“Michael” Nanocarriers Mimicking Transient-Binding Disordered Proteins

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Supporting Information

ABSTRACT: We report herein a very efficient synthesis strategy for the construction of artificial transient-binding protein-mimic nano-objects. Michael addition-mediated multidirectional self-assembly of individual polymeric chains at r.t. leads to “Michael” nanocarriers that in solution resemble disordered multidomain proteins, as revealed by a combination of small angle neutron scattering (SANS) measurements and coarse-grained molecular dynamics (MD) results, whereas in the dry state adopt a collapsed, globular morphology, as observed by transmission electron microscopy (TEM). This extended-to-compact morphology transition taking place upon solvent removal is of paramount importance, among other applications, for the construction of efficient biosensors based on immobilized protein-mimic nano-objects and for the development of transient vitamin-binding systems. As a proof of concept, we show the controlled delivery of vitamin B9 from these novel transient-binding nanocarriers.

Nature is a continuous source of inspiration for scientists across different disciplines. In particular, the specific –native– conformation of proteins allowing these large biomolecules to carry out sophisticated tasks such as catalysis inspired the construction of a first generation of artificial enzymes based on a variety of molecular and macromolecular structures such as macrocyclic compounds,2 star and helical polymers,3 dendrimers4 and micelles.5 Single-chain polymer nanoparticles are emerging soft nano-objects showing unique and remarkable physicochemical, rheological and sensing properties,6 as a result of their locally collapsed structure and its ultra-small size. We7 and others8,9 have demonstrated that catalytic properties can be imparted to single-chain nanoparticles through polymer folding/collapse accompanied by efficient catalyst immobilization. The potential use of single-chain nanoparticles as drug / siRNA nanocarriers and photostable bioimaging agents relies on the improved construction of functional folded/collapsed single polymer chains. Moreover, controlled synthesis of single-chain nanocarriers becomes critical to the elucidation of useful structure-property relationships in materials science and to deepen our current understanding of complex cooperative folding events taking place in synthetic single polymer chains emulating protein folding.10 Unfortunately, most of the current synthesis routes to stable single-chain nanoparticles suffer from different shortcomings, such as use of extremely high temperatures, requirement of severe anhydrous conditions, involvement of metallic catalysts or necessity of exotic, non-commercial monomers, which severely limit their potential applications in some promising fields (e.g., nanomedicine).11 Despite all recent advances, the development of a natural route to well-defined single-chain nano-objects allowing to investigate how far, or close, are the structure and function of these synthetic nanocarriers from those of globular, or intrinsically disordered, proteins is still a challenging issue.

In this letter, we report a simple and highly-efficient bio-inspired method for transient-binding nanocarrier construction that relies on multidirectional self-assembly of individual polymeric chains at r.t. driven by multiple intrachain Michael addition reactions involving external multifunctional acrylate-based cross-linkers (see Scheme 1). Single-chain nano-object formation through multidirectional self-assembly is inspired by protein assembly to the folded state. For proteins, the driving force for folding depends on the sequence of amino acids, their mutual interactions and their interactions with solvent molecules, whereas for “Michael” nanocarriers the driving force is replaced by multiple, cooperative, omnidirectional chemical (Michael addition) reactions taking place across the collapsing polymer chains under good solvent conditions. The Michael reaction has been previously employed in the synthesis of linear, graft, hyperbranched, den-
direct monitoring in solution of the self-organized, cooperative folding/collapse process, iv) versatility for obtaining nano-objects resembling in solution transient-binding disordered proteins, and v) promising nanocarrier properties of the resulting multidomain nano-objects.

“Michael” nanocarriers were prepared from polymeric precursors synthesized through reversible addition-fragmentation chain-transfer (RAFT) polymerization by starting with commercially available methyl methacrylate (MMA) and (2-acetoacetoxy)ethyl methacrylate (AEMA) monomers (SI, Schemes S1-S2). Statistical copolymers were obtained due to the similar reactivity ratios of MMA ($r = 0.90$) and AEMA ($r = 0.95$) providing materials of high molar mass, relatively narrow size dispersity and random distribution of β-ketoester functional groups (i.e., Michael donors) along the polymer chain. As external cross-linkers playing the role of Michael acceptors, we selected commercially available low-molecular-weight multi-functional (bi-, tri-, tetra-, penta / hexa-) acrylate compounds (see SI, Scheme S1). The multidirectional self-assembly of the individual polymeric precursor to Michael nanocarriers was performed in tetrahydrofuran (THF), at r.t. under potassium hydroxide catalysis and air atmosphere, at a concentration of polymeric precursor of 1 mg / ml by using equimolar amounts of β-ketoester and acrylate functional groups. As an example, Figure 1A shows the systematic shift in size exclusion chromatography (SEC) traces towards longer retention time during the synthesis of single-chain Michael nanocarriers in the presence of a tri-functional cross-linker (trimethylolpropane triacrylate, TMT). The noticeable shift observed is a consequence of the progressive reduction in hydrodynamic size and, consequently, in apparent molar mass ($M_w$) with reaction time (open circles in Figure 1A). The absence of significant inter-particle aggregation was confirmed through simultaneous light scattering (SLS) measurements from which, as expected, a nearly constant value of actual molar mass was observed ($M_w \approx 293$ kDa, closed circles in Figure 1A, Figures S1-S3). Under identical reaction conditions (i.e., solvent, temperature, reagent concentration) the kinetics of the Michael addition reaction for multidirectional self-assembly was ca. 50-fold slower than that corresponding to condensation via Michael addition of mixtures of low-molecular-weight model compounds (e.g., methyl acetoacetate / TMT mixtures) as determined by $^1H$ NMR measurements (Figure 1B, Figures S4-S5). The main reason for the significantly slower kinetics seems to be the macromolecular character of the polymeric precursor which imposes diffusional constrains and significantly restricted intrachain accessibility to the low-molecular-weight acrylate cross-linker units.

We have determined the kinetics of the folding/collapse process in solution through the direct monitoring of the value of $z$-average radius of gyration ($R_g$) of the polymeric precursor as a function of reaction time, by means of static light scattering (SLS) measurements (Figure 1C). The kinetics of the folding/collapse process was found to be ca. 5-fold slower than that of the Michael-mediated intrachain cross-linking reaction, which can be attributed to the coexistence of non-efficient and efficient folding events. In this sense, the first anchoring step of a multifunctional cross-linker molecule through reaction of a single acrylate group can be considered as a non-efficient folding event that does not contribute to chain collapse. Conversely, the delayed, subsequent intrachain Michael reactions of the remaining acrylate groups are truly folding/collapse-promoting events. As illustrated in Figure 1C, the kinetics of folding/collapse is slightly faster upon decreasing the molar mass of the polymeric precursor. Even displaying much slower folding kinetics when compared to that of natural proteins (from microseconds to hours), the resulting Michael nanocarriers in the dried state showed a compact spherical morphology as illustrated in Figure 1D.

Scheme 1. Analogy between protein folding (A) and multidirectional self-assembly during the synthesis of “Michael” nanocarriers (B). The driving force for nanocarrier formation in B is intramolecular Michael addition reaction taking place in a multiple, cooperative and omnidirectional manner at room temperature (C) involving external multifunctional acrylate-based cross-linkers that react with β-ketoester groups of the unfolded polymer chain under appropriate solvent, stoichiometric and dilution conditions (D).
Figure 1. “Michael” nanocarrier construction through multidirectional self-assembly: (A) The shift in SEC traces upon reaction time (from left to right: 0 h, 17 h, 48 h and 72 h) is due to a progressive reduction in hydrodynamic volume and, consequently, in apparent molar mass, \( \left( M_{\text{app}} \right) \) (open symbols), whereas the nearly constant value of actual molar mass \( (M_p \approx 293 \text{ kDa}, \text{closed symbols}) \) points to the collapse of individual polymer chains without significant inter-particle aggregation. (B) Pseudo-first order kinetics plots corresponding to the consumption of bi-functional (closed circles) and tri-functional (closed triangles) cross-linker units (X). Also displayed for comparison are the corresponding kinetics plots for low-molecular-weight model compound mixtures (open symbols). (C) Pseudo-first order kinetics plots corresponding to the folding/collapse process in solution at r.t. in the presence of tri-functional cross-linker units. (D) TEM image showing the morphology of Michael nanocarriers in the dry state.

Upon isolation by precipitation, drying under dynamic vacuum and further solubilization, Michael nanocarriers were found to form transparent, colorless dispersions in common organic solvents (tetrahydrofuran, chloroform, dimethyl formamide). Unfortunately, all our attempts to synthesize well-defined Michael nanocarriers by means of tetra- or penta/hexa-acrylate cross-linker molecules failed suggesting that under our reaction conditions the optimum cross-linker functionality is \( f = 3 \). We suspect that the increased steric hindrance for cross-linkers of \( f > 3 \) and the higher tendency to form insoluble macroscopic gels due to inter-particle coupling events could be responsible of this behavior.

Valuable information about the actual structure in solution of single-chain Michael nanocarriers synthesized by means of bi- and tri-functional cross-linker units was obtained through small angle neutron scattering (SANS) measurements in deuterated solvent and complementary molecular dynamics (MD) simulations. Figure 2A shows Kratky plots for the unfolded polymeric precursor and the resulting nano-objects are drawn in Figure 2B. It is worth noting the qualitative agreement between SANS and MD simulation results. For both kinds of nanoparticles, MD simulation data showed increased compact conformations on decreasing the quality of the solvent, which is consistent with the globular conformation observed by TEM for the Michael nanocarriers in the dry state (Figure 1D).
Inspired by the behavior of natural transient-binding disordered proteins, we have investigated the drug-delivery properties of Michael nanocarriers synthesized with tri-functional cross-linker units for potential application in dermal supply of vitamin B\textsubscript{9} (i.e., folate or folic acid). Several in vitro and in vivo studies have indicated that this essential vitamin may offer a treatment option for photo-aged skin when incorporated in topical formulations.\textsuperscript{21} Hence, vitamin B\textsubscript{9} nanocarriers based on single-chain Michael nanoparticles were placed in distilled water at neutral pH to investigate their controlled delivery properties. Figure 2C illustrates the progressive delivery of vitamin B\textsubscript{9} from the single-chain Michael nanocarriers as determined from UV/Vis spectroscopy measurements at 283 nm. The continuous line is a best-fit to the well-known power law model ($C_t / C_f = K t^n$, $C_t =$ concentration of drug released at time $t$, $C_f =$ total concentration of drug released, $K =$ constant, $n =$ release exponent).\textsuperscript{22} The value obtained for the release exponent, $n = 0.5$, suggests that the delivery process proceeds through a Fickian diffusion mechanism. Complete delivery of vitamin B\textsubscript{9} from Michael nanoparticles with a drug loading content of 41 wt % was observed to take place in 5-6 h. These promising results pave the way for mimicking the multiple transient-binding behavior of intrinsically disordered proteins (IDPs)\textsuperscript{23} using Michael nanocarriers.

In summary, we have developed a highly-efficient strategy for the permanent multidirectional self-assembly of unfolded polymeric chains to Michael nanocarriers that in solution resemble disordered multidomain proteins with semiflexible linkers. Conversely, in the dry state they adopt a collapsed, globular morphology. This natural synthesis route proceeds under very mild reaction conditions (without metal catalysts, at room temperature, under air atmosphere) by starting with commercially available reagents, allowing the efficient preparation of novel “Michael” nanocarriers showing multiple, locally compact domains in solution that turn to a collapsed, globular morphology in the dry state. By using single-chain Michael nano-objects as vitamin B\textsubscript{9} nanocarriers, sustained release of vitamin B\textsubscript{9} in water at neutral pH following a Fickian diffusion mechanism has been demonstrated. Both the multidirectional self-assembly approach, as a synthesis strategy to nano-objects resembling partially folded multidomain proteins in solution, and the own Michael nanocarriers, resembling transient-binding IDPs, are expected to find applicability in different fields, such as nanomedicine, biosensing / bioimaging uses, and heterogeneous catalysis.

ASSOCIATED CONTENT

Supporting Information. Materials, methods, characterization techniques, molecular dynamics (MD) simulations and supporting data. This material is available free of charge via the Internet at http://pubs.acs.org.

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The authors declare no competing financial interest.

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(19) We are not claiming that “Michael” nanocarriers have similarity to the precise sequence of proteins but that their form factor in solution resembles that of disordered proteins, as a consequence of the multidirectional self-assembly process which leads to the formation of local globules along the individual chains. In resemblance to transient-binding disordered proteins, the resulting Michael nanocarriers are able to bind temporally vitamin B9 that can be delivered in a controlled manner (Figure 2C).
Multidirectional Self-Assembly

IN DRY STATE

Multidirectional Self-Assembly

IN DRY STATE

Globular

IN DRY STATE

Globular

IN DRY STATE

SANS

Disordered

IN SOLUTION

Disordered

IN SOLUTION

Vitamin B$_9$

Vitamin B$_9$

0 1 2 3 4 5 6

0 1 2 3 4 5 6

t (h)

t (h)