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Enhancement of high concentration aqueous iron oxide nanoparticle ink printability through well-defined polymer additives

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Polymer additives play a crucial role in modifying the stability and rheology of ceramic nanoparticle suspensions. A library of anionic, cationic and non-ionic polymer additives were prepared via reversible addition-fragmentation chain-transfer (RAFT) solution polymerisation, and the impact of these polymer additives on the stability, rheology and printability of aqueous iron oxide nanoparticle (IOP) suspensions was investigated. Zeta potential measurements, particle size characterisation and sedimentation experiments at a range of pH values revealed that the polymer additives significantly altered IOP suspension stability. Specifically, poly(glycerol monomethacrylate) (PGMA), quaternised poly(2-(dimethylamino)ethyl methacrylate) (q-PDMAEMA), poly(2-(dimethylamino)ethyl methacrylate) (PDMAEMA), and polyethyleneimine (PEI) enhanced stability in acidic conditions. At neutral and alkaline pH, the stability was significantly improved with the addition of poly(methacrylic acid) (PMAA) and PGMA. Subsequently, rheological assessments on IOP suspensions with 0.5% w/w of polymer additive demonstrated that PGMA₄₈ reduced the dispersion viscosity at all pHs studied. In contrast, PDMAEMA₄₈ and PEI reduced the viscosity at pH 3 but increased it at pH 7 and 10. Poly(potassium 3-sulfopropyl methacrylate) (PKSPMA₄₉) consistently raised the viscosity at all pH values studied. The practical application of these findings was demonstrated through the direct ink writing (DIW) of polymer additive-containing IOP inks to form 11-layered thin-walled square structures, which showed enhanced shape retention and crack-free drying on aluminium substrates. These findings underscore the potential of precise polymer additives to refine ceramic ink rheology at minimal polymer loadings, paving the way for the development of tailored polymer additives for ceramic ink formulation and 3D printing technologies.

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Introduction

Iron oxide nanoparticles (IOPs), known for their soft magnetic properties and high surface area, are extensively employed in various applications including water treatment,^{1,2} diagnostic imaging,^{3,4} drug delivery,^{4,5} and inductive structures.⁶⁻⁸ However, the machining of IOP-based ceramics into complex structures presents significant challenges due to their inherent hardness and brittleness.⁹⁻¹² Traditional ceramic manufacturing techniques, such as tape casting,¹³ gel casting,^{14,15} slip casting,^{16,17} and injection moulding,^{18,19} facilitate the creation of intricate designs. However, these methods are prone to defects in the demoulding stage, which is critical for producing structures with thin walls, high aspect ratios, or intricate

cornering. As an innovative solution, additive manufacturing (AM) has emerged as a potential alternative. This technology builds near-net-shape objects layer-by-layer directly from three-dimensional (3D) model data, eliminating the need for moulds.^{9,10,20,21} These attributes of AM make it an exceptionally promising approach for overcoming the manufacturing challenges associated with crafting ceramic items.

Extrusion-based direct ink writing (DIW), a sub-branch of AM, extrudes concentrated suspension inks through a printing nozzle to form desired shapes.^{9,22-26} While DIW and other technologies address demoulding issues, they introduce other challenges, such as delamination between printed layers and entrapped air bubbles. These defects can be mitigated through pre-printing preparation such as centrifugation or sonication to remove trapped bubbles, and through optimising processing parameters including print nozzle moving speed, extrusion speed, nozzle size and layer height to prevent delamination between layers. Fundamentally, the success of all these optimisations depends on achieving suitable rheological properties and good ink stability. These inks should have shear

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thinning behaviour to allow them flow during extraction and a large enough storage modulus (G'), yield stress (σ_y) and flow stress (σ_f) to prevent the collapse and distortion of printed structures.^{21,22,24,26-30} Additionally, these inks should be stable and without large ceramic particle aggregates to prevent clogging of the printing nozzle. Organic additives are commonly utilised during formulation to adjust the rheological properties of DIW inks. Depending on their effect on rheology, additives can be divided into two categories: those that thicken the ink,^{6,31-35} and those that thin it.^{7,9,36} Currently, most of the work in the literature selects one kind of polymer additive and demonstrates its effect,^{7,9,33,34} rather than performing a systematic exploration of the effects of a range of additives on ink rheology. One key parameter affecting additive properties is the ionisation of the additive. Water soluble polymers can be non-ionic, cationic, anionic and zwitterionic. In addition, they can have a permanent charge or be pH responsive. These parameters influence the interactions between polymers and ceramic particles, thus the suspension stability and rheology.^{22,36,37} Consequently, there is a pressing need for conducting a systematic exploration of the effects of polymer additives with different charges on the rheology of DIW ceramic inks to establish clearer guidelines for selecting additives to optimise ink rheology.

Recent studies have explored the impact of various additives on the rheology of ceramic slurries and suspensions. Yaghtin *et al.*³⁸ investigated the effects of polyethyleneimine (PEI), 2-phosphonobutane-1,2,4-tricarboxylic acid, and alpha-terpineol on the rheology of highly concentrated aqueous yttria-stabilised zirconia slurries at different pHs. Similarly, Lakhdar and Goodridge *et al.*³⁶ examined the influence of a range of commercial polymer dispersants on the stability and rheology of colloidal boron carbide suspensions. The scope of this research encompassed a diverse array of additives, including cationic branched PEI of varying molecular weights, non-ionic polyoxyalkyleneamine derivatives, acrylic copolymers, anionic poly(acrylic acid ammonium salt), and poly(methacrylic acid sodium salt). Both research groups primarily focused on how these additives affected suspension rheology. However, they did not control additive molecular weight variations, and the effect of polymers with differing degrees of ionisation on rheology was not examined. This limitation arose because the polymer additives were procured from commercial manufacturers, rather than being custom produced to meet specific research requirements. Moreover, most research in this field has been directed towards additives that reduce viscosity and flow stress, thereby enabling higher maximum ceramic loading.^{7,9,36,39,40} However, for additives that increase ink thickness, other viscosifiers or binders are typically added alongside dispersants. This practice complicates ink composition, leading to high cumulative polymer additive loads.^{34,41,42} In some instances, this necessitates further curing processes, posing additional challenges for 3D printer setup and extending printing times.^{6,43-45}

IOPs typically have surfaces containing hydroxyl groups which serve as anchoring points for polymers containing car-

boxyl, amine, hydroxyl, and other functional groups.^{7,9,46-48} Reported (co)polymers containing these groups have been used as additives to adjust the rheology of IOP suspensions.^{7,9,49,50} However these studies have the same limitations as mentioned above, such as focusing on polymer additives which can thin IOP inks, and neglecting the influence of degree of ionisation and molecular weight on rheology. It is therefore necessary to conduct a systematic study on how polymer additives with different charge, degree of ionisation, and molecular weight affect the rheology of IOP suspensions, and subsequently investigate their different performance for DIW.

Herein, a range of well-defined polymer additives with different functionalities and molecular weights were synthesised *via* reversible addition-fragmentation chain-transfer (RAFT) solution polymerisation and investigated as polymer additives for the formulation of aqueous IOP inks (Fig. 1). The stability of IOP dispersions containing these different polymer additives was methodically evaluated, employing Zeta potential, dynamic light scattering (DLS), disc centrifuge photosedimentometry (DCP) and sedimentation measurements. Subsequent investigations focused on the impact of these polymer additives on the rheology of IOP suspensions, utilising oscillatory and steady-state analysis. Finally, the enhancement of ink printability through the addition of these polymer additives was demonstrated by 3D printing IOP inks into thin-walled squares and quantitatively assessed using several printability criteria.⁵¹⁻⁵³

Experimental

Materials

Potassium 3-sulfopropyl methacrylate (KSPMA, 98%), methacrylic acid (MAA), 2-(dimethylamino) ethyl methacrylate (DMAEMA, 98%), quaternised 2-(dimethylamino) ethyl methacrylate solution (q-DMAEMA, 75% in H_2O), and 4,4'-azobis(4-cyanovaleric acid) (ACVA) were purchased from Sigma-Aldrich (UK). Glycerol monomethacrylate (GMA) was generously donated from GEO Specialty Chemicals (UK). 4-Cyano-4-(((dodecylthio)carbonothietyl)thio)pentanoic acid (CDTPA) and 4-(((2-carboxyethyl)thio)carbonothietyl)thio)-4-cyanopentanoic acid (CCC) were purchased from Boron Molecular (Australia). Ethanol (95%) was obtained from Fisher Scientific (UK). Iron (III) oxide nanoparticle powder (20–40 nm average particle size) was purchased from Alfa Aesar (UK). Deionised (DI) water with a resistivity of 18.2 M Ω cm was used in all experiments. All reagents were used as received unless otherwise specified.

Synthesis of polymer additives

Polymer additives were synthesised *via* RAFT solution polymerisation, as described in detail in the SI (Fig. S1–S5). A typical example protocol for the synthesis of PKSPMA₄₈ is as follows. KSPMA (2.0 g, 8.12 mmol), CCC (0.05 g, 0.16 mmol) and ACVA (0.009 g, 0.033 mmol) were dissolved in DI water (8 g) within a 24 mL glass vial. This vial was subsequently



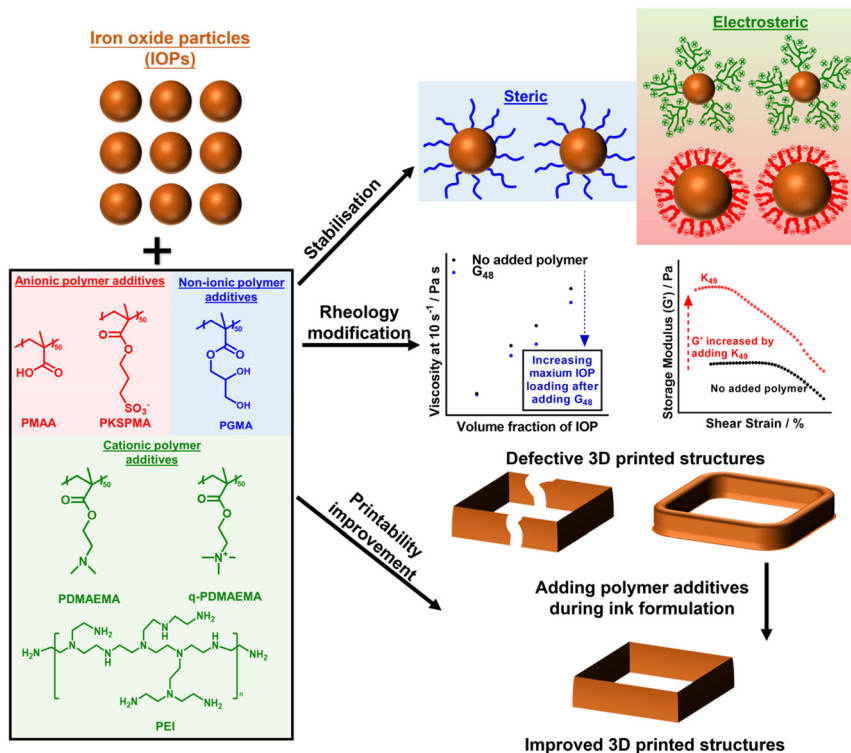


Fig. 1 Polymer additives synthesised via RAFT polymerisation were utilised for stabilising, rheology modification and printability improvement of aqueous IOP suspensions. Without polymer additives, the printed structures exhibited significant defects, attributed to a low storage modulus and insufficient IOP loading capacity. Adding poly(glycerol monomethacrylate) (G_{48}) notably enhanced the maximum IOP loading capacity, mitigating crack formation. Similarly, the addition of poly(potassium 3-sulfopropyl methacrylate) (K_{49}) significantly improved the storage modulus for low IOP loading (50% w/w) inks, effectively preventing structural collapse.

sealed and purged with N_2 for 30 min. The vial was then immersed in a preheated oil bath at 70 °C for 1.5 h. Subsequently, the vial was taken out from the oil bath and immersed in an ice bath to stop the polymerisation.

The obtained polymer solution was purified using dialysis (MWCO = 1000 g mol⁻¹) against DI water for 2 days, and then freeze-dried to dryness. The final degree of polymerisation was determined by ¹H NMR spectroscopy using deuterium oxide (D_2O) as solvent (Fig. S6) and the molar mass distribution was measured using gel permeation chromatography (GPC) (Fig. S11).

Dynamic light scattering (DLS) and aqueous electrophoresis

DLS and aqueous electrophoresis studies were performed using a Malvern Zetasizer Ultra instrument to measure both intensity-average hydrodynamic diameter ($D_{\text{intensity}}$) and Zeta potential. The instrument was equipped with a He-Ne solid-state laser operating at 633 nm, detecting back-scattered light at a scattering angle of 173°. All samples were diluted to 0.1% w/w in presence of 1 mM KCl and sonicated in an ultrasonic bath for 1 h before measurements were taken. Data were averaged over three consecutive runs at 25 °C. Plastic cells (Malvern DTS0012) were used for measuring $D_{\text{intensity}}$ and capillary cells (Malvern DTS1070) were used for measuring Zeta potential.

Disc centrifuge photosedimentometry (DCP)

DCP analyses were conducted using a Centrifugal Photo Sedimentation (CPS) Disc Centrifuge Model 24 000 to measure weight-average diameter (D_{weight}). A density gradient that ranged from 24 to 8% w/w sucrose solution in DI water was constructed and allowed to stabilise for approximately 30 min. A 483 nm diameter poly(vinyl chloride) calibration standard was injected prior to the analysis of each sample. All samples were diluted to 0.1% w/w in presence of 1 mM KCl and sonicated in an ultrasonic bath for 1 h before measurements were taken. The run time was approximately 50 s with the centrifuge set at 11 000 rpm.

Sedimentation experiments of IOP dispersions

Sedimentation experiments were performed to determine the sedimentation behaviour and stability of IOP dispersions with different polymer additives (1% w/w of the mass of IOPs) using the following protocol. 0.01 g of polymer additive was dissolved in 34 g DI water, then 1 g of IOPs were added to form 2.86% w/w IOP dispersions. The dispersion pH was adjusted to the required value using 0.025 and 0.25 M HCl and KOH solutions. The dispersion was sonicated for 1 h, vortex mixed for 1 min and transferred to a volumetric cylinder which was then sealed using parafilm to prevent evaporation. The sample was then left undisturbed for 120 h. After 120 h, the volume of



sedimented IOPs was recorded as the sedimentation volume. After sedimentation, 2 mL of the supernatant from selected samples was collected for UV-vis spectrophotometry analysis, as detailed in the SI.

Preparation of iron oxide suspensions

The preparation of a suspension with 50% w/w IOP loading and 0.5% w/w PDMAEMA₅₀ loading based on IOP concentration at pH 10 is as follows. 2 g of IOPs were transferred into a 10 mL jar. 0.01 g PDMAEMA₅₀ was dissolved in 2 g DI water and the pH adjusted to 10 by adding 0.25 and 0.025 M KOH solution. This solution was then injected into the jar containing the IOPs. The jar was mixed using a speed mixer (Synergy Devices Ltd, Bucks, UK) at 480 rpm for 1 min, 1500 rpm for 1 min, 1200 rpm for 2 min, 1800 rpm for 2 min, 2000 rpm for 1 min and 400 rpm for 1 min to form a homogenous IOP suspension. Other reported IOP suspensions were prepared through the same procedure by changing the pH, type & loading of polymer additive, and the loading of IOPs.

Rheology of iron oxide suspensions

Rheological measurements were performed using a HAAKE MARS iQ Rheometer, equipped with a flat 35 mm diameter titanium plate and a solvent trap to minimise solvent evaporation. In dynamic oscillation testing, the oscillation frequency was set to 1.592 Hz, and the strain was changed from 0.035% or 0.1% to 300%. During steady-state viscosity measurements, the shear rate was changed from 0.1 s⁻¹ to 100 s⁻¹. The temperature was set to 25 °C for all rheology measurements.

Direct ink writing

11-Layer thin-walled square structures were printed onto aluminium substrates by a robot printer (I&J7300R-LF Robots, I&J Fisnar Inc. Wayne, NJ, USA). Inks were loaded into 5 mL dispensing syringes (FIS8001002, FISNAR) with dispensing pistons for syringe barrels (FIS8001007, FISNAR). Loaded syringes were centrifuged at 800 rpm for 2 min before printing.

Straight, flexible dispensing tips (FIS5601087, FISNAR) with 0.84 mm diameter and 12.7 mm length were used as the print nozzle head. The nozzle head speed was fixed at 8 mm s⁻¹. After printing, green bodies were dried in air for at least 12 h before being removed from the substrate.

Results and discussion

Polymer additives studied

A series of well-defined homopolymers with different functionalities and molecular weights were prepared by RAFT solution polymerisation, as described in the SI. Specifically, poly(methacrylic acid) (PMAA or MA_x), poly(potassium 3-sulfopropyl methacrylate) (PKSPMA or K_x), poly(2-(dimethylamino)ethyl methacrylate) (PDMAEMA or D_x), quaternised poly(2-(dimethylamino)ethyl methacrylate) (q-PDMAEMA or q-D_x), and poly(glycerol monomethacrylate) (PGMA or G_x) were prepared and characterised by ¹H NMR and GPC (Fig. S1–S13), as summarised in Table 1. As expected, these polymers had degrees of polymerisation (DPs) close to the targeted value and relatively narrow molar mass distributions. These polymers and a commercial branched PEI were investigated as additives for the formulation of aqueous IOP inks. Initial studies were conducted on low concentration IOP dispersions before investigating how these observations translated to high concentration dispersions suitable for DIW.

Zeta potential, DLS and DCP studies

The stability and size distribution of low concentration (0.1% w/w) IOP dispersions, both with (1% w/w, based on IOP concentration) and without polymer additives, were studied at pH 3, 7 and 10. The Zeta potential values, which are indicative of surface charge and colloidal stability,^{54–56} along with particle aggregation measurements obtained by DLS and DCP are given in Fig. 2.

Table 1 Summary of polymer additives

Acronym	[Monomer] : [CTA] ^a	Monomer conversion ^b	Polymer composition ^b	<i>M_n</i> /g mol ⁻¹	<i>M_w</i> / <i>M_n</i>
K ₄₉	50	97%	PKSPMA ₄₉	9900 ^c	1.09 ^c
K ₂₈₄	300	94%	PKSPMA ₂₈₄	36 380 ^c	1.21 ^c
MA ₄₈	55	87%	PMAA ₄₈	4800 ^d	1.17 ^d
MA ₂₅₀	330	76%	PMAA ₂₅₀	22 100 ^d	1.11 ^d
q-D ₄₈	50	95%	q-PDMAEMA ₄₈	10 000 ^e	
q-D ₂₈₈	300	96%	q-PDMAEMA ₂₈₈	59 600 ^e	
D ₅₀	55	92%	PDMAEMA ₅₀	7900 ^e	
D ₂₈₈	300	96%	PDMAEMA ₂₈₈	45 200 ^e	
G ₄₈	50	95%	PGMA ₄₈	7700 ^f	1.30 ^f
G ₂₅₇	330	78%	PGMA ₂₅₇	41 100 ^f	1.23 ^f
PEI				60 000 ^g	12.5 ^g

^a The ratio between the molar concentration of monomer and chain-transfer agent (CTA) during the RAFT polymerisation. ^b Determined by ¹H NMR analysis. ^c Measured by aqueous GPC using polyethylene oxide/glycol (PEO/PEG) standards. ^d Measured by THF GPC using polystyrene standards (PS) standards, PMAA samples were methylated using trimethylsilyl diazomethane to afford poly(methyl methacrylate) before measurement. ^e Calculated using equation $M_{n,NMR} = (\text{repeat unit molar mass} \times \text{degree of polymerisation determined by NMR analysis}) + \text{CTA molar mass}$. Further details are given in the SI. ^f Measured by DMF GPC using poly(methyl methacrylate) (PMMA) standards. ^g Provided by the manufacturer.



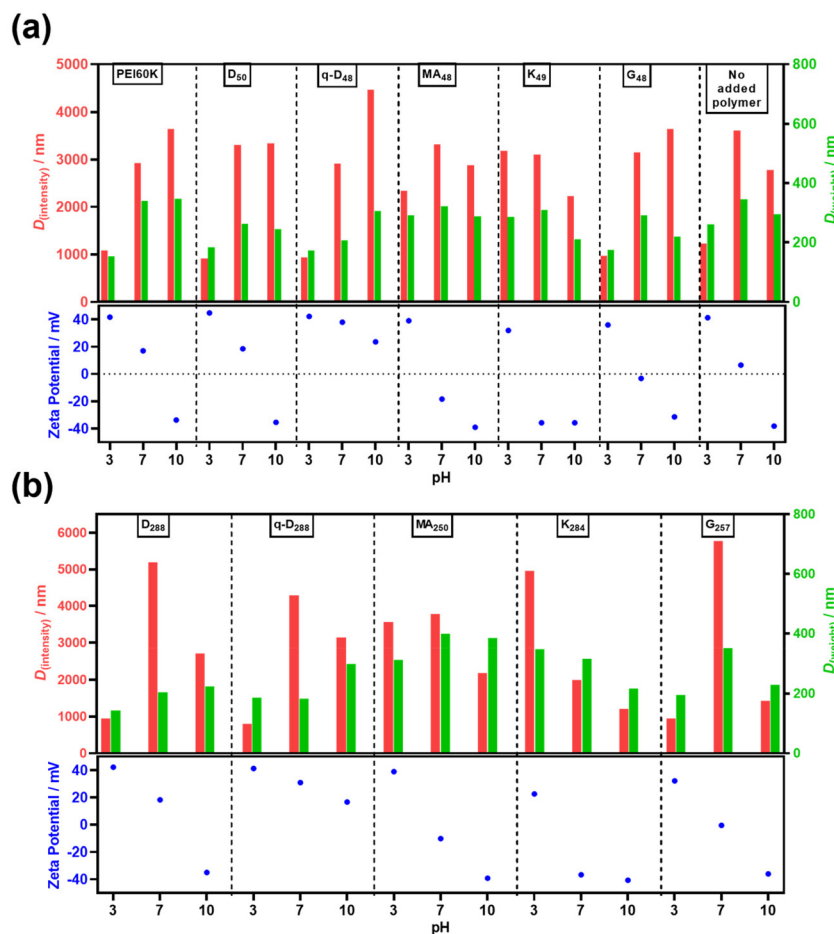


Fig. 2 Zeta potential (blue dots), intensity-average diameter $D_{\text{intensity}}$ (red columns) and weight-average diameter D_{weight} (green columns) of IOP dispersions with different added polymers at pH = 3, 7 and 10. The IOP concentration was 0.1% w/w based on the mass of the dispersion, and the added polymer concentration was 1% w/w based on the mass of the IOPs.

IOPs typically have surfaces containing hydroxyl groups ($-\text{OH}$)^{4,47,48,57} which become protonated ($-\text{OH}_2^+$) at pH 3 and deprotonated ($-\text{O}^-$) at pH 10. This produces a pronounced pH-dependent variation in the Zeta potential of the pristine IOPs. At pH 3, the pristine IOPs demonstrated a highly positive Zeta potential of 41 mV as a result of hydroxyl group protonation. The Zeta potential exhibited a modestly positive value of 7 mV as the pH increased to pH 7 because of the reduced degree of protonation. When the pH was increased from 7 to 10, the hydroxyl groups became deprotonated, which yielded a negative Zeta potential value (-38 mV). The primary particle size of these pristine IOPs was 20–40 nm, as provided by the supplier and confirmed by electron microscopy (Fig. S14). The primary IOPs aggregated to form aggregates with various sizes at different pH values. As illustrated in Fig. 2a, $D_{\text{intensity}}$ (measured by DLS) of IOPs in the absence of polymer indicated the presence of relatively small aggregates at pH 3 (1230 nm). This can be attributed to the relatively high Zeta potential (41 mV) providing strong electrostatic repulsion. $D_{\text{intensity}}$ significantly increased at pH 7 to 3600 nm due to

increased aggregation as a result of the low Zeta potential value (7 mV) providing weak electrostatic repulsion. As the pH increased from 7 to 10, the large negative Zeta potential (-38 mV) resulted in a decrease in aggregate size ($D_{\text{intensity}} = 2800$ nm). D_{weight} (measured by DCP, Fig. S15) showed a similar trend to $D_{\text{intensity}}$, increasing from 140 nm at pH 3 to 250 nm at pH 7, and decreasing to 200 nm at pH 10 (Fig. 2a). It is noteworthy that, in all cases, $D_{\text{intensity}}$ was much larger than D_{weight} . This difference is due to several reasons. First, DCP measures weight-average aggregate size, whereas DLS reports intensity-average aggregate size. Thus, dispersions of IOP aggregates with large size polydispersity are substantially oversized by DLS, especially with the backscattering detector used herein.⁵⁸ Second, DLS measures the dispersion without performing any separation, and light scattering is dominated by the presence of any large aggregates. In contrast, DCP separates out the particle size distribution centrifugally during measurement and is therefore less biased towards the presence of the larger species present. Nevertheless, both of these techniques indicated the presence of IOP aggregates, rather



than a uniform dispersion of primary particles (Fig. 2, and Fig. S15) and confirmed the effect of pH on the relative size of these aggregates.

For PEI-containing IOPs dispersions, the Zeta potential was 42, 17 and -34 mV at pH 3, 7 and 10, respectively. Meanwhile, the PEI-containing IOP dispersions demonstrated decreased amounts of particle aggregation at pH 3 and 7 in comparison to pristine IOPs, and increased particle aggregation at pH 10 (Fig. 2a). PEI is protonated at pH 3 and 7 and thus this cationic polymer provides electrosteric repulsion between particles, reducing aggregation. At pH 10, the residual positive charge of PEI partially shields the negative charge of the IOPs, leading to a decrease in electrostatic repulsion and resulting in more aggregation. The Zeta potential and aggregation behaviour of the D_{50} -containing IOP dispersions were similar to PEI-containing dispersions (Fig. 2a). This is because PDMAEMA₅₀ is also a pH responsive cationic polymer with a pK_a of approximately 7.⁵⁹ q-D₄₈-containing IOP dispersions exhibited positive Zeta potentials at pH 3, 7 and 10 (42, 38 and 24 mV, respectively) due to the permanent cationic charge of q-D₄₈ imparting this behaviour when adsorbed to the surface of IOPs. In this case, D_{weight} increased from 172 nm at pH 3 to 207 nm at pH 7 and 305 nm at pH 10. Similarly, $D_{\text{intensity}}$ increased with increasing pH from 940 nm at pH 3 to 4462 nm at pH 10 (Fig. 2a). The reason for the increase in aggregate size for q-D₄₈-containing IOP dispersions is not fully apparent. However, it can be hypothesised to be caused by the underlying IOP charge resulting in increased aggregation of the IOPs before the q-D₄₈ was able to provide sufficient stabilisation.

For MA₄₈-containing IOP dispersions, the Zeta potential was observed to be 39 mV at pH 3, which was slightly lower than the Zeta potential of pristine IOP dispersions. This is because the carboxylic acid groups on PMAA may have shielded the $-\text{OH}^+$ groups on the surface of the IOPs. At pH 7 and 10, the Zeta potential of MA₄₈-containing IOP dispersions was -18 mV and -39 mV. This reversal in Zeta potential suggested the carboxylic acid groups on MA₄₈ were deproto- nated and adsorbed onto the IOPs, causing the particles to become anionic. For K₄₉-containing IOP dispersions, the Zeta potential was 32, -36 and -36 mV at pH 3, 7 and 10, respectively. The Zeta potential was lower at pH 3 and 7 than the pristine IOP dispersions because of the negatively charged PKSPMA adsorbed to the surface of the IOPs. The values of D_{weight} and $D_{\text{intensity}}$ suggested an increase in stability of K₄₉ containing IOPs with increasing pH due to a decrease in measured aggregate size. However, during these experiments, K₄₉ containing samples were unstable at all pHs, aggregating and sedimenting relatively rapidly. This was confirmed in subsequent sedimentation experiments, discussed below.

PGMA was selected as a non-ionic polymer to investigate whether steric stabilisation alone (rather than electrosteric stabilisation) would provide benefits to IOP ink formulation. The Zeta potential of the G₄₈-containing IOP dispersions did not change significantly compared to the pristine IOP dispersion due to the non-ionic nature of PGMA. Notably, D_{weight} and $D_{\text{intensity}}$ decreased significantly (Fig. 2a) and was attribu-

ted to the stabilisation imparted by adsorbed G₄₈. In addition to the polymers described above, cationic, anionic and non- ionic polymers with larger molar masses were prepared to assess the effect of molecular weight on the rheological properties and printability of polymer containing IOP dispersions. The Zeta potential, D_{weight} and $D_{\text{intensity}}$ of IOP dispersions with 1% w/w added polymer, based on IOP concentration, with larger M_n are shown in Fig. 2b. Interestingly, increased molecular weight did not significantly affect the measured Zeta potential, D_{weight} and $D_{\text{intensity}}$ values when comparing between IOP dispersions containing the same type of polymer with smaller M_n (Fig. 2a). However, it was expected that the molecular weight would have an effect on the rheological properties of high concentration IOP suspensions, discussed below.

In summary, the Zeta potential and aggregation behaviour of IOP dispersions were markedly influenced by the pH and the presence of differently charged polymer additives. Pristine IOPs showed maximum stability at pH 3 due to strong electrostatic repulsion. Cationic polymers (PEI and PDMAEMA) enhanced stability at low pH through electrosteric repulsion. However, at alkaline pH, these cationic polymers were less effective at stabilising IOPs against aggregation. Carboxylic acid functional anionic polymers (PMAA) provided effective stabilisation of IOPs at neutral and alkaline pH whereas sulfonate functional PKSPMA seemed to act as a poor stabiliser. The non-ionic polymer PGMA demonstrated a stabilising effect at all pH values but did not significantly change the Zeta potential when compared to pristine IOPs.

Sedimentation experiments

Sedimentation experiments were conducted to obtain insights into the effects of different polymers on the stability of moderately concentrated IOP dispersions (2.86% w/w). The effect of polymer additive on the settling behaviour of IOPs was demonstrated by the volume of sedimented IOPs, known as the sedimentation volume (Fig. 3). As illustrated in Fig. 3d, a small sedimentation volume indicated good stability and less aggregation of IOPs. On the other hand, a large sedimentation volume indicated low stability and large amounts of IOP aggregation.³⁶

At pH 3, IOP dispersions with G₄₈, D₅₀, q-D₄₈, PEI and no polymer resulted in a low sedimentation volume (<5 mL, Fig. 3). The low sedimentation volume resulted from the relatively high colloidal stability and small degree of IOP aggregation, as indicated by DLS and DCP studies (Fig. 2). IOP dispersions containing MA₄₈ and K₄₈ formed sediments with large sedimentation volumes of approximately 11 mL due to relatively low stability and formation of large IOP aggregates. The observed sedimentation behaviour of the MA₄₈-containing IOP dispersion at pH 3 presented an intriguing anomaly in that both D_{weight} and $D_{\text{intensity}}$ measurements were unexpectedly small. This discrepancy was because of a complex sedimentation dynamic that was not fully captured by the DLS and DCP size measurements. At pH 3, aggregation of IOPs rapidly occurred (within approximately 10 s) in the MA₄₈-containing



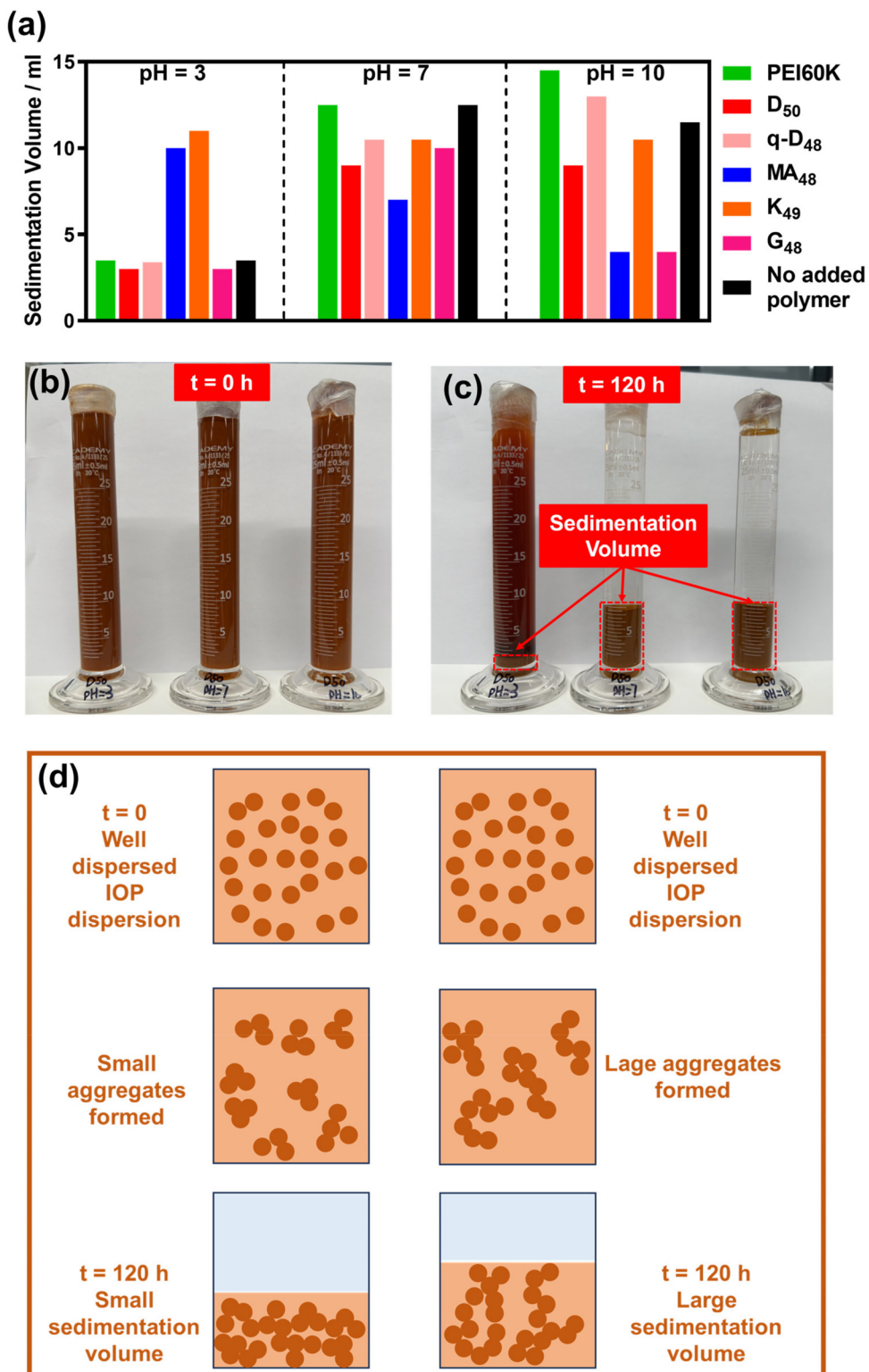


Fig. 3 Sedimentation tests for 2.85% w/w IOP dispersions with and without 1% w/w added polymer (based on IOP concentration), at pH 3, 7, and 10. Dispersions were sonicated for 1 h, vortex mixed for 1 min, and then left to stand undisturbed for 120 h, after which the sedimentation volume was measured, as summarised in (a). Photographs of 2.85% w/w IOP dispersions with 1% w/w D₅₀ at pH 3, 7 and 10, at (b) $t = 0$ and (c) $t = 120\text{ h}$. (d) Schematic showing the formation of small (left) and large (right) sedimentation volumes.



IOP dispersion forming sizable clusters. These agglomerates exceeded the set test range of DCP (50–1000 nm) and settled too swiftly to be accurately sized by DLS, resulting in the recorded D_{weight} and $D_{\text{intensity}}$ values being much lower than anticipated.

At pH 7, the IOP dispersions containing D₅₀, q-D₄₈, K₄₈, PEI and G₄₈, as well as the pristine IOP dispersion, exhibited sedimentation volumes of 10 mL or more (Fig. 3a), signifying reduced colloidal stability and notable particle aggregation. This trend was corroborated by the increased value of D_{weight} and $D_{\text{intensity}}$. In contrast, the MA₄₈-containing dispersion demonstrated moderately better stability, evidenced by a sedimentation volume of only 7 mL.

At pH 10, the MA₄₈ and G₄₈-containing IOP dispersions had a reduced sedimentation volume of 4 mL (Fig. 3a), indicative of improved colloidal stability and minimal particle aggregation. The enhancement in stability for these two dispersions was primarily ascribed to the amplified electrostatic and steric repulsion. Conversely, the pristine IOP dispersion and the IOP dispersions containing the other polymers studied showed considerable sedimentation volumes (≥ 9 mL), pointing to a lower stability as corroborated by large D_{weight} and $D_{\text{intensity}}$ values in Fig. 2. An exception was observed for the K₄₈-containing IOP dispersion, which, despite its large sedimentation volume, recorded unexpectedly low D_{weight} and $D_{\text{intensity}}$ values at pH 10 compared to those at pH 3 and 7. The reason for this anomaly is same as that previously discussed for the MA₄₈-containing IOP dispersion at pH 3, where large aggregates were not effectively recorded by DLS and DCP.

In addition to the considerations above, swelling of the added polymer may also have affected the final sedimentation volume. In theory, polymer swelling can reduce interactions between colloidal particles after sedimentation, resulting in open sediment structures with larger volumes. However, in this research, the sedimentation volume was not primarily controlled by swelling behaviour. For instance, as the pH increased, MA₄₈ was expected to swell, which would typically lead to a gradual increase in sedimentation volume from pH 3 to 7 and 10. Contrary to this expectation, the sedimentation volume of the MA₄₈-containing IOP dispersion decreased with increasing pH (Fig. 3a), indicating that the sedimentation volume was mainly dominated by aggregate size, as discussed above, rather than by polymer swelling. This conclusion also extends to the pH-responsive D₅₀ and PEI-containing IOP dispersions.

Overall, the findings from the sedimentation experiments were found to be consistent with the Zeta potential and size distribution measurements described above. The two exceptions were MA₄₈ and K₄₈-containing dispersions at pH 3 and 10, respectively, and were due to the rapid aggregation and the formation of sizeable clusters not detected by the particle sizing methodology used herein. This highlights the importance of comprehensive sedimentation experiments in assessing the true effects of polymer additives and pH on IOP stability, offering insights beyond those provided by Zeta potential and size distribution measurements alone.^{36,38} The following

rheology investigations provide even further insights into suspension behaviour, especially when considering highly concentrated suspensions.

Rheology of different polymer-containing IOP suspensions

Initially, different polymer additives (Table 1) were each mixed with 50% w/w IOP aqueous suspensions at 0.5% w/w polymer based on IOP concentration using a high-speed mixer. The pH was varied (pH 3, 7 and 10) and the resulting viscosity compared to pristine IOP dispersions.

The pristine IOP suspensions demonstrated shear thinning behaviour. Their viscosity decreased from ~ 40 Pa s at 1 s^{-1} to ~ 1 Pa s at 100 s^{-1} (Fig. 4f). There were negligible differences in the measured viscosity of the pristine IOP suspensions at the three different pH values studied. This is perhaps surprising based on the particle size, Zeta potential and sedimentation investigations above for less concentrated IOP dispersions. In addition, previous reports typically assume the stability of low-concentration ceramic dispersions to be related to the viscosity of high-concentration ceramic suspensions, where poor dispersion stability leads to high viscosity, and good stability results in low viscosity. The pristine IOP dispersions demonstrated different stability and aggregate sizes at pH 3, 7, 10, but the viscosity of the pristine IOP suspensions was not affected by the pH. One plausible explanation is that the hydroxyl groups on pristine IOPs are non-ionic under the conditions studied, and thus the thickness of electrical double-layer was relatively thin. Under these circumstances, the pristine IOP suspensions could be assumed to be a hard-sphere system, where interparticle interactions are not experienced until they closely approach each other, meaning that the viscosity is dominated primarily by the solid loading, primary particle/aggregate size and shape.⁵⁵

The addition of D₅₀ resulted in pH-sensitive viscosity changes in the IOP suspensions. At pH 3, protonated D₅₀ provided electrostatic repulsion between IOPs, significantly decreasing the viscosity (Fig. 4a) to 25 Pa s at 1 s^{-1} , approximately half the viscosity of the pristine IOP suspension (Fig. 4f). At pH 7, close to the pK_a of PDMAEMA,⁵⁹ D₅₀ chains collapsed due to deprotonation, increasing their hydrophobic character. The hydrophobic association between the D₅₀ polymer chains induced aggregation, which led to a significant increase in viscosity (390 Pa s at 1 s^{-1}). At pH 10, the viscosity increased to 1270 Pa s at 1 s^{-1} because of further deprotonation of D₅₀ further increasing the amount of aggregation.

The q-D₄₈-containing IOP suspensions had viscosities of 30 , 40 and 60 Pa s at 1 s^{-1} , at pH 3, 7 and 10, respectively (Table S1). There was a slight decrease in viscosity at pH 3 and 7 when compared to pristine IOP suspensions due to strong electrostatic repulsion caused by the surface-adsorbed q-D₄₈. The slight increase in viscosity at pH 10 for q-D₄₈-containing IOP suspensions was potentially caused by the decrease in particle stability as discussed above (Fig. 2 and 3a). Similarly, the addition of G₄₈ (Fig. 4e) slightly decreased the viscosity when compared with pristine IOPs suspensions (Fig. 4f), with the non-ionic G₄₈ reducing viscosity through steric repulsion.^{9,60–62}



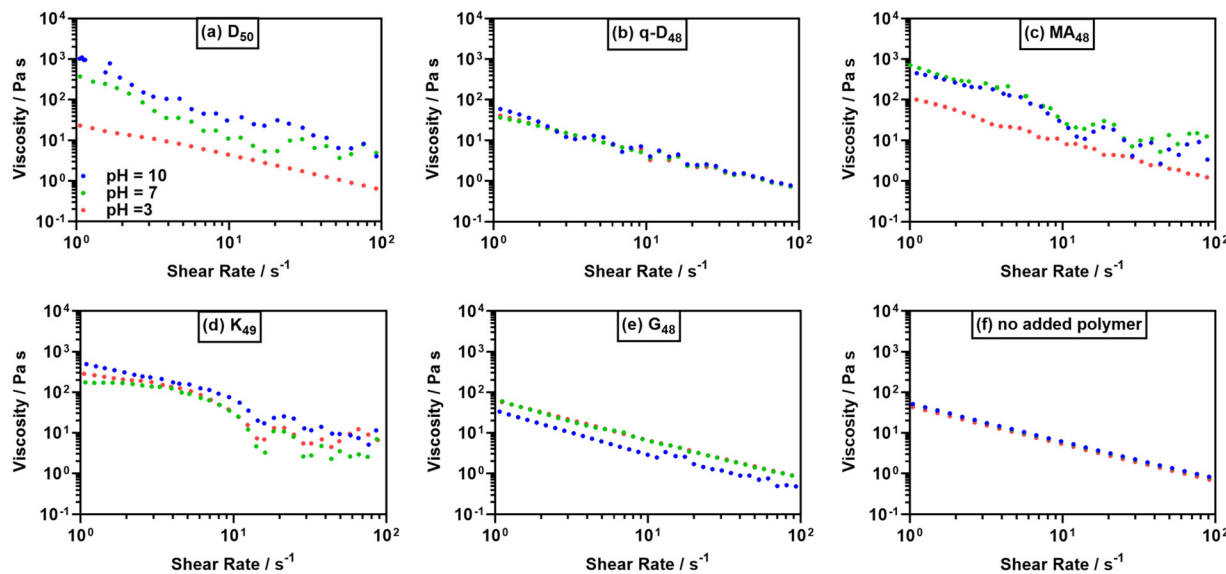


Fig. 4 Viscosity vs. shear rate for 50% w/w IOP suspensions with 0.5% w/w (a) D₅₀, (b) q-D₄₈, (c) MA₄₈, (d) K₄₉, (e) G₄₈, and (f) without added polymer at pH = 3 (red dots), pH = 7 (green dots) and pH = 10 (blue dots). The 0.5% w/w polymer additive concentration is based on IOP concentration.

MA₄₈-containing IOP suspensions displayed a distinct pH-dependant viscosity profile (Fig. 4c). At pH 3, the viscosity was 110 Pa s at a shear rate of 1 s⁻¹ and this value significantly increased to 710 and 500 Pa s at pH 7 and 10, respectively. The pK_a of PMAA is approximately 5.5, thus MA₄₈ was deprotonated (and anionic) at pH 7 and 10. At pH 3 MA₄₈ was protonated, and the chains were therefore collapsed resulting in a decrease in viscosity.⁶³ Similarly, anionic K₄₈-containing IOP suspensions (Fig. 4d) had high viscosities at shear rates of 1 s⁻¹ (300, 200, 530 Pa s at pH 3, 7, 10, respectively), which were 4-to-9 times those of the pristine IOP suspensions. These observations demonstrate that IOP suspensions could be significantly thickened using anionic polymer additives, as expected. To confirm this thickening behaviour was caused by adsorbed polymer, UV-Vis spectrophotometry studies were conducted (Fig. S19 and Table S2). For example, nearly all of the added K₄₈ was adsorbed onto the IOPs (see Fig. S19), leaving little free K₄₈ in solution.

One interesting observation is that the use of cationic additives tended to reduce the viscosity of IOP suspensions whereas anionic additives caused thickening. This is commonly observed for ionic rheology modifiers and is often attributed to differences in their electrostatic interactions with the surrounding medium, where anionic polymers adopt extended conformations due to intramolecular charge repulsion and enhanced hydration, thereby increasing viscosity, while cationic polymers tend to collapse or aggregate through charge neutralisation and bridging interactions, resulting in reduced thickening effects.⁶⁴⁻⁶⁶ The relationship between viscosity and shear rate within the specific range of 1 to 100 s⁻¹ was fitted using power law equation $\eta = a \times \gamma^{b-1}$ (see Table S1),^{67,68} where η is the viscosity, a is the flow consistency

index, γ is the shear rate, and b is the flow behaviour index. Low flow consistency index indicates a low resting viscosity of the IOP suspensions, while a low flow behaviour index indicates stronger shear thinning behaviour. Notably, the use a Herschel-Bulkley model²⁴ to fit this shear stress data would have been preferable. However, noise and the limited shear rate range in our dataset made it unfeasible to use this model. The pristine IOP suspensions exhibited significant shear thinning behaviour ($b \leq 0.1$) at pH 3, 7 and 10. This significant shear thinning was also observed for the G_{48/257}, PEI, q-D₂₈₈, and MA_{48/250}-containing IOP suspensions at pH 3, 7 and 10, as well as K₂₈₄-containing IOP suspensions at pH 3 and 7. In contrast, the D_{50/288} and q-D₄₈-containing IOP suspensions demonstrated high flow behaviour index ($b \geq 0.25$) at pH 3, 7, 10, indicating lower shear thinning behaviours.

Fig. 5 shows the storage modulus (G'), loss modulus (G''), and shear stress as functions of oscillation strain for IOP suspensions containing various polymer additives in amplitude sweep tests. In all plots shown in Fig. 5, three regions are clearly distinguishable. The first region, known as linear viscoelastic region (LVR), is where the storage modulus (G') is larger than the loss modulus (G'') and remains constant. In this region, the shear stress shows a linear response and the storage modulus in LVR (G'_{LVR}) is equal to the elastic modulus of a solid-like material. The second region is the yield region, where G' decreases as the shear strain increases due to irreversible microstructure evolution. The yield point marks the transition from linear viscoelastic behaviour to nonlinear viscoelastic behaviour. However, in this region the elastic behaviour still dominates the viscoelastic performance over the viscous behaviour ($G' > G''$). The final region is the flow region, which starts from the point where ($G' = G''$).^{9,24} In this region the



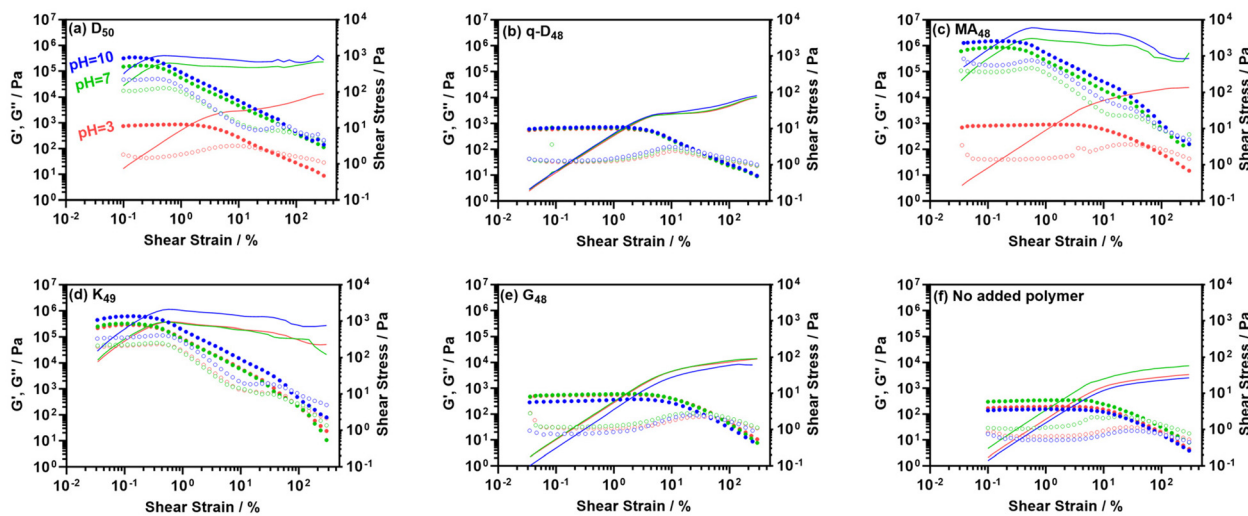


Fig. 5 Storage modulus (G' , solid dots), loss modulus (G'' , hollow dots) and shear stress (solid lines) as a function of shear strain for 50% w/w IOP suspensions with 0.5% w/w (a) D_{50} , (b) $q\text{-}D_{48}$, (c) MA_{48} , (d) K_{49} , (e) G_{48} , and (f) without added polymer at pH = 3 (red), pH = 7 (green) and pH = 10 (blue). The 0.5% w/w polymer additive concentration is based on IOP concentration.

viscous behaviour dominates over the elastic behaviour and the suspension flows.^{22,24,67} The relationship between shear strain and shear stress varied across different suspensions. For soft suspensions ($G'_{LVR} < 10^4$ Pa and shear stress < 200 Pa), shear stress continued to increase with shear strain beyond the yield point, which is typical behaviour for shear thinning suspensions. In contrast, more solid-like samples ($G'_{LVR} > 10^4$ Pa and shear stress > 200 Pa) reached a maximum stress near the yield point, followed by a decrease in stress with increasing strain. This behaviour could be caused by network break during yielding or some issues during measurement such as fracture, slip, and shear banding.^{28,30,52,69}

For pristine IOP suspensions (Fig. 5f), the low G'_{LVR} values of 190, 349, and 150 Pa at pH 3, 7, and 10, respectively, indicated a weak elastic modulus, suggesting that these suspensions are too weak to retain their shape *e.g.*, after printing. Additionally, they had low flow stress values of 30, 50, and 20 Pa at pH 3, 7, and 10. The G'_{LVR} of D_{50} -containing IOP suspensions were 830, 133 200 and 340 000 Pa, at pH 3, 7 and 10, respectively and the flow stress values were 80, 420 and 710 Pa (Fig. 5a and Table S1). At pH 7 and 10, the high value of G'_{LVR} ($>10^5$ Pa) and flow stress (>400 Pa) were attributed to the formation of a bridging network of aggregates through hydrophobic associations.^{59,70,71} This network structure enhanced resistance to deformation and prevented collapse. When the pH decreased to 3, D_{50} became protonated and hydrophilic. Consequently, the hydrophobic associations were no longer present, resulting in very low values of storage modulus and flow stress. The G'_{LVR} of $q\text{-}D_{48}$ -containing IOP suspensions was 610, 660 and 710 Pa at pH 3, 7 and 10, respectively (Fig. 5b and Table S1). The slight increase of G'_{LVR} compared to pristine IOP suspensions was potentially caused by the entanglement of $q\text{-}D_{48}$ chains. This entanglement no longer affected the flow of IOPs at high shear strain, which was confirmed by the low

flow stress of $q\text{-}D_{48}$ -containing IOP suspensions (40–70 Pa) at pH 3, 7 and 10. For MA_{48} -containing IOP suspensions, the values of G'_{LVR} and flow stress were strongly pH dependent (Fig. 5c and Table S1). At pH 7 and 10, G'_{LVR} was 672 300 and 1 233 000 Pa, respectively, and the flow stress was 820 and 900 Pa. Similar to the viscosity behaviour described above, the values of G'_{LVR} (820 Pa) and flow stress (100 Pa) were low at pH 3 as a result of PMAA protonation. K_{49} -containing IOP suspensions demonstrated high G'_{LVR} values of 216 300, 296 900 and 471 200 at pH 3, 7 and 10, respectively (Fig. 5d and Table S1) due to strong polymer hydration and entanglement. These factors also contributed to high flow stresses of 300, 200 and 530 Pa at pH 3, 7 and 10, respectively. The G'_{LVR} of G_{48} -containing suspensions were 540, 580 and 310 Pa at pH 3, 7 and 10, respectively (Fig. 5e and Table S1). The G'_{LVR} was pH independent due to the non-ionic nature of the G_{48} adsorbed onto the IOPs, providing steric repulsion at all pHs. This steric repulsion also decreased the flow stress at pH 10 (Table S1).

A series of homopolymers with higher M_n (DPs > 250) were also synthesised (Table 1) and their effect on IOP suspension rheology assessed (Fig. S16, S17 and Table S1). With increasing M_n , the G'_{LVR} and flow stress of IOP suspensions containing PKSPMA and PMAA decreased. Conversely, the G'_{LVR} and flow stress of IOP suspensions containing $q\text{-}PDMAEMA$ and PDMAEMA increased with increasing M_n . However, the M_n of PGMA did not significantly affect the rheology of PGMA-containing IOP suspensions.

The PEI-containing ($M_n \sim 60\,000$ g mol $^{-1}$) IOP suspensions demonstrated a rise in viscosity (Fig. S16g), G'_{LVR} and flow stress (Fig. S17g) with increasing pH, as summarised in Table S1. This pH-responsive performance correlated with the multiple pK_a values for PEI, which are approximately 9.0, 8.0 and 6.0 for primary, secondary and tertiary amines,⁷² respectively. At pH 3, all amine groups are fully protonated and the

PEI chains become fully extended, providing strong electrosteric repulsion and halving the viscosity (20 Pa s at 1 s⁻¹) compared to the pristine IOP suspension (40 Pa s at 1 s⁻¹). At pH 7, the viscosity increased significantly, to ten-fold higher than the value at pH 3, and increased further at pH 10. This was due to the formation of hydrophobic associations caused by deprotonation of the amine groups, which has been discussed above for D₅₀-containing IOP suspensions.⁷⁰ The G'_{LVR} and flow stress of PEI-containing IOP suspensions showed the same trend as the viscosity response to changes in pH.

In summary, the addition of polymer additives increased the G'_{LVR} in all cases compared to pristine IOP suspensions. In particular, the formation of hydrated polymer networks, entanglements and hydrophobic association significantly enhanced the G'_{LVR} , flow stress and viscosity of these suspensions. In addition, G₄₈, D₅₀ and q-D₄₈ demonstrated their ability to decrease the flow stress at pH 10, 3 and 7, respectively, due to either steric or electrosteric repulsion. These rheological studies therefore elucidated the effects of various polymers on the potential formulation of 3D-printable IOP inks. It is also worth noting that the rheological behaviour of bare IOP suspensions and IOP suspensions containing q-D₄₈, K₄₉ and G₄₈ was not significantly affected by aggregate size at different pHs, compared to IOP suspensions containing the other additives. This observation can be attributed to two possible mechanisms. First, in these suspensions, the rheological properties are primarily governed by the ionisation state of the polymer additives and the resulting polymer–polymer and polymer–particle interactions. Since q-D₄₈, G₄₈, K₄₉ and the hydroxyl groups on bare IOP surfaces exhibit minimal pH-dependent ionisation changes,^{62,73–79} their interactions remain relatively constant across the pH range. Second, the applied shear stress during rheological measurements may cause the breakup of aggregates, leading to a dynamic equilibrium in aggregate size regardless of pH.⁶¹ However, we lack direct evidence to confirm this hypothesis. Future studies employing techniques such as rheomicroscopy⁶⁹ and Rheo-SANS⁸⁰ could provide insights into aggregate size and shape evolution during rheology measurements.

Application of polymer additives for 3D printing of the IOPs inks

For pristine IOP suspensions, a relatively low IOP loading (50% w/w, based on mass of the suspension) results in G'_{LVR} and flow stress values that are too low to maintain structures post-printing or resist sagging and collapse. The addition of a small amount of polymer additive such as PKSPMA, PMAA, PEI and PDMAEMA at ~0.5% w/w, relative to the mass of IOPs, and adjusting the pH appropriately, can markedly enhance the G'_{LVR} and shear stress, thereby potentially improving shape retainability and increasing resistance to sagging or collapse of the printed filament. In addition, G₄₈ was promising for the preparation of high loading inks as it decreases IOP dispersion viscosity at pH 10. The addition of D₅₀, q-D₄₈ and PEI also decreased viscosity at pH 3.^{40,72}

Unexpectedly, upon increasing the IOP loading from 50 to 70% w/w or higher, the viscosity-reducing effect of adding D₅₀

was not observed (Fig. S18a). This phenomenon was also noted for q-D₄₈ pH 3 (Fig. S18b). In contrast, G₄₈ consistently reduced viscosity across all IOP loadings (Fig. S18a), suggesting that the steric repulsion provided is more effective in decreasing suspension viscosity than the cationic electrosteric repulsion of q-D₄₈ and D₅₀, particularly at high IOP loadings. Additionally, G₄₈ lowered the flow stress of formulations with IOP loadings over 70% w/w (Fig. 6f). This is important as inks with ultra-high flow stress (>1000 Pa) can prevent ink extrusion during DIW. For stable ink extrusion during DIW, the flow stress of the ink should be lower than the maximum shear stress at the wall of the printhead to ensure controllable flow of the ink. Theoretically, the maximum shear stress is determined using $\tau = (\Delta P/2L)r$,^{7,9,23} where τ is the maximum shear stress, ΔP is the pressure applied at the nozzle, L is the length of the nozzle, and r is the radius of the nozzle. For the printer used in this work, the maximum shear stress at the wall of the nozzle was calculated to be ~723 Pa ($\Delta P = 43\,750$ Pa, $r = 0.42$ mm, $L = 12.7$ mm). Practically, inks with flow stresses slightly above the theoretical value can be printable due to the non-Newtonian nature of inks and dynamic printing conditions. The usable flow stress was therefore limited to approximately 1000 Pa for the printer used in this study. Consequently, without added polymer, the highest feasible IOP loading for 3D printing was 65% w/w and introducing G₄₈ enabled an increase in printable IOP loading to at least 70% w/w.

Thin-walled squares (dimensions: 20 × 8 × 0.84 mm, width × height × wall thickness) were printed onto aluminium substrates using IOP inks formulated with various IOP concentrations, polymer additives and pH (summarised in Table 2). As shown in Fig. 6a, the squares printed using 50% w/w IOP ink without added polymer at pH 7 collapsed after printing due to its low G'_{LVR} (340 Pa) and low flow stress (50 Pa). The squares printed using 50% w/w IOP suspension containing 0.5% w/w K₄₉ at pH 7, demonstrated good shape retainability (Fig. 6b) due to its increased G'_{LVR} (~600 kPa) and shear stress (340 Pa). 50% w/w IOP suspensions containing 0.5% w/w PEI (Fig. S20b) or MA₄₈ (Fig. S20c) at pH 10 also had good shape retainability thanks to the high G'_{LVR} and shear stress, respectively. However, collapse of the PEI-containing IOP ink at pH 3 was observed (Fig. S20a) due to its low G'_{LVR} and shear stress. At the same time, due to the hydrophilic nature of K₄₉, the drying process for objects formulated with this polymer was more uniform and slower,^{82,83} which resulted in crack-free dried green bodies (Fig. S21).

As summarised in Fig. 7a and Table 2, K₄₉, D₅₀, PEI and MA₄₈ could be used at a suitable pH to improve the ink strength of IOP inks by improving the G'_{LVR} and flow stress. K₄₉ was the stand-out from these polymer additives because its thickening effect was not significantly affected by pH, which is promising as these inks could therefore be used under various pH conditions. However, K₄₉ was not suitable for high IOP loading inks (>50% w/w) because adding K₄₉ strongly enhanced the ink strength, making the ink challenging to be mixed homogenously and smoothly extruded from the printing nozzle.



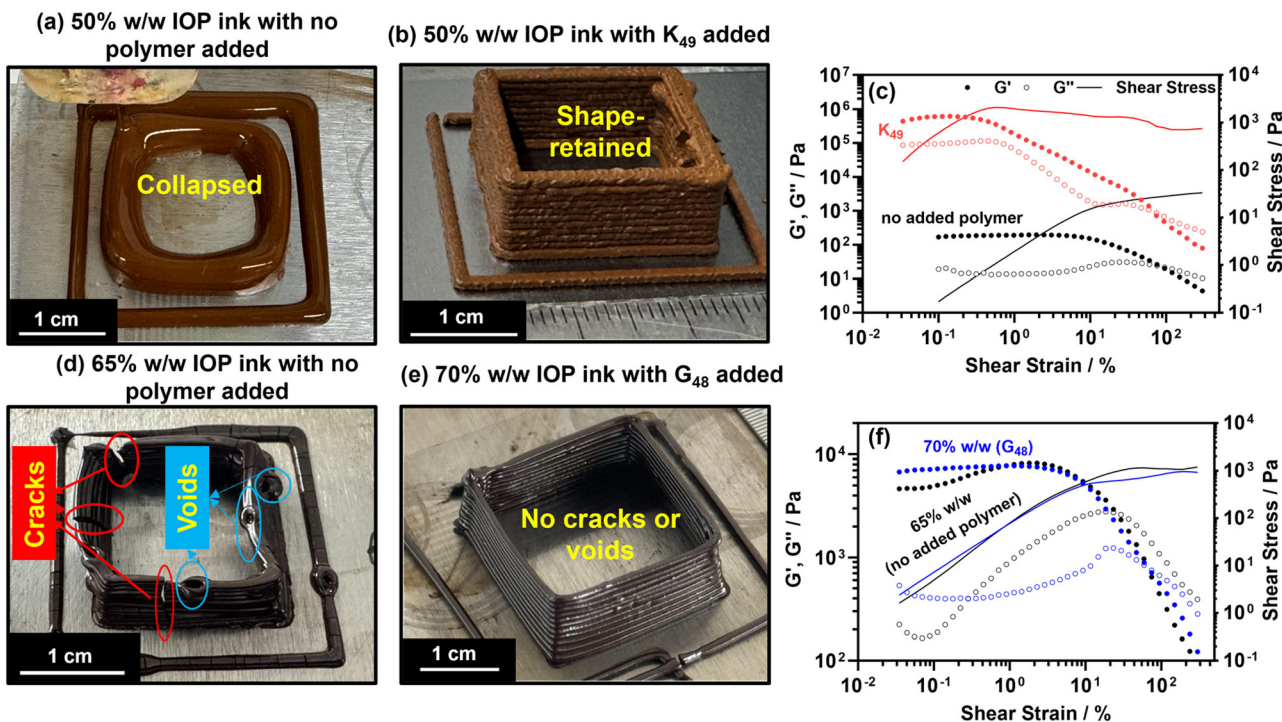


Fig. 6 Linking rheology with direct ink writing: photographs of (a and b) just-printed 11-layer squares using 50% w/w IOP ink with (a) no polymer added at pH 7 and (b) 0.5% w/w K_{49} added at pH 7; (d and e) printed and dried 11-layer squares using (d) 65% w/w IOP ink with no polymer added at pH 10 and (e) 70% w/w IOP ink with 0.5% w/w G_{48} added at pH 10; (c and f) Plot of G' (solid dots), G'' (hollow dots) and shear stress (solid lines) as a function of shear strain for (c) 50% w/w IOP suspensions with 0.5% w/w K_{49} added (red) and 50% w/w IOP suspensions with no polymer added to demonstrate how K_{49} thickened the ink at pH 10; (f) 70% w/w IOP suspensions with 0.5% w/w G_{48} added (blue) and 65% w/w IOP suspensions with no polymer added to show how G_{48} thinned the ink at pH 10.

Table 2 Summary of printability and drying conditions of inks in this work

Ink formulation	Printability evaluation	Drying conditions
50% w/w pristine IOP ink at pH 7	Collapsed, poor shape retention	Cracking observed
50% w/w + 0.5% K_{49} IOP ink at pH 7	Good shape retention	Crack free
65% w/w pristine IOP ink at pH 10	Collapsed, poor shape retention	Cracking observed
70% w/w IOP + 0.5% G_{48} ink at pH 10	Good shape retention	Crack free
50% w/w IOP + 0.5% PEI ink at pH 3	Collapsed, poor shape retention	Cracking observed
50% w/w IOP + 0.5% PEI ink at pH 10	Good shape retention	Crack free
50% w/w IOP + 0.5% MA ₄₈ ink at pH 10	Good shape retention	Crack free
70% w/w pristine IOP ink at pH 10	Blocked the nozzle, unprintable	

Comparing Fig. 7a and b, while some of the additives enhanced ink strength and maximum printing height, they also decreased the Flow Transition Index (FTI)⁸⁶ to <1 . This indicated that the inks became more brittle with an abrupt yield to flow transition. This corresponds to the stress overshoot observed for these formulations. Interestingly, the PEI 60K at pH 10 formulation was an exception as it enhanced the ink strength without making it brittle.

Printed thin-walled squares using 65% w/w pristine IOPs formed voids during printing due to the non-continuous flow of the ink⁵² and collapsed due to low ink stiffness, with $G'_{LVR} < 10\,000$ Pa (Fig. 6d and Fig. 7a). In contrast, adding G_{48} allowed IOP suspensions of 70% w/w to be printed by increasing

stiffness and decreasing the flow stress at the same time. Although the G'_{LVR} of G_{48} containing 70% w/w IOP ink was lower than 10 000 Pa, which is the lowest limitation in printability map (Fig. 7a), this ink still showed good printability and shape retention. The drying of the ink after extrusion from the nozzle swiftly enhanced the storage modulus, which improved the shape retention of the printed green body. In the meantime, the increased solid loading reduced shrinkage of the green body after drying (Fig. 6e). This effect is highly desired for fabricating dense ceramic components,^{9,37,84,85} where investigations into the drying behaviour of green bodies printed with different inks and optimised sintering profiles would be required to tailor these inks for different applications.



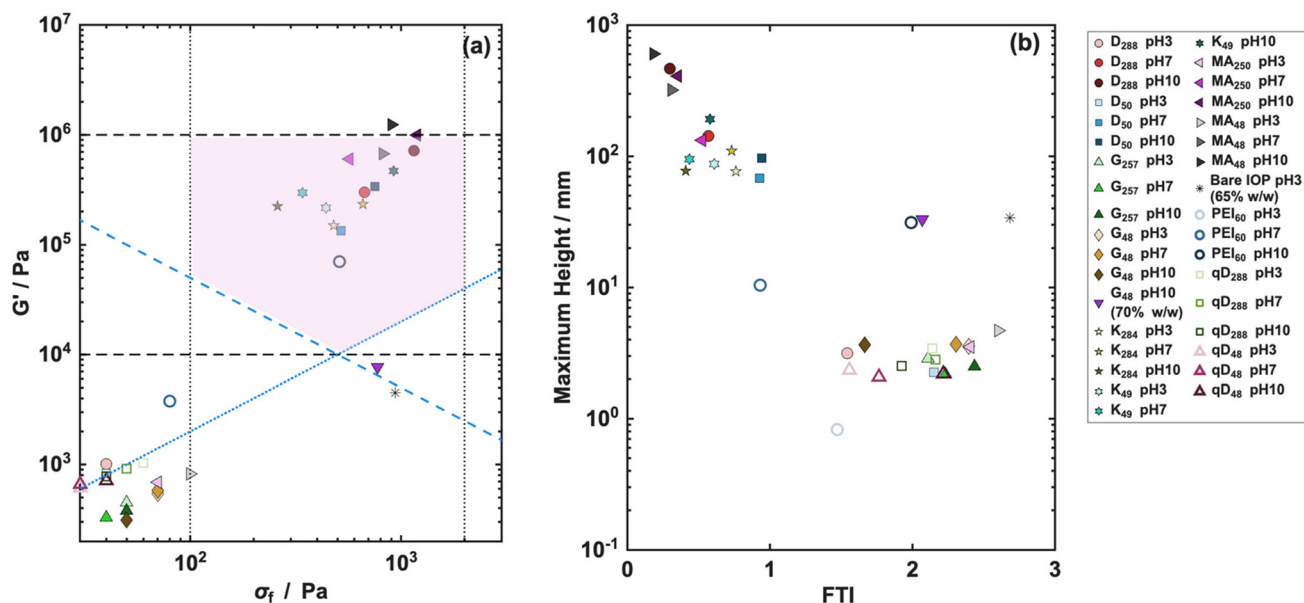


Fig. 7 Comparison of IOP suspension rheology against established printability.^{51,53,81} (a) storage modulus at LVR (G') vs. flow stress (σ_f) map to demonstrate suspension strength. The two vertical black dotted lines indicate σ_f limits and the two horizontal black dashed lines show G' limits for ceramic DIW. The blue dotted line is the figure of merit ($FOM = G'_{LVR}/\sigma_f = 20$) and the blue dashed line indicates Chan's printability criteria.⁸¹ The pink area is the optimised printability window. (b) The theoretical maximum printable height vs. Flow Transition Index ($FTI = \sigma_f/\sigma_y$),⁸⁶ where maximum height = $2\sigma_y/\rho g$ (where ρ is the ink density and g is gravitational acceleration),^{51,87} demonstrating suspension yield to flow transition, and printing height limitations. The solid loading of suspensions was 50% w/w, except for two samples with different solid loadings indicated in the legend.

Conclusions

A library of anionic, cationic, and non-ionic polymers with varying degrees of polymerisation were synthesised *via* RAFT polymerisation. The effects of these polymer additives on IOP dispersion stability were assessed through Zeta potential, DLS for hydrodynamic diameter, DCP for weight-intensity average diameter, and sedimentation tests. IOP dispersions without additives were stable at pH 3 but not at pH 7 and 10. Introducing polymers such as non-ionic G_{48} and cationic q-D₄₈, D₅₀, and PEI improved stability at pH 3 *via* electrosteric and steric repulsion. At pH 7 and 10, MA₄₈ and G_{48} significantly bolstered stability through electrosteric and steric repulsion, respectively, while other polymers had negligible or adverse effects. Rheological assessment of 50% w/w pristine IOP suspensions, and G_{48} and qD₄₈-containing 50% w/w IOP suspensions indicated a low viscosity, G'_{LVR} and flow stress at all pH values studied. D₅₀, PEI and MA₄₈-containing IOP suspensions demonstrated low viscosity, G'_{LVR} and flow stress at pH 3. These rheological properties were significantly increased at pH 7 and 10 due to hydrophobic association for D₅₀ and PEI-containing IOP suspensions and polymer network formation for MA₄₈-containing IOP suspensions. K₄₉-containing IOP suspensions consistently showed high viscosity at all pHs and was attributed to polymer networks preventing the flow of IOPs. Adding G_{48} , D₅₀, qD₄₈ and PEI reduced the viscosity of 50% w/w IOP suspensions at suitable pHs. However, as IOP loading rose from 50% to 70% w/w, charged polymers q-D₄₈,

D₅₀, and PEI unexpectedly increased in viscosity. Conversely, G_{48} addition consistently lowered the viscosity and flow stress of the suspensions, even at 70% w/w IOP loading.

These polymers played a pivotal role in improving the printability, shape retention and drying behaviour of 3D-printed IOP objects, demonstrated by printing thin-walled squares using various ink formulations onto aluminium substrates. The addition of PEI, K₄₉, or MA₄₈ at the appropriate pH notably enhanced the G'_{LVR} , improving shape retention and preventing cracking during drying. Addition of G_{48} facilitated increased IOP loadings in printable inks from 65% to 70% w/w, yielding low-shrinkage and crack-free green bodies. This multifaceted strategy encompassing polymer synthesis, stability evaluation, rheological characterisation, and 3D printing, underscores the potential of well-defined polymer additives to fine-tune ceramic ink rheology with minimal polymer content (0.5% w/w based on the nanoparticle loading), and lays the groundwork for the development for polymer additives tailored for ceramic ink formulations and advanced 3D printing applications.

Author contributions

The manuscript was written through contributions of all authors & all authors have given approval to the final version of the manuscript.



Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information: additional experimental methods; schematics of polymer additives synthesis; NMR of polymer additives; GPC of the polymer additives; SEM and TEM of the pristine IOPs; DCP size distribution of IOPs; rheology data; UV-Vis spectrophotometry data and photograph of 3D printed samples. See DOI: <https://doi.org/10.1039/d5lp00232j>.

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