{g,r=0}^3$.

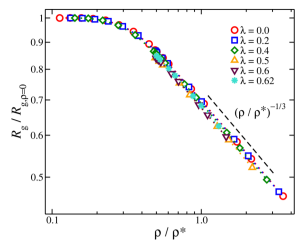


Fig. 6 Average radius of gyration normalized by its value of infinite dilution, $R_{\rm g}/R_{{\rm g},\rho=0}$ as a function of the reduced segmental density, ρ/ρ^{\star} , where $ho^*=rac{4\pi}{3}R_{\mathrm{g},
ho=0}{}^3$

The nanogel particles exhibit two distinct behaviors. In dilute solutions $\rho/\rho^* < 0.2$, we find $R_g/R_{g,r=0} \approx 1$ meaning that in these dilute concentrations the size of the nanogel is not concentration dependent. Above the threshold $\rho/\rho^* \approx 0.2$, we find a sharp crossover to a power-law $R_g/R_{g,r=0} \sim (\rho/\rho^*)^{-\mu}$, where $\mu \approx 1/3$, see Fig. 6. This scaling exponent provides information on how the nanogel particle's size changes once each particle starts to interact with each neighbors in solution. An effect that has been observed in polymer solutions, the corresponding exponent is $\mu \approx 0.12$ for linear chain, $^{48}\mu \approx 0.29$ for ring polymers, 48 and $\mu \approx 0.22$ for single-chain nanoparticles solutions. 49 The starting point in understanding this effect is based on the observation that the conformational properties of linear polymer chains in a melt are essentially the same as those for 'unperturbed' chains without excluded volume interactions, consistent with arguments given by Flory.⁵⁰ Freed and Edwards⁵¹ theoretically explained this behavior from the screening of excluded volume interactions with increasing polymer concentration and it has been confirmed by Small Angle Neutron Scattering (SANS) with labeled linear chains.⁵² For nanogel particles, at concentrations where we observe this emergence of $R_{\rm g}/R_{\rm g,r=0} \sim (\rho/\rho^*)^{-1/3}$ scaling, the interstitial space is filled by the nanogel particles. This is consistent with recent modeling of dense microgels suspensions.⁵³ Thus any volume changes (δV) are correlated by changes in the size of the nanogel particle, suggesting $\delta V \sim \rho^{-1} \sim R_{\rm g}^{3}$. This behavior is observed for all the solvent quality range as seen Fig. 6. The exponent in nanogel particles is much larger than in the case of linear chains but closer to ring polymers.

C. Osmotic pressure and second virial coefficients of compact nanogel particles

The osmotic pressure Π can be estimated from the virial coefficients in the virial equation, a formulation developed by McMillan and Mayer. 54,55 For a monodisperse polymer solution, the compressibility factor Z can be expanded in

powers of ρ ,

$$Z = \frac{M_{\rm w}\Pi}{k_{\rm B}T\rho} = 1 + \sum_{n=1} B_{n+1}\rho^n,$$
 (6)

where ρ is the polymer number density, M_w is the molecular mass of the polymer, Π is the osmotic pressure, and the coefficients B_n are the virial coefficients. The osmotic pressure is one of the most easily accessible quantities in polymer physics, and knowledge of B_n provides the estimation of Π in the dilute regime in which $\rho R_{\rm g}^{\ 3} \ll 1$. The B_n coefficients also contain interparticle information on the nature of the interaction between the particles in the solution, which are temperature-dependent. In particular, B_2 provides information of the pair-wise interactions, B_3 the 2- and non-additive 3-body interactions. The second virial captures the deviation from the ideal behavior $(M_w\Pi\rho = 1)$ and it is related to the effective interaction u(r) between two polymers,

$$B_2 = -2\pi \int f(r)r^2 \mathrm{d}r,\tag{7}$$

where f(r) is the Mayer function defined as $f(r) = \exp(-u(r)/kT)$ – 1.55 Higher-order terms if known provide a wider range in which eqn (6) is accurate. For polymer solutions and gels, the nature of the virial coefficients deviates from the McMillan and Mayer framework, and their meaning remains an active topic of research. 56,57 Specifically, the determination of the values of the second and third virial coefficients (or even higher-order terms) requires to be determined solely on the interaction among two and three (or more for high order terms) solvated particles, respectively.⁵⁸ From an experimental perspective, this is challenging. However, in practice, knowledge of the second and third virial coefficients provides a reliable prediction for the osmotic pressure in the dilute regime where the expression is accurate up to $B_2\rho \approx 1$, even for relatively small values of the degree of polymerization. The value of coefficients B_n depends on the polymer solution and temperature. Even in the context of the soft sphere model, where the polymer is viewed in an average sense as a soft interpenetratable Gaussian 'blob' representing a segmental density cloud, 59-61 the nature of the virial coefficients is not clear.

To estimate the virial coefficients, we consider the virial expansion for ρ as power series in terms of a reduced concentration that is defined in terms of the second virial coefficient. 56,62,63 In particular, we have,

$$\frac{M_{\rm w}\Pi}{k_{\rm B}T\rho} = 1 + \hat{\rho} + \gamma\hat{\rho}^2 + \dots \tag{8}$$

where $\gamma \approx 0.25$ is the dimensionless virial ratio, $\hat{\rho} = \rho M_{\rm w} B_2$. Concentrated polymer solutions and gels exhibit similar osmotic pressure behavior at the high concentrations where packing interactions of the polymers becomes predominant and we correspondingly hypothesize that this universality also applies in the semi-dilute regime between the dilute and concentrated regimes. In particular, we hypothesize that eqn (8) applies as an approximation to both polymer solutions of different architectures and polymer gels.

The θ -temperature or solvent is defined when the temperature and/or solvent results in $B_2 = 0$, 64,65 requiring the evaluation of

the integral in eqn (7) at $\rho \to 0$. This calculation can be done analytically for simple models like hard spheres and square well potential, but it becomes challenging, requiring extensive computation/sampling resources, with increasing the complexity of the molecular architecture. There are a few studies of the estimation of the second virial of different polymers having molecular topology near the θ -point than in the good solvent conditions, such as stars 47,66,67 and rings 68,69 that offer some insight into this problem. We calculate the pair-correlation of the center-of-mass of star polymers in dilute solutions, see Fig. 7, and by utilizing $g(r) = e^{-u(r)/kT}$, we obtain the concentrationdependent second virial by eqn (7). We note that by doing this calculation in a polymer solution, we effectively mimic the experimental difficulties in estimating B_2 solely based on the interaction of two particles/polymers. We minimize this problem by repeating the process by reducing the polymer concentration and identifying the second virial by extrapolating to $\rho \to 0$, see inset of Fig. 7. As we decrease the solvent quality (increasing λ), the adjustment needed by extrapolating to $\rho \rightarrow 0$ becomes smaller. Moreover, the dependence of B_2 in our star solution model as a function of λ suggest that $\lambda_{\theta,\text{star}} \approx 0.7$. This value corresponds to a lower θ -temperature than linear chains, consistent with experimental observations of the star and ring^{68,70} polymers. Now that we have determined the values of B_2 at different quality solvents, we use them to compact nanogel particles composed of star polymers.

The osmotic pressure of compact nanogel particles is presented in Fig. 8. We find that the osmotic pressure of both systems (gels and star polymer solutions) in the high-density regime are in agreement even though we used the same value for B_2 in both cases. This agreement confirms our initial hypothesis that these two systems exhibit approximately the same osmotic behavior at high densities. The deviation is expected at lower densities because the nanogel particles cease to behave as a unified macrogel structure or a star polymer

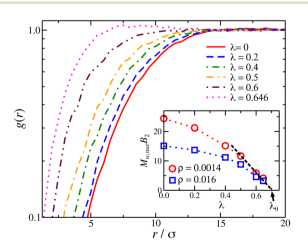


Fig. 7 Pair correlation function, g(r), for center of mass star polymer (f = 4 arms and M = 25 arm length) solution at a segmental density $\rho\sigma^3 \approx 0.016$. Inset: Second virial of star polymer solutions as a function of λ ; results at different polymer concentrations are also presented. The dashed line is an extrapolation to identify the θ -point at $B_2 = 0$.

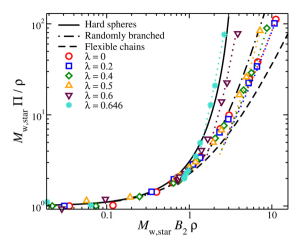


Fig. 8 Compressibility factor, $M_{w,star}\Pi/\rho$, for compact nanogel particles (dotted lines) and star polymer solutions (symbols), as a function of $B_2 M_{\text{\tiny MM}} \rho$ for different values of solvent quality parameter, λ . The cases of hard spheres based on the Carnahan-Starling, 71 flexible chains, 72 and randomly branched structures⁵⁶ are also presented.

solution. Another consequence of our assumption is that $\lambda_{\theta, \text{gel}}$ = $\lambda_{\theta, \rm star} \approx 0.7$, which is in agreement with our estimation of the θ -point by the R_g mass scaling as we discussed above. In the case of star polymer solutions, T_{θ} in gels is also expected to be lower than in linear chains.

D. Structure factor

To probe the structure of nanogel particles, we focus on calculating the spatial correlations between the polymer segments. The structure factor, S(q), is a suitable property for this purpose and describes the mean correlations in the positions of a collection of point particles distributed in space. 73 S(q) is defined 55 as:

$$S(q) = \frac{1}{N_s} \left\langle \sum_{j=1}^{N_s} \sum_{k=1}^{N_s} \exp\left[-i\mathbf{q} \cdot \left(\mathbf{r}_j - \mathbf{r}_k\right)\right] \right\rangle, \tag{9}$$

where $i = \sqrt{-1}$, $q = |\mathbf{q}|$ is the wave number, \mathbf{r}_i is the position of particle j, $\langle \rangle$ denote the time average, and N_s is the total number of polymer segments defined as $N_s = N_x N_y N_z (fM + 1)$.

The scattering profiles generated by our model can be described by the following functional form,

$$S(q) = Aq^{-\mu} + \frac{S_0}{1 + (\xi q)^2}. (10)$$

The heterogeneous structure formation resulting in a steep upturn in S(q) is captured in the first term. Both parameters are A and μ are fitting parameters, where μ describes the fractal nature of the clusters formed at length scales larger than the size of the polymer chains and typically, in experiments, the scaling exponent μ is found to be in the range 2 to 5.⁷⁴ In polymer systems, clustering often plays an important role, however, large clusters do not contribute significantly to the thermodynamics of the solution but strongly modify the scattering spectrum in the low-q range.⁷⁴ The second term, given by Ornstein-Zernike expression⁷⁵ (Lorentzian function), captures the local structure and interactions between the polymer chains composing the

nanogel structure. The parameter ξ is a 'correlation length' that describes that local packing of the chain segments and S_0 is the estimated height of the plateau in the absence of heterogeneous structure formation. We emphasize that eqn (10) is phenomenological and we are following common experimental practice in defining the correlation length as prescribed by eqn (10).

At highly dilute concentrations of compact nanogel particles, the form factor P(q) is typically described by the "fuzzy sphere" model, 76 in which the particles are described by a dense homogeneous core and an outer loose corona.77-79 At length scales larger than the size of the nanogel particle, there is a sharp increase in P(q) that results in $P(q \to 0) \to M_w$. At length scales near R_g , and sufficiently high values of $N_b > 5$, there is a primary scattering peak characteristic of particle-like scattering. A powerlaw behavior is observed at high q-values corresponding to the fractal nature of the polymer chains. We proposed an extension to the fuzzy sphere model to improve the description of the scattering profile at higher q-values, for more details see ref. 20. A typical example of S(q) data for a nanogel particle and a fit to the "extended fuzzy sphere model" is presented in Fig. 9(a).

The scattering features of an isolated nanogel particle disappear in concentrated solutions of nanogel particles because the interface surrounding the nanogel particles vanishes once the nanogel particles start to interpenetrate. In other words, the scattering features of nanogel particles become indistinguishable at moderate and high concentrations of these particles. In the athermal solvents, the polymer segments are relatively uniformly spatially distributed and fill the space resulting in a plateau at lower q-values in S(q), see Fig. 9(a). The height of this plateau is often in experiments extrapolated to q = 0 to determine the S(q = 0)value, which provides information about the long-range density fluctuations. In equilibrium, $S(0) = kT\rho\kappa_T$, where κ_T is the isothermal compressibility (we will discuss more about κ_T below). The height of this plateau decreases with an increase in Π . At $\Pi \approx$ 0.1, the structure factor becomes equal to unity over a wide range of length scales ($q\sigma < 1$), and this is analogous to an ideal gas that exhibits S(q) = 1 for all q. In other words, there is a pressure threshold where the gel effectively loses its structural features.

Near the θ -point, the chains in the nanogel structure begin to shrink and partially 'collapse'. This effect creates denser regions in the gel structure than an assumed homogeneous spatial segmental distribution. The contrast between these denser regions, which occur randomly within the gel structure, and the voids (regions in which there are no gel segments) results in the formation of heterogeneous structures. This is evident in the structure factor profiles at low-q regime, where a clustering term $S_c(q) = Aq^{-\mu}$ is needed to describe the emergence of this behavior. The emergence of clusters/heterogeneous structure formation has been observed experimentally.80 The heterogeneous structures become more dominant because the collapsed structures increase in size as the solvent quality worsens (increasing λ) below but near the θ -point, $\lambda_{\theta} < \lambda \approx 0.7$. Eventually, these collapsed structures increase in size and start to merge as the solvent quality worsens, $\lambda > 0.7$. Moreover, any voids in the gel structure shrink, resulting in a more homogeneous and dense structure. An increase in the osmotic pressure reduces the impact of heterogeneous structure formation in S(q). We note that the influence of the solvent quality is also expected to influence the scaling exponent in the high qregime in S(q) scattering profiles.⁸¹ For linear chains and other regular polymers, it is expected to scale as $S(q) \sim q^{1/\nu}$. For the chains in the nanogel particles, we find that $S(q) \sim q^{-1.3}$ for $\lambda <$ 0.7, suggesting that the chains are significantly stretched.

E. Osmotic modulus

The osmotic modulus, *K*, is defined as the inverse of isothermal compressibility, $\kappa_{\rm T}$,

$$K \equiv \kappa_{\rm T}^{-1} = \rho \frac{\partial \Pi}{\partial \rho} = \left(-\frac{1}{V} \frac{\partial V}{\partial \Pi} \right)_{T}^{-1} = \left(\frac{\langle V^2 \rangle - \langle V \rangle^2}{kT \langle V \rangle} \right)_{NPT}^{-1}. \tag{11}$$

We find that for concentrated solutions, the osmotic modulus of the nanogel particles scales with ρ for the whole range

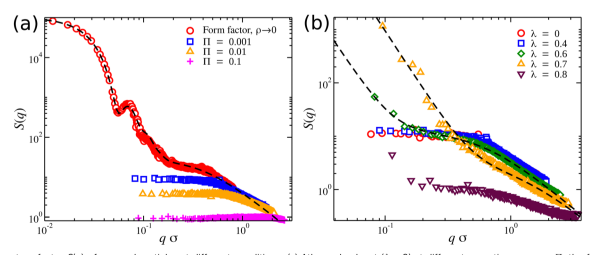


Fig. 9 Structure factor S(q) of nanogel particles at different conditions. (a) Athermal solvent ($\lambda = 0$) at different osmotic pressures Π ; the form factor (dashed line) of an isolated nanogel particle is also presented along with a fit to the "extended fuzzy sphere" model.²⁰ (b) Different solvent qualities at osmotic pressure Π = 0.0001; the dashed lines are fits with eqn (10)

Paper Soft Matter

of osmotic pressures. As the osmotic pressure increases, the material's resistance to volumetric changes increases with ρ raised to a scaling exponent that describes the fractal distribution of polymer segments in the gel structure. It is established in experiments and in theory that $K \sim \rho^{\delta}$, where $\delta = 3\nu/(3\nu - 1)$ corresponding to $\delta = 9/4$ for $\nu = 0.588$ and $\delta = 3$ for $\nu = 1/2$. ^{75,82,83} Alternative functional forms for osmotic modulus based on empirical observations have been developed. 56,84,85 Linear chain solutions under good solvent conditions exhibit $K \sim$ $\rho^{9/4}$ as expected. However, in our nanogel particle solutions, the scaling exponent was found to be $K \sim \rho^3$ in good solvent, see Fig. 10. This highlights the difference in the fractal nature of the gel from that of linear chain solution. We also note that the experimental procedure of gelation through the cross-linking of long linear chains may lead to the formation of gel structures in which the chains are trapped in a non-equilibrium state where the linear chain scaling of the osmotic modulus is apparently preserved. Recent measurements of polymers cross-linked to form a solution network have shown a tendency to exhibit the osmotic scaling exponents of the parent linear polymer chains. It has been observed that diffusion of randomly branched polymers and the viscosity of branched polymer solutions in which the branched polymers are formed from polymerizing oligomers in solution can be quite different from solutions in

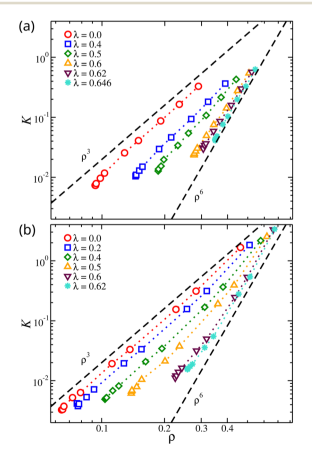


Fig. 10 Osmotic modulus, K, of compact nanogel particles as a function of segmental density, ρ . Results for different solvent qualities are also presented. The dashed lines are guides for the eye.

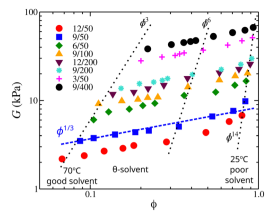


Fig. 11 Elastic modulus of poly(vinyl acetate)/isopropyl alcohol gels as a function of the polymer volume fraction, ϕ . The dotted lines approximately correspond to isotherms with a temperature range is (from left to right) 70 °C, 60, 55 °C, 52 °C (identified as the θ -temperature), 50 °C, 45 °C, 37 °C, 30 °C, and 25 °C. Samples are identified by their polymer concentration at the time of introduction of crosslinks and with their degree of crosslinking; for example, 12/50 means that the concentration of PVA solution at which the crosslinks were introduced was 12 mass% and the degree of crosslinking was 50. For more details see ref. 88.

which the branched polymers are formed by cross-linking high molecular mass polymers.86,87 There is evidently a 'memory effect' of the conditions of cross-linking that requires further investigation. The fractal nature of our gel model is also apparent in our solutions of network polymers. In particular, we find a scaling exponent of $K \sim \rho^6$ near the θ -point, suggesting $\nu_{\theta} \approx 2/5$, which is in agreement with our previous calculations on R_g mass scaling. This scaling exponent is expected to increase further in the poorer solvent conditions but is not expected to change with the variation of the chain length, see Fig. 10. However, the range of *K* for the same range of Π was significantly different. Longer chains provide the gel with a wider range of K variation. The gel can initiate its resistance to volumetric changes at lower segmental densities and reach higher values of K at higher segmental densities.

To evaluate these predictions, we examine the elastic modulus G of poly(vinyl acetate)/isopropyl alcohol gels as a function of the polymer volume fraction. The elastic modulus defines the elastic pressure that contracts the gel while the osmotic pressure expands it. The shear moduli of osmotically deswollen gel samples at constant temperature exhibited a scaling exponent of 1/3 with the polymer concentration in both good and θ -solvents, see Fig. 11.⁸⁸ At equilibrium and an excess amount of solvent, the elastic pressure balances the osmotic pressure, suggesting that the scaling trends found in osmotic modulus will be expected in elastic modulus. The elastic modulus of poly(vinyl acetate)/isopropyl alcohol gels presented in Fig. 11 confirms the predictions for the scaling exponents of our model.

IV. Conclusions

We investigated the structure and osmotic properties of polymer gels composed of compact nanogel particles over a wide

range of solvent qualities by molecular dynamics simulations with a bead-spring model. We characterized the osmotic pressure changes in the gel by estimating the second virial coefficients. Moreover, we identified the conditions at which the θ -point emerges. We found that the location of θ -point is the same in both dilute and concentrated conditions, however, the θ point occurred at a lower temperature than linear chains. Based on the location of the θ -point, we developed an empirical master curve to describe the swelling of the nanogel particles in dilute concentrations. Even at small values of osmotic pressures, the nanogel particles fill the interstitial space so that the particle-like scattering is no longer apparent at moderate concentrations. We also find that improving the solvent quality enhances spatial segmental uniformity in an average sense, while significant heterogeneous structure formation occurs near the θ -point. Well below the θ -point, these heterogeneous structures start to diminish as the gels have nearly collapsed at high osmotic pressures. We also investigate the degree of swelling and structure of compact nanogel particles with the variation of solvent quality at both infinite dilution and concentrated conditions. Finally, we calculated the osmotic modulus and the scaling exponents in good and θ -point conditions found are characteristic of lattice animals.37 These findings provide useful guidelines for the design of gels and a better understanding of the role of gels in biological function.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 J. P. Rolland, B. W. Maynor, L. E. Euliss, A. E. Exner, G. M. Denison and J. M. DeSimone, Direct fabrication and harvesting of monodisperse, shape-specific nanobiomaterials, *J. Am. Chem. Soc.*, 2005, 127, 10096–10100.
- 2 X. Zhang, S. Malhotra, M. Molina and R. Haag, Micro-and nanogels with labile crosslinks-from synthesis to biomedical applications, *Chem. Soc. Rev.*, 2015, 44, 1948–1973.
- 3 K. S. Soni, S. S. Desale and T. K. Bronich, Nanogels: An overview of properties, biomedical applications and obstacles to clinical translation, *J. Controlled Release*, 2016, 240, 109–126.
- 4 K. Y. Lee and D. J. Mooney, Hydrogels for tissue engineering, *Chem. Rev.*, 2001, **101**, 1869.

- 5 R. Langer and D. A. Tyrrell, Designing materials for biology and medicine, *Nature*, 2004, 428, 487.
- 6 S. Lv, D. M. Dudek, Y. Cao, M. M. Balamurali, J. Gosline and H. Li, Designed biomaterials to mimic the mechanical properties of muscles, *Nature*, 2010, 465, 69.
- 7 N. Morinloto, X. P. Qiu, F. M. Winnik and K. Akiyoshi, Dual stimuli-responsive nanogels by self-assembly of polysaccharides lightly grafted with thiol-terminated poly(Nisopropylacrylamide) chains, *Macromolecules*, 2008, 41, 5985–5987.
- 8 E. B. Dickerson, W. H. Blackburn, M. H. Smith, L. B. Kapa, L. A. Lyon and J. F. McDonald, Chemosensitization of cancer cells by siRNA using targeted nanogel delivery, *BMC Cancer*, 2010, 10, 10.
- 9 P. Srisawat, M. Higuchi-Takeuchi, R. Honda, T. Shirai, A. Kondo and Y. Hoshino, *et al.*, Engineered nanogel particles enhance the photoautotrophic biosynthesis of polyhydroxyalkanoate in marine photosynthetic bacte, *ACS Sustainable Chem. Eng.*, 2022, 10, 4133–4142.
- 10 G. V. Kolmakov, K. Matyjaszewski and A. C. Balazs, Harnessing labile bonds between nanogel particles to create self-healing materials, ACS Nano, 2009, 3, 885–892.
- 11 T. Liang, J. Hou, M. Qu, M. Zhao and I. Raj, High-viscosity α -starch nanogel particles to enhance oil recovery, *RSC Adv.*, 2020, **10**, 8275–8285.
- 12 A. M. Atta, H. A. Al-Lohedan and K. A. Al-Haddad, Epoxy coating with embedded self-healing networks formed by nanogel particles, *RSC Adv.*, 2016, **6**, 41229–41238.
- 13 D. Keskin, G. Zu, A. M. Forson, L. Tromp, J. Sjollema and P. van Rijn, Nanogels: A novel approach in antimicrobial delivery systems and antimicrobial coatings, *Bioact. Mater.*, 2021, 6, 3634–3657.
- 14 W. Wu, N. Mitra, E. C. Yan and S. Zhou, Multifunctional hybrid nanogel for integration of optical glucose sensing and self-regulated insulin release at physiological pH, ACS Nano, 2010, 4, 4831–4839.
- 15 J. A. Buckwalter, H. J. Mankin and A. J. Grodzinsky, Articular cartilage and osteoarthritis, *Instr. Course Lect.*, 2005, 54, 465.
- 16 A. D. Pearle, R. F. Warren and S. A. Rodeo, Basic science of articular cartilage and osteoarthritis, *Clin. Sports Med.*, 2005, 24, 1–12.
- 17 J. Martel-Pelletier, C. Boileau, J. P. Pelletier and P. J. Roughley, Cartilage in normal and osteoarthritis conditions, *Best Pract. Res., Clin. Rheumatol.*, 2008, 22, 351–384.
- 18 D. Heinegard and T. Saxne, The role of the cartilage matrix in osteoarthritis, *Nat. Rev. Rheumatol.*, 2011, 7, 50–56.
- 19 F. Horkay and P. J. Basser, Composite hydrogel model of cartilage predicts its load-bearing ability, *Sci. Rep.*, 2020, **10**, 1–7.
- 20 A. Chremos, J. F. Douglas and F. Horkay, Structure and conformational properties of ideal nanogel particles in athermal solutions, *J. Chem. Phys.*, 2021, **155**, 134905.
- 21 A. Chremos, J. F. Douglas and F. Horkay, Influence of network defects on the conformational structure of nanogel particles: From "closed compact" to "open fractal" nanogel particles, *J. Chem. Phys.*, 2022, **156**, 094903.
- 22 J. D. Weeks, D. Chandler and H. C. Andersen, Role of repulsive forces in determining the equilibrium structure of simple liquids, *J. Chem. Phys.*, 1971, **54**, 5237–5247.

Paper Soft Matter

- 23 M. O. Steinhauser, A molecular dynamics study on universal properties of polymer chains in different solvent qualities. Part I. A review of linear chain properties, J. Chem. Phys., 2005, 122, 094901.
- 24 A. Chremos, E. Glynos, V. Koutsos and P. J. Camp, Adsorption and self-assembly of linear polymers on surfaces: A computer simulation study, Soft Matter, 2008, 5, 637-645.
- 25 A. Chremos, E. Glynos, V. Koutsos and P. J. Camp, Adsorption of star polymers: Computer simulations, Soft Matter, 2010, 6, 1483-1493.
- 26 A. Chremos, A. Nikoubashman and A. Z. Panagiotopoulos, Flory-Huggins parameter chi, from binary mixtures of Lennard-Jones particles to block copolymer melts, J. Chem. Phys., 2014, 140, 054909.
- 27 J. Dudowicz, K. F. Freed and J. F. Douglas, New patterns of polymer blend miscibility associated with monomer shape and size asymmetry, J. Chem. Phys., 2002, 116, 9983-9996.
- 28 J. Dudowicz, K. F. Freed and J. F. Douglas, Beyond Flory-Huggins theory: New classes of blend miscibility associated with monomer structural asymmetry, Phys. Rev. Lett., 2002, 88, 095503.
- 29 B. D. Butler, G. Ayton, O. G. Jepps and D. J. Evans, Configurational temperature: Verification of Monte Carlo simulations, J. Chem. Phys., 1998, 109, 6519-6522.
- 30 W. R. Krigbaum and D. O. Geymer, Thermodynamics of polymer solutions. The polystyrene-cyclohexane system near the Flory theta temperature, J. Am. Chem. Soc., 1959, 81, 1859-1868.
- 31 A. Y. Grosberg and D. V. Kuznetsov, Single-chain collapse or precipitation? Kinetic diagram of the states of a polymer solution, Macromolecules, 1993, 26, 4249-4251.
- 32 X. Wang and C. Wu, Light-scattering study of coil-to-globule transition of a poly(N-isopropylacrylamide) chain in deuterated water, Macromolecules, 1999, 32, 4299-4301.
- 33 F. A. Escobedo and J. J. de Pablo, Simulation and theory of the swelling of athermal gels, J. Chem. Phys., 1997, 106, 793-810.
- 34 A. Karim, J. F. Douglas, F. Horkay, L. J. Fetters and S. K. Satija, Comparative swelling of gels and polymer brush layers, Phys. B, 1996, 143, 221.
- 35 J. F. Douglas and K. F. Freed, Renormalization and the twoparameter theory, Macromolecules, 1984, 23, 2344-2354.
- 36 K. F. Freed and J. F. Douglas, Comparison between Borel resummation and renormalization group descriptions of polymer expansion, J. Chem. Phys., 1988, 88, 2764-2768.
- 37 J. F. Douglas, Swelling and growth of polymers, membranes, and sponges, Phys. Rev. E: Stat. Phys., Plasmas, Fluids, Relat. Interdiscip. Top., 1996, 54, 2677-2689. The scaling exponent ν for polymer without excluded volume is $\nu = 1/4$ and it is not equivalent to branched polymers at their θ -point or having screened excluded volume interactions where $\nu \approx 2/$ 5. "Percolation clusters" are branched polymers with screened excluded volume interactions.
- 38 E. Guth and M. James, Elastic and thermoelastic properties of rubber like materials, Ind. Eng. Chem., 1941, 33, 624.
- 39 M. James and E. Guth, Statistical thermodynamics of rubber elasticity, J. Chem. Phys., 1953, 21, 1039.

- 40 G. Parisi, Hausdorff dimensions and gauge theories, Phys. Lett., 1979, 81B, 357.
- 41 M. E. Cates, The Liouville field theory of random surfaces: When is the bosonic string a branched polymer?, EPL, 1998, 7, 719.
- 42 U. Glaus, Monte Carlo simulation of self-avoiding surfaces in three dimensions, Phys. Rev. Lett., 1996, 56, 1996.
- 43 U. Glaus and T. L. Einstein, On the universality class of planar self-avoiding surfaces with fixed boundary, J. Phys. A, 1987, 20, L105.
- 44 E. V. Patton, J. A. Wesson, M. Rubinstein, J. C. Wilson and L. E. Oppenheimer, Scaling properties of branched polyesters, Macromolecules, 1989, 22, 1946-1959.
- 45 H. Yamakawa, On the theory of the second virial coefficient for polymer chains, Macromolecules, 1992, 25, 1912-1916.
- 46 B. J. Cheravil, J. F. Douglas and K. F. Freed, Effect of residual interactions on polymer properties near the theta point, J. Chem. Phys., 1985, 83, 5293-5310.
- 47 B. J. Cherayil, J. F. Douglas and K. F. Freed, Effect of residual interactions on polymer properties near the theta point. II. Higher moments and comparison with Monte Carlo calculations, J. Chem. Phys., 1987, 87, 3089-3098.
- 48 S. Y. Reigh and D. Y. Yoon, Concentration dependence of ring polymer conformations from Monte Carlo simulations, ACS Maro Lett., 2013, 2, 296-300.
- 49 A. J. Moreno, F. L. Verso, A. Arbe, J. A. Pomposo and J. Colmenero, Concentrated solutions of single-chain nanoparticles: A simple model for intrinsically disordered proteins under crowding conditions, J. Phys. Chem. Lett., 2016, 7,838-844.
- 50 P. J. Flory, Statistical Mechanics of Chain Molecules, Interscience, New York, 1969.
- 51 K. F. Freed and S. F. Edwards, Polymer viscosity in concentrated solutions, J. Chem. Phys., 1974, 61, 3626-3633.
- 52 M. Daoud, J. P. Cotton, B. Farnoux, G. Jannink, G. Sarma and H. Benoit, et al., Solutions of flexible polymers. Neutron experiments and interpretation, Macromolecules, 1975, 8, 804-818.
- 53 S. V. Nikolov, A. Fernandez-Nieves and A. Alexeev, Behavior and mechanics of dense microgel suspensions, Proc. Natl. Acad. Sci. U. S. A., 2020, 117, 27096-27103.
- 54 W. G. McMillan and J. E. Mayer, The statistical thermodynamics of multicomponent systems, J. Chem. Phys., 1945,
- 55 J. P. Hansen and I. R. McDonald, Theory of simple liquids, Academic Press, Cambridge, 2006.
- 56 T. Aberle and W. Burchard, Universality in behaviour of branched macromolecules in semidilute solutions, Comput. Theor. Polym. Sci., 1997, 7, 215-225.
- 57 S. Vafaei, B. Tomberli and C. G. Gray, McMillan-Mayer theory of solutions revisited: Simplifications and extensions, J. Chem. Phys., 2014, 141, 154501.
- 58 W. Burchard, Particle scattering factors of some branched polymers, Macromolecules, 1977, 10, 919.
- 59 P. J. Flory and W. R. Krigbaum, Statistical mechanics of swelling of network structures, J. Chem. Phys., 1950, 18, 1086.

- 60 W. Gobush, K. Sole and W. H. Stockmayer, Statistical mechanics of random-flight chains. V Excluded volume expansion and second virial coefficient for linear chains of varying shape, J. Chem. Phys., 1974, 60, 12.
- 61 C. N. Likos, H. Löwen, M. Watzlawek, B. Abbas, O. Jucknischke and J. Allgaier, et al., Star polymers viewed as ultrasoft colloidal particles, Phys. Rev. Lett., 1998, 80, 4450-4453.
- 62 I. Noda, N. Kato, T. Kitano and M. Nagasawa, Thermodynamic properties of moderately concentrated solutions of linear polymers, Macromolecules, 1981, 14, 668-676.
- 63 T. Yasuda, N. Sakumichi, U. I. Chung and T. Sakai, Universal equation of state describes osmotic pressure throughout gelation process, Phys. Rev. Lett., 2020, 125, 267801.
- 64 P. J. Flory, The statistical thermodynamics of multicomponent systems, J. Chem. Phys., 1949, 7, 303.
- 65 H. A. G. Chermin and J. W. Kennedy, The statistical thermodynamics of multicomponent systems, Macromolecules, 1972, 5, 655-656.
- 66 J. Batoulis and K. Kremer, Residual 3-body interactions of a θ -polymer, *EPL*, 1988, 7, 683.
- 67 S. Huissmann, R. Blaak and C. N. Likos, Star polymers in solvents of varying quality, Macromolecules, 2009, 42, 2806-2816.
- 68 J. Roovers, Dilute-solution properties of ring polystyrenes, J. Polym. Sci., 1985, 23, 1117-1126.
- 69 A. Takano, Y. Kushida, Y. Ohta, K. Masuoka and Y. Matsushita, The second virial coefficients of highlypurified ring polystyrenes in cyclohexane, Polymer, 2009, **50**, 1300-1303.
- 70 J. Roovers and P. M. Toporowski, Synthesis of high molecular weight ring polystyrenes, Macromolecules, 1983, 16, 843-849.
- 71 N. F. Carnahan and K. E. Starling, Equation of state for nonattracting rigid spheres, I. Chem. Phys., 1969, 51, 635.
- 72 P. J. Flory, Principles of Polymer Chemistry, Cornell University Press, Ithaca, 1953.
- 73 K. Mortensen, Structural studies of polymer systems using smallangle neutron scattering, in Advanced functional molecules and polymers processing and spectroscopy, ed. H. Nalwa, Overseas Publishers Association, 2001, pp. 223-269, vol. 2.
- 74 F. Horkay and B. Hammouda, Small-angle neutron scattering from typical synthetic and biopolymer solutions, Colloid Polym. Sci., 2008, 286, 611-620.

- 75 P. G. de Gennes, Scaling Concepts in Polymer Physics, Cornell University Press, Ithaca, 1979.
- 76 M. Stieger, W. Richtering, J. S. Pedersen and P. Lindner, Small-angle neutron scattering study of structural changes in temperature sensitive microgel colloids, J. Chem. Phys., 2004, 120, 6197.
- 77 N. Gnan, L. Rovigatti, M. Bergman and E. Zaccarelli, *In silico* synthesis of microgel particles, Macromolecules, 2017, 50, 8777-8786.
- 78 S. Bergmann, O. Wrede, T. Huser and T. Hellweg, Superresolution optical microscopy resolves network morphology of smart colloidal microgels, Phys. Chem. Chem. Phys., 2018, 20, 5074-5083.
- 79 A. J. Moreno and F. L. Verso, Computational investigation of microgels: Synthesis and effect of the microstructure on the deswelling behavior, Soft Matter, 2018, 14, 7083-7096.
- 80 A. M. Hecht, F. Horkay, S. Mallam and E. Geissler, Scattering properties of polymer gels at the theta temperature, Macromolecules, 1992, 25, 6915-6920.
- 81 S. Goossen, A. R. Bras, W. P. A. Wischnewski, M. D. Richter, M. Rubinstein and J. Roovers, et al., Influence of the solvent quality on ring polymer dimensions, Macromolecules, 2015, 48, 1598-1605.
- 82 E. Geissler, A. M. Hecht, F. Horkay and M. Zrinyi, Compressional modulus of swollen polyacrylamide networks, Macromolecules, 1988, 21, 2594-2599.
- 83 F. Horkay, A. M. Hecht and E. Geissler, The effects of crosslinking on the equation of state of a polymer solution, J. Chem. Phys., 1989, 91, 2706-2711.
- 84 G. Merkle, W. Burchard, P. Lutz, K. F. Freed and J. Gao, Osmotic pressure of linear, star, and ring polymers in semidilute solution. A comparison between experiment and theory, Macromolecules, 1993, 26, 2736-2742.
- 85 G. Galinsky and W. Burchard, Starch fractions as examples for nonrandomly branched macromolecules. 2. Behavior in the semidilute region, Macromolecules, 1996, 29, 1498-1506.
- 86 J. E. Martin, J. Wilcoxon and D. Adolf, Critical exponents for the sol-gel transition, Phys. Rev. A: At., Mol., Opt. Phys., 1987, 36, 1803.
- 87 J. A. M. Smit, J. A. P. P. Van Dijk, M. G. Mennen and M. Daoud, Polymer size exponents of branched dextrans, Macromolecules, 1992, 25, 3585-3590.
- 88 M. Zrinyi and F. Horkay, On the elastic modulus of swollen gels, Polymer, 1987, 28, 1139-1143.