



Cite this: *Soft Matter*, 2025, 21, 2759

Recent advances in nanoprecipitation: from mechanistic insights to applications in nanomaterial synthesis

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Nanoprecipitation is a versatile, low-energy technique for synthesizing nanomaterials through controlled precipitation, enabling precise tuning of material properties. This review offers a comprehensive and up-to-date perspective on nanoprecipitation, focusing on its role in nanoparticle synthesis and its adaptability in designing diverse nanostructures. The review begins with the foundational principles of nanoprecipitation, emphasizing the impact of key parameters such as flow rate, mixing approach, injection rate, and Reynolds number on nanomaterial characteristics. It also discusses the influence of physicochemical factors, including solvent choice, polymer type, and drug properties. Various nanoprecipitation configurations—batch, flash, and microfluidic—are examined for their specific advantages in controlling particle size, morphology, and internal structure. The review further explores the potential of nanoprecipitation to create complex nanostructures, such as core-shell particles, Janus nanoparticles, and porous and semiconducting polymer nanoparticles. Applications in biomedicine and other fields highlight nanoprecipitation's promise as a sustainable and tunable method for fabricating advanced nanomaterials. Finally, the review identifies future directions, including scaling microfluidic techniques, expanding compatibility with hydrophilic compounds, and integrating machine learning to further enhance the development of nanoprecipitation.

Received 4th January 2025,
Accepted 2nd March 2025

DOI: 10.1039/d5sm00006h

rsc.li/soft-matter-journal

Introduction

Nanoprecipitation, a simple solution-based approach for the preparation of nanoparticles, was first introduced by Fessi *et al.*^{1,2} Recently, it has gained increasing attention as a versatile technique for the synthesis of novel nanomaterials, owing to its simplicity, low energy requirements, ease of implementation, tunability, reproducibility, and versatility. Nanomaterials produced *via* nanoprecipitation include polymer particles,^{3–6} cellulose acetate particles,⁷ protein nanoparticles,⁸ semiconductor nanoparticles,^{9,10} Janus nanoparticles,^{11,12} nanocrystals,¹³ and mesoporous particles.¹⁴ One significant advantage of nanoprecipitation is its ability to form nanoparticles without requiring surfactants or toxic, undesirable organic solvents during synthesis. Recent advancements in nanoprecipitation techniques have further enhanced its efficiency, enabling the development of new functional nanomaterials.

A typical nanoprecipitation process involves mixing a polymer solution dissolved in an organic solvent with an aqueous solution. When the organic solution containing polymers rapidly and uniformly mixes with the aqueous non-solvent, it crosses the solubility barrier, resulting in the precipitation or phase separation of the solute into nanoparticles within the continuous phase. The resulting oversaturation and chain collapse lead to the formation of polymer nanoparticles (PNPs), which can range in size from a few nanometers to several micrometers. The formation of these nanomaterials involves three key processes: (1) mixing of the organic solution (containing solute molecules) with an anti-solvent (aqueous phase), (2) nucleation of solute molecules, and (3) aggregation and growth into nanomaterials.^{7,15–17} This same process can also nucleate liquid droplets through spontaneous emulsification, a phenomenon known as the “ouzo effect” or solvent exchange in flow systems.^{18–21} Alternatively, similar effects can occur in evaporating ternary liquid mixtures.^{22,23}

Several review articles have summarized specific aspects of preparing polymer nanoparticles (PNPs).^{24–27} In this review, we aim to provide an updated and holistic perspective on the physicochemical aspects and hydrodynamics that significantly influence nanoprecipitation. We place particular emphasis on

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the solution and flow conditions in various mixing configurations used during synthesis and their impact on controlling nanomaterial formation. Additionally, we discuss the fundamental principles of different mixing configurations for nanoprecipitation. Special attention is given to key parameters, such as mixing methods and conditions, including flow rate, stream velocity, injection rate, and the properties of fluids and their compositions – such as solvent, non-solvent, polymer, and the physicochemical conditions of the liquid mixtures.²⁸ The general principles discussed in this review provide a framework for understanding how the final size, structure, and properties of nanomaterials are determined. While the field of nanoprecipitation is rapidly evolving, and it is beyond the scope of this review to cover all recent articles, we aim to present

representative examples in each section. These examples are intended to inspire further exploration and advancements in future studies.

Basic principles of nanoprecipitation

The nanoprecipitation technique relies on the transition from miscibility to immiscibility, achieved by mixing an organic solution of the solute (e.g., polymers or small molecules) with an anti-solvent (miscible with the organic phase but a non-solvent for the solute), as illustrated in Fig. 1(a). Nanoparticle formation occurs by reducing the solvent quality through the addition of the anti-solvent. This technique is also known as

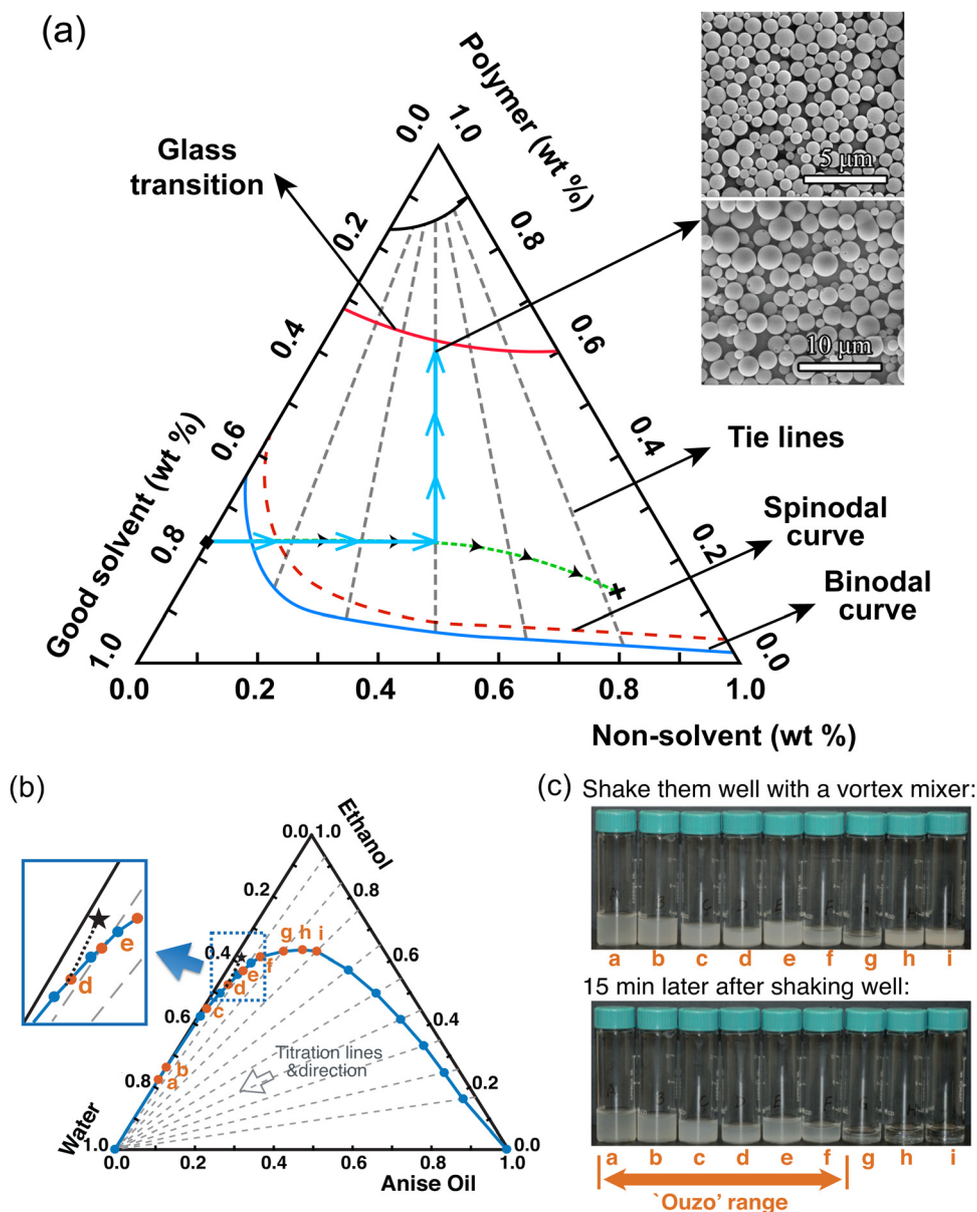


Fig. 1 Nanoprecipitation in ternary systems. (a) Ternary phase diagram for a polymer in a binary solvent and the SEM images of the synthesized polymer nanoparticle.^{15,32} (b) Solubility diagram and (c) photos of the ternary mixtures of ethanol, water, and anise oil.²²



solvent shifting, solvent displacement, anti-solvent precipitation, or solvent exchange, the latter referring to the anti-solvent displacing the solution in a directional flow.^{19,29,30} When the organic and aqueous phases are mixed, the process leads to rapid saturation of the solute to its critical nucleation concentration. This occurs due to the miscibility between the solvent and anti-solvent, combined with the solute's immiscibility in the anti-solvent. Fig. 1(b) shows the ternary diagram of water, ethanol, and anise oil, with the blue solid line representing the measured phase-separation curve. The black star and the black dotted line in the inset indicate the initial composition of the Ouzo drop and its trajectory over time based on numerical simulations. The gray dashed lines represent the paths of various composition coordinates from titration experiments. Fig. 1(c) shows the stability of the macrosuspension for compositions labeled (a)–(i) in the ternary diagram. This comparison reveals that the curve connecting the solid circles (a)–(f) in Fig. 1(b) marks the boundary of the Ouzo region, corresponding to the critical composition at which the Ouzo effect occurs. The process of nanoparticle formation can be understood through thermodynamic principles. As the solvent quality decreases, the solute experiences a supersaturation state, driving nucleation. According to classical nucleation theory, the critical nucleus size is determined by the balance of volumetric free energy change and interfacial energy. High supersaturation lowers the energy barrier for nucleation, promoting the rapid formation of numerous small particles.³¹

Many ternary mixtures consist of both miscible pairs and immiscible pairs. The solutes in such systems can include polymers, oils,^{33–36} or even gases.^{37,38} When the solute is oil, the nanoprecipitation process is equivalent to what is commonly referred to as the Ouzo effect.^{22,23,29} The most common solvent and anti-solvent combinations for nanoprecipitation are polar organic solvents with water (or an aqueous solution). However, other combinations are also feasible, including organic solvent–organic solvent systems,^{34–36} ionic liquid–ionic liquid systems,^{39,40} and deep eutectic solvents.⁴¹

The nucleation and growth of supersaturated solute molecules lead to the formation of polymer nanoparticles. Once nucleation is initiated, the growth of nanoparticles proceeds *via* the diffusion of solute molecules from the surrounding medium. Fick's laws of diffusion describe the transport of solute molecules, where the flux is proportional to the concentration gradient. Fast diffusion ensures uniform particle growth, while variations in local concentrations can lead to particle size heterogeneity. Both kinetic and thermodynamic factors play a role in the particle structure and size distribution.^{6,42–44} Both mixing and diffusion processes occur simultaneously, depleting the solute molecules in the mixture. Additionally, the anti-solvent dilutes the solute, reducing its concentration. As the solute concentration falls below the critical nucleation threshold, nucleation and growth cease. The formed nanoparticles remain dispersed in the mixture, trapped in a thermodynamically metastable state. The rate of oversaturation plays a critical role in determining nanoparticle size. A high nucleation rate leads to the formation of smaller particles with a narrow size distribution. A key complexity in this

process is that nanoparticle formation arises from local and temporal oversaturation of the solute during mixing. Oversaturation evolves over time and may also vary spatially. Both the physicochemical properties of the compounds in the mixture and the mixing dynamics between the solution and the antisolvent are critical in creating the out-of-equilibrium oversaturation, which determines the size, structure, and properties of the nanoparticles. Interfacial tension plays a crucial role in the stabilization of nanoparticles. The presence of stabilizers or surfactants reduces interfacial energy, preventing coalescence.

Mixing processes in nanoprecipitation

The properties of synthesized nanoparticles depend on various factors, including mixing conditions, solvent composition, stabilizing agents, and physicochemical parameters.^{16,45} Efficient mixing is critical for achieving uniform particle size and morphology. The hydrodynamic flow regime, characterized by the Reynolds number (Re), governs the mixing dynamics. Laminar flow ($Re < 2100$) provides controlled mixing suitable for precision nanoprecipitation, while turbulent flow ($Re > 4000$) improves mixing rates but may introduce shear stress, affecting particle stability. Additionally, the Damköhler number (Da), which compares the timescales of reaction and mixing, is crucial in determining the uniformity of the synthesized nanoparticles. When ($Da < 1$), mixing dominates, ensuring homogeneity, whereas ($Da > 1$) can result in heterogeneities due to reaction dominance.⁴⁶

There are three primary nanoprecipitation techniques: batch, flash, and microfluidic methods, as illustrated in Fig. 2. In batch nanoprecipitation (BNP), the polymer solution and solvent are mixed rapidly, either through dropwise addition⁴⁷ or by injecting the entire polymer solution directly into the aqueous medium, as shown in Fig. 2(a).⁴⁸ In batch processes, the degree of mixing significantly influences the oversaturation dynamics. Rapid mixing reduces local concentration gradients, promoting uniform nucleation and growth. The shear forces generated during mixing also play a pivotal role, as they impact the particle size distribution and morphology. Controlled addition can also be achieved using a syringe pump or slow diffusion across a dialysis membrane. Several factors—such as the mixing method,⁴⁹ the polymer's molecular weight,⁵⁰ and the choice of organic solvent⁴⁵—influence the size and morphology of polymer nanoparticles (PNPs) synthesized *via* BNP. Pervyazko *et al.* demonstrated the fabrication of nanoparticles from solutions of poly(methyl methacrylate) and its copolymers. The particle characteristics strongly depended on the polymer's chemical structure and preparation method. In the studied cases, particle sizes ranged from 6–680 nm, with polydispersity indices (d_w/d_n) varying between 1.02 and 1.40. Their findings showed that nanoparticles of a desirable size range could be synthesized using solvent–nonsolvent methods. Fig. 3(a)–(f) shows the formation of nanoparticles by the batch process, achieved by varying the acetone-to-water solvent ratio.



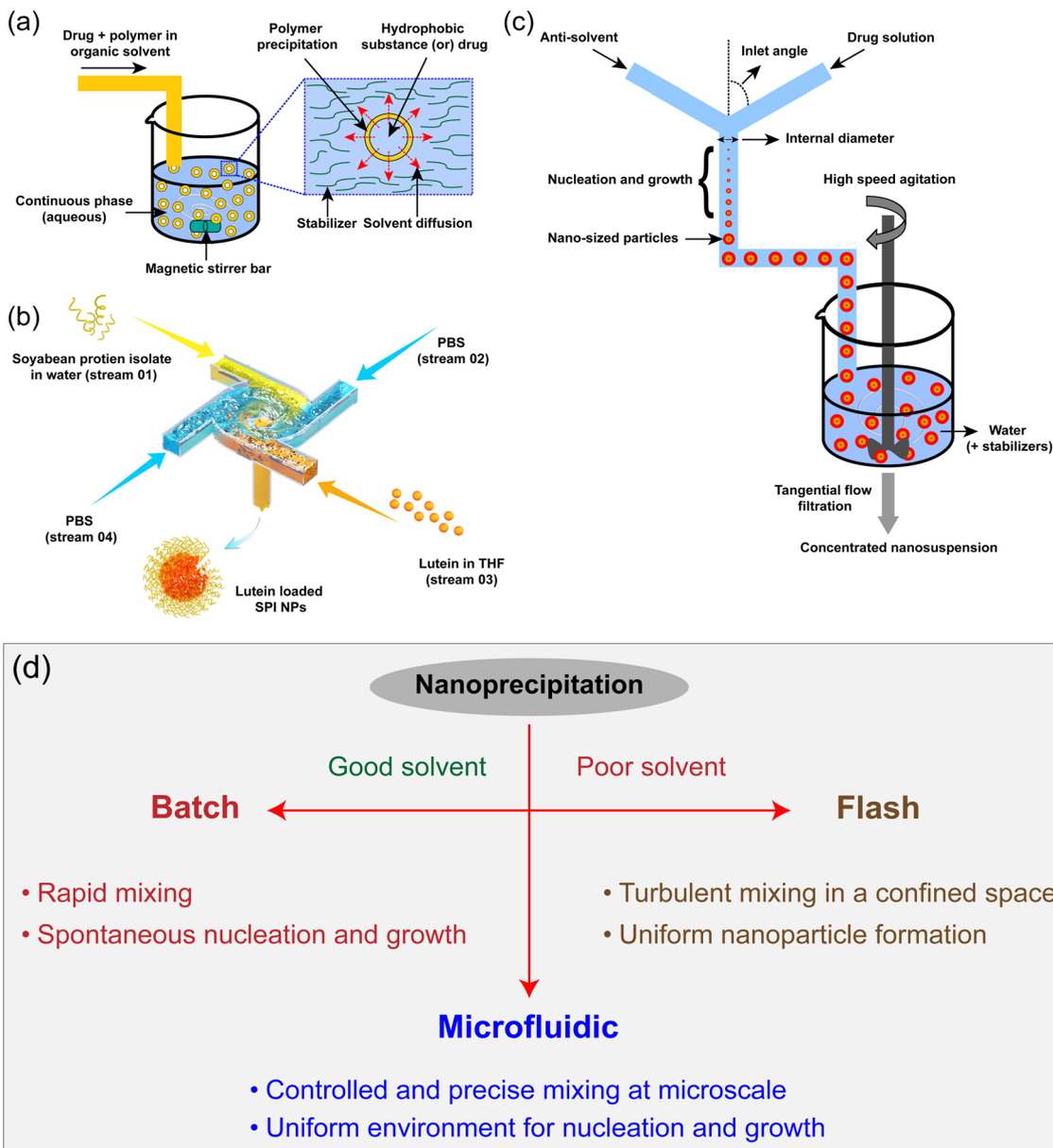


Fig. 2 Major types of mixing processes for nanoprecipitation. (a) Batch nanoprecipitation (BNP),⁴⁸ (b) flash nanoprecipitation (FNP),⁵¹ and (c) microfluidic nanoprecipitation (MNP),⁵² (d) principles of batch, flash, and microfluidic nanoprecipitation techniques.

Flash nanoprecipitation (FNP) utilizes a high-pressure injection pump to rapidly mix streams of organic solution and nonsolvent within a confined chamber, typically for milliseconds. The high shear forces generated during FNP facilitate rapid diffusion and mixing at the molecular level. The confinement within a narrow chamber ensures consistent mixing, minimizing spatial concentration gradients. The interplay between shear rates and mixing efficiency directly impacts particle size and uniformity, with higher shear rates promoting smaller, more homogeneous particles. This promotes the rapid formation of polymer colloids with specific morphologies and compositions.^{17,51,54,55} Wang *et al.* fabricated lutein-loaded nanoparticles (NPs) using FNP. In their process (Fig. 2(b)),

SPI was dissolved in water at a fixed concentration of 0.8 mg mL^{-1} , which represents the maximum solubility of SPI under the designed conditions. The properties of the kinetically controlled SPI NPs, including particle size, size distribution, drug loading efficiency, stability, and bioavailability, were investigated as well.⁵¹

Microfluidic nanoprecipitation (MNP) has been described as a simple method for drug nanosizing.^{56,57} Microfluidic systems rely on precise control of flow dynamics, which are dictated by the dimensions and geometry of the channels. The mixing efficiency in microfluidics is enhanced by chaotic advection and rapid diffusion-driven processes. By adjusting flow rates and channel configurations, it is possible to achieve highly



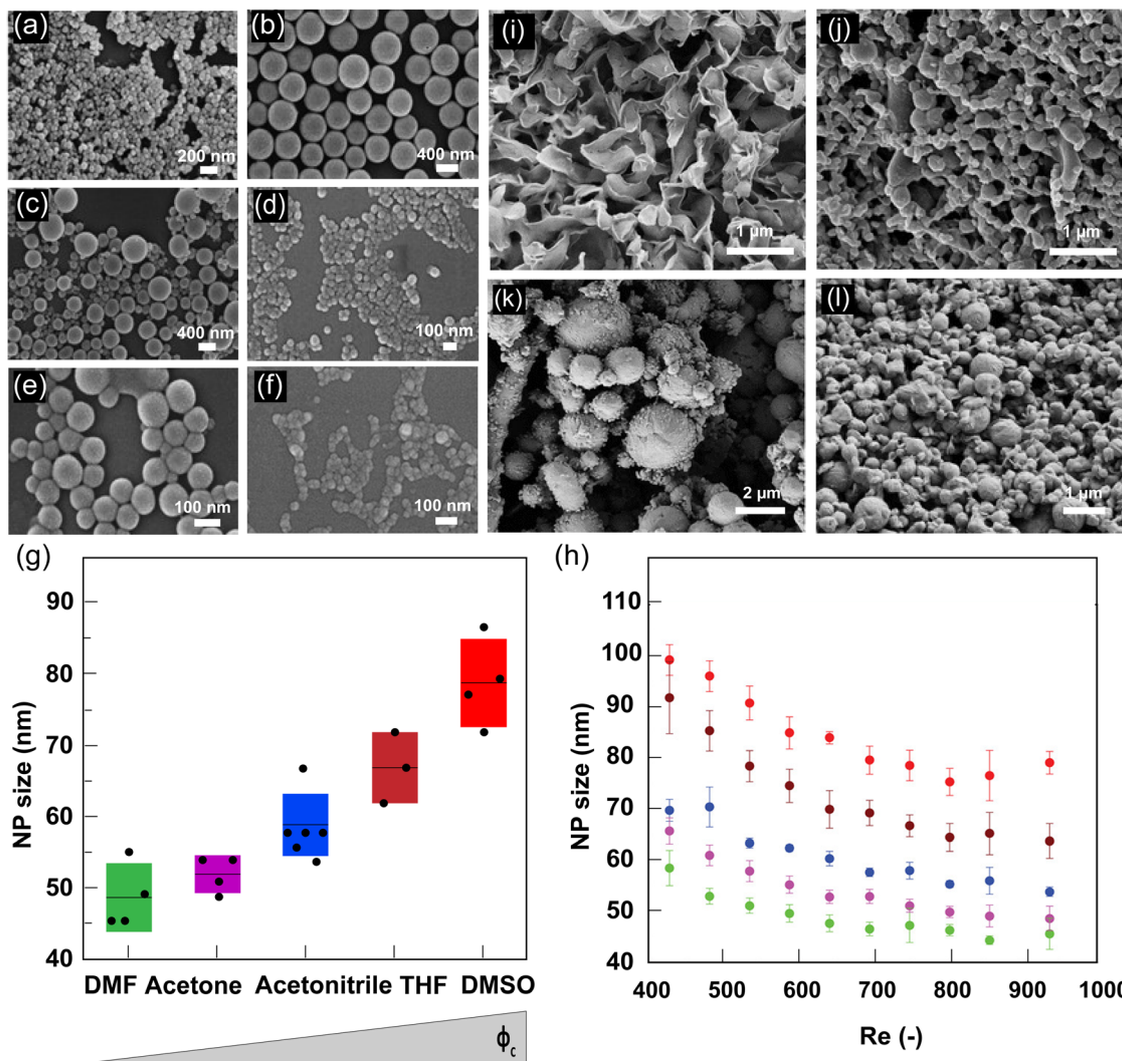


Fig. 3 Polymer nanoparticles (PNPs) prepared by nanoprecipitation. (a)–(f) SEM image of the PNPs prepared using two polymer poly(MMA-*stat*-pyMAA) (PMMA) and copolymer poly(MMA-*stat*-EA). (a)–(c) PMMA; (d)–(f) PMMA and copolymer. The mixing conditions are by dripping acetone to water (a) and (d), by dialysis of *N,N*-dimethylacetamide (c) and (f), or by dripping water to acetone (b) and (e).⁴⁹ Variation of the PNP size as a function of (a) solvents under constant mixing kinetics ($Re = 749$), and (b) Reynolds number ($Re = 400$ – 1000).⁴⁵ (i)–(l) Field emission scanning electron microscopic image representing the variation of PNP size on the final polymer concentrations (i) 0 mg mL^{-1} , (j) 1 mg mL^{-1} , (k) 2 mg mL^{-1} , and (l) 5 mg mL^{-1} from an initial polymer concentration of 5 mg mL^{-1} .⁵³

controlled nanoprecipitation, with particle sizes being inversely proportional to the flow rate due to enhanced nucleation at higher mixing velocities.⁵⁸ The results show that stable aqueous hydrocortisone NPs can be obtained using a bottom-up approach with microfluidic reactors. Particle size can be controlled by modifying the processing conditions and the design of the microfluidic reactors, such as internal diameters and inlet angles. Changes in flow rates were found to have a dominant effect on the size of the generated particles.^{44,59} The setup typically includes a central channel squeezed by two vertical channels, which allows for rapid diffusion-driven mixing (Fig. 2(c)).⁵² The dimensions of the channel, such as length, height, and structure, are key determinants of the properties of the particles in the microfluidic process.^{56,60,61}

Slater *et al.* reported the preparation of hydrophobic branched NPs *via* rapid nanoprecipitation. The resulting aqueous nanoparticle dispersions were robust and stable to dilution, solvent addition, sonication, and temperature changes. The addition of small amounts of NaCl led to nanoparticle destabilization, suggesting that electrostatic repulsion is a key factor in maintaining stability. The presence of NaCl likely screens surface charges, reducing repulsive interactions and promoting aggregation, thereby emphasizing the role of charge stabilization.⁶² Variation in particle size and fluorescent intensity at different $\text{H}_2\text{O}/\text{THF}$ ratios (Fig. 4(i) and (j)) for EDP NPs ranged from 28 nm to 55 nm, while BDP NPs ranged from 20 nm to 80 nm (Fig. 4(k) and (l)). In the following sections, we will discuss FNP and MNP in more detail.



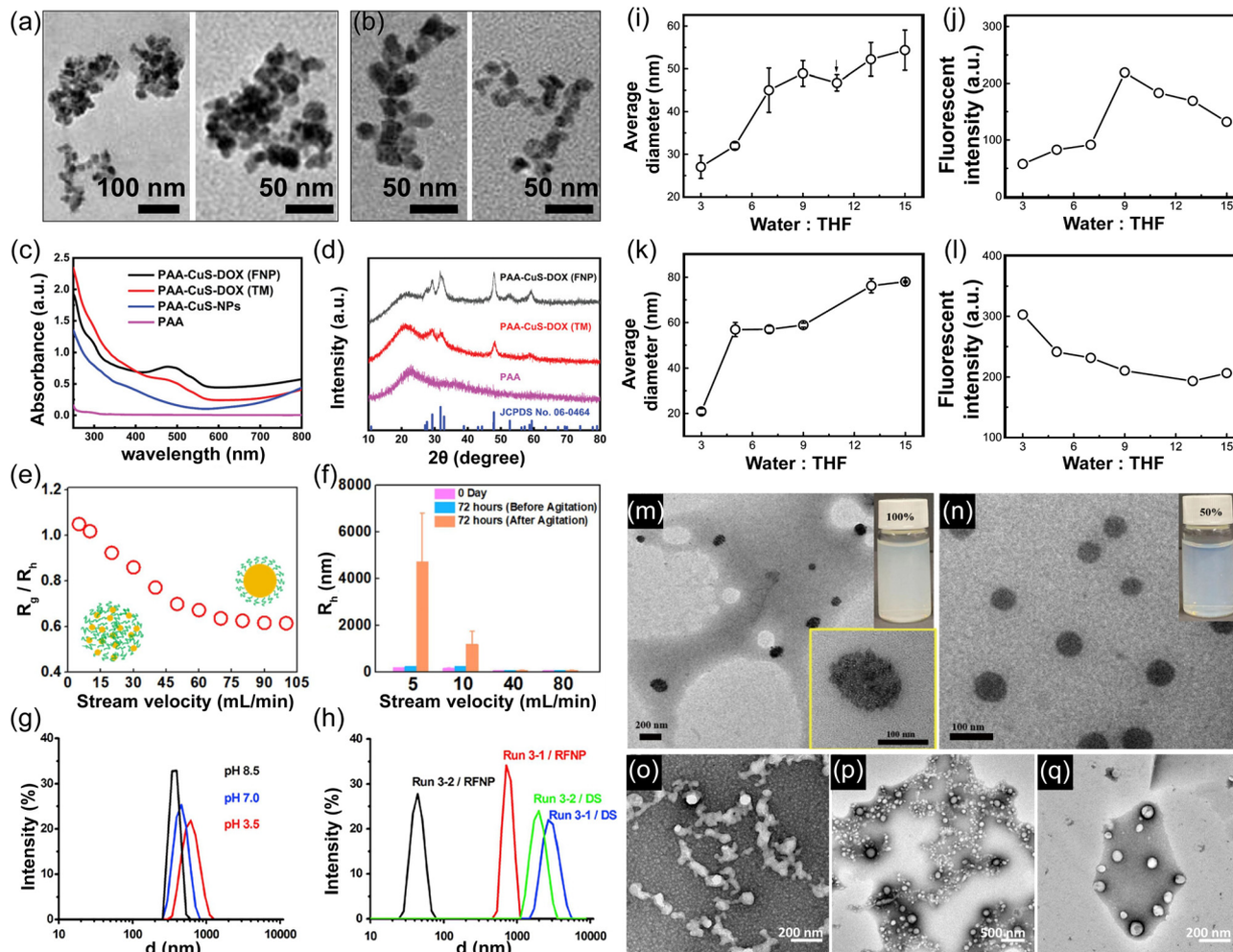


Fig. 4 TEM images of CuS-NPs-FNP with different AA/Cu ratios and corresponding hydrodynamic diameters of PAA-CuS-DOX NPs: (a) 1 (173 nm) and 2.5 (117 nm), (b) 5 (85 nm) and 20 (50 nm). Variation of hydrodynamic size distributions of CuS-NPs prepared *via* FNP and thermal method (TM) method (c) UV-vis spectra of NPs with and without drug (DOX) molecules prepared *via* FNP and TM. (d) XRD patterns of the NPs prepared *via* FNP and TM methods with and without DOX molecules.⁶³ (e) The ratio of the radius of gyration to the hydrodynamic radius (R_g/R_h) of the PNPs at stream velocities. (f) Stability of the prepared PNPs as a function of time.⁸ (g) and (h) Variation of PNP sizes at different solution pH.⁶⁴ RFNP: reactive nanoprecipitation. Run 3-1: without chitosan; run 3-2: with chitosan. DS: drip and stir method, in contrast to nanoprecipitation. (i)–(k) Variation of particle size and fluorescent intensity at different H₂O/THF ratio (i) and (j) for EDP NPs (k) and (l) BDP NPs.⁶⁵ (m) and (n) TEM images and the corresponding nanopesticide dispersion prepared using sophorolipids of mass concentrations (m) 100% acidic sophorolipids, and (n) 50% acidic sophorolipids and 50% lactic acid sophorolipids.⁶⁶ (o)–(q) TEM images of the PNPs obtained using block copolymers of different molecular weights.⁶⁷

Flash nanoprecipitation

Flash nanoprecipitation (FNP) enables the rapid, continuous production of polymer nanoparticles (PNPs) with smaller and more uniform particle sizes compared to batch nanoprecipitation (BNP).^{68,69} Both flow conditions and material properties jointly influence the size, morphology, and loading efficiency of PNPs, underscoring the importance of parameter control for precise nanoparticle synthesis.^{51,70}

Flow dynamics and mixing conditions, such as stream velocity, Reynolds number, and injection rate, are crucial in achieving desired particle characteristics. Wang *et al.* showed that increasing the Reynolds number over 1400 reduced the hydrodynamic size of PNPs from 122 nm to 80 nm, with minimal size change beyond this threshold.⁵¹ Conversely, Zhao *et al.*

observed that higher flow rates in a buffer solution gradually increased particle size, with a significant size jump at very high flow rates, demonstrating the need for careful control over flow to maintain size uniformity.⁷¹ Bhutto *et al.* investigated the internal structure of β -carotene-loaded protein PNPs, showing that increasing stream velocity resulted in densely packed core-shell structures, transitioning from laminar to turbulent flow. As stream velocity further increased, particle size stabilized, highlighting how flow dynamics affect internal particle organization and density, as shown in Fig. 4(e) and (f).⁸

Material properties, such as surfactant composition and polymer characteristics, also influence particle formation in FNP. Ma *et al.* examined the effect of surfactant ratio on lambda-cyhalothrin-loaded nanopesticides, finding that acidic



sophorolipids alone produced spindle-like particles while adding lactonic sophorolipids led to spherical particles due to altered hydrophobic interactions. This demonstrates how specific surfactant compositions can direct particle shape and stability (Fig. 4(m) and (n)⁶⁶). Jia *et al.* used FNP to create stable, low-toxicity copper sulfate nanoparticles for chemotherapy. They observed that as stream velocity increased from 6 to 30 mL min⁻¹, particle size decreased from 64 nm to 50 nm, stabilizing beyond that point. Additionally, Jia *et al.*⁶³ demonstrated the effect of varying the AA/CuS ratio, as shown in Fig. 4(a) and (b), where increasing the ratio from 1 (173 nm) and 2.5 (117 nm) in Fig. 4(a) to 5 (85 nm) and 20 (50 nm) in Fig. 4(b) led to a significant reduction in particle size. This is due to PAA acting as a capping agent, inhibiting growth through steric hindrance and electrostatic stabilization. At lower ratios, limited PAA results in larger nanoparticles, while higher concentrations restrict growth, leading to smaller sizes. Beyond an AA/CuS ratio of 10, the particle size remains unchanged, indicating surface saturation with PAA. In Fig. 4(c) and (d), Jia *et al.*⁶³ demonstrated the successful loading of DOX into the PAA-CuS-DOX NPs, as confirmed by the absorbance peak at 500 nm in the UV-vis spectra, corresponding to the characteristic peak of DOX. XRD analysis was also performed to characterize the crystal structure of the PAA-CuS-DOX NPs. Despite the large background signal from the amorphous PAA polymer, intense characteristic crystalline CuS peaks were clearly observed, confirming the presence of the CuS crystal structure.

Zhu *et al.* reported the production of lead(II) sulfate (PbSO₄) nanosuspension, with an average particle diameter of ~50 nm, *via in situ* reactive flash nanoprecipitation. Fig. 4(h) shows chitosan as a pH-sensitive surface stabilizer whose hydrophobicity can be tuned by varying its pH (Fig. 4(g) and (h)). By increasing the pH of the suspension, the chitosan/PbSO₄ nanoparticles rapidly aggregated and settled down. After filtration and drying, the particles were easily separated from water, with significantly reduced size enlargement due to Ostwald ripening and recrystallization.⁶⁴ Pustulka *et al.*⁶⁷ present TEM images in Fig. 4, depicting (o) 5k–5k PEG-*b*-PLA, (p) 5k–10k PEG-*b*-PLGA, and (q) 10k–10k PEG-*b*-PLGA nanoparticles, which demonstrated stability in suspension for at least 10 days.

The influence of drug loading on particle properties has been explored in several studies. Zhao *et al.* used an FNP setup with a multi-inlet vortex mixer to prepare curcumin-loaded zein nanoparticles, finding that higher drug concentrations initially increased particle size and yield, with particles exhibiting narrow size distribution at higher drug levels.⁷¹ Wang *et al.* similarly found that at low drug-to-polymer ratios, particle size initially decreased before increasing at higher ratios, indicating an optimal ratio range for controlling particle dimensions in drug-loaded systems.⁵¹

Nanoprecipitation in microfluidics

Microfluidic nanoprecipitation (MNP) allows precise control over polymer nanoparticle characteristics, including size, morphology,

and surface properties.^{61,72} The microfluidic setup uses ultra-low volumes (pico- to nanoliters) to regulate fluid flow, reduce energy consumption, and lower costs. Various configurations—such as hydrodynamic flow focusing, staggered herringbone micromixers, bifurcating mixers, T-junction mixers, and baffle mixers—enhance control over the mixing environment by tailoring flow in narrow channels.²⁵ These configurations are particularly advantageous for precise formulations, even for expensive drugs, although the small channel sizes can limit scalability.

Flow dynamics and mixing conditions are critical in MNP. The physics of surface and interface interactions plays a pivotal role in MNP. At the nanoscale, the high surface-to-volume ratio amplifies interfacial forces, significantly impacting particle formation and stabilization. Surface tension (γ) governs the interfacial energy, while the interfacial curvature determines the Laplace pressure, influencing nanoparticle size and shape. The balance between cohesive forces (within the liquid) and adhesive forces (between liquid and solute) drives nucleation and growth processes.⁷³

In the confined microchannels of MNP, the interplay of diffusion and interfacial tension creates a uniform solute distribution, enhancing the nucleation rate. Additionally, wetting properties, characterized by the contact angle (θ), influence solvent and anti-solvent interactions, affecting particle morphology. Hydrophilic channel walls promote spreading and mixing, while hydrophobic surfaces may induce localized aggregation due to poor wetting. The wettability of microfluidic channel walls significantly influences nanoprecipitation by affecting fluid mixing, solute diffusion, and particle formation. Hydrophilic surfaces ($\theta < 90^\circ$) enhance solvent spreading, promote uniform solute distribution, and improve mixing efficiency, resulting in smaller and more homogeneous nanoparticles.⁷⁴ Conversely, hydrophobic surfaces ($\theta > 90^\circ$) can lead to poor wetting, localized aggregation, and increased particle polydispersity due to inefficient mixing.⁵⁸ Furthermore, excessive hydrophobicity may cause nanoparticle adhesion to channel walls, leading to fouling and flow disturbances.⁷⁵ Optimizing wettability through surface modifications, such as plasma treatment or chemical coatings, can enhance process stability and ensure consistent nanoparticle synthesis.⁷⁶ Optimizing surface energy and flow conditions is crucial to ensuring uniform particle formation. Increasing the flow rate ratio between the solvent and anti-solvent promotes diffusion, generating more nucleation sites and resulting in smaller, more uniform PNPs.⁶¹ Studies by Heshmatnezhad *et al.* and Chiesa *et al.* confirmed that higher total flow rates reduce particle size due to intensified shear forces, causing oversaturation and greater nucleation.^{77–80} However, excessive flow rates can induce particle aggregation, leading to an increase in particle size at channel junctions, as demonstrated in the TEM images and DLS plot (Fig. 5(a)–(f)).⁸¹ The Reynolds number (Re) also influences particle size; Liu *et al.* found that increasing Re enhanced turbulence, improving mixing until particle size stabilized. At lower drug-to-polymer ratios, thinner polymer shells formed around drug cores, while higher Re led



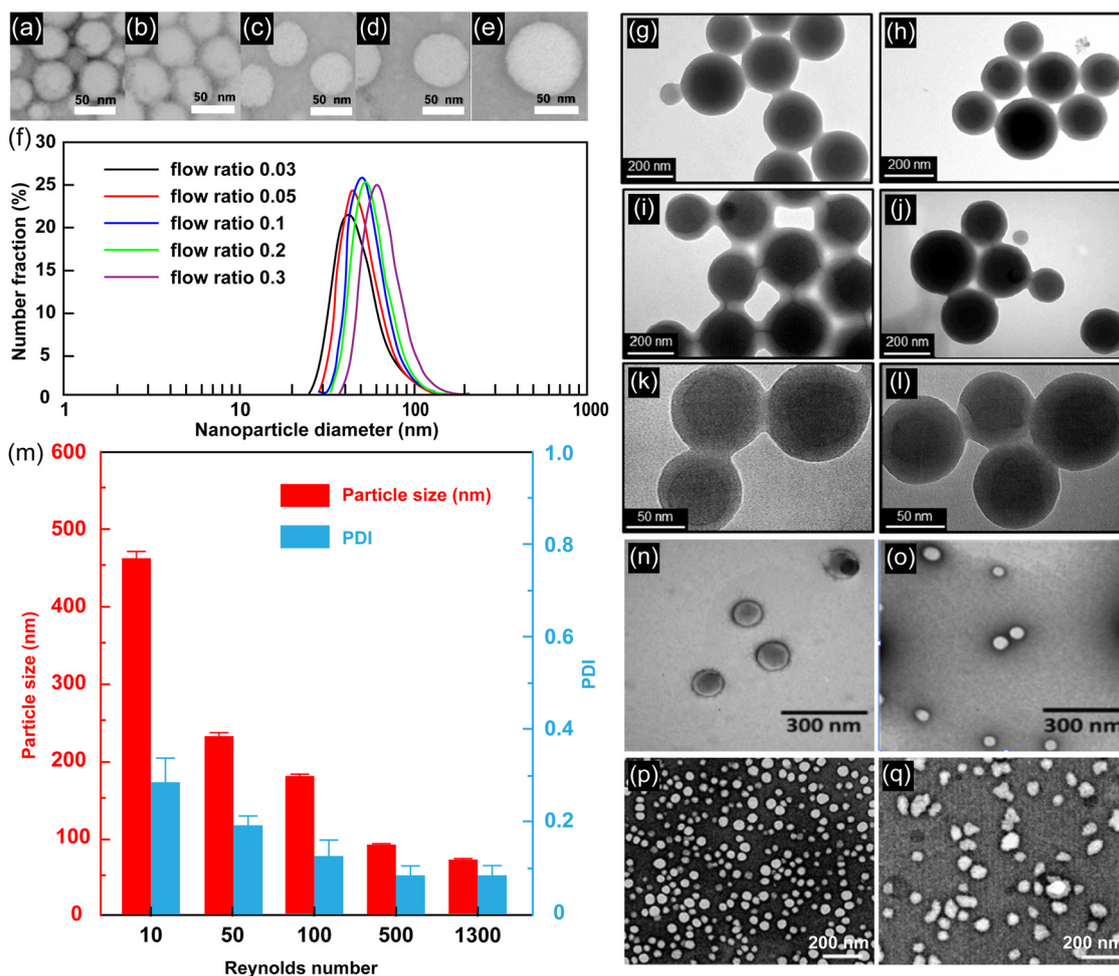


Fig. 5 (a)–(e) TEM images of PNPs prepared at flow rate ratios. The PNPs of mPEG-PLGA from the flow ratios of (a) 0.03, (b) 0.05, (c) 0.1, (d) 0.2, (e) 0.3. (f) Corresponding dynamic light scattering plots of PNPs.⁸¹ (g)–(l) TEM images of the core-shell PNP synthesized by varying drug-polymer weight ratio and Reynolds number. (g) and (h) Drug-polymer weight ratio of 1:1 and Re of 100. (i) and (j) Drug-polymer weight ratio of 2:1 and Re of 100. (k) and (l) Drug-polymer weight ratio of 1:1 and Re 1300. (m) Particle size distribution and polydispersive index (PDI) as a function of Reynolds number.⁸² (n) and (o) TEM images of PNPs prepared under varying flow rate ratios of (n) 1:1 and (o) 10:1.⁵⁹ (p) and (q) TEM images of PNPs as a function of drug (curcumin) to polymer ratio at (p) 0.5 and (q) 10.⁵

to smaller core-shell particles across different charged polymers (Fig. 5(g)–(m)).⁸²

The material properties of microfluidic channels play a pivotal role in determining the outcomes of MNP by influencing fluid dynamics, mixing efficiency, and nanoparticle characteristics. Key material attributes such as wettability, surface charge, and mechanical properties directly impact the nucleation and growth processes of nanoparticles. The wettability of channel materials, defined by their surface energy, affects fluid flow patterns and mixing behavior within microchannels. Hydrophilic materials, characterized by high surface energy, promote effective solvent spreading and facilitate uniform solute distribution. This leads to enhanced mixing efficiency and results in the formation of smaller, more homogeneous nanoparticles. In contrast, hydrophobic materials with low surface energy can cause inadequate wetting, leading to poor mixing and localized solute concentration gradients. These conditions may result in the formation of larger, polydisperse

nanoparticles due to uneven nucleation rates. For instance, the use of PDMS in microfluidic devices has been shown to influence nanoparticle synthesis outcomes due to its inherent hydrophobicity.⁸³

The surface charge of channel materials influences electrostatic interactions between the channel walls and the solute particles. Materials with surface charges that are compatible with the solute can prevent unwanted adsorption of nanoparticles onto the channel walls, thereby reducing fouling and ensuring a stable nanoprecipitation process. Conversely, incompatible surface charges can lead to particle adhesion, affecting the yield and quality of the nanoparticles produced. The chemical compatibility between the channel material and the solvents used is also crucial, as interactions at the interface can alter the physicochemical properties of the forming nanoparticles. Additionally, the mechanical strength and thermal conductivity of channel materials determine their ability to withstand operational conditions without deforming



or degrading. Materials with high mechanical strength maintain structural integrity under various flow rates and pressures, ensuring consistent nanoparticle synthesis. Thermal properties influence heat dissipation within the microchannels, affecting solvent evaporation rates and, consequently, the supersaturation levels critical for controlled nanoprecipitation. Materials like PDMS, commonly used in microfluidic devices, offer flexibility and ease of fabrication but may have limitations in mechanical and thermal stability under certain conditions.⁵⁸ To optimize MNP outcomes, surface modification techniques are employed to tailor the material properties of microfluidic channels. Methods such as plasma treatment, chemical grafting, or coating with hydrophilic polymers can enhance surface wettability, improving fluid mixing and nanoparticle uniformity. These modifications can also introduce desired surface charges or functional groups, reducing particle adhesion and fouling. For example, treating PDMS surfaces to increase hydrophilicity has been shown to improve nanoparticle synthesis by promoting better mixing and reducing aggregation.⁸³

Precise control over flow dynamics and channel design allows for fine-tuning of the nanoprecipitation process. The ratio of flow rates between solvent and anti-solvent governs mixing efficiency and interfacial area generation, both critical for particle nucleation. For instance, high flow rates create thinner diffusion layers, reducing interfacial resistance and promoting rapid saturation of solute molecules. At the same time, the *Re* dictates whether flow remains laminar or transitions to turbulent, with laminar flow providing the most reproducible conditions for nanoprecipitation.⁸⁴ Channel geometry, including T-junctions, staggered herringbone micromixers, and bifurcating designs, determines shear rates and residence times. Shear rates at the interface influence interfacial stability, where excessive shear can disrupt particle formation or cause aggregation.^{85,86} Scaling up these designs while maintaining optimal mixing conditions remains a challenge due to the complex interplay of surface forces and flow dynamics. Fig. 5(n) and (o) illustrates the effect of flow rates on NP size, showing that increasing the flow rate reduces NP size.⁵⁹ Silverman *et al.* demonstrated that increasing drug-to-polymer ratios in poly(caprolactone)-*block*-poly(ethylene glycol) NPs primarily affected polydispersity. At the same time, particle size changed only slightly due to curcumin's plasticizing effect, as shown in TEM images (Fig. 5(p) and (q)).⁵ Ma *et al.* showed that surfactant ratios affect particle morphology in nanopesticides.⁶⁶ The chemical composition of surfactants and stabilizers significantly influences interfacial properties during MNP. Surfactants reduce interfacial tension, stabilizing newly formed nanoparticles by preventing coalescence or Ostwald ripening. For example, ionic surfactants provide charge stabilization, while non-ionic surfactants rely on steric hindrance to maintain particle dispersion. The choice of surfactant must balance these stabilization mechanisms to achieve desired particle properties, particularly under varying flow rates and solvent compositions. Additionally, interfacial rheology—describing the deformation and flow of the interface—determines the stability of emulsions formed during nanoprecipitation. For instance, interfacial elasticity resists deformations under shear,

preventing coalescence in high-shear environments such as those in microfluidic channels.

FNP and MNP both enable controlled production of PNPs but differ significantly in their methodologies and applications. FNP is characterized by rapid mixing in a confined space, producing particles with narrow size distributions and high loading efficiency, particularly suitable for hydrophobic drugs. However, FNP has limitations in processing water-soluble biomolecules and often requires high-pressure pumps, which increase operational complexity. In contrast, MNP leverages ultra-low volumes and precise control over flow rates within microchannels, allowing for highly reproducible particle sizes and morphologies through laminar flow and diffusion-driven mixing. MNP excels in its fine-tuning capabilities, accommodating various solvent compositions and complex surfactant systems, making it ideal for applications requiring small-scale, precise formulations. While MNP offers high precision and low energy consumption, it faces challenges in scalability due to the small channel size. FNP, by contrast, can more readily accommodate larger production scales, albeit with less precise control over certain particle properties. Both methods are versatile tools in nanoparticle synthesis, with FNP favored for rapid production and MNP for applications demanding intricate control over nanoparticle characteristics. From an interfacial physics perspective, FNP operates under conditions of high interfacial tension and rapid mixing, which can lead to kinetic control over particle size but may limit precision. In contrast, MNP emphasizes interfacial control through diffusion-driven mixing, allowing finer tuning of particle characteristics.³⁶ The lower interfacial tension achieved in MNP setups, often facilitated by surfactants or solvent composition, supports the formation of more uniform nanoparticles. For instance, in MNP, the relationship between surface energy and flow velocity can be optimized to achieve a balance between nucleation and growth, ensuring monodispersity. Meanwhile, FNP relies on intense mixing to overcome interfacial resistance, often resulting in broader size distributions.⁸⁷

Other mixing techniques for controlling nanoprecipitation

Recently, new techniques have emerged to enhance the mixing between solvent and anti-solvent phases in nanoprecipitation processes.⁸⁸ One such method, electrohydrodynamic (ED) mixing, achieves turbulent mixing through an applied voltage that triggers the nanoprecipitation process.^{89,90} Unlike flash nanoprecipitation (FNP), which relies on high flow rates of solvent and anti-solvent to induce turbulence and achieve homogeneous mixing, ED mixing uses an electric field to create similar mixing conditions at significantly lower flow rates. The applied voltage generates an electric field that promotes turbulent mixing, resulting in nanoparticle formation with precise control over particle size.

Microfluidic platforms, such as Y-junction mixers, have demonstrated precise control over mixing dynamics, leading



to uniform NPs synthesis with high reproducibility.⁹¹ The application of staggered herringbone micromixers further enhances mass transfer efficiency, facilitating the synthesis of metal–organic frameworks (MOFs) with tunable porosity and controlled drug release.⁹² Beyond microfluidic devices, high-speed homogenization remains an effective approach for achieving rapid solvent-exchange-driven nanoprecipitation, yielding lipid–polymer hybrid nanoparticles with enhanced colloidal stability and encapsulation efficiency.⁵⁸ Additionally, ultrasonic-assisted nanoprecipitation has been employed for the synthesis of bioactive metal–organic frameworks, enabling targeted and responsive drug delivery applications.⁹³ Electrohydrodynamic mixing techniques, including jetting-based nanoprecipitation, have further optimized particle formation by leveraging charge-induced forces, leading to highly monodisperse NPs for biomedical applications. The potential of these techniques is evident in diverse fields, from drug delivery to energy storage, where precise NPs control is critical.⁹⁴ These advancements in nanoprecipitation mixing techniques highlight the growing role of controlled microenvironment engineering in NPs synthesis, offering new opportunities for enhanced functionality and efficiency.

Membrane nanoprecipitation is another method for producing polymer-based nanoparticles, where mixing occurs directly at the pores of a membrane.⁹⁵ In this method, an organic solvent and anti-solvent are introduced on opposite sides of a membrane with defined pore structures. Mixing occurs at these pores, enabling nanoparticle formation by regulating solvent diffusion. For example, the organic solvent may occupy the shell side of the membrane, mixing with the anti-solvent in the lumen side, or *vice versa*, driven by a pressure gradient. Membrane properties, particularly pore size and wettability, significantly influence particle size and polydispersity. Despite its precision, membrane nanoprecipitation faces challenges in scaling up due to the limited availability of suitable membranes with defined pore characteristics.

Electrospray nanoprecipitation combines solvent-shifting techniques with electrospray to synthesize polymer nanoparticles (PNPs).^{96,97} In this process, the polymer solution is pumped through a nozzle connected to a high-voltage power supply, generating charged droplets that are subsequently mixed with a non-static aqueous phase under stirring. The electric field controls droplet size, facilitating rapid solvent evaporation, oversaturation, and nanoparticle formation. This technique produces monodisperse, surfactant-free nanoparticles, allowing for precise control over particle characteristics.

Techniques such as UV-vis absorption, Raman spectroscopy, and fluorescence can be employed in microfluidic channels to monitor the particle formation process in real-time. By coupling these spectroscopic methods with machine learning algorithms, we can dynamically adjust parameters like flow rates, solvent composition, and temperature to maintain consistent particle size and morphology. Additionally, dynamic light scattering (DLS) and *in situ* light scattering or particle tracking microscopy offer real-time monitoring of particle size distribution and growth dynamics. Temperature control and monitoring within

the microfluidic system ensures consistent reaction kinetics, while solvent composition monitoring provides further precision in maintaining optimal conditions for nanoprecipitation. This integration enhances reproducibility and scalability, providing a precise and adaptable framework for controlling nanoprecipitation and producing nanoparticles with tailored properties.

For systems involving metal-based nanoparticle synthesis, electrochemical techniques such as cyclic voltammetry (CV) or electrochemical impedance spectroscopy (EIS) can be employed to monitor ion reduction and redox processes. These methods are particularly effective in controlling the nucleation and growth of metal nanoparticles, enabling real-time adjustments to ion concentrations, pH, and other critical parameters in metal nanoprecipitation processes.

Physicochemical properties of polymers and solvents in nanoprecipitation

The interaction between the organic phase, aqueous phase, and the solute molecules plays a crucial role in determining the size and properties of polymer nanoparticles (PNPs) in nanoprecipitation.¹⁶ This can be understood by considering the diffusivity of the organic phase in the aqueous phase, which influences the distribution of solute molecules. A decrease in the diffusion rates of the organic and aqueous phases results in an increase in particle size. Moreover, as the affinity or interaction between the aqueous and organic phases increases, the resulting PNPs tend to have smaller sizes.^{98–100}

Additionally, the interaction between the organic phase and the solute molecules determines the quality of the solvent for a given polymer, whether it is a good or poor solvent. In the case of a good solvent for a polymer, the resulting particle size tends to be larger due to the extended conformation of the solute molecules. Conversely, in a poor solvent, the solute molecules collapse, resulting in smaller particle sizes. Since these interactions vary for different solvents, the size and properties of the PNPs also change with variations in the organic phase.

This section will discuss the effects of polymer characteristics (such as concentration, molecular weight, and degree of hydrophobicity) and solvents on the size and properties of nanomaterials. The variation in particle properties with changes in solution parameters in nanoprecipitation techniques is summarized in Table 1.

The interactions between the organic phase, aqueous phase, and solute molecules are crucial in determining the size and characteristics of polymer nanoparticles (PNPs) in nanoprecipitation.¹⁶ The diffusivity between phases influences the distribution of solute molecules: slower diffusion typically results in larger particles, while stronger affinity between the phases tends to reduce particle size. Furthermore, the interaction between the organic phase and solute molecules determines the solvent quality for a given polymer. In the case of good solvents, which extend the polymer chains, larger particles are typically



Table 1 Comparison of mixing techniques: batch, flash, and microfluidic

Mixing technique	Advantages	Disadvantages	Ref.
Batch	<ul style="list-style-type: none"> – Simple setup and easy to scale up – Suitable for large volumes – Cost-effective 	<ul style="list-style-type: none"> – Longer mixing times – Poor control over mixing conditions – Higher batch-to-batch variability 	101 and 102
Flash	<ul style="list-style-type: none"> – Rapid mixing – Suitable for high-throughput processes – Reduces aggregation in nanoparticle synthesis 	<ul style="list-style-type: none"> – Requires precise control of flow rates – Limited to specific reaction conditions – Equipment may be expensive 	67 and 103
Microfluidic	<ul style="list-style-type: none"> – Excellent control over mixing parameters – High reproducibility – Suitable for small volumes and lab-on-a-chip applications 	<ul style="list-style-type: none"> – Low throughput – Complex fabrication and operation – Higher initial costs 	58 and 104

Table 2 Summary of particle size variation under different conditions

Condition	ACN/water flow rate ratios	Particle size (nm)	DLS mean diameter (nm)	PDI
Fig. 5(a)	0.03	47 ± 2.5	49 ± 2.9	0.131
Fig. 5(b)	0.05	48 ± 2.0	52 ± 1.7	0.097
Fig. 5(c)	0.1	54 ± 1.9	56 ± 2.6	0.080
Fig. 5(d)	0.2	62 ± 2.5	59 ± 1.2	0.069
Fig. 5(e)	0.3	74 ± 2.2	71 ± 1.2	0.092

formed, whereas poor solvents cause the polymer chains to collapse, resulting in smaller particles (Tables 2 and 3).

Polymer properties, such as concentration, molecular weight, and architecture, also significantly affect PNP size and stability. For instance, Slater *et al.* observed that increasing polymer concentration led to a reduction in PNP size and polydispersity for vinyl polymers, while polysaccharide nanoparticles exhibited the opposite trend, with particle size increasing as polymer concentration increased.^{62,112} Additionally, J. H. Lee and colleagues found that higher lignin concentrations resulted in larger PNPs with increased polydispersity and surface charge.⁵⁰ Polymer architecture also influences nanoparticle stability. For example, Slater *et al.* found that branched polymers produced more stable PNPs than linear polymers, which tended to precipitate after synthesis.⁶² Furthermore, Pustulka *et al.* reported that higher polymer hydrophobicity, measured by the water–octanol partition coefficient, enhanced PNP stability. Specifically, coefficients above 7 led to more stable particles that were resistant to rapid aggregation.⁶⁷

Solvent choice plays a critical role in controlling PNP properties, influencing size, stability, and morphology. Rao and Geckeler¹¹³ examined various solvent-based methods, highlighting how solvent evaporation and nanoprecipitation impact particle size through solvent diffusion rates and polymer–solvent interactions. Dwivedi *et al.*¹¹⁴ explored nanoprecipitation and emphasized that the choice of solvent and its miscibility with water dictate the final nanoparticle size, with polar solvents like acetone yielding smaller particles due to rapid diffusion into the aqueous phase. Huang and Zhang¹⁰⁵ systematically studied factors affecting PLGA nanoparticle size and found that solvent diffusion coefficient strongly dictates size distribution, with solvents like acetonitrile producing finer nanoparticles compared to acetone or THF.

Additionally, Aubry *et al.* demonstrated that lower polymer concentrations in PMMA, combined with a high aqueous phase

volume, resulted in PNPs with narrow size distributions. However, higher polymer concentrations led to the formation of a mixture of micro- and nanoparticles, in line with the Smoluchowski kinetic model.¹¹⁵ Similarly, Bovone *et al.* observed that solvent type influences PNP growth dynamics. Initial dynamic aggregates, formed through polymer exchange, stabilize once the solvent-specific water fraction is reached, ultimately determining the final particle size.⁴⁵ Solvent effects have also been explored in microfluidic nanoprecipitation (MNP) systems; Donno *et al.* found that increasing polymer molecular weight and flow rate ratio in MNP reduced PNP size and increased surface charge, with larger particles forming in the presence of surfactants at higher viscosities.¹¹⁶

Surfactants and other additives further modulate PNP size and morphology by controlling particle interactions during formation. Luque-Alcaraz *et al.* studied chitosan-based PNPs and found that surfactant presence, polymer concentration, and solvent-to-non-solvent ratios significantly influenced particle characteristics. For example, the presence of Tween 80 as a surfactant reduced particle size, while the absence of surfactant led to larger, more irregular PNPs due to coalescence.¹¹⁷ Heshmatnezhad *et al.* showed that combining surfactants like PVA and Tween 80 improved particle uniformity and prevented aggregation, resulting in smoother, smaller, and more stable PNPs. In contrast, PNPs formed without surfactants were larger, with rougher morphology and non-uniform distributions.⁷⁷

Latest developments in green solvents and their impact on nanoprecipitation processes

Nanoprecipitation, a widely adopted technique for nanomaterial synthesis, depends on the rapid mixing of solvents and non-solvents to trigger supersaturation and nanoparticle formation. Traditionally, organic solvents like acetone, THF, and DMSO have dominated due to their miscibility with water and ability to dissolve diverse precursors. However, their toxicity, volatility, and environmental footprint have driven a shift toward greener alternatives, reshaping nanoprecipitation in recent years. The integration of green solvents into nanoprecipitation processes has garnered significant attention due to their potential to enhance sustainability and reduce environmental impact in



Table 3 BNP: effects of nanoprecipitation conditions on the size and morphology of the PNPs

Polymer	Organic/aqueous phase	Parameter varied	Size	Morphology
PEG- <i>b</i> -PLA, PEG- <i>b</i> -PLGA, or PEG- <i>b</i> -PCL	DMF, acetone, acetonitrile, THF, or DMSO/water	Organic solvents	50–100 nm	Spherical ⁴⁵
Poly(vinyl alcohol)	Methanol, ethanol, propanol, <i>tert</i> -butanol/water	Polymer concentrations, interaction parameters	50–300 nm	Spherical ¹⁶
p(HPMA50-EGDMA)	Acetone/water	Polymer structure (linear and branched), temperature, volume of organic phase, NaCl	50–800 nm	Spherical ⁶²
Poly(lactic- <i>co</i> -glycolic acid)	Acetonitrile, acetone and tetrahydrofuran (THF)/PVA aqueous solution	Polymer concentration, organic solvent, ionic strength of aqueous phase and temperature	80–3500 nm	Spherical ¹⁰⁵
Polystyrene	Acetone, chloroform, tetrahydrofuran and acetonitrile/Tween-40, Pluronic F-68	Polymer, surfactant, non-solvent	100 nm–3 μm	Spherical ¹⁰⁶
Cellulose acetates	Acetone/water	Concentrations of cellulose acetate	160–400 nm	Spherical and bean-shaped particles ¹⁰⁷

FNP: effects of nanoprecipitation conditions on the size and morphology of the PNPs.

Polymer	Organic/aqueous phase	Parameter varied	Size	Morphology
Polystyrene	Tetrahydro-furan/water	Time, molecular weight, solution concentrations, stirring rate, solvent/non-solvent ratio	60–200 nm	Spherical ¹⁵
Poly(ethylene glycol)- <i>block</i> -poly(ε-caprolactone)	Tetrahydro-furan and dimethyl sulfoxide (DMSO)/water	Solvent/non-solvent ratio, stream velocity	20–80 nm	Spherical ⁶⁵
Poly(3-hydroxybutyrate- <i>co</i> -3-hydroxyvalerate)	Dichloro-methane/SDS and PVA solutions	Polymer and surfactant concentrations	100–600 nm	Spherical ⁵³
PEG- <i>b</i> -PLGA, PS(10k)- <i>b</i> -PEG, PEG- <i>b</i> -PLA	Tetrahydro-furan/water	Nanoparticle concentrations	60–180 nm	Spherical ⁶⁷
PEO- <i>b</i> -PS, PEG- <i>b</i> -PCL	Tetrahydro-furan/water	Reynolds number	25–500 nm	Spherical ¹⁰⁸
PS- <i>b</i> -PI	Tetrahydro-furan/water	Polymer concentrations, chain length	30–300 nm	Spherical particles with concentric shells or a disordered lamellar ¹⁰⁹

MNP: effects of nanoprecipitation conditions on the size and morphology of the PNPs.

Polymer	Organic/aqueous phase	Parameter varied	Size	Morphology
Poly(methyl methacrylate) (PMMA)	Tetrahydrofuran/water	Polymer concentrations, volume flow rate ratio between water and the polymer solution	100–200 nm	Spherical ⁷²
Poly(lactic- <i>co</i> -glycolic acid)	Dimethylformamide/water	Micro-channel geometry, aspect ratio	50–300 nm	Spherical ⁵⁹
Pluronic F127	Tetrahydrofuran/water	Drug to polymer concentrations, flow rate, mixing time	70–200 nm	Spherical ¹¹⁰
PLGA	Dimethylformamide (DMF) and tetrafluoroethylene/water	Flow rate	50–250 nm	Spherical ⁶⁰
PLGA-lipid-NPs	Tetrafluoroethylene (TFE) and 0.7 mL dimethylformamide (DMF)/water	Flow rate	60–90 nm	Spherical ¹¹¹
Polycaprolactone	THF and DMF/PVA solution	Flow rate ratio and total flow rate	100–300 nm	Spherical ⁷⁷

nanomaterial synthesis. These eco-friendly solvents not only minimize the use of hazardous chemicals but also offer unique physicochemical properties that can influence the characteristics of the resulting nanoparticles.

Deep eutectic solvents (DESSs) have emerged as promising alternatives to traditional organic solvents in nanoprecipitation. Comprising a mixture of hydrogen bond donors and acceptors, DESSs exhibit low volatility, biodegradability, and tunable solubility parameters. Their unique properties facilitate the efficient extraction of bioactive compounds from natural sources, which can subsequently act as reducing and stabilizing agents in nanoparticle synthesis. For instance, Vorobyova *et al.* demonstrated the use of a DES-based plant extract for the biosynthesis of

silver nanoparticles, highlighting the solvent's role in enhancing nanoparticle stability and antibacterial efficacy.¹¹⁸ Similarly, ionic liquids (ILs) particularly those based on imidazolium cations, have been extensively studied for their role in stabilizing nanomaterials during synthesis. Their negligible vapor pressure, thermal stability, and ability to dissolve a wide range of compounds make them suitable candidates for green nanoprecipitation processes. ILs can act as both solvents and stabilizing agents, influencing nanoparticle size, morphology, and dispersion stability. Tshemese *et al.* reviewed the application of imidazolium-based ILs in nanomaterial stabilization, emphasizing their potential to replace conventional, more hazardous solvents.¹¹⁹



The utilization of solvents derived from renewable biological sources, such as plant extracts and natural surfactants, aligns with the principles of green chemistry. These solvents often contain bioactive compounds capable of reducing metal ions and stabilizing nanoparticles. For example, sophorolipids, which are glycolipid biosurfactants, have been employed in flash nanoprecipitation techniques to construct nanodelivery systems. Their natural origin and biodegradability make them attractive alternatives to synthetic surfactants in nanoparticle formulation.⁶⁶ The incorporation of green solvents into nanoprecipitation processes offers several advantages such as (i) environmental sustainability: green solvents reduce the reliance on toxic organic solvents, thereby decreasing environmental pollution and health hazards,¹²⁰ (ii) enhanced biocompatibility: nanoparticles synthesized using bio-based solvents often exhibit improved compatibility for biomedical applications due to the absence of harmful residues,¹²¹ and (iii) controlled nanoparticle characteristics: the unique properties of green solvents, such as viscosity and polarity, can be tailored to control nanoparticle size, morphology, and dispersion, which are critical parameters in various applications.¹²² The adoption of green solvents in nanoprecipitation processes represents a significant advancement in the sustainable synthesis of nanomaterials. Ongoing research in this area is expected to further optimize these processes, leading to environmentally friendly and efficient production of nanoparticles for diverse applications.

Advanced nanostructures synthesized via nanoprecipitation

Nanoprecipitation is a highly advantageous technique for synthesizing drug delivery nanocarriers, making it a preferred method in pharmaceutical applications. Its simplicity, scalability, and ability to produce nanoparticles with controlled size and high drug loading make it suitable for encapsulating a wide range of active pharmaceutical ingredients, from small hydrophobic drugs to complex biomolecules. Additionally, nanoprecipitation is a solvent-efficient process with minimal energy requirements, which lowers production costs and reduces environmental impact. However, the technique does have limitations: it is primarily effective for hydrophobic drugs, as hydrophilic molecules tend to diffuse rapidly into the aqueous phase, leading to poor encapsulation efficiency and rapid drug leakage.¹²³ To address this challenge, various strategies have been explored. One common approach is to modify the polymer matrix by introducing amphiphilic copolymers, such as PEG, which enhances the solubility and interaction of hydrophilic drugs with the nanoparticle core.¹²⁴ Another strategy involves forming polyelectrolyte complexes, where oppositely charged polymers interact with hydrophilic drugs to stabilize encapsulation.¹²⁵ Additionally, the use of hydrophobic ion pairing has been employed, wherein hydrophilic drugs are complexed with counterions to temporarily enhance their hydrophobicity, improving retention within the nanoparticle matrix.¹²⁶ For example, hydrophobic ion pairing of heparin with surfactant-like cations has

been shown to significantly improve its encapsulation efficiency in polymeric nanoparticles.¹²⁷ Despite these advancements, optimizing formulation parameters, such as solvent selection, polymer–drug compatibility, and mixing conditions, remains crucial for achieving stable and efficient encapsulation of hydrophilic molecules.

Nanoprecipitation is particularly advantageous for hydrophobic drugs, improving their solubility and bioavailability. Yang *et al.* utilized salt-induced precipitation to achieve high drug loading (66.5% w/w) for hydrophobic drugs. By varying salt concentrations, they controlled particle size and aggregation, resulting in uniformly dispersed nanoparticles (50 nm) with high loading efficiency in high-salt conditions (Fig. 6(d)–(g)).¹²⁸ Liu *et al.* created stable core–shell PNPs with loading up to 58.5% by adjusting polymer precipitation timing, allowing for either single-core or multi-core formations depending on the desired release profile.¹²⁹ For the antioxidant drug astaxanthin, Azaman *et al.* encapsulated it in PLGA nanoparticles, producing stable and homogeneous PNPs (142 nm), optimized for enhanced oral bioavailability and antioxidant efficacy.¹³⁰

Batch nanoprecipitation (BNP) has proven effective for encapsulating proteins and peptides, which benefit from controlled release. Chopra *et al.* developed insulin-loaded PLGA–PEG PNPs using BNP, achieving a tenfold increase in insulin loading while maintaining small particle size. The encapsulated insulin retained its structure, promoting long-term stability (Fig. 6(a)–(c)).¹³¹ Zada *et al.* used BNP in a non-aqueous setup for nasal insulin delivery, yielding rapid release rates with nearly 50% of the drug released within the first hour, ideal for nasal administration where quick absorption is advantageous.¹³²

Flash nanoprecipitation (FNP) has shown success in stabilizing hydrophobic drugs with complex release profiles. A “complex release profile” refers to a non-uniform or multi-phase drug release behavior over time, characterized by an initial burst followed by sustained, delayed, controlled, extended, or environment-responsive release. This behavior is influenced by factors such as drug–polymer interactions, nanoparticle composition, and external environmental conditions (*e.g.*, pH, temperature), making it critical for optimizing therapeutic efficacy. In this context, FNP enables precise control over nanoparticle formation, facilitating tailored drug release kinetics. Li *et al.* numerically studied the effect of solution/water flow rate ratios and microfluidic device geometry on mixing time. Fig. 6(h)¹³³ shows the microfluidic nanoprecipitation of curcumin nanoparticles. In the first stage, most of the polymer precipitates, forming polymeric NPs, followed by curcumin NPs. Some are stabilized by mPEG–PLGA, creating drug-loaded NPs. However, at higher curcumin concentrations, there is insufficient mPEG–PLGA to stabilize the drug NPs, leading to aggregation and microchannel blockage. To improve drug loading and stability, it is ideal to precipitate the drug first, ensuring enough polymer is available to stabilize the NPs. Thus, the precipitation times for the polymer and drug should be aligned.¹³³ Qi *et al.* used FNP to encapsulate celastrol, achieving tunable drug loading (11–63%) in dextran-based PNPs. The formulation exhibited controlled release and reduced cytotoxicity toward liver cells,



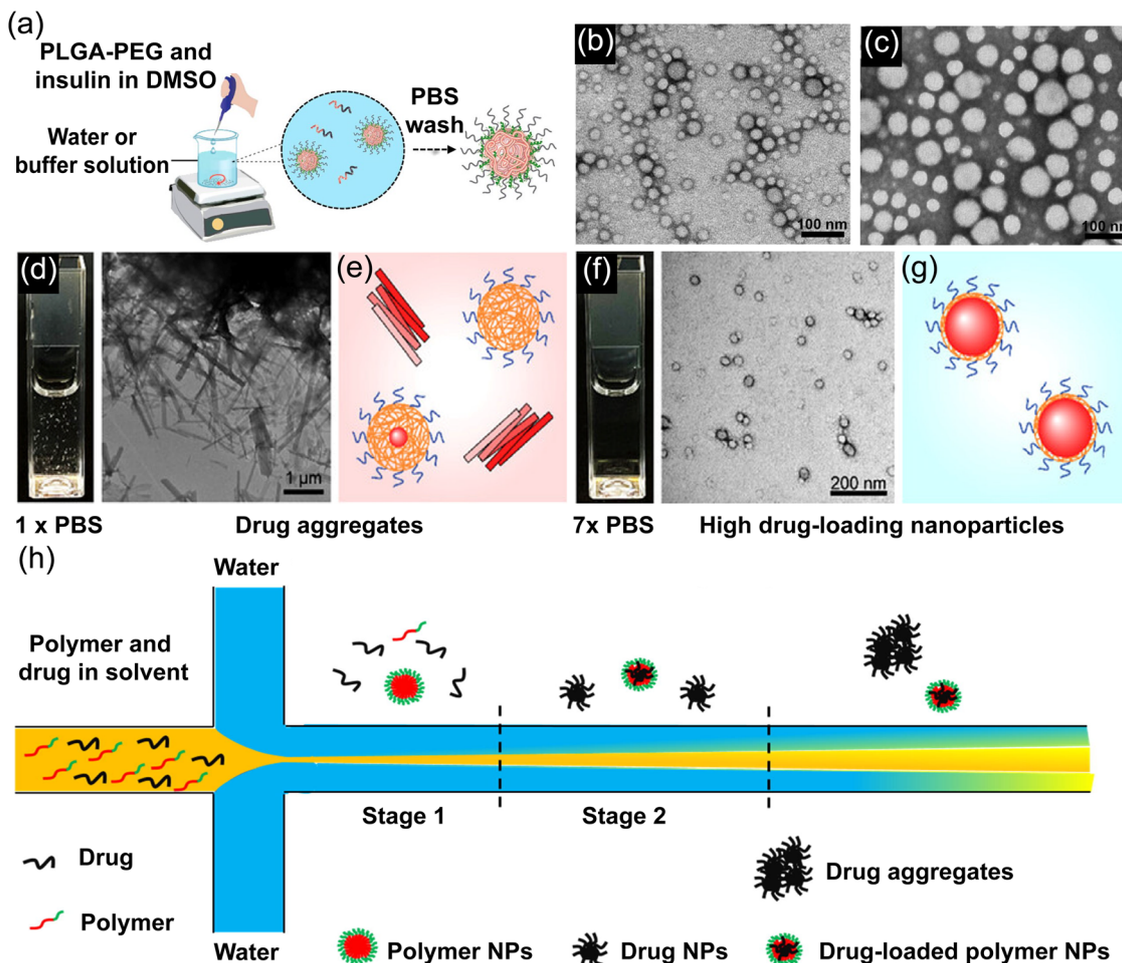


Fig. 6 Nanoprecipitation for fabrication of nanocarriers of pharmaceutical active ingredients. (a) Schematic representation of the preparation of the insulin-loaded PNPs (Ins-NPs) and schematic for the nanoparticles produced by (d) and (e) bad salt concentration and (f) and (g) good salt concentration in producing 50 wt% docetaxel-loaded PLGA10k-PEG5k nanoparticles.¹²⁸ (h) Schematic representation of the formation of the polymer-stabilized peroxide antimalarial drug nanoparticle and the drug release profile of the nanoparticle and unencapsulated powder in the bio media.⁸¹

with effective inhibition of lung cancer cells, demonstrating potential for targeted cancer therapies.¹³⁴

Caggiano *et al.* employed FNP with confined impinging jets to co-encapsulate cannabidiol (CBD) and iron oxide. This design increased particle density, improving sedimentation for controlled release studies. Stabilized with HPMCAS or lecithin, the nanoparticles demonstrated enhanced dissolution, with HPMCAS-coated particles releasing six times faster in intestinal media compared to bulk CBD as shown in Fig. 7(a).¹³⁵ Zeng *et al.* utilized a scalable approach to prepare lipid-coated solid drug (methotrexate) nanoparticles by combining flash nanoprecipitation and extrusion techniques (Fig. 7(b)), optimizing individual steps and providing flexibility in selecting nanoparticle surface functionalities.¹³⁶ Ye *et al.* encapsulated the cancer drug paclitaxel within PEG-PLA/zein nanoparticles using FNP, achieving high encapsulation efficiency (up to 78.1%) and controlled, sustained release. The presence of zein promoted hydrophobic interactions, allowing slow release at acidic pH, advantageous for tumor-targeted delivery.¹³⁷

Microfluidic nanoprecipitation (MNP) enables precise control over particle size, ideal for creating pH-responsive nanocarriers. Li *et al.* synthesized PEG-PLGA PNPs for curcumin delivery, achieving a loading capacity of 2.6% and 77.3% encapsulation efficiency. However, they observed particle aggregation at higher curcumin concentrations, highlighting the importance of concentration control for stable dispersions.⁸¹ Baby *et al.* developed pH-responsive shellac nanoparticles for curcumin delivery using MNP, achieving up to 50% drug loading. At neutral pH, the nanoparticles released 28% of the drug in 4 hours, increasing to 51% over 51 hours, demonstrating suitability for sustained release in response to pH changes.⁸⁰ The pH-responsive capabilities of FNP were further explored by Qi *et al.*, who used dextran-based PNPs to encapsulate celestrol, an anti-cancer drug. The PNPs achieved adjustable drug loading (11–63%) with effective release, reduced liver toxicity, and significant inhibition of lung cancer cells. This adaptable release profile shows promise for personalized cancer therapies.¹³⁴



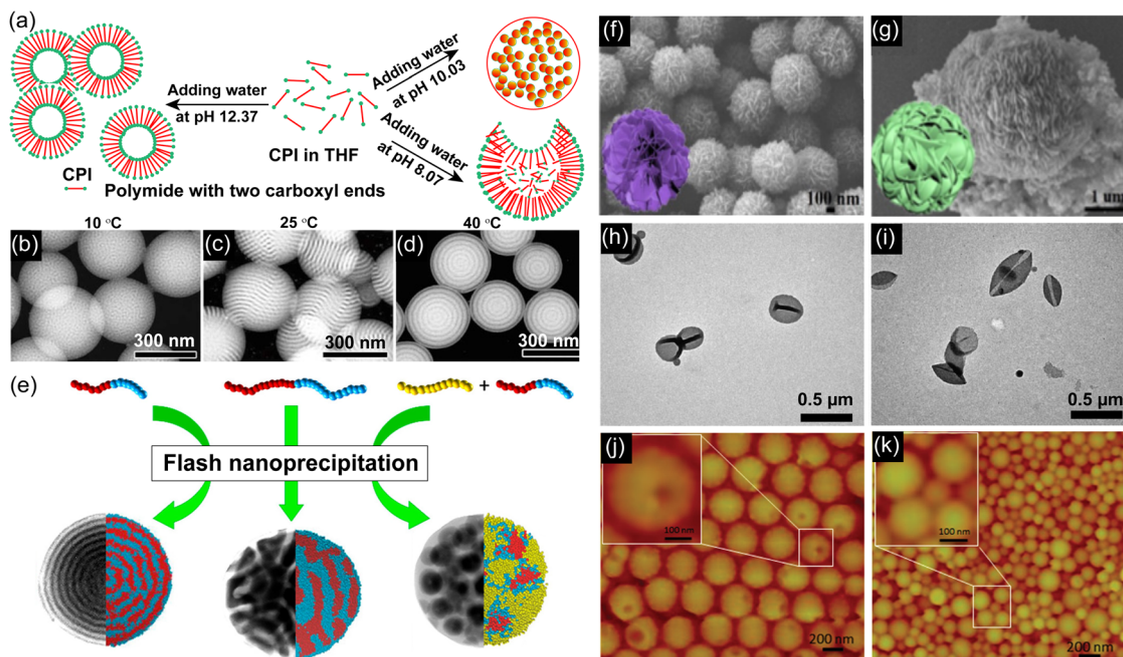


Fig. 8 (a) Complex morphologies through mono-component PNPs obtained by varying the solution pH.¹⁴⁰ (b)–(d) Effect of temperature on morphology of the PNPs.¹⁴¹ (e) Schematic illustration of the Janus colloid formed from a blend of homo and block co-polymers solutions.¹⁰⁹ (f) and (g) SEM images of the particles of different morphologies (f) MoDo-SiO₂ and (g) CTS-MoDo-SiO₂.¹⁴² (h) and (i) TEM images of organosilica nanoparticles (h) collapsed hollow structure with intrinsic flexibility and deformation. (i) Ultra-thin shells with deformable hollow structure¹⁴ (j) and (k) AFM images of the polymer Janus nanoparticles obtained using (j) thiol-PLGA (k) PLA/PGA solutions.¹⁴³

fields ranging from opto-electronics and biosensing to bioimaging, catalysis, and drug delivery.^{9,10,12,14,138,139} Nanoprecipitation simplifies the synthesis of nanomaterials that benefit from controlled solvent-based assembly, particularly where low energy input and single-step synthesis are required. This makes it an ideal method for producing nanostructures that would otherwise be difficult or inefficient to achieve using traditional techniques.

Nanocrystals are crystalline nanoparticles with a wide range of potential applications in opto-electronics, biosensing, bioimaging, and catalysis due to their enhanced dissolution rates and tunable physicochemical properties.^{13,144,145} Their properties can be customized by selecting appropriate stabilizing agents, and nanoprecipitation offers a simple yet effective synthesis method.^{146,147} For example, Xu *et al.*¹⁴⁷ used nanoprecipitation to create aggregation-induced emission (AIE) activated nanocrystals for super-resolution imaging. These AIE nanocrystals exhibited enhanced brightness and photostability, making them ideal for stimulated emission depletion microscopy. This technique improved the resolution of lysosomal imaging and enabled dynamic tracking of lysosomal motion over extended periods, surpassing standard confocal microscopy capabilities. Thakkur *et al.*¹⁴⁸ employed electrospray nanoprecipitation to produce docetaxel nanocrystals, which showed a significant increase in dissolution rates (77% *vs.* 30% for bulk docetaxel) due to their high surface area. *In vivo* studies demonstrated effective tumor load reduction in a lung cancer model, underscoring the therapeutic potential of nanocrystals in oncology.

Semiconducting polymer nanoparticles (SNPs) are emerging materials with applications in sensing, imaging, and diagnostics, owing to their fluorescence, biocompatibility, and superior optical and photothermal properties compared to traditional dyes.^{12,139,149} The molecular structure of the semiconducting polymers largely determines the properties of the SNPs. Holmes *et al.*¹² synthesized Janus SNPs with electron- and hole-accepting faces using batch nanoprecipitation. These Janus SNPs were employed to fabricate organic field-effect transistors, demonstrating efficient charge transport across thin films, highlighting their potential in optoelectronic applications. He *et al.*⁵⁴ used flash nanoprecipitation (FNP) to synthesize ultra-small polymer dots (less than 10 nm) with narrow size distribution and tunable optical properties. By selecting polymers such as MEH-PPV, PFBT, and PFPV, they achieved highly bright and stable polymer dots suitable for high-resolution imaging. The size of these polymer dots was controlled by adjusting the polymer type and precursor concentration, showcasing the flexibility of FNP for tailoring nanoparticle properties.

Pu *et al.*¹⁵⁰ developed low-bandgap diketopyrrolopyrrole (DPP) SNPs for *in vivo* photoimaging. DPP, known for its photostability and thermal stability, was copolymerized with electron-donating monomers to fine-tune band gaps, resulting in stable SNPs (45 nm) with modifiable fluorescence and photoacoustic properties. By adjusting the donor-acceptor composition, they optimized SNPs for photothermal applications, enhancing photothermal conversion efficiency while reducing fluorescence. This tunable property makes DPP SNPs particularly promising for biomedical imaging and therapy.



Wang *et al.*¹⁴⁰ demonstrated that vesicles, porous spheres, and dimpled beads could be synthesized by controlling the pH of an aqueous solution of carboxyl-terminated polyimide (Fig. 8(a)). At pH 8.07, the polymers aggregated into dimpled beads (100–600 nm), while increasing the pH to 10 produced smooth, spherical particles (100–200 nm). This pH-dependent assembly highlights how solution conditions can direct nanoparticle morphology (Fig. 8(b)–(d)). Similarly, Higuchi *et al.* used thermal annealing to induce disorder–order and order–order phase transitions in block copolymer nanoparticles (Fig. 8(b)–(d)). The unusual thermal behaviors suggest that the nanoparticle effect lowers the glass transition temperature of the block copolymer, likely due to the increased surface-area-to-volume ratio.¹⁴¹

Grundy *et al.*¹⁰⁹ synthesized colloids with complex internal structures by blending poly(styrene) and poly(styrene-*b*-isoprene) block copolymers in an FNP process. Low-molecular-weight copolymers produced concentric, onion-like shells, while higher molecular weights resulted in disordered, lamellar structures, demonstrating control over internal nanoparticle architecture through polymer selection (Fig. 8(e)). Shahnava *et al.*¹⁵¹ developed pH-sensitive core–shell nanoparticles composed of PLGA and carboxymethyl chitosan, allowing dual drug loading. Doxorubicin was loaded in the shell, while docetaxel was encapsulated in the core, creating a system responsive to environmental pH. The core–shell structure enabled controlled drug release: doxorubicin was released rapidly from the shell, while docetaxel in the core showed slower release, allowing for layered, sequential delivery.

Ma *et al.*¹⁴ used a modified FNP setup to develop deformable hollow mesoporous organosilica nanoparticles (HMONs) with high surface area and tunable mechanical properties (Fig. 8(f)–(i)). By introducing disulfide bonds into the silica network, they created nanoparticles with lower Young's modulus, enabling deformation. Loaded with the model nanopesticide abamectin, these HMONs demonstrated improved insecticidal efficacy, showing their potential for applications in agriculture where controlled-release carriers are needed. In another work, Xie *et al.*¹⁴³ designed a fluidic nanoprecipitation system capable of fabricating biocompatible Janus polymeric nanoparticles made from the FDA-approved polymer poly(lactic-*co*-glycolic acid) (PLGA), as shown in AFM images (Fig. 8(j) and (k)). The system features dual inlets, each for one-half of the particle, which is inserted into the precipitation stream.

The possible future scope of nanoprecipitation includes AI and ML integration for predictive control over nanoparticle synthesis, automated and scalable processes using continuous-flow and microfluidic systems, and hybrid/multi-functional nanoparticles for advanced applications in drug delivery, sensing, and catalysis. These advancements will enhance precision, reproducibility, and scalability, making nanoprecipitation more efficient and versatile.

Conclusions

Nanoprecipitation is a simple, versatile, low-energy, and cost-effective method for synthesizing a wide variety of nanomaterials.

Its adaptability, coupled with precise control over physicochemical properties and mixing hydrodynamics, enables customized size, morphology, and internal structures of particles. This solution-based approach has evolved significantly, enabling the production of nanomaterials with distinct morphological characteristics. Through these capabilities, nanoprecipitation holds great promise for developing high-performance, multifunctional nanomaterials. As highlighted in this review, nanoprecipitations effectiveness across batch, flash, and microfluidic mixing configurations provides fine control over drug loading capacity, encapsulation efficiency, and release kinetics, making it a powerful approach for tailoring drug delivery systems.

Despite its potential, challenges persist, particularly in large volume of solvents, scaling microfluidic techniques and optimizing formulations for hydrophilic compounds. Future advancements in nanoprecipitation could address these limitations through several innovative approaches. First, the development of fast and efficient methods for concentrating and separating nanoparticles from large liquid volumes would streamline production, making the process more time-efficient. Second, extending nanoprecipitation techniques to include hydrophilic compounds could broaden its applicability across biomedical and industrial fields. Finally, the automation of nanoprecipitation through robotic systems, coupled with machine learning for optimizing formulation and flow conditions, could accelerate the discovery and development of new nanomaterial formulations.

List of abbreviations

BNP	Batch nanoprecipitation
CBD	Cannabidiol
DMF	Dimethylformamide
DMSO	Dimethyl sulfoxide
FNP	Flash nanoprecipitation
HMONs	Hollow mesoporous organosilica nanoparticles
HPMCAS	Hydroxypropyl methylcellulose acetate succinate
MNP	Microfluidic nanoprecipitation
PBS	Phosphate buffer solution
PDI	Polydispersive index
PEG- <i>b</i> -PLA	Poly(ethylene glycol)- <i>block</i> -poly(DL-lactide)
PNP	Polymer nanoparticle
PMMA	Poly(methyl methacrylate)
PLGA	Poly(lactic- <i>co</i> -glycolic acid)
PLGA	Poly(lactide- <i>co</i> -glycolide)
PVA	Poly(vinyl alcohol)
Re	Reynolds number
THF	Tetrahydrofuran
PbSO ₄	Lead(II) sulfate

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.



Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We acknowledge funding support from the Natural Sciences and Engineering Research Council of Canada (NSERC)-Discovery project, and Alliance Grant Alberta Innovates-Advanced program. This manuscript was prepared with the assistance of AI-based tool, ChatGPT 4o, which was used to refine language and improve clarity. All intellectual and technical contributions to the research and analysis were made by the authors, who take full responsibility for the content and conclusions presented in this work.

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