Sequence Patterning, Morphology, and Dispersity in Single-chain Nanoparticles: Insights from Simulation and Machine Learning

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Abstract

Single-chain nanoparticles are intriguing materials inspired by proteins that consist of a single precursor polymer chain that has collapsed into a stable structure. In many prospective applications, such as catalysis, the utility of a single-chain nanoparticle will intricately depend on the formation of a mostly specific structure or morphology. However, it is not generally well understood how to reliably control the morphology of single-chain nanoparticles. To address this knowledge gap, we simulate the formation of 7,680 distinct single-chain nanoparticles from precursor chains that span a wide range of, in principle, tunable patterning characteristics of cross-linking moieties. Using a combination of molecular simulation and machine learning analyses, we show how the overall fraction of functionalization and blockiness of cross-linking moieties biases the formation of certain local and global morphological characteristics. Importantly, using the total volume of simulation data, we illustrate and quantify the dispersity of morphologies that arise due to both the stochastic nature of collapse from a well-defined sequence as well as from the ensemble of sequences that correspond to a given
specification of precursor parameters. Moreover, we also examine the efficacy of precise sequence control in achieving morphological outcomes in different regimes of precursor parameters. Overall, this work critically assesses how precursor chains might be feasibly tailored to achieve given SCNP morphologies and provides a platform to pursue future sequence-based design.

1 Introduction

Single-chain nanoparticles (SCNPs) are an intriguing class of materials obtained by collapsing or folding a polymer chain into a stable nanostructure. The formation and stability of an SCNP is driven by intra-chain interactions, which are characteristically non-covalent (e.g., hydrophobic, electrostatic, polar), covalent, or dynamic covalent. Inspired by proteins, SCNPs have prospective applications in catalysis, nanomedicine, and biosensing. Ultimately, the functionality and utility of an SCNP depends on its morphology, which may determine factors such as hydrophobic character nearby catalytic active sites. Consequently, fundamental study regarding the structures formed by single polymer chains has implications for the technological advancement of SCNPs and understanding of single-chain polymer physics.

Significant progress has been made in both developing facile chemical pathways for synthesis of possible precursors (i.e., the initial, unfolded polymer chains) and the characterization of resultant morphologies for SCNPs. Typically, precursors are obtained by first synthesizing a polymer with specific functional groups and then performing post-polymerization reactions at dilute conditions to promote structure-formation. Experimental scattering techniques (e.g., small-angle X-ray and neutron scattering, dynamic light scattering, etc.) have shown that the conformational behavior of SCNPs ranges between a self-avoiding random walk ($\nu \approx 0.56$) and globular ($\nu \approx 1/3$) behavior, depending on the nature of the linking chemistry. Reaction conditions may also be varied or advanced synthetic techniques employed to achieve more precise monomer and functional-group placement and bias forma-
tion of target structures in SCNP.\textsuperscript{21–25} Substantial structural dispersity, though, can exist amongst SCNP that formed from identical precursors in even polymers with relatively small numbers of chemical moieties.\textsuperscript{26} Knowledge relating a precursor to structure formation (and its reliability) in resulting SCNP will be crucial in achieving target functional properties of these materials.

Molecular dynamics (MD) simulation provides a useful tool to examine the formation of SCNPs under well-controlled conditions and with exquisite structural resolution. Prior MD studies have supported experimental observations while also generating insight regarding both the detailed nanoscopic structure of SCNPs and the mechanisms by which they might form.\textsuperscript{11,15,16,20,27–30} For example, good solvent conditions limit contact between distal portions of the polymer chain, thereby promoting intra-chain linkages over shorter backbone contour distances and resulting in pearl-necklace-like structures.\textsuperscript{16,20,28,29} MD simulations have also highlighted new strategies, such as the use of ring-polymer architectures for the precursor or manipulation of solvent quality\textsuperscript{29} to bias formation of more compact, globular structures,\textsuperscript{11} which have been experimentally challenging to achieve. Nevertheless, most simulations probe either random or regular sequence patterning of cross-linking moieties on SCNP structure formation.\textsuperscript{29} Moreover, although structural dispersity is often noted in simulations of SCNP formation, it is rarely characterized, and its implications in technological applications is largely unexplored. Thus, a comprehensive view of the impact of sequence patterning of cross-linking moieties, a thorough assessment of structural dispersity in SCNP, and examination of their interplay, is needed.

Exploring and characterizing the structure-function landscape of polymeric materials is generally non-trivial given the multitude of behaviors enabled by a large chemical and architectural space.\textsuperscript{31–35} To contend with this challenge, machine learning (ML) techniques have been increasingly utilized to probe and understand structure-function relationships in soft materials.\textsuperscript{36–44} In the context of single polymer chains,\textsuperscript{45–47} supervised ML models have been proven effective at relating polymer chain characteristics to average conformational behav-
ior, thereby expediting targeted sequence- and composition-based design tasks. Meanwhile, unsupervised ML algorithms have usefully discriminated amongst morphological structures formed in many-chain soft materials assembly by non-covalent and supramolecular interactions.\textsuperscript{48,49} Collective variables obtained from unsupervised ML can also form the basis for predicting and designing morphology.\textsuperscript{50} Overall, these works illustrate the promise of ML to help understand the distribution of morphologies that are accessible by a given precursor and to reveal potential strategies to control that distribution.

In this study, we use molecular simulation in tandem with machine learning analyses to assess how the patterning of cross-linking moieties on precursor chains impacts the morphology and properties of SCNPs. To comprehensively probe the sequence-structure space, we simulate the formation of 7,680 SCNPs from precursors chains that are predominantly distinguished by the fraction and distribution of reactive cross-linking groups along the backbone. These simulations are analyzed to characterize resulting SCNPs via structural analysis and unsupervised manifold learning. These analyses not only reveal the landscape of possible SCNP morphologies but also the dispersity of structures arising from given precursor specifications, which has not been previously well-characterized. We further examine the variability in morphological outcomes for different sequences with the same overall precursor specifications. In aggregate, this work provides a critical assessment of how precursors might be feasibly tailored to achieve given SCNP morphologies and a platform to pursue SCNP design with specific consideration of structural fidelity.
Figure 1: Overview of model and precursor chain parameters. (a) A simulation snapshot of a precursor chain. Annotations highlight the presence of backbone beads (pink) and linker beads (teal). Single-chain nanoparticles are formed by allowing the linker beads to react based on specific geometric criteria. (b) A schematic depiction of the precursor parameters blockiness $b$. (c) Visualization of the precursor chain parameter space. Ten distinct precursor chains are generated for each $(b, f)$ pair indicated by the markers. The underlying shaded region visualizes the accessible space subject to the bound on $b$ imposed by $f$. (d) Visualization of the precursor chain parameter space in the normalized blockiness $\beta$-$f$ plane. For a given $f$, points are uniformly distributed with respect to $\beta$. In panels (c) and (d), the markers are colored according to $f$ to illustrate the mapping from $b$ to $\beta$.

2 Methods

2.1 Systems

2.1.1 Model of Precursor Chains and Single-chain Nanoparticles

We adopt a coarse-grained phenomenological model to study the formation of SCNPs from precursor polymer chains. Within the model, precursor chains are comprised of backbone beads as well as pendant beads that can function as cross-linking moieties (Fig. 1a); these pendant beads are colloquially referenced as “linkers.” SCNPs are thus obtained from precursor chains by allowing linkers to react and form bonds. A modified Kremer-Grest model\textsuperscript{51} describes the interactions amongst beads in both precursor chains and SCNPs. The standard elements include (i) excluded-volume interactions characteristic of good-solvent conditions (captured by a Weeks-Chandler-Andersen pair potential) and (ii) covalent-bond interactions (captured by the typical finite extensible nonlinear elastic potential function with standard
parameters); good-solvent conditions are often experimentally used to avoid intermolecular linkages between different precursor molecules.\textsuperscript{2} The modified components include (i) the introduction of chain semiflexibility, which is common,\textsuperscript{52,53} and (ii) the reactivity of linker beads, which has been handled similarly in prior studies of SCNPs.\textsuperscript{11,16,20,28} All beads (backbone and linker) are of the same characteristic size $\sigma$, the repulsive interaction from the Weeks-Chandler-Andersen pair potential is set by $\varepsilon = k_b T$, and all bonds are treated equivalently, irrespective of whether they are between backbone-backbone, backbone-linker, or linker-linker beads. Interaction parameters are kept uniform in this study to emphasize the role of patterning of cross-linking moieties, although manipulation of monomer chemistry and sequence effects to bias structure-formation\textsuperscript{45} may be of future interest.

Chain semiflexibility is introduced via 1-3 bending interactions of the form

$$E(\theta_{ijk}) = K(1 - \cos \theta_{ijk}),$$  \hspace{1cm} (1)

for which $\theta_{ijk}$ is the angle formed by two consecutive bonds between polymer backbone beads ($i$ with $j$ and $j$ with $k$) and $K$ controls the local bending rigidity, which we set to $K = 5k_b T$. Future phenomenological models may benefit from additional bending and torsional interactions that include the linker beads, but these are not included in the present work.

Reactions between linkers to form SCNPs are handled in the following manner. First, reactions between two linkers results in the formation of a covalent bond; this bond is modeled equivalently as with any other covalent bond in the model. Second, reactions between two linkers can and will proceed only if certain geometric criteria are satisfied. In particular, the distance between two linkers must be within a “capture” radius of $1.3\sigma$, and the angle formed by bond vectors adjoining the linker and their respective backbone bead must be $180 \pm 30^\circ$. Third, reactions may not proceed between linkers on adjacent backbone beads. Fourth, reactions are irreversible (i.e., covalent bonds are preserved for the duration of the subsequent simulation). Fifth, a linker may only participate in one reaction (i.e., its valency
is one and saturates upon formation of a covalent bond with another linker). With respect to prior literature on modeling SCNPs, the utilization of a capture radius is common and follows work from Moreno and collaborators, whereas the addition of an angle-based criterion is not ubiquitous but has been employed in atomistic simulations. Here, this constraint and the null reactivity amongst topologically adjacent linkers is motivated by considering steric and electronic factors of conventional cross-linking chemical moieties. Without this addition, we find that linkers on adjacent backbone beads can undergo facile reaction, which seems generally unphysical given the disposition of cross-linking agents.

### 2.1.2 Precursor Chain Parameters

Many facets of precursor chains (chain length, chemistry, patterning, architecture, etc.) might be feasibly manipulated to control SCNP morphology. Here, only particular parameters of precursor chains are varied while others are common to all precursor chains. Specifically, all precursors consist of \( N_{bb} = 400 \) backbone beads arranged in a purely linear topology; each backbone bead may be considered as a monomer or constitutional unit. We choose this number of constitutional units inspired by Verso et al.\(^{29}\) who noted apparent differences in structures formed by precursors with random sequence patterning versus those formed from regular sequences at this length. In principle, the number of backbone units is another physical parameter that can be tuned and experimentally realized by controlling the extent of polymerization. The primary variable parameters herein are (i) the extent of functionalization or linker fraction \( f = N_l/N_{bb} \) and (ii) a descriptor \( b \), which relates to the “blockiness” of linker beads as distributed along the polymer chain (Fig. 1b). The parameter \( b \) is given by

\[
b = \frac{1}{N_{bb}} \left( 1 + \sum_{k=0}^{N_{bb}-2} 1_{k,k+1} \right),
\]  

(2)
where \( k \) and \( k + 1 \) are indices of backbone beads along the polymer chain (\( \in [0, N_{bb} - 1] \)) and \( 1_{i,j} \) is an indicator function, such that

\[
1_{i,j} = \begin{cases} 
1, & \text{if } i, j \text{ both bonded to linker} \\
1, & \text{if } i, j \text{ both not bonded to linker} \\
0, & \text{otherwise}
\end{cases}
\]  

(3)

In this fashion, a precursor sequence with backbone beads perfectly alternating between having and not having a linker would yield \( b = N_{bb}^{-1} \), while a sequence with all backbone beads possessing linkers followed by all backbone beads without linkers would yield \( b = 1 \). This blockiness parameter bears some similarity to but is not equivalent to the correlation parameter used to describe the statistics of random copolymers.\(^{54,55}\)

An overview of precursor chain parameter combinations examined in this study is shown in Fig. 1c. Notably, Eq. (2) implicitly depends on the linker fraction \( f \), such that the minimum accessible \( b \) varies with \( f \) (i.e., \( \min b(f) \approx |2(f - 0.5)| \)). Consequently, precursor chains are studied for \( f \in \{0.10, 0.15, 0.2, 0.3, 0.4, 0.6, 0.8\} \), and values of \( b \) at each composition are chosen such that \( \beta \in \{0.2, 0.4, 0.6, 0.8\} \) where

\[
\beta(f) = \frac{b(f) - \min b(f)}{1 - \min b(f)}
\]  

(4)

is a normalized blockiness parameter that can be sampled uniformly irrespective of composition (Fig. 1d). We emphasized study at lower \( f \) with the expectation that heterogeneity in SCNP morphologies would diminish at higher \( f \).

Based on prior work,\(^{40,46}\) we hypothesized that these parameters would enable study of a vast set of precursor chains that would result in distinct SCNPs. Although a precise connection of \( f \) and \( \beta \) to experimental synthetic conditions may be non-trivial, we expect that such parameters would meaningfully relate to aspects of monomer concentration and reactivity ratios.\(^{56,57}\) In the long term, advances in sequence-level polymerization may enable
more precise definitions.21–25,58

2.1.3 Precursor Chain Sequence Generation

Ten unique sequences are generated for each combination of \((f_i, \beta_j)\) to assess dispersity of SCNP morphologies as a function of precursor parameters. Thus, in total, 320 unique precursor chains are considered. To generate a given precursor chain sequence at a specified \((f_i, \beta_j)\), a fixed number of linkers (set by \(f_i\)) are first randomly distributed across the polymer backbone, and the resulting initial normalized blockiness \(\beta_j^{(0)}\) is computed. Supposing that \(\beta_j^{(0)} \neq \beta_j\), a random pair of backbone beads (one with a linker and one without) is selected, and the linker position is swapped if it would yield \(|\beta_j^{(1)} - \beta_j| < |\beta_j^{(0)} - \beta_j|\). This process is repeated until the trial \(|\beta_j^{(k)} - \beta_j| < 0.001\). All sequences generated by this process can be found in the supplementary information.

2.1.4 Simulation

All MD simulations are performed using a modified version of the 3 Mar 2020 distribution of the LAMMPS simulation package.59 Simulations are performed in reduced units with characteristic quantities of \(m\), \(\sigma\), and \(\varepsilon\) for mass, distance, and energy, respectively; the reduced time unit is \(\tau = (m\sigma^2/\varepsilon)^{1/2}\). Simulations correspond to a single polymer chain in implicit solvent. There are no periodic boundary conditions to prevent self-interaction. The simulations therefore correspond to SCNPs forming at infinite dilution. Additional details regarding general simulation protocols, precursor preparation, precursor equilibration, SCNP formation, and SCNP simulation are provided in the Supporting Information.

2.2 Analysis

2.2.1 SCNP Structural Descriptors

Topological descriptors of the SCNP are assessed via its representation as a molecular graph\(^{60,61}\) \(G = (\mathcal{V}, \mathcal{E})\). Here, the beads of the SCNP comprise the set of vertices \(\mathcal{V} = \)
(v_1, \ldots, v_{N_{bb}}, v_{N_{bb}+1}, \ldots, v_{N_{bb}+N_i}) and bonds amongst beads comprise the set of undirected edges, or bonds between beads, \( \mathcal{E} \); coordinates of beads are not used for this analysis. Using this framework, we compute two topological descriptors: the number of free backbone beads \( n_f \) and the number of topological domains \( n_d \); both of these descriptors have been previously examined in simulation studies to assess SCNP structure.\textsuperscript{28,62} For their computation, we adopt the labeling algorithm of Moreno et al. reported in Ref. 62. In brief, for any pair of bonded linkers \( v_m, v_n \), the backbone vertices along the directed contour path comprise a set \( \mathcal{D}_{m,n} \); if there are \( n_b \) linker-linker bonds, then there will be \( n_{bb} \) such sets. Then, a domain \( \mathcal{D} \) is defined as the union of sets, for which every set has non-empty intersection with at least one other set in \( \mathcal{D} \). Backbone beads that are not found in any path set (i.e., the complement of the set of all backbone beads with the union of all domains) constitute a set of free segments \( \mathcal{F} \), such that \( n_f = |\mathcal{F}| \) is the cardinality (number of members) of the free-segment set.

SCNP structures are also characterized by shape descriptors derived from its gyration tensor:

\[
\mathbf{S} = \frac{1}{N_{bb}} \sum_{i=1}^{N_{bb}} (\mathbf{r}_i - \mathbf{r}_{cm})(\mathbf{r}_i - \mathbf{r}_{cm})^T, \tag{5}
\]

where \( \mathbf{r}_i \) is a column vector of the position of the \( i \)th backbone bead, \( \mathbf{r}_{cm} \) is the center-of-mass position of the SCNP, and \( T \) denotes the transpose. Subsequent diagonalization of Eq. (5) yields \( \mathbf{S} = \text{diag}(\lambda_1^2, \lambda_2^2, \lambda_3^2) \), where the diagonal elements are the principal moments of the gyration tensor such that \( \lambda_1^2 \leq \lambda_2^2 \leq \lambda_3^2 \). From these, conventional shape descriptors\textsuperscript{63} are computed, including the radius of gyration

\[
R_g = \sqrt{\lambda_1^2 + \lambda_2^2 + \lambda_3^2}, \tag{6}
\]

which indicates the overall size of the SCNP; the acylindricity

\[
c = \lambda_2^2 - \lambda_1^2 \tag{7}
\]
which measures departure from cylindrical symmetry with respect to the eigenvectors of $\lambda_2$ and $\lambda_1$; the asphericity

$$b = \lambda_3^2 - \frac{1}{2}(\lambda_1^2 + \lambda_2^2),$$

which measures the overall symmetry of the distribution of beads with respect to three coordinate axes; and the relative shape anisotropy

$$\kappa^2 = \frac{3}{2} \frac{\lambda_1^4 + \lambda_2^4 + \lambda_3^4}{(\lambda_1^2 + \lambda_2^2 + \lambda_3^2)^2} - \frac{1}{2} \in [0, 1],$$

which assesses departure from spherical symmetric morphologies ($\kappa^2 = 0$). Notably, both $b$ and $\kappa^2$ will yield zero for spherically symmetric configurations, while $b$ will yield zero for any Platonic solid. Ensemble averages of the shape parameters are computed using the final $10^7$ timesteps of the production trajectory.

### 2.2.2 Unsupervised Machine Learning

SCNP morphologies are also identified, distinguished, and compared with the aid of unsupervised ML. The essence of the approach is to discriminate between SCNP morphologies in a data-driven fashion on the basis of collective differences in the local environments of the composite backbone beads; this is largely inspired by methods and analyses by Reinhart, Statt, and coworkers in the context of colloidal crystals, ice crystals, binary mesophases, and model polymer aggregation.\cite{48,50,64} Here, we consider the local density of backbone beads $n(r_c) = N/\sigma^3$ to be descriptive of the local environment of a bead, such that a given configuration of an SCNP can be numerically represented by a histogram of such local densities. In this study, local densities are measured only for backbone beads using a spherical cutoff of $r_c = 6\sigma$, and histograms possess 40 evenly-spaced bins over the range $[6.63 \times 10^{-3}, 0.239]\sigma^{-3}$, which spans observed values across all configurations.

The histograms can be constructed for any given configuration of an SCNP to provide a 40-dimensional feature vector $x_i$. To obtain a representative ensemble, twenty distinct
configurations for each SCNP (dumped at a frequency of $2.5 \times 10^6$ timesteps) are featurized. This results in $320 \times 24$ independently formed SCNPs $\times 20$ configurations $= 153,600$ total feature vectors; the 20 configurations taken for each SCNP accounts for structural fluctuations, which may be of interest in certain applications. Subsequently, the uniform manifold approximation and projection (UMAP) algorithm is used to learn a low-dimensional numerical embedding of the SCNP morphologies and the manifold over which they are distributed. To facilitate visualization and interpretation, we simply target a two-dimensional embedding. Hyperparameters related to UMAP include the size of the local neighborhood (set to 200), the overall density of the embedding (minimum distance between points set to 1), and the distance metric for points in the feature space (Euclidean). In effect, this approach generates a mapping $\mathbb{R}^{40} \rightarrow \mathbb{R}^2 : \text{UMAP}(x_i) = Z_i$ where $Z_i$ is a coordinate vector in the learned low-dimensional representation of the set of local densities around monomers. As such, $Z_i$ provides information of the nanostructural characteristics of SCNP. We note that additional descriptions of the local environment, including that used by Reinhardt$^{64}$ as well as those equivalent to atom-centered symmetry functions,$^{65}$ were also examined for the embedding described above (see supporting information, Fig. S2). However, all approaches yielded qualitatively similar organization of morphologies, and local density was thus employed for its ease of interpretability, despite it being least the descriptive of the local environment.

### 2.2.3 Metrics of Dispersity and Sequence Variability

To quantify morphological dispersity in SCNPs and assess the importance of precise sequence control, we introduce a series of so-called dispersity and sequence variability metrics. Dispersity metrics primarily address variability in outcomes of SCNPs for a given precursor chain, while sequence variability metrics address the range of possible outcomes for a given specification of $f$ and $\beta$, which is associated with a set of realizable precursor chains. In the following, a given SCNP is characterized by a tuple $(c(f, \beta), s_c)$ where $c$ denotes a precursor...
chain from the set $C$ of all possible precursor chains with characteristics $f$ and $\beta$, and $s_c$ is an element of the set $S_c$ of all possible SCNP structures that can form from precursor $c$. In addition, the notation $x \sim Y$ indicates that the variable $x$ was sampled from the set $Y$. In the present study, relevant sets practically correspond to (i) the 24 independent trajectories that yield unique SCNP s from a given precursor chain or (ii) the ten distinct precursor chains generated for each combination of $f$ and $\beta$; these sets are respectively used in the calculation of expectation values indicated by $E_{s_c \sim S_c}$ and $E_{c \sim C}$.

Dispersity in the size of the SCNP s arising from a given precursor chain is given by

$$D_{R_g}(c(f, \beta)) \equiv \mathbb{E}_{s_c \sim S_c}[(\langle R_g \rangle_{s_c} - \mathbb{E}_{s_c \sim S_c}[\langle R_g \rangle_{s_c}])^2];$$

(10)

where $\langle R_g \rangle_{s_c}$ is the ensemble-average $R_g$ for a given SCNP; this is the variance in $\langle R_g \rangle_{s_c}$ sampled over $S_c$. Dispersity in $\langle Z \rangle$ (related to the density of local environments) for SCNP s arising from a given precursor chain is given by

$$D_Z(c(f, \beta)) \equiv \mathbb{E}[-\log(P(\langle Z \rangle|c))|c];$$

(11)

where ‘$x|c$’ denotes conditioning the variable $x$ on $c$. This quantity is the entropy of the distribution of $\langle Z \rangle$ and measures the differences in the nanostructures of the SCNP that are formed from a given precursor $c$.

Because not all manifestations of $c(f, \beta)$ are expected to yield the same distribution of SCNP s, we define four sequence variability metrics. Sequence variability on the outcome of $R_g$ at a particular $f$ and $\beta$ is defined as

$$SV_{R_g}(f, \beta) \equiv \mathbb{E}_{c \sim C}[(\mathbb{E}_{s_c \sim S_c}[\langle R_g \rangle_{s_c}] - \mathbb{E}_{c \sim C}[\mathbb{E}_{s_c \sim S_c}[\langle R_g \rangle_{s_c}])^2];$$

(12)

which probes the variance of the mean SCNP sizes formed from $c$ across all $c$ in $C$. Sequence
variability on the dispersity of chain sizes is similarly defined as

\[ \text{SV}_{\mathcal{D}_{R_g}}(f, \beta) \equiv \mathbb{E}_{c \sim \mathcal{C}}[\mathcal{D}_{R_g}(c) - \mathbb{E}_{c \sim \mathcal{C}}[\mathcal{D}_{R_g}(c)]]^2; \]

(13)

this probes the variance of the dispersities of SCNP sizes (Eq. (10)) formed from \( c \) across all \( c \) in \( \mathcal{C} \). Because the position on the manifold \( \langle \mathbf{Z} \rangle \) is two-dimensional, sequence variability of \( \mathbf{Z} \) is instead defined as

\[ \text{SV}_{\mathbf{Z}}(f, \beta) \equiv \mathbb{E}[-\log(P(\langle \mathbf{Z} \rangle|f, \beta))|f, \beta], \langle \mathbf{Z} \rangle \equiv \mathbb{E}_{s \sim \mathcal{S}_c}[\langle \mathbf{Z} \rangle_s]; \]

(14)

this is the entropy of the distribution formed by the expectation values of \( \langle \mathbf{Z} \rangle_s \) over \( \mathcal{S}_c \) for all \( c \) in \( \mathcal{C} \). Lastly, we define the sequence variability in the dispersity of positions on the manifold:

\[ \text{SV}_{\mathcal{D}_{\mathbf{Z}}}(f, \beta) \equiv \mathbb{E}_{c \sim \mathcal{C}}[(\mathcal{D}_{\mathbf{Z}}(c) - \mathbb{E}_{c \sim \mathcal{C}}[\mathcal{D}_{\mathbf{Z}}(c)])^2]; \]

(15)

this is related to the variance in the dispersity of nanostructures (Eq. (11)) formed from \( c \) across all \( c \) in \( \mathcal{C} \).

3 Results & Discussion

3.1 Characterization of Morphologies

3.1.1 Unsupervised learning facilitates understanding of SCNP morphologies

To effectively process and identify morphological archetypes of all simulated SCNPs, we utilized unsupervised machine learning to construct a low-dimensional manifold of SCNP morphologies (Section 2.2) wherein the structure of every SCNP can be represented via a two-dimensional coordinate vector \( \mathbf{Z} = (Z_1, Z_2) \). Importantly, the morphologies of SCNPs are expected to share similar features when closely positioned on the manifold (e.g., when \( |Z_i - Z_j| \) is small).
Figure 2: Visualization and analysis of single-chain nanoparticle (SCNP) morphologies. (a) A two-dimensional manifold organization of morphologies for SCNPs based on the Uniform Manifold Approximation and Projection (UMAP) unsupervised learning algorithm. For visual clarity, only a fraction of the overall data is shown (see also Fig. S3). Each marker corresponds to a single configuration of a SCNP and is colored by the average local density around backbone beads, $\bar{n}$, revealing bands of constant $\bar{n}$ distributed across the manifold. Five additional markers (warm colors) that correspond to selected coordinates that span the manifold for each band are also shown; these coordinates are used for analysis in panel (b). Particular examples are noted within the figure axes for $\bar{n} = 80$. (b) The standard deviation of local densities around backbone beads, $\sigma(n)$, as a function of fractional distance $\lambda$ across each band. The distance across each band is estimated by selection of two distal reference points $z_0(\bar{n})$ and $z_1(\bar{n})$, such that intermediate positions can be selected as $z_\lambda = (1 - \lambda)z_0(\bar{n}) + \lambda z_1(\bar{n})$. Each marker corresponds to the average of $\sigma(n)$ for a collection of structures found within 0.3 units of the corresponding marker in (a); the line colors reflect the band for each $\bar{n}$. Error bars report the standard deviation of $\sigma(n)$ across the aforementioned collection of structures. (c) Simulation snapshots of 103 representative configurations across the latent space manifold. Each rendering is positioned above its coordinate position in the manifold. In the renderings, only backbone beads are shown and are colored by the local density of other backbone beads within a $6\sigma$ radius.
Fig. 2a shows that the manifold of SCNP morphologies, as determined by the unsupervised machine learning, is primarily organized by the average local density of backbone beads $\bar{n}$. Moving from left-to-right (increasing $Z_1$), SCNPs transition from compact (high $\bar{n}$, green colors) to expanded (low $\bar{n}$, blue colors) structures. Fig. 2b reveals a secondary level of organization, moreso along $Z_2$ wherein structures are differentiated by the distribution of local environments, which is quantified by the standard deviation of local densities, $\sigma(n)$. Moving from the bottom towards the top of the manifold within a band of constant $\bar{n}$, morphologies transition from having relatively homogeneous local environments towards having increasingly heterogeneous nanostructures. Thus, the unsupervised learning approach distinguishes between, for example, different manifestations of similarly compact SCNP morphologies (i.e., dense core and less confined shell versus more homogeneous structure).

Fig. 2c illustrates how representative SCNP structures are distributed across the low-dimensional manifold, revealing several archetypal morphologies that arise in congruence with the specific local environments. Specifically, structures on the left side of the manifold ($Z_1 < 5$) are globular, typified by relatively large $\bar{n}$. Structures on the top-right side of the manifold ($Z_1 \approx 5$, $Z_2 > 5$) resemble pearl-necklace and tadpole-like morphologies, typified by larger $\sigma(n)$. Structures on the bottom-right side ($Z_1 > 10$, $Z_2 \approx 0$) are populated with diffuse globular, worm-like, and rod-like morphologies, with lesser $\sigma(n)$. Therefore, it becomes possible to make a distinction between stringy versus clumpy pearl-necklace-like structures based on the heterogeneity of local environments. Collectively, these results indicate that (i) the precursor chains over the $\beta$-$f$ parameter space yield a rich array of SCNP morphologies and (ii) the manifold coordinates $Z$, which broadly correlate to the average and heterogeneity of nanostructural environments within an SCNP, provide a reasonable organization of those structures.

Observed archetypes also resemble those of prior studies, albeit with some distinctions. While Moreno and coworkers observe many of the same archetypes, they noted an abundance of pearl-necklace-like structures,\textsuperscript{16,20,28} and these are represented to a much lesser degree
across our morphologies. We suggest that this difference arises due to our use of angular constraints for linker-bond formation, which inhibits linkers forming bonds over short contour distances of the polymer backbone. In another study, Tulsi and Simmons found SCNPs that also adopt pearl-necklace, worm-like, and dense globular structures.\(^{30}\) They further identified “gnarled” structures, which are similar to the more diffuse globules observed in this study. Interestingly, their SCNPs arise from a model with different physical characteristics and distinct mechanism for assembly. This suggests that the present and prior theoretical studies may be homing in on a general set of readily realizable SCNP morphologies. In addition, many of our structures resemble tadpole morphologies which are similar to morphologies observed in experiments.\(^{66–68}\)

3.1.2 Sequence characteristics of precursors biases structure formation

To assess how linker patterning impacts SCNP structure, we examine how precursor parameters \(f\) and \(\beta\) dictate position on the manifold of SCNP morphologies (Figs. 3a,b). While \(f\) and \(\beta\) both impact morphology, they have different effects. Roughly, trends with \(f\) and
track with $\bar{n}$. This suggests that structures formed from high-$f$ precursors (left) predominantly display more compact, globular morphologies typified by high $\bar{n}$, while structures formed from low-$f$ precursors (right) result in more diffuse, low-$\bar{n}$ morphologies. The impact of $\beta$ is less striking but evident at low $f$ (i.e., $f \lesssim 0.3$). In particular, $\beta$ impacts how cross-links are distributed over the SCNP topology: at fixed and low $f$, high-$\beta$ precursors yield SCNPs with few centralized “hubs” of cross-links, while the low-$\beta$ precursors result in SCNPs with cross-links that are more evenly distributed over the chain. As a result, increasing $\beta$ biases morphologies from worm- or rod-like (bottom right) to more pearl-necklace- and tadpole-like structures (top right); the nature of this effect is further resolved in Fig. S4.

We characterized 7,680 unique SCNPs across the $\beta - f$ space by the number of free backbone beads $n_f$, the number of domains $n_d$, and the radius of gyration $R_g$ to quantitatively examine how particular morphologies arise as a result of $\beta$ and $f$. Figs. 3c-f show the averages of these structural descriptors over the ensemble of SCNPs generated from precursors with specific blockiness and linker-fraction combinations; precise numerical values are reported in the supporting information. All four descriptors possess identifiable trends that resonate with the qualitative observations from Figs. 2c and 3a,b. Fig. 3c shows that decreasing $f$ and increasing $\beta$ typically increases the extent of polymer chain that is not present in any cross-linked domain. Meanwhile, Fig. 3d indicates that decreasing $\beta$ and $f$ tends to result in SCNPs with more domains that possess fewer beads. Together, these results suggest that $\beta$ and $f$ can be manipulated to tune aspects of SCNP topology, although not arbitrarily. For example, at $f = 0.1$, decreasing $\beta$ of precursors will manifest in the formation of more domains separated by shorter free segments. Additionally, increasing $f$ tends to result in fewer and larger domains that can be tempered by shifting $\beta$. However, beyond a certain $f$, it appears these structural descriptors are not strongly affected by $\beta$. This is evident by the flat color saturation at $f \geq 0.6$, indicating that all SCNPs are essentially one large topological domain in this precursor regime. The nature of these topological structures has clear implications on the size of the SCNP, as seen in Fig. 3e. To first order, SCNP size,
as given by $R_g$, is controlled by the number of free segments, which is strongly biased by $f$; only at low $f$ does $\beta$ appear to have subtle impact on $R_g$; we observe near-identical trends for shape descriptors derived from the gyration tensor (Fig. S5). Rather, $\beta$ mostly controls the extent of heterogeneity of SCNP nanostructures (Fig. 3f). In particular, decreasing $\beta$ biases from morphologies that feature more heterogeneous local environments around beads towards those with more homogeneous ones, even while $R_g$ is largely unaffected. Collectively, these results illustrate how sequence patterning of precursors can be manipulated to bias the morphological characteristics of SCNP.

3.2 Characterization of Structural Dispersity

3.2.1 Morphological dispersity depends on sequence patterning

We next investigate the impact of precursor patterning on the consistency of forming SCNP morphologies. Fig. 4 illustrates that a single precursor sequence can give rise to a set of SCNP with diverse morphological characteristics. This is first demonstrated in Fig. 4a, which compares $P(\langle R_g \rangle)$ for ten SCNP originating from precursor chains with $f = 0.1$ and $\beta = 0.2$ and ten originating from precursor chains with $f = 0.1$ and $\beta = 0.8$. Selected sequences (e.g., 5, 6, and 9 for $f = 0.1$ and $\beta = 0.8$) are noted to have particularly broad distributions, which result from averaging over the 24 replicate simulations. Across both precursor conditions, many distributions are non-Gaussian, bordering on bimodal or heavy-tailed, which would indicate that the different trajectories produce disparate morphologies rather than just fluctuate about a dominant archetypal morphology. Figs. 4b,c shows that there can also be substantial nanostructural diversity in the ensemble SCNP formed by a given precursor sequence; this would be evidenced by a diffuse $P(\langle Z \rangle)$ for a given sequence. Such diffuse distributions are particularly identifiable for sequence 0, 1, and 3 in Fig. 4b, which derive from low-blocky precursors. However, sequence patterning can significantly impact the tightness of $P(\langle Z \rangle)$, as sequence 2 for $\beta = 0.2$ is much more strongly peaked than that of sequence 0. Diffuse $P(\langle Z \rangle)$ for $\beta = 0.8$ (Fig. 4c) are not quite as evident, and
Figure 4: Morphological dispersity of single-chain nanoparticles originating from given precursor sequences. (a) A comparison of distribution of radius of gyration $P(\langle R_g \rangle)$ for distinct single-chain nanoparticles. The distributions are obtained from 24 independent replicate simulations of the same precursor chain sequence. The data are for SCNPs formed from ten precursor sequences each from $f = 0.1$ and $\beta = 0.2$ (yellow, left) and $f = 0.1$ and $\beta = 0.8$ (green, right). The width of violins correspond to the density obtained from Gaussian kernel density estimation; the edges of boxplots in the violin depict the inner quartile range, while white dots indicate the median value. A comparison of the distribution of manifold-coordinate vectors $P(\langle Z \rangle)$ for single-chain nanoparticles formed from selected precursor chain sequences with parameters of (b) $f = 0.1$ and $\beta = 0.2$ and (c) $f = 0.1$ and $\beta = 0.8$. In panels (b) and (c), the color reflects Gaussian kernel density estimation over the $\langle Z \rangle$ for each of the 24 distinct SCNPs formed by each precursor sequence. The color scheme reflecting precursor parameters and sequence labels is the same across panels.
the $P(\langle Z \rangle)$ are generally similar in shape, hinting at the possible role of precursor parameters like $\beta$ on dispersity.

### 3.2.2 Precursor parameters have disparate effects on different measures of dispersity

Fig. 5 elucidates the role of $f$ and $\beta$ on both dispersity in $P(\langle R_g \rangle)$ and dispersity in $P(\langle Z \rangle)$. Intriguingly, we find that these measures of dispersity are generally anticorrelated. This is evident by examining trends with increasing $f$, which tends to decrease dispersity in $P(\langle R_g \rangle)$ and increase dispersity in $P(\langle Z \rangle)$. This can be understood by considering the high-$f$ regime, for which well-connected globular morphologies would have similar overall size and limited capacity to change (low dispersity in $P(\langle R_g \rangle)$) but can nonetheless have disparate populations of local environments (low dispersity in $P(\langle Z \rangle)$), such as observed in Fig. 2. At low $f$ but high $\beta$, dispersity in $P(\langle R_g \rangle)$ is high yet dispersity in $P(\langle Z \rangle)$ is diminished. Combined with the observation that SCNP formed from precursors with these characteristics resemble pearl-necklace-like morphologies, diminished dispersity in $P(\langle Z \rangle)$ suggests that the formation of dense (pearl) and diffuse (lace) nanostructures is relatively consistent. Furthermore, heightened dispersity in $P(\langle R_g \rangle)$ suggests that depending on how these nanostructures are arranged or connected (i.e, the precise topology) could result in relatively large changes $P(\langle R_g \rangle)$. In addition, we observe an apparent maximum in $P(\langle Z \rangle)$ dispersity at intermediate $f$ (0.25,0.3) and intermediate $\beta$ (0.4,0.6), suggesting relative greater diversity in the set of nanostructures between SCNP formed from these precursors. Moving away from this maximum towards higher $f$ decreases the entropy of the distributions but to a lesser degree than moving away from the maximum towards lower $f$. This reflects the consistent, high-density core formed by structures at high $f$ but high variability in local densities around the outside of the structure that depends on the precise folded shape of the SCNP even when broadly characterized as a compact globule.
Figure 5: Effect of precursor parameters on morphological dispersity. Heat maps at a given linker fraction \( f \) and normalized blockiness \( \beta \) for the average (a) dispersity in \( \langle R_g \rangle \) given by \( D_{R_g} \) in Eq. (10) and (b) dispersity in manifold coordinates given by \( D_Z \) in Eq. (11). For both panels, average effects are estimated by considering the ten unique precursor chains generated at a given \( f \) and \( \beta \).

### 3.2.3 Specific sequences enable more precise control over morphology

Differences in the distributions of \( \langle R_g \rangle \) and \( \langle Z \rangle \) across sequences with the same \( \beta \) and \( f \) exhibit differences (Fig. 4) suggest that sequences might be precisely crafted to fine-tune characteristics of formed morphologies. To examine this, we calculate four “sequence variability” metrics that report how distinct aspects of SCNP morphologies are across different sequences for the same \( f \) and \( \beta \). In particular, we specifically examine the sequence variability in the mean of \( P(\langle R_g \rangle) \), the dispersity in \( P(\langle R_g \rangle) \), the mean of \( P(\langle Z \rangle) \), and the dispersity in \( P(\langle Z \rangle) \). We anticipate that understanding sequence variability may be useful to ascertain whether tailoring sequences would offer prospective advantages over an ensemble of sequences defined by a given \( f \) and \( \beta \).

Fig. 6a,b shows that sequence variability is highest for the mean value and dispersity of \( P(\langle R_g \rangle) \) at low \( f \) and high \( \beta \) values. By contrast, sequence variability in the mean and dispersity of \( P(\langle R_g \rangle) \) is diminished at high \( f \), suggesting that precise sequence control would
Figure 6: Analysis of sequence effects on morphological control. Heatmaps show sequence variability (Seq. Var.) as a function of linker fraction $f$ and normalized blockiness $\beta$ calculated as (a) variance in the means of $P(\langle R_g \rangle)$, (b) variance in dispersities of $P(\langle R_g \rangle)$, (c) entropy of the means of $P(\langle Z \rangle)$, and (d) variance in the dispersity of $P(\langle Z \rangle)$. Quantities are computed from 10 independently generated sequences for each specified $f$ and $\beta$. 
be less valuable; this observation is consistent with prior results (Fig. 3 and Fig. 5) that indicated negligible impact of $\beta$ on structure formation at high $f$. Furthermore, Fig. 6c indicates that sequence variability in the mean of $P(\langle Z \rangle)$ follows a similar trend, peaking at small $f$ and intermediate values of $\beta$ and decreasing at large $f$. There are no discernible trends in sequence variability for the dispersity of $P(\langle Z \rangle)$ (Fig. 6d). Together, these results suggest that precisely tailoring sequences to manipulate the distribution of formed SCNPs, whether it be aspects of the SCNP size or its average local environment, is likely to be most effective for low $f$ and intermediate to high $\beta$.

4 Conclusions

In this work, we examined how sequence patterning of intramolecular cross-linking moieties (linkers) in polymer chains impacts the formation of single-chain nanoparticles (SCNPs). To do so, we simulated the formation of 7,680 unique SCNPs from precursor chains, which altogether comprehensively spanned a parameter space defined by the number of linkers in the chain, $f$, and the blockiness of the chain, $\beta$. The morphologies of the SCNPs were subsequently characterized using unsupervised machine learning and several structural descriptors (e.g., the radius of gyration, the number of free beads, and the number of topological domains) to elucidate the general roles of $\beta$ and $f$ on morphological outcomes of SCNPs. Finally, we assessed how $\beta$ and $f$ impact dispersity in the sizes of SCNPs ($P(\langle R_g \rangle)$) and its characteristic nanostructural environments ($P(\langle Z \rangle)$), and we explored how effectively precise sequence control might influence these behaviors.

Overall, we found several major trends with respect to SCNP topology, morphology, and precursor parameters. Using manifold-learning, we showed that SCNP morphologies could be distinguished principally by the mean-density of beads and secondarily by the heterogeneity of local nanostructural environments. For example, we identified two distinct globular morphologies (one with a dense core-flexible shell structure and one with more
homogeneous density throughout). Specific archetypal morphologies were also found to be strongly associated with certain regimes of precursor parameters. Low-$f$ precursors tend to adopt topologically diffuse structures with several domains and large chain segments between domains; in this regime, increasing $\beta$ biases structure-formation towards fewer and larger domains but a higher proportion of the polymer chain that is not within any topological domain. By contrast, precursors with high $f$ consistently give rise to SCNPs with globular structures that show weak sensitivity to $\beta$. In addition, while low-$\beta$ precursors tend to result in SCNPs with rod-like morphologies (i.e., several domains of dense nanostructures), high-$\beta$ precursors result in tadpole-like morphologies (i.e., fewer domains with disparate nanostructures). By examining the dispersity of structures formed from every precursor, we found that low-$f$ precursors yield SCNPs with substantial diversity in size but overall consistent local environments, while high-$f$ precursors generate SCNPs with the opposite trend. Furthermore, comparing the distributions of SCNPs generated from specific sequences that all correspond to the same $\beta$ and $f$ highlighted the potential to leverage sequence patterning for morphological control, manifest in either specific average values or distribution characteristics. Investigation of new “sequence variability” metrics revealed that precursor chains with low $f$ and large $\beta$ illustrate the widest range of possible morphology and dispersity outcomes. Therefore, we hypothesize that this regime could benefit most from precise sequence control to tailor the morphological properties of SCNPs.

Ultimately, the methods and analyses herein may spawn several directions of future research. First, while the present and many prior studies emphasized the final SCNP morphologies, future work may aim to quantify the physical implications of SCNP structure and understand their relationship to other SCNP properties (e.g., response to shear flow, mechanical unfolding, etc.). Furthermore, there is significant opportunity to establish how precursor patterning dictates the mechanism(s) or pathway(s) of SCNP formation. Examination of formation pathways may facilitate additional understanding as to how SCNP morphology might be more tightly controlled. Finally, our results point to the possibility of
tuning the dispersity of SCNP morphologies in numerous ways. While prior work has lever-
aged assembly protocols to bias structure-formation (e.g., towards globular morphologies\textsuperscript{29}),
here we show that dispersity outcomes (either SCNP size or nanostructural environments)
might be manipulated by either setting particular precursor parameters or by tailoring se-
quences. Furthermore, with many demonstrations of using statistical ensembles of chains in
functional materials,\textsuperscript{69–74} it will be interesting to consider how more precise sequence pat-
terning might enhance materials performance. Approaches to navigate such design tasks,
particularly in the context of experimentally verifiable systems will be needed.

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**Supporting Information**

Information regarding simulation protocols. Additional strategies explored for unsupervised
learning. Correlation between topological descriptors. Correlation between topological and
morphological descriptors. A complete dataset including SCNP properties. List of SCNP
precursor sequences. All data associated with single-chain nanoparticles, their characteris-
tics, and accompanying Python scripts are also available for download at

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