

Cite this: *RSC Appl. Polym.*, 2024, **2**, 701

Poly(L-glutamic acid) augments the transfection performance of lipophilic polycations by overcoming tradeoffs among cytotoxicity, pDNA delivery efficiency, and serum stability†

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Polycations are scalable and affordable nanocarriers for delivering therapeutic nucleic acids. Yet, cationicity-dependent tradeoffs between nucleic acid delivery efficiency, cytotoxicity, and serum stability hinder clinical translation. Typically, the most efficient polycationic vehicles also tend to be the most toxic. For lipophilic polycations—which recruit hydrophobic interactions in addition to electrostatic interactions to bind and deliver nucleic acids—extensive chemical or architectural modifications sometimes fail to resolve intractable toxicity—efficiency tradeoffs. Here, we employ a facile post-synthetic polyplex surface modification strategy wherein poly(L-glutamic acid) (PGA) rescues toxicity, inhibits hemolysis, and prevents serum inhibition of lipophilic polycation-mediated plasmid (pDNA) delivery. Importantly, the sequence in which polycations, pDNA, and PGA are combined dictates pDNA conformations and spatial distribution. Circular dichroism spectroscopy reveals that PGA must be added last to polyplexes assembled from lipophilic polycations and pDNA; else, PGA will disrupt polycation-mediated pDNA condensation. Although PGA did not mitigate toxicity caused by hydrophilic PEI-based polycations, PGA tripled the population of transfected viable cells for lipophilic polycations. Non-specific adsorption of serum proteins abrogated pDNA delivery mediated by lipophilic polycations; however, PGA-coated polyplexes proved more serum-tolerant than uncoated polyplexes. Despite lower cellular uptake than uncoated polyplexes, PGA-coated polyplexes were imported into nuclei at higher rates. PGA also silenced the hemolytic activity of lipophilic polycations. Our work provides fundamental insights into how polyanionic coatings such as PGA transform intermolecular interactions between lipophilic polycations, nucleic acids, and serum proteins, and facilitate gentle yet efficient transgene delivery.

Received 7th March 2024,
Accepted 27th April 2024

DOI: 10.1039/d4lp00085d

rsc.li/rscapppolym

Introduction

Nucleic acids are transforming therapeutic landscapes, not only for relatively rare inherited disorders such as Duchenne's muscular dystrophy, but also for more prevalent diseases such as diabetes, hypercholesterolemia, or cardiovascular

diseases.^{1–3} Anionic, bulky, and hydrophilic, nucleic acids struggle to cross cell membranes and enter cellular targets, making engineered viral vectors indispensable. Although viral vectors are safe, efficient, and tissue-specific,^{4,5} they are prohibitively expensive to manufacture at scale, curtailing their utility in highly prevalent diseases where patient populations may span hundreds of millions.^{6–8} To overcome scale-up bottlenecks, synthetic nanocarriers such as lipid nanoparticles, polypeptides, polymers, or inorganic nanoparticles have been extensively investigated for the intracellular delivery of messenger RNA, plasmids (pDNA) or ribonucleoprotein payloads.^{9–12} Polymers, in particular, can be precisely tailored to fulfill design requirements imposed by diverse nucleic acid payloads and cellular targets.^{13,14}

Positive charge centers, typically installed as ionizable primary, secondary, or tertiary amines, are ubiquitous design

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†Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4lp00085d>



motifs. Polycations exploit electrostatic interactions to bind and condense nucleic acids into polyplexes. Subsequently, polycations must safeguard their payloads from degradation, improvise cellular uptake, and navigate endolysosomal interrogation.¹⁵ Long considered indispensable for polymer-mediated nucleic acid delivery, cationicity is also a double-edged sword. For instance, non-phagocytic cells internalize cationic nanocarriers at higher rates than anionic ones.^{16,17} But polycations disrupt the integrity of cell membranes, depolarize mitochondria, and accelerate autophagy-mediated degradation of their nucleic acid payloads.^{16,18} To juggle the seemingly conflicting demands of efficient nucleic acid delivery and the preservation of cellular viability and phenotype, several researchers modulated the distribution of positive charges along polymer backbones.^{19,20} Others employed charge conversion,^{21–23} fluorination,^{24–27} aromatic residues,^{28–32} or other lipophilic functionalities.^{33,34} Despite these time-consuming and iterative chemical or architectural modifications to promising polycations, tradeoffs between delivery efficiency and cytotoxicity still persist.^{35,36}

Cytotoxicity apart, polycations are also susceptible to non-specific protein adsorption,³⁷ which triggers rapid polyplex clearance by the reticuloendothelial system.^{38,39} Cationicity aggravates the deleterious effects of non-specific protein adsorption⁴⁰ especially since plasma proteins are predominantly negatively charged.⁴¹ Even during *ex vivo* gene delivery, non-specific adsorption of serum proteins inhibits the cellular uptake of polyplexes.⁴² Cationic polyplexes are generally serum-intolerant, requiring either serum-free or reduced serum media formulations such as OptiMEM. However, serum starvation reduces transgene expression in clinically valuable cell types, requiring us to engineer serum-compatible polycations that transfect diverse cellular targets.⁴³ Recognizing that careful control over cationicity is essential to resolve tradeoffs among nucleic acid delivery efficiency, toxicity, and serum compatibility, researchers incorporated anionic moieties within polycations, creating amphoteric polymers.^{44,45} Others employed electrostatic self-assembly rather than covalent conjugation to append anionic moieties that mask polyplex cationicity.⁴⁶ In this “layer-by-layer” approach, anionic glycosaminoglycans⁴⁷ such as heparin,⁴⁸ or hyaluronic acid,^{49–52} polypeptides,⁵³ or other polyanions are added to enhance polyplex interfacial properties.

Prior work on polyanionic surface modifiers focused overwhelmingly on poly-ethylene-imine (PEI), a hydrophilic (log *P* of -0.97 ^{54,55}) polycation and various polyanionic surface modifiers suppressed non-specific protein adsorption by PEI polyplexes.⁵³ In these studies, polyplex stoichiometry (the C/N/P ratio or the molar ratio of carboxylates in polyanions to amines in polycations to phosphate groups in nucleic acids) defined the magnitude and polarity of polyplex surface charge. For instance, Pack and colleagues found that if PGA incorporation exceeded a critical C/N/P threshold, PEI polyplexes became anionic,¹⁷ causing them to be internalized *via* clathrin-mediated rather than caveolin-mediated endocytosis

pathways. Clathrin-trafficked polyplexes would subsequently suffer entrapment within lysosomes, ultimately sabotaging their transfection efficiency. Similarly, for PEI polyplexes coated with poly(ethylene-*alt*-maleic acid), Pearson and colleagues employed N/P ratios as high as 30 to maintain polyplex cationicity.³⁸ Previous studies explored the roles of polyanion pK_a , polyplex stoichiometry, architectural and chemical modifications (such as acetylation) of PEI, focusing almost exclusively on hydrophilic polycations such as PEI rather than lipophilic polycations.^{56–61} Lipophilic polycations effect highly efficient transfection even at N/P ratios as low as 1, underscoring the importance of non-electrostatic pDNA binding interactions, particularly hydrophobicity.^{62–65} Because lipophilic polycations are less reliant on electrostatic interactions to bind and deliver pDNA,⁶⁶ it is unclear whether electrostatically appended polyanionic coatings will conformally coat polyplex surfaces or substantially improve transfection outcomes. In this work, we demonstrate that surface modification with poly(L-glutamic acid) (PGA) helps lipophilic polycations navigate toxicity–efficiency tradeoffs, prevents polycation-induced hemolysis, and promotes plasmid (pDNA) delivery in serum-containing media.

Owing to its dense negative charge, hydrophilicity, and biodegradability, poly(L-glutamic acid) or PGA is an attractive stealth coating for nanocarriers.^{67–70} In a head-to-head comparison between PGA, alginate, heparan, and poly(aspartic acid), PGA was most effective in inhibiting the non-specific adsorption of serum proteins.⁷¹ Hammond and coworkers reported that PGA-coated nanoparticles bind strongly and specifically to ovarian cancer cells^{67,72} and established that PGA reconfigures intracellular trafficking pathways to favor endosome-evading caveolar pathways over clathrin-mediated pathways.^{67,72} We chose PGA to augment the interfacial properties and transfection performance of lipophilic polycations due to its anti-inflammatory properties, endosome-evading intracellular trafficking patterns, and potential for receptor-mediated uptake.^{73,74}

We selected *p*(DIPAEMA₅₆-*st*-HEMA₄₈) (DIP50H50) as a model lipophilic polycation (Fig. 1), since toxicity–efficiency tradeoffs undermine its utility as a pDNA carrier.^{75,76} Although this contribution focuses on DIP50H50 and PGA as a model system, our findings will be broadly valuable for other combinations of lipophilic polycations and polyanionic surface modifiers. Our paper informs the design of ternary polyplexes by contributing three fundamentally valuable insights. First, we thoroughly investigated the sequence of addition of the lipophilic polycations (DIP50H50), pDNA payloads, and PGA during the assembly of ternary polyplexes—a variable that has not been explored previously—and found that the addition sequence governed pDNA spatial distribution and conformations. Second, although PGA mitigated toxicity–efficiency tradeoffs for DIP50H50, the toxicity of PEI-based polyplexes remained unaffected. Third, we employed quantitative confocal microscopy to map differences in nuclear localization between uncoated and PGA-coated polyplexes. Our work illuminates binding interactions between polycations, nucleic



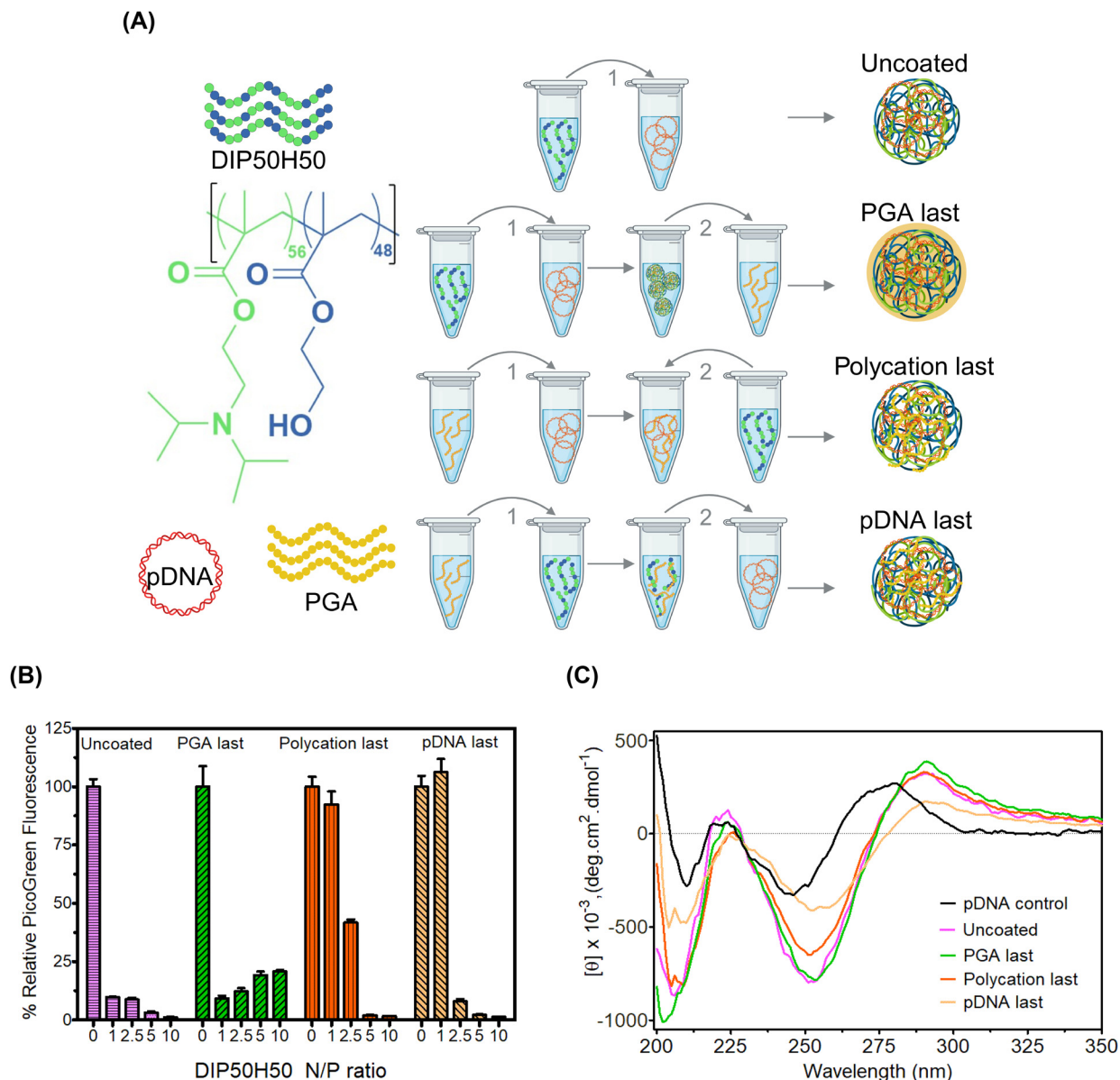


Fig. 1 We varied the sequence of addition of polycations (DIP50H50), pDNA, and poly(L-glutamic acid) or PGA. We compared pDNA encapsulation efficiency and pDNA conformations across four addition sequences. (A) Overview of addition schemes. (B) PicoGreen dye exclusion assays compared DIP50H50–pDNA binding across uncoated polyplexes and three groups of PGA-coated polyplexes (PGA last, polycation last, and pDNA last). At lower N/P ratios, PGA disrupted DIP50H50–pDNA interactions in polycation last and pDNA last polyplexes. (C) CD spectroscopy revealed that the pDNA conformations in polycation last and pDNA last polyplexes resemble those of unbound pDNA; we did not observe this conformation in PGA last polyplexes.

acids, and serum proteins, facilitating rational design of surface modification strategies for lipophilic polycations.

Results and discussion

Here, we demonstrate that PGA mitigates tradeoffs between cytotoxicity, pDNA delivery efficiency, and serum stability for polyplexes formed from the lipophilic polycation, DIP50H50.

First, CD spectroscopy identified the optimal sequence of addition of DIP50H50, pDNA, and PGA to mask cationicity without triggering pDNA leakage. Subsequently, we performed pDNA delivery assays, establishing that PGA is a highly effective surface modifier that triples the population of transgene-expressing cells arising from DIP50H50-mediated transfection. PGA also assists DIP50H50 in forming stable polyplexes in serum-containing media, potentiating serum-tolerant transfection. Confocal analysis revealed that, despite lower cel-



lular uptake, PGA-coated polyplexes have a higher propensity to occupy nuclear interiors than uncoated polyplexes. Finally, hemolysis assays demonstrated that PGA augments the hemocompatibility of DIP50H50 polyplexes.

Addition sequence dictates polycation–pDNA binding configurations

Our rationale is that for PGA to overcome tradeoffs among pDNA delivery efficiency, toxicity, and serum stability without perturbing DIP50H50–pDNA binding interactions, the addition sequence must be optimized first. PGA molecules must occupy polyplex exteriors instead of competing with pDNA for polycation binding sites in polyplex cores. The sequence of addition of polycations and pDNA modulates exerts dramatic effects on polyplex size distribution and encapsulation efficiency.^{77,78} However, we do not understand how the sequence in which polycations, pDNA, and PGA are combined (Fig. 1) impacts polyplex properties and performance. We tested four addition sequences (Fig. 1A): (1) for uncoated polyplexes, we added the polycation, DIP50H50 to pDNA and realized desired N/P ratios (molar ratio of protonatable amines in DIP50H50 to phosphates in pDNA), (2) for “PGA last” polyplexes, we added PGA to uncoated polyplexes, (3) for “polycation last” polyplexes, we added DIP50H50 to a mixture of PGA and pDNA, (4) for “pDNA last” polyplexes, we added the pDNA payload to a mixture of DIP50H50 and PGA. We kept the C/N ratio (molar ratios of carboxylates in PGA to amines in DIP50H50) fixed at 1 for sequences (2)–(4). During preliminary experiments, we discovered that a C/N ratio of 1 was sufficient to realize polyplex charge reversal from cationic to anionic, and that higher C/N ratios did not offer any benefits.

For all three addition sequences, PGA reverses the polarity of polyplex surface charge from positive to negative, confirming that PGA effectively masks cationicity (Fig. S2(a) in the ESI†). Further, the addition sequence had no impact whatsoever on polyplex size distributions (Fig. S2(b) in the ESI†). Using dye exclusion assays we asked whether the addition sequence influenced pDNA encapsulation efficiency (Fig. 1B). In PicoGreen assays, fluorescence quenching estimates the strength of DIP50H50–pDNA binding. The fluorescence intensity of uncoated polyplexes decreased steeply upon complexation with DIP50H50; strong DIP50H50–pDNA binding was observed at N/P ratios as low as 1. In PGA last polyplexes, results mirrored that of the uncoated polyplexes at an N/P value of 1. However at this N/P value, polycation last and pDNA last polyplexes exhibited four- to eight-fold higher fluorescence intensities than uncoated polyplexes. However, a large excess of polycation counteracted the destabilizing influence of PGA in polycation last and pDNA last polyplexes. For instance, the fluorescence intensity at the highest N/P ratios (N/P of 5 and 10) in the polycation last and pDNA last polyplexes was on par with values from the uncoated and the PGA last polyplexes.

To deepen our understanding of how PGA addition sequence impacts DIP50H50–pDNA binding, we performed circular dichroism (CD) spectroscopy, a highly sensitive tool that detects helicity changes as pDNA is condensed. CD tracked

pDNA conformational changes as a function of addition sequence (Fig. 1C). Neither DIP50H50 nor PGA generated CD signals above 220 nm (Fig. S3 in the ESI†), so any CD signal above 220 nm is from pDNA alone. CD spectra were normalized with respect to relative pDNA concentration, which was determined from the absorbance maximum (between 250–275 nm). Uncomplexed pDNA exhibited a positive band around 280 nm and a negative band around 250 nm, consistent with its native B-type helical conformation. When DIP50H50 is added to pDNA, CD spectra shifted to the right and the molar ellipticity increased. This increased ellipticity is consistent with polycation-induced changes in pDNA conformation, wherein DIP50H50 collapses the native B-type helical structure of pDNA, triggering a conformational shift to C-type structures.⁵³

CD spectra for PGA last polyplexes and uncoated polyplexes were near-identical, suggesting the PGA forms an anionic coating around polyplexes without perturbing DIP50H50–pDNA interactions in polyplex cores. As for polycation last polyplexes, the ellipticity of the negative band at 250 nm was lower than uncoated and PGA last polyplexes. We attribute this reduced ellipticity to competition between pDNA and PGA for binding sites on DIP50H50. In polycation last polyplexes, pDNA has fewer opportunities to complex with DIP50H50, hindering pDNA condensation and depressing ellipticity. Finally, the CD spectra of pDNA last polyplexes closely resembled uncomplexed pDNA. In pDNA last polyplexes, polycations bind PGA preferentially over pDNA, leaving fewer polycation binding sites available for pDNA payloads. Unencumbered by polycations, uncondensed pDNA occupies the polyplex exterior, adopting conformations nearly identical to that of uncomplexed pDNA. Previously, CD spectroscopy tracked the unfolding of α -helices in PGA and identified an isodichroic point at 204 nm, where folded and unfolded PGA have the same CD signal.⁷⁹ This 204 nm signal is far more prominent in polyplexes where PGA is added last compared to those where polycation or pDNA is added last. Cumulatively, our data suggests that the sequence of addition influences the organization of pDNA and PGA within the polyplexes.

The order in which polycations, PGA, and pDNA are combined governs how tightly DIP50H50 binds pDNA payloads and the extent to which PGA interferes with DIP50H50–pDNA complexation. PGA will be most effective as a surface modifier when the PGA last addition sequence is employed; in other sequences, PGA weakens DIP50H50–pDNA binding.

PGA resolves cationicity-dependent tradeoffs between pDNA delivery efficiency and cytotoxicity

Based on CD spectroscopy results, we decided to focus on PGA last polyplexes in the rest of the paper; results from other addition sequences are furnished in the ESI (Fig. S4 and S8–S10†) Having identified PGA last as the optimal addition sequence to modify polyplex surfaces without triggering pDNA leakage, we next asked whether PGA helps DIP50H50 overcome toxicity–efficiency tradeoffs (Fig. 2). We formulated uncoated and PGA-coated DIP50H50 polyplexes with GFP-encoding pDNA and compared transfection efficiency and cell viability



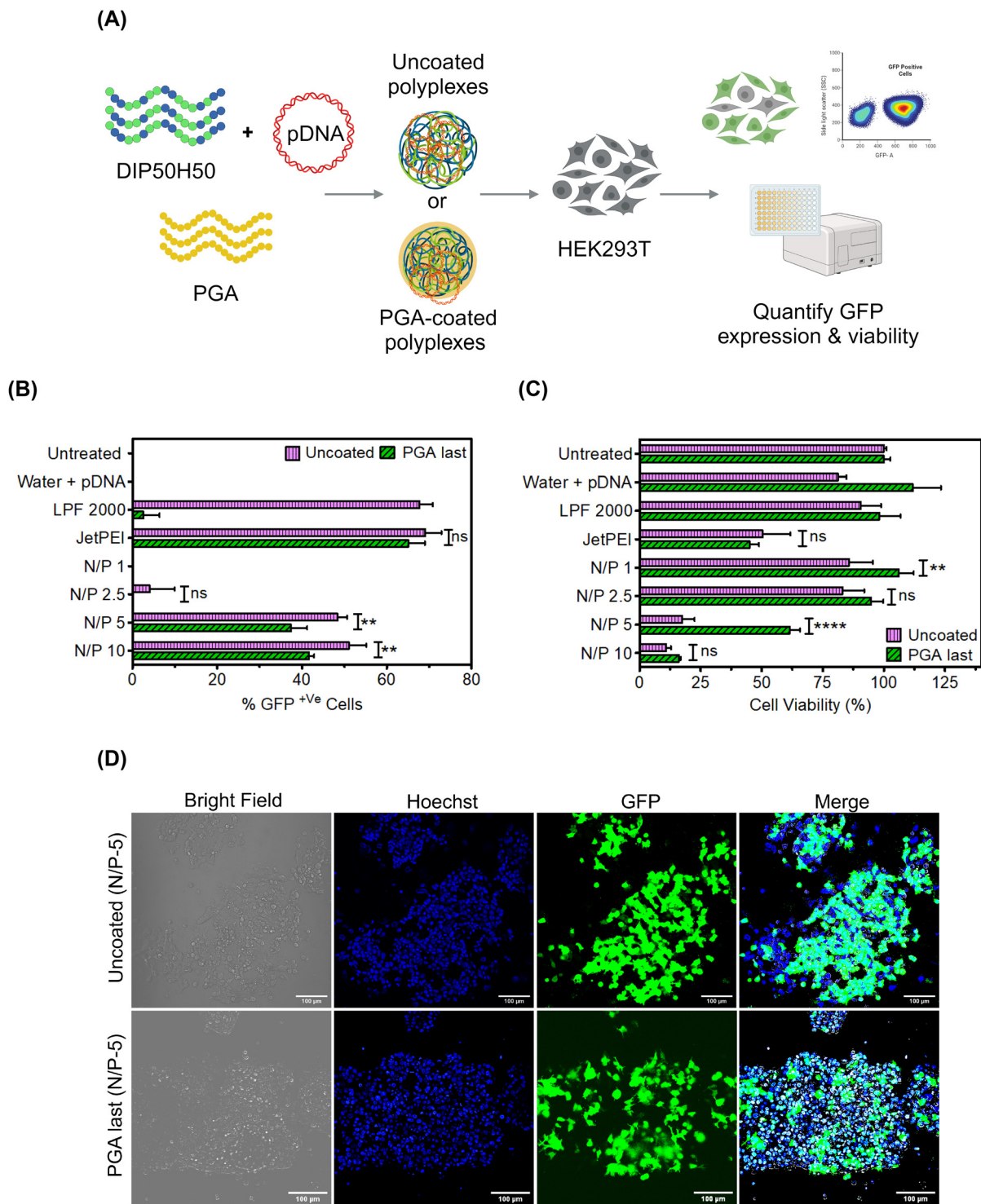


Fig. 2 PGA-coated polyplexes mediate gentle yet efficient transfection. (A) We complexed GFP-encoding pDNA with DIP50H50 and administered uncoated or as PGA-coated polyplexes to HEK293T cells. (B) The percentage of GFP⁺ cells was slightly lower among PGA-coated than uncoated polyplexes but the population of GFP⁺ cells tripled. (C) Although PGA alleviated toxicity induced by DIP50H50, PGA coating did not rescue cytotoxicity triggered by JetPEI. (D) Representative fluorescent micrographs. Scale bar is 100 μ m.

(Fig. 2A). Lipid- and PEI-based commercial reagents for pDNA delivery, Lipofectamine 2000 and JetPEI, were employed as positive controls. Although uncoated polyplexes exhibited

marginally higher transfection efficiency than PGA-coated polyplexes (Fig. 2B and D), cell viability was three-fold higher for PGA-coated polyplexes (Fig. 2C).



The percentage of GFP⁺ cells—the fraction of viable cells expressing GFP—can be a misleading measure of transfection efficiency. Since we assign equal priority to cellular viability and transfection efficiency, the number of GFP⁺ cells is a more useful metric than either the percentage of GFP⁺ cells or cell viability alone. PGA-coated polyplexes exhibited lower transfection efficiency (by 10–15%) and higher viability (by 10–35%) than uncoated polyplexes. PGA tripled the population of GFP⁺ cells (Fig. S5 in the ESI†) relative to uncoated polyplexes and helped DIP50H50 mediate gentle yet efficient pDNA delivery. From our correlation analysis (Fig. S6 and Table S1 in the ESI†), we observed a strong negative correlation (-0.935 ± 0.55) between cell viability and transfection efficiency in uncoated polyplexes but PGA coating weakened this correlation considerably (-0.84 ± 0.13). We concluded that surface modification with PGA mitigates cationicity-dependent tradeoffs between cytotoxicity and delivery efficiency for lipophilic polycations like DIP50H50.

Interestingly, PGA exerted divergent effects on the transfection performance of lipophilic (DIP50H50, $\log P$ of 6.4⁷⁵) and hydrophilic polycations (JetPEI, $\log P$ of $-0.97^{54,55}$). While PGA blunted the toxicity of DIP50H50, JetPEI was largely indifferent to PGA, exhibiting similar toxicity and transgene expression levels with or without PGA. We measured the reduction in ζ -potential for JetPEI and DIP50H50 polyplexes after PGA coating (Fig. S7(a) in the ESI†), and observed striking differences. PGA transformed the surface character of DIP50H50 polyplexes from moderately cationic ($+17 \pm 1$ mV) to strongly anionic (-30 ± 4 mV), suggesting that PGA conformally covered polyplex surfaces. In contrast, PGA did not reduce the surface charge of JetPEI polyplexes ($+27 \pm 3$ mV to -2.5 ± 15 mV) to the same extent as it did with DIP50H50, suggesting that the surface coverage of JetPEI polyplexes by PGA was patchy and incomplete. We attribute the contrasting effects of PGA on the transfection performance DIP50H50 and JetPEI to incomplete surface coverage of JetPEI polyplexes by PGA. Overall, PGA proved to be a more effective surface modifier for polyplexes formed from lipophilic polycations than hydrophilic polycations.

PGA rescues polyplexes from serum-induced aggregation without exacerbating serum-mediated pDNA unbinding

We anticipated that PGA-coated DIP50H50 polyplexes will resist aggregation in serum-containing media (Fig. 3A), potentiating serum-tolerant transfection. Since serum proteins may displace pDNA from polyplexes by competing with pDNA for binding sites on DIP50H50, we tested whether PGA-coated polyplexes were more susceptible to serum-mediated pDNA release than uncoated polyplexes. Uncoated polyplexes and three sets of PGA-coated polyplexes were incubated in either a 10% or 50% solution of serum extracted from whole human blood. Size distributions, electrokinetic analyses, and pDNA encapsulation efficiencies for all addition sequences and

serum concentrations are furnished in Fig. S8–S10 in the ESI,† respectively.

We measured the size distributions of uncoated and PGA-coated polyplexes in water and in serum using dynamic light scattering (DLS). In water, the hydrodynamic diameters of uncoated and PGA-coated polyplexes were comparable (<300 nm). In human serum, uncoated polyplexes aggregated severely, with hydrodynamic diameters approaching 7–8 × their values in water (Fig. 3B). Further, serum-induced aggregation of uncoated polyplexes was strongly N/P-dependent with the highest sizes (diameters approaching 3 μm) observed for N/P 10 polyplexes. In contrast, the hydrodynamic diameters of PGA-coated polyplexes were nearly identical in water and in serum (Fig. 3B).

To substantiate conclusions from DLS, we performed transmission electron microscopy (TEM) of uncoated and PGA-coated polyplexes (N/P ratio of 5) in water and in 10% serum. Both uncoated and PGA-coated polyplexes formed well-defined sub-100 nm assemblies in water but micron-scale aggregates appeared prominently when uncoated polyplexes were incubated in serum (Fig. 3C). Consistent with observations from DLS, PGA-coated polyplexes exhibited comparable size distributions in water and in serum in TEM images. Serum exposure transformed uncoated polyplexes from cationic to anionic (Fig. 3D). PGA-coated polyplexes were already negatively charged in water; anionic serum proteins may have displaced PGA from polyplex surfaces, modifying the magnitude of negative charge.

Serum-triggered pDNA release was comparable across PGA-coated and uncoated polyplexes (Fig. 3C), suggesting that PGA does not exacerbate competitive unbinding of pDNA in serum. Physicochemical characterization of polyplex–serum interactions established that PGA augments polyplex colloidal stability in serum without aggravating pDNA unbinding.

Despite depressing cellular uptake, PGA helps lipophilic polycations mediate pDNA delivery in serum-containing media

After verifying that PGA prevented polyplex aggregation in 10% serum without exacerbating pDNA unbinding, we evaluated pDNA delivery efficiency of PGA-coated polyplexes in serum-containing cell culture media. Serum proteins inhibit polycation-mediated pDNA delivery, forcing experimenters to perform transfection in reduced serum media (*e.g.*, proprietary formulations such as OptiMEM) or in serum-free media.⁸⁰ We compared five media formulations (Fig. 4A): (1) OptiMEM, a proprietary reduced-serum formulation that is ubiquitously used in polycation-mediated delivery, (2) serum-free DMEM without fetal bovine serum (FBS) and (3–5) DMEM with FBS supplementation of 2.5, 5, or 10% v/v. We compared the transfection efficiency of uncoated and PGA-coated polyplexes in each of the above five media formulations.

PGA-coated polyplexes fared worse than uncoated polyplexes (Fig. 4B) in OptiMEM but performed better in DMEM (across all serum concentration levels). Serum concentration



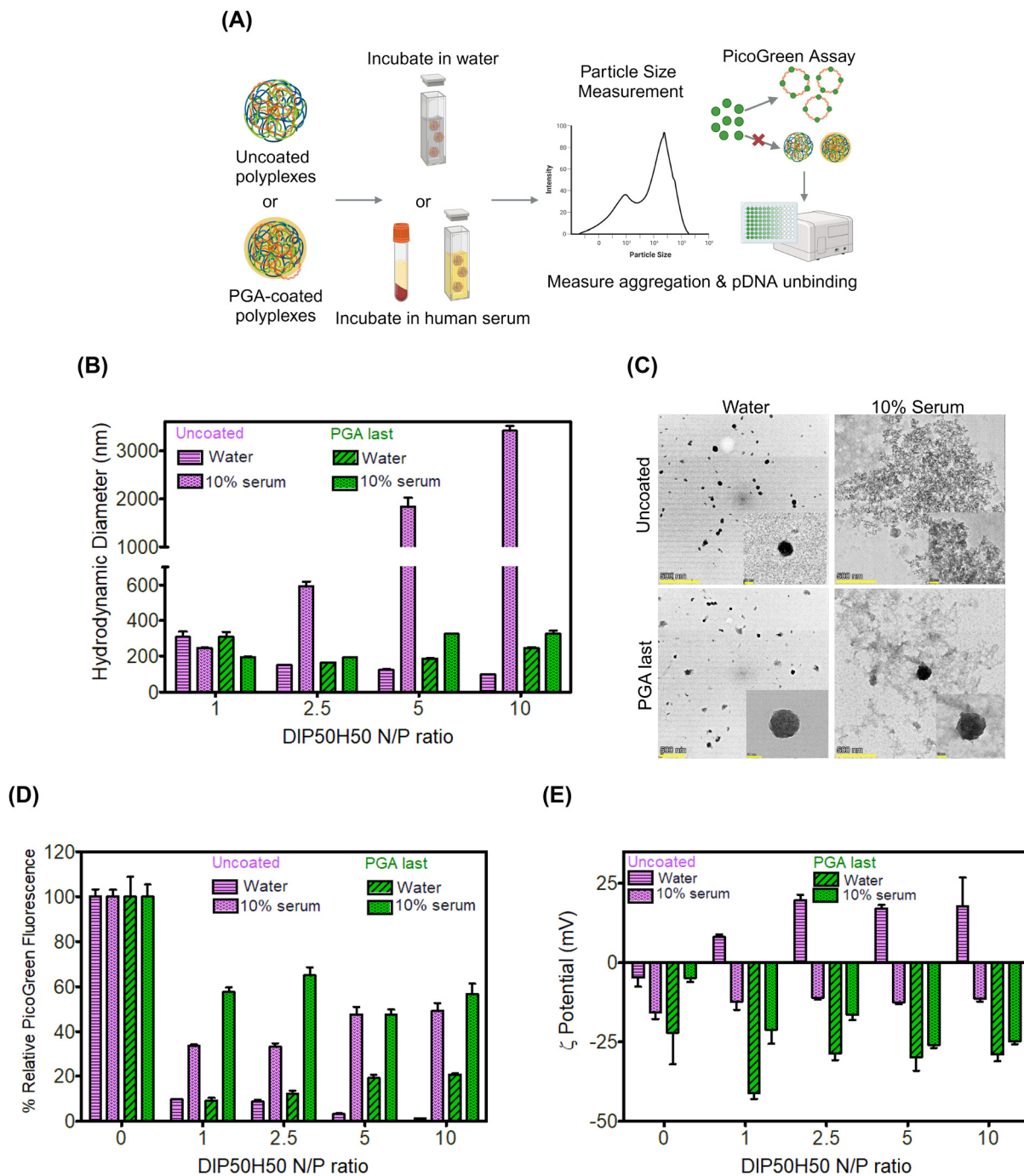


Fig. 3 PGA stabilizes lipophilic polyplexes against aggregation in serum. (A) We tracked changes in hydrodynamic size, pDNA binding, and polyplex charge after serum exposure. (B) Uncoated polyplexes aggregated severely but PGA-coated polyplexes remained unchanged in size after serum incubation. (C) TEM confirms that PGA rescued polyplexes from aggregation in serum (scale bar 500 nm and 50 nm in image and inset respectively). (D) Serum-triggered pDNA release was comparable for uncoated and PGA-coated polyplexes. (E) Serum incubation reversed the surface charge polarity of uncoated polyplexes from cationic to anionic.

levels as low as 2.5% completely abolished the transfection of uncoated polyplexes but only halved the transfection efficiency of PGA-coated polyplexes. At higher serum concentration levels (5% and 10%), the gulf in transfection performance between uncoated and PGA-coated polyplexes grew wider.

Interestingly, the transfection efficiency of uncoated polyplexes was lower in serum-free DMEM (0% FBS) than in OptiMEM. We attribute this result to differences in glucose concentration between OptiMEM and DMEM; DMEM contains more glucose (which has previously been shown to impact cell metabolism⁸¹) than OptiMEM.⁸² However, DMEM may lack



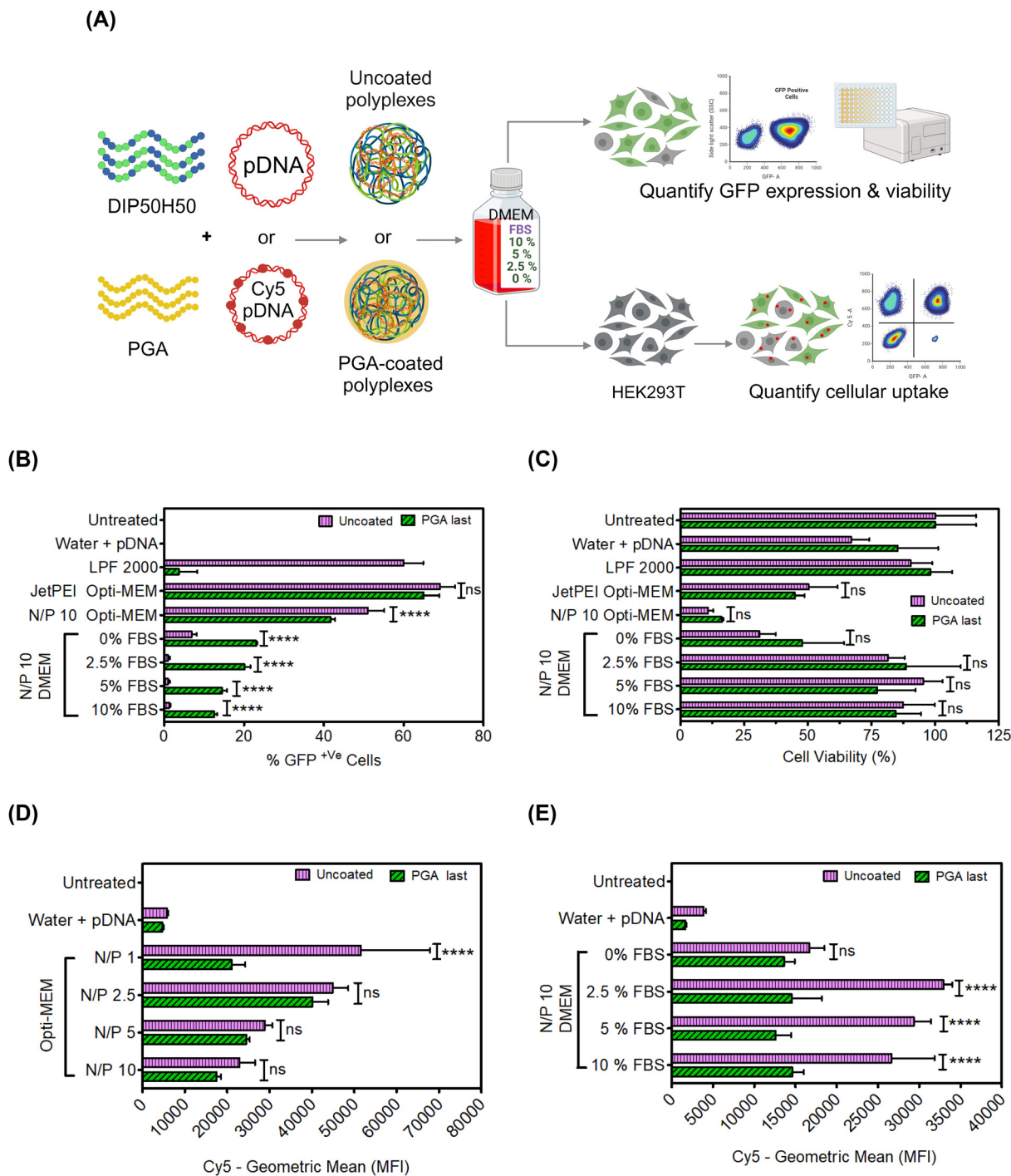


Fig. 4 Despite lowered cellular uptake, PGA-coated polyplexes facilitate pDNA delivery more efficiently than uncoated polyplexes in serum-containing media. (A) Overview of cell culture media formulations tested. (B) Uncoated polyplexes exhibited high transfection efficiency in OptiMEM, a reduced serum media formulation but serum supplementation abolished GFP expression. In contrast, PGA coated polyplexes promoted intracellular pDNA delivery even at serum concentrations as high as 10%. (C) Polycation-induced toxicity was far lower in serum-containing media than in OptiMEM or in serum-free DMEM. In Opti-MEM (D) and in serum-supplemented DMEM (E), PGA-coated polyplexes exhibited lower cellular uptake than uncoated polyplexes.

other biomolecules found in OptiMEM; OptiMEM is a proprietary formulation, whose components remain a matter of conjecture, preventing us from digging deeper into this result. Another interesting albeit unexpected result was that uncoated

polyplexes underperform PGA-coated polyplexes in serum-free DMEM (0% FBS) but exhibit higher transfection efficiencies in the presence of FBS. We speculate that serum proteins from FBS exert divergent effects on uncoated and PGA-coated



polyplexes, with some adsorbed serum proteins hindering transfection and others promoting uptake and transfection. To fully understand how serum proteins from FBS or OptiMEM components interact with uncoated and PGA-coated polyplexes, we must characterize the composition of protein coronae formed around polyplexes.^{83–85} Protein corona investigations of polyplexes are expensive and challenging, requiring

careful sample preparation and extensive method development.^{86,87} We opted to pursue proteomics investigations in a future study.

Cell viability was much higher in serum-containing media than in OptiMEM (Fig. 4C), particularly for PGA-coated polyplexes (Fig. S11 in the ESI†). By potentiating rescuing cells from toxicity associated with serum starvation, PGA delays the

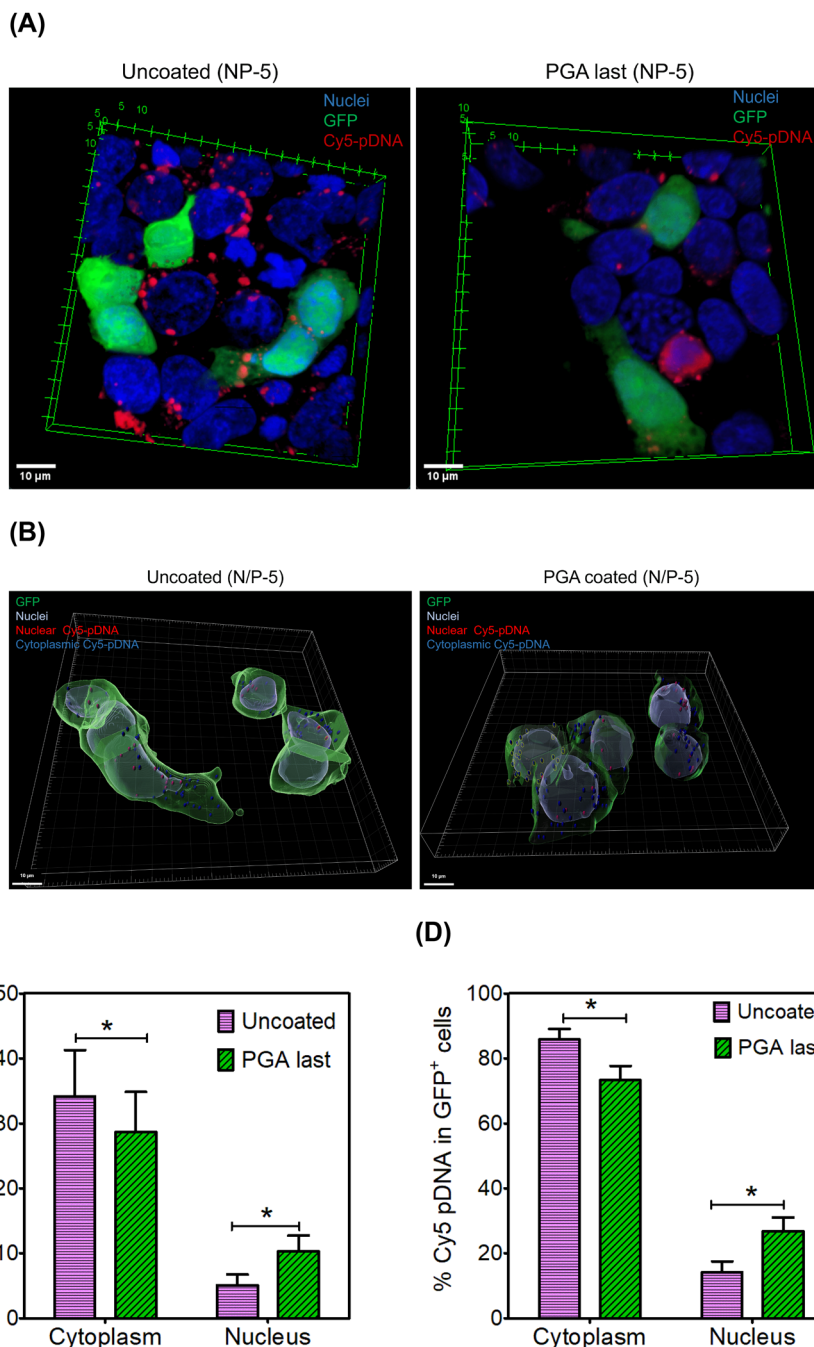


Fig. 5 PGA-coated polyplexes are imported into nuclei at higher rates than uncoated polyplexes. (A) HEK293T cells were transfected with uncoated and PGA-coated polyplexes at an N/P ratio of 5. Nuclei are stained with DAPI (blue), pDNA payloads were labeled with Cy5 (red), and transgene expression visualized using GFP (green). Scale bar is 10 μ m. (B) Three-dimensional reconstructions highlight differences in pDNA distribution (between nuclear and cytoplasmic regions) for GFP⁺ cells. Scale bar is 10 μ m. In terms of both absolute polyplex numbers (C) and relative distribution (D), PGA-coated polyplexes display a higher propensity for nuclear localization than uncoated polyplexes.



onset of cellular stress responses that inhibit GFP expression.^{88–90}

Next, we compared cellular internalization for PGA-coated and uncoated polyplexes in diverse media formulations (reduced serum OptiMEM and DMEM with serum concentration levels spanning 0–10% v/v). We labeled pDNA with Cy5 fluorophores,⁷⁶ and measured Cy5 fluorescence intensity in OptiMEM and DMEM (Fig. 4D and E respectively). In

OptiMEM, PGA-coated polyplexes were taken up by cells at lower levels than uncoated polyplexes. In the presence of serum, PGA-coated polyplexes are smaller than uncoated polyplexes, which in turn lowers their sedimentation velocity and lowers the frequency of cell–polyplex contact.⁹¹ Despite lower cellular uptake, PGA-coated polyplexes displayed higher transfection efficiency than uncoated polyplexes in serum-containing media (Fig. 4D and E). Cellular uptake, therefore, cannot

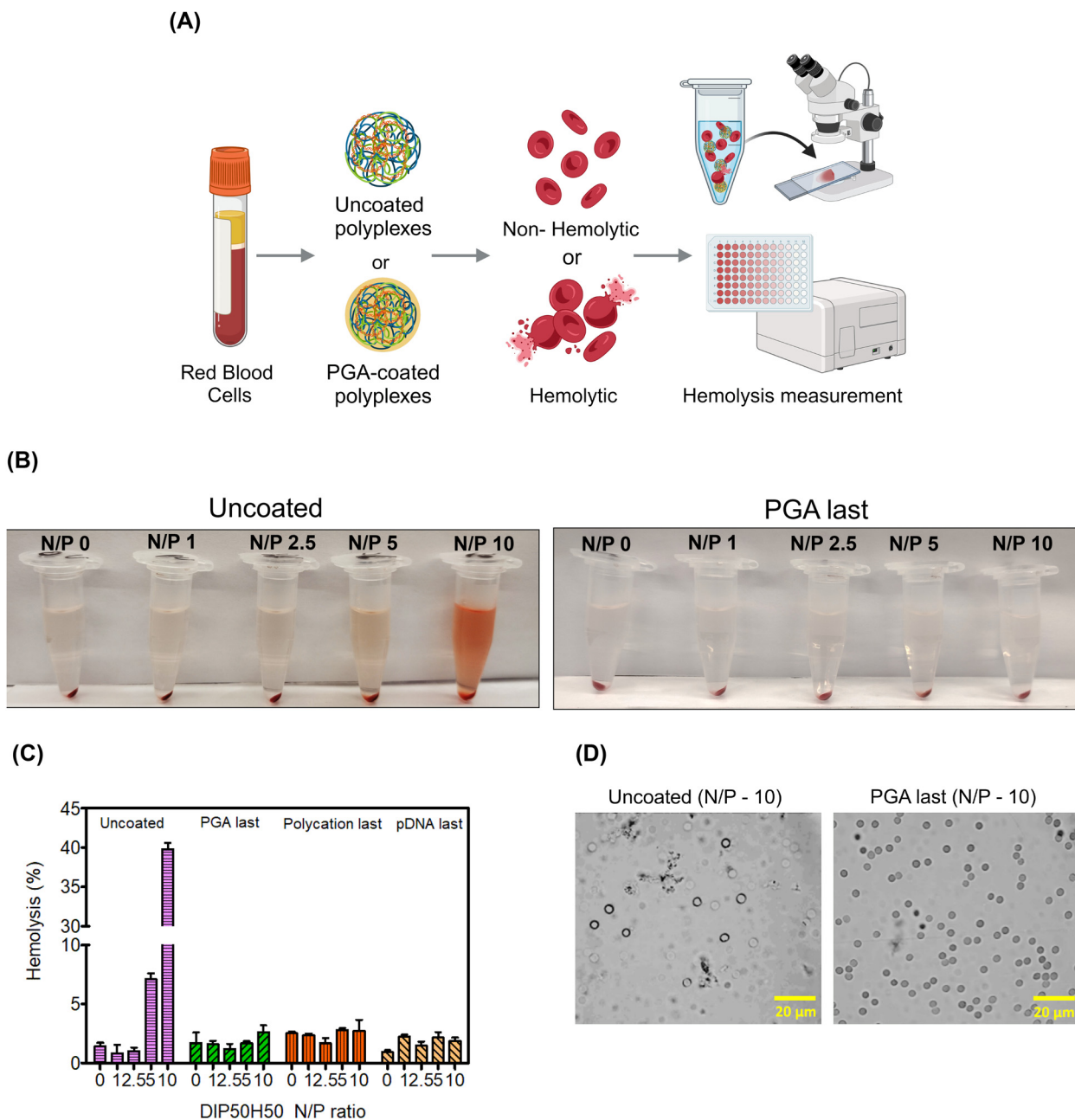


Fig. 6 PGA shields red blood cells (RBCs) from polyplex-mediated hemolysis. (A) Overview of hemolysis assay. (B) Hemolysis is accompanied by pink colouration in the supernatant whereas a colorless supernatant indicates that RBCs remain intact. Visual inspection of supernatant color reveals that PGA coating inhibits hemolysis even at N/P ratios as high as 10 whereas uncoated polyplexes trigger hemolysis even at lower N/P ratios (N/P of 5). (C) PGA-coated polyplexes induced hemolysis in fewer than 5% of RBCs (across all 3 addition sequences and all N/P ratios) whereas uncoated polyplexes caused up to 40% of RBCs to lyse at an N/P ratio of 10 (D) Micrographs of RBC suspensions treated with uncoated and PGA-coated polyplexes. Scale bar is 20 μm .



explain PGA-mediated enhancements in transfection efficiency in serum-containing media. To verify whether PGA-coated polyplexes compensate for depressed cellular uptake by routing polyplexes along intracellular trajectories culminating in nuclear import, we performed confocal microscopy.

PGA coating promotes nuclear import of pDNA

We hypothesized that PGA will reroute polyplexes along intracellular itineraries that promote nuclear import. To test this hypothesis, we mapped the intracellular distribution of Cy5-labeled polyplexes using confocal microscopy. We formulated uncoated and PGA-coated polyplexes with Cy5-labeled pDNA at an N/P ratio of 5. Twenty-four hours after polyplex administration, we fixed and stained cells with Hoechst (Fig. 5A). Interestingly, Cy5 intensity was noticeably lower in the PGA-coated condition than in the uncoated group (Fig. S12 and S13 in the ESI[†]), corroborating uptake measurements (Fig. 4D).

Internalized polyplexes were classified as cytoplasmic (cyan) and nuclear (white) based on the spatial location of Cy5 signals (Fig. 5B). Thereafter, we enumerated polyplexes (Fig. 5C) and computed the fraction of pDNA partitioning between cytoplasmic and nuclear regions (Fig. 5D). Among GFP⁺ cells, pDNA is far more likely to accumulate within the nuclear periphery when polyplexes are modified with PGA (Fig. 5D). Although uncoated polyplexes are internalized more efficiently than their PGA-coated counterparts, nuclear import was significantly higher for PGA-coated polyplexes. PGA might be rerouting polyplexes along caveolar rather than clathrin-mediated pathways, consistent with reports by¹⁷ Consequently, polyplexes trafficked along caveolar pathways (PGA-coated) enjoy a higher likelihood of nuclear import than clathrin-trafficked polyplexes (uncoated). Although we found fewer polyplexes per cell in the PGA-coated condition than in the uncoated condition, PGA-coated polyplexes were more likely to inhabit nuclear regions than uncoated polyplexes. One limitation of our study is that fluorescence labeling is poorly suited for accurate quantification of pDNA within nuclei. This is because fluorescent labeling alters the hydrophobicity of pDNA, potentially impacting their intracellular trajectories. In future work, we will compare nuclear import using label-free techniques such as the qPCR method developed by Szoka *et al.*⁹²

PGA silences the hemolytic activity of polyplexes

Since red blood cells (RBCs) are the most abundant cell type in the blood stream,⁹³ studying RBC lysis will illuminate the role of PGA in augmenting polyplex hemocompatibility.⁹⁴ Polyplex-induced toxicity in HEK293T cells does not predict hemolysis since RBCs possess distinct cellular characteristics. For instance, RBC plasma membranes are unusually thin to facilitate oxygen uptake, RBCs are also smaller and more discoid, and lack organelles that may initiate cellular repair following polycation-mediated lysis.^{93,95}

We determined whether PGA improves RBCs' tolerance to polyplexes. We harvested RBCs from healthy human donors and exposed them to uncoated and PGA-coated polyplexes.

Then, we measured RBC lysis as a function of N/P ratio and surface treatment (Fig. 6A). To quantify hemolysis, we measured the absorbance of hemoglobin liberated from lysed cells into the supernatant. For RBCs treated with uncoated polyplexes (Fig. 6B), the pinkish red colour in the supernatant intensified progressively with increasing N/P ratio, confirming that uncoated polyplexes are detrimental to RBC health. On the other hand, PGA-coated polyplexes maintained a transparent colorless supernatant even at the highest N/P ratio tested (N/P = 10), suggesting that PGA protected RBCs from polycation-triggered lysis. Plate reader measurements (Fig. 6C) confirmed visual observation, with the highest absorbance detected for uncoated polyplexes (N/P = 10). Interestingly, whether PGA protects RBCs from polyplex-induced hemolysis is independent of addition sequence, with PGA silencing the hemolytic activity of DIP50H50 for all three PGA-coated polyplexes (PGA last, polycation last and pDNA last).

PGA effectively inhibits lipophilic polycations from triggering hemolysis, potentially augmenting polyplex hemocompatibility. In future work we will study polyplex interactions with platelets and white blood cells, and elucidate how PGA impacts complement activation, platelet aggregation, and blood clotting.

Conclusions

Our study demonstrates the utility of polyplex surface modification in navigating tradeoffs among toxicity, transfection efficiency, and serum stability during polycation-mediated gene delivery. Instead of pursuing chemical and architectural modifications of a promising lipophilic polycation for pDNA delivery (DIP50H50), we augmented cytocompatibility and serum stability through post-synthetic surface modification with PGA. The conformations adopted by pDNA payloads and the degree to which polycations condensed pDNA was highly sensitive to the sequence of addition of PGA, polycations, and pDNA—a variable that to the best of our knowledge has never been studied before. Although PGA dramatically expanded the population of transfected cells in the DIP50H50 treatment condition, PGA had no impact on JetPEI-induced toxicity. Interestingly, PGA-coated polyplexes lagged uncoated polyplexes in cellular uptake but still exhibited higher pDNA delivery efficiency in serum-containing media. This suggested that intracellular transport, rather than cellular uptake, is the key delivery bottleneck. To understand differences in the intracellular fates of uncoated and PGA-coated polyplexes, we studied how each polyplexes partition between nuclear and cytoplasmic regions using quantitative confocal microscopy, and found that PGA-coated polyplexes were more likely to undergo nuclear import than uncoated polyplexes. Uncoated polyplexes triggered lysis in almost 50% of red blood cells, but PGA inhibited hemolysis by masking polyplex cationicity. PGA also shielded polyplexes from non-specific protein adsorption and aggregation in serum-containing media. By carefully characterizing binding interactions between polycations, pDNA



payloads, serum proteins, and cells, our paper provides a useful framework to rationally design polyplex surface modification protocols. In the future, we foresee that the protein corona and cell type specificity of polyplexes can be rationally tuned by individually tailoring polyanionic coating protocols for each polycation.

Experimental section

Materials

2-(diisopropylamino)ethyl methacrylate (DIPAEMA), 2-hydroxyethyl methacrylate (HEMA), 4-cyano-4-[(ethylsulfanylthiocarbonyl)sulfanyl]pentanoic acid (CEP), 4,4'-azobis(4-cyanovaleic acid) (V501), dimethyl formamide (DMF) were purchased from Sigma Aldrich (St Louis, MO). Poly-L-glutamic acid sodium salt (PGA) (MW 50 000–100 000 g mol⁻¹) was purchased from Biosynth. The plasmid, pZsGreen N-1 (4708 bp), was obtained from Aldevron (Fargo, ND) and the stock solution was further diluted in ultrapure water. DMEM (Dulbecco's modified Eagle's medium), fetal bovine serum, antibiotic-antimycotic (100×), Trypsin-EDTA (0.25%), phenol red, Ultrapure™ DNase/RNase-free distilled water, phosphate-buffered saline (PBS, 1×), Opti-MEM™ I reduced serum medium, FluoroBrite™ DMEM, Hoechst 33342 solution (20 mM), trypan blue solution, 0.4%, and Lipofectamine™ 2000 were purchased from Thermo Fisher (Waltham, MA), jetPEI®, HTS DNA (catalog #89129-916) was purchased from Polyplus. Pierce™ 16% formaldehyde (w/v), methanol-free, ProLong™ glass antifade, were purchased from Invitrogen, Thermo Fisher (Waltham, MA). Label IT® nucleic acid labeling kit was purchased from Cy5®, Mirus bio (Madison, WI). Calcein Deep Red™ AM ester was procured from ATT Bioquest (Pleasanton, CA). Human serum and red blood cells (RBCs) were shared by Dr Nanette Boyle's lab at the Colorado School of Mines. Dialysis membrane tubing (MWCO 3 kDa) was purchased from Spectrum® Laboratories (Reseda, CA).

Polymer synthesis and characterization

p(DIPAEMA₅₆-*st*-HEMA₄₈) was synthesized using RAFT polymerization.⁷⁵ Briefly, DIPAEMA (0.0025 mol), HEMA (0.0025 mol), CEP (0.00007140 mol) and V-501 (0.000007140 mol) were weighed, dissolved in DMF (4 mL) and transferred into scintillation glass vials (20 mL). The reaction mixture was degassed by bubbling nitrogen through the solution for 30 min, heated to 80 °C, and magnetically stirred overnight under nitrogen environment. After 18 h, the reaction mixture was quenched with liquid nitrogen followed by addition of 1 M HCL to dissolve precipitated polymer. The product was purified using dialysis in acidified milliQ water over 5 days, lyophilized, and stored at 4 °C. Purified polymer was designated as DIP50H50. Molecular weight distribution was determined by size exclusion chromatography (Agilent, Santa Clara, CA) with multi-angle light scattering (SEC-MALS) using refractive index and multiple-angle light-scattering detectors (Wyatt, Santa Barbara, CA). ¹H NMR spectra were

recorded using a JEOL ECA 500 instrument (JEOL, Peabody, MA). Polymer ζ-potential was measured using Brookhaven NanoBrook-90 Plus PALS (Brookhaven Instruments, Holtsville, NY). Characterization data is furnished in Fig. S1 in the ESI.†

Polyplex formulation

To obtain DIP50H50 stock solutions, 4.95 mg of DIP50H50 was dissolved in 1 mL of ultrapure water and sterile-filtered. pDNA stock solutions were prepared at 1.04 mg mL⁻¹. Polyplexes were formed by adding equal volumes of diluted polymer stock solutions (diluted to realize targeted N/P ratios) to a 50 ng μL⁻¹ pDNA solution. After gentle pipette-mixing, polyplexes were incubated at 23 °C for 45 min. Polyplexes were further diluted with ultrapure water prior to dye exclusion, dynamic light scattering, and electrokinetic characterization.

A stock solution of PGA (1 mg mL⁻¹) was prepared and sterile-filtered, and diluted to achieve targeted C/N/P ratios. Three PGA coating methods were compared (Fig. 1). In PGA last, PGA was added to pre-made polyplexes and incubated for 45 min. In polycation last, PGA was added to pZsGreen and incubated for 10 min before adding DIP50H50, and incubating for 45 min. In pDNA last, PGA was added to DIP50H50 and incubated for 10 min. Then pZsGreen was added and ternary polyplexes incubated for an additional 45 min. Ternary polyplexes were further diluted with ultrapure water before physical characterization (dye exclusion, DLS, electrokinetic characterization).

Polyplex characterization

To determine whether PGA coating attenuated DIP50H50–pDNA binding, we performed PicoGreen assays. Unbound pDNA will bind PicoGreen, generating a fluorescence signal whereas polymer-bound pDNA will be inaccessible to the PicoGreen dye. We expect the reduction in fluorescence signal to scale with the strength of DIP50H50–pDNA binding. Uncoated or PGA-coated polyplexes were prepared at a pDNA concentration of 50 ng μL⁻¹ and further diluted with ultrapure water. Consistent with the manufacturer's protocol, PicoGreen was diluted to 1 : 200 ratio with ultrapure water. Equal volumes of diluted PicoGreen and polyplexes were mixed and incubated at 23 °C for 30 min. Samples were transferred into black bottom 96 well plates. The fluorescence intensity was measured with monochromator settings of excitation/emission 485/520 nm. Particle size and ζ-potential of polyplexes were analyzed by using Brookhaven NanoBrook 90Plus PALS. The samples were transferred into plastic disposable cuvettes and three measurements performed. The mean diameter, PDI, and ζ-potential values were averaged and reported.

Effect of human serum on polyplex size distribution and DIP50H50–pDNA binding. Serum was separated from whole human blood without addition of anticoagulant. Human serum was further diluted to 10% v/v with sterile PBS (1×). Polyplexes and PGA-coated polyplexes were prepared and the final concentration of pDNA in all samples were fixed at 50 ng μL⁻¹. Equal volumes of diluted serum (10% v/v) and polyplexes were mixed and incubated for 60 min at 23 °C. Samples were



diluted with ultrapure water for prior to dye exclusion assays, electrokinetic characterization, and DLS studies.

Transmission electron microscopy. Polyplex morphology was examined by transmission electron microscopy (TEM). Two microliters of polyplex samples were deposited on copper coated formvar grids (200 mesh), incubated for 5 min, and excess polyplexes were removed using blotting paper. Then grids were washed in milliQ water twice and dipped in 2% uranyl acetate for 10 s. To remove the excess stain, grids were washed with milliQ water and dried in vacuum desiccator overnight. Images were captured using the FEI Talos F200X TEM.

Circular dichroism. Circular dichroism spectra were obtained at 25 °C using a JASCO 815 CD Spectrometer and 1 mm path length quartz cuvette. Polyplexes were prepared as previously described at a pDNA concentration of 125 $\mu\text{g mL}^{-1}$ and a C/N/P ratio of 1/5/1. Three scans were performed and averaged from 200–350 nm at a scan rate of 100 nm min^{-1} and a resolution of 1 nm. Each sample spectrum was normalized to the maximum absorbance between 250–275 nm to adjust for concentration differences.

Cellular assays

Cell culture. Human embryonic kidney cells (HEK 293T) CRL-3216 were procured from American Type Culture Collection (ATCC). HEK 293T cells were grown in T-75 tissue culture flasks using DMEM with 10% heat-inactivated fetal bovine serum (FBS) and 1% antibiotic-antimycotic solution and incubated at 37 °C and 5% CO_2 in a humidified atmosphere. The cell culture media was changed every 2–3 days until the cells reached 80% confluence. Subsequently cells were seeded in 24-well plates for transfection.

pDNA delivery. HEK293T cells were cultured (T-75 flasks) until they reached a confluence of 70–80% and seeded at a density of 50 000 cells per well in 24-well plates. The cells were then incubated for 24 h at 37 °C and 5% CO_2 . Polyplexes were formed at 4 N/P ratios (0, 1, 2.5 & 10) in sterile conditions and incubated for 45 min. Positive controls, jetPEI and Lipofectamine 2000 (LPF 2000) were used in accordance with the manufacturer's directions. Polyplexes (uncoated and PGA-coated) and jetPEI controls were diluted in OptiMEM and added to the HEK293T cells and incubated at 37 °C and 5% CO_2 for 4 h. The concentration of pDNA was fixed as 1 μg per well across all treatment conditions. After 4 h of incubation, 1 mL of DMEM media was added to polyplex and JetPEI sample groups and incubated for 24 h at 37 °C and 5% CO_2 . At the 24 h time point, the media was replaced with fresh DMEM media. After 48 h, the cells were trypsinized, centrifuged at 1000 rpm for 5 min and pellets were washed with PBS. The cells were stained with Calcein Red viability dye (400 nM) in PBS for 30 min at 23 °C. The cells were transferred into FACS tubes and GFP expression was measured using flow cytometry (Cytek, Fremont, CA). Gating strategies are furnished in Fig. S15 and S16 in the ESI.† Transfection efficiency was computed by determining the proportion of GFP⁺ cells in each treatment group.

Effect of serum concentration on transgene expression. The serum tolerance of polyplexes was investigated by increasing the FBS concentration (0, 2.5, 5, or 10%) in DMEM. Before adding polyplexes to cells, polyplexes were resuspended in FBS-supplemented DMEM media rather than OptiMEM. The transfection process and flow cytometry analysis for GFP positive cell measurement were carried out as described earlier. Gating strategies are furnished in Fig. S17 and S18 in the ESI.†

Cell viability assays. Polyplex-induced cytotoxicity was examined 48 h after transfection. Cell culture media was replaced with Fluorobrite-DMEM media supplemented with CCK-8 solution (CCK-8 diluted according to manufacturer instructions). Cells were incubated in CCK-8 containing media for 4 h at 37 °C and 5% CO_2 . Subsequently, the supernatant was transferred into 96-well plates and absorbance was measured at 460 nm using a micro plate reader (Synergy H1, BioTek, Winooski, VT). The background from CCK-8 solution was subtracted and absorbance values normalized to that of untreated cells.

Cellular uptake. Polyplex uptake was measured using Cy5-labeled pDNA. pZsGreen was labeled with Cy5 (Label IT Nucleic Acid Labeling Kit Cy5, Mirus, Madison, WI) and purified using ethanol precipitation. pDNA concentration was quantified using spectrophotometry. Polyplexes (uncoated or PGA-coated) were formulated with Cy5-labeled pDNA at different N/P ratios. Twenty-four hours after pDNA delivery, cells were trypsinized and pelleted. Cells were washed sequentially with PBS, Cell scrub (Genlantis, San Diego, CA), and PBS. Finally, cells were resuspended in 200 μL of PBS with 2% FBS. The geometric mean of Cy5 fluorescence intensity was computed. Gating strategies are furnished in Fig. S19–S22 in the ESI.†

Confocal imaging. The intracellular uptake of polyplexes was analyzed by laser scanning confocal imaging. HEK cells (50 000 cells per well) were seeded in black 24-well glass bottomed plates and incubated for 24 h at 37 °C and 5% CO_2 . Transfection experiments were conducted as described above. Twenty-four hours after pDNA delivery, cells were washed with PBS (1 \times) and stained with Lysobrite Red and incubated for 30 min in dark and washed with cold PBS (3 \times 5min). Cells were counter-stained with Hoechst 3342, incubated for 20 min, washed with cold PBS (3 \times 5 min), and fixed with 4% v/v paraformaldehyde. Cells were imaged under laser scanning confocal microscopy (Nikon A1R) with the following laser emission filter combinations for each fluorophore: Hoescht 3342 - 405 nm laser, Chroma ET450/50m emission filter, GFP - 488 nm laser, Chroma ET525/50m emission filter, Lysobrite Red - 561 nm laser, Chroma 600/50m emission filter, Cy5 - 640 nm laser, Chroma ET685/70m emission filter. The images were captured using 20 \times 0.75 NA Plan Apo Lambda (air) and 60 \times 1.4 NA Plan Apo Lambda (Oil) objective lenses.

The images were analyzed using the Imaris software (Bitplane, version 10.0.0). An initial surface was generated using the Cell (Nucleus and Cytoplasm) tool in Imaris. The DAPI channel was used to segment the nuclei and the GFP channel was used to identify cell bodies. We set a nucleus dia-



meter of 5 μm and a smoothing filter width of 0.5 μm was used for both channels. Intensity thresholds were chosen visually. Nuclei in non-GFP labelled cells were erased. We then manually corrected the surfaces generated by this tool to obtain individual surfaces for the nucleus and cell body for each cell. Spots were then identified using the Spot tool, with an estimated spot size of 0.866. Point spread function elongation along the z-axis was estimated at 2 μm . The spots were then filtered by the “quality” metric, which was set visually between 9.43–16.9.

Finally, the spots were filtered for individual cells and nuclei using the distance to the surface. Spots were initially filtered to identify spots within individual cell bodies, then within individual nuclei (distance to the corresponding surface was set to below 0 μm in the corresponding filters). The total number of spots within the cell body only and within the nucleus was then tallied.

Hemolysis measurements. Hemolysis assays were performed using previously reported procedures.^{96–98} Briefly, human blood was collected in EDTA coated vacutainers. Red blood cells (RBCs) were separated from the whole blood by centrifuging at 1500 rpm for 5 min. Sedimented RBC pellets were separated and supernatant was discarded. RBC pellets were washed thrice with normal saline (150mM). Then RBC pellets were dispersed in PBS (pH 7.4) and diluted to 10-fold from initial concentration. Diluted RBCs were counted using a Neubauer hemocytometer and the diluted RBCs (3×10^7 cells) were added to 100 μL of polyplexes. The concentration of pDNA in the RBC-polyplex mixture was maintained at 50 $\text{ng } \mu\text{L}^{-1}$ for all the samples. Finally, the sample volume was adjusted to 1 mL using PBS (pH 7.4). Negative and positive control RBCs were prepared at the same dilution without polyplexes by dispersing in PBS and water respectively. All the samples were incubated for 1 h at 37 $^\circ\text{C}$ and centrifuged at 1500 rpm for 5 min. The supernatant was collected, and absorbance measured at 540 nm using a microplate reader (Synergy H1, BioTek, Winooski, VT). The hemolysis percentage was calculated from absorbance values (A) as shown below:

$$\text{Hemolysis} = \frac{A_{\text{Sample}} - A_{\text{Water}}}{A_{\text{PBS}}} \times 100$$

Conflicts of interest

The authors do not have any conflicts of interest to disclose.

Acknowledgements

Research reported in this publication was supported by the National Institute of Biomedical Imaging and Bioengineering of the National Institutes of Health under award number 1 R21 EB034464-01. J. L. L. acknowledges support from the National Science Foundation Graduate Research Fellowship Program. This material is based upon work supported by the National Science Foundation Graduate Research Fellowship

Program under Grant No. 2137099. Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation. Authors are grateful to Prof. Kevin Cash and his group for access to a plate reader. Authors acknowledge Prof. Nanette Boyle and her group for sharing human blood samples. Confocal imaging was performed at the BioFrontiers Institute Advanced Light Microscopy Core (RRID: SCR018302). Laser scanning confocal microscopy was performed on an Nikon A1R microscope supported by NIST-CU Cooperative Agreement award number 70NANB15H226. The analysis workstation and the software package Imaris were supported by NIH 1S10RR026680. TEM was performed in the following core facility, which is a part of Colorado School of Mines’ Shared Instrumentation Facility (Electron Microscopy: RRID:SCR022048). Schematics were created using biorender.com.

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