DNA Release Dynamics from Reducible Polyplexes by Atomic Force Microscopy

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DNA condensation by polycations has been studied in the context of polyelectrolyte complexes, phase transitions, DNA packing in human sperms, viral capsids, chromosomes, and gene therapy. DNA condensation is driven by an entropy increase associated with the release of counterions upon the formation of DNA/polyion complexes, or polyplexes. Manning’s counterion condensation theory predicts that 90% of the DNA charges must be neutralized for condensation to occur. When condensed in an aqueous environment, semiflexible DNA chains with a persistence length of ~50 nm tend to display well-defined toroid or rod structures.

Atomic force microscopy (AFM) is used in this study to probe the disassembly dynamics of polyplexes in order to better understand the molecular origin of the difference in in vitro gene delivery performance of polypeptide vectors revealed by a previous study. AFM is a useful tool for the understanding of DNA condensate nanostructures and dynamics. AFM is capable of revealing the degree of heterogeneity among individual condensates that often eludes the more traditional methods such as gel electrophoresis, light scattering, fluorescence microscopy, circular dichroism, and absorption spectroscopy. While the bulk methods are necessary for providing structure and dynamics information without the influence of a surface, the bulk data are averaged over a large number of molecular ensembles. AFM as a direct imaging tool of biochemical processes has several advantages: (1) it is label-free, (2) it can be conducted under near-physiological conditions, and (3) it provides near single-molecule resolution. Specific to this study, the topographical data from AFM consist overwhelmingly of individual nonaggregated polyplexes. Therefore, the common problems associated with aggregation and excess free polycations are largely avoided. In addition, polyplex dynamics on a 2-D surface may resemble more closely its in vivo movement in intracellular protein scaffolds, in the nuclear matrix, and at the biomembrane surface.

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while polyplexes are thought to enter the nucleus most efficiently in reducing environments. The AFM results will be discussed in the context of the physicochemical and transfection properties of the redox active polyplexes. We apply dynamic AFM imaging protocols to observe DNA decondensation and release from polyplexes triggered by dithiothreitol (DTT). A DTT concentration of 20 mM is used to simulate the effect of reducing agents such as glutathione in vivo. The concentration of glutathione has been reported to be ∼4 mM in the cytoplasm and ∼20 mM in the nucleus. The thiol/disulfide exchange reaction between DTT and the disulfide bonds in the polypeptides cleaves the high molecular weight polypeptides into smaller oligopeptides. Consequently, the oligopeptides unbind from the DNA chain. Continuous AFM scanning captures DNA undergoing decondensation within minutes in the DTT solution. We report different DNA release dynamics between NLS and HRP polyplexes in the rate and degree of DNA release and DNA conformational transition pathways.

Experimental Section

Materials. Plasmid DNA vector, gWiz high-expression luciferase, containing the luciferase reporter gene was purchased from Aldevron. Polyplexes containing disulfide bonds in the backbone were synthesized as described previously.45,46 Here, DNA solution (20 mg/mL) in 30 mM sodium acetate buffer (pH 5) was used to prepare polyplexes at the desired basic amino acid (K, H, R) residue-to-DNA phosphate molar ratio (N/P). Acetate buffer (pH 5) was used to increase the protonation of histidine residues in poly(HRP). The polypeptide was added to the DNA solution, and they were mixed by vortexing at 3200 rpm (Fisher Scientific vortex mixer) for 10 s, and then the polyplexes were incubated at room temperature for 30 min. Twenty μL of polyplex solution was placed on 1 cm² freshly cleaved mica. Excess solution was removed after 2 min, and the surface was rinsed with deionized water.

AFM Characterization. AFM imaging was conducted using a Nanoscope III MultiMode atomic force microscope from Digital Instruments. In order to image pure DNA, tapping mode in air was used to increase the protonation of histidine residues in poly(HRP). The polypeptide was added to the DNA solution, and they were mixed by vortexing at 3200 rpm (Fisher Scientific vortex mixer) for 10 s, and then the polyplexes were incubated at room temperature for 30 min.

Sample Preparation. The structure of polyplexes has been found to be sensitive to preparation conditions.45,46 Here, DNA solution (20 mg/mL) in 30 mM sodium acetate buffer (pH 5) was used to prepare polyplexes at the desired basic amino acid (K, H, R) residue-to-DNA phosphate molar ratio (N/P). Acetate buffer (pH 5) was used to increase the protonation of histidine residues in poly(HRP). The polypeptide was added to the DNA solution, and they were mixed by vortexing at 3200 rpm (Fisher Scientific vortex mixer) for 10 s, and then the polyplexes were incubated at room temperature for 30 min. Twenty μL of polyplex solution was placed on 1 cm² freshly cleaved mica. Excess solution was removed after 2 min, and the surface was rinsed with deionized water.

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to always image the adsorbed polyplexes in the same buffer (30 mM pH 5 sodium acetate buffer) and at the same ionic strength as the reductive solution prior to reduction. This will determine the polyplex stability in nonreductive environments and minimize tip disturbance. Usually, the polyplexes were imaged in 40 μL of buffer solutions before 10 μL of DTT solution was injected. The DTT concentration in the liquid cell (VEECO) was maintained at 20 mM. While maintaining the tip and substrate separation to a minimum distance, the DTT solution was injected through silicone rubber tubing into the liquid cell, which was sealed by an O-ring. AFM imaging ensued immediately after solution injection. The surface was imaged continuously at an average rate of 1.5–2 Hz on a 2 × 2 μm² area until no significant changes were occurring at the surface. The ranges of frequency, amplitude, integral, and proportional gains used were 8 kHz, 0.5–1 V, 0.5–2, and 0.75–3 respectively. The AFM images were analyzed using Nanoscope software version 5.12b by Veeco.

We used both sectional height and bearing image analysis commands to determine particle size. The lateral dimension of a particle was determined at the full-width at half-maximum height in order to minimize tip convolution. The bearing analysis command in the Nanoscope software was used to measure the particle volume and area in which the substrate was used as the threshold bearing plane. The particle volume above the threshold plane is calculated by integrating the depth histogram over the entire area above the threshold plane. The DNA contour length was measured using the WSxM 4.0 software. The dynamic results are reproducible with particle size data exhibiting a typical error of 30%.

Light Scattering Characterization. The static light scattering measurements were performed with a DAWN EOS instrument (Wyatt Technology) equipped with a 30 mW linearly polarized GaAs laser. The analysis of the polyplex solutions was conducted at an angular range θ = 22.5°–147° in 20 mL glass scintillation vials at 25 °C. The static light scattering data were analyzed by the second-order Debye fit, \( Kc/R(\theta) \) vs sin\(^2\)θ/2, to obtain polyplex particle density and z-average of the root-mean-square radius (radius of gyration, \( R_g \)) of the DNA polyplexes. \( K \) is the optical constant, which includes the square of the refractive index increment \( n \); \( R(\theta) \) is the Rayleigh ratio, which is proportional to the intensity of the light scattered from solutions; and the polyplex concentration \( c \) is in g/mL. Extrapolation to zero concentration was not performed due to very low concentrations of the polyplexes (<10\(^{-2}\) g/mL). To determine polyplex concentration for \( M_w \) determination required to calculate particle density, complete complexation between DNA and polycations was assumed at N/P ≤ 1 and 1:1 stoichiometry at N/P > 1. The refractive index increments of the DNA polyplexes were calculated as weight-average values using the same assumptions about the polyplex stoichiometry (\( \rho_{DNA} = 0.185 \) and \( \rho_{polyplexes} = 0.188 \)).

Salt Dissociation Assay. Polyplexes at N/P ratio 4 were formed by adding a predetermined amount of polypeptide to a 20 mg/L solution of gWiz-luciferase plasmid DNA in 30 mM sodium acetate (pH 5). Polyplexes were then incubated in 20 mM DTT for 30 min at 37 °C, following which NaCl was added at the indicated concentrations. Samples were then loaded onto 0.8% agarose gel containing 0.5 mg/L ethidium bromide and run for 75 min at 120 V. Polyplexes were then incubated in 20 mM DTT for 30 min. Polyplexes were then imaged in 40 μL of buffer solutions before 10 μL of DTT solution was injected. The DTT concentration in the liquid cell (VEECO) was maintained at 20 mM. While maintaining the tip and substrate separation to a minimum distance, the DTT solution was injected through silicone rubber tubing into the liquid cell, which was sealed by an O-ring. AFM imaging ensued immediately after solution injection. The surface was imaged continuously at an average rate of 1.5–2 Hz on a 2 × 2 μm² area until no significant changes were occurring at the surface. The ranges of frequency, amplitude, integral, and proportional gains used were 8 kHz, 0.5–1 V, 0.5–2, and 0.75–3 respectively. The AFM images were analyzed using Nanoscope software version 5.12b by Veeco.

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Results

In order to avoid confusion, we define depolymerization as the process of converting a polymer into a monomer. In this paper, it specifically describes the degradation of the reducible polypeptides into their oligopeptide building blocks by breaking disulfide bonds. Decondensation is defined as the alteration of DNA structure from the condensed form to a relaxed disperse

Figure 1. Static light scattering measurements of \( R_g \) and particle density variations in solutions of (a) poly(NLS)/DNA (N/P = 4) and (b) poly(HRP)/DNA (N/P = 4) as functions of time in 20 mM DTT. Data were acquired every 5 s.

form. In AFM imaging, it appears as the transition from spherical/rodlike particles to supercoiled DNA chains. Desorption describes the process of removing an adsorbed material from the solid on which it is adsorbed. In this paper, it means that the decondensed DNA molecules (or fragments) detach from the substrate and thus are “invisible” to AFM. Disassembly describes both decondensation and dissociation of the polycations from the polyplexes.

First, static light scattering was used to follow changes in \( R_g \) and structural density (calculated as \( 3M_w/(4\pi N A R_g^3) \)) of the polyplexes in 20 mM DTT (Figure 1). \( N_A \) is Avogadro’s number. Abrupt changes for poly(NLS) polyplexes in solution were observed by static light scattering (Figure 1a). There was an initial period after addition of DTT when no changes in particle density (∼650 s) and in \( R_g (∼730 s) \) were observed. This initial period is then followed by a rapid increase of both of the measured parameters, suggesting aggregation of the polyplexes and internal rearrangement leading to the increased structural density. The increase in particle density suggests that the initial degradation stage tends to compact the polyplexes prior to the DNA release stage. It is possible that better packing is achieved by rearrangement of shorter polymer chains. The earlier onset of the changes in particle density compared to particle size suggests that, in the first phase of the reduction, internal rearrangement of polyplexes takes place. The observed aggregation is consistent with the removal of the surface loops of free polycations that otherwise stabilize the polyplexes against aggregation. Poly(HRP) formed polyplexes with lower structural density than those of poly(NLS), and unlike the case of poly(NLS) changes in \( R_g \) were observed almost immediately after DTT addition (Figure 1b). The initial decrease in size of poly(HRP)/DNA is followed by aggregation similar to that of the poly(NLS) polyplexes.

The light scattering data show that while this technique is valuable for the study of polyplex dynamics in solution, it is not suitable...
to follow the polyplex dissociation dynamics above the minimum salt concentration required for polyplex dissociation, $c_d$. This is due to extensive aggregation of the polyplexes above $c_d$ even under oxidizing conditions before DTT addition. The limitation of the light scattering technique is a motivation for carrying out the in situ AFM study above $c_d$.

Second, gel electrophoresis was used to follow DNA release in the reducing and nonreducing environments. Representative results for poly(NLS) polyplexes are shown in Figure 2. The results confirm the stability of the polyplexes to dissociation in a range of increasing salt concentrations (lanes 5–8). In the presence of DTT, reduction of disulfide bonds results in an almost complete release of DNA as expected from the polyplexes (lanes 9–12). However, the fact that the DNA mobility is not completely restored to that of control noncomplexed DNA (lane 2–4) suggests the possibility of a residual binding of reduced NLS monomers.

Lastly, we describe the primary results from AFM investigation. Before presenting the results of the complexes, pure plasmid DNA was imaged by AFM. Plasmid DNA prior to condensation with the polypeptides shows the typical relaxed open-loop structure$^{51}$ with little twisting of the strands and a typical height of 2 nm (Figure 3a). Since negatively charged DNA does not adsorb on negatively charged mica, plasmid DNA was anchored on mica by the addition of a small amount of Mn$^{2+}$ ($10^{-6}$ M). The average contour length ($W_{sxM}$ 4.0 image analysis software), 1.9 ± 0.2 nm, is less than the contour length calculated for B-form DNA with 6732 bp, 2.3 μm. The measured contour length is usually limited by the spatial resolution of AFM.

Our previous paper reported positive ζ potential values, 55 and 47 mV, for polyplexes prepared with excess poly(HRP) and poly(NLS), respectively.$^7$ The positively charged polyplexes adsorb readily on negatively charged mica and can be imaged by in situ AFM in the fluid-tapping mode. Both poly(NLS) and poly(HRP) were capable of condensing DNA at N/P ≥ 2 (Figure 3b). When N/P increased from 2 to 8, AFM showed negligible variation in polyplex morphology (data not shown). The majority of the DNA condensates are either spheroids (Figure 3c) or rods (Figure 3d) with a small population, ∼5%, of larger flowerlike objects (Figure 3e). The typical dimensions of spheroids were measured to be 7.0 ± 2.5 nm in height and 50.3 ± 7.8 nm in diameter. The typical dimensions of the rods were 3.8 ± 1.5 nm in height, 36.3 ± 11.5 nm in width, and 163 ± 50 nm in length. The lateral dimensions were measured at half-height (see sectional height profiles in Figure 3c,d) in order to minimize tip convolution.

All the values are within the range reported previously.$^{2,11,52}$

Another typical morphology of polyplexes, toroids, was not observed with these polyplexes. Figure 4a plots the volume histograms of spheroids and rods from AFM bearing analysis for poly(NLS)/DNA and poly(HRP)/DNA. The number-average volumes of poly(NLS) and poly(HRP) condensates are 2.71 × 10$^4$ and 2.92 × 10$^4$ nm$^3$, respectively. The volume of a closely packed polyplex containing a single plasmid DNA with 6732 bp is estimated to be 1.4 × 10$^5$ nm$^3$, by assuming an interhexagonal separation between a neighboring peptidene/DNA chain of 2.7 nm.$^{55}$ Therefore, according to the volume histograms, approximately 50% of the condensates contain only one DNA plasmid per particle. The volume histograms also indicate that rods are more uniform in size than spheroids. Figure 4b shows that over 80% of rods are in the smallest size range while the spheroids have a wider size distribution. AFM morphological analysis of the condensates concludes that poly(NLS) and poly(HRP) share a similar condensate structure.

The controlled release nature of the polyplexes permits further visualization of the DNA molecular distribution among the

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condensates. AFM images of the same area before and after the introduction of DTT allow a direct count of the number of DNA molecules in each condensate. The open loop structure and unique contour length of DNA makes it possible to count their numbers. In order to immobilize DNA on mica, 0.2 M MgCl₂ was added to 20 mM DTT solution. Divalent ions such as Mg²⁺ have been shown to attract negatively charged DNA to negatively charged mica by counterion correlations. Figure 5 shows before and after AFM images of DNA condensates with poly(NLS). In the absence of Mg²⁺, DNA chains from poly(NLS) condensates were directly released to the solution (see results below). Therefore, Mg²⁺ is proven effective in allowing DNA to be imaged clearly on mica after it is no longer bound to NLS. We were able to determine that indeed condensates with an initial volume of 1.4 × 10⁴–2.0 × 10⁴ nm³ (represented by A–D) contain only one DNA molecule per particle using the Mg²⁺-assisted method. Similarly, condensates with a volume of 3.7 × 10⁴–4.1 × 10⁴ nm³ (represented by E and F) contain two DNA molecules. The rodlike condensate (represented by G) with a slightly higher initial volume of 3.1 × 10⁴ nm³ also contains only one DNA plasmid. In our case, poly(NLS) and poly(HRP) are effective in condensing DNA into the closest packing structure.

Prior to the injection of the reducing DTT solution, the stability of the condensates was studied in aqueous buffer solutions containing up to 1 M NaCl. The structure of the condensates was unchanged for at least 1 h under repeated AFM scanning. It was concluded that salt alone was not able to induce decondensation due to the high molecular weight of the polypeptides in agreement with the gel electrophoresis results. Minimum scanning force magnitudes were maintained in the subsequent measurements.

In the presence of 20 mM DTT, the stability of the polyplexes was found to be dependent on the salt concentration. The minimum salt concentration required for polyplex dissociation, c₄₅, provides a means to assess the binding affinity of polycation to DNA, and it has been found to be strongly dependent on the molecular weight when the degree of polymerization is low. AFM measurements determined the c₄₅ value for both poly(NLS) and poly(HRP) condensates to be in the range of 0.1–0.2 M NaCl. No changes in the appearance of the poly(NLS)/DNA (N/P = 4) condensates were observed below 0.1 M NaCl (data not shown). In the case of poly(HRP) condensates (N/P = 4), changes in the form of a gradual height reduction from over 10 to ∼6.5 nm were observed in the first 2.3 × 10³ s in DTT and 0.1 M NaCl (Figure 6). No further changes were observed after 2.3 × 10³ s. The cause of particle flattening is hypothesized to be the removal of excess cations in the particle shell region and/or molecular rearrangement as a consequence of polypeptide degradation into oligopeptides. DTT cleaves the disulfide bonds to reduce the high molecular weight polypeptide into shorter α,ω-dithiol-oligopeptides. Below c₄₅, shorter oligopeptides are capable of closer packing in the particle than the polypeptide counterpart. Gel dissociation assays of the same systems determined a lower

limit of $c_d$ to be 0.3 M NaCl. Here, AFM offers another means of DNA dissociation assays. It shows high accuracy, requires minimum amount of sample, and minimizes the contribution from free polycations.

DNA decondensation from poly(NLS) polyplexes occurred at salt concentrations significantly below 1 M in the presence of DTT. Figure 7 presents time-lapse AFM images of DNA release from poly(NLS) polyplexes (N/P = 4) in 20 mM DTT and 0.4 M NaCl solution captured immediately after the addition of DTT. The most significant events occurred after $1.28 \times 10^3 - 1.50 \times 10^3$ s. The start of the nanoassembly unraveling can be gauged by the appearance of fuzzy edges ($480 - 800$ s). Immediately after the complete release and desorption of DNA (1.50 $\times 10^3$ s), the peptide remnants, akin to the “footprints” left by the condensates, were visible in the AFM image. The DNA release event is marked by a sudden decrease in the average volume per particle (Figure 8a). It shows that on average a particle has lost 50–60% of its total volume after $1.50 \times 10^3$ s for N/P = 2, 4, and 8 (see Figures S1 and S2 in the Supporting Information for in situ AFM images of DNA release from condensates with N/P = 2 and 8). The data suggest that poly(NLS) polyplexes release DNA simultaneously after an induction period with little dependence on the initial particle size, shape, or the N/P ratio. Presumably, the initial ∼20% volume reduction is due to the gradual loss and rearrangement of the $\alpha,\omega$-dithiol-NLS residues; the abrupt 50–60% volume reduction corresponds to the release of DNA, which can still carry up to 90% of its counterion charges; and the last 20–30% are the NLS residues remaining on mica. We calculate the N/P ratio of the initial poly(NLS) polyplex composition using the formula $(0.54 \text{ nm}^3 \times P + 0.75 \text{ nm}^3 \times 0.90 \times P)/(0.54 \text{ nm}^3 \times P + 0.75 \text{ nm}^3 \times N) = 0.6$, which yields an N/P ratio close to 2.0. The values 0.54 nm$^3$ and 0.75 nm$^3$ are the unit volume for P and N repeat unit, respectively. In this calculation, we assume that the released DNA occupies roughly 60% of the total complex volume based on the AFM data. We further assume that the released DNA is 90% neutralized by the NLS peptide residues. The same volume loss percentage for N/P = 2, 4, and 8 suggests a maximum degree of overcharging was reached when the solution N/P mixing ratio was increased from 2 to 8.

The disassembly of poly(NLS) polyplexes (N/P = 4) was studied as a function of salt concentration. Figure 8b shows that DNA is released at a faster rate as measured by the particle volume reduction with increasing salt concentration. The volume percentage was obtained from AFM bearing analysis averaged over 10 particles at each time interval. Therefore, the salt effect is mainly to shorten the time leading to the sudden DNA release event from $1.3 \times 10^3$ s in 0.2 M NaCl to almost instantaneous release in 0.8 M NaCl solution. Depolymerization, loss of excess oligocations, and rearrangement of remaining oligocations within the particle could occur during the induction period. Depolymerization is unlikely to be the rate-limiting step. Control $\alpha,\omega$-dithiol-oligocations showed the same salt dissociation rate as those of the polyplexes. Rearrangement of NLS could be the
chains continued to emerge and expand until 1.2 im immediately after the addition of DTT (160 s). Loose DNA the unraveling of the nanoassemblies, were observed almost increase due to DNA decondensation. Fuzzy edges, signaling dynamics of poly(HRP) polyplexes is quite different from that cause the change near the end of the sequence. No significant changes occurred after 3.0 × 10^3 s. It is clear that the release dynamics of poly(HRP) polyplexes is quite different from that of poly(NLS) polyplexes. DNA plasmids were released from HRP polyplexes, gradually resulting in partially decondensed intermediates, and the plasmids never fully desorbed from the mica substrate.

The AFM study also captured intriguing size-dependent disassembly behavior of poly(HRP) condensates. Particles with an initial diameter (measured at half-height) of 50 nm or more underwent decondensation while particles below this critical size, marked as A–F, were more resistant to reductive degradation (Figure 9). Exceptions marked as G–I can be found to be close to larger particles. These particles with a center-to-center distance to nearest larger particles of less than 100 nm seem to be destabilized by the nearby chain unraveling events. The momentary bridge by a DNA chain from a neighboring particle to the smaller particle J can be seen at 850 s.

**Discussion**

This study demonstrates the use of AFM in capturing the dynamics and heterogeneity during DNA release from polypeplexes. The effects of physical attributes of polypeplexes such as particle size on gene delivery performance are complicated because the size effect is different at different stages of gene delivery. While it is unlikely that we can predict DNA release in vivo solely based on the DNA release behavior observed by AFM conducted in a highly simplified environment, we discuss the AFM results in the context of a previous in vitro study of the physicochemical and physiological properties of the poly(NLS) and poly(HRP) DNA delivery vectors. The hydrodynamic diameters and ζ potentials of the polypeplexes (pH = 5, 20 mg/L DNA, N/P = 4) were determined by dynamic light scattering to be 105 ± 3 nm, 47 ± 1 mV for poly(NLS) and 96 ± 1 nm, 55 ± 1 mV for poly(HRP). AFM morphological analysis indicates that the two form condensates similar in shape and size distribution. AFM results further indicate that the condensates are highly compact and approximately half of them contain only one DNA plasmid.

Destabilization experiments by a polyanion exchange reaction found that compact DNA condensates are more stable against polyanion exchange. While all the polycations bind equally well to DNA, they show selectivity toward dissociation. In this study, correlation between disassembly dynamics and physical attributes of as-formed polypeplexes such as size could not be evaluated, since there were no significant differences between sizes of poly(NLS) and poly(HRP) polypeplexes. The AFM results provide a partial explanation for the apparent lack of NLS effect on the transfection efficiency. AFM results suggest that NLS dissociates abruptly and completely from DNA. It is possible that the disulfide bonds in poly(NLS) are cleaved in the reducing environment of the cytosol so that NLS are no longer associated with DNA to promote nuclear transport. Bulk experiments tend to show strong influence of the N/P ratio on biophysical properties and transgene efficiency. AFM experiments conducted in polymer-free buffer solutions focus on surface-adsorbed polypelexes. Therefore, it is not surprising that we did not observe the N/P effect caused by excess polycations.

The issue of molecular weight effect on polypeplex packing is unresolved in the literature. Some have argued that high molecular weight polymers prevent close packing of DNA in the condensates. For example, the volume of a toroid or rod containing a single plasmid DNA (6832 bp) and polylysine-asialooso-

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**Figure 9.** *In situ* AFM sequence of DNA release from poly(HRP)/DNA polypeplexes (N/P = 4) in 20 mM DTT and 0.2 M NaCl solution (no Mg^2+ was added). Time zero corresponds to the addition of DTT. All images are 2 × 2 µm² in scan size and 10 nm in z range.
mucoid was greater than $2 \times 10^4$ nm$^3$. While others have reached an opposite conclusion, that is, high molecular weight favors close packing. Still others found no direct correlation between molecular weight and packing density. Our study suggests in the first place that the high molecular weight of the polypeptides does not prevent close packing of the polyplex, and in the second place that, as the molecular weight is decreased by chemical reduction, the polyplex undergoes further compaction before the irreversible expansion sets in. It points to an optimal molecular weight range for molecular packing in the condensate.

The intermediates observed during DNA decondensation largely resemble partially condensed DNA reported in the literature. Partially condensed DNA with 8%–83% charge neutralization formed bundles, folded loops of DNA surrounding central cores to lose coils with isolated nodes. The high-density nodes have been speculated to be the possible intermediate structures between random coils and compact globule condensates. AFM and simulation studies by Iwataki et al. illustrated the first-order transition from single chain compaction and multichain bundle formation. The AFM results show that, during decondensation, structural heterogeneity develops within the particle. Correlation-induced charge inversion gives rise to overcharge of DNA by polycations. The polyplexes may consist of a neutral core and a shell of the excess polycation. The reduction starts from the shell to the core, giving rise to the fuzzy edge at the onset of the polyplex disassembly. Heterogeneity during decondensation means that DNA segments in different parts, that is, the less compact region versus the highly condensed center region (nucleation site), may show different stability against enzymatic degradation.

AFM shows that particles during decondensation display interesting interparticle dynamics. Ono et al. obtained in situ AFM images that demonstrate the DNA condensate mobility when attached to the surface by electrostatic interaction. Two nearby condensates were capable of interacting with each other and formed parallel strands as an intermediate to the final multimeric aggregate. The parallel strand formation was attributed to hydrophobic minification from the bound polycation. The hydrophobic interaction is responsible for the formation of parallel and staggered bundles of DNA strands, which then leads to crystallization after spermne neutralizes the DNA surface.

We found evidence of interparticle interaction in DNA decondensation. The interparticle interaction is facilitated by a reduction in the positive charges or local charge reversal due to polymer degradation. The same local charge reversal mechanism results in intersegment attraction in polyplex formation. The less compact condensate starts to recover phosphate groups that make part of the chain negatively charged. The negatively charged DNA interacts electrostatically with the positively charged DNA and mica and hinders DNA release. The fact that DNA strands after decondensation are still bound to mica indicates that HRP must remain on mica to provide an attractive charge correlation. The counterion valence has been shown to affect the mobility of DNA on mica. In one study, DNA chains bound to Mg$^{2+}$ were free to condense into toroids while multivalent protonate with 22 arginine residues preadsorbed on mica trapped DNA in various uncondensed to partially condensed states. DNA can adsorb on mica at relatively high ionic strength in the presence of a low concentration of multivalent cations. It is interesting that NLS with a charge valence of 6 is ineffective in trapping DNA while 0.2 M MgCl$_2$ is effective. It is possible that the concentration of NLS$^{6+}$ is too low for it to be effective in this case. However, salt dissociation assays carried out in agarose gel and by AFM do not show a significant difference in salt dissociation between HRP and NLS. The $c_d$ value determined by AFM, 0.1–0.2 M NaCl, is in agreement with that of the salt dissociation assay, which identified an upper limit of 0.3 M. The AFM results suggest that while it is likely that the difference in binding affinity is responsible for the different DNA release dynamics, it may not be reflected in the simple salt dissociation assays.

### Conclusions

The AFM study demonstrates the release of DNA from reducible polyplexes under simulated physiological reducing conditions. The reducible polyplexes provide an ideal system to study DNA decondensation dynamics by real-time AFM. The polyplexes are stable in the oxidizing environment representative of the nonreducing extracellular space. Rapid release of DNA from both NLS and HRP polyplexes in the presence of DTT and moderate salt concentration simulates the conditions of reducing cytosolic and nuclear compartments. The goal of this study was to understand DNA release dynamics from polyplexes in mildly...
reducing conditions. In order to mimic conditions in cytosol, future experiments will be conducted in conditions more closely resembling the cytosol, for example, in glutathione instead of DTT. The polyplex stability, the intermediate partially destabilized structure, and their interactions impact its delivery efficiency and its effect on cell viability. Despite the similarity in the physical properties and nanostructures of the two types of polyplexes, their disassembly dynamics is quite different. DNA is released from NLS abruptly regardless of the initial polyplex morphology, while HRP-bound DNA displays a gradual decondensation that is size dependent. The disassembly rate is higher for larger HRP polyplexes. The results reveal potentially rich DNA release dynamics that can be controlled by subtle variation in multivalent counterion binding affinity to DNA and the surrounding cellular matrix. The tendency of partially disassembled complexes to interact and aggregate with each other cannot be overlooked. The results point to a potential challenge in regulating the stability of gene delivery vectors during disassembly due to the heterogeneous structure and charge distribution of the various intermediates.

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**Supporting Information Available:** In situ AFM images of DNA release from poly(NLS)/DNA polyplexes with N/P = 2 and 8. This material is available free of charge via the Internet at http://pubs.acs.org.