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A Critical Appraisal of RAFT-Mediated Polymerization-Induced Self-Assembly

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ABSTRACT: Recently, polymerization-induced self-assembly (PISA) has become widely recognized as a robust and efficient route to produce block copolymer nanoparticles of controlled size, morphology, and surface chemistry. Several reviews of this field have been published since 2012, but a substantial number of new papers have been published in the last three years. In this Perspective, we provide a critical appraisal of the various advantages offered by this approach, while also pointing out some of its current drawbacks. Promising future research directions as well as remaining technical challenges and unresolved problems are briefly highlighted.

INTRODUCTION

The self-assembly of surfactant amphiphiles has been studied for over 100 years;1 McBain was the first to discuss the formation of micelles within soap solutions in 1913.2 However, the study of block copolymer self-assembly only began in the early 1960s3–6 following the discovery of living anionic polymerization by Szwarc et al., enabling access to well-defined block copolymers for the first time.7,8 Traditionally, block copolymer self-assembly has been achieved via two steps: (i) initial molecular dissolution of the copolymer chains and (ii) reduction of the solvency for one of the blocks to drive microphase separation. For example, the Eisenberg group dissolved poly(4-vinylpyridine)–polystyrene (P4VP–PS) diblock copolymers in N,N-dimethylformamide (DMF) and gradually added either water or methanol (non-solvents for PS) to induce the formation of spherical micelles.9 They later showed that more complex morphologies (e.g., spheres, rods, or vesicles) could be produced by the same approach using poly(acrylic acid)–PS (PAA–PS) diblock copolymers with varying degrees of polymerization (DPs) of the two blocks.10,11

The development of living radical polymerization (LRP) chemistries12–15 over the past two decades has enabled the synthesis of many new functional diblock copolymers. A wide range of diblock copolymer nano-objects has been prepared using post-polymerization processing routes, including cylindrical (or worm-like) micelles,16–18 vesicles (or polymersomes),19,20 shell cross-linked micelles,21,22 toroids,23 schizospheric micelles24,25 and vesicles,26 and micellar gels27–29 as well as more complex morphologies.30–32 However, final copolymer concentrations are rather low (<1.0% w/w) in almost all cases, which precludes many potential commercial applications.

Over the past eight years, considerable attention has been focused on developing an alternative route to produce block copolymer nano-objects known as polymerization-induced self-assembly (PISA). Typically, a soluble homopolymer (A) is chain-extended using a second monomer in a suitable solvent such that the growing second block (B) gradually becomes insoluble, which drives in situ self-assembly to form AB diblock copolymer nano-objects. The A block is usually prepared via solution polymerization and acts as a steric stabilizer, while the insoluble B block is prepared via either dispersion or aqueous emulsion polymerization (depending on the monomer solubility in the continuous phase). This process is shown schematically in Scheme 1. By varying the DPs of the two blocks, either spheres or higher order morphologies (e.g., worms or vesicles) can be obtained. In principle, PISA syntheses can be conducted using any type of living polymerization,38–43 but in practice, the majority of literature
examples are based on reversible addition−fragmentation chain transfer (RAFT) polymerization. This radical-based chemistry enables PISA syntheses to be conducted with many functional monomers in a wide range of solvents, including water, lower alcohols, non-polar solvents (such as n-alkanes, mineral oil, and poly(α-olefins)), and also more exotic media such as ionic liquids. One very important advantage of such PISA formulations is that reactions can be conducted at relatively high solids (25−50% w/w). The versatility of this approach is illustrated in Scheme 2, which shows PISA formulations in water, ethanol, and n-dodecane. The same poly(glycerol monomethacrylate) (PGMA) stabilizer block can be used for either the RAFT aqueous dispersion polymerization of 2-hydroxypropyl methacrylate (HPMA) or the RAFT aqueous emulsion polymerization of benzyl methacrylate (BzMA). BzMA can also be used as the core-forming block for RAFT dispersion polymerization in polar solvents, such as ethanol, using a PHPMA stabilizer or for RAFT dispersion polymerization in non-polar solvents, such as n-heptane, n-dodecane, or mineral oil, using a poly(lauryl methacrylate) (PLMA) stabilizer. These particular literature examples serve to demonstrate that just four blocks can provide the basis for four different PISA formulations.

In many cases the final copolymer morphology is dictated primarily by the relative volume fractions of the two blocks, as described by the packing parameter (P). In addition to spheres, worms, and vesicles, other unusual morphologies have also been produced by PISA, such as lamellae, framboidal vesicles, spaced concentric vesicles, and yolk/shell particles. Examples of these morphologies are shown in Figure 1. Other examples of unusual morphologies produced by RAFT-mediated PISA include large compound vesicles and doughnuts. This wide range of well-defined morphologies illustrates the versatility and remarkable control afforded by PISA. However, the painstaking construction of phase diagrams is essential for the reproducible targeting of desired pure copolymer morpholo-
This rational approach has enabled the synthesis of various types of well-defined spherical nanoparticles and also low-polydispersity vesicles. Block copolymer worms are invariably well-defined in terms of their mean widths but typically exhibit a relatively broad distribution of worm lengths. Nevertheless, PISA syntheses remain the best synthetic route to produce concentrated dispersions of block copolymer worms, which is highly desirable for studying their rheological behavior. In this Perspective, the various advantages offered by RAFT-mediated PISA formulations are discussed along with some of the current drawbacks and problems associated with this platform technology.

## COMPARISON OF AQUEOUS PISA FORMULATIONS WITH CONVENTIONAL AQUEOUS EMULSION POLYMERIZATION

Conventional aqueous emulsion polymerization involves free radical polymerization rather than RAFT polymerization; it is highly efficient and can be conveniently conducted at high solids. Hence, it is worth asking whether RAFT-mediated PISA formulations offer any advantages over such a well-established, commercially successful technology. If near-monodisperse spherical particles of 100–1000 nm diameter are desired, then conventional aqueous emulsion polymerization is clearly superior to aqueous RAFT-mediated PISA formulations. Both approaches enable high monomer conversions to be achieved within 1−2 h at 60−70 °C, but significantly narrower particle size distributions and a much wider range of mean particle diameters can be produced using conventional aqueous emulsion polymerization. However, if relatively small spheres of (say) 20−50 nm diameter are desired, then aqueous PISA offers a potentially decisive advantage because it does not require high levels of added surfactant. In this context, it is worth emphasizing that aqueous emulsion polymerization formulations often suffer from excess surfactant, which is known to compromise performance and whose removal via either centrifugation or dialysis is not normally cost-effective. Moreover, if sterically stabilized particles are required, then an optimized aqueous PISA formulation usually offers high blocking efficiencies (and hence effective steric stabilization) via surfactant-free formulations. This is in striking contrast to the relatively...
low grafting efficiencies usually achieved when using either macromonomers, block copolymers, or graft copolymer stabilizers for aqueous emulsion polymerization.\textsuperscript{97−100} Possibly the most important advantage offered by RAFT aqueous emulsion polymerization over conventional aqueous emulsion polymerization is the ability to prepare diblock copolymer worms and vesicles. However, this has only been achieved for a small minority of PISA syntheses,\textsuperscript{55,79,80,101} whereas many RAFT aqueous dispersion polymerization syntheses only result in the formation of kinetically-trapped spheres, even when targeting highly asymmetric diblock compositions.\textsuperscript{35,95,96,104,105} These observations are currently not properly understood and surely warrant further work. In contrast, most RAFT dispersion polymerization formulations typically exhibit the expected range of copolymer morphologies (spheres, worms, and

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**Figure 2.** Chemical structures of various types of steric stabilizer blocks utilized for (a) RAFT-mediated aqueous dispersion polymerization and (b) RAFT-mediated aqueous emulsion polymerization.
vesicles) provided that such syntheses are conducted at relatively high solids (>20% w/w) while using a sufficiently short stabilizer block as a macromolecular chain transfer agent (macro-CTA). The latter aspect is important because a relatively long stabilizer block leads to highly effective steric stabilization immediately after micellar nucleation. This prevents efficient sphere—sphere fusion, which is the essential first step in the production of anisotropic worms.54

It should be noted that targeting a spherical morphology via PISA can offer important advantages. First, the solution viscosity is dramatically reduced for spherical nanoparticles compared to the equivalent synthesis of molecularly-dissolved block copolymer chains via solution polymerization. Additionally, the onset of micellar nucleation during the production of spherical nanoparticles is typically followed by a significantly faster rate of polymerization.69,102,106,107 This is because the unreacted monomer diffuses into the nascent nanoparticle cores in order to solvate the growing insoluble block. This leads to a higher local monomer concentration and enables aqueous PISA syntheses to be completed within 2 h at 70 °C. Similar, albeit less pronounced, rate enhancements are also observed for n-alkane PISA formulations.37 However, alcoholic PISA syntheses appear to be significantly slower, often requiring 24 h for 95% conversion85,56,57 and sometimes considerably longer.60 Moreover, in at least some cases no rate enhancement is observed to achieve such formulations.72,108 Currently, this striking difference is not understood. It is known that substantially faster polymerizations can be achieved for alcoholic RAFT PISA formulations simply by adding water as a co-solvent.61,106 This is most likely because (i) water is a non-alcoholic RAFT PISA formulation simply by adding water as substantially faster polymerizations can be achieved for syntheses appear to be significantly faster in dilute aqueous solution compared to the radical polymerization of various vinyl monomers is significantly faster in dilute aqueous solution compared to bulk polymerization.109,110 This suggests that the water co-solvent could be playing an additional role at the molecular level in PISA syntheses conducted in alcohol/water mixtures.

■ SURFACE CHEMISTRY OF BLOCK COPOLYMER NANO-OBJECTS

In principle, the nature of the steric stabilizer block should dictate the surface chemistry of the resulting block copolymer nano-objects. For example, aqueous RAFT-mediated PISA can be conducted with a wide range of steric stabilizers under both dispersion (Figure 2a) and emulsion conditions (Figure 2b). This approach enables the design of a wide range of spheres, worms, or vesicles exhibiting nonionic,54,89,111 zwitterionic,81,103,112 anionic,102 or cationic56,71,113 character in the case of RAFT aqueous dispersion polymerization. Similarly, RAFT aqueous emulsion polymerization has been used to produce non-ionic,55,114 anionic,105,115−117 or cationic spheres.118 As expected, the chemical nature of the stabilizer block directly influences the colloidal stability of the nanoparticles. Thus, choosing a zwitterionic polysulfobetaine (PSBMA) block confers enhanced salt tolerance,103 a nonionic poly(glycerol monomethacrylate) (PGMA) block enables pH-modulated selective adsorption of nanoparticles onto a micropatterned planar substrate55 and an anionic poly-(ammonium 2-sulfatoethyl methacrylate) (PSEM) block promotes efficient occlusion within ZnO host crystals.115

Since the chemical nature of the stabilizer block determines the surface wettability of the particles (or particle contact angle199), hydrophilic diblock copolymer nanoparticles can be designed to stabilize oil-in-water Pickering emulsions.35,73,120 Similarly, hydrophobic diblock copolymer nanoparticles can be designed to stabilize water-in-oil emulsions.122 By judiciously combining these two types of nanoparticles (and optimizing the homogenization conditions), Thompson and co-workers exploited PISA to produce Pickering double emulsions.122 In this case, both types of Pickering emulsifier possessed a worm-like morphology. Block copolymer worms were found to adsorb much more strongly at the oil—water interface than the more commonly employed spheres because the former nanoparticles exhibit a relatively high surface area per unit mass.120 Similarly, Mable and co-workers recently reported that frambooidal PGMA−PHPMA−PBzMA vesicles are much more efficiently adsorbed at the oil—water interface than the equivalent smooth PGMA−PHPMA vesicles.120 This model system illustrates the importance of surface roughness on Pickering emulsifier performance.

■ STIMULUS-RESPONSIVE BLOCK COPOLYMER NANO-OBJECTS

There are various literature reports describing thermoresponsive block copolymer nano-objects prepared via PISA. Typically, these syntheses are based on RAFT aqueous dispersion polymerization rather than RAFT aqueous emulsion polymerization. Presumably, this is because the less hydrophobic core-forming block is more readily plasticized by water in the former case.35 For PGMA−PHPMA block copolymer worms prepared in water, a worm-to-sphere transition can be induced simply by cooling from around 20−25 °C to 5−10 °C123,124. In contrast, for PLMA−PBzMA worms prepared via PISA in n-dodecane, Fielding et al. showed that heating up to 90 °C was required to induce the same order−order transition.56 The temperature-dependent gel moduli for the two systems are shown in Figure 3. Similar worm-to-sphere transitions were subsequently reported by Pei et al. on heating various methacryl diblock copolymer worms prepared in either n-tetradecane or ethanol.125,126 In each case, the thermally-triggered transformation of highly anisotropic worms into isotropic spheres results in degelation. This is because multiple inter-worm contacts in the initial gel are lost, which results in the formation of free-flowing dispersions.

According to Fielding et al., the switch in copolymer morphology is a direct result of surface plasticization of the core-forming block, which leads to a reduction in the effective packing parameter.60 Reasonably good thermoreversibility can be observed, provided that the copolymer concentrations are in the 5−20% w/w range. However, on returning to the original temperature, the complementary sphere-to-worm transitions are not observed at lower copolymer concentrations (< 1% w/w), presumably because the one-dimensional (1D) fusion of multiple spheres is less probable under these conditions. The concentration dependence of the (ir)reversibility of such thermal transitions warrants further attention and is the subject of ongoing research in our laboratory.

Thermoresponsive core−shell nanoparticles have also been reported by An et al.127 Chain extension of a poly(ethylene glycol) (PEG)-based macro-CTA with 2-methoxethyl acrylate (MEA), PEG methyl ether acrylate, and a small amount of PEG
diacrylate (PEGDA) cross-linker produced spherical nanogels, whose dimensions decreased almost linearly as the solution temperature was increased from 20 to 60 °C. Other examples of thermoresponsive nanogels from the same group include a PEG-based macro-CTA chain-extended with MEA and a small amount of PEGDA cross-linker, which formed well-defined spherical nanogels up to 32% solids and core–shell nanogels composed of either linear or branched PEG-based shells and methacrylic cores. Core dehydration was observed by 1H NMR on heating above 40 °C. Spectroscopy studies indicated subtle differences in hydrogen bonding between the core-forming blocks and the surrounding water molecules.

Similar thermoresponsive nanogels were also prepared by Rieger et al. In this case, PEG-based macro-CTAs were chain-extended with a mixture of N,N'-diethylacrylamide (DEAam) and N,N'-methylene bis(acrylamide) via RAFT aqueous dispersion copolymerization, with in situ cross-linking resulting in the formation of thermosensitive nanogels.

There are also recent reports of block copolymer worms prepared using a thermoresponsive stabilizer block. Monteiro and co-workers used a range of chain-end functional poly(N-isopropylacrylamide) (PNIPAM) macro-CTAs to polymerize styrene at 70 °C, well above the lower critical solution temperature (LCST) of PNIPAM, in a RAFT aqueous emulsion polymerization. On cooling to 23 °C, below the LCST, worms were formed with multifunctional groups located at the surface, allowing further coupling reactions or chemical transformations to be made. It is, however, important to note that the addition of toluene was required to plasticize the PS cores. Moreover, strictly speaking this is not an example of a conventional PISA formulation as the PNIPAM block is above its LCST during the polymerization of styrene and hence not able to act as a steric stabilizer for the PS block. Instead, colloidal stability is maintained via addition of an anionic surfactant. These thermoresponsive worms were used in combination with PNIPAM functionalized with cell-binding vitronectin protein to bridge and aggregate human embryonic stem cells (hESCs), allowing 3D cell growth and exploiting the thermoresponsive properties to allow breakdown and subsequent reformation of the hESC aggregates. Similarly, Davis et al. examined the RAFT aqueous emulsion polymerization of styrene but, in this case, used a poly(di(ethylene glycol) ethyl ether methacrylate-co-N-(2-hydroxypropyl)methacrylamide) (P(DEGMA-co-HPMAc)) stabilizer.

Cooling from 70 to 23 °C, below the cloud point temperature of the thermoresponsive stabilizing block, led to restructuring of the copolymer assemblies from spheres to worm-like nanoparticles or vesicles. However, this again required the addition of toluene to plasticize the PS cores.

pH-responsive diblock copolymer nano-objects have been prepared via PISA by Lovett et al. and Penfold et al. In the first case, a carboxylic acid-functionalized RAFT CTA was used to prepare anionic PGMA–PHPMA worms via RAFT aqueous dispersion polymerization at around pH 3. Under these conditions, the terminal carboxylic acid group is in its protonated neutral form and a soft aqueous worm gel was obtained at 10% solids. Adjusting the solution pH to 6 led to ionization of the weakly acidic end-groups, thus increasing the relative volume fraction of the stabilizer block. This subtle change was sufficient to induce a worm-to-sphere transition, which caused in situ degelation. This order–order transition proved to be reversible: a worm gel was reformed on lowering the solution pH. Because there is only one ionizable acid group per copolymer chain, relatively little acid or base is required to induce the morphological transition compared to traditional pH-responsive block copolymers. However, this subtle pH-responsive behavior is suppressed in the presence of salt because of charge screening. The second example of pH-responsive diblock copolymer nano-objects prepared by PISA is again based on PGMA–PHPMA worms, but in this case they possess tertiary amine end-groups arising from a morpholine-functionalized RAFT CTA. A worm gel prepared at pH 7–7.5 at 15% solids underwent a worm-to-sphere transition on acidification to pH 3, causing in situ degelation as the morpholine end-groups became protonated. This order–order transition is fully reversible in salt-free solutions, but in the presence of added electrolyte, the terminal cationic charge is screened, which enables the worm gels to remain intact. These complementary examples illustrate that the judicious

Figure 3. Temperature-dependent gel moduli as a result of worm-to-sphere transitions observed for (a) a PLMA16–PBzMA27 worm gel in n-dodecane at 20% w/w solids and (b) an aqueous PGMA54–PHPMA150 worm gel at 10% w/w solids. (a) Reproduced with permission from ref 66. (b) Reproduced with permission from ref 124. Copyright 2012 The Royal Society of Chemistry.
selection of an appropriate RAFT CTA can be used to confer pH-responsive behavior on ostensibly non-ionic diblock copolymers while requiring minimal amounts of added base or acid.

Another example of stimuli-responsive nano-objects prepared by PISA has been recently reported by Zetterlund et al.\textsuperscript{137} In this case the stimulus is gaseous CO\textsubscript{2}, which enables the copolymer morphology to be fine-tuned. Alcoholic RAFT dispersion polymerization of styrene from a poly(4-vinylpyridine) (P4VP) macro-CTA was used to produce P4VP–PS diblock copolymer spheres, worms, or vesicles in the absence of CO\textsubscript{2} and spheres or worms in the presence of CO\textsubscript{2}. This weakly acidic gas interacted with the basic pyridine groups on the P4VP stabilizer block, which increased the relative volume of the block and hence lowered the effective packing parameter; thus, the introduction of CO\textsubscript{2} was used to make the formation of higher order morphologies less energetically favorable. Additionally, introduction of CO\textsubscript{2} led to (i) lower chain mobility in the core, (ii) reduced solvent polarity, leading to better solvation of the solvophobic block, hence shifting morphology transitions to higher DPs, and (iii) lower effective block copolymer concentrations, shifting the copolymer morphology toward spheres. Moreover, the phase space for pure worms, which is typically rather narrow and hence somewhat elusive,\textsuperscript{34} proved to be much more readily accessible in the presence of CO\textsubscript{2}.

### CHARACTERIZATION TECHNIQUES

Much of the PISA literature has focused on core-forming blocks based on either methacrylic monomers or styrene. Such polymers have relatively high glass transition temperatures ($T_g$),\textsuperscript{138} which aids their characterization by transmission electron microscopy (TEM). In contrast, there are far fewer studies involving acrylic formulations. The low $T_g$ of acrylic polymers compared to their methacrylic counterparts results in a strong tendency toward film formation during TEM grid preparation, producing images that are not representative of the true copolymer morphology in solution.\textsuperscript{139} In this case rigorous morphological characterization requires cryo-TEM, which is a much more expensive, time-consuming, and less widely available technique. Nevertheless, it seems likely that all-acrylic PISA formulations will be explored in more detail in the near future, particularly if film-forming nano-objects offer a decisive advantage for commercial applications.

Another powerful characterization tool is small-angle X-ray scattering (SAXS). Unlike TEM, this technique enables systems to be characterized directly as dispersions. Moreover, statistically robust particle dimensions can be obtained since X-ray scattering is averaged over many millions of nano-objects. In contrast, TEM studies typically sample only a few hundred to a few thousand particles and are hence prone to sampling bias. Provided that a physically realistic model is employed, SAXS studies enable the dimensions of nano-objects to be calculated with precision. In contrast, techniques such as dynamic light scattering (DLS) merely report diffusion coefficients, which are indirectly related to the particle size via the Stokes–Einstein equation (which assumes a spherical morphology). Nevertheless, selection of an appropriate scattering model for SAXS analysis is usually informed by some prior knowledge regarding the particle size and morphology of the nano-objects. This is normally provided by imaging techniques, such as TEM. Static light scattering (SLS) has also been used to characterize diblock copolymer nano-objects prepared via PISA. This technique reports the radius of gyration ($R_g$) and the weight-average particle mass, with the latter parameter typically being used to calculate the particle aggregation number, $N_{agg}$. In two recent studies $N_{agg}$ data obtained via SLS has been compared to that calculated from SAXS analyses, with good agreement providing greater confidence in the model used for the latter technique.

Mykhaylyk and co-workers have demonstrated that SAXS can be used to characterize spheres, worms, and vesicles prepared via PISA.\textsuperscript{55,66,73,90,121,123,144,145} For example, SAXS analysis of vesicles formed during the RAFT aqueous dispersion polymerization of HPMA (Figure 4) has revealed an unexpected growth mechanism for this morphology. As the DP of the core-forming PHPMA block increases, the vesicle membrane thickens, but the outer vesicle diameter is conserved, which leads to a gradual reduction in the vesicle lumen volume.\textsuperscript{34} Geometric considerations confirm that this is actually the only mechanism by which the growing vesicles can minimize their total interfacial area and hence their free energy. Moreover, this mechanism places an important constraint on vesicle growth. As longer core-forming blocks are targeted, the vesicle morphology eventually becomes
unstable as a result of increasing steric congestion of stabilizer chains within the inner leaflet combined with greater solvent plasticization of the vesicle membrane. It is rather difficult to imagine any other single characterization technique providing such detailed mechanistic insights.

### POTENTIAL APPLICATIONS AND OPPORTUNITIES FOR PISA FORMULATIONS

There have been various reports of highly efficient “one-pot” PISA syntheses based on RAFT aqueous emulsion polymerization,\(^\text{101,104,105,107,116}\) RAFT aqueous dispersion polymerization,\(^\text{70}\) and RAFT n-alkane dispersion polymerization.\(^\text{64}\) Such advances seem to be particularly promising for potential commercial applications, which have begun to emerge as this subdiscipline has matured. In this context, there has been significant recent progress in the efficient removal of RAFT end-groups from copolymer chains using various reagents,\(^\text{146−149}\) although it remains to be seen whether such strategies are equally effective for copolymer nanoparticles (as opposed to soluble chains). While such post-polymerization modification undoubtedly adds both cost and complexity to PISA syntheses, this approach may yet be cost-effective for certain high-value biomedical applications suggested for block copolymer nanoparticles.\(^\text{123,133,50−152}\) Nevertheless, the relatively high cost, intrinsic color, and malodorous nature of the sulfur-based RAFT CTAs may well prove to be detrimental to the development of next-generation paints and coatings or even relatively “high-value” cosmetics additives. On the other hand, it is perhaps worth emphasizing that a series of RAFT-synthesized star copolymers have already been commercialized by the Lubrizol Corporation as high-performance thickeners for automotive engine oils.\(^\text{153,154}\) Thus, RAFT chemistry is commercially viable for at least some industrial sectors.

RAFT polymerization is not a prerequisite for PISA syntheses, which have also been conducted using nitroxide-mediated polymerization (NMP)\(^\text{38−40}\) and atom transfer radical polymerization (ATRP).\(^\text{41−43}\) However, although the downsides of RAFT chemistry are thereby avoided, such formulations often suffer from either incomplete conversions (NMP) or metal catalyst contamination (ATRP). One recent synthetic development is single electron transfer living radical polymerization (SET-LRP).\(^\text{155}\) SET-LRP proceeds under mild reaction conditions in various solvents\(^\text{156−160}\) and can be used for a wide range of monomers.\(^\text{156,157,161−166}\) Very recently, SET-LRP has been utilized by Cunningham and co-workers for growing either one or two poly(methyl methacrylate) (PMMA) chains from an alginate-based macroinitiator in methanol/water mixtures to produce alginate-stabilized PMMA-core spheres.\(^\text{167}\)

PISA syntheses using organotellurium-mediated living radical polymerization (TERP)\(^\text{168,169}\) have also been attempted.\(^\text{170}\) For example, tert-butyl acrylate and styrene were polymerized using a PMMA-based macroTERP agent producing triblock copolymer nanoparticles,\(^\text{171}\) although significant improvements in control over \(M_\text{w}/M_\text{n}\) and/or blocking efficiency seem to be desirable to warrant wider use of this chemistry.

In 2011, Blanazs et al.\(^\text{69}\) monitored the RAFT aqueous dispersion polymerization of HPMA using TEM for a formulation targeting vesicles as the final copolymer morphology. More specifically, an evolution in copolymer morphology from spheres to worms to vesicles was observed, with jellyfish being identified as a key intermediate structure between the latter two nano-objects. Given the “open” structure of such jellyfish (see Figure 1), one interesting question that certainly warrants exploration is whether encapsulation can be achieved in situ when targeting vesicles as the final copolymer morphology. In this context, it is worth emphasizing that small molecules are likely to permeate through such vesicle membranes rather quickly, but soluble macromolecules or nanoparticles should be retained much more efficiently. Indeed, significant progress toward this goal has just been reported by Zhang, Sumerlin, and co-workers.\(^\text{172}\) Photoinitiated RAFT polymerization of HPMA was conducted using a PEG macro-CTA in the presence of an aqueous silica sol, with TEM studies providing reasonable evidence for nanoparticle encapsulation within the resulting PEG−PHPMA vesicles. Moreover, a model globular protein (bovine serum albumin, BSA) could also be incorporated within the same vesicles, although no direct visual evidence for this encapsulated species was possible. Unusually, these PISA syntheses were conducted at 25 °C, which makes in situ protein denaturation highly unlikely. In closely related work, we have shown that the aqueous RAFT dispersion polymerization of HPMA using a PGMAs macro-CTA in the presence of silica nanoparticles enabled their in situ encapsulation inside the resulting PGMAs−PHPMA vesicles, as confirmed by TEM and SAXS studies (Figure 5).\(^\text{173}\)

![Figure 5](image-url)
was followed by thermally-triggered release of the silica payload on cooling to 0–10 °C, since this induces a vesicle-to-sphere transition. Furthermore, BSA could be encapsulated intact by conducting the polymerization at 37 °C using a low-temperature initiator, thus avoiding its denaturation. The BSA loading efficiency was determined to be 11% by fluorescence spectroscopy, although TEM suggested protein flocculation.

In the absence of any attractive interactions between the block copolymer chains and the encapsulated species, the theoretical maximum encapsulation efficiency is simply given by the ratio of the total vesicle lumen volume to the total solution volume; this suggests that the majority of the silica nanoparticles or protein molecules cannot be encapsulated within the vesicles. Nevertheless, if these two studies can be extended to include enzymes or antibodies, then this in situ encapsulation approach suggests potential biomedical applications, especially if triggered release could be achieved under biologically relevant conditions. In related work, Davis et al. reported that Nile Red could be encapsulated within poly(oligoethylene glycol methacrylate)–polystyrene (POEGMA–PS) vesicles during the one-pot RAFT alcoholic dispersion polymerization of styrene. However, the final monomer conversions were relatively low (10%). Moreover, the amount of encapsulated dye was stated to be more than the original dye concentration. This suggests that light scattering from the vesicles artificially increased the apparent absorbance of the dye, which would invalidate the encapsulation assay.

It is well-known that the target DP for a RAFT polymerization is equal to the monomer concentration divided by the macro-CTA concentration. Given that the upper limit of concentration for PISA syntheses is of the order of 50% solids, lowering the macro-CTA concentration under such conditions becomes the only available means of increasing the target DP of the core-forming block. However, the initiator concentration must be reduced accordingly, because RAFT polymerizations are typically conducted at [macro-CTA]/[initiator] molar ratios of 3 to 10 so as to minimize the proportion of dead chains while maintaining an acceptable polymerization rate. Thus, targeting very high DPs eventually leads to either gradual loss of RAFT control or no polymerization at all (because there is insufficient initiator to generate the required radical flux). Clearly, the precise upper limit DP will vary significantly depending on (i) the monomer class (e.g., methacrylate vs acrylate vs styrene), (ii) initiator type (due to the differing characteristic half-lives at a given temperature), (iii) nature of the CTA (dithiobenzoate vs trithiocarbonate vs xanthate), and (iv) the precise PISA formulation (e.g., RAFT aqueous emulsion polymerization vs RAFT aqueous dispersion polymerization).

Nevertheless, RAFT-mediated PISA formulations possess some intrinsic advantages over RAFT solution polymerization. It is well-recognized that PISA syntheses can be very efficient, with very high monomer conversions achieved within short reaction times. The marked rate acceleration that typically occurs during PISA syntheses, as discussed previously, usually coincides with the onset of micellar nucleation and, as noted by Blanazs et al., can lead to a five-fold increase in the rate of polymerization. In addition, copolymer chains are produced in the form of low-viscosity nanoparticle dispersions rather than high-viscosity solutions. Hence it seems likely that RAFT-mediated PISA should enable higher DPs to be targeted for the core-forming block within reasonable time scales. There are already several examples of PISA formulations with relatively high DP core-forming blocks. For example, we have previously reported DPs of up to 1000 for the RAFT aqueous dispersion polymerization of HPMA. In as-yet unpublished work, we have recently achieved DPs of up to 4700 when utilizing a highly polar methacrylic monomer. Similarly, Davis et al. recently reported polystyrene-core block copolymer spheres with molecular weights above 10<sup>6</sup> g mol<sup>−1</sup> (corresponding to polystyrene DPs around 14 000) via RAFT aqueous emulsion polymerization at 80 °C. High conversions (>90%) were attained in 6 h with polydispersities remaining at 1.40 or below.

Blanazs et al. reported that PGMA–PHPMA worms prepared via RAFT aqueous dispersion polymerization of HPMA can form soft, free-standing biocompatible hydrogels. Gelation is believed to be the result of multiple inter-worm contacts, rather than worm entanglements. Moreover, these worm gels proved to be thermoresponsive: cooling from 20–25 °C to around 5–10 °C induced a reversible worm-to-sphere transition, which led to in situ degelation (Figure 4b). Unlike other PHPMA-based diblock copolymers, this morphological switch is fully reversible and provides a convenient route to sterilization via cold ultrafiltration. Very recently, such worm gels have been examined as 3D matrices for human stem cell colonies, with protein assays indicating that the stem cells enter stasis (G<sup>0</sup> state) within 16 h of immersion within the worm gel. Such quiescent cells can survive for up to 2 weeks at 37 °C without passaging. On cooling to 5–10 °C, gelation occurs, and the stem cell colonies can be readily removed from the copolymer aqueous dispersion. On returning to 37 °C, the stem cells slowly emerge from stasis over a 16–24 h period while retaining their original pluripotent character. Thus, these worm gels may offer a cost-effective alternative to cryo-preservation for global stem cell transportation. Additionally, next-generation <i>thiol-functional</i> PGMA–PHPMA worm gels have been recently reported by Warren and co-workers. These are currently being evaluated as potential muco-adhesive gels, while the gel reinforcement conferred by inter-worm disulfide cross-links has just been demonstrated to be critical in the context of thermoresponsive 3D hydrogels for “cells-in-gels-in-paper” applications. Such wholly synthetic worm gels offer an interesting alternative to animal-derived products such as Matrigel.

### LIMITATIONS OF PISA FORMULATIONS

Armes and co-workers demonstrated that when targeting linear diblock copolymer chains, the worm and vesicle morphologies that can be accessed via PISA cannot tolerate the presence of surfactants. In particular, addition of ionic surfactants to colloidal stable aqueous vesicular dispersions led to rapid dissociation, producing either spheres or molecularly dissolved copolymer chains. In principle, this problem can be overcome by cross-linking the copolymer chains, either during their PISA synthesis by addition of a bifunctional monomer such as ethylene glycol dimethacrylate or by post-polymerization derivatization. This typically requires the cross-linker to be added as a third block rather than via statistical copolymerization. This is because the latter approach tends to result in a loss of colloidal stability. The former cross-linking protocol works well for spheres and vesicles and can also work for worms, although it is somewhat less reliable for this morphology. Unfortunately, such covalent cross-linking also eliminates the desirable stimulus-responsive behavior, which most likely precludes certain potential...
applications for worms (as smart thickeners) and vesicles (for encapsulation/release applications) in home and personal care applications.

Another obvious limitation of the PISA approach is for monomers that are non-solvents for the corresponding core-forming block. For example, in our early exploratory PISA syntheses we attempted to prepare diblock copolymer nanoparticles consisting of a core-forming polyacrylonitrile block in water. Acrylonitrile monomer has appreciable aqueous solubility, and its polymerization initially proceeded smoothly in homogeneous solution. However, once the critical DP for nucleation was attained (as judged by the appearance of Tyndall scattering), essentially no further polymerization occurred because the remaining unreacted monomer does not solvate the polyacrylonitrile cores. In principle, this problem could be addressed by adding a suitable co-solvent. However, this would necessarily produce a non-aqueous formulation, which would negate the various advantages conferred by using water as a polymerization medium. Fortunately, there are very few vinyl monomers that are not good solvents for their corresponding homopolymer, so this problem is rather rare in practice.

It is well-documented that reproducible PISA syntheses require the construction of detailed phase diagrams. This is particularly true if the worm phase is desired, since this morphology typically occupies rather narrow phase space. This is usually the case regardless of the PISA formulation, with strikingly similar core-forming block DP vs copolymer concentration phase diagrams being observed for RAFT dispersion polymerizations conducted in water, alcohols, or n-alkanes. Typical phase diagrams for PGMA$_{78}$−PHPMA$_x$ and PGMA$_{75}$−PHPMA$_x$ syntheses via RAFT aqueous dispersion polymerization of HPMA are shown in Figure 6. In each case, a large batch of macro-CTA is first prepared, since it is difficult to reproducibly target precise stabilizer DPs at the intermediate conversions required to ensure chain-end fidelity. Hence a single large batch of macro-CTA is used to carry out a series of subsequent multiple small-scale PISA syntheses. For most methacrylic or styrene-based PISA formulations, assignment of the final copolymer morphology at approximately full monomer conversion can be made via conventional TEM studies. Generally speaking, it is good practice to examine a partially constructed phase diagram in order to inform further PISA syntheses, with the aim being to minimize uncertainty in the positions of the various phase boundaries.

Unlike traditional block copolymer phase diagrams depicting equilibrium morphologies, the lower concentration regions of PISA phase diagrams often correspond to kinetically trapped morphologies (typically spheres). This is certainly the case for the PGMA$_{78}$−PHPMA$_x$ phase diagram shown in Figure 6a. In contrast, the PGMA$_{75}$−PHPMA$_x$ phase diagram shown in Figure 6b has little or no concentration dependence. Presumably, this is because this shorter PGMA block is a less effective steric stabilizer, which makes the multiple 1D fusion of spheres much more likely to occur on the time scale of the PISA synthesis. This is the critical event that enables access to higher order morphologies, in addition to the spheres initially formed during nucleation.

Currently, there are no PISA syntheses that provide good control over the mean worm length. This appears to be a formidable technical challenge, but recent success in the rational design of low-polydispersity vesicles suggests that there may be some scope in this regard. One possibility may be to take advantage of a thermoreversible sphere-to-worm transition and introduce an “initiator” type nanoparticle to ensure that the 1D fusion of multiple spheres occurs from a predefined number of nucleation sites. A similar concept has been recently reported by the Manners group for the construction of well-defined cylinders based on the principle of crystallization-driven self-assembly. Alternatively, Monteiro and co-workers have recently described an interesting strategy for generating tadpole-like morphologies via a so-called “temperature-directed morphological transformation”. In principle, this approach may also have some merit for the production of low-polydispersity worms.

Another important constraint lies in our current inability to use the packing parameter $P$ in order to predict final copolymer morphologies for PISA formulations. This is not particularly surprising because this simple geometric concept simply cannot accommodate the relative degrees of solvation (and hence effective volume fractions) of the stabilizer and core-forming blocks. This problem is further exacerbated because $P$ is also likely to be sensitive to an unknown degree of monomer solvation of the core-forming block once nucleation has occurred. Moreover, the packing parameter concept cannot account for the known concentration dependence of certain PISA syntheses. Furthermore, we are currently unable to explain, even qualitatively, why many (but not all) RAFT aqueous emulsion polymerization formulations lead solely to kinetically-trapped spheres. Unfortunately, even for favorable situations where equilibrium copolymer morphologies are...
produced, it is not yet possible to predict the positions of phase boundaries for PISA syntheses. Clearly, theoretical advances would be particularly welcome for enhancing our understanding of these fascinating and versatile formulations.

**SUMMARY AND PROSPECT**

For the synthesis of a wide range of block copolymer nano-objects, PISA formulations offer decisive advantages in terms of versatility, efficiency, and cost-effectiveness. This generic approach is beginning to transform the subdiscipline of “polymer colloids” while also offering a remarkably diverse range of potential commercial applications. However, in several important aspects, our fundamental understanding of various PISA formulations remains frustratingly incomplete. Nevertheless, there are now numerous literature examples whereby PISA has provided a highly convenient route to new types of block copolymer nanoparticles. This has enabled fundamental scientific advances to be made in the design of bespoke Pickering emulsifiers, nanosized vehicles for intracellular delivery of fluorescent probes, rational design of transparent dispersions, stimulus-responsive gels, nanoparticle-loaded vesicles prepared directly at high solids, model organic nanoparticles for occlusion within inorganic host crystals, nanosized lubricants for engine oils, nanoparticles for catalysis applications, efficient encapsulation of pigment particles, and sterilizable 3D hydrogels for various biomedical applications, including the preservation of human stem cells and red blood cells.

In summary, PISA is now widely recognized as an important platform technology for the design of bespoke nano-objects of controllable size, shape, and surface chemistry. This fast-maturing subdiscipline is expected to become an important component in the toolbox of the synthetic polymer chemist.

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**Notes**

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Gregory N. Smith studied chemistry at the University of Bristol (UK) and graduated with a MSci in 2007. He continued at the same institution and was awarded a PhD under the supervision of Professor Julian Eastoe in 2015. He is currently working as a research associate in synthetic polymer chemistry at the University of Sheffield (UK), investigating polymerization-induced self-assembly. His research interests concern the self-assembly and surface interactions of soft matter from small molecules to macromolecules. He has a particular interest in colloids in non-polar media, electrokinetics, and applications of small-angle scattering.

Prof. Steven P. Armes graduated from the University of Bristol (BSc 1983 and PhD 1987). After a two-year postdoc stint at Los Alamos National Laboratory in New Mexico, he accepted a lectureship at Sussex University. He was promoted to Professor in 2000 and moved to the University of Sheffield in 2004. He has published more than 540 papers (H-index = 94), received the Tilden medal from the Royal Society of Chemistry in 2013, and was elected as a fellow of the Royal Society in 2014. His current research activities are focused on polymerization-induced self-assembly (PISA).

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