Chemical Science



EDGE ARTICLE

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View Journal | View Issue



Cite this: Chem. Sci., 2018, 9, 3986

Cluster-mediated assembly enables step-growth copolymerization from binary nanoparticle mixtures with rationally designed architectures†

Xianfeng Zhang, ‡a Longfei Lv, ‡b Guanhong Wu, b Dong Yang and Angang Dong b*b

Directed co-assembly of binary nanoparticles (NPs) into one-dimensional copolymer-like chains is fascinating but challenging in the realm of material science. While many strategies have been developed to induce the polymerization of NPs, it remains a grand challenge to produce colloidal copolymers with widely tailored compositions and precisely controlled architectures. Herein we report a robust colloidal polymerization strategy, which enables the growth of sophisticated NP chains with elaborately designed structures. By quantifying NP assembly statistics and kinetics, we establish that the linear assembly of colloidal NPs, with the assistance of PbSO₄ clusters, follows a step-growth polymerization mechanism, and on the basis of this, we design and fabricate NP chains structurally analogous to random, block, and alternating copolymers, respectively. Our studies offer mechanistic insights into cluster-mediated colloidal polymerization, paving the way toward the rational synthesis of colloidal copolymers with quantitatively predicted architectures and functionalities.

Received 15th January 2018 Accepted 1st April 2018

DOI: 10.1039/c8sc00220g

Introduction

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In spite of the inherent difference between nanoparticles (NPs) and atoms, the do bear resemblance to each other in many ways, especially when NPs self-assemble into ordered structures such as superlattices.1-4 Co-assembly of two types of NPs into binary superstructures provides a low cost yet efficient way to design metamaterials with tailored properties that arise from the organization and interactions of the constituent NP components.3,4 Many self-assembly techniques, including solvent-evaporation-induced assembly at solid-2,5 or liquid-air surfaces, 6,7 have been applied to build binary NP superlattices (BNSLs) with three-dimensional (3D) or 2D structures for multiple applications.8 In comparison with their 3D and 2D counterparts, the fabrication of 1D binary superstructures, with widely tuned NP components and precisely controlled architectures, is fascinating but more complicated because of the lack of definitive directional interactions between isotropic NPs. Fortunately, the analogy between NP linear assembly and molecular polymerization has led to the concept of colloidal polymerization with NPs behaving like monomers,9 enabling

With the development of colloidal polymerization, one would naturally expect that more complicated NP chains reminiscent of copolymers are also obtainable by exploiting the concept of molecular copolymerization. Indeed, many groups have reported the growth of colloid copolymers by co-assembly of binary NPs. 16-23 Existing colloidal copolymerization methods typically rely on dipole-16 or ligand-directed assembly with multiple driving forces (e.g., electrostatic and hydrophobic interactions).17-23 For instance, Chen and co-workers have demonstrated the linear assembly of Au NPs with different sizes by adjusting the dipole moment of Au NPs with the addition of salts.16 Kumacheva and co-workers have reported the stepgrowth copolymerization of block copolymers comprising polystyrene-functionalized Au and Pt nanorods.21 Apart from the limited adjustability of NP compositions (normally noble metal NPs)16-23 and tedious surface functionalization procedures,16-23 the colloidal copolymers resulting from these methods usually suffer from low polymerization degrees¹⁶⁻²¹ and poor structure control. 16,19-21 Moreover, it is still challenging

the design and growth of chain-like superstructures (denoted as colloidal polymers) based on a molecular approach. Although the forces that govern the assembly and linkage of NP monomers are dramatically different in nature from the case of molecular polymerization, the assembly kinetics and structural features of NP chains can be described by borrowing the knowledge learned in polymer chemistry. ^{2,9,10} Over the past few years, a number of strategies have been proposed to initiate colloidal polymerization, resulting in a variety of 1D NP superstructures with tailored compositions, structures, and functionalities. ⁹⁻¹⁵

[&]quot;State Key Laboratory of Molecular Engineering of Polymers, Department of Macromolecular Science, Fudan University, Shanghai 200433, China

biChem, Shanghai Key Laboratory of Molecular Catalysis and Innovative Materials, Department of Chemistry, Fudan University, Shanghai 200433, China. E-mail: agdong@fudan.edu.cn

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c8sc00220g

 $[\]ddagger$ These authors contributed equally to this work.

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Open Access Article. Published on 02 April 2018. Downloaded on 12/10/2025 11:48:00 PM.

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to quantitatively predict the structure of the resulting colloidal copolymers. Therefore, it is highly desirable to develop generalized colloidal copolymerization strategies that enable binary NP chains with widely tailored compositions and rationally designed architectures.

In a previous study, we have shown that colloidal NPs can assemble linearly in solution with the assistance of PbSO₄ clusters.24 As has been reported previously, molecular clusters ligated with long-chain ligands such as oleylamine and octylamine themselves tend to self-assemble into lamellar mesostructures. 13,25 With the addition of NPs capped by the similar long-chain ligands, we have found that these ultrasmall clusters have the tendency to co-assemble with the added NPs, presumably due to the strong van der Waals interactions between the two components.24 This co-assembly process, driven by the minimization of the system energy, leads to the stepwise end-on attachment of NPs, resulting in polymer-like chains encapsulated by a half-cylindrical PbSO₄ cluster shell.²⁴ Although this cluster-mediated strategy has been demonstrated to be capable of inducing the linear assembly of a variety of NPs including metals, metal oxides, semiconductors,24,26 the assembly kinetics and its application to the controlled assembly of binary NP systems have yet to be established.

In this study, by quantifying NP chain growth statistics and kinetics, we establish that this cluster-mediated self-assembly process follows the modes consistent with molecular step-growth polymerization. The validation of the polymerization characteristic of cluster-mediated assembly allows us to apply the synthetic strategies developed in molecular polymerization to the growth of colloidal copolymers. High-quality NP chains with structures analogous to those of random, block, and alternating molecular copolymers, respectively, have been realized by optimizing assembly conditions. This work not only offers mechanistic insights into cluster-mediated assembly, but also opens a step-growth copolymerization route for constructing 1D sophisticated NP superstructures with quantitatively predicted architectures.

Results and discussion

Monodisperse Fe_3O_4 NPs,²⁷ capped with oleic acid, are chosen as the model monomer system to illustrate colloidal polymerization. The self-assembly of polymer chains is triggered by simply mixing Fe_3O_4 NPs with the freshly prepared, oleylamineligated PbSO₄ clusters in hexane followed by incubation under ambient conditions for a certain period of time.²⁴ To gain insights into the chain growth kinetics, we monitor the temporal evolution of NP assemblies by *ex situ* transmission electron microscopy (TEM). Fig. 1a–d show a series of TEM images captured at different time intervals based on a sample with an initial monomer concentration of $[M]_0 = 2.0 \times 10^{-7}$ M (the details for calculating Fe_3O_4 NP molar concentrations were provided in ESI†).

Starting with NP and cluster random mixtures (Fig. 1a), we observe the emergence of oligomers consisting of 2 to 6 NPs after 3 h of incubation (Fig. 1b), and only a few free monomers are observable at this stage, suggesting that most NPs have been assembled into oligomers. PbSO₄ cluster shells that encapsulate NP arrays are also discernible in TEM (Fig. 1b, inset), consistent with previous observations.24 As time elapses, these oligomers gradually evolve into longer chains (Fig. 1c) and finally grow into linear polymers consisting of a batch of closed-packed NPs after 24 h of incubation (Fig. 1d). Presumably, individual chains are grown by end-to-end coupling of NP oligomers, as many coupled chains are found to have a kinked morphology (Fig. 1c, inset and S1†). These kinked chain intermediates are expected to undergo nonlinear-to-linear conformational transition during the subsequent chain growth process, considering the nearly straight morphology exhibited by the final polymer chains. A schematic illustrating the chain growth process is given in Fig. 1e. It should be noted that chain branching is not observed during the entire assembly process. We attribute this to the encapsulation effect of the cluster shell,24 which essentially inhibits the side-on attachment of NP monomers or oligomers during incubation.

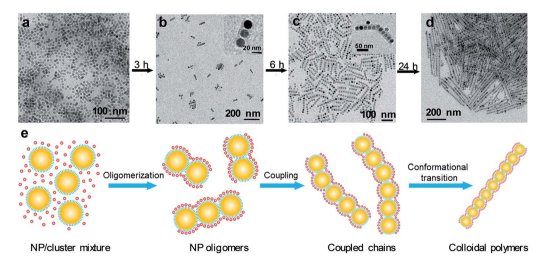


Fig. 1 (a-d) TEM images of Fe_3O_4 NP ensembles captured at different time intervals. Inset in (b) shows an oligomer consisting of three NPs. Inset in (c) shows a kinked NP chain resulting from oligomer coupling. (e) Schematic illustration of the chain growth process.

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The disappearance of NP monomers in the early assembly stage and the subsequent oligomer coupling events are the key signatures of reaction-controlled step-growth polymerization.^{9,23} In order to verify this assumption, a quantitative analysis of chain growth kinetics is desirable. For molecular stepgrowth polymerization, the reaction of monomers with two functional groups proceeds via second-order kinetics when the functional group reactivity is independent of the chain length, which can be described by:9,23

$$\bar{X}_n = 4k[M]t + 1 \tag{1}$$

where \bar{X}_n is the number-average degree of polymerization, k is the reaction rate constant, [M] is the monomer concentration, and t is the reaction time. Likewise, the reaction kinetics of colloidal polymerization can also be described using the same equation. Here, \bar{X}_n is defined by:^{9,23}

$$\bar{X}_n = \sum n_i X_i / \sum n_i \tag{2}$$

where X_i is the number of NPs constituting a single chain and n_i is the number of polymer chains containing X_i NPs. To analyze cluster-mediated assembly kinetics, four samples with different monomer concentrations were examined, and for each sample, aliquots of NP assemblies were taken out the incubation solution at specified time intervals. For each aliquot, the \bar{X}_n value, determined by statistical analysis, was plotted as a function of the assembly time. As shown in Fig. 2a, the \bar{X}_n value increases linearly with the assembly time within the explored time range regardless of the initial monomer concentrations, which is characteristic of molecular step-growth polymerization.^{9,23} Also evident from Fig. 2a is that the chain growth rate, $d\bar{X}_n/dt$, increases monotonically with the increased monomer concentration, which is also consistent with step-growth polymerization. 9,21,23 Plotting the chain growth rate as a function of the monomer concentration yielded a straight line after fitting (Fig. 2b), from which the reaction rate constant k was determined to be $6.5 \times 10^2 \, \text{M}^{-1} \, \text{s}^{-1}$.

It is noteworthy that the polymerization rate constant can be modulated by varying the amount of the free ligands existing in the incubation solution. It has been found that chain growth can be accomplished within 30 min when a small amount of

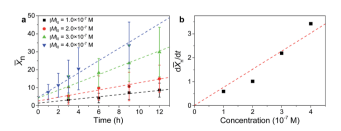


Fig. 2 (a) Dependence of the degree of polymerization (\bar{X}_n) on assembly time at different monomer concentrations. For each data point, about 200 chains were counted for statistical analysis. Error bars indicate the standard deviation. The dashed lines are the respective linear fit to the experimental data. (b) Dependence of the polymerization rate $(d\bar{X}_n/dt)$ on the monomer concentration. The dashed line is the linear fit to the experimental data

oleylamine is introduced prior to self-assembly.²³ The increased polymerization rate constant could be attributed to the stronger interactions between PbSO₄ clusters and Fe₃O₄ NPs brought by the added oleylamine ligands, which facilitate the co-assembly of the two components. We also note that the linear assembly of other types of NPs (i.e., metals and semiconductors) should also follow the modes for step-growth polymerization, as this cluster-mediated assembly process, which is independent of NP composition, size, and shape, is mainly dictated by the van der Waals interactions between the ligands (i.e., oleylamine and oleic acid) stabilizing PbSO4 clusters and NPs.24

Having established the step-growth polymerization characteristic of cluster-mediated assembly, we proceed to the fabricolloidal copolymers. cation of sophisticated microstructure of a molecular copolymer comprising monomers i and j can be characterized by a microheterogeneity coefficient, K, which is defined by:21,28

$$K = P_{ii}/(P_{ii} + 2P_{ii}) + P_{ii}/(P_{ij} + 2P_{ji})$$
(3)

where P_{ii} , P_{ii} , and P_{ii} are the fractions of the pairwise monomer units i-j, i-i, and j-j, respectively. For molecular copolymers, a Kvalue corresponding to block, random, and alternating copolymers is 0, 1, and 2, respectively. 21,28 In order to test the feasibility of cluster-mediated assembly for growing copolymers, we begin with the fabrication of random copolymers by simply coassembling two NP monomers, namely 16 nm Fe3O4 NPs and 15 nm CsPbBr₃ nanocubes, with the assistance of PbSO₄ clusters, as illustrated in Fig. 3a. Fig. 3b and c show the TEM and the corresponding high-angle annular dark-field scanning TEM (HAADF-STEM) images of a typical chain resulting from coassembly, in which the spherical and cubic NP monomers can

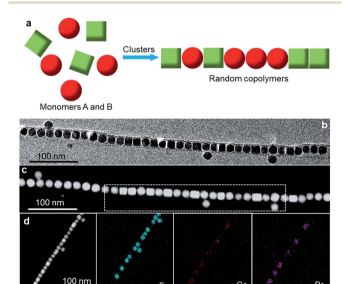


Fig. 3 (a) Illustration of the synthesis of random copolymers. For clarity, the NP surface-coating ligands and clusters were omitted. (b) TEM and (c) the corresponding HAADF-STEM images of a hybrid chain self-assembled from 16 nm Fe₃O₄ NPs and 15 nm CsPbBr₃ nanocubes. (d) HAADF-STEM image and the corresponding elemental mapping of the segment indicated by the rectangle in (c).

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be clearly distinguished. Energy-dispersive X-ray spectroscopy (EDS) elemental mapping provides further evidence for the formation of a hybrid chain (Fig. 3d), where the disordered monomer arrangement is indicative of a structure analogous to random copolymers. Statistical analysis gave rise to a K value of \sim 0.92, further confirming the random copolymer-like structure. The formation of random copolymers by co-assembly of binary NPs is not surprising, given the generality of this clustermediated assembly method.24

We next seek to fabricate colloidal block copolymers, which possess greater structural regularity compared with their random counterparts. Considering the resemblance between NP assembly and molecular polymerization, we expect that the prepolymerization technique,21 which is conventionally used for molecular block copolymer synthesis, may also be applicable to the growth of colloidal block copolymers. To test this hypothesis, we first investigate the feasibility of the "one-prepolymer" strategy,21 in which one NP monomer is assembled to form prepolymers followed by the addition of the second monomer. Specifically, 16 nm Fe₃O₄ NPs were selected to prepare prepolymers with desired lengths by incubation for a certain period of time. After that, 15 nm CsPbBr₃ nanocubes and PbSO₄ clusters were added into the incubation solution containing the asprepared Fe₃O₄ prepolymers to initiate copolymerization, as illustrated in Fig. 4a. The morphology of the resulting Fe₃O₄-CsPbBr₃ copolymers was examined with TEM, HAADF-STEM, and EDS elemental mapping.

Fig. 4b shows a low-magnification TEM image of the resulting Fe₃O₄-CsPbBr₃ block copolymers, in which the intrachain segments consisting of Fe₃O₄ NPs and CsPbBr₃ nanocubes were highlighted in red and yellow, respectively. Impressively, many chains are several hundreds of nanometers long and display a structure reminiscent of AB diblock or ABA triblock copolymers. Note that similar to molecular copolymerization, the homopolymerization of Fe₃O₄ oligomers or CsPbBr₃ monomers cannot be completely inhibited in this "one-prepolymer" approach. Despite the inherent challenge to achieve a high degree of structural homogeneity, statistical analysis shows that di- and tri-block copolymers constitute the majority (~65%) of the species (Fig. 4c). The microheterogeneity coefficient K was determined to be \sim 0.1, thus corroborating the high-yield synthesis of block copolymers. Magnified TEM on a single diblock chain reveals the seamless connection between the two segments (Fig. 4d), while HAADF-STEM and the corresponding elemental mapping show distinct spatial distribution of Fe and Cs/Br segments (Fig. 4e), providing further evidence for the formation of a perfect block copolymer-like structure.

Such Fe₃O₄-CsPbBr₃ block copolymers are highly luminescent under UV excitation (Fig. 4b, inset), with a photoluminescence (PL) spectrum similar to that of CsPbBr₃ monomers (Fig. 4f). The PL quantum yield is in the range of 45-50%, lower than that (~65%) of the original CsPbBr₃ monomers. As PbSO₄ clusters have been demonstrated to have a negligible influence on the luminescent properties of CsPbBr₃ NPs,26 the decreased fluorescence might be caused by the reabsorption effect or energy transfer processes between two NP monomers. The long tail at longer wavelengths in the extinction

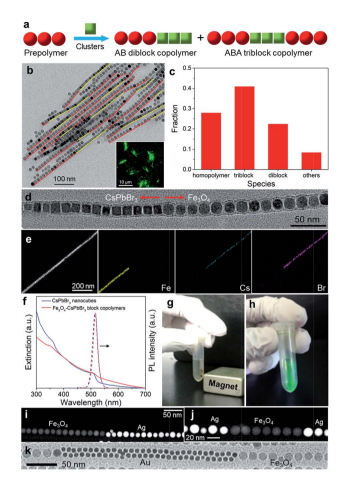


Fig. 4 (a) Illustration of the synthesis of block copolymers by the "oneprepolymer" approach. (b) Low-magnification TEM image of Fe₃O₄-CsPbBr₃ block copolymers. For each copolymer chain, the Fe₃O₄ and CsPbBr₃ segments were highlighted in red and yellow, respectively. Inset shows a fluorescence image of Fe₃O₄-CsPbBr₃ block copolymers under UV irradiation. (c) Distribution of the species resulting from copolymerization. About 500 chains were counted for statistical analysis. Others refer to multiblock chains. (d) Magnified TEM image of a single Fe_3O_4 -CsPbBr $_3$ diblock chain. (e) HAADF-STEM image and the corresponding elemental mapping of another diblock chain. (f) Extinction and PL (dashed) spectra of Fe₃O₄-CsPbBr₃ block copolymers and CsPbBr₃ nanocubes. (g) Photograph showing the attraction of Fe₃O₄-CsPbBr₃ block copolymers by a magnet. (h) Photograph of the dried Fe₃O₄-CsPbBr₃ block copolymers under UV irradiation. (i and j) HAADF-STEM images of di- and triblock copolymers of Fe₃O₄-Ag, respectively. (k) TEM image of Fe_3O_4 -Au block copolymers.

spectrum (Fig. 4f) could be caused by light scattering arising from the long polymer chains. In addition to the intriguing optical properties, these Fe₃O₄-CsPbBr₃ block copolymers also display strong magnetic response, such that they can be readily separated from the reaction solution by a magnet (Fig. 4g). Notably, the bright green emission was well retained after separation (Fig. 4h). As isolated Fe₃O₄ NPs are known to respond weakly to a magnet, the greater response of Fe₃O₄-CsPbBr₃ block copolymers should be attributed to the collective dipolar interactions of close-packed Fe₃O₄ NP segments.

Importantly, this "one-prepolymer" strategy can be readily extended to the growth of colloidal block copolymers of diverse compositions by simply selecting desired NP monomers for **Chemical Science Edge Article**

copolymerization. For instance, Fig. 4i and j show the HAADF-STEM images of Fe₃O₄-Ag di- and triblock copolymers, respectively, which are obtained by co-polymerization of 16 nm Ag monomers with Fe_3O_4 prepolymers. Likewise. polymerization of 8 nm Au monomers with the same Fe₃O₄ prepolymers also leads to block copolymers (Fig. 4k). Interestingly, unlike the aforementioned Fe₃O₄-CsPbBr₃ and Fe₃O₄-Ag copolymers, the Au segments in Fe₃O₄-Au copolymers are composed of double lines of closed-packed Au NPs, probably due to the size-matching effect exerted by the cluster shell, that is, the effective width of two lines of small Au NPs is comparable to that of a single line of larger Fe₃O₄ NPs.

In addition to the "one-prepolymer" strategy, we have also attempted to grow colloidal block copolymers by the "two-prepolymer" approach.²¹ In this approach, block copolymers are realized by a coupling reaction between two independently prepared prepolymers (Fig. S2a†). A typical example was shown in Fig. S2b and c,† where Fe₃O₄-Ag block copolymers were obtained by incubating 16 nm Fe₃O₄ and 16 nm Ag prepolymers in the presence of PbSO₄ clusters. Although block copolymers can be grown by this approach, most prepolymers remain uncoupled after incubation. The low-yield synthesis could be attributed to the relatively low reactivity between distinct NP chains.

It is interesting to note that in addition to the stepwise approaches, colloidal block copolymers can also be achieved in high yield by "one-pot" synthesis by exploiting the intrinsic selfassembly behaviors of colloidal NPs. It is found that coassembling two types of NPs of distinct sizes often leads to block copolymer-like chains rather than random copolymers, as illustrated in Fig. 5a. Fig. 5b shows a typical TEM image of block copolymers obtained by incubating 16 nm Fe₃O₄ and 8 nm Au NPs in the presence of PbSO₄ clusters, following the procedure used for growing random copolymers. Similar to the Fe₃O₄-Au block copolymers synthesized by the "one-prepolymer" approach (Fig. 4k), the copolymers grown by this approach also contain Au-blocks composed of two lines of Au NPs. The

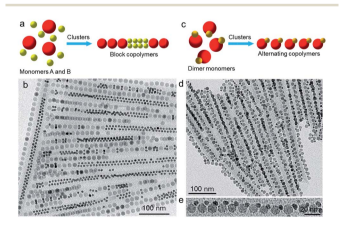


Fig. 5 (a) Illustration of the synthesis of block copolymers by coassembly of binary NPs of distinct sizes. (b) Typical TEM image of Fe₃O₄-Au block copolymers resulting from direct co-assembly. (c) Illustration of the synthesis of alternating copolymers by the selfassembly of NP heterodimers. (d) TEM image of alternating copolymer-like assemblies made from Ag-Fe₃O₄ NP heterodimers. (e) Magnified TEM image of a perfect alternating chain segment.

evolution of block copolymer-like structures under this situation is attributed to a depletion-induced phase separation process,29 where the larger Fe₃O₄ NPs tend to assemble while excluding the smaller Au NPs. Co-assembly of 16 nm Fe₃O₄ and 6 nm CoFe₂O₄ NPs under similar conditions also leads to block copolymers (Fig. S3†), further demonstrating the generality of the approach.

Finally, we extend this cluster-mediated assembly method to the fabrication of alternating copolymers. Compared with random and block copolymers, alternating copolymers are more difficult to achieve due to the challenge to inhibit homopolymerization, and to our knowledge, the assembly of binary alternating NP chains has been rarely explored. 17,18 One conceptually straightforward, albeit experimentally challenging, way to achieve alternating copolymers is to engineer the structure of NP building blocks. Fortunately, recent advances in colloidal synthesis allow us to synthesize a wide range of NPs, among which heterogeneous dimers30 could be a promising candidate for building NP chains structurally analogous to alternating copolymers by exploiting shapedirected assembly (Fig. 5c). Fig. 5d shows a low-magnification TEM image of such an example, where the chains, selfassembled from Ag-Fe₃O₄ NP heterodimers (Fig. S4†),³¹ show an alternating copolymer-like structure (Fig. 5e). We should note that these NP chains are conceptually different from real alternating copolymers, as there is only one actual monomer. Nonetheless, the microheterogeneity coefficient K was determined to be \sim 1.77, only slightly deviating from the value (i.e., 2) expected for an ideal alternating structure, confirming the formation of an alternate copolymer-like structure. Presumably, the small K value is caused by the presence of a fraction of imperfect alternating chains. Further modification of the dimer structure by tuning the size and/or shape of the two components could help to improve the quality of alternating copolymer-like chains.

The inclined alignment of Ag-Fe₃O₄ dimers in chains can be explained by the templating effect of PbSO₄ cluster shells. As we and others have demonstrated previously, 24,26 the halfcylindrical PbSO₄ cluster shells, having a width dimension of \sim 30–40 nm, behave like a self-template to confine the assembly of NPs. Our Ag-Fe₃O₄ dimers have an overall size of ~28 nm (considering the presence of organic ligands), close to the width of the cluster shells. As a consequence, Ag-Fe₃O₄ dimers will tend to assemble with certain inclination within PbSO4 cluster shells. In this context, the close packing of Ag-Fe₃O₄ dimers can proceed via either AB-AB or AB-BA modes (Fig. S5†), yet we mainly observe the AB-AB configuration in the resulting assemblies (Fig. 5e). We suspect that the preferential AB-AB alignment is energetically more favourable compared with the AB-BA mode. However, further studies are needed to elucidate the underlying mechanism.

Conclusions

In summary, we have verified a mechanism of step-growth polymerization for the 1D cluster-mediated NP assembly process. Based on guidelines in macromolecular chemistry, **Edge Article Chemical Science**

colloidal copolymers made up of binary NP mixtures with structures reminiscent of those of random, block, and alternating copolymers, respectively, have been designed and fabricated. Apart from the robust process, the generality of this copolymerization strategy allows the ready tuning of NP combinations. This provides unique possibilities for tailoring the properties of these multifunctional colloidal copolymers. This work not only offers mechanistic insights into clustermediated colloidal polymerization, but also paves the way toward the synthesis of colloidal copolymers with quantitatively predicted architectures and functionalities.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

A. D. acknowledges the financial support from MOST (2017YFA0207303, 2014CB845602), Key Basic Research Program of Science and Technology Commission of Shanghai Municipality (17JC1400100), Shanghai International Science and Technology Cooperation Project (15520720100), and NSFC (21733003). D. Y. thanks the financial support from NSFC (51573030, 51573028, 51773042).

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