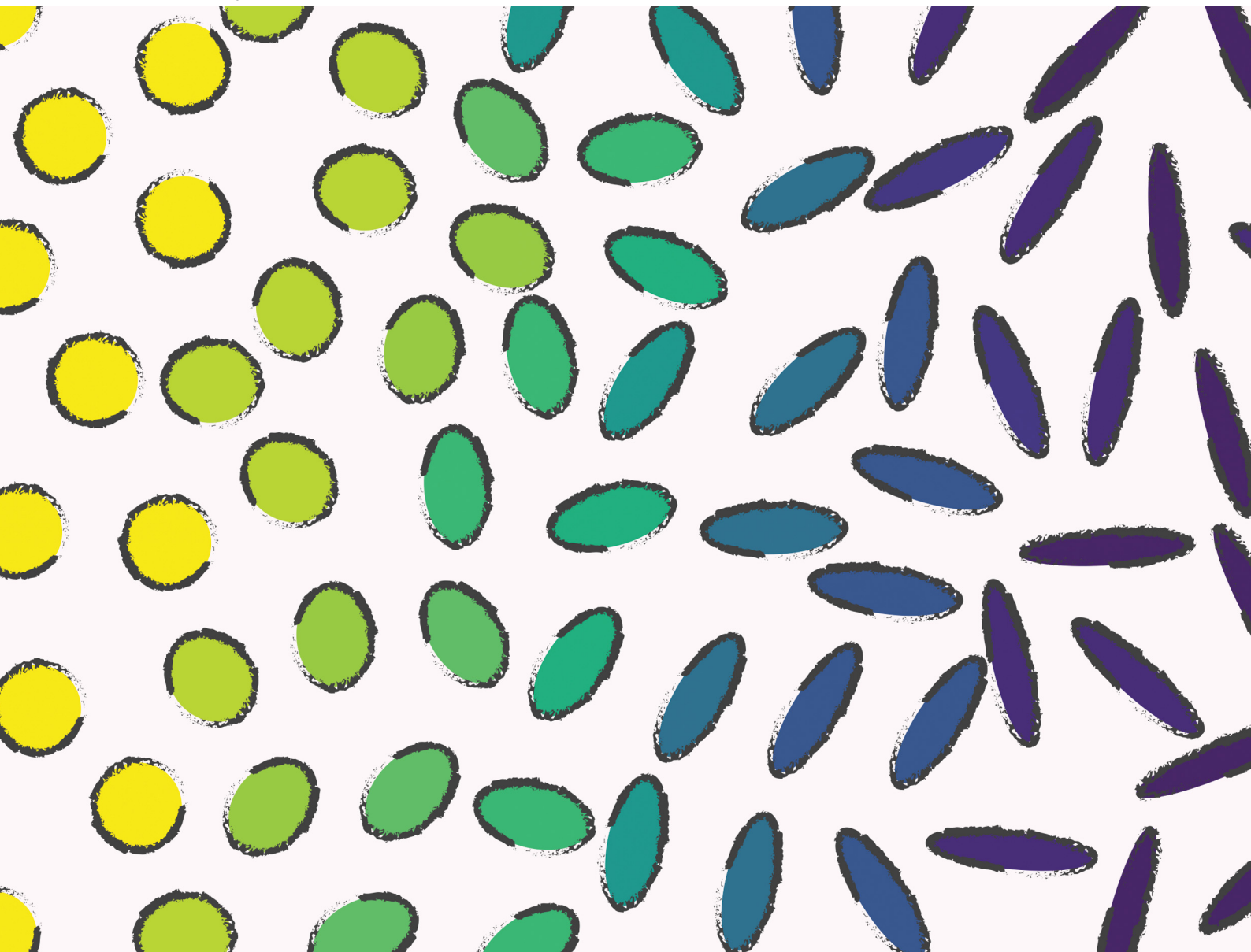


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Reshapable magnetic particles for morphology-controlled soft systems†

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Spherical polymeric particles are essential in a wide range of applications, from fundamental self-assembly to bioseparation technologies, with well-established synthetic methodologies. However, incorporating functional components into polymeric matrices to enhance their properties remains a significant challenge, especially when uniformity is required. In this study, we introduce a simple and versatile emulsion evaporation method to fabricate magnetic-loaded polymeric microparticles with exceptional malleability. These composite particles maintain their magnetic functionality while being reshaped into ellipsoids through mechanical stretching. This scalable and straightforward approach offers precise control over particle morphology, offering broad potential for applications in soft robotics, drug delivery, and other magnetically responsive systems.

Introduction

Functional microparticles embedded with inorganic materials find broad applications, from cosmetics¹ to optics² and biotechnology.^{3–5} Of particular interest are magnetic particles, due to their ability to be manipulated by external magnetic fields.^{5–8} Several approaches have been developed for the preparation of magnetic polymeric particles. A common method involves polymerizing a monomer in the presence of magnetic nanoparticles,⁵ such as *via* dispersion or (mini-)emulsion-based polymerization. These processes typically produce two-phase systems, where polystyrene and magnetic nanoparticles form distinct phases, resulting in Janus-like or core-shell structures depending on the polymerization method.^{6,9–13} Additional challenges such as phase separation, nanoparticle aggregation, and low nanoparticle loading significantly limit the effectiveness of these techniques.

Physical-based methods provide viable alternatives to polymerization. For instance, Ugelstad *et al.*^{14,15} developed a technique that involves swelling polymeric microparticles with an organic solvent to allow diffusion of reagents and direct nucleation of inorganic nanoparticles throughout the polymer matrix. In another approach, magnetic micron-size particles are embedded in a polymer matrix by exploiting opposite surface charges and using a plasticizer to physically trap the magnetic particles.¹⁶ Similarly, magnetic nanoparticles can be embedded

onto the surface of a polymeric particle with opposite surface charge.^{17,18}

Emulsion droplet evaporation has emerged as a promising technique to prepare polymeric microparticles. This method involves evaporation of solvent from emulsion droplets containing dissolved polymers, producing robust polymeric microparticles.¹⁹ When paired with block copolymers, microparticles with different morphologies can be formed under controlled evaporation conditions and surfactant selection.²⁰ Adjustment of the emulsification method and evaporation time can also produce dimpled and crumpled microparticles, systems with higher surface areas compared to their spherical counterpart.²¹ This is advantageous for specific applications such as drug and cargo delivery.²² The most promising synthetic methods primarily yield spherical polymeric particles, while anisotropic shapes, such as ellipsoids, donuts, discs, and rods are generally limited to techniques that lack scalability.²³

An alternative to produce anisotropic shapes is the thermal stretching method, developed by Ho *et al.*,^{24,25} where polymeric films containing polystyrene microspheres are heated above the glass transition temperature of the film and stretched, resulting in ellipsoidal particles with controlled axial ratios. The stretching of the particle is in part due to the fact that the polymer is uncrosslinked allowing for easier deformation. Champion *et al.*²⁶ refined this technique to achieve a wide variety of shapes, including rods, disks, and even UFO-like structures. More recently, Lo *et al.*^{27,28} further simplified this technique, generating ellipsoids with sharp or blunt ends by simply stretching the particle embedded films with a weight linked to a binder clip. Despite these advances in particle synthesis, achieving precise control over both nanoparticle loading percentage and overall particle shape remains challenging.

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In this work, we introduce a straightforward two-step synthesis technique to expand the design space for colloidal particles. Using uncrosslinked polystyrene, and cobalt ferrite nanoparticles, we employ an emulsion evaporation method to produce a variety of magnetic-loaded composite microparticles. These particles are embedded into PVA films and stretched to achieve controlled anisotropic shapes, demonstrating the versatility of this approach in the preparation of anisotropic colloid with controlled magnetic properties.

Methods

Materials

Styrene (with 4-*tert*-butylcatechol as a stabilizer, $\geq 99\%$), dibenzoyl peroxide (BPO, $>75\%$), cobalt(II) chloride hexahydrate ($\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$, $\geq 95\%$), iron(III) chloride hexahydrate ($\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$, $\geq 98\%$), iron(III) nitrate nonahydrate ($\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$, $\geq 98\%$), sodium hydroxide (NaOH, $\geq 98\%$), hydrochloric acid (HCl, 37%), nitric acid (HNO_3 , 60%), oleic acid (90%), sodium lauryl sulfate solution (SDS, 10%), poly(vinyl alcohol) (PVA, 145 000 mW), and tetrahydrofuran (THF, 99.5%) were used as received and purchased from Sigma Aldrich or TCI Europe.

Chemical synthesis

Polystyrene was synthesized by free radical polymerization of styrene monomers.²⁹ Styrene (27 g), toluene (45 mL), and BPO (0.28 g) were combined in a 100 mL round bottom flask, deoxygenated with argon for 30 minutes, and heated to 80 °C for 37 hours. Polystyrene was obtained at 84% conversion (determined by ^1H NMR). The product was diluted with 30 mL chloroform and precipitated in 500 mL ice-cold methanol, then dried under vacuum to achieve 99% purity (Fig. S1, ESI†).

Magnetic cobalt ferrite nanoparticles were prepared *via* a coprecipitation method.^{30,31} $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$ (2.40 g) was dissolved into a 5 mL aqueous solution of 7.5% HCl, and 5.50 g of $\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$ was dissolved in 40 mL of water. The two solutions were briefly stirred together and then rapidly added to 200 mL of boiling NaOH (1 M) solution where a black precipitate was quickly formed. The mixture was vigorously stirred with a magnetic stirrer at 100 °C for 1 hour under reflux. The dispersion was cooled to room temperature and washed *via* magnetic sedimentation once. The particles were redispersed in 50 mL of water, and 30 mL of 2 M HNO_3 was added. Under stirring, 30 mL of a 0.35 M $\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$ solution was added to the particle suspension and heated at 100 °C for 1 hour. The dispersion was cooled again to room temperature, washed thrice *via* magnetic separation, and redispersed into 50 mL of water (Fig. S1, ESI†). Surface modification was achieved by adding 15 mL oleic acid dropwise to the dispersion, stirring for 18 hours. Surface modification was confirmed upon the successful phase transfer of the cobalt ferrite particles to the oil phase. The aqueous phase was removed using a separatory funnel, and the particles were washed three times with ethanol through magnetic separation. Excess ethanol was then

removed, and the particles were dried overnight at 80 °C to yield a powder.

Evaporation-based particle formation

Loaded magnetic polystyrene microparticles were prepared by emulsion evaporation.³² In a typical synthesis, 1 mL of 0.1% SDS solution was mixed with 50 μL DCM solution containing 1.5–10 wt% dissolved polystyrene and 0–1 wt% dispersed cobalt ferrite. The mixture was vortexed for 30 seconds, sonicated up to 5 minutes until it appeared milky white, and diluted with 3 mL water. The dispersion was placed under a vacuum on a rotary evaporator at 40 °C for 25 minutes to remove DCM. Particles were washed thrice *via* gravitational sedimentation overnight and redispersed in 1 mL of water. Plasticization of the particles was employed to eliminate voids within their structure. To 1 mL of the prepared particles, 3 mL of a 40% THF solution was added. The mixture was gently agitated by hand for approximately 20 seconds, diluted with 20 mL of water, and subsequently washed using magnetic separation or centrifugation.

Stretching

Films with embedded polymeric particles were prepared following a modified method from ref. 27. A 20 mL aqueous dispersion containing 0.01 wt% (loaded) polystyrene particles was heated to 80 °C. At temperature, 1 g of PVA was added and magnetically stirred until the PVA dissolved completely, typically 1 hour. The solution sat without stirring for at least 3 hours to remove bubbles. Inside a round 8.5 cm plastic Petri dish, 10 mL of the solution was poured and placed on a level surface overnight for the water to evaporate. The films prepared as described resulted in films with thicknesses around 50 μm devoid of bubbles and lumps. After drying, PVA films were cut into strips ($3.5 \times 1 \text{ cm}^2$) and stretched to specific draw ratios using a Universal Testing Machine (Instron 3365) at 140 °C (Fig. S2, ESI†). Uniform stretching was confirmed by marking films every 0.25 cm before stretching. Films stretched under a magnetic field were first inserted into the machine and clamped with two neodymium magnets placed to apply a magnetic field perpendicular to the stretching direction. Stretched films were cut, rinsed with IPA, and submerged in a 20% v/v IPA–water mixture at 60 °C to dissolve the PVA. The now dispersed stretched particles were washed *via* centrifugation at 13 000 rpm for 1 hour and redispersed in distilled water.

Material characterization

Transmission electron microscopy (TEM) measurements were performed on a JEOL JEM-1400 plus TEM at 120 kV. Samples were prepared by placing 5 μL of a diluted particle dispersion onto carbon-coated copper grids. Scanning electron microscopy (SEM) measurements were performed on a JEOL JSM-6010LA with secondary electron, back-scattered electron detectors, and energy dispersive X-ray spectroscopy (EDX) capabilities. Samples were prepared by drying 5 μL of a diluted particle dispersion onto a cut silica wafer. Electron images and elemental distribution maps were collected. Particle (and droplet) size



distributions were calculated from the microscopy images using a custom python script. Particle dynamics were followed using an inverted optical microscope (Nikon Eclipse E600pol) equipped with a 50 $\times$ objective and a CCD camera. Optical microscopy samples were prepared inside flat rectangular capillaries (Vitrocom 0.05 $\times$ 1.00 mm) and sealed with wax at each end to a microscope slide. A uniform magnetic field was applied to the sample with a custom magnetic set-up (Fig. S2, ESI†). Image analysis was performed with ImageJ Fiji.³³ We fit images of the stretched particles to an ellipse with a major axis, a , and minor axis, b , and determine the axial ratio by $\frac{a}{b}$. The reported angle is defined as the angle between the major axis of the ellipses and a line parallel to the x -axis of the image.

Results and discussion

Magnetic polystyrene microparticles were synthesized using an emulsion evaporation method. In this process, polystyrene (PS) and cobalt ferrite nanoparticles were emulsified and subsequently dried to form solid particles. As illustrated in Fig. 1A, the process involves three main stages: (i) mixing of the dichloromethane (DCM) solvent phase containing PS and cobalt ferrite nanoparticles with the aqueous surfactant solution, (ii) emulsification to encapsulate nanoparticles and PS within DCM droplets, and (iii) evaporation of the solvent to solidify the droplets into particles. Two mechanisms are proposed for particle formation during solvent evaporation

within an emulsion.^{34,35} In both, the polymer migrates to the droplet interface, forming a region of high polymer concentration. As the solvent evaporates, the polymer at the interface begins to aggregate and solidify, creating a shell. This shell formation occurs either through a receding mechanism, where the polymer layer contracts inward with the evaporating solvent, or through a stationary mechanism, where the shell remains at the original interface while the solvent diffuses outward through the polymer layer.

To assess the baseline effects of the emulsion evaporation method without the influence of magnetic nanoparticles, we prepared PS microparticles at various initial PS concentrations (1.5, 2.5, 5.0, and 7.5 wt%) in DCM. Higher concentrations (≥ 10 wt%) produced viscous polymer solutions that failed to fully emulsify, resulting in large, phase-separated polystyrene regions. Conversely, lower concentrations (< 1.5 wt%) did not yield sufficient amount of particles. Within the specified range, spherical particles were easily prepared. SEM images in Fig. 1B show representative samples. Generally, most particles exhibited spherical shapes, with occasional bucket-like and dimpled or donut-like morphologies, with sizes ranging from 0.3 to 15 μm . The proportion of non-spherical particles decreased as the initial PS concentration increased. For example, at 1.5 wt% PS, 21% of particles had non-spherical shapes, whereas this number decreases to 2% at 7.5 wt% PS, based on a sample size of over 500 particles. The buckling observed at lower PS concentrations is likely caused by significant pressure differences between the inside and the outside of the particles during

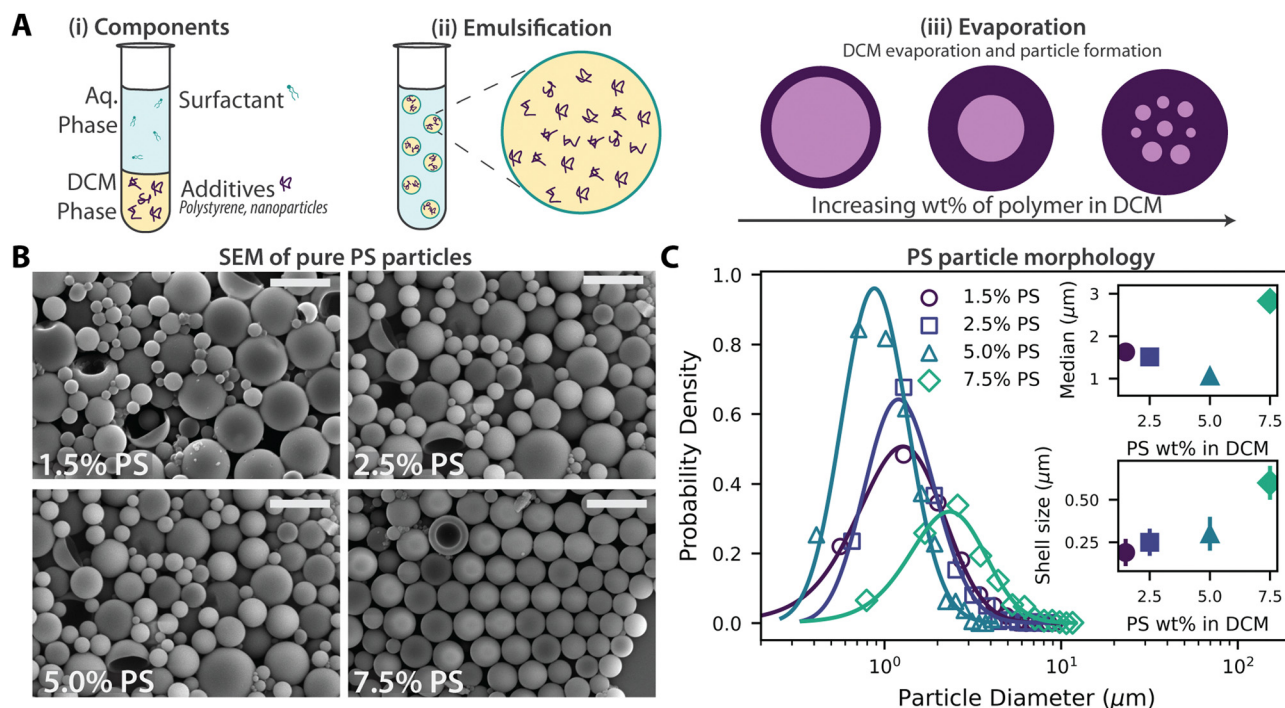
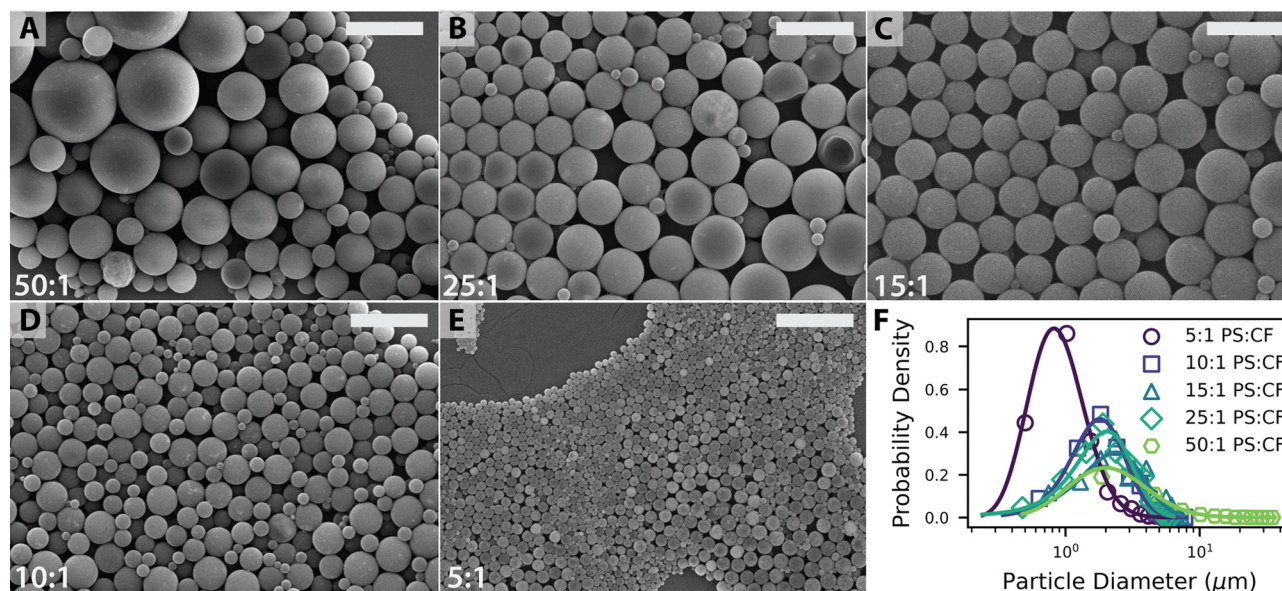


Fig. 1 Preparation and morphology of magnetic polystyrene microparticles via the emulsion evaporation method. (A) Schematic of the emulsion evaporation process: (i) initial components include polystyrene (PS), cobalt ferrite nanoparticles, and a surfactant solution. (ii) Emulsification results in PS droplets with encapsulated nanoparticles. (iii) Evaporation produces particles with thinner shells at low PS wt% and thicker shells at higher PS wt%. (B) SEM images of particles formed from varying PS concentrations: 1.5 wt%, 2.5 wt%, 5.0 wt%, and 7.5 wt%. (C) Particle size distributions for each PS concentration, with insets showing the median particle size and corresponding shell thickness.



To investigate the effect of nanoparticle encapsulation on the composite particles, we prepared samples in the presence of cobalt ferrite (CoFe) nanoparticles. Fig. 2A–E shows SEM images of particles prepared with 2.5 wt% PS in DCM and PS/CoFe nanoparticle weight ratios of 50:1, 25:1, 15:1, 10:1, and 5:1. These ratios correspond to theoretical CoFe loading values of 2–20%. Across all samples, most particles appear spherical, although a small number of bucket-like and dimpled morphologies were also observed. Fig. 2F shows the probability density plots of particle diameters for different loading ratios. As the

To investigate the uniformity of the cobalt ferrite distribution within the microparticles, we utilize an SEM that detects back-scattered electrons (BSE) and is equipped with



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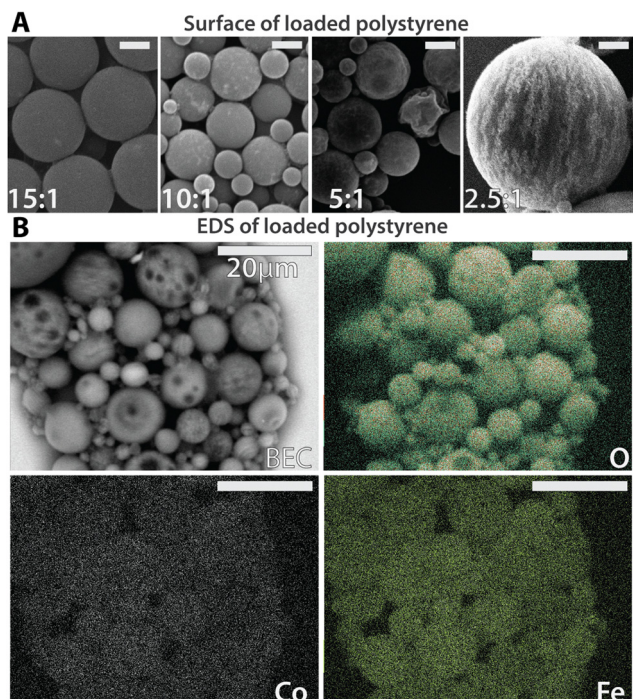


Fig. 3 Morphology of polystyrene microparticles loaded with cobalt ferrite. (A) Scanning electron microscopy (SEM) images of particle surfaces at polystyrene-to-cobalt ferrite ratios of 15:1, 10:1, 5:1, and 2.5:1. Scale bars represent 1 μm . (B) Energy-dispersive X-ray spectroscopy (EDS) analysis of particles prepared at a 2.5:1 polystyrene-to-cobalt ferrite ratio, with corresponding back-scattered electron composition (BEC) images and elemental maps for carbon, iron, and cobalt. Scale bars represent 20 μm .

energy-dispersive X-ray spectroscopy (EDS). In back-scattered electron composition (BEC) images, changes in apparent intensity correspond to variations in material density, with darker

regions indicating lower density and brighter regions indicating higher density. This, combined with EDS, allows us to determine the elemental composition of the samples, as shown in Fig. 3B for a 2.5:1 PS/CoFe sample. In the elemental maps, the PS matrix is identified in the carbon signal, while the cobalt ferrite nanoparticles appear in the iron and cobalt maps. Since the sample is deposited on a silica wafer, particle edges are clearly visible across all maps. The elemental maps confirm that cobalt ferrite nanoparticles are evenly distributed throughout the PS matrix rather than aggregating into distinct, phase-separated domains. However, we observe significant density variations in the BEC-SEM image that are not reflected in the elemental maps, indicating that these density differences arise from voids within the composite particles rather than variations in cobalt ferrite concentration. Additional EDS maps for 100:1 and 10:1 PS/CoFe samples are provided in Fig. S7 (ESI[†]), further confirming the cobalt ferrite distribution across samples even at low concentrations.

To remove internal voids and ensure the particles are solid, we added tetrahydrofuran (THF) as a plasticizer to the dispersion. The addition of THF causes the particles to swell, allowing the polymer strands to minimize the interfacial energy with the surrounding medium,²⁷ thereby forming uniform spherical particles. After diluting the dispersion and removing the THF, the particles deswell into uniform spheres. Non-destructive BEC-SEM imaging was used to explore the internal morphology of the particles. Fig. 4A shows PS/CoFe particles before and after THF treatment. The bright areas in the SEM image likely correspond to CoFe nanoparticle aggregates, as iron and cobalt are heavier elements compared to the carbon and oxygen atoms in the polymer. Before THF treatment, 94% of particles with diameters $>3\text{ }\mu\text{m}$ displayed hollow cores, visible as darker centers compared to the edges in the SEM images. Some examples are highlighted with white arrows in the image.

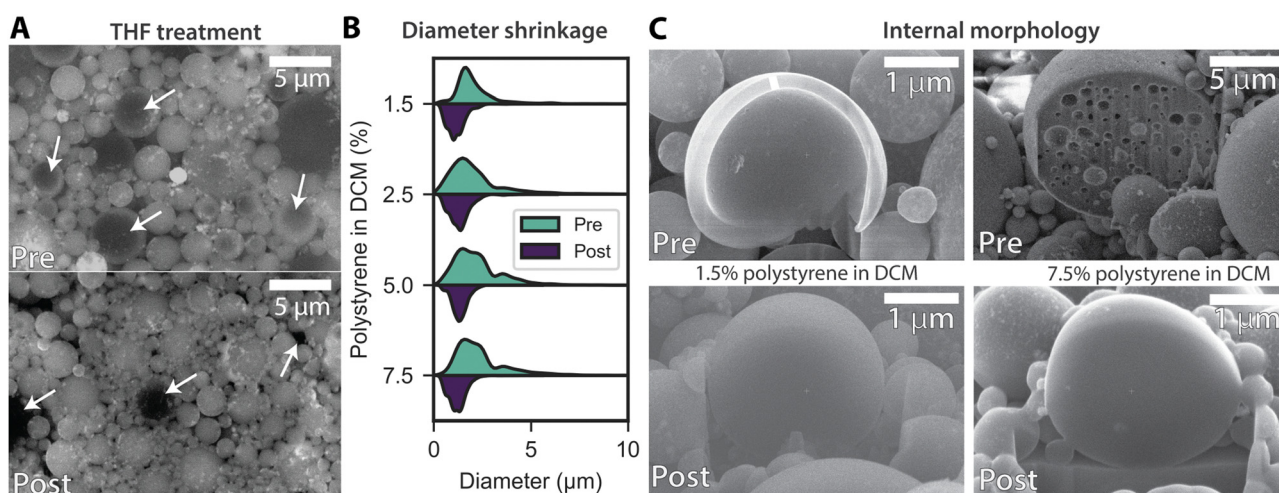
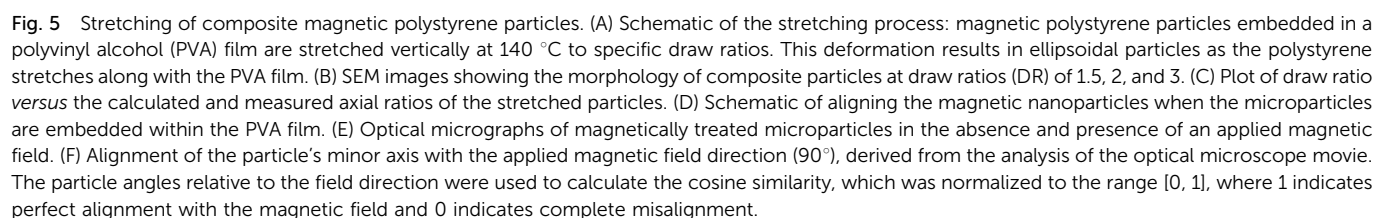


Fig. 4 Densification of particles through void removal using a plasticizer. (A) Back-scattered electron composition (BEC) and scanning electron microscopy (SEM) images of polystyrene (PS) and cobalt ferrite particles in a 10:1 ratio, starting with an initial PS concentration of 2.5% in dichloromethane (DCM), shown before and after plasticization with tetrahydrofuran (THF). (B) Violin plot illustrating the particle size distribution changes before and after THF treatment at a 10:1 PS-to-cobalt ferrite ratio. (C) Focused ion beam (FIB)-cut SEM images revealing internal particle structures at low (1.5%) and high (7.5%) initial PS concentrations in DCM, both before and after plasticization.



To further modify the particle morphology, PS spheres treated with THF were embedded in polyvinyl alcohol (PVA) films, which were stretched under controlled conditions. To perform this procedure, PVA films loaded with 10:1 PS/CoFe particles were dried, clamped, and subjected to stretching 140 °C for 25 minutes. During stretching, the PS particles deformed along with the PVA film, owing to their glass transition temperature of approximately 100 °C. After the stretching process, the particles were recovered by selectively dissolving the PVA film in an isopropanol–water solution. The stretching process was controlled to achieve specific draw ratios (DRs), defined as the final film length divided by the initial film length. Fig. 5B shows SEM of particles stretched to draw ratios of 1.5, 2, and 3. We clearly observe the presence of ellipsoidal particles with sharp tips throughout the samples. This is in contrast with previous reports by Lo *et al.*,²⁷ in which



configuration. Furthermore, the resulting particle shape could be tailored by stretching in multiple planes, as demonstrated in prior work on anisotropic particle fabrication.²⁶ Together, these approaches would enable the fabrication of customizable microparticles with tunable shape and magnetic directionality.

These results highlight the versatility of the magnetic alignment process during particle elongation and complement previous methods for anisotropic particle fabrication. For example, Güell *et al.*⁴⁹ produced Janus-like ellipsoids using a non-uniform magnetic field applied during stretching in a solvent bath. Their solvent-based approach was particularly effective for commercial magnetic polystyrene spheres, which are coated with polymer layers that hinder adhesion during thermal stretching.^{50,51} In contrast, our method leverages the thermal plasticization of PS/CoFe particles in a PVA film, enabling the formation of sharp-tipped ellipsoids with high magnetic loadings. This approach provides a scalable and efficient route for fabricating magnetically anisotropic particles with tunable shapes and magnetic properties, opening avenues for applications in advanced colloidal assemblies and responsive materials.

Conclusion

In this study, we developed a general and versatile method to produce magnetic polystyrene microparticles embedded with varying weight percentages of cobalt ferrite nanoparticles. This technique involves creating a solvent-in-water emulsion where the solvent phase (DCM) contains various concentrations of polystyrene and cobalt ferrite nanoparticles, while the continuous phase is an aqueous SDS solution. To eliminate porosity and improve size distributions, we employed a plasticizer to condense the structure while maintaining the desired magnetic loading. Additionally, embedding the particles in a polyvinyl alcohol (PVA) film and thermally stretching them produced anisotropic shapes.

Future applications could leverage the inherent magnetic properties of these composite particles for remote manipulation. We showed that by aligning magnetic nanoparticles within a polymer matrix during stretching induces a net magnetization, with the magnetic response depending on factors like magnetocrystalline anisotropy,^{46,49} which governs magnetic moments reorientations and relaxation behaviors. Cobalt ferrite nanoparticles, with their long Néel relaxation time,⁵² are particularly promising in this regard, offering stable magnetic retention that could be advantageous in creating permanently magnetized active materials.

Currently, controlling the orientation of magnetic particles is achieved on the millimeter scale through 3D printing, where magnetic fields applied at the extruder nozzle align particles in specific configurations.^{53–56} Such approaches have been used to create millimeter-sized robots capable of controlled locomotion.^{57–59} However, scaling these capabilities down to the micrometer level remains challenging due to fabrication limitations. Alternative lithographic techniques have been

explored to induce magnetic alignment on a smaller scale, with methods such as confining a photopolymer in a mold under a magnetic field to create micro-actuators.⁶⁰ In contrast, we show that our method offers a new avenue for producing uncrosslinked, anisotropic magnetic particles embedded in a solid matrix. This opens possibilities for microscale systems, such as soft robotics, where anisotropically aligned magnetic particles could enable precise and responsive movement. Similarly, these particles hold potential in targeted drug delivery, where magnetic guidance could direct them to specific sites in the body. As research progresses, embedding techniques like ours may help overcome microfabrication constraints, paving the way for the development of remotely controlled, magnetically responsive materials at the microscale.

Author contributions

S. S.: conceptualization, methodology, investigation, and writing – original draft. N. R. M.: validation. L. R.: resources, writing – review & editing, and supervision.

Data availability

The data that support the findings of this study are openly available in 4TU.ResearchData at <https://doi.org/10.4121/f0c6a28c-9f22-430c-8c43-f2199150192b>, ref. 61.

Conflicts of interest

There are no conflicts to declare.

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