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Solvent-driven sod-ZIF-8 \leftrightarrow ZIF-C phase transformation preserves nucleic acid functionality for gene delivery

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Metal-organic frameworks (MOFs) built from zinc ions and 2-methylimidazole (HmIM) are widely explored carriers for gene delivery. Although preliminary reports show phase-dependent properties, often studies label distinct crystalline phases as “ZIF-8” and overlook how routine washing steps influence the property-to-function relationship. Here, we examine in depth how solvent history tunes the crystal phase and biological performance in plasmid DNA (pDNA) encapsulated in Zn-mIM carriers. An aqueous biomimetic mineralization at 25 °C (HmIM:Zn²⁺ = 4:1) yields defect-rich sodalite (sod) ZIF-8 with an ~80% encapsulation efficiency of the input plasmid and four times the loading obtained from an identical ethanolic synthesis. Post-synthetic solvent washes or aging govern phase transformations: media containing $\geq 70\%$ water trigger a solution-mediated transformation into a carbonate-imidazolate framework (ZIF-C), whereas absolute ethanol stabilizes sod ZIF-8 topology. Despite the extensive recrystallization processes into chemically distinct solids, pDNA loss remains $\leq 12\%$ for sod \rightarrow ZIF-C and $\leq 20\%$ for ZIF-C \rightarrow sod transformation. The green fluorescent protein (GFP) assays confirm that the recovered pDNA retains full transcriptional activity. Comparative cytotoxicity tests on PC-3 cells show that water-aged ZIF-C sustains $\geq 85\%$ viability and superior colloidal stability, while ethanol-stabilized sod-ZIF-8 possesses a higher pDNA loading and an on-demand burst release upon first contact with water. By examining how post-processing with ethanol/water yields pure sod, mixed sod/ZIF-C, and pure ZIF-C carriers, this work provides the first solvent-phase guide for MOF gene carriers and establishes simple washing protocols as tools to tune release kinetics, stability, and biocompatibility without altering the ZIF precursors or the reaction temperature. By enabling phase-engineered DNA@ZIF carriers with predictable gene-delivery performance, this study advances MOF-mediated nucleic acid therapies toward reproducible biomedical applications.

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1. Introduction

An ideal nucleic acid delivery system must protect the biotherapeutic cargo from degradation before it reaches its intracellular targets. Engineered viral vectors meet this criterion and have been widely used for nucleic acid delivery; however, they are costly to manufacture and are burdened by immunogenicity concerns.¹ These limitations have driven intensive research into non-viral delivery platforms.^{2,3} The

literature reports different particle systems used for the delivery of biotherapeutics,⁴ and, among these, metal-organic frameworks (MOFs) show strong potential in biomedicine.^{5–7} MOFs are porous hybrid materials composed of metal ions (or clusters) coordinated to organic bridging ligands.⁸ Their high surface areas, tunable pore apertures, crystallinity, and chemical versatility enable diverse biological applications.⁸ Specifically, within the MOF family, zeolitic imidazolate frameworks (ZIFs) have emerged as materials of interest for biological and biomedical uses: Zn-based ZIFs can form in water under mild aqueous conditions around biomolecules, possess thermal and chemical stability, are generally biocompatible, and the release of the biotherapeutic cargo can be triggered on demand.⁹ However, identical Zn²⁺ and methylimidazole (HmIM) precursors can yield different Zn-mIM-based crystalline structures.

Despite the use of identical building blocks, different Zn-mIM based crystalline structures possess distinct physicochemical

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properties, porosity, accessible surface area, specific volume, and chemical stability.¹⁰ Examples include ZIF-8 sodalite (**sod**), diamondoid (**dia**), and a carbonated derivative (**ZIF-C**) that, for biomedical applications, exhibit diverse encapsulation efficiencies, degradation kinetics, and release profiles, cytotoxicity, and cellular uptake.^{11,12} For example, the **sod** phase is an open, highly porous framework that can support high biomolecule loading but is comparatively less stable in aqueous or physiological media.^{13,14} In contrast, the denser **ZIF-C** phase that incorporates carbonate anions in the nonporous framework offers stability and cargo protection; these distinct properties are desirable in customized delivery systems.^{15,16} Therefore, for ZIF-based biomedical systems, fine-tuning of ZIF phases enables the precise control over crucial functional properties.

Despite growing awareness of ZIF polymorphism, the literature often labels distinct Zn-mIM crystalline materials as “ZIF-8”, overlooking fundamental structure–function relationships. In addition, post-synthesis washing steps – commonly performed with alcohols (*e.g.*, ethanol) or water to remove unreacted precursors before microscopy, X-ray diffraction (XRD), or surface area measurements – can themselves induce crystal phase changes. When this goes unnoticed or unreported, the characterized material no longer matches the material structure employed in biological studies, leading to inaccurate conclusions. Because phase governs properties central to gene and drug delivery (loading, stability, release, cytocompatibility), a clearer understanding of how routine solvent washes influence Zn-mIM biocomposites is needed.

For gene delivery, liposomes have been widely used in mRNA vaccines and siRNA therapeutics. However, colloidal instability, rapid degradation, and cargo leakage behavior in physiological fluids underscore the need to find alternative nonviral carriers. The tunable physicochemical properties of MOFs enable controlled drug release and improved storage stability.¹⁷ A concise comparison is provided in SI (see Table S1). ZIF-based carriers have been explored in cancer therapy, gene therapy, photodynamic/photothermal therapy, theranostic nanomedicines, and antimicrobial applications.^{6,18–20} However, successful translation depends on reliable synthesis and processing conditions. Prior studies have shown that phase outcome can depend on precursor concentrations, biomolecule identity, and washing solvent (water *versus* ethanol). The mechanistic understanding of solvent-induced transitions has remained incomplete. In DNA-loaded ZIFs (DNA@ZIF), precursor mass fractions and metal-to-ligand molar ratio influence crystal topology and DNA loading.^{21,22} While the washing solvent was preliminarily observed influencing the phase of DNA@ZIF systems,²¹ a systematic study of how ethanol/water wash mixtures control phase formation, recrystallization, nucleic acid retention, and biocompatibility in plasmid-loaded ZIF biocomposites is still lacking. As highlighted in recent critical review studies, progress toward clinical use will depend on closing the reproducibility gap: developing robust synthetic

protocols, standardized preparation/characterization, and systematic structure–function studies that validate MOF carriers under biologically relevant conditions.^{23,24}

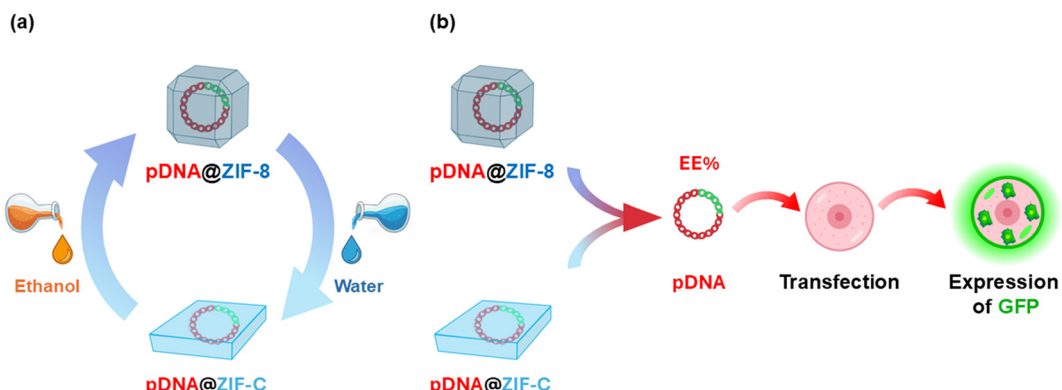
Here, for the first time, we systematically investigate the influence of mixtures of ethanol/water (EtOH/H₂O) in the synthesis of plasmid DNA-loaded ZIF particles (pDNA@ZIF) and as wash mixtures. Interestingly, while the **ZIF-C** to **sod** ZIF-8 is the main recrystallization process documented in the literature, here we observe the unnoticed recrystallization from **sod** ZIF-8 to **ZIF-C**. The concept of solvent-dependent phase transitions is summarized in Scheme 1. We observed the effect of carbonate ion and solvent polarity in directing the transformation of ZIF phases. Specifically, we (i) systematically map the crystalline phase mixtures as a function of EtOH/H₂O wash ratio, (ii) identify and quantify a mixed **sod/ZIF-C** phase window, (iii) show that exchanging wash solvents post-synthesis can drive reversible **sod** ⇌ **ZIF-C** transformations (Scheme 1a) and (iv) assess plasmid DNA loading/retention/integrity (Scheme 1b) and cytotoxicity across **sod** and **ZIF-C** phases. Importantly, we found that plasmids recovered from these recrystallized framework materials retain functional integrity for subsequent transfection into mammalian cells. The PC-3 prostate cancer cell line was selected as a representative *in vitro* model to evaluate the cytocompatibility and gene delivery efficiency. Collectively, our results demonstrate that solvent washing, as a post-synthetic treatment, is a simple and effective procedure for tuning phase and properties in pDNA@ZIF systems relevant to gene delivery and broader nanomedicine applications. A final section summarizes the results and provides guidelines addressing the development of the ZIF-based materials.^{23,24}

2. Results and discussion

The molar ratio between 2-methylimidazole (HmIM) and Zn²⁺ and the surface charge of biomolecules play a significant role in the formation of ZIF.²⁵ A typical biomimetic mineralization approach was employed to encapsulate plasmid DNA into two distinct ZIFs (*i.e.*, **sod** ZIF-8 and **ZIF-C**) according to our previous studies.^{16,26,27} A fixed 4:1 ligand-to-metal ratio was used (160 mM HmIM to 40 mM Zn(OAc)₂, total volume 200 μL) to encapsulate 5 μg of pDNA. Two distinct batches were prepared to perform the synthesis: (i) water (aqueous conditions from HmIM and Zn²⁺ dissolved in deionized (DI) H₂O, see Scheme 1a) and (ii) ethanol, EtOH (HmIM dissolved in absolute EtOH, and Zn²⁺ dissolved in 50% v/v EtOH/H₂O, see Scheme 1b).

Here, pDNA was encapsulated within ZIF-based materials in two different conditions: H₂O and EtOH. The MOF precursors (zinc acetate and HmIM) were dissolved in H₂O and EtOH, respectively.²⁵ Right after mixing the Zn²⁺ solution with the DNA/ligand solution, the suspension turned turbid, indicating rapid coordination-driven self-assembly of the Zn-mIM network.²⁸ After 10 minutes, high-speed centrifugation was used to collect the biocomposites (see SI for details). To



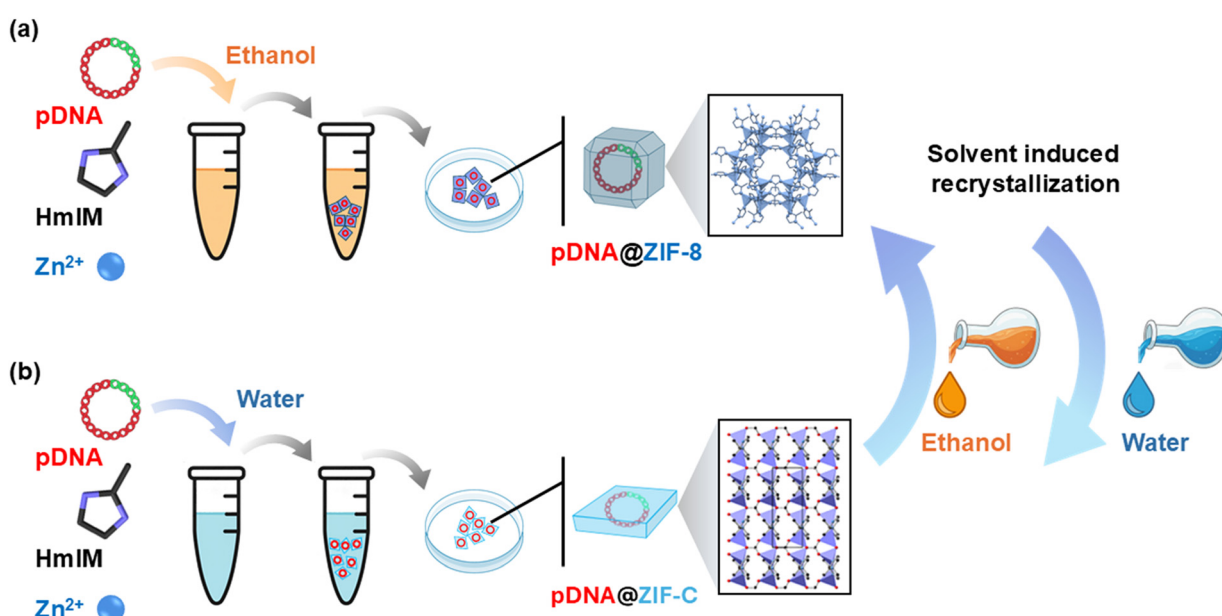


Scheme 1 Conceptual representation of the study: a) influence of the solvent (water and ethanol) on the crystalline structure of Zn-imidazolate frameworks (sod) ZIF-8 and ZIF-C and DNA plasmid (pDNA) fraction loss over $\text{sod} \rightleftharpoons \text{ZIF-C}$ re-crystallization. b) The pDNA encapsulation efficiency (EE%) and plasmid integrity via transfection and expression of green fluorescent protein (GFP) were studied.

investigate the effect of post-synthesis solvent processing, the as-formed pellets were washed three times with EtOH/H₂O mixtures containing 0–100% H₂O (v/v), using the same solvent composition for all three washes. The washed biocomposites were then vacuum-dried and are referred to in the text by the final wash solvent composition (e.g., 50% H₂O wash). This systematic preparation of solvent-wash samples (see Scheme 2) allowed us to identify the distinct contributions of (i) the solvent present during the *in situ* formation of pDNA@ZIF and (ii) the solvent during washes as post-synthetic treatments before characterization and biological testing. The sections below examine how solvent exposures can tune the crystal phase, morphology, DNA loading, and cytotoxicity.

2.1. Crystallinity and morphology analysis

The crystalline nature of the pDNA@ZIF was studied by XRD. pDNA@ZIF was prepared from an aqueous synthesis (Zn(OAc)₂ + HmIM in H₂O; L:M = 4:1). When this powder sample was washed with anhydrous ethanol (i.e., 0% H₂O), diffraction patterns characteristic of pure **sod** were obtained, confirming results from prior reports.²⁶ To examine the previously unexplored relative contribution of EtOH/H₂O mixtures on the pDNA@ZIF (**sod**) samples, these samples were washed with different EtOH/H₂O solutions with a progressive increase of H₂O (10% v/v increments). During this screening, an unexpected **sod** → **ZIF-C** phase transition was observed (Fig. 1a). Biocomposites washed with up to



Scheme 2 Schematic illustration of pDNA encapsulation within MOFs via two different methods. (a) Ethanol-mediated synthesis of pDNA@ZIF-8: pDNA is mixed with HmIM and Zn²⁺ in ethanol, leading to the formation of crystalline **sod** ZIF-8 structures encapsulating pDNA. (b) Water-mediated synthesis of pDNA@ZIF-C: pDNA is combined with HmIM and Zn²⁺ in water, resulting in ZIF-C crystals around the pDNA. The recrystallization process suggests the modular MOF crystal formation driven by ethanol and water exchange.



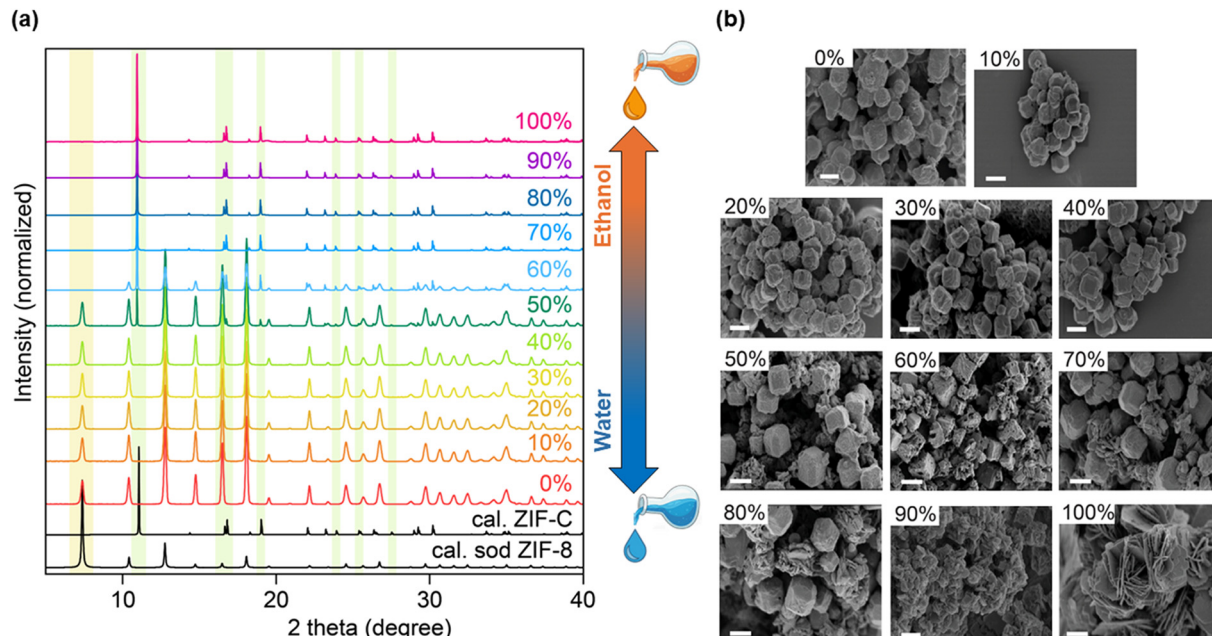


Fig. 1 X-ray diffraction patterns (a) and scanning electron microscopy images (b) of pDNA@ZIF synthesized in aqueous conditions and subsequently washed with ethanol-solvent mixtures of varying water percentages. Scale bar 500 nm.

60% water retained the characteristic **sod** topology of ZIF-8. This indicates that the **sod** phase is preserved when the washing solvent is predominantly EtOH. However, a small fraction of **ZIF-C** phase began to emerge for EtOH/H₂O = 50/50, indicating a partial **sod** → **ZIF-C** transformation and a mixed-phase structure. Starting from a H₂O = 70% or higher, all pDNA@ZIF composites transform completely into **ZIF-C**. These data indicate that the washing solvent composition regulates the ZIF crystal structure in H₂O-synthesized pDNA@ZIF (**sod**) samples. For comparison, pDNA@ZIF (**sod**) samples were synthesized by using ethanolic precursors (Scheme 1b) and exposed to the same EtOH/H₂O wash series. For these samples, the **sod** diffraction patterns indicate that the original topology was retained as the dominant topology for all washes (Fig. S1). Even at the previously observed most aggressive wash (100% H₂O), there is no appearance of the **ZIF-C** (110) peak at 11.0°. The relative **sod** peak intensities and full-width-at-half-maximum (FWHM) remain almost unchanged; only a slight overall intensity drop is evident at ≥80% H₂O, implying a trace loss of crystallinity, but the phase transition is suppressed. SEM analyses showed that pDNA@ZIF particles prepared by using EtOH as a solvent possess the classical rhombic dodecahedral morphology (Fig. S2).

SEM micrographs provide complementary morphological evidence of the phase transition (Fig. 1b). Truncated cube particles, as a typical shapes that eventually result in rhombic dodecahedral particles,²⁹ have **sod** topology that dominates at ≤40% H₂O; at 50–60% H₂O their facets become roughened and tiny plate-like crystallites nucleate on the surface, consistent with the nascent **ZIF-C** phase detected by XRD analysis. When the H₂O fraction reaches 70% or higher, the

original cubic morphology disappears, and the crystals are replaced entirely by larger, plate-like particles, which show the typical **ZIF-C** morphology.

Combined XRD and SEM analyses suggest a solution-mediated recrystallization mechanism. Peak-width evolution indicates that at 50% H₂O the nascent **ZIF-C** (110) reflection is broad (FWHM ≈ 0.43°, calculated average crystallite size: 18 nm) with a low intensity, then the (110) reflection sharpens to 0.27° (calculated average crystallite size: 30 nm) as the H₂O fraction reaches 70%, while **sod** peaks disappear. SEM shows plate-like **ZIF-C** crystallites growing on dissolving the rhombic dodecahedral **sod** particles. These data support the dissolution of the original **sod** framework followed by reprecipitation of **ZIF-C**, rather than a topotactic solid-state rearrangement.³⁰ As a weaker coordinating solvent, ethanol promotes faster nucleation by limiting Zn²⁺ solvation and yields small crystals. In contrast, water strongly coordinates with Zn²⁺ through its lone pair electrons and favors the crystal growth of ZIF-8.

2.2. Phase quantification and reactivity under washes

A web application named “ZIF Phase Analysis” was used to quantify the relative amounts of **sod** and **ZIF-C** in the biocomposites.³¹ The tool deconvolutes XRD patterns and ascribes diffraction peaks to the proper phases, examining the related intensities.³² With this, we analyzed all XRD patterns collected from the different pDNA@ZIF samples (Fig. 2). Mixed **sod/ZIF-C** profiles appeared only for the aqueous syntheses washed with 50% or 60% H₂O, whereas ≥70% H₂O yielded phase-pure **ZIF-C** and ≤40% H₂O preserved phase-pure **sod**. In contrast, particles grown in



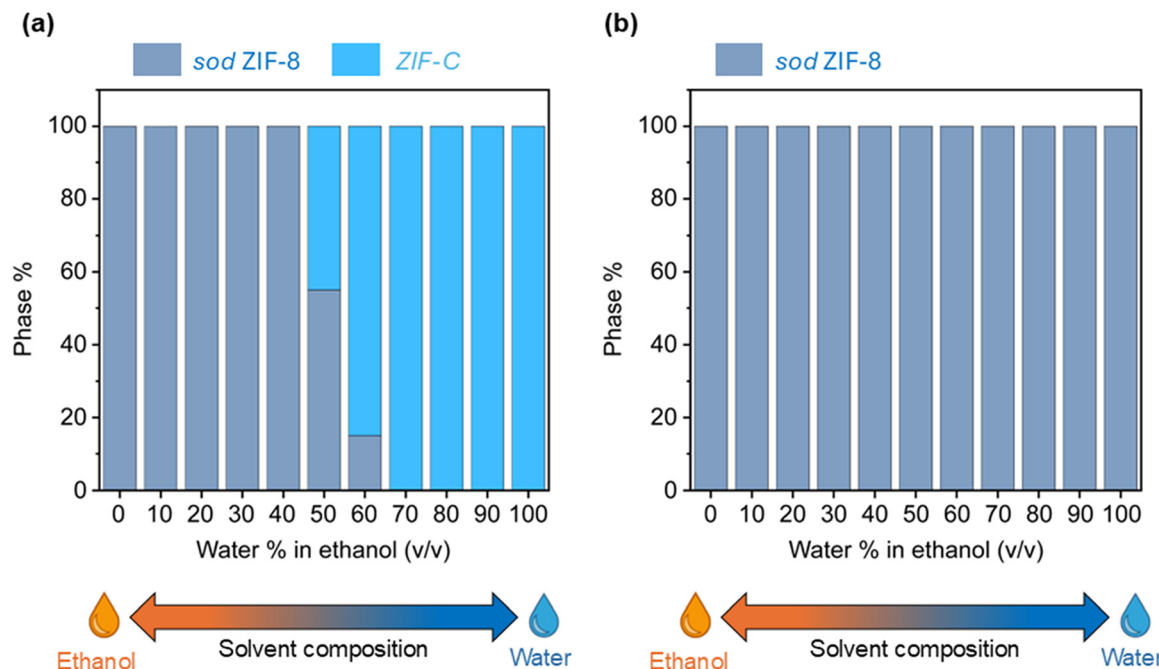


Fig. 2 Calculated phase of pDNA@ZIF prepared in (a) ethanol and (b) aqueous ZIF precursors and subsequent washing with different % of water in ethanol.

ethanol remained >99% **sod** regardless of wash composition, confirming the qualitative XRD trends discussed in section 2.1.

The markedly higher reactivity of the aqueous-route **sod** toward the **sod** \rightarrow **ZIF-C** transformation can be explained by a greater density of structural defects (e.g., missing linkers and/or under-coordinated Zn nodes) originating during rapid

nucleation in water. Specifically, water-based pDNA@ZIF (**sod**) (hereafter simply pDNA@ZIF-8) samples show broader reflections, a diffuse hump at 18–20° 2θ , and dodecahedra with a rougher surface, all supporting the presence of defects (see Fig. 2a). MOF lattice defects enhance the material network hydrophilicity and reduce the framework stability, providing vacant sites for carbonate uptake, lowering the

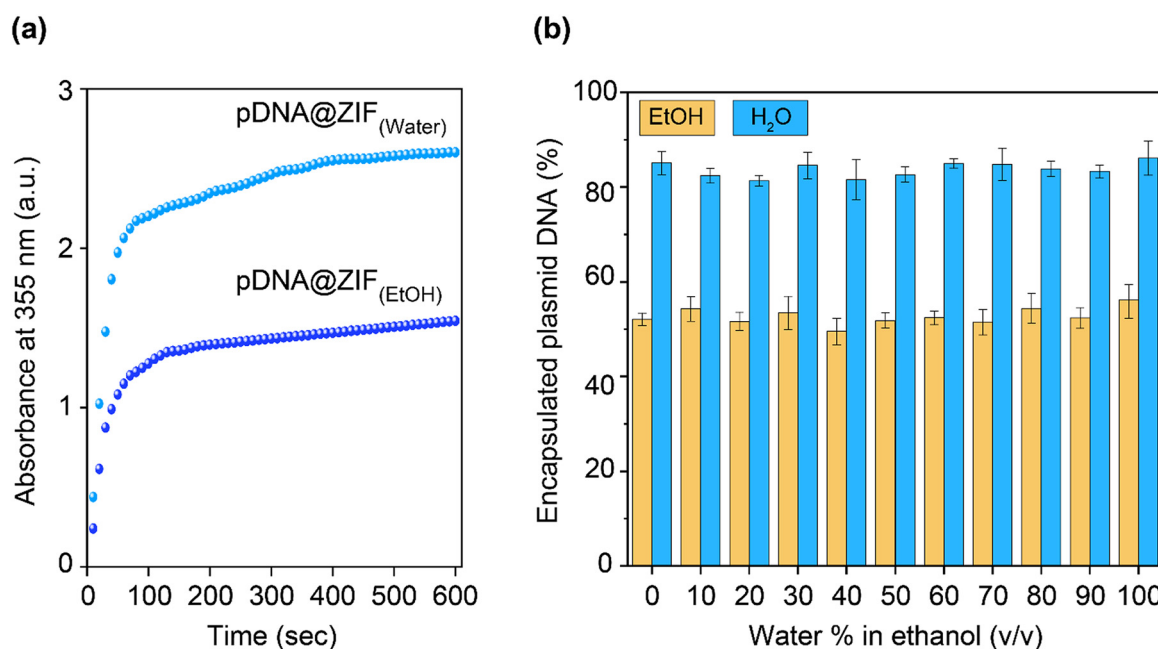


Fig. 3 (a) Growth kinetics and (b) DNA encapsulation efficiency in ZIF under aqueous and ethanolic conditions. The respective precipitates were washed with 0–100% of H₂O in EtOH (v/v). Three independent experiments were performed to test plasmid encapsulation.



barrier for **sod** → **ZIF-C** recrystallization.^{33–35} We note that defect-assisted phase transitions in crystalline materials have been documented.³⁶

By comparison, EtOH-based syntheses yield **sod** with narrower diffraction peaks and higher relative intensities. SEM micrographs show sharp, truncated cubes (Fig. 2b and S2). Collectively, these data indicate the formation of crystalline particles with fewer defects. The enhanced lattice regularity, combined with the hydrophobic nature of **sod** ZIF-8,^{37,38} further retards water penetration, carbonate diffusion in porous **sod**, and Zn ion leaching. This is consistent with the literature that experimentally demonstrates how **sod** ZIF-8 prepared in organic solvents is robust to water exposure.³⁹ Overall, our data support a defect-mediated, solution-assisted recrystallization mechanism for the **sod** → **ZIF-C** transition in pDNA@ZIF-8 particles prepared in water. In contrast, an EtOH-based synthesis yields particles with stable **sod** topology.

2.3. Particle growth kinetics and encapsulation efficiency

A spectroscopic study was performed to compare the growth kinetics of pDNA@ZIF prepared in H₂O and EtOH (Fig. 3a). Immediately after mixing precursors with pDNA, the solution turbidity at 355 nm was recorded to examine the particle formation.^{16,40} Real-time absorption measurements reveal that particles nucleate rapidly in both solvents, but a clear change in slope separates an initial rapid nucleation phase from a slower particle growth phase. The higher maximum turbidity and steeper initial slope in water indicate faster nucleation/growth and a greater particle concentration. This solvent effect is synergistic to the enhanced pDNA solubility in water: specifically, H₂O fully solvates the plasmid.⁴¹ Under these conditions, all charged seeds (pDNA) can promote the biomimetic mineralization of ZIF materials. Conversely, nucleic acids are rather insoluble in EtOH,⁴² thus the partial precipitation of pDNA reduces the heterogeneous nucleation of ZIF particles. The obtained precipitates were collected and digested with 2% nitric acid to analyze the Zn content of the biocomposites *via* microwave plasma-atomic emission spectrometer (MP-AES). A significant four-fold increase in yield was observed when biomimetic mineralization was conducted under aqueous conditions compared to ethanolic conditions, confirming the beneficial role of choosing a H₂O/pDNA system to increase the yield of pDNA@ZIF (Fig. S2).

The encapsulation efficiency (EE) of pDNA in both alcoholic and aqueous conditions was verified by agarose gel electrophoresis. After washing with different % of H₂O in EtOH, the biocomposites were digested with 20 mM EDTA to release the loaded pDNA completely. The intensity of the nucleic acid band was quantified using ImageJ and then plotted for comparison alongside other data points (Fig. 3b and S3). pDNA@ZIF prepared in H₂O possesses an average EE of 80 ± 4% across the full EtOH/H₂O wash series

(Fig. 2b and S4). Ethanolic syntheses showed consistently lower EE (~45–55%); this observation agrees with the limited DNA solubility during nucleation. H₂O is therefore the preferred medium for the preparation of pDNA@ZIF, providing higher yield and plasmid content. While the subsequent EtOH washes control the ZIF phase, this post-synthetic procedure does not influence the cargo content. With these experiments, we therefore identify guidelines specifically relevant for the preparation of pDNA@ZIF with controlled properties.

2.4. Impact of washing solvent on ZIF crystallinity

Given that pDNA@ZIF composites are routinely isolated, purified, or resuspended through multiple washes and/or solvent exchanges,^{43–45} we hypothesized that each cycle may further promote lattice reorganization. To examine possible phase transitions, we vacuum-dried EtOH-washed pDNA@ZIF-8 and H₂O-washed pDNA@ZIF-C and subjected each to five additional washes performed with pure EtOH or pure H₂O. After each washing cycle, the stability of the crystalline phase and particle morphology were investigated by PXRD and SEM, respectively (Fig. 3 and S5–S8).

The diffraction pattern of pDNA@ZIF-8 (initially washed with EtOH three times) matched the theoretical **sod** ZIF-8 diffraction pattern (see characteristic peaks at 7.34°, 10.4°, 12.8°, 16.5°, and 18.1°).^{46–48} The diffraction pattern remained unchanged after five additional washes with 100% EtOH (Fig. 4a and S4a). However, the diffraction pattern changed when pDNA@ZIF-8 was washed with H₂O (Fig. 4a and S4b). While the **sod** topology withstands the first wash, beginning with the second water rinse and continuing thereafter, a new peak at the 2θ value of 11.04° emerged. The position of this new peak can be ascribed to the (110) **ZIF-C** crystalline plane.^{15,49,50} While the intensity of the (110) **ZIF-C** diffraction peak increased with subsequent washes, the intensity of the (110) **sod** diffraction peak showed an opposite trend, indicating that **sod** progressively re-crystallized into **ZIF-C**. Thus, the diffraction patterns indicate that multiple H₂O washes result in the coexistence of **sod** and **ZIF-C**. Next, the morphologies of pDNA@ZIF-8 after five washes with pure EtOH or H₂O were investigated by SEM (Fig. 4b and S5). EtOH washes (from one to five) do not influence the truncated cube morphology, confirming the stability of pDNA@ZIF-8 particles. However, in the case of H₂O washing, the SEM images revealed that, for five washes, some original truncated cubes became holed and mixed with plate-like particles typically ascribed to **ZIF-C**. This supports the hypothesis that the phase transformation occurs *via* dissolution-recrystallization mechanisms. During synthesis, 2-methylimidazolate (mIM[–]) coordinates with Zn²⁺ cations to form the framework structure. Here, the polarity of the solvent facilitates the deprotonation of HmIM and promotes the hydrolysis of Zn²⁺ coordination. Carbonate ions of water play the role of phase transformation. Falcaro and co-workers showed that washing treatment induces phase



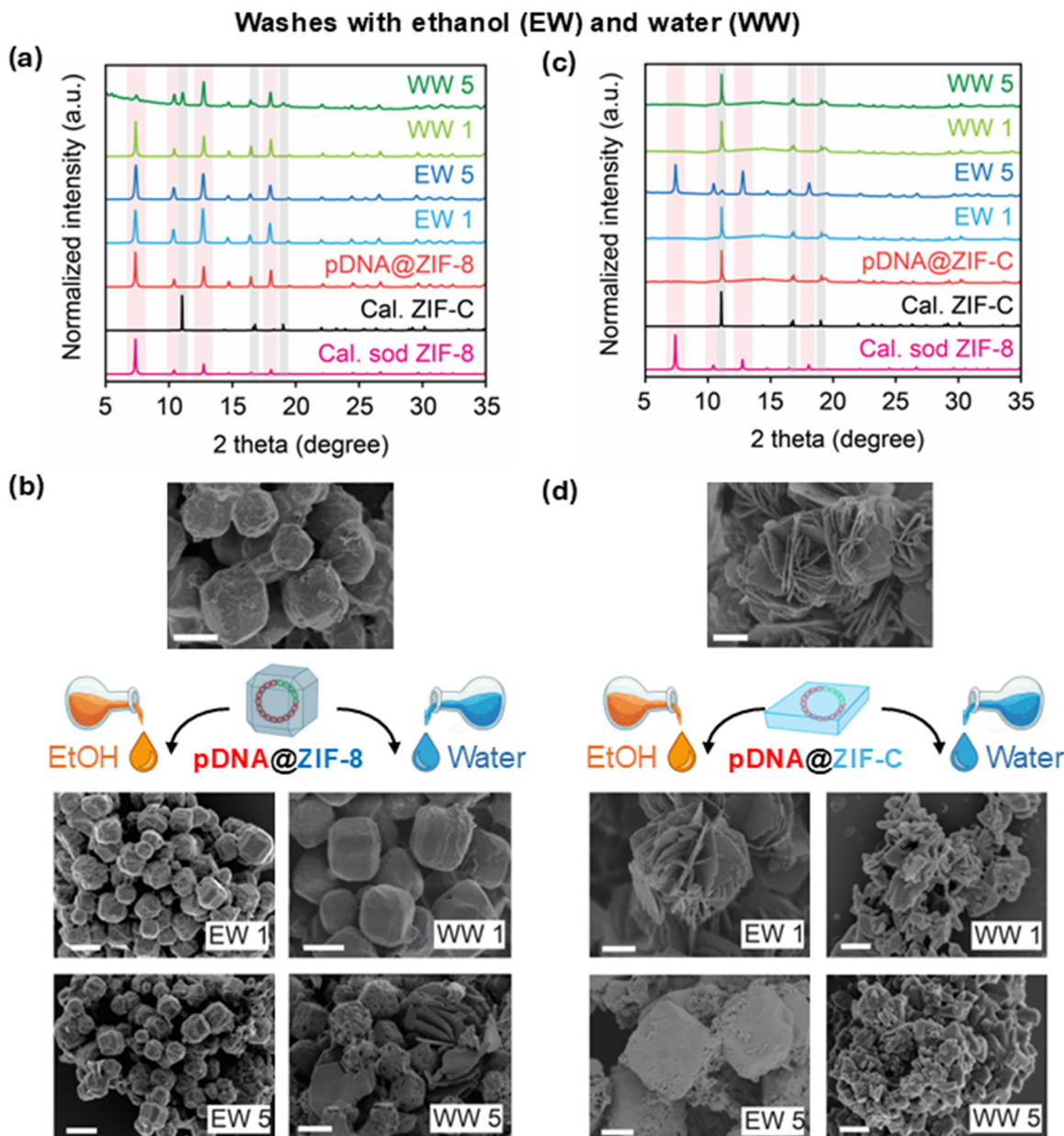


Fig. 4 Crystallinity and morphology assessment of DNA@ZIF-8 (a and b) and DNA@ZIF-C (c and d) after subsequent ethanol wash (EW) and water wash (WW). The numbers represent the washing steps. Scale bar: 500 nm.

transformation of ZIF biocomposite.³⁰ Low CO₂ solubility in water allows the incorporation of one CO₃²⁻ with one Zn²⁺ per formula unit to form ZIF-C (Zn₂(mIM)₂CO₃). On the other hand, solubilization of atmospheric CO₂ during ethanol washing modifies the coordination equilibrium (Zn-mIM-CO₃²⁻) in ZIF-C by removing CO₃²⁻ to yield a carbonate-free **sod** framework (Zn(mIM)₂).

Then, we examined pDNA@ZIF-C (initially washed with water, 3 times) by using the same procedure. During EtOH washing, the diffractogram of pDNA@ZIF-C did not show changes until the 2nd wash (Fig. 4c and S6). However, after the 3rd EtOH diffraction peaks of **sod** ZIF-8 emerged, subsequent EtOH washes resulted in a progressive increase

of the intensity of the **sod** peaks and a decrease of (110) ZIF-C. This trend highlights how EtOH washes can trigger the progressive phase transformation of pDNA@ZIF-C into pDNA@ZIF-8. Conversely, water washes do not trigger any effect on the original ZIF-C phase (Fig. S6). Then, the morphology of pDNA@ZIF-C after this washing procedure was examined *via* SEM (Fig. 4d and S7). The pDNA@ZIF-C maintained typical plate-like morphology up to 2nd EtOH wash, while subsequent washes yielded a drastic change in morphology (formation of truncated cubes), indicating the ZIF-C → **sod** recrystallization. We note that this phase transformation is followed by a variation of the original particle size, a particle property that in drug delivery systems

should be judiciously related to the planned administration route.⁵¹ The plasmid encapsulated **sod** ZIF-8 and ZIF-C followed phase transformation due to the dissolution-recrystallization mechanism.³⁰ Competitive coordination of water and carbonate ions can destabilize the **sod** framework and promote the formation of the denser ZIF-C phase.

2.5. Stability of ZIF crystalline phase

After synthesizing pDNA@ZIF particles, the investigation of their stability is critical to ensure the consistency of their

properties over time. Despite the standardization of protocols being an urgent matter in MOF biomedicine^{23,24} and the wide attention given to ZIF systems as drug carriers, at the moment there are no studies examining the stability of pDNA@ZIF systems. Motivated by this gap and by some early reports showing how some pure **sod** ZIF-8 is converted by water to particles with an unknown crystalline phase with leaf-like morphology,⁵² we examine the stability of pDNA@ZIF-8 and pDNA@ZIF-C when stored in EtOH and DI H₂O (static conditions in sealed vials) for hours (*i.e.*, 1, 3, 5, and 24 h). After incubation, the particles were

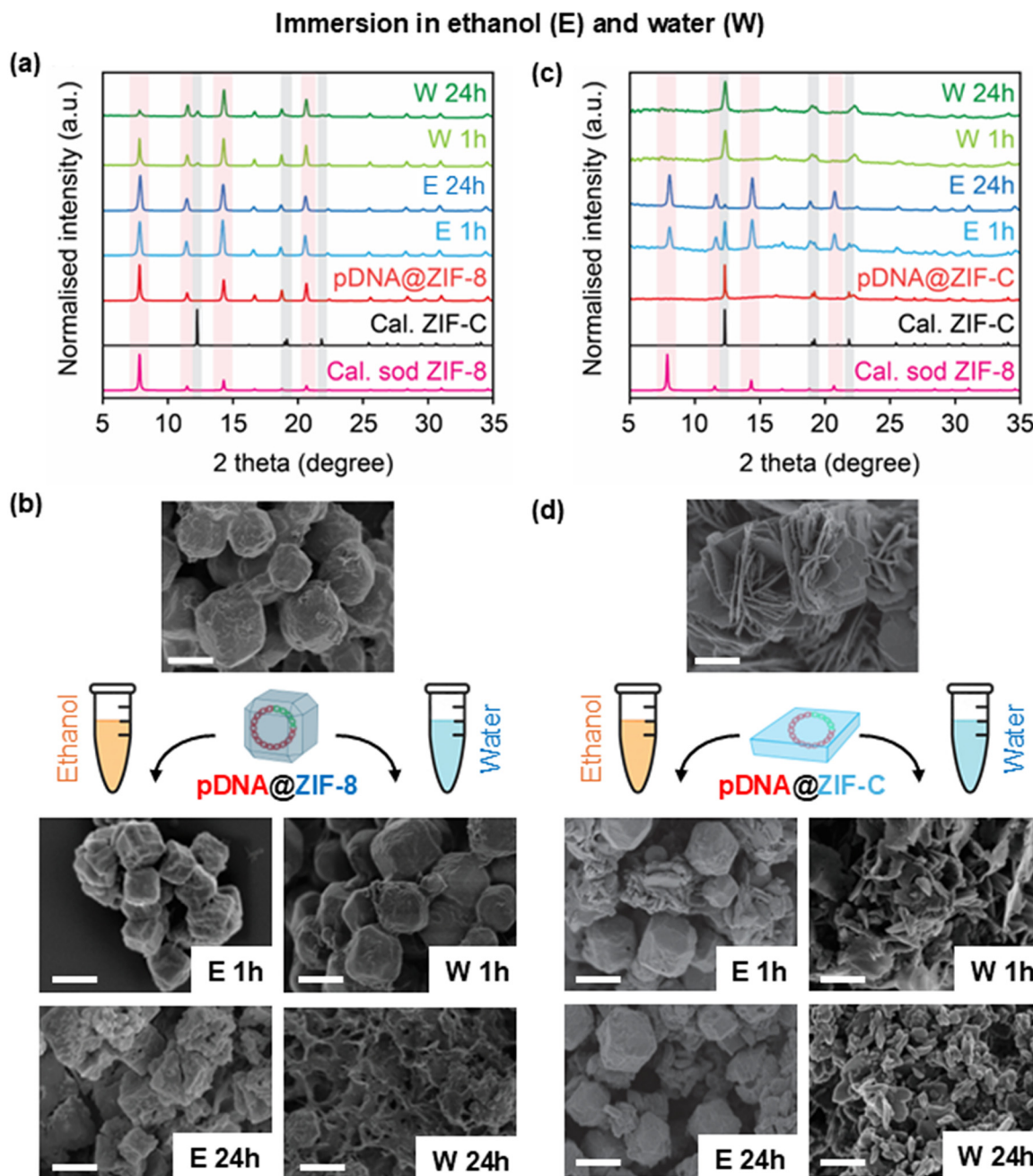


Fig. 5 Stability assessment of pDNA@ZIF-8 (a and b) and pDNA@ZIF-C (c and d) after incubation in ethanol (EtOH, E) and water (H₂O, W). The powder material incubated at different times was collected and analyzed via (a and c) XRD and (b and d) SEM. Scale bar: 500 nm.



centrifuged, collected, and dried for XRD analysis (Fig. 5a and S9). When immersed in EtOH (Fig. 5a and S9a), the diffraction pattern of pDNA@ZIF-8 remained invariant, thus fully retaining **sod** phase within 24 h. However, when pDNA@ZIF-8 was immersed in H₂O (Fig. 5b and S9b), a 1-hour exposure produced a weak peak of (110) **ZIF-C** ($2\theta = 11.04^\circ$), indicating the beginning of a **sod** \rightarrow **ZIF-C** transformation. The intensity of such a peak increases over time and becomes 20-times more intense after 24 h immersion.

The electron micrograph of representative pDNA@ZIF-8 particles exposed to EtOH revealed a truncated cube morphology (Fig. 5b and S10a). This observation suggests the long-term stability of the **sod** phase in EtOH. Conversely, when the same pDNA@ZIF-8 particles were incubated in H₂O for more than 3 h (Fig. 5b and S10b), the emergence of the typical plate-like **ZIF-C** shape appeared alongside the feedstock cubic **sod** ZIF-8. This observation is consistent with the changes observed in the XRD patterns.

Next, we focused on examining the effect of EtOH on pDNA@ZIF-C particles and we performed identical solvent/time exposures (pure H₂O and EtOH for 1, 3, 5, and 24 h). While pDNA@ZIF-C maintained its crystallinity in H₂O for the entire period (*i.e.*, 24 h), during EtOH incubation (Fig. 5c and S11b) the (110) **ZIF-C** peak showed a gradual intensity reduction with the immersion time. The decrease in intensity coincided with a broadening of the FWHM, indicating a reduction of crystalline domain size. The appearance of rising peaks at 7.34° and 33° indicated the formation of **sod** (ZIF-8) and zinc hydroxide.^{53,54} The detailed mechanism behind this phase transformation and the formation of Zn byproducts requires further investigation. Electron micrographs show a change in pDNA@ZIF-C morphology that is triggered by EtOH exposure (Fig. 5d and S12a). Plate-like **ZIF-C** particles decreased in number and size, transforming into cubic crystals. Interestingly, pDNA@ZIF-C in water rearranged into an interconnected particle network (Fig. 5d and S12b).

2.6. Encapsulated DNA content in the phase transformed ZIF

ZIF has been confirmed to be an effective therapeutic carrier for DNA delivery.⁵⁵ However, as we show here that **sod** \rightleftharpoons **ZIF-C** transformations occur *via* dissolution and recrystallization, it is crucial to determine whether the carrier reorganization compromises the pDNA payload and integrity. Thus, we tested both washed and prolonged exposures to EtOH and H₂O.

Washes. To quantify any loss, we recovered pDNA@ZIF-8 (**sod**) and pDNA@ZIF-C after five washes that, according to the experiments described above, triggered the phase conversions (*i.e.*, H₂O washes for **sod** and EtOH washes for **ZIF-C**), digested the frameworks with 20 mM EDTA, and analyzed the released plasmid *via* agarose electrophoresis (Fig. S13). The loading content was determined *via* densitometry using ImageJ software. These analyses revealed

that for pDNA@ZIF-8, the partial **sod** \rightarrow **ZIF-C** conversion removed $\sim 8\%$ of the pDNA, whereas the reciprocal pDNA@ZIF-C with **ZIF-C** \rightarrow **sod** recrystallization reduced the pDNA by 7% (Fig. S13). Additionally, electrophoresis showed single, unsmeared bands in each lane. This indicates that the plasmid remained intact under these partial dissolution and recrystallization cycles with minimal pDNA reduction measured (Δ pDNA $< -8\%$).

Extended exposures to solvent (up to 24 h). pDNA@ZIF-8 and pDNA@ZIF-C were incubated in H₂O and EtOH for up to 24 h. In pDNA@ZIF-8 exposed to water for 3 h, $\sim 10\%$ of the pDNA was lost from the solid material (Fig. S14a and b) and it plateaued to $\sim 12\%$ loss for the remaining examined times (5 and 12 h). For pDNA@ZIF-C, the exposure to EtOH resulted in a decrease of $\sim 15\%$ after a 5 h incubation and $\sim 20\%$ after 24 h (Fig. S14c and d). The finding confirms that some ZIF recrystallization processes can influence the final fraction of pDNA encapsulated in ZIF systems; however, the 80% DNA retention indicates that the vast majority of functional nucleic acid is available for delivery applications if the ZIF biocomposite is administered within 24 h from its preparation.

Phase transformation. We hypothesize that the conversion of **sod** ZIF-8 to **ZIF-C** occurs under aqueous conditions, where H₂O molecules facilitate a phase transformation by interacting with the coordination bonds between zinc ions and HmIM ligands. Atmospheric CO₂ yields CO₃²⁻ dissolved in DI H₂O that serves as a building block for the formation of a carbonated framework. The H₂O interaction and the presence of carbonate anions promote ligand rearrangement and result in the formation of **ZIF-C**, a denser and more thermodynamically stable phase.⁴⁹ Conversely, exposure of **ZIF-C** to EtOH induces a reverse transformation back to **sod** ZIF-8. EtOH disrupts the stabilized coordination environment in **ZIF-C**, enabling re-coordination of zinc ions into the ZIF-8 framework.³⁰ This reversible phase transition is driven by the contrasting polarities and coordination abilities of water and ethanol, as well as the markedly different solubilities of CO₂ in the two solvents. Interestingly, the recrystallization occurs without a significant loss of the encapsulated payload, making it suitable for stimuli-responsive delivery applications.

Long-term stability. The stability of pDNA@ZIF-8 and pDNA@ZIF-C was assessed after 7 days of storage under identical experimental conditions. At room temperature, the dried powder samples were stored in a sealed container to prevent exposure to moisture and contaminants. XRD of the aged samples confirms that the material retains the structure after phase transformation (Fig. S15). Both phase-transformed **sod** and **ZIF-C** structures remained unchanged after 7 days. This finding addresses an interesting point, as ZIF acts as a stable carrier for DNA delivery. Further comprehensive studies, such as validating the performance of loaded cargo, are required to advance clinical translation of ZIF-based non-viral DNA delivery.



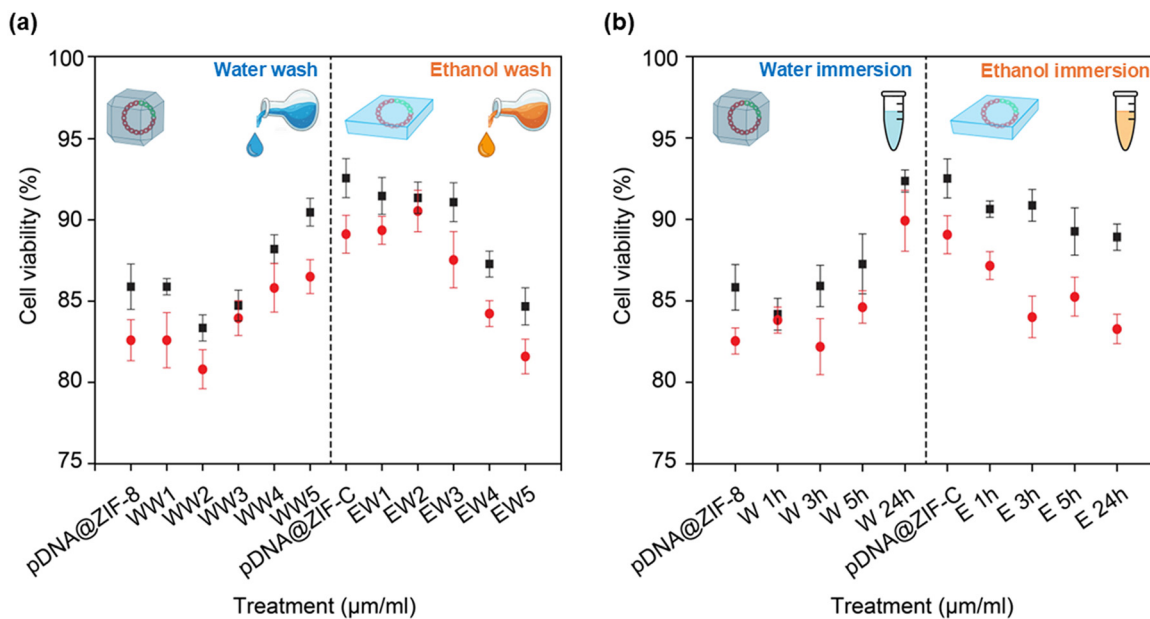


Fig. 6 Cell viability of PC-3 cells after 24 h of incubation. (a) pDNA@ZIF-8 and pDNA@ZIF-C particles were collected after incubation in water (W) and ethanol (E) at 1, 3, 5, and 24 h. (b) pDNA@ZIF-8 and pDNA@ZIF-C particles were collected after water wash (WW) and ethanol wash (EW) up to 5 times. Here, two different concentrations (e.g., ■ 25 and ● 50 µg mL⁻¹) were used for concentration-dependent effect analysis. Data represent the average of three independent experiments. Untreated cells were considered 100% viable. No statistical significance was found.

2.7. Cytotoxicity

A viable gene-delivery vehicle combines efficient payload delivery with minimal cytotoxicity. We therefore evaluated the cytotoxicity of the pDNA-loaded biocomposites against PC-3 cells using the MTT reduction assay. In living cells, the MTT, a yellow tetrazole, is reduced to purple formazan by the activity of mitochondrial dehydrogenases. Therefore, the MTT reduction assay reflects the redox activity of living cells. Fig. 6 shows the percentage of cell viability after a 24 h exposure to pDNA@ZIF-8 and pDNA@ZIF-C. This duration is used in preliminary cytotoxicity assessments of MOFs, and it is sufficient for materials to interact with cells for transfection studies that typically use 1–3 h of exposure time for cellular uptake.^{16,56}

Fig. 6a shows the effect of successive washes with H₂O and EtOH. When pDNA@ZIF-8 was synthesized and repeatedly washed in H₂O, cell viability remained $\geq 85\%$ through all five cycles. An almost identical profile was obtained for the EtOH-washed series, with the lowest value of 82% recorded after the fifth wash. When referring to the aforementioned structural characterization, the 5th wash correlates with **sod** ZIF-8 with fewer defects. Thus, when considering preliminary reports on the cytotoxicity of pure ZIF-C and **sod** ZIF-8,^{15,49} we hypothesize that the higher the content of highly crystalline **sod** ZIF-8 with respect to ZIF-C, the higher the cytotoxicity.

Fig. 6b examines the effect of incubation time on cytotoxicity. When for pDNA@ZIF-8 the process of synthesis, wash, and aging was entirely performed in H₂O, cell viability remained $\geq 85\%$ across all steps. A slightly enhanced cell

viability was observed after the third and fifth washes (**sod** \rightarrow ZIF-C), while an additional 5% drop was observed for the samples stored in EtOH (more defective **sod** \rightarrow less defective **sod**). These data further support the lower cytotoxicity of ZIF-C compared to **sod** ZIF-8 in mixed ZIFs phases and show how defects in **sod** ZIF-8 can slightly enhance cell viability. Despite these measurable changes in viability, all the ZIF-based composites tested in Fig. 6 exhibited viability values above 70%, a threshold defined by ISO 10993-5 standard to formally distinguish non-cytotoxic from cytotoxic materials.⁵⁷

2.8. Bioactivity analysis

To verify that pDNA retained its functional integrity after encapsulation, solvent handling, and phase conversion, we performed a transfection in PC-3 cells to express green fluorescent protein (GFP). Because the different MOF matrices (**sod** ZIF-8, ZIF-C, or their mixtures) could themselves influence cellular uptake, we first released pDNA from the ZIF-based particles with EDTA and then loaded pDNA in Lipofectamine to use a standard transfection protocol. This release and reload approach aims at the assessment of the pDNA functionality by removing any carrier-specific effects.

The transfection efficiency was quantified by examining GFP-positive PC-3 cells 24 h after transfection with a confocal laser scanning microscope (CLSM). Fluorescence data for transfections by plasmid recovered from particles that had undergone (i) one to five sequential washes in H₂O or EtOH and (ii) 1, 3, 5 or 24 h aging in the same solvents is shown in Fig. 7. Untreated cells (negative control) showed no green



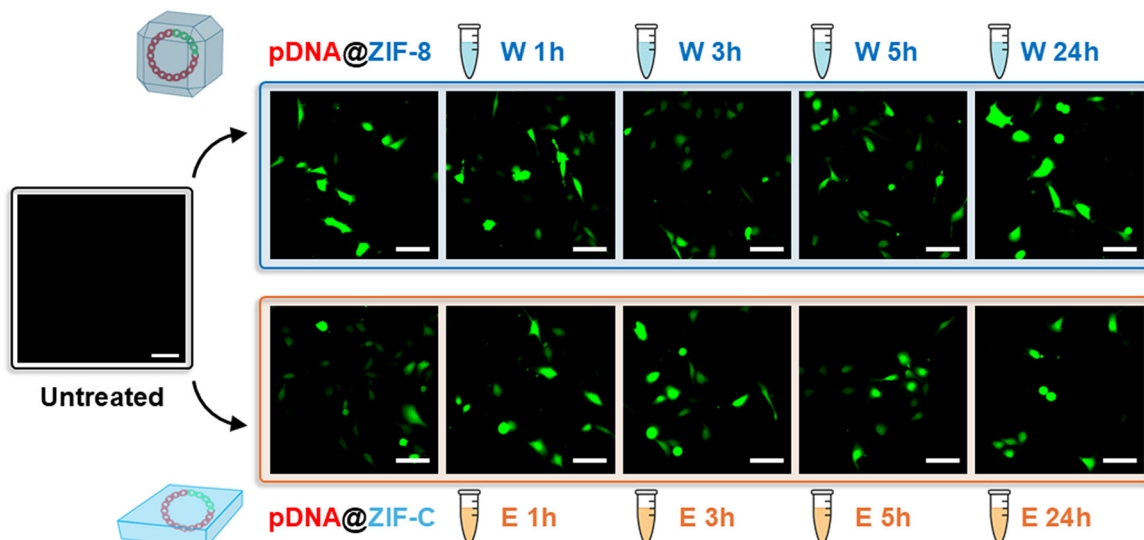


Fig. 7 PC-3 cell image transfected with GFP-expressing plasmid DNA. Both pDNA@ZIF-8 and pDNA@ZIF-C particles were subjected to water (W) and ethanol (E) incubation (1–24 h). Cells treated with only media were used as untreated control. Scale bar 100 μ m.

fluorescence, whereas all cells receiving released plasmid emitted the typical GFP signals (Fig. 7 and S16). The fluorescence intensity and the fraction of positive cells ($\approx 40\%$ across the data set) were statistically indistinguishable from those obtained with pDNA extracted from freshly synthesized and never-washed pDNA@ZIF-8. The experiment demonstrates that the encapsulated plasmid retained full transcriptional and translational functionality irrespective of solvent history. Here, the release-and-reload strategy confirmed the functional integrity of encapsulated nucleic acid, which remained transcriptionally competent after solvent-mediated phase change. Therefore, this approach provides an unambiguous indication that solvent treatments did not cause loss of function of encapsulated nucleic acid.

2.9. Comparative summary and recommendations

When comparing water- and ethanol-based syntheses, an EtOH solvent yields pDNA@ZIF-8 with a higher crystallinity, a much lower encapsulation efficiency (EE $\approx 45\text{--}55\%$), and an inferior yield of solid material. Thus, the following comparison focuses on carriers prepared in water, where fully dissolved pDNA acts as an effective nucleation seed for biomimetic mineralization. When pDNA@ZIF is prepared in water at a 4:1 = L:M ratio (*i.e.*, 40 mM Zn(OAc)₂·2H₂O and 640 mM HmIM added to 1 mL pDNA solution), a defect-rich, cubic **sod** ZIF-8 biocomposite with EE = $80 \pm 4\%$ is obtained. Storage in water (≤ 1 h) preserves the **sod** topology and yields the maximum plasmid density per gram of ZIF. If the suspension is then brought into contact with a water predominant environment ($\text{H}_2\text{O} \geq 70\%$), defects favor a solution-mediated re-crystallization into the denser **ZIF-C** polymorph. In 20 h the recrystallization is complete. The phase change releases no more than 12% of the initially

captured pDNA (payload retained $\approx 88\%$). However, such a pDNA release that accompanies **sod** \rightarrow **ZIF-C** can afford the delivery of higher absolute amounts of pDNA during the first hours after administration. On the other hand, **sod**-based carriers that remain suspended in pure water without completing the transition lose up to 20% pDNA over the same interval, most likely through enhanced diffusion in defects rather than wholesale lattice rearrangement. Exposure to ethanol converts ZIF-C ($\text{Zn}_2(\text{mIM})_2\text{CO}_3$) into the carbonate-free **sod** phase ($\text{Zn}(\text{mIM})_2$) by removing Zn^{2+} and CO_3^{2-} from the framework, a process influenced by how carbonate ions interact with each solvent. Ethanol, despite the higher CO_2 solubility compared to water, does not generate carbonate ions due to its weaker basicity, which helps to drive the recrystallization toward the **sod** phase.³⁰ On the other hand, water dissolves CO_2 to form carbonate and bicarbonate ions. In this case, the thermodynamics of the system is driving the **sod** to ZIF-C conversion. In fact, Navrotsky and co-workers evaluated the thermodynamic driving force for **sod** and **ZIF-C** topologies.⁵⁸ ZIF-C ($\Delta H = -87.88 \text{ kJ mol}^{-1}$) is thermodynamically more stable than the **sod** ($-21.04 \text{ kJ mol}^{-1}$) phase and the transformation between ZIF phases (*e.g.*, **sod** \rightleftharpoons **ZIF-C**) is driven by Gibbs free energy.

Immediate solvent exchange to anhydrous EtOH (three consecutive 1 mL washes, same centrifugation parameters) removes water from the porous frameworks, recovers defects, and locks the framework in the **sod** topology. The resulting particles are smaller, with sharper cubic features, and exhibit a slightly lower PC-3 cell viability floor ($\approx 82\%$) than their water-aged, **ZIF-C** counterparts ($\geq 85\%$), yet they retain the full plasmid payload as long as ethanol remains the continuous phase. Notably, re-introducing $\geq 70\%$ water at any later point restarts the **sod** \rightarrow **ZIF-C** conversion and again generates a rapid plasmid-release pulse, demonstrating full reversibility of the solvent trigger.



3. Conclusion

This study addresses reproducibility issues in ZIF-based gene delivery by demonstrating that the crystal structure, defects, and biological properties of plasmid-loaded zinc-imidazolate frameworks (pDNA@ZIF composites), for the same initial stoichiometry, can be controlled by the solvent exposure protocol applied. This includes the post-preparation washes and storage conditions. Using a single, scalable aqueous biomimetic mineralization protocol at a fixed 4:1 HmIM:Zn²⁺ ratio, we establish a solvent-phase map in which routine washing and storage solvents alone define the final material: media that remain $\leq 60\%$ water preserve a defect-rich **sod**-ZIF-8, whereas $\geq 70\%$ water drives a previously unreported, vacancy-assisted reconstruction into carbonate-containing ZIF-C, and ethanol reverses the process on demand. These **sod** \rightleftharpoons ZIF-C transitions occur with low plasmid loss ($\leq 12\%$ during **sod** \rightarrow ZIF-C and $\leq 20\%$ for the reverse step), whereas pDNA maintains transcriptional activity. By correlating X-ray patterns, SEM micrographs, and MP-AES data, we show that the higher vacancy density from the aqueous synthesis facilitates the solution-mediated transformation, while ethanol-grown **sod** particles with fewer defects remain phase-invariant. Biological assays reveal complementary advantages: post-synthesis ethanol-washed **sod**-ZIF-8 offers the greatest pDNA density and a solvent-triggered burst release, whereas H₂O-aged ZIF-C provides superior colloidal stability and slightly higher cell viability. Because both Zn-mIM polymorphs originate from the same precursors and reaction conditions, researchers can now choose the washing solvent or procedure as a facile approach to finely tune the particle properties (e.g., morphology, DNA loading, cytocompatibility) without altering the synthesis. Therefore, these findings shed light on solvent washes, as a step often relegated to a one-line procedural detail that is here revealed to be a critical design parameter for ZIF biocomposites. Reporting details about washing conditions is therefore essential for the reproducible development, comparison, and clinical translation of future ZIF-based nucleic-acid delivery systems.

4. Materials and methods

4.1. Required chemicals

Zinc acetate dihydrate (Zn(OAc)₂·2H₂O) and HmIM were purchased from Sigma-Aldrich (Australia). PureLink™ plasmid kit, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), and Lipofectamine 3000 were purchased from ThermoFisher Scientific. PC-3 prostate cancer cells were kindly provided by Prof John Mariadoson's lab in the Olivia Newton-John Cancer Research Institute. A 6.5 kbp length plasmid DNA was collected from Sir Ian Potter NanoBiosensing Facility, RMIT University, Melbourne, Australia.

4.2. Preparation of pDNA@ZIF

Organic preparation includes the addition of 5 μ g of plasmid DNA into 100 μ L of HmIM (160 mM) in ethanol. A separate solution of Zn(OAc)₂ (100 μ L, 40 mM) in 50% ethanol (v/v) was also prepared and added. The mixture was aged for 10 minutes at room temperature. The obtained precipitate was recovered by centrifugation at 13 000 rcf for 12 minutes.

Similarly, inorganic preparation includes the addition of 5 μ g plasmid DNA to 100 μ L HmIM (160 mM) in deionized water. A separate solution of Zn(OAc)₂ (100 μ L, 40 mM) in water was also prepared and added. The mixture was aged for 10 minutes at room temperature. The obtained precipitate was recovered by centrifugation at 13 000 rcf for 12 minutes.

4.3. Preparation of washing solvents and washing treatment

A series of washing solutions were prepared by increasing the water% in ethanol (v/v). The DNA, HmIM, and Zn(OAc)₂ mixtures were aged for 10 minutes at room temperature. The formed biocomposites were recovered by centrifugation at 13000 rcf for 12 minutes. The supernatant was separated to remove unreacted precursors and ions. Then the pellet was repeatedly washed with 200 μ L of washing solvents (0–100% H₂O) and centrifuged thrice in the same solvents to examine the influence of water content in the washing solvent on determining the crystal fate of the plasmid@ZIF biocomposite. Finally, the biocomposites were vacuum-dried overnight.

4.4. Characterization

X-ray diffraction (XRD) was performed in Bruker D8 General Area Detector Diffraction System (GADDS) using Cu K α ($\lambda = 1.54056 \text{ \AA}$) radiation at 40 kV generator intensity and 40 mA current. The 2θ scanning range was 5–60° and the step size was 0.01°. The vacuum-dried samples were mounted on a surface clean silicon wafer attached to a sample holder and placed into the instrument. All the spectra were recorded at room temperature. The acquired (.raw) data was converted to a UXD file using the File Exchange Program XCH (Ver. 5.0.10, 2004, Bruker AXS, Socabim, Karlsruhe, Germany). Following that, the file was converted to Text Documents (.txt) and plotted using the OriginPro software. Crystal structure data (.cif) of simulated **sod** ZIF-8 and ZIF-C patterns were collected from previous literature and processed in Mercury (3.8).

For electron microscopy, a diluted biocomposite solution was dropped on a clean silicon wafer surface and air-dried to remove the solvent. Then 5 nm iridium coating was applied using Leica EM ACE600 Sputter Coater to enhance conductivity under Scanning Electron Microscope (SEM) measurements. FEI Verios 460 L SEM was used to characterise the morphology of samples. The accelerating voltage and current of the electron beam were 2 kV and 0.2 nA, respectively. All tests were performed in triplicate to quantify the variability and reproducibility.



4.5. Growth kinetics study

To gain additional insight into the growth kinetics of pDNA@ZIF, we conducted turbidity measurements using UV-vis absorption spectroscopy. The optical turbidity of the mixture, both in organic (*e.g.*, ethanol) and inorganic (*i.e.*, aqueous) conditions, was measured by recording the absorption value at 350 nm position for up to 600 seconds.

4.6. Encapsulation efficiency determination

The DNA loading in the water- and ethanol-based preparation of ZIF was investigated by agarose gel electrophoresis. The prepared pDNA@ZIF (both in ethanol and aqueous ZIF precursors) was washed with a range of solvents with different water % in ethanol (v/v). The final biocomposites were dissolved in 200 μ L of aqueous EDTA (20 mM) to break the ZIF shell and release DNA in the solution. All samples were loaded on pre-heated 1% agarose gel containing SYBR Safe stain alongside an equal quantity of DNA used during synthesis as control. The electrophoresis was carried out in 1 \times TAE buffer for 120 min at 90 V and bands were visualized in the GelDoc® system (BioRad, Hercules, USA). The intensity of DNA bands was calculated using densitometry analysis by ImageJ software, and encapsulation efficiency (EE) of DNA was determined by the following formula:

$$\text{EE (\%)} = \frac{\text{Intensity of encapsulated pDNA}}{\text{Intensity of total pDNA added}} \times 100$$

4.7. Zn yield determination

The centrifuged pDNA@ZIF pellet was separated and then dried under a vacuum. Next, 2% nitric acid was applied to dissolve the ZIF carrier and to quantify the Zn content *via* an Agilent 4200 Microwave Plasma-Atomic Emission Spectrometer.

4.8. Effect of washing solvent on ZIF phases

The effect of the washing solvent on ZIF-8 particles was investigated by rinsing the prepared pDNA@ZIF-8 five times with either 500 μ L ethanol or deionized water. Similarly, the effect of washing solvent on ZIF-C particles was investigated by washing the prepared pDNA@ZIF-C five times with either deionized water or ethanol. At each washing step, the ZIF pellet was recovered by centrifugation and dried for further analysis.

4.9. Stability of ZIF phases

ZIF biocomposites were prepared by mixing 5 μ g plasmid DNA into 100 μ L aqueous HmIM (160 mM) and 100 μ L aqueous Zn(OAc)₂ (40 mM). After 10 minutes the obtained precipitate was recovered by centrifugation and washed with absolute ethanol and deionized water to form pDNA@ZIF-8 and pDNA@ZIF-C. The stability of the biocomposites was investigated by keeping the ZIF particles in 500 μ L ethanol and deionized water in static conditions. At each time

interval (*i.e.*, 1, 3, 5, and 24 h) the ZIF pellet was recovered by centrifugation and dried for further analysis.

4.10. Cytotoxicity assessment

PC-3 cells were seeded in a density of 10 000 cells per well in 96 well plates and incubated for 24 h at 37 °C. Later, cells were treated with 25 and 50 μ g mL⁻¹ of pDNA@ZIF in a fresh RPMI medium containing 10% fetal bovine serum and 1% penicillin-streptomycin. The cells were then incubated at 37 °C and then the media was aspirated after 24 h. Immediately, 100 μ L of serum-free media containing MTT reagent were further incubated at 37 °C in the dark. After 4 h, the MTT media was aspirated carefully without disturbing purple formazan crystals formed at the bottom of the well. The purple formazan crystals were dissolved in 100 μ L DMSO, and the absorbance was measured by a microplate reader at 570 nm. Pristine sod ZIF-8 and ZIF-C were used as negative controls. The percentage of cell viability was calculated by the formula:

$$\text{Cell viability (\%)} = \frac{\text{Absorbance in sample}}{\text{Absorbance in control}} \times 100$$

4.11. Bioactivity assessment

The 6.5 kbp plasmid DNA used in this study contains a gene responsible for producing a green fluorescent protein (GFP). While the plasmid itself doesn't glow, once it undergoes transcription and translation processes within the cell's machinery, it generates a harmless GFP. When exposed to blue light, this GFP emits a vibrant green fluorescence. Thus, expression of green fluorescence under blue light indicates successful uptake of material followed by transcription and translation of the coded gene. A cancerous human prostate cell line (PC-3) was used in this study to investigate the bioactivity of the encapsulated nucleic acid in ZIF. The PC-3 cells were treated with pDNA@ZIFs and were examined live to investigate the expression of GFP in cells using CLSM (NSTORM SuperResolution/Confocal microscope, Nikon). Prior to treatment, the pDNA@ZIFs were treated with sterile EDTA (0.5 M, pH 8.0) and the released DNA was encapsulated in the Lipofectamine 3000 reagent. After 24 h of transfection, cells were washed with sterile 1 \times PBS (pH 7.4) thrice. Live cell imaging was employed to record the expressed GFP in untreated and treated cells using CLSM.

4.12. Statistical analysis

All tests were performed in triplicate. Statistical differences in the properties of different samples were analyzed using One-Way ANOVA in GraphPad Prism.

Conflicts of interest

There are no conflicts to declare.



Data availability

On behalf of all authors, I'm confirming that all the data presented in this work are original and the data supporting this article have been included as part of the supplementary information (SI). Supplementary information is available. See DOI: <https://doi.org/10.1039/d5lf00306g>.

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